RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2018, 8, 22676

A series of new polyoxometalate-based metal—organic complexes with different rigid pyridyl-bis(triazole) ligands: assembly, structures and electrochemical properties†

Xiuli Wang, ** Xue Bai, Hongyan Lin, Junjun Sun, Xiang Wang and Guocheng Liu

Five new polyoxometalate (POM)-based metal-organic complexes (MOCs) with different rigid pyridylbis(triazole) ligands, namely, $H\{Co_2(Hpyttz-I)_2(H_2O)_6[CrMo_6(OH)_6O_{18}]\}\cdot 8H_2O$ (1), $\{Co_2(H_2pyttz-I)_2(H_2O)_6[CrMo_6(OH)_6O_{18}]\}\cdot 8H_2O$ (1), $I_{2}(H_{2}O)_{4}[TeMo_{6}O_{24}]\}[Co(H_{2}O)_{6}] \cdot 3H_{2}O$ (2), $\{Co_{3}(Hpyttz-II)_{2}(H_{2}O)_{6}[\gamma-Mo_{8}O_{26}]\} \cdot 10H_{2}O$ (3), $\{Ni_{3}(Hpyttz-II)_{2}(H_{2}O)_{6}[\gamma-Mo_{8}O_{26}]\} \cdot 10H_{2}O$ $II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O \ \ \textbf{(4)}, \ \ \{Ni_3(Hpyttz-III)_2(H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O \ \ \textbf{(5)} \ \ (H_2pyttz-I=3-(pyrid-2-1)^2(H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O \ \ \textbf{(6)} \ \ (H_2pyttz-I=3-(pyrid-2-1)^2(H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O \ \ \textbf{(7)} \ \ \textbf{(8)} \ \ \textbf{(8)} \ \ \textbf{(9)} \ \ \textbf{(9)} \ \ \textbf{(9)} \ \ \textbf{(19)} \ \$ yl)-5-(1H-1,2,4-triazol-3-yl)-1,2,4-triazolyl, H_2 pyttz-II = 3-(pyrid-3-yl)-5-(1H-1,2,4-triazol-3-yl)-1,2,4triazolyl, H_2 pyttz -III = 3-(pyrid-4-yl)-5-(1H-1,2,4-triazol-3-yl)-1,2,4-triazolyl), were successfully synthesized and structurally characterized by single-crystal X-ray diffraction, IR spectra, powder X-ray diffraction (PXRD) and thermogravimetric analyses (TGA). Complex 1 is a two-dimensional (2D) supramolecular network based on the binuclear complex unit: [Co₂(Hpyttz-I)₂(H₂O)₆ [CrMo₆(OH)₆O₁₈]]. Complex 2 is a 1D supramolecular chain derived from the binuclear cobalt complex: {Co₂(H₂pyttz- $I_{2}(H_{2}O)_{4}TeMo_{6}O_{24})^{2-}$, the discrete $[Co(H_{2}O)_{6}]^{2+}$ units act as counter cations. Complexes 3 and 4 are isostructural with different center metals (M = Co or Ni), the adjacent γ -Mo₈O₂₆⁴⁻ anions are linked by the M II ions to form a 1D M- γ -Mo $_8O_{26}$ inorganic chain. Then 1D M- γ -Mo $_8O_{26}$ inorganic chains are linked together by the 1D metal-organic M-(Hpyttz-II) chains to form a 3D framework. In complex 5, γ - $Mo_8O_{26}^{4-}$ anions are bridged by the Ni^{II} ions to give a 1D Ni- γ - Mo_8O_{26} inorganic chain, the adjacent 1D $Ni-\gamma-Mo_8O_{26}$ chains are connected through $[Ni(Hpyttz-III)_2]$ units to form a 2D layer. The effect of POM type and coordination site of the ligands on the structures of the title complexes were discussed. The title complexes 1, 2 and 5 exhibit excellent bifunctional electrocatalytic activities toward the reduction of bromate/hydrogen peroxide and the oxidation of ascorbic acid. In addition, the redox potentials of complexes 1, 2 and 5 are highly sensitive to pH and may be used as a kind of potential pH sensor.

Received 17th April 2018 Accepted 12th June 2018

DOI: 10.1039/c8ra03277g

rsc.li/rsc-advances

Introduction

Polyoxometalates (POMs) have been attracting extensive interest in the areas of chemistry and materials, owing to their physicochemical properties and important potential applications in medicine, catalysis, electrochemistry and materials science. Many POMs-based metal-organic complexes (MOCs) with different organic ligands have been reported, in which POMs were used as building subunits or templates. In recent years, considerable progress has been made on designing POMbased MOCs materials, due to their potential applications in the fields of biology, materials chemistry, catalysis, medicine, electronics, electrochemistry, and so on.

Department of Chemistry, Bohai University, Jinzhou, 121000, P. R. China. E-mail: wangxiuli@bhu.edu.cn; Fax: +86-416-3400158; Tel: +86-416-3400158

Electrocatalytic materials have received considerable attention owing to their potential applications in electrochemical sensors and the hydrogen evolution reaction (HER) or oxygen evolution reaction (OER).5 Recently, the introduction of POMs into electrocatalytic materials represents an exciting direction based on their ability to undergo reversible redox process, which is the most important properties of POM anions.6a For example, Das' group has shown that Keggin anion $[CoW_{12}O_{40}]^{6-}$ turned into an efficient and robust water oxidation electrocatalyst upon its confinement in the well-defined void space of a metal organic framework ZIF-8, ([H₆CoW₁₂O₄₀]@ZIF-8).6b Wang's group obtained a new POM-based MOCs with polyrotaxane structure $\{Ag_8O(Htrz)_4(4,4'-bpy)_2\}\{AgPMo_{12}O_{40}\}$ (Htrz = 1,2,4-triazole, 4,4'-bpy = 4,4'-bipyridine), which possessed electrocatalytic activity for the reduction of hydrogen peroxide and bromate.7 Usually, the reduction processes of POMs are accompanied by concomitant protonation. 6a For this reason, the redox properties of many POMs are pH sensitive.8 Liu's group reported the pH dependence of redox properties of the

[†] Electronic supplementary information (ESI) available: IR Spectra, TG, and additional figures. CCDC 1586442–1586446. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8ra03277g

Paper RSC Advances

'sandwich' type cobalt(Π)-substituted POM-based compound $[Co^{\Pi}_4(H_2O)_2P_4W_{30}O_{112}]^{16-}$ in Layer-by-layer (LbL) assemblies.⁹

In fact, the organic moieties are crucial for the formation of the desired POM-based MOCs. 10-12 In other words, the organic ligands, especially the N-donor ligands including bis(pyridlyl)-,13 bis(imidazole)-,14 bis(triazole)-,15 and bis(tetrazole)16 based derivatives, have become popular for the construction of coordination architecture and a variety of POM-based MOCs have been constructed.17 However, the POM-based MOCs constructed from the pyridyl-bis(triazole) ligands have been reported very limited. 18-20 Herein, three pyridyl-bis(triazole), 3-(pyrid-2-yl)-5-(1H-1,2,4-triazol-3-yl)-1,2,4-triazolyl, 3-(pyrid-3-yl)-5-(1H-1,2,4-triazol-3-yl)-1,2,4-triazolyl, 3-(pyrid-4-yl)-5-(1H-1,2,4triazol-3-yl)-1,2,4-triazolyl (hereafter noted as H2pyttz-I, H2pyttz-II and H₂pyttz-III; see Scheme 1), were selected as the organic ligands to assemble with transition metal ions and POM anions. The ligands H₂pyttz-I, H₂pyttz-II and H₂pyttz-III possess same bis(triazole) and pyridyl groups, but different pyridyl N-sites. The multiple N-donors and various pyridyl N-sites may play a positive role in the construction of diverse structures and stable POM-based frameworks. It is worth mentioning that the introduction of the pyridyl-bis(triazole) ligands into the Anderson-type POM-based MOCs were reported rarely.21 To our knowledge, the reports on the combination of pyridylbis(triazole) ligand with the $[\gamma-Mo_8O_{26}]^{4-}$ anions systems have not been found up to now. Additionally, the Co(II)/Ni(II) ions are chosen in this work, which exhibit versatile coordination geometries and may construct novel topological structures in POMs' system.22,23

In this paper, we reported the synthesis and characterization of five POM-based metal–organic complexes based on the three pyridyl-bis(triazole) ligands above and Anderson-type POMs/octamolybdate, that is, $H\{Co_2(Hpyttz-I)_2(H_2O)_6[CrMo_6(OH)_6-O_{18}]\}\cdot 8H_2O$ (1), $\{Co_2(H_2pyttz-I)_2(H_2O)_4[TeMo_6O_{24}]\}[Co(H_2O)_6]\cdot 3H_2O$ (2), $\{Co_3(Hpyttz-II)_2](H_2O)_6[\gamma-Mo_8O_{26}\}\cdot 10H_2O$ (3), $\{Ni_3(-Hpyttz-II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (4) and $\{Ni_3(Hpyttz-III)_2(-H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (5). In complex 3, the Anderson-type CrMo_6 anions were *in situ* transformed to γ -Mo_8O_{26}^4– anions. The pH-dependent electrochemical behaviours and the bifunctional electrocatalytic properties of the title compounds have been reported.

HN—N
$$H_2$$
pyttz-II

HN—N H_2 pyttz-III

HN—N H_2 pyttz-III

Scheme 1 The ligands used in this work.

Experimental

Materials and characterization

 $(NH_4)_6[TeMo_6O_{24}]\cdot 7H_2O$ and $Na_3[CrMo_6(OH_6)O_{18}]\cdot 8H_2O$ were prepared according to a reference and confirmed by the IR spectrum. Other chemicals were of reagent grade and were used without further purification. FT-IR data were taken on a 60 Scimitar 2000 Near FT-IR Spectrometer by using KBr pellets in the range of 4000–500 cm $^{-1}$. Thermogravimetric analyses (TGA) were carried out on a Pyris Diamond TG instrument under a flowing N_2 atmosphere with a heating rate of 10 °C min $^{-1}$. Powder X-ray diffraction data were recorded on an Ultima IV with D/teX Ultra diffractometer at 40 KV and 40 mA with Cu Ka ($\lambda=1.5406$ Å) radiation. C, H, and N elemental analyses were carried out on a Perkin-Elmer 240C elemental analyser. A CHI 760 electrochemical workstation was used for the electrochemical experiments at room temperature.

X-ray crystallography

Data for complexes 1–5 were performed on a Bruker SMART APEX II diffractometer with Mo K α radiation ($\lambda=0.71069$ Å) in ω and θ scan modes at 293 K. The structures were solved by direct methods, and non-H atoms were refined anisotropically by a least-squares method on F^2 using the SHELXL-2014/7 package. For these complexes, the H atoms on C and N atoms were fixed in calculated positions. Crystal data as well as details of the data collection and refinement for the complexes are summarized in Table 1. Selected bond distances and angles are shown in Table S1.† CCDC contains the supplementary crystallographic data for this paper with deposition numbers 1586442–1586446 for 1–5.

Preparation of complexes 1-5

Synthesis of H{Co₂(Hpyttz-I)₂(H₂O)₆[CrMo₆(OH)₆O₁₈]}·8H₂O (1). A mixture of $CoCl_2 \cdot 6H_2O$ (0.12 g, 0.5 mmol), H_2 pyttz-I (0.027 g, 0.10 mmol), Na_3 [CrMo₆(OH₆)O₁₈]·8H₂O (0.3 g, 0.24 mmol) and H_2O (10 mL) was stirred for 30 min at the room temperature, and the pH value was then adjusted to about 4.0 by using 1.0 M NaOH. The suspension was put into a Teflon lined autoclave (25 mL) and heated at 120 °C for 4 days. After slow cooling to room temperature, pink block crystals 1 were obtained, washed with distilled water and dried in a desiccator at room temperature to give a yield of 23% based on Co. Anal. calcd (found %) for 1: C, 11.91 (11.83); H, 2.59 (2.55); N, 10.81 (10.73). IR (KBr, cm⁻¹) for complex 1: 3316 (m), 1651 (m), 1557(m), 1451 (m), 1228 (w), 937 (s), 913 (s), 668 (s), 580 (w).

Synthesis of $\{Co_2(H_2pyttz-I)_2(H_2O)_4 [TeMo_6O_{24}]\}[Co(H_2O)_6] \cdot 3H_2O$ (2). The synthesis of 2 was similar to that of 1, except that Na₃[CrMo₆(OH)₆O₁₈] \cdot 8H₂O was replaced by $(NH_4)_6[TeMo_6O_{24}] \cdot 7H_2O$ (0.3 g, 0.23 mmol). The pH value was then adjusted to about 3.9 using 1.0 M NaOH. Pink block crystals of complex 2 were filtered off, washed with distilled water, and dried in a desiccator at room temperature to give a yield of 36% based on Co. Anal. calcd (found %) for 2: C, 11.22 (11.17); H, 2.08 (2.02); N, 10.18 (10.09). IR (KBr pellet, cm⁻¹): 3156 (m), 1650 (m), 1294 (m), 994 (m) 944 (s), 881 (s), 668 (s), 611 (s), 550 (m).

RSC Advances

Table 1 Crystallographic data for complexes 1-5

Complex	1	2	3	4	5			
Empirical formula	$C_{18}H_{47}Co_{2}CrMo_{6}N_{14}O_{38} \\$	$C_{18}H_{40}Co_{3}TeMo_{6}N_{14}O_{37}$	$C_{18}H_{44}Co_{3}Mo_{8}N_{14}O_{42} \\$	$C_{18}H_{44}Ni_{3}Mo_{8}N_{14}O_{42} \\$	$C_{18}H_{40}Ni_{3}Mo_{8}N_{14}O_{40} \\$			
Formula weight	1810.17	1924.67	2072.98	2072.32	2036.29			
Crystal system	Triclinic	Triclinic	Triclinic	Triclinic	Triclinic			
Space group	$P\bar{1}$	$P\bar{1}$	$P\bar{1}$	$P\bar{1}$	$P\bar{1}$			
a (Å)	10.8960(17)	10.9220(6)	10.9150(4)	10.8810(4)	10.5650(7)			
b (Å)	11.1160(18)	11.0630(5)	12.4010(5)	12.3290(5)	11.4200(8)			
c (Å)	12.346(2)	21.2740(10)	12.5140(5)	12.4580(5)	12.1260(8)			
α (°)	106.385(3)	100.8430(10)	107.4950(10)	106.8430(10)	94.4490(10)			
β (°)	108.807(3)	97.7960(10)	113.3740(10)	113.5710(10)	114.0790(10)			
γ (°)	105.418(3)	94.7390(10)	102.0980(10)	102.6890(10)	95.0830(10)			
$V(\mathring{A}^3)$	1248.8(3)	2485.8(2)	1374.33(9)	1354.63(9)	1320.03(15)			
Z	1	2	1	1	1			
$D_{\rm c}~({ m g~cm}^{-3})$	2.407	2.571	2.505	2.540	2.562			
$\mu (\mathrm{mm}^{-1})$	2.430	3.130	2.758	2.922	2.994			
F(000)	884	1854	1003	1006	986			
Reflection collected	9053	18 586	7755	10 123	7580			
Unique reflections	6322	12 274	4967	6702	4668			
Parameters	366	712	385	385	407			
$R_{ m int}$	0.0215	0.0258	0.0387	0.0182	0.0122			
GOF	1.017	0.989	1.008	1.029	1.045			
$R_1^a [I > 2\sigma(I)]$	0.0390	0.0392	0.0441	0.0252	0.0214			
wR_2^b (all data)	0.1011	0.0864	0.0988	0.0594	0.0529			
$^{a}R_{1}=\Sigma F_{\mathrm{o}} - F_{\mathrm{c}} /\Sigma F_{\mathrm{o}} .$ b w $R_{2}=\Sigma[\mathrm{w}(F_{\mathrm{o}}{}^{2}-F_{\mathrm{c}}{}^{2})^{2}]/\Sigma[\mathrm{w}(F_{\mathrm{o}}{}^{2})^{2}]^{1/2}.$								

Synthesis of $\{Co_3(Hpyttz-II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (3). A mixture of Co(NO₃)₂·6H₂O (0.12 g, 0.4 mmol), H₂pyttz-II (0.02 g, 0.1 mmol), $Na_3[CrMo_6(OH_6)O_{18}] \cdot 8H_2O$ (0.12 g, 0.1 mmol) and H₂O (10 mL) was stirred for 30 min at the room temperature. The pH value was adjusted to 5.8 with 1.0 M NaOH. The suspension was transferred into a Teflon-lined autoclave (25 mL) and kept at 120 °C for 4 days. After slow cooling to room temperature, pink block crystals of 3 were obtained in a yield of 54% based on Co. Anal. calcd (found %) for 3: C, 10.41 (10.36); H, 2.12 (2.08); N, 9.45 (9.37). IR (KBr pellet, cm⁻¹): 3403 (w), 3137 (w), 1657 (s), 1557 (m), 1249 (s), 906 (s), 849 (m), 668 (s),

Synthesis of $\{Ni_3(Hpyttz-II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (4). The preparation of complex 4 was prepared in the same way as 3 except that the NiCl₂·6H₂O (0.12 g, 0.5 mmol) and (NH₄)₆-Mo₇O₂₄·4H₂O (0.12 g, 0.1 mmol) were used instead of $Co(NO_3)_2 \cdot 6H_2O$ and $Na_3[CrMo_6(OH_6)O_{18}] \cdot 8H_2O$. The pH value was then adjusted to about 4.2 using 1.0 M HCl. After slow cooling to room temperature, green block crystals of 4 were obtained and then dried in a desiccator at room temperature to give a yield of 62% based on Ni. Anal. calcd (found %) for 4: C, 10.42 (10.35); H, 2.12 (2.09); N, 9.46 (9.41). IR (KBr pellet, cm⁻¹): 3391 (w), 1657 (m), 1557 (s), 1300 (m), 949 (s), 906 (s), 849 (s), 662 (s), 555 (w).

Synthesis of $\{Ni_3(Hpyttz-III)_2(H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (5). The preparation of complex 5 was similar to that of 4, except that the H₂pyttz-III was used instead of H₂pyttz-II. The pH value was then adjusted to about 2.8 using 1.0 M HCl. After slow cooling to room temperature, green block crystals of 5 were obtained and then dried in a desiccator at room temperature to give a yield of 65% for 5 based on Ni. Anal. calcd (found %) for 5: C, 10.80 (10.74); H, 1.80 (1.77); N, 9.80 (9.76). IR (KBr pellet, cm⁻¹): 3854 (m), 1632 (s), 1563 (m), 1507 (m), 1200 (m), 1087 (s), 949 (s), 849 (s), 636 (s).

Preparation of complexes 1, 2 and 5 bulk-modified carbon paste electrode

Complex 1 bulk-modified carbon paste electrode (1-CPE) was fabricated by mixing 0.010 g of complex 1 and 0.10 g of graphite powder in an agate mortar for approximately 60 min to achieve a uniform mixture, and then 0.16 mL paraffin oil was added and stirred with a glass rod. The homogenized mixture was packed into a 3 mm inner diameter glass tube and the tube surface was wiped with weighing paper. The electrical contact was established with the copper wire through the back of the electrode. In a similar procedure, 2- and 5-CPEs were prepared with complexes 2 and 5, respectively. The bare CPE was prepared by similar process without any complexes.

Results and discussion

Syntheses

In the process of hydrothermal synthesis, several factors can influence the formation of crystal phases, such as initial reactants, molar ratio, pH value, reaction time, temperature, etc. In this work, pH value of the reaction is crucial for the formation of the title complexes. Only at the pH value about 4.0 for 1, 3.9 for 2, 5.8 for 3, 4.2 for 4 and 2.8 for 5, the crystals can be obtained. There also are some precipitates in the final products. The crystals suitable for single-crystal structure determination were manually isolated, and washed with distilled water.

Paper

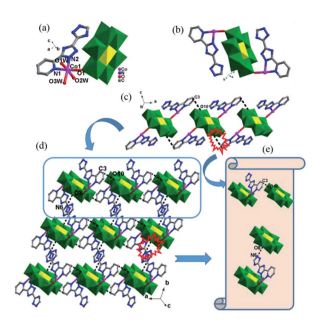


Fig. 1 (a) Coordination environment of the Co II ion in 1. The hydrogen atoms and lattice water molecules are omitted for clarity. (b) The discrete binuclear complex $[Co_2(Hpyttz-I)_2(H_2O)_6[CrMo_6(OH)_6O_{18}]]$ in 1 (c) view of the 1D supramolecular chain in complex 1. (d) The 2D supramolecular layer of 1. (e)Two types of hydrogen bonding interactions.

Structural description

 $H\{Co_2(Hpyttz-I)_2(H_2O)_6[CrMo_6(OH)_6O_{18}]\}\cdot 8H_2O$ (1). X-Ray crystal structure analysis reveals that complex 1 is a twodimensional (2D) supramolecular network based on the binuclear complex unit: [Co₂(Hpyttz-I)₂(H₂O)₆[CrMo₆(OH)₆O₁₈]]. The asymmetric unit of 1 contains two CoII ions, one [CrMo6-(OH)₆O₁₈]³⁻ (CrMo₆) anion, two deprotonated Hpyttz-I anions, six coordinated water molecules and eight lattice water molecules. To balance the charge, one proton is added in 1. The Co1 ion is six-coordinated by two nitrogen atoms from one deprotonated Hpyttz-I ligand, one terminal oxygen atom from one CrMo₆ polyoxoanion and three oxygen atoms from three water molecules in a slightly distorted octahedral geometry (Fig. 1a). The bond lengths around the Co1 ion are 2.139(4) Å for Co1-N1, 2.103(4) Å for Co1-N2, 2.072(4)-2.114(4) Å for Co1-O, the bond angles are 90.17(19)-176.70(15)° for O-Co1-N and 84.45(16)- $171.25(19)^{\circ}$ for O-Co1-O, respectively.

In complex 1, the Hpyttz-I ligand adopts a chelating mode to coordinate with one $\mathrm{Co^{II}}$ ion and each $\mathrm{CrMo_6}$ polyoxoanion acts as a bidentate ligand that coordinating to two $\mathrm{Co^{II}}$ ions by its two terminal oxygen atoms to form the binuclear unit (Fig. 1b). Interestingly, the discrete binuclear units are linked together through hydrogen bonding interactions between the oxygen atoms from the $\mathrm{CrMo_6}$ polyoxoanions and carbon atoms of the Hpyttz-I ligands $\left[\mathrm{C(3)-H(3A)\cdots O(10)},\ 3.2540\ \mathring{\mathrm{A}}\right]$ to form a 1D supramolecular chain (Fig. 1c).

The adjacent chains are further joined together by hydrogen bonding interactions $[N(6)-H(6A)\cdots O(8), 2.7150~\text{Å}]$ between the oxygen atoms of the CrMo₆ polyoxoanions and the nitrogen atoms of the Hpyttz-I ligands to generate a 2D supramolecular network, as shown in Fig. 1d and e.

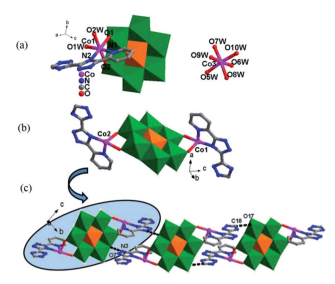


Fig. 2 (a) The coordination geometry of the Co^{II} ions in **2**. The hydrogen atoms and lattice water molecules are omitted for clarity; (b) the binuclear complex $\{Co_2(H_2pyttz-I)_2(H_2O)_{10}[TeMo_6O_{24}]\}^{2-}$; (c) the 1D supramolecular chain linked by the hydrogen bonding interaction $[N(3)-H(3A)\cdots O(7), 2.7180 \text{ Å}]$ and $[C(18)-H(18A)\cdots O(17), 3.0125 \text{ Å}]$.

 $\{Co_2(H_2pyttz-I)_2(H_2O)_4[TeMo_6O_{24}]\}[Co(H_2O)_6] \cdot 3H_2O$ (2). Single-crystal X-ray diffraction analysis shows that complex 2 is a 1D supramolecular chain based on the binuclear complex unit: $\{Co_2(H_2pyttz-I)_2(H_2O)_4[TeMo_6O_{24}]\}^{2-}$. There are three Co^{II} ions, two H2pyttz-I ligands, one TeMo6 anion, ten coordinated water molecules, three lattice water molecules in the asymmetric unit of 2. The Co1 ion is six-coordinated in a distorted octahedral coordination geometry, defined by two N atoms from one H₂pyttz-I ligand, two O atoms from one [TeMoO₂₄]⁶⁻ (TeMo₆) polyoxoanion and two O atoms from two coordinated water molecules (Fig. 2a). The bond lengths around the Co1 ion are 2.118(4) Å for Co1-N1, 2.142(4) Å for Co1-N2, 2.046(3)-2.129(4) Å for Co1–O, the bond angles are 89.80(15)–176.60(15)° for O-Co1-N and 83.05(14)-175.63(14)° for O-Co1-O. The coordination environment of Co2 is similar to that of Co1, which is also coordinated by two N atoms from one H₂pyttz-I ligand, two O atoms from one TeMo6 polyoxoanion and two O atoms from two coordinated water molecules. It is noted that there is a separate $[Co(H_2O)_6]^{2+}$ unit as a counter cation, in which the Co3 ion is six-coordinated by six O atoms from six coordinated water molecules, showing a distorted octahedral environment. The bond lengths around the Co3 ion are 2.052(4)-2.139(4) Å for Co3-O, the bond angles are 85.11(16)-176.14(17)° for O-Co3-O.

The H_2 pyttz-I ligand in 2 also adopts a bidentate chelating coordination mode and only coordinates to a Co1 ion. It should be noted that the coordination mode of H_2 pyttz-I in 2 is a slightly different from that in complex 1 (Table 2). In compound 1, Co ion is coordinated by the nitrogen atoms of the pyridine and the 1-position nitrogen atom of triazole, while in compound 2 is the nitrogen atom of the pyridine and the 4-position nitrogen atom of triazole. In addition, each TeMo₆ anion exhibits a bidentate coordination mode through its four

Table 2 Coordination modes of the metal ions (Co^{II} or Ni^{II}), POM anions and pyridyl-bis(triazole) ligand in complexes 1−5 (the colors represent:

Complex	1	2	3	4	5
Metal ions	*	**	**	**	**
Organic ligands	000	000	M°	down.	M _o
POM anions	-		N N	N	N

terminal oxygen atoms to coordinate with two Co1 ions to form a binuclear complex $[\text{Co}_2(\text{H}_2\text{pyttz-I})_2(\text{H}_2\text{O})_4 \ [\text{TeMo}_6\text{O}_{24}]]^{2-}$ (Fig. 2b), which is different from **1**. The discrete binuclear units are linked together through the hydrogen bonding interactions between the oxygen atoms of the TeMo $_6$ polyoxoanions and the nitrogen atoms, carbon atoms of the H $_2$ pyttz-I ligands $[\text{N}(3)-\text{H}(3\text{A})\cdots\text{O}(7), 2.7170 \ \text{Å}], [\text{C}(18)-\text{H}(18\text{A})\cdots\text{O}(17) \ 3.0100 \ \text{Å}]$ to form a 1D supramolecular chain (Fig. 2c).

 $\{Co_3(Hpyttz-II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (3) and $\{Ni_3(Hpyttz-II)_2(H_2O)_6[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ II)₂(H₂O)₆[γ -Mo₈O₂₆]}·10H₂O (4). Complexes 3 and 4 are isostructural (Fig. S1†), so only the structure of 3 is described in detail. It should be noted that the CrMo₆ anions were in situ transformed to $[\gamma - Mo_8O_{26}]^{4-}$ $(\gamma - Mo_8O_{26})$ anions at pH = 5.8 in 3. However, in 4, the γ -Mo₈O₂₆ was constructed from the (NH₄)₆Mo₇O₂₄·4H₂O. The asymmetric unit of 3 consists of three CoII ions, two deprotonated Hpyttz-II ligands, one γ-Mo₈O₂₆ anion, six coordinated water molecules and ten lattice water molecules (Fig. 3a). There are two crystallographically independent CoII ions in 3. The Co1 ion is sixcoordinated by two O atoms from two coordinated water molecules and four N atoms from the triazole groups of two Hpyttz-II ligands. The bond lengths around Co1 ion are 2.053(5)-2.135(5) Å for Co1-O, 2.134(7)-2.152(7) Å for Co1-N and the bond angles are 87.4(2)-176.8(2)° for O-Co1-O, 169.6(2)° for N-Co1-N and 80.7(2)- 99.1(2)° for N-Co1-O. The Co2 ion is six-coordinated by two N atoms from the pyridyl and triazole groups of two Hpyttz-II ligands, two O atoms from two γ -Mo₈O₂₆ polyoxoanions and two O atoms from two coordinated water molecules. The bond lengths and angles around the Co2 ion are 2.084(6)-2.194(6) Å for Co2-N, 2.071(6) Å for Co2-O and the bond angles are 87.7(3)-92.3(3)° for O-Co2-N, 180.0(3)° for O-Co2-O and 76.7(2)-180.0(3)° for N-Co2-N, respectively.

Each Hpyttz-II ligand provides five nitrogen atoms, in which the four nitrogen atoms of the bis-triazole coordinate to two Co ions (Co1, Co2) and one Mo atom of $\gamma\text{-Mo}_8O_{26}$ and the pyridyl nitrogen atom coordinates to Co2 ion, respectively (Table 2). Thus the Hpyttz-II ligands connect adjacent Co1 and Co2 ions forming a 1D Co-Hpyttz-II metal–organic chain (Fig. 3c), which contains the [Co2(Hpyttz-II)2] metal–organic loop with dimension of 5.901 \times 9.773 Ų (Fig. 3b).

In 3, the γ -Mo $_8$ O $_{26}$ anions firstly provided two terminal oxygen atoms (O1) to coordinate with two Co2 ions from two adjacent 1D Co-Hpyttz-II chains, constructing a 2D network via Co2–O2 bonds (Fig. 3d). It should be noticed that a N-atom of the Hpyttz-II ligand coordinated to one γ -Mo $_8$ O $_{26}$ anion through Mo–N bond, which consolidate the 2D network.

 $\{Ni_3(Hpyttz-III)_2(H_2O)_8[\gamma-Mo_8O_{26}]\}\cdot 10H_2O$ (5). Complex 5 consists of three crystallographically independent Ni^{II} ions, two deprotonated Hppty-III ligands, one coordinated $[\gamma-Mo_8O_{26}]$ 4-anion, eight coordinated water molecules and ten lattice water molecules (Fig. 4a). In complex 5, the coordination modes of Ni1 and Ni2 display octahedral geometry.

Finally, each γ -Mo₈O₂₆ anion provided four terminal atoms (O1, O2) to coordinate with Co2 ions to form a Co-γ-Mo₈O₂₆ inorganic chain (Fig. 3e), which extended the adjacent 2D layers into a 3D framework (Fig. 3f). The Ni1 is surrounded by three O atoms of three coordination water molecules, one N atom from the triazole of Hppty-III ligand, two O atoms from two [γ-Mo₈O₂₆]4-anions. The bond lengths around Ni1 ion are 2.000(3)-2.082(2) Å for Ni1-O, 2.054(3) Å for Ni1-N and the bond angles are 79.03(12)-180.0(13)° for O-Ni1-O and 79.03(12)–180.0(13)° for N-Ni1-O. Nevertheless, the Ni2 atom is coordinated by two O atoms of two coordination water molecules, four N atoms from triazole groups of two H2ppty-III ligands, which is similar to that of Co1 in complex 3. The bond lengths around Ni2 ion are 2.092(3) Å for Ni2-O, 2.055(3)-2.057(3) Å for Ni2-N and the bond angles are 180.000(12)° for O-Ni2-O, 79.03(12)-180.00(12)° for N-Ni2-N and 86.68(13)-93.32(13)° for N-Ni2-O, respectively.

In 5, each $\gamma\text{-Mo}_8O_{26}$ anions provided four terminal oxygen atoms to coordinate with four Ni^{II} ions to form a $Ni\text{-}\gamma\text{-Mo}_8O_{26}$ inorganic chain (Fig. 4b). Each Hpyttz-III ligand provides four nitrogen atoms of the bis-triazole groups to coordinate with two Ni^{II} ions and one Mo atom, in which the two nitrogen atoms (N3 and N4) adopts a bidentate chelating coordination mode to coordinate with Ni2 ion, the other two nitrogen atoms (N1 and N2) coordinate with the Ni1 ion and one Mo atom of $\gamma\text{-Mo}_8O_{26}$ respectively. Different from the Hpyttz-II in 3 and 4, the pyridyl N atom of the Hpyttz-III is noncoordinated, thus two

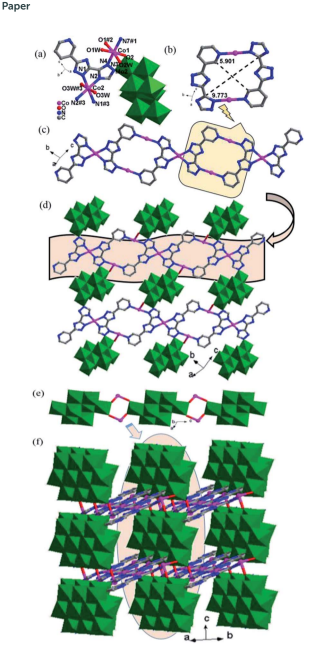


Fig. 3 (a) The coordination geometry of the Co^{II} ions in 3. The hydrogen atoms and lattice water molecules are omitted for clarity; symmetry code for 3: #1 – x + 2, –y + 2, –z + 1; #2 – x + 1, –y + 2, –z + 1; #3 – x + 1, –y + 1, –z + 1. (b) [Co₂(H₂pyttz-II)₂] loop in 3; (c) The 1D Co–H₂pyttz-II chain in 3; (d) view of the 2D layer in 3; (e) a view of Co- γ -Mo₈O₂₆ inorganic chain in complex 3; (f) the 3D Mo₈O₂₆-based metal–organic framework of 3.

Hpyttz-III ligands in 5 chelating a Ni2 ion, forming a [Ni(Hpyttz-III)₂] mononuclear unit (Fig. 4c), which link the adjacent 1D Ni- γ -Mo₈O₂₆ inorganic chains conducing to the formation of the 2D network (Fig. 4d). It is worthy of mentioning that there also exist a Mo–N bond, which consolidate the 2D network.

Influences of the POM anions type and the rigid pyridylbis(triazole) ligands on the architectures

The title complexes represent the novel examples of POM-based MOCs constructed from three rigid pyridyl-bis(triazole) ligands

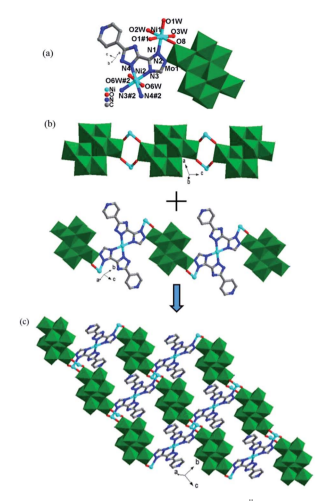
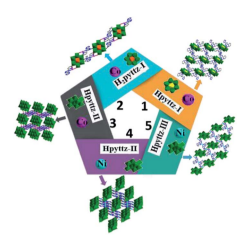


Fig. 4 (a) The coordination geometry of the Ni^{II} ions in 5. The hydrogen atoms and lattice water molecules are omitted for clarity; Symmetry codes for 5: #1 –x, –y, –z + 2; #2 –x, –y + 1, –z + 2. (b) A view of Ni- γ -Mo₈O₂₆ inorganic chain (up), the [Ni(Hpyttz-III)₂] unit and its connection with γ -Mo₈O₂₆ (down) in complex 5; (c) the 2D γ -Mo₈O₂₆-based 2D layer of 5.

and three different polyoxoanions, in which the polyanions and pyridyl-bis(triazole) ligands with different coordination sites show various coordination modes, exhibiting important effect on the whole structures (Table 2).

In 1, the A-type Anderson anion CrMo₆ was used, and each CrMo₆ polyoxoanion acts as a bidentate ligand to coordinate with two Co^{II} ions through its two terminal oxygen atoms, forming the binuclear unit. In 2, the B-type Anderson anion TeMo₆ was employed, which provided four terminal oxygen atoms to coordinate with two Co1 ions, constructing a binuclear complex $[\text{Co}_2(\text{H}_2\text{pyttz-I})_2(\text{H}_2\text{O})_4[\text{TeMo}_6\text{O}_{24}]]^{2-}$. In addition, the coordination modes of the rigid pyridyl-bis(triazole) ligands in complexes 1 and 2 are slightly different (Table 2). As a result, 1 is 2D supramolecular network, while 2 is a 1D supramolecular chain, which indicate that POMs' type show an obvious influence on the final structures. In complexes 3 and 4, each Hpyttz-II ligand provides five nitrogen atoms, in which the four nitrogen atoms of the bis-triazole coordinate to two Co^{II}/Ni^{II} ions and one Mo atom of γ -Mo₈O₂₆ and the pyridyl nitrogen

RSC Advances Paper



Scheme 2 Effect of POMs type and the rigid pyridyl-bis(triazole) ligands on the architectures.

atom coordinates to Co2/Ni2 ion, respectively. In complex 5, each Hpyttz-III ligand provides four nitrogen atoms from triazole groups, coordinating with Ni ions and one Mo atom of γ -Mo₈O₂₆. While the pyridyl N atom is noncoordinated. Thus, the Hpyttz-II and Hpyttz-III ligands with different coordination sites show completely different coordination modes, which finally result in various architectures: 3D frameworks for 3 and 4, 2D network for 5. Various structures of 1–5 confirmed the important effects of the rigid pyridyl-bis(triazole) ligands with different coordination sites (Scheme 2).

As discussed above, these results strongly suggest that the various coordination modes of the organic ligands and POM anions have assignable influences on the construction of the POM-based complexes. Therefore, an appropriate combination of different types of organic ligands and POMs may lead to the various structures with different dimensions.

IR spectra

The IR spectra of the title complexes are shown in Fig. S2.† The characteristic bands observed at 937, 906, 665 cm⁻¹ for 1, 944, 887, 668 cm⁻¹ for 2, 944, 843, 668 cm⁻¹ for 3, 949, 837, 662 cm⁻¹ for 4, 944, 856, 643 cm⁻¹ for 5 are attributed to ν (Mo–Ot), ν (Mo–Ob–Mo), ν (Mo–Oc–Mo), respectively.²⁶ Bands in the regions of 1657–1234 cm⁻¹ for 1, 1657–1056 cm⁻¹ for 2 and 1657–1025 cm⁻¹ for 3, 1657–1022 cm⁻¹ for 4 and 1632–1006 cm⁻¹ for 5 are attributed to the H₂pyttz-I, Hpyttz-II and Hppty-III, respectively.²⁷ The bands around 3400 cm⁻¹ can be ascribed to the water molecules.

Powder X-ray diffraction (PXRD) and thermal analyses

The powder X-ray diffraction (PXRD) patterns for complexes 1–5 are presented in the Fig. S3 in the ESI.† The diffraction peaks of both simulated and experimental patterns match well, thus indicating that the phase purities of the compounds are good. The intensity differences may be owed to the different orientations of the crystals in the powder samples.²⁸

Thermogravimetric analyses (TGA) of the title complexes were performed under flowing N_2 atmosphere with a heating

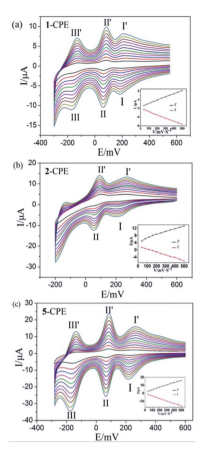


Fig. 5 Cyclic voltammograms of the (a) 1-CPE (b) 2-CPE (c) 5-CPE in 0.1 M $H_2SO_4 + 0.5$ M Na_2SO_4 aqueous solution under scan rates from inner to outer: (scan rates: 40, 80, 120, 160, 200, 250, 300, 350, 400, 450, 500 mV s⁻¹). Insert: the dependence of cathodic peak and anodic peak currents on scan rates of 1-, 2- and 5-CPEs.

rate of 10 °C min⁻¹ from room temperature to 800 °C (Fig. S4†). The TG curve of 1 shows two distinct weight loss steps. The first weight loss step from room temperature to 393 °C corresponds to the loss of water molecules 12.93% (calcd 13.90%). The second weight loss at 395-800 °C is ascribed to the decomposition of organic molecules 24.67% (calcd 25.25%). For complex 2, the TGA gives an initial loss of 11.77% (calcd 12.16%) below 228 °C, representing the loss of water molecules. The second weight loss occurs in the temperature range of 230-800 °C, corresponding to the loss of H₂pyttz-I ligands (calcd 22.14%, obsd 21.07%). For complex 3, the weight loss 13.23% attributed to the gradual release of water molecules is observed when the temperature is increased to 167 °C (calcd 13.89%), and the organic molecules decompose in the temperature range of 432 °C to 800 °C (calcd 19.77%, obsd 20.49%). For compound 4, the TG curve exhibits two steps of weight loss: the first loss is 13.33% from the room temperature to 273 °C corresponding to the loss of all water molecules (calcd 13.94%), and the second loss is 20.12% (calcd 19.46%) at 396-800 °C arising from the decomposition of the Hpyttz-II molecules. For 5, the first step starts from room temperature to 370 °C corresponding to the loss of lattice and coordination water molecules (calcd 10.83%, obsd 11.08%); and the second step begins from approximately

425 $^{\circ}$ C and is related to the decomposition of organic ligands (calcd 20.17%, obsd 22.38%).

Electrochemical behaviours of complexes 1-CPE, 2-CPE and 5-CPE

In order to investigate the electrochemical properties of these complexes, the title complexes bulk-modified CPEs are optimal choice, owing to their insolubility in water and classical solvents. Considering the electrochemical behaviors of complexes 3–5 are similar apart from some light potential shift, complexes 1, 2 and 5 were taken as the examples. The electrochemical behaviours of complex 1 bulk-modified carbon paste electrode (1-CPE) in 0.1 M $\rm H_2SO_4$ + 0.5 M $\rm Na_2SO_4$ aqueous solution at different scan rates are presented in Fig. 5a. The 1-CPE displays three pairs of reversible redox peaks in the potential range of +550 to -300 mV and the mean peak

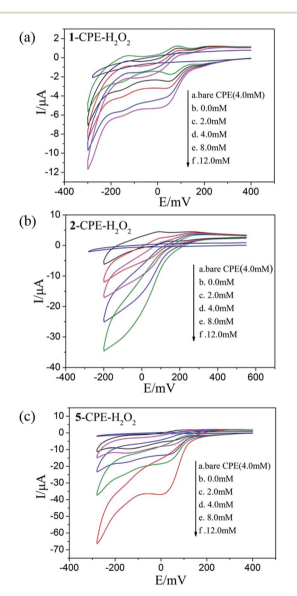


Fig. 6 Cyclic voltammograms of the 1-CPE, 2-CPE and 5-CPE in 0.1 M $\rm H_2SO_4\,+\,0.5\,$ M $\rm\,Na_2SO_4$ aqueous solution containing 0.0–12.0 mM $\rm\,H_2O_2.$

potentials $E_{1/2} = (E_{pa} + E_{pc})/2$ are 225 mV (I-I'), 73 mV (II-II') and -143 mV (III-III') at 40 mV s⁻¹, which belong to the three consecutive two-electron redox processes of Mo centres of CrMo₆ polyanions.²⁹ The 2-CPE displays two pairs of reversible redox peaks in the potential range of +600 to -200 mV (Fig. 5b), and the mean peak potentials $E_{1/2} = (E_{pa} + E_{pc})/2$ are 239 mV (I– I') and 79 mV (II–II') at 40 mV $\rm s^{-1}$, which correspond to two consecutive two-electron redox processes of Mo centres of TeMo₆ polyoxoanions in complex 2.30 The 5-CPE also displays three pairs of reversible redox peaks in the potential range of +600 to -300 mV and the mean peak potentials $E_{1/2} = (E_{pa} +$ E_{pc})/2 are 244 mV (I–I'), 76 mV (II–II') and -144 mV (III–III') at 40 mV s^{-1} (Fig. 5c), which can be attributed to three consecutive two-electron redox processes of Mo centers in the Mo₈O₂₆ polyoxoanion.31,32 The peak potentials change gradually with the increasing of scan rates. In the scan rate range of 40-500 mV s⁻¹, all the cathodic peak potentials shift toward the negative direction gradually, while the corresponding anodic peak potentials shift to the positive direction with the increase of the

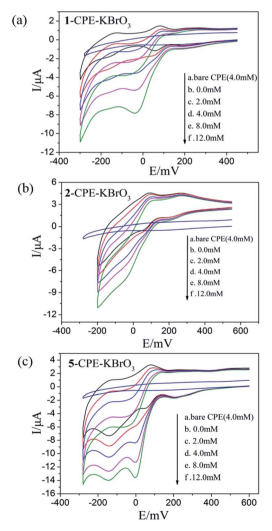


Fig. 7 Cyclic voltammograms of the 1-CPE, 2-CPE and 5-CPE in 0.1 M $H_2SO_4\,+\,0.5\,$ M $Na_2SO_4\,$ aqueous solution containing 0.0–12.0 mM $KBrO_3.$

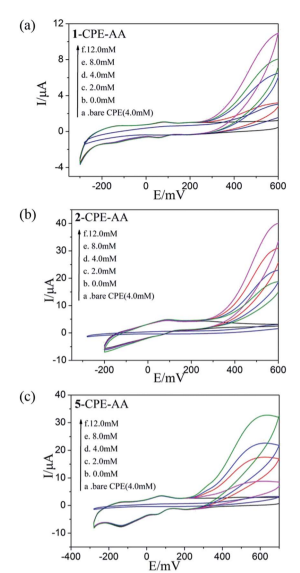


Fig. 8 Cyclic voltammograms of the 1-CPE, 2-CPE and 5-CPE in 0.1 M $H_2SO_4 + 0.5$ M Na_2SO_4 aqueous solution containing 0.0-12.0 mM AA.

scan rates. As shown in the inset of Fig. 5, the peak currents are proportional to the scan rates up to 500 mV s⁻¹, showing that the redox processes of 1-, 2-, and 5-CPEs are surface-controlled.

It is well known that some POMs usually show electrocatalytic activities in the reduction process of nitrite, bromate, hydrogen peroxide and other substances in aqueous solution.³¹ In this work, the electrocatalytic activities of the 1-, 2- and 5-CPEs toward the reduction of bromate/hydrogen peroxide and the oxidation of ascorbic acid (AA) were investigated.

As shown in Fig. 6a–c and 7a–c, with the addition of hydrogen peroxide or bromate, the reduction peak currents of 1-, 2- and 5-CPEs increase gradually and the corresponding oxidation peak currents gradually decrease, which indicate that the 1-, 2- and 5-CPEs possess good electrocatalytic activity toward the reduction of hydrogen peroxide and bromate and the reduction products are $\rm H_2O$ and $\rm Br^-$, respectively. 33,34 As can be seen from Fig. 8a–c, with the addition of AA, the oxidation peak currents of 1-, 2- and

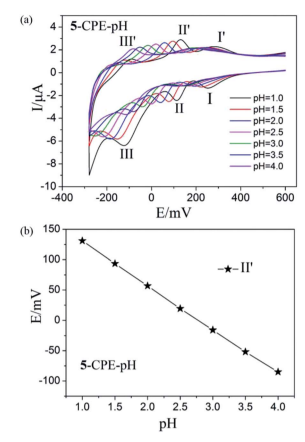


Fig. 9 (a) Cyclic voltammograms of 5-CPE at different pH values (Scan rate: 40 mV s^{-1}); (b) Variation of anodic peak potentials of the Mo^{VI}-based wave (II') with different pH for 5-CPE.

5-CPEs gradually increase, which suggests that 1-, 2- and 5-CPEs present excellent electrocatalytic activities for the oxidation of AA and the oxidation product is g-lactone.³⁵ Alternatively, no obvious voltammetric response is observed at the bare CPE in the presence of hydrogen peroxide, bromate or AA in the same potential range, suggesting that the reduction of bromate/hydrogen peroxide and the oxidation of AA is electrocatalyzed by the reduced species of the POM anions in the title complexes.

In order to probe the influence of the variation of acidity on the electrochemical response of the 5-CPE, H₂SO₄/Na₂SO₄ aqueous solution was used to adjust the pH values of the medium. The results indicated that electrochemical behavior of 5-CPE is pH dependent, as shown in Fig. 9. With the pH value increase from 1.0 to 4.0, the redox waves shift gradually to the negative potential direction and the peak current intensity gradually decrease. The 1- and 2-CPEs also show similar electrochemical responses (Fig. S5†). The successive redox processes versus pH for the 1-, 2- and 5-CPEs are shown in Fig. 9b and S6.† Good linearity in the pH range from 1.0 to 4.0 was obtained, which indicated that the 1-, 2- and 5-CPEs may be used as a kind of pH potential sensor. Such an electrochemical behaviors of 1-, 2-, and 5-CPEs is consistent with the previous reports, which highlights the significance of protons in the electrochemistry of POMs.36 The results indicate that the title

complexes may be used as pH sensor materials and have

Conclusions

potential application prospects.

Paper

In conclusion, we present five new Anderson-type POM and octamolybdate-based metal-organic complexes with different rigid pyridyl-bis(triazole) ligands, POMs ions and transition metals Co^{II}/Ni^{II}. The organic ligands with different coordination sites and the polyoxoanions with various coordination modes play important roles in tuning the dimensionalities and structures of the title complexes. Complexes 1–5 show highly electrochemical sensitive of pH in the range from 1.0 to 4.0. In addition, the title complexes exhibit excellently electrocatalytic activities for the reduction of BrO₃⁻/H₂O₂ and the oxidation of AA, which may be potential candidates as multifunctional materials.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The supports of the National Natural Science Foundation of China (No. 21471021, 21671025, 21501013 and 21401010), Program for Distinguished Professor of Liaoning Province (No. 2015399) and Key Laboratory of Polyoxometalate Science of Ministry of Education are gratefully acknowledged.

Notes and references

- 1 (a) D. L. Long, E. Burkholder and L. Cronin, Chem. Soc. Rev., 2007, 36, 105; (b) J. T. Rhule, C. L. Hill and D. A. Judd, Chem. Rev., 1998, 98, 327; (c) N. Mizuno and M. Misono, Chem. Rev., 1998, **98**, 199; (*d*) M. T. Pope and A. Müller, *Angew. Chem.*, *Int.* Ed. Engl., 1991, 30, 34; (e) Y. Hou, L. Zakharov and M. Nyman, J. Am. Chem. Soc., 2013, 135, 16651; (f) A. Merca, S. Garai, H. Bögge, E. Haupt, A. Ghosh, X. Lépez, J. Poblet, F. Averseng, M. Che and A. Müller, Angew. Chem., Int. Ed., 2013, 52, 11765; (g) Y. Zhu, P. C. Yin, F. P. Xiao, D. Li, E. Bitterlich, Z. C. Xiao, J. Zhang, J. Hao, T. B. Liu, Y. Wang and Y. G. Wei, J. Am. Chem. Soc., 2013, 135, 17155; (h) J. Zhou, J. W. Zhao, Q. Wei, J. Zhang and G. Y. Yang, J. Am. Chem. Soc., 2014, 136, 5065; (i) Z. G. Jiang, K. Shi, Y. M. Lin and Q. M. Wang, Chem. Commun., 2014, 50, 2353; (j) U. Kortz, A. Müller, J. vanSlageren, J. Schnack, N. S. Dalal and M. Dressel, Coord. Chem. Rev., 2009, 253, 2315.
- 2 (a) C. Qin, X. L. Wang, L. Yuan and E. B. Wang, Cryst. Growth Des., 2008, 8, 2093; (b) X. L. Wang, C. Qin, E. B. Wang and Z. M. Su, Chem. Commun., 2007, 41, 4245.
- 3 (a) X. J. Dui, W. B. Yang, X. Y. Wu, X. F. Kuang, J. Z. Liao,
 R. M. Yu and C. Z. Lu, *Dalton Trans.*, 2015, 44, 9496; (b)
 J. Z. Liao, H. L. Zhang, S. S. Wang, J. P. Yong, X. Y. Wu,
 R. M. Yu and C. Z. Lu, *Inorg. Chem.*, 2015, 54, 4345.
- 4 (a) Z. Peng, Angew. Chem., Int. Ed., 2004, 43, 930; (b) A. K. C. Gallegos, M. L. Cant'u, N. C. Pastor and

- P. G. Romero, Adv. Funct. Mater., 2005, 15, 1125; (c) M. Carraro, M. Gardan, G. Scorrano, E. Drioli, E. Fontananova and M. Bonchio, Chem. Commun., 2006, 0, 4533; (d) M. Bonchio, M. Carraro, M. Gardan, G. Scorrano, E. Drioli and E. Fontananova, Top. Catal., 2006, 40, 133; (e) J. M. C. Juan and E. Coronado, Coord. Chem. Rev., 1999, 193, 361; (f) H. Lv, Y. V. Geletii, C. Zhao, J. W. Vickers, G. Zhu, Z. Luo, J. Song, T. Lian, D. G. Musaev and C. L. Hill, Chem. Soc. Rev., 2012, 41, 7572.
- 5 (a) X. J. Yang, X. J. Feng, H. Q. Tan, H. Y. Zang, X. l. Wang, Y. H. Wang, E. B. Wang and Y. G. Li, J. Mater. Chem., 2016, 4, 3947; (b) Z. H. Pu, C. T. Zhang, I. S. Amiinu, W. Q. Li, L. Wu and S. C. Mu, Mater. Interfaces, 2017, 9, 16187.
- 6 (a) M. Sadakane and E. Steckhan, Chem. Rev., 1998, 98, 219;
 (b) S. Mukhopadhyay, J. Debgupta, C. Singh, A. Kar and S. K. Das, Angew. Chem., Int. Ed., 2018, 130, 1936.
- 7 C. J. Wang, T. T. Wang, Q. Lan, S. Yao, H. L. Wu, Y. Y. Zhou, Z. M. Zhang and E. B. Wang, RSC Adv., 2016, 6, 15513.
- 8 (a) X. Ma, K. F. Song, J. Gao, P. J. Gong, H. L. Li, L. J. Chen and J. W. Zhao, *Inorg. Chem. Commun.*, 2015, 60, 65; (b) J. Luo, X. Ma, L. J. Chen and J. W. Zhao, *Inorg. Chem. Commun.*, 2015, 54, 25.
- S. Q. Liu, D. G. Kurth and D. Volkmer, *Chem. Commun.*, 2002,
 976.
- 10 J. X. Meng, Y. Lu, Y. G. Li, H. Fu and E. B. Wang, *Cryst. Growth Des.*, 2009, 9, 4116.
- 11 P. P. Zhang, J. Peng, H. J. Pang, J. Q. Sha, M. Zhu, D. D. Wang, M. G. Liu and Z. M. Su, *Cryst. Growth Des.*, 2011, 11, 2736.
- 12 X. L. Wang, H. L. Hu and A. X. Tian, *Cryst. Growth Des.*, 2010, **10**, 4786.
- 13 B. K. Tripuramallu and S. K. Das, *Cryst. Growth Des.*, 2013, 13, 2426.
- 14 Y. Q. Lan, S. L. Li, X. L. Wang, K. Z. Shao, D. Y. Du, H. Y. Zang and Z. M. Su, *Inorg. Chem.*, 2008, **47**, 8179.
- 15 J. Q. Sha, M. T. Li, J. W. Sun, Y. N. Zhang, P. F. Yan and G. M. Li, *Dalton Trans.*, 2013, 42, 7803.
- 16 X. L. Wang, H. L. Hu, G. C. Liu, H. Y. Lin and A. X. Tian, Chem. Commun., 2010, 46, 6485.
- 17 X. L. Wang, J. J. Sun, H. Y. Lin, Z. H. Chang, A. X. Tian, G. C. Liu and X. Wang, *Dalton Trans.*, 2016, 45, 2709.
- 18 X. Y. Yang, T. Wei, J. S. Li, N. Sheng, P. P. Zhu, J. Q. Sha, T. Wang and Y. Q. Lan, *Inorg. Chem.*, 2017, **56**, 8311.
- 19 P. P. Zhu, N. Sheng, G. D. Liu, J. Q. Sha and X. Y. Yang, *Polyhedron*, 2017, **131**, 52.
- 20 X. L. Wang, J. Luan, F. F. Sui, H. Y. Lin, G. C. Liu and C. Xu, Cryst. Growth Des., 2013, 13, 3561.
- 21 X. L. Wang, J. J. Sun, H. Y. Lin, Z. H. Chang, G. C. Liu and X. Wang, *CrystEngComm*, 2017, **19**, 3167.
- 22 Y. Duan, J. M. Clemente-Juan, C. Giménez-Saiz and E. Coronado, *Inorg. Chem.*, 2016, **55**, 925.
- 23 L. Zhang, L. Liu, C. Huang, X. Han, L. Guo, H. Xu, H. W. Hou and Y. T. Fan, *Cryst. Growth Des.*, 2015, 15, 3426.
- 24 (a) Z. X. Zhang, T. Murayama, M. Sadakane, H. Ariga, N. Yasuda, N. Sakaguchi, K. Asakura and W. Ueda, *Nat. Commun.*, 2015, 6, 7731; (b) M. Filowitz, R. K. C. Ho, W. G. Klemperer and W. Shum, *Inorg. Chem.*, 1979, 18, 93;

RSC Advances

- (c) H. T. Evans, Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem., 1974, 30, 2095.
- 25 G. M. Sheldrick, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2008, **64**, 112.
- 26 C. R. Deltcheff, M. Fournier, R. Franck and R. Thouvenot, *Inorg. Chem.*, 1983, 22, 207.
- 27 P. P. Zhu, L. J. Sun, N. Sheng, J. Q. Sha, G. D. Liu, L. Yu, H. B. Qiu and S. X. Li, *Cryst. Growth Des.*, 2016, 16, 3215.
- 28 A. Z. Tian, J. Ying, J. Peng, J. Q. Sha, H. J. Pang, P. P. Zhang, Y. Chen, M. Zhu and Z. M. Su, *Cryst. Growth Des.*, 2008, 8, 3717.
- 29 C. H. Gong, X. H. Zeng, C. H. Zhu, J. H. Shu, P. X. Xiao, X. Hao, L. C. Liu, J. Y. Zhang, Q. D. Zeng and J. L. Xie, RSC Adv., 2016, 6, 106248.
- 30 S. M. Wang, W. L. Chen and E. B. Wang, *J. Cluster Sci.*, 2010, **21**, 133.

- 31 (a) A. Dolbecq, P. Mialane, B. Keita and L. Nadjo, J. Mater. Chem., 2012, 22, 24509; (b) P. Wang, X. P. Wang and G. Y. Zhu, Electrochim. Acta, 2000, 46, 637.
- 32 (a) P. Wang, X. P. Wang, Y. Yuan and G. Y. Zhu, J. Non-Cryst. Solids, 2000, 277, 22; (b) H. Y. Liu, H. Wu, J. F. Ma, Y. Y. Liu, J. Yang and J. C. Ma, Dalton Trans., 2011, 40, 602.
- 33 J. Taraszewska, G. Rosłonek and W. Darlewski, *J. Electroanal. Chem.*, 1994, 371, 223.
- 34 L. D. Li, W. J. Li, C. Q. Sun and L. S. Li, *Electroanalysis*, 2002, 14, 368.
- 35 P. Wang, X. P. Wang, X. Y. Jing and G. Y. Zhu, *Anal. Chim. Acta*, 2000, 424, 51.
- 36 M. Ammam and E. B. Easton, *Electrochim. Acta*, 2011, **56**, 2847.