



Navigating Alkaline Hydrogen Evolution Reaction Descriptors for Electrocatalyst Design

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Abstract: The quest for efficient green hydrogen production through Alkaline Water Electrolysis (AWE) is a critical aspect of the clean energy transition. The hydrogen evolution reaction (HER) in alkaline media is central to this process, with the performance of electrocatalysts being a determining factor for overall efficiency. Theoretical studies using energy-based descriptors are essential for designing high-performance alkaline HER electrocatalysts. This review summarizes various descriptors, including water adsorption energy, water dissociation barrier, and Gibbs free energy changes of hydrogen and hydroxyl adsorption. Examples of how to apply these descriptors to identify the active site of materials and better design high-performance alkaline HER electrocatalysts are provided, highlighting the previously underappreciated role of hydroxyl adsorption-free energy changes. As research progresses, integrating these descriptors with experimental data will be paramount in advancing AWE technology for sustainable hydrogen production.

Keywords: green hydrogen; alkaline water electrolysis; descriptor; density functional theory; electrocatalyst design

1. Introduction

The shift towards hydrogen gas as an alternative energy source is a significant step in combating climate change. Green hydrogen via electrocatalytic water splitting is directly competitive with fossil fuels, which decreases overall greenhouse gas emissions [1]. The intricacies of electrocatalytic water splitting involve the hydrogen evolution reaction (HER) and the oxygen evolution reaction (OER) [2]. There are several types of water electrolyzers (WE): solid oxide water electrolyzer (SOWE), proton exchange membrane water electrolyzer (PEMWE), and alkaline electrolyzer (AWE) [3]. SOWE can be adapted for continuous operation in industrial areas under high temperatures. However, the technology has seen limited applications due to long start-up times, mechanical compatibility, and chemical instabilities [4,5]. PEMWE in acidic environments is the most efficient for green hydrogen production due to the abundance of the proton reactant in the solution [6]. However, the main challenge associated with PEMWE is the high cost of the electrocatalysts, which are based on platinum and platinum group metals (PGMs) to resist acid corrosion [7]. In contrast, alkaline HER in AWE offers a promising alternative allowing for the exploration of earth-abundant transition metal-based catalysts because their hydro-oxide counterparts are stable in this environment [8-10]. Shifting towards an alkaline environment will reduce reliance on expensive noble metals and reduce the hydrogen production cost. Green



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Copyright: © 2024 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). hydrogen via electrocatalytic water splitting is directly competitive with fossil fuels, which decreases overall greenhouse gas emissions.

The operational principle of AWEs involves the conduction of hydroxide anions through a liquid electrolyte between two electrodes [8]. The main challenges of the current AWE are its low current densities, the inability to operate with intermittent renewable energy sources, and slow kinetics [11]. These limitations are associated with early versions of atmospheric electrolyzers designed for continuous operation at industrial sites with stable and inexpensive electricity sources. Modern pressurized high-temperature AWEs have shown the capability to reach performance and adaptability on par with PEMWEs [12]. The adaptability to various energy sources, coupled with improved design and control systems, positions pressurized AWEs as a promising solution for sustainable and flexible hydrogen generation in the evolving energy technology landscape [12].

Moreover, the anion exchange membrane water electrolyzer (AEMWE) replaces the conventional diaphragm of AWE with an anion exchange membrane [13]. A membrane reduces the gas cross-over and the need for pressure difference operation between the anode and cathode. Additionally, the thickness of the anion exchange membrane is less than the diaphragm, which leads to a lower ohmic over-voltage. The AEMWE uses an alkaline solution with low concentration. Conventional electrolyzers use concentrated potassium hydroxide electrolytes, which increase corrosion. Therefore, AEMWE has several benefits, including no leakage, ease of installation, and control. However, the low energy conversion efficiency of AWEs due to the slow kinetics remains a bottleneck, requiring high-performance electrocatalyst design.

The pursuit of an effective catalyst for AWE is multifaceted, necessitating a blend of experimental and theoretical studies to unlock the full potential of electrocatalysts [8,11,14]. The challenges for the alkaline HER electrocatalyst design are twofold: the inherently lower reaction rates of catalysts and a limited understanding of the complex kinetics involved. The HER process varies significantly depending on the electrolyte medium, which can be either acidic or alkaline. In acidic media, the HER mechanism involves the electrochemical reduction of a proton (H⁺), leading to the release of hydrogen gas (H₂) [15–17]. The HER efficiency is associated with the adsorption of hydrogen atom intermediate (H^{*}), which consists of the Volmer/Heyrovsky or Volmer/Tafel steps as delineated below [18]:

Volmer reaction :
$$H^+ + e^- \rightarrow H^*$$
 (1)

Heyrovsky reaction : $H^* + H^+ + e^- \rightarrow H_2$ (2)

Tafel reaction :
$$H^* + H^* \rightarrow H_2$$
 (3)

Conversely, alkaline HER operates through the electrochemical reduction of water molecules, resulting in the formation of hydrogen gas and hydroxide ions (OH⁻) [19].

Water adsorption :
$$H_2O + * \rightarrow H_2O^*$$
 (4)

Volmer reaction (I):
$$H_2O^* + e^- \rightarrow H^* + OH^* + e^- \rightarrow H^* + OH^-$$
 (5)

Volmer reaction (II):
$$H_2O^* + e^- \rightarrow H^* + OH^-$$
 (6)

Heyrovsky reaction :
$$H_2O + e^- + H^* \rightarrow H_2 + OH^-$$
 (7)

Tafel reaction :
$$H^* + H^* \rightarrow H_2$$
 (8)

The HER pathway in alkaline media involves complex Volmer-Heyrovsky or Volmer-Tafel steps, with the intermediate hydrogen formed after an initial water adsorption step (Equation (4)). Additionally, the adsorbed hydroxyl intermediate may form during the Volmer step if its binding energy with the active site is strong enough (Equation (5)). Otherwise, the solution-phase hydroxide (OH⁻) can directly form after the water dissociation (Equation (6)) during the Volmer reaction. The H_2 product can be produced through an adsorbed H atom intermediate and water in the Heyrovsky reaction (Equation (7)) or through the combination of two adsorbed H* atom intermediates in the Tafel reaction (Equation (8)). Figure 1 indicates the possible alkaline HER pathway. It shows that the performance of alkaline HER can be influenced by four main factors: (1) water adsorption strength on active sites, (2) water dissociation ability, (3) hydrogen binding energy, and (4) OH adsorption ability. To this end, the properties relevant to these four factors have been used to screen the alkaline HER electrocatalysts.



Figure 1. Possible hydrogen evolution reaction processes in alkaline media.

The step-by-step process that governs HER needs to be theoretically revealed by delving into the intricate details of reaction mechanisms through atomic-scale simulation [8]. Theoretical investigations can offer insights into the energetic aspects of water adsorption, water dissociation, intermediates adsorption, and hydrogen desorption [15,20–22]. This granular view allows researchers to identify and optimize the performance of active sites that influence both catalytic efficiency and stability. Such knowledge is instrumental in recognizing the rate-determining steps and potential energy barriers, which are critical for designing catalysts with optimal performance [15]. Furthermore, theoretical models serve as a blueprint for predicting the performance of different materials under various reaction conditions. Therefore, computational studies enable the fine-tuning of working conditions to enhance the overall efficiency of the process. Additionally, these studies contribute significantly to the green hydrogen field by offering a cost-effective strategy for material screening, circumventing the need for laborious and expensive experimental procedures [20,23]. Ultimately, the insights gained from theoretical research facilitate the development of innovative solutions for alkaline hydrogen production [8].

Recent advances have highlighted the importance of atomically precise electrocatalysts, which utilize theoretical descriptors to correlate the properties of catalysts with their alkaline HER performance [8,19,24]. Since different descriptors related to the different reaction steps shown in Figure 1 have been proposed and previously utilized [25–30], it is necessary to devise guidelines for selecting descriptors that can accurately predict the performance of HER electrocatalysts in alkaline conditions. Density functional theory (DFT) calculations are essential in computing the proposed energy descriptors related to water adsorption, water dissociation, intermediates adsorption [18,31–33], and product desorption. These descriptors are crucial to understanding alkaline HER electrocatalysts and electrocatalytic processes in the alkaline electrolyzer (AEL). Additionally, identifying important descriptors can improve computational accuracy while minimizing costs [23,34]. While several reviews on the recent progress of alkaline HER electrocatalysis have been presented [8,11,35], a summary and discussion of the most appropriate descriptors for guiding the design of efficient AEL is still rare. To fill this gap, this review will provide a framework by which the desired theoretical descriptor can be selected via DFT computations to design electrocatalysts rather than a comprehensive overview of AWEs.

2. Theoretical Descriptors

As indicated by Equations (4)–(8) and Figure 1, the performance of alkaline HER electrocatalysts is a complex interplay of various factors. The water adsorption strength on active sites, the ability of the catalyst to dissociate water molecules, the hydrogen binding energy, and the OH adsorption processes are all critical in determining the efficiency of the HER process. These factors are intricately linked to the intrinsic properties of the electrocatalysts and their interaction with the operational reaction environment, e.g., electrolyte. Therefore, different theoretical energy-related descriptors have been employed as indicators to screen electrocatalysts for alkaline HER [25–30,36].

2.1. Water Adsorption and Dissociation

The first step of the electrocatalytic water splitting in an alkaline solution is water adsorption, followed by water dissociation to provide the H⁺ cation for hydrogen production (Equations (4)–(6)). HER performance in alkaline or neutral media is primarily determined by the equilibrium between water dissociation (Volmer step) and the subsequent chemisorption of water-splitting intermediates (OH and H) on the surface of the electrocatalysts [37]. To this end, it is critical to strengthen water adsorption at the active sites and decrease the water dissociation activation energy during the Volmer step to increase the overall HER activities of these electrocatalysts. Consequently, the adsorption energy of water ($\Delta E_{H_2O^*}$) and its dissociation energy barriers (ΔE_a) have been used as descriptors to evaluate the performance of alkaline HER electrocatalysts. The $\Delta E_{H_2O^*}$ value can be calculated as shown below:

$$\Delta E_{H_2O*} = \frac{1}{n} \left(E_{H_2O*} - E_* - E_{H_2O} \right) \tag{9}$$

 E_{H_2O*} and E_* are the energies of the surface with and without adsorbed water, respectively. E_{H_2O} and *n* represents the energy of an isolated water molecule and the number of adsorbed water molecules in each surface cell, respectively. According to Equation (9), a more negative value of ΔE_{H_2O*} corresponds to stronger water adsorption.

To calculate the water dissociation energy barrier on the active site, the nudged elastic band (NEB) is widely used [38,39]. The NEB method is highly effective for locating the maximum energy point between specified initial and final configurations of a given transition. The NEB approach for climbing images involves finding a transition path where the saddle point corresponds to the image with the highest energy. The image does not depict the spring forces acting on the band. The image attempts to optimize its energy along the band while minimizing it in all other directions. Once this image reaches convergence, it can be precisely located at the saddle point [40]. The climbing image NEB method enhances the possibility of finding the transition state (TS) for the reaction at the highest energy point. The ΔE_a value is the energy difference between the TS state and the surface with the adsorbed water.

2.2. Gibbs Free Energy Change of Hydrogen Atom Adsorption

Hydrogen atoms adsorbed at the active site are the HER intermediates. The adsorption and desorption of hydrogen atoms on the catalyst surface are commonly viewed as competing processes. During the HER process, the active catalytic site must have a strong affinity for a hydrogen atom. However, a weak adsorption between the active catalytic site and the hydrogen atom is required [41,42]. The high adsorption strength of hydrogen atoms can promote hydrogen atom intermediate formation but impede the formation of

hydrogen molecules. Comparatively, weak adsorption can benefit the formation of H_2 but hinder the formation of hydrogen atom intermediates. A balance must be achieved where the hydrogen intermediate must form but not compete with/prevent the generation and release of the hydrogen molecule. As a result, the optimal adsorption strength of the hydrogen atom intermediate is essential for improving the HER. The adsorption strength of hydrogen atom intermediates on active sites in catalysts can be evaluated using the Gibbs free energy change of hydrogen atom adsorption (ΔG_{H^*}) [43].

The ΔG_{H^*} is obtained by using the computational hydrogen electrode (CHE) method:

$$\Delta G_{\mathrm{H}^*} = \Delta E_{\mathrm{H}^*} + \Delta Z \mathrm{PE}_{\mathrm{H}^*} - T \Delta S_{\mathrm{H}^*} \tag{10}$$

where ΔE_{H^*} describes the binding energy, which is calculated as follows:

$$\Delta E_{\rm H^*} = E_{\rm H^*} - E_* - \frac{1}{2} E_{\rm H_2} \tag{11}$$

Here, E_{H^*} is the total energy of the system with one adsorbed H atom and E_{H_2} represents the energy of an isolated H₂ molecule. ΔZPE_{H^*} is calculated by $\Delta ZPE_{H^*} = ZPE_{H^*} - \frac{1}{2}ZPE_{H_2}$, where the ZPE_{H^*} denotes the vibrational energy of the adsorbed H atom on the active site and the value of ZPE_{H_2} is 0.27 eV. Further, ΔS_{H^*} is the entropy change of H* adsorption, which is obtained by $1/2 \Delta S_{H_2}$. The $T\Delta S_{H^*}$ value is equal to $1/2 T\Delta S_{H_2}$, which is -0.20 eV at 300 K, obtained by Nørskov et al. [32]. The ideal ΔG_{H^*} value is 0 eV.

2.3. Gibbs Free Energy Change of Hydroxyl Adsorption

While hydrogen adsorption energy ($\Delta G_{\rm H}$) primarily determines the alkaline HER activity, the Gibbs free energy change of OH adsorption energy ($\Delta G_{\rm OH^*}$) is found to be equally important since it is responsible for the water dissociation and the amount of hydrogen available [44]. Therefore, the adsorption and desorption of both reaction intermediates (OH* and H*, see Equation (2)) dictate the overall alkaline HER efficiency [45].

The ΔG_{OH^*} is calculated by using the computational hydrogen electrode (CHE) method:

$$\Delta G_{\rm OH^*} = \Delta E_{\rm OH^*} + \Delta ZPE - T\Delta S \tag{12}$$

where ΔE_{OH^*} describes the binding energy of an OH group at the active site, which is calculated as follows:

$$\Delta E_{OH*} = \left(E_{OH*} - E_* - E_{H20} + \frac{1}{2} E_{H_2} \right) \tag{13}$$

where E_{OH^*} is the total energy of the system with one adsorbed OH intermediate, the $\Delta ZPE - T\Delta S$ value is 0.35 eV at 300 K, obtained by Nørskov et al. The ideal ΔG_{OH^*} value is -0.3 eV for the alkaline HER [46].

3. Applications of Descriptors

3.1. Water Adsorption and Dissociation

Figure 1 shows that water adsorption is the first step in generating hydrogen atom intermediates for alkaline HER. The H₂O dissociation becomes a rate-determining step that regulates the catalytic HER activity in alkaline media [47,48]. To this end, the adsorption energy of water and its dissociation energy barriers have been used as descriptors in some research to explain the performance of alkaline HER electrocatalysts.

For example, Chen et al. experimentally synthesized a Ni-doped Pt catalyst on carbon support at 500 °C (termed as NiPt-C-500), which has a low overpotential of 14.0 mV at 10 mA/cm² in the alkaline media [49], using the adsorption energy of water molecules and their dissociation energy barriers as descriptors through DFT computations at the PBE-GGA level to identify the active sites of the NiPt-C-500 catalyst. Figure 2a shows that H₂O can be adsorbed on the Ni sites with an adsorption energy of -0.48 eV. As a comparison, the water adsorption energy on Pt is +0.37 eV. The positive value suggests that the adsorption on Pt is energetically unfavorable. Moreover, a lower H₂O-dissociation barrier on Ni-doped Pt (111) of 0.17 eV was theoretically revealed compared to that on Pt(111), as illustrated in Figure 2b. This suggests a rapid dissociation of H_2O catalyzed by surface Ni atoms on the Pt surface in NiPt-C-500 electrocatalysts, which can increase the rate of H* generation for alkaline HER. Using the same descriptors, Yao et al. resorted to DFT computations to explain the high performance of Cr-doped Co_4N nanorods on carbon cloth (Cr- Co_4N) for alkaline HER [50]. Figure 2c shows that the calculated H_2O adsorption energy on the top of the Cr dopant within $Cr-Co_4N$ is -0.63 eV, which is much lower than that on the surface Co of Co_4N (-0.18 eV). Therefore, the water adsorption on Cr dopants is substantially stronger than on undoped Co₄N. This is because the oxygen atom of water carries a negative Bader charge of -1.13 |e|. As a result, Cr atoms with a higher positive charge $(0.72 \mid e \mid)$ than surface Co $(0.21 \mid e \mid)$ can more strongly adsorb water molecules via higher electrostatic interactions. The enhanced adsorption strength of water can further facilitate the dissociation of the adsorbed water and lower the dissociation barrier, as illustrated in Figure 2d. It explains the experimental observation that introducing Cr dopants can boost the alkaline HER activity.



Figure 2. (a) DFT-calculated water adsorption energies (ΔE_{H_2O*}) and (b) water dissociation energy barriers (ΔE_a) on Pt (1 1 1) and Pt (1 1 1)-Ni surfaces. Reprinted with permission from Ref. [49]. Copyright 2023 Elsevier. (c) DFT-calculated water adsorption energies (ΔE_{H_2O*}) and (d) water dissociation energy barriers (ΔE_a) on Co₄N Cr-Co₄N. Reprinted with permission from Ref. [50]. Copyright 2019 Wiley.

More examples of studies using the $\Delta E_{H_2O^*}$ to identify the active sites of the alkaline HER electrocatalysts can be found in Table 1. Similarly, some examples of using the ΔE_a as the descriptor to explain the performance of alkaline HER electrocatalysts are listed in Table 2.

Catalysts	DFT Method	Adsorption Site	$\Delta E_{H_2O^*}(\mathrm{eV})$	Overpotential@ Current Density of 10 mA cm ⁻²	Conclusion	Reference
Co-doped WO ₂ /Amorphous Co _x W	PBE	Co WO ₂ (001) Co _x W	0.12 - 0.66 - 0.88	- 49 25.0	Amorphous CoxW is an active site for water adsorption.	ACS Appl. Mater. Interfaces (2019), 11, 38771 [51]
Cr-doped Ni(111)	-	Ni(111) Cr-doped Ni(111)	-0.21 -0.58	322.0 203.0	Co-dopants can strengthen water adsorption.	J. Am. Chem. Soc. (2020), 143, 1399 [52]
CuCoMo-doped Ni(111)	PBE	Ni-top Co-top Cu-top Mo-top	-0.23 -0.25 -0.15 -0.67	- - - -	Water is mainly adsorbed on Mo dopants.	Electroanal. Chem. (2019), 839, 224 [53]
N-doped Ni	PBE	Surface Ni Surface Ni next to N dopants	-0.30 -0.41	- 64.0	N dopants can strengthen the water adsorption on their neighboring Ni atoms.	J. Am. Chem. Soc. (2017), 139, 12283 [54]
Cu-Ru/RuSe ₂	PBE	Cu-RuSe ₂ Cu-Ru/RuSe ₂	$-0.40 \\ -1.28$	61.0 23.0	Cu-doped Ru/RuSe ₂ displays a much stronger affinity to water.	Adv. Mater. (2023), 35, 2300980 [55]
Ru _{SA} @Ti ₃ C ₂ O ₂	PBE	$\begin{array}{c} Ti_3C_2O_2\\ Ru_{SA}@Ti_3C_2O_2 \end{array}$	$-0.18 \\ -1.47$	40.3	Enhanced H_2O adsorption by single Ru atom (Ru _{SA}) on MXenes.	EcoMat (2023), 5, 12274 [56]

Table 1. Water adsorption energies as descriptors to identify the active site of the alkaline HER electrocatalyst.

The overpotential values are reported based on the experimental results in the corresponding reference.

Table 2. The dissociation energy barrier of the adsorption water as descriptors to identify the active site of the alkaline HER electrocatalyst.

Catalyst	DFT Method	Active Site	ΔE_a (eV)	Overpotential (mV@ Current Density of 10 mA cm ⁻²)	Conclusion	Reference
Co-doped WO ₂		Co-WO ₂	1.25			ACS Appl. Mater.
/Amorphous	PBE	WO ₂ (001)	1.03	49	Amorphous CoxW is an active site for water	& Interface. (2019),
Co _x W		Co_xW	0.46	25	dissociation.	11, 38771 [51]
		W	0.84	183		NAG
W/WO ₂	PBE-D3	WO ₂	0.06	106	W/WO ₂ interface is the active site that allows water	Nat. Commun. (2023), 14, 5363 [57]
		W/WO ₂	0.02	35	dissociation.	
ENI C	PBE + U	Ni ₃ S ₄	1.05	112	$F\text{-}Ni_3S_4$ allows a better H_2O adsorption.	Adv. Funct. Mater.
F-IN1354		F-Ni ₃ S ₄	0.55	29		(2021), 31, 2008578 [58]
	PBE	Co ₂ P	0.88	247	Oxygen incorporation may induce a higher positive	Adv. Mater. (2017),
Co ₂ P/O-Co ₂ P		O-Co ₂ P	0.51	160	charge state Co, which could benefit water adsorption and weaken the O-H bond in adsorbed H_2O .	29, 1606980 [59]
C. 1 1 N [*] (111)	-	Ni(111)	0.78	322		J. Am. Chem. Soc.
Cr-doped Ni(111)		Cr-doped Ni(111)	0.43	203	Co-dopant can facilitate water dissociation.	(2020), 143, 1399
Bimetallic		NiCu	1.02	85	Partial oxidization of CuNi allov can facilitate water	Angew. Chem. Int.
Nickel-Based Alloys	PBE	E O-NiCu 0.65 23	dissociation.	Ed. (2022), 61, 202202518 [60]		
N-modified Ni	PBE	Ni	0.84	-	N dopant can greatly facilitate water dissociation on	J. Am. Chem. Soc.
		N-Ni	0.42	64	its neighbouring active site.	(2017), 139, 12283 [54]
Cu-Ru/RuSe ₂	DDE	Cu-RuSe ₂	0.31	61	Cu-dopant can promote water dissociation on $Ru/RuSe_{2}$	Adv. Mater. (2023),
	PBE	Cu-Ru/RuSe ₂	0.10	23		35, 2300980 [55]

Catalyst	DFT Method	Active Site	$\Delta E_a(\mathrm{eV})$	Overpotential (mV@ Current Density of 10 mA cm ⁻²)	Conclusion	Reference
NiP ₂ -FeP ₂ /Cu	PBE	NiP2 FeP2 NiP2-FeP2 NiP2-FeP2/Cu F-Ni3S4	0.52 0.49 0.40 0.16 0.55	37 - - 23.6 -	The coupling between interface-rich NiP ₂ –FeP ₂ and metallic Cu can synergistically accelerate water dissociation.	ACS Energy Lett. (2021), 6, 354 [61]
High-entropy alloys (HEAs) Pt ₁₈ Ni ₂₆ Fe ₁₅ Co ₁₄ Cu ₂₇	PBE	Fe	0.11	11	Adsorption of H ₂ O located on the Fe sites, which activates the dissociation of water molecules.	Nat. Commun. (2020), 11, 5437 [62]
2D transition-metal dichalcogenide	PBE	$\begin{array}{c} MoS_2\\ MoSe_2\\ WS_2\\ WSe_2 \end{array}$	0.62 0.74 0.60 0.84	- - - -	MoS_2 and WS_2 have similar activation energy.	J. Phys. Chem. C. (2022), 126, 5151 [63]

Table 2. Cont.

The overpotential values are reported based on the experimental results in the corresponding reference.

3.2. Hydrogen Atom Adsorption

The Gibbs free energy change of the intermediate H atom adsorption (ΔG_{H^*}) was proposed as a descriptor for HER by Nørskov and his co-workers. This descriptor has successfully identified the active site of acidic HER electrocatalysts and the relevant reaction mechanism [32]. A volcano plot is a valuable tool for understanding and optimizing catalyst performance in acidic HER. It is usually a plot of ΔG_{H^*} against activities (such as overpotential, specific current density, and turnover frequency) [64,65]. The highest point on the volcano map is the ideal ΔG_{H^*} value at which the catalytic activity is maximized. Catalysts located on the left side of the volcano plot exhibit excessive hydrogen binding (very strong), while those on the right side have insufficient hydrogen binding (very weak). According to the Sabatier principle, an optimum catalyst should exhibit adsorption energies that are neither too strong nor too weak. In the alkaline HER, the adsorption of H atoms at the active site is also important since it is related to the water dissociation, Volmer, Heyrovsky, and Tafel steps, as suggested by Equations (5)–(8) and Figure 1. To this end, the ΔG_{H^*} was also widely used in the analysis of the performance of the alkaline HER electrocatalysts [66].

For example, Chen et al. adopted a dual approach to simultaneously promote water dissociation and hydrogen desorption kinetics with Co-doped WO₂/amorphous Co_xW hybrid catalysts using DFT [51]. As shown in Figure 3a, the ΔG_{H^*} on amorphous Co_xW is -1.88 eV, indicating a strong hydrogen-binding interaction that prevents H₂ production and desorption (Figure 4a). The most catalytically active site on $Co-WO_2(011)$ has a ΔG_{H^*} of -0.06 eV, closer to thermoneutral than Pt's -0.09 eV, indicating significant hydrogen adsorption capability. According to DFT calculations, the HER on Co-doped WO₂/amorphous CO_xW hybrid catalyst follows the Volmer-Tafel step, where water molecules are activated and cleaved to form H atoms on the surface of amorphous $Co_x W$. These H atoms then rapidly combine to form H_2 on the surface of Co-doped WO_2 . He et al. incorporated a single Ru atom into Ni_5P_4 to achieve an effective electrocatalyst for alkaline HER [67]. The DFT calculations were conducted to comprehend the impact of single-atomic Ru incorporation into Ni_5P_4 on its catalytic and structural properties. The structures of Ni_5P_4 were determined, both with and without Ru incorporation, and they revealed several potential catalytic sites. Experimental analysis and the established model structural parameters were remarkably congruent. In the case of Ni₅P₄-Ru, the computed ΔG_{H^*} value at the Ru-doped sites was -0.30 eV, which was comparatively higher than the value of -0.42 eV observed at the site of P of pristine Ni_5P_4 (see Figure 3b). This finding indicated that the Ru dopant exhibited favorable energetics for the desorption of absorbed H atoms to form the H_2 gas.



Figure 3. (a) DFT-calculated Gibbs free energy change of the intermediate H atom adsorption (ΔG_{H^*}) on WO₂, Co-WO₂, CoxW, and Pt. Reprinted with permission from Ref. [51]. Copyright 2019 American Chemical Society. (b) DFT-calculated Gibbs free energy change of the intermediate H atom adsorption (ΔG_{H^*}) on Ni₅P₄ and Ni₅P₄: Ru. Reprinted with permission from Ref. [67]. Copyright 2020 Wiley.

Table 3 lists more examples of using ΔG_{H^*} as the descriptor to identify the active site of the alkaline HER electrocatalysts and evaluate their performance.

Table 3. Gibbs free energy change of H intermediate adsorption as descriptors to identify the active site of the alkaline HER electrocatalyst.

Catalyst	DFT Method	Active Site	$\Delta G_{H^*}(\mathrm{eV})$	Overpotentials (mV@ Current Density of –10 mA cm ^{–2})	Conclusion	Reference
		W	-0.51	183.0		Nat. Commun. (2023) 14, 5363 [57]
W/WO ₂	DFT-D3	WO ₂	-1.24	106.0	W/WO_2 interface is the active site for HER.	
		W/WO ₂ interface	-0.41	35.0		
Co-doped WO ₂		Co-WO ₂	-0.06	-		ACS Appl. Mater. &
/Amorphous	PBE	WO ₂ (001)	-0.30	49.0	Co-WO ₂ is the active site for H_2 formation.	Interface (2019), 11, 38771 [51]
Co _x W		Co _x W	-1.88	-		
C 1 1N((111)		Ni(111)	-0.23	322.0	Co-dopant can reduce the hydrogen desorption rate.	J. Am. Chem. Soc. (2020), 143, 1399 [52]
Cr-doped Ni(111)	-	Cr-doped Ni(111)	-0.38	203.0		
		Ni ₃ -fcc	-2.76 *	-	The preferential adsorption site is a Mo-coordinated fcc site.	Electroanal. Chem. (2019), 839, 224 [53]
CuCoMo-doped	DDE	Cu ₂ Co-fcc	-2.51 *	-		
Ni(111)	PBE	Co ₂ Cu-fcc	-2.67 *	-		
		CuCoMo-fcc	-2.78 *	-		
		Ni	-0.31	-	Cu can weaken the adsorption strength of hydrogen to benefit the H ₂ production.	Angew. Chem. Int. Ed. (2022), 61, 202202518 [60]
Ni/NiCu/O- NiCu	PBE	NiCu	-0.11	85.0		
		O-NiCu	-0.035	23.0		
Cu-Ru/RuSe ₂		Cu-RuSe ₂	0.50	61.0	Cu can weaken the adsorption strength of hydrogen to benefit the $\rm H_2$ production.	Adv. Mater. (2023), 35, 2300980 [55]
	PBE	Cu-Ru/RuSe ₂	0.19	23.0		

Catalyst	DFT Method	Active Site	$\Delta G_{H^*}(\mathrm{eV})$	Overpotentials (mV@ Current Density of -10 mA cm ⁻²)	Conclusion	Reference
		NiP ₂	-0.358	37.0		
NERE R. (C		FeP ₂	-0.203	-	NiP-FeP2/Cu interface effectively adsorbs	ACS Energy Lett.
NiP-FeP ₂ /Cu	PBE	NiP-FeP2	-0.043	-	generated H intermediate.	(2021), 6, 354. [61]
		NiP-FeP ₂ /Cu	-0.03	23.6		
O-Co ₂ P	PBE	Co ₂ P	0.70	247.0	Oxygen incorporation may induce a higher positive charge state Co to benefit water adsorption and	Adv. Mater. (2017), 29,
		O-Co ₂ P	0.19	160.0	weaken the O-H bond in adsorbed H_2O .	1606980 [39]
		Ti ₃ C ₂ O ₂	-0.33	-	Dec. Generation	EcoMat (2023), 5, 12274
$Ru_{SA} @ II_3 C_2 O_2$	PBE	$Ru_{SA}@Ti_3C_2O_2$	-0.07	40.3	Ku_{SA} ravours H_2 formation.	[56]
ENLO		Ni ₃ S ₄	0.12	112.0	F-doping can promote water dissociation to	Adv. Funct. Mater.
F-1N1354	PBE	F-Ni ₃ S ₄	-0.034	29	increase the rate of Hads formation.	(2021), 31, 2008578 [58]
		Pt/Mo ₂ C interface	0.01	-	The $\Delta G_{\mathbf{H}^*}$ has a strong correlation with the alkaline HER performance.	ACS Catal. (2019), 9, 2415 [29]
		Pt/NbC	-0.28	-		
		Pt/TaC	-0.07	-		
		Pt/TiC	-0.03	-		
Metal-modified		Mo2C(0001)	-0.75	-		
transition metal	PW91	NbC	-0.90	-		
carbides		TaC	-0.99	-		
		TiC	-0.96	-		
		VC	-0.84	-		
		W ₂ C(0001)	-0.60	-		
		WC(0001)	-0.81	-		
		MoSe ₂	1.49	301	Partial substitution of Ni or Co atoms for Mo	Angew. Chem. (2020),
N1/Co-modified	PBE	Ni-MoSe ₂	-0.43	98		
MoSe ₂		Co-MoSe ₂	-0.20	183	atoms can optimize the ΔG_{H^*} value.	132, 15344 [27]
		Ni	-0.08	436		
		Fe	0.17	509		
FeCoNiCu _{0.5}	PBE	Со	-0.18	361	ΔG_{H^*} value of -0.08 eV, close to that of the Pt catalyst.	J. Alloys Compd.
		Cu	0.23	252		(2024), 175356 [68]
		FeCoNiCu _{0.5}	-0.08	71		
		MoS ₂	0.03	-	Except for WSe2, the calculated ΔG_{H^*} of the rest is close to zero.	J. Phys. Chem. C. (2022), 126, 5151 [63]
2D		MoSe ₂	0.09	-		
transition-metal PBE dichalcogenides	PBE	WS ₂	-0.03	-		
		WSe ₂	0.24	-		
		-				

Table 3. Cont.

* The authors calculated the adsorption energy of the H atoms here. The overpotential values are reported based on the experimental results in the corresponding reference.

3.3. Hydrogen and Hydroxyl Adsorption

In recent studies, the volcano plot has been adopted to describe HER activities in alkaline media by plotting ΔG_{OH^*} against activities. In contrast to the volcano-shaped association observed with ΔG_{H^*} , Zhang et al. demonstrated that ΔG_{OH^*} exhibits a significantly weaker correlation with alkaline HER activity than the volcano-shaped relationship established with ΔG_{H^*} [29]. Au-modified TMCs exhibit OHBEs comparable to those of Pt and Pt-modified TMCs; however, the alkaline exchange current densities of Au/TMCs are two to three orders of magnitude lower than those of their Pt counterparts. Furthermore, carbides modified with Ag and Cu exhibit a broad spectrum of ΔG_{OH^*} , but there is no discernible HER trend within this spectrum. According to the research of Zhang et al., ΔG_{OH^*} is not an appropriate descriptor for alkaline HER on TMCs and metal-modified TMCs. It also implies that the adsorbed hydroxyl group does not directly participate in the rate-determining step of alkaline HER kinetics on these surfaces [29].

Markovic et al. were the first to study the impact of adsorbed OH on Pt-based electrocatalysts, emphasizing its poisoning effect on surface sites and its influence on the kinetics of the alkaline HER [69]. The hydrogen binding energy (HBE) and hydroxyl binding energy (OHBE) values for pure Ru and various Mo-Ru compounds were computed (Figure 4a) by Zhao et al. Through DFT, the optimal H and OH adsorption sites on distinct catalysts for alkaline HER were identified [70]. In the Mo-Ru-1 structure, one of the Ru

atoms is doped with Mo, while Mo-Ru-2 and Mo-Ru-3 are doped with two and four atoms of Mo, respectively. Figure 4a displays the HBE (ΔG_{H^*}) values of Ru, Mo-Ru-1, Mo-Ru-2, and Mo-Ru-3, which are -0.64, -0.69, -0.67, and -0.76 eV, respectively. These results indicate that the adsorption energy of H is similar for all catalysts and nearly identical to experimental values [25,71,72]. The OHBE (ΔG_{OH^*}) values of Mo-Ru-1 (-0.53 eV), Mo-Ru-2 (-0.68 eV), and Mo-Ru-3 (-0.97 eV) exhibited a substantial increase after Mo-doping, in comparison to native Ru (-0.31 eV). Consequently, the OHBE progressively rose as the Mo atom content increased. Furthermore, the increased OHBE after Mo-doping can be ascribed to the augmented HOR/HER activity of Mo-Ru.

Zhang et al. designed high-efficiency alkaline HER electrocatalysts using a unique dual descriptor of optimal free energies (ΔG_{H^*} and ΔG_{OH^*}). N_{i3}N surface reactivity was tailored using theory to balance the adsorption energies of hydrogen and hydroxyl species. Nickelbased materials are highly promising non-noble metal electrocatalysts for the hydrogen evolution reaction (HER) [73,74]. In this study, the metallic Ni₃N was considered a case study to illustrate the application of dual-descriptor-driven design. Ni₃N has been regarded as a suitable electrocatalyst for cleaving OH-H bonds in the Volmer phase. According to Figure 4b, the DFT calculations demonstrate that the Ni_3N (111) surface exhibits a facile hydroxyl adsorption free energy of 0.03 eV. This characteristic is advantageous for the energetics of water dissociation and subsequent hydroxyl desorption [75]. Such findings align with prior studies indicating that Ni₃N can function as a promoter of water dissociation [76]. Nevertheless, the DFT calculations indicate that the average G_{H^*} value observed on the surface of Ni_3N (111) is approximately -0.31 eV, which surpasses the ideal value of $\Delta G_{H^*} = 0$ eV by a wide margin. The excessive reactivity of the surface results in unfavorable desorption of hydrogen and subsequent formation of H₂. This approach has successfully enhanced the hydrogen evolution reaction (HER) performance by incorporating various transition-metal dopants into Ni3N, including Mo-Ni₃N, W- Ni_3N , and V- Ni_3N [30]. The hydrogen and hydroxyl binding energy results on Mo- Ni_3N effectively balance the dual descriptors.



Figure 4. (a) DFT-calculated Gibbs free energy change of the intermediate H atom adsorption (ΔG_{H^*} , HBE) and the Gibbs free energy change of OH adsorption energy (ΔG_{OH^*} , OHBE) on Ru, Mo-Ru-1, Mo-Ru-2, Mo-Ru-3. Reprinted with permission from Ref. [70]. Copyright 2022 American Chemical Society. (b) DFT-calculated Gibbs free energy change of the intermediate H atom adsorption (ΔG_{H^*}) and the Gibbs free energy change of OH adsorption energy (ΔG_{OH^*}) on Ni₃N, Mo-Ni₃N, W-Ni₃N, and V-Ni₃N. Reprinted with permission from Ref. [30]. Copyright 2019 American Chemical Society.

More examples of using ΔG_{H^*} and ΔG_{OH^*} as the descriptor to identify the active site of the alkaline HER electrocatalysts and evaluate their performance are listed in Table 4.

Catalyst	DFT Method	Active Site	ΔG_{OH^*}	Overpotentials (mV@ Current Density of -10 mA cm ⁻²)	Conclusion	Reference
Cr-doped Ni(111)	-	Ni(111)	-3.23 *	322	Co-doping allows for a balance between facilitating	J. Am. Chem. Soc.,
		Cr-doped Ni(111)	-3.83 *	203	the dissociation of water and preventing the poisoning effect.	(2021), 143, 1399 [52]
Burnetti C. O. /Ti C. O.	DDE	RuSA@Ti3C2O2	-0.49	40.3	Ru improved adsorption ability of Ti ₃ C ₂ O ₂	EcoMat (2023), 5, 12274 [56]
Kusa@113C2O27113C2O2	PDE	$Ti_3C_2O_2$	2.00	-	toward HO*.	
Ni/Comodified		Mo-MoSe ₂	-1.74	301		Angew. Chem. (2020), 132, 315344 [27]
Ni/Co-modified	PBE	Co-MoSe ₂	0.67	183	Co-MoSe ₂ is favorable for OH ⁻ desorption in alkaline HER.	
MoSe ₂		Ni-MoSe ₂	1.22	98		
	PW91	Pt	-2.39	-	The ΔG_{OH^*} does not show a strong correlation with the alkaline HER performance.	ACS Catal. (2019), 9, 2415 [29]
		Nb	-2.41	-		
		Ta	-2.51	-		
		Мо	-4.6	-		
N + 1 110 1		Nb	-4.85	-		
Metal-modified		Ta	-4.6	-		
transition metal		Ti	-4.85	-		
carbides		V	-4.96	-		
		W	-4.91	-		
		W	-4.19	-		
		RuSA@Ti3C2O2	-0.49	-		
		Ni	0.17	-		
		Pt	1.05	28.9	The strong adsorption of OH on Ni indicates a fast	J. Colloid Interface Sci.
Ni-modified Pt (111)	PBE	Ni-Pt	-0.51	14	H_2O dissociation.	(2023) 650, 1715 [49]

Table 4. Gibbs free energy change of OH intermediate adsorption as descriptors to identify the active site of the alkaline HER electrocatalyst.

* The authors calculate the adsorption energy of the OH radical at the active site. The overpotential values are reported based on the experimental results in the corresponding reference.

4. Discussion

The selection of the right descriptor for screening alkaline HER electrocatalysts is a long-standing debate, especially using the ΔG_{OH^*} descriptor. For example, Zhang et al. demonstrated that ΔG_{OH^*} exhibits a significantly weaker correlation with alkaline HER activity than the volcano-shaped relationship established with ΔG_{H^*} (Figure 5a), which implies that the adsorbed hydroxyl group does not directly participate in the rate-determining step of alkaline HER kinetics on these surfaces [29]. Contrastingly, McCrum et al. found that the performance of alkaline HER electrocatalysts shows a volcano relationship between the natural logarithm of the experimentally measured rate of hydrogen evolution and the DFT-calculated ΔG_{OH^*} [19]. The reaction is bifunctional as it involves both ΔG_{H^*} and ΔG_{OH^*} on the too-strong OH binding side of the volcano. On the too-weak OH binding side of the volcano, ΔG_{H^*} is only apparently bifunctional; ΔG_{OH^*} is not a useful descriptor for evaluating the performance of catalysts. The different conclusions of these two studies may be ascribed to the scopes of ΔG_{OH^*} values. All the ΔG_{OH^*} values from Zhang et al. shown in Figure 5a are less than -1.0 eV, which belong to the too-strong OH binding side suggested in Figure 5b. As a result, no volcanic relationship can be identified in Figure 5a. Since all ΔG_{OH^*} values are far from the optimal value of -0.3 eV, the performance of these metal-modified transition metal carbides is more affected by the ΔG_{H^*} value, as suggested by Zhang et al. [29].



Figure 5. (a) The logarithm of the experimentally measured rate of hydrogen evolution of metalmodified transition metal carbides in 0.1 M KOH and the corresponding DFT-calculated ΔG_{OH^*} . Reprinted with permission from Ref. [29]. Copyright 2019 American Chemical Society. (b) The natural logarithm of the experimentally measured rate of hydrogen evolution of Pt (553), Pt(553) with Mo*, Re*, Ru*, Rh*, and Ag* adsorbed at the step, and Pt(111) in 0.1 M NaOH and the corresponding DFT-calculated ΔG_{OH^*} . Reprinted with permission from Ref. [19]. Copyright 2020 Springer Nature.

To better screen the alkaline HER electrocatalysts, McCrum et al. adopted a threedimensional (3D) volcano plot depicting the rate of hydrogen evolution in relation to the ΔG_{OH^*} and ΔG_{H^*} , as illustrated in Figure 6a. Catalysts exhibiting minimal hydrogen and hydroxide binding energies yield the lowest rates, as indicated by the purple area on the upper right of Figure 6a. The catalysts show the highest performances when they bind hydrogen at an intermediate strength at 0 V_{RHE} (near 0 eV) and hydroxide strongly (near -0.3 eV) (yellow in the middle of Figure 6a). The too-strong H and OH binding leads the H₂ or OH* desorption to become the rate-determining step. Consequently, Figure 6a qualitatively represents trends in alkaline HER kinetics and provides design guidelines for efficient catalysts. For example, to improve the hydrogen evolution rate of the well-studied Pt(111) electrocatalysts in an alkaline solution, the ΔG_{H^*} must be increased by approximately 0.2 eV. In comparison, the ΔG_{OH^*} must be significantly reduced by approximately 0.9 eV through appropriate manipulation strategies such as doping, surface engineering, and defect engineering [19].

Following these guidelines, we employed both ΔG_{H^*} and ΔG_{OH^*} descriptors to investigate the catalytic HER performance of 1T' transition metal dichalcogenides (TMDs such as MoSe₂, MoS₂, WSe₂, and WS₂) in an alkaline solution using DFT [77]. Our findings indicate that the pristine sulfides exhibited superior alkaline HER performance compared to their selenide counterparts. Nevertheless, the activities of all pristine 1T' TMDs are insufficient to dissociate water (See Figure 6b). Defect engineering techniques were employed to improve the reactivity of TMD-based electrocatalysts. Our DFT results indicate that the reactivities of TMD materials can be enhanced by introducing single S/Se vacancy defects, as shown in Figure 6c. However, the rate-determining phase is the desorption of OH. The reactivities of active sites were further regulated to achieve optimal OH desorption by doping defective MoS₂ with late 3d transition metal (TM) atoms, particularly Cu, Ni, and Co, as illustrated in Figure 6d. Consequently, the TM-doped defective 1T' MoS₂ can substantially improve the alkaline HER performance, which matches the recently reported experimental observations.



Figure 6. (a) Logarithm of the rate of hydrogen evolution (contours) as a function of DFT-calculated ΔG_{H^*} and ΔG_{OH^*} on Pt(111), Pt(553), Ru* adsorbed at the step of Pt(553), PtRu(111) alloy and Ru* clusters on Pt(111). Reprinted with permission from Ref. [19]. Copyright 2020 Springer Nature. (b) Plot of ΔG_{OH^*} against ΔG_{H^*} of pristine MX₂ TMDs (M = Mo, W and X = S, Se), (c) plot of ΔG_{OH^*} against ΔG_{H^*} of MX₂ TMDs with a single X vacancy; and (d) plot of ΔG_{OH^*} against ΔG_{H^*} of MX₂ TMDs with a single X vacancy. Reprinted with permission from Ref. [77]. Copyright 2024 IOPscience.

It is crucial to remember that each descriptor has limitations and challenges when studying alkaline HER in alkaline media. Descriptors such as water adsorption energy and water dissociation energy barriers serve as indicators of material reactivity. They are closely linked to the Volmer step, which encompasses the adsorption of water molecules and their subsequent dissociation into adsorbed hydrogen (H*) and hydroxide ions (OH⁻). These steps are fundamental to the overall HER process, yet they do not provide a complete picture of the catalytic site's capabilities, especially concerning the Heyrovsky and Tafel steps. The Heyrovsky step involves the electrochemical desorption of H* to form hydrogen gas.

The ΔG_{H^*} has been the most prevalent descriptor for assessing HER performance at the Heyrovsky and Tafel steps. It offers a measure of the free energy change when H* is adsorbed on the catalyst surface, which is a critical factor in determining the rate of the HER. However, ΔG_{H^*} alone is insufficient for evaluating the Volmer step or the desorption efficiency of adsorbed OH, which are also essential for a complete understanding of the HER mechanism.

The Gibbs free energy of hydroxide adsorption (ΔG_{OH^*}) can be used to evaluate the desorption efficiency of adsorbed OH, providing insights into the potential for water dissociation, as suggested by the Bronsted-Evans-Polanyi (BEP) relationship [19]. This relationship posits a linear correlation between the activation energy of a reaction and the reaction enthalpy, allowing for the prediction of reaction barriers based on thermodynamic

parameters. However, it does not offer information on the catalyst's performance during the Heyrovsky or Tafel steps. Therefore, relying solely on ΔG_{OH^*} would give an incomplete assessment of a catalyst's overall activity and efficiency.

Given these considerations, it is evident that no single descriptor can fully encapsulate the complexities of the HER process. Researchers must carefully select and combine multiple descriptors to gain a comprehensive understanding of the catalytic activity and to design more efficient catalysts. This approach allows for the evaluation of catalysts across all steps of the HER, ensuring a more accurate prediction of their performance in real-world applications.

It is also worth noting that different descriptors may be required to understand the performance of alkaline HER composite electrocatalysts. For example, Chen et al. designed a complicated composite electrocatalyst including a Co dopant on WO₂/amorphous Co_xW hybrid materials [51]. Using the water adsorption energy and dissociation energy barrier energy as the descriptor, they found that the amorphous CoxW is the active site. However, the analysis of the ΔG_{H^*} descriptor indicates that Co-doped WO₂ is the active site for H₂ formation. The DFT calculations then suggest that the individual component of this composite electrocatalyst has a different function. The synergy between them enables the high alkaline HER performance of this Co-Doped WO₂/amorphous Co_xW hybrid electrocatalyst.

5. Conclusions and Outlook

The hydrogen evolution reaction (HER) in alkaline media is a prominent method for large-scale hydrogen production from an electrolyzer. Alkaline Water Electrolysis (AWE) has demonstrated superior cost-effectiveness to acidic proton exchange membrane (PEM). One of the biggest challenges of AWE is ascribed to the relatively low energy conversion efficiency of their electrocatalysts. To address this issue, selecting the right computational tools for theoretical studies is essential in designing high-performance alkaline HER electrocatalysts. The descriptors used to evaluate the energy conversion efficiency of electrocatalysts include water adsorption energy, water dissociation barrier, Gibbs free energy change of hydrogen adsorption, and Gibbs free energy change of hydroxyl adsorption from DFT calculations. In the review, some of the latest examples have been used to illustrate the applications of different descriptors. It reveals that the hydroxyl (ΔG_{OH^*}) adsorption process is one of the important parameters often ignored in many previous theoretical studies. When the adsorption strength of hydroxyl is weak, it suggests a high water dissociation energy barrier due to the BEP relationship [19]. The water cannot effectively interact with the active site to provide an H atom intermediate. At the same time, the weak adsorption of OH also indicates that the dissolved hydroxide is via the Volmer (II) mechanism (Equation (6)), as shown in Figure 1. In this case, the ΔG_{H^*} becomes a more important descriptor to analyze and predict the efficiency of an active site. It explains the great matches between some experimental measurements and the trend of the ΔG_{H^*} descriptor in some combined studies.

The CHE model has evolved to include previously overlooked factors such as solvent and electrolyte effects, reflecting its adaptability [18]. Despite these advancements, a knowledge gap persists, particularly in the dynamic aspects of water splitting. A significant hurdle remains in developing a more accurate model of the electrode–electrolyte interface, which is crucial for pinpointing active sites in HER and connecting microscopic interactions with macroscopic observations. The electrified solid–liquid interface in alkaline HERs largely determines the charge transfer rate of electrochemical redox reactions [78]. Innovative approaches, such as engineering a localized acid-like environment within an alkaline medium, have significantly boosted HER performance significantly [79]. This is achieved by tailoring the local reaction conditions, which are critical for the process. Moreover, understanding the impact of variables like electrolyte concentration on HER allows for further refinement of reaction conditions [16]. The descriptors described in this review can only provide limited dynamic information about the electrochemical processes at the electrolyte interface [15,21,22]. This is because the atomistic models used

in most of these studies make it difficult to comprehend the thermodynamic state and dynamic properties of interfacial processes [21]. The interaction between water molecules and the electrified surface must be investigated by examining scenarios in which water molecules are introduced to the catalyst surface in various configurations [80]. Insights into the hydrogen bonding network and structural characteristics of the water layer can be derived from the average dipole orientation of water molecules relative to the surface normal [80]. Furthermore, the complex relationship among adsorbed hydrogen, water molecules, and the electrochemical environment, which influences the behaviors of adsorption and the overall characteristics of the interface between solid and water, needs to be investigated [81]. To comprehend the activities of HER intermediates in alkaline media, it is critical to know the potential and pH-dependent adsorption energies of these intermediates [82]. To this end, it is clear that theoretical studies on alkaline HER are still an area ripe for further research.

The intersection of machine learning (ML) with density functional theory and multiscale modeling is a burgeoning field that holds great promise for advancing materials science and chemistry [83–86]. By leveraging ML's ability to analyze vast datasets and identify patterns, researchers can significantly reduce the computational resources required for DFT calculations. This synergy enables the prediction of complex chemical behaviors and the design of new materials with tailored properties. However, the success of the ML methods hinges on the quality and consistency of the data fed into the ML models [84]. As such, creating comprehensive and reliable databases is crucial for training algorithms that can accurately predict electrochemical behaviors and guide the development of efficient alkaline HER electrocatalysts. This integrated approach is set to transform the landscape of computational chemistry, offering a more streamlined and precise method for exploring the vast potential of chemical space. They can predict the performance of electrocatalysts, thereby informing the design of next-generation materials with enhanced catalytic properties. These concerted efforts in theoretical research are essential for the progression of clean energy technologies.

In sum, the exploration of electrocatalytic mechanisms in alkaline HER is a complex field that necessitates a comprehensive approach. Advancements beyond the DFTcalculated descriptors are critical for a deeper understanding. Operando simulations offer a dynamic perspective by considering actual working conditions, providing insights into the real-time structural and chemical changes during the reaction process. Meanwhile, MLbased force fields for classical molecular dynamics and Monte Carlo simulations represent a significant leap in mesoscale modeling, enabling simulations that capture the nuanced interactions within molecular systems. These ML models can bridge the gap between classical and quantum mechanical accuracy, offering a more detailed view of the catalytic processes. Lastly, ML-driven high-throughput screening is revolutionizing the way electrocatalysts are discovered and optimized. By analyzing vast datasets, ML algorithms can predict performance, stability, and efficiency, thereby accelerating the development of new materials for HER. Together, these methodologies form a multi-faceted approach that could significantly advance the field of electrocatalysis.

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