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Defects and disorder in metal organic frameworks

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The wide-ranging properties of metal organic frameworks (MOFs) rely in many cases on the presence of defects within their structures and the disorder that is inevitably associated with such defects. In the present work we review several aspects of defects in MOFs, ranging from simple substitutional defects at metal cation or ligand positions, to correlated defects on a larger length scale and the extreme case of disorder associated with amorphous MOFs. We consider both porous and dense MOFs, and focus particularly on the way in which defects and disorder can be used to tune physical properties such as gas adsorption, catalysis, photoluminescence, and electronic and mechanical properties.

1. Introduction

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It is widely recognized in physics and chemistry that the presence of both point and extended scale defects in materials can have a dramatic impact on their properties. In the case of inorganic structures, for example, simple lattice and interstitial vacancies control diffusion and ionic conductivity, while sub-

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Tony Cheetham obtained his D.Phil. at Oxford in 1971 and joined chemistry faculty in 1974. In 1992, he took up the Directorship of the new Materials Research Laboratory at the University of California at Santa Barbara, before moving to Cambridge in 2007 to become the Goldsmiths' Professor of Materials Science. Tony is a Fellow of the Royal Society (1994) and has received numerous major awards for his work in

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focuses on the formation of liquids and glasses from crystalline metal-organic framework materials.

View Article Online View Journal | View Issue about the identity or propensity of their defects, though a very recent review by Fang *et al.*¹ gives a good coverage of the literature from a crystal engineering perspective, with a particular emphasis on porous MOFs and their applications. The purpose of this short review is to explore the interplay between defects and disorder in both porous and dense MOFs.

We shall focus primarily on defects that are introduced artificially, since these have been quite widely studied. We know, for example, that defects can be used to introduce new functionality into MOFs, *e.g.* photoluminescence, active sites for catalysis, and enhanced selective adsorption, as well as for fine-tuning properties such as magnetic and ferroelectric phase transitions. It is however also known that the presence of defects may undermine the performance of MOFs, as in the case of porous solids where selective adsorption capacity can be reduced by poor crystallinity.

It is important to recognize that defects are normally associated with disorder, and so entropic factors are a major driving force in determining the levels of defects in solids. All materials under ambient conditions – however pure they may be – contain intrinsic defects according to the partition function:

$$n_{\rm d} \sim e^{-e_{\rm d}/RT} \tag{1}$$

where n_d is the number of defects of a particular type and e_d is the energy required to form them. The concepts set out for simple ionic solids by Schottky and Wagner in 1930² pointed to the importance of Schottky (equal numbers of complementary cation and anion vacancies) and Frenkel (equal numbers of cation or anion vacancies and interstitials) defects in stoichiometric inorganic solids. These are known as intrinsic defects. Each particular solid normally has a preferred intrinsic, point defect incorporation mechanism; in the case of sodium chloride, for example, the dominant defect is of the Schottky type, and we can estimate that a mole of NaCl contains approximately 10⁴ cation and anion vacancies at room



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temperature (and as many as 10^{17} at 1000 K). A solid will also contain extrinsic defects due to impurities, and the relative proportions of intrinsic and extrinsic defects will depend on the purity of the sample as well as the temperature. By contrast, we do not know the nature of the intrinsic defects in MOFs.

The scope of this mini-review will span both porous and dense MOFs, recognizing that the ease of defect formation may be different. We shall also consider the correlation of such defects, along with the relationship between defects and disorder (section 4), and in particular the extreme case of disorder that is amorphization (section 5). Defects and disorder caused by extra-framework guests or solvent molecules, which are ubiquitous in porous MOFs, and extended defects such as dislocations and stacking faults, which are covered in ref. 1, are avoided. The following two sections will give examples of defects on the cation and anion sub-lattices, respectively.

2. Cation defects in metal–organic frameworks

2.1 Examples with metal cation substitution

There are two contrasting synthetic approaches to the formation of substitutional cation defects in MOFs. One approach aims to crystallize the materials from starting solutions containing the two cations in the desired proportions. An appropriate example is taken from calcium and strontium tartrates ($C_4H_4O_6^{2-}$), where crystallization from a solution containing the two cations, in the presence of the appropriate tartrate ligand, is possible.

$$(1 - x)Ca^{2+}(aq) + xSr^{2+}(aq) + C_4H_4O_6^{2-}(aq)$$

$$\rightarrow Ca_{1-x}Sr_x(C_4H_4O_6)$$
(2)

Alkaline earth tartrates can be made in this manner and provide one of the simplest cases of cation substitution in



Fig. 1 (a) The X-ray patterns of $Ca_{1-x}Sr_x(L-C_4H_4O_6)$ show systematic shifts as a function of composition. (b) The unit cell volume obeys Végard's Law. (c) The structure of Ca ($L-C_4H_4O_6$) viewed along the *a* axis. Ca – blue, O – red, C – black, H – white. Adapted with permission from L. Appelhans, *et al., J. Am. Chem. Soc.*, 2009, **131**, 15375. Copyright 2009 American Chemical Society.

dense MOFs.³ Calcium and strontium L-tartrates are isomorphous and form a complete solid solution, $Ca_{1-x}Sr_x(L-C_4H_4O_6)$, which obeys Végard's Law, *i.e.* the cell volume is a linear function of composition (Fig. 1).

Complexities arise where, as in the corresponding strontium–barium system, the two end-member structures are different. In the case of the *meso*-tartrate phases, there is a solid solution over a very limited range of composition, $Sr_{1-x}Ba_x(meso-C_4H_4O_6)$ for 0 < x < 0.1, followed by a two phase region (0.1 < x < 0.2) in which this anhydrous *meso*-tartarte, $Sr_{0.9}Ba_{0.1}(meso-C_4H_4O_6)$ co-exists with a monohydrate phase, $Sr_{0.8}Ba_{0.2}(meso-C_4H_4O_6)(H_2O)$. This monohydrate then forms a solid solution across the range 0.2 < x < 1.0. It is not unusual to find examples of this type where phases with different compositions will crystallize from mixed cation starting solutions.

Reactions such as those illustrated in eqn (2) works well in some instances, but can be problematic when one of the endmember phases crystallizes more rapidly than the solid solution, resulting in an inhomogeneous product. In such circumstances, a second approach involving post-synthetic cation exchange (PCE) might be adopted. There are two variations on this theme. In the first case, the two end-member MOFs are prepared separately and the two solids are then allowed to undergo cation exchange in solution:

$$(1-x)\mathbf{M}(\mathbf{L}) + x\mathbf{M}'(\mathbf{L}) \to \mathbf{M}_{1-x}\mathbf{M}'_{x}(\mathbf{L})$$
(3)

In a second variation on the PCE approach, only one of the end-members is prepared and this is allowed to undergo ionexchange with a solution containing the second cation:

$$(1-x)\mathbf{M}(\mathbf{L}) + x\mathbf{M}'(\mathbf{aq}) \to \mathbf{M}_{1-x}\mathbf{M}'_{x}(\mathbf{L}) + x\mathbf{M}(\mathbf{aq})$$
(4)

These two PCE approaches has been successfully used to synthesize mixed (Al/Fe) samples of MIL-53 and mixed (Zr/Ti) samples of UiO-66, respectively.⁴

The use of cation solid solutions has been used to control adsorption properties in porous MOFs.⁵ By exploiting the reactivities of different precursors to form mixed cation systems

according to eqn (2), it has been possible to form both homogeneous solid solutions on the one hand, and phase separated materials on the other. This was achieved in a MOF system based upon manganese and zinc with 5-nitroisophthalate and 4,4'-bipyridyl as linkers. Classical solid solutions are obtained when using sodium 5-nitroisophthalate as a precursor because it reacts very rapidly, but phase-separated products are found when the 5-nitroisophthalic acid is used (Fig. 2). The two different samples showed strikingly different adsorption behaviour with methanol.

A second approach to the realization of mixed-metal MOFs leverages the ability to substitute one cation for another into an existing MOF, a process called transmetalation,⁶ that can create new functionality. It has been exploited quite extensively for optical materials where the levels of substitution that are needed are usually quite small. Rare-earth ions, especially Eu³⁺ and Tb^{3+} , have been inserted at low levels (typically <3%) into Y³⁺, Gd³⁺ and Bi³⁺ containing host frameworks to create photoluminescent materials that are suitable for use as phosphors. In an early example in glutarate $(C_5H_6O_4^{3-})$ frameworks, Eu³⁺ was substituted for Gd^{3+} in a $Gd(C_5H_6O_4)\cdot nH_2O$ phase and the photoluminscent lifetime studied as a function of the level of water in the MOF channels.⁷ One of the attractions of MOFs for photoluminescence applications is that the organic ligands themselves can be used as sensitizers to absorb the excitation, as in the case of the Eu³⁺- and Tb³⁺-doped sodium/bismuth 1,4-benzenedicarboxylates, which show very good quantum efficiencies (Fig. 3).8

Transmetalation, sometimes also called "post-synthetic metal exchange",⁹ has also been used in the area of gas separation and storage to control adsorption and transport properties in porous MOFs. One such example is the use of Ti-exchanged UiO-66 [$Zr_6O_6(OH)_4(BDC)$] to increase gas permeability in mixed matrix membranes.¹⁰ It has also been shown that metal exchange can be used to obtain materials with topologies unreachable through solvothermal synthesis. One example is the case of MOFs based on the *N*,*N*'-bis(3,5-dicarbo-xyphenyl)pyromellitic diimide (H₄BDCPPI) linker. Solvothermal



Fig. 2 (a, b) Different distributions can be seen in the X-ray powder patterns. (c) Schematic showing the formation of solid solutions and phase-separated products. Adapted with permission from T. Fukushima, *et al., J. Am. Chem. Soc.*, 2012, **134**, 13341. Copyright 2012 American Chemical Society.



Fig. 3 (a) Unit cell of sodium/bismuth 1,4-benzenedicarboxylate, which, when doped with Eu^{3+} - and Tb^{3+} -shows (b) excellent luminescence in the red and green, respectively, when excited in the ultra violet. Adapted with permission from A. Thirumurugan, J.-C. Tan and A. K. Cheetham, *Cryst. Growth Des.*, 2010, **10**, 1736. Copyright 2010 American Chemical Society.



Fig. 4 The cobalt/nickel succinates form four different structures with entirely different compositions when crystallized under the same conditions. Reprinted with permission from C. Livage, *et al.*, *Angew. Chem., Int. Ed.*, 2007, **46**, 5877. Copyright 2007. John Wiley and Sons.

synthesis yield materials with nbo and pts nets respectively, but metal exchange from the copper-based material allowed Prasad *et al.* to obtain a $Zn_2(BDCPPI)$ framework which retained pts topology.¹¹

However, many systems that are superficially similar may show complex phase behaviour in reality. Transition metal succinates, for example, are notoriously fickle and yield complex results. In the case of the cobalt and nickel succinates $(C_4H_4O_4^{2-})$, the two end-members have different compositions and structures, $Co_5(OH)_2(C_4H_4O_4)_4$ and $Ni_7(OH)_2(H_2O)_2$ $(C_4H_4O_4)_6\cdot 2H_2O$, respectively, when crystallized under the same conditions. At intermediate Co/Ni ratios, yet other compositions and structures are obtained: $M_4(OH)_2(H_2O)_2$ $(C_4H_4O_4)_3\cdot 2H_2O$ and $M_7(OH)_2(H_2O)_2(C_4H_4O_4)_6\cdot 3.5H_2O$ for 25% and 75% nickel, respectively (Fig. 4), with disordered cations in each case.¹²



Fig. 5 (a) Structure of the fully ordered hybrid perovskite, $[(CH_3)_2NH_2]Cu^{II}(HCOO)_3$. Reproduced from ref. 18 with permission from The American Physical Society (2013). (b) The hydrogen bonding between the amine cation and the iodide anions in the A-site cavity of $[CH_3NH_3]Pb^{II}I_3$. Reproduced from ref. 19 with permission from The Royal Society of Chemistry (2015).

2.2 Amine cations in hybrid perovskites: substitution and disorder

Cations that can undergo substitution are not restricted to metallic ions. The hybrid perovskites, which some may consider MOF-type materials, of general formula $[AmH]M^{II}X_3$, where Am is an organic amine cation and X can be a wide range of anions, *e.g.* formate, azide, halide, borohydride, provide important examples (Fig. 5). The $[AmH]Pb^{II}I_3$ frameworks are of great current interest because of their outstanding performance as semiconductors in solar cells.^{13,14} The most widely studied system is $[CH_3NH_3]Pb^{II}I_3$, and it is possible to tune the band gap by partially substituting the methylammonium cation by formamidinium, thereby increasing the efficiency to greater than 14%.¹⁵ This fine-tuning is normally achieved during the initial synthesis according to eqn (2), rather than by PSE.

Halide perovskites also exhibit disorder of a different nature: order–disorder transitions associated with hydrogen bonding between the amine hydrogens and the iodide anions. $[CH_3NH_3]Pb^{II}I_3$ itself, for example, is cubic above 330 K, tetragonal between 330 K and 160 K, and orthorhombic below 160 K, due to progressive ordering of the methylammonium cations on lowering the temperature.¹⁶ However, unlike the analogous formate perovskites, the low temperature, orthorhombic phase of $[CH_3NH_3]Pb^{II}I_3$ is centrosymmetric and is therefore not a ferroelectric.¹⁷

Aside from the lead halide systems, the most widely studied class of the hybrid perovskites are the formates of general formula $[AmH]M^{II}(HCOO)_3$,²⁰ which show a range of interesting multiferroic behaviour.²¹ Order–disorder transitions associated with hydrogen bonding again (Fig. 5) play a central role in determining the ferroelectric properties of these interesting systems.²² It has been found recently that the formation of amine cation solid solutions of the type $[AmH_{1-x}Am'H_x]]M^{II}(HCOO)_3$, can be used to tune the ferroelectric phase transition, though there are outstanding questions concerning the distribution of the two amines within individual grains and crystals.²³

2.3 Mixed valence and cation vacancies in metal–organic frameworks

Classical mixed valence behaviour is rare in MOFs, and those cases that are known tend to have rational compositions where valence ordering can be achieved. The perovskite-related metal formate composition [NH₂(CH₃)₂]Fe^{II}Fe^{III}(HCOO)₆ is typical and has interesting magnetic properties due to the charge ordering of Fe(II) and Fe(III).²⁴ Similarly, it appears to be difficult to obtain non-stoichiometric MOFs where charge balance might be achieved by forming simple cation vacancies (vac). Thus, while mixed valence $Fe_{1-x}vac_xO$, which adopts the NaCl structure, is well known and achieves charge balance by combining cation vacancies with Fe(II) and Fe(III), there are no well-documented examples of analogous MOFs, e.g. $Fe_{1-x}vac_xL^{2-}$. Nor does it seem possible to create cation vacancies by partial replacement of, say, a monovalent cation by a divalent one to form frameworks of the form $Li_{2-x}Mg_{x/2}vac_{x/2}L$. This is unfortunate because strategies such as this are used in lithium solid electrolytes to enhance cation mobility by creating vacancies into which the Li⁺ ions can move.

3. Ligand defects in metal–organic frameworks

The replacement of one organic ligand by another in MOFs is often more challenging than cation substitution because it is generally more difficult to match the coordinative and spatial requirements of the substituted ligand than it is to match a simple cation. In this respect, porous MOFs are likely to be more accommodating towards ligand substitution and there has been extensive work in this area. Dense MOFs, on the other hand, have less space to accommodate differences in size.

3.1 Examples with anion or ligand substitution in porous MOFs

As with cation substitutions, partial anion or ligand substitution can be carried out during the initial synthesis or postsynthetically. We shall focus first on the latter category, and in particular on reactions that do not involve the replacement of the main linkers. Not surprisingly, it can be straightforward to replace ligands that are pendant rather than bridging. Thus, early work on layered phosphonates showed that it was possible to replace up to 60% of the phenylphosphonate ligands by phosphite without changing the layer spacing but nevertheless modulating the intercalation chemistry.²⁵ More recently, work on HKUST-1, a MOF that is formed from copper(II) paddlewheel dimers linked by 1,3,5-benzenetricarboxylate ligands, showed that the axial water molecules on the copper dimers could be replaced post-synthetically by pyridine, while the pyridine form could not be obtained by direct reaction.²⁶ The removal of water ligands to create under-coordinated cation centers can be particularly advantageous for binding dihydrogen in hydrogen storage applications.²⁷

Some of the first systematic studies of ligand functionalization in MOFs involved work by Cohen on post-synthetic modification (PSM) of 1,4-benzenedicarboxylate linkers in a series of common MOFs.^{28,29} The linker itself was not replaced, but aromatic substituents such as $-NH_2$ were exchanged for $-CONH_2$ by means of PSM (Fig. 6). Alternatively, additional functionality can be added to the aromatic linkers by introducing other reactants into an activated MOF. For example, Cr(CO)₆ will react with benzene-containing linkers to attach Cr(CO)₃ groups which can then undergo further reactions and create gas binding sites by decarbonylation.³⁰

In a recent and very interesting variation on the above theme, the complete organic linker was exchanged in a process termed "Solvent Assisted Linker Exchange" (SALE).³¹ It was applied to the zeolitic imidazolate framework, ZIF-8 $[Zn(C_4H_5N_2)_2]$. The use of SALE enabled the exchange of the organic linker (2-methylimidazolate) for a smaller one (imidazolate, $C_3H_3N_2$). The resultant framework, SALEM-2, possessed very different sorption properties due to the opening of the pore-connecting apertures within the framework. Further functionalization using *n*-butyl lithium created active sites for Brønsted base catalysis that could not be achieved in the parent ZIF-8 (Fig. 7). Here again, as in transmetalation, postsynthetic linker substitution allows the synthesis of novel MOF



Fig. 6 A schematic representation of post synthetic modification in porous MOFs. Reproduced from ref. 29 with permission from The Royal Society of Chemistry.



Fig. 7 Conversion of ZIF-8 to SALEM-2, followed by its functionalization with *n*-butyl lithium. Reprinted with permission from O. Karagiaridi, *et al., J. Am. Chem. Soc.,* 2012, **134**, 18790. Copyright 2012 American Chemical Society.

materials with desirable function-property combinations, which could not be attained through direct solvothermal synthesis. Indeed, it has been shown, by computational studies on the family of zeolitic imidazolate framework (ZIF) materials, that the thermodynamic stability of various polymorphs does not uniquely determine their experimental feasibility and accessibility through solvothermal synthesis.^{32,33}

While the above examples have focused on cases where the modification of the MOF is done post-synthesis, there has been extensive work on systems where mixed ligands are incorporated during the initial synthesis (eqn (2)). One of the most detailed studies concerns the substitution of multiple combinations of substituted benzene dicarboxylate (BDC) – based ligands into the MOF-5 structure by Yaghi and co-workers.³⁴ The scope is illustrated in Fig. 8, which shows some of the many combinations of these so-called multivariate MOFs (in one case, as many as 8 different substituted linkers were introduced). As with other strategies for ligand substitution, one of the objectives is to create novel functionalities within a known structure type. In this particular work, it was possible to substantially enhance the CO_2 adsorption capacity by appropriate choice of the substituents on the 1,4-BDC linkers.

Work by Kitagawa and others focused specifically on flexible frameworks, and endeavored to tune the flexibility in order to optimize sorption performance.^{35,36} In one example, two doubly interdigitated frameworks - [Zn(5-NO₂-ip)(bpy) (0.5DMF · 0.5MeOH)] [Zn(5-MeO-ip)(bpy)](0.5DMFand 0.5MeOH), where ip = 5-nitroisophthalate and bpy = 4,4'-bipyridyl were prepared, giving two closely related structures with similar lattice parameters. However, the guest-free structures were very different after desolvation. The former shows a porous to non-porous transition on degassing (and gateopening on adsorption), whereas the latter remains in the open form. Ligand-based solid solutions were then prepared and the two substituents appeared to be homogeneously distributed in the as-synthesized samples. The adsorption properties of the degassed samples were then explored and it was found that the gate-opening transition could be fine-tuned by adjusting the composition of the solid solutions, giving rise to enhanced performance for the adsorption of CO₂ from CO₂/ CH₄ mixtures.



Fig. 8 Various permutations of 1,4-BDC derivatives in the MOF-5 structure type. From H. X. Deng, et al., Science, 2010, 327, 846. Reprinted with permission from AAAS.



Fig. 9 (a) Ternary phase diagram shows solid solutions from lithium succinate and related phases. (b) The succinate-tetrafluorsuccinate system obeys Vegard's Law throughout its composition range. Reprinted with permission from H. H. M. Yeung, *et al., Angew. Chem., Int. Ed.,* 2013, **52**, 5544. Copyright 2013. John Wiley and Sons.

3.2 Examples of anion substitution in dense MOFs

Recent work on lithium succinates and related systems³⁷ illustrate what can be achieved in this area. The three dense MOFs – lithium succinate, lithium hydroxysuccinate (*i.e.* malate) and lithium methylsuccinate – turn out to be isostructural, in spite of the different sizes of their ligands. There is no solvent-accessible volume, but the different ligands can still be accommodated; indeed, the Young's modulus of the succinate is only 20 GPa, which is comparable to many porous MOFs. Mixed samples were synthesized by mechanical milling in order to avoid the problems with differential crystallization rates, and complete solid solution was found throughout the ternary phase diagram (Fig. 9a). This was confirmed Le Bail refinement of the 1/3, 1/3, 1/3 composition using high resolution synchrotron X-ray diffraction.

Surprisingly, even the tetrafluorinated succinate ligand can be incorporated into the succinate structure and a complete solid solution can be obtained (Fig. 9b). A 50:50 sample in the lithium succinate-tetrafluorosuccinate solid solution was analyzed in greater detail in order to confirm that the two ligands are homogeneously distributed. Both full Rietveld refinement with synchrotron PXRD data (Fig. 9b), infrared analysis, and spin diffusion solid-state NMR measurements confirmed that the two ligands were well mixed at the molecular level.³⁷

Interestingly, the flexibility of lithium succinate and related phases provides a rare example of solid solution formation by post-synthesis modification in a dense MOF. Thus, in the specific case of lithium hydroxysuccinate (*i.e.* malate), it is possible to perform a topotactic dehydration across the C–C bond to form the corresponding fumarate phase.³⁸ In fact, 80% of the water can be removed from a crystal without loss of crystallinity, and the process has been followed by *in situ* single crystal X-ray diffractometry. Density functional theory (DFT) calculations show that the fumarate thus formed is a metastable phase, that is 12.7 kJ mol⁻¹ less stable than the normal form.

3.3 Other types of anion defects in MOFs

The above sections have dealt with conventional ligand defects where one ligand, typically a bridging anion, is replaced by another one. There are, however, other possibilities, such as replacing a bridging ligand by two monodentate ones, and this type of defect has been incorporated into porous, pillared diphosphonates, including the widely studied zirconium diphosphonates.³⁹ For example, in the case of $Al_2[O_3PC_2H_4PO_3](H_2O)_2F_2$, it has proved possible to replace some of the ethylene diphophonic acid pillars by two phosphite groups to form Al₂[(O₃PC₂H₄PO₃)_{1-x}(HPO₃)_{2x}](H₂O)₂F₂ solid solutions. Up to 32% of the pillars can be replaced and the distribution of the defects appears to be homogeneous. Such tuning enables the adsorption and desorption properties of the MOF to be modulated.⁴⁰

The same strategy has been used to replace a dicarboxylate such as 1,4-BDC by monocarboxylate anions, thus removing the bridge between the cations and creating two dangling ligands.^{41,42} This has been reported for UiO-66(Zr or Hf), a MOF that contains octahedral zirconium (or hafnium) clusters that are linked by 1,4-BDC ligands to form a highly porous structure. It has recently been shown that pairs of trifluoroacetate (TFA) groups can replace individual 1,4-BDC linkers and that such substitution results in enhanced Lewis acidity for catalytic reactions.⁴³ In an interesting extension of this theme, it has also been demonstrated that the removal of linkers by such substitution can enhance the mechanical stability of UiO-66, as shown by the improved resistance to amorphization during milling.44 This rather counter-intuitive behaviour is ascribed to the strengthening of the zirconium-carboxylate bonds due to the local electron withdrawing effect of the TFA.

4. Domain structures in MOFs

To a first approximation, the various defects described above are generally thought to be randomly-distributed throughout a given framework. In conventional inorganic solids, such a picture is only sometimes true. For example, Schottky defects do not correlate strongly in NaCl but can be so robustly ordered in transition metal oxides that in the case of NbO they even break the lattice symmetry.45 Understanding the extent to which defects are correlated in MOFs may prove important for a number of reasons. Wherever defects are associated with transport properties (e.g. acid sites for proton conductivity, or vacancy sites for gas storage), percolation will depend critically on the existence and nature of correlated defect positioning. Moreover, the dynamical and elastic properties of MOFs are also sensitive to the presence of domain structures, which influence directly the underlying phonon dispersion. And, although a significant challenge, the controlled arrangement of multiple defects might eventually enable design of protein-like "active sites" for catalyticallyactive MOFs.46

Perhaps the clearest indication that MOFs can support correlated defects comes from the structural chemistry of the UiO-66 family.⁴⁷ The parent compound is usually prepared under basic conditions from a mixture of $ZrCl_4$ and H_2bdc , but addition of monocarboxylate "modulators" leads to a defective framework structure in which some of the bdc linkers have been replaced by modulator pairs [Fig. 10].⁴⁸ Varying the modulator chemistry and concentration determines the density of defects introduced. Because these defects must always occur in pairs, it has always been known that there must be at least local correlations between defect sites. However, most of the experimental and computational



Fig. 10 The structure of non-defective UiO-66 is comprised of $[ZrO_4(OH)_4]$ clusters connected by terephthalate linkers. The metal-containing clusters of the resulting framework are arranged on a face-centred cubic lattice. (b) Inclusion during framework synthesis of mono-carboxylate modulators, such as formate as shown here, can lead to correlated linker vacancies where a single terephthlate linker is replaced by two monocarboxylates in an opposing geometry.

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approaches used to characterise these defects are insensitive to the longer-range defect correlations associated with domain formation.48,49 Even detailed single-crystal X-ray diffraction studies have determined only the total defect concentration and the geometry of individual defect sites.⁵⁰ This is because conventional single-crystal analysis is insensitive to long-range correlations if those correlations are not sufficiently strong to force periodic patterning (e.g. of defects). Instead the experimental methods most sensitive to defect correlations are diffuse scattering measurements - X-ray, neutron, or electron; powder or single-crystal and electron microscopy.⁵¹ These are precisely the methods which have informed much of our understanding of collective defect structures in conventional inorganic materials such as zeolites and transition-metal oxides.^{52,53} We note that electron microscopy is the least translatable of these methods to the study of MOFs since so many systems are terminally sensitive to the electron beam.⁵⁴

Structured diffuse scattering characteristic of defect nanodomains was first observed in the Hf-containing analogue UiO-66(Hf), prepared using high concentrations of formic acid modulator. The X-ray diffraction patterns of these samples include a series of diffuse scattering peaks at positions forbidden by the symmetry of the parent UiO-66 structure (space group $F\bar{4}3m$).⁵⁵ That this diffuse scattering is associated with defects rather than solvent ordering as originally suggested is evident in their persistence on thermal desolvation. Intriguingly, defect structure models which consider only linker/ modulator substitution cannot reproduce the observed diffuse scattering intensities. Instead a model in which vacancies at the entire $[Hf_6O_4(OH)_4]$ cluster site are correlated to form defect nanoregions with reduced symmetry (space group $P\bar{4}3m$) quantitatively accounts for the diffuse scattering [Fig. 11].⁵⁶ There is a strong analogy to the Schottky defect structure of the transition-metal monoxides with the cluster vacancy arrangement in defect nanoregions mapping directly onto the Nb vacancy sites in NbO.57 This model was supported by electron diffraction, X-ray pair distribution function and anomalous X-ray diffraction measurements, the latter showing conclusively that defects in UiO-66(Hf) involve modulation in Hf concentrations.⁵⁶

Cluster vacancies are not unique to the Hf-containing UiOs. Surface area measurements for a number of defective Zr-containing UiO-66 variants are consistent only with the existence of [Zr₆O₄(OH)₄] cluster vacancies.⁴¹ Likewise, doping of UiO-66(Zr) with Ce gives rise to a series of defective frameworks whose diffraction patterns contain the same characteristic diffuse scattering peaks observed for UiO-66(Hf).⁵⁸ As in the Hf MOFs, this scattering persists on thermal desolvation suggesting that it also arises from nanodomains of correlated cluster vacancies. An extreme example of correlated defects in this structural family has recently been discovered in a 2-sulfoterephthalate-bridged Zr-containing UiO-66 analogue.59 The structure of this sulfonated MOF is characterised by a $(\sim 40 \text{ Å})^3$ unit cell in which 3/16 of the Zr cluster sites are vacant. The vacancy ordering observed experimentally can be considered a periodic intermediate lying between the defect-free UiO-66



Fig. 11 (a) Reproduced from ref. 56. The X-ray powder diffraction pattern of d-UiO-66(Hf) contains a series of weak diffuse reflections that are forbidden by the symmetry of the parent structure. (b) These reflections are evident in selected area electron diffraction patterns of individual d-UiO-66(Hf) crystallites. (c) The fundamental unit of the domain structure in this material has cluster vacancies at the corner of the unit cell and has primitive cubic symmetry. (d) These primitive cells coexist with regions of the parent face-centred cubic structure, facilitated by the close match in unit cell metric. (d) Because the face-centred cubic lattice can be understood in terms of four interpenetrating primitive cubic lattices, there are four possible orientations for the defect nanodomains. In the bulk d-UiO-66(Hf) structure all four of these occur (here shown using different colours). This model of the domain structure was used to calculate the diffraction pattern shown in panel (a).

parent and the $P\bar{4}3m$ domain structure of UiO-66(Hf), where the cluster vacancy fraction is 1/4 [Fig. 12].

Already there are indications that these correlated defect structures in UiO frameworks result in interesting physical and chemical properties. In Ce-doped UiO-66(Zr), linker vacancies are coupled to $Ce^{4+/3+}$ reduction, which in turn promotes catalytic performance in methanol oxidation.⁵⁸ A similar conclusion was reached in ref. 41, where defect sites promoted uptake of catalytically-active Cr^{3+} centers during post-synthetic modification. The fractal-like network of exposed defect sites in the sulfonated MOF of ref. 59 is thought to be responsible for its extremely high proton conductivity. And, finally, the concentration of correlated cluster vacancies in defective UiO-66(Hf) samples was shown found to influence their thermal expansion behaviour, which can include the desirable property of "colossal" negative thermal expansion.⁶⁰

It is not yet clear whether strongly-correlated defects are unique to the UiO family of MOFs. Certainly the strong X-ray scattering contrast of absent vs. present $[M_6O_4(OH)_4]$ clusters means that defect correlations have a clearer experimental sig-



Fig. 12 (a) The face-centred cubic structure of defect-free UiO-66. (b) The 2-sulfoterephthalate UiO-66 derivative reported in ref. 59 has a bodycentred cubic structure with a lattice parameter twice that of the parent structure shown in (a). Cluster vacancies order at the edges of this enlarged unit cell. (c) The primitive cubic structure of the defect domains in d-UiO-66(Hf), which is related to the cell shown in (b) on removal of the clusters at the corners of that unit cell (shown here in blue).

nature for this system than is likely for other defective MOF families. Nevertheless there are at least three reasons why the UiO family might be particularly prone to defect inclusion and then to the existence of correlations between those defects to form domain structures. First, the parent network has a very high connectivity, which means that linker/cluster vacancies can be incorporated while maintaining mechanical stability. Second, the unit cell parameters calculated for defect-free and defect structures differ by only 0.05%, and so defects can be incorporated without significant lattice strains.⁵⁹ Third, the core $[M_6O_4(OH)_4(OR)_{12}]$ building unit is susceptible to symmetry-lowering distortions that may play a role in driving defect correlations. The idealised $T_{\rm d}$ point symmetry of the cluster lowers spontaneously to D_{2d} (\equiv tetragonal $\bar{4}2m$) in the crystal structures of non-bridging cluster packings (e.g. M = Hf, $R = CH_3$; ref. 56 and 61) in a way that is reminiscent both of symmetry lowering in the transition-metal acetate paddlewheels and indeed of Jahn Teller effects in general [Fig. 13].⁶² This susceptibility to symmetry lowering suggests that modulator substitution at the four equatorial sites of a $[M_6O_4(OH)_4(OR)_{12}]$ cluster during MOF growth allows local



Fig. 13 (a) Representation of the Zr_6 cluster geometry found in the zirconium formate compound $Zr_6O_4(OH)_4(HCOO)_8$.⁶¹ The equatorial oxygen atoms belong to dangling formate anions, and are disordered over closely-related sites. The two crystallographically-distinct Zr atoms are shown in different shades of green. (b) The central Zr_6 octahedron geometry in this cluster, showing the spontaneous symmetry-lowering of Zr…Zr distances.

relaxation. Then, because this local distortion can only be propagated on the NbO-type defect lattice,⁶³ the experimentallyobserved domain structure emerges naturally.

Our understanding of the microscopic driving forces responsible for defect correlations in MOFs would be improved enormously by an increased diversity of experimentally-characterised systems for which domain structures occur. The increased application of diffuse scattering and electron microscopy methods to the study of MOFs will be critical in this sense. Indeed, experience suggests that the presence of structured diffuse scattering in MOF single crystal diffraction studies is probably more widespread than has been reported in the literature. A recent study of the relationship between correlated defect structures relevant to MOF networks and the corresponding diffuse scattering patterns⁶⁴ demonstrates that the signature of domain structures in the parent Bragg reflections can often be subtle, but the existence of characteristic diffuse scattering may prove invaluable in diagnosing correlated disorder of the type discussed here. A number of known MOF structures, such as the intriguing example of "partial interpenetration" in NOTT-202,65 may benefit from reanalysis in this context.

5. Amorphous solids, liquids and melt-quenched glasses

Arguably the clearest examples of disordered materials, those which lack any long range order (*i.e.* those which are amorphous) are easily identifiable from the absence of Bragg peaks in their X-ray diffraction patterns. In the context of MOFs and coordination polymers, such structures can, like cation substitution in solid solutions, be formed through (i) alteration of initial synthetic conditions, or (ii) through post synthetic introduction of disorder.

5.1 Metal-organic frameworks

Fast crystallization kinetics lie behind the propensity of MOF precursors to self-assemble into highly ordered networks during synthesis.⁶⁶ Highly disordered products, such as amor-



Fig. 14 Dimensionality of coordination polymers and MOFs known to undergo structural melting. (a) $[Zn(HPO_4)_2]\cdot 2C_3H_5N_2$. (b) $[Zn(H_2PO_4)_2(HTr)_2]_n$.⁷² (c) $[Zn(C_3H_3N_2)_2]$. Adapted with permission from D. Umeyama, S. Horike, M. Inukai, T. Itakura, S. Kitagawa, J. Am. Chem. Soc., 2015, **137**, 864, and D. Umeyama, S. Horike, M. Inukai, T. Itakura, S. Kitagawa, J. Am. Chem. Soc., 2012, **134**, 12780. Copyright 2015 and 2012 American Chemical Society respectively.

phous solids, are hence difficult to form directly, though differential formation of crystalline and amorphous states of a zeolite-related MOF of composition $Zn(C_4H_3N_2O)_2$ can be effected by altering the rate of addition of a structure directing agent to the initial metal and organic containing solution.⁶⁷ More commonly, heating or mechanical impact are used to irreversibly introduce disorder into crystalline hybrid materials.

The application of mechanosynthesis to crystalline, nonsolvated zinc imidazolate frameworks (ZIFs) results in formation of stable amorphous network arrangements.⁶⁸ However, a recent example on the in situ ball-milling of ZIF-8 in an aqueous acidic environment,69 found not only the expected amorphization, but subsequent recrystallization of the amorphous network upon continued milling, to a previously unidentified topology. The metastable nature of some amorphous MOF phases is also evidenced in the crystallineamorphous-crystalline transition sequence upon heating of ZIF-4,⁷⁰ a three-dimensional framework of $Zn(C_3H_3N_2)$ composition formed of tetrahedral Zn^{2+} nodes connected by bidentate imidazolate ligands. Here, after formation of an amorphous solid at 300 °C by heating of the crystalline phase, recrystallization to the dense 'ZIF-zni' MOF is observed at 450 °C. In this case, knowledge of the chemical composition of the amorphous phase (the transition sequence not being accompanied by mass loss), allowed total scattering data on the amorphous phase to be modeled by Reverse Monte-Carlo techniques.70

Furthermore, whilst MOFs are usually observed to decompose to oxides at high temperatures, the dense ZIF-zni was observed to undergo melting at 580 °C;⁷¹ the first example of the melting of a three-dimensional MOF. This observation was rationalized in terms of the accessibility of the intrinsic melting point of ZIF-zni, made possible by the raising of the point of thermal decomposition to over 600 $^{\circ}\mathrm{C},$ by excluding air from the heating process.

Cooling of the liquid back to room temperature yielded a glass, which contained both tetrahedral Zn^{2+} centers and Zn–Zn pairs separated by bidentate ligands, reminiscent of the crystalline phase (Fig. 14). The different network connectivity of crystal and glass phases suggested a reconstructive transition during melting and cooling, due to Zn–N dissociation in the disordering mechanism. Such dissociation would, in turn, infer ionic liquid formation upon melting, though no investigations into this phase were performed. An alternative explanation, analogous to that used to explain the melting (at much higher temperatures) of inorganic zeolites, involves dihedral angle changes.

5.2 1D and 2D coordination polymers

Lower dimensionality hybrid structures have also been observed to melt. The one dimensional coordination polymer, [Zn(HPO₄)(H₂PO₄)₂]·2C₃H₅N₂, consists of chains of four-fold coordinated Zn²⁺ centers linked by phosphate ligands, along with charge-compensating protonated imidazole species. At the same time as the example presented above, work on this system revealed a solid-liquid transition at 150 °C, in a process set off by the excessive disordering and eventual dissociation of one Zn–O bond in the Zn²⁺ coordination sphere.⁷³ Cooling led to glass formation, with dynamic mechanical analysis, ³¹P nuclear magnetic resonance spectroscopy, pair distribution function (PDF) and X-ray absorption spectroscopy studies (XAS) on the crystalline and glass solids confirming the absence of Zn-O association in the liquid state. Furthermore, the glass itself was deduced to have a one-dimensional structure, like that of the crystal. Work on another 1D polymer incorporating octahedral Zn^{2+} centers, $[Zn_3(H_2PO_4)_6(H_2O)_3]$.

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 $C_7H_7N_2$, also led to liquid formation, though at a lower temperature of 100 °C. In this case however, significantly faster recrystallization rates of the resultant ionic liquid prevented glass formation at room temperature.

In possessing similar immediate inorganic coordination environments in both crystal and glasses, the above systems share common features with other glass formers, including zeolites.⁷⁴ A system of intermediate two dimensional connectivity, $[Zn(H_2PO_4)_2(HTr)_2]_n$,⁷⁵ which melts at 184 °C, does however undergo network connectivity changes on the solidglass transformation. In the structure, each organic ligand equatorially bridges Zn^{2+} ions to form layers, with an octahedral coordination environment completed by non-bridging phosphate ions in the axial positions. The process again involved bond dissociation, however, unusually, reformation of an extended network was not observed upon glass formation. PDF and XAS studies were used to derive a molecular model for the end glass state, a transition driven by the preference of Zn^{2+} ions for a 4-fold coordination environment.

6. Concluding remarks

The concepts of defects and disorder are inextricably linked with one another. Point vacancies, substitutional defects and long-range disorder may all be introduced into MOFs pre-, or post synthetically. Whilst at present still system dependent, notable studies of changing both the physical and chemical behaviour of known frameworks using the methodologies presented here exist. Such work hints that perhaps we might make equal, if not greater, advances in the field by looking to change the properties of existing structures in a controlled manner, as opposed to synthesizing new structures and characterizing their properties.

Conflict of interest

The authors declare no competing financial interests.

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