Tuning the local reaction environment is an important and challenging issue for determining electrochemical performances. Herein, we propose a strategy of intentionally engineering the local reaction environment to yield highly active catalysts. Taking Pt\(^{6-}\) nanoparticles supported on oxygen vacancy enriched MgO nanosheets as a prototypical example, we have successfully created a local acid-like environment in the alkaline medium and achieve excellent hydrogen evolution reaction performances. The local acid-like environment is evidenced by operando Raman, synchrotron radiation infrared and X-ray absorption spectroscopy that observes a key H\(_3\)O\(^+\) intermediate emergence on the surface of MgO and accumulation around Pt\(^{6-}\) sites during electrocatalysis. Further analysis confirms that the critical factors of the forming the local acid-like environment include: the oxygen vacancy enriched MgO facilitates H\(_2\)O dissociation to generate H\(_3\)O\(^+\) species; the F centers of MgO transfers its unpaired electrons to Pt, leading to the formation of electron-enriched Pt\(^{6-}\) species; positively charged H\(_3\)O\(^+\) migrates to negatively charged Pt\(^{6-}\) and accumulates around Pt\(^{6-}\) nanoparticles due to the electrostatic attraction, thus creating a local acidic environment in the alkaline medium.
The development of a highly-efficient and low-cost pathway to produce renewable energy resources has been one of the top priorities in the science community in recent decades. Electrochemistry approaches play essential roles in this field due to the low reaction activation energy and high reaction rate and energy efficiency. The kinetics of electrode reactions depend strongly on both the nature of electrode materials and the local concentrations of solution constituents in the vicinity of the catalytic sites (hereafter, called the local reaction environment). Thus far, the efficiency of the electrode reactions is mainly increased by ameliorating the catalyst materials through various methods, such as controlling the crystal facet, doping heteroatoms, creating dual active sites, engineering defects, and strains, and so on. In most cases, these conventional strategies can only tailor their atomic structures, electronic states, and thereby catalytic properties in a gradual or mild way. To seek breakthroughs in electrode reactions, one needs to recall the elementary steps in electrochemistry. All the electrochemical processes involve the non-covalent interactions between the atoms and molecules of the electrocatalyst surface and the reaction intermediates generated by the solvent or solute molecules, leading to the formation of intermediate complexes or nonuniform distribution of the ionic species. Therefore, the local reaction environment around catalysts also plays a vital role in electrode processes. Tuning the local reaction environment through multiple physicochemical effects between the substrate, metal, and reaction intermediate thus provides an alternative way to promote the electrocatalytic performance and guide the higher efficiency electrocatalyst design. This task is challenged by the lack of facile and practical strategies to engineer the local reaction environment, as well as by the difficulties in identifying the weak signals arising from interfacial structures under complex reaction conditions.

The hydrogen evolution reaction (HER) is one of the most classic electrochemical reactions in both academic research and industry applications. Industrial plants prefer to use water-based alkaline solution instead of acidic solution as hydrogen sources; the reason is that the alkaline HER can both avoid using the high-cost proton exchange membrane and alleviate the problem of slow electron-transfer kinetics of the oxygen evolution reaction (OER) on the anode within an electrolyzer under acidic conditions. However, the alkaline HER also has its drawbacks. The principal one is the low conversion efficiency. Taking Pt-based catalysts, which are regarded as the most efficient HER catalysts as an example, the HER conversion efficiency in alkaline condition is two or three orders of magnitude lower than that in acidic solutions. The alkaline HER kinetics are still elusive, and several schools of thought on the slow kinetics of the alkaline HER have been proposed, including the hydrogen binding energy (HBE) theory, water-dissociation theory, and interface water and/or anion transfer theory. Nevertheless, the general consensus is that during the alkaline HER, the sluggish Volmer step directly or indirectly impacts the rate-determining step, but this step is unnecessary in an acidic solution. One can expect that if we could create a local acid-like environment for the HER in an alkaline medium where the corresponding OER occurs, the adverse factors of the alkaline HER and acidic OER will be solved fundamentally, and both the HER and OER could proceed at a high rate. The feasibility of this strategy is illuminated by recent studies showing that the interfacial microstructure of the electrolyte can be modified by the physicochemical properties of the solid surface. Therefore, modulating the local reaction environment by selecting a suitable system to create a local acid-like environment, will provide a route to design highly-efficient HER catalysts. Previous studies have revealed that MgO, Al2O3, and Ni(OH)2 surfaces facilitate the dissociation of H2O molecules, giving rise to H+ groups. However, these free protons cannot accumulate spontaneously around the catalytic sites to form a concentrated acidic region. Thus, an electrocatalyst system capable of driving the aggregation of H+ groups should be developed.

In this work, we propose a practical pathway for engineering a local acid-like reaction environment to design highly-efficient alkaline HER catalysts. By virtue of multiple physicochemical interactions between the substrate, metal active site, and reaction intermediate, we selected Pt/MgO as the prototypical example to construct an acid-like reaction environment in an alkaline medium. Operando Raman spectroscopy, synchrotron radiation Fourier transformed infrared spectroscopy (SR-FTIR) spectroscopy, and X-ray absorption near-edge spectroscopy (XANES) confirm the generation of massive amounts of H3O+ intermediates on the MgO surface and accumulation around negatively-charged Pt (Ptδ−). The local acid-like reaction environment leads to an extraordinary HER performance, with a very low overpotential of 39 mV at 10 mA cm−2, which is much better than the value of 62 mV for 20 wt% Pt/C and close to the acidic HER behavior of 20 wt% Pt/C (33 mV). The system also has a tenfold higher mass activity than that of the 20 wt% Pt/C electrodes in an alkaline medium and 2.5-fold higher than that of 20 wt% Pt/C in acidic medium at −39 mV vs. RHE. Experimental characterizations and first-principles calculations suggest that the oxygen vacancy-rich MgO is favorable for water dissociation, and the electronic interaction between the MgO and Pt nanoparticles drives electron transfer from VO−MgO to Pt, giving rise to the formation of negatively-charged Ptδ− species. Then the Ptδ− accelerates H3O+ migration and an acid-like environment is formed around the Ptδ− in an alkaline medium, thus boosting the HER in this alkaline medium. We believe that this finding will contribute to future explorations in other important solution-dependent fields, such as surface, energy and environmental science.

Results

Structural and morphological characterizations. To synthesize Pt/MgO catalysts with strong electronic metal-support interactions (EMSIs) and an abundance of oxygen vacancies, we chose MgMOFs as the precursor for the MgO support (Supplementary Fig. 1), and immersed Pt ions into the pores of MgMOFs, followed by annealing at 700 °C in an atmosphere of nitrogen and oxygen. As revealed by the thermogravimetric analysis shown in Supplementary Fig. 2, the formation of Pt nanoparticles on MgO consists of three steps. The successful synthesis of Pt/MgO is confirmed by powder X-ray diffraction (XRD) as shown in Fig. 1a. The diffraction peaks at 20 angles of 39.8° and 46.4° are indexed to the (111) and (200) planes of fcc Pt nanocrystals, respectively, and the other diffraction peaks correspond to the MgO support. Figure 1b presents a transmission electron microscopy (TEM) image of Pt/MgO, indicating the uniform dispersion of Pt nanoparticles on the MgO nanosheet support (the MgO nanosheet is shown in the inset and Supplementary Fig. 3). The sizes of the Pt nanoparticles are ~5 nm (Supplementary Fig. 4). From the high-resolution TEM (HRTEM) image in Fig. 1c, the lattice fringes of 0.196 and 0.209 nm are ascribed to the Ptδ− plane of MgO, respectively. The high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) image (inset) in Fig. 1d shows the well-defined Pt nanocrystals on MgO, with a size distribution of ~10 nm (Table 1).
RHE, a new peak at ~1750 cm⁻¹ (1600 cm⁻¹) was observed, which is ascribed to the stretching vibrations of the O-H group in H₃O⁺ when the applied potential is reduced to ~40 mV vs. RHE (Fig. 3b and Supplementary Fig. 8)³⁰,³¹; similar phenomena are observed on MgO nanosheets, but not on Pt/C (Supplementary Fig. 8), indicating the critical role of MgO in facilitating water dissociation and creating the H₂O²⁺ intermediate. More strikingly, the potential-dependent signal of H₂O²⁺ is reversible, manifesting that the local acid-like environment is formed only during the process of HER (Supplementary Fig. 9). In addition, it is worth noting that the detection limits of SR-FTIR spectroscopy and Raman spectroscopy are approximately ppm levels for hydroxyl groups or hydration molecules, which is much higher than that of the content of H⁺ (~10⁻¹⁴ M) ionized from water in the 1 M KOH solution. Such a high level of H₂O²⁺ cannot be ubiquitous in KOH but can accumulate only within a local region. The generation of H₂O²⁺ species under specific alkaline conditions (pH = 13) was also discovered by Wang et al. on an ill-defined Pt site of a commercial Pt/C catalyst, however, how to intentionally mediate and make good use of the H₂O²⁺ species remains a major challenge. Our aim in this work is to engineer this unique H₂O²⁺ species to create a desired local acidity under widespread alkaline conditions (Supplementary Fig. 10).

To obtain further information on the coordination and interactions between H₂O²⁺ and catalysts and to reveal the essential role of the Pt nanoparticles, we resorted to the operando XAFS measurements to observe the local coordination environment of the catalysts during the reaction. Operando XAFS is a very powerful tool for this purpose due to its element specificity and local structure sensitivity. Supplementary Fig. 11 shows the Pt L₂-edge X-ray absorption near-edge structure (XANES) spectra of Pt/MgO under the operando conditions. A comparison of the XANES spectral features of Pt/MgO with those of Pt foil and PtO₂ reveals that the Pt in Pt/MgO under the ambient conditions is in a metallic state, instead of a partially-oxidized state as reported for many other supported Pt nanoparticles²⁴–³⁶. In addition, the absence of Pt-O bonds in the k²-weighted Fourier transformed (FT) EXAFS spectra further excludes the surface oxidation of the Pt particles in Pt/MgO (Supplementary Fig. 12). To magnify the spectral changes induced by the applied potentials and to identify the adsorbates on Pt, the XANES difference, or their XANES intensities, we calculated the XANES spectra of Pt/MgO on the basis of the XANES spectra of Pt foil, PtO₂, and MgO (see Supplementary Information and Table 1).

Confirmation and formation mechanism of local acidic environment. We speculate that our synthesized Pt/MgO nanosheets can create a local acid-like environment under alkaline conditions. To test this speculation, we employed operando Raman spectroscopy and synchrotron radiation Fourier transform infrared spectroscopy (FTIR) under HER reaction conditions. These measurements will enable us to identify the key acidic intermediates of H₂O°/Pt/MgO during the HER process and to monitor their dynamic evolution. The operando measurements were conducted under three different conditions: open-circuit (immersed in KOH electrolyte), near the onset potential at a current density of –0.5 mA cm⁻² (~10 mV), and the overpotential at 10 mA cm⁻² (~40 mV). Figure 2a shows the Raman spectra of Pt/MgO against the applied potential. Interestingly, when the applied potential is reduced to ~40 mV vs. RHE, a new peak at ~1750 cm⁻¹, which is assigned to H₂O²⁺ intermediate species, is observed²⁹, whereas the peak of H₂O (1600 cm⁻¹) becomes weaker and the G-band of graphite (1580 cm⁻¹) remains unchanged, indicating the facilitated water dissociation on the surface of MgO and thereby the generation of abundant H₂O°/intermediates. This observation is also supported by the cutting-edge operando synchrotron Fourier transform infrared (SR-FTIR) spectroscopy, which is highly sensitive to the functional groups within ~3 molecular monolayers. The FTIR spectra of Pt/MgO display a progressively intensified absorption band at 3525 cm⁻¹ that is ascribed to the stretching vibrations of the O-H group in H₂O° when the applied potential is reduced to ~40 mV vs. RHE (Fig. 3b and Supplementary Fig. 8)³⁰,³¹; similar phenomena are observed on MgO nanosheets, but not on Pt/C (Supplementary Fig. 8), indicating the critical role of MgO in facilitating water dissociation and creating the H₂O²⁺ intermediate. More strikingly, the potential-dependent signal of H₂O²⁺ is reversible, manifesting that the local acid-like environment is formed only during the process of HER (Supplementary Fig. 9). In addition, it is worth noting that the detection limits of SR-FTIR spectroscopy and Raman spectroscopy are approximately ppm levels for hydroxyl groups or hydration molecules, which is much higher than that of the content of H⁺ (~10⁻¹⁴ M) ionized from water in the 1 M KOH solution. Such a high level of H₂O²⁺ cannot be ubiquitous in KOH but can accumulate only within a local region. The generation of H₂O²⁺ species under specific alkaline conditions (pH = 13) was also discovered by Wang et al. on an ill-defined Pt site of a commercial Pt/C catalyst, however, how to intentionally mediate and make good use of the H₂O²⁺ species remains a major challenge. Our aim in this work is to engineer this unique H₂O²⁺ species to create a desired local acidity under widespread alkaline conditions (Supplementary Fig. 10).

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that for OH-adsorbed Pt. When the applied potential is reduced to ~40 mV versus RHE, the magnitude of the ΔXANES results increases remarkably. This change is caused by the progressively increased adsorption coverage of H$_3$O$^+$ on Pt, indicating that the generated H$_3$O$^+$ intermediates are enriched in the proximity of Pt nanoparticles. Summarizing the above operando Raman, FTIR, and XANES spectra, we conclude that an abundance of H$_3$O$^+$ has been generated and forms a local H$_3$O$^+$ enriched, acid-like reaction environment around Pt nanoparticles in Pt/MgO.

To investigate the origin of the local acid-like environment created around these Pt nanoparticles, electron paramagnetic resonance (EPR), X-ray photoelectron spectroscopy (XPS), and XANES spectroscopy measurements were conducted on Pt/MgO in comparison to the Pt/C and MgO references. The EPR spectra of both Pt/MgO and MgO (Supplementary Fig. 13) show a strong signal at g = 2.00 arising from oxygen vacancies (V$_O$), suggesting a high concentration of V$_O$ in both the MgO and Pt/MgO. The abundance of oxygen vacancies as n-type doping could significantly improve the conductivity of MgO nanosheets. In addition, the oxygen vacancies of MgO are occupied by unpaired electrons (F centers), and the negative charge will transfer to the metal sites when the metal is trapped by the F centers, giving rise to the formation of electron-enriched Pt$_{\delta^-}$ species. This deduction is verified by the Pt 4f/XPS spectra as shown in Fig. 2d. The Pt 4f$_{5/2}$ XPS spectrum of Pt/MgO can be deconvoluted into two peaks at 71.4 and 70.6 eV. The 71.4 eV peak is ascribed to the Pt$^0$ state as seen in Pt foil. The 70.6 eV peak in the lower binding energy region can arise only from negatively-charged Pt atoms (Pt$_{\delta^-}$). More importantly, the Pt 4f XPS spectra is highly sensitive to the surface, which excludes surface oxidation of the Pt particles in Pt/MgO. The formation of Pt$_{\delta^-}$ in Pt/MgO is further confirmed by the Pt $L_3$-edge XANES spectra as shown in Fig. 2e. The main characteristic peak located at ~11568 eV in the Pt $L_3$-edge XANES spectra is the so-called white-line peak, which arises from an electron transition from the occupied Pt 2$p_{3/2}$ orbital to the empty 5d orbital, and thus is indicative of the Pt 5d occupancy. The slightly weaker white line of Pt/MgO relative to that of Pt foil thus indicates the higher 5d occupancy of Pt in agreement with the existence of Pt$_{\delta^-}$ therein. A similar phenomenon was also reported in TiO$_2$-supported metal catalysts due to strong metal-support interactions and in Pt-based alloys. In contrast, in the commercial Pt/C, electron transfer occurs from Pt nanoparticles to the carbon support, as evidenced by XPS (Fig. 2d) and XANES (Fig. 2e) spectra showing the Pt$_{\delta^+}$ states. All of these results suggest the strong electronic interactions between Pt nanoparticles and MgO at the interface, where electrons transfer from V$_O$ in MgO to the supported Pt nanoparticles (Supplementary Figs. 14, 15), resulting in the formation of negatively-charged Pt$_{\delta^-}$ at the interface (Fig. 2f). Due to the electrostatic attraction between the negatively-charged Pt$_{\delta^-}$ atoms and the positively-charged H$_3$O$^+$ species whose formation is facilitated by the V$_O$-enriched MgO, a local acid-like environment around Pt$_{\delta^-}$ nanoparticles is created.

**Electrocatalytic performance toward HER.** From the above operando Raman, SR–FTIR, and XANES characterizations, it is evidenced that a local acid-like reaction environment is created around Pt$_{\delta^-}$ nanoparticles in Pt/MgO, which is expected to be able to substantially boost HER activity. Thus, the electrocatalytic HER activity of Pt/MgO, commercial Pt/C, and MgO catalysts was evaluated in 1.0 M KOH. As a reference, the HER performance of commercial Pt/C was also evaluated under acidic conditions (the MgO is easily dissolved in acidic media). The Pt/MgO catalyst exhibits remarkable activity, with an overpotential of 39 mV at a current density of 10 mA cm$^{-2}$ (Fig. 3a), which is significantly lower than the 62 mV overpotential of 20 wt% Pt/C. Notably, the HER activity of the MgO support is almost negligible. To estimate the HER catalytic reaction kinetics, the Tafel slope is derived from the LSV curves, with values of 352, 68, and 39 mV dec$^{-1}$ for MgO, Pt/C, and Pt/MgO, respectively (Fig. 3b), indicating that Pt/MgO affords faster HER kinetics than Pt/C and follows the Volmer–Heyrovsky mechanism. More importantly, by comparing the LSV curves and the Tafel slopes of Pt/MgO with those of Pt/C under acidic and alkaline conditions, one can find that the alkaline HER performance of Pt/MgO is close to the acidic HER behavior of Pt/C, suggesting the acid-like HER behavior of Pt/MgO in an alkaline medium. The EIS curves of Pt/MgO (Supplementary Fig. 16) show an electrochemical resistance...
of 7 Ω, which is smaller than that of Pt/C (15 Ω), indicating the substantially facilitated interfacial electron-transfer kinetics in Pt/MgO. In addition, the durability and stability of Pt/MgO in 1.0 M KOH were implemented by an accelerated durability test (ADT) and chronoamperometry test as depicted in Fig. 3c. The ADT performance displays a slight drop after 10,000 cycles compared with the initial curve in 1.0 M KOH. Furthermore, the current density at 10 mA cm$^{-2}$ (the inset of Fig. 3c) of Pt/MgO remains virtually unchanged after 20 h of operation. More importantly, the TEM image (Supplementary Fig. 17) and XRD pattern (Supplementary Fig. 18) of Pt/MgO after continuous operation also maintain the original features, and the chemical states of Pt, Mg, and O as inferred from the XANES spectra (Supplementary Fig. 19) and XPS spectra (Supplementary Fig. 20) remain unchanged, confirming that Pt/MgO is stable in the long-term electrochemical process and that the negatively-charged Pt$^{\delta-}$ nanoparticles could not be oxidized during the HER. The EPR spectrum of Pt/MgO after the HER is also shown in Supplementary Fig. 21. It is obvious that the signal at $g = 2.00$ arising from oxygen vacancy still remains significant, suggesting the persistence of oxygen vacancies in Pt/MgO during HER. To further quantitatively compare the electrochemical activity of Pt/MgO with that of 20 wt% Pt/C, the mass activities at various potentials were determined, as illustrated in Fig. 3d. The mass current density of Pt/MgO reaches up to 0.53 mA µgPt$^{-1}$ at $-39$ mV vs. RHE, tenfold higher than that of 20 wt% Pt/C (0.05 mA µgPt$^{-1}$) in alkaline medium. Moreover, the ECSA-normalized HER polarization curves (Supplementary Fig. 22) and the turnover frequency (TOF) (Supplementary Fig. 23) suggest that the improvement in HER catalytic performance is mainly due to the increased catalytic activity of the intrinsic active sites. Reducing the catalyst loading on the cathodes of Pt/C and Pt/MgO has no impact on the interaction between H* and the catalyst surfaces (Supplementary Fig. 23), further confirming the high intrinsic HER activity of Pt/MgO. It is worth noting that the mass current activity of Pt/MgO is 2.5-fold higher than that of similarly sized 20 wt% Pt/C (0.2 mA µgPt$^{-1}$) in an acidic medium. This performance is due to the existence of the electron-enriched Pt$^{\delta-}$ as will be discussed later. In addition, the Pt/MgO shows nearly 100% Faradaic efficiency in base (Supplementary Fig. 24). More importantly, a series of experiments and simulations, including the synthesis of negatively-charged Pt$^{\delta-}$ nanoparticles supported on TiO$_2$ (without strong water-dissociation ability), reducing the quantity of oxygen vacancies on MgO, and the calculations of the water-dissociation ability of negatively-charged Pt$^{\delta-}$ (Supplementary Figs. 25–29), show that the high HER activity comes from the synergistic effect of the local reaction environment and the negatively-charged Pt$^{\delta-}$, rather than only from the modified electronic structure of Pt. These results confirm that Pt/MgO possesses the high intrinsic HER activity, in accord with the expectations based on the local acidic environment. In addition, we synthesized other noble metal (Au, Ru, and Ir) nanoparticles supported on MgO to replace Pt (Supplementary Figs. 30–33). Compared with that of Ru/C and Ir/C, the HER performance of Ru/MgO and Ir/MgO is much better, indicating that an abundance of H$^+$ created by MgO can also improve the alkaline HER activity of other noble metals. Structural characterizations show that negatively-charged metal sites are also formed on the Ir surface (Supplementary Fig. 34). Nevertheless, the HER performance of Ir/MgO is still not as good as that of Pt/MgO (Supplementary Fig. 35), probably due to the inherent differences between Pt and Ir such that Pt has the most approachable zero hydrogen absorption energy.
Density functional theory (DFT) calculations. Based on the aforementioned operando spectroscopic observations that MgO can create a local acid-like reaction environment around Pt\(\delta^+\), DFT calculations were conducted to understand how the oxygen vacancies in MgO influences the charge density distribution of supported Pt and how the various electronic structures of Pt species affects the local reaction environment and HER activity. As shown in Fig. 4a–c, compared with the cases of Pt and Pt supported on oxygen vacancy-free MgO, in Pt supported on oxygen vacancy-rich MgO, strong electron injection occurs from oxygen vacancies occupied by unpaired electrons (F centers) into Pt nanoparticles, which leads to the formation of electron-enriched Pt species (Pt\(\delta^–\)). The Bader charge analysis results further shows that approximately 0.3e electrons are transferred from oxygen vacancy-rich MgO to the Pt atoms adjacent to the oxygen vacancy, and no significant electron transfer occurs between oxygen vacancy-free MgO and Pt nanoparticles. Next, we explored the dissociation of H\(_2\)O molecules followed by a proton migration process on the MgO (001) surface as shown in Figs. 4d, 5e. A water molecule is dissociated into an OH\(^–\) and a chemisorbed H\(^+\) species, with an energy barrier of \(-1.7\) eV on the MgO (001) surface, which is significantly lower than that on Pt (0.06 eV)\(^{44}\), RhO\(_2\) (0.35 eV)\(^{45}\), and so on (Supplementary Fig. 36). This finding is demonstrated by the carbon monoxide (CO) stripping tests (Supplementary Fig. 37). The CV curves show that the stripping peak for CO\(_{ad}\) oxidation of Pt/MgO has a more negative value than that for Pt/C, indicating a better water-dissociation ability for Pt/MgO over that of Pt/C\(^{29}\). The negligible energy barrier for breaking the OH–H bonds suggests that the water-dissociation step on MgO is energetically favorable on Mg sites. The large distance between H\(^+\) adsorbed on the lattice O\(_2–\) atoms and OH\(^–\) on the lattice Mg\(_2+\) atoms stabilizes the dissociated structure due to the weak intermolecular interactions. Next, the free energy diagram was constructed to gain further thermodynamic insight into the migration process of H\(^+\) dissociated from water molecules, as shown in Fig. 4d, e and Supplementary Fig. 38. Along the path indicated by the arrows, the migration energy of H\(^+\) decreases gradually, indicating the strong tendency of the H\(^+\) to accumulate around Pt\(\delta^+\) nanoparticles. Considering the fact that V\(_{O}\)-rich MgO creates a local acid-like environment for negatively-charged Pt\(\delta^–\), we calculated the Gibbs free energy of hydrogen adsorption (\(\Delta\)G\(_{H^+}\)), which is widely accepted as a universal descriptor for the HER in acid, to judge the activity of Pt in the step of hydrogen adsorption. We examined the effects of three different charge states (Pt\(\delta^–\), Pt\(\delta^0\), and Pt\(\delta^+\)) of the Pt nanoparticles on the HER activity (Supplementary Fig. 39). The calculated partial density of states (PDOS) of Pt d orbitals with different charge states are shown in Fig. 4f, where the d-band center position is explicitly labeled. Apparently, Pt\(\delta^–\) has the lowest d-band center position (\(-2.74\) eV), as compared with that of Pt\(\delta^0\) (\(-2.49\) eV) and Pt\(\delta^+\) (\(-2.06\) eV). According to the d-band theory, a lower d-band center (farther away from Fermi level) of metal-based catalysts leads to weaker interaction between the 1s orbital of hydrogen and the unfilled d orbitals of the metals\(^{46}\). To verify this point, we calculated the Gibbs free energy (\(\Delta\)G\(_{H^+}\)) of H\(^+\) adsorption on the surfaces of Pt\(\delta^–\), Pt\(\delta^0\), and...
alkaline electrolyte (Fig. 5). The oxygen vacancy-rich MgO molecules to form H3O+ for the H+ lowering the energy barrier. Beneath the potential, and then promote the hydrogen evolution reaction by the environment will shift the pzfc of Pt toward the HER equilibrium.

Based on the above experimental and theoretical results, we propose an HER mechanism for the Ptδ− nanoparticles in an alkaline electrolyte (Fig. 5). The oxygen vacancy-rich MgO facilitates the water dissociation and the surface Ptδ− species promote H2O+ migration and accumulation to achieve the acid-like HER activity and kinetics in alkaline media. The specific reaction mechanism is as follows: First, a water molecule diffuses to an oxygen vacancy site of the MgO surface, where it is adsorbed and dissociated into an OH* and an H*. As the water-dissociation reaction proceeds, a great many H* species are generated which subsequently combine with adjacent H2O molecules to form H2O+. Due to the lower migration energy for the H* and the additional electrostatic attraction between the positively-charged H2O+ and the negatively-charged Ptδ−, the H* easily migrates throughout the rigid water network of the double layer caused by the positive shift in the potential of zero free charge (pzfc) to form a local acid-like environment around Ptδ−. Within the framework of pzfc, the local acid-like environment will shift the pzfc of Pt toward the HER equilibrium potential, and then promote the hydrogen evolution reaction by lowering the energy barrier. Benefiting from the local acid-like reaction environment, afterward, H2O+ then adsors hydrogen atoms H* on the Ptδ− surface and later combines with another H* to generate H2 following the Tafel step. Meanwhile, the remaining OH* species in the oxygen vacancy will combine with a hydrated K+ ion to form a hydroxyl-water-caton (OHad−(H2O)xK+−) adduct. The existence of OHad−(H2O)xK− was confirmed by the deuterium (2H, D) nuclear magnetic resonance (NMR) and CO stripping experiments (Supplementary Fig. 41). Then, the adduct quickly transfers to the electrolyte across the electrical double-layer, causing no change in the local OH− concentration24. After the release of OH*, the oxygen vacancy is restored, and another water molecule diffuses to the oxygen vacancy site and continues the next reaction cycle. As a result, the V0−-rich MgO/Pt catalyst can continuously and stably produce and accumulate a large amount of H2O+, thus creating a local acid-like environment. The synergistic effect of the local acid-like reaction environment and electron-enriched Ptδ− species endow the catalyst with excellent intrinsic HER activity.

**Discussion**

In summary, we have developed a facile pathway to create a local acid-like reaction environment in an alkaline medium. Taking the V0−-MgO nanosheet supported Pt as a prototypical example, this catalyst is favorable for water dissociation and the formation of electron-enriched Ptδ− species. A combined study of operando Raman, SR-FTIR, and XAFS spectra evidences that negatively-charged Ptδ− species accelerate positively-charged H2O+ migration and accumulation, resulting in a local acid-like environment. This acid-like environment provides Pt with a favorable reaction condition for the HER in the alkaline electrolyte. As a result, the Pt/MgO catalyst shows an overpotential of 39 mV at a current density of 10 mA cm−2, which is significantly lower than the 62 mV of 20 wt% Pt/C in an alkaline medium and close to the acidic HER behavior of Pt/C (33 mV). Our study provides insight into tailoring the local reaction environment to design high-performance electrocatalysts in a more rational and precise way.

**Methods**

**Material characterizations.** The TEM images, the HRTEM, and EDS mapping analyses were performed on a JEM-2100F field emission transmission electron microscope at an acceleration voltage of 200 kV. XPS spectra were acquired on an ESCALAB MKII with Mg Ka (hν = 1253.6 eV) as the excitation source and were corrected for specimen charging by referencing C 1s to 284.5 eV. The EPR measurements were performed in a JES-FA200E EPR spectrometer at X-band (−9 GHz) with a resolution of 2.35 μT at 300 K. The Pt L-edge X-ray absorption near-edge (XANES) spectra were measured at the 1W1B beamline of Beijing Synchrotron Radiation Facility (BSRF). The O K-edge XANES spectra were collected at the BL12B beamlines of NSRL (Hefei, China), respectively.

**Synthesis of MgMOF-74 precursor.** Mg(NO3)2·6H2O (0.4 g) and 2,5-dihydroxy benzene carboxylic acid (0.09 g) were mixed and dissolved in 40 mL of N,N-dimethylformamide (DMF). A mixture of ethanol, water, and triethylamine mixtures (9:9:1, v/v/v) was added to the above solution and stirred for 3 h at room temperature. The resulting MgMOFs were collected by centrifugation and then re-dispersed in DMF. The suspension was treated at 100 °C for 2 h to dissolve other amorphous impurities.

**Synthesis of Pt/MgO and MgO.** In a typical procedure, a mixture of 50 mg of MgMOF and 3 mL of aqueous H2PtCl6 solution (10 mg/mL) was dissolved in 8 mL of deionized water and ultrasonically treated for 10 min. After magnetic stirring for 24 h at room temperature, the precipitates were collected by centrifugation and then freeze-dried to obtain the precursor. The precursor was then placed into a tube furnace and heated to 700°C in a mixture of oxygen and nitrogen (O2:N2 = 5:95) for 4 h at a heating rate of 10°C/min to obtain the desired Pt/MgO. MgO was synthesized in a similar way without adding an aqueous H2PtCl6 solution.

**DFT calculation details.** The DFT calculations were carried out with the Quantum Espresso software package48. The Generalized gradient approximation (GGA) in the Perdew–Burke–Ernzerhof (PBE) parametrization and projected augmented
Data availability
All data generated in this study are provided in the Supplementary Information/Source Data file. Source data are provided with this paper.

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