Review

Smart Designs of Mo Based Electrocatalysts for Hydrogen Evolution Reaction

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Abstract: As a sustainable and clean energy source, hydrogen can be generated by electrolytic water splitting (i.e., a hydrogen evolution reaction, HER). Compared with conventional noble metal catalysts (e.g., Pt), Mo based materials have been deemed as a promising alternative, with a relatively low cost and comparable catalytic performances. In this review, we demonstrate a comprehensive summary of various Mo based materials, such as MoO₃, MoS₂ and MoC. Moreover, state of the art designs of the catalyst structures are presented, to improve the activity and stability for hydrogen evolution, including Mo based carbon composites, heteroatom doping and hetero-structure construction. The structure–performance relationships relating to the number of active sites, electron/ion conductivity, H/O binding and activation energy, as well as hydrophilicity, are discussed in depth. Finally, conclusive remarks and future works are proposed.

Keywords: hydrogen evolution reaction; Mo based materials; electrocatalyst; doping; binding energy; active sites

1. Introduction

Considering the environmental pollution and greenhouse effect caused by the excessive utilization of fossil fuels, the exploitation of renewable and clean energy sources is imminent and imperative in order to realize the sustainable development of industries and economies [1,2]. Characterized by a carbon free nature, green product (H₂O) and high energy density, hydrogen has been regarded as a promising alternative to conventional fossil fuels, prior to other renewable energies (e.g., wind and solar energy) that suffer from an intermittent nature and low energy density [3,4]. Thus far, general routes of producing hydrogen include the gasification of coal and biomass, conversion of hydrocarbons (e.g., methane and tar molecules) and electrolytic water splitting [2]. The former two methods require a high reaction temperature (i.e., intense energy input) and exhibit a low purity of hydrogen with additional emissions of carbon dioxide [5]. In comparison, the production of hydrogen via the hydrogen evolution reaction (HER) in electrolytic water splitting enjoys zero emissions of greenhouse gases and a high hydrogen yield [6]. Particular properties of an ideal HER electrocatalyst consist of a low overpotential, low cost (scalability), high conductivity and structural stability [7]. Up to now, Pt based materials have exhibited the highest efficiency of hydrogen production; however, the limited reserves and high price essentially inhibit their utilization at an industrial level [8,9]. Recently, nonnoble transition metals have presented great potentials as an alternative to the Pt based catalysts in HER [10,11]. Among them, molybdenum (Mo)
based electrocatalysts have drawn increasing interest due to their admirable activities at a wide range of pH values [12–14]. For example, NiMo nanowires were constructed on Ni foam and exhibited a comparable activity to Pt/C commercial catalysts due to the metal–metal synergy and abundant active sites in the hierarchical porous structure [15]. In another case, a MoS2 bicontinuous network was synthesized with a nanoscale size and mesoporous structure, providing more active sites for hydrogen evolution [16]. Moreover, 3 nm MoC nanoparticles were highly dispersed on the carbon matrix, creating a large contact surface with the reactants and presenting considerable HER activity [17]. Other Mo based compounds (e.g., MoSe2, MoO3, MoN, MoP, MoB) also exhibit a good potential for producing hydrogen in high efficiencies [18–22]. Nevertheless, the sluggish reaction kinetics, because of their intrinsically low activity and poor conductivity, limit their large scale applications in HER [23]. To address the issues, various modifications have been developed to improve the catalytic performances of Mo based materials, such as combination with carbon matrix, doping with heteroatoms and construction of heterostructures [24–27]. So far, most of the reviews are focused on specific Mo based compounds; a comprehensive review based on the modification strategies of Mo based electrocatalysts rather than the materials is rarely reported in HER [1–5]. Therefore, in this review, in addition to the introduction of the reaction mechanism, all the types of Mo based materials used in HER will be summarized. Moreover, three strategies for enhanced activity and stability will be discussed in depth, including carbon/Mo based composites, heteroatom doping and the construction of heterocatalysts. The structure–performance relationships will be illustrated by referring to the physicochemical properties (e.g., active site, conductivity, hydrophilicity) and catalytic performances (e.g., overpotential and Tafel slope). Finally, conclusive remarks and possible solutions to the existing challenges will be proposed.

2. HER Mechanism

Two mechanisms are commonly recognized for HER: the Volmer–Tafel and Volmer–Heyrovsky mechanisms [1]. In acidic solutions (Equations (1)–(3) where * denotes a free site on the surface and H* denotes a hydrogen atom adsorbed on the surface), the adsorbed hydrated protons (H2O*) are reduced to produce H* (intermediate) via a Volmer step; subsequently, two H* combine to form one H2 molecule via a Tafel route or one H* reacts with the hydrated proton to form hydrogen through a Heyrovsky step.

\[ \text{H}_3\text{O}^+ + e^- \rightarrow \text{H}^* + \text{H}_2\text{O} \quad \text{(Volmer step)} \]  
\[ \text{H}^* + \text{H}^* \rightarrow \text{H}_2 \quad \text{(Tafel step)} \]  
\[ \text{H}^* + \text{H}_2\text{O}^+ + e^- \rightarrow \text{H}_2 + \text{H}_2\text{O} \quad \text{(Heyrovsky step)} \]  

In comparison, Equations (4)–(6) present the hydrogen evolution mechanism in neutral and alkaline electrolytes. In detail, H* is generated from the reduction of water molecules, followed by either a combination of 2 H* (a Tafel step) or the reaction between H* and H2O (a Heyrovsky step) to produce H2.

\[ \text{H}_2\text{O} + e^- \rightarrow \text{H}^* + \text{OH}^- \quad \text{(Volmer step)} \]  
\[ \text{H}^* + \text{H}^* \rightarrow \text{H}_2 \quad \text{(Tafel step)} \]  
\[ \text{H}^* + \text{H}_2\text{O} + e^- \rightarrow \text{H}_2 + \text{OH}^- \quad \text{(Heyrovsky step)} \]  

For both mechanisms, HER is a two step reaction, where either the Volmer step (adsorption) or the Heyrovsky/Tafel step (desorption) determines the reaction rate. Particularly, the pH of the electrolytes exerts a profound influence on the mechanism. In acidic solutions, the adsorption and electrochemical conversion of protons dominate the reaction kinetics in the Volmer step; while in neutral or alkaline conditions, the H2O dissociation and OH− desorption also play a crucial role in determining the rate.
Besides, a Tafel slope is strongly related to reaction kinetics and acts as an indicator of the rate-determining step. In particular, the Volmer, Heyrovsky and Tafel steps are reflected from the Tafel slope of 120, 40 and 30 mV·dec⁻¹. Interestingly, under a high hydrogen coverage over 0.6, 120 mV·dec⁻¹ a Tafel slope is also seen for the Heyrovsky step, suggesting the potential and coverage dependence of Tafel slope [28].

3. Mo Based HER Electrocatalysts

Considering the limited adsorption and desorption of hydrogen for the metallic Mo, alloying or compounding with other elements (e.g., alloys, carbides, oxides, selenides, borides, nitrides, sulfides and phosphides) effectively adjusts the electronic structure and promotes H adsorption. In addition, these Mo based materials enjoy an enhanced stability at various pH values and good electronic conductivity, benefiting the reaction kinetics and lifespan of the electrolytic HER [29,30]. In the following, different Mo based electrocatalysts will be briefly introduced by category.

The hydrogen adsorption and desorption energy of Mo metal can be adjusted by introducing other metallic elements, thus tailoring the catalytic activity for HER. Brown et al. [31] compared a series of Mo based alloys in HER and found that NiMo outperformed CoMo and FeMo. Recently, by the calcination of the NiMoO₄ precursor, the formed MoNi alloy possessed a facilitated H₂O dissociation in a Volmer step in 1 M KOH electrolytes, thus presenting only a 15 mV overpotential at 10 mA·cm⁻² and a 30 mV·dec⁻¹ Tafel slope [32]. More efforts are recommended for the study of mass loading effects.

Apart from the alloys, other types of Mo based compounds have been successfully applied in HER. Owing to its good electronic conductivity, structural stability and abundant active sites (O edge and Mo edge), MoO₃ exhibits great potential as an HER catalyst. Various morphologies are synthesized to enhance the exposed active sites and inhibit agglomeration, such as nanowires, nanosheets and nanoflowers [33-37]. Another transition metal dichalcogenide—MoS₂—is widely utilized as an alternative to Pt catalysts because of earth abundance, rich active sites and chemical stability [5]. Similar with MoO₂, nearly zero differential free energy for H adsorption is achieved at the Mo and S edge [38]. A further improvement of the exposed active sites is expected for better HER activity [3]. The other dichalcogenide proven efficient in HER is MoSe₂, where the abundant exposed edge sites exhibit a higher HER activity than the inert basal plane.

In addition to combination with group VI elements (O, S, Se), a Pt-like electronic structure is formed along with the d-band contraction derived from the N atom (group V element) insertion in the Mo atom interstices [39,40]. Featured with metallic behavior and thermal stability, MoN nanosheets synthesized by liquid exfoliation or the NaCl template strategy exhibited a low overpotential in both alkaline and acidic electrolytes [21]. MoP, as another representative material combining Mo and group V elements, possesses a wide range of Mo/P compositional ratios in the compounds, including Mo rich and P rich structures [41]. The electronic, structural and physicochemical properties vary with the composition. In particular, electron transfer is accelerated through the Mo–P and Mo–Mo bonds in the Mo rich MoP [42].

Mo can also form Mo:C compounds with C (a group IV element), where the d-band of Mo is widened due to the orbital optimization of Mo and C atoms [30]. To enhance hydrophilicity and mass diffusion, a hierarchical porous structure is preferred due to the high surface area provided by micropores and fast ion transfer via the meso- and macropores [43]. In a comparison of four Mo:C phases, the HER activity followed the order: β-Mo₅C > γ-MoC > η-MoC > α-MoCₙ₋₁, suggesting the different intrinsic electrochemical activities of Mo carbides [44]. Further explorations are expected to balance the adsorption and desorption of hydrogen, in order to promote the reaction kinetics in HER.

The last Mo based compounds are Mo borides (a group III element), including MoB₃, α-MoB, β-MoB and MoB [45]. Interestingly, the Mo rich boride MoB exhibited a
lower HER activity than α-MoB and β-MoB, indicating an enhanced performance along with the increase in boron content, which was further proven by the comparable activities of MoB₄ and MoB₂ [20,46]. Significantly, Mo borides with a graphene like boron layer outperformed those with a phosphorene like boron layer, especially in acidic electrolytes. However, more investigations are suggested for catalytic behavior in alkaline solutions for Mo borides [1].

4. Smart Designs of Mo Based HER Catalysts

Despite the wide application of Mo based materials in HER, their activity and stability are still unsatisfactory due to limited active sites and sluggish electron/ion transfer. To address the issues, Mo based materials can be coupled with a conductive carbon matrix to enhance electric conductivity, the dispersion of Mo sites and specific surface area [17,47–50]. In addition, doping with heteroatoms effectively improves the intrinsic activity, lowers the kinetic energy barrier for dissociation, facilitates the charge transport and optimizes the H binding energy [26,51–54]. Moreover, the construction of heterostructures forms a synergy and benefits the HER performances [34,55]. In the following sections, the three modification strategies mentioned above for Mo based electrocatalysts will be illustrated in detail, with an in depth discussion of the structure–performance relationships in HER and a summary of the key data for representative HER catalysts in Table 1.

Table 1. Summary of the representative Mo based electrocatalysts for hydrogen evolution reaction.

| Catalyst                  | Electrolyte | η₁₀/(mV at 10 mA·cm⁻²) | Tafel Slope/(mV·dec⁻¹) | Ref. |
|---------------------------|-------------|------------------------|-------------------------|------|
| NP-MoC                    | 0.5 M H₂SO₄ | 210                    | 64                      | [56] |
| MoC/C                     | 0.5 M H₂SO₄ | 117                    | 60.5                    | [57] |
|                           | 1 M KOH     | 121                    | 73.5                    |      |
| MoC/CNFs                  | 0.5 M H₂SO₄ | 160                    | 66                      | [58] |
| MoP/MoC@C                 | 1 M KOH     | 92                     | 63                      | [59] |
| MoC-MoP/N-CNFs            | 1 M KOH     | 137                    | 65                      | [60] |
| MoC/GNR                   | 1 M KOH     | 121                    | 54                      | [61] |
|                           | 0.5 M H₂SO₄ | 155                    | 73                      | [62] |
| MoC/N-C                   | 1 M KOH     | 78                     | 64                      | [63] |
| MoC/C                     | 1 M KOH     | 165                    | 63.6                    | [64] |
| Co/Ni-MoO₂                | 1 M KOH     | 103                    | 80                      | [65] |
| MoO₂/MoS₃/P                | 1 M KOH     | 45                     | 64.2                    | [66] |
|                           | 0.5 M H₂SO₄ | 69                     | 31                      |      |
| MoO₂-Ni                   | 1 M PBS     | 84                     | 75.3                    | [67] |
|                           | 1 M KOH     | 46                     | 56.9                    |      |
| MoP@NPSC                  | 0.5 M H₂SO₄ | 71                     | 75                      | [68] |
| MoP@NCHSs                 | 1 M KOH     | 92                     | 62                      | [69] |
|                           | 1 M KOH     | 42                     | 127                     |      |
| CoP–MoO₂/MF               | 0.5 M H₂SO₄ | 65                     | 85                      | [70] |
| MoS₂/MoO₂                 | 1 M KOH     | 157                    | 119                     |      |

Note: η₁₀, overpotential at 10 mA·cm⁻²; CNTs, carbon nanotubes; CNFs, carbon nanofibers; GNR, graphene nanoribbon; NCHSs, N doped carbon hollow spheres; NPSC, N,P,S codoped carbon; MF, Mo foil.
4.1. Carbon/Mo Based Composites

Coupling with a carbon matrix or additives can increase surface area, improve Mo dispersion and control particle size due to a mesoporous structure, nanoscale morphology and protective shell [56,57,71–74]. In addition, integration with conductive carbons (e.g., graphene, carbon nanotubes, and carbon black) facilitates charge transport and accelerates reaction kinetics [33,75–77]. The promotional effects of carbons on the HER performances of Mo based catalysts are elucidated in depth below.

4.1.1. Coupling with Nanoscale Carbons

Various carbon materials with nanostructures can be introduced to Mo based catalysts to achieve a high surface area and fast electron transfer, including nanotubes [76], nanosheets [78], quantum dots [79] and nanofibers [58]. For example, by carbonizing the Mo–melamine polymer precursor on carbon nanotubes (CNTs), a high dispersion of β-Mo2C nanoparticles and enhanced electric conductivity were both realized, delivering a much lower onset potential vs. reversible hydrogen electrode (RHE) (~195 mV for 10 mA·cm⁻²) than the counterpart without CNTs (~340 mV for same current density), suggesting superior electrochemical activity [74]. Moreover, a smaller Tafel slope of 75 mV·dec⁻¹ was obtained for Mo2C/CNT than the 110 mV·dec⁻¹ for the control sample, indicating a lower activation energy and faster reaction kinetics (Table 1). The outstanding HER performances could be attributed to the abundant active sites on the uniformly dispersed Mo2C for H adsorption and activation, and the facilitated charge transport through the Mo2C–CNT interfaces [59]. Owing to its merits, no decay was presented after 1000 cycles of cyclic voltammograms, demonstrating great potential as a highly durable HER catalyst in practical applications. To further immobilize the Mo2C particles on the CNTs upon thermal treatment, oxalic acid was added to replace the ethoxy by forming oxalate groups with Mo ions (Figure 1a). Benefiting from the steric hindrance of the complex and interaction with CNTs, the Mo2C agglomeration was considerably inhibited during the 800 °C calcination, achieving a highly dispersed and small particle (4–8 nm) (Figure 1b) [80]. Owing to the reduced internal charge transfer resistance, the improved dispersion of active sites and large electrochemical surface area, the Mo2C/CNTs composite catalyst exhibited an overpotential of 110 mV for 10 mA·cm⁻² and stable activity over 15 h and 1000 potentiodynamic sweeps (Figure 1c and Table 1). Additionally, the low Tafel slope of 51.34 mV·dec⁻¹ suggested a Volmer–Heyrovsky mechanism, where the Heyrovsky desorption step determined the reaction kinetics [81]. Apart from the promotional effect on active site exposure by enhancing the dispersion and electric conductivity by increasing the Fermi level, the addition of CNTs exerted a profound influence on the Mo valence state. In particular, more Mo³⁺ were reduced to Mo²⁺, so as to generate a small Mo⁵⁺/Mo²⁺ ratio in the presence of CNTs, which strengthened the Mo–H bond, accelerated the H:O reduction in the Volmer step and facilitated the H⁺ adsorption [82].
Figure 1. (a) Growth mechanism of Mo₂C on CNTs surface. (b) TEM images of oxalate derived Mo₂C/CNTs. (c) Time dependent profile of current density achieved under a constant 110 mV overpotential, with the inset figure of the polarization curves initially recorded and after 1000 cycles. Reproduced with permission from [81]. Copyright 2019, Elsevier.

Similar to CNTs, well defined 1D carbon nanofibers (CNFs) enjoy a moderate strength Mo–H bond, facilitated mass transport and abundant active sites [82]. Via a facile electrospinning followed by pyrolysis strategy, ultrasmall hexagonal phase Mo₂C nanocrystals (5 nm in average) were anchored in the ultralong CNFs with a diameter of 200 nm. Interestingly, a synergy was generated between the Mo₂C and CNFs. Mo₂C was well confined in the CNFs to maintain a small size during the thermal treatment and the Mo–H bonding energy was adjusted; also, the electron transfer and hydrophilicity were both promoted due to the CNF coupling [83,84]. On the other hand, the embedded Mo₂C nanoparticles enhanced the thermal stability of CNFs, which maintained their original structure after high temperature calcination, in sharp contrast to the curly morphology in the absence of an Mo source. Benefiting from synergistic effects, the Mo₂C/CNFs catalyst delivered a low overpotential at 10 mA·cm⁻² (160 and 92 mV) and a Tafel slope (66 and 63 mV·dec⁻¹) in 0.5 M H₂SO₄ and 1 M KOH, respectively (Table 1). Moreover, only a negligible overpotential shift was presented after 2000 cycles at 100 mA·cm⁻² [58]. Similarly to another work, where MoC and MoP were both embedded in the CNFs, ultrafine nanoparticles with a size of 2–7 nm were obtained after 850 °C Ar calcination, indicating a considerable confinement effect on the controlled growth of active sites [60].

In addition to 1D nanostructures, carbon quantum dots (CQDs) (less than 10 nm) act as a protective layer of MoP, to inhibit agglomeration and surface oxidation due to admirable electronic conductivity, good solubility and easy functionalization [85]. For instance, a negatively charged H₃PMo₁₂O₄₀ (HPM) precursor and positively charged ethanediamine modified CQDs (ECQDs) interacted electrostatically with each other (Figure 2a) [86]. Upon calcination, the migration, growth and oxidation of MoP were considerably alleviated by the steric hindrance of the oxo clusters and immobilization effect of ECQDs, thus achieving a small MoP size (20 nm) encircled by the CQDs (Figure 2b). Moreover, the electron transfer was promoted based on the charge polarization of Mo³⁺ (3.1 eV peak separation in XPS Mo 3d spectra) and P⁵⁻ (lower binding energy than P⁰) and low charge transport resistance (13.7 Ω), potentially benefiting the HER reaction.
As a result, a low overpotential of 210 mV at 20 mA·cm$^{-2}$ and small Tafel slope of 56 mV·dec$^{-1}$ were exhibited for MoP/CQDs (Table 1), outperforming those of pristine MoP (280 mV and 87 mV·dec$^{-1}$) (Figure 2c). Furthermore, negligible current degradation was presented without morphology change over 24 h and 1000 cycles [79].

A hierarchical nanostructure can form a 3D carbon network, which increases the active material loading and accelerates the charge/mass transfer through the abundant interfaces between ultrasmall Mo based particles and electrolytes [88]. For example, 3D networks consisting of carbon nanorod arrays (10 µm in length) supported on the carbon nanosheets were derived from carbonizing the biomass natural agaric, where Mo:C nanoparticles were embedded in the carbon rod and lamellar matrices. The resulting low overpotentials of 350, 100 and 82 mV at 10 mA·cm$^{-2}$ in neutral, alkaline and acidic electrolytes, respectively, were ascribed to the enhanced diffusion of electrolytes (based on the good hydrophilicity), the abundant active sites (relating to the high surface area and Mo:C contents in 3D complex networks), and the promoted electron/ion transfer.

### 4.1.2. Coupling with Graphene Based Materials

Graphene based materials, such as graphene (G), graphene oxide (GO) and reduced graphene oxide (rGO), possess a 2D nanostructure and specific properties, including a high surface–volume ratio, chemical stability and good conductivity [89–91]. Therefore, Mo/graphene based composites are introduced separately from the nanocarbon in previous sections.

By simply mixing the GO powder and Mo precursors followed by Ar calcination, an MoO$_2$/Mo$_2$C/G composite structure was successfully prepared. The agglomeration of Mo species was effectively prevented and electron transfer was promoted with the addition of graphene, exhibiting a small Tafel slope of 59 mV·dec$^{-1}$. Similar promotional effects of graphene based matrices are reported for Mo$x$C [92], MoS$_2$ [75], MoO$_2$-Mo$_2$C [93], and MoP [94, 95], which presented stable electrolytic HER activity in both alkaline and
acids electrolytes. To further control the size of active sites and promote exposure to reactants, a conducting MoO\textsubscript{2}/GO membrane was derived from the carbonization of nanoballs synthesized in a template free hydrothermal method. In detail, initially, MoO\textsubscript{2} nanoparticles were dispersed on the GO sheets (a layer for MoO\textsubscript{2} nucleation and growth [90]); subsequently, the MoO\textsubscript{2} decorated GO sheets gradually self-assembled into hollow spheres via hydrogen bonding, π–π interactions and van de Waals force [96]. The minimized MoO\textsubscript{2} aggregation, abundant exposed active sites, high surface area, superior structural rigidity and fast transfer of H\textsuperscript{+} and electrons were contributed by the formation of MoO\textsubscript{2}/GO conducting membranes, exhibiting a much smaller Tafel slope of 52 mV·dec\textsuperscript{−1} compared with the pristine MoO\textsubscript{2} nanoparticles (130 mV·dec\textsuperscript{−1}) and only a 5% current decrease after 6 h [89].

Differing from the physical mixing of the active Mo species and graphene based matrices, the in-situ formation of the composites further strengthens the interaction, and effectively anchors the active sites. In the preparation of MoO\textsubscript{2}/rGO composites (Figure 3a), graphene as a template was first modified by poly(diallyldimethylammonium chloride) (PDDA) to be positively charged, electrostatically interacting with the negatively charged MoO\textsubscript{2} and facilitating the attachment of MoO\textsubscript{2} nanobelts on the electron rich graphene surface [97]. After annealing, MoO\textsubscript{2} nanobelts were in-situ reduced by H\textsubscript{2} and graphene, to generate a MoO\textsubscript{2}/rGO nanocomposite. Meanwhile, rGO as an electron acceptor facilitates the charge transfer. As a consequence, a small Tafel slope of 68 mV was obtained with 70% activity retention after 1000 cycles [34]. Apart from the in-situ reduction of MoO\textsubscript{2} by graphene and H\textsubscript{2}, in another case, highly dispersed MoC nanoparticles were anchored in the graphene nanoribbons (GNRs) and protected by a few graphite layers via hot filament chemical vapor deposition (HF-CVD) [61]. Particularly, H atoms derived from the H\textsubscript{2} and CH\textsubscript{4} dissociations reduced the MoO\textsubscript{2} directly into Mo metals without the formation of MoO\textsubscript{2}. Subsequently, MoC nanocrystals with a size of 3–5 nm were generated by the carburization of metallic Mo. The assembled hierarchical nanostructure, where 1D MoC was vertically aligned on the 2D graphene nanosheets, possessed the following merits: a small MoC size, loose arrangement of active sites, high electrochemical surface area, good conductivity and moderate Mo–H binding strength. In detail, a small MoC particle size exposed a high active surface to the reactants, increasing the electrocatalytic activities in HER [98]. Additionally, MoC nanocrystals were loosely distributed and covered in the graphene nanosheets, enabling H\textsubscript{2} release from the catalyst without harm. Moreover, the enhanced active surface area could be measured based on the capacitance (Figure 3b) that the MoC/GNRs exhibited a much higher capacitance of 23.34 mF·cm\textsuperscript{−2} than that of the pristine MoC (0.42 mF·cm\textsuperscript{−2}). Correspondingly, MoC/GNRs possessed a significantly rougher surface, with a factor of 1060, than the naked MoC (19). Furthermore, d-band centers shift downwards along with the charge transfer from MoC to graphene, which reduced the Mo–H bonding energy and promoted the desorption of H\textsuperscript{+}, thus improving the reaction kinetics of HER [99]. Notably, the relative inert graphene could be activated by MoC that the H adsorption free energy reduced from 1 to 0.3 eV (Figure 3c). Owing to the above merits, MoC/GNRs delivered a lower overpotential in alkaline (121 mV) and acidic (152 mV) electrolytes at 10 mA·cm\textsuperscript{−2} than those of the counterpart (266 and 275 mV, respectively) (Table 1). On the other hand, a smaller Tafel slope was also presented in both solutions for MoC/GNRs, with a negligible current density drop shown after 30,000 s in both electrolytes, demonstrating a superior electrocatalytic activity and stability over a long term operation [61].
4.1.3. Coupling with Mesoporous Carbons

By combination with mesoporous carbon matrices, the contact surface between the Mo based electrode and electrolyte was enlarged, improving the transfer of charge and mass [50]. Generally, a mesoporous structure presents a type IV isotherm of N₂ adsorption/desorption, with a large slope at high pressure regions [100]. In addition, the average pore size ranges from 2–50 nm. With such a specific structure, the electrocatalyst possesses a fast diffusion of electrolytes and high active surface for the adsorption and activation of HER [101]. For example, via the thermal carbonization of sunflower seed shells, a mesoporous carbon matrix with a high specific surface area (401.5 m²·g⁻¹) and average pore size (3.9 nm) was produced for the homogeneous distribution of Mo₂C nanoparticles (5–8 nm) [102]. The higher HER performances were attributed to the boosted electron transfer rate and interfacial reaction kinetics, which resulted from the large surface area and abundant mesopores [103]. To further enhance the reaction kinetics, a mesoporous carbon matrix with a 3D architecture was prepared by the calcination of ammonium heptamolybdate and rice paper. Owing to the fast diffusion of charge/mass through the channel rich interfaces, large exposed surface area and abundant active sites for adsorption and reaction, the Mo₂C embedded in 3D mesoporous carbons exhibited a low overpotential of 155 and 78 mV at 10 mA·cm⁻² and a small Tafel slope of 73 and 64 mV·dec⁻¹ in 0.5 M H₂SO₄ and 1 M KOH, respectively, with a stable catalytic activity over 10 h and 2000 runs (Table 1) [62].

In addition to the preparation of mesoporous carbons from natural products and textiles, metal–organic frameworks (MOFs), as a precursor, can offer a confined environment with abundant pores generated by organic ligands. In particular, an Al-MOF (MIL53) was hydrothermally synthesized and possessed a microporous structure based on the type I curves of adsorption/desorption [104]. Benefitting from the large surface area (1165 m²·g⁻¹), MoCl₅ was homogeneously impregnated in the MOF frameworks.
During the carbonization, Mo₂C was in-situ formed from the reduction of Mo species under CH₄ decomposition (Figure 4a). Meanwhile, the microporous MOFs collapsed into a rigid and mesoporous structure (type IV behavior in BET and pore size of 3.6 nm). As a result, the nucleation and growth of β-Mo₂C nanocrystals were considerably inhibited by the mesoporous texture and confined space of the MOF derived carbon matrix, producing a small and highly dispersed Mo₂C with the size of 5–10 nm, while those prepared with commercial carbon support possessed a much larger size of 20–40 nm. The carbon layers not only protected the Mo₂C from agglomeration during high temperature calcination, but also facilitated electron transfer and H₂ gas release from the system (Figure 4b). Following the Volmer–Heyrovsky mechanism, the β-Mo₂C/C composite electrocatalyst presented a 165 mV overpotential at 10 mA·cm⁻² and a Tafel slope of 63.6 mV·dec⁻¹, outperforming the Mo₂C/commercial carbon (229 mV and 74.5 mV·dec⁻¹) (Figure 4c) [63].

![Figure 4.](image)

Figure 4. (a) Formation scheme of Mo₂C/C electrocatalyst. (b) TEM of Mo₂C/C. (c) Current–potential profiles of Pt/C, Mo₂C/C and Mo₂C/XC72. Reproduced with permission from [63]. Copyright 2016, The Royal Society of Chemistry.

Compared with mesopore alone, a hierarchical porous structure generates a synergy of macropore and mesopores. Particularly, mass diffusion is facilitated through the macropores and abundant active sites are exposed in the mesopores [105]. By the facile impregnation carbonization of the ammonium heptamolybdate (Mo precursor), F127 (surfactant), resorcinol and formaldehyde, Mo₂C was formed in-situ since the carbon source was partially consumed, which was reflected by the increase in mesopore size from 4.8 to 5.6 nm. Moreover, homogeneous macropores (9 µm) and ordered mesopores (5.6 nm) coexisted after the thermal treatment, proven by an Hg intrusion test [106]. Anchored in the wall of mesopores, the migration and growth of Mo₂C nanocrystals were limited, producing an ultrasmall particle size of 5 nm. In turn, the embedded Mo₂C nanoparticles improved the hydrophilicity based on the reduced water contact angle (80.4° vs 119.8°). The combination of macropores and good wettability enabled a fast and efficient electrolyte diffusion and the escape of gas products. As a consequence, a low overpotential of 230 mV was delivered to generate a high current density of 500 mA·cm⁻². The small Tafel slope of 63 mV·dec⁻¹ indicated an outstanding HER activity and Volmer–Heyrovsky mechanism [107].
4.1.4. Core Shell Structure

To protect Mo species from agglomeration, oxidation and corrosion (etching), a core shell structure is developed with carbon layers as the shell covering the Mo based compounds [57,83,108]. Meanwhile, the presence of the carbon shell promotes the charge transfer through the electrode–electrolyte interfaces [109]. For example, when Mo2C particles were embedded in the 3D carbon network and encapsulated by the carbon shell, they interacted intimately with the carbon matrix and were separated from direct contact with the electrolytes, thus facilitating electron transfer and inhibiting corrosion [57]. Moreover, the size of the Mo2C was only 2–2.5 nm due to the confinement effect of the carbon shell, possessing a large amount of active sites exposed to the reactant. Thus, in acidic and alkaline electrolytes, a low overpotential of 117 and 121 mV were needed to obtain 10 mA·cm$^{-2}$ current density (Table 1). In addition, the small Tafel slope of 73.5 mV·dec$^{-1}$ indicated a fast kinetics in HER, benefiting from the high electrochemical active area based on capacitance [110] and low internal charge transfer resistance [57].

In the above work, the application of Polyoxometalates (POMs) has been proven a good precursor to generate highly distributed Mo based particles due to their diverse structures and unique composition [108]. To realize a better dispersion of the POMs, dicyandiamide (DCA) was mixed with the precursor; after calcination, graphitic carbon layers were formed, covering MoP/Mo2C nanoparticles to protect them from aggregation (Figure 5a) [108]. Moreover, the carbon shells reduced the charge transfer resistance at the interface between the electrocatalyst and electrolyte, reflected from the small $R_{ct}$ (7.08 Ω) based on the EIS test (Figure 5b). Owing to the above merits, an MoP/Mo2C@C core shell catalyst exhibited a low overpotential of 75 mV to achieve 10 mA·cm$^{-2}$ current density at 1 M KOH (Table 1). In addition, the admirable stability over 1000 cycles and 14 h operation was attributed to the strong resistance to agglomeration and etching with the protective carbon layers (Figure 5c) [59]. Similar findings were also reported in MoC@C, which possessed a superior performance in HER [111].

![Figure 5](image-url)

**Figure 5.** (a) Preparation scheme of MoP/Mo2C@C. (b) Nyquist plots of electrochemical impedance spectra (EIS). (c) Polarization curves of MoP/Mo2C@C. Inset: Time dependent current density curve under a constant overpotential of 60 mV. Reproduced with permission from [59]. Copyright 2017, American Chemical Society.
4.2. Heteroatom Doping

Nonmetallic (e.g., N, P, S, C) and metallic (e.g., Ni, Co) elements can be doped in the Mo compounds and carbon matrices to enhance the electrocatalytic properties in HER by increasing the number of active sites, tuning electron structure, facilitating electron transfer, improving hydrophilicity, promoting H^+ adsorption/desorption and stabilizing the Mo sites [56,64,65,75,112–114]. In the following sections, the doping effects of heteroatoms will be introduced by category, including nonmetallic atom doping (single element [74,115–119], dual elements [103,117,120,121] and triple elements [122]) and metallic atom doping [64,66,72].

4.2.1. Nonmetallic Atom Doping

Single Atom Doping

Owing to the strong electronegativity and lone pairs, nonmetallic atoms (e.g., N,S,P,C) can activate the carbon matrix, introduce structural defects and weaken Mo–H bonds, thus stabilizing the Mo species, increasing the active center amount, enhancing electronic conductivity and facilitating H2 release [56,57,67,112,113,119,123,124]. As for the N doping, abundant Mo^2+ sites are produced and Mo–H binding is moderated by Mo–N interactions [60]. Electrons are withdrawn by N atoms and H2 is easily activated [125]. In addition, the surface electronic structure is reconstructed to form an activated carbon matrix [126]. Specifically, the carbon nanolayers coupled with N atoms become wrinkled and rough, and sp2 hybrid orbitals are changed, which generates a nearly zero Gibbs free energy, thus promoting H adsorption [88]. Moreover, the carbon substrate doped with N possesses n-type and metallic character, benefiting electron transfer [127]. For example, MoC nanosheets doped with N were synthesized via chemical vapor reduction with dicyandiamide as the N source [123]. Based on the characteristic peaks of Mo^2+ at 231.6 and 228.6 eV (Mo 3d spectra) and pyridinic N at 398.5 eV (N 1s spectra), the successful introduction of N and the electron rich environment surrounding Mo were proven [4]. Benefiting from the lone pairs of N atoms, the adsorption of active H was promoted and electronic conductivity was enhanced with an increased spin density and electron density [125]. In addition, more active sites were exposed with the manipulation of electron density states by N atoms. Thus, a low overpotential of 99 mV at 10 mA·cm⁻² and a small Tafel slope of 44.5 mV·dec⁻¹ were realized for N doped MoC nanosheets [123]. The detailed mechanism was elucidated based on a DFT calculation [112]. The lone pair electrons of the N atom modulated the electronic structure of the neighboring Mo and C atoms and enabled an intimate interaction with protons and water [128]. Compared with the −1.57 eV Gibbs free energy of H atom adsorption for undoped MoC, the ΔGrs of N doped MoC was only −0.32 eV, much closer to zero, which suggested a facilitated electron/proton transfer and hydrogen release [112]. However, excessive N doping possibly inhibited electron transport from MoC to the underlying materials during H2O activation, and subsequent OH− adsorption [129].

Different N configurations were compared regarding their contributions to the HER activity of N doped MoP@carbon, including pyrrolic N, pyridinic N and graphitic N [68]. In the preparation, N atoms were introduced by the oxidative polymerization of pyrrole initiated by H3POMO6O19·nH2O (an Mo precursor). After calcination and phosphidation with the help of polystyrene spheres as the sacrificial template, MoP nanoparticles were encapsulated within the hollow N doped carbon spheres (Figure 6a). After peak fitting, the N 1s spectra were deconvolved into four configurations (pyridinic N at 397.8 eV, pyrrolic N at 400.5 eV, graphitic N at 401.4 eV and pyridinic N–O at 402.5 eV) (Figure 6b) [130]. Compared with graphitic carbon, H2O adsorption was facilitated in pyridinic and pyrrolic N atoms due to their stronger electron-withdrawing property (Figure 6c). Moreover, MoP coupled with pyridinic N possessed a lower dissociation energy barrier for H2O than the pyrrolic N, promoting the proton generation in Volmer step (Figure 6d). Notably, a combined doping with pyridinic and pyrrolic N exhibited
the smallest ΔG_{aw}, enhancing the evolution of H₂ in the subsequent Heyrovsky step. To further elucidate the dominated contribution of pyridinic N [131], the adsorption energy of OH* was compared. It turned out that the ΔE_{aw} of the pyridinic N was 2.84 eV, much higher than that of graphitic N (−0.41 eV) and pyrrolic N (1.84 eV), significantly facilitating the −OH desorption and preventing the active sites from being covered by the readsoberd water molecules (Figure 6e) [132]. Owing to the electron rich N atoms, lower d-band center, weakened Mo−H bond, more structural defects at grain boundaries, abundant Mo−N bonds on the surface and proper balance between OH* adsorption/desorption, the highly active pyridinic-N-MoP sites resulted in a low overpotential of 92 mV at 10 mA·cm⁻² [68,133].

Figure 6. (a) Fabrication schematic of MoP@NCHS. (b) XPS spectra of N 1s. (c) Average Bader charge of N doped carbon. (d) Free energy diagram of the water dissociation at pyridinic-N-MoP. (e) Chemisorption energies of OH* (right). Reproduced with permission from [68]. Copyright 2020, Wiley.
Similar to N doping, the introduction of P atoms can modulate the electronic structure and surface electron state of a carbon matrix and Mo compounds, which improves conductivity and optimizes H adsorption, thus enhancing reaction kinetics [65,134]. When P was doped into MoS$_2$, defects were generated in the disordered hexagonal structure of the MoS$_2$ basal plane, acting as the active sites for HER. Meanwhile, Mo bridges (e.g., O–Mo–P and O–Mo–S) with a shorter bond length benefited the electron transfer at the interface [65]. The doping concentration of P greatly influenced the intrinsic activity and HER kinetics. At a P/S ratio of 0.14, the Gibbs free energy of H adsorption was only 0.24 eV, consistent with the better HER performances. Specifically, the electron density of S was reduced because of the electron transfer from Mo to P, so as to regulate the adsorption and desorption of H* [135]. In comparison, excessive P doping lowered the intrinsic activity of MoS$_2$ by breaking the structure. Thus, the appropriate amount of P atoms enabled a balance between the H$_2$O dissociation and H desorption. On the other hand, a narrower bandgap of 1.58 eV was obtained by P doping, compared with the pristine MoS$_2$ (1.84 eV). Meanwhile, more charge carriers were generated based on more electron states near the Fermi level and the appearance of new bands. Owing to the merits above, a small overpotential of 45 mV was delivered, to achieve 10 mA·cm$^{-2}$ in alkaline electrolytes (Table 1) [65].

Dual Atoms Doping

Compared with single nonmetallic atom doping, dual atoms doping enjoys the synergy between the two dopants in HER. When N,P or N,S were codoped into MoC/C by precipitation, carbonization and hydrothermal treatment with phosphomolybdic acid (P source), dicyandiamide (DCA, N source) and thioacetamide (TAA, S source) (Figure 7a), the newly formed hybrids possessed a larger active area for reaction, higher intrinsic activities, lower internal electronic resistance and enhanced wettability by increasing the polarity and modulating the Mo:C d-orbitals [56,136]. Benefiting from the modulated electronic structures and improved HER kinetics, the dually doped sample (NP- and NS-doped) exhibited a lower overpotential than a single N doped counterpart (210 and 213 vs. 265 mV at 10 mA·cm$^{-2}$) (Figure 7b) [56]. In addition, based on the Tafel slope (64 and 44 vs. 78 mV·dec$^{-1}$) (Figure 7c and Table 1), the reactions all followed the Volmer–Heyrovsky mechanism and the Volmer step determined the overall rate. More importantly, the smaller slopes for the dual atoms doping suggested a faster HER kinetics resulting from moderate Mo–H bonding and abundant active centers for diffusion, adsorption and reaction [137].
The synergistic effects of dual doping are also reported in other works. In a series of couples (P–S, P–N, P–C), the electron transfer from P to the Mo sites via the other dopant was greatly promoted, stabilizing the MoP, offering more adsorption sites of H⁺ and presenting better electronic conductivity [75,92,138]. In another case, MoC possessed a higher Fermi level with the codoping of N and P, facilitating the reduction in hydrated protons at the beginning step of HER [139]. In comparison to the single N doped MoC, N,P- or N,S codoped MoC exhibited a lower ΔG_H₂ [140]. Lan et al. [57] also doped N and P into MoC nanoparticles embedded in a carbon matrix, and found that the doping enhanced the reaction kinetics and provided more adsorption sites for reactants. Based on the C_{dl} values, dually doped Mo₂C/C possessed a much higher electrochemical active surface area (20.1 mF·dec⁻¹) than the single doped and undoped samples (15.3, 10.5 and 7.3 mF·dec⁻¹).

### Triple Atoms Doping

It is well known that doping with nonmetallic heteroatoms (e.g., N, P, S) onto carbon matrices enables a fast electron and ion transport, modulates electron density and adjusts hydrogen binding energy [141–143]. Simultaneous doping of the above elements could further modify the electronic structure and enhance the reaction kinetics in HER [142]. By codoping N, P and S into the porous carbon matrix for MoP, abundant active sites were also generated for adsorption, thus leading to a low overpotential of 70 mV at 10 mA·cm⁻², with a Tafel slope of 87.2 mV·dec⁻¹ and no obvious shift of current density over 12 h [122].

To further improve the morphology of Mo species and interaction with the carbon matrix, N, P and S heteroatoms were doped with egg white (S and N source), phosphomolybdic acid (Mo and P source) and NaH₂PO₂ (P source) via hydrothermal, followed by thermal calcination [67]. In the preparation process, proteins in the egg white promoted the assembly with [PMO₁₂] clusters through hydrogen bonds, forming an inorganic–organic hybrid precursor; after being treated with NaH₂PO₂, small MoP flakes were coated by a thin layer of N,P,S doped carbon, proven by the XPS spectra of C–N, C–S and C–P bonds [144]. With this multiple doping, the spin density was adjusted and
the adsorption energies of H and –OH were tailored [145]. Meanwhile, the intimate contact between the MoP flakes and multiatom doped carbon enhanced the electron transfer [146]. Owing to the good electronic conductivity, abundant active centers and efficient ion transfer, low overpotentials were delivered in acid (71 mV), neutral (76 mV) and alkaline (50 mV) electrolytes at 10 mA·cm\(^{-2}\) with a nearly constant operation over 40 h [67].

4.2.2. Metallic Atom Doping

Similar to nonmetallic atom doping, various metal elements (e.g., Ni, Co) can be doped in Mo based catalysts to improve HER activity and stability by modulating the electronic structure, enhancing conductivity, tuning the binding energy of H and producing more active sites [147–149]. For example, Hu et al. [72] doped Ni into an MoxC/carbon fiber paper hybrid and found that the resistance to charge transfer decreased from 12.7 to 9 Ω compared with an undoped counterpart. More importantly, a good balance was struck between the adsorption and desorption of H on/from the MoxC by changing the valence state of Mo [137]. Thus, a low overpotential of 121.4 mV at 10 mA·cm\(^{-2}\) was exhibited with a 10-h stable operation [72]. To promote the electronic interaction between Ni and Mo, Ni was first electrodeposited on carbon cloth (CC) followed by preparing the Ni–Mo precursor by the hydrothermal treatment of molybdate; after calcination, an Ni-MoO\(_2\)/CC hybrid catalyst was formed [66]. Based on the positive shift to 855.8 eV (Ni-MoO\(_2\)) from 854.7 eV (Ni) in a Ni 2p XPS spectra, a strengthened interaction was suggested by the electron transfer from Ni to Mo [148]. Due to the strong Ni–Mo interaction, the O 2p orbitals in MoO\(_2\) were upraised, thus facilitating the hydrogen adsorption [150]. In the meantime, water adsorption and dissociation were improved on the positive charged Ni species, further enhancing the H adsorption on MoO\(_2\). The modified metal–H bonds could interact with H\(_2\)O molecules and electrons to release H\(_2\) via the Heyrovsky step (neutral and alkaline electrolytes) and Tafel recombination step (acidic electrolyte) [151]. As a result, this robust HER catalyst presented a low overpotential of 84, 69 and 46 mV at 10 mA·cm\(^{-2}\) in neutral, acidic and alkaline electrolytes with a superior stability over 36 h in all pH ranges [66].

Apart from Ni doping, the dual atoms doping of Ni and Co synergistically optimizes the metal–H binding energy and facilitates electron transport. Via a facile physical mixing and calcination, Co, Ni codoped MoO\(_2\) possessed a homogeneous dispersion of Co, Ni, Mo and O elements. The weak Co–H and Ni–H bonds modulated the hydrogen adsorption and desorption behaviors. In addition, a larger electrochemical active area was enabled for codoped MoO\(_2\), based on the higher C\(_{\text{dl}}\) value of 7.7 mF·cm\(^{-2}\), than the single doping (5.5 and 5.6 mF·cm\(^{-2}\) for Co- and Ni-MoO\(_2\), respectively). Moreover, benefiting from the optimized electronic structure, charge transfer resistance was reduced with the dual doping, favoring a fast HER kinetics [152,153]. As a consequence, the overpotential to achieve 10 mA·cm\(^{-2}\) was smaller for Co, Ni codoped MoO\(_2\) (103 mV) in comparison to Co-MoO\(_2\) and Ni-MoO\(_2\) (Table 1) [64].

4.3. Heterostructure Construction

Compared with the single Mo based materials applied in HER (e.g., MoP, MoS\(_2\), MoN, Mo:C and MoO\(_3\)), heterostructures constructed by different Mo species possess a synergy, such as balanced H adsorption and desorption, promoted H\(_2\)O adsorption and dissociation, mass/electron transfer ability, abundant active sites, high electrochemical surface area and enhanced intrinsic activities [59,65,93,154–156]. For example, MoO\(_3\) is featured with good electronic conductivity, high chemical stability and abundant active sites (Mo and O edges), thus being widely studied as a HER catalyst [93,157,158]. By forming a heterostructure with CoMo, the positive \(\Delta$$G_r$$ (0.15 eV) of MoO\(_3\) was complemented by the negative values of Co (−0.27 eV) and Mo (−0.37 eV) [159], improving the intrinsic HER activity at the metal/metal oxide interface [160]. Benefiting from the stabilized Mo\(^{4+}\) by the insoluble Mo species and the abundant active sites, a low overpotential of 76 mV at 50 mA·cm\(^{-2}\) was delivered [161]. Apart from CoMo, coupling with CoP and
Mo foil enabled an optimized hydrogen adsorption capability in the presence of P sites and charge redistribution at the interface of heterostructures. Owing to the ultralow $\Delta G^*$ (0.02 eV) and promoted H$_2$O adsorption, the overpotential to drive 10 mA·cm$^{-2}$ was only 42 mV (Table 1) [69]. Based on a DFT analysis, the possible mechanism was proposed that H$_2$O dissociated from the hydrophilic MoO$_2$ surface; subsequently, the produced H adsorbed onto the nearby P sites to combine with H$_2$O to release H$_2$ from CoP surface. Both the Volmer and Heyrovsky steps were accelerated at the heterointerfaces [162,163].

Besides integration with Mo metals and CoP, since MoS$_2$ enjoys a low cost and special structure, with S edges as the active centers [164], MoO$_2$–MoS$_2$ heterojunction may generate a synergy in the HER process. By vertically growing MoS$_2$ nanosheets onto MoO$_2$ nanobelts, the stacking of S–Mo–S interlayers was alleviated, exposing more active sites and facilitating electron transport [165]. Compared with MoS$_2$ (230 mV) and MoO$_2$ (280 mV), a much lower onset potential of 120 mV (vs. RHE) was delivered for MoS$_2$/MoO$_2$ heterocatalyst [119]. Notably, the ratio of MoO$_2$ and MoS$_2$ determined the HER activity. In particular, with the optimal ratio of 10:1, the smallest Tafel slope of 46 mV·dec$^{-1}$ and a negligible activity drop over 5000 cycles were achieved owing to the lowest internal electron transfer resistance ($121 \Omega$) and largest electrochemical surface area/amount of active sites ($C_{dl} = 7.22 \text{ mF·cm}^{-2}$) [155,166]. In comparison, a higher ratio (excessive MoO$_2$) inhibited the adsorption/desorption of molecules and ions and a lower ratio (excessive MoS$_2$) deteriorated the HER activity due to hindered electron transfer, thus slowing down H$_2$ production via proton recombination [155].

Differing from the sulfidation of Mo oxides, Huang et al. [70] partially oxidized MoS$_2$ to form a MoS$_2$/MoO$_2$ hybrid HER electrocatalyst with Fe$_3$O$_4$ as the redox agent and hard template in the solid phase reaction route. In particular (Figure 8a), tiny Fe$_3$O$_4$ nanoparticles were initially anchored on the MoS$_2$ basal plane; under thermal treatment in N$_2$, oxygen atoms were released from Fe$_3$O$_4$ to convert the MoS$_2$ into MoO$_2$ and SO$_2$ with the simultaneous formation of Fe$_2$O$_3$; after acid etching, porous MoS$_2$/MoO$_2$ heterostructure was generated with abundant active centers (e.g., Mo, O, S), exhibiting a synergy for reactant adsorption, dissociation, activation and desorption in a fast kinetics [36,167]. Interestingly, both the calcination temperature and Mo/Fe ratio significantly affected the catalytic performances. As shown in Figure 8b, the lowest overpotential was delivered at an annealing temperature of 550 °C; at a higher temperature (e.g., 600 °C), the drop in catalytic property might result from the exceedingly high crystallinity (Table 1). On the other hand, an appropriate oxygen concentration was obtained with a Mo/Fe ratio of 7:3, which presented the highest current density and largest electrochemical surface area (ECSA) of 196 cm$^{-2}$ (Figure 8c) [70].
Figure 8. (a) Preparation scheme of porous MoS$_2$/MoO$_2$ hybrid. (b) HER polarization curves of the MoS$_2$/MoO$_2$ samples annealed at different temperatures. (c) Electrochemical surface areas (ECSA) of MoS$_2$/MoO$_2$ with different Mo/Fe molar ratios (9:1, 8:2, 7:3, and 6:4). Reproduced with permission from [70]. Copyright 2020, Elsevier.

To facilitate electron transport and increase the ECSA of MoS$_2$, coupling with MoN was proven effective in creating more active sites and modulating the hydrogen binding energy [168]. With the help of C$_3$N$_4$, an MoS$_2$/MoN heterostructure was constructed via hydrothermal and thermal annealing [154]. The strengthened interface interaction was confirmed by XPS spectra where more apical and bridging S ligands were generated than the edge/terminal ones [169]. Following the Volmer–Heyrovsky mechanism based on the Tafel slope (59 mV·dec$^{-1}$), the heterocatalyst exhibited a low overpotential of 57 mV at 10 mA·cm$^{-2}$ [154]. Similarly, integrating MoS$_2$ with MoC and CoS$_2$ generated tremendous amounts of interfaces with abundant active sites and higher inherent conductivity for electron transfer. Based on the DFT calculation, ΔG$^{\text{H}^+}$ was greatly reduced, from 5.31 eV (MoS$_2$) to 1.99 eV (heterostructure), promoting hydrogen evolution with a 174.2 mV overpotential at 10 mA·cm$^{-2}$ [76].

In addition to the MoS$_2$–MoC couple, MoO$_2$ and MoC can also be integrated to form a heterostructure, combining the unique properties of MoC at all pH ranges and good conductivity of MoO$_2$ [170]. Abundant active sites and electron transport paths were generated from the MoO$_2$/MoC heterojunction, benefiting hydrogen adsorption and desorption behaviors [171]. The resulting enhanced intrinsic activity was reflected from the large exchange current density (6.8 × 10$^{-2}$ mA·cm$^{-2}$) [73]. MoC was also coupled with MoP to generate a strong interfacial interaction, which promoted the formation of ultrafine heterostructure MoC/MoP nanoparticles with a good antioxidation property. Due to the modulated Mo$^{2+}$ electron density and H binding energy, the inherent activity was considerably enhanced, thus delivering a high exchange current density of 8.5 × 10$^{-3}$ mA·cm$^{-2}$. With the optimal ratio of MoC and MoP, the desorption of H and reduction of proton were well balanced [82], obtaining a low overpotential of 158 and 137 mV to drive 10 mA·cm$^{-2}$ in acidic and alkaline electrolytes, respectively [60].
5. Conclusions and Prospects

In this review, a comprehensive summary of the recent development of Mo based electrocatalysts for the hydrogen evolution reaction (HER) is demonstrated. The fundamentals of HER and representative Mo compounds catalysts are introduced at first, and main contents are state of the art modifications of the Mo based structures. Particularly, by coupling with carbon nanostructures, doping with heteroatoms (nonmetallic and metallic elements), and constructing heterostructures, HER activity and stability are both enhanced. The improved performances can be attributed to the modulated adsorption, conversion and desorption of reactants and intermediates, the faster transport of electrons and ions, improved wettability and increased number of active sites. Despite existing progress, several issues are still waiting for a possible solution.

First, more efforts are expected to further enhance the HER performances of Mo based catalysts to satisfy large scale applications in industry. Recent advanced preparation strategies, such as MOF assisted synthesis and single atom structures, might be a promising option. Meanwhile, to realize the industrialized production of hydrogen, the cost of raw materials and simplicity of the synthesis procedures need special consideration.

Second, the detailed catalytic mechanisms that have been proposed for Mo based heterostructures with/without doping are not clear enough. More efforts are recommended to elucidate the structure–performance relationships and the origin of active sites, relying on theoretical studies and advanced characterization techniques.

Third, to achieve a balanced and optimal water splitting performance, the electrocatalyst design should enable a robust activity in all pH ranges, especially for HER in alkaline solutions and OER in acidic electrolytes.

Author Contributions: X.G.: conceptualization, data curation, investigation, writing—original draft, writing—review and editing; H.D.: conceptualization, data curation, investigation, writing—original draft, writing—review and editing; Q.D.: data curation, writing—original draft; Q.Z.: data curation; S.Q.: funding acquisition, resources, project administration, supervision, validation; X.L.: funding acquisition, resources, project administration, supervision, validation. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by Natural Science Foundation of Guangdong Province (2018A030310301), Guangzhou Basic and Applied Basic Research Project in China (202102020134; 202102020690) and Youth Innovation Talents Project of Guangdong Universities (natural science) in China (2019QKNCX098).

Data Availability Statement: All data included in this study are available upon the permission from the publishers.

Conflicts of Interest: The authors declare no conflict of interest.

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