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Robust acid–base Ln-MOFs: searching for efficient catalysts in cycloaddition of CO₂ with epoxides and cascade deacetalization–Knoevenagel reactions†

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A family of microporous and robust Ln(III)-based metal–organic frameworks (**1-Ln**, Ln = Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb) have been obtained using 4,4',4''-nitrotribenzoic acid (H₃NTB) in NMP–HCl solvent. Both single-crystal and powder X-ray diffraction analyses demonstrate that **1-Ln** are isostructural and possess 3D frameworks with permanent porosity for Ar and CO₂ adsorption. Strikingly, the incorporation of both Lewis acidic lanthanide ions and the basic triphenylamine group into **1-Ln** makes them efficient acid–base catalysts for both cycloaddition of epoxides with CO₂ and one-pot cascade deacetalization–Knoevenagel reactions. The systematic catalytic studies show that **1-Tb** and **1-Yb** possess the best catalytic activities for both reactions, indicating the catalytic activities of these Ln-MOFs are strongly dependent on metal Lewis acid sites embedded in the frameworks.

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Introduction

Recently, lanthanide based metal–organic frameworks (Ln-MOFs) have been intensely investigated as heterogeneous catalysts because of their unique features, such as inherent porosity, tunable pore sizes, reusability, and a broad range of catalytic sites.^{1–3} In principle, the active sites of Ln-MOF catalysts can be (i) lanthanide ions with unsaturated coordination environments, (ii) organic or organometallic functional sites realized using pre-designed ligands, and (iii) other catalytic species encapsulated in the pores. The integration of these catalytic sites into one material would promise the Ln-MOFs a powerful platform for heterogeneous catalysis. Currently, Ln-MOFs as Lewis acids have been applied for a number of organic transformations, including cycloaddition of CO₂, Knoevenagel condensation, and Heck reactions.^{4–6}

Carbon dioxide (CO₂), as the primary greenhouse gas, is also a nontoxic, economic, and renewable C1 building block. CO₂ capture and conversion into value-added chemicals are attractive in terms of green chemistry.^{7,8} Particularly, the cycloaddition of epoxides and CO₂ into cyclic carbonates is considered to

be one of the most promising approaches for CO₂ utilization, given the wide industrial applications of the target cyclic carbonates.^{9,10} In recent years, it has been reported that some porous Ln-MOFs have shown preeminent catalytic performances on cycloaddition reactions of epoxides and CO₂, mainly because these MOFs feature selective CO₂ capture and lanthanide ions as Lewis acid sites to activate epoxide.^{11–15}

On the other hand, among various catalytic processes, cascade reactions are sequentially proceeded with two or multiple transformations in one-pot, which eliminate the time-consuming isolation and purification of intermediates.^{16–18} However, some cascade reactions require the catalysts with antagonistic catalytic sites (e.g., Lewis acidic sites for the first step and basic sites for the second step in deacetalization–Knoevenagel reaction or deacetalization–Henry reaction). Therefore, the catalyst design toward these one-pot cascade reactions is challenging.^{19–23} In this respect, Ln-MOFs offer the unique possibility of combining isolated Lewis acidic and basic active sites for cascade or cooperative catalysis. To date, several acid–base bifunctional MOFs have shown to be high-performance catalysts for one-pot cascade reactions.^{4,11,24–26}

The Knoevenagel reaction as one important synthetic method for the C–C bond formation, is typically catalyzed by base,²⁷ and triphenylamine groups in a MOF as a weak base for the Knoevenagel reaction have been reported by Duan's and Wen's group.^{28,29} Meanwhile, the lanthanide ions embedded in Ln-MOFs have proven to be valuable Lewis acid sites for a number of organic reactions, as mentioned before.^{4–6} Bearing these considerations in mind, one would expect a Ln-MOF incorporating triphenylamine group to be an acid–base

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bifunctional catalyst for both cycloaddition of epoxides with CO_2 and one-pot deacetalization–Knoevenagel reaction.

In this work, using 4,4',4''-nitritotribenzoic acid (H_3NTB , a typical triphenylamine ligand) and lanthanide nitrates dissolved in NMP–HCl mixture, we have solvothermally synthesized a new family of 3D microporous Ln-MOFs (denoted **1-Ln**, $\text{Ln} = \text{Sm}, \text{Eu}, \text{Gd}, \text{Tb}, \text{Dy}, \text{Ho}, \text{Er}, \text{Yb}$). The structural analyses reveal **1-Ln** are intrinsically isostructural, which offer the opportunity to determine the effect of the metal center on catalytic performance. All **1-Ln** solids exhibit remarkable robustness and permanent porosity evaluated by Ar adsorption analysis. Their CO_2 adsorption at 273, 298 K/1 bar are also investigated. Interestingly, the incorporation of both Lewis acidic lanthanide ions and basic triphenylamine group into one single framework makes them highly efficient catalysts for both cycloaddition of epoxides with CO_2 and one-pot cascade deacetalization–Knoevenagel condensation reactions. Among them, **1-Tb** and **1-Yb** possess the best catalytic activities for both two reactions. More strikingly, the systematic catalytic studies reveal that the catalytic activities of these Ln-MOFs are strongly dependent on Lewis acidic lanthanide ions embedded in the framework, uncovering the possibilities to develop Tb- and Yb-MOFs as high-performance Lewis acid catalysts.

Results and discussion

Structure analyses of **1-Ln** crystals

Solvothermal reactions of different lanthanide salts with H_3NTB in NMP–HCl mixture harvest a family of crystalline Ln-MOFs, **1-Ln** ($\text{Ln} = \text{Sm}, \text{Eu}, \text{Gd}, \text{Tb}, \text{Dy}, \text{Ho}, \text{Er}, \text{Yb}$), which are isostructural as checked by powder X-ray diffraction (XRD, Fig. 2) and infrared spectroscopy (Fig. S3†). Under the same synthetic conditions, $\text{LaCl}_3 \cdot 6\text{H}_2\text{O}$ affords a different structure that would be reported in due course. Single crystal X-ray diffraction (SCXRD) analyses of **1-Tb** and **1-Yb** show that they crystallize in the triclinic space group $P\bar{1}$ (Table S1†) and have the formula of $\{[\text{Ln}(\text{NTB})(\text{NMP})] \cdot (\text{NMP})\}_m$, which are in good accordance with their TG analyses (Fig. S4†). SCXRD analyses reveal **1-Ln** have a porous architecture (Fig. 1 and 2). Therefore, only the structure of **1-Yb** is briefly discussed herein. The asymmetric unit of **1-Yb** contains one Yb^{3+} cation, one NTB^{3-} ligand, one coordinated and one free guest NMP molecule (Fig. S1†). As shown in Fig. 1a, each Yb^{3+} is coordinated by seven oxygen atoms from one NMP molecule and six $\mu_2\text{-}\eta^1\text{:}\eta^1$ carboxylate groups that belong to six different ligands. Interestingly, every two Yb^{3+} ions are four-fold bridged by four $\mu_2\text{-}\eta^1\text{:}\eta^1$ carboxylate groups to form a dimeric unit. These dimeric units are further two-fold bridged by another two $\mu_2\text{-}\eta^1\text{:}\eta^1$ carboxylate groups to generate a 1D Yb-COO inorganic chains running along [100] (Fig. 1b). These Yb-COO inorganic chains are connected by NTB^{3-} ligands to define the three-dimensional porous framework (Fig. 2). The framework possesses two kinds of rhombus channels, one with diagonals of $7.1 \times 5.9 \text{ \AA}^2$ (measured between the opposite atoms of $\text{N}\cdots\text{N}$ and $\text{Yb}\cdots\text{Yb}$, and considering the van der Waals radii) along [100] which are aligned by coordinated NMP molecules, and the other one with diagonals of $8.8 \times 5.8 \text{ \AA}^2$ (measured between the opposite atoms of $\text{N}\cdots\text{N}$ and

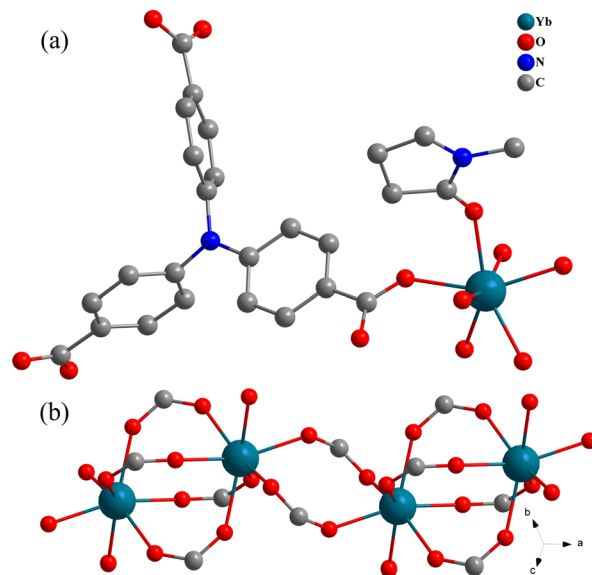


Fig. 1 (a) The coordination environment around Yb^{3+} in the **1-Yb**; (b) 1D Yb-COO inorganic chain.

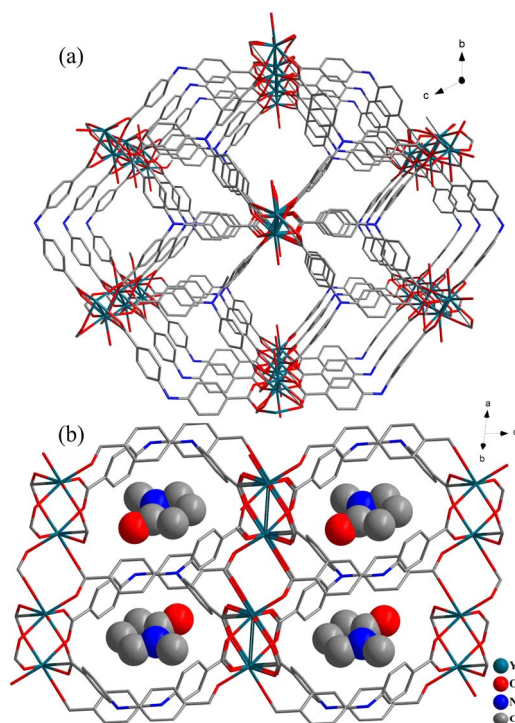


Fig. 2 (a) 3D porous framework of **1-Yb** showing channels along [100], with removal of all coordinated and guest NMP molecules for clarity; (b) perspective view of channels along [110], with NMP guest molecules occupying the channels.

$\text{Yb}\cdots\text{Yb}$) along [110] which are filled by guest NMP molecules. After removal of all NMP molecules, the free void calculated by PLATON is up to 47.4% of the crystal volume. Thus, **1-Yb** is a highly microporous framework, which would be suitable for gas adsorption. Moreover, the incorporation of Lewis acidic



lanthanide centers and Lewis basic triphenylamine sites in the channels indicates that **1-Yb** could be a potential acid–base heterogeneous catalyst. With these structures (**1-Sm**, Eu, Gd, Tb, Dy, Ho, Er, Yb) in hand, comparative experiments were performed to determine the effect of the metal centers on catalytic performance and to search for more efficient catalysts.

PXRD analyses, thermal stability of the frameworks, Ar and CO₂ adsorption isotherms

The PXRD profiles of as-synthesized **1-Ln** solids match well with their simulated ones, verifying the phase purity of the as-synthesized bulk samples (Fig. S2†). Furthermore, all **1-Ln** solids display similar and excellent thermal stability (Fig. S4†). As shown by thermogravimetric analyzer (TGA), the guest-free phases of **1-Ln** are thermally stable up to 500 °C. This exceptional robustness of the frameworks is also observed in other Ln-MOFs structurally built on Ln-COO inorganic chains.^{30–33}

Ar adsorption experiments were performed at 87 K to assess the permanent porosity of all compounds. The adsorption isotherms (Fig. 3 and S5†) demonstrate type-I adsorption which is typical of microporous materials, with the BET (Brunauer–Emmett–Teller) surface area of 113.7 (**1-Sm**), 128.1 (**1-Eu**), 222.0 (**1-Gd**), 299.0 (**1-Tb**), 268.8 (**1-Dy**), 261.2 (**1-Ho**), 274.7 (**1-Er**), and 300.7 (**1-Yb**) m² g^{−1}, respectively. The total pore volumes are 0.105 (**1-Sm**), 0.111 (**1-Eu**), 0.152 (**1-Gd**), 0.204 (**1-Tb**), 0.176 (**2-Dy**), 0.182 (**1-Ho**), 0.202 (**1-Er**), and 0.209 (**1-Yb**) cm³ g^{−1}, respectively. These values are comparable to other porous lanthanide MOFs.^{33–35}

Encouraged by the robustness and permanent porosity of **1-Ln**, we investigated their CO₂ adsorption behaviors. It is clearly demonstrated that all **1-Ln** solids are readily accessible to CO₂ (Fig. S5†). For **1-Tb** and **1-Yb**, their CO₂ adsorption isotherms at 273 and 298 K are shown in Fig. 3. The maximal CO₂ capacities

is 27.5, 15.9 cm³ g^{−1} (for **1-Tb**) and 29.4, 16.5 cm³ g^{−1} (for **1-Yb**) respectively. In order to gain more insights into the framework–CO₂ interactions, adsorption heat of CO₂ (Q_{st}) was calculated by using the viral equation based on the isotherms at 273 K and 298 K. As shown in Fig. S6,† the Q_{st} value at zero coverage is around 22.5 kJ mol^{−1} (for **1-Tb**) and 25.0 kJ mol^{−1} (for **1-Yb**), respectively, which are comparable to some MOFs such as [Zn₂(L)(dpe)₂](NO₃)·(DMF)₃·(H₂O)₂ (20.5 kJ mol^{−1}; L = 1,3-bis(3,5-dicarboxyphenyl)imidazolium, dpe = 1,2-di(4-pyridyl) ethylene)³⁶ and FJU-14-BF₄ (18.8 kJ mol^{−1}),³⁷ indicating moderate interactions between the framework and guest CO₂ molecules.

Catalytic cycloaddition of CO₂ and epoxides

On account of the considerable CO₂ adsorption quantity, the accessible channels, and the coexisting Lewis acid–base sites, the catalytic performances of **1-Ln** towards the cycloaddition of CO₂ and epoxides into cyclic carbonates are investigated. We wonder the effect of the metal center on catalytic performance among the eight Ln-MOF candidates (**1-Sm**, Eu, Gd, Tb, Dy, Ho, Er, Yb). Firstly, with **1-Yb** and *n*-Bu₄NBr (tetra-*n*-tertbutylammonium bromide) as catalytic system, the cycloaddition between CO₂ and epichlorohydrin under ambient temperature has been used as a model reaction to screen the optimal conditions. After detailed investigations, the optimal conditions are set as follows: epichlorohydrin (4 mmol), CO₂ (1 atm), **1-Yb** catalyst (0.02 mmol based on Ln metal sites; 0.5 mol%) and cocatalyst *n*-Bu₄NBr (0.2 mmol; 5 mol%), room temperature, 36 hours. As seen in Table 1, in the case of the reaction without any catalyst (entry 1), the reaction does not proceed. Without

Table 1 Cycloaddition of CO₂ and epichlorohydrin with different **1-Ln** catalysts^a

Entry	Catalyst	Catalyst (mol%)	<i>n</i> -Bu ₄ NBr (mol%)	Yield ^b (%)
1	—	0	0	<1.0
2	1-Yb	0.5	0	12.0
3	—	0	5	4.8
4	1-Yb	0.5	5	97.0
5	1-Sm	0.5	5	89.0
6	1-Eu	0.5	5	83.0
7	1-Gd	0.5	5	95.0
8	1-Tb	0.5	5	99.0
8	1-Dy	0.5	5	79.0
9	1-Ho	0.5	5	81.0
10	1-Er	0.5	5	87.0

^a Reaction conditions: 4 mmol of epichlorohydrin, solvent free, CO₂ (1 atm). ^b The yields were determined by ¹H NMR spectrum of the crude reaction mixtures.

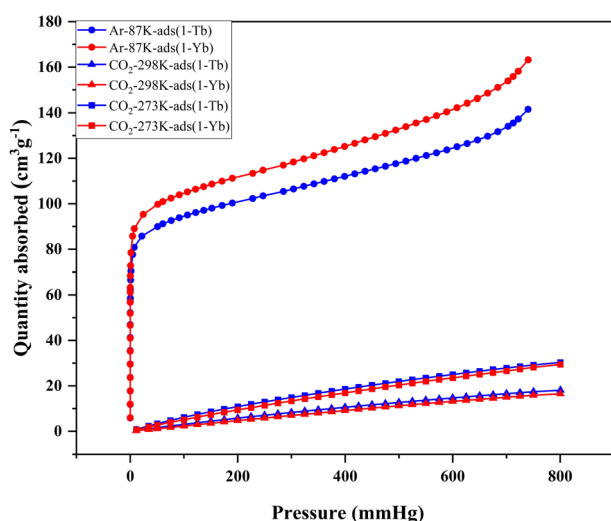


Fig. 3 The Ar adsorption isotherms at 87 K and the CO₂ adsorption isotherms at 273 K and 298 K of **1-Tb** and **1-Yb**.



cocatalyst $n\text{-Bu}_4\text{NBr}$, pure **1-Yb** catalyst shows a low conversion of 12.0% (entry 2); while without **1-Yb**, $n\text{-Bu}_4\text{NBr}$ alone lead to almost no reactivity (4.8%, entry 3). However, when **1-Yb** catalyst and cocatalyst $n\text{-Bu}_4\text{NBr}$ were added as a binary catalytic system (entry 4), the conversion of epichlorohydrin is up to 97%, highlighting the remarkable synergistic effect of the binary catalytic system.

Then using the optimized conditions, we extend this reaction to other seven isomorphous MOF candidates to search for efficient catalysts and determine the effect of the metal centers on catalytic performances. As illustrated in Table 1, the comparative catalytic results reveal that **1-Ln** solids are all valuable catalyst candidates for cycloaddition reaction of CO_2 and epoxides. Among them, **1-Gd**, Tb and Yb demonstrate the highest catalytic activity, comparable to the most effective MOF catalysts reported in literature (see Table S3†). Fig. 4 demonstrates the relationship between the catalytic yield of each Ln-MOFs and atomic number of Ln (Z_{Ln}). It is found that the catalytic yield shows two increase stages, one with a gradual increase from Sm (89%) to Tb (99%) and then an abrupt decrease at Dy (79%), and the other one with gradual increase from Dy to Yb (97%).

Generally, the strength of a Lewis acid is a critical factor in determining its catalytic activity, because a certain threshold of Lewis acidity is required to induce catalytic chemical transformations. In Fig. 4, we also plot the empirical scale for Lewis acid strength of Ln^{3+} along with Z_{Ln} based on the reported data, showing an almost linear increase of Lewis acid strength of Ln^{3+} against Z_{Ln} ; the heavier Ln^{3+} , the more Lewis acidic.³⁸ Obviously, the trend of the catalytic yield (activity) of Ln-MOFs vs. Z_{Ln} does not coincide with that of the Lewis acid strength vs. Z_{Ln} . These analyses demonstrate that the catalytic activity of Ln-MOFs, at least for **1-Tb**, is not solely dependent on Lewis acid strength of Ln^{3+} , but also on some other factors. We tend to presume that, compared to Yb^{3+} , the lighter lanthanide ions like Tb^{3+} possesses high and variable coordination number allowing

easy rearrangements for substrate activation.^{39,40} Therefore, we attribute the anomalously high activity of **1-Tb** to the combining effect of the modest Lewis acid strength and the labile coordination environment of Tb^{3+} during the catalysis. Note that the underlying mechanistic path of **1-Tb** and **1-Yb** may be different, and new systems or experiments should be designed to elucidate the exact catalytic mechanism in the future. Importantly, the present study not only confirms the available Lewis acidity of Ln^{3+} ions in MOFs, but also point out the way to develop Tb- and Yb-MOFs as high-performance Lewis acid catalysts.

Lastly, with the optimal catalysts in hand, we turn our attention to the recyclability, stability, and catalytic generality of **1-Yb** as a heterogeneous catalyst. **1-Yb** can be readily recovered, washed thoroughly with CH_2Cl_2 , and used for next runs. Within five catalytic runs, high yields of 97–99% are maintained (Fig. S7†), suggesting that **1-Yb** possessed high recyclability to the cycloaddition reaction. In addition, a PXRD pattern of **1-Yb** recovered from the fifth run is consistent with the one of the as-synthesized, confirming the structural integrity of **1-Yb** during the catalytic process (Fig. S8†). Furthermore, the catalytic generality over **1-Yb** can be extended to several typical epoxide substrates with product yields from moderate to good (Table S2†). These results show that stability and heterogeneous nature of the **1-Yb** catalyst. Based on structural analysis and catalytic performances of **1-Yb**, a tentative synergistic catalytic mechanism for the cycloaddition of CO_2 and epoxides over **1-Yb**/ $n\text{-Bu}_4\text{NBr}$ is proposed in Fig. S9†. First, dissociation of coordinated NMP of **1-Yb** occurs and the open Lewis Yb active site coordinates with oxygen atom of the epoxide. This coordination bond polarizes the C–O bond to activate the epoxide. The activated epoxide is opened by Br^- ion of TBAB to generate a metal-ordinated bromoalkoxide intermediate. Next, CO_2 was nucleophilically attacked by the O atom of bromoalkoxide to form a metal carbonate, which is finally converted to cyclic carbonate by intramolecular ($\text{S}_{\text{N}}2$ -type nucleophilic) ring-closure reaction, regenerating the catalyst and co-catalyst.¹⁰

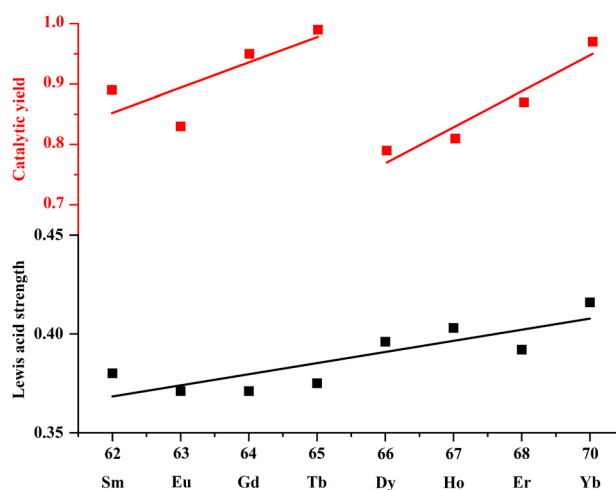


Fig. 4 The trend chart showing catalytic yield (activity) of Ln-MOFs vs. Z_{Ln} in catalyzing CO_2 -cycloaddition reaction and the Lewis acid strength of Ln^{3+} vs. Z_{Ln} .

Catalytic deacetalization–Knoevenagel reactions

Owing to the incorporation of both lanthanide Lewis acidic sites and triphenylamine Lewis basic sites within their porous frameworks, **1-Ln** could be viewed as a kind of potential bifunctional acid–base catalysts. In order to prove this point, **1-Ln** crystals were used to study their catalytic activities on one-pot cascade deacetalization–Knoevenagel reactions. The reaction between benzaldehyde dimethylacetal (BD) and malononitrile (MA) under solvent-free condition was taken as the template reaction to investigate the catalytic activities of these Ln-MOF solids (Table 2). At the first step of the catalysis, Ln ions as a Lewis acid can catalyze the deprotection of BD to give benzaldehyde (BA), and at the second step, triphenylamine as a weak Lewis base can catalyze Knoevenagel condensation reaction between BA and MA to afford benzylidene malononitrile (BM).



Table 2 One-pot cascade deacetalization–Knoevenagel reaction between benzaldehyde dimethylacetal (BD) and malononitrile (MA) catalyzed by different 1-Ln catalysts^a

Entry	Catalyst	Conversion of BD ^b (%)	Conversion of BA ^c (%)	Yield of BM ^b (%)
1 ^d	1-Yb	0	0	0
2 ^e	—	0	0	0
3	1-Sm	83	81	68
4	1-Eu	93	83	78
5	1-Gd	96	96	93
6	1-Tb	98	100	98
7	1-Dy	79	98	77
8	1-Ho	88	96	84
9	1-Er	86	97	86
10	1-Yb	97	100	97

^a Reaction conditions: BD (1 mmol), H₂O (3 mmol), MA (2 mmol), catalysts (0.5 mol%), 60 °C, 6 h, N₂ atmosphere. ^b The yields were determined by ¹H NMR spectrum of the crude reaction mixtures. ^c The conversion of intermediate BA is defined by the ratio of (yield of BM)/(conversion of BD). ^d Control experiment was performed without water in the presence of **1-Yb**. ^e Control experiment was done without **1-Yb** in the presence of water.

The catalytic reactions were performed using the following conditions: BD (1 mmol), H₂O (3 mmol), MA (2 mmol), catalysts (0.5 mol%), 60 °C, 6 h, N₂ atmosphere. Experimental results are given in Table 2, showing that **1-Ln** solids are all valuable catalyst candidates for one-pot cascade reaction. Among them, **1-Tb** and **1-Yb** demonstrate the highest catalytic activity. Control experiments were also performed to figure out the role of components during the catalytic procedure. H₂O was necessary in the deacetalization (step I) process, as the reaction was sluggish without water even in the presence of **1-Yb** (entry 1). As shown in entry 2, the reaction did not proceed without **1-Yb**.

In order to clarify the active catalytic site for each step, we calculate and plot the conversion of BD and intermediate BA, and the yield of BM against Z_{Ln} (the coordination number of Ln) in Table 2 and Fig. 5. The conversion of BD to BA in first step (Fig. 5a) clearly show that all Ln³⁺ cations in MOFs have the Lewis acidity enough to efficiently catalyze the deprotection of BD, with two maxima at **1-Tb** (98%), **1-Yb** (97%) and two minima at **1-Sm** (83%) and **1-Dy** (79%). At the first step of the one-pot catalysis, the trend of the catalytic activity in Ln-MOFs is not regularly increased along with Z_{Ln} but very similar to that found in catalytic cycloaddition of CO₂ and epoxides, demonstrating Ln³⁺ cations embedded in MOF as Lewis acids is distinct from homogeneous species, possible due to their unique coordination environment in MOF solid. In second step, the conversion of intermediate BA to the product BM, which is defined by the ratio of (yield of BM)/(conversion of BD), is increased from 81% for Sm to almost 100% for Gd, Tb, Dy, Ho, Er, Yb (Fig. 5b). Generally, Knoevenagel condensation reactions are considered to be base-catalyzed.^{27–29} However, for comparison, control experiments show that the conversion of BA to BM catalyzed by sole triphenylamine or H₃NTB is only 70 or 46%, respectively. The present study confirms that Ln³⁺ especially for Tb³⁺, Yb³⁺ in MOFs plays an important role in accelerating the Knoevenagel

reaction. This synergistic effect of the Lewis acidic metal centers and Lewis base sites in porous MOFs on catalyzing Knoevenagel reaction is also observed in other scientific studies.^{6,29,31,41}

The above analyses reveal that both steps are strongly correlating with the catalytic activity of lanthanide ions embedded in the framework. As a consequence, the total yields of BM show **2-Tb** and **2-Yb** have the highest catalytic activity towards one-pot reaction. Interestingly, similar to that found in Fig. 4, the trend of catalytic activity of each Ln-MOFs vs. Z_{Ln}

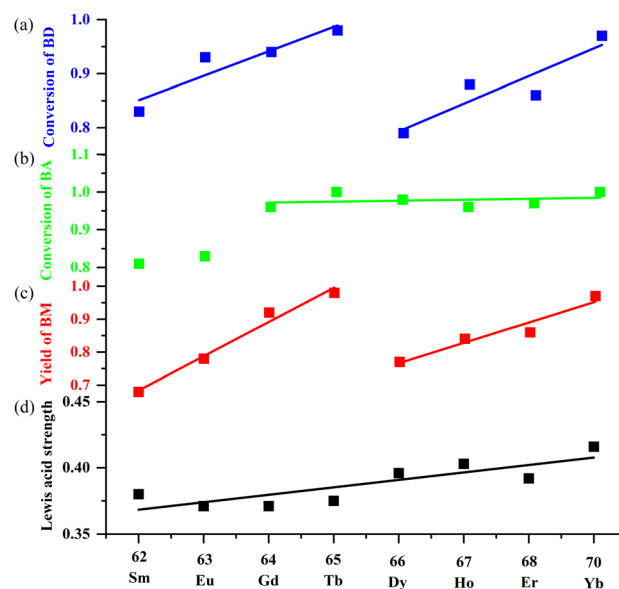


Fig. 5 The trend chart showing (a) the conversion of BD, (b) the conversion of intermediate BA, and (c) the yield of BM vs. Z_{Ln} (the coordination number of Ln) in catalyzing one-pot cascade reaction and (d) the Lewis acid strength of Ln³⁺ vs. Z_{Ln} .



(Fig. 5c) also shows two increase stages, one with a gradual increase from Sm (68%) to Tb (98%) and then an abrupt decrease at Dy (77%), and the other one with gradual increase from Dy (77%) to Yb (97%). Once again, the present study confirm the feasibility to develop Tb- and Yb-MOFs as high-performance Lewis acid catalysts, comparable to the most effective MOF catalysts reported in literature (see Table S5†).

Finally, we investigate the recyclability, stability, and catalytic generality of **1-Yb**. After five consecutive catalytic runs under optimized conditions, **1-Yb** kept high catalytic activity (Fig. S10†). Moreover, the PXRD patterns between the recovered and the original one are almost identical, confirming the structural integrity of **1-Yb** (Fig. S11†). The recovered **1-Yb** was also evaluated by Ar adsorption, showing the retention of the porous framework (Fig. S12†). In addition, **1-Yb** had the general high catalytic activity on a series of epoxides with various substitutes under the optimal reaction conditions (Table S4†). All results show the stability and heterogeneous nature of the **1-Yb** catalyst. Based on the structure and catalytic behaviors of **1-Yb**, a possible catalytic mechanism for one-pot cascade deacetalization–Knoevenagel reaction on **1-Yb** is proposed in Fig. S13.†

Conclusions

In summary, we have synthesized a family of robust and microporous Ln-MOFs (**1-Ln**, Ln = Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb) based on H₃NTB, demonstrating permanent porosity for Ar and CO₂ adsorption abilities. Importantly, the combination of Lewis acidic lanthanide ions and basic triphenylamine group into the framework of **1-Ln** makes them efficient acid–base catalysts for both cycloaddition of CO₂ with epoxides and one-pot cascade deacetalization–Knoevenagel reactions. Among them, **1-Tb** and **1-Yb** demonstrates the best catalytic activities for both two reactions. More strikingly, the systematic catalytic studies reveal that the catalytic activities of these Ln-MOFs are strongly dependent on Lewis acidic lanthanide ions embedded in the framework, opening up the possibilities to develop Tb- and Yb-MOFs as high-performance Lewis acid catalysts. Work is underway to synthesize new Ln-MOFs in our lab.

Experimental

Materials and instruments

Materials and instruments can be seen in ESI S1† part.

Synthesis

Synthesis of {[Sm(NTB)(NMP)]·(NMP)}_n (1-Sm**).** A mixture of Sm(NO₃)₃·6H₂O (16.0 mg), H₃NTB (23.3 mg), NMP (1.0 mL), HCl (0.01 M, 0.2 mL), H₂O (0.3 mL) was sealed in a 23.0 mL Teflon-lined stainless-steel container, which was heated at 85 °C for 21 hours to afford block crystals. The collected crystals were dried in air at room temperature. Yield: 11.24 mg. IR/cm^{−1} (KBr): 3424 (br), 3045 (w), 2925 (w), 2873 (w), 1687 (m), 1627 (m), 1584 (m), 1534 (m), 1393 (m), 1332 (m), 1178 (m), 1144 (w), 1117

(w), 1015 (w), 925 (w), 831 (m), 783 (s), 719 (m), 659 (w), 628 (w), 580 (m), 529 (w), 506 (m), 470 (w), 441 (w).

Synthesis of {[Eu(NTB)(NMP)]·(NMP)}_n (1-Eu**).** **1-Eu** was synthesized by using the same procedure except by using Eu(NO₃)₃·6H₂O (16.6 mg). Yield: 9.51 mg. IR/cm^{−1} (KBr): 3430 (br), 3044 (w), 2926 (w), 2873 (w), 1687 (m), 1628 (m), 1585 (m), 1535 (m), 1392 (s), 1332 (m), 1262 (w), 1177 (m), 1144 (w), 1015 (w), 831 (m), 784 (s), 719 (m), 659 (w), 628 (w), 559 (m), 529 (w), 506 (m), 470 (w), 441 (w).

Synthesis of {[Gd(NTB)(NMP)]·(NMP)}_n (1-Gd**).** **1-Gd** was synthesized by using the same procedure except by using Gd(NO₃)₃·6H₂O (14.5 mg). Yield: 11.49 mg. IR/cm^{−1} (KBr): 3393 (br), 2926 (w), 1686 (w), 1655 (w), 1625 (w), 1585 (w), 1535 (w), 1401 (s), 1315 (w), 1275 (w), 1170 (m), 1106 (w), 1014 (w), 924 (m), 582 (m), 784 (s), 710 (m), 659 (w), 628 (w), 565 (m), 525 (w), 505 (m), 470 (w), 442 (w).

Synthesis of {[Tb(NTB)(NMP)]·(NMP)}_n (1-Tb**).** **1-Tb** was synthesized by using the same procedure except by using Tb(NO₃)₃·6H₂O (27.2 mg). Yield: 25.65 mg. IR/cm^{−1} (KBr): 3319 (br), 1654 (m), 1596 (m), 1532 (m), 1401 (s), 1316 (w), 1281 (w), 1176 (m), 1150 (w), 1105 (w), 1013 (m), 954 (m), 859 (m), 783 (s), 710 (m), 672 (w), 628 (w), 568 (m), 521 (w), 496 (w), 438 (w).

Synthesis of {[Dy(NTB)(NMP)]·(NMP)}_n (1-Dy**).** **1-Dy** was synthesized by using the same procedure except by using Dy(NO₃)₃·6H₂O (18.3 mg). Yield: 19.97 mg. IR/cm^{−1} (KBr): 3434 (br), 3044 (w), 2916 (w), 2873 (w), 1687 (m), 1630 (m), 1585 (m), 1537 (w), 1393 (s), 1331 (w), 1307 (w), 1260 (w), 1178 (w), 1144 (w), 1116 (w), 1015 (w), 865 (w), 831 (w), 783 (s), 719 (w), 659 (w), 628 (w), 584 (w), 561 (w), 530 (w), 506 (w), 470 (w), 442 (w).

Synthesis of {[Ho(NTB)(NMP)]·(NMP)}_n (1-Ho**).** **1-Ho** was synthesized by using the same procedure except by using Ho(NO₃)₃·6H₂O (17.6 mg). Yield: 20.51 mg. IR/cm^{−1} (KBr): 3428 (br), 3044 (w), 2925 (w), 2874 (w), 1687 (m), 1632 (m), 1586 (m), 1539 (m), 1514 (w), 1401 (s), 1331 (m), 1261 (w), 1178 (w), 1145 (w), 1117 (w), 1105 (w), 866 (w), 832 (m), 784 (s), 719 (m), 659 (m), 628 (w), 585 (m), 563 (w), 531 (w), 507 (w), 470 (w), 443 (w).

Synthesis of {[Er(NTB)(NMP)]·(NMP)}_n (1-Er**).** **1-Er** was synthesized by using the same procedure except by using Er(NO₃)₃·6H₂O (30.0 mg). Yield: 21.32 mg. IR/cm^{−1} (KBr): 3424 (br), 3044 (w), 2919 (w), 2873 (w), 1687 (m), 1632 (m), 1586 (m), 1539 (m), 1512 (w), 1401 (s), 1307 (m), 1261 (w), 1178 (w), 1145 (w), 1117 (w), 1015 (w), 866 (w), 832 (m), 784 (s), 719 (m), 659 (m), 627 (w), 585 (m), 563 (w), 531 (w), 507 (w), 470 (w), 442 (w).

Synthesis of {[Yb(NTB)(NMP)]·(NMP)}_n (1-Yb**).** **1-Yb** was synthesized by using the same procedure except by using YbCl₃·6H₂O (15.5 mg). Yield: 9.62 mg. IR/cm^{−1} (KBr): 3415 (br), 3044 (w), 2921 (w), 2874 (w), 1687 (m), 1633 (m), 1588 (m), 1540 (m), 1512 (w), 1401 (s), 1331 (m), 1261 (w), 1178 (w), 1145 (w), 1117 (w), 1105 (w), 925 (w), 866 (w), 833 (w), 784 (s), 719 (m), 659 (m), 628 (w), 588 (m), 565 (w), 531 (w), 509 (w), 471 (w), 443 (w).

Single-crystal structure characterization of the 1-Ln crystals

SCXRD data for **1-Tb** and **1-Yb** were performed on a Rigaku Oxford X-ray Diffractometer with graphite-monochromated MoK α radiation (λ = 0.71073 Å) at 293 K. All absorption corrections were applied using multi-scan technique. The



crystal structures of **1-Ln** were solved by the direct method and refined through full-matrix least-squares techniques method on F^2 using the Olex2 crystallographic software package.^{42,43} Full crystallographic data for **1-Tb** and **1-Yb** has been deposited with the CCDC 2210635 and 2210636. The crystallographic data for **1-Tb** and **1-Yb** are summarized in Table S1.†

Ar and CO₂ adsorption isotherms

Ar adsorption isotherm was recorded at 87 K on a micromeritics ASAP2460 analyzer. The sample was activated under vacuum by heating at 200 °C overnight. The specific surface area was calculated from the data in the adsorption branch at $p/p_0 = 0.05$ – 0.30 . The total pore volume was calculated from the uptake at p/p_0 of 0.950. CO₂ adsorption isotherms was performed at 273 and 298 K respectively.

General procedure for cycloaddition of CO₂ and epoxides

In a typical catalytic reaction, epichlorohydrin (4 mmol) and cocatalyst *n*-Bu₄NBr (0.2 mol; 5 mol%) were added to the Schlenk tube, to which was further added **1-Ln** (Ln = Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb) (2 mol% based on metal ion), and then slowly stirred under 1 atm CO₂ pressure for 36 hours at room temperature.

General procedure for one-pot cascade deacetalization–Knoevenagel reactions

In a typical catalytic reaction, benzaldehyde dimethylacetal (1 mmol), H₂O (3 mmol) and malononitrile (2 mmol) were added to the Schlenk tube, to which was further added **1-Ln** (Ln = Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb) (2 mol% based on Ln³⁺), and then slowly stirred for 6 hours at 60 °C in a nitrogen atmosphere.

Author contributions

Qingxia Yao, Yi Qiu, Jun Li: project administration, supervision. Xuezhen Si, Qingxia Yao: writing original draft, methodology, data collection and analysis. Xuze Pan, Jintang Xue, Minglei Cao, Wenzeng Duan, Xianqiang Huang: data collection and analysis. Yi Qiu, Jie Su: X-ray single-crystal structural analysis.

Conflicts of interest

There are no conflicts to declare.

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