

Cite this: *J. Mater. Chem. A*, 2025, **13**, 19717

Cohesive Co and Mo₂C heterostructure catalysts strongly confined to hollow carbon support for enhanced kinetics and durability in the alkaline hydrogen evolution reaction†

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Water electrolysis technologies in alkaline environments have drawn considerable interest for their potential in low-cost hydrogen production without noble metals. However, it remains challenging to achieve high performance and durability in the alkaline hydrogen evolution reaction (HER) using non-noble metal catalysts because of their intrinsically poor water dissociation kinetic properties. Here, we address this challenge by introducing a cohesive Co and Mo₂C heterostructure catalyst strongly confined to N-doped carbon hollow polyhedron (NCHP) supports. Computational analysis and X-ray spectroscopic analysis results reveal that the charge redistribution between metallic Co and Mo₂C not only promotes water dissociation on Mo₂C sites but also accelerates hydrogen gas evolution kinetics at their interfaces, leading to enhanced HER performance. The Co–Mo₂C/NCHP catalyst reduces the HER overpotential by 66% at 10 mA cm^{−2} compared to the single-active-site catalyst (Co/NCHP). Moreover, the Co–Mo₂C/NCHP catalyst demonstrates enhanced durability, exhibiting a 38% lower HER overpotential than commercial Pt/C at 10 mA cm^{−2} after 1000 cycles. By forming a cohesive structure of Co and Mo₂C within the hollow carbon framework, the formation of interfaces is maximized, and physical and chemical degradation is prevented. This study presents key strategies for designing interfaces to overcome the limitations of alkaline HER kinetics and durability.

Received 2nd April 2025
Accepted 22nd May 2025

DOI: 10.1039/d5ta02607e

rsc.li/materials-a

Introduction

Hydrogen energy is one of the most important alternatives to fossil fuels because of its abundance and lack of harmful emissions.^{1,2} Among hydrogen production methods, water electrolysis, particularly anion exchange membrane water electrolysis (AEMWE), has attracted significant interest in recent years.^{3–7} AEMWE offers a promising solution by addressing the high material costs of polymer electrolyte membrane water electrolysis (PEMWE) and overcoming the low operating current

density and limited dynamic response of liquid alkaline water electrolysis.^{8,9}

A key challenge in advancing AEMWE is achieving efficient hydrogen evolution reaction (HER) with low-cost materials, comparable to Pt.¹⁰ In alkaline environments, water dissociation significantly impacts HER kinetics.^{11–13} Thus, water dissociation energy, along with the hydrogen adsorption energy, has been identified as a key activity descriptor for the alkaline HER.¹⁴ While some 3d transition metals, like Co and Ni, have shown potential as cost-effective catalysts for alkaline HER,^{15–18} their intrinsic activity remains low^{19–22} primarily due to unfavorable water dissociation energetics.^{23,24} Consequently, designing catalysts that can effectively modulate both water dissociation and hydrogen adsorption energies is crucial for enhancing HER activity in alkaline environments.

A feasible solution to overcome this limitation is the construction of synergistic dual active sites for water dissociation and hydrogen evolution,¹⁴ particularly to improve the water dissociation activity on the 3d transition metal catalyst surfaces. Furthermore, heterogeneous structures incorporating these dual active species can modulate their electronic structure at the interface, thereby further enhancing the intrinsic catalytic performance for each reaction.^{25–27} Among various candidates,

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d5ta02607e>

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Mo-based compounds, particularly Mo₂C, have been reported to achieve remarkable improvements in HER activity due to their ability to alter the Volmer reaction kinetics relating to water dissociation.^{28–31} This enhanced water dissociation capability is attributed to the unique Pt-like electronic structure of Mo₂C, which optimizes the interaction between water molecules and active sites, thereby lowering the energy barrier for hydrogen evolution. For instance, S. Sun *et al.* demonstrated that protons generated by the Mo₂C sites *via* water dissociation can be immediately reduced by adjacent MoO_x sites to promote overall HER activity.²⁸ Y. Zhan *et al.* also demonstrated the synergistic effect between highly conductive metallic Ni and Mo₂C–MoN for improving HER activity.²⁹

Despite these advantages, the primary limitation of Mo compounds is the poor stability of Mo₂C in alkaline environments, which leads to its transformation into less active phases, such as MoO_x, thereby degrading long-term performance.^{32–34} To achieve an effective dual active site for alkaline HER with Mo₂C, structural strategies are needed to address the instability and to foster the formation of a synergistic interface.

Metal–organic frameworks (MOFs) offer a versatile platform for catalyst design by forming well-defined structures through the coordination of metal ions with organic ligands.³⁵ Moreover, the catalytic properties of MOFs can be further tailored through a carbonization process, which enables the formation of unique carbon structures with embedded metal components. The strong confinement of metallic particles within MOF-derived carbon can create heterogeneous structures with synergistic properties,³⁶ while also effectively protecting them from the harsh dissolution conditions often encountered in alkaline environments.^{37,38} Additionally, MOF-derived carbon provides the numerous nanopores adjacent to metallic particles, which can enhance gas bubble detachment and improve catalyst performance during HER.^{39,40} Evolved gas bubbles can block active sites and hinder ion transport, reducing catalyst performance.^{41–43} Therefore, the unique ability of carbonized MOF to create heterogeneous structures, provide strong confinement of active sites, and facilitate gas bubble detachment makes it a promising strategy for developing high-performance and durable catalysts for alkaline HER.

In this study, we propose a catalyst design strategy that employs two distinct MOFs to construct a unique hollow catalyst structure with cohesive and integrated dual active sites of Co and Mo₂C. The strategy utilizing MOFs as a framework introduces and effectively confines Co and Mo₂C nanoparticles within an N-doped carbon hollow polyhedron (NCHP) after the carbonization process. This cohesive structure generates abundant Co–Mo₂C dual active sites with strong interfacial interactions, enhancing both water dissociation and hydrogen evolution kinetics. Additionally, the carbon-embedded catalyst structure has beneficial effects in suppressing chemical degradation of active phases, such as Mo leaching.²⁹ Furthermore, the NCHP, with sub-nanometer-sized pores, can facilitate early and efficient bubble desorption, mitigating the physical degradation of active phases.⁴⁴ Consequently, the Co–Mo₂C/NCHP catalyst exhibited enhanced alkaline HER performance and durability, compared to the single-site catalysts and commercial

Pt/C. These strategies hold immense promise for advancing AEMWE research by addressing the key challenges to achieving efficient HER with non-noble metal catalysts, and paving the way for the development of next-generation high-performance and durable catalysts.

Experimental section

Chemicals and materials

All chemicals were purchased from commercial sources and directly used without further pretreatment; zinc nitrate hexahydrate (Zn(NO₃)₂·6H₂O, 96%) and cobalt nitrate hexahydrate (Co(NO₃)₂·6H₂O, 97%) were obtained from Junsei Chemical. 2-Methylimidazole (C₄H₆N₂, 99%), ammonium molybdate tetrahydrate ((NH₄)₆Mo₇O₂₄·4H₂O, 99%), polyvinylpyrrolidone (PVP, (C₆H₉NO)_n, M_w = ~55 000), and *N,N*-dimethylformamide (DMF, 99.8%) were obtained from Sigma-Aldrich. Commercial Pt/C catalyst (TEC10V20E, 20 wt% Pt) was purchased from Tanaka Kikinzoku Kogyo. Pt/C (TEC10E50E, 47 wt% Pt) and IrO₂ (Premion®, Ir 84.5%) catalysts for single-cell electrode fabrication were purchased from Tanaka Kikinzoku Kogyo and Alfa Aesar, respectively. Ethanol (C₂H₅OH, 95%) was purchased from Samchun Chemicals. Deionized water with a specific resistance of 18.2 MΩ cm was obtained using a Millipore-DQ5 purification system.

Synthesis of ZIF-8. An amount of 5.95 g of Zn(NO₃)₂·6H₂O was dissolved in 150 mL of methanol. Then, the solution was mixed with 150 mL of methanol solution containing 6.15 g of 2-methylimidazole and stirred at room temperature for 24 hours. After that, the product was collected by centrifugation, washed with methanol three times, and dried at 50 °C.

Synthesis of ZIF-8@ZIF-67. Then, 0.5 g of ZIF-8 was dispersed in 100 mL of methanol. Then, the solution was mixed with 100 mL of methanol solution containing 4.07 g of Co(NO₃)₂·6H₂O, and the solution of 4.31 g of C₄H₆N₂ was slowly added dropwise into the above mixture solution through a burette. After the solutions were mixed and stirred at room temperature for 24 hours, the product was collected by centrifugation and washed with methanol three times and dried at 50 °C.

Synthesis of Mo-ZIF-8@ZIF-67. 240 mg of ZIF-8@ZIF-67, 200 mg of PVP, and 43.8 mg of (NH₄)₆Mo₇O₂₄·4H₂O were dissolved in 10 mL of DMF. Then, the mixture was heated at 150 °C for 12 hours. The precipitate was separated by centrifugation, washed with DMF, water, and ethanol, and dried at 50 °C.

Synthesis of Co–Mo₂C/NCHP and Mo₂C/NCHP. The Mo-ZIF-8@ZIF-67 was placed in a tube furnace, heated to 900 °C with a ramp rate of 5 °C min⁻¹, and kept for 3 hours in flowing Ar. The obtained black powder was washed in 1.0 M H₂SO₄ solution at room temperature for 8 hours to remove any accessible Zn species on its surface, yielding Co–Mo₂C/NCHP. For the synthesis of Mo₂C/NCHP, the remaining conditions were kept the same, except that the amount of Mo precursor was doubled, and acid leaching was performed in a 2.5 M H₂SO₄ solution for 12 hours.

Synthesis of Co/NCHP. The ZIF-8@ZIF-67 was placed in a tube furnace, heated to 900 °C at a ramp rate of 5 °C min⁻¹,



and kept for 3 hours in flowing Ar. The obtained black powder was washed in 1.0 M H₂SO₄ solution at room temperature for 8 hours to remove any accessible Zn species on its surface, yielding Co/NCHP.

Synthesis of Mo₂C/Co/NCHP. 12 mg of (NH₄)₆Mo₇O₂₄·4H₂O and 120 mg Co/NCHP were mixed in DI and then these slurries were dried at 100 °C. For carbonization, the obtained powder was placed in a tube furnace, heated to 700 °C at a ramp rate of 5 °C min⁻¹, and kept for 2 hours under flowing H₂ (3.9%)/Ar.

Materials characterizations. Crystallographic investigations were performed using a Powder X-ray Diffractometer (XRD, SmartLab, RIGAKU) with a 1.5406 Å Cu-Kα radiation source. Transmission electron microscopy (TEM) images were obtained using Tecnai G2 F30 S-Twin and Talos F200X devices equipped with an energy dispersive X-ray (EDX) spectrometer and high-angle annular dark-field scanning TEM (HAADF-STEM). The catalysts were analyzed semi-quantitatively using X-ray photoelectron spectroscopy (XPS, K-alpha, Thermo VG Scientific) and inductively coupled plasma optical emission spectroscopy (ICP-OES, Agilent ICP-OES 720, Agilent). The Brunauer–Emmett–Teller (BET, 3Flex, Micromeritics) surface area and pore size were estimated by measuring the N₂ isotherms using a surface area and pore size analyzer. Thermal stability was confirmed using thermogravimetric analysis (TGA, TG209 F1 Libra, Netzsch) at a heating rate of 10 °C min⁻¹ under N₂ flow. Raman spectra were measured using a dispersive Raman spectrometer (ARAMIS, Horiba Jobin Yvon) with a 514 nm excitation source. FT-IR spectra were collected on a FT-IR spectrophotometer (Nicolet iS50, Thermo Fisher Scientific Instrument) using the KBr pellet method. All characterizations mentioned above were carried out at the KAIST Analysis Center for Research Advancement (KARA). X-ray absorption fine structure (XAFS) spectra were measured using an 8C nanoprobe XAFS beamline (BL8C) of the Pohang Light Source (PLS-II) in the 3.0 GeV storage ring, with a ring current of 360 mA. The XAFS spectra of the powder samples were collected in both the transmission and fluorescence modes. The acquired data were processed using the Athena module in the IFEFFIT software package.

Electrochemical measurements. The electrochemical performance was measured in a conventional three-electrode cell using PGSTAT 302N (Metrohm AUTOLAB). For electrochemical measurements, working electrodes were prepared by uniformly spreading slurries (8.8 μL) of the catalysts onto a glassy carbon electrode (GCE, 0.196 cm²) to achieve a loading of 0.45 mg cm⁻². Each slurry was composed of a catalyst powder (10 mg), 80 μL of 5% Nafion solution (1100 W, Aldrich), and 920 μL of ethanol. The counter electrode was a Pt mesh, and the reference electrode was a Hg/HgO electrode filled with 20% sodium hydroxide (NaOH) solution. All potentials were converted into reversible hydrogen electrode (RHE) measured potentials and are reported in this work. Linear sweep voltammetry (LSV) curves were obtained in a H₂-purged 1.0 M KOH solution at a scan rate of 5 mV s⁻¹. All polarization curves were IR corrected according to the equation $E = E_m - iR$ (where E is the corrected potential, E_m is the measured potential, and R is the resistance of the solution). Tafel slopes were calculated by plotting the overpotential as a function of the logarithm of the

current density, providing valuable insights into the mechanism of the electrocatalytic reaction. For an accelerated durability test (ADT), 1000 potential cycles were carried out ranging from 0.05 V_{RHE} to -0.50 V_{RHE} in a 1.0 M KOH solution at a scan rate of 50 mV s⁻¹. Electrochemical impedance spectroscopy (EIS) testing was performed at -0.3 V_{RHE},⁴⁵ in the frequency range of 10 kHz to 0.1 Hz. A single cell was assembled with catalyst-coated electrodes and a commercial Fumasep FAA-3-50 membrane (Fumatech). The electrodes were fabricated by ultrasonic spray coating (Nadetech) of catalyst materials onto a carbon gas diffusion layer (JNTG, JNT30-A3) for the cathode, and Ni fiber felt (Bekaert, 2NI 18-0.25) for the anode. The active catalyst-coated area is 4 cm². Catalyst loading of Pt and Ir on each electrode was 0.20 ± 0.3 and 0.85 ± 0.3 mg cm⁻², respectively. The i - V curves were measured at 60 °C with a supply of 1 M KOH solution.

Computational details. Density functional theory (DFT) calculations were conducted using the Vienna *Ab initio* Simulation Package (VASP) with projector augmented wave (PAW) potentials.⁴⁶ The spin-polarized generalized gradient approximation of Perdew, Burke, and Ernzerhof (GGA-PBE) was utilized for the exchange–correlation function, and van der Waals correction of Grimme was applied.⁴⁷ The FCC Co (111) and the β-Mo₂C (101) models were constructed with a Monkhorst–Pack k -point grid of 3 × 3 × 1. The Co–Mo₂C model was calculated with a Monkhorst–Pack k -point of 1 × 1 × 1. In addition, Co–Mo₂C was adopted as a lateral heterostructure since both Co and Mo₂C were exposed on the surface as shown in the HRTEM in Fig. 1c.

In this model, Co(111) $p(3 \times 2)$, which had lattice parameters of 14.93 Å × 9.95 Å × 26.10 Å with $\alpha = 90^\circ$, $\beta = 90^\circ$, and $\gamma = 120^\circ$, was combined with Mo₂C (101) $p(2 \times 1)$, which had lattice parameters of 15.28 Å × 6.03 Å × 25.83 Å with $\alpha = 90^\circ$, $\beta = 90^\circ$, and $\gamma = 113.3^\circ$, through the fully relaxed optimization. The optimized Co–Mo₂C model had lattice parameters of 14.93 Å × 16.69 Å × 26.10 Å with $\alpha = 90^\circ$, $\beta = 90^\circ$, and $\gamma = 120^\circ$. Mo₂C experienced a 2.33% compressed strain along with the a -axis and a 5.22% tensile strain in the gamma angle, which was the interaxial angle between a -axis and b -axis. The model where C was directly bonded with Co showed the most stable surface energy of 0.26 eV Å⁻². As shown in Fig. S1c,† the Co–Mo₂C heterostructure was divided into two regions based on the interface: the Co/C interface, where C atoms of Mo₂C bonded with Co, and the Co/Mo interface, where Mo atoms of Mo₂C bonded with Co. The models were isolated with a vacuum of 20 Å, and the bottom half of the layers were fixed to simulate bulk properties.

The DFT models are shown in Fig. S1a–c,† and the d-band center of metals in Co–Mo₂C exhibited similar values to the individual Co and Mo₂C models in Fig. S1d.† This similarity in the d-band centers indicated that the heterostructure did not undergo excessive distortion during the modeling process. For density of states (DOS) calculations, a Monkhorst–Pack k -point grid of 6 × 6 × 1 was used for the Co and Mo₂C, and 2 × 2 × 1 was used for Co–Mo₂C. The Methfessel–Paxton smearing with a width of 0.2 eV and a cutoff energy of 450 eV were applied. Ionic relaxations were converged until the energy change was less than 10⁻⁵ eV, and electronic optimizations were converged



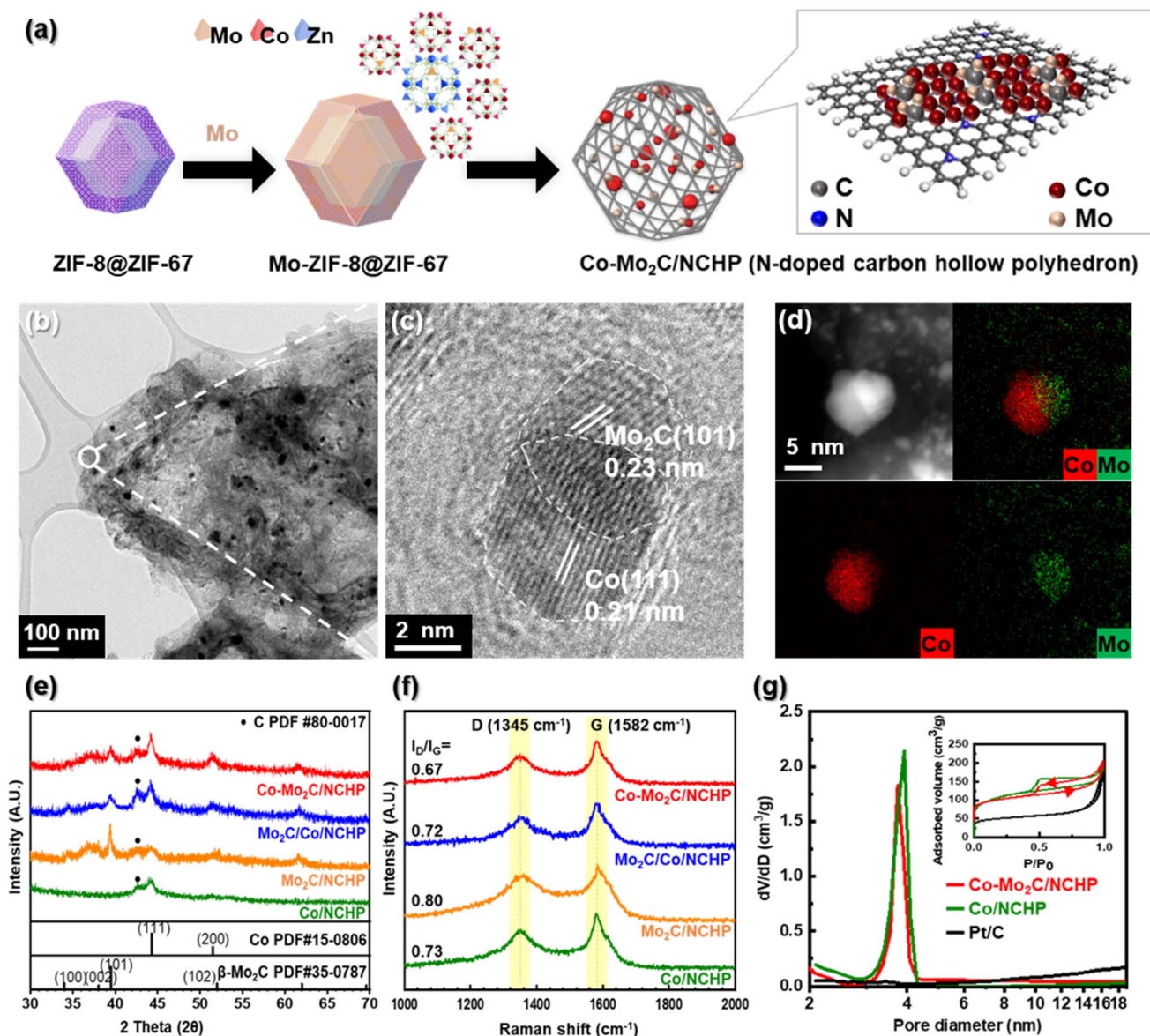


Fig. 1 (a) Schematic illustration of the preparation, (b) TEM image, (c) HRTEM image, and (d) HAADF-STEM elemental mapping results of Co-Mo₂C/NCHP. (e) XRD patterns and (f) Raman spectra of Co/NCHP, Mo₂C/NCHP, Mo₂C/Co/NCHP, and Co-Mo₂C/NCHP. (g) BJH desorption pore size distribution and N₂ adsorption/desorption isotherms (inset) of Pt/C, Co/NCHP, and Co-Mo₂C/NCHP.

until the forces were smaller than 0.03 eV \AA^{-1} . VESTA was utilized to visualize the results of calculations and models.⁴⁸ Activation energies for water dissociation were calculated using the climbing image-nudged elastic band (CI-NEB) method.^{49,50} The transition states (TS) were confirmed by a single imaginary mode. The Gibbs free energy was calculated with $\Delta G = \Delta E_0 + \Delta ZPE - T\Delta S$ at 298.15 K and 1 atm.

The charge of each model (ρ) was calculated by Bader charge analysis.⁵¹ The charge density difference ($\Delta\rho$) was calculated using the following equation.

$$\Delta\rho_{\text{diff}} = \rho_{\text{Co-Mo}_2\text{C}} - \rho_{\text{Co}} - \rho_{\text{Mo}_2\text{C}}$$

A positive sign indicates electron depletion, while a negative sign indicates electron accumulation.

Results and discussion

Fig. 1a schematically illustrates the Co-Mo₂C/NCHP catalyst prepared *via* a two-step approach. The synthesis begins with the formation of a core-shell metal-organic framework (MOF) structure, where a cobalt-organic framework (ZIF-67) is grown on a zinc-organic framework (ZIF-8) (Fig. S2†).⁵² This core-shell ZIF-8-ZIF-67 (ZIF-8@ZIF-67) structure is then subjected to Mo ion incorporation. The XRD analysis (Fig. S3†) indicates that the Mo incorporation process into the ZIF-8@ZIF-67 structure does not alter the overall crystal structure, as there are no significant changes in peak positions compared to the ZIF-8@ZIF-67 structure. This observation confirms that the crystal structure remains intact during the Mo incorporation process, but it does not directly confirm the presence of Mo ions.



Subsequently, the Mo incorporated ZIF-8@ZIF-67 (Mo-ZIF-8@ZIF-67) structure undergoes pyrolysis under an inert atmosphere at 900 °C. This pyrolysis process leads to the decomposition of the thermally less stable ZIF-67 shell and the collapse of the ZIF-8 core (Fig. S4a–c†). As the shell decomposes, it forms graphitized carbon that contributes to the creation of N-doped carbon polyhedron (NCHP). During this process, cobalt nanoparticles and Mo₂C are embedded within the NCHP structure, and they form interfaces that play a key role in enhancing the catalytic activity. These dual active sites—Co and Mo₂C—supported on NCHP are essential for the observed catalytic performance. The utilization of Zn as the sacrificial metal in the core-shell Mo-ZIF-8@ZIF-67 structure offers several advantages. During pyrolysis and subsequent acid leaching, Zn is effectively removed due to its volatility and solubility, which suppresses metal sintering and contributes to the formation of porous NCHP. This is further supported by X-ray photoelectron spectroscopy (XPS) of Co–Mo₂C/NCHP (Fig. S5†), which confirms negligible Zn content after thermal treatment.

To elucidate the synergetic effects of the Co and Mo₂C, catalysts with different metal compositions and configurations were synthesized for comparison (Fig. S6†). The Mo₂C supported on NCHP (Mo₂C/NCHP) catalyst was prepared through the selective acid leaching of Co from Co–Mo₂C/NCHP, resulting primarily in Mo₂C within the NCHP. The Co supported on NCHP (Co/NCHP) catalyst was synthesized by the pyrolysis of the core-shell ZIF-8@ZIF-67 without Mo addition. The Mo₂C supported on Co/NCHP (Mo₂C/Co/NCHP) catalyst was synthesized using an additional heat treatment of the Co/NCHP catalyst mixed with Mo precursor. This additional Mo introduction method may lead to a different Mo distribution pattern within the NCHP compared to the Co–Mo₂C/NCHP catalyst prepared with the one-step heat treatment. Furthermore, the XPS Zn 2p spectra (Fig. S5†) indicate effective removal of Zn from Co/NCHP, Mo₂C/NCHP, and Mo₂C/Co/NCHP during the synthetic process.

Fig. 1b and c depict the characterization of the Co–Mo₂C/NCHP catalyst after thermal treatment. The transmission electron microscope (TEM) images reveal a uniform distribution of both Co and Mo throughout the NCHP structure, with their interfaces formed between them. The Co and Mo₂C particles are embedded in the graphitized carbon structure, as evidenced by TEM images that show the layering of graphite (Fig. S7a†). This uniform distribution can be attributed to the inherent properties of the MOF precursor, where metal ions are arranged regularly alongside organic ligands.⁵³ Notably, the high-resolution TEM (HRTEM) image further showcases crystalline domains within the catalysts with d-spacings of 0.21 nm and 0.23 nm, corresponding to the (111) planes of metallic Co and the (101) planes of Mo₂C, respectively. As illustrated in Fig. 1d, S7b and c,† the sizes of the Co nanoparticles range from single atoms to ~5 nm, while the Mo₂C particles also exist as small nanoparticles, several nanometers in size. The EDX spectrum (Fig. S7d†) confirms the presence of both Co and Mo, and the inset of Fig. S7d† displays the elemental compositions determined by inductively coupled plasma optical emission spectrometry (ICP-OES), which shows that Mo is present at 5.5 wt%

and Co at 4.5 wt%. The cohesive Co–Mo₂C heterostructure embedded within the NCHP framework can lead to the formation of numerous interfaces.

Fig. 1e presents the X-ray diffraction (XRD) patterns of the Co/NCHP, Mo₂C/NCHP, Co–Mo₂C/NCHP, and Mo₂C/Co/NCHP catalysts. Peaks observed at 42.5°, 44.3°, and 51.5° correspond to the diffraction angles of C (101) and Co (PDF#15-0806), confirming the successful reduction of Co ions to metallic Co nanoparticles within the NCHP support after pyrolysis. Additionally, peaks at 34.0°, 38.0°, 39.5°, 52.0°, 62.1°, and 69.8° correspond to the diffraction angles of β-Mo₂C (PDF#35-0787), a molybdenum carbide known for its exceptional hydrogen evolution reaction (HER) activity.⁵⁴ This observation confirms the successful incorporation of Mo into the catalyst structure as Mo₂C.

For comparison purposes, TEM images (Fig. S8†) and high-angle annular dark-field scanning TEM (HAADF-STEM) elemental mapping and line profiles (Fig. S9†) were obtained for the Co/NCHP, Mo₂C/NCHP, and Mo₂C/Co/NCHP catalysts. In the case of the Co/NCHP catalyst, Co nanoparticles are observed in the NCHP structure. The Mo₂C/NCHP catalyst exhibits a lower Co content and a relatively larger amount of Mo in the form of larger nanoparticles compared to the Co–Mo₂C/NCHP catalyst. The Mo₂C/Co/NCHP catalyst displays a noticeable depletion of Mo in specific regions, as evidenced by the elemental line profile. This observation suggests there is a non-uniform Mo distribution pattern compared to the Co–Mo₂C/NCHP catalyst.

The Raman spectrum presented in Fig. 1f provides insights into the structural disorder and graphitic nature of the carbon support. The spectra display two characteristic bands at 1345 cm⁻¹ and 1582 cm⁻¹, corresponding to the D and G bands, respectively.⁵⁵ Notably, the intensity ratio (*I*_D/*I*_G) of Co–Mo₂C/NCHP catalyst (0.67) is lower compared to the Co/NCHP (0.73), Mo₂C/NCHP (0.80), and Mo₂C/Co/NCHP (0.73) catalysts. This suggests a higher degree of graphitization in the Co–Mo₂C/NCHP catalyst, potentially catalyzed by the carbonization of Mo.⁵⁶ Enhanced graphitization translates to improved electrical conductivity, potentially benefiting HER catalytic activity.⁵⁷

In contrast, the *I*_D/*I*_G for the Mo₂C/Co/NCHP catalyst remained unchanged compared to Co/NCHP. This indicates that Mo introduction after the carbonization process of the MOFs does not influence the further graphitization of carbon. The highest *I*_D/*I*_G observed for the Mo₂C/NCHP catalyst can be attributed to the introduction of surface defects caused by an additional acid-leaching step, which consequently reduces the degree of graphitization.⁵⁸ Fourier transform infrared spectra (FT-IR) (Fig. S10†) show peaks corresponding to activated carbon⁵⁹ Co–O,⁶⁰ and Mo–O⁶¹ bonds, indicating the presence of these components in the catalyst.

Fig. 1g displays the Barrett–Joyner–Halenda (BJH) desorption pore size distribution, revealing that the NCHP support possesses a network of fine nanoscale pores. The inset graph shows the N₂ adsorption/desorption isotherms of the measured catalysts. The Co/NCHP (390 m² g⁻¹) and Co–Mo₂C/NCHP (351 m² g⁻¹) catalysts exhibited a significantly larger specific surface



compared to the commercial Pt/C ($186 \text{ m}^2 \text{ g}^{-1}$), highlighting the enhanced surface area provided by the NCHP support.

X-ray absorption fine structure (XAFS) analysis was conducted to investigate the coordination environment of the Co and Mo ion centers in the Co/NCHP, $\text{Mo}_2\text{C}/\text{NCHP}$, $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$, and $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalysts. Fig. 2a presents the X-ray absorption near-edge structure (XANES) spectra at the Co K-edges of the investigated catalysts. Reference spectra of commercial Co foil, CoO, and Co_3O_4 are included for comparison. The absorption edge energy is known to be sensitive to the oxidation state of Co. It can be obtained by performing a second-derivative analysis of the XANES spectral edge.⁶² The inset in Fig. 2a depicts the correlation between the threshold energy (E_0) and the Co oxidation state in the reference compounds. A monotonic increase in E_0 is observed with increasing Co oxidation states. The E_0 values for the Co/NCHP, $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$, and $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalysts are 7721.69 eV, 7721.39 eV, and 7708.78 eV, respectively. These values suggest an effective oxidation state of Co atoms between 0 and +2 (the E_0 values for Co and CoO are 7709.09 eV and 7721.1 eV, respectively). Notably, the XANES spectrum of the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst exhibits a pre-edge absorption feature similar to that of the Co foil, indicating that Mo introduction during the ZIF-8@ZIF-67 stage effectively facilitates the reduction of Co to metallic form. In contrast, the Co K-edge in the Co/NCHP and $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ catalysts is shifted to a higher energy compared to the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst, implying a partially oxidized state for Co in these catalysts.

Extended X-ray absorption fine structure (EXAFS) analysis at the Co K-edge was performed to investigate the configuration

and local coordination environment of the Co ion center using Fourier transform (FT) analysis (Fig. 2b).⁶³ The Co foil spectrum exhibits a strong peak at 2.2 Å corresponding to Co–Co bonds, while the peak at 1.5 Å is associated with Co–O bonds in the CoO_x clusters. Interestingly, the peak at 1.4 Å, indicative of the Co–O(N) bonds, is absent in the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst but is present in the Co/NCHP and $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ catalysts. Additionally, the Co–Co bond peak appears at 2.0 Å in Co/NCHP and $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$, while it is observed at 2.2 Å in $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$. These differences likely stem from variations in the oxygen content within the catalysts. This observation suggests that Co in the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst exists primarily in the metallic state, whereas Co in the Co/NCHP and $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ catalysts is partially oxidized.⁶⁴

To further elucidate the distinct coordination environments of the Co atoms, EXAFS fitting based on the experimental data is presented in Fig. S11 and Table S1.† As expected for a metallic Co phase, the Co foil exhibits a Co–Co bond length of 2.5 Å. Similarly, the Co/NCHP and $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ catalysts show a Co–Co bond length of 2.5 Å. In contrast, the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst displays a unique coordination environment with a slightly shorter Co–Co bond length compared to 2.5 Å. This reduction in the Co–Co bond length can be attributed to the smaller ionic radius of Co induced by electron donation from Co to Mo.⁶⁵ The presence of a Co–Mo scattering path with a coordination number (CN) of 0.5 further supports the existence of an interface facilitating these interactions.

Unlike the $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$ catalyst, the $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ catalyst does not exhibit a decrease in bond length. This is likely because the additional introduction of Mo_2C to Co/NCHP,

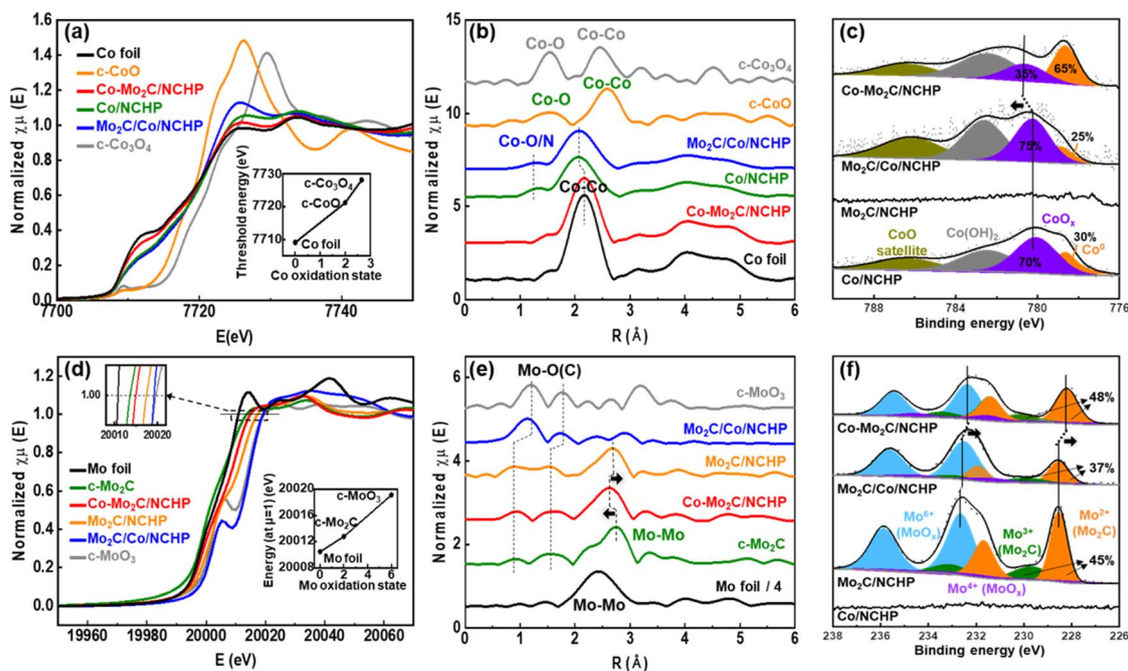


Fig. 2 Normalized XANES and FT-EXAFS spectra (a and b) at the Co K-edge and (d and e) Mo K-edge of different catalysts and reference samples (Co foil, CoO, Co_3O_4 , Mo foil, Mo_2C , and MoO_3). XPS patterns of (c) Co 2p and (f) Mo 3d of Co/NCHP, $\text{Mo}_2\text{C}/\text{NCHP}$, $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$, and $\text{Co}-\text{Mo}_2\text{C}/\text{NCHP}$.



does not influence the pre-existing Co–Co bonds. Fig. 2c displays the Co 2p XPS spectra of the Co/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts. In the Co 2p_{3/2} region of the Co/NCHP spectrum, four peaks at 778.8 eV, 781.2 eV, 782.5 eV, and 786.4 eV correspond to the Co metal, CoO_x, Co(OH)₂, and satellite peaks, respectively.⁶⁶ Consistent with the XANES data, a prominent metallic Co peak is evident in the spectrum of the Co–Mo₂C/NCHP catalyst. Additionally, the Co²⁺ peak of the Co–Mo₂C/NCHP catalyst is observed at 781.9 eV, which is shifted positively compared to those of the Co/NCHP and Mo₂C/Co/NCHP catalysts. These results reveal that the Mo plays a crucial role in facilitating the formation of the Co–Mo₂C interface, which reduces the Co oxidation state to its metallic state. This metallic Co–Mo₂C interface promotes charge redistribution between Co and Mo₂C, which alters their local electronic environment and strengthens interfacial interactions. As revealed by XAFS and XPS analyses, the formation of this interface supports the design strategy of creating dual active sites within the NCHP framework, underscoring the critical role of Co–Mo₂C interactions.

Fig. 2d displays the XANES spectra at the Mo K-edges of the Mo₂C/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts. Reference spectra of commercial Mo foil, Mo₂C, and MoO₃ are included for comparison. The inset in Fig. 2d depicts the dependence of the K-edge position at $\mu = 1$ on the Mo oxidation states in the reference compounds.⁶⁷ These compounds exhibit a monotonic increase in the K-edge position at $\mu = 1$ with increasing Mo oxidation states. The K-edge positions at $\mu = 1$ for the Mo₂C/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts are 20 016.5 eV, 20 019 eV, and 20 015.8 eV respectively. These values suggest an effective oxidation state of Mo atoms between +2 and +6 (the K-edge position at $\mu = 1$ for Mo₂C and MoO₃ are 20 012.8 eV and 20 019.1 eV, respectively).

For the Mo₂C/Co/NCHP catalyst, Mo₂C is primarily exposed on the surface, since it was introduced after the construction of the NCHP. Consequently, it undergoes oxidation in air, exhibiting an oxidation state similar to MoO₃. Conversely, in the Co–Mo₂C/NCHP catalyst, where Mo₂C resides in close proximity to metallic Co within the NCHP structure, the Mo is significantly protected from oxidation. Mo₂C in the Mo₂C/NCHP catalyst, after Co leaching, shows a relatively increased oxidation. Therefore, the following trend can be observed for the Mo oxidation state: Mo₂C/Co/NCHP > Mo₂C/NCHP > Co–Mo₂C/NCHP.

EXAFS analysis at the Mo K-edge (Fig. 2e) reveals that the peaks at 1.50 Å and 2.74 Å in commercial Mo₂C correspond to Mo–O(C), Mo–Mo bonds, respectively.⁶⁸ In the Co–Mo₂C/NCHP catalyst, a Mo–Mo scattering peak is observed at a shorter bond distance (around 2.60 Å) compared to commercial Mo₂C (around 2.74 Å). Conversely, in the Mo₂C/NCHP catalyst, where Co was removed by harsh acid leaching, the Mo–Mo scattering peak shifts back at a longer bond distance (around 2.70 Å) similar to that of commercial Mo₂C. This peak shift suggests the presence of charge redistributions between metallic Co and Mo₂C. Additionally, the Mo₂C/Co/NCHP catalyst exhibits predominantly Mo–O scattering peaks. This indicates that molybdenum oxide was probably in an amorphous state.

The EXAFS fitting based on the experimental data is presented in Fig. S12 and Table S2.† The Co–Mo₂C/NCHP catalyst shows a CN of 0.5 for the Mo–Co scattering path, 5.3 for the Mo–Mo scattering path, and 1.8 for the Mo–C scattering path, confirming the presence of an interface for Co–Mo charge redistributions. The presence of both Mo–Mo₁ and Mo–Mo₂ bonds in the Mo based structure complicates the interpretation of the EXAFS fitting results. In the Co–Mo₂C/NCHP catalyst, due to the electron donation from Co, a large number of electrons are involved in Mo bonding, leading to an elongation of the Mo–Mo₁ bond length (from 2.89 Å to 2.98 Å).⁶⁵

Furthermore, Fig. 2f displays the Mo 3d XPS spectra of Mo₂C/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts. In the Mo 3d_{5/2} region, four peaks are observed at 228.7 eV, 229.5 eV, 231.4 eV, and 232.6 eV, corresponding to Mo²⁺, Mo³⁺, Mo⁴⁺, and Mo⁶⁺ from Mo₂C and MoO_x, respectively.⁶⁹ The MoO_x peak likely originates from the oxidation of Mo₂C upon exposure to air. This suggests a reduced state for Mo in the Co–Mo₂C/NCHP catalyst, as evidenced by the higher Mo₂C to MoO_x ratio. In the Co–Mo₂C/NCHP catalyst, consistent with the results from the Co²⁺ peak shift observed in Fig. 2c, charge redistributions between the metallic Co and Mo₂C lead to a negative shift in the Mo peaks. These findings are consistent with the observations from XANES and EXAFS analyses.

Fig. 3a presents polarization curves of the HER performance of electrodes utilizing various catalysts: Co/NCHP, Mo₂C/NCHP, Co–Mo₂C/NCHP, Mo₂C/Co/NCHP, and commercial Pt/C. At the current density of 10 mA cm^{−2}, the overpotentials for HER follow this trend: Mo₂C/NCHP (highest, 397 mV) > Mo₂C/Co/NCHP (266 mV) > Co/NCHP (182 mV) > Co–Mo₂C/NCHP (125 mV) > Pt/C (63 mV). The Co–Mo₂C/NCHP catalyst exhibits significantly enhanced HER activity due to the synergistic presence of metallic Co for the hydrogen evolution sites, Mo₂C for the water dissociation sites, and abundant interfaces facilitating charge redistribution between metallic Co and Mo₂C.^{70,71} Notably, the Co–Mo₂C/NCHP catalyst outperforms other reported powder-based HER catalysts at a current density of 10 mA cm^{−2} (refer to Fig. S13† for overpotential values). To facilitate a direct LSV comparison of both intrinsic and practical HER activities, LSV curves with and without *iR* correction are presented in Fig. S14.†⁷²

However, the Mo₂C/Co/NCHP catalyst exhibits higher overpotentials compared to the Co–Mo₂C/NCHP catalyst. This can be attributed to Co oxidation and MoO_x formation. Oxidation of Co weakens its interaction with Mo₂C, while surface exposure of the Mo leads to the formation of MoO_x, which has lower HER activity (as discussed in Fig. 2f).

Fig. 3b illustrates the Tafel slopes of the catalysts. Pt/C exhibits a slope of 45 mV dec^{−1}, indicating that the rate-determining step (RDS) is the Tafel reaction step (H* + H* ↔ H₂).⁷³ In contrast, the Mo₂C/NCHP, Co/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts display a slope of approximately 120 mV dec^{−1}. This similarity suggests that the Volmer reaction step (H₂O + e[−] ↔ H* + OH[−]) is the RDS for these non-platinum group metal catalysts. Interestingly, introducing Mo, which is known to lower water dissociation energy compared to Co,^{14,74} improves the Volmer reaction kinetics and decreases Tafel



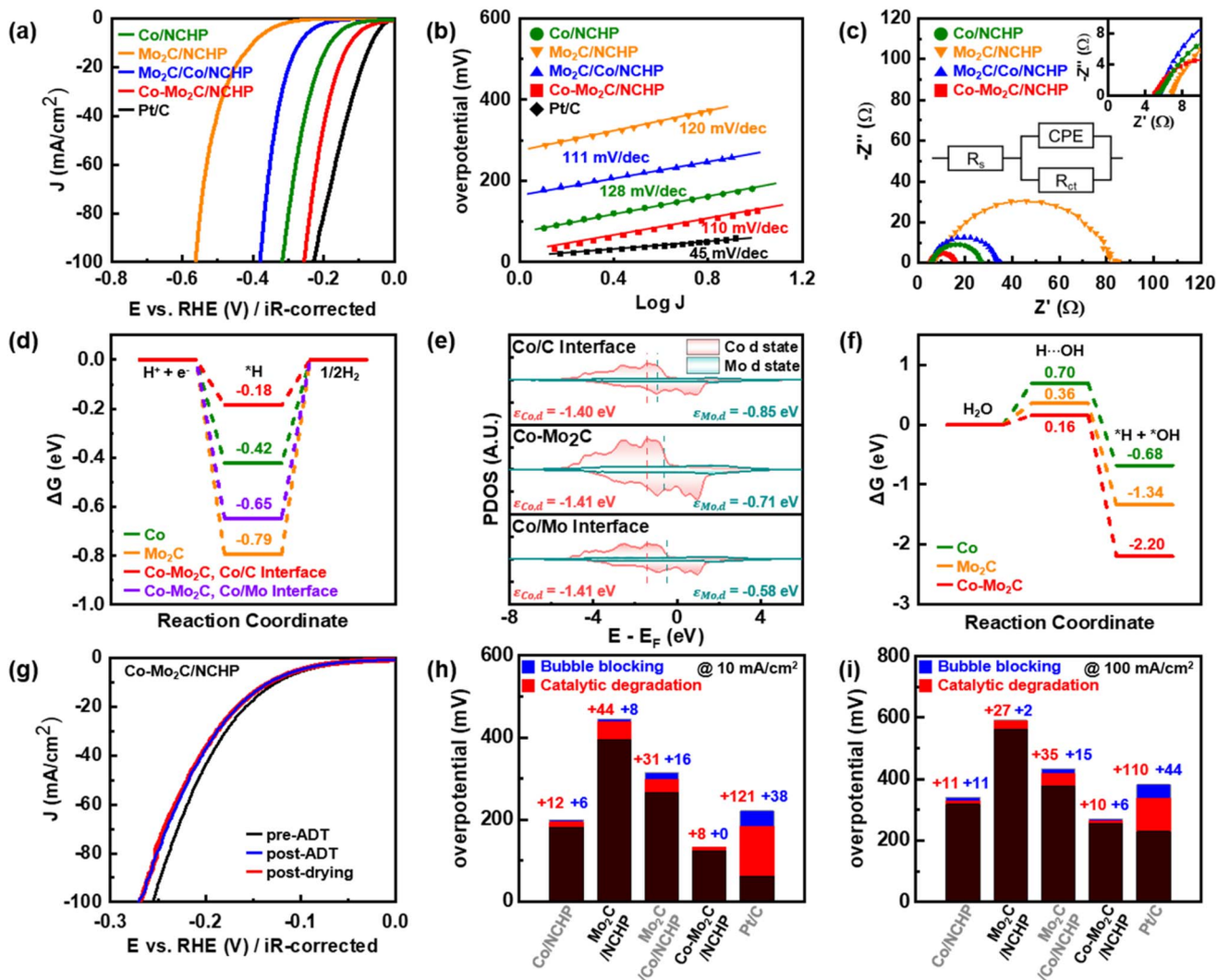


Fig. 3 (a) IR-corrected HER polarization curves, (b) Tafel slopes, (c) Nyquist plots at -300 mV. (d) The Gibbs free energy of H adsorption on Co, Mo_2C , Co/C interface in Co- Mo_2C , and Co/Mo interface in Co- Mo_2C . (e) The d-band center of Co and Mo in Co- Mo_2C , the Co/C interface, and the Co/Mo interface. (f) The activation energy for water dissociation on Co, Mo_2C , and Co- Mo_2C . (g) HER polarization curves of Co- $\text{Mo}_2\text{C}/\text{NCHP}$ before (black) and after (blue) 1000 potential cycles and after bubble removal (red) from 0.05 to -0.55 V_{RHE} at a scan rate of 0.05 V s^{-1} in 1.0 M KOH. (h and i) Overpotential at 10 and 100 mA cm^{-2} .

slopes. Notably, the Co- $\text{Mo}_2\text{C}/\text{NCHP}$ catalyst exhibits the lowest slope (110 mV dec^{-1}).

The Nyquist plots (Fig. 3c) show the electrochemical impedance characteristics of the catalysts. In the high frequency region, the system resistance is solely the solution resistance (R_s). Conversely, the low frequency range reflects the sum of R_s and charge transfer resistance (R_{ct}), represented by the diameter of the semicircle. A smaller diameter signifies a faster reaction rate. Among the catalysts, Co- $\text{Mo}_2\text{C}/\text{NCHP}$ exhibits the lowest R_{ct} , indicating the fastest electron transfer and most favorable HER kinetics at the electrolyte interface. This aligns with the observations from the polarization curves (Fig. 3a).

As an estimate of the electrochemical active surface area (ECSA) of the catalysts, the double-layer capacitance (C_{dl}) was determined from cyclic voltammetry (CV) measurements,

performed in the non-faradaic region (1.0 – 1.1 V_{RHE}) at various scan rates ranging from 10 mV s^{-1} to 50 mV s^{-1} in H_2 -saturated 1 M KOH solution (Fig. S15†). At 1.05 V_{RHE} , the capacitive current density was calculated as the sum of the anodic and cathodic current densities and plotted as a function of scan rate (Fig. S16†). The C_{dl} was calculated based on the following relationship between the capacitive current (i) and the scan rate (ν):

$$i = C_{dl} \cdot \nu$$

The calculated C_{dl} values followed the order: $\text{Mo}_2\text{C}/\text{NCHP}$ (0.32 mF cm^{-2}) < $\text{Mo}_2\text{C}/\text{Co}/\text{NCHP}$ (0.99 mF cm^{-2}) < Co/NCHP (1.16 mF cm^{-2}) < Co- $\text{Mo}_2\text{C}/\text{NCHP}$ (1.67 mF cm^{-2}), suggesting that the Co- $\text{Mo}_2\text{C}/\text{NCHP}$ catalyst possesses the largest



accessible electrochemical surface area among the samples. These results further support the superior HER activity of Co-Mo₂C/NCHP, as a larger surface area offers more active sites for catalytic reactions.

Fig. S17† shows the single-cell performance of the Co-Mo₂C/NCHP catalyst-coated cathode assembled with commercial AEM and iridium(IV) oxide-coated anode (catalyst loading: $0.85 \pm 0.30 \text{ mg cm}^{-2}$), compared with the commercial Pt/C-coated cathode. A 1 M KOH solution at 60 °C was supplied to both the anode and cathode inlets. Although only a small amount of active material (Co + Mo₂C loading: $0.14 \pm 0.03 \text{ mg cm}^{-2}$) was loaded onto the electrode, a voltage of 2.02 V was observed at a current density of 1 A cm^{-2} . Interestingly, as the current density increases, the single-cell performance difference compared to the Pt/C electrode decreases, showing only a 30 mV voltage difference at 3 A cm^{-2} . This result indicates that the Co-Mo₂C/NCHP material possesses potential characteristics suitable for high-current-density hydrogen production. To achieve better performance, it will be necessary to increase the amount of metallic active materials within the NCHP support and further optimize the membrane electrode assembly (MEA). However, the present results demonstrate the potential applicability of this approach to practical cells. A 50-hour stability test at 0.5 A cm^{-2} showed a voltage increase of $\sim 0.09 \text{ V}$, indicating the need for further optimization to improve durability under practical AEMWE conditions (Fig. S18†).

The origin of the improved HER activity of Co-Mo₂C/NCHP was elucidated through DFT calculations. A theoretical model of Co-Mo₂C was simulated to identify the active sites for hydrogen desorption and water dissociation. To compare hydrogen desorption capability, the Gibbs free energy of hydrogen adsorption was calculated.⁷⁴ Specifically, the Co-Mo₂C heterostructure presented unique electronic structures at the interfaces between the Co and Mo₂C. The division of the Co/C and Co/Mo interfaces, as shown in Fig. S1,† helps to identify the favorable electronic properties of the specific interfaces in the Co-Mo₂C. Fig. 3d and S19† illustrate that Mo₂C possesses the strongest H binding energy (-0.79 eV), followed by the Co/Mo interface (-0.65 eV), Co (-0.42 eV), and the Co/C interface (-0.18 eV).

From these results, the Co/C interface in the Co-Mo₂C is the most thermoneutral result, facilitating hydrogen gas production. Meanwhile, the Co/Mo interface exhibits an intermediate adsorption strength between individual Co and Mo₂C.

To explain why the Co-Mo₂C/NCHP catalyst has the highest HER activity, the charge density difference was calculated in the Co-Mo₂C model, where sufficient charge transfer was observed at the interfaces, as shown in Fig. S20.† At the Co/C interface, charges are transferred from Co to Mo₂C, enabling moderate adsorption of hydrogen atoms. This charge transfer direction at the Co/C interface is consistent with a previous study of Co/Mo₂C@C.⁷⁵ Conversely, at the Co/Mo interface, charges are transferred from Mo₂C to Co. The detailed charge density difference values are summarized in Table S3.†

Further, the d-band centers of metals at different interfaces within Co-Mo₂C were analyzed, as shown in Fig. 3e. While Co presents similar d-band centers across interfaces, Mo₂C

displays notable differences. The average d-band center of Mo in the Co-Mo₂C is -0.71 eV . Compared to this value, at the Co/C interface, the Mo d-band center shifts downward to -0.85 eV . In contrast, at the Co/Mo interface, the Mo d-band center shifts upward to -0.58 eV . The lower and more distant Mo d-band center from the Fermi level at the Co/C interface supports the thermoneutral Gibbs free energy of hydrogen adsorption.

Additionally, Fig. S21† illustrates that the Co and Mo₂C sites within Co-Mo₂C have hydrogen adsorption energies similar to that of individual Co and Mo₂C, as depicted in Fig. 3d. Consequently, these results highlight that the interfaces in Co-Mo₂C are key active sites of the improved HER activity, based on the synergistic effects between Co and Mo₂C. Especially, the accumulation of charge on Mo₂C and the lower d-band center of Mo promote moderate hydrogen adsorption and thus accelerate the hydrogen evolution.

The water dissociation ability is also crucial for generating proton sources for hydrogen gas in alkaline HER.¹⁴ The Gibbs free energy profile of water dissociation indicates the activation energies of 0.70 eV for Co, 0.36 eV for Mo₂C, and 0.16 eV for Co-Mo₂C in Fig. 3f and S22.† The lowest energy barrier of Co-Mo₂C highlights its excellent water splitting capability, which surpasses that of individual Co and Mo₂C. While Co is advantageous for hydrogen desorption, as indicated in Fig. 3d, it has poor water dissociation ability compared to Mo₂C and Co-Mo₂C.¹⁹ On the other hand, Mo₂C exhibits strong hydrogen adsorption strength but demonstrates a lower activation barrier for water dissociation than Co.²⁸ This combination of properties in Co-Mo₂C, with efficient hydrogen desorption by Co and enhanced water dissociation by Mo₂C, results in the improved HER performance.

To further evaluate the pH-dependent catalytic behavior, we measured the HER performance of all catalysts in neutral electrolyte (1.0 M PBS), and the *iR*-corrected polarization curves are presented in Fig. S23.† It is well established that HER kinetics in neutral media are significantly influenced by both the Gibbs free energy of hydrogen adsorption and the efficiency of water dissociation, particularly due to the sluggish Volmer step under near-neutral pH conditions. Consistent with this, all non-precious metal catalysts showed a notable decrease in activity in PBS compared to 1.0 M KOH, while the commercial Pt/C catalyst retained high activity with only a slight decline, owing to its intrinsic ability to efficiently adsorb protons and dissociate water even at low proton concentrations.

Among the catalysts, Mo₂C/NCHP exhibited the lowest HER performance in neutral media, followed by Mo₂C/Co/NCHP, Co/NCHP, and Co-Mo₂C/NCHP, which demonstrated the highest activity. This catalytic activity trend strongly correlates with the hydrogen adsorption and water dissociation, as shown in Fig. 3d-f. These results highlight the synergistic effect of the Co-Mo₂C interface, which facilitates both hydrogen adsorption and water dissociation, thereby maintaining favorable HER kinetics even under proton-limited neutral conditions.

In addition, the charge redistribution between Co and Mo₂C further improves the water splitting performance beyond that of Mo₂C. This can contribute to the lower Tafel slopes experimentally observed in both Co-Mo₂C/NCHP and Mo₂C/Co/



NCHP. As both catalysts exhibit similar Tafel slopes in Fig. 3b, it can be interpreted that the charge redistribution between Co and Mo₂C promotes water dissociation through the heterostructure strategy. Furthermore, the order of activation energies for water dissociation correlates well with that of the Tafel slopes in Fig. 3b. These results suggest that both thermoneutral hydrogen adsorption at the Co/C interface and the lowest water dissociation barrier contribute to the noteworthy HER activity of Co–Mo₂C/NCHP, driven by the synergistic effects between Co and Mo₂C. In line with this, the theoretical results further support that the Co–Mo₂C interface facilitates optimal hydrogen adsorption and efficient water dissociation. The synergistic effects between Co and Mo₂C, combined with the structural confinement within the NCHP support, enhance HER performance by overcoming kinetic limitations at each single active site.

Following the accelerated durability test (ADT) experiments, an increase in overpotential for the HER was observed. This can be attributed to two factors: catalytic degradation and the adverse effect of bubbles on the catalysts.^{32,76} To distinguish between these factors, durability testing was conducted. Linear sweep voltammetry (LSV) polarization curves were recorded at the three stages: pre-ADT (before the test), post-ADT with surface microbubbles, and post-drying (after removing surface microbubbles). The post-drying process was performed by separating the RDE and vacuum drying it at room temperature for 20 minutes. After this drying step, LSV measurements were recorded again to evaluate the changes in the catalytic performance after the removal of surface microbubbles. As a representative example, Fig. S24† illustrates the polarization curves for commercial Pt/C during the ADT experiments. The initial overpotential at 10 mA cm⁻² is 63 mV. After ADT, the overpotential increased significantly to 222 mV. However, drying to remove surface microbubbles led to a slight reduction in overpotential (to 184 mV).

This observation suggests that microbubbles blocking active sites is a major contributor to the overpotential increase after ADT for Pt/C. The detachment of Pt nanoparticles from the carbon support during ADT can also contribute to the remaining overpotential increase after drying. Therefore, the change in overpotential between post-ADT and post-drying stages is attributed to bubble blocking, while the difference between pre-ADT and post-drying stages reflects the true catalytic degradation.

This phenomenon can be accelerated in commercial carbon supported Pt catalysts by the destruction of Pt anchoring sites by the harsh ADT conditions.⁷⁷ Additionally, delayed bubble desorption during HER can lead to the formation of larger bubbles, which can cause extensive peeling of Pt nanoparticles from the surface.

Similar trends in overpotential changes are observed for the Co/NCHP, Mo₂C/NCHP, Mo₂C/Co/NCHP, and Co–Mo₂C/NCHP catalysts (Fig. 3g and S25†). To quantify the contributions of catalytic degradation and bubble blocking, the overpotentials of these catalysts and Pt/C at current densities of 10 mA cm⁻² and 100 mA cm⁻² were investigated and are presented in Fig. 3h and i, respectively. The red region represents the overpotential

increase attributed to catalytic degradation, while the blue region corresponds to the overpotential increase resulting from bubble blocking. In the red region (catalytic degradation), the overpotential increase for NCHP-supported catalysts is relatively small, with a maximum increase of 44 mV compared to Pt/C (121 mV). Similarly, in the blue region (bubble blocking), the overpotential increase for NCHP-supported catalysts is minimal, with a maximum increase of 16 mV compared to Pt/C (38 mV). The Co–Mo₂C/NCHP catalyst reduces the HER overpotential by 66% at 10 mA cm⁻² compared to the single active site catalyst (Co/NCHP). Moreover, the Co–Mo₂C/NCHP catalyst demonstrates enhanced durability, exhibiting a 38% lower HER overpotential than commercial Pt/C at 10 mA cm⁻² even after 1000 cycles. These findings are consistent with the ADT results obtained at a current density of 100 mA cm⁻². Overall, both catalytic degradation and bubble blocking are significantly lower for all catalysts supported on NCHP, compared to Pt/C. These durability results highlight the effectiveness of the NCHP support in mitigating performance degradation during prolonged HER cycles through efficient bubble desorption.

Particle agglomeration and dissolution are known to be the main mechanisms leading to catalytic degradation and performance loss during HER. TEM is a valuable tool for investigating the morphological changes of catalysts after undergoing ADT.⁷⁸ To understand the factors contributing to the high durability of NCHP-supported catalysts, TEM images of the Co–Mo₂C/NCHP and Pt/C catalysts were obtained. Fig. 4 presents TEM images and corresponding particle size distributions for Co–Mo₂C/NCHP (Fig. 4a and b) and Pt/C (Fig. 4c and d) before and after ADT.

Interestingly, the Co–Mo₂C/NCHP catalyst exhibits minimal change in particle size, with an increase from 4.08 ± 1.10 nm to 4.16 ± 0.92 nm. In contrast, Pt/C particles show a more significant increase, from 2.57 ± 0.53 nm to 3.01 ± 0.61 nm. This slight increase in particle size for the Co–Mo₂C/NCHP catalyst correlates with the observed 8 mV increase in catalytic degradation after the durability test at 10 mA cm⁻². However, it is important to note that this increase is not attributed to bubble blocking. This exceptional durability of the Co–Mo₂C/NCHP catalyst can be attributed to the presence of micro-pores (Fig. 1g) and its three-dimensional structure. This unique structure facilitates easy access of the electrolyte to active particles and allows for the early desorption of generated hydrogen gas. Consequently, it reduces both catalytic degradation and the negative impact of bubbles on HER performance.^{79,80}

Fig. S26† shows the TEM images and the amounts of Mo measured in the electrolyte using ICP analysis, which helps to quantify the Mo dissolution from the Co–Mo₂C/NCHP and Mo₂C/Co/NCHP catalysts. Both catalysts exhibit minimal structural changes before and after ADT. However, the Co–Mo₂C/NCHP catalyst displays a lower concentration of Mo in the electrolyte compared to the Mo₂C/Co/NCHP catalyst. This indicates that the Mo loss is associated with the surface exposure of Mo₂C. As a result, Co–Mo₂C/NCHP exhibits the lowest catalytic degradation due to the protected Mo₂C, in conjunction with metallic Co within the NCHP structure.



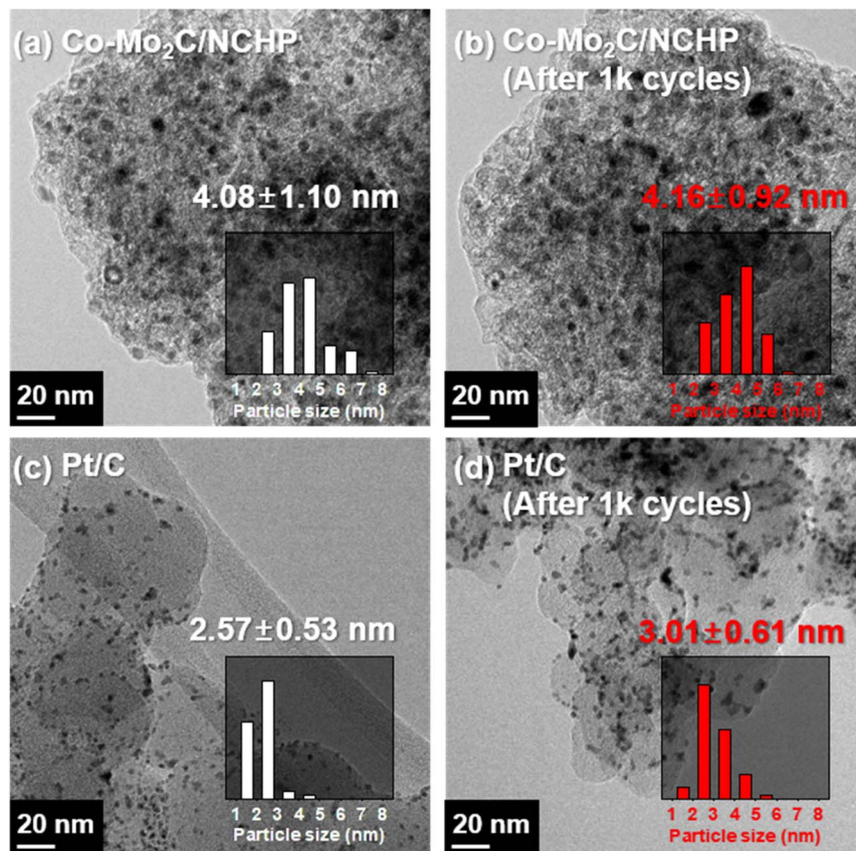


Fig. 4 TEM images of (a and b) Co–Mo₂C/NCHP and (c and d) Pt/C, taken before and after 1000 potential cycles. Insets show the corresponding particle size distributions, along with the average particle sizes.

Even after 1000 cycles of ADT testing, the overpotential of the Co–Mo₂C/NCHP catalyst increased by only 16 mV at 100 mA cm⁻², indicating minimal bubble blocking and catalytic degradation. In contrast, the Mo₂C/NCHP (+44 mV @ 10 mA cm⁻², +27 mV @ 100 mA cm⁻²) and Mo₂C/Co/NCHP (+31 mV @ 10 mA cm⁻², +35 mV @ 100 mA cm⁻²) catalysts exhibit considerable catalytic degradation compared to the Co–Mo₂C/NCHP. This can be attributed to the presence of surface exposed Mo₂C (as showed in Fig. 2f), which is susceptible to leaching. The slight increase in particle size and the low Mo dissolution observed after ADT suggest that the NCHP structure effectively prevents the common issues of particle agglomeration and Mo leaching. These findings emphasize the importance of structural design in preventing the loss of active material, thereby enhancing catalyst durability. These results demonstrate the critical role of the NCHP support in enhancing the operational durability of Co–Mo₂C/NCHP catalysts in an alkaline HER environment.

The actual catalytic substances in the catalytic process are of great significance for the design of efficient and practical electrodes. Fig. S27a and b† shows the XRD patterns and Raman spectra of the Co–Mo₂C/NCHP before and after the ADT testing. The results indicate that the crystalline Co and Mo₂C nanoparticles, as well as the *I*_D/*I*_G ratio of the sample, remained unchanged, demonstrating the robust nature of the

nanoparticles and the porous carbon framework during the stability test.

After the HER ADT testing in alkaline solution, the XPS spectra of Co–Mo₂C/NCHP were measured to investigate changes in surface properties (Fig. S27c and d†). After ADT testing, the metallic Co signal was weakened and the peaks corresponding to cobalt hydroxides became increased as exposed to alkaline solution, coinciding with previous reports.^{81,82} Meanwhile, the peaks corresponding to MoO_x decreased, indicating the dissolution of air-formed MoO_x on the Mo₂C surface. Even after prolonged HER operation, a significant portion of metallic Co and Mo remains, and the Co–Mo₂C/NCHP catalyst maintains strong interfacial interactions between Co and Mo₂C, contributing to high HER durability.

Overall, the cohesive integration of Co and Mo₂C dual active sites within the NCHP addresses key limitations in alkaline HER kinetics, offering improvements in both catalytic efficiency and long-term stability. This strategy dual-active sites and provides a clear path forward for advancing non-noble metal catalysts in hydrogen evolution systems, with potential for further optimization in future studies.

Conclusions

To overcome the challenges of sluggish water dissociation kinetics associated with non-noble metal catalysts, we have



introduced a catalyst design featuring cohesive Co and Mo₂C dual active sites, which are strongly confined within an N-doped carbon hollow polyhedron (NCHP). Carbonizing a core-shell ZIF structure containing Co and Mo facilitates the formation of abundant metallic Co–Mo₂C interfaces. Charge redistribution between the metallic Co and Mo₂C at active Co–Mo₂C interfaces was corroborated by several observations: a shorter peak in the Mo K-edge EXAFS spectra, distinct coordination environments of Co–Co and Mo–Mo revealed by EXAFS fitting, and the peak shifts in the Co 2p and Mo 3d XPS spectra. These findings demonstrate that strategic incorporation of Mo effectively maximizes the formation of metallic Co–Mo₂C interfaces, promoting efficient HER performance.

DFT calculations reveal that the Co–Mo₂C interface can promote the optimal hydrogen binding strength and facilitate water dissociation. The Co–Mo₂C/NCHP catalyst also demonstrated superior durability, compared with the commercial Pt/C catalyst. This was attributed to the strongly confined Co–Mo₂C structure within the microporous NCHP, which not only enhanced bubble desorption but also mitigated Mo leaching. Consequently, the Co–Mo₂C/NCHP catalyst exhibits a low overpotential of 125 mV at a current density of 10 mA cm⁻², outperforming many other reported non-noble metal catalysts, along with improved durability. This catalyst design strategy paves the way for the development of next-generation non-noble metal catalysts with enhanced efficiency, stability, and durability for large-scale hydrogen production by addressing key challenges such as sluggish water dissociation kinetics, Mo leaching, and bubble desorption.

Data availability

All data supporting this article, including raw electrochemical measurements, TEM images, and XRD patterns, are provided in the ESI.†

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

H. L. and M. K. contributed equally to this work. This work was partly supported by Korea Institute of Energy Technology Evaluation and Planning (KETEP) grant funded by the Korea government (MOTIE) (20214000000650, Energy Innovation Research Center for Fuel Cell Technology) and the National Research Foundation of Korea (NRF) Grant funded by the Ministry of Science and ICT, Republic of Korea (RS-2024-00466554).

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