

## RESEARCH ARTICLE

View Article Online  
View Journal | View IssueCite this: *Inorg. Chem. Front.*, 2023,  
10, 158

## Multivariate indium–organic frameworks for highly efficient carbon dioxide capture and electrocatalytic conversion†

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It is crucial to design highly selective and cost-effective materials for efficient CO<sub>2</sub> uptake and the desirable transformation of CO<sub>2</sub> into fuels or high-value chemical products. Here, we synthesize a series of trinuclear indium cluster-based metal–organic frameworks (MOFs) for highly efficient CO<sub>2</sub> capture and electrocatalytic conversion into C<sub>1</sub> products. By adjusting the metal type and replacing the ligand, the mixed metal-based MOF InCo-ABDBC-HIN shows high CO<sub>2</sub> capture performance (101.7 cm<sup>3</sup> g<sup>-1</sup> at 273 K under low pressure) and superior activity (a maximum FE<sub>C<sub>1</sub></sub> of 81.5% at –2.2 V vs. Ag/Ag<sup>+</sup>) for electrocatalytic CO<sub>2</sub> reduction in organic electrolytes due to its modified electronic structure. Theoretical calculations suggest that the substitution of In by Co significantly modified the electronic structure of In-based clusters, which provides rich unpaired electrons for the CO<sub>2</sub>RR and enhances the charge transfer efficiency. This work provides a specially coordinated trinuclear indium cluster-based MOF and sheds new light on the precise structural design of both CO<sub>2</sub> adsorbents and electrocatalysts for efficient CO<sub>2</sub> reduction.

Received 19th September 2022,  
Accepted 8th November 2022

DOI: 10.1039/d2qi02020c

rsc.li/frontiers-inorganic

## Introduction

Reducing the atmospheric CO<sub>2</sub> concentration is one of the most important scientific research goals at present.<sup>1</sup> Besides reducing CO<sub>2</sub> emissions *via* replacing fossil fuels using low-carbon or non-carbon clean energy, CO<sub>2</sub> capture and conversion is considered as a promising strategy to reduce the CO<sub>2</sub> concentration. Compared with the direct reduction of CO<sub>2</sub> emissions *via* CO<sub>2</sub> capture and storage, converting CO<sub>2</sub> into fuels,<sup>2,3</sup> such as carbon monoxide,<sup>4,5</sup> methane,<sup>6–8</sup> formate,<sup>9,10</sup> ethylene<sup>11–13</sup> and ethanol<sup>14–17</sup> is considered as a promising strategy to mitigate the greenhouse effect. Nevertheless, the conversion of CO<sub>2</sub> into high value-added products suffers from the slow kinetic reaction, the competitive hydrogen evolution reaction (HER), and poor Faraday efficiency (FE) and product selectivity.<sup>18</sup> Thus, the exploration of highly efficient and highly selective catalysts for the CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) is extremely urgent.<sup>19–22</sup>

Metal–organic frameworks (MOFs), as porous crystalline materials composed of multifarious organic linkers and metal-

rich nodes,<sup>23–26</sup> provide abundant active sites for CO<sub>2</sub> adsorption and electrocatalytic reaction. The tailorable nature and their structural diversity make them ideal candidates for CO<sub>2</sub> capture and conversion. Compared to other adsorbents, the ultra-high surface area of MOFs provides an excellent ability to absorb CO<sub>2</sub> by physical adsorption methods. The tailorable chemical environments, such as the coordination number and oxidation values, provide insight into the precise design of a structure for a highly efficient electrocatalytic CO<sub>2</sub>RR and high selectivity for the final products.<sup>27,28</sup> However, there are a number of important limitations that need to be addressed when applying MOFs to electrochemical applications, primarily their poor stability and conductivity.<sup>29</sup>

Among various MOF-based catalysts, indium (In)-based catalysts with low toxicity and environmental friendliness show high selectivity towards formate production.<sup>30–34</sup> Substantial works demonstrated that the electrochemical performance can be improved *via* the modification of the chemical environment of In-based MOFs *via* the precise control of the metal species and organic ligands. For example, Ding *et al.* reported an original In-MOF by the incorporation of redox-active organic linkers, which enhanced the electrocatalytic activity for the CO<sub>2</sub>RR to HCOO<sup>-</sup> with a highly improved FE<sub>HCOO<sup>-</sup></sub> (from 54.7% to 89.6%).<sup>35</sup> Xia *et al.* reported a peer-to-peer strategy for modular assembly of In-based catalysts by amino functionalization, which demonstrated a high FE of 94.4% for CO<sub>2</sub> electroreduction.<sup>36</sup> In addition, incorporating a second

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2qi02020c>

metal to form bimetallic compounds has been proved to be an efficient strategy to promote the electrocatalytic activity of various catalysts (*e.g.* metal oxides, metal nitrides, and single-atom catalysts). For example, Rasul *et al.* constructed a Cu–In alloy by electrodepositing a thin layer of In on oxide-derived copper, which exhibited a high selectivity of 95% towards CO.<sup>37</sup> Thus, the bimetallic regulation of In-based MOFs is promising to regulate the electrocatalytic CO<sub>2</sub> reduction performance. As a matter of fact, heterometallic In-MOFs have been reported for a wide range of applications, including N<sub>2</sub> reduction,<sup>38</sup> catalytic hydrogenation reduction of CO<sub>2</sub>,<sup>39</sup> multiple ion detection<sup>40</sup> and photocatalytic fixation.<sup>41</sup> However, to the best of our knowledge, bimetallic engineering on In-based MOFs as catalysts for the CO<sub>2</sub>RR has not been reported to date.

Here, we report a unique trinuclear cluster In-based MOF (InCo-ABDBC-HIN), in which each trinuclear cluster is eight-coordinated, as an efficient CO<sub>2</sub>RR catalyst and precise structural control was carried out to optimize the activity. Compared with the widely reported nine-coordinated structure, the unusual eight-coordinated environment provides a non-coordinate site for the adsorption of intermediates, which is promising to demonstrate enhanced electrocatalytic activity. The metal (In-ABDBC-HIN) and the functional groups in ligands (In-ABDBC-HAIN) are further regulated to boost the CO<sub>2</sub>RR performance. The results demonstrate that the introduction of Co to a partially substituted indium atom (InCo-ABDBC-HIN) enables efficient production of C<sub>1</sub> products from the CO<sub>2</sub>RR on account of the modified electronic structure of the active sites. The current density under an applied potential of –2.2 V *vs.* Ag/Ag<sup>+</sup> reached up to 21.2 mA cm<sup>–2</sup>, while a high FE of 81.5% for C<sub>1</sub> products is achieved. Density functional theory (DFT) calculations uncover that the InCo<sub>2</sub>-based structure exhibits improved unpaired electronic distribution and enhanced electrical conductivity, while the amino group suppresses the electrical conductivity, thereby resulting in the high CO<sub>2</sub>RR activity of InCo-ABDBC-HIN. The reported novel trinuclear indium cluster-based MOF is also promising as an efficient catalyst for photocatalysis, thermo-catalysis, and other electrocatalytic reactions.

## Experimental

### Chemicals

The following chemicals were used as received without further purification: indium nitrate hydrate (In(NO<sub>3</sub>)<sub>3</sub>·xH<sub>2</sub>O, >99%), cobalt nitrate hexahydrate (Co(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O, >99%), isonicotinic acid (HIN, >99%) and 2-aminoisonicotinic acid (HAIN, >97%) were purchased from Aladdin Bio-Chem Technology Co., Ltd (Shanghai, China). *N,N*-Dimethylformamide (DMF), sulfuric acid (H<sub>2</sub>SO<sub>4</sub>, 95%–98%) and acetonitrile (CH<sub>3</sub>CN) were purchased from Sinopharm Chemical Reagent Co. Ltd (Shanghai, China). The Nafion D-520 dispersion (5 wt% in lower aliphatic alcohols and water), the Nafion 117 proton exchange membrane and a gas diffusion layer (GDL) were purchased from Shanghai Hesen Electric Co.

### Synthesis

**Synthesis of 4,4'-azobenzene dicarboxylic acid (H<sub>2</sub>ABDBC).** H<sub>2</sub>ABDBC was synthesized in accordance with the previously reported method.<sup>42</sup> 4-Nitrobenzoic acid (7.5 g) and sodium hydroxide (25 g) were dissolved in 200 ml deionized water, then added to a 500 ml flask and heated at 50 °C. 3.7 M glucose is stirred and dissolved in an aqueous solution (75 mL), and then added to a round-bottomed flask. The reaction was stirred for 10 minutes, and then cooled to room temperature (RT). After stirring the solution in air for 12 hours, a proper amount of acetic acid solution was added to precipitate H<sub>2</sub>ABDBC (Fig. S4†).

**Synthesis of In-ABDBC-HIN.** In-ABDBC-HIN was synthesized following the previously reported method.<sup>43</sup> In(NO<sub>3</sub>)<sub>3</sub>·xH<sub>2</sub>O (0.060 g), H<sub>2</sub>ABDBC (0.027 g), and HIN (0.012 g) were sequentially added to a 20 mL vial, followed by 5 mL DMF, which was stirred until complete dissolution. The vial was heated at 80 °C for 3 days. The orange polyhedral crystals were washed three times with fresh DMF and acetonitrile.

**Synthesis of InCo-ABDBC-HIN.** In (NO<sub>3</sub>)<sub>3</sub>·xH<sub>2</sub>O, Co (NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O (0.029 g), H<sub>2</sub>ABDBC (27.0 mg), and HIN (0.012 mg) were mixed with the solvent of 5 mL of DMF. The vial was heated in an 80 °C oven for 3 days. The red polyhedral crystals were washed with fresh DMF and acetonitrile three times. The molecular formula is (In<sub>3</sub>O)(OH)(ABDBC)<sub>2</sub>(HIN)<sub>2</sub>H<sub>2</sub>O.

**Synthesis of In-ABDBC-HAIN.** A mixture of In(NO<sub>3</sub>)<sub>3</sub>·xH<sub>2</sub>O (0.060 g), H<sub>2</sub>ABDBC (0.027 g), and HAIN (0.014 g) was heated at 80 °C for 3 days. Sequentially, the orange polyhedral crystals were collected and washed with DMF and acetonitrile three times.

### Sample activation

The as-synthesized samples of InCo-ABDBC-HIN were immersed in acetonitrile for 3 days and acetonitrile was refreshed every 12 hours. Then, the InCo-ABDBC-HIN samples exchanged with acetonitrile were activated under vacuum at 373 K for 8 h. The sample activation conditions of In-ABDBC-HIN and In-ABDBC-HAIN were the same as those of InCo-ABDBC-HIN.

### Characterization

X-ray diffraction (XRD) was performed on a MiniFlex 600 (Rigaku Co. Ltd, Japan) with Cu K $\alpha$  radiation from 4 to 80° ( $\lambda$  = 1.5406 Å). Field emission scanning electron microscopy (FE-SEM) images were obtained on an SU8220 spectrometer (Hitachi Co. Ltd, Japan) under a voltage of 10 kV. Thermogravimetric analysis (TGA) experiments were conducted on a HCT-1 (Hengjiu Co. Ltd, Beijing). Thermo analyzer tested in the atmosphere of N<sub>2</sub> (RT–800 °C). Fourier transform infrared (FT-IR) measurements were performed on a TENSOR II spectrometer (Bruker Co. Ltd, Germany) with KBr pellets. The metallic elements in the samples and the electrolyte were obtained *via* ICP-MS using an Agilent 7800 (Agilent Co. Ltd, US).

## Electrochemical measurements

The electrodes In-ABDBC-HIN, InCo-ABDBC-HIN and In-ABDBC-HAIN were prepared using the same protocol. 20 mg of MOF was suspended in 900  $\mu\text{L}$  isopropanol with 100  $\mu\text{L}$  Nafion D521 dispersion (5 wt%) assisted by ultrasound sonication for 30 minutes to form ink. 100  $\mu\text{L}$  of ink (containing  $\sim 2$  mg MOF) was spread on the surface of the gas diffusion layer (GDL,  $1 \times 1 \text{ cm}^2$ ) to obtain the working electrode.

Electrochemical measurements were conducted in a three-electrode system (CHI660E, CH Instrument, Shanghai, China) at room temperature. Cyclic voltammetry (CV) scans were conducted in an H-type cell with a three-electrode configuration, which consisted of an Ag/Ag<sup>+</sup> (0.01 M AgNO<sub>3</sub> in MeCN) reference electrode, a platinum plate electrode as a counter electrode and a working electrode ( $1 \times 1 \text{ cm}^2$ ). The electrochemical CO<sub>2</sub>RR performances of these samples were tested in a catholyte with 0.5 M EmimBF<sub>4</sub> in MeCN and an anolyte with 0.5 M H<sub>2</sub>SO<sub>4</sub>, using an air-tight H-cell separated by a Nafion 117 proton exchange membrane (CO<sub>2</sub>-saturated). The experiments were performed in the potential range from  $-0.6 \text{ V}$  to  $-2.5 \text{ V}$  vs. Ag/Ag<sup>+</sup> at a potential scan rate of  $10 \text{ mV s}^{-1}$ . The gas products were quantitatively identified by gas chromatography (GC) and the liquid products were analyzed by <sup>1</sup>H nuclear magnetic resonance (<sup>1</sup>H-NMR) spectra. EIS spectra were recorded at an open-circuit potential of  $5.0 \text{ mV}$  ( $10^{-2}$ – $10^5 \text{ Hz}$ ) using a CO<sub>2</sub>-saturated electrolyte. Tafel plots were generated from the variation, which was obtained *via* step potential electrolysis at the partial current density against the applied over-potential.

## FE calculations

$$FE_{\text{gas}} = \frac{Z \times F \times V(\text{mL min}^{-1}) \times v(\text{vol}\%) \times P}{T \times R \times I_{\text{total}} \times 60(\text{s min}^{-1})} \quad (1)$$

where  $F$  is the Faraday constant ( $96485 \text{ C mol}^{-1}$ ),  $Z$  is the electron transfer number,  $V$  is the gas flow rate obtained using a flow meter ( $10 \text{ ml min}^{-1}$ ),  $P$  is one atmosphere ( $1.013 \times 10^5 \text{ Pa}$ ),  $R$  is the universal gas constant ( $8.314 \text{ J mol}^{-1} \text{ K}^{-1}$ ),  $I$  is the total steady-state cell current, and  $T$  is the room temperature ( $298.15 \text{ K}$ ).

$$FE_{\text{HCOOH}} = \frac{Z \times F \times n_{\text{products}}}{Q} \times 100\% \quad (2)$$

where  $Q$  denotes the total charge (C) and  $n_{\text{products}}$  denotes the moles of the obtained formate calculated by <sup>1</sup>H-NMR.

## Turnover frequency (TOF)

The turnover frequency (TOF) describes the level of catalytic activity and the calculation formula is expressed as below:

$$\text{TOF} = \frac{j_{\text{products}} \times \text{FE}}{Z \times F \times n} \times 3600 \quad (3)$$

where FE is the Faraday efficiency of the product,  $n$  is the molar weight of the catalytic sites, and  $j_{\text{products}}$  is the current density.

## Electrochemical surface area (ECSA)

The electrochemical surface area can be used to calculate the number of electrochemically active sites of the electrode, which can be calculated as follows:

$$\text{ECSA} = \frac{C_{\text{dl}}}{C_s} \quad (4)$$

where  $C_{\text{dl}}$  is the double-layer capacitance obtained by cyclic voltammetric stripping at different scanning rates in the non-Faraday region and  $C_s$  is the specific capacitance of the corresponding smooth metal or oxide electrode. The slope of the linearity between  $j$  and  $v$ ,  $C_{\text{dl}} = j/v$ . CV curves from  $-0.60 \text{ V}$  to  $-0.65 \text{ V}$  vs. Ag/Ag<sup>+</sup> were measured at different scan rates, and then the electrochemical surface area was calculated.

## DFT calculations

We performed the DFT calculations using the Quantum-ESPRESSO package as implemented in the PWscf code.<sup>44</sup> The projector augmented wave pseudopotentials were used to describe the interactions between the valence electrons and ionic cores.<sup>45</sup> We used 50 and 500 Ry cutoffs for the kinetic energy of the plane-wave basis and charge density, respectively. The Perdew–Burke–Ernzerhof form of the generalized gradient approximation was employed to describe the electronic exchange and correlation.<sup>46</sup> For simplification, the trinuclear metal clusters (In<sub>3</sub>-cluster and InCo<sub>2</sub>-cluster) with eight-coordinated ligands, which were simplified from the experimental single-crystal structure, were used for DFT calculations.

## Results and discussion

### Material characterization

Single-crystal X-ray diffraction analysis (Tables S1 and S2†) reveals that InCo-ABDBC-HIN crystallizes in the  $I\bar{4}3d$  space group. Its asymmetric structure contains a half  $\mu_3$ -OH, one half indium atom, one cobalt atom, one ABDBC ligand, one HIN ligand and one coordination hydrone (Fig. S5†). Both Co<sup>2+</sup> ions of the InCo<sub>2</sub>O cluster exhibit six-coordination by one  $\mu_3$ -OH, four O atoms and one N atom. [InCo<sub>2</sub>( $\mu_3$ -OH)(O<sub>2</sub>C)<sub>6</sub>(H<sub>2</sub>O)(py)<sub>2</sub>] is surrounded by one coordination hydrone, two pyridyl groups of HIN, two  $-\text{COO}^-$  from HIN and four  $-\text{COO}^-$  from H<sub>2</sub>ABDBC ligands (Fig. 1a). Interestingly, only two apical sites are occupied by HIN ligands and the third one is coordinated by a hydrone in the trinuclear cluster, which provides active sites for the interaction between adsorbents and the host framework (Fig. 1b). An SEM image shows that InCo-ABDBC-HIN exhibits an irregular 3D polyhedron structure (Fig. S6†).

The X-ray diffraction (XRD) results of the samples are basically consistent with the simulation results, indicating the successful synthesis of the target MOFs. The XRD patterns of InCo-ABDBC-HIN are consistent with that of In-ABDBC-HIN, indicating that the entire framework has not changed after the introduction of Co (Fig. 2a). ICP-MS (Fig. S7†) suggests that the atomic ratio of In:Co in InCo-ABDBC-HIN is approxi-



**Fig. 1** (a) Structure of the coordination environment of the metal atom and (b) the 3D framework structure along the *a*-axis.



**Fig. 2** (a) XRD patterns, (b) FTIR spectra and (c) In 3d XPS spectra for In-ABDBC-HIN, InCo-ABDBC-HIN and In-ABDBC-HAIN. (d) Co 2p XPS spectra of InCo-ABDBC-HIN.

mately 1 : 1.6, which is close to the theoretical value based on the single-crystal structure. In FT-IR, the peak at  $3449\text{ cm}^{-1}$  is due to the antisymmetric stretching modes of the amino group, suggesting the successful introduction of the amino group (Fig. 2b). TGA curves (Fig. S8†) show that a series of single indium site MOF frameworks can remain stable up to about  $370\text{ }^{\circ}\text{C}$ , implying that these In-based MOFs are thermally robust.

The electronic structures of In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN were further employed for X-ray photoelectron spectroscopy (XPS). All the resulting XPS spectra were calibrated with the C ( $\text{sp}^2$ ) peak position at  $284.6\text{ eV}$  in the C 1s spectra (Fig. S9†). The peak of In 3d ( $445\text{ eV}/452\text{ eV}$ ) indicates that  $\text{In}^{3+}$  species dominate (Fig. 2c).<sup>47</sup> For InCo-ABDBC-HIN, due to the effects of Co, the binding energies of both In  $3d_{5/2}$  and In  $3d_{3/2}$  are left shifted (Fig. 2c). The above results imply that the replacement of In by Co atoms alters the entire electronic structure of the active sites in InCo-ABDBC-HIN.<sup>48</sup> The peak profile of Co 2p ( $445\text{ eV}/452\text{ eV}$ ) indicates the existence of  $\text{Co}^{2+}$  species in InCo-ABDBC-HIN (Fig. 2d).<sup>49</sup> The charge difference between the In ions and Co ions modifies the charge distribution in the  $\text{InCo}_2$ -cluster,

which is promising to enhance both the  $\text{CO}_2$  adsorption capacity and the electrocatalytic  $\text{CO}_2\text{RR}$  performance.

**Carbon dioxide capture.**  $\text{CO}_2$  adsorption isotherms of activated In-MOF materials were investigated to determine the effect of a polymetallic strategy on the porous environments of MOFs.  $\text{N}_2$  adsorption–desorption isotherms at  $77\text{ K}$  were first profiled to reveal the pore structure of these samples. As shown in Fig. 3a, nitrogen sorption curves show completely reversible type-I isotherms, representative of the microporous structure of the samples. In addition, the Brunauer–Emmett–Teller (BET) surface areas of In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN are  $958$ ,  $1355$ , and  $1106\text{ m}^2\text{ g}^{-1}$ , respectively, which shows that the introduction of cobalt and the functional groups greatly increased the specific surface area (SSA). Based on the  $\text{N}_2$  adsorption–desorption isotherm, the pore size distribution of the samples was calculated using the Horvath–Kawazoe model, which suggested that the pore sizes are mainly in the range  $0.6\text{--}0.8\text{ nm}$  for these samples (Fig. S10†).

The  $\text{CO}_2$  uptake performance under both low pressure and high pressure was further studied. (Fig. 3b, c and Fig. S11, 12†). The maximum  $\text{CO}_2$  uptakes of In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN under low pressure at  $273\text{ K}$  are  $68.5$ ,  $101.7$ , and  $94.8\text{ cm}^3\text{ g}^{-1}$ , respectively. In-ABDBC-HAIN shows higher  $\text{CO}_2$  uptake than In-ABDBC-HIN, implying that the amino functionalization can enhance the  $\text{CO}_2$  uptake ability.<sup>50</sup> InCo-ABDBC-HIN shows the highest  $\text{CO}_2$  uptake capacity, suggesting that the incorporated Co is efficient to promote the gas uptake ability due to the modified electronic structure. For high-pressure  $\text{CO}_2$  adsorption (Fig. 3c), due to the fact that the effect of amino modification on the adsorption of  $\text{CO}_2$  molecules is greater than that of Co replacement, the maximum  $\text{CO}_2$  uptakes of In-ABDBC-HIN, InCo-



**Fig. 3** (a)  $\text{N}_2$  adsorption and desorption isotherms at  $77\text{ K}$ , (b) low-pressure  $\text{CO}_2$  adsorption isotherms at  $273\text{ K}$  and (c) high-pressure  $\text{CO}_2$  adsorption isotherms at  $273\text{ K}$  for In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN. (d) Isothermic heat of adsorption ( $Q_{\text{st}}$ ).



ABDBC-HIN, and In-ABDBC-HAIN at 273 K are 193.9, 222.6, and 234.2  $\text{cm}^3 \text{g}^{-1}$ , respectively.

To determine the interaction between the  $\text{CO}_2$  molecules and these In-MOF materials, the virial model was used to analyze the isosteric adsorption enthalpy ( $Q_{\text{st}}$ ) (Fig. 3d). The enthalpy of adsorption at zero-coverage ( $Q_{\text{st}}^0$ ) values on In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN were 19.6, 17.3, and 20.4  $\text{kJ mol}^{-1}$ , respectively, indicating that the interaction between the MOF skeleton and the  $\text{CO}_2$  molecules is weaker because of the incorporation of Co, and skeleton charge is the main factor affecting the  $\text{CO}_2$  adsorption performance, which is in good agreement with the XPS spectra. The results of  $\text{CO}_2$  uptake performance demonstrate the great potential of InCo-ABDBC-HIN for efficient  $\text{CO}_2$  conversion.

**Electrochemical  $\text{CO}_2$  reduction.** The electrochemical  $\text{CO}_2$ RR performances of these samples were tested in a catholyte with 0.5 M EmimBF<sub>4</sub> in MeCN and an anolyte with 0.5 M  $\text{H}_2\text{SO}_4$ , using an air-tight H-cell separated by a Nafion-117 proton exchange membrane ( $\text{CO}_2$ -saturated).<sup>51</sup> A gas diffusion layer (GDL) with a coarse surface was used as the support to fabricate the working electrodes. The potentials mentioned in this work are all related to  $\text{Ag}/\text{Ag}^+$  and there is no IR compensation.

The cyclic voltammetry curve of InCo-ABDBC-HIN was plotted to determine its electrocatalytic activity for the  $\text{CO}_2$ RR (Fig. S13†). Obviously, the current density is higher in the  $\text{CO}_2$ -saturated electrolyte than in the Ar-saturated electrolyte, especially as the voltage is lower than  $-1.5$  V, suggesting that their activities resulted from the  $\text{CO}_2$ RR. The potential range for the electrochemical  $\text{CO}_2$ RR was from  $-2$  to  $-2.4$  V vs.  $\text{Ag}/\text{Ag}^+$ . Fig. 4a shows the representative linear scanning voltammetry (LSV) curves of In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN. The current density from high to low is: InCo-ABDBC-HIN > In-ABDBC-HIN > In-ABDBC-HAIN in the  $\text{CO}_2$ -saturated electrolyte, indicating that metal substitution is

beneficial for the improvement of catalytic activity, while the modification of functional groups in ligands reduces the  $\text{CO}_2$ RR activity.

Electrolysis was performed at a given potential to further determine their reduction products. The gas products were quantitatively identified by GC (Fig. S14†) and the liquid products were analyzed by  $^1\text{H-NMR}$  spectra (Fig. S15†). The electrocatalytic process continued for 1 hour and the results are summarized in Fig. 4b and c. As the applied potential increases, the current density correspondingly increases (Fig. S16†). The FE for all three samples reached the maximum at  $-2.2$  V vs.  $\text{Ag}/\text{Ag}^+$ . Fig. 4b compares the FE for various catalysts at various potentials. In-ABDBC-HIN shows a higher current density than In-ABDBC-HAIN at  $-2.2$  V vs.  $\text{Ag}/\text{Ag}^+$  (Fig. 4c), indicating that the amino functional groups in In-ABDBC-HAIN are not conducive to the catalytic capability, which conflicts with the results on  $\text{CO}_2$  adsorption tests and the conclusion in some of the previous work.<sup>36</sup> We further measured the  $\text{CO}_2$ RR activity of InCo-ABDBC-HAIN to study the effects of amino modification. As shown in Fig. S17,† compared with InCo-ABDBC-HIN, InCo-ABDBC-HAIN shows reduced  $\text{CO}_2$ RR performance. The above results suggest that amino modification reduces the activity for the  $\text{CO}_2$ RR. We infer that the poor performance of In-ABDBC-HAIN is attributed to the increased steric hindrance caused by the amino modification that prevents  $\text{CO}_2$  molecules from approaching the catalytic site. The subsequent DFT calculations suggest that the amino modification suppresses the electrical conductivity, which may be a major factor for the poor  $\text{CO}_2$ RR activity of the HAIN-modified samples. Among all samples, InCo-ABDBC-HIN exhibits the highest electrocatalytic activity for the  $\text{CO}_2$ RR. The FE of  $\text{C}_1$  products ( $\text{FE}_{\text{C}_1}$ ) reaches 81.5% with a current density of 21.20  $\text{mA cm}^{-2}$  under a potential of  $-2.2$  V vs.  $\text{Ag}/\text{Ag}^+$  for InCo-ABDBC-HIN, suggesting that the incorpor-

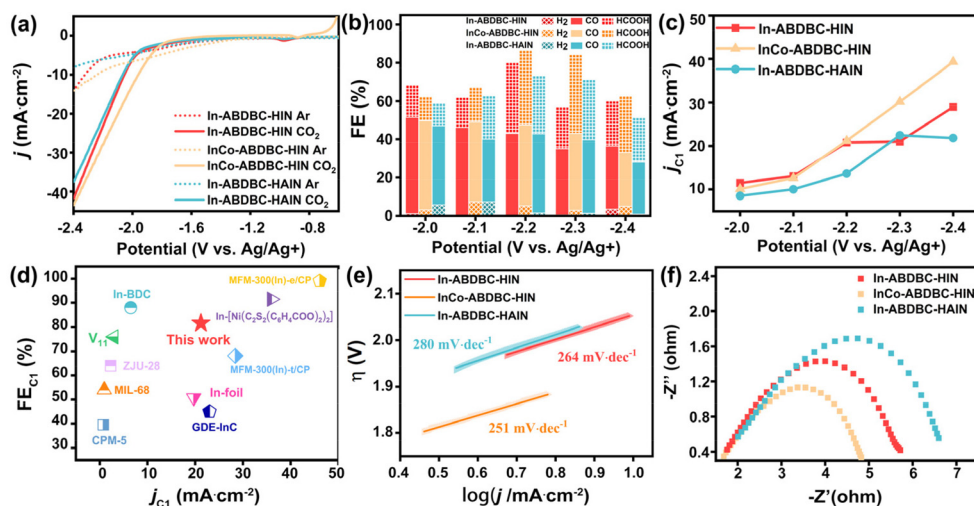


Fig. 4 (a) LSV curves, (b) FE and (c) current densities at different operation voltages of In-ABDBC-HIN, InCo-ABDBC-HIN and In-ABDBC-HAIN. (d) Comparison of InCo-ABDBC-HIN performance with other indium-based catalysts. (e) Tafel slopes and (f) Nyquist plots of In-ABDBC-HIN, InCo-ABDBC-HIN, and In-ABDBC-HAIN.

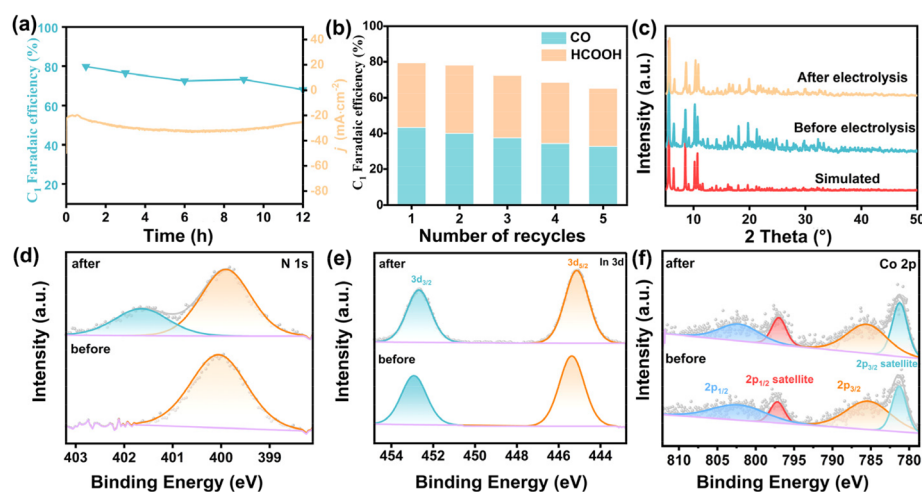
ation of Co into the framework greatly enhanced the electrocatalytic performance. Moreover, we calculated the turnover frequency (TOF, Fig. S18<sup>†</sup>) of these samples and also found that the catalytic performance of InCo-ABDBC-HIN exceeds that of In-ABDBC-HIN and In-ABDBC-HAIN when the potential is  $< -2.2$  V vs. Ag/Ag<sup>+</sup>. The electrochemical surface area (ECSA) of InCo-ABDBC-HIN exhibited a slightly larger  $C_{dl}$  (72 mF cm<sup>-2</sup>) value than In-ABDBC-HIN (63 mF cm<sup>-2</sup>) and In-ABDBC-HAIN (60 mF cm<sup>-2</sup>) (Fig. S19<sup>†</sup>). The very small difference in the ECSA implies that the high activity of InCo-ABDBC-HIN is mainly attributed to the high intrinsic activity resulting from the incorporation of Co atoms. The bimetal InCo cluster modified the electronic structure and promoted the adsorption of CO<sub>2</sub> and the activation of CO<sub>2</sub>, thereby leading to superior electrocatalytic CO<sub>2</sub>RR activity. Fig. 4d shows that the CO<sub>2</sub>RR activity of InCo-ABDBC-HIN is superior to most of the MOF-based catalysts (Table S3<sup>†</sup>).

To research the kinetics of CO<sub>2</sub> reduction to C<sub>1</sub> products over different electrocatalysts, the Tafel slope was measured (Fig. 4e). The resulting Tafel slope of InCo-ABDBC-HIN was 251 mV dec<sup>-1</sup>, which is smaller than that of In-ABDBC-HIN (264 mV dec<sup>-1</sup>) and In-ABDBC-HAIN (280 mV dec<sup>-1</sup>), implying the fast charge transfer in InCo-ABDBC-HIN. The production of OCHO\* or \*COOH is a rate-controlling step, which is formed by adsorbed carbon dioxide accepting an electron and binding to H<sup>+</sup> in the electrolyte.<sup>10,52,53</sup> The kinetics of the electrode/electrolyte interface in the electrocatalytic CO<sub>2</sub> process is represented by electrochemical impedance spectroscopy (EIS). The Nyquist plot diagram shows that the charge transfer resistance ( $R_{ct}$ ) of InCo-ABDBC-HIN (3.24 Ω) is significantly lower than that of In-ABDBC-HIN (3.94 Ω) and In-ABDBC-HIN (4.59 Ω), indicating that the charge transfer kinetics of InCo-ABDBC-HIN is the fastest (Fig. 4f).

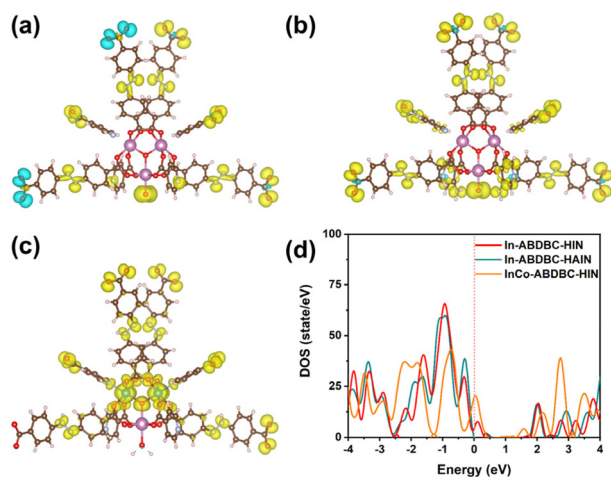
Long-term durability and cycling reproducibility were further studied to show the stability of InCo-ABDBC-HIN. The catalyst was subjected to the electrocatalytic CO<sub>2</sub>RR for con-

secutive 12 hours at  $-2.2$  V vs. Ag/Ag<sup>+</sup> to evaluate long-term durability (Fig. 5a). Although there was a slightly decrease in FE<sub>C1</sub>, the current density of InCo-ABDBC-HIN was maintained at 26.5 mA cm<sup>-2</sup>, indicating the high continuous durability. The repeatability of the In-MOF was demonstrated by five times successive repeatability tests (Fig. 5b). The FE<sub>C1</sub> retention after five times cyclic reuse reaches as high as 83.6%. The XRD patterns after long-term electrolysis show that the structure has been well preserved, implying high structural stability (Fig. 5c). In the N 1s XPS spectra after electrolysis with the InCo-ABDBC-HIN catalyst, an additional peak was observed at 401.7 eV (Fig. 5d), which was due to the N element in the electrolyte.<sup>54</sup> In 3d is slightly shifted to a low binding energy as part of In<sup>3+</sup> is reduced to In<sup>0</sup> during the electroreduction of CO<sub>2</sub> (Fig. 5e).<sup>55</sup> The Co 2p XPS spectra of the InCo-ABDBC-HIN catalyst did not change obviously before and after electrolysis (Fig. 5f). ICP-MS suggests that after continuous electrocatalysis, In and Co metals were slightly dissolved into the electrolyte, which is responsible for the reduced activity (Fig. S20<sup>†</sup>).<sup>56</sup> All these results demonstrate that the obtained InCo-ABDBC-HIN is an efficient and stable catalyst for the CO<sub>2</sub>RR.

**DFT calculations.** To reveal the effects of bimetal engineering and amino functionalization on the electronic structure, DFT theoretical calculations were further performed on In-ABDBC-HIN, In-ABDBC-HAIN, and InCo-ABDBC-HIN, which were simplified from the experimental single-crystal structures (Fig. 6a–c and Fig. S21<sup>†</sup>). As shown in Fig. 6a and b, no unpaired electronic distribution can be observed on metal atoms in both In-ABDBC-HIN and In-ABDBC-HAIN. Fig. 6a and b also imply that the amino modification nearly does not affect the charge distribution on the In<sub>3</sub>-cluster. After the incorporation of Co into the In<sub>3</sub>-cluster (Fig. 6c), InCo-ABDBC-HIN exhibits an obvious unpaired electronic distribution on Co metals. These unpaired electrons are promising for participating in the CO<sub>2</sub>RR, leading to the enhanced activity for the CO<sub>2</sub>RR. Fig. 6d shows that the DOS intensity at



**Fig. 5** (a) Long-term durability of InCo-ABDBC-HIN. (b) FE after five times electrolysis at  $-2.2$  V vs. Ag/Ag<sup>+</sup>. (c) Comparison of XRD spectra, (d) N 1s, (e) In 3d and (f) Co 2p XPS of InCo-ABDBC-HIN before and after electrolysis.



**Fig. 6** The distribution of unpaired electrons in (a) In-ABDBC-HIN, (b) In-ABDBC-HAIN, and (c) InCo-ABDBC-HIN. (d) DOS of In-ABDBC-HIN, In-ABDBC-HAIN, and InCo-ABDBC-HIN. The pink dotted line represents the location of the Fermi energy.

the Fermi energy level of InCo-ABDBC-HIN is improved compared with that of the other two structures, implying that the substitution of Co improves the electrical conductivity.<sup>57,58</sup> Moreover, the DOS intensity at the Fermi energy level of In-ABDBC-HAIN is extremely poor, indicating that the electrical conductivity is greatly suppressed due to the introduction of HAIN, thereby leading to the poor CO<sub>2</sub>RR activity. Overall, InCo-ABDBC-HIN demonstrates improved unpaired electronic distribution and enhanced electrical conductivity, and improves the CO<sub>2</sub>RR activity.

## Conclusions

In summary, we report a unique trinuclear cluster In-based MOF for efficient CO<sub>2</sub> capture and the electrocatalytic CO<sub>2</sub>RR. The precise structural control *via* bimetal engineering and ligand modification was carried out to optimize the activity. Due to the modified electronic structure and surface charge modification, InCo-ABDBC-HIN, in which In atoms are partially substituted by Co, demonstrates the highest CO<sub>2</sub> uptake performance and CO<sub>2</sub>RR activity among all the investigated samples. The CO<sub>2</sub> uptake capacity at 273 K under low pressure and high pressure is 101.7 and 222.6 cm<sup>3</sup> g<sup>-1</sup>, respectively. The current density under a potential of -2.2 V *vs.* Ag/Ag<sup>+</sup> reaches up to 21.2 mA cm<sup>-2</sup>, while a high FE<sub>C1</sub> of 81.5% is achieved. InCo-ABDBC-HIN also shows high long-term stability, good durability, and structural immobility during electrocatalysis. DFT calculations suggest that the substitution of In by Co significantly modified the electronic structure of In-based clusters, which provides rich unpaired electrons for the CO<sub>2</sub>RR and enhances the charge transfer efficiency. Our work demonstrated a novel trinuclear indium cluster-based MOF for efficient CO<sub>2</sub> capture and electrocatalytic conversion.

## Author contributions

Shuo Wang: methodology, data collection, and writing original draft. Ying Wang: writing original draft. Jia-Min Huo: data analysis. Wen-Yu Yuan: supervision, writing, checking & editing. Peng Zhang: data analysis – single-crystal analysis. Shu-Ni Li: investigation and resources. Quan-Guo Zhai: conceptualization, supervision, and funding acquisition.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by the National Natural Science Foundation of China (22071140 and 21901151), the Natural Science Foundation of Shaanxi Province (2021JLM-20), the China Postdoctoral Science Foundation (2022TQ0200) and the Fundamental Research Funds for the Central Universities (GK202101002). We acknowledge the support of Dr Shiyao Wang (State Key Laboratory of Solidification Processing, Northwestern Polytechnical University, 710072 Xi'an, China) on the theoretical calculations.

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