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## Asymmetric linker generates intrinsically disordered metal–organic framework with local MOF-74 structure<sup>†</sup>

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Here, we report an intrinsically disordered MOF in the MOF-74 family,  $Mg_{2-x}(as\text{-}dobpdc)$  ( $as\text{-}dobpdc^{4-} = 3',4\text{-dioxidobiphenyl-3,4'\text{-dicarboxylate}}$ ). Despite the absence of crystallinity, this material exhibits local ordering consistent with that of its crystalline isomers, maintains porosity, and exhibits a high density of open metal sites.

Metal–organic frameworks (MOFs) are typically valued for their combination of crystallinity, porosity, and tunability.<sup>1</sup> However, amorphous MOFs can retain local order while introducing advantages, such as broader distribution of active sites types, reduced thermal conductivity, and emergent modes of tunability *via* control over the type of disorder.<sup>2,3</sup> Conventional amorphization strategies rely on external stimuli including pressure, temperature, stress, electrical discharge, or fast precipitation.<sup>4–10</sup> Recently, amorphous MOFs have been accessed directly using low symmetry organic linkers.<sup>11</sup>

Here, we employed a symmetry-reduction strategy to access an amorphous isomer within the MOF-74 structural family. In the benzene-based series, there are two isomers: 2,5-dihydroxybenzene-1,4-dicarboxylic acid and 4,6-dihydroxybenzene-1,3-dicarboxylic acid. These combine with transition metals to form the crystalline MOFs  $M_2(dobdc)$  (also known as MOF-74, CPO-27, and  $M_2(dhbdc)$ ) and  $M_2(m\text{-}dobdc)$ , respectively.<sup>12,13</sup> For the biphenyl-derived series, two isomers have been reported:  $M_2(pc\text{-}dobpdc)$  (also referred to as IRMOF-74-II;  $pc\text{-}dobpdc^{4-} = 3,3'\text{-dioxidobiphenyl-4,4'\text{-dicarboxylate}}$ ) and  $M_2(mc\text{-}dobpdc)$  ( $mc\text{-}dobpdc^{4-} = 4,4'\text{-dioxidobiphenyl-3,3'\text{-dicarboxylate}}$ , where *pc* and *mc* refer to *para*- and *meta*-carboxylate substitution,

respectively).<sup>14,15</sup> Here, we investigate a third isomer,  $3',4\text{-dioxidobiphenyl-3,4'\text{-dicarboxylate}}$  ( $as\text{-dobpdc}^{4-}$ ), which yields a new porous material,  $Mg_{2-x}(as\text{-dobpdc})$ , where “*as*” refers to asymmetric substitution. Even though the secondary building units for  $Mg_2(mc\text{-}dobpdc)$  and  $Mg_2(pc\text{-}dobpdc)$  have similar connectivity, differences in bond angles introduce geometric frustration in  $Mg_{2-x}(as\text{-dobpdc})$ , leading to amorphization and loss of long-range order (Fig. 1). Despite this,  $Mg_{2-x}(as\text{-dobpdc})$  retains porosity and has pore size distributions and local order (as probed by X-ray pair distribution function) comparable to those of its crystalline counterparts. Additionally,  $Mg_{2-x}(as\text{-dobpdc})$  retains open metal sites, with  $\sim 0.9$  sites per Mg.

Synthetic conditions for  $Mg_{2-x}(as\text{-dobpdc})$  were adapted from previously reported procedures for  $Mg_2(mc\text{-}dobpdc)$  and  $Mg_2(pc\text{-}dobpdc)$ . Most synthesis attempts yielded either amorphous solids or metal salts, as confirmed by powder X-ray diffraction (PXRD) (Fig. S20–S22, ESI<sup>†</sup>). However, an optimized synthesis condition using  $Mg(NO_3)_2\cdot 6H_2O$  and  $H_4(as\text{-}dobpdc)$  in a 1:1 (vol) mixture of *N,N*-diethylformamide and methanol at 120 °C in a silanized vial produced a pale yellow solid, denoted  $Mg_{2-x}(as\text{-dobpdc})$ . PXRD of this material exhibited broad, low-intensity diffuse peaks centered at 2θ values of  $\sim 5^\circ$  and  $\sim 8^\circ$  (Fig. 2), indicative of a largely amorphous material with some local order. These reflections align with the characteristic {100} and {110}/{110} peaks of crystalline  $Mg_2(mc\text{-}dobpdc)$  (space group  $P3_121$ ; CCDC 1827449) and the {110} and {300} reflections of  $Mg_2(pc\text{-}dobpdc)$  (space group  $R\bar{3}$ ; CCDC 841642), suggesting structural similarity at the local scale. A model structure of an idealized  $Mg_2(as\text{-}dobpdc)$  framework (Fig. S18, ESI<sup>†</sup>) also shows simulated PXRD peaks at these positions (Fig. S19, ESI<sup>†</sup>), providing a plausible depiction of local connectivity. The PXRD of  $Mg_{2-x}(as\text{-dobpdc})$  remained unchanged after washing with *N,N*-dimethylformamide (DMF) and methanol, or after activation under dynamic vacuum at 180 °C (Fig. S23, ESI<sup>†</sup>). Thermogravimetric analysis (TGA) of  $Mg_{2-x}(as\text{-dobpdc})$  did not show significant mass loss of material until temperatures  $> 200$  °C (Fig. S37, ESI<sup>†</sup>).

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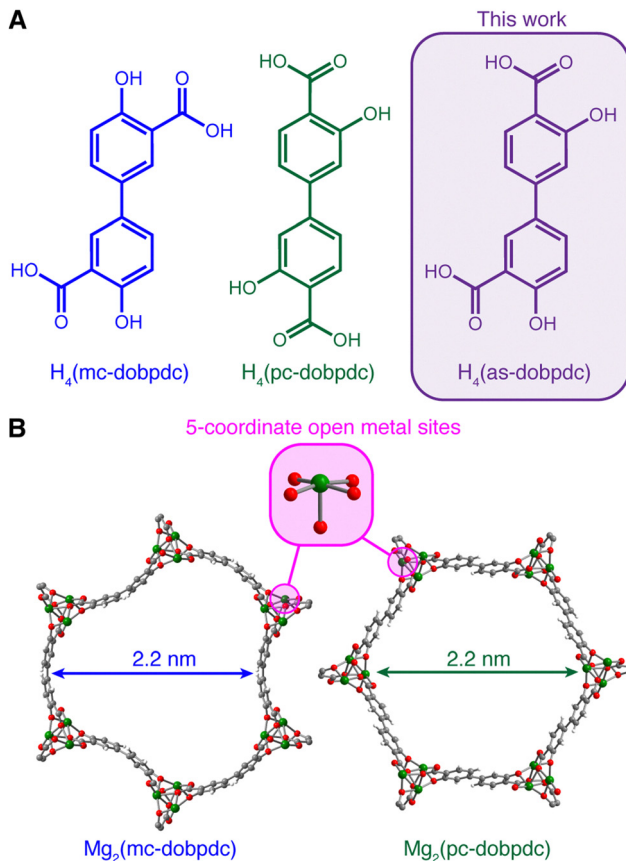


Fig. 1 (A) Comparison of  $H_4(\text{mc-dobpdc})$  and  $H_4(\text{pc-dobpdc})$  linkers used to form previously reported MOF-74-type structures with  $H_4(\text{as-dobpdc})$  linker used in this work. (B) Depiction of structures of  $Mg_2(\text{mc-dobpdc})$  and  $Mg_2(\text{pc-dobpdc})$  which both contain hexagon-shaped pores and coordinatively unsaturated metal sites.

We next attempted to determine the composition of  $Mg_{2-x}(\text{as-dobpdc})$ .  $^1\text{H}$  NMR analysis of digested activated  $Mg_{2-x}(\text{as-dobpdc})$  (using  $\text{D}_2\text{SO}_4$  and  $\text{d}_6\text{-DMSO}$ , in accordance with procedures used for other MOF-74 materials<sup>16–18</sup>) showed the presence of  $H_4(\text{as-dobpdc})$  and residual DMF in a ratio of  $\sim 0.45$  DMF per  $H_4(\text{as-dobpdc})$ . This residual solvent could not be removed under the activation conditions used. Thermo-gravimetric analysis (TGA) was employed to quantify the

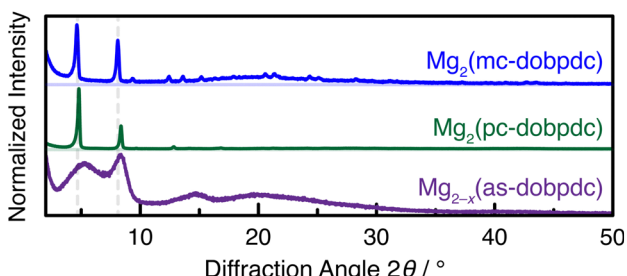


Fig. 2 Powder X-ray diffractograms (Cu  $\text{K}\alpha$  radiation,  $\lambda = 1.5418 \text{ \AA}$ ) of  $Mg_2(\text{mc-dobpdc})$ ,  $Mg_2(\text{pc-dobpdc})$ , and  $Mg_{2-x}(\text{as-dobpdc})$  obtained by dropcasting the solids from the synthesis suspension. Dashed lines at  $\sim 5^\circ$  and  $\sim 8^\circ$  are shown to guide the eye.

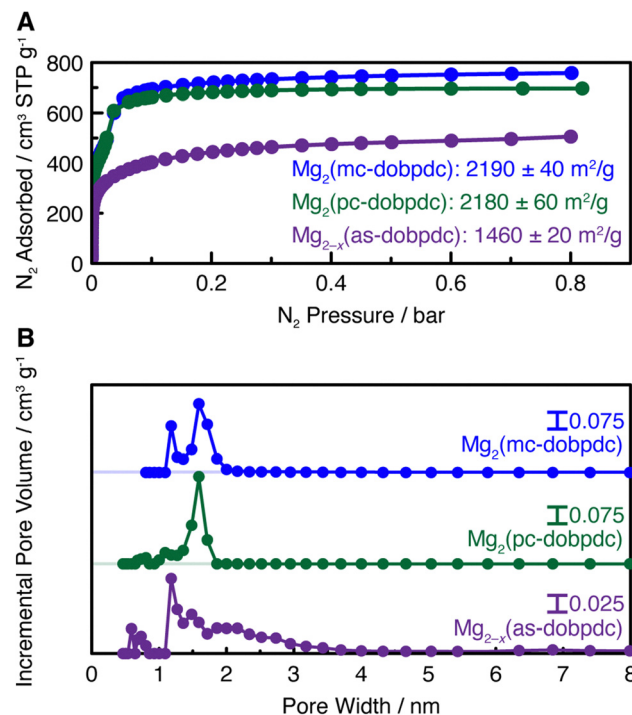


Fig. 3 (A) 77 K  $\text{N}_2$  adsorption isotherms of activated  $Mg_2(\text{mc-dobpdc})$ ,  $Mg_2(\text{pc-dobpdc})$ , and  $Mg_{2-x}(\text{as-dobpdc})$  with fitted BET surface areas. (B) Pore size distributions obtained from fitting the  $\text{N}_2$  adsorption isotherms.

inorganic content: the sample was first activated at  $180^\circ\text{C}$ , then oxidized in air at  $800^\circ\text{C}$  (Fig. S38, ESI†), yielding  $\text{MgO}$  as the final residue (Fig. S40, ESI†). Combining TGA and NMR data yields a formula of  $\text{Mg}_{1.9}\text{H}_{0.2}(\text{as-dobpdc})[\text{DMF}]_{0.45}$ . A comparable empirical formula,  $\text{Mg}_{1.76}\text{H}_{0.48}(\text{as-dobpdc})[\text{DMF}_{0.45}]$  [ $(\text{H}_2\text{O})_{0.69}$ ], was independently derived from elemental analysis by fitting C, H, N, and Mg content (Section S10, ESI†). Both formulas ( $Mg_{2-x}(\text{as-dobpdc})$  with  $x = 0.1\text{--}0.25$ ) suggest that although the ratio of Mg to linker is close to the ideal value of 2, there is an excess of linker relative to Mg that is consistent with the low crystallinity of the material.

To understand the structure of  $Mg_{2-x}(\text{as-dobpdc})$ , we collected  $\text{N}_2$  adsorption isotherms at 77 K and compared with those of  $Mg_2(\text{mc-dobpdc})$  and  $Mg_2(\text{pc-dobpdc})$  (Fig. 3A). The Brunauer–Emmett–Teller (BET) surface areas were  $2190 \pm 40 \text{ m}^2 \text{ g}^{-1}$  for  $Mg_2(\text{mc-dobpdc})$ ,  $2180 \pm 60 \text{ m}^2 \text{ g}^{-1}$  for  $Mg_2(\text{pc-dobpdc})$ , and  $1460 \pm 20 \text{ m}^2 \text{ g}^{-1}$  for  $Mg_{2-x}(\text{as-dobpdc})$ . Although the BET surface area for  $Mg_{2-x}(\text{as-dobpdc})$  is lower than that of the crystalline analogues, the material still displays remarkable porosity despite its largely amorphous character.

The pore size distributions were calculated with the “ $\text{N}_2$  – DFT Model” as implemented in Micromeritics software (Fig. 3B). The fitted distributions for  $Mg_2(\text{mc-dobpdc})$  and  $Mg_2(\text{pc-dobpdc})$  predominantly show micropores, consistent with the expected crystallographic structures. Interestingly, the fitted distribution for  $Mg_{2-x}(\text{as-dobpdc})$  shows significant microporosity, with most pores falling below 4 nm in width. Although the calculated values are approximate, owing to the use of an idealized slit-pore model, they clearly show that

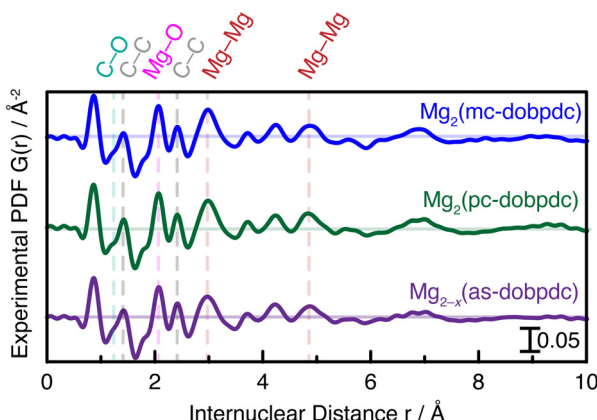


Fig. 4 Pair distribution functions (PDFs) of  $\text{Mg}_2(\text{mc-dobpdc})$ ,  $\text{Mg}_2(\text{pc-dobpdc})$ , and  $\text{Mg}_{2-x}(\text{as-dobpdc})$ . Labelled vertical dashed lines correspond to distances between particular element pairs.

$\text{Mg}_{2-x}(\text{as-dobpdc})$  contains a large fraction of micropores, similar to the other analogues.

To further probe the local structure, we performed X-ray pair distribution function (PDF) analysis to determine interatomic distances in the three materials. While PDF does not provide an exact global structural model for  $\text{Mg}_{2-x}(\text{as-dobpdc})$ ; comparison with the crystalline analogues provides information for the local structure and the ordering length scaling. Interestingly, all three PDFs are almost identical for internuclear distances  $< 10 \text{ \AA}$  (Fig. 4), suggesting their local structures are preserved despite differences in long-range order. At larger distances ( $10\text{--}30 \text{ \AA}$ ), the PDF intensity for  $\text{Mg}_{2-x}(\text{as-dobpdc})$  decays faster (Fig. S43, ESI†), as expected from the lower crystallinity.

Using previously obtained crystal structures for  $\text{Zn}_2(\text{mc-dobpdc})$  and  $\text{Zn}_2(\text{pc-dobpdc})$  (isostructural to the Mg analogues), we simulated PDFs in order to assign specific element pairs to the peaks in the experimental PDF (Fig. S42, ESI†). From this analysis, we found that in addition to C–O and C–C pairs occurring at similar internuclear distances (expected as they originate solely from the framework linker), we also found that Mg–O and Mg–Mg pairs occur at similar internuclear distances in all three frameworks, providing further evidence that the local metal–ligand geometry is preserved in  $\text{Mg}_{2-x}(\text{as-dobpdc})$ , despite its lack of long-range order.

As an additional test of local structure, we sought to determine if any of the Mg atoms in  $\text{Mg}_{2-x}(\text{as-dobpdc})$  contained open metal sites. As all of the Mg sites formed in  $\text{Mg}_2(\text{mc-dobpdc})$  and  $\text{Mg}_2(\text{pc-dobpdc})$  are coordinatively unsaturated, we hypothesized that if  $\text{Mg}_{2-x}(\text{as-dobpdc})$  had a similar local structure to the other analogues, then it should also contain a high density of open metal sites. Previous studies have demonstrated that these open sites can be functionalized with diamines for  $\text{CO}_2$  capture, providing a strategy for quantifying their accessibility.<sup>16–18</sup> Using established protocols, we appended 2-methyl-1,2-diaminopropane (dmen) to  $\text{Mg}_{2-x}(\text{as-dobpdc})$ , targeting a 1:1 dmen:Mg stoichiometry as observed in the crystalline analogues.<sup>16–18</sup> To quantify the Mg-bound dmen, we activated dmen– $\text{Mg}_{2-x}(\text{as-dobpdc})$  at  $130^\circ\text{C}$  under

$\text{N}_2$  via TGA to remove physisorbed dmen, followed by oxidation to  $\text{MgO}$  by heating to  $800^\circ\text{C}$  under air, where the dmen content could be calculated by the difference in inorganic:organic ratios before and after appending dmen to the framework. We also used the same activation procedure via TGA on a different batch of dmen– $\text{Mg}_{2-x}(\text{as-dobpdc})$ , after which we performed  $^1\text{H}$  NMR after digestion to quantify the ratio of diamine to linker. This  $^1\text{H}$  NMR did not contain DMF, indicating that dmen displaced DMF coordinated to the Mg sites during appending. Both the TGA method ( $\sim 0.93$  dmen:Mg, Fig. S46, ESI†) and the  $^1\text{H}$  NMR method ( $\sim 0.9$  dmen:Mg, Fig. S47, ESI†) provided similar dmen:Mg ratios of  $\sim 90\%$ , suggesting that most Mg sites in the framework strongly bind dmen, and were likely coordinatively unsaturated. Of these open sites, elemental analyses suggest that  $\sim 80\%$  can be accessed after activation, with the rest remaining bound to DMF, which can be displaced by dmen.

To probe if the local environment of these coordinatively unsaturated metal sites was similar in the three frameworks, we performed transmission Fourier transform infrared spectroscopy (IR) using acetone as a probe molecule. Specifically, the  $\text{C}=\text{O}$  stretching frequency,  $\nu(\text{C}=\text{O})$ , of acetone is a well-established reporter of Lewis acidity, with red-shifted frequencies corresponding to stronger Lewis acids.<sup>19</sup> In addition, this vibrational frequency is sensitive to local electric fields, which can stabilize the C–O bond dipole and similarly lead to red shifts.<sup>20</sup> Therefore, we hypothesized that if the  $\nu(\text{C}=\text{O})$  frequencies of Mg-bound acetone were significantly different between the three frameworks, it could indicate either a difference in Lewis acidity of the Mg or a difference in the local environment generating an electric field; either case would reflect changes in the local structure.

We collected IR spectra by first activating pelletized samples under flowing  $\text{N}_2$  at  $130^\circ\text{C}$ , followed by cooling to  $30^\circ\text{C}$  under the same atmosphere. The gas stream was then switched to  $\text{N}_2$  saturated with acetone vapor to fill the framework pores, and subsequently returned to pure  $\text{N}_2$  to monitor desorption. Spectra were recorded every 2 minutes relative to the acetone-free baseline at  $30^\circ\text{C}$  to isolate features associated with strongly bound acetone, which is expected to desorb more slowly than the physisorbed species. For both  $\text{Mg}_2(\text{mc-dobpdc})$  (Fig. S53, ESI†) and  $\text{Mg}_2(\text{pc-dobpdc})$  (Fig. 5A), desorption spectra show a strongly bound peak in the  $\nu(\text{C}=\text{O})$  region at  $\sim 1727 \text{ cm}^{-1}$ , which we attribute to Mg-bound acetone, in addition to a quickly desorbing peak at  $\sim 1710 \text{ cm}^{-1}$ , which we attribute to intraporous acetone. For  $\text{Mg}_{2-x}(\text{as-dobpdc})$ , we observe a similar strongly bound acetone feature at  $\sim 1725 \text{ cm}^{-1}$ , suggesting that the local environment of the Mg-bound acetone is similar in all three frameworks. We note that there is some baseline fluctuation for  $\text{Mg}_{2-x}(\text{as-dobpdc})$  between  $1640\text{--}1480 \text{ cm}^{-1}$ , due to perturbation of the  $\nu(\text{C}=\text{O})$  stretching frequency from residual DMF interacting with acetone as well as the DMF vibrational modes (Fig. S52, ESI†).

Here, we have demonstrated that the amorphous MOF  $\text{Mg}_{2-x}(\text{as-dobpdc})$  has little to no global translational symmetry, yet preserves porosity. The framework has similar pore size



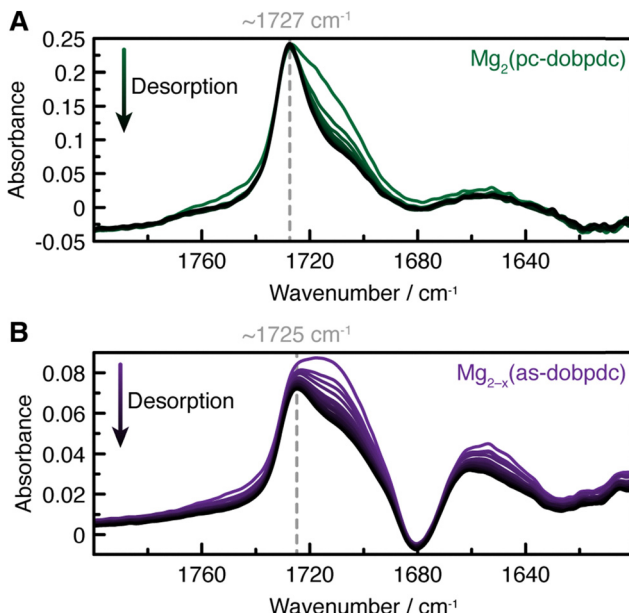


Fig. 5 Transmission-mode Fourier transform infrared spectra of acetone desorption from (A)  $\text{Mg}_2(\text{pc-dobpdc})$  and (B)  $\text{Mg}_{2-x}(\text{as-dobpdc})$ , background-subtracted from the bare framework spectra. The peaks marked by dashed gray lines ( $\sim 1727 \text{ cm}^{-1}$  and  $\sim 1725 \text{ cm}^{-1}$ ) are assigned to the  $\text{C}=\text{O}$  mode of acetone bound to the Mg sites.

distribution and local order to both  $\text{Mg}_2(\text{mc-dobpdc})$  and  $\text{Mg}_2(\text{pc-dobpdc})$ . Moreover, it maintains a high degree ( $\sim 0.9$  sites per Mg) of open metal sites, suggesting that even without crystallinity, the framework may possess similar functionality. We expect that the Mg deficiency, as compared to the crystalline isomorphs, will result in defects that have potential applications in catalysis and gas sorption/separation. Similarly, the difference in pore size distribution could potentially be exploited for adsorbing larger molecules than the ones accessible for crystalline isomorphs. While no crystalline phase of  $\text{Mg}_{2-x}(\text{as-dobpdc})$  has yet been isolated, we cannot rule out the possibility that appropriate synthetic conditions may yield a crystalline phase. We envision that this strategy of linker symmetry reduction might lead to a new class of amorphous MOFs that may have emergent functionality.

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## Conflicts of interest

There are no conflicts to declare.

## Data availability

The data supporting this article have been included as part of the ESI.<sup>†</sup>

## Notes and references

- 1 S. R. Batten, N. R. Champness, X.-M. Chen, J. Garcia-Martinez, S. Kitagawa, L. Öhrström, M. O'Keeffe, M. P. Suh and J. Reedijk, *Pure Appl. Chem.*, 2013, **85**, 1715–1724.
- 2 T. D. Bennett and A. K. Cheetham, *Acc. Chem. Res.*, 2014, **47**, 1555–1562.
- 3 J. Fonseca, T. Gong, L. Jiao and H.-L. Jiang, *J. Mater. Chem. A*, 2021, **9**, 10562–10611.
- 4 W. M. W. Winters, C. Zhou, J. Hou, M. Diaz-Lopez, T. D. Bennett and Y. Yue, *Chem. Mater.*, 2024, **36**, 8400–8411.
- 5 Y. Feng, M.-Y. Zou, H.-C. Hu, W.-H. Li, S.-L. Cai, W.-G. Zhang and S.-R. Zheng, *Chem. Commun.*, 2022, **58**, 5013–5016.
- 6 N. L. Baun, S. S. Mortensen, H. Bordallo, K. Jensen and A. Sapnik, *ChemRxiv*, 2025, preprint, DOI: [10.26434/chemrxiv-2025-psg4n](https://doi.org/10.26434/chemrxiv-2025-psg4n).
- 7 H. Wang, Y. Liu, L. Li, J. Zhang, M. Luo, Z. Sun, Y. Xiao, X. Zhai, L. Wu, H. Zhang, B. Ye, C. Yang, X. Zhang and M. Zhou, *Angew. Chem., Int. Ed.*, 2025, **137**(28), e202506960.
- 8 R. Zacharia, D. Cossement, L. Lafi and R. Chahine, *J. Mater. Chem.*, 2010, **20**, 2145–2151.
- 9 Y. Zhou and C. Liu, *Plasma Chem. Plasma Process.*, 2011, **31**, 499–506.
- 10 Y. H. Hu and L. Zhang, *Phys. Rev. B: Condens Matter Mater. Phys.*, 2010, **81**, 174103.
- 11 E. G. Meekel, E. M. Schmidt, L. J. Cameron, A. D. Dharma, H. J. Windsor, S. G. Duyker, A. Minelli, T. Pope, G. O. Lepore, B. Slater, C. J. Kepert and A. L. Goodwin, *Science*, 2023, **379**, 357–361.
- 12 N. L. Rosi, J. Kim, M. Eddaoudi, B. Chen, M. O'Keeffe and O. M. Yaghi, *J. Am. Chem. Soc.*, 2005, **127**, 1504–1518.
- 13 M. T. Kapelewski, S. J. Geier, M. R. Hudson, D. Stück, J. A. Mason, J. N. Nelson, D. J. Xiao, Z. Hulvey, E. Gilmour, S. A. FitzGerald, M. Head-Gordon, C. M. Brown and J. R. Long, *J. Am. Chem. Soc.*, 2014, **136**, 12119–12129.
- 14 T. M. McDonald, W. R. Lee, J. A. Mason, B. M. Wiers, C. S. Hong and J. R. Long, *J. Am. Chem. Soc.*, 2012, **134**, 7056–7065.
- 15 H. Deng, S. Grunder, K. E. Cordova, C. Valente, H. Furukawa, M. Hmадех, F. Gándara, A. C. Whalley, Z. Liu, S. Asahina, H. Kazumori, M. O'Keeffe, O. Terasaki, J. F. Stoddart and O. M. Yaghi, *Science*, 2012, **336**, 1018–1023.
- 16 R. L. Siegelman, T. M. McDonald, M. I. Gonzalez, J. D. Martell, P. J. Milner, J. A. Mason, A. H. Berger, A. S. Bhowm and J. R. Long, *J. Am. Chem. Soc.*, 2017, **139**, 10526–10538.
- 17 P. J. Milner, J. D. Martell, R. L. Siegelman, D. Gygi, S. C. Weston and J. R. Long, *Chem. Sci.*, 2017, **9**, 160–174.
- 18 B. Dinakar, A. C. Forse, H. Z. H. Jiang, Z. Zhu, J.-H. Lee, E. J. Kim, S. T. Parker, C. J. Pollak, R. L. Siegelman, P. J. Milner, J. A. Reimer and J. R. Long, *J. Am. Chem. Soc.*, 2021, **143**, 15258–15270.
- 19 K. Shimizu, T. Higuchi, E. Takasugi, T. Hatamachi, T. Kodama and A. Satsuma, *J. Mol. Catal. A: Chem.*, 2008, **284**, 89–96.
- 20 S. D. E. Fried, C. Zheng, Y. Mao, T. E. Markland and S. G. Boxer, *J. Phys. Chem. B*, 2022, **126**, 5876–5886.

