Using Supercritical CO₂ in the Preparation of Metal-Organic Frameworks: Investigating Effects on Crystallisation

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Abstract: In this report, we explore the use of supercritical CO₂ (scCO₂) in the synthesis of well-known metal-organic frameworks (MOFs) including Zn-MOF-74 and UiO-66, as well as on the preparation of [Cu₃(OH-mBDC)₃]ₙ metal-organic polyhedra (MOPs) and two new MOF structures \([\text{Zn}_n\text{L}](\text{DPE})\text{-}4\text{H}_2\text{O})_n\) and \([\text{Zn}_n\text{L}(4,4’\text{-azopy})]\text{-}7.5\text{H}_2\text{O})_n\), where BTC = benzene-1,3,5-tricarboxylate, BDC = benzene-1,4-dicarboxylate, L₁ = 4-carboxy-phenylene-methyleneaminobenzoate, DPE = 1,2-di(4-pyridyl)ethylene, 4,4’-azopy = 4,4’ azopyridine, and compare the results versus traditional solvothermal preparations at low temperatures (i.e., 40 °C). The objective of the work was to see if the same or different products would result from the scCO₂ route versus the solvothermal method. We were interested to see which method produced the highest yield, the cleanest product and what types of morphology resulted. While there was no evidence of additional meso- or macroporosity in these MOFs/MOPs nor any significant improvements in product yields through the addition of scCO₂ to these systems, it was shown that the use of scCO₂ can have an effect on crystallinity, crystal size and morphology.

Keywords: metal-organic framework; supercritical CO₂; crystallisation

1. Introduction

Use of supercritical carbon dioxide (scCO₂) has been extensively studied as a way to incorporate stable permanent porosity in materials during various processing steps including crystallisation [1–3], impregnation [4], dispersion [5], drying [6], and activation [7,8]. The study of porous materials, which can be either ordered structures (including zeolites, metal-organic frameworks (MOFs) and silicates), or disordered materials (including activated carbons, ceramics, metals, and polymers), has been listed among the fastest growing research areas of recent years. In this field, MOFs, normally microporous or mesoporous crystallites constructed by the coordination of transition-metal nodes and organic linkers, have emerged as enabling materials for a wide variety of potential applications, showing promise for gas storage [9–11], gas separation [12–15], catalysis [16,17], carbon dioxide
capture [18–20], and as semiconductors [21,22]. Due to the high interest in these materials, various synthetic methods have been developed beyond the conventional solvothermal approaches which, though a straightforward way to achieve highly crystalline materials, have the potential to use large amounts of organic solvents. In addition, there has been a recent trend towards the creation of multiple porosities (so-called hierarchical pore structures) in these materials [23,24], leading to the development of a number of innovative synthetic strategies including macrostructural templating [25–29], gelation [30–33], acid etching [34–37], use of scCO2 [38–42], and three-dimensional (3D) printing [43–46].

Using scCO2 during the synthesis has been shown to be useful for introducing additional porosity to traditionally microporous or mesoporous MOFs [2,3,47,48]. Unlike other methods such as templating or gelation, using scCO2 can preclude the use of additional purification or activation steps to obtain hierarchical structures without pore collapse. It has been reported that introducing scCO2 into a system containing organic or ionic liquids to obtain a switchable solvent [39] or binary solvent [49,50], can result in additional mesoporosity in MOFs. Interestingly, it has been shown that particle sizes and porosity can also be tuned by varying the CO2 pressure during synthesis, or by using scCO2 to acid etch MOFs post-synthetically, which introduces a further method for producing hierarchical porous MOFs with improved diffusion rates in catalytic applications [48].

However, the mechanism for the introduction of the larger pores in the MOF crystallites via scCO2 synthesis is not well understood. It was thought that the expanded liquid volume at CO2 pressure resulted in the mesocellular formation of building blocks before the frameworks were assembled [39]. However, by changing to hydrogen-bonded complexes (using dimethyl sulfoxide (DMSO) instead of dimethylformamide (DMF) to form a precursor solution in HKUST preparation), Doan et al. reported that the formation of the MOF occurred before additional macropores were introduced by scCO2. The latter synthesis can be related to the etching mechanism, which was further studied using different acidic agents [36,37,51]. In addition, use of scCO2 as an anti-solvent (i.e., a solvent in which the crystals are less soluble) to trigger nucleation from a MOF stock solution [47] was shown to remove the need for the 50-times excess of methanol antisolvent used in the original syntheses [52]. Here, we report our investigations into the use of scCO2 in different MOF syntheses to provide information which could improve the understanding and utility of such approaches.

The scCO2 synthetic method which was successfully employed for HKUST-1 [39,47] and Zn-benzene-1,3,5-tricarboxylate (BTC) (in DMF) [38] has here been extended to the Zn-MOF-74, UiO-66 and Zn-BTC (in DMSO) MOF systems, [Cu2(OH-mBDC)2]n, MOPs, as well as to two new flexible MOF materials [53]. The first four MOF/MOP materials were examined to see the effect of scCO2 on the structure and yield of the products. Zn-BTC was chosen as an example of a HKUST-1 MOF analogue that could not be synthesised using the mixed DMSO and MeOH solvent, which was reported previously as a crucial factor to obtain additional macro pores from expanded solvent systems [53], even though this MOF was shown to be a successful candidate for scCO2 templating [38]. [Cu2(OH-mBDC)2]n (consisting of metal-organic polyhedral nanocages) was self-assembled from the coordination between copper ions and the angular bifunctional ligand benzene-1,3-dicarboxylate and N,N-dimethylacetamide, forming a discrete structure [54]. Zn-MOF-74 was also synthesised with scCO2 in the same manner as reported in the literature [47], in order to investigate the behaviour of CO2 towards different solvent systems. UiO-66 was chosen as an example of a well-known MOF whereby low-temperature syntheses (40 °C) affords gel-like UiO-66 with many defect sites and low crystallinity. We, therefore, sought to investigate if the presence of scCO2 during low-temperature synthesis would improve the crystallinity of the resulting UiO-66. In the syntheses of Zn-Zn-MOF-74, UiO-66, Zn-BTC and the [Cu2(3,5-dBDC)2]n, MOP, scCO2 was introduced to precursor solutions where either MOFs could be formed without additional antisolvents (hereafter referred to as “precursor solution type I”) or where the MOFs crystallised only if an antisolvent was added (hereafter referred to as “precursor solution type II”—see Figure 1). Introducing scCO2 to these different solutions was carried out to help to understand the role of this supercritical fluid in MOF/MOP formation.
Figure 1. Precursor solutions used to synthesise different metal-organic frameworks/metal-organic polyhedra (MOFs/MOPs) in supercritical CO$_2$ (scCO$_2$). Precursor solution where MOFs/MOPs were formed without additional antisolvents is referred to as “precursor solution type I” and precursor solution where MOFs/MOPs crystallised only in the presence of an antisolvent is referred to as “precursor solution type II”.

The two new MOFs made from the flexible reduced Schiff-base linker 4-carboxy-phenylene-methyleneamino-4-benzoic acid (L$^1$H$_2$) and zinc hydroxide ([Zn$_2$(L$^1$)](4,4'-azopy))$\cdot$7.5H$_2$O$_n$—hereafter referred to as ZnAzopy-MOF—and ([Zn$_2$(L$^1$)](DPE))$\cdot$4H$_2$O$_n$—hereafter called ZnDPE-MOF were synthesised to see the effect of scCO$_2$ on the crystallisation between transition metals and flexible reduced Schiff-base linkers in the presence of pyridine derivatives as ancillary ligands.

MOF syntheses in scCO$_2$ are normally carried out in the presence of organic liquids such as DMF and DMSO to increase the solubility of both polar and nonpolar compounds in the reaction. It has been shown that using ligands such as tert-butylpyridine (t-bpy) and 4,4'-bipyridine can increase the solubility of the reagents in the reaction media, resulting in more efficient use of scCO$_2$ [41,42,55]. Here, we employ the flexible reduced Schiff-base compound 4-carboxy-phenylene-methyleneamino-4-benzoic acid as a linker; we note that the biphenyl-4,4'-dicarboxylate linkers have been used with great success in MOF synthesis [56–59].

In addition, research conducted by Yang et al. revealed the variability of structures achievable through the use of the flexible linker (4-carboxy-phenylene-methyleneamino-4-benzoic acid) and the related reduced Schiff-base dicarboxylic acids in conjunction with the flexible neutral pillaring linker 1,4-bis(1H-imidazol-1-yl)butane (bbi) and late transition metals [56]. Pillaring linkers, including 1,2-di(4-pyridyl)ethylene (DPE) and 4,4'-azopyridine (4,4'-azopy), were added to build up 2D or 3D MOFs with enhanced uptakes for hydrogen [60,61] and carbon dioxide [62,63]. Thus, we also wanted to investigate whether different crystalline products could be achieved via the use of scCO$_2$ in the synthesis.

2. Materials and Methods

All reagents were purchased from commercial sources and used without further purification. Due to constraints on CO$_2$ pressure, MOF synthesis at low temperature was preferred. In this study, 40 °C was chosen as the synthesis temperature for Zn-MOF-74, Zn-BTC, [Cu$_4$(OH-$m$BDC)$_{24}$]$\cdot$12H$_2$O, metal-organic polyhedron, ([Zn$_2$(L$^1$)](DPE))$\cdot$4H$_2$O$_n$ and ([Zn$_2$(L$^1$)](4,4'-azopy))$\cdot$7.5H$_2$O$_n$. Structures of
linkers and pyridine derivatives used in these MOF syntheses are given in Table A1 in Appendix B1. The conventional syntheses of these materials were also reported to allow comparison with the scCO₂ method.

2.1. Synthesis of MOFs by Conventional Methods

2.1.1. Synthesis of Zn-MOF-74

Zn-MOF-74 (i.e., Znₙ(DOBDC), where DOBDC is 2,5-dioxido-1,4-benzenedicarboxylate) was synthesised at low temperature following the method reported by Yaghi et al. [64]. Typically, 0.24 g 2,5-dihydroxyterephthalic acid (H₄DOBDC, 2.4 mmol) and 0.69 g Zn(OAc)₂·2H₂O (6.24 mmol) were dissolved in 40 mL of DMF in a 200 mL glass vial, stirring magnetically for 30 mins until a clear solution formed. This vial was sealed with parafilm before being placed in an oven for crystallisation at 40 °C for 24 h. The product was separated by filtration and repeatedly washed with methanol, before drying at room temperature to obtain 0.69 g MOF-74 solid (giving a yield of 34.0% based on Zn). The reaction scheme for formation of this MOF at low temperature is shown in Scheme 1.

![Scheme 1](image1.png)

**Scheme 1.** Synthesis of Zn-MOF-74.

2.1.2. Synthesis of [Cu₄(OH-mBDC)₃]ₙ Metal-Organic Polyhedra

The synthesis of [Cu₄(OH-mBDC)₃]ₙ MOP was followed by the procedure reported for the synthesis of copper-based cuboctahedron metal-organic polyhedra by Lee et al. [65]. The synthesis of this MOF is illustrated in Scheme 2. Typically, 40 mL of 1.46 g OH-mBDC in MeOH was mixed with 120 mL of 1.60 g Cu(OAc)₂·2H₂O in MeOH (MeOH was used as a solvent in this system) to form a stock solution. No precipitation occurred after 10 days. After that, 12.5 mL N,N'-dimethylacetamide and 7.5 mL of MeOH were added to 80 mL of the stock solution, stirring at 200 rpm at 40 °C for 3 days. The blue solid was separated by filtration, repeatedly washed with methanol, yielding 1.9 g of solid after drying at room temperature.

![Scheme 2](image2.png)

**Scheme 2.** Synthesis of [Cu₄(OH-mBDC)₃]ₙ (copper clusters represented as blue polyhedra, C–C bonds are represented as dark grey sticks, C–O bonds are represented as orange sticks; all hydrogen atoms are omitted for clarity).
2.1.3. Synthesis of ZnAzopy-MOF

The reduced Schiff-base linker, 4-carboxy-phenylene-methylene-4-benzoic acid (named L·H₂), was synthesised following the method reported previously by Liu et al. [56]. In brief, 3.0 g 4-carboxybenzoic acid (20 mmol) and 2.7 g 4-aminobenzoic acid (20 mmol) were dissolved in MeOH and the formation of a brownish solid was observed, indicating the formation of the intermediate Schiff-base product. After stirring for 2 h, NaBH₄ (2.00 g) was added to reduce the intermediate Schiff-base, resulting in the formation of the intermediate Schiff-base product. The colour faded, and the precipitate dissolved. The solvent was removed from the filtrate by rotary evaporation, and then 100 mL of distilled water was added. The mixture was adjusted to pH ~ 5 with HCl (1.0 mol l⁻¹), and an off-white solid of H₂L¹ formed immediately. The solid was filtered off, washed with Et₂O, dried, and recrystallized from MeOH/H₂O to afford the diacid, which was used without further purification. The synthetic route for the preparation of this linker is given in Scheme 3.

\[ \text{Scheme 3. The synthetic route for the preparation of } L·H₂. \]

In the synthesis of the ZnAzopy-MOF, 0.1 g of prepared L·H₂ (0.4 mmol), 0.04 g Zn(OH)₂: (0.4 mmol) and 0.04 g 4,4'-azopyridine (0.2 mmol) were added to 20 mL of distilled water. The suspension was sonicated for 10 min., and then transferred to a 23 mL Teflon-lined steel reaction vessel, which was subsequently sealed and heated to 40 °C for 72 h. The product was centrifuged and washed with distilled water, repeating 3 times, before finally drying at room temperature to obtain 0.06 g of red ZnAzopy-MOF prisms (giving a yield of 37.9% based on Zn).

2.1.4. Synthesis of ZnDPE-MOF

ZnDPE-MOF was synthesised in a similar manner as the ZnAzopy-MOF, but using 1,2-di(4-pyridyl)ethylene (DPE, 0.2 mmol) as a pillaring linker instead of 4,4'-azopyridine. All of the subsequent steps were maintained to obtain 0.14 g of yellow ZnDPE-MOF prisms (giving a yield of 40.6% based on Zn).

2.2. Synthesis of MOFs Using Supercritical CO₂

In the scCO₂ routes (Scheme 4), MOF precursor solutions (a mixture of metal salt and acid linker dissolved in a solvent) were produced with the same concentrations as the conventional methods above and were placed in a 200 mL glass vial, then sealed inside a 250 mL cylindrical steel reactor pressure vessel, magnetically stirred at 200 rpm and heated to 40 °C in an oven. The vessel was connected to a scCO₂ rig equipped with a flow-controllable liquid pump. The vessel was pressurised to 75 bar at a flowrate of 5 g min⁻¹, keeping the reaction time the same as for the conventional synthesis, i.e., without scCO₂. After reaction, the reactor was depressurised slowly to atmospheric pressure. The resulting solution was centrifuged (10,000 rpm for 10 min), washed with methanol 3 times, and dried in air at room temperature to obtain a solid product.
As a comparative control experiment in the absence of scCO₂, a mixture of 0.1 g LiH₂, 0.04 g Zn(OH)₂, 0.04 g 1,2-di(4-pyridyl)ethylene and 20 mL distilled water was pressurised with N₂ at 75 bar and 40 °C for 72 h to confirm the effect of the different gases under supercritical conditions.

Scheme 4. The scCO₂ route employed to synthesise MOFs/MOPs.

3. Results and Discussions

The first part of this study focuses on some well-known MOFs, namely: Zn-MOF-74, UiO-66, Zn-BTC, as well as the [Cu₂₄(OH₄BDC)₂₄]ₙ, MOP (see Table 1). The second part of this study focuses on new MOF systems (ZnAzopy-MOF and ZnDPE-MOF) using flexible reduced Schiff-base linkers and pyridine derivatives as ancillary ligands.

Synthetic procedures of the well-known MOFs mentioned above with and without scCO₂ are given either in the Materials and Methods section or in the Appendix B. In the conventional method in the absence of scCO₂, Zn-MOF-74 was synthesised by using DMF and taken to the crystallisation step without the use of antisolvents. Using scCO₂ in these MOF syntheses was expected to form the same meso- or macroporous structures as seen in HKUST-1 and Zn-BTC reported by Peng et al. [38,39]. [Cu₂₄(OH₄mBDC)₂₄]ₙ metal-organic polyhedron with nanoscale polyhedral structure was synthesised in MeOH in which a precursor solution of OH-mBDC and Cu(OAc)₂ was formed. The results of these syntheses are further discussed below.

Table 1. Summary of MOFs/MOPs synthesised by conventional methods.

<table>
<thead>
<tr>
<th>MOFs</th>
<th>Metal Nodes</th>
<th>Linkers</th>
<th>Comments on the Formation Using Conventional Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn-MOF-74</td>
<td>Zn</td>
<td><img src="image" alt="Linker" /></td>
<td>DMF solvent The mixture of Zn²⁺, H₄DOBDC and DMF is so-called precursor solution type I.</td>
</tr>
<tr>
<td>[Cu₂₄(OH₄mBDC)₂₄]ₙ MOP</td>
<td>Cu</td>
<td><img src="image" alt="Linker" /></td>
<td>MeOH² and DMSO⁶ solvents This MOF could not be formed in MeOH alone. The mixture of Cu²⁺, OH-mBDC and MeOH is so-called precursor solution type II.</td>
</tr>
<tr>
<td>Zn-BTC (see Appendix B2)</td>
<td>Zn</td>
<td><img src="image" alt="Linker" /></td>
<td>DMF solvent The mixture of Zn²⁺, benzene-1,3,5-tricarboxylate (BTC) and DMF is so-called precursor solution type I. This MOF could not be formed from a mixture of DMSO⁶ and MeOH².</td>
</tr>
</tbody>
</table>
The use of DMSO in BTC to form hydrogen-bonded complexes as reported in the HKUST-1 synthesis [47] was extended to the Zn-BTC. However, this method did not seem to be successful for this MOF, with no precipitation occurring from the stock solution after adding up to ten volume equivalents of MeOH and introducing scCO₂ into the system after 3 days. This might be due to the metal–ligand coordination forces between Zn and BTC not being able to overcome the H-bonding, even in the presence of CO₂-expanded MeOH as an antisolvent at 40 °C. In the synthesis of UiO-66, 40 °C was chosen as a technically achievable temperature for the scCO₂ method. However, UiO-66 synthesised at this temperature via either conventional or scCO₂ synthesis, while still crystalline, showed much broader PXRD peaks than normal UiO-66, indicating small crystallite sizes which were not noticeably affected by the presence of scCO₂. Syntheses of Zn-BTC (in DMSO) and UiO-66 are described and further discussed in the Appendix B.

3.1. Using scCO₂ in Precursor Solution Type I

Zn-MOF-74 was synthesised via the methods given in the Materials and Methods section, enabling comparison of the MOFs synthesised using conventional solvothermal synthesis, without CO₂ (here called “normal Zn-MOF-74”), to the MOF synthesised with scCO₂ (here called “Zn-MOF-74 in scCO₂”). As seen in Figure 2a, normal Zn-MOF-74 samples show a similar powder X-ray diffraction (PXRD) pattern to the simulated Zn-MOF-74 with all the main peaks at 7°, 12°, 22°, 25° and 32° 2θ preserved, confirming that this MOF can be successfully synthesised at low temperature (40 °C), in addition to the higher temperatures of 110 or 100 °C as reported by others [66,67]. The PXRD pattern of Zn-MOF-74 in scCO₂ is almost identical to the normal Zn-MOF-74 sample (see Figure 2b), showing that the scCO₂ did not overly affect the MOF crystal structure during the synthesis. Yields in both syntheses were also comparable (34% for conventional synthesis and 36% in scCO₂). SEM analysis of these samples revealed that the typical morphology of normal Zn-MOF-74 is octahedral with smooth faces, while Zn-MOF-74 in scCO₂ shows a similar morphology but with very rough faces (Figure 2b), which might be due to an acidic etching effect of the CO₂ at 75 bar and 40 °C for 72 h. Effect of the acid etching on surface roughness have been previously investigated on amorphous silica and ceramic [68,69]. In addition, etching of MOFs in aqueous acidic solutions such as a mixture of hydroquinone and MeOH (up to 180 °C for 72 h) or a mixture of phosphoric acid, MeOH and DMSO (at 40 °C for up to 10 days) has been shown to result in remarkable effects such as interconnected geometrical macropores and etched hole features [37,51,70].

<table>
<thead>
<tr>
<th>Zn-MOF-74</th>
<th>scCO₂ solvent</th>
<th>DMF solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>a: solvent</td>
<td>b: antisolvent</td>
<td></td>
</tr>
</tbody>
</table>

(see Appendix B3)
Figure 2. (a) PXRD and (b) SEM results of normal Zn-MOF-74 and Zn-MOF-74 in scCO₂. The main peaks which are preserved (compared to simulated Zn-MOF-74 [71]) are marked by light blue, confirming that this MOF can be successfully synthesised at 40 °C.

Gas sorption experiments were carried out on these samples to investigate the etching effect on the micro/mesoporosity. The surface area observed in the normal Zn-MOF-74 in this study (BET surface area of normal Zn-MOF-74 is 201 m² g⁻¹, see Figure 3a) was lower than those reported in the literature, and the type III isotherm indicated a lower level of microporosity [72]. This may be due to incomplete solvent removal as solvent removal was achieved through scCO₂ depressurisation rather than the lengthy high-temperature activation used in previous studies [72]. Further drying these samples under scCO₂ flow at 40 °C, 120 bar and 12 h showed improved microporous area (see Appendix C4). It should be noted that the scCO₂ route also provides additional features on the surface of the sample (presumably due to an etching effect), which had the effect of increasing the BET surface area of this MOF (BET surface area of Zn-MOF-74 in scCO₂ is ~350 m² g⁻¹) compared to the normal Zn-MOF-74 synthesised at low temperature (~200 m² g⁻¹, see Figure 3a).

The pore size distributions by the BJH method, however, do not show any significant differences between these two samples, meaning that there were no additional meso- and macropores formed in Zn-MOF-74 through exposure to scCO₂, indicating that addition of scCO₂ during this MOF synthesis did not help to enlarge the pores as was the case for HKUST-1 [39,47]. Note that the HKUST-1 precursor solution was stabilised by strong O–H···O hydrogen bonds, forming a two-dimensional supramolecular network within each layer (see Appendix B1).
donor groups are the hydroxyls of the trimesic acid molecules, while the acceptors are the carbonyl or the sulfoxide O atoms [73].

**Figure 3.** Gas sorption results of normal Zn-MOF-74 and Zn-MOF-74 in scCO₂. (a) N₂ isotherm at 77 K and BET surface area shows that Zn-MOF-74 in scCO₂ has improved gas uptake capacity compared with normal Zn-MOF-74. Filled markers are adsorbed N₂ and unfilled markers are desorbed N₂. (b) Pore size distribution by the BJH method shows no additional mesopores in the Zn-MOF-74 synthesised using scCO₂.

3.2. Using scCO₂ in Precursor Solution Type II

Synthesis of the [Cu₂(OH-mBDC)₂]ₖ MOP is based on self-assembly of Cu₂(COO)₄ paddlewheels serving as square secondary building units (SBUs) bonded at a 120° angle to OH-mBDC and N,N’-dimethylacetamide serving as the ligands [54]. It was shown that the metal nodes and linkers in this metal-organic polyhedron remained unreacted in MeOH for up to 10 days, resulting in a stable precursor solution before DMSO is introduced. Stable precursors of this sort are promising for investigation of the effect of scCO₂ on this metal-organic polyhedron because the aggregation can happen at the same time the solvent is expanded. The crystallinity of [Cu₂(OH-mBDC)₂]ₖ MOP synthesised via the conventional method was confirmed by the similarity between the PXRD pattern of this sample to the simulated pattern reported by Li et al. [74]. In comparing the two [Cu₂(OH-mBDC)₂]ₖ MOP samples synthesised from the same starting materials and at the same temperature but with and without scCO₂ pressure (i.e., at 0 and 75 bar), some distortion occurred in the sample treated with scCO₂, as evidenced by differences in the intensities and shapes of the PXRD peaks at 6°, 7°, 11°, 14° and 17° 2θ (see Figure 4a). These two samples were analysed under inelastic neutron scattering (INS) spectroscopy (see Appendix C2), showing that all bond vibrations within these two structures are identical. In the SEM results (Figure 4b), it can be seen that they have identical morphologies. However, the crystallite size of the samples produced in scCO₂ is remarkably larger than the conventionally synthesised sample (~2 µm compared with ~1 µm), indicating the scCO₂ would lower the deprotonation rate of OH-mBDC, thus decreasing the growth rates of this MOP [75]. Dissolving 1.6 g Cu(OAc)₂·H₂O and 1.5 g OH-mBDC in 100 mL DMSO resulted in precipitation after 2 h (see Appendix B1).
Figure 4. (a) PXRD results of simulated \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\), normal \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\) and \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\) in scCO\(_2\). Peaks which are changed are marked by light orange. (b) SEM results of normal \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\) and \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\) in scCO\(_2\), showing that these samples have identical morphology but increased crystal size and do not appear to have additional meso- and macropores.

The results of the syntheses of Zn-MOF-74, UiO-66, \([\text{Cu}_{24}(\text{OH}-\text{mBDC})_{24}]_n\) and Zn-BTC (in DMSO) are summarised in Table 2.

Table 2. A summary of the results from both precursor solutions type I and II in the synthesis of MOFs/MOPs in the presence of scCO\(_2\).

<table>
<thead>
<tr>
<th>Precursor</th>
<th>MOFs</th>
<th>Solvents</th>
<th>Temperature</th>
<th>Comments on the Formation in scCO(_2):</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type I</td>
<td>Zn-MOF-74</td>
<td>DMF</td>
<td>40 °C</td>
<td>Product formed. Crystallinity and morphology preserved. Surface texture changed. Surface area increased.</td>
</tr>
<tr>
<td></td>
<td>UiO-66</td>
<td>DMF</td>
<td>40 °C</td>
<td>Same as without scCO(_2) at 40 °C but with crystal size smaller than normal UiO-66</td>
</tr>
<tr>
<td>Type II</td>
<td>[\text{Cu}<em>{24}(\text{OH}-\text{mBDC})</em>{24}]_n MOP</td>
<td>MeOH/DMSO</td>
<td>40 °C</td>
<td>Product formed. Crystallinity slightly changed. Morphology preserved. Crystal size increased.</td>
</tr>
<tr>
<td></td>
<td>Zn-BTC (in DMSO)</td>
<td>DMSO</td>
<td>40 °C</td>
<td>Product not formed.</td>
</tr>
</tbody>
</table>

3.3. Using scCO\(_2\) in the Presence of Pyridine Derivatives as Ancillary Ligands to Synthesise Pillared MOFs

Synthesis of MOF systems using flexible reduced Schiff base linkers and pyridine derivatives as ancillary ligands, including ZnAzopy-MOF and ZnDPE-MOF were also performed with and without scCO\(_2\), to determine the effect of these differing conditions on the nature of the final product.
3.3.1. In ZnAzopy-MOF Synthesis

ZnAzopy-MOF is conventionally synthesised by hydrothermal reaction between the flexible reduced Schiff base linker 4-carboxy-phenylene-methyleneamino-4-benzoic acid (L1H2) and zinc hydroxide and the rigid pillaring linker 4,4'-azopy. The inclusion of an element of rigidity provides a backbone around which the flexible linkers can bind, thereby allowing the assembly of 3D structures bearing novel network topologies. The molecular structure has been determined by single crystal X-ray diffraction (see Figure 5). The asymmetric unit comprises one Zn ion in a general position [Zn(1)], one Zn ion on a centre of symmetry [Zn(2)], half the bipyridyl ligand, one and a half dicarboxylate ligands and approx. 3.75 water molecules. Zn(1) is 4-coordinate, severely distorted tetrahedral or triangular based pyramidal, binding to O(1), O(4), O(5'), and N(1), whilst Zn(2) is 6-coordinate, distorted octahedral, binding to six carboxylate oxygens. Three Zn ions form a double lantern arrangement with carboxylates bridging pairs of Zn ions and pyridyl groups capping both ends. The coordination environment around the metal centres in this structure is reminiscent of the paddlewheel SBU, which comprises three zinc centres forming an hour-glass shape by the coordination of carboxylate moieties from six L1 units in the di-mono-dentate bridging mode in the equatorial positions, with 4,4'-azopy units coordinated to the axial positions of the hour glass, (Figure 5a). The L1 units extend in six directions in the a/b plane, adopting a bent conformation through the rotation of the amine and methylene bridges, whilst the 4,4'-azopy ligands extend along the c axis, linking hour-glass SBUs in to an extended 3D framework, bearing roughly cylindrical 1D channels of 4.75 Å diameter observed parallel to the c axis (Figure 5b). Elemental analysis of this MOF is given in Appendix C3, together with alternative views of the structure.

![Figure 5. Structure of ZnAzopy-MOF. (a) The coordination environment around the Zn^2+ metal centres in ZnAzopy-MOF. (b) Diagram of ZnAzopy-MOF as viewed along the c axis. Carbon, nitrogen, oxygen and zinc atoms are represented by empty, blue, red and green circles, respectively. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) and angles (°): Zn(1)–N(1) 2.061(8), Zn(1)–O(1) 1.995(8), Zn(1)–O(5) 1.975(6), Zn(2)–O(2) 2.256(12), Zn(2)–O(6) 2.194(10); O(1)–Zn(1)–O(5) 113.5(3), O(1)–Zn(1)–N(1) 96.3(3), O(2)–Zn(2)–O(6) 81.4(4).](image-url)
of the MOF synthesised in scCO₂, including the disappearance of low-angle peaks at 7°, 8° and 13° 2θ, the appearance of a new peak at 17° 2θ and a shift in the peak at 33° 2θ, showing that the addition of scCO₂ during the synthesis had a small but noticeable effect on the crystal structure. This might be due to the change in dilution of 4,4’-azopyridine, which was highly soluble in scCO₂ [76]. Note that this ancillary ligand is crucial for formation of this MOF. In general, large crystals were precipitated from both syntheses, as can be seen in Figure 7. The increase in the z-dimensional diameter of ZnAzopy-MOF in scCO₂ can be referred back to the higher solubility of the reagents in the media with 4,4’-azopyridine used as a co-solvent and a pillaring linker with respect to scCO₂, thus supporting crystal growth and nucleation. Gas sorption results (Appendix C4), however, showed that these samples are non-porous with surface areas less than 20 m² g⁻¹.

![Figure 6. PXRD results of normal ZnAzopy-MOF and ZnAzopy-MOF in scCO₂. The main peaks which are preserved are marked by light blue. Peaks which are changed are marked by light orange.](image1)

![Figure 7. SEM images of normal ZnAzopy-MOF and ZnAzopy-MOF in scCO₂.](image2)
3.3.2. In ZnDPE-MOF Synthesis

ZnDPE-MOF is conventionally formed by the hydrothermal reaction of equimolar quantities of Zn(OH)$_2$ and L$^1$Hz with half an equivalent of DPE, to produce the 3D structure[[Zn(L$^1$)(DPE)]-4H$_2$O]$_n$. The coordination environment around the metal centres in this structure, as determined by single crystal X-ray diffraction, is distorted octahedral with two chelating carboxylates and two monodentate DPE pyridinyl nitrogens in a cis conformation (see Figure 8a). Coordination of the pyridinyl nitrogens of the neutral DPE linkers in the cis conformation gives rise to zig-zag chains and considering also the L$^1$ connections an overall 3D network (see Figure 8b). Elemental analysis of this MOF is given in the Appendix C4.

![Figure 8](image_url)

**Figure 8.** Structure of ZnDPE-MOF. (a) The asymmetric unit of ZnDPE-MOF showing the anionic linker and the one unique Zn$^{2+}$ environment; H-bonding interactions as dotted lines. (b) Packing plot of ZnDPE-MOF showing the extended structure viewed parallel to c. Carbon, nitrogen, oxygen and zinc atoms are represented by empty circles and blue, red and green circles, respectively. Hydrogen atoms, except those involved in H-bonding, are omitted for clarity. Selected bond lengths (Å) and angles (°): Selected bond lengths (Å) and angles (°): Zn(1)–N(1) 2.062(4), Zn(1)–O(1) 2.038(4), Zn(1)–O(2) 2.434(4), Zn(1)–O(3A) 2.259(3), Zn(1)–O(4A) 2.061(4), Zn(1)–N(2A) 2.093(4); O(1)–Zn(1)–O(2) 58.28 (14), O(1)–Zn(1)–N(1) 107.34(15), N(2A)–Zn(1)–O(3A) 88.81(14).

Different gases (CO$_2$ with T$_c$ 31.1 °C and P$_c$ 73.9 bar; and N$_2$ with T$_c$ –147 °C and P$_c$ 34 bar) were introduced to the precursor solution to synthesise this MOF at 40 °C and 75 bar, forming supercritical systems (scCO$_2$ and scN$_2$) with increased solubility and decreased polarity. It can be seen that the yield of this MOF in scCO$_2$ is considerably higher than in scN$_2$ (an increase of 29.8 wt%). The crystallinity of samples was examined using PXRD (see Figure A6 in Appendix C1 and Figure 9). Most of the main peaks at 8°, 20°, 26°, 27° and 29° 2θ in the simulated ZnDPE-MOF were present in both samples (Figure A6), showing that the crystalline structure had formed and was preserved under high-pressure syntheses. There was a slight difference between the samples synthesised in scCO$_2$ and in scN$_2$. In the sample synthesised in scCO$_2$, there is an extra peak at 12° 2θ, and the peak appearing at 14° 2θ is more clearly a doublet in comparison to the sample in scN$_2$. The transformation in peak shape was extended to those at 15°, 17° and 24° 2θ. These differences show the evidence of the effect of CO$_2$ on the MOF crystal structure. At the same time, the difference in the nature of the supercritical fluids introduced had an effect on the morphology of this MOF during the crystallisation (Figure 10). In general, both samples appear to not have uniform shape, with the size varying between 3 and 15 μm. However, in more detail, the sample synthesised in scCO$_2$ is
represented by bulky crystal agglomerates compared with the smooth surfaces observed for the crystals of the MOF synthesised in N₂. This may be due to the etching effect of CO₂, which was shown to result in rougher, pitted crystal surfaces in the Zn-MOF-74 synthesis.

![Figure 9. PXRD results of ZnDPE-MOF synthesised in scCO₂ and scN₂ at 40 °C and 75 bar. Peaks which are changed are marked by light orange.](image)

### Figure 9.

**Figure 9.** PXRD results of ZnDPE-MOF synthesised in scCO₂ and scN₂ at 40 °C and 75 bar. Peaks which are changed are marked by light orange.

![Figure 10. SEM images of ZnDPE-MOF synthesised in scCO₂ and scN₂ at 75 bar.](image)

### Figure 10.

**Figure 10.** SEM images of ZnDPE-MOF synthesised in scCO₂ and scN₂ at 75 bar.

### 4. Conclusions

In this study, scCO₂ was introduced to different precursor complexes to synthesise various MOF/MOP systems. There was substantial evidence that scCO₂ had an effect on surface texturing and crystal growth in these MOF syntheses. Zn-MOF-74 was formed in scCO₂ and precursor solution type I, showing some changes in surface texture and surface area. [Cu₃{(OH-mBDC)}₃]ₙ MOP with increased crystal size was also formed in scCO₂ and precursor solution type II, indicating the growth rates of this MOF increased in the CO₂-expanded solvent system. However, there was no evidence showing that additional porosity appeared in these MOFs in scCO₂. Zn-BTC (in DMSO) and UiO-66 were formed but contained smaller crystallites in the scCO₂ case. The introduction of scCO₂ to the synthesis of two new MOF structures {[Zn(L²)(DPE)]·4H₂O}ₙ and {[Zn(L³)(4,4'-azopy)]·7.5H₂O}ₙ using pyridine derivatives as ancillary ligands additionally showed that changes in crystallinity and morphology could result. This was shown (through analogous experiments with N₂ under the same conditions) to be an effect of the presence of scCO₂, rather than simply a direct result of high-pressure synthesis. However, again this supercritical fluid did not
result in any further meso- or macroporosity in the resulting samples. While scCO₂ indeed showed some positive effects on the synthesis of Zn-BTC and HKUST-1 in introducing additional porosity, the mechanism of these effects needs to be further studied to extend the method to other MOF systems.

These findings provide new information on the effects of introducing scCO₂ into the synthesis of a broad range of different MOFs, and provides information to direct further application of scCO₂ as an approach for morphological control.

**Author Contributions:** Conceptualization, H.V.D.; Data curation, H.V.D. and F.C.; Formal analysis, H.V.D.; Funding acquisition, C.R. and V.P.T.; Investigation, H.V.D.; Methodology, H.V.D., T.D., M.R.J.E. and O.R.; Supervision, V.P.T.; Writing—original draft, H.V.D.; Writing—review & editing, H.V.D., J.C., C.R. and V.P.T. All authors have read and agreed to the published version of the manuscript.

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**Acknowledgments:** The X-ray Crystallographic Service at Southampton is thanked for data collection for ZnAzopy-MOF and ZnDPE-MOF. We thank the British Council Newton Fund for funding for a materials workshop at Northwest University in Xi’an. The INS spectra for [Cu₅(OH-mBDC)₃]·MOF herein, assisted by Svetmar Rudic, were collected on the TOSCA instrument at ISIS Neutron and Muon Source, Didcot, UK.

**Conflicts of Interest:** The authors declare no conflict of interest.

**Appendix A. Characterisation Methods**

Materials after synthesis were tested by powder X-ray diffraction (PXRD) on a BRUKER AXS D8-Advance instrument with Vantec-1 detector using Cu Kα (λ = 1.5418 Å) as the source of X-ray radiation, in flat plate geometry, spinner speed 15 rpm, at 21 °C. The 2θ range, between 2–60° was used with a 0.02° interval for collection and 20 min scans.

Scanning electron microscopy (SEM) images were taken using JEOL IT300 SEM with a SED detector at School of Chemistry, University of Bristol with the following settings, objective aperture: 1, acceleration voltage: 5 kV, probe current: 10.3 m, working distance: 10.8 mm and magnifications from 350 to 40,000 times. The samples were coated with 15 nm silver before conducting the experiment.

Gas sorption isotherms were determined using nitrogen sorption at 77 K with a Micromeritics 3-Flex volumetric gas sorption analysis system (nitrogen with purity of 99.9999% was purchased from Air Products). Samples were degassed at 120 °C under dynamic high vacuum (10⁻⁶ mbar) over 6 h prior to analysis. The total pore volume was taken at the end of the filling of the pore. Surface area was determined by the BET method according to British Standards, with relative pressure (P/P₀) selected considering the Rouquerol consistency criterion between values of 0 and 0.3.

Single crystal diffraction data for ZnAzopy-MOF and ZnDPE-MOF were collected on a Rigaku Saturn 724+ CCD diffractometer using a rotating anode X-ray source and 10 cm confocal mirrors monochromator at 100 K. Data were corrected for absorption and Lp effects. Structures were solved by direct methods and refined by full-matrix least squares on F² [77]. Further details are given in Table 1. H atoms were included in a constrained riding model except for those on water where coordinates were refined. In ZnAzopy-MOF the badly disordered solvent water in the void spaces was modelled as diffuse electron density by the Platon Squeeze procedure which recovered 2 × 73 and 2 × 78 electrons over four voids in the unit cell [78]. This equates to approx. 30 water molecules per unit cell or approx. 7.5 per void or Zn₆ unit. The CH₃NH linkers C(13)/N(3) and C(28)/N(4) in the dicarboxylate ligands could not be clearly distinguished and were each modelled as 50/50 C/N. For ZnAzopy-MOF, where disorder was modelled, restraints on both geometry and displacement parameters were applied. CCDC 935555 and 1919724 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

Analysis of [Cu₅(OH-mBDC)₃]·MOF with inelastic neutron scattering (INS) spectroscopy was achieved on the TOSCA indirect geometry spectrometer at ISIS Neutron and Muon Source, Didcot,
The dehydrated sample was prepared within a glove box under an inert argon atmosphere. The sample was first loaded into aluminium foil sachets, with the loaded sample mass recorded. The sachets were then sealed between two aluminium plates, using indium wire as a seal to keep the sample under an inert atmosphere. INS spectra were recorded within the energy transfer range −80–8050 cm⁻¹, at a temperature of 10 K for 5–7 h. Subsequent data analysis, visualisation and normalisation was achieved using the Mantid software [80].

Appendix B. Additional Syntheses and Discussion

Appendix B.1. Synthesis of [Cu₂₄(OH−mBDC)₂₄]n Metal-Organic Polyhedron in DMSO and MeOH

![Figure A1](image1)

*Figure A1.* A mixture of 1.6 g Cu(OAc)₂·H₂O and 1.5 g OH−mBDC in 100 mL DMSO after 2 h, showing a precipitation occurred in [Cu₂₄(OH−mBDC)₂₄]n stock solution.

![Figure A2](image2)

*Figure A2.* Hydrogen bonds formed between trimesic acid and DMSO. C atoms are represented by back spheres. O atoms are represented by red spheres. H atoms are represented by pink spheres. S atoms are represented by yellow spheres.

Appendix B.2. Synthesis of Zn-BTC in DMSO and MeOH

The coordination of Zn(II) metal nodes and benzene-1,3,5-tricarboxylic acid (BTC) linkers, resulting in a 3D open network as found in Zn-BTC MOF. This MOF was successfully synthesised by solvothermal reactions between BTC and Zn(NO₃)₂·6H₂O in a mixture of DMF and DMAC without any additives [81]. In fact, Peng et al. used this MOF to test the effect of CO₂ in the presence of N-EtFOSA/TMGt solution [38]. By changing the pressure of the formed ionic liquid system from 10
to 63 bar, tetrahedron-like Zn-BTC particles with some evidence of mesopores were observed. In some cases where a high number of bulky molecules get involved, macropores were more favoured to minimise diffusion barrier, hence were managed to achieve in this study. With the success of the creation of macroporous HKUST-1 (or Cu-BTC) in scCO2 [47], Zn-BTC synthesis was repeated in the same manner. In a typical experiment, 3.1 mg Zn(NO3)2·6H2O (10 mmol) and 1.5 mg BTC (5 mmol) were dissolved in 100 mL DMSO, resulting in a clear precursor solution. As demonstrated previously [82], H-bonding between DMSO and BTC dominated the metal-ligand coordination forces and hindered the aggregation process, resulting in a stable stock solution. This solution (1–5 mL) was added in MeOH at varied volume ratio between 1:1 and 1:10, and left for up to 10 days at 40 °C. However, no precipitation was observed, consistent with the precursor solution of HKUST-1 [47]. A mixture of 4 mL stock solution and 40 mL MeOH was pressurised with CO2 at 40 °C and 75 bar; however, no solid was formed after 3 days. This might be due to the metal–ligand coordination forces between Zn and BTC not being able to overcome the H-bonding even in the presence of CO2-expanded MeOH as an antisolvent at 40 °C.

Figure A3. Stock solution of Zn(NO3)2·6H2O (10 mmol), BTC (5 mmol) and DMSO in MeOH at different volume ratios. Stock solution/MeOH volume ratio (from left to right) is 1:1, 1:2, 1:4, 1:6, 1:8 and 1:10.

Appendix B.3. Synthesis of Uio-66 in DMF at Low Temperature

Normal UiO-66 was synthesised by dissolving 3.18 g ZrCl4 (0.014 mol) and 2.04 g benzene-1,4-dicarboxylic acid (H2BDC) (0.014 mol) in 80 mL anhydrous dimethylformamide (DMF) at room temperature. The reaction of this step is given in Scheme A1. The reaction mixture was heated in an autoclave at 120 °C for 24 h. After cooling in air to room temperature the white solid was separated by filtration, repeatedly washed with DMF and dried at room temperature. In the modified synthetic procedure, the autoclave was heated at 40 °C instead of 120 °C, with all other steps remaining the same. In the synthesis with scCO2, the stock solution was added in a glass vial and placed in a 250 mL cylindrical steel reactor pressure vessel and heated to 40 °C in an oven. The vessel was connected to a scCO2 rig equipped with a flow controllable liquid pump. The vessel was pressurised to 75 bar at a flowrate of 5 g min⁻¹ for 24 h. After that, the reactor was depressurised slowly to atmospheric pressure. The white solid was separated by filtration, repeatedly washed with DMF and dried at room temperature.

The sample synthesised at 40 °C and dried at room temperature appeared to be gel-like light yellow crystals which are very different from the solids in the sample synthesised at 120 °C. Tian et al. also reported that a gel-like UiO-66 monolith could be formed by a sol-gel process and mild drying conditions [83]. In the PXRD results, it can be seen that the low temperature synthesised sample has preserved peaks at 7°, 12°, 22°, 26°, 31°, 43° and 50° 2θ (see Figure A4), confirming the UiO-66 structure in this sample. However, these peaks appeared much broader than the peaks in normal UiO-66, showing that the sample synthesised at 40 °C has smaller crystallite size than those synthesised at 120 °C. It has been demonstrated that UiO-66 obtained by room-temperature synthesis had defect sites with the maximal number achieved (~1.3 missing linker per SBU) at a temperature of 45 °C [84]. In this study, highly-crystalline MOFs were the principal focus. The scCO₂ experiment was applied in this MOF synthesis, however, the PXRD patterns of both samples synthesised at 40 °C are almost identical, showing that scCO₂ did not enhance the crystallite size in this MOF.

Figure A4. UiO-66 synthesised at 40 °C (with and without scCO₂) and 120 °C.
Appendix B.4. Summary of MOFs Synthesised in the Presence of Pyridine Derivatives

Table A1. Reagents used for the synthesis of MOFs with pyridine derivatives.

<table>
<thead>
<tr>
<th>MOFs</th>
<th>Metal Nodes</th>
<th>Linkers</th>
<th>Pyridine Derivatives</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZnAzopy-MOF</td>
<td>Zn</td>
<td><img src="image" alt="ZnAzopy-MOF" /></td>
<td>4,4'-azopyridine</td>
</tr>
<tr>
<td>ZnDPE-MOF</td>
<td>Zn</td>
<td><img src="image" alt="ZnDPE-MOF" /></td>
<td>1,2-di(4-pyridyl)ethylene</td>
</tr>
</tbody>
</table>
Appendix C. Additional Results of MOFs Synthesised in the Presence of Pyridine Derivatives

Appendix C.1. PXRD Results

Figure A5. PXRD results of simulated ZnAzopy-MOF, normal ZnAzopy-MOF and ZnAzopy-MOF in scCO₂.

Figure A6. PXRD results of ZnDPE-MOF synthesised in scCO₂ and N₂ at 40 °C and 75 bar compared with simulated ZnDPE-MOF, showing that the primary crystallinity was maintained in the prepared samples (highlighted by light blue). Peaks for ZnO are highlighted by light orange, showing that residual ZnO from synthesised samples are insignificant.
Appendix C.2. INS Results

Figure A7. INS spectra of normal \([\text{Cu}_2(\text{OH}-m\text{BDC})_{24}]_n\) and \([\text{Cu}_2(\text{OH}-n\text{BDC})_{24}]_n\) in scCO\(_2\), showing bond vibrations within two structures are identical.

Appendix C.3. Method and Elemental Analysis of ZnAzopy-MOF and ZnDPE-MOF

Ligand L-H: (0.2 mmol, 0.05 g), Zn(OH)\(_2\) (0.2 mmol, 0.02 g) and 4,4’-azopy (0.1 mmol, 0.02 g) were added to 10 mL distilled water. The suspension was transferred to a 23 mL Teflon-lined steel reaction vessel which was subsequently sealed and heated to 110 °C for 72 h. The vessel was then cooled at a rate of 2 °C h\(^{-1}\) to room temperature, yielding 0.04 g red prisms (50.5% based on Zn).

Elem. anal. calcd. for ZnAzopy-MOF (C\(_{55}\)H\(_{41}\)N\(_7\)O\(_{12}\)Zn\(_3\)∙7.5H\(_2\)O): C 50.26; H 4.22; N 7.46; found C 50.61, H 4.29, N 7.08%.

IR/cm\(^{-1}\) (KBr): 3348(w), 2717(w), 2667(w), 2359(w), 2336(w), 1700(w), 1653(w), 1606(s), 1559(w), 1539(w), 1519(w), 1506(w), 1488(w), 1331(m), 1281(m), 1222(w), 1180(m), 1142(w), 1111(w), 1080(w), 1046(w), 1029(w), 987(w), 847(m), 782(m), 761(m), 721(w), 703(w).

Table A2. Crystallographic data for structures ZnAzopy-MOF and ZnDPE-MOF.

<table>
<thead>
<tr>
<th>Structure</th>
<th>ZnAzopy-MOF</th>
<th>ZnDPE-MOF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formula</td>
<td>C(<em>{55})H(</em>{41})N(<em>7)O(</em>{12})Zn(_3)∙7.5H(_2)O</td>
<td>C(<em>{27})H(</em>{20})N(_3)O(_7)Zn(_2)∙4(H(_2)O)</td>
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<tr>
<td>Formula weight (g mol(^{-1}))</td>
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<td>653.31</td>
</tr>
<tr>
<td>Crystal system</td>
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<td>Tetragonal</td>
</tr>
<tr>
<td>Space group</td>
<td>C2/c</td>
<td>P4(_1)</td>
</tr>
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<td>Unit cell dimensions</td>
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<td>a (Å)</td>
<td>21.2518(17)</td>
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<tr>
<td>c (Å)</td>
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<td>19.8120(5)</td>
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<td>$\gamma$ (°)</td>
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<td>2743.68(11)</td>
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<td>Calculated density (Mg m$^{-3}$)</td>
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<td>1.423</td>
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<tr>
<td>Absorption coefficient (mm$^{-1}$)</td>
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<td>$0.06 \times 0.05 \times 0.03$</td>
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<td>Unique reflections</td>
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<td>$R_{\text{int}}$</td>
<td>0.053</td>
<td>0.032</td>
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<tr>
<td>Reflections with $F^2 &gt; 2\sigma(F^2)$</td>
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<td>5040</td>
</tr>
<tr>
<td>Number of parameters</td>
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<td>362</td>
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<tr>
<td>$R_1$, $wR_2$: [$F^2 &gt; 2\sigma(F^2)$]</td>
<td>0.108, 0.321</td>
<td>0.043, 0.103</td>
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<tr>
<td>GOOF</td>
<td>1.07</td>
<td>1.03</td>
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<tr>
<td>Largest difference peak and hole (e Å$^3$)</td>
<td>0.86 and −2.11</td>
<td>0.32 and −0.45</td>
</tr>
</tbody>
</table>

**Figure A8.** Alternative models and views of ZnAzopy-MOF: ball and stick models in asymmetric unit (a) and packing plot (b) and (c), and space filling model (d). Carbon, nitrogen, oxygen and zinc atoms are represented by empty circles and blue, red and green circles, respectively. Selected bond
lengths (Å) and angles (°): Zn(1)–N(1) 2.061(8), Zn(1)–O(1) 1.995(8), Zn(1)–O(5) 1.975(6), Zn(2)–O(2) 2.256(12), Zn(2)–O(6) 2.194(10); O(1)–Zn(1)–O(5) 113.5(3), O(1)–Zn(1)–N(1) 96.3(3), O(2)–Zn(2)–O(6) 81.4(4).

Figure A9. Alternative models of ZnDPE-MOF: ball and stick models in packing plot (a) and (b), and space filling model (c). Carbon, nitrogen, oxygen and zinc atoms are represented by empty circles.
and blue, red and green circles, respectively. Selected bond lengths (Å) and angles (°): Zn(1)–N(1) 2.062(4), Zn(1)–O(1) 2.038(4), Zn(1)–O(2) 2.434(4), Zn(1)–O(3A) 2.259(3), Zn(1)–O(4A) 2.061(4), Zn(1)–N(2A) 2.093(4); O(1)–Zn(1)–O(2) 58.28(14), O(1)–Zn(1)–N(1) 107.34(15), N(2A)–Zn(1)–O(3A) 88.81(14).

Appendix C.4. Gas Sorption Results of ZnAzopy-MOF

Figure A10. 77 K nitrogen isotherms of Zn-MOF-74 in scCO₂ before and after dried with scCO₂ flow at 40 °C, 120 bar and 12 h.

Figure A11. Gas sorption results of ZnAzopy-MOF synthesised without scCO₂ (normal ZnAzopy-MOF) and with scCO₂ (ZnAzopy-MOF in scCO₂). (a) N₂ isotherm and BET surface area. (b) CO₂ isotherm. Filled markers are adsorbed gas molecules and unfilled markers are desorbed gas molecules.

References


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