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Regulating the coordination environment of a metal–organic framework for an efficient electrocatalytic oxygen evolution reaction†

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The oxygen evolution reaction (OER) plays a vital role in fuel cells, water splitting and metal–air batteries. Efficient electrocatalysts could overcome the higher overpotential of the OER, improve electron transfer efficiency, and promote water decomposition. In this work, a novel metal–organic framework (MOF) with efficient OER electrocatalytic performance (defined as $FeCo-L₁L₂$) was successfully prepared by a free assembly of metal ions (Fe and Co), 2,5-dihydroxyterephthalic acid (defined as L_1) and 4,6dihydroxyisophthalic acid (defined as L_2). The overpotential was only 283 mV at the current density conditions of 10 mA cm^{-2} with a Tafel slope of 31.6 mV dec⁻¹. Its excellent OER performance is attributed to the synergistic effect of the bimetals of FeCo-L₁L₂ and the coordination environment optimization created by the dual ligands. This work not only improved the catalytic performance of MOFs in the OER but also proposed a new strategy for the structural design of MOFs. PAPER
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Introduction

With the rapid increase in environmental pollution and energy crisis, it is essential to advance eco-friendly and efficient energy storage and conversion systems. Various electrochemical devices, including fuel cells, water electrolyzers and metal–air batteries, have achieved considerable attention because of their environmental friendliness and high energy conversion efficiency. Hydrogen energy is considered to be the most efficient green energy with development potential in the future. However, large-scale production and storage of hydrogen energy through low-cost and high-yield industrial production is still a considerable challenge.¹ Although hydrogen energy can be obtained from non-renewable fossil fuels through traditional petrochemical methods, at the same time, inefficient conversion methods and resource crisis of non-renewable fossil fuels are inevitable.² In contrast, electrochemical water splitting to produce H_2 has been considered to be the most promising approach.3–5 Electrochemical water splitting mainly includes the anodic oxygen evolution reaction (OER) and the cathodic hydrogen evolution reaction (HER).⁶ These are the two most

crucial core half reaction steps in electrochemical water splitting.⁷ To overcome the overpotential (the theoretical overpotential \sim 1.23 V) caused by the activation energy barrier, a higher potential is required to trigger the initial reaction.⁸ At present, electrocatalysts based on precious metals, such as $\text{IrO}_2/\text{RuO}_2$ (OER) and Pt (HER), have higher catalytic performance. However, precious metals have poor durability and high cost as catalysts,⁹ which significantly limits the development and application of precious metal-based electrocatalysts.¹⁰ Therefore, to improve the efficiency of electrocatalysis and reduce costs, researchers are actively seeking efficient nonnoble metal electrocatalysts.¹¹⁻¹³

The reports in recent years have proved conclusively that earth-abundant 3d transition metals (3d TMs) (e.g., Ni, Co, and Fe) have enormous catalytic potential.^{14–16} For example, the catalytic performance of transition metal-based oxides is comparable to that of $IrO₂$, and transition metal-based carbides have extremely excellent catalytic stability.^{17,18} In addition, there are transition metal oxyhydroxides,¹⁹ sulfides,^{20,21} phosphides, 2^{2-24} etc., and their good catalytic performance shows the possibility of replacing precious metal catalysts.²⁵ Metal–organic frameworks (MOFs) have emerged as unique porous materials, and have been applied to numerous fields.^{26–31} The coordination of pore structure and functional sites 32 can accelerate the reaction kinetics, 33 and makes them an ideal platform for excellent electrocatalysts.³⁴⁻³⁹ However, in the structure of MOFs, the metal nodes are surrounded by organic linking groups,⁴⁰ which limits their catalytic activity

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and conductivity;⁴¹ thereby, it is vital to construct these structures with a rational design.⁴²⁻⁴⁵ In terms of macrostructure control,⁴⁶ Lou et al. typically construct a unique core–shell structure through chemical etching methods and other methods,⁴⁷ which could provide a larger catalytic interface $area₁⁴⁸$ thereby significantly improving the electrocatalytic performance.⁴⁹ Li et al. used carbonization to precisely regulate Hofmann-type MOFs into different forms (including nanosheets, nanoflowers,⁵⁰ nanotubes and aggregates), which OER catalytic performance far exceeds that of commercial $\rm RuO_2$ catalysts. 51 In terms of microstructure design, doping heterogeneous metal atoms and different ligands into the crystal lattice can improve the electronic structure.⁵²⁻⁵⁴ The latter is more challenging,⁵⁵ and the electrocatalytic performance of MOFs needs to be enhanced to the next level.⁵⁶

To deal with the above challenges, we have synthesized a bimetal–organic framework (FeCo-L₁L₂) using a simple and cost-effective solvothermal method using the free assembly of Fe and Co ions with 2,5-dihydroxyterephthalic acid (defined as L_1) and 4,6-dihydroxyisophthalic acid (defined as L_2). The synthetic strategy is shown in Scheme 1. The synthesis process is simple and does not require additional modification and processing. In the structure of $FeCo-L₁L₂$, the isomeric ligand L_2 replaces the partial ligand L_1 to coordinate with the metal ion, causing the coordination environment of the metal center to change. Moreover, the synergistic effect of the bimetal also provides abundant metal active sites for the FeCo- L_1L_2 structure. Therefore, FeCo-L₁L₂ shows excellent OER activity in 1.0 M KOH. Under the ultra-low current density conditions of 10 mA cm^{-2} , the overpotential is only 283 mV, the Tafel slope is 31.6 mV dec $^{-1}$, and the long-lasting stability was excellent. This work not only improved the catalytic performance of MOFs in OER, but also proposed a new strategy for the structural design of MOFs.

Results and discussion

The scanning electron microscopy (SEM) images in Fig. 1 show that FeCo-L₁, FeCo-L₂, and FeCo-L₁L₂ have different surface morphologies. As shown in Fig. 1, the morphology of FeCo-L1 comprises about 1 μ m sized uniform spindle-shaped particles. $FeCo-L₂$ comprises nano-particles with no specific morphology,

Scheme 1 Schematic illustration of the synthesis of FeCo-L₁L₂ electrocatalysis.

Fig. 1 (a) SEM image of FeCo-L₁, (b) SEM image of FeCo-L₂, (c) SEM images and (d) (e) TEM images of FeCo-L₁L₂, (f) XRD patterns of the as-synthesized FeCo-L₁, FeCo-L₂, and FeCo-L₁L₂ with the corresponding simulated patterns.

to improving the OER performance. Thus, the nano-acicular structure does have advantages for enhancing the OER performance of FeCo- L_1L_2 .

The crystal structures of the prepared $FeCo-L_1$, $FeCo-L_2$ and FeCo- L_1L_2 were analyzed by powder X-ray diffraction (XRD). As shown in Fig. 1f, the diffraction peaks of $FeCo-L₁$, $FeCo-L₂$ and FeCo-L₁L₂ at 7.0° and 12.0° correspond to the main peak positions of the corresponding simulated MOF-74. The PXRD results confirmed that $FeCo-L_1$, $FeCo-L_2$ and $FeCo-L_1L_2$ all were the isomorphs of MOF-74. Comparing the XRD patterns of FeCo-L₁L₂ with FeCo-L₁ and FeCo-L₂, it is found that adding isomeric ligand L_2 did not change the original crystal structure. However, the diffraction peak intensity of FeCo-L₁L₂ at 7.0 $^{\circ}$ and 12.0 $^{\circ}$ decreased slightly. It might because that ligand L₂ replaced the position of partial body L_1 , which caused some structural defects in the original crystalline form, resulting in a decrease in crystal crystallinity. Comparing Fe- L_1L_2 and Co- L_1L_2 with the corresponding simulated MOF-74 (Fig. S5a, ESI†), shows that the structure composed of single metal and double ligand coordination is the same as the structure consisting of double metals and double ligands coordination. It proves that no matter the incorporation of a metal or a ligand, a simple coordination substitution has occurred, and the original crystal structure of MOF-74 has not been changed. The presence of surface functional groups in FeCo- L_1 , FeCo- L_2 and FeCo- L_1L_2 was demonstrated *via* Fourier transform infrared (FT-IR) spectroscopy (Fig. S5b, ESI†). The peak observed at 3700–3000 cm^{-1} can be ascribed to the O–H vibration. Due to the hydrogen bond formed between the carbonyl group and the hydroxyl group, the O–H stretching band range is enlarged. The $\nu_{\text{C-H}}$ peak appears at the tail of the $\nu_{\text{O-H}}$ broad peak, and the O–H and C–H stretching bands are superimposed. When the carbonyl group is conjugated with the benzene ring, the π electron cloud density in the system tends to be more uniform, making the single bond shorter and the double bond elongated, and the characteristic frequency appears to shift to a

lower band. Therefore, the $C=O$ vibration showed a strong absorption band at 1750–1480 cm^{-1} . FeCo-L₁ has a strong absorption band at 1580 cm^{-1} . FeCo-L₂ showed two similar symmetrical rabbit ear peaks at 1625 cm^{-1} and 1560 cm^{-1} . It may be due to the difference in the carboxyl substitution position on the benzene ring, which makes the peak shape different. The shape of the corresponding peak in the FeCo- L_1L_2 band combined the characteristics of FeCo- L_1 and FeCo-L2. The symmetrical rabbit ear peak is weakened, but it still maintains a good peak intensity. It indicates that both ligand L_1 and ligand L_2 exist in the structure of FeCo- L_1L_2 , further proving that $FeCo-L₁L₂$ was successfully synthesized.

To gain insight into the valence engineering of $FeCo-L₁$, $FeCo-L₂$ and $FeCo-L₁L₂$, the structural and electronic properties of FeCo-L₁, FeCo-L₂ and FeCo-L₁L₂ were compared by X-ray photoelectron spectroscopy (XPS) (Fig. S6, ESI†). The full range XPS spectra proved the presence of Fe, Co, C, O and N elements in FeCo- L_1 , FeCo- L_2 and FeCo- L_1L_2 . Moreover, the chemical states of Fe and Co in FeCo-L₁, FeCo-L₂ and FeCo-L₁L₂ are very similar in Fig. 2. The Fe 2p spectrum of $FeCo-L_1L_2$ shows two main peaks at 710.87 and 724.37 eV. $62,63$ Compared with FeCo- L_1 and FeCo- L_2 , the binding energy of Fe 2p_{3/2} in FeCo- L_1L_2 is reduced by about 0.3 eV (Fig. S7a, ESI†). In particular, the Co 2p spectrum of FeCo- L_1L_2 shows two main peaks at 781.08 and 796.80 eV. 64 Compared with FeCo-L₁ and FeCo-L₂, the binding energy of Co $2p_{3/2}$ in FeCo-L₁L₂ is also reduced by 0.16 eV (Fig. S8b, ESI†). The isomeric ligand L_2 replaces the partial ligand L_1 to coordinate with the metal ion, which causes the coordination environment of the metal center to change.⁶⁵ For further confirmation, the Fe 2p spectrum and the Co 2p spectrum of Fe- L_1L_2 , Co- L_1L_2 and FeCo- L_1L_2 were compared. It was found that the binding energy of Fe $2p_{3/2}$ and Co $2p_{3/2}$ did not decrease. Therefore, incorporating isomeric ligands can change the coordination environment of metal centers and enhance the coupling of Fe and Co in the bimetal FeCo- L_1L_2 . It is worth noting that the synergistic effect of bimetals has been

recognized by the researchers as having a vital role in improving the catalytic performance of the OER, $66-68$ and the substitution coordination of isomeric ligands still needs theoretical analysis.⁶⁹ On the other hand, quantitative analyses of FeCo- L_1 , FeCo-L₂ and FeCo-L₁L₂ from XPS are consistent with the results of EDS spectrum atomic analysis, which all prove that the atomic ratio of Fe and Co is approximately 1 : 1 (Table S1, ESI†).

Electrochemical measurements were carried out in a standard three-electrode system in an alkaline medium (1.0 M KOH solution), and the samples were utilized as catalysts directly by depositing onto a glassy-carbon electrode (GCE, geometric area: 0.07 cm^{-2}). Linear sweep voltammetry (LSV) tests were performed on the prepared samples and commercial precious metal catalysts (IrO₂) at a scanning speed of 10 mV $\rm s^{-1}$ to characterize the OER activity. Fig. 3a and b show that the overpotentials for FeCo-L₁, FeCo-L₂ and FeCo-L₁L₂, Co-L₁L₂, and commercial IrO₂ at 10 mA cm⁻² are 366 mV, 353 mV, 283 mV, 321 mV and 309 mV, respectively. In particular, the FeCo- L_1L_2 displayed the best OER performance, which is 26 mV better than commercial $IrO₂$. Moreover, the overpotential at 50 mV cm⁻² for FeCo-L₁L₂ is only 314 mV, which is far below those overpotentials of $FeCo-L_1$, $FeCo-L_2$, and $Co-L_1L_2$ at 50 mV cm⁻². Fig. S9 (ESI[†]) further supplements the LSV curves of a single metal and a single ligand. The comparison found that after the introduction of Co ions in Fe- L_1 and Fe- L_2 , a heterogeneous bimetal FeCo- L_1 and FeCo- L_2 . With the synergistic effect of mixed metals, its OER catalytic activity is optimized. Interestingly, after adding an isomeric ligand $L₂$ to Co-L₁, the substitution of ligand L₂ connects some metal ions to enhance the catalytic effect of $Co-L_1L_2$. Therefore, we found

Fig. 3 OER performance of various samples. (a) Polarization curves (5 mV s⁻¹) in 1.0 M KOH solution and (b) overpotentials required for 10 mA cm⁻² and 50 mA cm⁻². (c) Tafel plots. (d) Electrochemical impedance plots. (e) Capacitive currents as a function of the scan rate to give the double-layer capacitance (C_{dl}). (f) Chronopotentiometry curves of FeCo-L₁L₂ for 10 h at 10 mA cm⁻² in 1.0 M KOH solution. Inset: LSV curves before and after 1000 cycles for FeCo-L₁L₂

that whether it is the introduction of Co ions into $Fe-L_1L_2$ or the addition of ligand L_2 to FeCo- L_1 , FeCo- L_1L_2 can enhance the OER catalytic performance.

The Tafel slope further evaluated the OER kinetics of electrocatalysts, as shown in Fig. 3c. The results of the Tafel slope are consistent with LSV, and $FeCo-L₁L₂$ has a superior Tafel slope (31.6 mV dec⁻¹), which is lower than that of most of the reported OER catalysts. Electrochemical impedance spectroscopy (EIS) tests were performed to explore the charge transport dynamics further. Fig. 3d shows that the $R_{\rm ct}$ is about 10 Ω for FeCo- L_1L_2 , which the Nyquist semicircle is much smaller than other catalyst samples. It indicates that $FeCo-L₁L₂$ has a lower mass transfer resistance and a faster reaction rate. The doublelayer capacitance (C_{d1}) was calculated through the measured cyclic voltammogram (CV) to explore the electrochemical active surface area (ECSA) of the samples. Each sample was scanned at 20, 40, 60, 80, 100 and 120 mV $\rm s^{-1}$ to get different CV plots (Fig. S10, ESI†). As shown in Fig. 3e, it is clear that $FeCo-L₁L₂$ has the largest C_{dl} (57.5 mF cm^{-2}). The addition of another ligand could significantly improve the C_{d1} of FeCo-L₁L₂. It shows that under a specific catalytic area, the number of active sites in $FeCo-L₁L₂$ increases, which means the density of active sites increases. Fig. 3f shows the chronopotential curve tested at a constant current density (10 mA $\rm cm^{-2})$ to evaluate the durability and stability of FeCo-L₁L₂. After FeCo-L₁L₂ worked stably for 10 hours, the crystal structure and morphology showed very limited changes (Fig. S11, ESI†) and retained 98.9% of the initial potential. In addition, the LSV curve of FeCo-L₁L₂ only slightly changed after 1000 CVs, which depicts the superior and long-term stability of the obtained catalyst.

To optimize the ratio of bimetals to the dual ligands and design $\text{FeCo-L}_1\text{L}_2$ with the best catalytic performance, we performed linear sweep voltammetry (LSV) tests on FeCo- L_1L_2 samples with different metal ratios and ligand ratios (Fig. S12, ESI†). To more intuitively demonstrate the relationship in metal ratio, ligand ratio and catalytic performance, volcanotype plots of overpotential with different metal atom ratios and different ligand ratios at 10 mA cm^{-2} were obtained, as shown in Fig. 4. When a single metal or ligand $FeCo-L₁L₂$ is incorporated with heterogeneous metal atoms or isomeric ligands, the

OER catalytic performance of $FeCo-L₁L₂$ changes significantly. The incorporation of heterogeneous metal atoms improves the electronic structure, so the catalytic activity of $FeCo-L₁L₂ OER$, which has the synergistic effect of bimetal, is significantly enhanced. Unexpectedly, adding isomeric ligands replaced part of the in situ ligands, optimized the coordination environment, and further improved the OER catalytic performance of FeCo- L_1L_2 . According to Fig. 4, it can be found that the OER performance of $FeCo-L₁L₂$ is the best when the ratio of bimetal and the ratio of double ligand are close to $1:1$. At the ratio of 1 : 1, the morphology and electronic structures of the electrocatalysts were optimized, resulting to the best OER performance.

Conclusions

We successfully prepare a novel bimetal MOF with high-efficiency OER catalytic performance by the free assembly of metal ions Fe, Co, 2,5-dihydroxyterephthalic acid (L_1) , and 4,6-dihydroxyisophthalic acid (L_2) . In the structure of FeCo- L_1L_2 , the isomeric ligand L_2 replaces the partial ligand L_1 to coordinate with the metal ion, causing the coordination environment of the metal center to change. Moreover, the synergistic effect of the bimetals also provides abundant metal active sites for the FeCo- L_1L_2 structure. Therefore, $FeCo-L₁L₂$ shows excellent OER activity in 1.0 M KOH. Under the ultra-low current density conditions of 10 mA cm^{-2} , the overpotential is only 283 mV, the Tafel slope is 31.6 mV dec $^{-1}$, and the long-lasting stability is excellent. This work not only improved the catalytic performance of MOFs in OER, but also proposed a new strategy for the structural design of MOFs.

Conflicts of interest

There are no conflicts to declare.

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