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A zirconium squarate metal–organic framework with modulator-dependent molecular sieving properties†

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We report the first zirconium metal–organic framework based on squaric acid, representing the member with the smallest unit cell in the isorecticular UiO-66 family. Its molecular sieving properties are strongly influenced by the monocarboxylic acid modulator incorporated during synthesis.

Metal–organic frameworks (MOFs) are hybrid materials consisting of metal ions or clusters bridged by polytopic organic linkers.¹ MOFs often have large specific surface areas and porosities, which can easily be fine-tuned due to their chemical flexibility.^{2,3} In this regard the principle of isorectularity is frequently applied to construct isotopological frameworks with varying pore sizes by using linkers with the same connectivity but different lengths. Well-known examples are the IRMOF-series based on expanded or functionalized versions of MOF-5 and MOF-74,^{4,5} and the cubic MOFs based on the Zr-terephthalate UiO-66.⁶ Recently several publications have highlighted the possibility to engineer the porosity of the latter group of MOFs *via* the incorporation of monocarboxylate modulators or linker fragments such as acetate or benzoate into the framework.^{7–9} During synthesis these modulators aid the formation of the characteristic hexanuclear Zr cluster but also directly compete with the dicarboxylate linkers for binding to it, ultimately leading to a framework in which the linkers are partially replaced by the modulator. Additionally, some modulators can be post-synthetically removed from the lattice.⁹ Both the incorporation and the subsequent removal of modulators can be considered as strategies to generate defects within the framework, resulting in a material with larger pores.

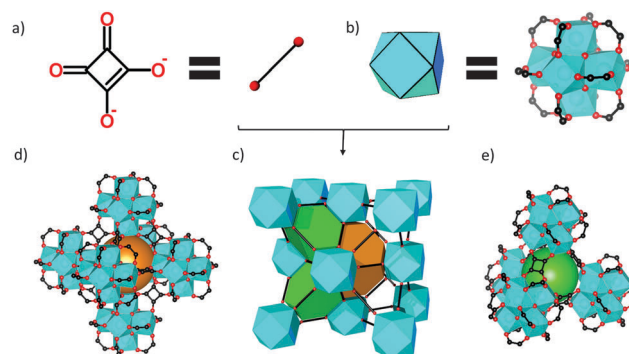


Fig. 1 Structural and topological representation of ZrSQU. SQU linkers (a) and $\text{Zr}_6\text{O}_4(\text{OH})_4(\text{C}_2\text{O}_2)_{12}$ clusters (b) stack in an **fcu**-lattice (c), delineating octahedral (orange, d) and tetrahedral (green, e) cages with triangular windows. (Zr = blue; O = red; C = black).

Here we demonstrate how modulation-based porosity engineering can be extended down the line of the isorecticular UiO-66 series to the point where the intrinsic pore size of the defect-free lattice would be too small to allow for the uptake of guest molecules, rendering such frameworks essentially non-porous. Controlling modulator-induced linker defects provides the key to influencing the porosity of such MOFs. To explore this concept, we applied squaric acid ($\text{C}_4\text{O}_4\text{H}_2$, SQA, Fig. 1a) as a linker for Zr-MOFs. SQA is a non-carboxylic, organic acid which is known to take on a variety of coordination modes with metal ions upon deprotonation to squarate ($\text{C}_4\text{O}_4^{2-}$, SQU).^{10,11} Due to the short distance between coordinating O-atoms (~ 3.2 Å vs. ~ 7.0 Å for terephthalate) only a small number of porous coordination polymers based on SQA have been reported,^{12–16} and no zirconium-based squarate has been described to date. The first Zr-based squarate MOF (ZrSQU), the MOF with the smallest unit cell isorecticular to UiO-66, is reported here. While no accessible porosity is expected based on the crystal structure, we show that the incorporation of small modulators effectively confers molecular sieving properties to this framework. The choice of modulator allows for a subtle control on the size of gas molecules that can be adsorbed.

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† Electronic supplementary information (ESI) available: Experimental details, structure solution and refinement data and crystal structure summary of ZrSQU and HfSQU, FTIR spectra, TGA data, EDX data, gas uptake comparison with UiO-66 and Zr-fumarate. CCDC 1003940 and 1003941. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c4cc03764b



ZrSQU crystallizes as a microcrystalline powder of intergrown aggregates (Fig. 2) upon reaction of squaric acid with ZrCl_4 in a mixture of dimethylformamide (DMF), aqueous HCl and a monocarboxylic modulator, either acetic acid or formic acid, at 110 °C for 2 h (92% yield based on Zr). Similar to several other Zr-MOFs, the presence of a modulator is required to form a crystalline material.^{17–24} By increasing the amounts of HCl and modulator during synthesis, the crystallinity of the formed product can greatly be increased (Fig. S1†). Materials of comparable crystallinity are obtained by using either acetic or formic acid (Fig. S2†). These compounds are hereafter named ZrSQU_A and ZrSQU_F respectively. No effect on particle size is observed when varying the modulator or its concentration, most likely due to the fast nucleation. The isostructural Hf-analogue can easily be obtained by replacing ZrCl_4 with equimolar amounts of HfCl_4 (Fig. S3†).

The structure of ZrSQU was solved from powder XRD data in the cubic space group $Fm\bar{3}m$ ($a = 15.784(3)$ Å) and it is isoreticular to that of UiO-66.⁶ The inorganic building unit consists of six Zr^{4+} -ions each occupying the vertex of an octahedron that is face-capped by $\mu_3\text{-O}$ or $\mu_3\text{-OH}$ groups, while its edges are bridged by twelve C_2O_2^- groups corresponding to half a squarate anion (Fig. 1b). Due to the twelve-fold connectivity the clusters arrange in a face-centered cubic stacking, resulting in a lattice with the **fcu**-topology (Fig. 1c). The two broad reflections around $5.5^\circ 2\theta$, which are forbidden for the $Fm\bar{3}m$ space group, have been observed for other Zr-MOFs. While their origin is not yet fully elucidated, these reflections are allowed for a primitive cubic phase with the same cell parameter as that of the face-centered phase. This in turn could hint at the presence of primitive defect regions within the ZrSQU lattice.^{25,26}

The ZrSQU framework features both octahedral and tetrahedral cages with diameters of 5.6 Å and 4.7 Å respectively (Fig. 1), enclosed by triangular windows. The diameter of the largest sphere that can pass through these windows is approximately 2.4 Å, taking into account the Van der Waals radii of the atoms lining the pores (Fig. S4†). Based on the crystal structure model a void volume of 32.6% (PLATON,²⁷ probe 1.2 Å) was calculated. Residual electron density was found by Fourier analysis in both types of voids and is most likely due to guest

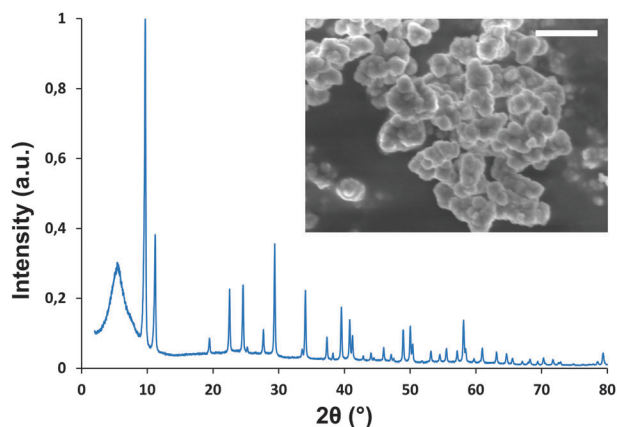


Fig. 2 Experimental powder pattern for ZrSQU. Inset: SEM image of aggregated ZrSQU crystallites (scale bar: 2 μm).

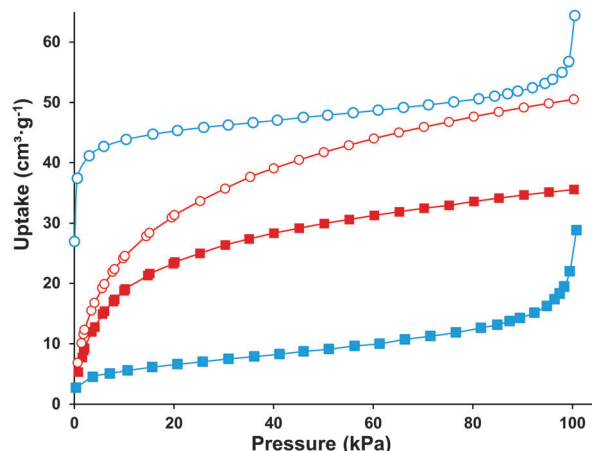


Fig. 3 Adsorption isotherms for N_2 (blue) and H_2 (red) measured at 77 K on ZrSQU_A (square) and ZrSQU_F (circle) outgassed at 10^{-4} bar at 373 K for 4 h.

molecules (water, DMF, modulator) occluded in the cages during synthesis. Following thermal activation under vacuum, N_2 physisorption (Fig. 3) revealed a type II isotherm for ZrSQU_A, which is indicative of a non-porous or macroporous material. ZrSQU_F on the other hand showed a clear type I isotherm reaching a micropore uptake of $40.4 \text{ cm}^3 \text{ g}^{-1} \text{ N}_2$. A multi-point BET surface area²⁸ of $179 \text{ m}^2 \text{ g}^{-1}$ was determined for the latter, taking into account the consistency criteria developed for MOFs.²⁹ Note that theoretically, based on Zeo++ calculations for the ideal crystal lattice (3.861 Å probe diameter), ZrSQU does not have any capacity for N_2 .³⁰ H_2 adsorption demonstrated that both ZrSQU_A and ZrSQU_F can adsorb this smaller probe molecule (kinetic diameter 2.82 Å), with a capacity of $35.6 \text{ cm}^3 \text{ g}^{-1}$ and $50.5 \text{ cm}^3 \text{ g}^{-1}$ respectively at 1 bar and 77 K (Fig. 3). The type I isotherms clearly show that hydrogen is adsorbed in the micropores of both materials.

The fact that ZrSQU is able to adsorb gasses that have kinetic diameters larger than the 2.4 Å window size of the perfect lattice can be explained by the presence of framework defects. To prove that the origin of these defects lies in the incorporation of modulator during synthesis, FTIR measurements were performed on both materials in KBr (Fig. S5 and S6†). Indeed, apart from the broad band centered around 1510 cm^{-1} , which is assigned to the combined $\nu_{\text{C}=\text{C}}$ and $\nu_{\text{C}=\text{O}}$ modes of SQU, clear proof for the incorporation of formate or acetate ions comes from the $\nu_{\text{C}=\text{O}}$ vibration at 1655 cm^{-1} as well as the $\nu_{\text{C}-\text{H}}$ modes at 2965 cm^{-1} and 2861 cm^{-1} . In order to quantify the amount of modulator and thus the composition of ZrSQU, a combined thermogravimetric (TG) and $^1\text{H-NMR}$ approach was used. The TG profiles of ZrSQU_A and ZrSQU_F (Fig. 4) show three distinct steps. A first step from the onset of the measurement up to 210 °C is attributed to the removal of occluded water, solvent and physisorbed modulator. A second step between 210 °C and 350 °C coincides with decomposition of the linker.³¹ The resulting phase was determined to be an amorphous Zr-carbonate by FTIR spectroscopy, which transforms to tetragonal ZrO_2 at 630 °C (Fig. S7 and S8†). While the actual framework decomposition starts at 210 °C, the material loses



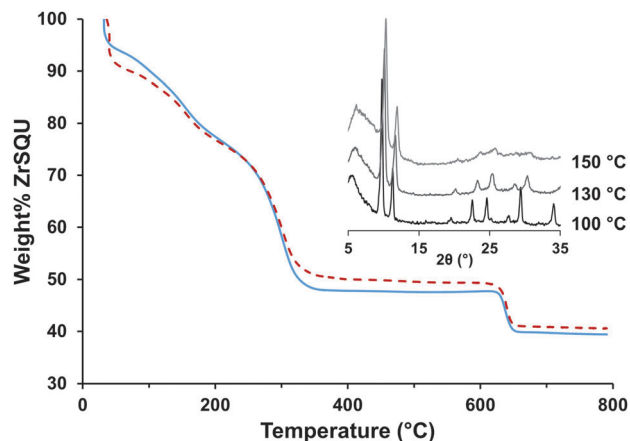


Fig. 4 TG profiles for as synthesized ZrSQU_A (blue solid) and ZrSQU_F (red dash) measured under O₂. The framework loses long range order in the 130–150 °C range with a decrease in cell volume (inset).

some of its crystallinity in the 130–150 °C range. We believe that in this range the pressure exerted by occluded water molecules irreversibly damages the lattice, for instance by partial hydrolysis of the SQU–Zr bond. Because of the strong delocalization of electrons in SQU, the bond between the soft electron-poor SQU anion and the hard Zr⁴⁺ is weaker than the Zr–carboxylate bonds in other Zr-MOFs,³² making such hydrolysis plausible. This partial collapse is irreversible and is accompanied by a decrease of the cell parameter to approximately 14.88 Å. Nevertheless, the 111 and 200 reflections remain clearly visible indicating the persistence of long range order. Pretreating the materials for sorption had a similar effect, with ZrSQU_F being less prone to collapse than ZrSQU_A (Fig. S9†). At room temperature however, ZrSQU is stable in water for at least 3 days (Fig. S10†).

In contrast with earlier observations on some other modulated Zr-MOFs,⁹ a separate step of modulator removal from the ZrSQU framework is not observed in the TG profile. Rather a combined decomposition and modulator volatilization step is observed because of the stronger modulator–cluster bonding. To accurately determine the composition of ZrSQU the activated solids were therefore dissolved in d₆-DMSO using CsF³³ and ¹H-NMR spectra were recorded. We found ZrSQU_A and ZrSQU_F to contain 6.4 wt% acetate and 10.5 wt% formate, respectively. Combining these results with the TG data gives the structural formulae Zr₆O₄(OH)₄(SQU)_{5.25}(CH₃COO)_{1.5} and Zr₆O₄(OH)₄(SQU)_{4.45}(HCOO)_{3.1}. These are approximate formulae which should be regarded as representative models for the actual materials. In EDX measurements, no Cl[–] was detected at all (Fig. S11†); hence it is justified to balance all positive charges of the Zr cations by O^{2–}, OH[–], SQU and modulator. The amount of incorporated modulator can be altered by changing the initial amount present in the synthesis mixture. For instance, a molar ratio of 360/1 acetic acid/ZrCl₄ leads to an incorporation of 9.5 wt% acetic acid in the activated solid, while a 330/1 formic acid/ZrCl₄-ratio leaves 17.1 wt% formate built into the MOF. Clearly, a significant amount of modulator is incorporated during the synthesis of ZrSQU. Formate is able to compete better with SQU than acetate for binding to the

cluster due to the fact that formic acid is a slightly stronger acid than acetic acid. Given the highly acidic synthesis conditions, it is therefore built in in larger amounts. Additionally, as formate is smaller than acetate (Van der Waals volume ~40 Å³ vs. ~57 Å³) it is expected that formate suffers less from sterical hindrance within the framework, allowing more molecules to be incorporated.

As the stoichiometric substitution of one SQU (~79 Å³) for two formate ions does not strongly affect the free volume within the unit cell, we believe that the increased porosity in ZrSQU_F is rather due to a change in size and shape of the triangular windows. The cluster-bound formates could also act as a trap-door by tilting out of plane around their O–O axis, further increasing the window size. The bulkier acetate ions are more sterically constrained, which could explain why ZrSQU_A cannot take up N₂, but is porous for the smaller H₂. This additional sterical bulk further rationalizes the lower H₂-uptake of ZrSQU_A compared to ZrSQU_F. The size-selective uptake of gas molecules in ZrSQU_A, which is absent in its formate-containing counterpart, is a clear demonstration of how selecting the appropriate modulator can have a distinct influence on the molecular sieving properties of the formed MOFs through a subtle alteration of the parent structure. ZrSQU_A could thus be exploited in the purification of small gasses, e.g. H₂ or He. A comparison with other Zr-MOFs is provided in the ESI.†

To conclude, we reported the synthesis and characterization of the first Zr squarate MOF, which is the smallest member of the isorecticular series based on UiO-66. Accessible microporosity is created in this theoretically non-porous material by the incorporation of different modulators during synthesis, the choice of which allows fine-tuning of ZrSQU's gas sorption properties.

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