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## PAPER



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# A photofunctional platform of bis-terpyridine ruthenium complex-linked coordination polymers with structural diversity<sup>†</sup><sup>‡</sup>

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Solvothermal syntheses of the bis(4'-carboxyl-2,2':6',2"-terpyridine) ruthenium complex [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] as a linker and different metal anions as nodes constructed a family of ten novel coordination polymers (CPs) with different topological types, ranging from one-dimensional (1D) chains and two-dimensional (2D) networks to three-dimensional (3D) frameworks. The structural diversity of CPs is achieved by the various coordinating geometries of metal nodes, including 3d transition (monometallic or bimetallic), lanthanide, and group 4 (zirconium and hafnium) metal anions. The Ru complex linker plays the role of a light-harvesting unit and renders multifarious photofunctions to the CP materials. While the ytterbium CP (Yb-Ru) exhibits near-infrared emission upon visible-light excitation, the 2D bimetallic Co/Ni CPs ( $Co_x-Ni_y-Ru$ ) and 3D interpenetrated copper CP (Cu-Ru) show high to modest efficiencies toward the photocatalytic hydrogen evolution reaction. We suggest the series of new CPs with versatile structural traits as a unique platform for the research of photocatalysis and other photonic functional applications.

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## Introduction

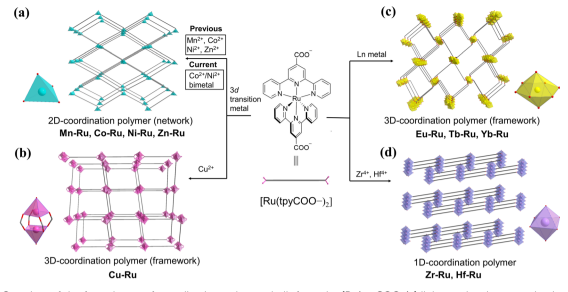
Coordination polymers (CPs), also known as metal-organic frameworks (MOFs), have emerged as a class of porous crystalline materials that attracted great interest across interdisciplinary research fields during the past decades.<sup>1-6</sup> One unique merit of CP materials is that their topological structures can be elaborately designed and modified at the molecular level by the choice of the building components-organic linkers and inorganic nodes. The chemical and physical properties of CPs at the macroscopic level, otherwise, are determined by their geometrical morphology as well as the intrinsic features of their inorganic and organic building units. Incorporation of photo- and redox-active units in the construction of CPs may not only enhance the original functionalities of these building components but also generate new functionalities, due to the potential synergistic effect between the components within the specifical spatial environments of CPs.7-15 Recent studies have shown promising applications of photosensitizer-incorporated CPs in the areas of photosynthesis,16-21 photocatalysis,9,22-27 lightemitting materials,<sup>28–30</sup> sensor materials,<sup>28,31–34</sup> and lightharvesting complexes.<sup>35–37</sup>

Polypyridyl ruthenium complexes are classic molecular photosensitizers with long-lived excited states and unique redox properties.<sup>38</sup> The porous architectures of CPs built from polypyridyl Ru( $\pi$ ) photosensitizers have shown outstanding performance in photocatalysis, especially reactions related to solar fuel production.<sup>39–46</sup> However, the tedious preparation of polypyridyl Ru( $\pi$ ) complexes and problematic crystallization of the frameworks, which might be attributed to the positive charge of the Ru( $\pi$ ) center and the obstacle of bulky polypyridyl ligands, have raised severe limitations over the research and development of functional CPs based on Ru( $\pi$ ) complex linkers.<sup>45,47–49</sup> Systematic synthesis of polypyridyl Ru( $\pi$ ) complex-linked CPs with various metal nodes and versatile topological structures is rarely reported.

We recently reported the syntheses of a series of twodimensional (2D) CPs comprising bis(4'-carboxyl-2,2':6',2''-terpyridine) ruthenium linker  $[Ru(tpyCOO^{-})_2]$ , which was readily prepared from commercially available compounds, and 3d transition-metal (Mn, Co, Ni, and Zn) nodes (Scheme 1a).<sup>50</sup> These 2D coordination networks are capable of driving highly efficient photocatalytic hydrogen evolution from water without the assistance of any co-catalysts or co-sensitizers. The current work describes our effort of systematically assembling the  $[Ru(tpyCOO^{-})_2]$  linker with various metal nodes and further outspreading a family of novel CPs. Besides the previous 2D monometallic coordination networks,<sup>50</sup> we report here four categories of coordination polymers based on the same

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<sup>†</sup> This paper is dedicated to Prof. Licheng Sun on the occasion of his 60<sup>th</sup> birthday.
‡ Electronic supplementary information (ESI) available. CCDC 2009804, 2024242, 2024244, 2024245, 2178964 and 2178965. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d2ta07219j



Scheme 1 Overview of the four classes of coordination polymers built from the [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linker and various metal anion nodes, with schematic topologies in their single crystal structures. (a) 2D bimetallic Co/Ni CPs, (b) 3D Cu CP, (c) 3D lanthanide CPs, and (d) 1D Zr and Hf CPs.

 $[Ru(tpyCOO^{-})_2]$  linker: mixed  $Co^{2+}/Ni^{2+}$ cations and [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] affording bimetallic 2D coordination networks  $Co_r-Ni_v-Ru$  (Scheme 1a),  $Cu^{2+}$ nodes bridged by [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] affording an interpenetrated 3D coordination framework Cu-Ru (Scheme 1b), Ln nodes affording isostructural 3D coordination frameworks Ln-Ru (Ln = Eu, Tb, or Yb, Scheme 1c), and group 4 Zr<sup>4+</sup> and Hf<sup>4+</sup> nodes affording 1D coordination chains Zr-Ru and Hf-Ru (Scheme 1d). The structural diversity of these [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>]-linked CPs explicitly demonstrates the viability of designing and tuning the crystalline structures of coordination polymers at the molecular level.

More importantly, integrating the light-harvesting [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] component with different metal nodes provides a unique platform of multifarious CP photofunctions.<sup>10,51-53</sup> While the photoluminescence spectrum of Yb-Ru shows emission at 976 nm upon excitation of the  $[Ru(tpyCOO^{-})_2]$  linker unit, the bimetallic  $Co^{2+}/Ni^{2+}$  coordination networks ( $Co_x - Ni_y - Ru$ ) are efficient catalysts for the visiblelight driven hydrogen evolution reaction. More interesting photofunctions are expected to be studied and exploited based on this CP platform via further constituent optimization or post-synthetic modification of the coordination frameworks.54,55

## **Results and discussion**

#### Syntheses and structures

(i) 2D bimetallic Co/Ni coordination polymers. In the previous work, we have reported the solvothermal reaction of  $[Ru(tpyCOOH)_2]^{2+}$  with  $Co(NO_3)_2$  or  $Ni(ClO_4)_2$  affording crystalline  $\{0.5Co\cdot[Ru(tpyCOO^-)_2]\cdot PF_6\}$  and  $\{0.5Ni\cdot[Ru(tpyCOO^-)_2]\cdot PF_6\}$  coordination polymers denoted as **Co-Ru** and **Ni-Ru**, respectively.<sup>50</sup> These two isomeric CPs possess a 2D coordination network topology (sql, 4<sup>4</sup>) and are readily exfoliated into ultrathin coordination polymer nanosheets that comprise several lattice layers.

By following the previous synthetic conditions for Co-Ru and Ni-Ru, we prepared 2D bimetallic Co/Ni coordination polymers as micro crystals denoted as  $Co_r - Ni_v - Ru$ , where the subscripts x and y represent the molar ratio of  $Co(NO_3)_2$  and  $Ni(ClO_4)_2$  salt precursors used in the syntheses (x/y = 1/1, 1/2, 1/4, and 1/6). The field-emission scanning electron microscopy (FE-SEM) images of Cox-Niv-Ru (Fig. S12,‡ taking Co1-Ni2-Ru as an example) display layered structures similar to the structural features of Co-Ru and Ni-Ru reported previously. The powder X-ray diffraction (PXRD) patterns of the bimetallic Co<sub>r</sub>-Ni<sub>v</sub>-Ru bulk samples (Fig. S7<sup>‡</sup>) with different x/y ratios show identical peaks to the PXRD patterns of Co-Ru or Ni-Ru, indicating the same 2D morphology of Co<sub>x</sub>-Ni<sub>y</sub>-Ru as that of Co-Ru or Ni-Ru.<sup>50</sup> Inductively coupled plasma mass spectrometry (ICP-MS) confirms the contents of both Co and Ni ions in the Cox-Niy-Ru products (Table S8<sup>‡</sup>). The measured molar metal ratios  $(n_{CO}/$  $n_{\rm Ni}$ ) in the bimetallic CPs range from  $n_{\rm Co}/n_{\rm Ni} = 15.2 (x/y = 1/1)$  to  $n_{\rm Co}/n_{\rm Ni} = 1.7 \ (x/y = 1/6).$ 

Ultrasonication in protic solvents allows exfoliation of the bimetallic Co/Ni CPs into thin nanosheets, which are shown by both the FE-SEM and transmission electron microscopy (TEM) images (Fig. 1 and S13–S15<sup>‡</sup>) of the ultrasonicated suspension.

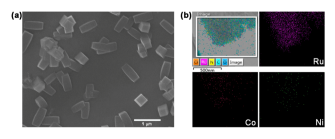


Fig. 1 (a) FE-SEM images of exfoliated  $Co_1-Ni_2-Ru$  nanosheets. (b) TEM image and elemental mapping of exfoliated  $Co_1-Ni_2-Ru$  nanosheets.

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At the same time, the element mapping identified the presence of both Ni and Co metal elements in the  $Co_x-Ni_y-Ru$  nanosheets (Fig. 1 and S15<sup>‡</sup>). The PXRD patterns of bimetallic CP nanosheets maintain all characteristic diffraction peaks of their networks prior to the exfoliation (Fig. S8,<sup>‡</sup> taking Co<sub>1</sub>–Ni<sub>4</sub>–Ru as an example), indicating the preserved 2D coordination lattices in the nanosheets. These nanosheets are estimated to consist of 3–7 layers of 2D lattices with a thickness of 2–6 nm.<sup>50</sup> The X-ray photoelectron spectroscopy (XPS) spectra of Co<sub>1</sub>–Ni<sub>4</sub>–Ru (Fig. S16 & S17<sup>‡</sup>) show binding energy peaks at 781.53 eV and 797.65 eV together with satellite peaks at 786.0 eV and 802.41 eV that are ascribed to the Co(n) ion.<sup>56,57</sup> Further the XPS Ni 2p scan of Co<sub>1</sub>–Ni<sub>4</sub>–Ru (Fig. S18<sup>‡</sup>) illustrates peaks at 855.98 eV and 873.34 eV, suggesting the presence of the Ni(n) ion.<sup>58</sup>

(ii) 3D Cu coordination polymer. The copper coordination polymer, Cu-Ru, was solvothermally prepared in DMF using  $CuCl_2$  and  $[Ru(tpyCOOH)_2]^{2+}$ . Single-crystal structure analysis of **Cu–Ru** revealed the two-fold interpenetration **cds**  $(6^5 \cdot 8)$ topology of this coordination framework, which is composed of paddlewheel cupric nodes (Fig. 2 and S1<sup>‡</sup>) and the Ru complex linker. Each dimeric copper node, with a Cu…Cu separation of 2.6389(11) Å, is coordinated by four  $\mu_2$ -bridging carboxylate groups of distinct [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linkers in a square bipyramid manner. The two axial sites of the Cu-Cu vector are both occupied by the DMF oxygen atoms. The equatorial Cu-O (carboxylate) bond lengths range from 1.947(4) to 1.986(4) Å while the axial Cu-O (DMF) distances are between 2.114(5) and 2.114(5) Å (Table S2<sup>‡</sup>). The elongation of these axial Cu–O bonds has been widely observed in the paddlewheel cupric centers due to the Jahn-Teller effect. The O-Cu-O angles in the Cucarboxylate coordination plane fall in the range of 88.37(17)° to 90.68(17)° (Table S5<sup>‡</sup>). The framework is electrically positively charged; both chloride (from the CuCl<sub>2</sub> precursor) and PF<sub>6</sub> (from the Ru complex) counterions can be identified by X-ray diffraction in the porous vacancy. The composition of Cu-Ru can be formulated as  $\{Cu \cdot [Ru(tpyCOO^{-})_2] \cdot DMF \cdot 0.5Cl \cdot 1.5PF_6\}$ .

(iii) 3D lanthanide coordination polymers. Lanthanide CPs, including Eu-Ru, Tb-Ru and Yb-Ru, were prepared *via* solvothermal methods using lanthanide nitrates and [Ru(tpyCOOH)<sub>2</sub>]<sup>2+</sup> as the reactants. The obtained crystalline precipitates of these three lanthanide CPs were resolved by single-crystal X-ray diffraction to be isostructural frameworks and therefore, only the structure of Tb-Ru was described here as an example (Fig. 3 and S3<sup>‡</sup>). There are two crystallographically

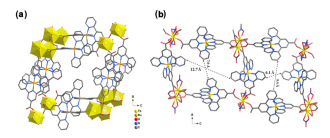


Fig. 3 (a) Stick-polyhedron representation of the Tb-Ru single crystal structure (used as an example for lanthanide CPs) viewed along the *b* axis direction, where the yellow polyhedron represents terbium nodes. (b) The sql  $(4^{4} \cdot 6^{2})$  coordination topology of Tb-Ru.

independent mononuclear Tb(III) nodes in the crystal structure. The Tb1 ion adopts a coordination geometry of capped square antiprism with nine oxygen atoms, of which four are donated by the carboxylates of Ru linkers, two by a  $\mu_2$ -nitrate, two by a  $\mu_2$ formate and one by a  $\mu_1$ -formate (Fig. S4<sup>‡</sup>). The Tb2 ion is coordinated by eight oxygen atoms from four  $\mu_1$ -carboxylates of the Ru linkers and two  $\mu_2$ -nitrates, of which one isotropic NO<sub>3</sub><sup>-</sup> plane is parallel or perpendicular to the other (Fig. S5<sup>‡</sup>). The Tb1 and Tb2 ions, both acting as 4-connected nodes, form two inorganic chains as the skeleton of a 3D framework. Accordingly, the framework of Tb-Ru, as well as Eu-Ru and Yb-Ru, possesses two types of one-dimensional channels with different diameters and can be abstracted into a sql  $(4^4 \cdot 6^2)$  topology (Fig. 3 and S3<sup> $\ddagger$ </sup>). Non-binding nitrate or  $PF_6^-$  counterions are trapped in the channels to balance the charge of lanthanide ions. The formula of Tb-Ru can be written as {1.5Tb·2  $[Ru(tpyCOO^{-})_2]$ ·3NO<sub>3</sub>·0.5PF<sub>6</sub>·HCOO<sup>-</sup>}. The compositions of counterions for Eu-Ru and Yb-Ru are slightly different (see the synthesis section in ESI<sup>‡</sup>).

(iv) 1D Zr or Hf coordination polymer. The zirconium and hafnium CPs, Zr–Ru and Hf–Ru, were prepared as dark red crystals under solvothermal conditions. The single-crystal X-ray diffraction analysis reveals isostructural coordination chains of these two CPs and thus only the structure of Zr–Ru is discussed here as an example. As shown in Fig. 4, each Zr(IV) is coordinated with six oxygen atoms from two  $\mu_1$ -carboxylates of discrete [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linkers and four hydroxy groups of solvent water molecules, forming a ZrO<sub>6</sub> octahedra node (Fig. S6‡). These linked Zr nodes and Ru complexes produce infinite one-dimensional chains parallel to each other along the *a*-axis. The individual chains align *via* the edge-to-face  $\pi$ - $\pi$  interaction and

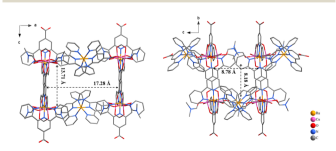


Fig. 2 Stick representation of the Cu–Ru single crystal structure, showing the cds ( $6^5 \cdot 8$ ) coordination topology.

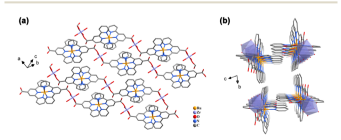


Fig. 4 (a) Stick-polyhedron representation of the Zr-Ru single crystal structure. (b) Selected adjacent coordination chains showing the interactions and void space in the structure.

DMF molecules are observed in the void space between the coordination chains. Elemental analysis (C, N, H) of **Zr-Ru** and **Hf-Ru** suggests the presence of disordered  $PF_6^-$  and  $Cl^-$  counterions trapped in the lattice of these coordination polymers.

#### Characterization and properties

The thermogravimetric analysis (TGA) demonstrates that all four categories of CPs begin to lose non-coordinating DMF and water molecules below 100 °C (Fig. S19–S22<sup>‡</sup>). A significant weight loss of these CPs occurs at higher temperatures ranging from 260 °C (for the bimetallic Co/Ni CPs) to 400 °C (for the Zr and Hf CPs), due to the degradation of the coordination frameworks. The N<sub>2</sub> sorption measurement of **Cu–Ru** at 77 K reveals a type II isotherm (Fig. S23<sup>‡</sup>), corresponding to a Brunauer–Emmett–Teller (BET) surface area of 1.9 m<sup>2</sup> g<sup>-1</sup>. The N<sub>2</sub> sorption isotherm of **Yb–Ru** (Fig. S24<sup>‡</sup>), as an example of the Ln CPs, indicates a BET surface area of 7.6 m<sup>2</sup> g<sup>-1</sup>. The weak N<sub>2</sub> gas adsorption capability of these 3D CPs is ascribed to the presence of counterions that occupy the porous space within the crystal frameworks. In addition, the interpenetrated structure of **Cu–Ru** makes it almost imporous.

The cyclic voltammogram (CV) of the bimetallic Co/Ni CP, taking **Co<sub>1</sub>-Ni<sub>2</sub>-Ru** as an example, displays redox waves at -1.6and -2.0 V *versus* ferrocenium/ferrocene (Fc<sup>+</sup>/Fc, Fig. S26<sup>+</sup><sub>4</sub>). Very similar redox features have been reported for the previously synthesized **Co-Ru** or **Ni-Ru** CPs and assigned to the redox events of [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linkers.<sup>50</sup> Such characteristic redox features at -1.6 and -2.0 V were also observed in the CVs of **Cu-Ru** and the lanthanide CPs (Fig. S27 and S28<sup>+</sup><sub>4</sub>). Despite the different metal nodes and various structures of these CPs, they all retain the redox properties of the bis-terpyridine ruthenium complex.

The solid-state UV-vis absorption spectra of these CPs all exhibit broad absorbance bands at about 300 nm and 500 nm, besides shoulder absorbance bands of some CP samples at 550 nm (Fig. S30-S33<sup>‡</sup>). These longer-wave bands in the visible light region are attributed to the characteristic metal-to-ligand charge transfer (MLCT) of [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linkers, while the short-wave bands at 300 nm originate from the  $\pi$ - $\pi$ \* excitation of tpy ligands. The solid-state emission bands of 3d transitionmetal-based CPs, Cox-Niv-Ru and Cu-Ru, are centered at approximately 670-690 nm (Fig. S36<sup>‡</sup>),<sup>50</sup> while the emission bands of lanthanide CPs are centered at about 670 nm (Fig. S37<sup>‡</sup>). The emission spectrum of [Ru(tpyCOOH)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> powder exhibits a broad band at  $\lambda_{max} = 720$  nm. These absorption and emission studies emphasize that the CPs preserve the photophysical properties of the Ru complex linker, whereas various coordinating metal nodes slightly alter the ground/excited states of the Ru linker unit.

Since the Yb( $\mathfrak{m}$ ) ion emits luminescence in the near-infrared (NIR) range distinctively separated from the emission of the Ru linker in the visible-light range, the photochemical properties of **Yb–Ru** were further investigated. Upon excitation of the Ru linker of **Yb–Ru** at  $\lambda_{ex} = 580$  nm, a sharp NIR emission band at 978 nm assigned to the  ${}^{2}F_{5/2} \rightarrow {}^{2}F_{7/2}$  transition of the Yb( $\mathfrak{m}$ ) ion<sup>59</sup> was observed in the spectrum (Fig. 5), besides the characteristic Ru

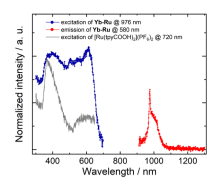


Fig. 5 Near-IR emission (red dot,  $\lambda_{ex} = 580$  nm) and excitation (blue dot,  $\lambda_{em} = 976$  nm) profiles of **Yb**-**Ru** solid powder, and the excitation profile (grey line,  $\lambda_{em} = 720$  nm) of the [Ru(tpyCOOH)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> complex in the solid state.

linker luminescent band at 670 nm (Fig. S37‡). Meanwhile, the excitation spectrum of **Yb–Ru** recorded upon the NIR luminescent band ( $\lambda_{em} = 976$  nm) shows very similar featured bands to the excitation spectrum of the [Ru(tpyCOOH)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> complex ( $\lambda_{em} = 720$  nm, Fig. 5). The excitation spectra of **Yb–Ru** collected upon the Yb(m) node emission ( $\lambda_{em} = 976$  nm) and the Ru linker emission ( $\lambda_{em} = 670$  nm) adopt similar profiles (Fig. S37‡). These spectroscopic results suggest that the Yb(m) node of **Yb–Ru** can be excited by the MLCT stated of the Ru linker, which plays the role of a sensitizer, tentatively *via* an energy transfer process.<sup>60–62</sup> The time-resolved photoluminescence experiments (Fig. S38‡) reveal a reduced decay time of the excited [Ru(tpyCOOH)<sub>2</sub>]<sup>2+</sup> complex upon coordination to the Yb(m) ions in **Yb–Ru**, consistent with the proposed energy transfer process throughout the CP framework.<sup>49,63</sup>

#### Photocatalytic hydrogen evolution

In the previous study, 2D Co-Ru and Ni-Ru nanosheets have been shown to effectively drive photocatalytic hydrogen evolution from an aqueous solution under the irradiation of visible light ( $\lambda > 420$  nm).<sup>50</sup> No co-catalyst or co-sensitizer was required for this photocatalytic hydrogen evolution process. The hydrogen evolution reaction is proposed to be initiated by quenching of the excited  $[Ru(tpyCOO^{-})_2]$  linker. The resulting formally monovalent Ru(I) species transfer electrons to the Co or Ni nodes at the edge of CP nanosheets, where the reduction of protons occurs. The driving force for photocatalytic hydrogen evolution originates from the energy gap between the LUMO level of the [Ru(tpyCOO<sup>-</sup>)<sub>2</sub>] linker and the hydrogen evolution potential. Because the bimetallic Co/Ni CPs, the Cu-Ru, and the lanthanide CPs maintain the photochemical and electrochemical properties of the  $[Ru(tpyCOOH)_2](PF_6)_2$  complex, they were assessed for photocatalytic hydrogen evolution after being dispersed in aqueous solution. Sonicating Zr-Ru and Hf-Ru in aqueous solution leads to decomposition of these two CPs and thus their photocatalytic activity towards hydrogen evolution was not investigated.

Under the irradiation of visible light ( $\lambda > 420$  nm), the exfoliated **Co<sub>x</sub>-Ni<sub>y</sub>-Ru** CP nanosheets effectively produced hydrogen from ascorbic acid solution without the presence of any co-catalysts or co-sensitizers (Fig. 6, S39 and S40<sup>‡</sup>). A pH range

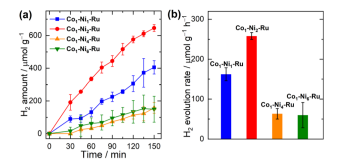


Fig. 6 (a) Photocatalytic hydrogen evolution by 2D bimetallic Co/Ni CPs under visible-light irradiation ( $\lambda > 420$  nm) from ascorbic acid solution (0.5 M, pH = 4.0). (b) Initial photocatalytic hydrogen evolution rates (µmol g<sup>-1</sup> h<sup>-1</sup>) for different 2D bimetallic Co/Ni CPs.

from 3 to 5 was tested and pH = 4 was found optimal for the photocatalytic performance of these CPs (Fig. S41<sup>‡</sup>). Among the four 2D bimetallic Co/Ni CPs, Co<sub>1</sub>–Ni<sub>2</sub>–Ru achieved the greatest initial reaction rate of 258 ± 8 µmol g<sup>-1</sup> h<sup>-1</sup>, producing the most hydrogen amount of 645 ± 8 µmol g<sup>-1</sup> in the 2.5 h reaction period (Fig. 6). More Ni content in the cases of Co<sub>1</sub>–Ni<sub>4</sub>–Ru and Co<sub>1</sub>–Ni<sub>6</sub>–Ru than Co<sub>1</sub>–Ni<sub>2</sub>–Ru significantly suppressed the photocatalytic hydrogen evolution activity, and the initial hydrogen evolution rates for Co<sub>1</sub>–Ni<sub>4</sub>–Ru and Co<sub>1</sub>–Ni<sub>6</sub>–Ru are merely 63 ± 13 and 60 ± 31 µmol g<sup>-1</sup> h<sup>-1</sup>.

Based on our previous investigation,<sup>50</sup> we propose that the photocatalytic hydrogen evolution by Cox-Niv-Ru nanosheets undergoes a reductive quenching pathway, where the excited Ru linker is first reduced by the sacrificial donor and then transfers one electron to the Co/Ni node. The driving force for the photocatalytic reaction is estimated to be about 0.7 eV at pH = 4.0, according to the redox potentials of Co/Ni CPs (Fig. S43<sup>‡</sup>). The critical step of the catalytic cycle is believed as proton association with the reduced and unsaturated Ni or Co node. Even though the previous findings concluded the more catalytically efficient Ni node at the edge of nanosheets than the edge Co node, a too large portion of Ni ions present in the bimetallic Co/ Ni CPs might create a significant number of defective sites and thus hinder the electron transfer within the bimetallic CP nanosheets. In addition, these defective sites are detrimental to the completeness of CP crystalline and reduce the amount of active catalytic nodes at the edge of nanosheets.

The **Cu–Ru** coordination polymer, after being dispersed by ultrasonic treatment, is also able to catalyze visible-light driven hydrogen evolution in the pH = 4 ascorbic acid solution with a modest reaction rate of 151 ± 19 µmol g<sup>-1</sup> h<sup>-1</sup> (Fig. S42‡). In contrast, the lanthanide CP **Eu–Ru**, **Tb–Ru** or **Yb–Ru** does not display photocatalytic activity under the same light-driven hydrogen evolution conditions described above. The type of metal nodes rather than the topologies of CPs is ascribed to the photocatalytic disparity.

## Conclusions

In the present work, we synthesized a family of ten coordination polymers composed of Ru(n) linker  $[Ru(tpyCOO^{-})_2]$  and

different metal cations, including the mixed bimetallic Co/Ni ion nodes. Single-crystal X-ray diffraction analysis revealed topological diversity of these porous coordination polymers depending on the type of metal ions. The series of CPs provide a promising platform for the development of multifarious functionalities based on the photochemical properties of the polypyridyl Ru linker. The obtained CPs were screened as catalysts for the photocatalytic hydrogen evolution reaction. Both the 2D bimetallic Co/Ni CP (Co<sub>x</sub>-Ni<sub>v</sub>-Ru) and 3D copper CP (Cu-Ru) showed notable catalytic performance without the addition of co-catalysts or co-sensitizers. Moreover, the 3D yttrium CP (Yb-Ru) exhibited characteristic near-infrared emission upon excitation of the ruthenium complex linker. These results elaborately demonstrate how various structures and functions of CPs derive from the interplay of their organic and inorganic components. Design and syntheses of CPs based on novel bis-terpyridine ruthenium complex derivatives are now in progress, aiming at further development of multifunctional materials.

## Author contributions

L. Tong and Z. Qiao conceived the concept and designed the experiments. L. Tong supervised this study. A. Wu, Y. Tang, and X. Li prepared the samples, conducted the characterization, and assessed the catalytic activity. B. Zhang carried out the photochemical investigation and processed the data. A. Zhou resolved the single crystal X-ray diffraction data and determined the CP structures. All authors discussed the results, contributed to the manuscript writing, and approved its final version.

## Conflicts of interest

There are no conflicts to declare.

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