The discovery of moiré superlattices (MSLs) opened an era in the research of ‘twistronics’. Engineering MSLs and realizing unique emergent properties are key challenges. Herein, we demonstrate an effective synthetic strategy to fabricate MSLs based on mechanical flexibility of WS2 nanobelts by a facile one-step hydrothermal method. Unlike previous MSLs typically created through stacking monolayers together with complicated method, WS2 MSLs reported here could be obtained directly during synthesis of nanobelts driven by the mechanical instability. Emergent properties are found including superior conductivity, special super-aerophobicity and superhydrophilicity, and strongly enhanced electro-catalytic activity when we apply ‘twistronics’ to the field of catalytic hydrogen production. Theoretical calculations show that such excellent catalytic performance could be attributed to a closer to thermo-neutral hydrogen adsorption free energy value of twisted bilayers active sites. Our findings provide an exciting opportunity to design advanced WS2 catalysts through moiré superlattice engineering based on mechanical flexibility.

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Moiré superlattices (MSLs) created by a twist inspire a hot area of ‘twistronics’ for advanced materials sciences. With the help of periodic moiré patterns, MSLs could optimize the structure and energy band, resulting in many phenomena, including moiré phonon1, moiré exciton2–5, magnetism6, topological edge states7,8, unconventional superconductivity9–11, Mott31 and so on. MSLs exhibit promising applications in electronics12–15, optoelectronics16,17, valleytronics18, photonic19–22, spintronics23–25, and electrocatalysis26–30.

The commonly used physical and chemical methods to fabricate MSLs through stacking together are quite complicated and require the use of specific substrate and experimental conditions11–15,31–33. Consequently, it is desirable to develop an easy and versatile strategy to construct MSLs and present an ideal model system for investigating the emergent properties. Recently, Eli Sutter et al. developed a method for the preparation of van der Waals chiral nanowires distorted by layered crystals, extending the path of interlayer distortion to achieve MSLs from two-dimensional planes to one-dimensional nanowires. However, this method still needs the use of substrates, which are difficult to produce on a large scale. It is very meaningful to explore a class of van der Waals one-dimensional (1D) nanostructures of layered crystals, in which MSLs evolve naturally during synthesis without substrate. TMDs nanobelts, combining both the flexibility and unidirectional properties of 1D nanomaterials would enable the production of MSLs easily through spontaneous deformation. Herein, we successfully synthesize large-scale homogeneous MSLs based on the mechanical flexibility of WS₂ by a facile and reproducible one-pot hydrothermal method. MSLs could be well introduced to WS₂ along with their controllable growth. Ultrathin WS₂ nanobelts with high flexibility can spontaneously bend and twist into the helix nanocones arising from mechanical instability. The bending and twisting could cause the S–W–S layer to slip for the production of MSLs.

Furthermore, we find the emergent properties of nanocone-like WS₂ MSLs, such as superior conductivity, special super-aerophobicity and superhydrophilicity, which brought unexpected catalytic hydrogen production performance by comparison with various other WS₂ based electrocatalysts. The as-synthesized WS₂ MSLs electrocatalysts display an over-potential of 60 mV at a current density of 10 mA cm⁻² and a Tafel slope of 40 mV dec⁻¹. Meanwhile, the unique nanostructures of WS₂ MSLs with the superhydrophilic property for the rapid access of the electrolyte and the underwater super-aerophobic property further facilitate the fast mass transfer characteristics of WS₂ MSLs. The experimental results are supported by theoretical calculations and the underlying mechanism is ascribed to much more appropriate ΔGₛ of twisted bilayers WS₂ active sites compared with that of normal bilayers WS₂.

**Results**

**Synthesis and structural characterization of WS₂ MSLs.** Owing to the excellent mechanical properties of ultrathin 1D and 2D materials, various specific topological structures such as ripples, bends, scrolls, helices, wrinkles, folds, and curls are shown in Supplementary Fig. 1, could spontaneously form by the thermodynamic and mechanical factors during the synthesis process. These unique topological structures may bring rich and excellent electronic properties. To achieve this goal, herein, a unique WS₂ topology deformed from nanobelts has been designed by a facile hydrothermal method.

Field-emission scanning electron microscope image (Fig. 1a) illustrated the uniformity of the as-prepared WS₂ MSLs at a large-scale view, consisting of numerous conical nanoarray with an average width of ~200 nm. The electrodes consist of 3D WS₂ MSLs with open space were in favor of electrolyte ion transport. The SEM-energy dispersive spectrometer element mapping images (Supplementary Fig. 2) showed the uniform coverage of W and S elements on the surface of the WS₂ nanoarray. As shown in Fig. 1b, the single conical tube was curled from nanobelts as indicated by scanning transmission electron microscopy. One end remains nanobelt, meanwhile, the other end has transformed into coin-like. In the synthesis process, the WS₂ nanobelts easily deform under the various unbalanced external forces and then twist into nanocones. Finite-element calculations of strain in a nanocone (Fig. 1c) showed the relative stress distribution with negligible strain at no twisted end and big strain at another largely twisted end. The strain introduced by twisting may contribute to activating the basal plane of the nanobelts by changing the electronic structure of catalytic active sites and facilitating mass transfer35. As shown in Fig. 1d, S–W–S layer slipping could be triggered by the mechanical instability, accompanying the generation of MSLs.

Moiré superlattice is created through stacking two monolayers together with respect to each other, along wavelength periodic modulation results36. The van der Waals force of multilayer WS₂ nanobelt is further weakened by the strain that is induced by the mechanical instability. WS₂ nanobelt would transform to a WS₂ nano-cone with layer slipping that could induce the formation of moiré patterns. The significant honeycomb-structured moiré patterns are found throughout the measured HRTEM images region in Fig. 2a, b. Figure 2c illustrated the simulated model of the WS₂ MSL atomic structure with a twisted θ (14°), which has high consistency with the structure shown in Fig. 2b. The corresponding FFT pattern of the HRTEM image in the sheet contains 12 (1100) spots, which constitute two hexagons (double sets of sixfold symmetry diffraction spots), as illustrated in Fig. 2d. The FFT patterns and IFFT images of HRTEM lattice images were performed, as shown in Fig. 2e, f, respectively. On the basis of the splitting spots in the FFT patterns (Fig. 2g), the moiré patterns in this region exhibited twist angles of 13.82°. We collected ten HRTEM images and their corresponding FFT images taken from different WS₂ nanocones to ensure the twist angle in Supplementary Fig. 3. All the values of the twist angles are in the range of 13°–14°. We build a model with the rotational stacking faults of 13.2° as shown in Supplementary Fig. 4, which is the calculable model closest to the rotational stacking faults of 13°–14° obtained by the experiment. The high consistency between the experimental HRTEM images and the simulated HRTEM images is presented in Supplementary Fig. 5. Dislocations of atomic planes and strain distributions of corresponding lattice planes were indicated by Geometric phase analysis images in Fig. 2h. i obtained from HRTEM image (Fig. 2a). It is obviously illustrated that the strain was introduced successfully by topology engineering based on mechanical flexibility.

2H (trigonal prismatic), 1T (octahedral) and 1T⁺ (clustered W) phase were different phases of WS₂, and their atomic structure models were shown in Supplementary Fig. 6. Bending and twisting could induce the glide of S atoms in basal planes of WS₂ MSLs to generate a 1T’/1T⁺ phase. X-ray photoelectron spectroscopy (XPS) and Raman spectroscopy were used to distinguish the 1T’/1T⁺ and 2H phase of WS₂. To make a better comparison, 1T’-WS₂ NSs were also successfully prepared and well-characterized by TEM, XRD, and XPS in Supplementary Figs. 7–9. The XPS characterization of 1T’-WS₂ NSs in Supplementary Fig. 9 showed that 1T’ content in 1T’-WS₂ NSs reaches nearly 100%. As shown in Supplementary Fig. 10, two peaks at around 34.7 and 32.7 eV were characteristic of the 2H-WS₂ features corresponding to W4f⁷/₂ and W4f⁵/₂, respectively37–39. The new peaks of the 1T’ phase clearly shifted toward lower binding.
energies (33.1 and 31.1 eV, corresponding to W4f5/2 and W4f7/2 of 1T-WS2 components). The result reveals the obvious formation of the 2H and metallic 1T phase in WS2 MSLs. Raman characterizations of all samples are shown in Supplementary Fig. 11. Two prominent peaks corresponding to the in-plane $E_{2g}$ and out-of-plane $A_{1g}$ modes of 2H-WS2 are observed in WS2 MSLs. The WS2 MSLs sample also exhibits small peaks in the lower frequency region that correspond to the active modes of distorted 1T-WS2 NSs which are not allowed in the 2H-WS2 NSs. The three peaks should be ascribed to the $I_1-I_3$ vibration modes of S–W–S bonds in 1T-WS2 phase, clearly demonstrating the coexistence of 1T-WS2 in WS2 MSLs38–41. In addition, the HRTEM analysis unambiguously confirmed the formation of 1T@2H-WS2 heterostructures (Supplementary Fig. 12). The 1T-WS2 structure could be locked by a collective elastic-deformation barrier from distortion against the transformation into the more stable 2H polymorph.

**Catalytic activity.** The as-prepared 2H-WS2 NSs, 1T-WS2 NSs, WS2 MSLs, and commercial Pt/C (20 wt%) were used to investigate the HER electrocatalytic performance. Polarization curves of these samples with a scan rate of 10 mV s$^{-1}$ in Ar-bubbled 0.5 M $\text{H}_2\text{SO}_4$ are shown in Fig. 3a and Supplementary Fig. 13. All electrochemical performance tests of catalysts were carried out on carbon fiber cloth (CFC). As shown in Supplementary Fig. 14, the bare CFC exhibits negligible electrocatalytic performance. As shown in Fig. 3a, the low overpotential of just 60 mV vs. RHE under the current density ($J = 10 \text{ mA cm}^{-2}$) was needed for the WS2 MSLs, which is smaller than those of other WS2 samples, such as the as-prepared 2H-WS2 NSs (248 mV vs. RHE) and 1T-WS2 NSs (212 mV vs. RHE), except for the commercial Pt/C (20 wt%). The Tafel slope suggests that the HER reaction of WS2 MSLs may follow a similar Volmer–Heyrovsky mechanism and is closely related to electrochemical desorption$^{12-45}$, unlike the Pt/C electrocatalyst (30 mV dec$^{-1}$) via the Volmer–Tafel mechanism (Fig. 3b). The as-prepared WS2 MSLs exhibited much more excellent HER performance (e.g., low overpotential at $J = 10 \text{ mA cm}^{-2}$ and small Tafel slope) than the reported representative non-precious HER electrocatalysts. (Supplementary Fig. 15 and Supplementary Tables 1 and 2).

To verify the long-term stability, the as-prepared WS2 MSLs were tested in a prolonged run by chronoamperometry test (Fig. 3c). Compared to the stability of 1T-WS2 NSs (Supplementary Fig. 16), WS2 MSLs have not weakened during the chronoamperometric response for 20 h.

Accurate determination of the electrochemically active surface area (ECSA) is vitally important to evaluate the electrocatalytic activity of catalyst. In general, the ECSA of each catalyst is estimated from measurements of the double-layer capacitance ($C_{dl}$). The $C_{dl}$ values of given electrocatalysts are determined by cyclic voltammetry (CV) in a non-faradaic region or by electrochemical impedance spectroscopy (EIS)$^{46–48}$ As shown in Supplementary Figs. 17–20, using the CV method, plotting the cathodic and anodic current as a function of the scan rate revealed a linear function, where the slope indicated the $C_{dl}$. As shown in Supplementary Fig. 21 and Supplementary Table 4, the fitting parameters of EIS further verified the $C_{dl}$ values of measured from the scan rate-dependent CVs (within 15% difference$^{46}$). We assume the general specific capacitance of 60 $\mu\text{F cm}^{-2}$ to estimate ECSA from the $C_{dl}$ values of catalysts$^{49–51}$ (Detailed calculation and analysis can be found in Supplementary Note 1). The ECSA value of WS2 MSLs (396.6 $\text{cm}^2$ ECSA) was much higher than that of 1T-WS2 NSs (253.3 $\text{cm}^2$ ECSA) and 2H-WS2 NSs (190.0 $\text{cm}^2$ ECSA), indicating that the WS2 MSLs possessed more enrichment of active sites for electrochemical hydrogen evolution (Fig. 3d and Supplementary Table 3).

Moreover, the fitted $R_s$ values for WS2 MSLs, 1T-WS2 NSs, and 2H-WS2 NSs are 1.6, 3.4, and 11.2 $\Omega$, respectively (Supplementary Fig. 22). The result suggests that the surface of...
WS₂ MSLs has excellent interfacial charge transfer kinetics for electrocatalysis. To demonstrate the superior conductivity of WS₂ MSLs, the total potentials of WS₂ MSLs with misorientation angles of 13.2° were investigated by density functional theory (DFT) calculations. As shown in Supplementary Fig. 23, apparently, the potential barriers of WS₂ MSLs with different phases reduced in contrast with normally stacked bilayer WS₂, indicating that electron orbitals coupling in WS₂ MSLs became much stronger²⁸. Thus, electrons transfer much more easily between two adjacent layers that would have good effect on HER catalytic properties of WS₂ MSL.

Exchange current density ($j_0$) was used to evaluate the HER activity of different WS₂ catalysts⁵² (Supplementary Fig. 24 and Supplementary Table 3). The $j_0$ of 2.13 μA cm⁻² ECSA for the WS₂ MSLs sample surpasses the values of 1.55 μA cm⁻² ECSA for 1T'-WS₂ NSs sample and 1.09 μA cm⁻² ECSA for 2H-WS₂ NSs sample, highlighting the electrochemical activity of WS₂ MSLs. The turnover frequency (TOF) was used to determine the intrinsic activity of WS₂ MSLs⁴⁹,⁵⁰,⁵³,⁵⁴. As shown in Supplementary Table 5, our results have demonstrated that the TOF (at −0.2 V vs. RHE) of WS₂ MSLs is 0.739 s⁻¹, much larger than that of 1T'-WS₂ NSs (0.090 s⁻¹) and 2H-WS₂ NSs (0.078 s⁻¹), indicating the significantly enhanced intrinsic activity of WS₂ MSLs (see the Supplementary Note 2 for details on the calculation of the TOF values). The excellent intrinsic activity of the WS₂ MSLs catalyst is likewise evidenced by its ECSA-normalized current density (Fig. 3d, Supplementary Fig. 25, Supplementary Table 3) and the comparison of mass activity with other WS₂-based electrocatalysts (Supplementary Table 6, Supplementary Note 3).

Excellent mass transfer performance has emerged as an essential factor to evaluate the property of high-efficiency electrocatalysts for HER. In the macro presentation, mass transfer is mainly the gas evolution and the contact between electrolytes and electrode surface, where it occurs at the solid–liquid–gas three-phase interface. Therefore, the wetting state of the electrode surface has become a significant factor to influence the whole mass transfer performance⁵⁵. The contact angle (CA), as a parameter to measure the wettability at the intersection of gas, liquid, and solid, is one of the important criteria for evaluating wettability and even mass transfer performance⁵⁶. Generally, solid surfaces with CAs < 90° are considered to be hydrophilicity, and those with CAs > 90° are hydrophobicity. Moreover, solid surfaces with CAs < 10° are considered to be superhydrophilicity⁵⁷–⁵⁹ (see Supplementary Fig. 26 for detail). The CAs on the electrode surface of bare CFC and WS₂ MSLs-CFC are 127.1° and 9.1°, respectively, indicating the significant hydrophilicity of WS₂ MSLs (Fig. 4a), which benefits from the
unique micro–nanostructure and metallic phase of WS$_2$. As shown in Supplementary Fig. 27, the hydrated cation preferentially adsorbs onto the 1T–WS$_2$ surface, evidenced by more favored adsorption energy (−3.45 eV) as compared to slightly weaker adsorption energy of the 2H phase (−1.82 eV). In addition, the surface wettability of the electrode material under the electrolyte was investigated by measuring the CA of the 5 μm diameter when they leave the surface of WS$_2$ MSLs as shown in Supplementary Fig. 28. As shown in Fig. 4f, the simplification of the as-generated gas bubbles releasing from the surface of WS$_2$ MSLs to demonstrate the morphological evolution of WS$_2$ nanoarray electrodes during the HER process. The circular three-phase contact line (TPCL) (white line) of as-formed bubbles on the electrode surface and exhibit higher transportation velocity on the electrode surface (Supplementary Fig. 28). As shown in Fig. 4d, compared with the continuous TPCL on the ideal flat electrode surface (Supplementary Fig. 28 and Supplementary Movie 1), the cut three-phase contact line by nanoarray makes the big bubble split into small ones more naturally (Fig. 4c and Supplementary Movie 2), maintaining rapid contact between the electrodes and electrolyte and deterring the formation of inactive sites (Fig. 4e). The relatively smaller bubbles have much lower adhesion force with electrode surface and exhibit higher transportation velocity on cones. As shown in Fig. 4f, the simplified stress analysis on a single bubble at the electrode surface indicated that the adhesion force ($F_a$) plays a pivotal part in gas bubble detachment. As expected, a small bubble adhesive force (10.4 ± 1.5 μN) was measured on the WS$_2$ MSLs–CFC surface underwater, accompanied with negligible shape change of the gas bubble (Fig. 4g and Supplementary Movie 3). The aerophilicity manifestation of bare CFC in the adhesion measurements also further verified the superaerophobic property of the WS$_2$ MSLs (Supplementary Fig. 29 and Supplementary Movie 4). The obtained results definitely demonstrate that the superhydrophilic and superaerophobic characteristics of the unique micro/nano surface structure of WS$_2$ MSLs play vital roles in accelerating HER kinetics.

**HER enhancement mechanism.** To reveal the origin of enhanced catalytic activity of WS$_2$ MSLs, the electronic properties of WS$_2$ MSLs were investigated by DFT in Supplementary Fig. 30. Obviously, the charge densities of WS$_2$ MSLs is bigger and clearer than that of normally stacked bilayer WS$_2$, indicating much stronger electron orbitals coupling in WS$_2$ MSLs. Supplementary Fig. 31 shows various active sites of 1T@2H WS$_2$ nanobelts for catalytic HER. The optimized structural model of monolayer WS$_2$ plane consisting of 2H and 1T phases (Supplementary Fig. 32a, b), shows that the WS$_2$ plane has undergone significant deformation with strain. The Δ$G_{H}$ has been demonstrated to be a successful descriptor of the HER activity, where a value of Δ$G_{H}$ closer to zero results in the higher activity. The Δ$G_{H}$ was calculated for H adsorption on sites as marked in Supplementary Fig. 32a, b. shows that the WS$_2$ plane has undergone significant deformation with strain. The Δ$G_{H}$ has been demonstrated to be a successful descriptor of the HER activity, where a value of Δ$G_{H}$ closer to zero results in the higher activity. The Δ$G_{H}$ was calculated for H adsorption on sites as marked in Supplementary Fig. 33. Obviously, the favorable sites for HER are distributed on the edges, polymorphs interface, and strained metallic phase surface. A linear relation between Δ$G_{H}$ and $p$ band center of S atom in Supplementary Fig. 32 indicated that the intrinsic activity of HER active sites was closely related to the p band center of S atom. The influence on Δ$G_{H}$ of interesting MSLs was investigated by theoretical calculations. We performed DFT calculations on non-twisted bilayers WS$_2$ and twisted bilayers WS$_2$ with 14° to study the influence of Δ$G_{H}$ induced by the twisted effect in Supplementary Figs. 34–38. Computational predictions for the MSLs effect on the HER activity indicated that the active sites of W-edge and S-edge of twisted bilayers WS$_2$ have much more appropriate Δ$G_{H}$ compared with normal bilayers WS$_2$ in.
Fig. 5. The atomic structure can fine-tune the electronic structure of the active sites by upshifting the d band center of W atoms and p band center of S atoms, which indicates the same trend of the enhanced hydrogen binding energy, thus promoting the HER performance (Supplementary Fig. 36c and Supplementary Fig. 38c). Accordingly, we evaluated the HER activity of 2H-WS2 MSLs and 1T'-WS2 MSLs using DFT by comparing \( \Delta G_H \) for hydrogen adsorption at both basal planes (Supplementary Fig. 39). The \( \Delta G_H \) close to zero at 1T'-WS2 MSLs basal plane site (−0.24 eV) validates the HER activity of WS2 MSLs relating to the base plane in this study. Therefore, the enhanced HER activity can be ascribed to the synergistic effect of phase and MSLs.

**Discussion**

Through a combined theory-experiment approach, we identify and develop highly active WS2 MSLs based on mechanical flexibility for the HER. We ascribe the total activity enhancement to a combination of electronic, geometric, superaerophobic, and superhydrophilic effects. The highly active under-coordinated sites at the edges, polymorphs interface, strained metallic phase surface of WS2 MSLs are more active than those of non-twisted bilayers WS2. \( \Delta G_H \) is sensitive to the MSLs, which implies that engineering MSLs of WS2 or other TMDs can be a route for their catalytic property engineering. This research extends twistronics and moiré fringe physics to HER catalysts and opens the way to new materials for energy conversion technologies.
possibility of designing the type of catalyst by topological physics engineering based on mechanical flexibility.

**Methods**

**Materials synthesis**

Synthesis of WS$_2$ MSLs. A facile one-step hydrothermal method was used to fabricate WS$_2$ MSLs. For the synthesis, 0.5 mmol (NH$_4$)$_2$WO$_4$·xH$_2$O and 30 mmol CH$_3$N$_2$S were dispersed in 35 mL distilled water (60 °C) by sonication for 1 h. The hydrothermal reaction was carried out in a 45 mL Teflon-lined stainless-steel autoclave. The above mixture solution was transferred into the autoclave and maintained at 300 °C for 100 h. After cooled to room temperature gradually, the as-obtained product was centrifuged and dried in a vacuum at 60 °C.

Synthesis of metallic 1T'-phase dominated WS$_2$ nanosheets, referred to as 1T'-WS$_2$ NSs. Typically, 0.2 mmol (NH$_4$)$_2$WO$_4$·xH$_2$O and 2.4 mmol thioureas were added to a 100 mL three-neck flask containing 40 mmol oleylamine (OM, 70%, Sigma-Aldrich) at room temperature. Then, vacuum the system for 5 min and inject of N$_2$ for 5 min. This process is cycled three times and carried out at 120 °C. The mixture solution was rapidly heated to 280 °C and vigorously stirred for 90 min under an N$_2$ atmosphere. After cooled to room temperature gradually, the black products were filtered and washed 5 times with cyclohexane and ethanol. The as-obtained 1T'-WS$_2$ nanosheets were centrifuged and dried in a vacuum at 60 °C.

Synthesis of 2H phase WS$_2$ nanosheets referred to as 2H-WS$_2$ NSs. For comparison, the 2H-WS$_2$ sample was prepared by heating the 1T'-WS$_2$ sample at 300 °C for 2 h in a vacuum.

**Material characterizations.** TEM images were measured by transmission electron microscope (Hitachi HT7700). High-resolution TEM images were operated at an acceleration voltage of 200 kV (FEI Talos F200X). SEM images were measured using an FE-SEM (S-4800). XRD patterns were recorded with an X-ray diffractometer (Bruker AXS D8 Advance A25), using Cu K$_\alpha$ radiation (λ = 1.5406 Å) over the range of 2θ = 5.0–80.0°. Data were collected using 2θ scan step of 0.02° at a rate of 2° min$^{-1}$. XPS measurements were conducted using an X-ray photoelectron spectrometer (KRATOS Axis Supra). A 200 W Mg X-ray excitation was used. All the samples were analyzed with reference to adventitious carbon 1 s peak. Raman spectra were recorded by a confocal Raman microscope (inVia, Renishaw, England) equipped with a 532 nm He–Ne laser as an excitation source.

Electrochemical measurements. All the electrochemical experiments were carried out using a conventional three-electrode system on an Electrochemical Workstation (CS310, Wuhan Kesite Instrument Co., Ltd.). All electrochemical performance tests of samples were carried out on CFR (Phychemi (HK) Company Limited-W051010). A typical three-electrode configuration was used to investigate all samples' HER performance with an Ag/AgCl electrode and a graphite rod as the reference and counter electrodes, respectively. All the electrochemical measurements were conducted in Ar-bubbled 0.5 M H$_2$SO$_4$ electrolyte at room temperature. All potentials were referenced to the reversible hydrogen electrode (RHE). Before the
electrochemical test, the fresh as-prepared 1T-WS2 NSs product and 2H-WS2 NSs were added into a 100 mL Erlenmeyer flask containing 3 mL thioglycolic acid and 50 mL ethanol, and vigorously stirred for 12 h under N2 atmosphere to partially remove the surfactant molecules. After that, the acid-treated 1T-WS2 NSs were separated from the solution by centrifugation (8500 rpm, 10 min), washed twice with ethanol. The catalyst dispersion was prepared by mixing 5.0 mg of catalyst in 1 mL ethanol. The Dataphysics DCAT25 system was used for iR compensation by the equation of corrected current density ($J - a$) plotted as a function of log current (log $I$) to obtain a Tafel plot for evaluating the HER kinetics of the electrocatalyst. The Tafel slope ($\eta$) to HER kinetics in the ECSA. For CV measurements, a series of CV curves were performed at various scan rates (10–140 mV s$^{-1}$) in the 0.25–0.30 V vs. RHE region. The cathodic (C) and anodic (A) charging currents tested at 0.275 V (vs. RHE) plotted as a function of log current ($J = b \log (I) + a$). By extrapolating the linear region back to zero overpotential, the exchange current density ($i_0$) can be obtained from the Tafel plots in Fig. 3b. CV and EIS were performed to evaluate the electrochemical double-layer capacitance ($C_D$) of the materials at non-Faradaic processes as the means of estimating the corresponding electrochemical active surface areas (ECSA). For CV measurements, a series of CV curves were performed at various scan rates (10–140 mV s$^{-1}$) in the 0.25–0.30 V vs. RHE. The cathodic (C) and anodic (A) charging currents corrected ($J_{\text{corr}} = J_{\text{actual}} - \frac{1}{R_C} \Delta V_{\text{fl}}$).

Bubble adhesion force test. The electrolyte used in all tests is Ar-bubbled 0.5 M H$_2$SO$_4$. The CAs between the electrolyte and electrode surface was tested by using a KRUS (DS2A0) system in ambient air. The Dataphysics DCA2T5 system was used to measure the ECSA of the catalyst. The CAs were used to evaluate the ECSA of catalysts. The ECSA values were obtained by calculating the average of the absolute value of the fitted line slope. The ECSA values were used to estimate the ECSA of catalysts. The ECSA values were calculated using the following equation:

$$\eta = b \log (I) + a$$

By extrapolating the linear region back to zero overpotential, the exchange current density ($i_0$) can be obtained from the Tafel plots in Fig. 3b. CV and EIS were performed to evaluate the electrochemical double-layer capacitance ($C_D$) of the materials at non-Faradaic processes as the means of estimating the corresponding electrochemical active surface areas (ECSA). For CV measurements, a series of CV curves were performed at various scan rates (10–140 mV s$^{-1}$) in the 0.25–0.30 V vs. RHE region. The cathodic (C) and anodic (A) charging currents. The Tafel slope ($b$) to HER kinetics in the ECSA. For CV measurements, a series of CV curves were performed at various scan rates (10–140 mV s$^{-1}$) in the 0.25–0.30 V vs. RHE region. The cathodic (C) and anodic (A) charging currents corrected ($J_{\text{corr}} = J_{\text{actual}} - \frac{1}{R_C} \Delta V_{\text{fl}}$).

Data availability

The data that support the findings of this study are available from the corresponding authors upon reasonable request. All source data underlying Figs. 3a–d, 4a, b, d, g, and 3b are provided as a Source Data file.

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Author contributions

Q.Z. and W.H. conceived the project. L.L.W. designed, supervised, and analyzed the whole project. L.B.X. carried out the materials synthesis and electrochemical test. L.L.W. and L.B.X. wrote the paper together. S.I.L. and W.W.Z. analyzed the data. All the authors contributed to the discussion during the whole project.

Competing interests

The authors declare no competing interests.

Additional information

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