



Using a flexible bis(pyrazol) ligand to construct four new Keggin-based compounds: syntheses, structures and properties†

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Using a flexible bis(pyrazol) ligand, four new compounds based on Keggin anions, namely $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{O})_2(\text{HPW}_{12}\text{O}_{40})]$ (1), $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{PW}_{12}\text{O}_{40})]_2 \cdot 2\text{H}_2\text{O}$ (2), $[\text{Cu}_2(\text{H}_2\text{bdpm})_3(\text{H}_2\text{O})_2(\text{SiW}_{12}\text{O}_{40})] \cdot 2\text{H}_2\text{O}$ (3), and $[\text{Cu}_2(\text{H}_2\text{bdpm})_4(\text{H}_2\text{O}) (\text{SiW}_{12}\text{O}_{40})] \cdot 16\text{H}_2\text{O}$ (4) (H_2bdpm = bis(3,5-dimethyl-1*H*-pyrazol-4-yl) methane), have been synthesized under hydrothermal conditions. Compound 1 is a zero dimension (0D) structure, showing a $[\text{Cu}(\text{H}_2\text{bdpm})_2]^{2+}$ mono-supporting $[\text{PW}_{12}\text{O}_{40}]^{3-}$ anion. Compound 2 shows a 1D supramolecular chain, with bi-supporting anions lined by discrete $[\text{PW}_{12}\text{O}_{40}]^{3-}$ anions. Compound 3 contains bi-nuclear Cu^{2+} cycles, which are linked by O_4W to form a cycle-connecting-cycle chain. The $[\text{SiW}_{12}\text{O}_{40}]^{4-}$ anions connect adjacent chains to construct a 2D structure. Compound 4 contains a nested cycle subunit $[\text{Cu}_4(\text{H}_2\text{bdpm})_6]^{8+}$, with two bi-nuclear cycles $[\text{Cu}_2(\text{H}_2\text{bdpm})_2]^{4+}$ linked by two H_2bdpm ligands. The nested cycle subunits are connected by H_2bdpm molecules to construct a 2D metal-organic layer. The Keggin anions link adjacent layers to form a 3D framework. The electrochemical and photocatalytic properties of compounds 1–4 are studied.

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Introduction

Recently, polyoxometalates (POMs) have attracted attention as a branch of inorganic chemistry, not only due to their diverse structures,¹ but also for their extensive applications in catalysis, medicine, magnetism, photochemistry and other fields.² POMs can provide abundant O donors to connect transition metal complexes (TMCs) to construct fascinating structures,³ which has become a booming field of POMs.

Among a large number of POM-TMCs, using cycle-connecting-cycle and nested cycle subunits to modify POMs is relatively rare.⁴ In the synthetic strategy of forming this series, the choice of appropriate organic ligands is essential and important. Observed from reports, flexible organic ligands can construct molecular cycles with ease, containing cycle-connecting-cycle and nested cycle subunits.⁵ The flexible ligands own conformational freedom and can rotate in some extent, conducting to fuse TM ions and construct cycle structures. In our previous work, we have utilized a series of symmetric flexible bis(triazole) ligands to build POM-based cycle structures.⁶ For example, by using 1,3-bis(1,2,4-triazol-1-yl)propane (btp) we have obtained a mono-substituted

Keggin-based tetra-nuclear metallamacrocyclic structure.⁷ Thus, the usage of symmetric flexible ligands to build POM-based cycle structures is rational and challenging. In this work, we utilize a flexible bis(pyrazole) ligand bis(3,5-dimethyl-1*H*-pyrazol-4-yl)methane (H_2bdpm) instead of bis(triazole) ligands to explore whether H_2bdpm can still build POM-based cycle subunits. The selection of H_2bdpm owns three advantages: (i) it has four N donors to link TM ions, inducing strong coordination capacity. (ii) The $-\text{CH}_2-$ group increases flexibility of ligand skeleton, conducting to rotate and fuse TM ions. (iii) It has four $-\text{CH}_3$ groups, which can increase steric hindrance to avoid interpenetrating structures and induce cycle structures with ease.⁸ Thus, the selection of H_2bdpm is a reasonable method to build metal-organic cycle modified POM compounds.

Hydrothermal technique has been proven an effective method to construct POM-TMC compounds.⁹ Commonly, in one-pot under hydrothermal conditions only one type of crystal could be obtained. However, two or more types of crystals obtained in one-pot are rarely observed.¹⁰ Namely, high efficient usage of hydrothermal technique to construct two or more POM-based compounds is challenging and appealing. In this work, we try to tune some influencing factors under hydrothermal conditions, aiming to form more crystals in one-pot.

Herein, by introducing H_2bdpm to Keggin- Cu^{2+} system, four new compounds were obtained, $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{O})_2(\text{HPW}_{12}\text{O}_{40})]$ (1), $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{PW}_{12}\text{O}_{40})]_2 \cdot 2\text{H}_2\text{O}$ (2), $[\text{Cu}_2(\text{H}_2\text{bdpm})_3(\text{H}_2\text{O})_2(\text{SiW}_{12}\text{O}_{40})] \cdot 2\text{H}_2\text{O}$ (3) and $[\text{Cu}_2(\text{H}_2\text{bdpm})_4(\text{H}_2\text{O}) (\text{SiW}_{12}\text{O}_{40})] \cdot 16\text{H}_2\text{O}$ (4). It is worth mentioning that compounds 3 and 4 were

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obtained in one pot. We also studied the electrochemical and photocatalytic properties of compounds **1–4**.

Experimental

Materials and methods

The reagents and solvents for syntheses were all purchased from commercial sources and used without further purification. Elemental analyses (C, H, and N) were achieved with a Perkin-Elmer 240C elemental analyzer, and the FT-IR spectra were taken on a Varian FT-IR 640 spectrometer (KBr pellets). UV-vis absorption spectra were obtained using a Lambda 750 UV/VIS/NIR spectrophotometer. Electrochemical measurements and data collection were performed with a CHI 440 electrochemical workstation. A conventional three-electrode system was used with a saturated calomel electrode (SCE) as reference electrode and a Pt wire as counter electrode. The title compounds modified carbon paste electrodes (CPEs) were used as the working electrodes.

X-ray crystallography

X-ray diffraction analysis data for compounds **1–4** were collected with a Bruker Smart Apex CCD diffractometer (Bruker Corporation, Germany) with Mo-K α ($\lambda = 0.71073$ Å) at 293 K. The structures were solved by direct methods and refined on F^2 by full-matrix least squares methods using the SHELXTL package.¹¹ For the compounds, all the hydrogen atoms attached to carbon atoms were generated geometrically. Furthermore, the hydrogen atoms attached to water molecules were not located but were included in the structure factor calculations. In compound **4**, the seventeen water molecules were highly disordered and could not be modeled properly, thus the SQUEEZE routine of PLATON was applied to remove the contributions to the scattering from the solvent molecules. The reported refinements are of the guest-free structures using the *.hkl files produced by using the SQUEEZE routine.¹² All crystallographic data and structural determination is provided in

Table 1. The data of selected bond lengths and angles of the title compounds are listed in Table S1.[†] Crystallographic data for the structures reported in this paper have been deposited in the Cambridge Crystallographic Data Center with CCDC Numbers of 1486618, 1486619, 1486620 and 1509083 for **1–4**.

Preparation of compounds **1–4**

Synthesis of $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{O})_2(\text{HPW}_{12}\text{O}_{40})]$ (1). A mixture of $\text{H}_3\text{PW}_{12}\text{O}_{40}$ (100 mg), CuCl_2 (80 mg), H_2bdpm (20 mg) and H_2O (10 mL) was stirred for 40 min in air at room temperature. The pH value was adjusted to about 2.5 with 1 M HCl and then the suspension was transferred to a 25 mL Teflon lined autoclave and kept at 160 °C for 5 days. After slow cooling to room temperature, green clavate crystals of **1** were obtained (final pH = 3.3). Elemental analysis (%) calc. for $\text{C}_{22}\text{H}_{37}\text{CuN}_8\text{O}_{42}\text{PW}_{12}$ (3386): C 7.79, H 1.09, N 3.30. Found: C 7.87, H 1.03, N 3.36. IR(KBr pellet, cm^{-1}): 3444(w), 1633(w), 1527(w), 1473(w), 1417(w), 1375(w), 1288(w), 1176(w), 970(m), 898(m), 819(s).

Synthesis of $[\text{Cu}(\text{H}_2\text{bdpm})_2(\text{H}_2\text{PW}_{12}\text{O}_{40})]_2 \cdot 2\text{H}_2\text{O}$ (2). A mixture of $\text{H}_3\text{PW}_{12}\text{O}_{40}$ (100 mg), CuCl_2 (80 mg), H_2bdpm (20 mg), H_2O (8 mL) and ethanol (2 mL) was stirred for 40 min in air at room temperature. The pH value was adjusted to about 2.5 with 1 M HCl and then the suspension was transferred to a 25 mL Teflon lined autoclave and kept at 160 °C for 5 days. After slow cooling to room temperature, yellow clavate crystals of **2** were obtained (final pH = 3.5). Elemental analysis (%) calc. for $\text{C}_{44}\text{H}_{72}\text{Cu}_2\text{N}_{16}\text{O}_{82}\text{P}_2\text{W}_{24}$ (6738): C 7.79, H 1.09, N 3.36. Found: C 7.84, H 1.05, N 3.31. IR(KBr pellet, cm^{-1}): 3432(m), 1731(m), 1646(m), 1581(m), 1546(s), 1500(w), 1463(w), 1398(w), 1346(w), 1247(m), 1081(m), 973(m), 806(s), 896(m).

Synthesis of $[\text{Cu}_2(\text{H}_2\text{bdpm})_3(\text{H}_2\text{O})_2(\text{SiW}_{12}\text{O}_{40})] \cdot 2\text{H}_2\text{O}$ (3) and $[\text{Cu}_2(\text{H}_2\text{bdpm})_4(\text{H}_2\text{O}) \cdot (\text{SiW}_{12}\text{O}_{40})] \cdot 16\text{H}_2\text{O}$ (4). A mixture of $\text{H}_4\text{SiW}_{12}\text{O}_{40}$ (100 mg), CuCl_2 (76 mg), H_2bdpm (20 mg) and H_2O (10 mL) was stirred for 40 min in air at room temperature. The pH value was adjusted to about 2.4 with 1 M HCl, then suspension was transferred to a 25 mL Teflon lined autoclave and kept at 160 °C for 5 days (final pH = 3.0). After slow cooling

Table 1 Crystal data and structure refinement for compounds **1–4**

Compounds	1	2	3	4
Formula	$\text{C}_{22}\text{H}_{37}\text{CuN}_8\text{O}_{42}\text{PW}_{12}$	$\text{C}_{44}\text{H}_{72}\text{Cu}_2\text{N}_{16}\text{O}_{82}\text{P}_2\text{W}_{24}$	$\text{C}_{33}\text{H}_{56}\text{Cu}_2\text{N}_{12}\text{O}_{44}\text{SiW}_{12}$	$\text{C}_{44}\text{H}_{98}\text{Cu}_2\text{N}_{16}\text{O}_{57}\text{SiW}_{12}$
F_w	3386	6738	3686	4124
Crystal system	Monoclinic	Triclinic	Monoclinic	Monoclinic
Space group	$P2_1/c$	$\bar{P}1$	$P2_1/n$	$C2/c$
a (Å)	11.3926(10)	12.444(2)	15.2377(13)	35.318(5)
b (Å)	28.008(2)	14.557(3)	18.6736(16)	23.532(5)
c (Å)	17.0586(14)	17.393(3)	25.291(2)	23.250(5)
β (°)	95.775(2)	105.644(4)	97.928(2)	117.176(5)
V (Å 3)	5415.4(8)	2947.5(10)	7127.7(11)	17 190(6)
Z	4	1	4	8
D_c (g cm $^{-3}$)	4.147	3.792	3.426	2.945
μ (mm $^{-1}$)	25.895	23.787	19.975	16.569
$F(000)$	5952	2960	6560	13 696
R_1^a [$I > 2\sigma(I)$]	0.0435	0.0705	0.0402	0.0487
wR_2^b (all data)	0.0896	0.2172	0.0806	0.1284
GOF on F^2	0.994	1.015	0.987	1.016

^a $R_1 = \sum ||F_0| - |F_c|| / \sum |F_0|$. ^b $wR_2 = \{\sum [w(F_0^2 - F_c^2)^2] / \sum [w(F_0^2)^2]\}^{1/2}$.



to room temperature, both blue clavate crystals of **3** and green clavate crystals of **4** were obtained. Elemental analysis (%) calc. for $C_{33}H_{56}Cu_2N_{12}O_{44}SiW_{12}$ (3686, **3**): C 10.75, H 1.53, N 4.56. Found: C 10.83, H 1.47, N 4.64. Elemental analysis (%) calc. for $C_{44}H_{98}Cu_2N_{16}O_{57}SiW_{12}$ (4124, **4**): C 12.81, H 2.38, N 5.43. Found: C 12.72, H 2.45, N 5.37. IR (KBr pellet, cm^{-1}) for **3**: 3399(m), 1731(m), 1647(m), 1581(m), 1548(s), 1500(w), 1463(w), 1400(w), 1348(w), 1247(m), 1149(w), 1078(w), 970(m), 919(s), 792(m), 528(m). IR(KBr pellet, cm^{-1}) for **4**: 3440(w), 1731(m), 1646 (m), 1581(m), 1548(s), 1500(w), 1463(w), 1398(w), 1346(w), 1247(m), 1079(w), 970(m), 919(s), 798(s), 526(m).

Preparation of 1- to 4-CPEs. Compound **1** modified CPE (1-CPE) was fabricated as follows: 0.10 g of graphite powder and 0.01 g of compound **1** were mixed and ground together by an agate mortar for approximately 40 min to achieve a uniform mixture. Then with stirring 0.15 mL of Nujol was added. The homogenized mixture was packed into a glass tube with a 1.5 mm inner diameter. The tube surface was wiped with weighing paper. The electrical contact was established with a copper rod through the back of the electrode. In a similar manner, **2**, **3** and **4**-CPEs were made with compounds **2**, **3** and **4**.

Results and discussion

Syntheses

Under hydrothermal conditions, many factors can influence the final structures, such as temperature, reaction time, reactants and pH.⁹ In this work, we try to explore the effect of reactants. In PW_{12} - Cu^{2+} - H_2bdpm system, we obtained a discrete mono-supporting structure of **1** and a supramolecular structure of **2**. However, when we changed POM precursor PW_{12} to SiW_{12} , two distinct compounds **3** and **4** were synthesized in one pot. So the POM precursors own influence on the structures. Furthermore, highly efficient usage of hydrothermal technique is interesting to obtain two or more crystals in one pot. Compounds **3** and **4** were obtained in one pot, which can support informative examples for highly efficient usage of hydrothermal technique.

Structural description

Crystal structure of compound 1. Crystal structure analysis reveals that compound **1** is composed of one Cu^{2+} ion, two H_2bdpm molecules, one $[PW_{12}O_{40}]^{3-}$ (abbreviated to PW_{12}) anion and two coordinated water molecules ($O1W$ and $O2W$). The valence sum calculations show that all the W atoms are in the +VI oxidation states,¹³ all the Cu atoms are in the +II oxidation states. In order to balance the charge, one proton has been added in the formula of **1**.

Compound **1** owns only one crystallographically independent Cu ion. The Cu1 is coordinated by two N donors (N2 and N5) from two H_2bdpm ligands, one O atom from a PW_{12} anion and two coordinated water molecules ($O1W$ and $O2W$). The Cu-O distances are in the range of 1.942(9)–2.553(10) Å, while the Cu-N distances range from 1.924(11) to 1.955(12) Å. The O-Cu-N angles are in the range of 90.7(4)–174.5(5)°, while the N-N-Cu angles are from 123.8(9) to 125.3(9)°.

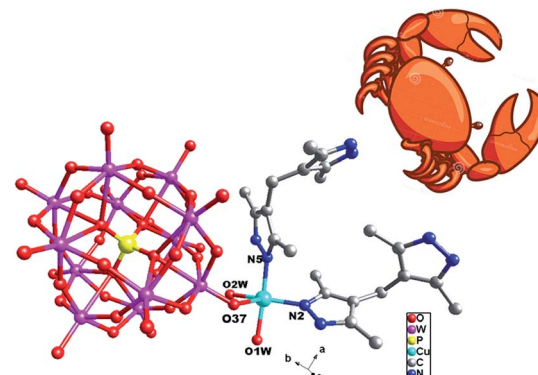


Fig. 1 Stick/ball view of the asymmetric unit of **1**. All H atoms are omitted for clarity.

In compound **1**, the H_2bdpm ligand only offers its one N atom to coordinate with one Cu^{2+} ion. Two H_2bdpm molecules are fused by one Cu1 atom to form a $[Cu(H_2bdpm)]^{2+}$ subunit, which links a PW_{12} anion through Cu1-O37 bond to generate a mono-supporting Keggin structure (Fig. 1). This mono-supporting anion is just like a “crab” and two H_2bdpm is just like its “pincers”. The adjacent mono-supporting anions further connect each other through hydrogen bonding interactions ($O31\cdots O34 = 3.238$ Å) to form a 1D supramolecular chain (Fig. S1†). The adjacent 1D supramolecular chains are still further linked by hydrogen bonding interactions ($O1W\cdots O24 = 2.80$ Å) to construct a 2D supramolecular layer (Fig. 2).

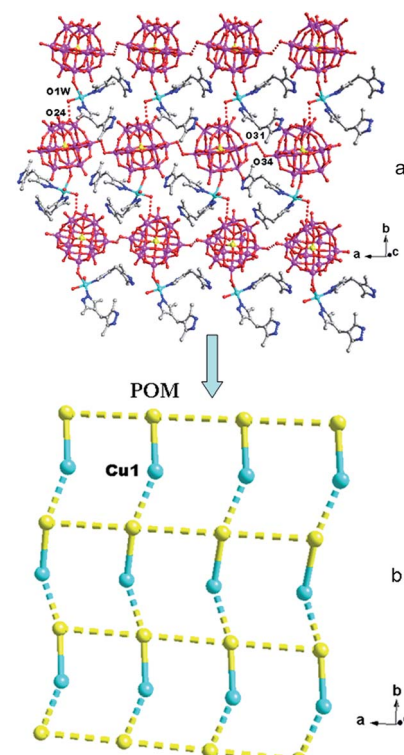


Fig. 2 The 2D supramolecular layer (a) and its schematic view (b) of **1**.



Crystal structure of compound 2. Crystal structure analysis reveals that compound 2 consists of two Cu^{2+} ions, four H_2bdpm molecules, two PW_{12} (POM1 and POM2) anions and two lattice water molecules, as shown in Fig. 3. There are two kinds of PW_{12} anions: bi-supporting anions (POM1) and discrete anions (POM2). The central P atom in PW_{12} anion is surrounded by eight oxygen atoms with each site half occupied, which is commonly observed in Keggin-based structures. The valence sum calculations show that all the W atoms are in the +VI oxidation states,¹³ all the Cu atoms are in the +I oxidation states. In order to balance the charge, two protons have been added in the formula of 2.

In compound 2, there is only one crystallographically independent Cu1 ion. Considering the long-range coordinative Cu–O bonds, the Cu1 atom adopts a “T”-type geometry, coordinated by two N atoms (N1 and N5) from two H_2bdpm ligands and one O atom from one POM1. The Cu–N distances are in the range of 1.86(2)–1.87(2) Å, while the Cu–O distance is 2.93(2) Å. The N–Cu–N angles are in the range of 123.1(18)–124.0(16)°, while the O–Cu–O angles are from 54.5(6)–146.1(6)° (Table S1†).

In compound 2, there are two different building blocks: (i) a POM1 anion links two $[\text{Cu}(\text{H}_2\text{bdpm})_2]$ segments to form a bi-supporting anion, like alphabet “H”. (ii) A discrete anion POM2. The two building blocks connect each other through hydrogen bonding interactions ($\text{N}2\cdots\text{O}42 = 3.216$ Å) to construct a 1D chain, as shown in Fig. 4.

Crystal structure of compound 3. Single-crystal X-ray diffraction analysis reveals that compound 3 consists of two Cu^{2+} ions, three H_2bdpm molecules, one $[\text{SiW}_{12}\text{O}_{40}]^{4-}$ (abbreviated to SiW_{12}) anion, two coordinated and two lattice water molecules, as shown in Fig. 5. The valence sum calculations show that all the W atoms are in the +VI oxidation states,¹³ all the Cu atoms are in the +II oxidation states.

In compound 3, there are two crystallographically independent Cu1 and Cu2 ions. Both Cu1 and Cu2 ions are five-coordinated with distorted square pyramid geometry, having τ value of 0.34 for Cu1 and 0.12 for Cu2 ($\tau = (\beta - \alpha)/60$). For Cu1 the angles are $\text{N}2-\text{Cu}1-\text{O}3\text{W} = 178^\circ$ (β) and $\text{N}12-\text{Cu}1-\text{O}4\text{W} = 156^\circ$ (α), while the angles for Cu2 are $\text{N}3-\text{Cu}2-\text{O}4\text{W} = 177^\circ$ (β) and $\text{N}7-\text{Cu}2-\text{N}9 = 170^\circ$ (α). The Cu1 ion is coordinated by two

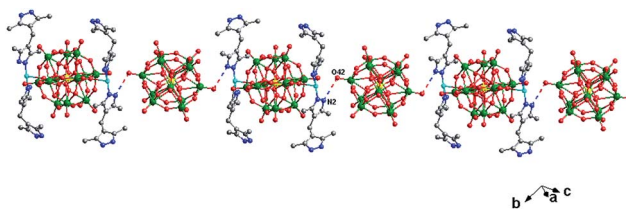


Fig. 4 The 1D supramolecular chain of 2.

nitrogen atoms (N2 and N12) from two H_2bdpm molecules, one terminal O37 atom from a SiW_{12} anion and two coordinated water molecules (O3W and O4W). The $\text{Cu}2$ ion is coordinated by three N atoms (N3, N7 and N9) from three H_2bdpm ligands, one terminal O35 atom from a SiW_{12} anion and one coordinated water molecule O4W. The Cu–O distances are in the range of 1.911(8)–2.451(10) Å, while the Cu–N distances range from 1.960(10) to 2.029(10) Å. The O–Cu–N angles are in the range of 87.4(4)–178.0(4)°, while the N–Cu–N angles are from 89.8(4) to 170.9(4)°.

In compound 3, the H_2bdpm shows two kinds of coordination modes: offering only one N donor to link one Cu^{2+} ion (type-I) and providing two N atoms from two dimethylpyrazole groups respectively to connect two Cu^{2+} ions (type-II). Two type-II H_2bdpm are fused by two Cu^{2+} ions and a bi-nuclear $[\text{Cu}_2(\text{H}_2\text{bdpm})_2]^{4+}$ cycle is built. The adjacent bi-nuclear cycles share the same O4W to construct a cycle-connecting-cycle chain (Fig. 6b). Furthermore, the type-I H_2bdpm hang this chain up and down by linking $\text{Cu}2$ ions (Fig. S2†). There also exists an inorganic chain constructed from SiW_{12} anions linked by $\text{Cu}1-\text{O}4\text{W}-\text{Cu}2$ units (Fig. 6a). The cycle-connecting-cycle metal-organic chains and POM-based inorganic connect each other alternately by sharing Cu ions to generate a 2D grid-like layer (Fig. 6c and d).

Crystal structure of compound 4. Single-crystal X-ray diffraction analysis reveals that the asymmetric unit of 4 is composed of two Cu^{2+} ions, four H_2bdpm ligands, one SiW_{12} anion, one coordinated and seventeen lattice water molecules (Fig. 7). The valence sum calculations show that all the W atoms

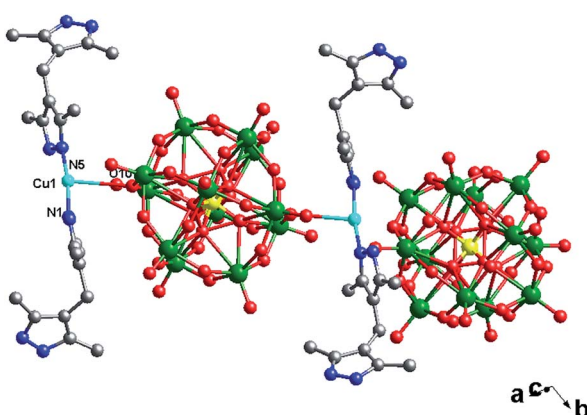


Fig. 3 Ball/stick view of the asymmetric unit of compound 2. The hydrogen atoms and lattice water molecules are omitted for clarity.

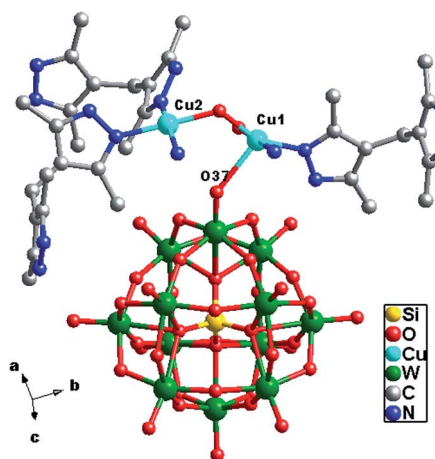


Fig. 5 Ball/stick view of the asymmetric unit of compound 3. The hydrogen atoms and lattice water molecules are omitted for clarity.



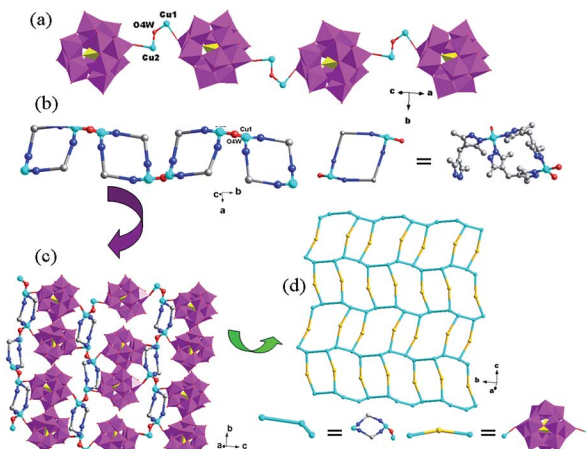


Fig. 6 (a) The 1D inorganic chain in **3** with SiW_{12} anions linked by $\text{Cu1}-\text{O4W}-\text{Cu2}$ units. (b) The 1D cycle-connecting-cycle metal-organic chain (some C atoms are omitted for clarity). (c) The 2D grid-like layer with inorganic chains and cycle-connecting-cycle metal-organic chains linking each other. (d) The schematic diagram of the 2D layer.

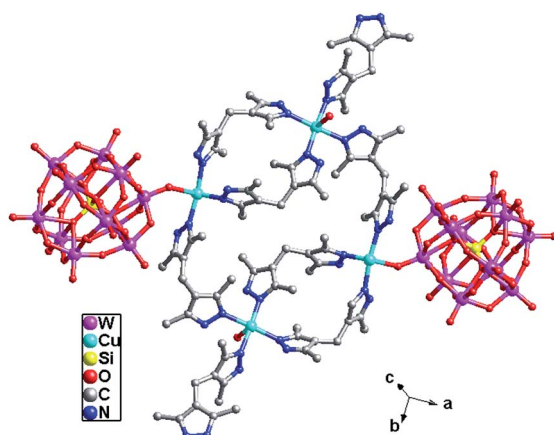


Fig. 7 Ball/stick view of the symmetric unit of compound **4**. The hydrogen atoms are omitted for clarity.

are in +VI oxidation state,¹³ all the Cu atoms are in +II oxidation state. In compound **4**, there are two crystallographically independent Cu1 and Cu2 ions. The Cu1 is six-coordinated with four N donors (N4, N6, N10 and N12) from four H_2bdpm ligands, one terminal O40 from one anion and one O1W molecule, showing an octahedral geometry. The Cu2 exhibits a distorted square pyramid geometry, having τ value of 0.26 ($\text{N}2-\text{Cu1}-\text{O}3\text{W} = 178^\circ$ (β) and $\text{N}12-\text{Cu1}-\text{O}4\text{W} = 156^\circ$ (α)). The Cu2 is coordinated with four N atoms (N1, N7, N14 and N15) from four H_2bdpm ligands and one terminal O30 from one anion. The Cu–O distances are in the range of 2.332(10)–2.601(10) Å, while the Cu–N distances range from 1.978(12) to 2.036(11) Å. The O–Cu–N angles are in the range of 86.7(4)–113.2(5)°, while the N–Cu–N angles are from 87.3(5) to 178.1(5)°.

In compound **4**, the H_2bdpm provides two N atoms from two dimethylpyrazole groups respectively to link two Cu^{2+} ions, which shows two functions: linking mode (type-A) and

cyclization mode (type-B) (Fig. S3†). Two type-B H_2bdpm molecules are fused by two Cu^{2+} ions to form a bi-nuclear $[\text{Cu}_2(\text{H}_2\text{bdpm})_2]^{4+}$ cycle. Two bi-nuclear cycles are further linked by two another type-B H_2bdpm molecules and a new tetra-nuclear cycle is formed (Fig. S4†). The $[\text{Cu}_4(\text{H}_2\text{bdpm})_6]^{8+}$ can be viewed as a nested cycle subunit. Furthermore, the type-A H_2bdpm offers two N donors to connect adjacent nested cycle subunits and a 2D metal-organic layer is generated (Fig. 8a and b). The POM anions link adjacent layers covalently to construct a 3D framework of **4** (Fig. 8c and d).

FT-IR spectra and powder X-ray diffractions

The IR spectra of compounds **1**–**4** are shown in Fig. S5†. The characteristic bands at 970(m), 898(m), 809(s) and 1072(m) cm^{-1} for **1** are attributed to $\nu(\text{W}-\text{O}_d)$, $\nu(\text{W}-\text{O}_b-\text{W})$, $\nu(\text{W}-\text{O}_c-\text{W})$ and $\nu(\text{P}-\text{O})$.¹⁴ The characteristic bands at 973(m), 806(s), 896(m) and 1081(m) cm^{-1} for **2** are attributed to $\nu(\text{W}-\text{O}_d)$, $\nu(\text{W}-\text{O}_b-\text{W})$, $\nu(\text{W}-\text{O}_c-\text{W})$ and $\nu(\text{P}-\text{O})$. In the spectra of **3** and **4**, characteristic bands 970(m), 919(s), 792(m), 528(m), cm^{-1} for **3**, 970(m), 919(s), 798(s), 526(m) cm^{-1} for **4**, are attributed to $\nu(\text{W}-\text{O}_d)$, $\nu(\text{Si}-\text{O})$ and $\nu(\text{W}-\text{O}_c-\text{W})$.¹⁵ The bands in the region of 1633–1176 cm^{-1} for **1**, 1731–1247 cm^{-1} for **2**, 1731–1078 cm^{-1} for **3**, 1731–1079 cm^{-1} for **4** can be attributed to the H_2bdpm ligand.¹⁶

Fig. S6† shows the powder X-ray diffraction patterns for compounds **1**–**4**. The diffraction peaks of both simulated and experimental patterns match well in positions, thus showing that the phase purities of the compounds **1**–**4** are good.

Electrochemical properties

We have investigated the electrochemical properties of compounds **1**–**4**. Owing to the similar electrochemical behaviors of compounds **1** and **2**, **3** and **4** modified carbon paste electrodes, the 1- and 3-CPEs have been taken as examples to study their electrochemical properties. The cyclic

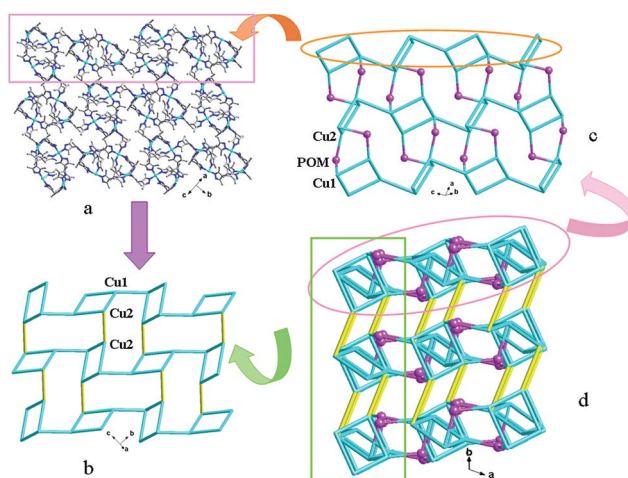


Fig. 8 (a) View of the 2D metal-organics layer of **4** with tetra-nuclear cycles linked by type-A H_2bdpm . (b) The schematic view of the 2D layer. (c) The details of the POM linking mode. (d) The 3D framework with adjacent 2D layers connected by anions.



voltammograms for **1**- and **3**-CPEs in 0.1 M H_2SO_4 + 0.5 M Na_2SO_4 aqueous solution at different scan rates are presented in Fig. 9. In the range of -850 to $+800$ mV for **1**-CPE, there are three reversible redox peaks $\text{II}-\text{II}'$, $\text{III}-\text{III}'$ and $\text{IV}-\text{IV}'$, with the half-wave potentials at $-171(\text{II}-\text{II}')$, $-464(\text{III}-\text{III}')$ and $-756(\text{IV}-\text{IV}')$ mV (scan rate: 100 mV s^{-1}). These three redox peaks should be ascribed to two consecutive one-electron and one two-electron process of PW_{12} .¹⁷ There exists one irreversible anodic peak I' with the potential of $+179$ mV assigned to the oxidation of the copper centers.^{6a} The **3**-CPE shows three reversible redox peaks $\text{II}-\text{II}'$, $\text{III}-\text{III}'$ and $\text{IV}-\text{IV}'$ in the potential range of -850 to $+800$ mV, with the half-wave potentials at $-200(\text{II}-\text{II}')$, $-463(\text{III}-\text{III}')$, $-754(\text{IV}-\text{IV}')$ mV (scan rate: 100 mV s^{-1}). Redox peaks $\text{II}-\text{II}'$ and $\text{III}-\text{III}'$ correspond to two consecutive one-electron processes of W centers, while $\text{IV}-\text{IV}'$ corresponds to a two-electron process.^{6a} The irreversible anodic peak I' with the potential of $+85$ mV is also assigned to the oxidation of the copper centers. With the scan rates from 40 to 500 mV s^{-1} , the peak potentials change gradually: the cathodic peak potentials shift towards the negative direction and the corresponding anodic peak potentials to the positive direction with increasing scan rates. Up to 500 mV s^{-1} , the peak currents are proportional to the scan rates (Fig. S7†), indicating that the redox processes of **1**- and **3**-CPEs are surface-confined.

Fig. 10 shows cyclic voltammograms for the electrocatalytic reduction of nitrite and bromate at **1**- and **3**-CPE in 0.1 M H_2SO_4 + 0.5 M Na_2SO_4 aqueous solution. With the addition of bromate, the second and third reduction peak currents increase gradually, while the corresponding oxidation peak currents gradually

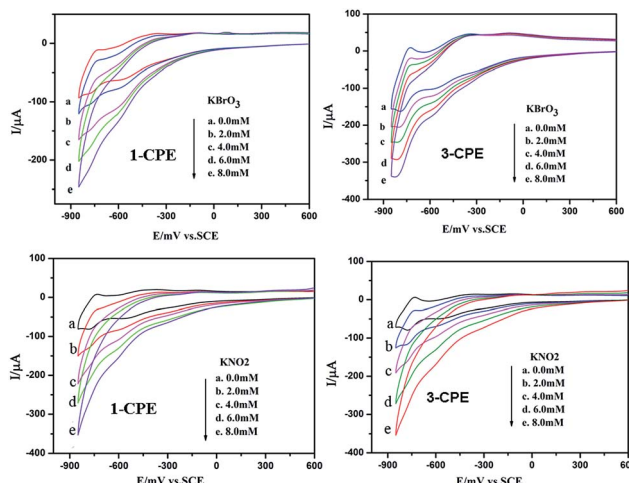


Fig. 10 Cyclic voltammograms of the **1**- and **3**-CPEs in $0.1 \text{ M H}_2\text{SO}_4$ + $0.5 \text{ M Na}_2\text{SO}_4$ aqueous solution containing 0.0 – 8.0 mM KBrO_3 and KNO_2 , respectively. Scan rate: 100 mV s^{-1} .

decrease. However, the first redox peak remains almost unchanged. Furthermore, it can be clearly seen that with adding nitrite, all three reduction peak currents gradually increased and the corresponding oxidation peak currents visibly decreased. In a word, the **1**- and **3**-CPE exhibit good electrocatalytic activity for the reduction of bromate and nitrite.

Photocatalytic activity

As is known, the POM-based hybrids can exhibit good effect in the degradation of some organic dyes.¹⁸ We selected two organic dyes methylene blue (MB) and Rhodamine B (RhB) as model pollutants in aqueous media to evaluate the photocatalytic effect of compounds **1**–**4** under UV irradiation. In the process of photocatalysis, 100 mg compound was suspended in 90 mL of 0.02 mmol L^{-1} aqueous solution of MB or RhB. Furthermore, under dark conditions the suspension was magnetically stirred for about 10 min to ensure the equilibrium. Every interval (10 min for MB and 25 min for RhB) 5.0 mL samples were taken out for analysis by UV-vis spectrophotometer. As shown in Fig. 11a–c, with increasing reaction time we can clearly see that the percentage of MB degradation photocatalyzed by **1**, **2** and **4** increased obviously. The conversions of MB are nearly 100% for **1** and **2**, 76% for **4** (Fig. S8a†). However, the absorption peaks of MB photocatalyzed by compound **3** almost show no photocatalytic effect. Fig. 11d–f shows the photocatalytic degradation of RhB with the conversions of 58% for **2**, 46% for **3** and 77% for **4** after 175 min (Fig. S8b†). But the absorption peaks of RhB with compound **1** as the catalyst shows tiny change. This result shows that compounds **1**, **2** and **4** own good photocatalytic activities for the degradation of MB and compounds **2**, **3** and **4** can act as good photocatalysts for the degradation of RhB. In order to confirm whether compounds **1**–**4** as catalysts were stable during the photocatalytic process, the IR spectra and XRD diffraction patterns of compounds **1**–**4** catalyst before and after catalytic reactions for MB and RhB were measured. The IR spectra (Fig. S9†) show that the characteristic bands of POMs

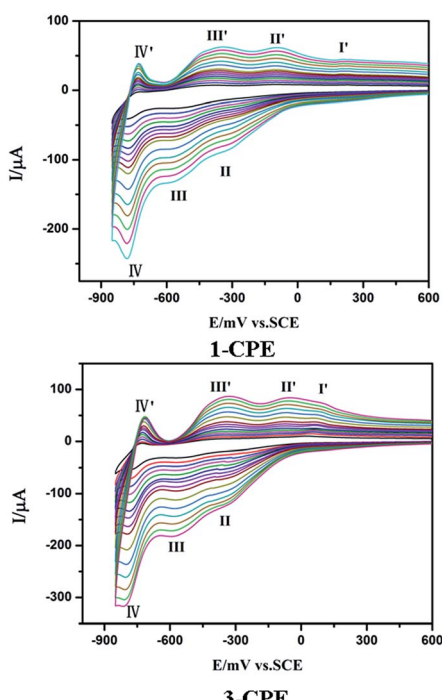


Fig. 9 The cyclic voltammograms of the **1**- and **3**-CPEs in $0.1 \text{ M H}_2\text{SO}_4$ + $0.5 \text{ M Na}_2\text{SO}_4$ aqueous solution at different scan rates (from inner to outer: 40 , 60 , 80 , 100 , 120 , 140 , 160 , 180 , 200 , 250 , 300 , 350 , 400 , 450 and 500 mV s^{-1} , respectively).



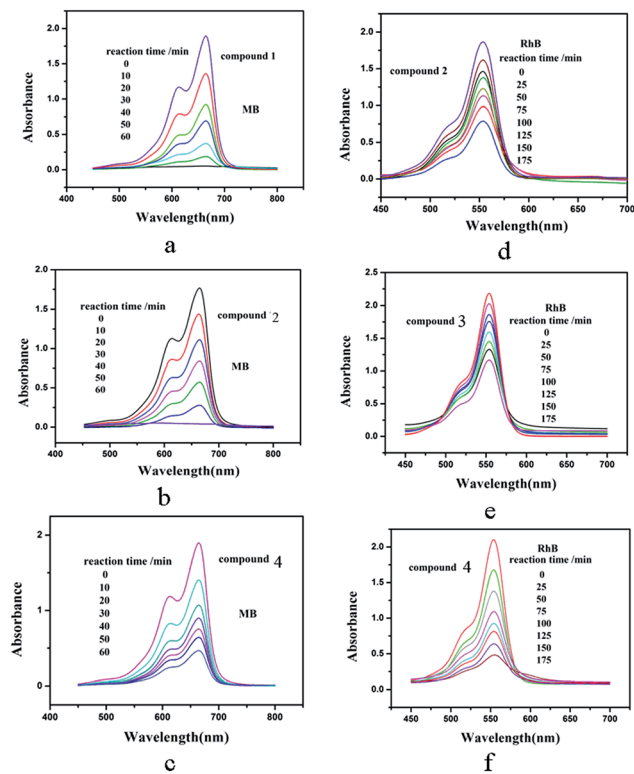


Fig. 11 Absorption spectra of the MB (a, b and c) and RhB (d, e and f) solution during the decomposition reaction under UV irradiation at the presence of title compounds 1–4.

and ligands for **1–4** catalyst after catalytic reactions for MB and RhB are maintained. The XRD diffraction patterns (Fig. S6†) of the catalyst before and after catalytic reactions show the diffraction peaks of experimental and recycled patterns match well in positions, proving that compounds **1–4** were stable during photocatalytic process.

Thermal gravimetric analyses

The TGA experiments were performed under N_2 atmosphere with a heating rate of $10\text{ }^\circ\text{C min}^{-1}$ in the temperature range of $50\text{--}850\text{ }^\circ\text{C}$, as shown in Fig. S10.† TG analyses of the title compounds all indicate two weight loss steps. The first weight loss below $350\text{ }^\circ\text{C}$ may be due to the release of water molecules. The second weight loss step in $400\text{--}850\text{ }^\circ\text{C}$ ascribes to the loss of organic H_2bdpm molecules 14.43% (calc. 12.0%) for **1**, 13.97% (calc. 12.05%) for **2**, 16.61% (calc. 16.52%) for **3**, 21.12% (calc. 21.17%) for **4**.

Conclusions

In summary, three new Keggin-based compounds have been constructed through using flexible H_2bdpm ligand. The H_2bdpm induces cycle subunits of the title compounds. Compound **1** shows a mono-supporting Keggin structure. Compound **2** shows a supramolecular structure. Compound **3** owns cycle-connecting-cycle chains, which are further linked by SiW_{12} anions to form a 2D layer. Compound **4** contains nested

cycle subunits, which are linked by linking H_2bdpm and anions to build a 3D framework. Further study on H_2bdpm to generate new POM-based cycle structures is underway.

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Notes and references

- (a) R. Yu, X. F. Kuang, X. Y. Wu, C. Z. Lu and J. P. Donahue, *Coord. Chem. Rev.*, 2009, **253**, 2872; (b) J. W. Zhao, Y. Z. Li, L. J. Chen and G. Y. Yang, *Chem. Commun.*, 2016, **52**, 4418; (c) A. Rubinstein, P. J. Lozano, J. J. Carbó, J. M. Poblet and R. Neumann, *J. Am. Chem. Soc.*, 2014, **136**, 10941.
- (a) J. J. Walsh, A. M. Bond, R. J. Forster and T. E. Keyes, *Coord. Chem. Rev.*, 2016, **306**, 217; (b) Z. J. Liu, X. L. Wang, C. Qin, Z. M. Zhang, Y. G. Li, W. L. Chen and E. B. Wang, *Coord. Chem. Rev.*, 2016, **313**, 94; (c) T. Hirano, K. Uehara, K. Kamata and N. Mizuno, *J. Am. Chem. Soc.*, 2012, **134**, 6425.
- J. Zhou, J. W. Zhao, Q. Wei, J. Zhang and G. Y. Yang, *J. Am. Chem. Soc.*, 2014, **136**, 5065.
- X. L. Wang, Y. F. Bi, B. K. Chen, H. Y. Lin and G. C. Liu, *Inorg. Chem.*, 2008, **47**, 2442.
- B. X. Dong, J. Peng, C. J. G. Garcíá, S. Benmansour, H. Q. Jia and N. H. Hu, *Inorg. Chem.*, 2007, **46**, 5933.
- (a) A. X. Tian, J. Ying, J. Peng, J. Q. Sha, H. J. Pang, P. P. Zhang, Y. Chen, M. Zhu and Z. M. Su, *Inorg. Chem.*, 2009, **48**, 100; (b) J. Ying, X. J. Liu, A. X. Tian and X. L. Wang, *Z. Anorg. Allg. Chem.*, 2011, **637**, 613; (c) A. X. Tian, X. J. Liu, J. Ying, D. X. Zhu, X. L. Wang and J. Peng, *CrystEngComm*, 2011, **13**, 6680.
- A. X. Tian, X. J. Liu, J. Ying, D. X. Zhu, X. L. Wang and J. Peng, *Inorg. Chem. Commun.*, 2011, **14**, 697.
- A. X. Tian, Y. Yang, J. Ying, N. Li, X. L. Lin, J. W. Zhang and X. L. Wang, *Dalton Trans.*, 2014, **43**, 8405.
- (a) K. Pavani and A. Ramanan, *Eur. J. Inorg. Chem.*, 2005, 3080; (b) K. Pavani, S. E. Lofland, K. V. Ramanujachary and A. Ramanan, *Eur. J. Inorg. Chem.*, 2007, 568.
- (a) A. X. Tian, Y. L. Ning, J. Ying, J. W. Zhang, X. Hou, T. J. Li and X. L. Wang, *Dalton Trans.*, 2015, **44**, 10499; (b) J. W. Zhang, W. Zhao, Q. L. Lu, J. Luan, Y. Qu and X. L. Wang, *J. Solid State Chem.*, 2014, **212**, 151.
- G. M. Sheldrick, *SHELXS-97*, 1997.
- A. L. Spek, *Acta Crystallogr., Sect. C: Struct. Chem.*, 2015, **71**, 9.
- I. D. Brown and D. Altermatt, *Acta Crystallogr., Sect. B: Struct. Sci.*, 1985, **41**, 244.
- (a) A. X. Tian, Y. L. Ning, J. Ying, G. C. Liu, X. Hou, T. J. Li and X. L. Wang, *CrystEngComm*, 2015, **17**, 5569; (b) A. X. Tian, Y. L. Ning, J. Ying, X. Hou, T. J. Li and X. L. Wang, *Dalton Trans.*, 2015, **44**, 386.
- (a) X. L. Wang, R. Zhang, X. Wang, H. Y. Lin and G. C. Liu, *Inorg. Chem.*, 2016, **55**, 6384; (b) Z. Y. Shi, Z. Y. Zhang,

J. Peng, X. Yu and X. Wang, *CrystEngComm*, 2013, **15**, 7199.

16 A. X. Tian, Y. Yang, J. Ying, N. Li, X. L. Lin, J. W. Zhang and X. L. Wang, *Dalton Trans.*, 2014, **43**, 8405.

17 M. Sadakane and E. Steckhan, *Chem. Rev.*, 1998, **98**, 219.

18 X. L. Wang, X. Rong, H. Y. Lin, D. N. Liu, X. Wang and G. C. Liu, *Inorg. Chem. Commun.*, 2016, **68**, 4.

