Reticular control of interpenetration in a complex metal–organic framework†

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Metal–organic frameworks (MOFs) are a new generation of crystalline porous materials covering a broad spectrum of research fields due to their high porosity and the feasibility of modification of their structure. Due to their large void spaces, MOF structures tend to be more stable when the crystals are grown to form interpenetrated frameworks. Control of interpenetrated frameworks in MOFs is not only useful for various kinds of practical application but also provides insight towards understanding the reaction process and cationed behavior in MOF chemistry. In this research, we sought to investigate and successfully achieve reticular control of the interpenetration of PCN-280, a complex MOF with a 2-fold interpenetrated framework containing heterogeneity building blocks (two kinds of polytopic organic linking unit based on carboxylate), to form a new isoreticular single-frame MOF, termed MOF-908 and isoreticular MOF-909. The crystal structure and underlying network of MOF-908 and MOF-909 are proven by X-ray diffraction analysis as well as other related characterizations. Indeed, the mechanism of reticular control of the interpenetration of PCN-280 is studied in detail using density functional theory (DFT) calculations.

Introduction

Reticular chemistry at its best opens the window toward the blue sky of metal–organic framework (MOF) chemistry, a highly-crystalline ordered material possessing permanent porosity and designable properties which can be potentially applied for gas/vapor uptake, catalysis, energy conversion and chemical sensing. Interpenetration behavior mainly comes from the huge void fraction of the large unit cells of MOFs, in which another framework can grow to occupy the vacancy resulting in the stabilization of the resulting structures. Given the fact that control of the reaction in which interpenetration occurs on increasing the lengths of organic linkers to obtain a single host framework is one of the most interesting studies for paving the way to understanding the mechanism of MOF growth, it has experienced a lack of full exploration in MOF research. It is noted that interpenetration reportedly occurs in MOFs generated through the reaction of single carboxylate-based linkers, which, in some cases, could be a mixture of carboxylate and bipyridine linking units or metal–oxo clusters. In order to entirely control the structures of interpenetrated frameworks, (i) synthetic conditions were varied, for example bulky molecules can be introduced into the reaction, which unpredictably produces a host single frame or another different structure; (ii) organic templates were utilized and somehow the interpenetration control is not clear; (iii) the organic linker was functionalized to modify the geometrical direction of the point of extension and the carboxylate moiety; (iv) the coordinated/uncoordinated solvent removal was additionally carried out after the structure formation. Here we report a strategy that utilizes the power of reticular chemistry in the control of the structure of a tertiary MOF. We define this methodology as “reticular control” and it involves independent 2-fold interpenetrated frameworks formed using triangular 4,4′,4″-benzene-1,3,5-triyl-tris(benzoic acid) (H3BTB) and linear biphenyldicarboxylic acid [H2BPDC] linkers and a trigonal prism Fe₂O₆(CO₂)₁₂ cluster, using a heterogeneity MOF reported by Zhou and co-workers termed PCN-280. From two complex MOFs, namely MOF-908 and MOF-909 (Scheme 1), the isoreticular single net of PCN-280 is successfully produced by reticular control of the rational input linkers using a mixture of H2BPDC and 3,3′-azobDC (azobenzene-3,3′-dicarboxylic acid). This is the first time the structure of a tertiary interpenetrated MOF has been controlled to achieve a non-interpenetrated framework. Indeed, the “deinterpenetration”, the process of removing the interpenetrated frameworks to obtain a single-frame structure, was not successful either using the reported methods, varying the synthetic conditions, using a template, using bulky organic solvents or in the absence

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d Electronic supplementary information (ESI) available: Full synthesis and characterization of MOF-908, MOF-909, crystallographic data, and additional gas uptake measurement details. See DOI: 10.1039/c8qm00368h
of 3,3'-azoBDC. The role of 3,3'-azoBDC in the reticular control is subsequently demonstrated by adopting density functional theory (DFT) calculations to explore the process of the reaction. Both 3,3'-azoBDC\(^{2-}\) and BPDC\(^{2-}\) are competitive in the first reaction stage, but BPDC\(^{2-}\) is more favorable for the substitution of 3,3'-azoBDC\(^{2-}\) in a subsequent process, which leads to the formation of MOF-908. In terms of mechanical stability, we also verify that MOF-908 is much more stable than MOF-909. Not only does our finding provide new insight into interpenetration control in MOFs by reticular synthesis, but it also verifies that MOF-908 is much more stable than MOF-909.

Results and discussion

To verify the role of the geometrical requirement of both organic linkers in a heterogeneity MOF,\(^{18}\) we previously reported the concept of “perfect fit” of the ratio lengths of the tritopic \(\text{H}_3\text{BTB}\) linker over various ditopic linkers and the number of links surrounding the triangular Fe-based cluster, which is more convenient for non-interpenetrated structure formation.\(^{19}\) On increasing the length of the linear ditopic linker in the mixed linker system complex MOFs have shown a tendency to become interpenetrated structures; in particular, a series of MIL-142 (MIL-142A to \(19\)) and PCN-280 (a complex MOF structure comprising two kinds of link, \(\text{BTB}^{2-}\) and \(\text{BPDC}^{2-}\), and trigonal prism \(\text{Fe}_2\text{O}(-\text{CO}_2)_{6}\) cluster) (Scheme 1) have shown this.\(^{17}\) However, by replacing \(\text{BPDC}^{2-}\) in the PCN-280 structure by azobenzene-4,4’-dicarboxylate (4,4’-azoBD\(\text{C}\)\(^{2-}\)), an isoreticular structure with a single net was formed that is termed PCN-285.\(^{17}\) This observation is still ambiguous and has not received any explanation from either experimental or calculation methodology.

Before controlling the interpenetration behavior of PCN-280, we start by synthesizing the isoreticular structure of PCN-280 and PCN-285 in which the 3,3’-azoBDC linker, whose length is longer than \(\text{H}_2\text{BPDC}\) and shorter than 4,4’-azoBD\(\text{C}\), is used and a single frame MOF is expected to be obtained. MOF-909 microcrystalline powder was isoreticularly produced under synthetic conditions different to those used for PCN-280 and PCN-285 preparation, in line with our speculation. The powder X-ray diffraction (PXRD) patterns of the as-synthesized and activated samples are coincident with the pattern calculated from structural modeling (ESI,\(^{+}\) Section 3). A single crystal of PCN-280 was then prepared by the same synthetic method utilized for MOF-909 manufacturing. The single-frame structure of PCN-280, MOF-909, cannot be obtained after exhausting efforts following the reported methodologies including synthetic modification, bulky organic solvent changing and use of templates. We hypothesize that there exists an energy barrier for the reaction when the mixed linker system, \(\text{H}_3\text{BTB}\) and \(\text{H}_2\text{BPDC}\), interacts with the triangular Fe–oxo cluster, preventing the final structure from the single network of MOF-909- or PCN-285-type topology which is unseen in MOF chemistry and was not acknowledged in the original publication.\(^{17}\) In order to prove that postulation, we then move forward to rational investigation of the reaction in which the \(\text{H}_3\text{BTB}\) linker was an independent reagent and was constrained to be unchanged in molar amount while the ratio of 3,3’-azoBDC to \(\text{H}_2\text{BPDC}\) was proportionally controlled from 0:100 mol% to 100:0 mol%. As depicted in Fig. 1, after 24 h of reaction the highly crystalline products were observed for all of the samples. The PXRD patterns clearly show that PCN-280 is only formed when the molar percentage of the 3,3’-azoBDC linker in the solution is in the range from 0% to 40%. Interestingly, increasing the molar ratio of 3,3’-azoBDC to \(\text{H}_2\text{BPDC}\) to 60:40 mol% leads to the achievement of the non-interpenetrated structure of PCN-280, MOF-908 (a zoomed in image of Fig. 1 at low angles is found in Section 4 of the ESI†).

The PXRD pattern of the 50:50 mol% as-synthesized sample exhibited 5 characteristic peaks coincident with the simulated pattern which belong to the Miller planes of (101), (003), (201), (211–3), and (3−11). It is noted that the (101) Bragg position of the 50:50 mol% sample obviously enhances in intensity after the solvent exchange and activation processes (ESI,\(^{+}\) Section 4), confirming the observation of a single frame structure. PCN-280 appeared to be a single crystal represented by very thin hexagon shape while the single crystals of MOF-908 appeared in form of trigonal prisms due to crystal growth primarily along the \(c\) direction (ESI,\(^{+}\) Section 2) resulting in a large unit cell in the crystal structure compared to PCN-280.

To further verify if the purely single phase of MOF-908 was obtained, the activated sample was digested for proton nuclear magnetic resonance spectroscopy (NMR). The post-digestion
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Due to the small size, large pore volume and nature of Fe-based MOFs, which leads to lowering of the resolution (a resolution of only 1.4 Å in the data collection was obtained by Mo X-ray radiation) of single crystal analysis, the MOF-908 crystall structure was finally elucidated via Rietveld refinement. This revealed that the structural single framework crystallized by Mo X-ray radiation) of single crystal analysis, the MOF-908 crystall structure was finally elucidated via Rietveld refinement. This revealed that the structural single framework crystallized in a trigonal lattice with an R3 space group (no. 146) and unit cell parameters of \(a = b = 32.9145\,\text{Å}\) and \(c = 46.0360\,\text{Å}\) (ESI† Section 3). MOF-909 is an isoreticular network of MOF-908 with similar unit cell parameters (\(a = b = 32.2658\,\text{Å}\) and \(c = 45.8488\,\text{Å}\)). The structure of MOF-908 is made up of two kinds of cage. The small primary cage with a pore size of 12 Å in diameter is a distorted octahedra constructed by six trigonal prism Fe–oxo clusters which link together through four BTB\(^{3−}\) linkers connecting the vertices along opposing equatorial edges, and three BPDC\(^{2−}\) moieties fill in the remaining equatorial edges composing the triangular window which is found to be 8 Å. The secondary cage is then built by the connection of eight primary cages with corners as nodes. In particular, those primary cages sharing four vertices, when translating a distance equal to the distance of two primary cages (approximately 23.2 Å), locate at the vertices of a cube and generate a cubic cage with a diameter of 23 Å. The square channel window of the secondary cage, which comprises four sharing vertices of the primary cage building units, was found to be 15.5 Å (Fig. 2b). Possessing the isoreticular structure of MOF-908, MOF-909, with the longer ditopic linking unit, had larger pore sizes but the triangular window size is smaller than that of MOF-908 due to the geometrical arrangement of 3,3′-azoBDC\(^{2−}\). In particular, the diameters of the distorted octahedra and cubic cage in MOF-909 are around 12.5 and 24 Å, respectively, while the triangular and square windows are calculated to be 7.5 and 16 Å, respectively. It is noted that all of the estimations are based on the model of the crystal structure verified by a Platon calculation, which is also applied to calculate the empty void space of approximately 82% for both MOF-909 and MOF-909.

The topological analysis of MOF-908 representing the underlying net of MOF-909, PCN-280, in PCN-285, which appeared to be isoreticular structures, was performed using the Topos Pro package.\(^{21}\) For the underlying net simplification, the Fe\(_3\)(O−CO\(_2\))\(_6\) SBU and the linkers containing BTB\(^{3−}\) and BPDC\(^{2−}\) were simplified as nodes which are 6 and 3 coordination points, corresponding to 6-c and 3-c, of the points of extension of the trigonal prismatic Fe–oxo cluster and the triangular node of the BTB moiety, respectively. The linear 2-c of the ditopic linker is reduced and no longer under consideration due to the formation of the 6-c point of extension for the Fe\(_3\)(O−CO\(_2\))\(_6\) nodes. It is interesting that the structural deconstruction revealed an unseen network, termed the nht net, in MOF chemistry. The transitivity of the nht topology is described by the number of vertices \(p\), the number of edges \(q\), the number of faces \(r\), and the type of tiling \(s\). Indeed, the p q r s transitivity of nht is 3553 and the natural tiling of this kind of network is \(3[4^{−7}] + 3[4^{−5}] + [3^{−5}]^{−7}\) (where the symbol \([\ldots m^{n}\ldots]\) denotes \(n\) faces with \(m\)-membered rings in a topological network) (Fig. 2c).

The solvent-exchange was carried out using an anhydrous acetone solvent and the full activation procedure was subsequently followed by heating the material under low pressure at 100 °C for 24 h. The chemical formula of MOF-908 is Fe\(_3\)(O−OH)(BTB)\(_{14/3}\)(BPDC), in which the mixed linker ratio of BTB:BPDC was found to be 4/3 : 1, as proven by \(^1\)H-NMR post-digestion (ESI†, Section 4). Thermal gravimetric analysis (TGA) measurements (ESI†, Section 4) further supported that formula because the observed residual mass (21.7%) perfectly matched with the expected mass of Fe\(_3\)O\(_3\) (22.8%) after fully decomposing...
soon after the disruption of the Fe–O bond in the first step between the two: the pore size of MOF-908 (~22 Å) is larger than that of PCN-280 (~12 Å) (ESI,† Section 4).

We subsequently carried out first-principles density functional theory (DFT) calculations to unveil the mechanism of interpenetration control of PCN-280 to formulate MOF-908. We proposed to look at two different theoretical aspects which are believed to be helpful in explaining the overall scenario. First the reaction mechanisms between \( \text{Fe}_3\text{O(OH)}(-\text{CO}_2)_{16} \) and \( \text{H}_2\text{BPDC}/3,3'\text{-azoBDC} \) were explored to clarify the energetic favorability of the two cluster-linker combinations. Second, by estimating elastic tensors of the abbreviated MOF-908 and MOF-909 structures, we were able to make conclusions on their relative stability in the condensed phase.

For the reaction mechanism investigation we employed the localized atomic-orbital-basis calculation method. The Perdew–Burke–Ernzerhof (PBE) functional was used to execute these calculations as well as the later condensed-phase calculations. The 6-31g* basis set was employed to construct the electronic wave-functions for C, H, O and N, while the LANL2dz basis set was utilized to describe the effective core potential for Fe. In both gas-phase and condensed-phase calculations we introduced a D3 correction to account for long-range van der Waals interactions. The quartet electronic spin state of the \( \text{Fe}_3\text{O(OH)}(-\text{CO}_2)_{16} \) cluster was revealed to be the most energetically-favored spin state; in other words, each Fe\(^{3+}\) site preferred to establish one unpaired electron. For the following investigation of reaction mechanisms we carried out all localized atomic-orbital-basis calculations at the quartet spin state.

As a linker (\( \text{H}_2\text{BPDC} \) or 3,3'-azoBDC) attempts to substitute an acetate (\( \text{CH}_3\text{COO}^- \)) residue, one O atom from the linker approaches closely and makes a coordination bond to one Fe cation in the cluster. Because of this connection the Mulliken charge of the involved Fe\(^{3+}\) site decreased to 0.57, which is less positive than that of the other two Fe sites (0.74) in the cluster. The binding energy of the \( \text{H}_2\text{BPDC} \)-cluster was 0.67 eV, while the binding energy of the \( 3,3'\text{-azoBDC} \)-cluster was 0.59 eV, which showed the slight favorability of \( \text{H}_2\text{BPDC} \) connection to the cluster. Such \( \text{Fe}_3\text{O(OH)}(-\text{CO}_2)_{16} \)-linker complexes were regarded as initial reactants. The proton (H\(^+\)) still remains linked with the linker molecule while attempting to make a weak bond with an O atom from acetate (Fig. 3a). To accomplish the cluster-linker connection, the system has to undergo two transformation stages: (i) an O atom from acetate first departs from Fe (a coordination bond is broken), and (ii) the O atom (from the –OH group of the linker) approaches and makes a connection with the other Fe cation linked to the same acetate residue. In the first process, \( \text{H}_2\text{BPDC} \) or 3,3'-azoBDC actually play a role as a catalyst to loosen the Fe-acetate linkage. Actually, we performed a relaxed scan of the Fe–O(acetate) dissociation in each case and found a barrier of 0.76 eV (17.5 kcal mol\(^{-1}\)) for both cases, as illustrated in Fig. 3b. Hence, there is competition between \( \text{H}_2\text{BPDC} \) and 3,3'-azoBDC in the first binding process to the Fe\(_3\text{O}\) cluster and the weakening of the Fe-acetate bond.

Soon after the disruption of the Fe–O bond in the first step the linker is more motivated to attack the second Fe site. In the
second process, we located the transition states when H$_2$BPDC/3,3'-azoBDC attempt to replace acetate. In those transition states, the Fe--O(H$_2$BPDC) distance was observed as 3.21 Å, while the Fe--O(3,3'-azoBDC) distance was similar (3.23 Å). However, the H$_2$BPDC transition state was found to be 0.43 eV (9.9 kcal mol$^{-1}$) above the initial reactant, while the 3,3'-azoBDC transition state was 0.76 eV (17.5 kcal mol$^{-1}$) above the initial reactant. The energetic scheme (Fig. 4) shows that 3,3'-azoBDC is less competitive in forming the framework.

Additionally, we also took into account a circumstance in which 3,3'-azoBDC successfully establishes connection to the cluster, and verified how H$_2$BPDC could come in and replace the 3,3'-azoBDC linker. We believe that the substitution should be identical to the mechanism described above when H$_2$BPDC (or 3,3'-azoBDC) substituted CH$_3$COO$^-$A. As a matter of fact, the transition state of H$_2$BPDC replacing 3,3'-azoBDC is 0.78 eV (18.0 kcal mol$^{-1}$) above the initial state when 3,3'-azoBDC is connected to the cluster. The final state, in which H$_2$BPDC makes a connection to the cluster and eliminates 3,3'-azoBDC, was found to be more stable than the initial state by 0.24 eV (5.5 kcal mol$^{-1}$). At this stage, by conducting localized atomic-orbital-basis calculations, we found that H$_2$BPDC and 3,3'-azoBDC are competitive to each other in the first stage of establishing bonding with one Fe cation. This first stage is very important because the participation of 3,3'-azoBDC is involved in the formulation of the single framework and avoided interpenetration. The second stage takes place rapidly after the first process, where H$_2$BPDC was shown to be preferable in bonding with the iron cluster.

We also performed PBE calculations for the condensed phases of MOF-908 and MOF-909 to evaluate their relative stability using the projector-augmented wave method implemented in the Vienna Ab Initio Package. Those two structures were extremely large and sophisticated and therefore, we chose to simplify the model by substituting three acetylene residues into the position of three phenyls in BTB (the crystallographic data of the original MOF-908 and MOF-909 as well as the modified structures are available in the ESI†). Such substitution was believed to alter the unit cell parameters insignificantly. To afford computational feasibility, planewave expansion was only done around the Γ point with a cut-off energy of 350 eV. After a simultaneous relaxation of unit cells and atomic positions, we calculated the $C_{ij}$ elastic tensors for both systems by adopting the stress–strain relationship. It was observed that the obtained $C_{ij}$ tensors of MOF-908 and MOF-909 satisfied the four conditions of mechanical stability of hexagonal unit cells. Moreover, the $C_{ij}$ tensors of MOF-908 were much larger than those of MOF-909. To make a more quantitative justification, we then computed the shear (G), bulk (K), and Young’s (E) moduli, as shown in Table 1 along with the $C_{ij}$ quantities. Remarkably, each of the moduli of MOF-908 was six times larger than the corresponding modulus computed for MOF-909. Moreover, all G, K, and E of MOF-909 were below 200 kbar, which is relatively low for a large unit cell and indicates that the framework of MOF-909 could suffer and collapse more easily. Such theoretical quantities clearly explain the experimental fact that MOF-909 actually partially collapsed after activation (the PXRD pattern shows the reduced intensity of the peaks located higher than 7.5° 2θ, ESI† Section 4). Additionally, MOF-908 is more mechanically stable than...
Table 1 Elastic tensors \( C_{ij} \) and shear (\( G \)), bulk (\( K \)), and Young’s (\( E \)) moduli of the abbreviated MOF-908 and MOF-909 structures (in kbar)

<table>
<thead>
<tr>
<th>MOFs</th>
<th>( C_{11} )</th>
<th>( C_{12} )</th>
<th>( C_{13} )</th>
<th>( C_{33} )</th>
<th>( C_{44} )</th>
<th>( C_{66} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>MOF-908</td>
<td>1460</td>
<td>400</td>
<td>387</td>
<td>1263</td>
<td>510</td>
<td>530</td>
</tr>
<tr>
<td>MOF-909</td>
<td>231</td>
<td>58</td>
<td>65</td>
<td>193</td>
<td>84</td>
<td>88</td>
</tr>
</tbody>
</table>

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<thead>
<tr>
<th>MOFs</th>
<th>( G )</th>
<th>( K )</th>
<th>( E )</th>
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</thead>
<tbody>
<tr>
<td>MOF-908</td>
<td>510</td>
<td>726</td>
<td>1241</td>
</tr>
<tr>
<td>MOF-909</td>
<td>82</td>
<td>114</td>
<td>199</td>
</tr>
</tbody>
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MOF-909, as proven by the pellet pressing test which clearly shows that the MOF-908 framework still remains up to 11 tons cm\(^{-2}\) of pressure while the MOF-909 structure collapses at a pressure of 5 tons cm\(^{-2}\) (ESI,† Section 4).

Conflicts of interest

There are no conflicts to declare.

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