Title: Post-Synthetic Selective Ligand Cleavage by Solid-Gas Phase Ozonolysis Fuses Micropores into Mesopores in Metal-Organic Frameworks

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ABSTRACT

Herein we report a novel, ozone-based method for post-synthetic generation of mesoporosity in metal-organic frameworks (MOFs). By carefully selecting mixed-ligand Zr-fcu-MOFs based on organic ligand pairs in which one ligand has ozone-cleavable olefin bonds and the other ligand is ozone-resistant, we were able to selectively break the cleavable ligand via ozonolysis to trigger fusion of micropores into mesopores within the MOF framework. This solid-gas phase method is performed at room-temperature and, depending on the cleavable ligand used, the resultant ligand-fragments can be removed from the ozonated MOF by either washing or sublimation. Compared to the corresponding highly-microporous starting MOFs, the highly-mesoporous product MOFs exhibit radically distinct gas sorption properties.
INTRODUCTION

Since the advent of metal-organic frameworks (MOFs) in the late 1990’s, these porous materials have shown promise for critical applications such as gas storage, catalysis, drug delivery, thermal energy storage, sensors, etc. MOFs have become benchmark materials for adsorption, surpassing most traditional adsorbents for hydrogen storage, methane storage, CO$_2$/N$_2$ separation, etc. By marrying the advantages of the inorganic and organic chemistries, MOFs exhibit both polyfunctionality and a high degree of tunability, as well as the opportunity to post-functionalize them. Among the main advantage of this tunability is the possibility to perform reticular chemistry through ligand functionalization, length/width variation, and/or metal ion substitution, which enables strategic design and synthesis of materials for specific applications.

A remaining challenge in MOF chemistry is the creation of mesoscale cavities, which are necessary for increased storage capacity, encapsulation of large molecules, etc. Reticular chemistry strategies to surpass previous limits on pore size and cavity diameter include increasing the distance between the organic molecular building blocks and using longer or wider organic ligands. However, many of the most well-known MOFs are associated to self-dual nets (pcu: MOF-5, IRMOFs series, MXFSIX MOFs) or nets that can easily interpenetrate upon ligand elongation (tbo: HKUST-1, fcu: UiO-66, acs: MIL-88’s, etc.). To avoid this problem, researchers have developed platforms based on topologies that cannot interpenetrate, such as IRMOFs-74 and rht-MOFs, which, thanks to the design of very ambitious organic ligands, currently hold porosity records.

Regrettably, systematic access to MOFs exhibiting hierarchical porosity from the micro to the meso range remains a challenge, as it currently demands use of complex organic ligands that are not commercially available. Although MOFs were initially considered as structurally ideal, defect-free materials, numerous recent studies have highlighted the non-negligible level of structural defects and irregularities in these materials, which strongly influence their porosity and catalytic properties. These findings suggest that access to hierarchically porous materials would require rational control over the structural defects within MOFs. Unfortunately, to date, only a few methods for generating and controlling such defects have emerged. One such strategy is to use large amounts of monotopic agents (known as modulators) in competition with the required polytopic ligand corresponding to the targeted MOFs. Alternatively, rational design strategies such as transversal reticular chemistry also have proven invaluable for generating MOFs with ordered defects. However, there are very few reports on engineering of defects and control of hierarchical porosity with commercially available and/or moderately sized ligands and some of the existing methods require harsh chemical or thermal treatments.

Our group recently reported the first solid-gas phase covalent post-synthetic modification of a MOF: under mild conditions, we achieved quantitative conversion of the constituent olefin groups in single crystals of UiO-66-like MOF into trioxolane moieties, aldehydes and/or carboxylates without compromising the crystallinity. Encouraged by the stability of the MOFs under these conditions, we envisioned exploiting this solid-gas phase ozonolysis method to cleave and remove selected organic ligands within MOFs. We hypothesized and later confirmed (vide infra) that ligands containing non-terminal olefin groups could be split into several parts (Scheme S1). When implemented in MOFs, these ligands can be selectively broken and subsequently removed from the framework, to provide an original and controllable method for post-synthetic fusion of micropores into mesopores.

Herein we report application of our post-synthetic strategy to selectively and quantitatively cleave and remove the organic ligands in two multivariate (MTV) Zr-fcu-MOFs (Figure 1), thereby affecting their adsorption performance in gas uptake. By controlling the ozone inert/active ratio of ligands in these MOFs, we were able to control the final number of defects in their structures.
Figure 1. Post-synthetic modification of multivariate Zr-fcu-MOFs via solid-gas ozonolysis. Reaction of a) 12-connected hexanuclear clusters (eo molecular building blocks) with b) pairs of organic ligands that are similar in length but show opposite reactivity to ozone (H_{2}sti & H_{2}azo or H_{2}pdac & H_{2}bpd) yields c) multivariate MOFs with the fcu topology and two types of cages. Post-synthetic modification of the resultant MOFs through ligand cleavage by solid-gas ozonolysis results in d) Zr-fcu-MOFs with a defective network that exhibits hierarchical porosity generated upon fusion of micropores into larger pores (up to the meso scale).

RESULTS AND DISCUSSION

Zr-fcu-azo/sti system.

System validation. Our ligand-removal strategy is only amenable to MOFs based on ligand pairs that exhibit the following characteristics: similar shape and length; similar reactivity under given reaction conditions; and opposite reactivity to ozone (Figure 1b). To this end, we selected two dicarboxylic acids: 4,4’-azobenzene dicarboxylic acid (H_{2}azo) and 4,4’-stilbene dicarboxylic acid (H_{2}sti) (with lengths ≈ 13.3 Å). The reactivity of each ligand to ozone was initially tested in solution, in N,N’-dimethylformamide (DMF), which was used to avoid π-π stacking of the ligands and to facilitate access of the ozone molecules to the olefin groups. Interestingly, the two acids showed diametric responses to ozone: H_{2}azo was unreactive, whereas H_{2}sti was fully converted into terephthalic acid (H_{2}bdc) and formylbenzoic acid (Hfba) (Scheme S1, Figure S18).

Having confirmed the suitability of H_{2}azo and H_{2}sti as a ligand pair, we then decided to choose an appropriate MOF for testing. We selected the robust Zr-fcu-MOF platform, which had already proven amenable to post-synthetic modification and ozone resistant. Thus, we embarked on the synthesis of the resultant MOFs (Zr-fcu-azo and Zr-fcu-sti) and testing of their response to ozonolysis. Both MOFs were synthesized by suspending ZrCl_{4} and the corresponding dicarboxylic ligand in DMF, in the presence of HCl and L-proline, and then heating the resulting mixture at 120 °C.
overnight in a scintillation vial. The resulting crystals (of Zr-fcu-azo or Zr-fcu-sti) were washed with DMF and then subjected to solvent exchange with acetone. As expected, N$_2$ sorption isotherms collected at 77 K confirmed comparable Brunauer-Emmett-Teller areas (A$_{BET}$) of 3025 m$^2$/g for Zr-fcu-azo and 3065 m$^2$/g for Zr-fcu-sti (Figure 2a). When submitted to ozone treatment for 30 min, Zr-fcu-azo retained its crystallinity and porosity, whereas Zr-fcu-sti collapsed into an amorphous, non-porous solid due to cleavage of all the ligands of the framework (Figures 2a, S2, S9).

**Synthesis and porosity of Zr-fcu-azo/sti-X%**. To prepare multivariate Zr-fcu-MOFs containing specific ratios of azo/sti (ranging from 90:10 to 40:60), we used similar synthetic conditions as for Zr-fcu-azo, but partially replaced the H$_2$azo with the appropriate amount of H$_2$sti. The phase purity of the resulting single crystals of Zr-fcu-azo/sti-X% (X ≈ 10, 20, 30, 40, 50 or 60 stilbene molar percentage) was confirmed by powder X-ray diffraction (PXRD; Figure 2b). The crystals varied in color from bright orange to colorless (Figure S1) according to their constituent azo/sti ratio, which was confirmed by proton nuclear magnetic resonance ($^1$H NMR) of digested crystals (Figure S19, Tables S1, S3). Finally, N$_2$ sorption analysis performed at 77 K showed comparable uptake and type I isotherm for all different activated samples (Figure 2e). Indeed, all apparent A$_{BET}$ values ranged from 3000 m$^2$/g to 3065 m$^2$/g and total pore volumes, from 1.24 cm$^3$/g to 1.28 cm$^3$/g (Table 1). As expected, and similarly to reported values for comparable MOF systems, the pore size distributions (PSD, Figure 2h) of all Zr-fcu-azo/sti-X% are comparable: with a main pore population of ca. 18.4 Å attributed to the intrinsic porosity of the framework and a minor population of larger pores of ca. 22.4 Å attributed to some modulator related defects.

<table>
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<th>initial% sti</th>
<th>A$_{BET}$ (m$^2$/g)</th>
<th>V$_p$ (cm$^3$/g)</th>
<th>V$_t$ (cm$^3$/g)</th>
<th>A$_{BET}$ (m$^2$/g)</th>
<th>V$_p$ (cm$^3$/g)</th>
<th>V$_t$ (cm$^3$/g)</th>
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<td>1.26</td>
<td>1.31</td>
<td>Non-porous</td>
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**Ozonolysis of Zr-fcu-azo/sti-X%**. To selectively break the sti ligands, we packed ca. 50 mg of Zr-fcu-azo/sti-X% into a L- shaped glass tube, connected on one side to the ozonator (through a CaCl$_2$ humidity trap), and on the other side, to a vacuum pump (through a KI trap), to ensure a continuous flow of ozone through the column. The reaction was run at room temperature for 30 min. The ozonated Zr-fcu-azo/sti-X% materials maintained their high crystallinity after ozonolysis, as confirmed by PXRD (Figures S2-S10). $^1$H NMR of digested, Zr-fcu-azo/sti-X%_OZ confirmed the full conversion of sti into bdc and fba for all samples (Figure S20, Table S4). The 1:1 bdc/fba ratio observed for all Zr-fcu-azo/sti-X%_OZ also indicated the stoichiometric character of the reaction (Figures 2d, 3a). As expected, based on our preliminary results with Zr-fcu-azo, the content of azo ligand in the framework was not affected by ozonolysis, thus confirming its high selectivity towards olefin bonds.
Washing of Zr-fcu-azo/sti-X%_OZ. To remove the organic and inorganic fragments trapped in the pores of the Zr-fcu-azo/sti-X%_OZ, we washed the ozonated crude product in a 0.5 M solution of acetic acid in DMF. The washed samples did not exhibit any marked difference in crystallinity compared to the corresponding crude products, as confirmed by PXRD (Figure S10), with one exception: Zr-fcu-azo/sti-60%_OZ. The washing step revealed that the latter had indeed lost crystallinity, which we attributed to a partial collapse of its framework upon loss of 60% of its ligands via ozonolysis. In all cases, $^1$H NMR (Figures S21, S22 and Tables S5, S6) of the washed Zr-fcu-azo/sti-X% revealed the absence of fba and the persistence of some bdc, which could not be fully removed (repeated washing did not remove any additional bdc).

Porosity properties of washed, ozonated Zr-fcu-azo/sti-X%. After confirming that in the washed, ozonated Zr-fcu-azo/sti-X%, all the sti ligands had been broken (Figure S20), most of the resulting organic fragments had washed out (Figure S21) and the high crystallinity had remained (Figure S10c), we submitted these materials to N$_2$ sorption tests at 77 K (Figure 2f, Table 1). As expected, Zr-fcu-azo/sti-10% performed very similarly to the pre-ozonated material, showing only slightly higher values for apparent $A_{BET}$ (3115 m$^2$/g vs. 3060 m$^2$/g) and total pore volume (1.32 cm$^3$/g vs. 1.28 cm$^3$/g). Based on the Dubinin-Radushkevich (DR) equation,$^{27}$ the relative proportion of micro-porosity to the total pore volume was estimated to be 96% (Figure 2g). Although the washed ozonated Zr-fcu-azo/sti-20% showed a nearly identical value for this parameter (95%), its isotherm exhibited a shoulder/step characteristic from bigger pores, giving insights into the modified nature of the microporosity. The isotherms for the washed ozonated Zr-fcu-azo/sti-30% and Zr-fcu-azo/sti-40% did not show the type I shape of the isotherms of the corresponding starting MOFs, but rather they exhibited a small hysteresis on the desorption branch, characteristic of mesoporosity (Figures S29, S30). Notably, ozonolysis treatment reduced the apparent $A_{BET}$: for the washed ozonated Zr-fcu-azo/sti-30%, 2680 m$^2$/g (vs. 3000 m$^2$/g); and for the washed ozonated Zr-fcu-azo/sti-40%, 2580 m$^2$/g (vs. 3040 m$^2$/g) (Table 1). However, in both cases, this decrease was offset by a major increase in the proportion of pores with mesoscale dimensions, as derived from the DR equation (Figure 2g and Table S9): for the washed ozonated Zr-fcu-azo/sti-30%, 15% (vs. 2%); and for the washed ozonated Zr-fcu-azo/sti-40%, 24% (vs. 3%). Likewise, the total pore volume also increased in both cases: for the washed ozonated Zr-fcu-azo/sti-30%, 1.36 cm$^3$/g (vs. 1.24 cm$^3$/g) and for the washed ozonated Zr-fcu-azo/sti-40%, 1.44 cm$^3$/g (vs. 1.27 cm$^3$/g). Similar results were observed in the washed ozonated Zr-fcu-azo/sti-50%, which exhibited the most extreme change relative to the corresponding starting MOF: the contribution of the mesoporosity reached 42% of the total porosity (Figure 2g and Table S9), and the hysteresis characteristic of mesoporosity was clearly visible (Figure S31); consequently, its $A_{BET}$ was only 1860 m$^2$/g (vs. 3065 m$^2$/g). Note that additional washing treatment did not affect the sorption properties (Figure S40a), excluding the hypothesis of structural etching sometimes observed through chemical treatment.$^{18e, 28}$

The modification of the micro- vs meso-porosity ratio was also reflected by studying the PSD (Figure 2i). If no noticeable change was observed for the washed ozonated Zr-fcu-azo, a subtle appearance of pores of ca. 25 Å and above was noted for the washed ozonated Zr-fcu-azo/sti-10%. The washed ozonated Zr-fcu-azo/sti-20% and 30% showed a clear decrease of the initial main pore size of ca. 18.4 Å and an increase of the second population at ca. 22.4 Å, confirming that micropores are merged into wider pores. For these two samples, the formation of pores bigger than 25 Å was also visible. Finally, the initial main pores of ca. 18.4 Å completely vanished in washed and ozonated Zr-fcu-azo/sti-40% and 50%. In these cases, the pores were mainly observed at ca. 22.4 Å and spread over a wide range up to 50 Å. Albeit non observed through PXRD, the pore population centered at 22.4 Å could be attributed to the existence of non-ordered reo motifs, resulting from the removal of inorganic clusters, as hypothesized in some comparable systems.$^{18e}$
Based on these pioneering results, we envisioned extending our method to a more challenging system. We considered that 4,4'-biphenyldicarboxylic acid (H₂bpdc) and 1,4-phenylenediacrylic acid (H₂pdac), found in the two Zr-fcu-MOFs UiO-67 and NU-801, would be a suitable ligand pair, given their geometric (length \( \approx 11.3 \) Å) and chemical properties (vide supra). A particularity of H₂pdac is its two olefin bonds (Figure 3b), which we reasoned would enable cutting of the ligand into three distinct parts, rather than two, as in H₂sti (Scheme S2, Figure S23). In addition, we hypothesized that removal of the central part of the ligand could generate terephthalaldehyde, which could be sublimated under vacuum at moderate temperature -for example, during ozonolysis and/or during a classical pre-sorption activation procedure (Figure 3). That turned out to be correct and, by obviating the washing step, represented a major improvement to our method.
Figure 3. Post-synthetic generation of mesoporosity in MOFs. a) In Zr-fcu-azo/sti MOFs, removal of the sti ligand requires two steps: ozonolysis (to break the ligand) and washing. b) In contrast, in Zr-fcu-bpdc/pdac MOFs, removal of the pdac ligand requires only one step: ozonolysis (to break the ligand) and concomitant sublimation (to remove the central core). This is made possible by the two olefin groups in this ligand.

Synthesis of Zr-fcu-bpdc/pdac-X%. Multivariate Zr-fcu-MOFs containing specific ratios of bpdc/pdac, from 100:0 (UiO-67) to 0:100 (NU-801) were prepared from mixtures of ZrCl₄, H₂bpdc and H₂pdac dissolved in DMF in the presence of trifluoroacetic acid, and heated at 120 °C for 72 h in a scintillation vial. The samples were washed with DMF overnight and then soaked in toluene for 3 days. The phase purity of the single crystals of Zr-fcu-bpdc/pdac-X% (X = 10, 20, 30, 40% pdac molar percentage) was confirmed by PXRD (Figure 4b), ¹H NMR of digested crystals (Figure S24, Table S7) and N₂ sorption analysis performed at 77 K (Figure 4e and Table 2). The sorption isotherms for Zr-fcu-bpdc/pdac-X% were comparable (2210 m²/g < A_BET < 2310 m²/g and 0.92 cm³/g < Vt < 0.96 cm³/g). We observed a slight shoulder at a P/P₀ value of ca. 0.1, which we attributed to a higher number of initial defects in the structure, likely due to our use of a slightly different synthetic procedure than for Zr-fcu-azo/sti-X%. This is in fact reflected by the estimation of the PSD (Figure 4h), in which all samples presented two main contributions at ca. 15.2 Å and 18.4 Å; the bigger size being attributed (similarly as for Zr-fcu-azo/sti system, vide supra) to modulator-related defects.

Table 2. Apparent BET area, microporous and total pore volume of Zr-fcu-bpdc/pdac-X%, before and after ozonolysis.

<table>
<thead>
<tr>
<th>% pdac initial</th>
<th>Starting MOF</th>
<th>Ozonated MOF</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>A_BET (m²/g)</td>
<td>Vµ (cm³/g)</td>
</tr>
<tr>
<td>0</td>
<td>2250</td>
<td>0.91</td>
</tr>
<tr>
<td>10</td>
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</tr>
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</tr>
<tr>
<td>30</td>
<td>2245</td>
<td>0.93</td>
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</tbody>
</table>
Ozonolysis of Zr-fcu-bpdc/pdac-X%. Analogously to the previous MOF pair (vide supra), the crystallinity and porosity of UiO-67 were unaffected by ozonolysis (Figures 4a, S11), but cleavage of the pdac ligands in NU-801 led to collapse of its framework and loss of its crystallinity and porosity (Figures 4a, S16). Having demonstrated this, we proceeded with ozonolysis of the Zr-fcu-bpdc/pdac-X% MOFs. \(^{1}\)H-NMR of all Zr-fcu-bpdc/pdac-X%_OZ highlighted the absence of any remaining pdac (Figures S25, Table S8). Moreover, after the Zr-fcu-bpdc/pdac-X%_OZ were activated at 85 °C, a white solid was recovered in the cold trap (77 K) placed between the MOFs and the pump. This solid was confirmed by \(^{1}\)H NMR to be terephthalaldehyde (Figure 4d), which, after cleavage of pdac, we were able to directly extract from the framework via sublimation, as anticipated.

Porosity of Zr-fcu-bpdc/pdac-X%_OZ. To investigate the effects of removing the central core of pdac on the porosity of the Zr-fcu-bpdc/pdac-X%_OZ, we performed N\(_2\) sorption experiments on these MOFs (Figure 4f, Table 2). Upon extracting terephthalaldehyde from the backbone of Zr-fcu-bpdc/pdac-10%, the porosity was enhanced, as demonstrated by an increase in the apparent \(A_{\text{BET}}\) (from 2295 m\(^2\)/g to 2400 m\(^2\)/g). In this sample, besides the evolution of microporous pore volume (\(V_\mu\)) (from 0.95 cm\(^3\)/g to 0.97 cm\(^3\)/g) was not significant, a more pronounced shoulder in the isotherm reflected a transformation of the microporosity. The emergence of some mesoporosity was revealed by the DR equation, with a contribution of 11 % (vs. 1%) to the total porosity. In contrast, upon further removal of pdac pillars, from Zr-fcu-bpdc/pdac-20%_OZ and Zr-fcu-bpdc/pdac-30%_OZ, the apparent \(A_{\text{BET}}\) decreased sharply: for Zr-fcu-bpdc/pdac-20%_OZ (\(A_{\text{BET}} = 1870\) m\(^2\)/g vs. 2210 m\(^2\)/g) and for Zr-fcu-bpdc/pdac-30%_OZ (\(A_{\text{BET}} = 1730\) m\(^2\)/g vs. 2245 m\(^2\)/g). The loss in microporosity for Zr-fcu-bpdc/pdac-20%_OZ (\(V_\mu = 0.73\) cm\(^3\)/g vs. 0.91 cm\(^3\)/g) and for Zr-fcu-bpdc/pdac-30%_OZ (\(V_\mu = 0.65\) cm\(^3\)/g vs. 0.93 cm\(^3\)/g) was offset by a major increase in the proportion of pores with mesoscale dimensions, as derived from the DR equation (Figure 4g, Table S10): for Zr-fcu-bpdc/pdac-20%_OZ, 26 % (vs. 4%) and for Zr-fcu-bpdc/pdac-30%_OZ, 38 % (vs. 1 %). Likewise, the total pore volume also slightly increased in both cases: for Zr-fcu-bpdc/pdac-20%_OZ, 0.99 cm\(^3\)/g (vs. 0.95 cm\(^3\)/g) and for Zr-fcu-bpdc/pdac-30%_OZ, 1.05 cm\(^3\)/g (vs. 0.94 cm\(^3\)/g). As expected, Zr-fcu-bpdc/pdac-40%, which, upon ozonation, lost its crystallinity (Figure S15) due to structural collapse caused by removal of 40% of its ligands, was non-porous. Overall, the change in the porosity in Zr-fcu-bpdc/pdac-X% paralleled that of Zr-fcu-azo/sti-X%. However, in the former, these changes occurred at lower percentages of ligand removal—presumably due to a greater number of defects in the starting MOFs.

The estimation of the PSD in this system also confirmed the evolution of the porosity towards mesopores (Figure 4i). While ozonolysis of UiO-67 did not affect the pore system of the pristine MOF, the appearance of pores distributed between 21.6 Å and 27.4 Å was clearly visible for the ozonated Zr-fcu-bpdc/pdac-10%, -20% and -30%. In addition, the two initial main pore populations decreased in ozonated Zr-fcu-bpdc/pdac-20% and Zr-fcu-bpdc/pdac-30%. In these two last samples, the porosity population was extended up to ca. 50 Å, further confirming the fusion of micropores into novel mesopores.
Figure 4. a) N₂ sorption isotherms for UiO-67 and NU-801 before and after ozonolysis. b) PXRD diagrams for Zr-fcu-bpdc/pdac-X%. c) PXRD diagrams for Zr-fcu-bpdc/pdac-X%_OZ. d) ¹H NMR spectra of the sublimated terephthalaldehyde recovered by heating at 85 °C under vacuum after ozonolysis. e) N₂ sorption isotherms (77 K) for Zr-fcu-bpdc/pdac-X%. f) N₂ sorption isotherms (77 K) for Zr-fcu-bpdc/pdac-X%_OZ. g) Plot showing the overall stability of the total pore volume and evolution of the relative contribution of mesoporosity vs. microporosity to total uptake, in Zr-fcu-bpdc/pdac-X%_OZ, as calculated with the Dubinin-Radushkevich equation. h) Pore size distribution estimated by DFT for Zr-fcu-bpdc/pdac-X%. i) Pore size distribution estimated by DFT for Zr-fcu-bpdc/pdac-X%_OZ%.

CONCLUSIONS

In conclusion, we have developed a selective and quantitative method for post-synthetic modification of MOF porosity via ozonolysis. Through removal of organic ligands, our approach causes micropores to fuse into larger (mesoscale) pores in a controlled fashion and without any loss in total porosity. It selectively breaks the olefin bonds in organic ligands without affecting other organic moieties and can be applied for short reaction times (ca. 30 min) under mild conditions (room temperature). Moreover, we have demonstrated that ozonolysis can be used as a novel tool to selectively remove, and recover, specific fragments of molecules/frameworks through solid-gas manipulation.

EXPERIMENTAL SECTION

Materials and Instruments

Reagents. 4,4′-stilbenedicarboxylic acid (H₂sti), 4,4′-biphenyl dicarboxylic (H₂bpdc), and azobenzene-4,4′-dicarboxylic acid (H₂azo) were purchased from TCI Europe. Acetic acid, 1,4-phenylenediacrylic (H₂pdac), trifluoroacetic acid (tfa), zirconium chloride (ZrCl₄), L-proline, hydrochloric acid, potassium iodide and anhydrous granular CaCl₂ (2-6 mm) were purchased from Sigma-Aldrich (Merck). N,N-dimethylformamide (DMF) and acetone were obtained from Fisher Chemical. Toluene was purchased from Acros Organics. All the reagents and solvents were used as
Pyrex tubes (ext \( \Omega = 7 \text{ mm} \); in \( \Omega = 5.5 \text{ mm} \); length = 150 mm) were purchased from Vidrasa S.A. The tubes were bent into an L-shape with a flame torch.

**Optical Microscopy** images were captured by a Zeiss Axio Observer Z1m Optical Microscope.

**PXRD diagrams** were recorded on an XPert PRO MPD analytical diffractometer (Panalytical) at 45 kV, 40 mA using CuK\( \alpha \) radiation (\( \lambda = 1.5406 \text{ Å} \)).

\(^1\text{H} \text{ NMR spectra} \) were recorded on a Bruker Avance 400 NMR spectrometer (static magnetic field: 11.7 T) operating at Larmor frequencies of 400 MHz (\(^1\text{H} \)).

**Ozonolysis treatment** was performed using an ozone generator (N1668A from Ozonotec) generating a constant flow of 10.4 mmol/h at room temperature.

**Volumetric N\(_2\) sorption isotherms** were collected at 77 K (N\(_2\)) using an ASAP 2020 HD (Micromeritics). Temperature was controlled by using a liquid nitrogen bath. Total pore volume (\( V_\text{t} \)) was calculated at \( P/P_0 = 0.95 \), and microporous volume (\( V_\mu \)) was calculated using the Dubinin-Radushkevich equation. Pore size distribution was estimated using a density functional theory (DFT) model (N\(_2\) – cylindrical pores – oxide surface) implemented in the Microactive 4.00 software with a regularization factor of 0.01.

**Synthesis.**

**Synthesis of Zr-fcu-azo/sti-X\%**. 4,4’-stilbenedicarboxylic acid (H\(_2\)sti, 0.44-x mmol) and azobenzene-4,4’-dicarboxylic acid (H\(_2\)azo, x mmol) were suspended in 10 mL of DMF and sonicated for 30 min in a 23-mL scintillation vial. Then 5 mL of a DMF solution of ZrCl\(_4\) (105 mg, 0.45 mmol), L-proline (250 mg, 2.17 mmol) and 45 µL conc. HCl were transferred into the vial and vortexed for 1 minute. After heating at 120 °C for 16 h, the reaction afforded orange octahedral crystals of Zr-fcu-azo/sti-X\% (see Supporting Information for more details about the H\(_2\)azo:H\(_2\)sti ratio in this synthesis).

**Synthesis of Zr-fcu-bpdc/pdac-X\%**. ZrCl\(_4\) (200 mg, 0.86 mmol), 1,4-phenylenediacrylic acid (H\(_2\)pdac, x mmol) and biphenyl-4,4’-dicarboxylic acid (H\(_2\)bpdc, y mmol) were dissolved by sonication in 10 mL of DMF in the presence of trifluoroacetic acid (tfa, 1 mL) in a 23-mL scintillation vial. After heating at 120 °C for 72 h, colorless octahedral crystals of Zr-fcu-bpdc/pdac-X\% were obtained (see Supporting Information for more details about the H\(_2\)bpdc/H\(_2\)pdac ratio in the synthesis).

**Sample activation.**

**Activation of Zr-fcu-azo/sti-X\%**. Crystals of Zr-fcu-azo/sti-X\% were soaked in 15 mL of DMF (refreshed after 4 h) and then subjected to solvent exchange in 15 mL of acetone (refreshed after 4 h). The supernatant was removed after decantation, and the crystals were dried at 65 °C. Sample purity was verified by PXRD (Figure 2b), and experimental azo/sti ratios were analyzed by \(^1\text{H} \text{ NMR} \) of the digested samples (Figure S19, Table S3). Crystals of Zr-fcu-azo/sti-X\% were then transferred into a glass sorption cell and activated under dynamic vacuum at 85 °C prior to sorption experiments.

**Activation of Zr-fcu-bpdc/pdac-X\%**. Crystals of Zr-fcu-bpdc/pdac-X\% were soaked in 10 mL of DMF for 24 and then subjected to solvent exchange for 72 h in 20 mL of toluene (refreshed twice per day). The supernatant was removed by filtration. The sample purity was verified by PXRD (Figure 4b), and the experimental bpdc/pdac ratios were analyzed by \(^1\text{H} \text{ NMR} \) of the digested samples (Figure S24, Table S7). Crystals of Zr-fcu-bpdc/pdac-X\% were then transferred into a glass sorption cell and activated under dynamic vacuum at 85 °C prior to sorption experiments.

**Ozonolysis treatment.**

**Ozonolysis of the organic ligands.** In a typical procedure, 30 mg of the H\(_2\)azo, H\(_2\)sti, H\(_2\)bpdc or H\(_2\)pdac were dissolved/dispersed in 20 mL of DMF. The solution was then transferred to a 50 mL
three-necked round bottom flask and was cooled to 0 °C using an ice-water bath. A constant ozone flux was blown into the solution through a glass pipette for 30 minutes under stirring. The crude reaction was then allowed to warm up to room temperature and the solvent was removed in vacuo. The crude residue was collected and analyzed by 1H NMR. 1H NMR analysis confirmed the inert character of H2azo and H2bpdc, while revealing complete conversion, within 30 minutes, of the active linkers H2sti and H2pdac.

**Ozonolysis of Zr-fcu-azo/sti-X% and Zr-fcu-bpdc/pdac_X%**. A column of Zr-fcu-azo/sti-X% or Zr-fcu-azo/sti-X% (50 mg to 100 mg) was loaded over a cotton stopper in an L-shaped Pyrex tube. The glass tube was connected to the solid-gas post synthetic modification set-up and purged under vacuum for 5 min. The ozone gas flow (10.4 mmol/h) was passed through the MOF column for 30 min. After 1-2 min, the aqueous potassium iodide solution turned yellow, indicating the whole MOF column had been exposed to ozone atmosphere. The recovered Zr-fcu-azo/sti-X%_OZ or Zr-fcu-bpdc/pdac-X%_OZ samples were further characterized by PXRD (Figures S10b, S17b) and 1H NMR (Figures S20, S25, Table S4, S8).

**Acetic acid wash.**

About 50 mg of Zr-fcu-azo/sti-X%_OZ were vortexed for a few seconds in 20 mL of a 0.5 M acetic acid solution in DMF and left at room temperature for 24 h. The supernatant was removed and analyzed by 1H NMR after decantation (Figure S22, Table S6). The crystals were washed twice with 20 mL of acetone. The acetone was removed after decantation, and the crystals were dried at 65 °C. The sample purity was verified by PXRD (Figure S10c), and the experimental organic content was analyzed by 1H NMR of the digested samples (Figures S21, Table S5). Crystals of washed ozonated Zr-fcu-azo/sti-X% were then transferred into a glass sorption cell and activated under dynamic vacuum at 85 ºC prior to the sorption experiments. Note that the Zr-fcu-bpdc/pdac-X% samples did not require this additional treatment to reveal their tuned porosity potential (Figures 3, S40b).

**HF-Digestion methodology.**

In a typical procedure, 10 mg of MOF were placed in an Eppendorf tube, to which 120 µL of a 5% aq. HF solution were added. The mixture was sonicated for 15 minutes, and then the slurry was placed in a 110 ºC sand bath inside a fume hood to evaporate overnight. The crude solid samples were highly soluble in DMSO-d6 without any need for addition of acid. Note that attempts with standard digestion methodologies were not successful, even when using the same quantities of HF and solvent. Therefore, the evaporation step is crucial for proper sample digestion. Moreover, the digested solutions obtained with our methodology did not require any specific (e.g. HF-resistant) PTFE NMR tubes.

**ASSOCIATED CONTENT**

**Supporting Information.** Detailed synthetic procedures, microscopy, PXRD, 1H NMR, additional sorption data. This material is available free of charge via the Internet at http://pubs.acs.org.

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The manuscript was written through contributions of all authors.
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ABBREVIATIONS

1H NMR, proton nuclear magnetic resonance; \textit{A}BET, Brunauer-Emmett-Teller area; bdc, terephthalic acid; DFT, density functional theory; DMF, \textit{N},\textit{N}'-dimethylformamide; DR, Dubinin-Radushkevich; fba, formylbenzoic acid; H\textsubscript{2}azo4,4'-azobenzene dicarboxylic acid; H\textsubscript{2}bpdc, 4,4'-biphenyldicarboxylic acid; H\textsubscript{2}pdac, 1,4-phenylenediacrylic acid; H\textsubscript{2}sti, 4,4'-stilbene dicarboxylic acid (H\textsubscript{2}sti); HKUST, Hong-Kong university of science and technology; IRMOF, isoreticular metal-organic framework; MIL, material institute Lavoisier; MOF, metal-organic framework; MTV, multivariate; OZ, ozonated; PSD, pore size distribution; PSM, post synthetic modification; PXRD, powder X-ray diffraction; tfa, trifluoroacetic acid; UiO, Universitetet i Oslo; W, washed.

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