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Tuning of the flexibility in metal–organic frameworks based on pendant arm macrocycles†

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An isostructural series of flexible metal–organic frameworks based on macrocycles having diverse pendant arms was developed to tune flexibility depending on functional groups. The pendant arms directing into the pores were found to play a key role in imparting different gate-opening behaviours in the threshold pressure and sorption capacity upon interaction with guest molecules.

The design and syntheses of metal–organic frameworks (MOFs) begin with the selection of the metal nodes and bridging organic ligands.¹ Types of metal and organic building blocks and the nature of their assembly determine the geometry and structure of the MOFs. Often, they allow a high degree of flexibility, resulting in so-called 'flexible MOFs' or 'soft porous crystals'.² Flexible MOFs can reversibly change their form upon the action of external stimuli, owing to the intermolecular degrees of freedom in their components and thus, have tremendous potentials in applications such as adsorption, separation, sensing, catalysis, and biomedical usage.³ Accordingly, researchers studying MOFs have been devoting considerable efforts to establish the fundamental principles that facilitate the syntheses of flexible MOFs. For instance, the O–O axis of a carboxylate group endows significant breathing effect in MIL-53 and MIL-88 by acting as a kneecap.⁴ Farha *et al.* recently reported a flexible MOF, NU-1400, based on four connected Zr_6 nodes, which afforded degrees of freedom by reducing the connectivity, compared with rigid NU-1000 and NU-901 having eight connected Zr_6 nodes.⁵

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However, strategies for the incorporation of selective switching in flexible MOFs, as demanded by the applications based on the flexibility, are still elusive and serendipitous. Hence, designing and controlling the flexibility in these materials are essential and yet, a challenging issue. Several strategies have been demonstrated in order to tune the structural dynamics and responsiveness of the MOFs such as crystal size adjustment, metal substitution, and linker functionalization with various functional groups or flexible side chains.⁶ **COMMUNICATION**
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Macrocycle-based MOFs have been reported as a subclass of MOFs; in particular, aza-macrocyclic complexes with square planar geometry can act as a linear linker, providing axial sites for carboxylate ligands.⁷ Furthermore, various types of functional groups can be easily introduced into the pores during the synthesis, by using macrocyclic moieties such as the 1,3,6,8,10,13 hexaazacyclotetradecane $Ni(n)$ complex with pendant arms at 1 and 8 positions.8 Previously, we reported a flexible MOF based on the $Ni(II)$ macrocycle having two propyl arms that behaved as a molecular gate.⁹ Experimental and theoretical studies found that $CO₂$ molecules mainly interacted with the propyl pendant arms to induce the gate-opening and breathing phenomena. Based on this, we postulated that such a MOF can be an excellent platform for easily securing different functional groups in the pendant arm macrocycles, which would render them different structural dynamics. Thus, we introduced nitrile, hydroxyl, and allyl pendant arms in $Ni(II)$ macrocycles in this study and successfully synthesised three isostructural flexible MOFs, flex- $MOF(CN)$, flexMOF(OH), and flexMOF(CH₂) (Fig. 1). Upon types of the guest molecules such as $CO₂$ and water, these MOFs showed distinguishable flexible behaviours, which were examined by sorption isotherms, in situ X-ray powder diffraction (XRPD), and single-crystal diffraction (SCD).

To synthesise an isostructural series of flexible MOFs decorated with different functional groups, we induced the self-assembly of a tetradentate ligand, 2,2',5,5'-biphenyltetracarboxylic acid (H_4BPTC) , with $[NiL_R]^{2+}$ as a metal building block (R = CN, OH, and CH2 for nitrile, hydroxyl, and allyl substitutes, respectively, Fig. 1a), yielding $\{$ [(NiL_{CN)2}(BPTC)]·4DMF·2H₂O} (*as-flex***MOF(CN**);

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Fig. 1 (a) Square planar Ni(II) macrocycles with different pendant arms as metal building blocks and H4BPTC as an organic building block. (b) Representation of the linear connectivity between two building units via coordinate and hydrogen bonds. Colour scheme: Ni, yellow; C, grey; via coordinate and hydrogen bonds. Colour scheme: Ni, yellow; C, grey; \overline{F} Fig. 2 Comparison between the as-synthesised (as) and closed-pore (cp)
C, red; N, blue; H, white. \overline{F} which is a street of (a) flamMOF(C

 $\text{DMF} = N_{\rm s}N$ -dimethylformamide), {[$\text{(NiL}_{\text{OH}})_{\rm 2}\text{(BPTC)}$] 1DMF 3H₂O} (as-flexMOF(OH)), and $\{$ [(NiL_{CH2})₂(BPTC)]·1CH₃CN·5H₂O} (as-flex- $MOF(CH₂)$). For all the compounds, the square planar Ni(II) macrocycles were coordinated by two $BPTC^{4-}$ ligands in the axial sites, resulting in a six-coordinated octahedral geometry with hydrogen bonds between the uncoordinated oxygen atoms of the carboxylate and secondary amines from the aza-macrocyclic complex (dotted lines in Fig. 1b). Since the BPTC 4^- moiety in the MOFs is not planar (dihedral angles between two phenyl rings are 52.780° in $as\text{-}flexMOF(CN)$, 50.318° in as-flexMOF(OH), and 57.267 \textdegree in *as-flex*MOF(CH₂)) and coordinates four macrocycles, 3D networks having interconnected pores are formed (top of Fig. S4–S6, ESI[†]), in which the void space is 52.5% for *as-flex*MOF(CN), 49.6% for $as\text{-}flexMOF(OH)$, and 41.2% for $as\text{-}flexMOF(CH_2)$ per unit cell volume, as obtained by PLATON calculations.¹⁰ Upon drying the MOFs, the 3D structures drastically changed and exhibited structural flexibility, as evidenced by the XRPD patterns (Fig. S7–S9, and details of activation are provided in ESI;† hereafter, dried compounds are designated as cp -flexMOF (cp = closed-pore)). The void volume is also greatly reduced to 6.1%, 6.6%, and 3.9% per unit cell volume for cp-flexMOF(CN), cp-flexMOF(OH), and $\mathit{cp\text{-}flexMOF}$ (CH₂), respectively. Despite such tremendous changes in the cell volume, all the samples maintained single crystallinity, which was sufficient to obtain SCD data. Fig. 2 shows that the rectangular dimension of as phases in all the three compounds considerably shrank in cp phases (42.6%, 34.1%, and 23.8% decreases for flexMOF(CN), flexMOF(OH), and $flexMOF(CH_2)$, respectively). During the dynamic movement, the dihedral angle between the two phenyl rings and those between the carboxylate planes and benzene rings of the ligand changed, and the pendant arms of the macrocycles blocked the pores (Fig. S10–S12, ESI†).

phases of (a) *flexMOF(CN)*, (b) *flexMOF(OH)*, and (c) *flexMOF(CH₂)* in rectangular dimension. The macrocycles and hydrogen atoms are omitted for clarity. Colour scheme: Ni, yellow; C, grey; O, red.

In order to explore the guest-dependent flexibility of these MOFs, N_2 and CO_2 gas sorption isotherms were obtained at 77 and 196 K, respectively. All cp phases showed typical type II N_2 sorption isotherms, indicating a nonporous nature (Fig. S13, $ESI⁺$). However, the characteristic of the $CO₂$ sorption isotherms was interesting (Fig. 3). cp-flexMOF(CN) exhibited gate-opening for $CO₂$ with an abrupt uptake at a threshold pressure of 0.42 bar, and its total adsorption capacity approached 40.1 wt% (204.8 cc g^{-1} or 9.14 mmol g^{-1} of the host) at 1 bar. When the allyl pendant arms are introduced, the resulting MOF, $cp\text{-}flexMOF(CH_2)$ displayed a significantly different threshold pressure of \sim 0.70 bar, which was higher than that in $cp\text{-}flexMOF(CN)$. This was also accompanied by a decrease in the $CO₂$ uptake capacity (16.7 wt%, 85.5 cc g^{-1} , or 3.82 mmol g^{-1}). The different response of

Fig. 3 $CO₂$ adsorption/desorption isotherms of flexMOF(CN), flexMO-F(OH), and $flexMOF(CH_2)$ at 196 K (left) and schematic of their flexible behaviours upon $CO₂$ adsorption (right). Filled and empty circles represent adsorption and desorption curves, respectively.

cp-flexMOF(CN) and cp-flexMOF(CH₂) toward N₂ and CO₂ can be explained by the high quadruple moment and polarizability of CO₂, which is in contrast to the inert nature of N_2 .¹¹ It is also noteworthy that as phases of $flexMOF(CN)$ and $flexMOF(CH_2)$ were estimated to potentially adsorb *ca*. 13.9 and 5.4 mmol $g^$ of $CO₂$, respectively (greater than the experimental value; Table S11, ESI†), as calculated by grand canonical Monte Carlo (GCMC) adsorption simulation performed with the molecular software package, RASPA 2.0 (details of simulation in ESI[†]).¹² This implies that those cp phases are not fully transformed into as phases under this condition (1 bar of $CO₂$ at 196 K), but are open up to form new phases, lp -flexMOF (Fig. 3; lp stands for large pore phase). Also, both the $flexMOF(CN)$ and $flex$ $MOF(CH₂)$ desorbed in a hysteretic and stepwise manner, which adsorption and desorption curves do not merge at the end even under vacuum $(<10^{-5}$ mbar) at 196 K. In contrast, hydroxyl group-containing cp-flexMOF(OH) showed no stepwise adsorption profile with negligible uptake.

The forementioned different responsive behaviours were studied by in situ XRPD under the synchronized condition of $CO₂$ sorption. The XRPD patterns of cp-flexMOF(CN) did not change up to 0.4 bar, and a drastic structural change was observed at 0.6 bar, at which the $CO₂$ uptake increased sharply (Fig. S14, ESI†), retaining up to 1 bar. As mentioned previously, this is not a same structure with a as phase, but a lp phase. During desorption, the XRPD patterns were retained up to 0.4 bar, but it changed into a new structure below this pressure, which had neither cp nor lp structure and thereby denoted as np - $flexMOF(CN)$ (Fig. 4a; np stands for narrow pore phase). Such a discovery of a new intermediate phase derived upon desorption has been very rarely reported in flexible MOFs.¹³

The intermediate structure, $np\text{-}flexMOF(CN)$ has its own permanent porosity. After one adsorption–desorption experiment, the *np* phase is stabilised even under vacuum at 196 K. Interestingly, the successive measurement of the $CO₂$ uptake beginning from np-flexMOF(CN) at 196 K (Fig. 4b) showed type IV adsorption isotherm, which was clearly different from the adsorption isotherm of the first cycle. In this profile, even at the low pressure region $CO₂$ uptake was occurred up to 12.0 wt%, yet the threshold pressure was observed at the same position of the first cycle. The total $CO₂$ uptake decreased from 40.1 to 33.7 wt%, in which the uptake difference is equal to the trapped amount of $CO₂$ after the first cycle, and the desorption curve retraced to the adsorption curve. As revealed by the *in situ* XRPD (Fig. 4c), the np transformed to the lp phase during the $CO₂$ re-adsorption, and then reverted to its np phase after re-desorption. The further consecutive cycle of $CO₂$ adsorption/desorption for $np\text{-}flexMOF(CN)$ fully reproduced this trend (Fig. S15, ESI†), indicating the reversible transition between the np and lp phases (Fig. 4a). Furthermore, heating at 110 °C under vacuum resulted in the transformation into the cp phase (Fig. 4c and Fig. S16, ESI†). Remarkably, at higher temperature (298 K), the np phase can be still identified upon $CO₂$ desorption (Fig. S19, ESI†). The gate-opening pressure of flexMOF(CN) dramatically shifts to a higher pressure upon the temperature increase to room temperature, while $flexMOF(CH_2)$ still opens the pores at a very low pressure.

Fig. 4 (a) Schematic illustration of the phase transitions of **flexMOF(CN)** upon guest removal/reintroduction and CO₂ adsorption/desorption. (b) Consecutive cycles of $CO₂$ adsorption/desorption isotherms in flexMOF(CN) at 196 K, and (c) corresponding XRPD data measured before $CO₂$ adsorption, after adsorption, desorption (vacuum $< 10^{-5}$ mbar), re-adsorption, and redesorption, and after re-activation by heating under vacuum. Filled and empty circles represent adsorption and desorption curves, respectively.

It should be noted that the discrepancy of the flexibility in flexMOF(OH) is attributed to the increased number of hydrogen bonds from *as-flexMOF(OH)* to *cp-flexMOF(OH)*, as revealed by the SCD analysis, owing to the single-crystal-tosingle-crystal transformation (Fig. 5b and Fig. S20 in ESI†). In cp-flexMOF(OH), the hydroxyl groups participate in hydrogen bonding with the carboxylate oxygens and the secondary amines from the macrocycles as well as the hydroxyl groups from other pendant arms to stabilize the cp structure by strong host-host interactions. Considering this aspect, while $CO₂$ molecules failed to open up the structure (Fig. S18, ESI†), certain guest molecules, which can make stronger host–guest interaction such as water, is expected to lead a structural flexibility, and water vapour sorption isotherm (Fig. 5c) shows that the introduction of polar water molecules can open the pores of *cp-flex*MOF(OH); it adsorbs 144 cc g^{-1} of water vapour at 298 K in two steps, at 0.08 and 0.65 relative humidity $(P/P₀)$, and desorbs with a large hysteresis. The corresponding XRPD results, measured after water adsorption and re-activation by heating at 110 \degree C under vacuum, also verifies the significant and reversible structural transitions of *flexMOF(OH)* (Fig. S21, ESI[†]). Likely, *flexMOF*(CN) and *flexMOF*(CH₂) also shows the stepwise adsorption profile for water vapour, which was similarly observed in the $CO₂$ sorption (Fig. S22, ESI[†]). Furthermore, methanol and benzene vapour isotherms of *flexMOF(OH)* were measured to prove the role of polar protic solvents in controlling

Fig. 5 (a) Schematic illustration of the phase transitions of *flexMOF(OH)* upon guest removal/reintroduction and adsorption/desorption of $CO₂$ and $H₂O$ molecules. (b) Hydrogen bonds of cp -flexMOF(CN) as represented by dotted lines, and (c) its water vapour isotherm at 298 K. Filled and empty circles represent adsorption and desorption curves, respectively.

the host–guest interaction and opening the closed pores. Resultingly, the pores of **flexMOF(OH)** was opened by methanol, but there was no adsorption by the non-polar benzene vapour (Fig. S25, ESI†).

In conclusion, we employed the $Ni(_{II})$ macrocycle moiety with different pendant arms containing the nitrile, hydroxyl, or allyl groups, to demonstrate that such a MOF platform can readily grant and tune the flexibility based on the functional groups in the pores. Three isostructural MOFs exhibited distinct flexibility, including the gate-opening threshold pressure, sorption capacities, step steepness, and hysteresis. For flexMOF(CN) and $flexMOF(CH₂)$, an intermediate new form was discovered from the CO_2 -trapped np phase. Moreover, the hydroxyl group in flexMOF(OH) induced the strong host-host interaction via numerous hydrogen bonds in the cp phase, which could be attributed to the structural stabilisation, and *flexMOF(OH)* showed the responsive flexibility upon water over $CO₂$. Therefore, we envision that this pendant arm azamacrocycle-based MOF would not only provide insights for the rational design and finetuning of flexible MOFs but also enable their uses in a variety of practical applications.

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Conflicts of interest

There are no conflicts to declare.

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