Cation Overcrowding Effect on the Oxygen Evolution Reaction

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ABSTRACT: The influence of electrolyte ions on the catalytic activity of electrode/electrolyte interfaces is a controversial topic for many electrocatalytic reactions. Herein, we focus on an effect that is usually neglected, namely, how the local reaction conditions are shaped by nonspecifically adsorbed cations. We scrutinize the oxygen evolution reaction (OER) at nickel (oxy)hydroxide catalysts, using a physicochemical model that integrates density functional theory calculations, a microkinetic submodel, and a mean-field submodel of the electric double layer. The aptness of the model is verified by comparison with experiments. The robustness of model-based insights against uncertainties and variations in model parameters is examined, with a sensitivity analysis using Monte Carlo simulations. We interpret the decrease in OER activity with the increasing effective size of electrolyte cations as a consequence of cation overcrowding near the negatively charged electrode surface. The same reasoning could explain why the OER activity increases with solution pH on the RHE scale and why the OER activity decreases in the presence of bivalent cations. Overall, this work stresses the importance of correctly accounting for local reaction conditions in electrocatalytic reactions to obtain an accurate picture of factors that determine the electrode activity.

KEYWORDS: Hydrogen production, Oxygen evolution reaction, Electrolyte cation effects, Local reaction condition, Surface charging effects, Electric double layer

INTRODUCTION

Electrocatalysis is concerned, in the broadest sense, with the transformation between energy stored in chemical bonds and electrical energy. A hallmark of electrocatalysis is that the reaction rate, i.e., the rate of the transformation between chemical energy and electricity, can vary up to 10 orders of magnitude on different electrode materials.1−2 The electrode dependence of the reaction rate is ascribed mainly to electronic interactions between the electrode and reaction intermediates.1−3 A quintessential descriptor of electronic interactions is the binding energy of reaction intermediates.1−4 It is an important and vibrant direction of research that the reaction rate can be enhanced by tuning this descriptor, e.g., via harnessing strain and ligand effects.5−6

Electrocatalytic reactions occur in a nanoscale interfacial region between the solid electrode and the electrolyte solution, termed electric double layer (EDL). Therefore, the reaction rate can also be tuned by modifying the composition of the electrolyte solution; relevant effects are termed electrolyte effects. Electrolyte effects have been observed in many electrocatalytic reactions, including hydrogen oxidation/evolution,7−10 oxygen evolution/reduction,7,11−14 methanol electro-oxidation,15 and carbon dioxide reduction.16,17 This work is concerned with the effects of electrolyte cations on the oxygen evolution reaction (OER), which is the performance-limiting anode reaction in electrochemical water splitting devices for hydrogen production.

The study of cation effects in the OER has a long history. Early studies were conducted by Erdey-Gruz and Shafarik in 1961 at platinum in 1 M H2SO4 with 0.3−1 M K+, Na+, Li+, Zn2+, Mg2+, Al3+, and NH4+18 and by Kozawa in 1964 at several metals in 1 M NaOH with 0.1−10 mM Ba2+, Cr3+, and Ca2+.19 Erdey-Gruz and Shafarik observed that the activity decreases in the order K+ < Al3+ < NH4+ < Zn2+ < Na+ < Mg2+ < Li+.18 Kozawa found that the overpotential increases by 15−90 mV in the presence of 0.1−10 mM Ba2+, Cr3+, and Ca2+.19 Subsequent interest in the cation effects on the OER came from the field of alkaline batteries in the 1990s.20−23 More recent interests are sparked by the study of cation effects on hydrogen oxidation reaction and oxygen reduction reaction by Strmcnik et al.7

Cation effects on OER at NiOOH thick films in both purified and Fe-containing alkaline electrolytes are of primary current interest, as Fe-doped NiOOH-based materials are promising catalysts for the OER. In 2015, Michael et al. observed the activity trend of Cs+ > K+ > Na+ ≈ Li+ in purified electrolyte solutions and K+ ≈ Na+ > Cs+ > Li+ in Fe-containing electrolyte solutions.24 In 2017, Zaffran et al. observed that...
alkaline earth cations (Mg$^{2+}$ and Ca$^{2+}$) dramatically decrease the OER activity of NiOOH thin films in both purified and Fe-containing alkaline electrolyte solutions, a modern echo of the results of Kozawa reported more than 50 years ago.25 Yang et al. further added tetraalkylammonium (TAA) cations into the comparison. They found an activity trend of K$^+$ ≈ Na$^+$ ≈ TBA$^+$ > TMA$^+$ > Li$^+$ for the OER at NiOOH thin films in purified alkaline electrolyte solutions and a different activity trend of K$^+$ ≈ Na$^+$ > TBA$^+$ > Li$^+$ > TMA$^+$ in the presence of Fe impurities.26 In 2019, Garcia et al. studied the effects of alkali metal cations on the OER at NiOOH thin films in both purified and Fe-containing alkaline electrolyte solutions. Their results show that the OER activity increases in the order of Cs$^+$ > Na$^+$ > K$^+$ > Li$^+$ in purified 0.1 M MOH (M = Cs, K, Na, Li), and Cs$^+$ > K$^+$ > Na$^+$ > Li$^+$ in the presence of Fe impurities.14

Perini and Ticianelli reported an activity order of K$^+$ > Na$^+$ > Cs$^+$ > Li$^+$ in purified 0.1 M MOH (M = Cs, K, Na, Li), and Cs$^+$ > K$^+$ > Na$^+$ > Li$^+$ in the presence of Fe impurities.14

In 2015, Michael et al. measured Raman spectra using in situ SERS.28 In addition, they found that the density of intermediates. In 2019, Garcia et al. studied the effects of solution pH and other electrocatalytic reactions.

**DFT Calculations**

Spin-polarized periodic first-principles calculations based on density functional theory (DFT) were performed with the projector-augmented-wave (PAW)16,37 method using the Vienna ab initio simulation package (VASP).38-40 The Perdew–Burke–Ernzerhof (PBE) functional within the generalized gradient approximation (GGA) was used for exchange-correlation effects.16 The wave functions were expanded by a plane wave basis with a kinetic cutoff energy of 750 eV. The DFT+U approach42,43 was applied to correctly reproduce the formation energies of TMA$^+$ with active oxygen-containing intermediates of the OER, which are the output of the microkinetic model. A consistent treatment of these models is a necessity for deciphering how electrocatalytics influence the OER and is also a unique feature of this work in comparison with similar models for the OER40-42 and other electrocatalytic reactions.

We adopt the concerted theory-computation framework, previously developed for oxygen reduction reaction by some of us,39 for the present case of the OER. The framework is composed of three parts. In the first part, we determine the reaction mechanism using quantum-mechanical DFT calculations. In the second part, we treat the multistep kinetics using microkinetic modeling. In the third part, we use a continuum EDL model to calculate the local reaction conditions. These parts are interrelated. The microkinetic model, on the one hand, is based on the reaction mechanism determined from DFT calculations and, on the other hand, requires knowledge of the local reaction conditions that are determined from the EDL model. In turn, the EDL is influenced by interfacial reactions in that the surface charge density is a function of the concentrations of reaction intermediates, which are the output of the microkinetic model. A consistent treatment of these models is a necessity for deciphering how electrocatalytics influence the OER.

To describe the energetics of the OER on the β-NiOOH surface, the grand canonical ab initio thermodynamics scheme47,48 was used to evaluate the formation enthalpy of intermediates in solution. The computational hydrogen electrode (CHE) concept49,50 was applied to calculate the electrochemical potential of H$^+$ in solution, $\mu_{H^+}$, at a given electrode potential:

$$\mu_{H^+} = \frac{1}{2}\mu_{H^0} - eU_{\text{SHE}} - k_B T \ln(10) \text{pH} = \frac{1}{2}\mu_{H^0} - eU_{\text{SHE}}. $$  

It is certainly fair to say that the understanding of cation effects in the OER is still incomplete. Most studies have been focusing on the interaction of electrolyte cations with the OER intermediates. In 2015, Michael et al. measured Raman spectra during the linear scanning voltammetry in purified LiOH and CsOH. However, they pointed out that the "precise mechanism by which CsOH promoted better catalytic performance than LiOH is unclear with results from this study." In 2017, Zaffran et al. assumed that the electrolyte cations are intercalated into the layered NiOOH structure. Their computational results showed that intercalated Mg$^{2+}$ and Ca$^{2+}$ bind strongly to water and OER intermediates, thus decreasing the OER activity.27 Yang et al. interpreted the cation effects from the perspective of interactions between cations and intermediates of the OER. They attributed the inhibition effect of TMA$^+$ on the OER activity to the specific interaction of TMA$^+$ with active oxygen-containing intermediates formed upon deprotonation.28 Later, Garcia et al. identified NiOOO$^-$ as the active oxygen-containing intermediate using in situ surface enhanced Raman spectroscopy (SERS). Further, they showed that NiOOO$^-$ is better stabilized by larger cations Cs$^+$, which is believed to promote the OER. Contra- 

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Figure 1. Gibbs free energy profile of OER on β-NiOOH (0001) surface calculated under standard conditions (298 K, pH = 0, 1 bar, 0 V SHE).

where \( \mu_i \) is the electrochemical potential of the electrons and \( \mu_{\text{H}_2\text{O}} \) that of hydrogen gas under standard conditions. \( U_{\text{RHE}} \) is the electrode potential on the reversible hydrogen electrode (RHE) scale. Using \( U_{\text{RHE}} \) instead of \( U_{\text{SHE}} \) versus the standard hydrogen electrode (SHE) aligns our results with the convention used in most experimental studies. The electrochemical potential of \( \text{OH}^- \) can be derived from

\[
\mu_{\text{OH}^-} = \mu_{\text{H}_2\text{O}} - \mu_{\text{H}^+}
\]

First, we calculated the Gibbs free energy change for each elementary step of the OER and explored the most favorable pathway under standard conditions \((T = 298 \, \text{K}, \, pH = 0, \, p_{\text{H}_2} = 1 \, \text{bar}, \, 0 \, \text{V SHE})\). Multiple experimental studies have revealed that the original surface undergoes surface deprotonation prior to the OER. \(^{14,51,52}\) Accordingly, the first step of the OER is thus

\[
\text{O–Ni–OH} + \text{OH}^- \leftrightarrow \text{O–Ni–O} + \text{H}_2\text{O} + e
\]

where \( \text{O–Ni–OH} \) represents the original surface and \( \text{O–Ni–O} \) is the \((2 \times 2)\) surface with \(1/4\) hydrogen atoms deprotonated.

The second step is the adsorption of a hydroxyl anion (\( \text{OH}^- \)) onto the deprotonated site, forming \( \text{O–Ni–OOH} \),

\[
\text{O–Ni–O} + \text{OH}^- \leftrightarrow \text{O–Ni–OOH} + e
\]

Afterward, another \( \text{OH}^- \) approaches and removes one \( \text{H} \) from \( \text{O–Ni–OOH} \),

\[
\text{O–Ni–OOH} + \text{OH}^- \leftrightarrow \text{O–Ni–OO} + \text{H}_2\text{O} + e
\]

followed by a chemical step releasing \( \text{O}_2 \), leaving the surface \( \text{O–Ni–} \) with an oxygen vacancy,

\[
\text{O–Ni–OO} \leftrightarrow \text{O} – \text{Ni–} + \text{O}_2
\]

Finally, the initial surface is recovered by filling the oxygen vacancy site with a \( \text{OH}^- \) in solution, completing the reaction cycle,

\[
\text{O–Ni–} \leftrightarrow \text{O–Ni–OH} + e
\]

The Gibbs free energy changes of the elementary steps as a function of the electrochemical control parameters were derived using the CHE method.\(^{49,50}\) Steps 3, 4, 5, and 7 proceed via \( \text{OH}^- \)-coupled electron transfer steps, for which we need the electrochemical potential of the electron–OH’ pair,

\[
\mu_{\text{OH}^-} - \mu_{\text{e}} = \mu_{\text{H}_2\text{O}}^0 - \frac{1}{2} \mu_{\text{H}_2\text{O}}^0 + eU_{\text{RHE}}
\]

where \( \mu_{\text{OH}^-} \) is the electrochemical potential of hydroxyl anions and \( \mu_{\text{H}_2\text{O}}^0 \) is that of liquid water under standard conditions. In order to compute \( \mu_{\text{H}_2\text{O}}^0 \) and \( \mu_{\text{H}_2\text{O}}^0 \), we calculated the total energy of a single water molecule and a single hydrogen molecule using DFT and corrected for entropic contributions using entropy data from the NIST Chemistry Webbook.\(^{53}\)

Consider, for example, the first deprotonation step. The Gibbs free energy change of this step at any potential \( U_{\text{RHE}} \) is calculated as

\[
\Delta G_1 = E_{\text{O–Ni–O}} - E_{\text{O–Ni–OH}} - \mu_{\text{OH}^-} - \mu_{\text{e}} = (E_{\text{O–Ni–O}} - E_{\text{O–Ni–OH}}) + \frac{1}{2} \mu_{\text{H}_2\text{O}}^0 - eU_{\text{RHE}}
\]

where we used eq 8 for \( \mu_{\text{OH}^-} - \mu_{\text{e}} \) and \( E_{\text{O–Ni–O}} \) and \( E_{\text{O–Ni–OH}} \) represent the total energies of the \( \text{O–Ni–O} \) and \( \text{O–Ni–OH} \) surfaces calculated using quantum-mechanical DFT. The Gibbs free energy change of other steps can be calculated in the same manner, expressed as

\[
\Delta G_2 = E_{\text{O–Ni–OOH}} - E_{\text{O–Ni–OH}} - \mu_{\text{H}_2\text{O}}^0 + \frac{1}{2} \mu_{\text{H}_2\text{O}}^0 - eU_{\text{RHE}}
\]

\[
\Delta G_3 = E_{\text{O–Ni–OO}} - E_{\text{O–Ni–OH}} + \frac{1}{2} \mu_{\text{H}_2\text{O}}^0 - eU_{\text{RHE}}
\]

\[
\Delta G_4 = E_{\text{O–Ni–OOH}} - E_{\text{O–Ni–OO}} - \mu_{\text{H}_2\text{O}}^0 + \frac{1}{2} \mu_{\text{H}_2\text{O}}^0 - eU_{\text{RHE}}
\]

where we used \( \mu_{\text{H}_2\text{O}}^0 = 4.92 \, eV + 2(\mu_{\text{H}_2\text{O}}^0 - \mu_{\text{H}_2\text{O}}^0) \), in eq 12.

The Gibbs free energy profile of the OER calculated under standard conditions \((298 \, \text{K}, \, pH = 0, \, 1 \, \text{bar}, \, 0 \, \text{V SHE})\) is shown in Figure 1. The first deprotonation step has the most positive Gibbs free energy change, \( \Delta G_1^0 = 1.91 \, eV \), which means that it corresponds to the potential determining step, which is consistent with the results of Govind-Rajan et al.\(^{74}\) The following four steps are energetically feasible at the equilibrium potential of the OER, \( E = 1.23 \, V_{\text{RHE}} \). Especially, the NIO state with an oxygen vacancy is highly unstable and inclined to adsorb a \( \text{OH}^- \) from solution. Therefore, we combine the step 4 and 5 into a single step in the microkinetic model.

**Microkinetic Model**

A widely assumed reaction pathway of the OER in alkaline media proceeds via four \( \text{OH}^- \)-coupled electron transfer steps,\(^{55–57}\)

\[
\text{OH} + \text{OH}^- \leftrightarrow \text{O} + \text{H}_2\text{O} + e
\]
The reaction mechanism expressed in eqs 3–7 can be considered a specific case of the general mechanism, with eq 14 corresponding to eq 3, eq 15 to eq 4, eq 16 to eq 5, and eq 17 to the combination of eqs 6 and 7 because the NiO state is highly energetically unstable.

For the sake of generality, the microkinetic model is formulated for the mechanism expressed in eqs 14–17. The reaction rates are given by

\[ v_i = k_i^f[\text{OH}^-]^f \theta_{\text{OH}} - k_i^b \theta_{O} \]  
\[ v_i = k_i^f[\text{OH}^-]^f \theta_{\text{OH}} - k_i^b \theta_{O} \]  
\[ v_i = k_i^f[\text{OH}^-]^f \theta_{\text{OH}} - k_i^b \theta_{O} \]  
\[ v_i = k_i^f[\text{OH}^-]^f \theta_{O} - k_i^b \theta_{\text{OH}} \]  
\[ v_i = k_i^f[\text{OH}^-]^f \theta_{O} - k_i^b \theta_{\text{OH}} \]  

Here, [OH\(^-\)\(^f\)] represents the concentration of OH\(^-\) at the reaction plane, which is to be determined from the EDL model; \( k^f \) represent the forward (+) and backward (–) rate constants of the \( i \)th step, which are dependent on the local reaction condition provided by the EDL model; \( \theta_{\text{OH}}, \theta_{O}, \theta_{\text{OH}} \) and \( \theta_{O} \) represent the coverages of reaction intermediates constrained by the conservation of surface sites,

\[ \theta_{\text{OH}} + \theta_{O} + \theta_{\text{OH}} + \theta_{O} = 1 \]  

According to transition state theory, the rate constants are given by

\[ k_i^\pm = \frac{k_B T}{h} \exp \left( -\frac{G_i^\pm}{k_B T} \right) \]  

where the activation barriers are related to the Gibbs free energies of reactions according to the Bronsted–Evans–Polanyi (BEP) relation,

\[ G_i^0 = G_i^\pm + \beta \Delta \Delta G_i \]  

where \( G_i^0 \) is the activation barrier under standard equilibrium conditions (298 K, pH = 0, 1 bar), denoted by a superscript “\( \Theta \)”. \( \Delta \Delta G_i \) represents the variation in the Gibbs free energies of reactions when the reaction condition deviates from the standard equilibrium conditions. We shall also specify the coverages of adsorbates under any reaction condition, subject to surface reactions, and 

\[ \Delta \Delta G_i = \theta_{\text{OH}} + \theta_{O} + \theta_{\text{OH}} + \theta_{O} = 1 \]  

Under any reaction condition, \( \Delta \Delta G_i \) is given by

\[ \Delta \Delta G_1 = -\varepsilon_0(\phi_M - \phi' - U_0^\Theta) + \Delta \Delta G_\text{O} - \Delta \Delta G_{\text{OH}} \]  
\[ \Delta \Delta G_2 = -\varepsilon_0(\phi_M - \phi' - U_0^\Theta) + \Delta \Delta G_{\text{OH}} - \Delta \Delta G_\text{O} \]  
\[ \Delta \Delta G_3 = -\varepsilon_0(\phi_M - \phi' - U_0^\Theta) + \Delta \Delta G_{\text{O}} - \Delta \Delta G_{\text{OH}} \]  
\[ \Delta \Delta G_4 = -\varepsilon_0(\phi_M - \phi' - U_0^\Theta) + \Delta \Delta G_{\text{OH}} - \Delta \Delta G_\text{O} \]  

where \( \phi_M \) represents the electrode potential on the SHE scale and \( \phi' \) is the potential at the reaction plane in solution relative to the potential in the solution bulk which is taken as the potential reference. \( U_0^\Theta \), the standard equilibrium potentials of the reaction steps in eqs 14–17 versus the RHE, are given by

\[ U_0^\Theta = \frac{1}{e_0} \left( E_{\text{O}-\text{Ni}-\text{O}} - E_{\text{O}-\text{Ni}-\text{OH}} - \frac{1}{2} \mu_{\text{H}_2}^0 \right) \]  
\[ U_0^\Theta = \frac{1}{e_0} \left( E_{\text{O}-\text{Ni}-\text{OH}} - E_{\text{O}-\text{Ni}-\text{O}} - \mu_{\text{H}_2}^0 - \frac{1}{2} \mu_{\text{H}_2}^0 \right) \]  

based on eqs 9–13.

\( \Delta \Delta G_i \) in eqs 25–28 accounts for the variation in Gibbs free energies of the reaction intermediates OH, O, OOH, and OO when \( \phi_M \) changes. There are multiple reasons for such variations. On the one hand, \( \theta_{\text{OH}}, \theta_{O}, \theta_{\text{OH}} \) and \( \theta_{O} \) change with \( \phi_M \) so the Gibbs free energy of adding a new intermediate changes due to lateral interactions of the new intermediate with existing ones. On the other hand, the local electric potential and electric field change with \( \phi_M \), altering the Gibbs free energy of intermediates that carry a net charge. The Frumkin adsorption isotherm uses an empirical coefficient, called the lateral interaction coefficient, to describe the linear relation between \( \Delta \Delta G_i \) and \( \theta_x \). The linear relation is a first approximation. In addition, the Frumkin adsorption isotherm usually neglects interactions among adsorbed intermediates of different kinds. Norskov and co-workers expressed \( \Delta \Delta G_i \) as a function of the electric field, whose coefficients are fitted from DFT calculations. In an empirical sense, the Frumkin adsorption isotherm and the relation of Norskov and co-workers are equivalent, as there exists a one to one correspondence between \( \theta_x \) and the electric field, both of which are uniquely determined by \( \phi_M \). Herein, we use

\[ \Delta \Delta G_j = \mu_X E_{\text{AP}} \]  

where \( \mu_X \) being an empirical dipole moment of intermediate X and \( E_{\text{AP}} \) being the electric field strength at the adsorbate plane (AP). As will be made clear in the EDL model, \( E_{\text{AP}} \) is co-determined by all existing adsorbed intermediates. Therefore, eq 33 effectively considers interactions among adsorbed intermediates of all kinds.

At steady state, we have

\[ v_i = v_3 = v_4 = v_5 = i_{\text{OER}} \]  

where \( i_{\text{OER}} \) is the current density of the OER and \( \rho \), the number density of active sites.

We can solve for the coverages,

\[ \theta_{\text{OH}} = \frac{1}{\Xi} \left( \frac{1}{k_4} + K_4 K_5 K_4 K_3 k_2 + K_4 K_5 K_4 K_3 k_1 \right) \]  
\[ \theta_{O} = \frac{1}{\Xi} \left( \frac{K_1}{k_4 k_3} + \frac{K_4 K_5 K_4 K_3}{k_2} \right) \]  
\[ \theta_{\text{OH}} = \frac{1}{\Xi} \left( \frac{1}{k_2} + \frac{K_1}{k_4 k_3} + \frac{K_4 K_5 K_4}{k_2} \right) \]  
\[ \theta_{O} = \frac{1}{\Xi} \left( \frac{1}{k_3} + \frac{K_1}{k_4 k_3} + \frac{K_4 K_5 K_4}{k_2} \right) \]  

where \( K_i = k_i^+ / k_i^- \) and \( k_i^- = k_i^+ [\text{OH}^-] \), and \( \Xi = 1 + K_1 + K_4 K_5 K_4 K_3 + \frac{1}{k_4} + \frac{1}{k_3} + \frac{1}{k_2} + \frac{1}{k_1} \)

The current density is transformed to,

\[ i_{\text{OER}} = \frac{4e_0}{k_4^*} \left( \frac{\theta_1}{k_4^*} + \frac{\theta_2}{k_2^*} + \frac{\theta_3}{k_3^*} + \frac{\theta_4}{k_4^*} \right) \]  

with the thermodynamic factors...
Electric Double Layer Model

We require an EDL model to compute the local reaction conditions, including \( \phi_i \) [OH\(^{-}\)] and \( E_{AP} \). The mean-field Helmholtz free energy per unit volume of the electrolyte solution is written as\(^{51,52} \)

\[
f_{es} = c_i \phi (N_i - N_i^c) - \frac{e_i^c E^2}{2} - \frac{1}{\beta} N_i \ln \left( \frac{\sinh(pE\beta)}{pE\beta} \right) + \frac{1}{\beta} \ln \frac{N_i!}{N_i!N_i^c!} \tag{39}
\]

where \( c_i \) is the elementary charge, \( e_i^c \) is the optical component of the dielectric constant, \( \phi \) is the potential, \( E = V \phi \) is the electrical field, \( \beta = 1/k_BT \) is the inverse thermal energy, \( p \) is the dipole moment of solvent molecules, \( N \) is the total number density of lattice sites, and \( N_i \) \((i = s, a, c)\) are the number densities of solvent molecules (s), anions (a), and cations (c). On the right-hand side of eq 39, the first term is the electrostatic free energy of ions, the second term is the self-energy of the electric field, the third term is the electrostatic free energy of solvated molecules, and the last term is the entropic free energy related to the configuration of solution species, which is calculated using a lattice-gas approach. The conservation of lattice sites leads to \( N_i = N - N_i^c - N_c \).

Based on the variational principle, we obtain the following Euler–Lagrange equation in terms of \( \phi \),

\[
\frac{\partial f_{es}}{\partial \phi} - \frac{\partial}{\partial \epsilon} \left( \frac{\partial f_{es}}{\partial \epsilon} \right) = 0 \tag{40}
\]

leading to a modified Poisson–Boltzmann (PB) equation,

\[
-\frac{\partial}{\partial \epsilon} \left( \epsilon_0 + \frac{e_i^c L(E)}{k_BT} \right) E = e_i (N_i - N_i^c) \tag{41}
\]

where \( L(x) = \coth(x) - 1/x \) is the Langvin function. Compared with the conventional PB equation, eq 41 features an electric field-dependent local dielectric constant. Equation 41 has been derived in the dipolar Poisson–Boltzmann model by Abrashkin et al.,\(^53\) and the Langvin–Poisson–Boltzmann model by Gongadze and Iglic.\(^54\)

The distributions of cations and anions are obtained from

\[
\frac{\partial f_{es}}{\partial N} = \frac{\partial f_{es}}{\partial N_i} \tag{42}
\]

\[
\frac{\partial f_{es}}{\partial N} = \frac{\partial f_{es}}{\partial N_i^c} \tag{43}
\]

Equations 42 and 43 say that electrochemical potentials of cations and anions in the EDL are uniform and thus equal to their bulk values. Substituting eq 39 into eqs 42 and 43 leads to one of

\[
N_i = \frac{N \chi^c \exp(-\beta \epsilon_i \phi_i)}{\chi_s \exp(-\beta \epsilon_i \phi_i) + \chi_a \exp(\beta \epsilon_i \phi_i) + (1 - 2\chi^c) \sinh(\beta \epsilon_i \phi_i) / \beta \epsilon_i \phi_i} \tag{44}
\]

where \( \chi \) is the bulk number density of cations/anions normalized to \( N_i \), and \( \gamma_c \) and \( \gamma_a \) denote relative sizes of solvated anions and cations, respectively, calculated as

\[
\gamma_{x,c} = \left( \frac{d_{x,c}}{d_x} \right)^3 \tag{46}
\]

with \( d_{x,c} \) being the effective diameters of solvated anions and cations and \( d_x \) being the diameter of solvent molecules, which is taken as the reference size of the lattice. We stress that eqs 44 and 45 take into account the ion size effect in a phenomenological manner, namely, by introducing two size factors \( \gamma_c \) and \( \gamma_a \). A more rigorous treatment of the size effect can be found in ref 65.

Substituting eqs 44 and 45 into eq 41 leads to a single differential equation governing the distribution of electric potential. Its boundary conditions are as follows:

\[
\phi = 0 \tag{47}
\]

in the solution bulk and

\[
\phi = \phi_M = \phi_{pzc} + \delta_A E_{AP} \tag{48}
\]

at the adsorbate plane (AP), which is designated as the coordinate origin, \( x = 0 \), of the EDL model. The space between the electrode surface and the adsorbate layer has a thickness of \( \delta_A \), a permittivity of \( \epsilon_A \), and an electric field of \( E_{AP} \). \( \phi_{pzc} \) is the potential of zero charge. The third term on the right-hand side (RHS) of eq 48 represents the potential drop across the adsorbate layer, where the uniform electric field \( E_{AP} \) is given by\(^{66,67} \)

\[
E_{AP} = \frac{\epsilon}{\epsilon_A} \frac{\partial \phi}{\partial x} + \frac{\mu_{chem}}{\epsilon_A \mu_{chem}} \tag{49}
\]

where the first term on the RHS represents the electric field caused by the excess free charge on the electrode surface and the second term represents the electric field generated by the dipole moment formed between charged intermediates and its compensating charge in the electrode, denoted by \( \mu_{chem} \).\(^{66,67} \)

\[
\mu_{chem} = \sum_{x=OH-OH,OOH} \frac{\xi_X}{\delta AP} \tag{50}
\]

where \( \xi_X \) is the net charge number of adsorbed intermediates.

Model Parameters and Numerical Implementation

Table 1 lists the model parameters which are categorized into five subgroups termed “general constants,” “electrode,” “electrolyte solution,” “interfacial structure,” and “reaction.” The electrode parameters correspond to \( \beta \)-NiOOH (0001). The (2 \times 2) supercell has a surface area of \( S_e = (5.85 \times 10.24 \text{ Å}) = 59.90 \text{ Å}^2 \), according to the DFT calculation. Therefore, the number density of reaction sites equals \( \rho = \frac{1}{S_e} = 6.67 \times 10^7 \text{ m}^{-2} \). The work function of the slab surface with one layer of water molecules is calculated to be \( \Phi = 5.6 \text{ eV} \).\(^{44} \) Therefore, the pzc versus the SHE is calculated as \( \phi_{pzc} = \Phi - 4.44 = 1.16 \text{ V}_{SHE} \), assuming that the absolute electrode potential of the SHE is 4.44 eV.

The electrolyte parameters correspond to 0.1 M MOH (M = Li, Na, Cs). From the expression of the effective dielectric constant given in eq 41, we obtain the bulk dielectric constant as \( \epsilon_b = \epsilon_\infty + \frac{1}{3} \frac{N_i^c}{\epsilon_b} \). Therefore, the effective dipole moment of water molecules is found as \( \mu_{H_2O} = \frac{\epsilon_b}{\epsilon_\infty} = 4.44 \times 1.16 \text{ V}_{SHE} \), assuming that the absolute electrode potential of the SHE is 4.44 eV.
Table 1. Model Parameters for the Base Case

| general constants | electrode | electrolyte solution |
|-------------------|-----------|----------------------|
| $k_B$, Boltzmann constant | 1.38 × 10^{-23} J/K | DFT calculation |
| $e_e$, elementary charge | 1.6 × 10^{-9} C | |
| $N_A$, Avogadro constant | 6.02 × 10^{23}/mol | |
| $e_0$, vacuum permittivity | 8.85 × 10^{-12} F/m | |
| $e_p$, permittivity of bulk water | 78.5$e_0$ | |
| $h$, Planck constant | 6.626 × 10^{-34} J/s | 298.15 K |
| $T$, temperature | | |
| $\rho_r$, the number density of reaction sites | 6.67 × 10^{13}/m^2 | |
| $\Phi$, work function of the solvated electrode | 5.6 eV | DFT calculation |
| $\phi_{\text{pzc}}$, potential of zero charge | $-4.44 + \ln\frac{1}{10}M\text{pH}^+$ | absolute electrode potential of the SHE is 4.44 eV |
| $d_r$, diameter of water molecules | 2.75 Å | |
| $d_{eff}$, effective diameter of solvated anions | 6.0 Å | |
| $d_{solv}$, effective diameter of solvated cations | 7.6, 7.2, 6.6, 6.6 Å for Li^+, Na^+, K^+, Cs^+ | |
| $\epsilon_{opt}$, optical permittivity | $\epsilon_0$ | |
| $p$, effective dipole moment of water molecules | | |
| $\epsilon_{b}$, bulk concentration of water | 55.5 M | |
| pH | 13 | |
| $\epsilon_{b,c}$, bulk concentration of anions/cations | 0.1 M | |
| $N$, total number density of lattice sites | $(\epsilon'_c + \epsilon'_s)/N$ | |
| $\gamma$, bulk number density of cations/anions normalized to $N$ | $\gamma_{b}/N$ | |
| interface structure | | |
| $\epsilon_{AB}$, permittivity of the adsorbate layer | $4\epsilon_0$ | |
| $\delta_{AB}$, thickness of the adsorbate layer | 1.5 Å | |
| reaction | | |
| $U_{eq}^{f}$, standard equilibrium potentials of the reaction steps versus RHE | 1.92, 1.22, 1.18, 0.60 V for eqs 14−17, respectively | DFT calculation |
| $\beta_{eff}$, transfer coefficient | 0.5 | |
| $\epsilon_{b,c}^{eq}$, activation barrier under the standard equilibrium conditions | 0.50, 0.33, 0.31, 0.44 eV for eqs 14−17, respectively | fitted |
| $\mu_{X}$, empirical dipole moment of intermediate $X$ | $\xi_{X}/\epsilon_{AB}$ | fitted |
| $\xi_{X}$, net charge number of adsorbed intermediates | $-0.06, -0.01, -0.04, -0.04$ | fitted for OH, O, OOH, OO, respectively |

7.6, 7.2, 6.6, and 6.6 Å for hydrated Li^+, Na^+, K^+, and Cs^+, respectively. The adopted values are close to the Stokes radii of hydrated rigid ions in water from ionic conductivity measurements. As a first approximation, we assume that all adsorbates line up rigidly in the adsorbate layer and are equidistant from the electrode surface. The thickness of the adsorbate layer is $\delta_{AB} = 1.5$ Å as the mean value of the sizes of OH, O, OOH, and OO. The permittivity of the adsorbate layer is usually assumed to be a multiple of the vacuum permittivity, and here we inherit $\epsilon_{AB} = 3\epsilon_0$ from ref 67. The standard equilibrium potentials of the reaction steps versus the RHE are calculated from eqs 29−32 with $E_{O_{2}/O_{2}}$, $E_{O_{2}/OH}$, $E_{OCOO}$, $E_{OC}$, $\mu_{H2O}$, $\mu_{H}$, and $\mu_{O}$ obtained from DFT calculations. The Kohn–Sham DFT approach implemented with PBE and RPBE exchange-correlation functionals has a typical root-mean-square deviation of $\sim0.3$ eV for chemisorbed systems. 76 The error shall grow in electrochemical systems with the presence of the electrolyte solution. In addition, we have neglected the zero-point energy corrections; see the discussion in section S3 in the SI. Therefore, the values of $U_{eq}^{f}$ calculated with DFT results are allowed to vary in the range of $\sim0.3$ V. The transfer coefficients of the reaction steps are taken as $\beta_{eff} = 0.5$. Microscopically, $\beta_{eff}$ shall vary as a function of $\phi_{\text{pzc}}$ and vary from reaction to reaction. In ref 70, Huang derived an analytical expression for $\beta_{eff}$ where different factors are discriminated. However, since we use the phenomenological Butler–Volmer equations for the elementary step kinetics, we should not be bothered by mechanistic complexities of $\beta_{eff}$. The nudged elastic band (NEB) method can be used to calculate activation barriers of electrocatalytic reactions. Herein, we choose to fit the activation barriers from experimental polarization data, considering that the error in DFT calculations at the transition state should be larger than that at the ground state, namely, $\sim0.3$ eV. The net charge numbers of adsorbed intermediates $\xi_{X}$ are also fitted from experimental data. The fitting details are provided in section S1 of the SI. The sensitivity of the fitted parameters will be gauged at a latter point in the Discussion section. The numerical implementation of the model was performed in MATLAB. The code script is available upon request. We note that all potentials are rescaled to the RHE in the model calculation.

### RESULTS

In this section, we describe how the local reaction conditions are shaped by the cations in the first subsection, which lays the basis for understanding the cation effects on the OER in the second subsection. Afterward, we discuss other means to tune the cation effects, including the solution pH in the third subsection and bivalent cations in the last subsection.

#### Local Reaction Conditions

The local reaction conditions computed by the model are shown in Figure 2, where the effective diameter of the cations $d_{eff}$ is varied between 4 and 8 Å with a step of 0.4 Å and 11 lines are included in each figure. Figure 2a–c exhibits the spatial distributions of OH$^-$ concentration ([OH$^-$]), cation concentration ([C$^+$]), and dielectric constant ($\epsilon$) in the EDL at $\phi_{\text{pzc}} = 1.6$ V$_{\text{RHE}}$, respectively. Let us first analyze general trends and then dissect differences among different lines corresponding to different $d_{eff}$.

OH$^-$ decreases from the bulk solution to the electrode surface because the surface charge density $\sigma$ is negative even at this high electrode potentials (see Figure 2d) and OH$^-$ ions are repelled as co-ions away from the electrode surface. $\sigma$ is negative in the considered potential range because the pzc is $\sim1.93$ V$_{\text{RHE}}$ for pH = 13, which is consistent with the fact that the point of zero charge of Ni(OH)$_2$ is $\sim11$ (namely, the solid will be negatively charged at pH $> 11$). Consequently, cations are attracted as counterions to the negative surface charge, resulting in an increasing profile of [C$^+$] toward the electrode surface. In addition, $\epsilon$ decreases toward the electrode surface, which is ascribed to the combined effect of polarization saturation due to the high interfacial electric field and density decrement due to the crowding of counterions. The monotonically decreasing trend of $\epsilon$ in the EDL has been revealed by Nakayama and Andelman, and Gongadze and Iglisch. In a previous work, Huang, Chen, and Eikerling showed nonmonotonic profiles of $\epsilon$ in the EDL due to the competition between ions and solvent molecules. The nonmonotonic behavior is more pronounced in the presence of specific ion adsorption.

As $d_{eff}$ increases, fewer cations can accumulate in the EDL and [C$^+$] decreases accordingly, as shown in Figure 2b. Moreover, the black line corresponding to $d_{eff} = 8$ Å levels off.
near the electrode surface, signifying the overcrowding phenomenon of cations. Due to the intensified steric repulsion of cations, both $[\text{OH}^-]$ and $\epsilon$ decrease for larger $d_c$; see Figure 2a and c, respectively. From now on, the cation overcrowding effects are referred to as effects caused by the substantial accumulation of cations near the electrode surface due to the negative surface charge, including decreased $[\text{OH}^-]$, lower $\epsilon$ and among others.

Figure 2d–f exhibits the variation of surface charge density ($\sigma$), $\text{OH}^-$ concentration at $x = 0$ ($[\text{OH}^-]$), and electric potential at $x = 0$ ($\phi'$) as a function of $\phi_{\text{M}}$, respectively. $\sigma$ increases toward positive values and $[\text{OH}^-]$ grows with increasing $\phi_{\text{M}}$. The larger the cation size, the smaller the magnitude of $\sigma$. The dependence of $\sigma$ on the cation size is readily understood from the effective thickness of the diffuse layer, which becomes thicker for larger cations, resulting in a smaller magnitude of $\sigma$. A similar analysis is provided by Ringe et al. for cation effects in carbon dioxide reduction. For the case of a larger $d_c$, $[\text{OH}^-]$ decreases due to greater steric repulsion because the size factor $\gamma_c$ is cubic with $d_c$, despite the decreasing magnitude of $\sigma$. Figure 2e shows that the local electric potential $\phi'$ follows the positive-shift trend of $\sigma$ with increasing $\phi_{\text{M}}$ and grows in magnitude with increasing $d_c$ due to diminished screening effect of water molecules, Figure 2c.
Model-Experiment Comparison

With a knowledge of cation effects on the local reaction conditions, we proceed to analyze cation effects on the polarization curve, namely, the voltage−current relation. Figure 3a shows that the exponential growth in $j_{\text{OER}}$ with increasing $\phi_M$ is deferred for Li$^+$ compared with Cs$^+$, which is consistent with experimental data. However, we also notice deviations in the details of the polarization curves between the model and experiments. Possible causes for the deviations are many, including the deviation between the idealized $\beta$-NiOOH(0001) slab and the actual surface states of the oxide, the deviation between the phenomenological Butler−Volmer equation and the complicated electron transfer kinetics, and the neglect of mass transport effects in this model.

In Figure S2, we compare the polarization curves of OER at NiOOH in purified 0.1 M OH (M = Li$^+$, Na$^+$, Cs$^+$) measured by two laboratories. Though the activity trend of Cs$^+$ > Na$^+$ > Li$^+$ is shared by both, noticeable deviations in the magnitude of the current density and the detailed profiles of the polarization curves are found. Nevertheless, the deviations in details of the polarization curves between the model and experiments in Figure 3 and between two experimental studies in Figure S2 shall not prevent us from understanding the trend of cation effects on the OER.

The microkinetic model enables us to gain deep insights into the reaction mechanism. Figure 3c shows the variation of different resistance terms $\Theta_i^{\kappa_i}$ (i = 1−4) in eq 37 as a function of $\phi_M$. The rate-determining term (RDT) transitions from the third term $i = 3$ to the second term and then to the first term as $\phi_M$ increases. The RDT above 1.6 $V_{\text{SHE}}$, namely, the first term, grows in the order Cs$^+$ < Na$^+$ < Li$^+$ (Figure 3d), leading to the catalytic activity decreasing in the order Cs$^+$ > Na$^+$ > Li$^+$ (Figure 3a). The reason is simply that larger cations occupy more volume near the negatively charged surface, leading to a smaller concentration of OH$^-$, the reactant (Figure 2a). In Figure S1, we analyze an extreme case where the activation energy of the first step is increased to 0.98 eV, around 0.6 eV higher than other steps, to provide the readers an example of how this model behaves in extreme cases.

pH Effects

For proton-coupled electron transfer (PCET) reactions, under thermodynamic equilibrium conditions, the reaction should be invariant on the RHE scale. Therefore, any observed pH effects on the RHE scale are usually ascribed to decoupled proton and electron transfer reactions. Note that our current model goes beyond such a purely thermodynamic analysis. We will show below that microkinetic considerations influenced by double-layer effects bring new insights into the pH effects in the OER.

Figure 4a shows that the OER activity on the RHE scale increases with increasing pH, consistent with experimental data. The total ionic strength is 1 M for all pH values considered. The pH-dependent trend in activity contrasts with a thermodynamic analysis that predicts a pH-independent activity on the RHE scale as all elementary steps are PCET reactions in our model. The discrepancy lies in the EDL effects. Figure 4b shows that $\sigma$ is more negative at higher pHs, which can be effectively understood by the fact that the pzc on the RHE scale increases with pH. Therefore, the cation overcrowding effect is more pronounced at higher pHs,
resulting in a lower \( [\text{OH}^-]' \) at potentials below 1.6 \( V_{\text{RHE}} \), even though the bulk \( [\text{OH}^-] \) concentration is greater at higher pHs. \( \sigma \) is less negative with increasing \( \phi_M \), alleviating the cation overcrowding effect and leading to larger \( [\text{OH}^-]' \) for higher pHs at potentials above 1.6 \( V_{\text{RHE}} \). Since the OER activity is limited by \( [\text{OH}^-]' \) that is dictated by the cation overcrowding effect, the OER activity thus grows with increasing pH at potentials above 1.6 \( V_{\text{RHE}} \), see Figure 4a.

**Bivalent Cation Effects**

Adding bivalent cations into the solution while retaining the same \( \text{OH}^- \) concentration, we observe a decrement in the OER activity, as shown in Figure 5a, which is consistent with the experimental observations.\(^{19,25}\) The underlying causes are related to the modifications to the EDL by the bivalent cations. Figure 5b exhibits the surface charging curves. Specifically, \( \sigma \) is slightly more negative above 1.6 \( V_{\text{RHE}} \) in the presence of \( \text{Ca}^{2+} \). Due to the larger size of \( \text{Ca}^{2+} \) (8.2 Å in diameter), \( [\text{OH}^-]' \) is lower in the presence of \( \text{Ca}^{2+} \), Figure 5c, leading to the lower OER activity in Figure 5a. Figure 5d and e show the spatial distributions of \( \text{OH}^- \) concentration and cation concentrations in the EDL at 1.6 \( V_{\text{RHE}} \). As bivalent cations are more effective in screening the electric field, the concentration of \( \text{Ca}^{2+} \) is greater near the electrode surface than that of \( \text{Cs}^+ \). Consequently, the \( \text{OH}^- \) concentration near the electrode surface is lower in the presence of \( \text{Ca}^{2+} \) (Figure 5d).

**DISCUSSION**

In this section, we first compare electronic and electrostatic factors regarding the cation effects. Then, we gauge the robustness of the model results in the vast parameter space by examining the sensitivity of the model output to the variation/uncertainty of the model parameters. Finally, we dissect the features and limitations of the model and discuss the extensions needed when applied to other catalysts and reactions.

**Electronic vs Electrostatic Factors**

The model considers both electronic and electrostatic factors. The former type is reflected in the binding energies under standard conditions and dipole corrections, namely, the \( \Delta \Delta G_i \) terms. The latter type includes local electric potential and concentrations that are influenced by electrostatic interactions. The critical property governing the electrostatic interactions is the surface charge density, which is modified by chemisorption-induced surface dipoles (\( \mu_{\text{chem}} \)). In this section, we compare the importance of electronic and electrostatic factors for cation effects in the OER.

In Figure 6a, we compare three cases, including the base case, the second case without dipole corrections for the binding energies (\( \Delta \Delta G_i = 0 \)), and the third case without chemisorption-induced surface dipoles. The x-axis is the potential of zero charge. The y-axis is the serial number of simulations in which \( U_1^0, U_2^0, U_3^0 \) are allowed to vary randomly within 0.2 V around their base values, with \( U_4^0 = 4.92 - U_1^0 - U_2^0 - U_3^0 \).
for \( d_c = 8 \) Å. All curves show that the OER current density decreases as \( d_c \) increases. The curves for the base case and the second case of \( \Delta \Delta G_i = 0 \) are indistinguishable, while the curve for the third case of \( \chi_i = 0 \) is markedly suppressed. This indicates that \( \Delta \Delta G_i \) is unimportant here and the cation size effect is mainly ascribed to the surface charging effects. The surface charging relation for the third case of \( \chi_i = 0 \) is shown in Figure 6b. Compared with Figure 2d for the base case, Figure 6b exhibits a reduced difference of \( \sigma \) for different \( d_c \)'s, thus leading to a much reduced \( d_c \)-dependence of the OER current in Figure 6a. In addition, compared with the base case, \( \sigma \) has a smaller magnitude for \( \chi_i = 0 \), because negatively charged adsorbates increase the pzc of the electrode. \( \chi_i \) As a result, the cation overcrowding effect is diminished for the third case of \( \chi_i = 0 \).

Our argument that the cation effect on the OER is mainly caused by surface charging effects instead of the shift in the field-dependent binding energies is further supported by the results presented in Figure 6c. The x-axis is the potential of zero charge which dictates the surface charging relation. The y-axis is the serial number of Monte Carlo simulations in which \( U_i^0, U_j^0, U_{i,j}^0 \) are allowed to vary randomly within 0.2 V around their base values. \( U_j \) is given by the constraint condition that \( U_j^0 = 4.92 - U_i^0 - U_{i,j}^0 - U_{j,k}^0 \). It is clearly shown that the OER current ratio between \( Cs^+ \) and \( Li^+ \) at 1.7 V SHE, denoted as \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \), increases significantly when the pzc grows, namely, when the electrode surface gets more negatively charged. On the contrary, \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \) is slightly affected by random variations in the equilibrium potentials of the elementary steps which are determined by the binding energies of the adsorbates. The insensitivity of the cation effects to binding energies is in accord with the robustness of the cation effects observed in a wide range of catalysts in experiments, including Ni(Fe)OOH, RuO\(_2\), Au, IrO\(_2\), and perovskites. \( J_{\Delta \Delta G_i} \)

Continuing in our line of reasoning, we deduce that the effect of cation size will be less significant for catalysts with a lower pzc. Titanium dioxide (TiO\(_2\)) has a work function of 5.4 eV, around 1 eV lower than that of nickel oxides. Based on the linear relation between the work function and the pzc, TiO\(_2\) shall carry a positive charge on its surface during the OER. Therefore, if electrostatic factors, more precisely, the surface-charge modulated cation overcrowding effects, dominate, we expect that the cation size effect on the OER is much weaker at TiO\(_2\) compared with NiO. A systematic comparison of the cation effects on different catalysts with different work functions is vital to examine the importance of surface charging effects in the cation effects on the OER.

**Parameter Sensitivity and Robustness of the Model Results**

Given the significant number of parameters in our model, we shall dissect the robustness of the model results against uncertainties and variations in model parameters. To this end, a parameter sensitivity study is conducted. Among model parameters in Table 1, those having significant uncertainties and variations include the pzc, the equilibrium potentials \( U_i^0 \) (equivalently, the binding energies), the activation barriers \( G_{\alpha \delta}^{\sigma \iota} \) the transfer coefficients \( \beta_i \), and the net charge numbers of adsorbed intermediates \( \xi_s \). In the preceding subsection, we have shown that the cation effects are highly sensitive to the pzc and robust against variations in \( U_i^0 \). In this subsection, we complement the parametric analysis by examining the influence of \( G_{\alpha \delta}^{\sigma \iota} \), \( \xi_s \) and \( \beta_i \). The output of the model used for the sensitivity analysis is the current density ratio between \( Cs^+ \) and \( Li^+ \) at 1.7 V SHE, denoted as \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \). Three groups of Monte Carlo simulations were conducted. Each group consists of 100 runs of the model. In each run, the four \( G_{\alpha \delta}^{\sigma \iota} \)'s are varied randomly in the range of [0.3, 0.6] eV in the first group, the four \( \xi_s \)'s are varied randomly in the range of \([-0.1, 0]\) in the second group, and \( \beta_i \) is changed randomly in the range of [0.3, 0.7] in the third group.

The histogram of the Monte Carlo simulation is exhibited in Figure 7. All data points of \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \) in the first group are above 2, indicating that the trend of higher OER activity at smaller \( d_c \) is robust against variations in \( G_{\alpha \delta}^{\sigma \iota} \). The data points in the second group are more dispersed, indicating that \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \) is highly sensitive to the four \( \xi_s \)'s because they determine the surface charging relation and thus the cation overcrowding phenomenon, as has been analyzed for the third case in Figure 6a. This gives further evidence to our claim in the preceding subsection that alkali metal cations influence the OER activity mainly by modulating the local reaction conditions determined by the surface charge density. The distribution of data points in the third group is much smaller, indicating that \( \frac{J_{\gamma_{V_{HHE}}}(Cs^+)}{J_{\gamma_{V_{HHE}}}(Li^+)} \) is less sensitive to the transfer coefficients.

**Features, Limitations, and Extensions**

In this subsection, we first dissect the features of the presented model compared with existing models. Then, we address the question: to what extent are the insights presented here valid for other cases. This requires us to scrutinize assumptions and limitations of the model. Afterward, we discuss modifications and extensions needed to implant the model for more general cases.

Taking a broader view, combining DFT calculations and microkinetic considerations in modeling electrocatalytic reactions is becoming increasingly popular. \( J_{\Delta \Delta G_i} \) Some studies...
have used mean-field EDL models to compute the local reaction condition for the ORR and carbon dioxide reduction. As for the OER, our model is, to the best of our knowledge, the first one that combines DFT calculations, microkinetic considerations, and an EDL model. Moreover, compared with similar models that were developed for other reactions, this model has two unique features. First, the EDL model considers nonlinear solvent polarization and ion size effect. Second, chemisorption-induced surface dipoles, significantly influencing the surface charging relation, are considered.

The assumptions and approximations of the model are fivefold. First, the microkinetic model is developed based on the OER mechanism consisting of four OH−-coupled electron transfer steps expressed in eqs 14−17. Second, the phenomenological Butler−Volmer equation is used to describe electron transfer kinetics. Third, the EDL is considered as a serial connection of a charged adlayer and a diffuse layer and treated at the mean field level for the one-dimensional case. Fourth, the electrolyte cations are restricted to be nonspecifically adsorbed. Fifth, macroscopic mass transport effects are neglected.

On the first point, the used reaction mechanism is common to many OER catalysts, though the active sites, the binding energies, and the activation energies are case-specific. We have demonstrated in the preceding subsection that our results are robust against variations in these parameters. For OER catalysts with decoupled OH− and electron transfer steps, the microkinetic model can be modified without major difficulties.

On the second point, the Butler−Volmer equation can be upgraded to more advanced electron transfer theories, however, at the cost of introducing additional microscopic parameters. For example, the Marcus−Hush−Chidsey theory requires the knowledge of interfacial solvent reorganization energies, and the Anderson−Newns model requires the knowledge of electronic interactions between the catalyst and the reactant in solution. Therefore, the Butler−Volmer equation best suits our purpose. In addition, the Marcus−Hush−Chidsey theory can be reduced to the Butler−Volmer equation with a potential-dependent transfer coefficient. Our sensitivity analysis shows that variations in the transfer coefficient do not alter the conclusion.

On the third point, the mean-field EDL model radically simplifies the atomistic picture of the EDL, which can be obtained from DFT-based first-principles calculations. Atomistic EDL models, such as in refs 86 and 87, can complement the mean-field EDL model by determining some of the model parameters, including the structure of the adlayer and the net charge numbers of adsorbed intermediates. We are aware that the ξ's are difficult to obtain accurately and depend on the theoretical method used. Through sensitivity analysis, we have shown that the cation effects are highly sensitive to ξ's.

On the fourth point, there is a controversy around whether electrolyte cations are nonspecifically adsorbed, specifically adsorbed, or even intercalated into the electrode material. As for alkali metal cations at a NiOOH surface, Garcia et al. showed that the electrochemical active surface area (ECSA) is not influenced significantly by electrolyte cations, invalidating a previous postulate that the cation-dependent activity is related to the ECSA change due to cation intercalation. The crystal truncation rod analysis of Rao et al. indicates that Li+ and K+ are nonspecifically adsorbed and retain their solvation shells near the RuO2 surface. This provides experimental evidence for the assumption of nonspecific adsorption in this model. For the case with specific adsorption of cations, its effect usually translates to a shift in binding energies. For such cases, we recall that the findings of the present model are robust against variations in binding energies, prompting us to conclude that the present model would remain applicable for the case of specific adsorption of cations.

On the fifth point, mass transport is one of the influencing factors of the polarization curves, and its importance is crucial in the high overpotential region where the current density is rather high. The presented study focused on the cation effect on the intrinsic catalytic activity in the low overpotential region. Therefore, the mass transport effects are secondary in this work. A comprehensive physicochemical model for electrocatalytic reactions considering multistep interfacial kinetics, EDL effects, and macroscopic mass transport effects has been developed recently.

## CONCLUSION

We have presented a physicochemical model for understanding how electrolyte cations influence the OER via shaping the local reaction conditions. The decreasing OER activity with larger cations was found to be correlated with cation overcrowding effects in the electric double layer. Specifically, the electrode surface is negatively charged due to the high pzc, resulting in cation accumulation and anion depletion near the electrode. Furthermore, cations with larger effective size occupy more volume, leading to smaller concentrations of OH− and thus lower OER activity. The same reasoning has been employed to understand why the OER activity increases with solution pH on the RHE scale and why the OER activity decreases in the presence of bivalent cations. The correction to the binding energies that depend on the local electric field plays a minor role in our case. A sensitivity analysis shows that the cation size effect is sensitive to properties related to the surface charging behavior, including the potential of zero charge and the net charge numbers of adsorbed intermediates. In addition, Monte Carlo simulations demonstrated that the key findings reported are robust against uncertainties and variations in binding energies, activation barriers, and transfer coefficients. This work adds to the understanding of how the local reaction environment influences electrocatalytic activities.

## ASSOCIATED CONTENT

 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00315.

 Model fitting details; results for an extreme case with one step having a much larger activation energy; zero-point vibrational corrections; projected magnetic moments for oxygen and nickel atoms; variations in experimental polarization curves on the same system measured by different laboratories (PDF)

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Notes

The authors declare no competing financial interest.

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