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Optimization of the overall water-splitting performance of N, S co-doped carbon-supported NiCoMnS_{x-10} at high current densities by the introduction of sulfur defects and oxygen vacancies†

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Designing an electrocatalyst with excellent performance for both the hydrogen evolution reaction and oxygen evolution reaction in overall water splitting is essential to promote the development of electrolyzed hydrogen production. Defect engineering is a promising method to improve the catalytic performance of target materials. In this study, we used a metal-organic framework as a precursor to establish oxygen-vacancy-rich metal oxide groups supported by a defect-rich carbon framework via calcination. Further, thermal ion exchange was performed to introduce sulfur ions to generate NiCoMnS_{x-10} with both oxygen vacancies and sulfur defects. Due to the synergistic effect between oxygen vacancies and sulfur defects as well as the good morphology, NiCoMnS_{x-10} exhibited good catalytic performance at high current densities. The NiCoMnS_{x-10} cell could achieve a current density of 10 mA cm⁻² at a cell voltage of 1.506 V and a current density of 100 mA cm⁻² at an overpotential of only 1.94 V, which exceeds that of most materials reported to date. Due to the supporting effect of N, S co-doped carbon, the NiCoMnS_{x-10} cell maintained stable performance during a 100 h operation at high current. This work provides an innovative method for the preparation of highly efficient electrocatalysts for overall water splitting.

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Introduction

Continuous attention on global energy and environmental issues has stimulated comprehensive research on efficient, natural, environment friendly, and sustainable alternate energy systems.^{1,2} It has been found that the electrocatalytic overall water splitting to H₂ and O₂ through the hydrogen evolution reaction (HER) and oxygen evolution reaction (OER) is a promising technology to expand the production of clean and renewable energy.³⁻⁵ This technology requires highly efficient HER and OER catalysts to overcome the high

overpotential generated by the two half reactions. Pt and RuO_x are used as traditional benchmark electrocatalysts for the HER and OER, respectively. However, their scarcity and high cost hinder their widespread application. Based on the current situation, the search for an efficient, stable, and noble metal-free bifunctional electrocatalyst is booming in the field of electrocatalytic overall water splitting.⁶⁻⁸

In recent years, transition metal chalcogenide materials have been widely studied for overall water splitting due to their remarkable electroactivity, lower metal-H adsorption bond energy, and higher hydrogen dissociation properties.^{9,10} However, structural problems, including single active sites and excessive order, are great challenges in the practical application of single metal sulfides as they significantly limit the catalytic performance. To solve this problem, researchers have tried to construct polymetallic hybrid sulfides to modify the interface and surface defects of sulfides and adjust their electronic structure.^{11,12} The introduction of a disordered structure into a catalyst may be a prospective way to improve the intrinsic catalytic performance. Oxygen vacancies can cause the redistribution of electrons on the catalyst surface during the catalytic process; this further improves the catalytic performance of the catalysts. Moreover, oxygen

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vacancies can reduce the adsorption barrier of the OH⁻ anions generated during the catalytic reaction and can significantly improve the reactivity of the catalytic active sites;^{13,14} accordingly, the OER activity of the catalyst is substantially improved rather than the HER activity. However, sulfur defects, which are common in transition metal chalcogenide materials, have a significant impact on the HER catalytic activity as they expose other active edges.^{15,16} Herein, we proposed a hypothesis that by simultaneously introducing oxygen vacancies and sulfur defects into a material, the hydrogen and oxygen evolution activities of the material can be excellently adjusted. Studies have shown that the oxide lattice breaks into a disordered distribution after the electronegative sulfur atom is incorporated into the material to produce more defect sites and enhance the active of low coordination oxygen.^{17,18} Therefore, we can choose a multi-metal hybrid oxide as a precursor for thermal ion exchange to introduce sulfur atoms instead of oxygen atoms for the preparation of multi-metal hybrid sulfides with high-density oxygen vacancies and sulfur defects.

In this study, we introduced a metal-organic framework (MOF) material with a specific structure and used its organic part to construct a well-structured carbon skeleton as a support for active materials by carbonization. Furthermore, its metal oxide groups were used to prepare oxygen-vacancy-containing metal oxide precursors. Then, along with the introduction of a sulfur source, thermal ion exchange was performed at high temperatures to prepare a polymetallic hybrid sulfide anchored by a N, S co-doped carbon skeleton. The obtained catalytic material not only retained the original structure of the MOF with N, S co-doped carbon skeleton and high-speed electron transport capability, but also had a high concentration of oxygen vacancies and sulfur defects that would synergistically promote the overall water-splitting reaction. Results shows good catalytic effect of this material, and it remained stable during operation at high current densities for a long time. Our study provides an innovative method for the preparation of highly efficient electrocatalysts.

Experimental

Materials

Cobalt acetate tetrahydrate (C₄H₆CoO₄·4H₂O), manganese acetate tetrahydrate (C₄H₆MnO₄·4H₂O), urea, ammonium fluoride (NH₄F), thioacetamide (TAA), 2-aminoterephthalic acid, and N,N-dimethylformamide (DMF) were supplied by Lantian Co., Ltd. (Nanning, China). Nickel foam (thickness: 1.7 mm, pores per linear inch: 120 ppi) was purchased from Suzhou Jialongde Foam Metals Co., Ltd. (Suzhou, China). All reagents were of analytical grade and directly used without further purification.

Synthesis

Pretreatment of NF. NF (2.0 cm × 3.0 cm × 0.17 cm) was ultrasonicated for 10 minutes with 1 M H₂SO₄, ethanol, and deionized water.

Synthesis of NiCoMn(OH)_x. C₄H₆MnO₄·4H₂O and 0.005 M C₄H₆CoO₄·4H₂O were dissolved in 20 ml deionized water followed by stirring for 20 minutes. After the solution was completely mixed, 100 mg NH₄F and 200 mg urea were added to the abovementioned mixture followed by stirring for another 20 minutes. The resulting mixture was transferred to a 50 ml Teflon-lined autoclave, and the treated NF was added followed by reaction at 120 °C for 12 h. The resulting product was labeled NCM(OH)_x.

Synthesis of NiCoMn(BDC)_x. Typically, 300 mg 2-aminoterephthalic acid was dissolved in 20 ml DMF followed by stirring for 20 minutes. Then, the abovementioned solution was transferred to a 50 ml Teflon-lined autoclave, and NCM(OH)_x/NF was added followed by reaction at 160 °C for 5, 10, 15, and 20 h. The resulting products were labeled NCM(BDC)_{x-5}, NCM(BDC)_{x-10}, NCM(BDC)_{x-15}, and NCM(BDC)_{x-20}.

Synthesis of NiCoMnO_x. The NCM(BDC)_x/NF electrodes prepared at different reaction times in the previous step were put in a quartz boat, and then, the quartz boat was placed in a tube furnace. Under a protective nitrogen atmosphere, the temperature was increased to 400 °C at the heating rate of 5 °C min⁻¹ and then maintained for 3 hours. After the reaction was completed, the reaction mixture was allowed to cool to room temperature and then taken out. The resulting products were labeled NCMO_{x-5}, NCMO_{x-10}, NCMO_{x-15}, and NCMO_{x-20}.

Synthesis of NiCoMnS_x. Herein, 300 mg TAA was dissolved in 20 ml ethanol followed by stirring for 15 minutes, and the resulting solution was transferred to a 50 ml Teflon-lined autoclave; the NCMO_x grown on nickel foam in the previous step was also added to the autoclave followed by reaction at 160 °C for 3 h. The resulting products were labeled NCMS_{x-5}, NCMS_{x-10}, NCMS_{x-15}, and NCMS_{x-20}.

Physicochemical characterizations

The synthesized materials were characterized by various techniques. The crystal structures of the materials were characterized by X-ray diffraction (XRD; Rigaku D/MAX-2500V, Rigaku Corporation) in the 2θ range of 5–90° at the scanning rate of 10° min⁻¹. The degree of graphitization of the prepared electrodes was determined using a Raman spectrometer (Renishaw inVia Reflex) at room temperature with a 532 nm laser and a source spectral range of 1000–2000 cm⁻¹. The samples were subjected to Fourier transform infrared (FTIR) spectroscopy using a Fourier transform infrared spectrometer (Nicolet iS 50, Somerfly, USA) with KBr as a diluent. X-ray photoelectron spectroscopy (XPS) was performed using an X-ray photoelectron spectrometer (ESCALAB 250XI+, Thermo Fisher Scientific) equipped with an Al (single) X-ray source and calibrated against a variable carbon signal (binding energy = 284.8 eV). The element content of the materials was analyzed using an ICP-AES inductively coupled plasma emission spectrometer (model ICPS-7510, Shimadzu

Corporation). Scanning electron microscopy (SEM) images were acquired using an SU8020 (Hitachi High-tech) instrument at an accelerating voltage of 10 kV. Transmission electron microscopy (TEM) images were obtained by a FEI TECNAI G2 F30 instrument. The hydrophilicity of the materials was tested using a contact-angle measuring instrument (model DSA100E, German KYUSS company). Using the OmniFluo-960 fluorescence spectrometer, photoluminescence (PL) spectra were obtained at the excitation wavelengths of 285 nm and 532 nm.

Electrochemical characterization

An electrochemical workstation (CHI 660E, China) with a typical three-electrode system was used for all the electrochemical tests. Herein, 1 M KOH was used as an electrolyte for all the electrochemical tests. The prepared catalyst materials were used as the working electrode, a graphite rod electrode was used as the counter electrode, and saturated $\text{Hg}|\text{HgO}|\text{OH}^-$ was employed as the reference electrode to examine the performance of HER and OER. The prepared electrode materials were simultaneously used as an anode and a cathode, and the performance of overall water splitting was obtained using a two-electrode system. Before the electrochemical tests, the electrolyte was purged with N_2 for 30 min to remove O_2 . For parallel comparison with literature values, the potentials were measured *vs.* $\text{Hg}|\text{HgO}|\text{OH}^-$ and converted to the reversible hydrogen electrode (RHE) scale according to the Nernst equation:

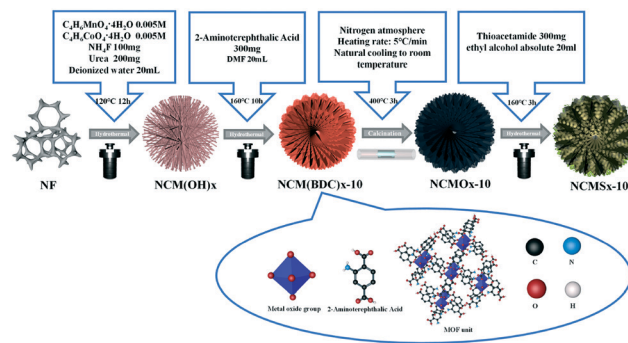
$$E(\text{RHE}) = E(\text{vs. Hg}|\text{HgO}|\text{OH}^-) + 0.059 \text{ pH} + 0.098 \text{ V}$$

For the HER performance, a CV negative scan was performed in the range from -2 to 0 V relative to $\text{Hg}|\text{HgO}|\text{OH}^-$ at the scan rate of 2 mV s⁻¹, and its polarization curve was obtained. The durability was tested at the current density of 100 mA cm⁻² for 100 h. For the OER performance, a CV negative scan was performed in the range from 0 to 1.2 V relative to $\text{Hg}|\text{HgO}|\text{OH}^-$ at the scan rate of 2 mV s⁻¹, and its polarization curve was acquired. The durability was tested at the current density of 100 mA cm⁻² for 100 h. Electrochemical impedance spectroscopy (EIS) was conducted in the frequency range of 100 kHz–0.01 Hz with an AC potential amplitude of 5 mV s⁻¹. The measured impedance data were fitted with an equivalent circuit to obtain the series and charge transfer resistances (two-time-constant parallel (2TP) circuit model using Zsimpwin). All the CV tests were performed with iR compensation without stirring. However, no iR compensation was performed during the chronopotentiometry test.

Results and discussion

Physical properties

As shown in Scheme 1, 2-aminoterephthalic acid acted as an organic ligand, and nickel foam acted as a nickel source;



Scheme 1 Using $\text{NiCoMn}(\text{BDC})_x$ as a precursor, an anion exchange method was used to synthesize highly disordered NCMS_x compounds.

cobalt and manganese salts were added to provide cobalt and manganese for the synthesis of $\text{NCM}(\text{BDC})_x$. Then, $\text{NCM}(\text{BDC})_x$ was carbonized to obtain a well-structured N-doped carbon skeleton-supported NCMO_x with oxygen vacancies. After this, the sulfur source was introduced for the thermal ion-exchange reaction, and N, S co-doped carbon skeleton-anchored NCMS_x rich in oxygen vacancies and sulfur defects was obtained.

As shown in Fig. 1 and S1,† the morphologies of the synthesized materials were characterized by a scanning electron microscope (SEM). It can be observed from Fig. S1a–j† that when the reaction time was 5 h, the resulting material $\text{NCM}(\text{BDC})_{x-5}$ retained the size of $\text{NCM}(\text{OH})_x$ and showed agglomeration. With an increase in the reaction time, dispersed flower-like $\text{NCM}(\text{BDC})_{x-10}$ was obtained. With a further increase in the reaction time, the petals continued to thicken, and $\text{NCM}(\text{BDC})_{x-15}$ and $\text{NCM}(\text{BDC})_{x-20}$ were achieved. Fig. 1a–d show the morphology of the pyrolysis products NCMO_x ; it can be observed that the pyrolysis products maintained the basic morphology of the original MOF; however, the petals became thinner. Among them, NCMO_{x-10} exhibited well-connected three-dimensional flower-like sheets, which are conducive for a full contact between the active sites and the electrolyte and the conduction of electrons inside the material.

Fig. 1e–h show the morphology of the product obtained after vulcanization. Many thin, transparent, and small-sized sheets were noticed on the petals after vulcanization. NCMS_{x-5} still showed an aggregated form (Fig. 1e), which was not conducive for the participation of deep active sites in the reaction. For NCMS_{x-15} and NCMS_{x-20} (Fig. 1g and h), as the petals of their precursors were too thick, only a few thin slices could be generated on the surface of these petals, and more sites were wrapped in them. The difference is that under the advantages of the precursor's just dispersion and proper thickness, NCMS_{x-10} (Fig. 1f) shows a uniform state of small sheets intertwined after vulcanization. This morphology significantly increases the contact area between the material and the electrolyte, leading to the participation of more active sites in the reaction. In addition, under the combined effect of the high winding degree of the sheet

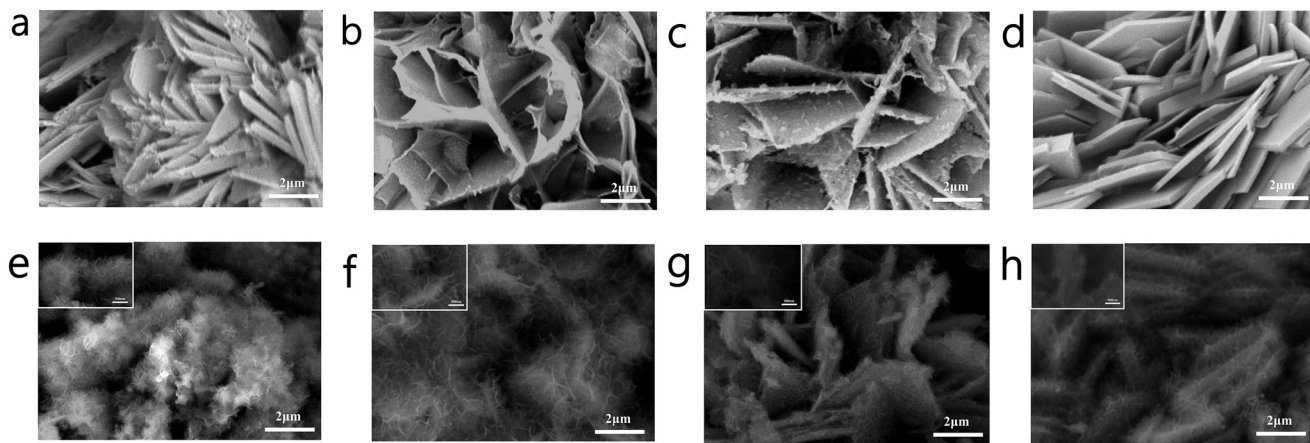


Fig. 1 SEM images of (a) NCMO_{x-5} ; (b) NCMO_{x-10} ; (c) NCMO_{x-15} ; (d) NCMO_{x-20} ; (e) NCMS_{x-5} ; (f) NCMS_{x-10} ; (g) NCMS_{x-15} ; and (h) NCMS_{x-20} .

structure and the carbon skeleton, the electron transfer within the material will be more convenient and fast.

In order to characterize the synthesis of the materials, XRD tests were performed on the materials, and the results are shown in Fig. S2a[†] and 2a–b. Fig. S2a[†] shows that the diffraction peaks in the XRD pattern of $\text{NCM}(\text{BDC})_x$ basically correspond to $\text{Ni}(\text{BDC})$, $\text{Co}(\text{BDC})$, and $\text{Mn}(\text{BDC})$. However, due to the interaction of different metal ions, a small shift was observed in some peaks of the polymetallic MOF as compared to the case of the monometallic MOF, which further verifies that the MOF synthesized herein is

polymetallic rather than a simple mixture of the three monometallic MOFs. As can be noticed in Fig. 2a and b, the characteristic diffraction peaks attributed to the MOF were retained in the patterns of both the oxide and sulfide; this indicates that even after pyrolysis and vulcanization, the characteristics of $\text{NCM}(\text{BDC})_x$ were retained. In addition, the products achieved after pyrolysis, as shown in Fig. 2a, were a mixture of NiO , CoO , and Mn_3O_4 ; this proves the successful synthesis of oxide precursors. Similarly, as shown in Fig. 2b, after the thermal ion-exchange reaction, the sulfur atoms successfully replaced the oxygen atoms. The products obtained were a mixture of Ni_3S_2 , Co_3S_4 , and MnS , which further proved the successful synthesis of NCMS_x .

In order to confirm the synthesis of MOFs and to characterize the functional groups in the precursor materials, FTIR was performed on $\text{NCM}(\text{BDC})_x$, and the FTIR spectra are shown in Fig. 2c. The peak observed at 1380 cm^{-1} corresponds to the symmetric tensile vibration of the coordinated carboxylate group, indicating the successful coordination of Co^{2+} ions with BDC ligands, proving the successful synthesis of $\text{Co}(\text{BDC})$.^{19,20} Furthermore, a peak corresponding to $\text{Ni}(\text{BDC})$ was observed at 508 cm^{-1} .²¹ In addition, the spectrum displayed a peak at 748 cm^{-1} , which is attributed to $\text{Mn}(\text{BDC})$.²² The peaks related to the functional groups of organic ligands were also obtained in the spectra, in which the two bands at 1032 and 808 cm^{-1} are attributed to the C–H tensile band.²³ Moreover, two obvious peaks were found in the range of 1410 – 1260 cm^{-1} , indicating the presence of $-\text{OH}$. The band at $1450 \pm 20\text{ cm}^{-1}$ is a sign of $\text{C}=\text{C}$ tensile vibration in the presence of an aromatic carbon network. The peak of the combination of N–H in-plane bending and C–N stretching was found at 1548 cm^{-1} , demonstrating the retention of amino groups in the ligand. The analysis of the FTIR spectra further proves the successful synthesis of the MOF precursors.

The concentration of surface defects has a non-negligible effect on the performance of the material. In order to characterize the defects of the materials and prove the high disorder of the materials, Raman, PL, and EPR spectroscopy

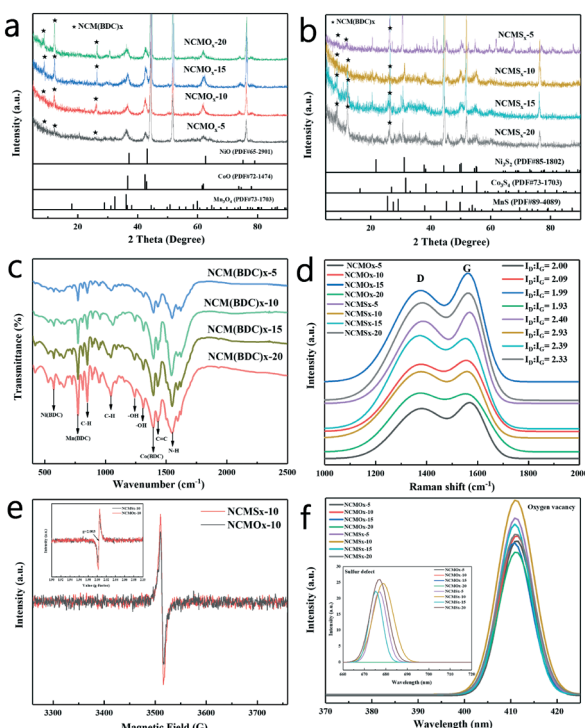


Fig. 2 (a) XRD patterns of NCMO_x ; (b) XRD patterns of NCMS_x ; (c) FTIR spectra of $\text{NCM}(\text{BDC})_x$; (d) Raman spectra of NCMO_x and NCMS_x ; (e) EPR spectra of NCMO_{x-10} and NCMS_{x-10} ; and (f) PL spectra of NCMO_x and NCMS_x .

tests were performed on the synthesized NCMO_x and NCMS_x . It can be noticed from the Raman spectrum shown in Fig. 2d that a defect-rich carbon skeleton with $I_D:I_G > 1$ was successfully prepared after calcination. Among them, the carbon defect degree of NCMO_{x-10} was greater than that of the other three oxides. After vulcanization, due to the doping effect of sulfur atoms, the defect degree of the carbon skeleton was further increased; this implies the successful synthesis of N, S co-doped carbon. It can also be observed that the defect degree of NCMS_{x-10} is slightly larger than those of the other materials.

In addition, as shown in Fig. 2e, the EPR curves prove that there are oxygen vacancies in NCMO_{x-10} , whereas there may be both oxygen vacancies and sulfur defects in NCMS_{x-10} .²⁴⁻²⁷ It can be verified that the synthesized NCMO_{x-10} and NCMS_{x-10} are rich in defects and show a high disorder. After this, the specific defects of the materials were characterized by PL spectroscopy. As can be observed from the small graph shown in Fig. 2f, at an excitation wavelength of 532 nm, the spectrum of NCMS_x showed a peak at 680 nm that was attributed to the S defects.²⁸⁻³⁰ However, the spectrum of NCMO_x does not have this peak, indicating that the NCMS_x compounds containing sulfur defects were successfully prepared. However, at the excitation wavelength of 285 nm, the spectra of all the NCMO_x and NCMS_x compounds showed a peak at 410 nm corresponding to oxygen vacancies.^{31,32} Interestingly, the peak intensity of the vulcanized product was higher than that of the oxide precursor; this shows that the concentration of the oxygen vacancies increases after vulcanization. This further verifies that after the introduction of negatively charged sulfur, the oxide lattice is decomposed into a disordered arrangement, thereby generating more defect sites and enhancing the active of low coordination oxygen. These characteristics of the materials rich in oxygen vacancies and sulfur defects are very beneficial for achieving efficient overall water splitting.

XPS was performed to explore the elemental characteristics of the material, and the results are shown in Fig. 3. As shown in Fig. 3a, the peaks at 284.5, 285.1, and 288.3 eV correspond to the C-C, C=C, and C=O bonds.^{33,34} Furthermore, the binding energy peaks at 284.1 and 285.5 eV can be assigned to C-S and C-N, respectively.^{33,35} This proves the successful synthesis of the N-doped carbon skeleton in NCMO_{x-10} and the successful synthesis of N, S co-doped carbon in NCMS_{x-10} . The peaks corresponding to pyridinic N (398.3 eV), pyrrolic N (399.4 eV), and graphitic N (400.7 eV) were found in the high-resolution N 1s spectrum (Fig. 3b),³⁵⁻³⁷ and the presence of the graphitic N peak further proves that N was doped into the carbon skeleton. We found that pyridinic N and pyrrolic N coexist in NCMS_{x-10} , whereas only pyridinic N exists in NCMO_{x-10} . The presence of pyrrolic N and pyridinic N proves that the material can exhibit improved electrocatalytic activity by interacting with H^+ for the HER or bind to OER intermediates (O^* , OOH^* , or OH^*) under appropriate force.^{38,39} Therefore, the performance of the material having both pyrrolic N and pyridinic N will be more excellent. Thus, it is inferred that the electrocatalytic

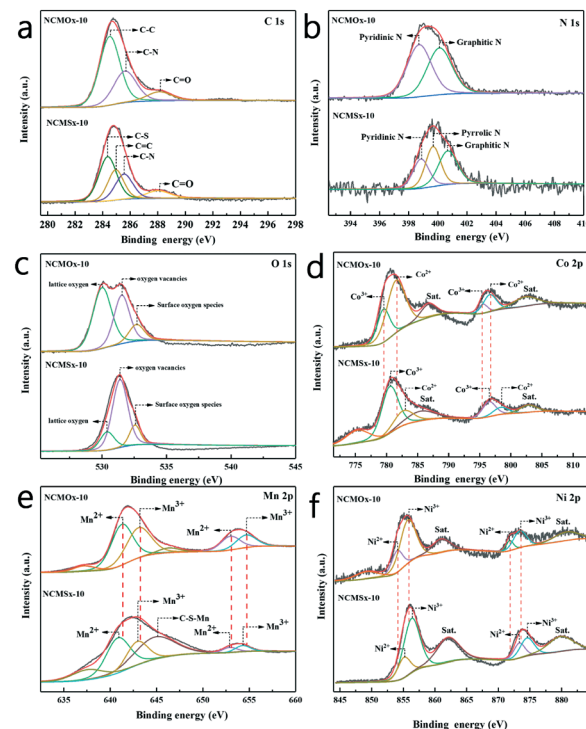


Fig. 3 (a-f) C 1s, N 1s, O 1s, Co 2p, Mn 2p, and Ni 2p spectra of NCMO_{x-10} and NCMS_{x-10} .

performance of NCMS_{x-10} will be better than that of NCMO_{x-10} . In addition, there are peaks at 531.4 eV, 530.2 eV, and 532.2 eV, attributed to lattice oxygen, oxygen vacancies, and surface oxygen adsorption, in the O 1s spectrum (Fig. 3c) of the materials.^{40,41} The analysis of the O 1s spectrum shows that there are oxygen vacancies in both NCMO_{x-10} and NCMS_{x-10} . From the peak areas, it can be analyzed that the concentration of oxygen vacancies in NCMS_{x-10} is even higher than that in NCMO_{x-10} , and this conclusion agrees well with the PL results.

In the Co 2p spectrum (Fig. 3d), the peaks at 779.5 and 795.1 eV are attributed to Co^{3+} ($2p_{3/2}$) and Co^{3+} ($2p_{1/2}$), respectively, and the two peaks at 781.3 and 795.9 eV correspond to Co^{2+} ($2p_{3/2}$) and Co^{2+} ($2p_{1/2}$). The two peaks obtained at 787.1 and 802.6 eV are satellite peaks.^{31,32,42} As shown in Fig. 3e, the two peaks at 641.0 and 652.5 eV can be assigned to Mn^{2+} , and the other two peaks at 642.2 and 653.7 eV can be attributed to Mn^{3+} .^{41,43} In addition, the peak at 644.3 eV can be assigned to C-S-Mn.^{44,45} The Ni 2p spectrum shown in Fig. 3f was analyzed, and the fitted peaks at 854.0 and 871.80 eV were attributed to the Ni^{2+} ions, whereas the fitted peaks at 855.70 and 873.40 eV were assigned to the Ni^{3+} ions.^{46,47} Via a comparative analysis of the XPS spectra of NCMO_{x-10} and NCMS_{x-10} , we found a regular phenomenon. When the sulfur atoms replace the oxygen atoms, the XPS peaks of different metals have different degrees of displacement; this shows that with the addition of sulfur atoms, a strong charge transfer occurs inside the material.

The morphology and lattice conditions of the materials were observed using a projection electron microscope, as shown in Fig. 4. As shown in Fig. 4a, NCMS_x exhibits a cross-linked film shape, which can provide more channels for electron transfer. In the HRTEM image (Fig. 4b), lattice fringes belonging to the MnS (201) crystal plane and Ni₃S₂ (210) crystal plane can be observed, and there are some parts without lattice spacing, which is due to the presence of N, S co-doped carbon. Moreover, the lattice fringes in Fig. 4c can be assigned to Co₃S₄(311), Ni₃S₂(200), and MnS(110). It can be observed from the HRTEM images (Fig. 4b and c) that the lattice fringes of NCMO_{x-10} are highly disordered, and this high disorder is also closely related to the defect-rich features of NCMO_{x-10}. In addition, the morphology of the other NCMS_x compounds was examined. It can be noticed that the NCMS_{x-5} (Fig. 4d) film is relatively dispersed and flat, and the electron transfer is limited by the number of channels. Moreover, NCMS_{x-15} and NCMS_{x-20} (Fig. 4e and f) exhibit aggregation phenomena, which is not conducive for the contact between the reaction site and the electrolyte, respectively. On the other hand, the morphology of NCMS_{x-10} is helpful for the electrocatalytic reaction.

In addition, the atomic ratios of Co and Mn in NCM(BDC)_x prepared at different reaction times were analyzed by the ICP-OES analysis (the proportion of the Ni element could not be accurately estimated due to the influence of the nickel substrate), and the results are shown in Table S1.[†] The atomic ratios of Co and Mn in NCM(BDC)_{x-5}, NCM(BDC)_{x-10}, NCM(BDC)_{x-15}, and NCM(BDC)_{x-20} are 1:0.88, 1:0.99, 1:1.01, and 1:0.82; it can be observed that the atomic ratios of Co and Mn in different

materials are close to 1. In addition, the hydrophilicity of different NCMS_x was investigated. As shown in Fig. S3,[†] the contact angle of NCMS_{x-10} is the smallest and its hydrophilicity is the best, which is most conducive for the full contact of the material with the electrolyte. This feature is also one of the reasons for the best catalytic performance of NCMS_{x-10}.

HER performance

Using a typical three-electrode system in 1.0 M KOH with the obtained materials as the working electrodes, the electrocatalytic performance of these materials for the HER was evaluated at a scan rate of 2 mV s⁻¹. In order to verify the applicability of these materials in the industry, we compared the performances of the materials at high current densities. It can be observed from Fig. 5a and Table S4[†] that when the current density was 10 mA cm⁻², the overpotential of NCMS_{x-10} was only -105 mV, which was far lower than those of other comparative materials. Similarly, when the current density reached the industrial application standard of 100 mA cm⁻², the required overpotential of NCMS_{x-10} (-330 mV) was substantially smaller than those of NCMO_{x-5} (-429 mV), NCMO_{x-10} (-401 mV), NCMO_{x-15} (-423 mV), NCMO_{x-20} (-448 mV), NCMS_{x-5} (-352 mV), NCMS_{x-15} (-365 mV), and NCMS_{x-20} (-372 mV). This performance gap is mainly because the well-crosslinked film morphology of the material provides a fast electron channel, and the rich sulfur defects are also beneficial for the HER.

Fig. 5b shows the EIS curves of the materials and the corresponding fitted circuit diagrams, and the EIS-related data of the materials is provided in Table S2.[†] From these data, we can observe that the R_{ct} (charge transfer resistance) of NCMS_{x-10} (37.37 Ω) is significantly smaller than those of NCMO_{x-5} (168.10 Ω), NCMO_{x-10} (129.80 Ω), NCMO_{x-15}

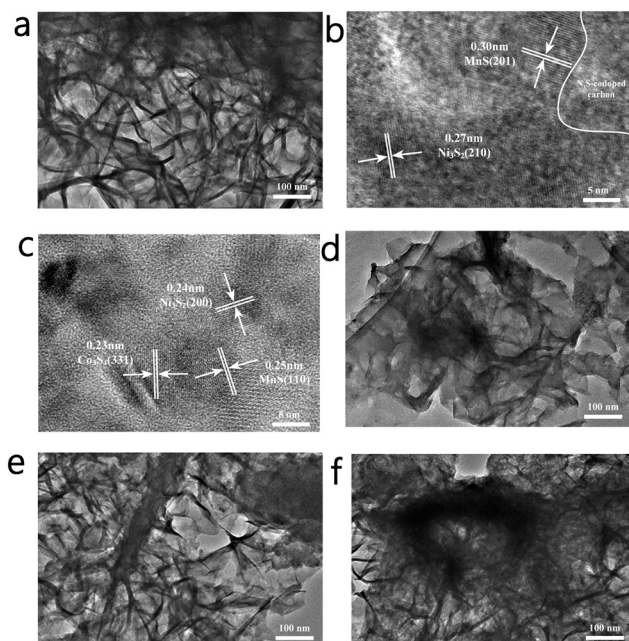


Fig. 4 (a) TEM image of NCMS_{x-10}; (b and c) HRTEM images of NCMS_{x-10}; (d) TEM image of NCMS_{x-5}; (e) TEM image of NCMS_{x-15}; and (f) TEM image of NCMS_{x-20}.

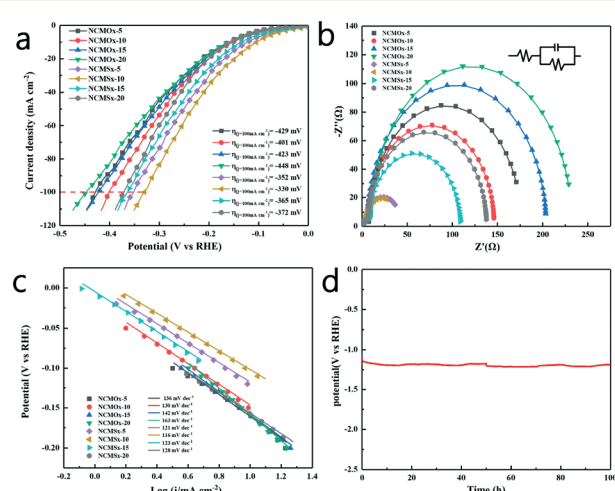


Fig. 5 (a) HER polarization curves (iR-corrected) obtained at a scan rate of 2 mV s⁻¹; (b) HER electrical impedance maps; (c) the Tafel slopes of the materials; and (d) results of the 100 h chronopotentiometry test of NCMS_x under HER.

(195.80 Ω), NCMO_{x-20} (222.1 Ω), NCMS_{x-5} (38.07 Ω), NCMS_{x-15} (99.24 Ω), and NCMS_{x-20} (129.20 Ω). This is due to the presence of the N, S co-doped carbon framework and the well-crosslinked morphology of the material. Fig. 5c presents the Tafel slopes of these electrodes. Compared with NCMO_{x-5} (136 mV dec^{-1}), NCMO_{x-10} (130 mV dec^{-1}), NCMO_{x-15} (142 mV dec^{-1}), NCMO_{x-20} (163 mV dec^{-1}), NCMS_{x-5} (121 mV dec^{-1}), NCMS_{x-15} (123 mV dec^{-1}), and NCMS_{x-20} (128 mV dec^{-1}), NCMS_{x-10} has the smallest Tafel slope (116 mV dec^{-1}). This indicates that the rate of apparent hydrogen formation rapidly increases with an increase in the applied potential. In order to explore the stability of the material at high current densities, NCMS_{x-10} was subjected to chronopotentiometry for 100 hours under the HER reaction. Fig. 5d shows that the change in the overpotential was negligible after 100 h, indicating that NCMS_{x-10} has good stability.

OER performance

Similarly, the OER performance of the catalyst was evaluated at a scan rate of 2 mV s^{-1} in a 1.0 M KOH solution. Consistent with the results of HER, NCMS_{x-10} exhibited better performance than other materials due to the synergistic effect of good morphology, more abundant defects, and N, S co-doped carbon. As shown in Fig. 6a and Table S4,† NCMS_{x-10} requires an overpotential of only 226 mV to achieve a current density of 10 mA cm^{-2} . Moreover, the current density could reach 100 mA cm^{-2} at a voltage of 417 mV, which was better than those of NCMO_{x-5} (564 mV), NCMO_{x-10} (557 mV), NCMO_{x-15} (568 mV), NCMO_{x-20} (579 mV), NCMS_{x-5} (440 mV), NCMS_{x-15} (453 mV), and NCMS_{x-20} (472 mV).

Fig. 6b shows the EIS curves of the materials and the corresponding fitted circuit diagrams, and the EIS-related data of the materials is provided in Table S3.† It can be noticed from the comparison that the R_{ct} (74.73 Ω) of

NCMS_{x-10} is smaller than those of NCMO_{x-5} (179.11 Ω), NCMO_{x-10} (142.70 Ω), NCMO_{x-15} (199.92 Ω), NCMO_{x-20} (243.84 Ω), NCMS_{x-5} (123.21 Ω), NCMS_{x-15} (124.22 Ω), and NCMS_{x-20} (127.80 Ω); this shows that NCMS_{x-10} has ultra-fast charge transfer capability and can lead to a higher reaction rate. It can be noticed in Fig. 6c that NCMS_{x-10} has a Tafel slope of 107 mV dec^{-1} , which is lower than those of NCMO_{x-5} (160 mV dec^{-1}), NCMO_{x-10} (139 mV dec^{-1}), NCMO_{x-15} (151 mV dec^{-1}), NCMO_{x-20} (165 mV dec^{-1}), NCMS_{x-5} (115 mV dec^{-1}), NCMS_{x-15} (118 mV dec^{-1}), and NCMS_{x-20} (128 mV dec^{-1}). This indicates that NCMS_{x-10} is kinetically advantageous for the OER. In order to explore the stability of NCMS_{x-10} at high current densities in the OER, a 100 h chronopotentiometry test was performed at a current density of 100 mA cm^{-2} , and the overpotential curve showed a slight change (Fig. 6d), confirming the stability of NCMS_{x-10} in the OER.

Overall water-splitting performance

By conducting the HER and OER using the proposed materials, we determined the feasibility of using NCMS_{x-10} as a bifunctional catalyst. In order to explore its overall water-splitting performance, we simultaneously used NCMS_{x-10} as a positive electrode and negative electrode in an alkaline electrolysis cell at a scan rate of 2 mV s^{-1} and conducted the overall water-splitting reaction. Fig. 7a and Table S4† show that the NCMS_{x-10} cell can achieve the current density of 10 mA cm^{-2} at the cell voltage of 1.506 V, obtaining a current density of 100 mA cm^{-2} at an overpotential of only 1.94 V. It can also be observed in Fig. 7a that when the current density is 100 mA cm^{-2} , the NCMS_{x-10} cell exhibits the smallest voltage value when compared with the NCMO_{x-5} cell (2.24 V), NCMO_{x-10} cell (2.23 V), NCMO_{x-15} cell (2.23 V), NCMO_{x-20} cell (2.32 V), NCMS_{x-5} cell (2.10 V), NCMS_{x-15} cell (2.05 V), and

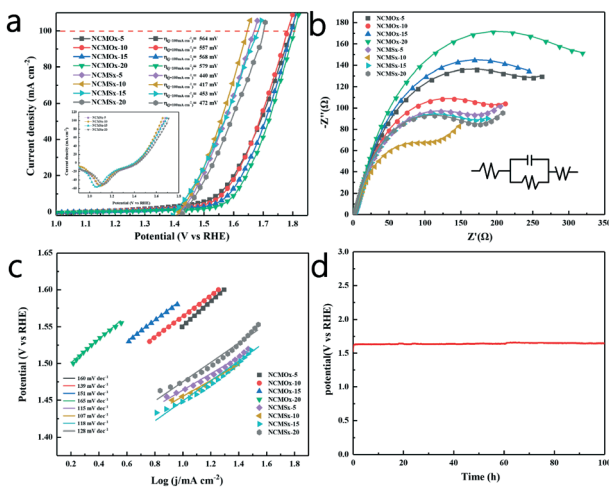


Fig. 6 (a) OER polarization curves (iR-corrected) at a scan rate of 2 mV s^{-1} ; (b) OER electrical impedance maps; (c) the Tafel slopes of the materials; and (d) results of the 100 h chronopotentiometry test of NCMS_{x-10} under the OER.

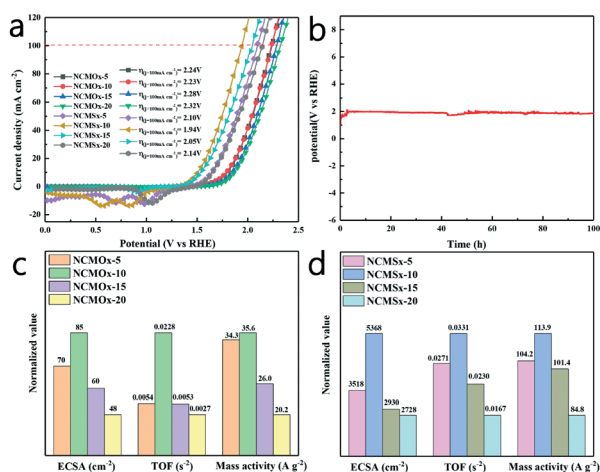


Fig. 7 (a) Polarization curves (iR-corrected) in the two-electrode configuration; (b) chronopotentiometry test of NCMS_{x-10} under overall water splitting; (c) comparison between the activities of different NCMO_x materials in 1.0 M KOH; and (d) comparison between the activities of different NCMS_x materials in 1.0 M KOH.

NCMS_{x-20} cell (2.14 V). Upon further comparing NCMS_{x-10} with the current noble-metal-free catalysts (Table S5†), it was found that NCMS_{x-10} has the best overall water-splitting catalytic performance. Thus, NCMS_{x-10} anchored by N, S co-doped carbon and rich in oxygen vacancies and sulfur defects exhibits excellent performance in catalyzing the overall water-splitting reaction. As shown in Fig. 7b, a 100 h chronopotentiometry test was performed at a current density of 100 mA cm⁻², and the overpotential curve showed a slight change, confirming the stability of NCMS_{x-10} in the overall water-splitting reaction.

In order to more comprehensively evaluate and compare the performances of these materials in overall water splitting, different electrochemical parameters were evaluated using different electrochemical parameter calculation methods (supporting materials). As shown in Fig. 7c and d, the mass activity, ECSA (electrochemically active surface area), and TOF (turn over frequency) of NCMO_x and NCMS_x were investigated to evaluate the properties of these materials. We calculated the double-layer capacitance (C_{dl}) of these materials based on their CV curves (Fig. S4†) and then evaluated the ECSA of these materials to determine their intrinsic activity (Fig. S5†). As shown in Fig. 7c and d, the ECSA of NCMS_{x-10} is the largest (5368 cm⁻²). The ECSAs of other materials are as follows: NCMO_{x-5} (70 cm⁻²), NCMO_{x-10} (85 cm⁻²), NCMO_{x-15} (60 cm⁻²), NCMO_{x-20} (48 cm⁻²), NCMS_{x-5} (3518 cm⁻²), NCMS_{x-15} (2930 cm⁻²), and NCMS_{x-20} (2728 cm⁻²). Moreover, *via* the values of mass activity, the performances of the electrocatalysts can be determined. The values of mass activity are as follows: NCMS_{x-10} (113.9 A g⁻¹), NCMO_{x-5} (34.3 A g⁻¹), NCMO_{x-10} (35.6 A g⁻¹), NCMO_{x-15} (26.0 A g⁻¹), NCMO_{x-20} (20.2 A g⁻¹), NCMS_{x-5} (104.2 A g⁻¹), NCMS_{x-15} (101.4 A g⁻¹), and NCMS_{x-20} (84.8 A g⁻¹). The ECSA and mass activity results are consistent with those obtained *via* polarization curves. In addition, the TOF of NCMS_{x-10} is also the largest, which reached 0.0331 s⁻¹, whereas the TOFs of NCMO_{x-5}, NCMO_{x-10}, NCMO_{x-15}, NCMO_{x-20}, NCMS_{x-5}, NCMS_{x-15}, and NCMS_{x-20} are 0.0054 s⁻¹, 0.0228 s⁻¹, 0.0053 s⁻¹, 0.0027 s⁻¹, 0.0271 s⁻¹, 0.0230 s⁻¹, and 0.0167 s⁻¹, respectively. Based on these parameters, we can conclude that NCMS_{x-10} has better performance than other materials in all aspects due to the synergistic effect of good morphology, abundant oxygen vacancies, and sulfur defects.

Discussion of stability

In order to verify the stability and explore the change in the valence band of the material after different reaction processes to infer the possible reaction active sites, we performed XRD, PL, SEM, and XPS characterizations of the material after 100 hours of OER and HER chronopotentiometry test. It can be observed that there is no obvious change in the XRD pattern of the material (Fig. S6a†) after the reactions; this shows that neither the OER nor the HER affected the crystal structure of the material; this further illustrates the stability of the material. In addition, it can be

noticed in Fig. S6b and c† that the intensity of the PL peaks of sulfur defects and oxygen vacancies slightly increased after the reaction; this may be caused by the leaching of a small amount of sulfide ions and oxidation reactions during the test. However, this change is so tiny that it can even be ignored. This indicates that the content of oxygen vacancies and sulfur defects in the material is basically maintained before and after the reaction, which further illustrates the stability of the material.

The XPS spectrum (Fig. S7†) shows the peaks of the elements with relatively obvious changes. It can be observed that compared with those before the reaction, the valence state of the metal elements and the ratio of different valence states had no obvious changes after the HER, and no new substances were produced; however, there are different degrees of deviation. This shows that the material could still maintain a relatively stable state after 100 hours of high-current hydrogen evolution reaction. After the OER, although the valence state of the metal element did not significantly change, the valence state ratio of the ion changed when compared with that before the reaction. Furthermore, the peak of CoO was observed in the Co 2p spectrum, which indicates that during the OER, the surface of the material was partially oxidized, and metal oxides with strong oxygen evolution properties were generated. This change could promote the OER.

Via the analysis of the S 2p spectrum, it can be found that the noise level of the S 2p spectrum increased after the HER and OER; this indicates that the leaching of the S element occurred during these two half reactions, and the leaching of S during the OER was most obvious. The stability test results show that the OER performance (Fig. 6d) is very stable, indicating that the leaching of sulfur ions does not affect the progress of the OER. Although there is leaching of only a small amount of S ions during the HER (Fig. 5d), the stability of the HER performance still significantly fluctuated. This shows that the S ion has a significant effect on the HER performance, whereas the effect on the OER performance is negligible.

As shown in Fig. S8a,† NCMS_{x-10} slightly aggregated after the HER; however, the basic morphology of NCMS_{x-10} was retained. Moreover, the SEM image (Fig. S8b†) obtained after the OER showed that the flower-like structure of NCMS_{x-10} was destroyed, the pore size of the material became larger, and the interlacing degree of the sheet structure reduced. By combining the analysis results of PL and XPS acquired after the reaction, it can be inferred that this phenomenon may be due to the leaching of the S element and the formation of an oxide film during the OER.

Conclusions

In summary, herein, we used an MOF material as a precursor and performed calcination and thermal ion exchange to successfully prepare N, S co-doped carbon framework-anchored NCMS_{x-10} with abundant oxygen vacancies and

sulfur defects. The oxygen vacancies and sulfur defects contributed to the improvement of the OER and HER performances, respectively. Due to the synergistic effect of oxygen vacancies and sulfur defects and morphology control, the material showed excellent performance. The NCMS_{x-10} cell could achieve a current density of 100 mA cm^{-2} at an overpotential of only 1.94 V. Moreover, at the high current density of 100 mA cm^{-2} , this cell could operate stably for 100 h, indicating that NCMS_{x-10} is an excellent electrocatalytic material. This study provides an available solution for further promoting the development of non-noble metal catalysts.

Conflicts of interest

There are no conflicts to declare.

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