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An unprecedented (3,4,24)-connected heteropolyoxozincate organic framework as heterogeneous crystalline Lewis acid catalyst for biodiesel production

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A novel 3D hexadecanuclear heteropolyoxozincate organic framework, IFMC-200, has been successfully synthesized based on a late transition metal-oxygen cluster. IFMC-200 not only represents the first example of (3,4,24)-connected framework but also contains the first 24-connected single metal cluster in a crystal structure. It exhibits superior thermal stability, good water-stability, and even insensitivity to the existence of acid and base within a certain range of pH values. Furthermore, it performs as a heterogeneous crystalline Lewis acid catalyst with good activity for the conversion of long-chain fatty acids rather than short-chain ones, and high recycling efficiency for esterification reaction of fatty acids with alcohols to produce biodiesel.

Polycometalates (POMs), soluble anionic metal oxide clusters, are comprised of early-transition metal with d^0 or d^1 electronic configurations (usually $Mo^{VI/V}$, W^{VI} , V^V , Nb^V or Ta^V), which have attracted intense attention during the last two decades not only for their remarkable structural and electronic properties, but also for their intriguing applications ranging from catalysis to medicine¹⁻⁸. The metal oxide clusters entirely consisted of late transition metal ions with d^{10} electronic configuration are rarely available owing to the instability of extremely high negative charge, though a series of addenda atoms substituted by late transition metal ions (such as Fe^{III}, Co^{II}, Cu^{II}, Mn^{II} and Zn^{II}) were commonly reported⁹⁻¹¹. Compared with conventional POM clusters, d^{10} metal-oxygen clusters may have some special properties resulting from the inherent characteristics of late transition metals. In this regard, how to lower the overall negative charge to obtain d^{10} metal-oxygen clusters is a significant and challenging task in synthetic chemistry and material sicence^{12,13}.

Biodiesel, a renewable source of energy, seems to be an attractive candidate for the solution of global energy demands due to its environmental and sustainable benefits^{14,15}. However, biodiesel is currently not competitive with the conventional diesel fuel because of its high costs of raw material and production processing^{16–18}. That is, the presence of free fatty acids (FFAs) and water in low quality feedstocks would result in two undesirable side reactions, saponification and triglyceride hydrolysis, making the production process complicated. One way to overcome this problem is to subject FFAs to esterification before the transesterification of triglycerides. Thus, esterification is an essential step in the production of biodiesel and the search for highly efficient catalysts becomes greatly desirable in esterification reaction. In this context, heterogeneous catalysts, which are insoluble, easily separated from products, and reusable, exhibit important advantages with respect to homogeneous catalysts and become particularly important^{19–22}. POM-based materials with Lewis and Brønsted acidity and reversible redox activity have been utilized as efficient solid acid and electron transfer catalysts for the production of industrially important chemicals since the late 1970s^{23–28}. Recently, POM-based materials were also used as acid catalysts for biodiesel production^{29–32}. In most cases, Lewis acidic POM-based catalysts could be obtained through the

following strategies: (i) exchanging protons with Lewis acidic cations, including Zn^{II}, Al^{III}, Sn^{IV}, Fe^{III} and so on³², (ii) incorporation of rare earth metal ions into the vacant sites of POM ions^{33,34}, and (iii) supporting POMs on Lewis acidic porous materials, for instance, Ta_2O_5 and zeolites^{31,35,36}. However, the above materials often encounter disadvantages, such as being homogeneous forms, difficulty in controlling synthesis, leaching of active sites from supports, hindering substrate accessing to catalytically active sites, and inferior activities compared to homogeneous analogues. Meanwhile, crystalline catalysts have caught increasing attention of researchers worldwide due to their well-defined crystalline structures, regular and tunable cavities and high catalytically active site density³⁷⁻³⁹. If organic ligands can directly bond to metal-oxygen clusters to achieve metal-oxygen clusters-based crystalline catalysts that meet the above-mentioned requirements, it may be a 'one stone two birds' method. So far, this kind of catalyst has not been reported for biodiesel preparation. Therefore, the search for design synthesis and development of environmentally benign catalysts which are highly active in water, higher thermodynamic stable, highly accessible to acid sites and recyclable is one of the most challenging issues in the production of biodiesel.

These inspired us to employ a semirigid ligand containing eight carboxyl groups, 5,5'-(2,2-bis((3,5-dicarboxyphenoxy)methyl)propane-1,3-divl)bis(oxy) diisophthalic acid (H₈L, Figure S1)⁴⁰, and H₃PO₃ for the isolation of crystalline catalysts containing d¹⁰ metal-oxygen clusters, which gives the following considerations: (i) It is difficult to isolate classical metal-oxygen clusters from late transition metal ions due to the instability of extremely high negative charge. The substitution of terminal -O and $\mu_2 - O$ of POM clusters with rich carboxyl groups of H₈L ligand would greatly reduce negative charge, which is beneficial to the formation of late transition metal-oxygen clusters. (ii) It is difficult to directly bond organic linkers to the traditional POM clusters owing to their oxygen-rich surface, so we intend to take the advantage of late transition metal ions to replace early ones to synthesize metal oxide cluster-organic frameworks. (iii) The introduction of H₃PO₃ is favorable to induce the formation of metal-oxygen clusters, which perhaps plays the role of heteroatom in the formation of hetero-POM anions. Fortunately, a novel heteropolyoxozincate containing hexadecanuclear zinc nanocluster, $[Zn_{16}(HPO_3)_4]L_3$ (IFMC-200, IFMC = Institute of Functional Material Chemistry), was successfully synthesized and structurally characterized. IFMC-200 is stable in air and common organic solvents, and remarkably insensitive to the presence of water, even in acidic and basic aqueous solution within a certain range of pH values. Furthermore, IFMC-200 was employed as a heterogeneous catalyst for the preparation of biodiesel, which is the first time to employ a crystalline material of entire d¹⁰ metal oxide cluster with Lewis acid nature.

Results

Single-crystal X-ray analysis. IFMC-200 crystallizes in the cubic space group F-43c (219, Table S1) and consists of two kinds of Zn^{II} cations (Figure S2), HPO_3^{2-} anion and deprotonated L^{8-} ion. Each Zn(1) ion is tetrahedrally coordinated by three carboxylic oxygen atoms derived from three bridging L8- fragments and one oxygen atom originated from HPO3²⁻ ion. And each Zn(2) atom shows a octahedral geometry, which is completed by six oxygen atoms originated from three bridging L⁸⁻ ions and three HPO₃²⁻ ions. The P element in HPO₃²⁻ ion is in +3 oxidation state, and exhibits a tetrahedral configuration with the lone pairs pointing into the tetrahedral cavity, which has been observed in arsenolite $(As_2O_3)^{41}$. In addition, twelve Zn(1) and four Zn(2) cations were connected by four HPO32- anions, resulting in a nanocluster subunit [Zn16 $(HPO_3)_4]^{24+}$ (noted as $\{P_4Zn_{16}\}$, Figure 1a, left). To a certain extent, it is similar to classical Keggin-type cluser⁴², which is composed of four internally edge sharing triads (M₃O₁₃) that share corners with



Figure 1 | (a) The structures of $\{P_4Zn_{16}\}$ (left) and Keggin-type cluster (right); (b) and (c) the cage structure in **IFMC-200**; and (d) the 3D framework of **IFMC-200**.

each other and are disposed tetrahedrally around a central heteroatom with T_d symmetry (Figure 1a, right). As a result of the expansion of space among four HPO₃²⁻ ions, the size of $\{P_4Zn_{16}\}$ cluster in IFMC-200 (with a diameter of \sim 13 Å) is clearly larger than that of Keggin-type anion (\sim 10 Å). For conventional fully oxidized POM anions, especially Keggin ions, the highest occupied molecular orbital (HOMO) is mostly from 2p orbitals of the bridging-O atoms, whereas the lowest unoccupied molecular orbital (LUMO) is from 4d orbitals of Mo or W atoms43. However, the frontier molecular orbitals of IFMC-200 show significant differences with those of traditional POM clusters owing to d¹⁰ electronic configuration of Zn^{II} cations. In IFMC-200, the carboxyl-O atoms that coordinate to Zn(2) atoms contribute to the HOMO, and the LUMO is mainly from p orbitals of carboxyl-C atoms (Figure S3). An interesting structural feature of **IFMC-200** is that each $\{P_4Zn_{16}\}$ cluster is stabilized by 24 carboxyl groups from twelve L8- moieties and the connection of eight $\{P_4Zn_{16}\}$ clusters and six L^{8-} fragments generates one nanocage with dimensions of $11 \times 11 \times 11$ Å (Figure 1b,c and S4). Such connectivity leads to a 3D open architecture (Figure 1d). PLATON calculations reveal that a free volume in IFMC-200 is about 7786.2 Å³ per unit cell (27.1% of the cell volume)44.

On the basis of topological analysis, the structure of IFMC-200 can be considered as a 3D (3,4,24)-connected framework, if the center of the L^{8-} ligand is regarded as a 4-connected node, the $C_6H_3(COO)_2$ moiety as a 3-connected node, and the {P₄Zn₁₆} cluster as a 24connected node with a distorted rhombicuboctahedral geometry (Figure 2). The underlying trinodal net, which has been assigned the reticular chemistry structural resource (RCSR) symbol⁴⁵, ddy, was analyzed with the program Systre⁴⁶ which shows that it has intrinsic symmetry Pm-3m and is one of a special group of nets with the minimum number of kinds of edge (link) - in this case two for a trindal net. A maximum symmetry embedding is presented as a systre file in the Supplementary Information. It is worthy of note that the 24-connected node is essentially different from that documented in the literatures⁴⁷. The previously-reported 24-connected node is really a tertiary building unit, constructed from twelve "paddle-wheel" dimetal tetracarboxylate secondary building units and may be further extended by functionalization at 5-position of 1,3benzenedicarboxylate ligand (Figure S5)⁴⁸⁻⁵⁰. It should be pointed out that {P₄Zn₁₆} cluster is the first real 24-connected node, which is an important discovery in topological crystal chemistry. From the data



Figure 2 | The (3,4,24)-connected structure of IFMC-200. (a) the center of H_8L ligand as a 4-connected node, the $C_6H_3(COO)_2$ moiety as a 3-connected node; (b) and (c) the { P_4Zn_{16} } cluster as a 24-connected node; and (d) the (3,4,24)-connected **ddy** topology.

in the RCSR, we found that IFMC-200 is the first example of (3,4,24)connected topology and the first undoubted 24-connected node constructed from a single cluster. Moreover, it was also not hypothesized in Euclidean Patterns in Non-Euclidean Tilings (EPINET)⁵¹. Offering further insight into this intricate architecture, the overall structure of IFMC-200 can also be considered as a (4,12)-connected ftw topology framework when L^{8–} fragments are considered as 4connected nodes and each {P₄Zn₁₆} cluster corresponds to a 12connected vertex (Figure S6)⁵². So far, only three topologies combine 4,12-connected components in the RCSR database, that is, **ith**, **sh**p, and ftw. Notably, our group has reported a (4,12)-connected **ith** topology network (Figure S7)⁵³.

Discussion

Crystals of IFMC-200 were isolated by hydrothermal reaction of ZnCl₂, H₃PO₃, Na₂MoO₄·2H₂O, Mo powder, H₈L, and TBAOH (Tetrabutylammonium hydroxide). The parallel experiments suggest that IFMC-200 cannot be obtained if there is no addition of Na2MoO4.2H2O and Mo powder. Meanwhile, H3PO3 not only plays the role of phosphorus source but also structure-directing agent in the formation of $\{P_4Zn_{16}\}$ cluster because there is no product if H₃PO₃ was replaced with H₃PO₄. It is clear to see that IFMC-200 formed a cubic shape under both optical microscopy and scanning electron microscopy (Figure S8). The corresponding EDS analysis reveals that there is no Mo element in IFMC-200 although sodium molybdate dihydrate and Mo powder were introduced to the reaction system (Figure S9), which is in agreement with the result of single crystal X-ray diffraction. Phase purity of the bulky crystals was confirmed by the similarity between the experimental and simulated Xray powder diffraction (XRPD) patterns (Figure S10). IFMC-200 was generated in aqueous media; theoretically, it is stable in water even at different temperatures, as confirmed by subsequent XRPD and FT/IR measurements (Figure S10 and S11). It was also found that

IFMC-200 is stable in acidic and basic aqueous solution in the pH range of 2–12 (Figure 3 and S12). The vapour adsorption experiments for IFMC-200 suggest that water is hardly adsorbed, but certain amounts of ethanol and cyclohexane are adsorbed (Figure S13). Thus, IFMC-200 possesses hydrophobic cages. Furthermore, the thermogravimetric analysis (TGA) suggests that IFMC-200 has exceptional thermal stability with decomposition starting at 450°C (Figure S14), which is much higher than that of Nafion NR50 (280°C), one of the most stable resins⁵⁴.

Generally, biodiesel can be prepared by transesterification of triglycerides or esterification of fatty acids with alcohols^{55,56}. In terms of both atomic economy and green synthesis, heterogeneous solid state acid catalysts that work actively with easily separation for recyclability are highly desirable. The acid value of 0.61 mmol \cdot g⁻¹ for IFMC-200 was detected by titration with aqueous ammonia, which is smaller than those of $ZnCl_2$ (7.96 mmol·g⁻¹) and Keggin structure $Zn_{1,2}H_{0,6}PW_{12}O_{40}$ (1.14 mmol·g⁻¹)³⁰, but higher than those of other Lewis solid state acid catalysts (e.g., Y-zeolites, 0.54 mmol $H^+ \cdot g^{-1}$)⁵⁷. In this regard, IFMC-200 was evaluated as a heterogeneous catalyst in the esterification of fatty acids with a series of alcohols (Figure 4). The reactions were performed in the presence of an excess amount of alcohols with oil/alcohol molar ratio of 1:10 at 65°C. Preliminary studies were carried out to optimize the reaction conditions, and the results are presented in Figure S15, Table S2 and S3. The esterification reaction of palmitic acid with ethanol catalyzed by IFMC-200 reached its highest conversion (44.7%) and turnover frequency (TOF, 26.4 h^{-1}) with catalyst amount of 1% wt total reactant weight in 10 h. In comparison, a blank experiment was also performed under the given conditions. The conversion of IFMC-200 catalyzed esterification of palmitic acid with ethanol was 10 times higher than that in the case of no catalyst (4.3%). Furthermore, we used the conventional Lewis acid ZnCl₂ with higher acid value than IFMC-200 as catalyst under the identical conditions, and the result suggested that IFMC-200 showed higher TOF than $ZnCl_2$ (8.6 h⁻¹). This may be attributed to the poor water-tolerance of ZnCl₂. It is known that ZnCl₂ requires strictly anhydrous conditions because it immediately reacts with water forming in esterification of acids with alcohols, resulting in its serious decomposition⁵⁸. IFMC-200 is much more stable than ZnCl₂ in esterification reaction and could not be decomposed by water. With the optimal reaction time and catalyst amount, a series of fatty acids and alcohols were then examined under this condition and the results are collected in Table 1 and Figure S16. As can be seen from Table 1, the biodiesel conversion increased as the chain length of free fatty acids was increased, maintaining other constants. This result is opposite to the previous report⁵⁹, in which Goodwin et al. illustrated that the porous silica support would diminish the likelihood of mass diffusion limitation, making it favorable for the esterification reaction of low-molecular weight carboxylic acids. This abnormal result perhaps can be attributed to the positive effects of inorganic-organic hybrid framework of IFMC-200 and the presence of hydrophobic cage structure in IFMC-200. It is well-known that the acidic catalysis depends on the acidic strength, as well as the concentration of substrates around the catalytic sites. IFMC-200 located its Lewis acidic sites on the backbone of framework, and hydrophobic surroundings inside the pores. This unique structure feature provides long chain fatty acid molecules more chance to access to the catalytic sites due to their large lipophilicity. In other words, the long chain fatty acid molecules could be concentrated around the catalytic sites compared to short-chain ones. In addition, the hydrophobic cages could promote the water generating during the reaction to leach from the catalytic sites, which avoids the deactivation of the acid sites by water. Up to now, there are only few reports on solid porous catalysts with hydrophobic cages, just Inumaru's group firstly demonstrated H₃PW₁₂O₄₀ immobilized in hydrophobic nanospaces of organomodified mesoporous silica to show water-tolerant properties⁶⁰. Therefore, IFMC-200 exhibited



Figure 3 | The XRPD patterns of IFMC-200 after heated in Teflonlined stainless steel container at $65^{\circ}C(a, b)$ and $140^{\circ}C(c, d)$ for 12 h at different pH values, respectively. Exp represents XRPD pattern of IFMC-200 without further treatment.

water-tolerant properties compared to conventional Lewis acid, ZnCl₂, generated by the existence of hydrophobic cages, which makes it favorable for esterification of long-chain carboxylic acids rather than short-chain ones. It might be the only one report on this opposite result. More importantly, **IFMC-200** showed very good

Table 1 | The Con. (%) and TOF (h^{-1}) of **IFMC-200** catalyzed esterification of FFAs with alcohols. Conditions: 1% wt catalyst; 65°C; oil/alcohol molar ratio = 1:10; 10 h

Entry	Alcohols	Fatty acids	Con./%	TOF/h ⁻¹
1	methanol	octanoic acid	8.8	5.3
2	methanol	lauric acid	20.4	12.3
3	methanol	palmitic acid	23.6	13.9
4	methanol	stearic acid	33.9	20.2
5	ethanol	octanoic acid	31.8	19.2
6	ethanol	lauric acid	46.1	25.6
7	ethanol	palmitic acid	44.7	26.4
8	ethanol	stearic acid	47.0	28.0
9	n-butanol	octanoic acid	62.8	37.9
10	2-methyl propanol	octanoic acid	62.2	37.6
11ª	ethanol	palmitic acid	40.5	8.6
12 ⁶	ethanol	palmitic acid	4.3	_
°ZnCl₂ w ⁵No cata	as used as catalyst. lyst was used.			

recyclability. For example, there was almost no activity loss over IFMC-200 after six times (Figure S17). This perhaps associated with the stable structure of IFMC-200 (Figure S18 and S19). The excellent stability and recyclability of IFMC-200 is potentially important for industrial applications.

In summary, we synthesized an unprecedented 3D (3,4,24)-connected heteropolyoxozincate organic compound, IFMC-200, which contains the first example of 24-connected node constructed from a single metal cluster. We have developed a new strategy with "carboxyl group-stabilized polyoxozincate cluster" mode that helps to lower the overall negative charge, and successfully proved that it is an efficient method to isolate late transition metal-oxygen cluster. IFMC-200 exhibits superior thermal stability, good water-stability, tolerance to acid and base, and Lewis acidity, which enable its cata-



Figure 4 | The equation of IFMC-200 catalyzed esterification of FFAs with alcohols.

lytic activity in the esterification of carboxylic acids with alcohols. Surprisingly, IFMC-200 exhibits abnormal catalytic result that is favor for the conversion of long-chain fatty acids under the same conditions compared to the previous reports. Further studies concerning this strategy for the synthesis of more late transition oxide cluster-based heterogeneous crystalline catalysts and their catalytic activities in the production of biodiesel are currently in progress.

Methods

Sample synthesis. 6 mL of distilled water was added to a beaker, which contains $Na_2MoO_4 \cdot 2H_2O$ (0.63 g, 2.6 mmol), Mo powder (0.05 g, 0.52 mmol), ZnCl₂ (0.14 g, 1.0 mmol), 50% H₃PO₃ (0.10 mL), and TBAOH with stirring for 15 min. Next, the pH value of the mixture was adjusted to 4.8 with 2 mol·L⁻¹ HCl and then the mixture was stirred for another 15 min followed by the addition of H₈L (0.12 g, 0.15 mmol). Then the mixture was transferred and sealed in a 23 mL Teflonlined stainless steel container, and cooled to room temperature at 5°C·h⁻¹ after heated at 180°C for 5 d. Cubic crystals of **IFMC-200** were collected and dried in air (yield 65%, based on H₈L). Elemental analysis: Anal. Calcd for C₁₁₁H₆4O₇₂P4Zn₁₆: C 35.84, H 1.73, Zn 28.13, P 3.33; found: C 35.78, H 1.82, Zn 28.05, P 3.26. IR (KBr, cm⁻¹): 3448 (s), 2939 (m), 1626 (s), 1561 (s), 1469 (s), 1425 (s), 1393 (s), 1266 (m), 1138 (m), 1082 (m), 1046 (m), 889 (w), 815 (w), 775 (m), 730 (m), 594 (w), 458 (w).

Sample characterization. The starting material $H_{s}L$ was synthesized according to modified procedure of the reported literature⁴⁰. All other chemicals were obtained from commercial sources, and were used without further purification. Elemental analyses (C, H and N) were measured on a Perkin-Elmer 2400 CHN elemental analyzer; Zn and P were determined with a Plasma-SPEC(I) ICP atomic emission spectrometer. IR spectrum was performed in the range 4000–400 cm⁻¹ using KBr pellets on an Alpha Centaurt FT/IR spectrophotometer. X-ray powder diffraction (XRPD) measurement was recorded radiation ranging from 3 to 60° at room temperature on a Bruker D8 Advance diffractometer with Cu-K_x (λ = 1.5418 Å). Thermogravimetric analysis (TGA) was performed using a Perkin-Elmer TG-7 analyzer heated from room temperature to 900°C under nitrogen at the heating rate of 5°C·min⁻¹. Field-emission scanning electron microscopy (FE SEM) images were obtained with a XL30 ESEM FEG microscope equipped with energy dispersive X-ray spectroscopy (EDS).

Determination of acid strength. Titration was used to evaluate the acidic characteristics of the solids. 100 mg of solid samples suspended in 25 mL of NaCl aqueous solution (0.05 mol·L⁻¹) was stirred for 30 h. Then, acid value was measured by titration with a solution of aqueous ammonia (0.01 mol·L⁻¹) using phenolphthalein as an indicator.

Catalytic experiments. A 100 mL three-necked glass flask charged with cooled condenser with ~5.0 g of FFAs, a certain amount of anhydrous alcohols and catalyst (IFMC-200) in it were vigorously stirred and refluxed (65°C) for the required reaction time. After reaction, the mixture was rotary evaporated at 50°C to separate the excess of alcohols. Then the mixture was left divided into two layers by centrifugation with 4000 rpm. The upper layer was esters of FFAs and the lower layer was **IFMC-200** with small amount of water. The conversion of esters was calculated by measuring the acid value of the products and the yield was detected by gas chromatography (GC). The catalyst was decanted at the bottom of centrifuge, which was easy to be separated, washed by methanol to remove the remaining reactants and water, then dried in air and reused for the next experiment.

Adsorption experiments. The adsorption isotherms of water, ethanol and cyclohexane are measured at 298 K for IFMC-200. Before the measurements, the crystalline sample of IFMC-200 was immersed in CH_2Cl_2 for three days, and then heated at 100°C for 12 h under a vacuum to evacuate the samples.

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Author contributions

D.Y.D. and J.S.Q. carried out the experiments and wrote the paper. Z.S. and X.H.W. performed the biodiesel preparation experiment. L.K.Y. calculated the frontier molecular orbitals. M.O. performed the topology analysis. Z.M.S., S.L.L., X.L.W. and Y.Q.L. supervised the research and revised the manuscript. All authors reviewed the manuscript.

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