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A Mesoporous Zirconium-Isophthalate Multifunctional Platform

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SUMMARY

Mesoporous materials suffer from limitations including poor crystallinity and hydrolytic stability, lack of chemical diversity, insufficient pore accessibility, complex synthesis and toxicity issues. Here the association of non-toxic Zr-oxo clusters and feedstock isophthalic acid (IPA) via a Homometallic-Multicluster-Dot strategy results in a robust crystalline mesoporous MOF, denoted as MIP-206, that overcomes the aforementioned limitations. MIP-206, built up from an unprecedented combination of Zr$_6$ and Zr$_{12}$ oxo-cluster inorganic building units into a single structure, exhibits accessible meso-channels of ca. 2.6 nm and displays excellent chemical stability under different hydrolytic and harsh conditions. Owing to the abundant variety of functionalized IPA linkers, the chemical environment of MIP-206 can be easily tuned without hampering pore accessibility due to its large pore windows. As a result, MIP-206 loaded with palladium nanoparticles acts as an efficient and durable catalyst for the dehydrogenation of formic acid under mild conditions, outperforming benchmark mesoporous materials. This paves the way towards the utilization of MIP-206 as a robust mesoporous platform for a wide range of potential applications.
INTRODUCTION

Mesoporous solids, a category of materials possessing structural voids of 2-50 nm in which large guest molecules can be accommodated, are of interest to sustainable development with potentially broad applications in fields related to energy, environment and health. Owing to their high availability and tunable pore size, traditional mesoporous materials, such as silica, metal oxides and activated carbons, are prevalent in current usage, particularly in heterogeneous catalysis\(^1\), energy conversion/storage\(^2\), analytical science\(^3\) and medical areas\(^4\), although they suffer from well-recognized limitations such as lack of crystallinity, chemical diversity, structural uniformity, stability and/or reproducibility. These limitations have inhibited pursuits to improve their performance, extend their use to other applicable fields and accumulate fundamental understanding\(^5\) of these materials. Considerable efforts have been devoted in the past two decades to develop substitutes, promote new strategies and develop alternative candidates\(^6\). Among them, mesoporous metal-organic frameworks (MOFs) are a promising family of materials that exhibits an ordered porosity and successfully addresses some limitations of conventional benchmarks. MOFs can be composed of an almost unlimited selection of inorganic building blocks (metal ions or clusters) and organic linkers in periodic structures which are merged efficiently, exhibiting high crystallinity with atomic precision, good reproducibility of preparation over control, and, most importantly, remarkable structural and chemical tunability\(^7; 8\).

Two strategies for preparing mesoporous MOFs are well-documented. Extending the
organic linker to construct mesoporous MOFs following the isoreticular chemistry strategy is the most straightforward route and illustrates well the ability to manipulate MOF structures to achieve diverse topologies and functionalities\textsuperscript{9-11}. The practical applicability is, however, severely limited by the costly and unscalable linkers, significantly reduced stability and, in some cases, structural interpenetration. On the contrary, the super-molecular building block (SBB) construction approach, which involves the use of small and rigid linkers (similarly to the construction of crystalline mesoporous zeolites from small inorganic building units\textsuperscript{12; 13}) to minimize structural interpenetration issues, leads to accessible mesoporous voids for utilization\textsuperscript{14-18}. For instance, terephthalic acid, a polymer industry feedstock, when assembled with homometallic monoclonal superoctahedra dots composed of Cr(III) trimers of octahedra, leads to the most chemically stable mesoporous MOF (MIL-101)\textsuperscript{14}. However, its practical use is limited by insufficient accessibility to the mesopores due to their microscale pore windows, toxic chromium species and harsh, poorly scalable synthesis conditions. Similarly, mesoporous MOFs based on heterometallic multicluster dots, such as the Zr/Cu carboxylate/azolate MOF-818 (Figure S1)\textsuperscript{17}, enhance the pore accessibility through the inclusion of a larger number of clusters within each pore window (12 versus 6 in benchmark mesoporous MOFs such as MIL-101 or NU-1000)\textsuperscript{19}, but suffer from limitations such as the formation of competitive byproducts, difficulty with the introduction of organic functional groups and limited chemical stability, thus making it still challenging to consider these mesoporous MOFs for most practical applications.
To date, there is no satisfying strategy to construct robust mesoporous MOFs with adequate pore accessibility and chemical tunability based on feedstock linkers and nontoxic metals. To address this challenge, we present here a Homometallic-Multicluster-Dot (HMD) strategy to efficiently blend the connection diversity of two different homometallic inorganic clusters into one structural dot for the construction of robust mesoporous MOFs. As a proof of concept, we selected the nontoxic Zr$_6$ and Zr$_{12}$ oxo-clusters to build the inorganic dot and isophthalic acid (IPA), a feedstock produced at billion-kilogram scale annually in industry, as the linker to synthesize a robust mesoporous Zr-IPA MOF, denoted as MIP-206 (MIP stands for the Materials of the Institute of Porous Materials of Paris). It displays excellent chemical and thermal stability as well as large accessible mesopores of ca. 2.6 nm in diameter. One can introduce diverse functional groups at the 5-position of the IPA moiety that faces the mesopores in the structure, allowing fine-tuning of the chemical environment of the pores by simply interchanging various cost-effective substituted IPAs as linkers. MIP-206 can thus act as an ideal mesoporous matrix to support palladium nanoparticles (Pd NPs) for the heterogeneous catalytic release of hydrogen from formic acid under mild conditions. The tailorable nature of the MIP-206 structure enables Pd(0)@MIP-206s catalysts to outperform current benchmark mesoporous catalysts, while exhibiting excellent stability and durability under experimental conditions. Hence, our work paves the way towards the design of robust mesoporous MOFs by applying the HMD strategy, which not only promotes the expanding structure diversity of MOFs, but also holds great promises in achieving
RESULTS AND DISCUSSION

Zr-MOFs assembled from Zr$_6$ or Zr$_{12}$ oxo-clusters separately with small linkers have been well-investigated in the past years despite the fact that only microporous structures could be achieved due to the short distances separating each cluster. While the Zr$_6$ oxo-cluster shows a flexible connectivity tolerance ranging from 4 to 12, the Zr$_{12}$ oxo-cluster displays an even wider connectivity range as a result of its intrinsic thirty-six terminal coordination sites. Thus, the combination of Zr$_6$ and Zr$_{12}$ oxo-clusters was anticipated to display considerable flexibility in forming homometallic multiclusters dots with large sizes suitable to construct new mesoporous MOFs. Following this expectation, MIP-206 was synthesized by simply heating a mixture of ZrCl$_4$, isophthalic acid and formic acid under solvothermal conditions, which is also suitable for the preparation of MIP-206s with substituted IPA linkers with diverse functional groups as well as multivariate MIP-206s based on mixed linkers. It is noteworthy that the syntheses of MIP-206 series are facilely scalable with, for instance, MIP-206 or MIP-206-OH at a 10 g scale production from a one-pot reaction.

The crystal structure of MIP-206 was determined by a computational reverse topological construction approach combined with synchrotron X-ray diffraction single crystal data and high-resolution powder X-ray diffraction (PXRD) data. The structural
model was fine-tuned until the calculated PXRD patterns of the DFT-derived minimum energy structure of MIP-206 was found to be consistent with the experimental structure (Figure S2). MIP-206 with a simulated structural formula of

$$\text{[Zr}_{12}\text{(\text{\textit{\mu}_3-O})_8\text{(\text{\textit{\mu}_3-OH})_8}\text{(formate})_6][\text{Zr}_6\text{(\text{\textit{\mu}_3-O})_4\text{(\text{\textit{\mu}_3-OH})_4}\text{(formate})_4]\text{IPA})_{18}}$$

crystallizes in a hexagonal $P-62C$ space group with unit cell parameters of $a = b = 33.7520(5) \text{ Å}$, $c = 21.4610(3) \text{ Å}$, $V = 21172.86(5) \text{ Å}^3$. There are two types of Zr oxo-cluster secondary building units (SBUs) in the structure, namely 8-connected Zr$_6$ and 12-connected Zr$_{12}$ oxo-clusters (Figure 1A and B).

Each Zr$_{12}$ oxo-cluster is centered between three adjacent Zr$_6$ oxo-clusters, through linkages formed by six pairs of bridging IPA molecules, forming a triangular HMD (Figure 1C). Each Zr$_6$ oxo-cluster is at one triangle vertex position, sharing a pair of IPA linkers which connect the nearest Zr$_6$ oxo-clusters to the neighboring HMDs, forming to a mesoporous window encircled by three Zr$_{12}$ oxo-clusters and six Zr$_6$ oxo-clusters (Figure 1D). This explains why a large window of ca. 2.6 nm of free aperture is formed, of a similar size as that of MOF-818 but significantly larger than those of topical mesoporous MOFs whose hexagonal windows involve only six oxo-clusters, such as MIL-101 or DUT-126. Noticeably, all IPA molecules share the same connection mode (Figure S3) and adopt a configuration facing the 5-position to the meso-void inner surface. This is of a great advantage for tailorable window size, shape and chemical environment by substitution with functionalized IPA linkers. A two-dimensional (2D) layer is generated via the interconnection of adjacent HMDs. In addition to the mesoporous window, there is a small triangular cavity with a diameter
of ca. 7 Å between the contiguous Zr$_6$ oxo-clusters and a slit-like void between the IPA linker pairs (Figure 1E). However, these small interspaces are largely inaccessible when the adjacent 2D layers are stacked together in an ABAB fashion, in which the Zr$_{12}$ oxo-clusters from the B layer occupy the space between the two small triangle voids from the two A layers above and below while the Zr$_6$ oxo-clusters occupy the gap between the bottom and top slits (Figure S4). As a result, a porous architecture with accessible meso-channels running along the c-axis is generated via the tight packing of each layer, with arrays of functional groups attached at the channel surface (Figure 1F).

Figure 1. Crystal structure detail of MIP-206 (A) A 8-connected Zr$_6$ oxo-cluster SBU. (B) A 12-connected Zr$_{12}$ oxo-cluster SBU. (C) A triangle HMD with a center Zr$_{12}$ oxo-cluster linked by three adjacent Zr$_6$ oxo-clusters. (D) A single mesoporous window constructed from the three neighboring
HMDs. (E) A single 2D layer. (F) Mesoporous 1D pore system viewed along the c-axis (Zr from $\text{Zr}_6$ oxo-cluster in purple, Zr from $\text{Zr}_{12}$ oxo-cluster in blue, C in gray, O in red, and functional group, such as OH, Br, I, F, Me, OMe and NO$_2$, in the 5-position of IPA marked with yellow ball).

Unlike benchmark mesoporous materials which display limited chemical diversity, MIP-206 is able to facilitate excellent tunability of the pore chemical environment through customization of the IPA linker, in particular as a considerable number of cost-effective functionalized IPA compounds are commercially available at large scales and more diverse, complex moieties can be generated through well-documented organic synthesis. To this end, a series of substituted IPA linkers were applied to the fabrication of isostructural MIP-206 analogues. As shown in Figure 2A, functional groups such as electron donating/withdrawing groups, halogens and heteroatoms, that are well-known functional groups enabling typically a careful tuning of the chemical environment of the pores of MOFs, did not cause a noticeable disruption to the synthesis, with all variations leading to highly crystalline isostructural materials despite slight variations of peak widths in the PXRD patterns due to different particle sizes (Figure 2B).

As a result of the tailorable chemical environment, the corresponding pores’ properties could be easily tuned. For instance, as a model characterization, the hydrophilicity/hydrophobicity of the mesochannels in MIP-206s can be easily altered as evidenced by water adsorption measurements (Figure 2C). While the non-functionalized MIP-206 displays an $\alpha$ value of 0.45, $\alpha$ decreased to 0.30 in MIP-206-OH, reflecting the influence of the hydrophilic hydroxyl group. In contrast,
the more hydrophobic bromo substituent in MIP-206-Br shifted the $\alpha$ value to 0.53, indicating the largely reduced hydrophilicity of the structural cavity. Similarly, the adsorption step and capacity of MIP-206s in alcohol adsorption isotherms could be adjusted as well (Figure S5), which is of interest for a wide range of applications including heat reallocation, separation and drug delivery$^{20,21}$.

**Figure 2.** Functional MIP-206s with tunable chemical environment and related performance in water sorption. (A) Simplified illustration of a structural segment of functional MIP-206s constructed with pure substituted IPA linkers marked in gold and MTV-MIP-206s containing secondary linkers marked in blue (red ball stands for carboxylate group of the IPA linker). (B) PXRD patterns ($\lambda_{Cu} \approx 1.5406$ Å) of selected MIP-206s with different functional groups. (C) Water sorption isotherms of
MIP-206, MIP-206-OH and MIP-206-Br collected at 30 °C showing their structural difference in hydrophilicity (The relative pressure (α) at which 50% of maximum capacity is achieved in water adsorption isotherm is a hydrophilicity indicator, which could be used to compare the hydrophilicity of the adsorbent). PXRD comparison between MIP-206 and selected MTV-MIP-206s.

However, some IPA-type compounds could not successfully assemble into the corresponding MIP-206 isostructures, such as 5-NH₂-IPA, 5-SO₃H-IPA, pyrazole-3,5-dicarboxylic acid (PDA) and furan-2,5-dicarboxylic acid (FDA). To further extend the chemical diversity of the pores in this case, we considered the construction of multivariate MIP-206 counterparts (MTV-MIP-206s) following the solid-solution mixed linker strategy. A mixture of the dominant compatible IPA linker with the minor, less compatible IPA-type compound functioned similarly with the major linker alone appearing in the MOF structure (Figure 2D). Phases other than MIP-206 obtained when using the secondary linkers separately were not observed in the mixed linker systems (Figure S6). It indicates the determining role of the major linker in the MOF structure formation process to avoid the generation of unexpected by-products. For instance, 5-SO₃H-IPA led to an unknown crystalline phase under the standard reaction conditions when used as a sole linker. In contrast, a mixture of IPA/5-SO₃H-IPA (4/1 molar ratio) yielded the mixed linker MIP-206/5-SO₃H-IPA compound with 60 mol% IPA and 40 mol% 5-SO₃H-IPA in the final structure (Figure S7 and Table S1). Different secondary linkers could also yield the MTV-MIP-206 phases with varied ratios of the two linkers involved, as evidenced by the ¹H NMR data collected from decomposed samples in KOH/D₂O solution. The linker content could be adjusted simply by altering the starting material ratios. For example, the
5-NH$_2$-IPA content in the MIP-206/5-NH$_2$-IPA product could be tuned from 9 mol% to 28 mol% when the amount of 5-NH$_2$-IPA added as starting material increased from 10 mol% to 25 mol% (Figure S7 and Table S1). This tailorable multivariate inclusion is very helpful to functionalize the material according to the application requirement.

Nitrogen porosimetry data of MIP-206 samples were collected at 77 K to characterize their porosity (Figure S8). MIP-206 displays a three-divisional adsorption isotherm with a notable step occurring in the 0.1-0.2 $P/P_0$ range, which is consistent with its mesoporous character. A nitrogen uptake of 290 cm$^3$g$^{-1}$ was recorded at the relative pressure point of $P/P_0$=0.9 associated with an accessible pore volume of 0.45 cm$^3$g$^{-1}$, conforming well with the value calculated from the crystal structure (Tables S2 and S3). The other samples exhibit isotherms of similar shapes to that of MIP-206 despite distinct nitrogen uptake profiles. It is important to note that the steps in the adsorption isotherms occur in a very narrow range of relative pressures (Figure S8, inset), which supports that the variations of the material have similar pore diameters, no matter the type of functional groups included. Though functional groups such as –OH and -OMe lead to a slightly larger size of the linker and more steric hindrance, the corresponding MOF samples show higher nitrogen uptake than that of the non-functionalized MIP-206, which could be ascribed to a larger number of structural defects and/or less residual impurities trapped in the functionalized structures other than the pure MIP-206 (Figures S9-S11, Tables S2-S6).

Pore Size Distribution (PSD) analysis of MIP-206 (Figure S8) suggests the presence of uniform meso-voids with a diameter of ca. 2.6 nm, which corresponds well to the
estimation from the crystal structure as well as the direct visualization of the hexagonal pore organization on the sample surface in high-resolution transmission electron microscopy (HR-TEM) images (Figure S12). As a result of the similar transition steps in the adsorption isotherms, samples tested for nitrogen sorption all possess similar mesopore diameters, within the range of 2.5-2.7 nm, which further highlights the negligible influence of bulky substitutes on the dimensions of the MOF porosity (Table S2).

Stability of the MOF is a very important parameter to be considered in assessing its suitability for practical use. Thermal stability of MIP-206 was first tested by thermogravimetric analysis (TGA) under an oxygen atmosphere and a temperature-dependent PXRD measurement carried out in open air. The long-range order within the 2D layer of the structural framework along the crystal b-axis is maintained up to ca. 450 °C, which corresponds to the maximum temperature point before the sharp drop in the TGA curve and the last crystalline PXRD pattern in the thermodiffraction profile (Figures S9 and S13). Hence MIP-206 is on the whole as stable as benchmark microporous Zr-MOFs such as UiO-66, and is significantly more thermally stable in the presence of oxygen than most of the carbon-based benchmarks and even comparable to some advanced molecular sieves^25.

Chemical stability of a material is required for industrial applications. In particular, hydrolytic resistance is highly prized as water is present in many processes. To assess its hydrolytic stability, a sample of MIP-206 was first soaked in liquid water either at room temperature (R.T.) or at boiling point. Neither cold water or hot water led to a
degradation of the MOF structural integrity even after long time periods (Figures S14 and S15, and Table S2). In this respect, MIP-206 clearly outperforms some conventional mesoporous benchmarks, such as MCM-4. MIP-206 also displays an excellent inertness in acidic conditions, such as concentrated HCl and even super acidic aqua regia, similar to the best microporous Zr-MOFs reported to date such as UiO-66 and MIP-200, highlighting a key advantage of this MOF that the vast majority of inorganic mesoporous materials do not share. In comparison with reported Zr-MOFs (microporous or mesoporous with micro-windows) constructed from similar small linkers with angled coordination sites, the superior stability of MIP-206 under acidic conditions might be related to the presence of the Zr\textsubscript{12} oxo-clusters SBU. Such an exceptional acidic resistance has indeed previously been documented for group 4 metal-carboxylate MOFs with high nuclearity SBUs, such as Hf\textsubscript{12} and Ti\textsubscript{12} oxo-clusters. In addition, its crystalline integrity can tolerate basic conditions until pH=12, which significantly extends the chemical stability of MIP-206 in aqueous solutions. To that end, it was reasonable to assume that MIP-206 could handle the harsh conditions related to hydrogen release from the catalytic decomposition of formic acid, which will be presented in the following sections. It should be noted that MIP-206 also displayed a good tolerance toward corrosive gas pollutants such as hydrogen sulfide (H\textsubscript{2}S) and sulfur dioxide (SO\textsubscript{2}) (Figures S15 and S16), which suggests that MIP-206 could also be a promising sorbent platform for potential toxic chemicals removal from contaminated environments such as polluted air and water.
Figure 3. Hydrogen release from formic acid catalyzed by MIP-206-OH supported Pd NPs. (A) and (B) HAADF-STEM images of Pd(0)@MIP-206-OH with different magnifications. (C) Catalytic performance comparison between Pd(0)@MIP-206-OH and Pd NPs immobilized on the other supports. (D) Cycling results of Pd(0)@MIP-206-OH in catalytic FA dehydrogenation reaction.

To demonstrate the potential of MIP-206 as a mesoporous support for heterogeneous catalysis, hydrogen generation from formic acid (FA) was selected as the model reaction. FA has been recognized as one of the most promising and safest hydrogen storage carriers for clean energy sources mainly due to its biomass origin, high hydrogen content, and easy transportation in its liquid state\textsuperscript{31,32}. Catalytic release of hydrogen from FA has been investigated and developed extensively\textsuperscript{31,33}. 
Heterogeneous catalysts have attracted particular attentions due to ease of separation and efficient recycling. As a class of promising heterogeneous catalysts, metal nanoparticles on porous supports (including oxides, silica, modified carbons and microporous MOFs) have performed adequately in the dehydrogenation of FA\textsuperscript{34; 35}. Though encouraging progress was achieved, notable flaws have been gradually noticed, including limitations in substrate diffusion and active species leaching and aggregation, resulting in inadequate catalytic activity and stability. Therefore, the design of supported metal catalysts featuring high activity, long-term stability, good reusability and facile process is highly sought after to advance the application of FA in hydrogen storage and generation. We expected that the unique features of MIP-206s could be suitable to match these requirements.

Due to their high activity toward this catalytic reaction\textsuperscript{36}, palladium nanoparticles (Pd NPs) templated by MIP-206 were tested initially to evaluate their corresponding performance. Encouraging activity of the catalyst was observed (Figure S17), which prompted us to further investigate its properties. MIP-206-OH was selected for all the following related investigations owing to its following features: 1) In comparison with the other functionalized MIP-206 family members, the hydroxyl-rich pore environment of MIP-206-OH is more comparable to those of mesoporous benchmarks, such as the silanol-rich silica, and hydroxyl or carboxylate-containing activated carbon. 2) MIP-206-OH was determined to be the optimal variation after taking into account MOF crystallinity, sample activation, pore accessibility and linker cost. After the systematic optimization of catalyst preparation and reaction conditions (Figures
S18-S21), MIP-206-OH loaded with 3.1 wt% of Pd NPs (Pd(0)@MIP-206-OH) exhibited the best catalytic performance at 60 °C under the optimized condition.

High-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images of the catalyst show a homogeneous distribution of small Pd NPs with a uniform size supported by the MIP-206-OH particles; the MOF template presents a similar morphology with no notable change compared to the pristine material (Figure 3A and B). Pd(0)@MIP-206-OH exhibits a complete release of H₂ in less than ten minutes under the optimized reaction condition, which is among the most active heterogeneous catalysts reported to date.³⁵ As shown in Figure 3C, the state-of-the-art mesoporous MOF support (MIL-101-Cr) suffers from an incomplete catalytic decomposition of FA due to the deactivation of Pd³⁷. Control experiments using the benchmark mesoporous Zr-MOFs, including NU-1000¹⁹ and PCN-224,³⁸ resulted in significantly lower activity and incomplete conversion of reactants due to the instability of the MOF structures and considerable Pd leaching under the catalytic conditions. The stable microporous UiO-66(Zr) displayed a poor performance, probably due to its micropores with limited accessibility. Benchmark mesoporous silica such as MCM-41 and SBA-15 demonstrated poor templating ability to form active Pd NPs. A commercially available mesocarbon material was also tested and exhibited moderate performance, mainly due to the considerable Pd leaching and eventual deactivation during catalysis. To further enhance the catalytic efficiency, we employed the aforementioned solid-solution-mixed-linkers strategy: the MTV-MIP-206-OH with partial PDA substitution was used as the Pd NPs support, as
the pyrazole moiety is known to interact specifically with Pd species, leading to the formation and stabilization of homogeneously distributed and smaller Pd NPs with even higher catalytic activity (Figure 4A and B).

Cycling ability is of utmost importance in heterogeneous catalysis. The durability of Pd(0)@MIP-206-OH was therefore evaluated by carrying out the catalysis for five consecutive runs, as shown in Figure 3D. It is worth noting that the catalytic performance is constant over the cycles with no obvious decrease, which indicates good reusability of the catalyst. Characterizations of the sample after cycling revealed that the characters of the catalyst, such as the Pd NP dispersion, the Pd loading content and the template integrity, was maintained (Figures S14, S15 and S18, and Tables S2 and S7). Therefore, both cycling catalytic performance tests and material characterizations fully support the good reusability of Pd(0)@MIP-206-OH in this reaction.

Furthermore, the influence of the MOF mesopore character on the catalytic performance was discussed. Firstly, catalytic activity of the composite increased with the decrease in size of the supported Pd NPs (Figure 4). For instance, fine Pd NPs loaded homogeneously on MTV-MIP-206-OH/PDA have a size distribution centered around 2 nm, and thus display an improved catalytic activity in comparison with that of the Pd NPs in a size of ca. 5 nm templated by MIP-206-OH when a similar pore accessibility of these two solid supports was observed. Similarly, lower activity of the Pd(0)@MIP-206 was noticed compared with Pd(0)@MTV-MIP-206-OH/FDA for the same reason (Figures S22 and S23). Secondly, the increased pore accessibility had a
positive effect on the catalytic activity, possibly due to the facilitation of reactant diffusion in larger pores. Pd NPs based on MIP-206-OH and MTV-MIP-206-OH/FDA have comparable size distributions but showed distinct catalytic performances, likely related to the differences in pore accessibility. Therefore, tunable catalytic activity of supported Pd NPs towards hydrogen generation from FA could be achieved via selecting a certain template candidate from the MIP-206s family with a suitable mesopore accessibility and a specific functional group attached.

Figure 4. Catalytic activity comparison between Pd NPs supported on different MIP-206s. (A) Different catalytic activity performance of Pd NPs templated by MIP-206, MIP-206-OH and MTV-MIP-206-OH/FDA and MTV-MIP-206-OH/PDA. (B) Comparison of Pd NPs size distributions between MIP-206-OH with MTV-MIP-206-OH/PDA.

In summary, we have demonstrated a Homometallic-Multicluster-Dot strategy to construct robust mesoporous MOFs materials from small linkers and non-toxic metals. MIP-206, a proof-of-concept example of this strategy that constructed from binary $\text{Zr}_6$ and $\text{Zr}_{12}$ oxo-cluster dot and feedstock isophthalic acid linker, possesses mesopores of
ca. 2.6 nm in free diameter with a good stability and tunability of pore chemical diversity. Our findings suggest that the MIP-206s not only function as ideal robust mesoporous supports for heterogeneous catalysts, but also hold considerable promises as a platform for a wide range of potential applications, such as heat reallocation, separation, removal of toxic chemicals and sensing.

**EXPERIMENTAL PROCEDURES**

**Synthesis of MIP-206**

MIP-206 was synthesized solvothermally by the reaction of ZrCl$_4$ and IPA in formic acid. IPA (1.1 g, 6.6 mmol) was weighed into a 23 mL Teflon reactor, formic acid (5 mL) was added followed by stirring at room temperature for 5 minutes until a homogeneous suspension was formed. ZrCl$_4$ (2 g, 8.6 mmol) was added to the suspension followed by 10 minutes stirring at room temperature to disperse the reactants uniformly. Afterwards, the reaction was sealed in an autoclave and heated to 180 °C in 2 hours and was kept at 180 °C for 24 hours. After cooling down to room temperature, the expected product of MIP-206 (1.98 g) was collected by filtration, washed with acetone and air dried. A scale-up synthesis could be achieved easily under the same reaction condition using a 125 mL Teflon reactor with a mixture of IPA (5.0 g, 30.0 mmol), ZrCl$_4$ (10.4 g, 44.4 mmol) and formic acid (30 mL) as reactants. 10.10 g of crude product of MIP-206 could be isolated. It is noteworthy that only high concentration of reactants could generate the MOF product. Otherwise clear solutions with no solid product were observed. The reaction duration was notably
shortened and the reaction yield was significantly improved along the increase of reaction temperature in the range of 120-180 °C.

**Synthesis of MIP-206-OH**

5-OH-IPA (0.72 g, 4.0 mmol) was weighed into a 23 mL Teflon reactor, formic acid (5 mL) was added followed by stirring at room temperature for 5 minutes until a homogeneous suspension was formed. ZrOCl₂ • 8H₂O (1.93 g, 6 mmol) was added to the suspension followed by 10 minutes stirring at room temperature to disperse the reactants uniformly. Afterwards, the reaction was sealed in an autoclave and heated to 180 °C in 2 hours and was kept at 180 °C for 20 hours. After cooling down to room temperature, the expected product of MIP-206-OH (2.01 g) was collected by filtration, washed with acetone and air dried. A scale-up synthesis could be achieved easily under the same reaction condition using a 125 mL Teflon reactor with a mixture of 5-OH-IPA (3.60 g), ZrOCl₂ • 8H₂O (9.65 g) and formic acid (25 mL) as reactants. 10.30 g of crude product of MIP-206-OH could be isolated.

**Synthesis of MTV-MIP-206/PDA**

IPA (0.88 g) and PDA (0.23 g, 20 mol%) were weighed into a 23 mL Teflon reactor, formic acid (5 mL) was added followed by stirring at room temperature for 5 minutes until a homogeneous suspension was formed. ZrCl₄ (2 g) was added to the suspension followed by 10 minutes stirring at room temperature to disperse the reactants uniformly. Afterwards, the reaction was sealed in an autoclave and heated to 180 °C in 2 hours and was kept at 180 °C for 24 hours. After cooling down to room temperature,
the expected product of MTV-MIP-206/PDA (1.83 g) was collected by filtration, washed with acetone and air dried.

**Synthesis of Pd(0)@MIP-206s**

10.8 mg of Pd(OAc)$_2$ was dissolved in 5 mL of acetone in a beaker under vigorous stirring at room temperature. 50 mg of MIP-206s was added and mixed for 1 hour for sufficiently absorbing the Pd precursor into the MOF pores. Subsequently, the solvent and residual metal precursor were removed by centrifugation. The collected solid was washed with anhydrous acetone twice. The sample was finally dried under vacuum at room temperature to give the Pd(II)@MIP-206s. Then the resulting solid was dispersed in 2.3 mL of H$_2$O. The required amount of NaBH$_4$ was dissolved in 1 mL of H$_2$O and added into the above mixture. After 5 min, the solid was collected by centrifugation, rinsed with H$_2$O for three times. The catalyst was dispersed in 0.6 mL of H$_2$O for dehydrogenation of formic acid (FA).

**Dehydrogenation of formic acid (FA)**

A mixture of as-synthesized catalyst and H$_2$O (0.6 mL) was placed in a two-necked round bottom flask (30 mL), which was placed in a water bath at 60 °C. A gas burette filled with water was connected to the reaction flask to measure the volume of released gas (room temperature kept at 25 °C). The reaction started when 0.33 mL of the mixed aqueous solution containing FA (46.7 mg) and sodium formate (SF) (68 mg) was injected into the mixture. The volume of the released gas was monitored by recording the displacement of water in the gas burette. In this reaction system, the SF
was used as a catalyst promoter. The durability test of Pd(0)@MIP-206-OH was performed at 60 °C. After each run, the catalyst was separated from the mixture and washed 3 times with H₂O. Then the catalyst was dispersed in 0.6 mL of H₂O and reused for next cycle. For each cycle, 0.33 mL of FA/SF solution (46.7 mg/68 mg) was injected to the reaction system. The experiment was repeated for 5 times.

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Author Contributions


Declaration of Interests

The authors declare no competing interests.
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