Isolating the Role of the Node–Linker Bond in the Compression of UiO-66 Metal–Organic Frameworks

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ABSTRACT: Understanding the mechanical properties of metal–organic frameworks (MOFs) is essential to the fundamental advancement and practical implementations of porous materials. Recent computational and experimental efforts have revealed correlations between mechanical properties and pore size, topology, and defect density. These results demonstrate the important role of the organic linker in the response of these materials to physical stresses. However, the impact of the coordination bond between the inorganic node and organic linker on the mechanical stability of MOFs has not been thoroughly studied. Here, we isolate the role of this node–linker coordination bond to systematically study the effect it plays in the compression of a series of isostructural MOFs, M-Uio-66 (M = Zr, Hf, or Ce). The bulk modulus (i.e. the resistance to compression under hydrostatic pressure) of each MOF is determined by in situ diamond anvil cell (DAC) powder X-ray diffraction measurements and density functional theory (DFT) simulations. These experiments reveal distinctive behavior of Ce-Uio-66 in response to pressures under one GPa. In situ DAC Raman spectroscopy and DFT calculations support the observed differences in compressibility between Zr-Uio-66 and the Ce-analogue. Monitoring changes in bond lengths as a function of pressure through DFT simulations provides a clear picture of those which shorten more drastically under pressure and those which resist compression. This study demonstrates that changes to the node–linker bond can have significant ramifications on the mechanical properties of MOFs.

INTRODUCTION

Over the past two decades, metal–organic frameworks (MOFs) have become a prominent class of porous materials for many applications, including gas adsorption,¹ ² chemical separations,³ ⁴ and heterogeneous catalysis.⁵ ⁷ These scaffolds consist of inorganic nodes connected by multitopic organic linkers to form 2- and 3-dimensional frameworks with exceptional porosity and tremendous structural diversity.⁸ While several classes of MOFs have demonstrated high chemical and thermal stability in recent years,⁹ a rigorous understanding of how these materials respond to mechanical stress is still emerging. Interesting phenomena, such as pressure-induced amorphization,¹⁰ pore collapse,¹¹ and structural transitions,¹² have been well documented in MOFs when exposed to pressures on the order of 1 GPa. Elucidating structure-property relationships of MOFs based upon their mechanical properties is challenging due to the sheer number of frameworks and the highly specialized equipment required for experimental measurements. On the other hand, high-throughput computational studies offer insight into the structural characteristics that relate to mechanical properties,¹³ though detailed design principles remain elusive. A combination of experimental and theoretical work provides a more nuanced picture of the factors that dictate the mechanical properties of MOFs.

UiO-66 is among the most well-studied MOFs due to its cubic space group, remarkable chemical and thermal stability,¹⁴ The structure consists of 12-connected Zr₆(µ₃-O)(µ₅-OH)₄ nodes joined by linear 1,4-benzenedicarboxylic acid linkers which assemble into the fcu topology (Figure 1).¹⁵ Several studies have demonstrated that UiO-66 is more resistant to mechanical stress than other prototypical MOFs,¹⁶ ¹⁷ though the presence of missing linker defects can significantly impact the mechanical properties.¹⁸ ¹⁹ In a recent study from our group, we demonstrated that systematic extension of the organic linker in the UiO family of MOFs leads to a decrease in the bulk modulus (that is, the resistance to hydrostatic pressure; K₀ = -V ∂P/∂V).²⁰ Given the breadth of knowledge regarding this framework, UiO-66 is a well-established platform for investigating the mechanism of structural changes at high pressures. While numerous reports support the notion that changes to the organic linker²¹ ²² and topology²³ ²⁴ can impact the compressibility of MOFs, systematic studies into the effect of changes to the metal node are less prevalent.²⁵
Herein, we investigate the role of the metal-carboxylate bond in the compression of UiO-66 by varying the identity of the metal node from Zr-Uio-66 to Hf-Uio-66 and Ce-Uio-66. By keeping the topology, linker, and experimental conditions constant, we isolate the coordination bond as the single structural variable that changes in this series of MOFs. While the Zr and Hf analogues exhibit almost identical behavior under pressure, Ce-Uio-66 is found to be much more compressible. Pressure-dependent Raman spectroscopy supports the observations made from high-pressure in situ powder X-ray diffraction (PXRD) measurements. Density Functional Theory (DFT) simulations of Zr- and Ce-Uio-66 are in good agreement with the experimental observations and reveal that the node–linker bond of Ce-Uio-66 compresses more readily than that of the Zr analogue. Together, these results indicate that the identity of the metal node in MOFs can have a significant impact on their response to high pressures by modulating the strength of the coordination bonds that hold these scaffolds together.

EXPERIMENTAL METHODS

All MOFs were synthesized and activated according to modified literature procedures. The materials were characterized using ambient pressure powder X-ray diffraction (PXRD) and isothermal nitrogen adsorption to verify their crystallinity and porosity, respectively. (Figures S1 and S2). X-ray photoelectron spectroscopy (XPS) was conducted with an Al Kα source. Five scans lasting 30 seconds each were collected. Ambient pressure Raman spectroscopy was conducted using a 532 nm excitation laser (see Supporting Information for details). High-pressure PXRD was conducted at the 17-BM-B beamline (λ = 0.45390 Å) at the Advanced Photon Source (APS) at Argonne National Laboratory (ANL) using a diamond anvil cell (DAC) sample environment. In a typical experiment, the MOF sample was gently ground with an internal standard, CaF₂, to ensure uniform mixing and to break up any large crystals. The mixture was loaded into a hole in a pre-indented stainless-steel gasket and sealed in the DAC. An ambient pressure diffraction pattern was collected, then a drop of Fluorinert™ FC-70 was added to the cell as a non-penetrating pressure transmitting fluid. In situ PXRD data were then collected as a function of pressure. Unit cell parameters were then extracted from the PXRD patterns using Le Bail refinement. Bulk moduli (K₀ = - V ᴦ P/ᴦV) were extracted by fitting a plot of unit cell volumes vs. pressure to a second-order Birch-Murnaghan equation of state. Pressure-dependent Raman spectroscopy experiments were conducted at the GSECARS offline Raman spectrometer located at the APS at ANL. Spectra were collected using a 532 nm excitation laser, and ruby fluorescence was used to monitor pressure inside the DAC. For further details regarding DAC diffraction and Raman experiments, see Supporting Information.

Each MOF was investigated through computational simulations using the Density Functional Theory (DFT) as implemented in the CRYSTAL14 code. All-electron localized basis sets were used for all the atoms except for hafnium and cerium, for which a pseudopotential approach was used. These basis sets can be found on the CRYSTAL online library with the corresponding acronyms and original references: C: C_G-31d1G_gatti_1994; O: basis set used by Vienzano et al.; H: H_3-1p1G_1994; Zr: Zr_all_electron_dovesi_unpub; Ce: Ce_ECP_Meyer_2009.

Several different functionals, at different levels of approximation, were considered to describe the exchange and correlation components of the energy. Among these, PBESOL0 hybrid functional was retained as it gives a good accuracy while keeping an acceptable computational cost. The effect of Grimme-type dispersion corrections was tested and found to be of a minor impact on the optimized structures and derived properties. Therefore, these corrections were not included in this study.

Mesh sampling of the reciprocal space was performed using the Monkhorst-Pack scheme. The k-point mesh of 2 × 2 × 2 was used for each MOF structure to obtain convergence of the properties of interest. This generated mesh was then used for all our calculations. Higher convergence criteria than the defaults...
proposed by CRYSTAL14 code were used for geometry optimization (a maximum of 0.0005 a.u. on atomic displacements during one optimization step and 0.0001 a.u. on forces). Optimized structures and representative input files for the calculations are available online at https://github.com/fxcoutert/citable-data.

RESULTS AND DISCUSSION

Ambient pressure PXRD patterns for each MOF are in good agreement with the simulated powder patterns (Figure S1) from reported crystal structures, indicating that the UiO-66 structure is retained when synthesized with different metals. Nitrogen physisorption isotherms (Figure S2) indicate that porosity of the UiO-66 MOFs is also maintained.

Upon applying pressure using a DAC, the PXRD peaks shift steadily to higher angles of diffraction, indicating a decrease in unit cell volume (Figures S4 and S5, Tables S1 and S2). We extracted the unit cell volumes of each MOF at each measured pressure. Plotting the unit cell volumes of Zr-Uio-66, Hf-Uio-66, and Ce-Uio-66 as a function of pressure reveals that the Zr and Hf analogues behave nearly identically over a pressure range of 0-0.4 GPa, followed by a slight divergence above 0.4 GPa (Figure 2). Remarkably, the unit cell volume of Ce-Uio-66 decreases much more rapidly than the other MOFs, with noticeable discontinuities around 0.1 GPa and 0.4 GPa. These discontinuities may indicate possible pressure-induced phase transitions, i.e., changes to the MOF structure upon achieving certain pressures that alter the mechanical properties of the material. Unfortunately, the PXRD patterns collected were of insufficient quality to determine the nature of these suspected structural transformations.

The initial data points for each MOF were fit to a second-order Birch-Murnaghan equation of state to estimate the bulk modulus (K_B = -V ∂P/∂V) of the material (Figure 2). The bulk modulus describes the pressure required to induce a given change in volume and is inversely related to compressibility. Zr-Uio-66 and Hf-Uio-66 exhibit nearly identical bulk moduli (K_B = 37.9 ± 0.6 GPa and 37 ± 1 GPa, respectively), while Ce-Uio-66 has a much lower bulk modulus (K_B = 16.9 ± 0.7 GPa) as evidenced by the steep decline in unit cell volume with pressure. These results indicate that while Zr-Uio-66 and Hf-Uio-66 resist compression to the same degree, the Ce framework is significantly less rigid. Because the three MOFs have similar properties at ambient conditions and share identical organic linker and topology, we attribute the distinct mechanical properties to differences in the bond between the node and linker.

Given the striking difference in the bulk modulus of Ce-Uio-66, we turned to pressure-dependent Raman spectroscopy to probe the coordination bond between the metal node and organic linker. Unfortunately, the node-linker bond cannot be directly monitored using this technique because this vibration is infrared active. To circumvent this challenge, we selected the C-C stretch between the aromatic core of the linker and the carboxylate carbon and the O-C-O symmetric stretch of the carboxylate (Figure S6) as target vibrations which provide insight into the strength of the metal-linker bond and have been assigned previously.

Ambient pressure Raman spectra were collected for each material to determine the frequencies that correspond to the vibrations of interest. Then, Raman spectra were collected within a DAC with incremental increases in pressure for each MOF (Figure 3). In each sample, significant peak broadening is observed with increasing pressure, which is a common phenomenon in high-pressure Raman spectroscopy. In Zr-Uio-66, both vibrations of interest exhibit a steady hypsochromic shift as pressure increases from 0-1.20 GPa. Hf-Uio-66 displays a similar behavior over a pressure range of 0-0.92 GPa, though the effect in the C-C stretch (starting at 1437.8 ± 0.7 cm⁻¹) is somewhat obscured by peak broadening. The Raman spectrum of Ce-Uio-66 changes dramatically between ambient pressure and the spectrum collected under pressure (0.24 GPa). Immediate broadening of the two peaks results in a single broad signal, precluding meaningful qualitative analysis. This significant change in the shape of the Raman spectrum may result from structural changes occurring at low pressures. The stark difference in behavior between Ce-Uio-66 and the Zr and Hf counterparts supports the observed contrast in compressibility from the PXRD data.

We then fit the region of the Raman spectra containing the vibrations of interest with two pseudo-Voigt functions in order to analyze the observed changes in a more quantitative manner. The frequencies corresponding to the center of each peak are shown in Figure 4 as a function of increasing pressure. In all three MOFs, the C-C stretch shifts steadily to higher frequencies in general, with a notable discontinuity present around 0.4 GPa in Ce-Uio-66 (Figure 4a). This jump in the Raman spectrum coincides with the dramatic change in compressibility observed in the DAC PXRD data. We assume that these discontinuities are a result of a structural change that occurs around 0.4 GPa, though further characterization is required to elucidate the details of such a transformation. For Zr-Uio-66 and Hf-Uio-66, the O-C-O symmetric stretch steadily increases in frequency with increasing pressure; however, in Ce-Uio-66 this vibration remains almost invariant after an initial increase in frequency upon pressurizing to 0.24 GPa. The consistency of this Raman shift indicates that the bond lengths and vibrational
frequency of the carboxylate group in Ce UiO-66 do not change as rapidly as the Zr and Hf analogues. The distinct behavior of the O–C–O vibration in Ce UiO-66 is evidence of the role of the metal-carboxylate bond in the pressure response of the material.

In order to better understand the compression of Zr-UiO-66 and Ce-UiO-66, we performed DFT calculations of their structures, their elastic properties, and their vibrational modes in the harmonic approximation. First, the experimental structures were optimized with full use of symmetry, relaxing both atomic positions and unit cell parameters. Very good agreement was found with the experimental cell parameters (Table S6). We then performed frequency calculations in the harmonic approximation in order to determine the characteristics of the MOFs’ vibration modes, in particular the C–C stretch and O–C–O symmetric stretch. Again, a good agreement with the experimental results was observed for both compounds (Table S7).

We note here a few negative frequencies are observed in the case of Ce–UiO-66; those vibration modes are related to the ligand bowing out of its average plane, also called “guitar string” modes. These vibrational modes have been observed in UiO-compounds, as well as in other MOFs, and arise from the fact that DFT calculations are performed at 0 K on a crystallographic structure with high symmetry, which is an average structure. At finite temperature, the
ligand would bow out of the plane in two directions, in this low-frequency mode, giving an in-plane average position which is the high-symmetry structure observed by X-ray diffraction.

Determination of the stiffness tensor (i.e., the second-order elastic constants) through linear response calculations allowed us to derive the bulk modulus for each material. Re-producing the experimental trend, simulations confirm a different compression between Zr–UiO-66 and its Ce analogue, with values of 42 and 37 GPa respectively. Although the difference is not as drastic as is observed in the experimental measurements, it demonstrates the softer nature of the Ce–UiO-66 framework.

We then conducted calculations of the structural evolution of the frameworks under pressure: geometry optimizations were performed in the pressure range of 0 to 2 GPa. Summary of the cell parameters for each pressure can be found in the supporting information (Table S6), as well as the optimized structures. As found experimentally, we observed a linear variation of the volume with respect to the pressure for Zr–UiO-66. However, our DFT calculations cannot reproduce the possible phase transition experimentally observed at 0.4 GPa for Ce–UiO-66 — indicating that it involves a symmetry-breaking transition to a lower-symmetry phase. In the case of Ce–UiO-66, computational results show a linear variation like that of the Zr analogue, though the slope is steeper, confirming the lower bulk modulus measured and calculated for this MOF.

From the stressed structures, we highlight a few structural variations of chemical importance. Considering first the ligand, a few similarities are observed between Zr- and Ce–UiO-66: variation of the O–C–O angle as well as the dihedral angle between ligands was identical in both materials, shifting around 1° in each case. However, using the cell parameter as a reference by plotting \( \frac{(l/l_0)}{(a/a_0)} \) (\( l \) = bond length, \( l_0 \) = bond length at ambient pressure, \( a \) = unit cell parameter, \( a_0 \) = unit cell parameter at ambient pressure) (Figure 5), we found that C–O bond decreases more slowly than the lattice parameter in both cases (\( (l/l_0)/(a/a_0) > 1 \) as pressure increases) whereas C–C bond decreases almost exactly at the same rate (\( (l/l_0)/(a/a_0) = 1 \) as pressure increases). Moreover, we saw that this C–O bond shortens faster in the case of Zr supporting the slower variation of the O–C–O symmetric stretch for Ce. This demonstrates the importance of the carboxylate group to understand the compression of this material. Metal–carboxylate distances were also investigated in both materials and we found that they show a faster decrease than the lattice parameter [\( (l/l_0)/(a/a_0) < 1 \) as pressure increases]. We also saw that this decrease is even more pronounced in the case of Ce–UiO-66 which could be due to the orbitals of Ce being more diffuse than those of Zr, leading then to a slightly higher flexibility of the Ce–carboxylate bond.

Given the apparent role of the metal–carboxylate bond in determining the compressibility of these MOFs, we considered the possibility of Ce\(^{3+}\) forming during the MOF synthesis due to the high reduction potential of Ce\(^{4+}\). This Ce\(^{3+}\) species could then incorporate into the Ce–UiO-66 structure. To quantify the portion of reduced Ce, we conducted XPS experiments, revealing ~10% Ce\(^{3+}\) and ~90% Ce\(^{4+}\) present in Ce–UiO-66 (Figure S7). This ratio corresponds to roughly 47% of Ce nodes containing at least one Ce\(^{3+}\) atom, assuming they are evenly distributed throughout the MOF. Even with a low proportion of Ce\(^{3+}\), a significant number of nodes likely contain a reduced ion. Ce\(^{3+}\) exhibits a longer ionic radius than Ce\(^{4+}\), which is often invoked to rationalize the well-documented unit cell expansion of bulk CeO\(_2\) upon reduction.\(^{40-41}\) The presence of Ce\(^{3+}\) in the nodes of Ce–UiO-66 may lead to distortions of the node structure of Ce–UiO-66, influencing the mechanical properties of the MOF, and may also contribute to the lowering of the elastic moduli of the Ce–UiO-66 due to the softer Ce–carboxylate coordination allowing easier compression and shearing deformations.

These rationales are in agreement with the difference observed between experimental and computational results on the bulk modulus of Ce–UiO-66, as calculations were performed on an ideal system with 100% Ce\(^{4+}\). We hypothesize that the prevalence of Ce\(^{3+}\) throughout Ce–UiO-66 contributes to the uniquely distinct behavior of the framework under pressure compared to the Zr- and Hf–analogues.

CONCLUSIONS

This study investigates the influence of metal identity on the compression of a prototypical MOF, UiO-66. We conducted in situ DAC PXRD experiments to quantify the bulk modulus for three materials: Zr–UiO-66, Hf–UiO-66, and Ce–UiO-66. These results indicate that Ce–UiO-66 compresses much more readily than the Zr and Hf analogues (Figure 2). We then conducted in situ DAC Raman spectroscopy to probe two vibrations involving the carboxylate group of the organic linker. While the Raman shifts of the C–C stretch between the carboxylate and aromatic core of the linker increase in frequency with pressure for all three MOFs, the O–C–O symmetric stretch of Ce–UiO-66 remains nearly constant from 0.24-0.84 GPa. This behavior contrasts the steady hypsochromic shift observed for the same vibration in Zr–UiO-66 and Hf–UiO-66 (Figure 4). DFT simulations reveal the important role of the inorganic node in influencing the compression of individual bond lengths upon exposure to high pressures.

Changing the metal that comprises the nodes of a MOF has been shown previously to influence the electronic properties,\(^42\) catalytic activity,\(^43\) and chemical stability\(^44\) of the material. Here, we have presented an interesting example in which altering the metal node of a MOF can have significant
impact on the mechanical properties of the framework. While the response of Zr-Uio-66 and Hf-Uio-66 to pressure is nearly identical, the Ce analogue exhibits drastically different behavior despite the isostructural nature of the three MOFs. These results indicate that the coordination bond between the linker and node of Uio-66 plays a role in the compression of the materials. This study provides insight into how these porous scaffolds respond to high pressures and the structural properties that can be adjusted to modulate the mechanical response of MOFs.

ASSOCIATED CONTENT
Supporting Information. Materials synthesis, ambient condition characterization, diamond anvil cell experimental details, and computational details. "This material is available free of charge via the Internet at http://pubs.acs.org."

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