

Electronic Supplementary Information

Hierarchical MoS₂@MoP Core-Shell Heterojunctions Electrocatalysts for Efficient Hydrogen Evolution Reaction over a Broad pH Range

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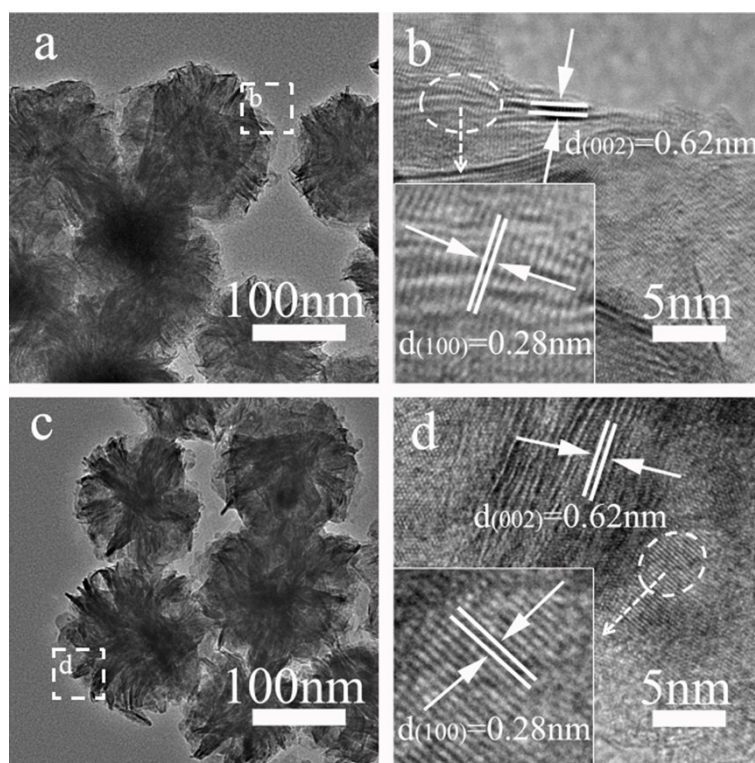


Figure S1. The low-magnification TEM and HRTEM images of HF-MoSP-800 (Figure S1a and b) and HF-MoSP-900 (Figure S1c and d).

Figure S1 shows the low-magnification TEM and HRTEM images of HF-MoSP-800 (Figure S1a and b) and HF-MoSP-900 (Figure S1c and d). The sphere-like structure have no large change before and after phosphorization reaction. The HRTEM test shows the formation of the thin sheet-like structure on outside surface of HF-MoSP-800, instead of the silk ribbon-liked structure in original HF-MoS₂ samples (Figure 1d). The presence of intimately contacted MoS₂ and MoP phase can be verified by HRTEM. The plane distance of 0.62 nm (002 for MoS₂) and 0.28 nm (100 for MoP) can be obviously seen (Figure S1c). The plane distance of 0.62 nm (002 for MoS₂) and 0.28 nm (100 for MoP) can be obviously seen (Figure S1d).

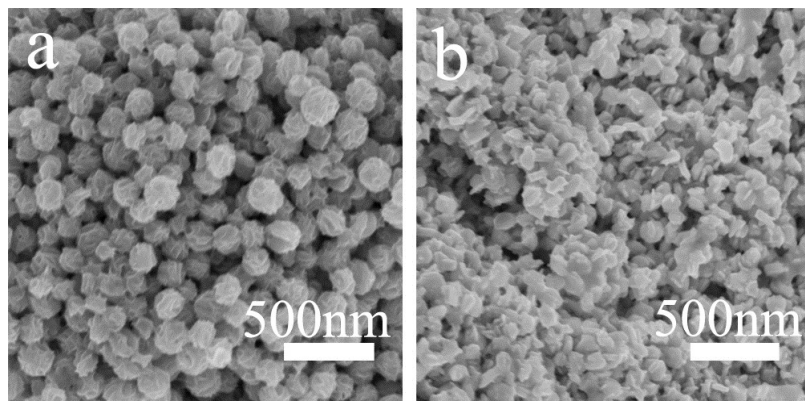


Figure S2. SEM images of HF-MoSP-700 (a) and **MoSP-1000** (b)

Figure S2 shows SEM images of HF-MoSP-700 (a) and **MoSP-1000** (b). We can see that the hierarchical spherical structure can be well remained for HF-MoSP-700 samples. The size and surface morphology have no obvious change after the phosphorization at 700 °C. However, for **MoSP-1000** sample, a larger change can happened. The low-magnification SEM image in Figure S2b shows the presence of irregular particles with size smaller than that of HF-MoS₂ sample. The irregular shape and smaller size should be due to destroying and shrink of HF-MoS₂.

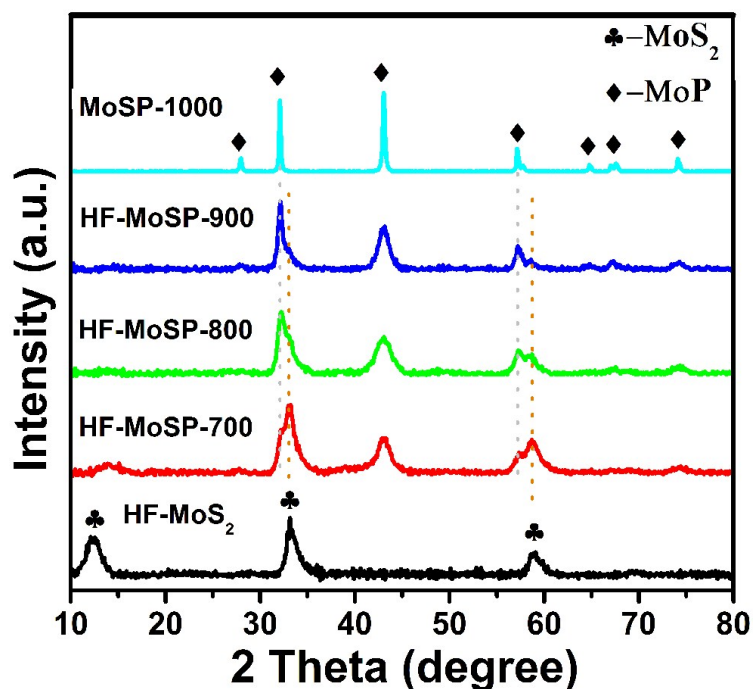


Figure S3. XRD patterns of HF-MoS₂, HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and **MoSP-1000**.

XRD patterns of different sample are shown in **Figure S3**. For HF-MoS₂, the intensive peaks about MoS₂ can be seen. The peaks located at about 13°, 33° and 59° can be indexed to the (002), (100) and (110) diffraction of hexagonal structured MoS₂. After the phosphorization at 700 °C, the 2 θ value of (002) diffraction peak was shifted to 14° revealing the decreased interlayer distance of (002) plane from 0.7 to 0.62nm and the new peaks at 32 °, 43° and 57° emerge, corresponding to the diffraction of (100), (101) and (110) of MoP, respectively (JCPDS No. 89–5110), besides the peaks belonging to MoS₂. The results indicate that the samples are composed of MoP and MoS₂ composite. The peak intensity of MoP phase increases gradually with the increase of phosphorization temperature. The intensity ratios of (100) diffraction of MoP/MoS₂ phase are about 0.63, 1.7 and 2.6 for HF-MoSP-700, HF-MoSP-800 and HF-MoSP-900 samples. For **MoSP-1000** sample, the intensive peaks belonging to MoP phase can be seen, and those corresponding to MoS₂ are not obvious. The results show that the microstructure (phosphorization degree) can be well tuned. Also, we can see that the hierarchical structure can be well remained at broad temperature range until the 900°C. The stability of the hierarchical, spherical

structures at broad range provides large space for the preparation of hierarchical MoSP with tuned structure, being favourable for tuning the performance of the materials.

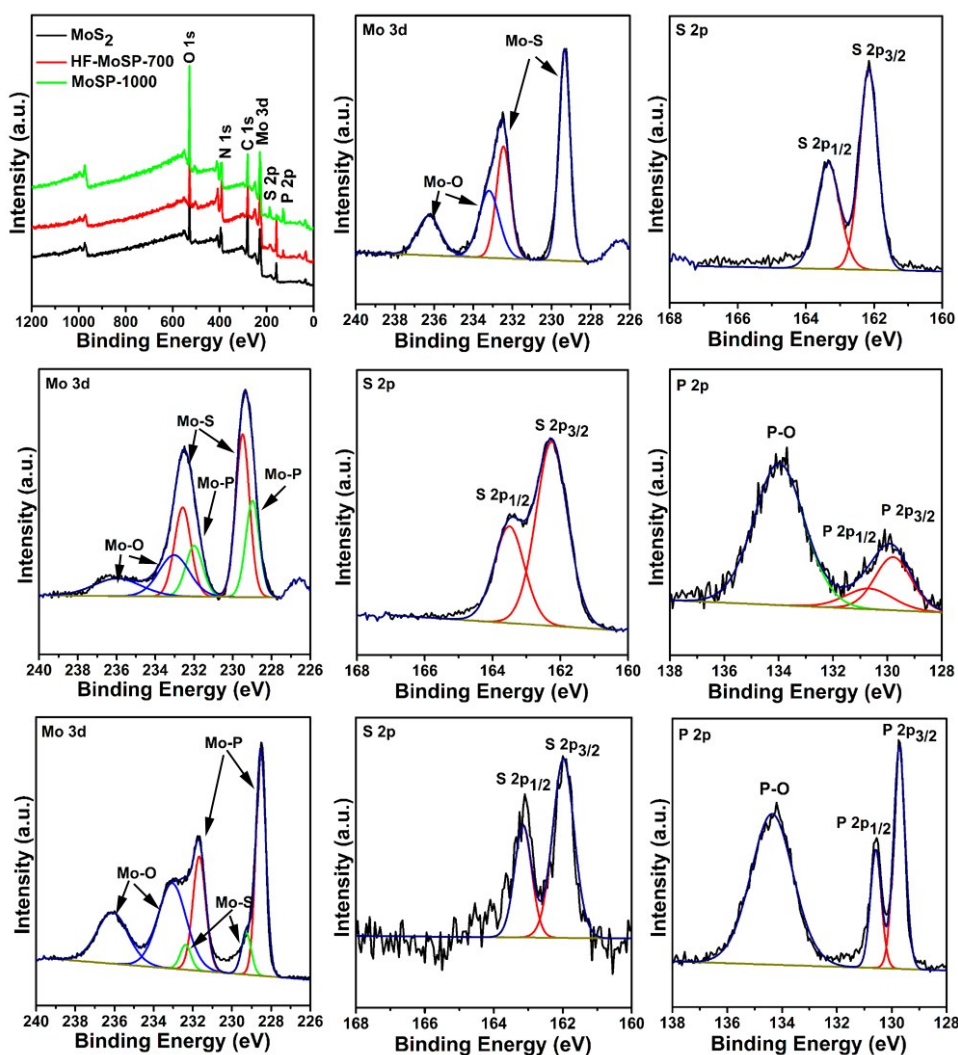


Figure S4. (a) XPS survey spectra of HF-MoS₂, HF-MoSP-700 and MoSP-1000; High resolution XPS spectra of (b) Mo 3d and (c) S 2p for HF-MoS₂ samples; High resolution XPS spectra of (d) Mo 3d and (e) S 2p and (f) P 2p for HF-MoSP-700 samples; High resolution XPS spectra of (g) Mo 3d and (h) S 2p and (i) P 2p for MoSP-1000 samples.

Figure S4a shows the XPS survey spectra of HF-MoS₂, HF-MoSP-700 and

MoSP-1000 samples. The XPS spectrum (Figure S4a) shows the presence of Mo, S elements and no P sign is observed. For the samples from phosphorization (HF-MoSP-700 and MoSP-1000), the obvious sign of P 2p can be detected. The Mo 3d spectra of all samples show the peaks corresponding to Mo-S (232.45 and 229.4eV) and Mo-O (233.1 eV and 236.05 eV). The appearance of Mo-O should due to the slight surface oxidation of samples after the exposure to air. For all samples, the S 2p_{1/2} and 2p_{3/2} of S 2p are located at 163.45 and 162.2 eV, which can be ascribed to the S²⁻ from MoS₂. For HF-MoSP-700 and MoSP-1000, the doublet peaks at about 129.55 eV and 130.4 eV can be indexed to P bonded to Mo in the form of a MoP. The P/S atom ratio is about 0.12, 0.24, 0.53 and 1.96 for HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and MoSP-1000, indicating an enhancement of MoP content with the increase of temperature. For HF-MoS₂, there are no detectable P sign. For MoSP-1000, only very low content of S can be observed. The results further indicate that the phosphorization degree can be well tuned just by tuning the phosphorization temperature.

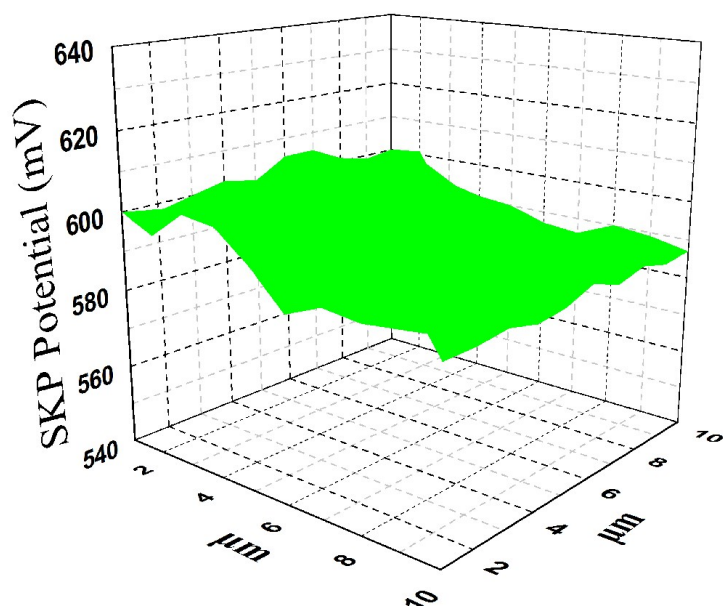


Figure S5. The work function (WF) drawings of MoSP-1000.

Figure S5 shows the work function drawings of MoSP-1000 samples. The work function value of MoSP-1000 is 5.93 eV, which is larger than other samples. Although the higher WF of MoSP-1000 than HF-MoSP-800 and HF-MoSP-900, the MoSP-1000 shows the poor performance for HER than HF-MoSP-800 and HF-MoSP-900, which should be ascribed to the low SBET of MoSP-1000.

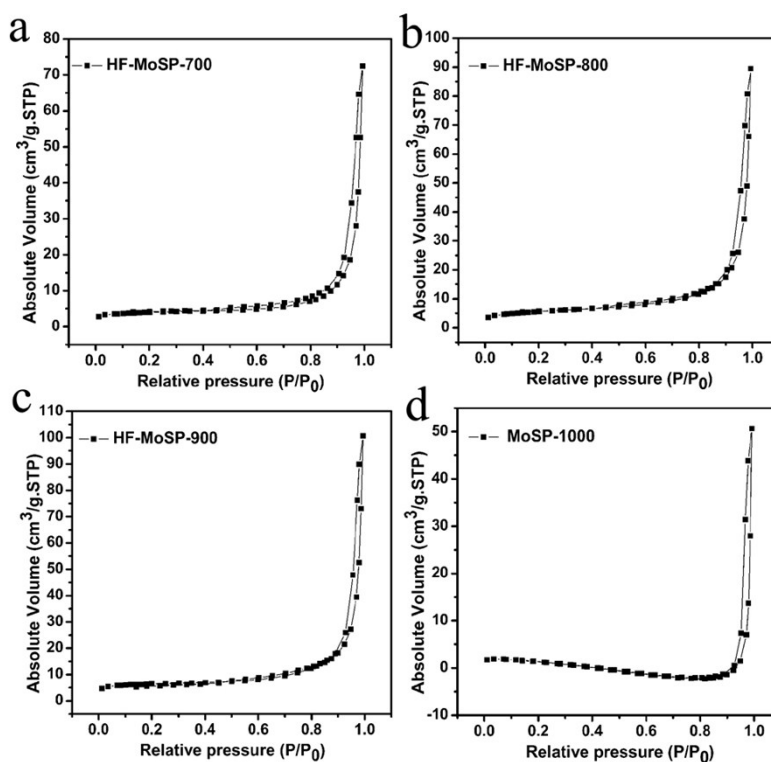


Figure S6. Nitrogen adsorption–desorption isotherm of HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and **MoSP-1000** samples.

The BET specific area of HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and **MoSP-1000** are about 13.9, 19.7, 22.1 and 4.0 m² g⁻¹. The small value of **MoSP-1000** implies the destroying the hierarchical structure, being consistent with TEM and SEM observation.

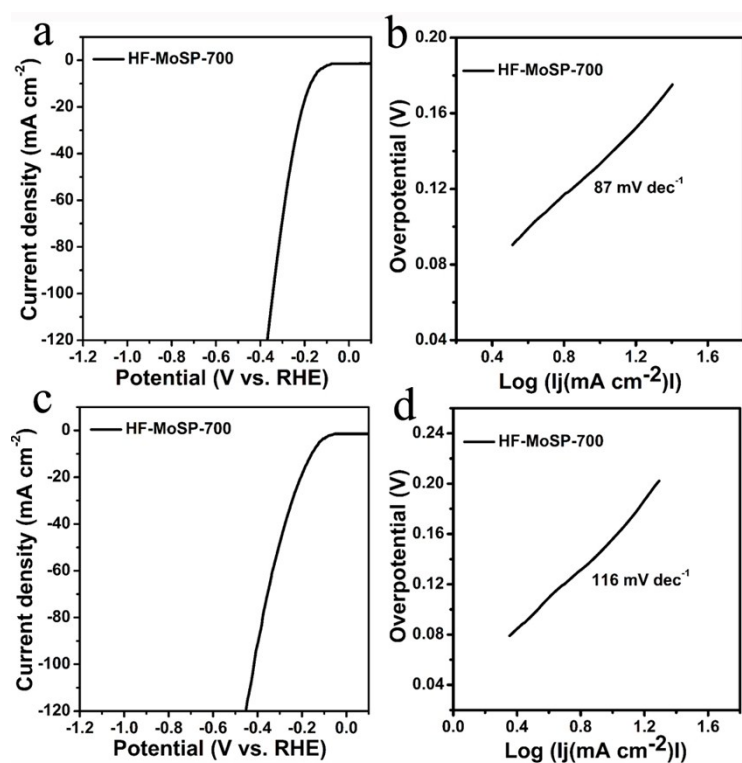


Figure S7. Polarization curves for HF-MoSP-700 in (a) 0.5 M H₂SO₄ and (c) 1M KOH with a scan rate of 5 mV s⁻¹. Tafel plots for HF-MoS-700 in (b) 0.5 M H₂SO₄ and (d) 1M KOH.

Figure S7 shows the polarization curves and onset potentials for HF-MoSP-700 in 0.5 M H₂SO₄ and 1M KOH. The onset potential are about (a) 89 mV in 0.5 M H₂SO₄ and (c) 76 mV in 1M KOH. Tafel plots for HF-MoSP-700 in (b) 0.5 M H₂SO₄ and (d) 1M KOH are about 87mV dec⁻¹ and 116mV dec⁻¹, respectively.

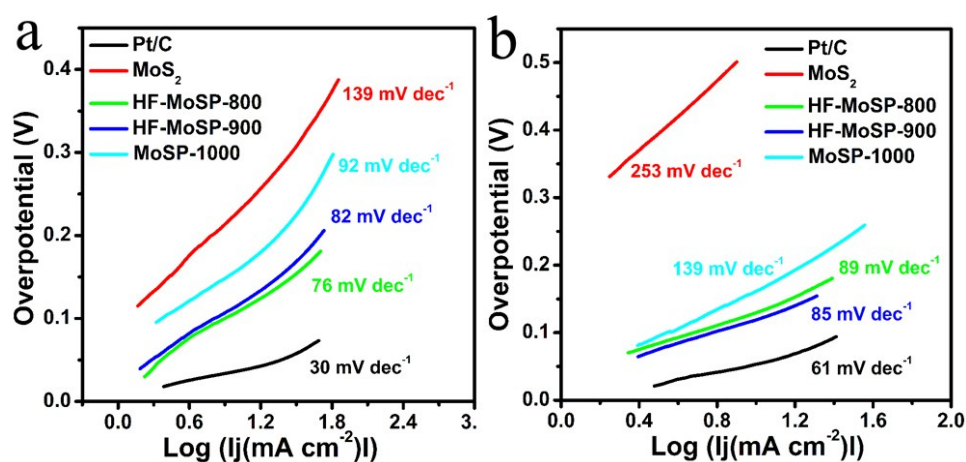


Figure S8. Tafel plots for HF-MoS₂, HF-MoSP-800, HF-MoSP-900 and MoSP-1000 and Pt/C.

The Tafel plots are fitted to the Tafel equation ($\eta = b \log j + a$, where j is the current density and b is the Tafel slope). The Tafel slopes of approximately 30, 139, 76, 82 and 92 mV dec⁻¹ for Pt/C, HF-MoS₂, HF-MoSP-800, HF-MoSP-900 and MoSP-1000 in 0.5M H₂SO₄ (Figure S8a), respectively. In 1M KOH, the Tafel slopes is about 61, 253, 89, 85 and 139 mV dec⁻¹ for Pt/C, HF-MoS₂, HF-MoSP-800, HF-MoSP-900 and MoSP-1000 (Figure S8b), respectively.

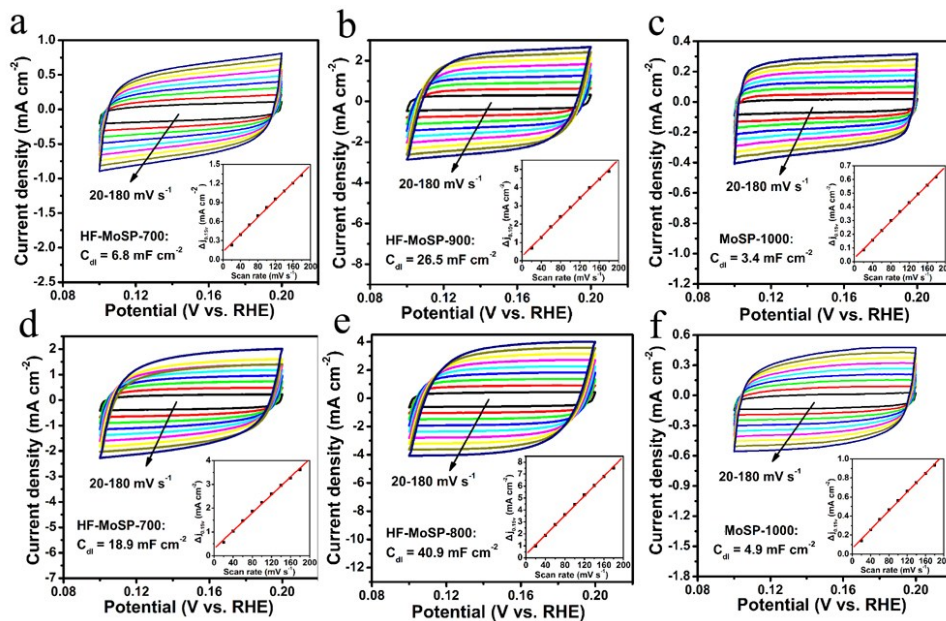


Figure S9. CVs for different samples with different rates from 20 to 180 mV s^{-1} . The inset is the capacitive current at 0.15 V as a function of scan rate ($\Delta j_0 = j_a - j_c$).

The capacitance is 6.8 mF cm^{-2} (Figure S9a), 41.6 mF cm^{-2} (Figure 6b), 26.5 mF cm^{-2} (Figure S9b) and 3.4 mF cm^{-2} (Figure S9c) in 0.5 M H_2SO_4 for HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and MoSP-1000 samples.

In 1M KOH, the capacitance is 18.9 mF cm^{-2} (Figure S9d), 40.9 mF cm^{-2} (Figure S9e), 45.3 mF cm^{-2} (Figure 6e) and 4.9 mF cm^{-2} (Figure S9f) for HF-MoSP-700, HF-MoSP-800, HF-MoSP-900 and MoSP-1000 samples.

We can see that the HF-MoSP-800 shows the highest capacitance in 0.5M H_2SO_4 , and HF-MoSP-900 gives the highest capacitance in 1M KOH, which should be responsible for the good performance of HF-MoSP-800 and HF-MoSP-900 in 0.5M H_2SO_4 and in 1M KOH.

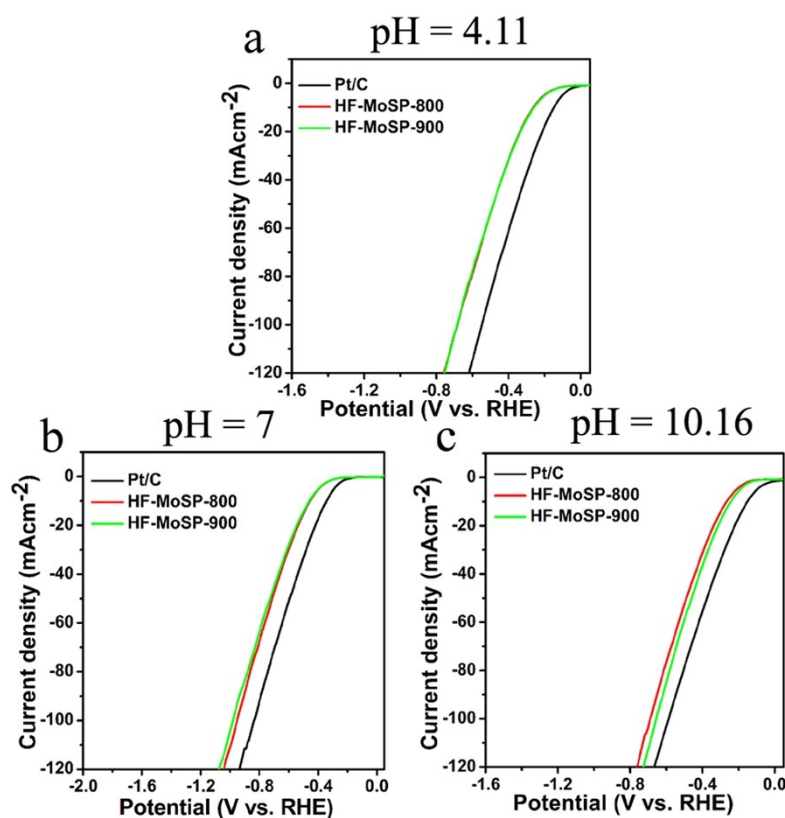


Figure S10. Polarization curves for HF-MoSP-800 and HF-MoSP-900 at different pH: (a) pH=4.11 (weak acid condition), (b) pH=7 (natural conditions) and (c) pH= 10.16 (weak alkaline conditions).

Figure S10 shows the polarization curves for HF-MoSP-800 and HF-MoSP-900 at pH=4.11 (weak acid condition), (b) pH=7 (natural conditions) and (c) pH=10.16 (weak alkaline conditions). The onset potential and η at 10 mA cm⁻² are listed at Table S1. We can see that the samples show the obvious catalytic activity for HER under different pH conditions. By combination of the test at pH=0 and pH=14, we can say that the HF-MoSP catalyst is promising catalyst for HER in all pH range.

Table S1. The performance summary of different catalyst at pH=0 (0.5M H₂SO₄)

Samples	HF-MoS ₂	HF-MoSP-700	HF-MoSP-800	HF-MoSP-900	MoSP-1000
Onset potential (mV)	121	89	29	40	68
η (mV) at 10mAcm ⁻²	228	164	108	116	158
Tafel slope (mV/dec)	139	87	76	82	92

Table S2. Comparison of HER performance of HF-MoSP with other non-noble metal HER electrocatalysts in acid and alkaline condition

Catalyst	Catalyst amount (mg cm ⁻²)	Onset potential (mV)	Overpotential at 10 mA cm ⁻²	Electrolyte solution	Ref.
HF-MoSP	0.35	29	108	0.5 M H ₂ SO ₄	This work
		40	116	1 M KOH	
metallic MoS ₂ nanosheets	-	150	187	0.5 M H ₂ SO ₄	1
MoS ₂ nanosheets	0.285	100	~180	0.5 M H ₂ SO ₄	2
defect-rich MoS ₂	0.285	120	~190	0.5 M H ₂ SO ₄	3
MoS ₂ @OMC	0.3	120	178	0.5 M H ₂ SO ₄	4
MoP-CA2	0.36	40	125	0.5 M H ₂ SO ₄	5
MoP	0.86	50	130	0.5 M H ₂ SO ₄	6
		55	140	1 M KOH	
MoP/Ti	1	-	110	0.5 M H ₂ SO ₄	7
MoP/FTO	0.1	80	150	0.5 M H ₂ SO ₄	8
		~95	190	1 M KOH	
MoP	0.071	100	246	0.5 M H ₂ SO ₄	9
MoP S-CB	0.3	70	150	0.5 M H ₂ SO ₄	10
MoP S-Ti	3	20	64	0.5 M H ₂ SO ₄	11
MoS ₂ (1-x) P _x	0.285	-	120	0.5 M H ₂ SO ₄	12
Mo ₂ C nanotubes	0.8	82	172	0.5 M H ₂ SO ₄	13
		37	112	0.1 M KOH	
MoCx nano-octahedrons	0.8	25	142	0.5 M H ₂ SO ₄	14
		80	151	1 M KOH	
MoDCA-5	0.25	6	78	0.5 M H ₂ SO ₄	15
Mo ₂ C@NC	0.28	< 60	124	0.5 M H ₂ SO ₄	16
		-	60	1 M KOH	
Mo _x C-Ni@NCV	1.1	-	68	0.5 M H ₂ SO ₄	17
		-	126	1 M KOH	
MoB	2.5	> 100	~215	0.5 M H ₂ SO ₄	18
		140	225	1 M KOH	
CoP/CC	0.92	38	67	0.5 M H ₂ SO ₄	19
		80	209	1 M KOH	
CoP/CNT	0.285	40	122	0.5 M H ₂ SO ₄	20
Ni ₅ P ₄ on Nickel foil	-	-	140	0.5 M H ₂ SO ₄	21
		-	150	1 M KOH	
Cu ₃ P NWs/CF	15.2	62	143	0.5 M H ₂ SO ₄	22
Co-NRCNTs	0.28	50	260	0.5 M H ₂ SO ₄	23
			370	1 M KOH	
NiO/Ni-CNT	0.28	-	80	1 M KOH	24
C ₃ N ₄ @NG	0.1	~150	~240	0.5 M H ₂ SO ₄	25
		-	> 600	1 M KOH	
Ni/NiO-NSA	-	34	~110	0.1M KOH	26

From Table S2, we can see that the HF-MoSP give good activity under both acid (pH=0) and alkaline (pH=14) conditions. It has shown enhanced comprehensive performance in comparison with Mo-based catalysts, such as MoS₂, MoP and MoC (Ref. 1, 2, 3, 4, 5, 6, 8, 9, 10, 12, 13, 14), and phosphide (CoP/CNT, Ni₅P₄ and Cu₃P NWs/CF) (Ref. 20, 21 and 22).

The catalytic performance of catalyst can also be affected by the catalyst loading. A high loading would be favorable to improve the performance of the catalysts. We think that the performance of HF-MoSP can be further improved by growth of them on suitable supports with high loading such as carbon cloth and Ti plate etc.

Table S3. The performance summary of different catalyst at pH=14 (1M KOH)

Samples	HF-MoS ₂	HF-MoSP-700	HF-MoSP-800	HF-MoSP-900	MoSP-1000
Onset potential (mV)	227	76	50	42	69
η (mV) at 10mAcm ⁻²	530	156	128	119	161
Tafel slope (mV/dec)	258	116	89	85	139

Table S4. The performance summary of HF-MoSP-800 and HF-MoSP-900 at different pH conditions

pH=4.11

Samples	HF-MoSP-800	HF-MoSP-900
Onset potential (mV)	55	48
η (mV) at 10mAcm ⁻²	263	261

pH=7

Samples	HF-MoSP-800	HF-MoSP-900
Onset potential (mV)	233	242
η (mV) at 10mAcm ⁻²	456	460

pH=10.16

Samples	HF-MoSP-800	HF-MoSP-900
Onset potential (mV)	85	72
η (mV) at 10mAcm ⁻²	266	244

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