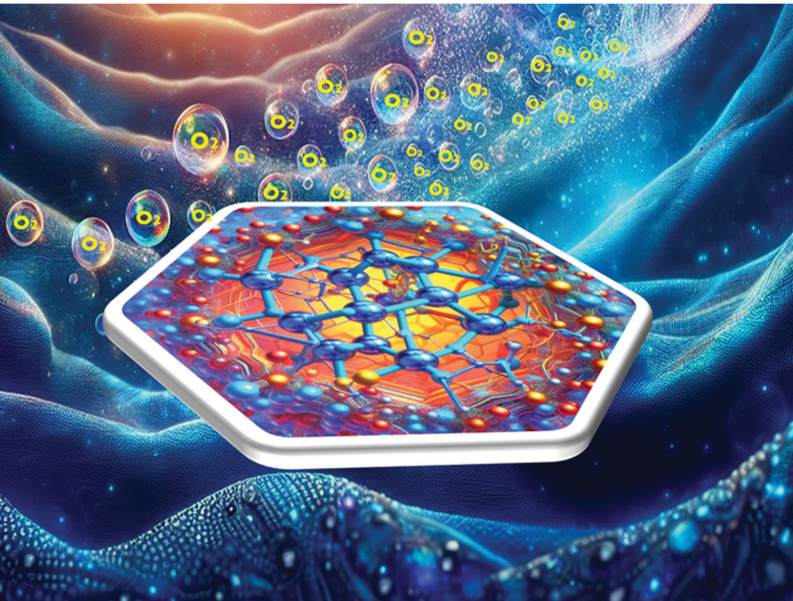
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# Catalyzing towards clean energy: tuning the oxygen evolution reaction by amide-functionalized Co(II) and Ni(II) pristine coordination polymers<sup>†</sup>

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We present the synthesis and characterization of two monometallic coordination polymers,  $[Co(L)_2(H_2O)_2]_n$ (Co-CP) and  $[Ni(L)_2(H_2O)_2]_n$  (Ni-CP), alongside a heterobimetallic counterpart, CoNi-CP, derived from an amide-based multifunctional pro-ligand 4-(pyrimidin-5-ylcarbamoyl)benzoic acid (HL), and discussed their electrocatalytic activity in the oxygen evolution reaction (OER). The CPs were characterized using various techniques, including elemental analysis, IR spectroscopy, X-ray diffraction, and thermal and powder XRD analyses. The low-cost amide-functionalized transition metal pristine coordination polymers Co-CP and Ni-CP were demonstrated to catalyze the OER in alkaline media, surpassing the benchmark  $IrO_2$  electrocatalyst performance. The heterometallic coordination polymer (CoNi-CP) displayed a lower Tafel slope value (and thus, faster kinetics) and higher long-term durability compared to its monometallic coordination polymer as a lowcost electrocatalyst of great promise and high performance for OER catalysis in alkaline media.

# Introduction

Fuel cells, metal-air batteries and water-splitting devices stand as bases in renewable energy, offering sustainable and eco-friendly solutions. Central to their efficiency, especially in water electrolysis and rechargeable metal-air batteries, lies the oxygen evolution reaction (OER).<sup>1–4</sup> However, the slow rate of reactions of the OER presents a formidable task, hindering the broader deployment of these technologies. While benchmark catalysts like RuO<sub>2</sub> and IrO<sub>2</sub> exist, their cost, availability, and stability issues in various electrolytes limit their practicality.<sup>5</sup> Hence, the quest for cost-effective, efficient OER catalysts is paramount.

In response to this challenge, considerable efforts have been channeled into developing transition metal-based electrocatalysts with enhanced OER properties.<sup>6,7</sup> Coordination polymers (CPs) and metal-organic frameworks (MOFs) have risen as notable structures in coordination chemistry, possessing tunable pore structures, large surface areas, and catalytically active metal centers.<sup>8–11</sup> By incorporating organic linkers and metals within these frameworks, we harness their dual role, providing redoxactive centers and proton transfer pathways vital for OER catalysis.<sup>8–11</sup>

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<sup>&</sup>lt;sup>†</sup> Electronic supplementary information (ESI) available: Experimental details, Scheme S1, Fig. S1–S12 containing ATR-IR, <sup>1</sup>H, <sup>13</sup>C- NMR, SXRD, topological, EIS, Nyquist plots, XPS, SEM and stability tests of **Co-CP**, **Ni-CP** and **CoNi-CP**. Tables S1–S3 present the single crystal XRD data of the crystal structures **Co-CP** and **Ni-CP**. Table S4 presents the EDX data of **CoNi-CP** and Table S5 contains OER comparison parameters. CCDC 2308477 and 2308478. For ESI and crystallographic data in CIF or other electronic format see DOI: **https//doi.org/ 10.1039/d4ce00179f** 

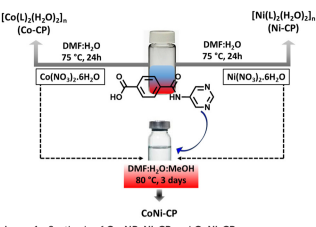
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Nonetheless, with a few notable exceptions, pristine MOFs and CPs' low conductivity is an intrinsic drawback.8,9,12 Consequently, only a limited number of these materials have been utilized as effective OER electrocatalysts.<sup>8,9,12</sup> Researchers have turned to MOF- and CP-derived materials, synthesized *via* high-temperature pyrolysis, as potential heterogeneous catalysts for the OER.8,9 However, this pyrolysis process can compromise the parent materials' intrinsic active sites and ordered structure, ultimately diminishing the catalytic activity. Strategies involving architectural modifications of MOFs and CPs have been explored to address these challenges. These modifications include introducing secondary transition metal ions and functionalized conductive linkers, among other innovative approaches.<sup>8,11,13-19</sup> Bimetallic MOFs/CPs in particular have been shown to exhibit better electrocatalytic performance than their monometallic counterparts because of the combined influence of various metals working together and the ability to adjust the engineering of the metal clusters.<sup>6,20,21</sup> Considering catalytic water splitting, heterobimetallic MOFs based on Ni-Fe,22,23 Ni-Co24 and Fe-Co<sup>25,26</sup> exhibited an improved electrocatalytic activity towards the OER when compared to monometallic MOFs. For example, nanosheets based on bimetallic Ni-Co MOFs, synthesized from mixed solutions of Ni<sup>2+</sup>, Co<sup>2+</sup> and benzenedicarboxylic acid, were reported as efficient electrocatalysts for the OER in alkaline conditions on account of a synergistic effect between the  $Ni(\pi)$  and  $Co(\pi)$ centers.27 Other heterobimetallic CPs of Co and Ni are known,<sup>6,7,21-23</sup> but there remains a significant untapped potential.

In recent years, amide-functionalized ligands have ushered in a new era of CP innovation.<sup>28</sup> These ligands offer unique binding possibilities with metal ions, leading to the assembly of diverse architectures.28 In a prior study, we reported several CPs derived from amide-functionalized ligands that exhibited remarkable activity in C-C coupling dve adsorption, supercapacitors reactions, and electrocatalysts for the OER and overall water splitting.<sup>29-32</sup> Distinguished by well-defined structures and tunable properties, these materials present a promising avenue to address challenges associated with conventional catalysts. The intricate interplay between metal centers and organic ligands in CPs/MOFs yields tailored active sites that can enhance OER kinetics. Their inherent porosity can further improve their electrochemical accessibility, facilitating efficient charge transfer.

In our previous work, we systematically changed the position of the pyridine nitrogen atom in amide-based ligands, creating CPs capable of catalysis,<sup>29</sup> dye adsorption,<sup>30</sup> water splitting,<sup>31,32</sup> and supercapacitance.<sup>31</sup> The present study presents a novel amide-based multifunctional proligand, 4-(pyrimidin-5-ylcarbamoyl)benzoic acid (HL, Scheme 1), with a pyridinium moiety.

Utilizing this pro-ligand, we successfully synthesized two pristine CPs, Ni-CP and Co-CP, and found their



Scheme 1 Synthesis of Co-NP, Ni-CP and CoNi-CP.

electrocatalytic activities in the OER. Further, we demonstrated that the heterometallic system (CoNi-CP), derived from the combination of the parent metal precursors ( $Co^{II}$  and  $Ni^{II}$ ) and the pro-ligand (HL), performs as a better OER electrocatalyst than Ni-CP and Co-CP.

# Results and discussion

#### Synthesis and characterization

The synthesis of the pro-ligand, 4-(pyrimidin-5-ylcarbamoyl) benzoic acid (HL), followed a multistep process detailed in The compound obtained underwent Scheme S1.† characterization via elemental analysis, and IR (Fig. S1<sup>†</sup>), <sup>1</sup>H and <sup>13</sup>C-NMR (Fig. S2 and S3<sup>†</sup>) spectroscopy. CPs Co-CP and Ni-CP were synthesized by subjecting a DMF solution of HL to a hydrothermal reaction in the presence of aqueous solutions of Co(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O and Ni(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O, respectively, as outlined in Scheme 1. To unravel their structural details, the synthesized CPs were subjected to elemental analysis, IR spectroscopy, and single crystal (SXRD) and powder X-ray diffraction (PXRD) analyses.

The FT-IR spectrum of the pro-ligand **HL** reveals a strong peak at 1705 cm<sup>-1</sup>, representing the v(OCO) group's stretching vibrations. Upon binding to the metal centers, this peak shifted noticeably to around 1648 cm<sup>-1</sup> (Fig. S1†). The <sup>1</sup>H NMR spectrum of **HL** reveals distinct resonances arising from the -COO(H) and -N(H) protons, evident at chemical shifts of  $\delta$  13.30 and 10.63 ppm, respectively (Fig. S2†). The aromatic protons of the phenyl ring exhibit resonances within the range of  $\delta$  8.94–7.44 ppm. In the corresponding <sup>13</sup>C NMR spectrum, the signal attributed to the  $-\underline{COOH}$  group is observed at  $\delta$  166.25 ppm, accompanied by other resonances at typical positions consistent with the formulation of the compound (Fig. S3†).

To gain insight into the thermal stability of **Co-CP** and **Ni-CP**, a thermogravimetric analysis (TGA) was conducted under a dinitrogen atmosphere, in the temperature range of 30–800 °C at a heating rate of 10 °C per minute, as depicted in Fig. S4.† In the case of **Co-CP** (Fig. S4a†), a weight loss of 6.6% within the temperature range of 150–180 °C corresponds to the elimination of the coordinated  $H_2O$  molecules. This value closely aligns with the calculated value of 6.2%. It displayed stability within the 180–360 °C temperature range, but upon further heating, the polymeric structure ultimately disintegrated.

A similar behavior was observed for Ni-CP, as depicted in Fig. S4b:† it exhibited a weight loss of 10.9% in the 150–190 °C range corresponding to the loss of the two coordinated H<sub>2</sub>O molecules (calculated value of 11.4%) and showed stability within the temperature range of 190–350 °C beyond which it underwent disintegration.

The surface area and pore characteristics of **Co-CP** and **Ni-CP** were studied by BET N<sub>2</sub> adsorption–desorption analysis, and the isotherm plots are presented in Fig. 1. According to the IUPAC classification, the nitrogen adsorption–desorption isotherm for **Co-CP** closely resembles a type I isotherm (BET surface area of 3 m<sup>2</sup> g<sup>-1</sup>) (Fig. 1a). The **Ni-CP** exhibited an isotherm reminiscent of a type III one (BET surface area of *ca.* 4 m<sup>2</sup> g<sup>-1</sup>) (Fig. 1b). To inspect the pore size distribution, non-local density functional theory (NLDFT) analysis was

employed, revealing the existence of micropores with diameters in the range of 1–1.5 nm in **Co-CP** (Fig. 1a) and both micropores and mesopores with diameters ranging from 1 to 2.5 nm in **Ni-CP** (inset, Fig. 1b). Furthermore, the N<sub>2</sub> adsorption isotherm of **CoNi-CP** exhibits characteristics typical of a type II isotherm, accompanied by an H2-type hysteresis loop as depicted in Fig. 1c. The BET surface area was found to be 16 m<sup>2</sup> g<sup>-1</sup> and its pore size distribution revealed the presence of mesopores with diameters ranging from 3 to 11 nanometers (inset, Fig. 1c).

X-ray photoelectron spectroscopy (XPS) was employed to analyze the materials' composition and oxidation states (Fig. 2 and 3). The survey spectra of **Co-CP** and **Ni-CP** revealed the presence of carbon (C), nitrogen (N), oxygen (O) along with Co in **Co-CP** and Ni in **Ni-CP** (Fig. 2). Further, the Co 2p spectrum was analyzed and deconvoluted into two primary components for **Co-CP**. The 780.4 eV and 785.7 eV peaks were assigned to  $\text{Co}^{3+}$  and  $\text{Co}^{2+}$  states, respectively (Fig. 3a). Notably, the peak originating from the Co  $2p_{1/2}$  part of the doublet was extremely weak, almost indistinguishable

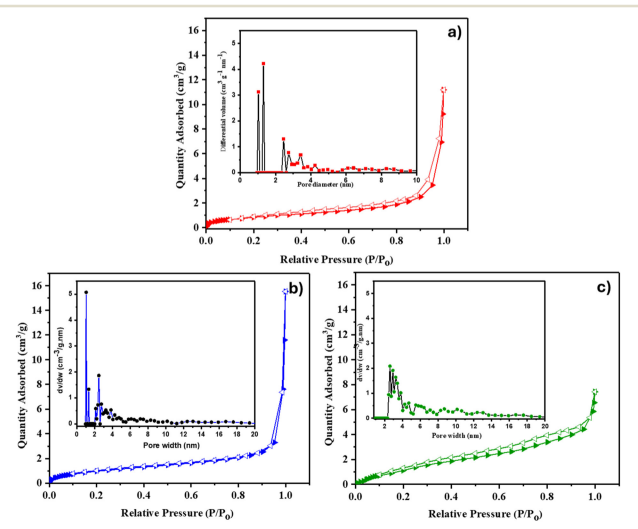
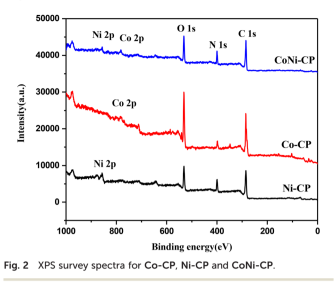


Fig. 1 N<sub>2</sub> adsorption-desorption isotherms at 77 K for Co-CP (a), Ni-CP (b) and CoNi-CP (c). Inset: Non-local density functional theory (NLDFT) pore size distribution. Filled arrows indicate adsorption and hollow arrows indicate desorption.



from the background noise, and therefore not presented. The Ni 2p spectrum for Ni-CP displayed a distinctive doublet

structure consisting of Ni  $2p_{1/2}$  and Ni  $2p_{3/2}$ , each further deconvoluted into three components (Fig. 3b). The peaks at 855.9 eV and 873.6 eV were attributed to Ni<sup>2+</sup>, while those at 860.4 eV and 877.4 eV corresponded to Ni<sup>3+</sup>. Additionally, two other peaks were identified as satellite features. XPS analysis of **CoNi-CP** is discussed below.

Atomic percentages of elements derived from the survey spectra are shown in Table 1. A low atomic percentage below 1% for Ni and Co is observed in all the samples. **Co-CP** is more oxidized than the other two samples with a correspondingly lower amount of N. The amount of C is constant in all samples.

SEM analysis of the CPs (**Co-CP** and **Ni-CP**) shows a polyhedral morphology composed of multiple stacking sheets (Fig. 4). Furthermore, EDX analysis was conducted to complement our findings. EDX allowed us to quantitatively determine the elemental composition of CPs **Co-CP** and **Ni-CP**, confirming the presence of C, N, oxygen O, and cobalt (Co) for **Co-CP**, and nickel (Ni) for **Ni-CP** (Fig. S5†).

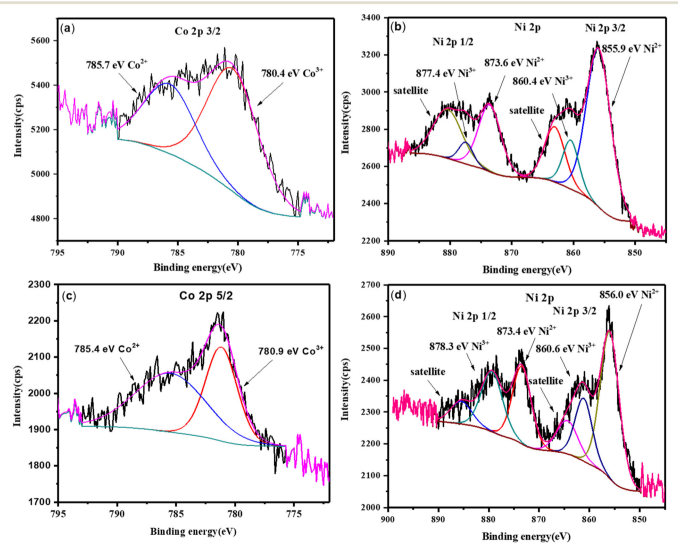


Fig. 3 High resolution XPS spectra: (a) Co 2p for Co-CP; (b) Ni 2p for Ni-CP; (c) Co 2p for CoNi-CP and (d) Ni 2p for CoNi-CP.

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Table 1 Atomic percentages of elements in samples as derived by XPS

	Ni	Со	С	О	Ν
Со-СР		0.4%	67.3%	27.7%	4.6%
Ni-CP	0.9%		67.0%	18.5%	13.6%
CoNi-CP	0.3%	0.5%	65.8%	19.8%	13.6%

#### X-ray crystallography

Polymers  $[Co^{II}(L)_2(H_2O)_2]_n$  (Co-CP) and  $[Ni^{II}(L)_2(H_2O)_2]_n$  (Ni-CP) are isomorphous and isostructural, with their asymmetric units containing one metal cation (Co<sup>II</sup> in Co-CP or Ni<sup>II</sup> in Ni-CP), a pyrimidine–carboxylate ligand (L) and a coordinated water molecule (Fig. 5 for Co-CP). Symmetry expansion uncovers the metals in nearly perfect octahedral geometries constructed from four L<sup>-</sup> moieties acting as  $\kappa N_{pyrimidine}$ -O<sub>carboxylate</sub> bridging ligands in the equatorial positions and

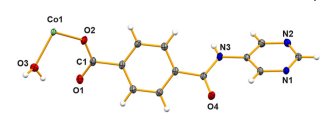


Fig. 5 The ORTEP representation (drawn at 30% probability level) of the asymmetric unit of  $[Co^{II}(L)_2(H_2O)_2]_n$  (Co-CP) with a partial atom labelling scheme, shown as a representative for Co-CP and Ni-CP.

two water molecules in the axial sites thus leading to N–M–N and  $O_{carboxylate}$ –M– $O_{carboxylate}$  angles of 180° and the  $O_{water}$ –M–N ones (values between 85.96(5) to 94.04(5)°, Table S2†). Despite having comparable covalent radii, the M– $O_{water}$ , M– $O_{carboxylate}$  and M– $N_{pyrimidine}$  bond distances are slightly different. The aromatic phenyl and pyrimidine rings are

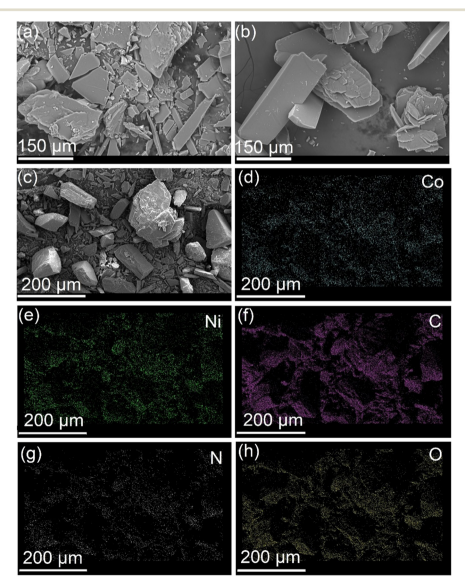


Fig. 4 SEM images of Co-CP (a), Ni-CP (b), CoNi-CP (c) and the corresponding elemental mapping of CoNi-CP using EDS: Co (d), Ni (e), C (f), N (g) and O (h).

Both polymers form elongated  $M_2L_2$  metallacycles (Fig. 6) assuming intermetallic distances of 13.470 (Co-CP) and 13.435 Å (Ni-CP) in loop-chain like one-dimensional network with base vector:  $[1\bar{0}1]$  (Fig. S6<sup>†</sup>).

Water ligands and the NH amide donors allow the formation of several H-bonding interactions in both CPs, which ultimately expand their 1D structures to the third dimension. The first level of such contacts in Co-CP is exemplified in Fig. S7a† and results from patterns between the N<sub>amide</sub> and the O<sub>carboxylate</sub> and the O<sub>water</sub> and the O<sub>amide</sub>; water molecules and pyrimidine describe a relation. In Ni-CP, the interaction between O<sub>water</sub> and the O<sub>amide</sub> was not detected, but the others described for Co-CP are active and shown in Fig. S7b.† Intramolecular interactions are common to both polymers. The highest second-level descriptors in Co-CP involve the Owater and the Oamide, or the Npyrimidine atoms (see Fig. S7a<sup>†</sup>). In Ni-CP, however, that category involves contacts engaging the Namide and the Ocarboxylate, as well as the Owater and the Npyrimidine. It is worth mentioning that H-bonds of such types are common in urea- and amide-based systems.<sup>33,34</sup> MOF The resulting three-dimensional supramolecular architecture is exemplified in Fig. 7 for the polymer Co-CP.

To enhance our comprehension of the crystal structures of **Co-CP** and **Ni-CP**, we delved into their topological characteristics by simplifying their complex multidimensional arrangements into more manageable nodeand-linker networks. The topological analysis unveiled the similarity in the framework architectures of both structures and identical 2,4-connected binodal networks (Fig. 8). However, when combining the 2D CPs and their hydrogenbonding interactions, we notice a 3,5,6 trinodal net in **Co-CP** (Fig. S8a†) and a 2,4,6 connected trinodal net in **Ni-CP** (Fig. S8b†).

We also conducted powder X-ray diffraction (PXRD) analyses of **Co-CP**, **Ni-CP** and **CoNi-CP**, which illustrate the similarity of the synthesized bulk materials (Fig. 9).

#### **Electrocatalytic studies**

The synthesized CPs **Co-CP** and **Ni-CP** were evaluated as electrocatalysts for the OER. Linear sweep voltammograms of

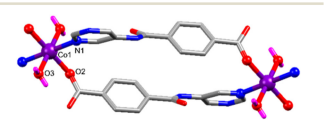
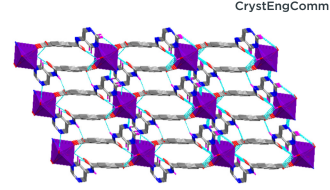


Fig. 6 The elongated  $M_2L_2$  metallacycle in  $[Co(L)_2(H_2O)_2]_n$  (Co-CP), shown as a representative for Co-CP and Ni-CP. H-atoms are omitted for clarity, except those of water ligands.



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**Fig. 7** H-bonds supported three dimensional architectures in **Co-CP**. All H-atoms except those that participate in H-bonds are omitted for clarity, and the metal environments are presented in polyhedra.

these materials were recorded under OER polarization conditions and compared to  $IrO_2$ , a benchmark catalyst for this reaction, as illustrated in Fig. 10. Both CPs exhibited nearly identical, yet superior, catalytic activity compared to the benchmark  $IrO_2$  catalyst.

The catalytic performance of these materials was evaluated based on several key parameters, including the onset potential ( $E_{onset}$ ), the overpotential required to achieve a current density of 10 mA cm<sup>-2</sup> ( $\eta_{10}$ ), the current at an overpotential of 500 mV ( $j_{500}$ ), and the Tafel slope. Both CPs displayed excellent OER activity in 1 M KOH solution, with **Ni-CP** exhibiting slightly higher performance than **Co-CP**, possibly due to the mesoporous structure enabling easier transfer of reactant/products to/from the active sites. Tafel analysis revealed a somewhat lower Tafel slope value for **Ni-CP** (110 mV dec<sup>-1</sup>) than for **Co-CP** (120 mV dec<sup>-1</sup>). **Ni-CP** also displayed a lower  $E_{onset}$  (1.549 V) and  $\eta_{10}$  (396 mV) compared to **Co-CP** ( $E_{onset} = 1.564$  V,  $\eta_{10} = 422$  mV).

Double-layer capacitance ( $C_{\rm dl}$ ) measurements were also performed in a non-faradaic region of 125 mV around the open circuit potential, as shown in Fig. S9.† The two CPs presented similar  $C_{\rm dl}$  values (Co-CP: 0.380 mF cm<sup>-2</sup>; and Ni-CP: 0.299 mF cm<sup>-2</sup>) suggesting a similar electrochemically active surface area (ECSA). Additionally, EIS measurements were conducted for both materials under OER polarization conditions (Fig. S10†), allowing the determination of both the electrolyte resistance ( $R_{\rm s}$ ) and the charge-transfer resistance ( $R_{\rm ct}$ ).  $R_{\rm s}$  showed similar values for both the CPs, with slight variations due to differences in cell geometry. In contrast,  $R_{\rm ct}$ , equivalent to the semicircle diameter observed in the Nyquist plots, exhibited a lower value of 10 Ω for Co-CP compared to 76 Ω for Ni-CP.

Motivated by the high performance of the CPs Co-CP and Ni-CP, we synthesized a heterobimetallic CP using the same

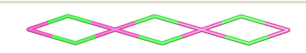
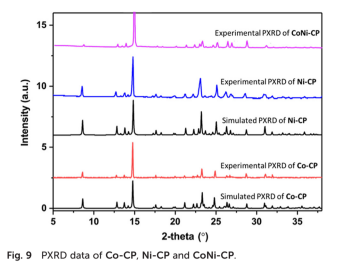


Fig. 8 Node-and-linker-type description of the 2D coordination frameworks in Co-CP and Ni-CP (metal centres indicated as pink and ligands in green).



set of starting compounds. One of the key driving factors behind the synthesis of the heterobimetallic CPs is the potential for synergy between the two different metals. The successful synthesis of CoNi-CP was achieved through the combination of equimolar amounts of Co(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O and  $Ni(NO_3)_2 \cdot 6H_2O$  with the organic linker HL during a hydrothermal process (as outlined in Scheme 1), and it represents a pivotal step in our research endeavor. The characterization studies conducted on CoNi-CP offer valuable insights into the composition and distribution of Ni and Co metals within this bimetallic CP. Inductively coupled plasma (ICP) analysis was employed to determine the global concentration of metal centers in the synthesized pink crystalline material, and the Ni to Co ratio of 1:1 was obtained. Similar PXRD patterns were observed for CoNi-CP to those of Co-CP and Ni-CP (Fig. 9). Moreover, the SEM images illustrate that the block morphology of CoNi-CP (Fig. 4c) closely resembles that of the monometallic CPs, Co-CP and Ni-CP (Fig. 4a and b). Furthermore, EDS mapping offers valuable insights into the distribution of Ni and Co

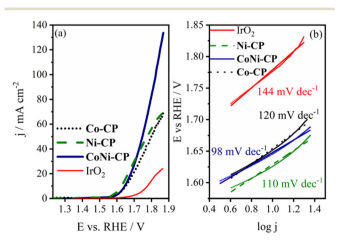


Fig. 10 OER LSV curves of Co-CP, Ni-CP and CoNi-CP in 1 M KOH electrolyte (a), and the corresponding Tafel plots and Tafel slope values (b).

metal centers. The co-localization of Ni and Co metal centers and their even distribution across the crystal surface provide further evidence of the homogeneous integration of these metals within the same material (Fig. 4d–h and Table S4†).

The elemental composition was further determined by XPS analysis (Fig. 3c and d). It confirms the presence of Ni, Co, O, N and C in the samples as expected. High resolution spectra of C 1s, O 1s and N 1s for CoNi-CP are shown in Fig. S11.† The spectrum for C 1s is deconvoluted into four components (Fig. S11a<sup>+</sup>). Peaks at 281.7, 284.1, 285.2 and 286.5 eV can be identified as carbon originating from the Cmetal (Co, Ni) bond, pyrimidine, amide and carboxylate structures, respectively.35-37 The spectrum of O 1s can be fitted with five components (Fig. S11b<sup>+</sup>). The peak at 527.5 eV can be assigned to the M-O bond (M = Co or Ni). Three peaks at 528.8, 530.8 and 532.1 eV belonging to pyrimidine, amide and carboxylate groups, respectively, are the same as for C 1s confirming the presence of these structures. Furthermore, an additional peak at 533.7 eV is attributed to the presence of adsorbed water oxygen molecules in the structure.<sup>37,38</sup> The spectrum for N 1s (Fig. S11c†) shows only two components, which originate from pyrimidine (398.9 eV) and amide (400.2 eV) structures.<sup>35-38</sup> Moreover, Co 2p and Ni 2p spectra for CoNi-CP (Fig. 3c and d) are deconvoluted analogously to the spectra of CPs Co-CP and Ni-CP with only slight differences in peak positions.<sup>39,40</sup> They confirm the presence of the same components as those of CPs Co-CP and Ni-CP.

The OER activity of CoNi-CP was studied under the same experimental conditions as in the case of its monometallic counterparts. It showed an excellent catalytic activity, surpassing the performance of the individual pristine CPs Co-CP and Ni-CP as well as of the benchmark IrO<sub>2</sub>, with a current density of 134 mA cm<sup>-2</sup> at a potential of 1.87 V  $\nu$ s. RHE (value *ca.*  $2 \times$  and  $5.4 \times$  higher than that for monometallic CPs and for commercial IrO<sub>2</sub>, respectively, Fig. 10a). Tafel analysis revealed a Tafel slope value for CoNi-CP of 98 mV  $dec^{-1}$  that is *ca.* 46 mV  $dec^{-1}$  lower than that of IrO<sub>2</sub>. *E*<sub>onset</sub> and  $\eta_{10}$  were determined to be 1.557 V and 416 mV (ca. 120 mV lower compared to IrO<sub>2</sub>), respectively. The difference between the bi- and monometallic CPs is further expressed in the current density at an overpotential of 500 mV, with CoNi-CP reaching 40.8 mA cm<sup>-2</sup> compared with 29.7 and 38.3 mA cm<sup>-2</sup> reached by CPs Co-CP and Ni-CP, respectively. The double-layer capacitance for the studied material was determined to be 1.33 mF cm<sup>-2</sup>, *i.e.*, one order of magnitude higher compared to Co-CP and Ni-CP, Fig. S9.† This indicates that the ECSA of CoNi-CP is significantly higher than those of the monometallic CPs, justifying the higher activity observed at more positive OER potentials.

The assessment of  $R_s$  and  $R_{ct}$  from the Nyquist plot (Fig. S10†), reveals the  $R_{ct}$  value for the bimetallic **CoNi-CP** of 35  $\Omega$ . The  $R_{ct}$  value of 35  $\Omega$  for the bimetallic **CoNi-CP** falls between the values of **Co-CP** and **Ni-CP** indicating a moderate charge-transfer resistance.

Investigation of the durability of the three CPs over 10 hours further supports the superior performance of the

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bimetallic CoNi-CP, Fig. S12.<sup>†</sup> An initial drop of the current density was observed in the case of all three CPs. However, the OER current densities recorded using the monometallic CPs fall close to zero by the end of the 10 hour study. Conversely, current densities recorded using the bimetallic CP stabilize and remain unchanged after 5 h. The superior electrocatalytic performance of the bimetallic CoNi-CP for the OER compared to the monometallic CPs (Co-CP and Ni-CP) may be attributed to a synergistic effect in the heterobimetallic materials.<sup>27</sup> A CoNi-CP working electrode was additionally analysed before and after the electrochemical tests by using SEM analysis with elemental mapping, depicted in Fig. S12.<sup>†</sup> The analysis revealed minor alterations in the electrode's morphology and the distribution of elements, without notable formation of oxides or oxyhydroxides, suggesting that the bimetallic CoNi-CP functions as hybrids combining attributes of both molecular and heterogeneous electrocatalysts, showcasing distinctive metal active sites.

We have also compared the electrocatalytic performance of our materials towards the OER with those of other MOFs in recent advancements in the field. For example, Bridha et al. introduced 2D MOFs with redox-active metal centers, Ni( $\pi$ ), Co( $\pi$ ) and Cu( $\pi$ ), paired with electron-rich linkers.<sup>41</sup> These MOFs displayed promising OER activity, with overpotentials in the range of 370-430 mV range comparable to ours, and Tafel slope values of 100–130 mV dec<sup>-1</sup> that are higher than (hence, not so favourable as) ours. In another study, Li et al. synthesized 2D CPs with Co(II) and Ni(II) centers using 3,5-(di(2',5'-dicarboxylphenyl)benozoic acid as the ligand source, showing overpotentials of 395 mV and 475 mV, with corresponding Tafel slopes of 68 and 143 mV dec<sup>-1</sup>.<sup>42</sup> Recently, Koner et al.<sup>15</sup> explored the role of mesaconate coordination in the OER with their Ni(II)-based CPs.  $[Ni(MCA)(bipy)(H_2O)_2]_n$  required 530 mV to reach a current density of 10 mA cm<sup>-2</sup>, while another CP without mesaconate exhibited an even higher overpotential of 630 mV. The Tafel slope values were 90 and 137 mV  $dec^{-1}$ , respectively. This comparative analysis highlights the comparable or superior activity of our CPs towards the OER, emphasizing their potential in advancing electrocatalytic applications. Further comparison with other relevant CoNi CPs (Table S5<sup>†</sup>) shows that our catalysts do not always exhibit a more favourable overpotential and Tafel slope, but it is noteworthy that such CPs, in contrast to ours, are mostly non-pristine.

# Conclusions

In this study, we have successfully synthesized and characterized two monometallic coordination polymers (CPs), **Co-CP** and **Ni-CP**, as well as a heterobimetallic CP, **CoNi-CP**, using a novel amide-based multifunctional proligand (**HL**) and assessed their electrocatalytic performance for the OER. The **Co-CP** and **Ni-CP** crystal structures were found to be isomorphous and isostructural, with octahedral metal

coordination geometries and the formation of elongated  $M_2L_2$  metallacycles. The 1D network structures were expanded into three dimensions through hydrogen-bonding interactions, resulting in unique supramolecular architectures. A topological analysis revealed the underlying framework architectures, which were characterized by 2,4-connected binodal networks in both structures.

Our findings demonstrate the promising electrocatalytic activity of these CPs, with CoNi-CP, in particular, outperforming both its monometallic counterparts and the benchmark catalyst, IrO2, with a low Tafel slope of 98 mV dec<sup>-1</sup> (ca. 46 mV dec<sup>-1</sup> lower than that of commercial  $IrO_2$ ), an overpotential of 416 mV at 10 mA cm<sup>-2</sup> (ca. 120 mV lower than that of IrO<sub>2</sub>), and current density of 40.8 mA cm<sup>-2</sup> at 500 mV overpotential (*ca.* 5x higher than that of IrO<sub>2</sub>), indicating it as one of the best-performing electrocatalysts for the OER. This heterobimetallic CP (CoNi-CP) not only exhibited an enhanced OER activity (due to improved electrochemically active surface area and lower charge transfer resistance), but also a higher longdurability, making it a highly promising term electrocatalyst. Our results also suggest the occurrence of a synergistic effect of the Co(II) and Ni(II) metals. Overall, this study provides valuable insights into the design and development of highly efficient electrocatalysts for the oxygen evolution reaction.

### Conflicts of interest

There are no conflicts to declare.

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