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## Introduction

Lewis acid catalysts are of considerable importance in modern organic chemistry because of their unique privilege in controlling reactivity and selectivity.<sup>1</sup> Numerous examples explored thus far have demonstrated that sufficient activation of substrates can not only accelerate reactions but also affect regio- and stereoselectivity. Although this arena has been expanded to encompass a diverse range of organic transformations, classical Lewis acids are moisture-sensitive and thus corrosive because of evolved acidic gases. In this context, the advent of water-compatible Lewis acids<sup>2</sup> was the cornerstone of green organic synthesis, accentuating their power in organic synthesis thereafter. In the meantime, notwithstanding the conspicuous prosperity of bioorthogonal catalysis,<sup>3,4</sup> creating biocompatible Lewis acidic analogues compatible within the sea of cellular nucleophiles remains arduous. Because of their high electrophilicity, Lewis acids may be poisoned by Lewis basic functionalities such as amines and thiols present in proteins or biogenic small molecules, glutathione (GSH) and carnosine. This dysfunction of Lewis acids in cellular environments downgrades their potential applicability. Although chiral Lewis

acids are too vulnerable to Lewis basic environments to deliver deteriorated yield and selectivity, nature executes Lewis acid-catalyzed enantioselective synthesis to function appositely in a living cell. While heterogeneous platforms, such as hydrophobic polymer and surfactant assemblies, embedding chiral Lewis acid complexes have been adopted for enantioselective reactions in aqueous environments,<sup>5–8</sup> they are inept for biomedical purposes because of their aggregation behavior. An aspirational goal would therefore be to set the stage for chiral Lewis acids that allow enantioselective reactions even in the presence of cellular debris. While there have been a handful of successes in precious metal-based enantioselective reactions within living cells,<sup>9</sup> successful applications of chiral Lewis acid catalysis rely absolutely on robust copper(II) complexes.<sup>10,11</sup>

We envisioned implementing nanoscale metal-organic frameworks (MOFs) as an alternative platform to this end. Owing to their high surface area, permanent porosity, highly tunable pore structures, and flexible functionality, MOFs have emerged as novel devices with a wide spectrum of potential applications: gas storage, chemical sensing, biomedical imaging, drug delivery, and catalysis.<sup>12,13</sup> Notwithstanding the brilliant successes of MOFs in diverse chemical transformations, two potential challenges that may limit their versatility can be identified: application to their chiral variant is often bottlenecked by the limited number of available chiral linkers and the limited techniques of postsynthetic modification.<sup>14</sup> In addition, the involvement of MOFs in organic synthesis in aqueous solutions has barely been reported.<sup>15</sup> This

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is surprising because of the inherent stability of nano-MOFs (NMOFs) in water<sup>16</sup> and their biomedical applications,<sup>17–20</sup> including drug delivery, cancer therapy, and medical diagnosis. In addition, we conceived that the confinement effects of NMOFs were fit for providing compartmentalized environments to conserve Lewis acidic sites from the sea of cellular nucleophiles. We herein report our efforts to address *ex vivo* the poisoning of chiral Lewis acid catalysts by biogenic nucleophiles.

## Results and discussion

To exploit the chemical and thermal stability of Zr-based UiOs comprising robust zirconium–carboxylate bonds between oxocluster nodes and organic linkers,<sup>21</sup> we intended to translate UiO-67 MOFs composed of a bipyridine linker to their chiral variant (Fig. 1a). 2,2'-Bipyridines have been extensively exploited for establishing numerous metal complexes in catalysis because of their robust redox stability, ease of functionalization, and entropically favored metal chelation.<sup>22</sup> Inspired by the simplicity and the ease of access to water-stable multidentate coordination of chiral 2,2'-bipyridine bearing 6,6'-dicarbinol substituents **1**,<sup>23</sup> our study commenced with identifying the best **1**-based linker for asymmetric catalysis. We opted for a divergent approach using an optically active intermediate **7** as a common intermediate to generate optically active NMOFs with UiO-67 topology (Fig. 1b). The length of candidate linkers is perceived to exhibit a sufficient size-exclusion effect to safeguard Lewis acidic metal ions encapsulated inside UiO-type NMOFs against nucleophilic residues of proteins. The scandium ion is a potent candidate to obtain proof-of-concept for

the hypothesis due to its unique catalytic performance to other d-block elements. Ueno *et al.* leveraged the chiral environment of the  $\beta$ -helical tubular motif found in bacteriophage T4 for scandium catalysis. A covalent linkage of maleimide-substituted 2,2'-bipyridine to a cysteine residue and subsequent metalation led to the scandium enzyme, which catalyzed the asymmetric ring-opening reactions of *meso*-epoxides (Fig. 1c).<sup>24</sup> The catalyst showed catalytic activity in the presence of nucleophilic residues, while the reaction suffered from low conversion and poor enantioselectivity even with the aid of organic cosolvent. In this work, we thus selected the asymmetric ring-opening reaction as a benchmark reaction to pursue both compatibility with biogenic nucleophiles and high catalytic performance.

The synthesis of optically active 2,2'-bipyridyl linkers is outlined in Fig. 2a. The synthesis of common intermediate **7** commenced with asymmetric transfer hydrogenation of compound **6** prepared from 2,6-dibromopyridine in 3 steps (Fig. 2a). Of note, adoption of type IIIb conditions<sup>25</sup> (aqueous suspension) proved to be pivotal for asymmetric transfer hydrogenation because canonical conditions (formic acid–triethylamine mixture) failed to produce the product. Homocoupling of **8** and ensuing deprotection processes provided the chiral 5,5'-diformyl-2,2'-bipyridine **10**. The chromium-based oxidation delivered 2,2'-bipyridine-5,5'-dicarboxylic acid (H<sub>2</sub>BPDC **2**). Notwithstanding the successful assembly of the MOF upon solvothermal treatment with ZrCl<sub>4</sub> and trifluoroacetic acid, this route faltered due to the severe degradation in the postsynthetic deacetylation as well as unsuccessful deprotection before assembly. The use of **4** proved unfeasible due to the deprotection matter as well, despite the successful installation of an alkyne functionality through the Corey–Fuchs homologation of **10** and subsequent reaction with CO<sub>2</sub> with alcohols protected by TBS. In the third plan, the Verley–Dobner modification of the Knoevenagel condensation and the subsequent deacetylation converted **10** to H<sub>2</sub>BPV **3**. The solvothermal synthesis of a Zr-based MOF from **3** became deadlocked by the unwanted intramolecular 1,4-addition of an alcohol group to a cinnamic acid moiety that was deduced from the <sup>1</sup>H NMR spectra of the digested sample. Thus, linker **5** emerged as an alternative way to bypass the issue. The formyl groups of **7** were elongated through Wittig olefination, which underwent a reductive homocoupling reaction and subsequent deprotection to afford H<sub>2</sub>BPVB **5** with perfect *E* configuration. A solvothermal reaction of **5** with ZrCl<sub>4</sub>, acetic acid, and H<sub>2</sub>O in *N,N*'-dimethylformamide (DMF) at 40 °C for 5 days afforded the targeted chiral BPVB-NMOF with a UiO framework as a pale yellow solid (Fig. 2b), despite the length of the organic linker being much larger than ever reported for Zr-MOFs.<sup>26,27</sup> In the structure, the 2,2'-bipyridyl moiety was deemed to adopt *cis* conformation considering the conformational preference of 2,2'-bipyridyl under moderately acidic conditions.<sup>28,29</sup> The presence of 6,6'-carbinols may restrict the inner rotational freedom after assembly, conferring benefits for postsynthetic metalation. The powder X-ray diffraction (PXRD) patterns of the as-synthesized MOF include the UiO-type structural fingerprint conformable with the reported pattern<sup>30</sup> with line broadening

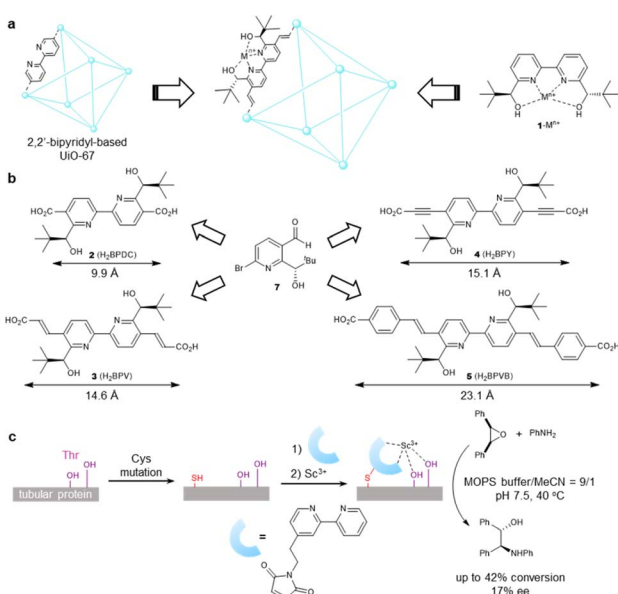
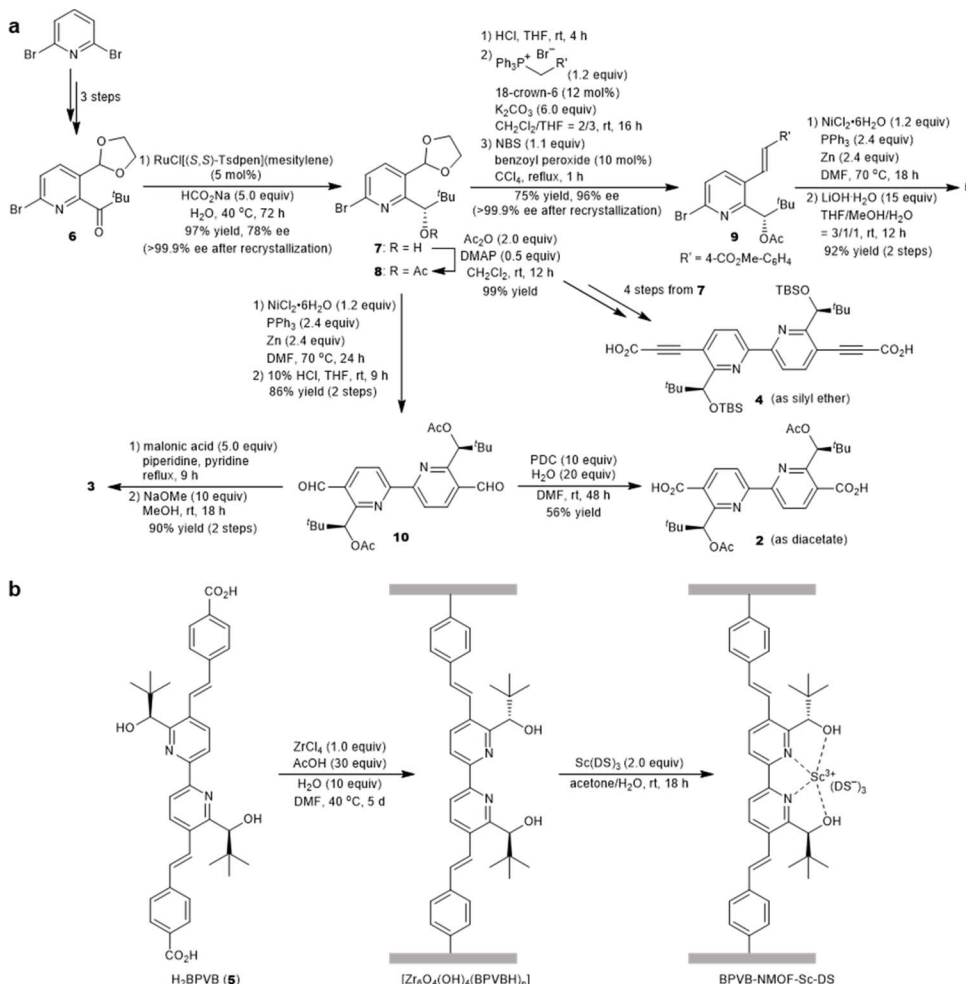


Fig. 1 Design of biocompatible Lewis acid catalysts. (a) Concept of catalyst design. (b) Optically active 2,2'-bipyridyl linkers surveyed in this work. The values represent the length between carbon atoms of dicarboxylate. (c) Previous example of scandium-based Lewis acid metalloenzyme.





**Fig. 2** Design and synthesis of chiral BPVB-NMOF-Sc-DS. (a) Divergent synthesis of optically active 2,2'-bipyridyl linkers from common intermediate **7**. (b) Preparation of BPVB-NMOF and postsynthetic metalation. DS stands for dodecyl sulfate. See the ESI† for details.

due to the small size of the nanoparticles<sup>31</sup> (Fig. 3c). The predominance of the (002) diffraction peak over (111) may originate from the richness of a hexagonal centered planar (hcp) structure over the usual face centered cubic (fcc) structure, indicating the presence of missing linker defects.<sup>32,33</sup> Because the increased activity in ring-opening reactions of epoxides was reported for the hcp UiO-66,<sup>34</sup> the presence of sufficient linker defects bodes well for exploration of chiral Lewis acid catalysis. The scanning electron microscopy (SEM) image also confirmed the nanostructure construction with an average diameter of 125 nm (Fig. 3a). Thermogravimetric analysis (TGA) revealed the missing linker defect to be 45% for a fcc structure and 25% for a hcp structure from the mass at 200 °C wherein the MOF is fully activated to the dehydroxylated compositions to the point of complete linker degradation (Fig. 3d). The open hysteresis found in the  $\text{N}_2$  adsorption–desorption isotherm may be attributed to irregular mesoporosity derived from such a high concentration of defects (Fig. S24†). To configure inner hydrophobic environments with increasing stability, we envisioned using hydrophobic anion, dodecyl sulfate (DS), for scandium installation. Because the length of dodecyl sulfate is 20.8 Å,

dodecyl sulfate anions would interact with one another to form a “pseudomicelle” aggregate within a confined space in the UiO framework. This “pseudomicelle” is expected not to form a larger aggregate due to the confinement effect. Incubation with scandium tris(dodecyl sulfate) ( $\text{ScDS}_3$ ) in aqueous acetone solution led to the formation of a yellow precipitate. The collected powder was dried under vacuum at room temperature to afford BPVB-NMOF-Sc-DS.

Inductively coupled plasma (ICP) analysis revealed the average Sc content to be 0.5 mmol g<sup>-1</sup>. The N loading is twice as high as the Sc loading and the average Sc/Zr ratio was 55%, suggesting quantitative installation of the scandium ion into the UiO framework whether it is fcc or hcp. A blue shift of the interfacial N 1s peak by 1.6 eV in X-ray photoelectron spectroscopy (XPS) analysis also confirmed the coordination of the scandium ion to 2,2'-bipyridyl moieties (Fig. S26 and S27†). Because the tetradentate coordination of  $\text{Sc}^{3+}$  with two nitrogen atoms of 2,2'-bipyridine and two oxygen atoms of alcohols has been reported to be important for sufficient reaction efficiency<sup>24</sup> and enantioselection,<sup>35</sup> we anticipated analogous coordination environments, as corroborated by

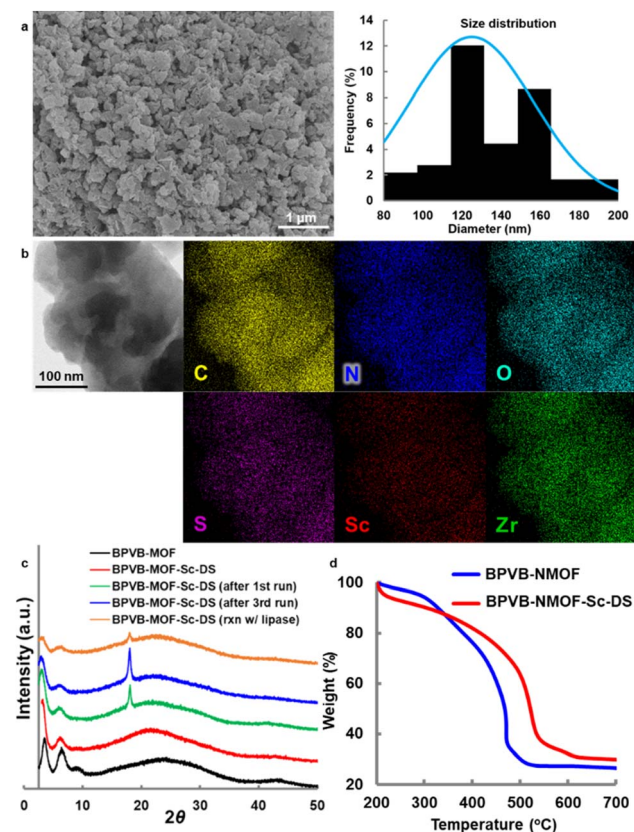


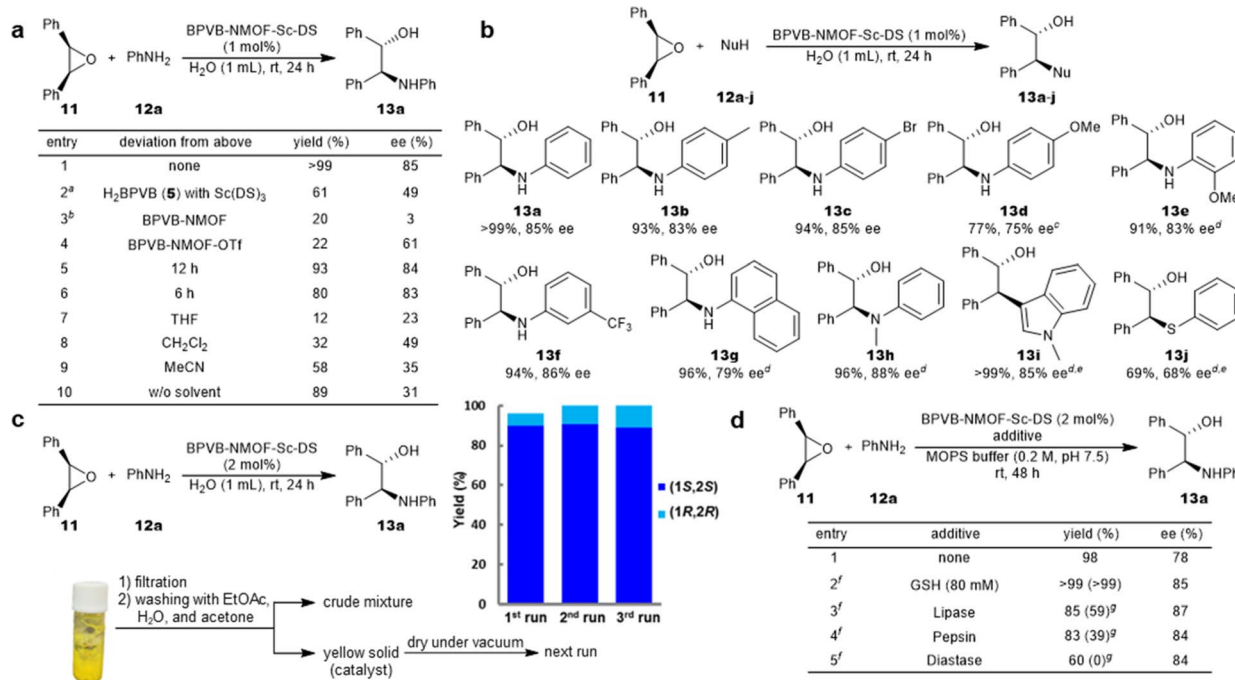
Fig. 3 Characterization of BPVB-NMOFs. (a) SEM image of BPVB-NMOF and size distribution histogram. (b) STEM and EDS analysis of BPVB-NMOF-Sc-DS. (c) PXRD patterns of BPVB-NMOFs. (d) TGA plot. See the ESI† for details.

relatively high stability in water, activity comparable to Ueno's result (Fig. S8†), and high enantioselectivity. Energy dispersive X-ray spectroscopy (EDS) tomography confirmed the uniform distribution of all elements in BPVB-NMOF-Sc-DS particles (Fig. 3b). The coordination of scandium lengthened the linker length by  $\sim 0.2$  Å due to an increased torsion angle, thereby provoking subtle changes in morphology, especially a relative decrease in the fcc structure, as evidenced by the PXRD pattern (Fig. 3c). The significant isotherm transition from type II to type III after uptake of  $\text{Sc}(\text{DS})_3$  suggested the occupancy of the interior space of a NMOF with dodecyl sulfate (Fig. S21†). The increased stability of BPVB-NMOF-Sc-DS particles was confirmed by TGA (Fig. 3d). This may be triggered by a loss of a certain amount of a linker, as shown in variability in Sc loadings. The scanning transmission electron microscopy (STEM) image of trifluoromethanesulfonate variant BPVB-NMOF-OTf manifested the presence of aggregates presumably derived from  $\text{Sc}(\text{OTf})_3$  (Fig. S19†), bespeaking the importance of dodecyl sulfate on the precise coordination with 2,2'-bipyridyl moieties within the framework. This is also corroborated by scandium loading that is more than 1.2 times higher than that of BPVB-NMOF-Sc-DS. The further area EDS analysis revealed the formation of scandium oxide or hydroxide clusters after hydrolysis (Fig. S23†).

The performance of the NMOF-Sc catalyst was evaluated in the asymmetric ring-opening reactions of epoxides with aniline in water (Fig. 4). A Sc loading of 1 mol% proved sufficient for the efficient reaction, delivering the desired product **13a** quantitatively with 85% ee (Fig. 4a, entry 1). The reaction outcome proved not dependent on the Sc loading (Table S11†). No leaching of  $\text{Sc}^{3+}$  and  $\text{Zr}^{4+}$  both in the aqueous and organic phases is noteworthy (Table S10†). It is also worth pointing out that the reaction was highly enantioselective, notwithstanding the decent level of activity of BPVB-NMOF (entry 3). The activity of UiO series can be explained using the Lewis acidity of zirconium<sup>36,37</sup> or the Brønsted acidity of zirconium-bound hydroxide,<sup>38</sup> which is supposed to affect the enantioselectivity when the NMOF-Sc catalyst is used. A summary of control experiments is outlined in entries 2–9. The significant decrease in both yield and enantioselectivity of the reaction using linkers  $\text{H}_2\text{BPVB}$  5 and  $\text{Sc}(\text{DS})_3$  supported the desired catalytic performance inside the framework (entry 2). The catalyst incubated with  $\text{Sc}(\text{OTf})_3$ , BPVB-NMOF-Sc-OTf, displayed poor activity along with moderate enantioselectivity (entry 4). This denotes that the framework alone fails to offer sufficient hydrophobicity for the reaction environments as well as that superfluous mounting of  $\text{Sc}^{3+}$  inside the framework is detrimental. This is consistent with our "pseudomicelle" hypothesis. Carrying out the model reaction with shorter reaction times still provided high product yields (entries 5 and 6). Organic solvents such as THF,  $\text{CH}_2\text{Cl}_2$ , and MeCN proved inappropriate (entries 7–9). Besides, the solvent-free reaction suffered from low enantioselectivity (entry 10), manifesting the pivotal role of water in controlling the stereoselectivity. With the optimal conditions in hand, the scope of the substrate was displayed as depicted in Fig. 4b. A variety of anilines could be employed irrespective of the electronic nature of substituents to afford optically active  $\beta$ -amino alcohols **13b–h** in excellent yields ( $>90\%$ ) with high selectivities (up to 88% ee). The reaction proved scalable without deterioration of the reaction outcome (**13a**). In addition, the range of applicable nucleophiles was expanded to include 1-methyl indole and thiophenol that underwent C–C and C–S bond formations (**13i** and **13j**), respectively. BPVB-NMOF-Sc-DS could be recovered by a simple filtration with no leaching of  $\text{Sc}^{3+}$  and then subjected to the next runs, resulting in excellent yields and high ee over three cycles (Fig. 4c). A facile separation of the NMOF catalyst from a post-reaction mixture probed our hypothesis of confined micelle formation. The gradual decline in the PXRD pattern of the fcc structure over the cycles may be ascribed to physical damage to the NMOF during the filtration (Fig. 3c). The emerging peak may be the impurities derived from the reactants, judging from the inconsistency with the reported peaks of scandium oxides or zirconium oxide.

The compatibility of the NMOF catalyst in the presence of biogenic nucleophiles was evaluated as shown in Fig. 4d. The reactions were run in MOPS buffer with comparison to micellar catalysis (type IIb conditions)<sup>25</sup> using  $\text{Sc}(\text{DS})_3$  and **1**. When GSH was loaded into the system, the reaction was promoted efficiently to provide excellent yield and high enantioselectivity in both cases (entry 2). The absence of negative effects can be explained by the absence of uptake of GSH in the interior of the





**Fig. 4** BPVB-NMOF-Sc-DS-catalyzed asymmetric ring-opening reactions of epoxides in aqueous media. (a) Control experiments. Standard reaction conditions: **11** (0.2 mmol), **12a** (0.3 mmol), BPVB-NMOF-Sc-DS (1 mol%, 0.002 mmol), H<sub>2</sub>O (1 mL), rt, 24 h. Isolated yields after chromatography are given. <sup>a</sup>H<sub>2</sub>BPVB (1 mol%) and Sc(DS)<sub>3</sub> (1 mol%) were premixed prior to the addition of reactants. <sup>b</sup>Zr loading was controlled as the same as that under the standard conditions (2.8 mol% Zr). (b) Substrate scope: **11** (0.2 mmol), **12** (0.3 mmol), BPVB-NMOF-Sc-DS (1 mol%, 0.002 mmol), H<sub>2</sub>O (1 mL), rt, 24 h. Isolated yields after chromatography are given. <sup>c</sup>Three equivalents of nucleophile were used. <sup>d</sup>BPVB-NMOF-Sc-DS (2 mol%) was used. <sup>e</sup>Run for 48 h. (c) Reuse experiments: **11** (0.4 mmol), **12a** (0.6 mmol), BPVB-NMOF-Sc-DS (2 mol%, 0.008 mmol), H<sub>2</sub>O (2 mL), rt, 24 h. The experimental procedure for catalyst recycling is briefly shown. The scale of the next run was calculated based on the Sc loading of the recovered catalyst. (d) Reaction in the presence of biogenic nucleophiles: **11** (0.2 mmol), **12a** (0.3 mmol), BPVB-NMOF-Sc-DS (2 mol%, 0.004 mmol), MOPS buffer (2 mL), rt, 48 h. Isolated yields after chromatography are given. <sup>f</sup>The additive (25 g L<sup>-1</sup>) was added. <sup>g</sup>Run with Sc(DS)<sub>3</sub> (2 mol%, 0.004 mmol) and **1** (2.4 mol%, 0.0048 mmol) instead of BPVB-NMOF-Sc-DS. DS = dodecyl sulfate. MOPS = 3-morpholino-1-propanol-1-sulfonic acid. See the ESI† for details.

micelle due to the small size of GSH. The reaction was therefore carried out in the presence of proteins to understand the tolerance to biogenic nucleophiles because the formation of SDS-protein composites was reported.<sup>39</sup> As a result, the activity of BPVB-NMOF-Sc-DS proved to be higher than that of the micellar catalysis in the presence of proteins. Of note, the activity of Sc(DS)<sub>3</sub> was completely inhibited in the presence of diastase, underpinning the proof of concept in this study. Because diastase is stable in acidic or neutral solutions,<sup>40</sup> the severe poisoning effects may have been exerted for the micellar catalysis. The confined environments of the NMOF are suited for the exclusion of these proteins: *e.g.* pepsin (48 Å × 64 Å,<sup>41</sup> the smallest among three).

## Conclusions

In summary, we have successfully developed a new UiO-67(BPY) MOF composed of a chiral 2,2'-bipyridyl-functionalized dicarboxylate linker (H<sub>2</sub>BPVB). The limited free rotation of 2,2'-bipyridyl by introduction of a 6,6'-carbinol moiety and fabrication of a defect-rich structure due to longer linker length resulted in a quantitative postsynthetic uptake of scandium salt. The use of Sc(DS)<sub>3</sub> was of vital importance for our design to precisely emplace Sc<sup>3+</sup> and to forge sufficiently hydrophobic

“pseudomicelle” environments sequestered inside the framework. Such a concept of the confined micelle is, to our knowledge, without precedent. The as-prepared BPVB-NMOF-Sc-DS efficiently catalyzed the asymmetric ring-opening reactions of epoxides in water without any organic cosolvent to deliver high yields and enantioselectivities with a broad range of nucleophiles. The impaired performance in pure organic solvents or under solvent-free conditions adorned this catalysis. The reusability of the catalyst by expedient filtration underscores its relative robustness even in water. Remarkably, this NMOF catalyst showed superior catalytic performance to the usual micellar catalysis in the presence of proteins, demonstrating the ability to prevent active Lewis acid sites from mortiferous deactivation. The ways of designing and preparing water-stable chiral NMOFs represent a unique direction for synthetic organic chemistry to build an abiological path without impairing homeostasis. As such, it warrants further study to build a bridge between organic chemistry and biochemistry.

## Data availability

All the data supporting this study are included in the main text and the ESI.†

## Author contributions

T. K. and S. K. conceived and directed the study. S. W. performed the experiments. T. K. and S. W. analyzed the data. T. K. performed DFT calculations to optimize the structure of chiral linkers. All the authors prepared the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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