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Control of Metal–Organic Framework Crystallization by Metastable Intermediate Pre-equilibrium Species†

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† Electronic Supplementary Information (ESI) available: detailed experimental methods, in situ XRD data, Pawley refinement results, Gualtieri and Sharp-Hancock plots, in situ pH and turbidity profile fits, rate constants, and SEM images.

Metal–organic frameworks (MOFs) show exceptional promise for gas separation,1 sensing,2 catalysis,3 and other applications, owing to their modular, porous architecture based on metal nodes and organic linkers.4,5 The processes of MOF nucleation and crystal growth are fundamentally important to particle size and shape,6 yield and efficiency in industrial scale-up,7 and phase selection.8 Yet, despite increasing awareness, the crystallization of MOFs is poorly understood.9,10 ZIF-8, [Zn(mIm)2] (mImH = 2-methylimidazole) shows potential for carbon capture, propene/propane separations and dielectric applications11 and is one of the few MOFs that have been investigated in detail as a model system for MOF crystallization.12–19 ZIF-8 nanoparticles were first synthesised by Cravillon et al. in 2009, using excess mImH to deprotonate linkers and terminate particle surfaces.12 Static light scattering revealed slow, continuous nucleation and fast crystal growth;14 in situ X-ray diffraction (XRD) suggested that the crystallization process is phase-boundary-limited,20,21 as evidenced by the Avrami exponent, n,22–24 taking values of n ≈ 1.25 However, other XRD measurements reported values ranging from n = 0.6 (diffusion-limited) in methanol:aqueous ammonia,26 to n = 1 in heated DMF27 and n = 4 (classical, 3-D monomer attachment) in pure methanol.28 Small-angle X-ray scattering (SAXS) showed that clusters approximately 1 nm in diameter coalesce to form intermediate amorphous particles,15 and electrospray ionisation mass spectrometry (ESI-MS) indicated that several different clusters co-exist in the reaction solution.16 However, in situ atomic force microscopy (AFM) revealed that the key species at the crystal surface are individual mImH linker and solvated Zn2+ ions.17 On the other hand, a recent total scattering investigation suggested that the solution state is dominated by [Zn(mImH)4]–like species.19 The apparent disparity between these observations indicates that we are far from understanding the chemistry of MOF nucleation and growth, even in such an ostensibly simple system as ZIF-8.

We report a tandem in situ XRD and pH investigation into ZIF-8 crystallization, in which we show that, under certain conditions, the rate of crystallization decreases with increasing reactant concentration. In order to explain this counterintuitive observation, we propose a model for ZIF-8 crystallization that involves a pre-equilibrium of metastable intermediate clusters with various Zn:mIm ratios and different degrees of protonation. The rate of crystallization depends on the position of equilibrium: owing to favourable Zn–mIm coordination, crystallization from over-stoichiometric clusters is dissociative and slower than crystallization from under-stoichiometric clusters. By considering how the equilibrium is perturbed by changes in crystal size and structure in ZIF-8.

In situ XRD29–33 data were collected at Beamline I12, Diamond synchrotron,34 on a series of room temperature reactions in methanol, in which the mImH:Zn ratio was kept constant at 4:1 and the Zn2+ concentration, [Zn2+]0, was varied from 0.02 M to 0.10 M in 0.02 M steps. Under these conditions, the crystallization rate was observed to decrease with increasing concentration (Fig. 1a, b). Integrated XRD peak intensities were extracted by Pawley refinements (see Sections S1 and S2 in the Supporting Information) and their changes were fitted to the expression of Gualtieri35 to give the rates of nucleation, kN, from 5.57(7) × 10−3 s−1 to 7.94(4) × 10−4 s−1 ([Zn2+]0 = 0.02 M to 0.10 M) and growth, kG, from 2.1(4) × 10−3 s−1 to 2.67(9) × 10−3 s−1, confirming that both nucleation and crystal growth are inhibited as concentration...
increases (Fig. 1c). The single exponent, \( n \), given by the Gualtieri expression\(^{35} \) ranges from 0.47(4) to 1.03(6), without an obvious trend with concentration. However, analysis by the Sharp-Hancock method\(^{31} \) shows that \( n \) changes with time, from \( n \approx 1 \) in the initial stages to \( n < 1 \) in the later stages, suggesting a progression from phase boundary conditions to diffusion-limited crystallization (see Fig. S28).

A decrease in the crystallization rate with increasing concentration cannot be understood by a linear mechanism of MOF crystallization according to the stoichiometric equation, \( \text{Zn}^{2+} + 2\text{mImH} \rightarrow [\text{Zn(mIm)}_2]_2 + 2\text{H}^+ \). We propose that crystallization is instead preceded by a pre-equilibrium of metastable intermediate coordination clusters with compositions \( [\text{Zn(mIm)}_x(\text{mImH})_y(S)]^{2-x+y+} \), where \( S = \text{coordinating solvent} \) (Fig. 2). Although purely solvated \( \text{Zn}^{2+} \) is six-coordinate, tetrahedral geometries are preferred for linker-coordinated clusters, i.e. \( x + y + z = 4 \).\(^{36} \) Each cluster exists in dynamic equilibrium with a counterpart of higher \( \text{mImH:Zn} \) ratio via solvent-exchange reactions, with equilibrium constant \( K_c \) and rate constants \( k_b \) and \( k_d \) for the forward and reverse reactions, respectively (Eq. 1). A second set of equilibria exists between each species and its deprotonated counterpart, with equilibrium constant \( K_d \) and rate constants \( k_d \) and \( k_d \) (Eq. 2). For simplicity, we assume that all coordination equilibria have the same constants, deprotonation likewise, \( S = \text{in large excess} \) (see Section S1.5). Following pre-equilibrium, the coordination clusters condense to give the stoichiometric crystalline product, \( [\text{Zn(mIm)}_2]_2 \). Although condensation might in principle occur between any neutral monomer with a deprotonated \( \text{mIm} \) ligand (i.e. \( x \geq 1 \)) and one with a site not coordinated by \( \text{mIm} (z \geq 1) \), in order to define a minimally-parameterized model that qualitatively captures the experimental trends, we use only the simplest neutral monomer species with the best condensation statistics (\( x = z = 2 \)), \( [\text{Zn(mIm)}_2(\text{S})_2]_2 \). Condensation is described by self-nucleation (Eq. 3) and autocatalytic growth (Eq. 4), with rate constants \( k_1 \) and \( k_2 \), respectively.\(^{38} \) We find good qualitative agreement with experiments using values of \( k_1 = 100 \text{ s}^{-1} \), \( k_2 = 10^6 \text{ s}^{-1} \), \( k_3 = 330 \text{ M}^{-1} \text{s}^{-1} \) \((k_c = 10^{-1} \text{ M}^{-1} \text{s}^{-1}) \) and \( k_d = 2800 \text{ M}^{-1} \text{s}^{-1} \) \((k_d = 10^6 \text{ M}^{-1} \text{s}^{-1}) \).

\[
[\text{Zn(mIm)}_x(\text{mImH})_y(S)]^{2-x+y+}_z + \text{mImH} \xrightarrow{k_{d}} [\text{Zn(mIm)}_x(\text{mImH})_y(S)]^{2-x+y+}_z + \text{S}, \quad K_c = \frac{k_c}{k_d}
\]

\[
[\text{Zn(mIm)}_x(\text{mImH})_y(S)]^{2-x+y+}_z \xrightarrow{k_{d}} [\text{Zn(mIm)}_x(\text{mImH})_y(S)]^{2-x+y+}_z + \text{S}, \quad K_d = \frac{k_d}{k_d}
\]

\[
[\text{Zn(mIm)}_2(\text{S})_2] \xrightarrow{k_1} [\text{Zn(mIm)}_2] + 2\text{S}
\]

\[
[\text{Zn(mIm)}_2(\text{S})_2] \xrightarrow{k_2} 2[\text{Zn(mIm)}_2] + 2\text{S}
\]

The important consequence of this model is that the crystallization rate and mechanism do not depend directly on the concentration of reagents but on the position of pre-equilibrium: when under-stoichiometric species (\( x + y < 2 \)) dominate, crystallization is associative and fast, whereas when over-stoichiometric species dominate, crystallization is limited by the rate of \( \text{mImH} \) dissociation and is therefore slower. Simulations in COPASI\(^{39} \) show that before pre-equilibrium is attained, under-coordinated species prevail (Fig. 3a), resulting in associative, nucleation-dominated crystallization (Fig. 3b). However, once pre-equilibrium is reached, clusters become over-coordinated, resulting in a reduction in nucleation relative to growth. At higher concentration, nucleation is initially faster (Fig. 3d) but pre-equilibrium is achieved faster (Fig. 3b). However, once pre-equilibrium is reached, clusters become over-coordinated, resulting in a reduction in nucleation relative to growth. At higher concentration, nucleation is initially faster (Fig. 3d) but pre-equilibrium is achieved faster (Fig. 3d) and is pushed further towards over-coordinated species (Fig. 3c), leading to a stronger suppression of crystallization overall (Fig. 3e). Sharp-Hancock analysis suggests that changes in \( n \) occur upon changes in the crystallization route (i.e. nucleation vs. growth), relative concentrations of intermediates and \( p\text{H} \) (see Fig. S29).

We test our model by monitoring \( p\text{H} \), which allows us to estimate the concentrations of protonated and neutral \( \text{mImH} \) species (see Sections S1.4 and S1.5 for details). Linear crystallization would give continuous release of \( \text{mImH}_2^+ \) and an exponential decay in \( p\text{H} \). However, we see a stepwise \( p\text{H} \) profile in the experimental and simulated data (Fig 4). A rapid decrease from \( p\text{H} 9–10 \) to \( 6–7 \) occurs upon mixing within the time resolution of our experiment (10 s). In the pre-equilibrium model this corresponds to a decrease in the concentration of neutral \( \text{mImH} \) species \((pK_a = 14.2 \text{ in water}) \)\(^{40} \) owing to rapid formation of \( \text{Zn(mImH)}_y \) coordination complexes \((pK_a = 10.3 \text{ in water}) \)\(^{41} \) and condensation of a few oligomers, which releases \( \text{mImH}_2^+ \). As pre-equilibrium is established, the rate of condensation decreases, leading to a plateau in \( p\text{H} \). Then autocatalytic crystal growth accelerates and is accompanied by the release of further \( \text{mImH}_2^+ \). The resulting second decrease in \( p\text{H} \) exhibits the same timescale as crystal growth observed by \textit{in situ} XRD (see Fig. S33), which suggests that condensation and crystallization processes are linked. If the rate of crystal growth is related to diffusion of \( \text{mImH}_2^+ \), this would be consistent with a value of \( n < 1 \) found towards the end of the reactions. Evidence that the pre-equilibrium exists in solution, rather than in amorphous particles, comes from \textit{in situ} measurements of turbidity, which display timescale and concentration dependence similar to nucleation, as measured by XRD (see Fig. S33), i.e. condensation to form particles occurs during the second \( p\text{H} \) decrease and not before. \textit{Ex situ} SEM shows that particle diameter increases with increasing concentration (Fig. 5), somewhat in contrast to the expectation of classical, linear crystallization. The pre-equilibrium model explains why this is the case: with increasing concentration, the number of nuclei produced remains constant but, by virtue of increased reactant availability, the average particle size increases (Fig. 5, inset).

Although the simplicity of the condensation process in our model precludes direct fitting to what are undoubtedly very complex crystalization curves, which arise from e.g. surface area and diffusion effects, we emphasise here that the trends in our data are qualitatively well captured. The pre-equilibrium model also helps to rationalise the diversity of previous observations. Highly basic solutions, e.g. a large excess of \( \text{mImH} \) or \( \text{NH}_3 \)\(^{12,14–16,26} \) promote deprotonation and a fast, associative route. Low values of \( n \) reflect the diffusion-limit of such rapid crystallization. Increasing concentration favours over-coordinated species; the model correctly predicts the dominance of \( [\text{Zn(mImH)}_4]_1 \)-like species.
in such crystallizations. The use of water in the same study also slows crystallization, via faster coordination, which allows pre-equilibrium conditions to be more rapidly reached: solvent:ligand exchange at labile ions, such as Zn$^{2+}$, is limited by solvent dissociation; water exchange is at least an order of magnitude faster than methanol. Finally, in extremely dilute, mildly acidic conditions, the pre-equilibrium favours low coordination numbers and even isolated mlmH and [Zn(S$_2$C$_2$O$_4$)$_2$]$^{3-}$ species (see Fig. 3a at extended times), resulting in extremely slow crystallization and phase-boundary conditions.

The qualitative importance of pH in the thermodynamics and kinetics of MOF formation is already widely known. For example, the synthesis of many MOFs requires decomposition of DMF to release dimethylamine, which acts as a pH modifier to induce nucleation. On the other hand, absolute concentration has rarely been used as a synthetic handle to control MOF formation. These results show that, somewhat counterintuitively, increasing concentration can inhibit nucleation by promoting over-coordination, and lead to changes in the growth mechanism. Whilst further verification of the pre-equilibrium model could be demonstrated by performing in situ ESI-MS, SAXS or TEM, our results provide a new framework in which to interpret MOF syntheses, predict the effects of tuning different reaction parameters, and relate them to the solution chemistry of the metal and linkers. For instance, pH control at high concentrations represents an obvious means to inhibit nucleation of metastable precursor solutions, prior to seeding or addition of base, in order to control particle size or film thickness. In addition, by diversifying the composition of pre-equilibrium species, e.g. via dynamic combinatorial chemistry, we may be able to access new phases with specific compositions, local structures, or correlated disorder.

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References

**Fig. 1** *In situ* XRD of ZIF-8 formation, showing a) variation of the 110 peak as a function of time at different concentrations; b) an extended view for \([\text{Zn}^{2+}]_0 = 0.10 \text{ M}\); and c) rate constants for crystal nucleation \((k_N, \text{purple closed diamonds})\) and growth \((k_G, \text{cyan open diamonds})\), calculated from Gualtieri fits to the changes in integrated XRD intensity, \(\alpha\), at five concentrations from \([\text{Zn}^{2+}]_0 = 0.02 \text{ M}\) to 0.10 M (inset).
Fig. 2 Pre-equilibria in ZIF-8 crystallization resulting from coordination (right-to-left) and deprotonation (top-to-bottom), described by constants $K_c$ and $K_d$, respectively, and self-nucleation and autocatalytic growth rates $k_1$ and $k_2$, respectively.
Fig. 3 Simulated evolution of pre-equilibrium intermediate monomer species with formula $[\text{Zn(mImH)}_y (\text{S})_{2-y}]^{(2-y)^+}$ (a, c) and flux of self-nucleation (labelled N) and autocatalytic growth (labelled G) reactions (b, d) for reactions with initial Zn concentrations of 0.02 M and 0.10 M. Regions shaded in grey mark periods during which pre-equilibrium is attained. Extent of crystallization, $\alpha$, is shown as a function of time and concentration (e; inset shows the same data on a linear time axis c.f. Fig. 1c).
**Fig. 4** Evolution of pH as a function of time and concentration, showing the rapid decrease that accompanies initial condensation events upon mixing, the pre-equilibrium plateau, and the second decrease that accompanies crystal growth. Curves are offset for clarity and bounded by standard deviations calculated from three independent measurements. Inset shows pH evolution calculated from the pre-equilibrium model.

**Fig. 5** Variation of particle diameter measured by SEM, $d$, with initial Zn$^{2+}$ concentration, [Zn$^{2+}$]$_0$. Scale bar = 2 µm; open circles show statistics from images judged to be non-representative of the bulk sample. Inset shows nuclei concentration, $\varepsilon$ (closed grey circles), and number of monomers per particle, $\nu$ (open grey circles), as a function of [Zn$^{2+}$]$_0$, calculated from the pre-equilibrium model.