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The Effect of Benzoic Acid as Modulator in UiO-66 Structure: an Experimental and Computational Study Cesare Atzori^a, Greig C. Shearer^b, Lorenzo Maschio^c, Bartolomeo Civalleri^c, Francesca Bonino^a, Carlo

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Abstract

The identification and quantification of defects is undoubtedly a thorough challenge in the characterization of "defect engineered" metal-organic frameworks (MOFs). UiO-66, known for its exceptional stability and defect tolerance, has been a popular target for defect engineering studies. Herein, we show that synthesizing UiO-66 in the presence of an excess of benzoic acid is a reliable method for obtaining UiO-66 samples with a very high concentration of missing cluster defects, allowing to modulate specific properties (i.e. surface area and hydrophobicity). This was elucidated by a multitechnique marriage of experimental and computational methods: a combination of PXRD, dissolution/¹H NMR spectroscopy, and N₂ sorption measurements were used to quantify the defect loading, while vibrational spectroscopies (FTIR and Raman) allowed us to unequivocally identify the defect structure by comparison with DFT simulated spectra and visual analysis of the computed vibrational modes.

1. Introduction

Among the numerous classes of porous materials, metal-organic frameworks (MOFs) stand out for their structural and chemical diversity, exhibiting a wide range of well-defined pore size distributions and chemical functionalities^{1,2}. This variety endows MOFs with incredible potential in many areas of technological relevance e.g. adsorption³ and catalysis⁴, but their real-world applicability in these areas has thus far been hindered by serious drawbacks such as poor stability, particularly to moisture^{5–9}. Much attention has therefore been given to relatively stable subclasses of MOFs, e.g. zirconium (IV) MOFs, particularly those whose clusters are based on a Zr₆ octahedral core^{10–15}. The most widely studied material among these "Zr₆ MOFs" is UiO-66 (Figure 1), whose Zr₆-oxyhydroxide clusters are 12-fold connected by 1,4-benzenedicarboxylate (BDC, a linear ditopic linker), resulting in a highly symmetric face-centered cubic (fcu) structure¹².

While UiO-66 is indeed exceptionally thermally, chemically, and mechanically stable, its performance in commonly proposed applications (e.g. adsorption, separation, and catalysis) is limited by its relatively modest porosity and lack of chemical functionality. In recognition of these shortcomings, many researchers have opted to instead focus on isoreticularly expanded and/or functionalized UiO-66 derivatives (e.g. UiO-67, UiO-66-NH₂)

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which offer improvements in these areas^{16–33}. However, these derivatives generally harbor two rather significant drawbacks with respect to UiO-66: lower stability^{18,21,23,24,34,35} and higher cost. The bulk of their expense can be attributed to the functionalized and/or extended organic linker(s), many of which are not even commercially available, necessitating the development of in-house synthesis procedures³⁶ which can take months (or longer) to optimize unless the molecule has already been reported in the literature. It is therefore clear that an alternative approach for tuning UiO-66's porosity and functionality (and thus, performance) is desired.

One such alternative approach is "defect engineering", a term often used as shorthand for the practice of manipulating defects via controlled, synthetic means^{37–46}. This approach has proven to be particularly effective for UiO-66^{16,37,38,43–72}, whose structure is famed for its ability to tolerate an extraordinarily high concentration of defects while retaining much of its stability³⁷ (except for certain cases where the thermal stability has been shown to be heavily compromised⁷³, and one sample (50Benz-HA in ref. [⁴⁹]) which was found to have reduced water stability). We stress that this fascinating trait is not just an academic curiosity; many studies have shown that defects significantly enhance UiO-66's porosity^{37,38,43,44,46,48,49,53,58,59,61,73,74} and reactivity^{44,47,49,50,57,60,62}, leading to better performances in a wide variety of applications^{38,48–50,53,56–59,62–67,75,76}. Such positive demonstrations have encouraged many researchers to seek out strategies to gain synthetic control over the defects in the UiO-66 framework.

To this end, the most common method is to synthesize UiO-66 in the presence of additives known as "modulators^{68,69,77–80}". Many types of modulator (e.g. HCl^{61,63}, HF⁶⁸, NH₄OH⁷⁹, amino acids⁶⁹) are known to promote the formation of defects, but monocarboxylic acid "coordination modulators" (e.g. acetic acid, formic acid, benzoic acid, etc.) are the most widely studied and thus, best understood^{37,38,46–50,52–55,58,59,77}. In recent work^{37,49}, we proved that monocarboxylic acid modulation promotes the formation of "missing cluster defects" whose charge and coordination deficiencies are compensated by modulator ligands (in their carboxylate form, see benzoate ligands in Figure 1). More importantly, we showed that these defects can massively enhance the porosity and reactivity of the material^{49,73}, and that new, otherwise elusive functionalities can be incorporated by post-synthetically exchanging the defect-compensating ligands⁴⁹.

Another important (albeit less striking) aspect of our systematic investigations is that they have enabled us to objectively compare different monocarboxylic acid modulators for their ability to introduce defects to the UiO-66 framework, a property which we hereafter refer to as their "defect-inducing potency". Based on such comparisons, we have identified trifluoroacetic acid³⁷ and benzoic acid⁴⁹ as the most potent modulators we have investigated thus far, and have ascribed their effectiveness to a combination of acidity³⁷, solubility⁴⁹, and steric⁴⁹ effects. While the defect-inducing potency of the two modulators are roughly equivalent, we now harbor a strong preference towards benzoic acid for the following reasons: (1) trifluoroacetic acid (a strong acid) readily catalyzes the hydrolysis of DMF during UiO-66 synthesis, generating formic acid which then goes on to act as a modulator in its own right, resulting in a material whose defects are compensated by a combination of trifluoroacetate and formate ligands³⁷. This increases the complexity and heterogeneity of the material in a manner which is probably

not entirely reproducible, and is much less of an issue when benzoic acid (a much weaker acid) is employed as the modulator⁴⁹; (2) trifluoroacetic acid (B.P. = 72.4 °C) is a relatively volatile molecule which may therefore evaporate from the solution during UiO-66 syntheses. We speculate that this volatilization could occur to different extents and rates from one synthesis batch to the next, potentially affecting reproducibility - a situation which is completely avoided by using the non-volatile benzoic acid (B.P. = 249.2 °C) as the modulator; (3) In our experience, trifluoroacetic acid modulated UiO-66 syntheses often provide poor yields, a negative outcome is much less severe when benzoic acid is employed as the modulator; (4) trifluoroacetic acid modulated syntheses tend to yield UiO-66 samples with very poorly defined crystal morphology and size^{37,50}, while benzoic acid modulated syntheses provide reasonably monodisperse octahedral crystals^{20,49,59,77}; (5) benzoic acid is much cheaper than trifluoroacetic acid.

It is important to keep in mind that all of the findings discussed thus far were backed up by a large amount of convincing experimental evidence, something which we feel is especially vital for defect engineering pursuits since disorder is notoriously challenging to characterize. In acknowledgement of this challenge, researchers have sought to develop methods that are able to detect, quantify, and evaluate the nature of the defects in different UiO-66 samples, preferably with routine characterization techniques available in most home labs. To this end, most research groups (including our own) employ some combination of PXRD^{37,49,52,54,73}, TGA^{37,49,50,53–55,58,61,68,69,73}, dissolution/ 1 H NMR spectroscopy 37,47,49,53,59,69 , and N_{2} adsorption measurements $^{37,38,47-49,53,58,59,61,63,68,69,73}$ (sometimes in conjunction with simulations^{37,49,61,67,73}). While indirect, this approach has proven to be valuable for detecting defects and assessing their concentration via simple semi-quantitative data analysis methods^{37,49}. Conversely, vibrational spectroscopies (FTIR, ATR, Raman) have thus far had only a peripheral, qualitative role for characterizing the defectivity of UiO-66. For example, they have been used to detect the presence of different defect compensating ligands (i.e. deprotonated modulator molecules^{37,49}), superfluous hydroxyl groups^{81,82} (also thought to terminate defects in some cases^{57,60,61,65,70-72}), and to speculate on defectivity by noting differences between experimental spectra and that simulated from the ideal, defect-free UiO-66 structural model⁷³. This rather limited use is somewhat surprising given that vibrational spectroscopies are traditionally very well suited to characterizing defects^{83,84}, and it was this realization that motivated us to investigate these techniques further herein.

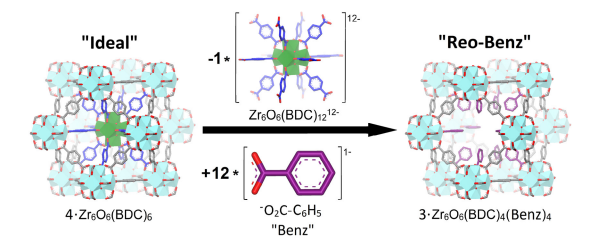


Figure 1 Illustration showing how the structure of a "Reo-Benz" defect (see main text) differs from that of ideal, defect-free UiO-66. Hydrogen atoms have been omitted for the sake of clarity.

To this end, we synthesized a series of 5 defective UiO-66 samples, each obtained in the presence of different amounts of benzoic acid (our preferred defect engineering modulator, see reasons above). All of the samples (given the general name "xBenz", where x is the molar equivalents of benzoic acid used in the synthesis, with respect to ZrCl₄) were then characterized by means of Raman and FTIR spectroscopies, as well as the methods more commonly used for characterizing defects in UiO-66 (PXRD, dissolution/¹H NMR, and nitrogen sorption measurements). In accordance with our previous studies^{37,49}, the results from the conventional characterization methods show that: 1) using benzoic acid as a modulator yields UiO-66 samples with a high concentration of missing cluster defects with benzoate as the compensating ligand (so-called "Reo-Benz" defects (see Figure 1) due to their RCSR⁸⁵ denoted topology (reo)), and; 2) the concentration of these defects systematically increases as increasing amounts of benzoic acid are used in the synthesis. Although we are fully convinced with these conclusions, we speculate that some of the MOF community may still be skeptical due to the indirect manner in which these methods probe defectivity. This potential criticism is opposed by our vibrational spectroscopy results, which, when compared with simulated spectra, provide direct and unequivocal evidence for our conclusions (and those of our previous studies^{37,49}). Perhaps more importantly, this paper demonstrates that vibrational spectroscopies can be a powerful tool for characterizing defects in MOFs, especially when the experimental results are compared with spectra simulated from models of proposed defect structures^{86,87}.

2. Experimental

2.1 Materials synthesis

2.1.1 Defective UiO-66 Samples

The 5 defective UiO-66 samples (named 10Benz, 20Benz, 30Benz, 40Benz, and 50Benz) were synthesized by the same method, albeit with different amounts of benzoic acid as a modulator (see Table 1). First, the synthesis solutions were prepared by sequentially adding 1.680 g ZrCl₄ (7.209 mmol), 0.173 mL H₂O (9.603 mmol), benzoic acid (quantities given in Table 1) and 1.198 g 1,4-benzenedicarboxylic acid (H₂BDC, 7.211 mmol) to a 1 L Erlenmeyer flask containing 413.2 mL of warm (ca. 70 °C) N,N'-dimethylformamide (DMF, 5336 mmol) under constant magnetic stirring.

Table 1. Quantities of benzoic acid used in the synthesis of the 5 defective UiO-66 samples. The names of the samples are derived from the benzoic acid:ZrCl₄ molar ratio in their respective synthesis solutions.

Sample	Amount of benzoic acid used in synthesis		
10Benz	8.807 g (72.12 mmol)		
20Benz	17.613 g (144.23 mmol)		
30Benz	26.420 g (216.34 mmol)		
40Benz	35.226 g (288.45 mmol)		
50Benz	44.033 g (360.57 mmol)		

Once the reagents had fully dissolved, the stirring bars were removed and watch glasses were placed over the mouths of the flasks as a loose cover. The covered synthesis solutions were then placed in an oven preheated to 120 °C and were allowed to react over a period of 24 hours before applying the purification steps given in Section 2.1.3.

2.1.2 "Defect-free" UiO-66 (Ref.)

The "defect-free" UiO-66 sample (Ref.) was synthesized via an optimized version^{37,49} of a procedure originally outlined by Serre and co-workers⁸⁸ and later promoted by us⁷³. First, the synthesis mixture was prepared by sequentially dissolving 3.781 g ZrCl₄ (16.22 mmol), 2.865 ml 35 % HCl (32.45 mmol), and 5.391 g H₂BDC (32.45 mmol) in 97.40 ml of warm (ca. 70 °C) N,N'-dimethylformamide (DMF, 1258 mmol) under constant magnetic stirring. Once all reagents had fully dissolved, the mixture (sans the magnetic stirrer) was transferred to a 200 mL Teflon liner, sealed in a stainless steel autoclave, and placed in an oven preheated to 220 °C. After 24 hours of reaction at this temperature, the autoclave was removed from the oven and rapidly cooled by dropping it into a bucket of cold tap water for 30 minutes before applying the purification steps given in Section 2.1.3.

2.1.3 Purification procedures

After all reactions, the resulting microcrystalline powder precipitates were separated from their synthesis solutions via centrifugation and washed by shaking them overnight in 80 mL of fresh DMF. Three further

separation/washing cycles were performed the next day, albeit for a shorter duration (ca. 2 hours per wash) before the washed products were separated by centrifugation, dried overnight in an oven set to 60 °C, and ground with a mortar and pestle. The samples were then subjected to a further "activation" procedure to remove the DMF guest molecules from their pores. This was achieved by heating them at 200 °C for 24 hours in a conventional oven, a simple treatment which we have found to be very effective for UiO-66^{37,49}. Ref. sample was subjected to a further heat treatment (dubbed "calcination"), in which unidentified non-volatile organic impurities³⁷ are removed via a 2-step program on a multi-step muffle furnace. In the first step, a 5 °C min⁻¹ ramp was used to heat the material from room temperature to 200 °C, at which it was held for 10 minutes. A 0.5 °C min⁻¹ ramp was then used between 200 to 270 °C, where the temperature was held for 70 hours. After calcination, the product was allowed to cool to room temperature naturally.

One should note that the activated/calcined samples are not completely empty as they adsorb a small amount of atmospheric water vapor when allowed to cool to room temperature in air. The theoretical yields of the procedures (discounting adsorbed water) is 2 grams for the defective samples and 4.5 grams for the defect-free one.

2.2 Experimental methods

2.2.1 Powder X-Ray Diffraction (PXRD)

Samples were prepared for measurement by dispersing 30 mg of sample on a flat, glass plate PXRD sample holder with a diameter of 2.5 cm. A stretched piece of plastic film was then used to spread, flatten and hold the sample in position for measurement. The plastic film is evident in the PXRD patterns, appearing as two broad peaks covering 2θ ranges of ca. 20- 22° and 23- 24° .

PXRD patterns (Cu-K α radiation, $\lambda = 1.5418$ Å, 2θ range = 2-50°, timescale = 1, resulting in a *d-spacing* down to 1.82 Å) were collected in reflectance Bragg-Brentano geometry with a Bruker D8 Discovery diffractometer equipped with a focusing Ge-monochromator and a Bruker LYNXEYE detector.

The relative intensity of the broad peak ($Rel(I)_{B.P.}$) have been extracted from the patterns by fitting with a peak-shaped function (a Pseudo-Voigt or a Pearson VII, depending on which one gave the lowest error) and normalizing the area of the fitted peak dividing by the intensity (measured in the same manner of the broad one) of 3 UiO-66 key reflections (the (100), (200) and (600)). A detailed description of this method can be found in the SI, Section 4 of the present paper and in two precedent works^{37,49}.

2.2.2 Dissolution / ¹H NMR Spectroscopy

Samples were prepared by weighing 20 mg of the MOF sample of interest into an NMR tube and adding 600 µL of 1M NaOH (in D₂O). The NMR tube was then capped and the contents were mixed by inverting the tube 2-3 times. The MOF was then left to digest over a period of 24 hours before measurement. This hydroxide based procedure dissolves only the organic portion of the MOF (linker, modulator, solvent etc.), while the inorganic content is converted into ZrO₂ which sinks to the bottom of the NMR tube and does not influence the spectra.

Liquid ¹H NMR spectra were obtained with a Bruker Avance DPX-300 NMR Spectrometer (300 MHz). The relaxation delay (d1) was set to 20 seconds to improve the reliability of integration and thus, the accuracy of the benzoic acid:BDC molar ratios $(\frac{n_{Benz}}{n_{BDC}})$ determined from the results. The number of scans (ns) was 64. Molar ratios were extracted from these spectral data using a procedure reported in the supporting information of the present manuscript (Section 6) and of two precedent works^{37,49}.

2.2.3 Nitrogen Sorption Measurements at 77K

Nitrogen sorption measurements were performed on a BelSorp mini II instrument at 77 K. In each experiment, approximately 50 mg of the sample of interest was weighed into a 9.001 cm³ glass cell. Prior to measurement, guest molecules were removed from the MOF pores via simultaneous vacuum and heat treatment; first for 1 hour at 80°C, then for 2 hours at 200°C. The sample cells were then immersed in a dewar of liquid nitrogen for the duration of the nitrogen adsorption measurement. BET areas were extracted from the nitrogen adsorption isotherms via the method described in the supporting information (Section 5) of the present work and in two of our previous papers^{37,49}.

2.2.4 Attenuated Total Reflection Infrared (ATR-IR) Spectroscopy

Loose MOF powder was introduced to a Bruker Vertex70 FTIR spectrometer equipped with a Bruker OPTIK Platinum ATR accessory with a diamond internal reflection element. ATR-IR spectra (2 cm⁻¹ resolution, 256 scans) were then recorded in the 4000-600 cm⁻¹ range and detected with an MCT detector. The spectral intensity was corrected for the change in the effective thickness value as a function of the incident wavelength.

2.2.5 Transmission IR Spectroscopy

Samples were measured in the form of small (ca. 2 cm²), thin self-supporting wafers, which were prepared by pellet-pressing ca. 30 mg of pure, undiluted MOF powder with a force of ca. 10 kN. In separate experiments, wafers were placed in a home-built airtight quartz cell with mid-IR transparent KBr windows. Before measurement, guest molecules were removed from the MOF pores by attaching the quartz cell to a custom-made vacuum line and applying simultaneous vacuum and heat treatment (1 h at 80°C then 2 h at 200°C). The transmission FTIR spectra were then recorded in the 4000-600 cm⁻¹ range with a Bruker Vertex 70 FTIR spectrometer equipped with a MCT detector (2 cm⁻¹ resolution, 32 scans).

2.2.6 Raman Spectroscopy

Samples were prepared and pretreated in the same manner as for the transmission FTIR measurements, albeit using another type of locally constructed airtight cell featuring a suprasil quartz cuvette. Raman spectra were recorded over a wavenumber range of 1800-500 cm⁻¹ on a Renishaw inVia Raman microscope spectrometer in backscattering mode. A 785 nm diode laser was used for the excitation, and the scattered photons were dispersed by a 1200 lines/mm grating monochromator and collected on a CCD camera whose collection optic was set at 20X objective summing up 66 scans with an exposition time of 20 s per step.

2.3 Computational methods

2.3.1 Used models

Two atomistic models were used for both DFT calculations and isotherm simulations: the crystallographic structure of UiO- $66^{12,82}$ for the defect-free model (named "Ideal") and a defective benzoate-containing structure (dubbed "Reo-Benz") that was created by starting from the perfect UiO-66 framework. First, from the primitive unit cell of UiO-66 (114 atoms, $Zr_6O_4(OH)_4[C_6H_4(COO)_2]_6$) the crystallographic cubic cell was built that corresponds to a four times larger supercell (456 atoms). Then, to create the defect, a $Zr_6O_4(OH)_4$ unit was removed along with the carboxylate groups of the neighbouring linkers and, finally, the dangling bonds on the aromatic rings were saturated with hydrogen atoms. The resulting structure was comprised of a 414-atom cubic unit cell (3· $Zr_6O_4(OH)_4[C_6H_4(COO)_2]_4[C_6H_5COO]_4$) that has 12 benzoic moieties in the defective site, as shown in Figure 1.

2.3.2 DFT calculations

All calculations were performed by means of the CRYSTAL14 *ab-initio* code⁸⁹. The hybrid B3LYP^{90,91} method combined with an all-electron Gaussian basis set were employed. The basis set consists of the following contractions of Gaussian functions: (8s)-(7631sp)-(621d) for Zr, (8s)-(411sp)-(1d) for O, (6s)-(31sp)-(1d) for C, and (31s)-(1p) for H (see ref. [82] for details). All computational parameters were the same as in the previous work of Valenzano et al. 82, except for the shrinking factor used in reciprocal space sampling (here raised to 3 from the original value of 2). Both the perfect (Ideal) and defective (Reo-Benz) UiO-66 models were fully relaxed by keeping the symmetry of the systems (24 symmetry operators). The presence of the defect leads to a negligible expansion of the unit cell volume (i.e. less than +0.1%), which is probably due to the well-known rigidity of the UiO-66 framework. Vibrational frequencies were computed at the Γ point on the optimized geometries 92. Infrared and Raman intensities were calculated via an entirely analytical procedure based on linear response 93,94. A Lorentzian broadening of 5 cm⁻¹ was adopted for both the infrared and Raman spectra. Vibrational frequencies were shifted by a factor of 0.98 in order to compensate for the well-known DFT overestimation effect.

The choice of B3LYP functional is justified by consistency with previous works on the UiO-66 crystal. The introduction of empirical dispersion corrections (Grimme's D3 with three-body contribution) leads to a mild contraction of cell volumes (1.4% for the perfect structure, less than 2% for the defective one) resulting in shifts for the individual frequencies not larger than 10 cm⁻¹. Since the discussion in the following focuses on isolated features of the vibrational spectra relative to the benzoate defect rather than collective modes, we consider the choice of the functional of minor importance.

2.3.3 Isotherm simulation

Adsorption isotherms were simulated with the "adsorption isotherm" task within the "Sorption Tools" menu in Accelrys Materials Studio version 8.1. The simulations were performed at 77 K, using a geometry optimized N_2 molecule as the adsorptive and a single, geometry optimized unit cell of the model structure of interest as the

adsorbent. We adopted the Metropolis method and the COMPASS force field. The isotherms were simulated over a pressure range of 0.01-100 kPa, in which 50 fugacity steps were distributed logarithmically in order to increase the number of data points in the steep initial portion of the isotherm. The "Fine" quality setting was used for all simulations. Charges were force field assigned, while electrostatic forces were calculated with the Ewald method. The "atom based" method was adopted for the calculation of Van der Waals forces. No constraints were assigned.

3. Results and Discussion

Figure 2 illustrates the complete set of PXRD patterns obtained on the full series of the defective samples over a 2θ range of $2\text{-}35^\circ$. All the samples are highly crystalline and with the expected patterns, except for the region preceding the first reflection of the UiO-66 phase. This region is emphasized in the left plot of Figure 2, where it can be seen that a very broad peak (spanning a 2θ range of ca. $2\text{-}7^\circ$) is observed in the PXRD patterns obtained on the samples synthesized in the presence of benzoic acid. In a couple of previous works^{37,49} we unambiguously assigned this "broad peak" to very tiny "nanoregions" of missing cluster defects, and showed that the relative intensity of the broad peak ($Rel(I)_{B,P}$) is correlated with the concentration of these defects. The absence of this peak in the pattern obtained on the reference UiO-66 sample ("Ref.", black line in Figure 2) proves its very low defectivity. In contrast, the broad peak is already clearly present in the pattern obtained on 10Benz, and its intensity (and thus, the concentration of missing cluster defects) systematically increases in 20Benz and 30Benz, at which point it essentially plateaus. This is more visually and quantitatively evident in Figure 3, where the $Rel(I)_{B,P}$ values (calculated via the method outlined in Section 4 of the SI, see Table S1 for numerical data) are plotted against the molar equivalents of benzoic acid used in the MOF syntheses. These observations suggest that benzoic acid is very effective at creating missing cluster defects in UiO-66, even at relatively low concentrations.

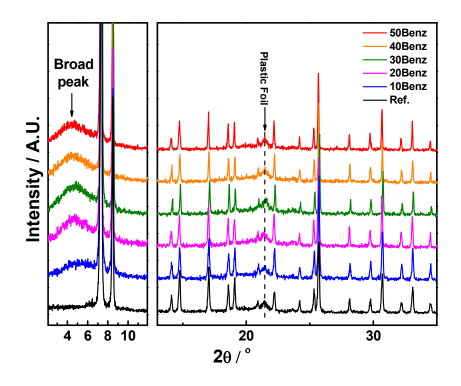


Figure 2 PXRD patterns obtained on the 6 UiO-66 samples. The same y-scale is applied to both plots.

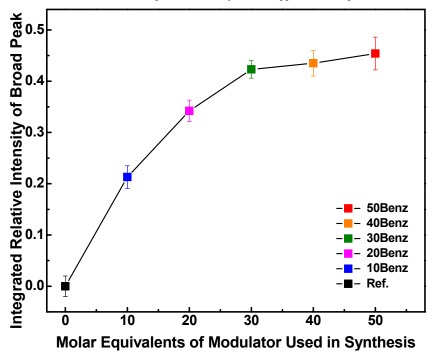


Figure 3 Integrated relative intensity of the broad diffraction peak $(Rel(I)_{B.P.})$ vs. the molar equivalents of modulator (benzoic acid) added to the UiO-66 synthesis mixture.

Liquid ¹H NMR spectroscopy was used to identify and determine the molar ratios between the organic components in the UiO-66 samples. Figure 4 compares the dissolution/¹H NMR spectra obtained on all 6 samples, and clearly illustrates the appearance of signals attributed to benzoate in the samples synthesized in the presence of benzoic acid. Since benzoate ligands are thought to terminate the missing cluster defects in these materials, their benzoate content is essentially a measure of their defectivity. However, differences in the intensity of the benzoate signal throughout the series are not particularly apparent upon visual inspection of the spectra, making it difficult to discern qualitative trends in their benzoate content by observation alone.

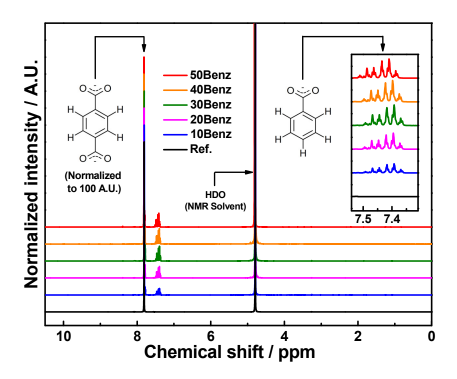


Figure 4 Dissolution/¹H NMR spectra obtained on the 6 UiO-66 samples.

Regardless, the real strength of the dissolution / 1 H NMR is that one can quantify the molar ratios between the organic components in MOF samples by integrating the spectra and performing simple calculations. Thus, we determined the benzoate/BDC molar ratios $\left(\frac{n_{Benz}}{n_{BDC}}\right)$ in our samples, which can be considered as a quantitative descriptor for their benzoate content. Moreover, a very small concentration of formate moieties (originating from the *in-situ* hydrolysis of DMF during the MOF synthesis³⁷) was determined via integration of the faint signal falling at ca. 8.4 ppm, yielding the formate to BDC molar ratio $\left(\frac{n_{Form.}}{n_{BDC}}\right)$. This contribution was added to $\frac{n_{Benz.}}{n_{BDC}}$ to get the "total modulator to BDC molar ratio" $\left(\frac{n_{Mod.}}{n_{BDC}}\right)$, which is a quantitative descriptor for the amount of monocarboxylate ligands in the samples (and thus, the concentration of defects in the UiO-66 framework³⁷). These

calculations were somewhat complicated by the overlap of the BDC and benzoate signals, as discussed in Section 6 of the supporting information where full details of the method are provided.

Figure 5 is the plot obtained when the $\frac{n_{Mod.}}{n_{BDC}}$ values of the 6 samples (see Table S2 for the numerical data) are plotted against the molar equivalents of benzoic acid added to their syntheses. The plot clearly shows that the extent of monocarboxylate incorporation (and thus, the defectivity of the samples) systematically increases throughout the series, but essentially plateaus for the last 2 samples (40Benz and 50Benz). This again emphasizes benzoic acid's ability to induce a high level of defectivity to the UiO-66 framework, even when used at relatively low concentrations. If we consider that the Reo-Benz model misses the 25% of the clusters (three Zr-bricks instead of four in the unit cell), the 50Benz sample (the most defective material) is missing the 19.5% of clusters.

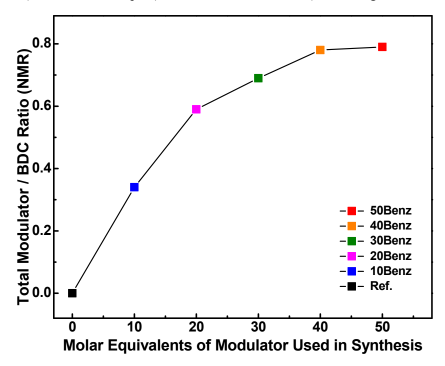


Figure 5 Graph obtained when the total modulator to BDC molar ratios $\left(\frac{n_{Mod.}}{n_{BDC}}\right)$, see Table S2) are plotted against the molar equivalents of modulator (benzoic acid) added to the UiO-66 synthesis mixture.

Nitrogen sorption measurements were used to assess the porosity of all the UiO-66 samples. All the experimentally obtained isotherms are presented in the right plot of Figure 6, while those simulated from the "Ideal" and defective "Reo-Benz" UiO-66 structural models (see Figure 1) are shown in the left plot for comparison. Starting with the simulated isotherms, one can clearly see that the "Reo-Benz" model's nitrogen uptake capacity (and thus, porosity) is significantly higher than that of ideal UiO-66. This essentially proves that increasing the concentration of missing cluster defects terminated by benzoate ligands should result in an increasingly porous material.

With this in mind, let us turn attention to the experimentally obtained isotherms shown in the right plot of Figure 6. Before delving into the trends, we feel it is noteworthy to comment on the contour (profile) of the isotherms. For example, the isotherm obtained on the reference UiO-66 sample (Ref.) has a textbook Type I profile, featuring a well-defined knee and perfectly flat plateau at higher relative pressures, emphasizing the uniformity of the sample's micropores. On the other hand, the isotherms obtained on the rest of the samples feature less well-defined knees and plateaus, as is expected for defective UiO-66 materials with a broader distribution of pore sizes. Nevertheless, these cavities do not appear to enter into the mesoporous size range as there is no evidence of hysteresis in any of the isotherms, whose adsorption and desorption branches are perfectly superimposed over the full pressure range.

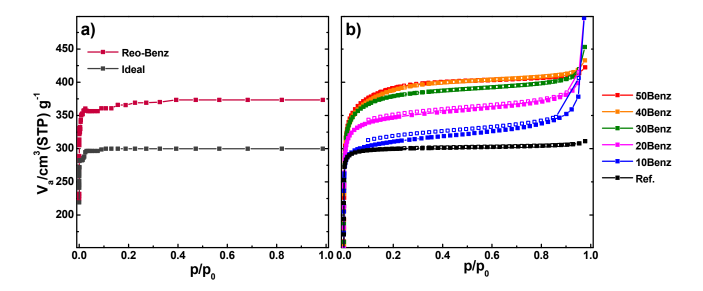


Figure 6: Simulated and experimental nitrogen adsorption isotherms obtained on all UiO-66 samples at 77 K. a) Simulated isotherms for the Ideal (dark grey) and Reo-Benz (dark red) models. b) Experimental isotherms Adsorption is depicted by filled squares, desorption by open squares. The same y-scale is applied to both plots.

Of greater relevance to this work is the clear trend which emerges when comparing the nitrogen uptake capacity (and thus, defectivity) of the samples. As one can see, the uptake capacity of the materials systematically increases as the amount of benzoic acid used in the synthesis was increased, eventually plateauing for the last two samples in the series (40Benz and 50Benz). This trend in porosity is quantitatively confirmed by Figure 7, in which the BET areas of the samples (calculated by the method outlined in Section 6 of the SI, see Table 2 for numerical data) are plotted against the molar equivalents of benzoic acid added to their syntheses.

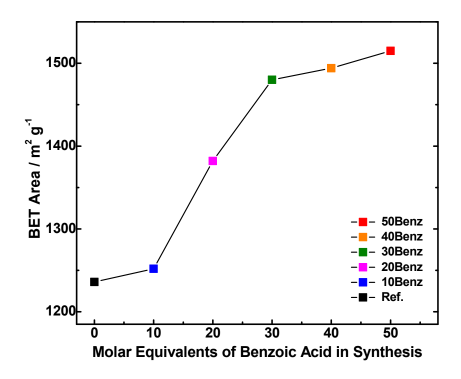


Figure 7 BET areas of the 6 UiO-66 samples (see also Table 2) plotted against the molar equivalents of modulator (benzoic acid) used in their synthesis.

Finally, it is of interest to compare the simulated and experimental data to contextualize our results. For example, one can clearly see that the qualitative (isotherm contour) and quantitative (nitrogen uptake capacity, BET areas) data obtained on the reference material (Ref.) matches very closely with that simulated from the Ideal UiO-66 model, strongly suggesting that the sample is close to defect-free. On the other hand, the data obtained on the 3 most defective samples (30Benz, 40Benz, and 50Benz) is similar to that simulated from the Reo-Benz model, suggesting that the 3 samples are extremely defective, which is an especially striking revelation when one considers that the Reo-Benz model represents an extreme case in which one quarter of the clusters are missing from the UiO-66 framework.

As was the case for the equivalent plots in previous work³⁷, the trends observed in Figures 3, 5, and 7 appear to be qualitatively similar, prompting us to explore the relationship between the data plotted therein $(Rel(I)_{B.P.}, \frac{n_{Mod.}}{n_{BDC}})$, and BET area). To this end, we plotted the $Rel(I)_{B.P.}$ values against the $\frac{n_{Mod.}}{n_{BDC}}$ and BET area values of the samples (see Table 2 for numerical data), with the results shown in Figure 8.

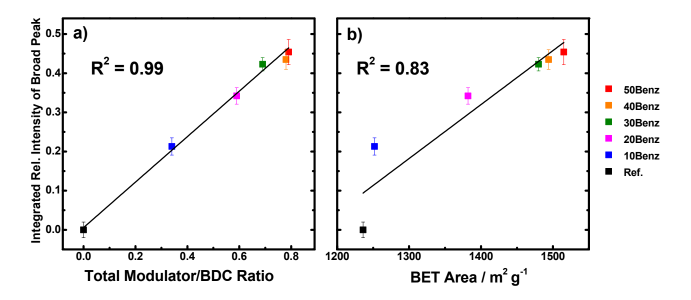


Figure 8 Graphs and linear fits obtained when the relative intensity of the PXRD broad peak $(Rel(I)_{B.P.})$ is plotted against: a) the total modulator to BDC ratio $\left(\frac{n_{Mod}}{n_{BDC}}\right)$; b) the BET area.

As one can see from the fitted straight lines and R^2 values on the figure, the data is very well correlated. This makes sense when one considers that all three parameters are all in some way correlated with the concentration of defects in the UiO-66 framework, further validating their branding as "defectivity descriptors" in our previous work³⁷. As also mentioned in said previous work, only one of the defectivity descriptors (the relative intensity of the broad peak, $Rel(I)_{B,P}$) is exclusively associated with one type of defect (missing cluster defects). The fact that the other two descriptors are linearly correlated with $Rel(I)_{B,P}$ is thus a strong indication that missing cluster defects (terminated by monocarboxylate ligands, mostly benzoate in these samples) are the predominant defect in the samples under investigation herein. This conclusion is further backed by the similarity between the nitrogen adsorption isotherms obtained on the most defective samples and the isotherm simulated from Reo-Benz, a highly defective UiO-66 structural model in which a quarter of the clusters are missing and compensated by benzoate ligands (see Figure 6 for isotherms and Figure 1 for the structural model).

Table 2. Summary of the quantitative experimental data extracted from the PXRD, dissolution/¹H NMR, and nitrogen adsorption and results obtained on the UiO-66 samples. Theoretical values from the models (see Figure 1) are also provided.

Sample	$Rel(I)_{B.P.}$ / Error	n _{Mod.}	BET Area (m ² g ⁻¹)	Reference
		n _{BDC}		
Ref.	0.000 / 0.020	0.00	1236	37
10Benz	0.213 / 0.022	0.34	1252	This work
20Benz	0.342 / 0.021	0.59	1382	This work
30Benz	0.423 / 0.017	0.69	1480	This work
40Benz	0.435 / 0.025	0.78	1494	This work
50Benz	0.454 / 0.032	0.79	1515	This work
Model	$Rel(I)_{B.P.}$ / Error	n _{Mod} .	BET Area (m ² g ⁻¹)*	Reference
Ideal	_	0.00†	1241	37
Reo-Benz	_	1.00:	1532	49

^{*}Calculated from the simulated nitrogen adsorption isotherms (see Figure 6a).

While the combined evidence from these results point strongly towards the predominance of Reo-Benz type defects in the samples under investigation herein, the individual methods each detect said defects in a rather indirect manner. We therefore speculate that some of the MOF community may be skeptical to the conclusions drawn from these results (and those of our previous papers^{37,49}). Thus, we employed vibrational spectroscopic techniques (FTIR, Raman, and simulations thereof) to more directly probe the short range order of these materials. The O-H and C-H stretching region of the transmission FTIR spectra obtained on the full set of samples after thorough activation (see experimental section) is reported in Figure 9a. As one can see, the spectrum obtained on the Ref. sample is characterized by an intense and sharp band at 3672 cm^{-1} , attributed to the isolated $\mu_3(\text{OH})$ groups on the hydroxylated $\text{Zr}_6\text{O}_4(\text{OH})_4^{12+}$ cluster ^{82,95}. Its spectrum also features a broad band centered at 3250 cm^{-1} , which is due to hydrogen bonded species. The barycentre of this band is rather low in frequency, testifying that the hydrogen bonding is rather strong. The aromatic C-H stretching region of the spectrum features a clear band at 3065 cm^{-1} , which we assign to C-H stretching modes of the BDC linkers.

Moving on to the set of samples made with benzoic acid as modulator, we observe that all the absorptions due to $\nu(OH)$ modes drastically decrease in intensity; both the hydrogen bonded species (the broad band centered at 3250 cm⁻¹) and the isolated $\mu_3(OH)$ groups at 3672 cm⁻¹ (especially in the 3 most defective samples; 30Benz, 40Benz, and 50Benz). This interesting observation signifies that the samples made with benzoic acid as modulator are more easily activated and dehydroxylated, indicating that they adsorb water less strongly. Indeed, the larger pores of these defective samples may be expected to provide lesser dispersion forces, thus weakening their adsorptive strength for small molecules. Part b of Figure 9 compares the ATR-IR spectra obtained on the full set of samples,

[†]There are no monocarboxylate ligands in the ideal UiO-66 composition, $Zr_6O_4(OH)_4(BDC)_6$, so $\frac{n_{Mod.}}{n_{BDC}}=0$.

 $^{^{\}ddagger}$ The composition of Reo-Benz is Zr₆O₄(OH)₄(BDC)₄(Benz)₄, where "Benz" is benzoate, a monocarboxylate ligand. Thus, $\frac{n_{Mod.}}{n_{BDC}} = \frac{4}{4} = 1$

which are presented together with ATR-IR spectra obtained on benzoic acid (in DMSO solution) and pure DMSO. The benzoic acid spectrum is included to determine whether the benzoate detected in the samples (see ¹H NMR results in Figure 4) can simply be attributed to free benzoic acid molecules in the MOF pores. However, it is clear that this is not the case since benzoic acid's most intense signals at 1700 cm⁻¹ and 1250 cm⁻¹ (due to v(C=O) and v(C-O), respectively highlighted by the hashes in Figure 9b) are only very faintly visible in the spectra obtained on the UiO-66 samples, and don't have a clear relationship with the amount of benzoic acid used in their syntheses. Instead, they feature bands at 1580 cm⁻¹ and at 1394 cm⁻¹, due to the v_{asym} (OCO) and v_{sym} (OCO) modes of carboxylate groups, respectively. One may have expected to see a trend in which the intensity of these bands follows the amount of benzoic acid used in the synthesis, but this is not the case since the carboxylate groups are not unique to benzoate (they are also present in the linker and formate ligands). However, a trend is clearly observed in the intensity of the weak signals falling at 1178, 1070 and 1025 cm⁻¹ (each highlighted by the asterisk in Figure 9b), eventually plateauing in the spectra obtained on 40Benz and 50Benz. An enlarged view of this spectral range is reported in the Supporting Information (Figure S1). This trend is very similar to that of the defect descriptors discussed earlier, strongly suggesting that these bands are associated with the presence of defect-compensating benzoate moieties in the UiO-66 framework..

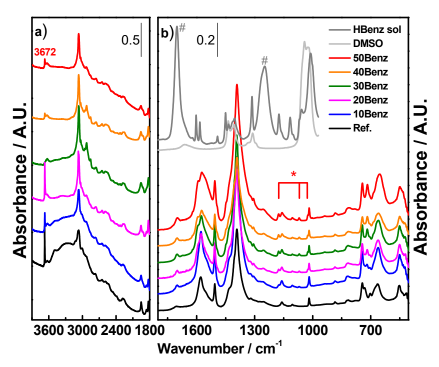


Figure 9 Part a): Transmission FTIR spectra obtained on UiO-66 samples activated under dynamic vacuum for 2 h at 200 °C. Part b) ATR-IR spectra obtained on the samples in air and on benzoic acid (in DMSO solution) and DMSO are also reported for comparison.

The Raman spectra obtained on the full set of samples (after thorough activation, see experimental section) are presented in Figure 10, together with those recorded on solid benzoic acid, benzoic acid in DMSO solution, and pure DMSO.

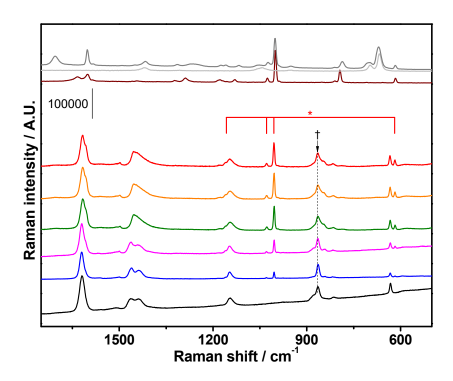


Figure 10 Raman spectra obtained on the full set of samples (activated under dynamic vacuum for 2 h at 200 °C), solid benzoic acid, benzoic acid in DMSO solution, and DMSO. The UiO-66 spectra were normalized to the intensity of the band at 866 cm⁻¹ (emphasized with a dagger in the figure). UiO-66-Ref (black), 10Benz (blue), 20Benz (magenta), 30Benz (green), 40Benz (orange), 50Benz (red). Solid Benzoic acid (brown) Benzoic acid in DMSO (grey) and DMSO (light grey).

As one can see, the Raman spectra of these materials feature much less bands than their FTIR counterpart, allowing for an easier identification of specific fingerprints ascribable to the presence of the benzoate ligands. Spectra obtained on benzoic acid (both as a pure solid and as a DMSO solution) have again been included for comparison to determine whether any free benzoic acid (evidenced in previous studies⁷⁷) is present in the MOF pores. To this end, it is clear that benzoic acid was completely removed by washing since key bands (such as the v(C=O) at 1705 cm⁻¹ and v(C-O) at 1260 cm⁻¹, among others) are completely absent from the UiO-66 spectra. A Figure that compares IR and Raman spectra of the MOFs with benzoic acid references is reported in the Supporting Information (Figure S2). Conversely, new bands (highlighted with an asterisk in the figure) appear at 1161 cm⁻¹ (shouldering the band at 1147 cm⁻¹), 1029 cm⁻¹, 1005 cm⁻¹, and 618 cm⁻¹ in the spectra obtained on the benzoic acid modulated UiO-66 samples, systematically increasing in intensity as the amount of benzoic acid used in the synthesis was increased. Much like the defectivity descriptors discussed throughout this work, the intensity of

these bands reach a plateau in the spectra obtained on 30Benz, 40Benz, and 50Benz, strongly suggesting that these new bands are associated with the presence of defect-compensating benzoate moieties in the UiO-66 framework. In order to assign all these spectroscopic features, a parallel computational study on the perfect "Ideal" UiO-66 framework and on the Reo-Benz defective counterpart (see the experimental section for the details) has been conducted. Figure 11 parts a) and b) compares the experimental data obtained in case of 50Benz UiO-66 sample with the computed ones. Thanks to the possibility to animate each vibrational frequency, the assignments of major bands was possible. Table 3 summarizes the most relevant vibrational features with the correspondent assignment based on the animations visualized for both Raman and IR computed spectra.

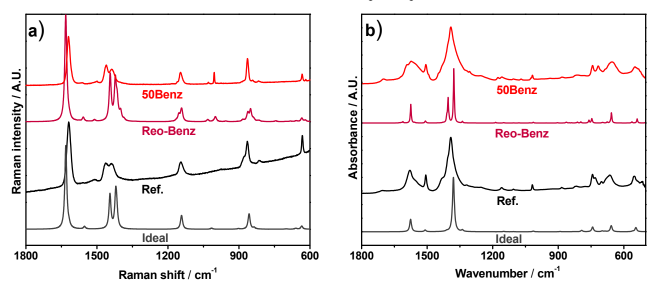


Figure 11 Comparison between simulated (dark grey and dark red) and experimental (black and red) spectra of Ref. (black) and 50Benz MOFs (red) samples with Ideal (dark grey) and Reo-Benz (dark red) models by the means of Raman (a) and IR (b) spectroscopies.

Table 3. Assignment of a selection of experimental and calculated vibrational modes compared for the perfect "Ideal" and defective "Reo-Benz" models. Scaling factor of 0.98 has been adopted in order to correct for the DFT overestimation issue.

Experi	imental	Com	puted	Description of the mode
IR	RAMAN	IR	RAMAN	
3672	Not visible	3812	3812	OH stretching of isolated hydroxyls
3065	Not active	3181, 3176	Not active	CH stretching in terephthalate and benzoate
3030	Not visible	3146, 3139	3146, 3139	CH stretching in benzoate
1700	1705	Not calc.	Not calc.	C=O stretching of benzoic acid
1580	Not visible	1575	1575	OCO asymmetric stretching in carboxylate
Not active	1620	Not active	1632	C=C stretching of aromatic rings
Not	1452, 1434	1445, 1422	1445, 1422	OCO symmetric stretching in carboxylate in-phase
detectable				
1394	Not active	1378	Not active	OCO symmetric stretching in carboxylate out-of-phase
1252	1260	Not calc.	Not calc.	C-O stretching of benzoic acid
1178	Not active	1187	Not active	In-plane CH bending in benzoates
1158	Not active	1150	Not active	In-plane CH bending in terephthalates and benzoates
Not active	1161	Not active	1165	In plane CCH bending in benzoates
Not active	1147	Not active	1154	Terephthalates rings breathing + benzoate rings deformation
1070	Not active	1080	Not active	In-plane CCH bending in benzoates
1025	1029	1031	1031	Benzene ring deformation in benzoates
1019	Not active	1014	Not active	Benzene ring deformation in terephthalates
Not active	1005	Not active	1001	Benzene ring deformation in benzoates
Not active	633	Not active	634	Benzene ring deformation in terephthalates
Not active	618	Not active	619	Benzene ring deformation in benzoates

At least four signals of the Raman spectra can be unambiguously assigned to the presence of modelled defects: the modes falling at 1029 cm⁻¹, 1005 cm⁻¹ and 618 cm⁻¹, present only in the defective samples, are due to the deformation of the benzene rings belonging to benzoate moieties (vibrational modes that are prohibited for symmetry in terephthalates). The 1161 cm⁻¹ signal is similarly assigned to a CH bending mode involving the proton in para position in respect to the carboxylate in benzoates.

Whereas in the IR spectra three very weak signals falling at 1178 cm⁻¹, 1070 cm⁻¹ and 1025 cm⁻¹ (shoulder) are ascribed in the same way to ring deformations and CH bending modes.

The computed intensity of all these signals in respect to the measured ones is slightly higher as the Reo-Benz model has been constructed with a molar ratio between benzoates and terephthalates of 1:1, whereas liquid NMR measurements (see Table S2) has found only a molar ratio of 0.78:1 for the 50Benz sample. It should be noticed

that as the broadening of the computed spectral signals is arbitrarily set to 5 cm⁻¹ only the integrated intensity should be taken into account.

The v(C-H) region (reported only for experimental infrared spectroscopy in Figure 9a), which shows two different signals belonging to terephthalates and benzoates (falling respectively at 3065 and 3030 cm⁻¹), is again nicely modelled by the DFT calculations.

4. Conclusions

A systematic and detailed study about the synthetic insertion of missing-cluster defects (via benzoic acid additions) inside the UiO-66 framework has been reported through the present work together with their characterization with a selection of techniques. While the long-range crystalline order of the original framework has been preserved, as demonstrated by the PXRD patterns, tiny nanoregions where defects order themselves were identified by the growth of a broad symmetry-prohibited diffraction peak which intensity is strongly correlated with the number of introduced defects.

The true concentration of Reo-Benz defects has been accurately measured with dissolution liquid ¹H/NMR spectroscopy: the outcome reveals that even massive excesses of benzoic acid during synthesis (up to 50 equivalents in the 50Benz sample) are only able to insert about 0.78 benzoic moieties per terephthalic ligand: this number is like to be a plateau as also 30Benz and 40Benz samples almost reached this value.

Coming to N_2 physisorption measurements we noticed that defectivity is able to effectively increase the accessible surface area inside the framework of about 30% in respect to the reference sample with a change in the shape of the isotherm which probably points out differences in the dimensions of the micropores now accessible thanks to the presence of defects. Vibrational spectroscopies (IR and Raman) measurements gave a beautiful set of spectra with a series of signals which relative intensity is growing with the concentration of defects; the identity of every vibrational band has been fully assigned with ab-initio DFT calculations and the graphical analysis of the vibrational modes. The effectiveness of the constructed atomistic models, representative of both ideal and defective structures, is confirmed by both isotherm simulation and simulated IR and Raman vibrational spectra which are in nice agreement with experimental results. Nevertheless in-situ IR spectroscopy of vacuum-treated samples pointed out that defective samples are more hydrophobic than their ideal counterpart, a result that is in agreement with the non-polar nature of benzoate moieties.

This work also demonstrates that these techniques can be used as a "tool-box" to probe the presence and the concentration of dangling benzoate moieties inside the MOF; this approach could potentially be extended to different chemical species (i.e. other monodentate ligands) if they also present a well-defined spectroscopic fingerprint.

Empowered by these results we presented in the present paper strong evidences of the possibility to include benzoate decorated defects inside the UiO-66 framework and to specifically characterize them. This "defect-engineering" approach is able to increase the sorptive performance of this material by a significant degree, as discussed in the introduction, and can be used to tune the properties of these materials for specific applications.

Supporting information

Quantitative data extracted from PXRD and ¹H/NMR measurements; vibrational spectra comparison between MOFs and Benzoic acid; detailed description of the method used for calculating the relative intensity of the broad PXRD peak, the BET area and the molar ratios between linker and modulator from ¹H/NMR data.

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