

Topochemical Synthesis of Single-Crystalline Hydrogen-bonded Crosslinked Organic Frameworks and Their Guest-induced Elastic Expansion

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ABSTRACT: Covalently linked single-crystalline porous organic materials are highly desired for structure-property analysis, however, periodically polymerizing organic entities into high dimensional networks is challenging. Here, we report a series of topologically divergent single-crystalline hydrogen-bonded crosslinked organic frameworks (H_cOFs) with visible guest-induced elastic expansions, which mutually integrate high structural order and high flexibility into one framework. These H_cOFs are synthesized by photo-crosslinking molecular crystals with alkyldithiols of different chain lengths. Their detailed structural information was revealed by single-crystal X-ray analysis and experimental investigations of H_cOFs and their corresponding single-crystalline analogues. Upon guest adsorption, H_cOF-2 crystals comprised of a 3D self-entangled polymer network undergo anisotropic expansion to more than twice their original size, while the 2D-bilayer H_cOF-3 crystals exhibit visible, layered sorption bands and form delaminated sheets along the plane of its 2D layers. The dynamic expansion of H_cOF networks creates guest-induced porosity with over 473 % greater volume than their permanent voids, as calculated from their record-breaking aqueous iodine adsorption capacities. Temperature-gated DMSO sorption investigations illustrated that the flexible nature of crosslinkers in H_cOFs provides positive entropy from the co-existence of multiple conformations to allow for elastic expansion and contraction of the frameworks.

Introduction

Covalently-linked single-crystalline porous organic materials,¹⁻⁴ compared to the coordination-driven assembly of their metal-organic framework (MOF) counterparts,⁵ are rare due to the synthetic challenges of polymerizing monomers into higher-dimensional networks (i.e. 2D and 3D) with precise long-range order. Sustained research efforts have been devoted to the development of single-crystalline porous organic polymers, since their resolved single-crystal structures will provide an atomic-level understanding of the structure-property relationship between the architectures of porous organic polymers and their sorbates. One approach, demonstrated in the syntheses of single-crystalline covalent organic frameworks (COFs), is the ultra-fine optimization of the reversible polymerization-crystallization evolution (e.g. nitrosyl dimerization,² boronic ester formation,³ and imine condensation⁴), despite the reversible nature of the crosslinking reactions posing limitations in their chemical stability. Another approach to the

development of porous materials which has not yet been demonstrated, is the topochemical polymerization of monomers into single-crystalline 1D^{6,7} and 2D^{8,9} polymers through a single-crystal to single-crystal (SCSC) transformation. The precise crystal engineering prerequisite of placing reactive motifs in close proximity, however, limits the scope of available monomers. Since rigid molecular entities connected by covalent bonds are critical to a predictable high dimensional polymerization, these single crystalline COFs and 2D polymers only tolerate a limited degree of structural variation,^{10,11} unlike flexible MOFs¹²⁻¹⁵ constructed with more pliable coordination bonds.

While high structural order and high flexibility are not mutually exclusive,¹⁶ simultaneous integration of these features into porous polymers becomes particularly challenging.^{17,18} To this end, a different approach that polymerizes reactive motifs in a monomer single crystal with crosslinkers that are heterogeneously diffused into the host lattice will provide a unique opportunity to overcome this obstacle

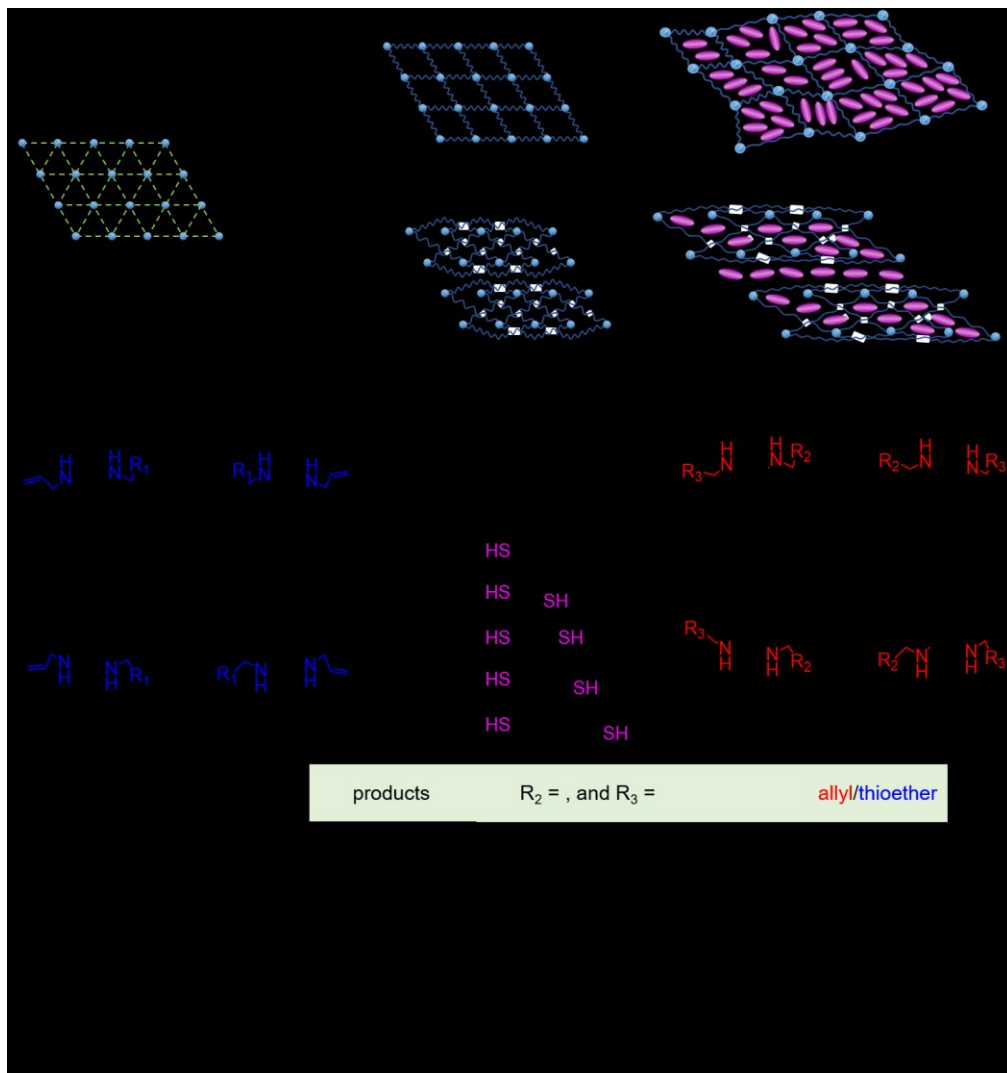


Figure 1. Design and topochemical synthesis of HcOFs and polymers via SCSC thiol-ene polymerization. (a) Schematic representation of crosslinker-dependent topological divergence and corresponding differences in guest sorption in HcOFs-2-4. (b) General synthetic scheme for topochemical SCSC synthesis of HcOFs-2-4, **P5-P8**, and tabulation of products with allyl/thioether ratios determined by NMR spectroscopy and elemental analysis.

and generate single-crystalline porous organic polymers with designed dynamic sorption behaviors. These monomers are pre-organized by hydrogen bonding interactions in the solid state similar to those hydrogen-bonded organic framework¹⁹⁻²³ (HOF) materials, which are photo-cross-linked subsequently via thiol-ene¹ and thiol-yne²⁴ reactions. The feasibility of this approach was first demonstrated¹ by Wuest *et al.*, and later the dynamic sorption behavior²⁴ of the obtained porous crystals was discovered by us. Notably, such SCSC transformations can employ flexible crosslinkers, such as alkyldithiols, in contrast to conventional COF synthesis²⁵⁻²⁸ or topochemical polymerization,²⁹ which utilize rigid building blocks. The flexibility of these dithiols encapsulates a design paradox: on one hand, their flexibility enables extensive diffusion and crosslinking; on the other hand, the resulting flexible linkages pose challenges for single crystal X-ray diffraction (SCXRD) analysis. In essence, this method demands careful balance of the properties that facilitate both dynamic structural variation and the ability to characterize the detailed polymer network architecture and crosslinking topology. So far, the lack of clearly defined

polymer architectures has limited the understanding of their dynamic sorption behavior to little more than empirical observation.

In this report, we demonstrate a comprehensive investigation on the development of single crystalline hydrogen-bonded crosslinked organic frameworks (HcOFs) by photo-crosslinking of molecular crystals with alkyldithiols of different chain lengths (Figure 1). We reveal the first topologically divergent topochemical synthesis of HcOFs synthesized from the same crystalline monomer. The synthesis of a series of oligomeric and polymeric analogues (i.e. **P5-P6**) with a decreased number of crosslinking motifs, and comparison to the corresponding HcOFs, revealed detailed structural information which allowed for the determination of the crosslinking topologies of the HcOFs for the first time. Remarkably, different crosslinking topologies of these HcOFs lead to notably distinctive guest sorption properties (Figure 1a). Upon I₂ and temperature-gated DMSO adsorption, HcOF-2 crystals comprised of a 3D self-entangled polymer network undergo anisotropic expansion to more than

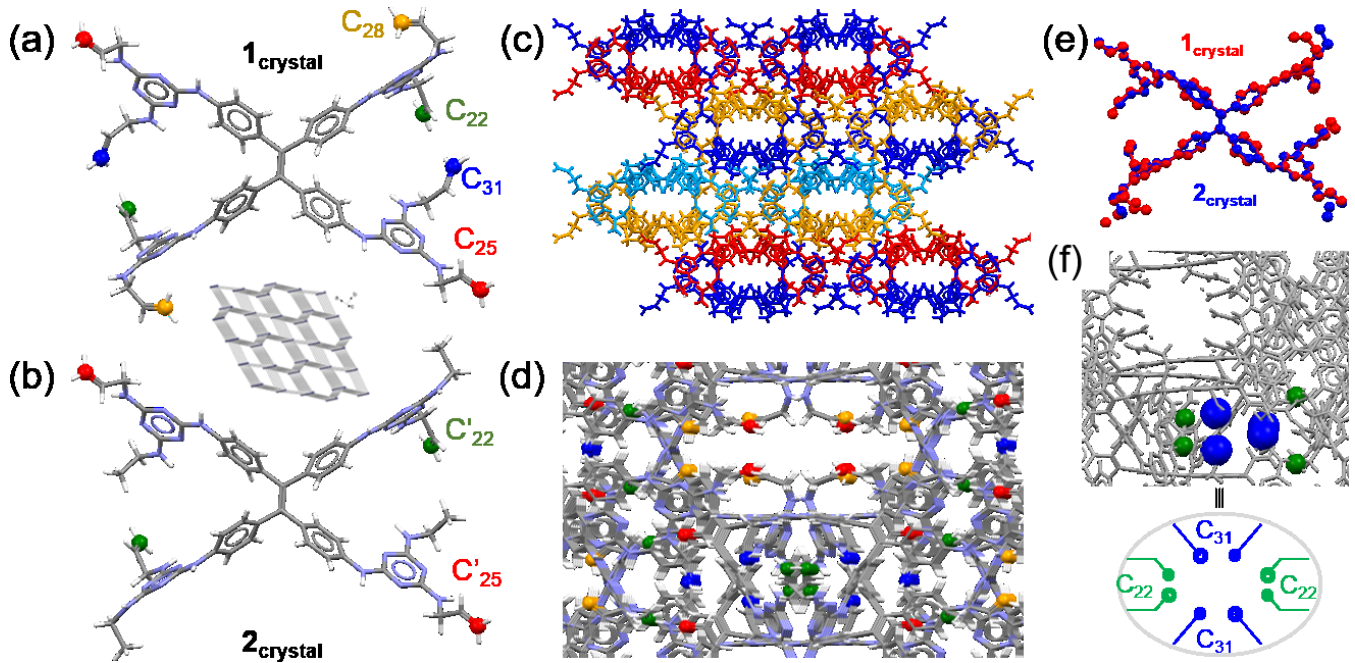


Figure 2. (a-b) Crystal structures and packing diagram of isomorphous (a) **1_{crystal}** and (b) **2_{crystal}** viewed along the [110] direction with simplified underlying six-connected (6-c) net topology (**snw**, inset). The terminal alkene atoms are color coded to highlight their positions in the crystal lattice. (c) Extended packing of **1_{crystal}** viewed along the *b*-axis. (d) Expansion of C₂₅ and C₂₈ allyl groups lining hexagonal pore and congested C₂₂/C₃₁ junction. (e) Structural overlay of **1_{crystal}**/**2_{crystal}** based on calculated RMSD of atomic positions. (f) Highlight of congested C₂₂/C₃₁ crosslinking junction with simplified cartoon representation.

twice their original size, induced by strong substrate-framework interactions. The 2D-bilayer HcOF-3 crystals exhibit visible, layered sorption bands and form delaminated sheets along the plane of its 2D layers. The dynamic expansion of HcOF networks creates guest-induced porosity, which are calculated from their record-breaking iodine adsorption capacities measured in aqueous environment, of more than 473% greater than their permanent voids. We also reveal that the flexible nature of crosslinkers in HcOFs-2-4 provides positive entropy from the co-existence of multiple conformations to allow for elastic contraction of the framework during desorption. These results not only provide a practical approach to achieve high structural order and flexibility in topologically diverse porous organic frameworks, but also enlighten a path for the design of porous organic materials with large guest-induced voids that promise high guest adsorption capacity and selectivity.

Results and Discussions

Monomers **1** and **2**, possessing tetraphenylethylene (TPE) cores with four melamine hydrogen bonding moieties, are synthesized from **S1** through a stepwise nucleophilic substitution with allylamine and ethylamine²⁴ (Figure 1b, Scheme S1). Compared to monomer **1** with eight allyl groups, four ethyl groups on the melamine arms of **2** reduce the number of reactive allyl groups while maintaining a similar spatial occupancy. Slow vapor diffusion of MeCN into DMF solutions of these monomers produced isomorphous **1_{crystal}** and **2_{crystal}** in the orthorhombic space group *Fddd* with nearly identical unit cell parameters (0.8 % difference, Table S3) and packing conformations (Figure 2a-b). In the SCXRD analysis, each monomer interacts with six neighboring monomers through a total of 20 N–H···N hydrogen

bonds formed between their melamine moieties to construct a 3D hydrogen-bonded framework (Figures 2c-d, S37-38). The solvent-filled void spaces of **1_{crystal}** and **2_{crystal}** comprise 21.4 % and 23.0 % of the unit cell volumes, respectively, with 3D open channels exhibiting hexagonal cross sections along the [110] and [1̄10] directions (Figure 2d). The inter-connected 3D channels (Figure S39), suitable pore sizes, and NH-decorated pore surfaces of **1_{crystal}** and **2_{crystal}** allow for extensive diffusion and assembly of linear alkylidithiols throughout the crystals via van der Waals and hydrogen bonding interactions.

From the [110] and [1̄10] perspectives, there are two types of allyl terminal atoms (C₂₅, red and C₂₈, yellow, Figure 2d) located along the hexagonal pore surfaces in **1_{crystal}** (see Table S6 for C···C distances). From the [010] perspective, symmetrically related terminal allyl atoms (C₃₁, blue, Figure 2f) are packed tightly in a clustered junction and are slightly removed from the pore surface. Allyl terminal atoms C₂₂ (green, Figure 2f) are found surrounding the clustered junction and are the least accessible set of allyl groups of monomer **1** as suggested by SCXRD analysis. The spatial distribution of allyl groups of **2_{crystal}** are similar to those of **1_{crystal}**, in which allyl terminal atoms (Figure 2b, C'₂₅, red) are located along the hexagonal pore surfaces and the C'₂₂-allyl groups (Figure 2b, green) in **2_{crystal}** are packed nearly identically to those of the C₂₂-allyl groups in **1_{crystal}**.

To quantitatively assess the accessibility and kinetic reactivity differences of these allyl groups in the solid state, ethanethiol (ET) was used as a thiol-ene probe, which was diffused into **1_{crystal}** and **2_{crystal}** followed by UV irradiation. At different ET diffusion times, averages of 4.8 ± 0.2 (3 d) and 6.7 ± 0.2 (14 d) of the eight allyl groups of **1** were converted

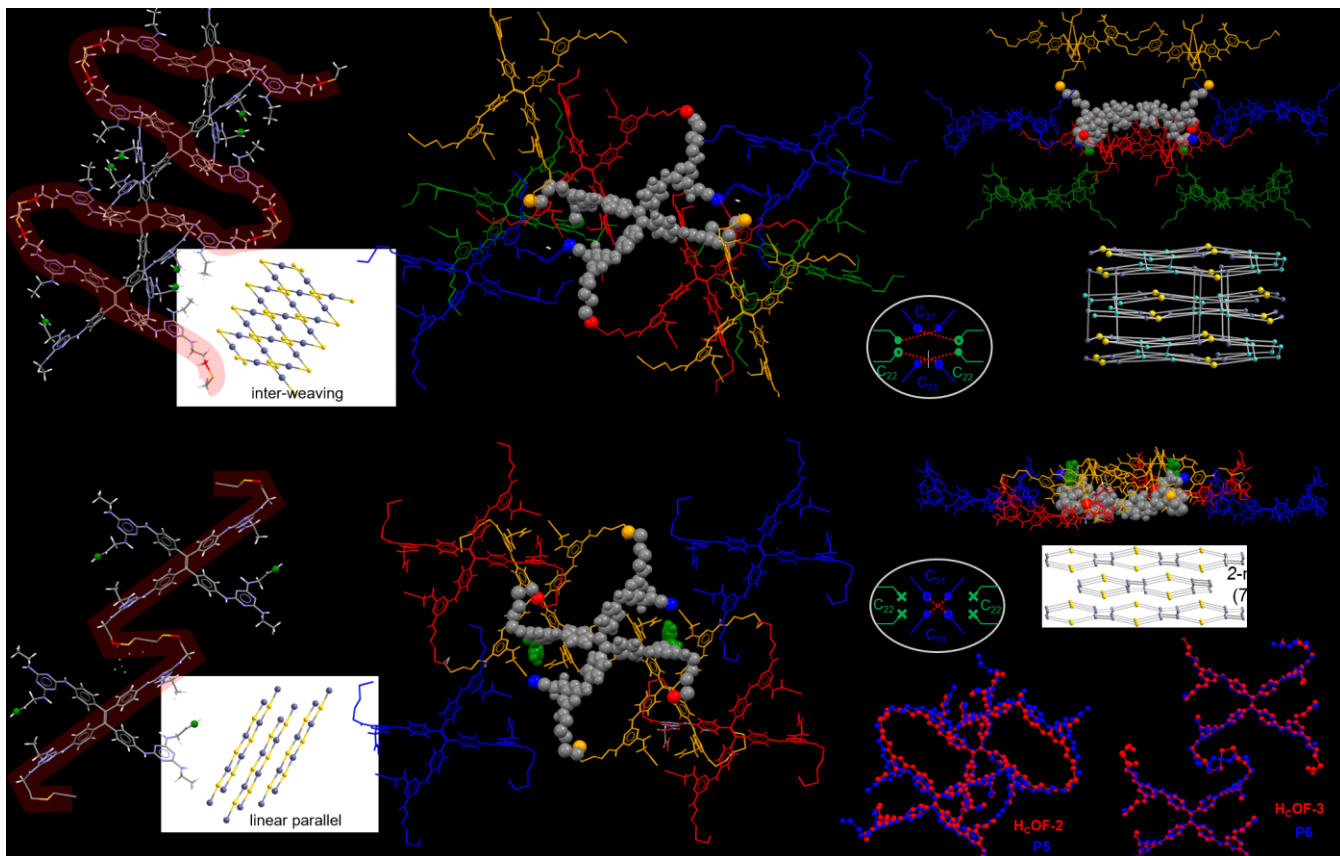


Figure 3. (a-b) Crystal structures of (a) **P5** and (b) **P6** with polymer backbones highlighted in red and polymer topology inset. (c) Packing diagrams of HcOF-2 highlighting the local environment of a tetramelamine-TPE repeating unit (spacefill) viewed along the *b*- and *c*-axes and schematic of crosslinking junction (cartoon inset). Neighboring repeating units (sticks) are colored according to the corresponding allyl carbon to which they are crosslinked (red – C₂₅, yellow – C₂₈, blue – C₃₁, green – C₂₂). (d) Calculated topology of HcOF-2 with TPE (yellow ball) and two topologically distinct melamines (cyan and blue balls). (e) Local packing environment of HcOF-3 repeating units viewed along the *b*- and *c*-axes with schematic of crosslinking junction (cartoon inset). (f) Calculated topology of HcOF-3 with TPE (yellow ball) and melamine moieties (blue ball). (g-h) Structural overlays of (g) HcOF-2/**P5** and (h) HcOF-3/**P6** based on calculated RMSD of atomic positions.

to thioethers (Figure S23), as determined by ¹H NMR spectroscopy. Products with four to eight ET substitutions ([**1**•(4-8)ET]) were identified in the mass spectrum (Figure S21), which revealed a trend of decreasing relative abundance with greater number of substitutions. When **2**_{crystal} was subjected to ET substitution, an average of 3.0 ± 0.1 out of four allyl groups were consumed (Figure S26), producing a mixture of species with two to four ET substitutions ([**2**•(2-4)ET], Figure S22). These experiments confirmed the ET accessibility of all allyl groups in **1**_{crystal} and **2**_{crystal} and suggested that C₂₅/C'₂₅-, and C₂₈/C'₂₈-allyl groups along the pore surfaces are more reactive. When the ET substitution was kinetically controlled (Table S2), crystal samples composed of largely [**1**•(3-4)ET], and trace amounts of other derivatives with an average of 3.8 ± 0.4 allyl-to-thioether conversion were being consistently produced.

Flexible alkyldithiols ($(HS(CH_2)_nCH_2SH)$, including ethanedithiol ($n = 1$, EDT), propanedithiol ($n = 2$, PDT), and butanedithiol ($n = 3$, BDT), are employed as crosslinkers for the synthesis of HcOFs. To mitigate variation in diffusion kinetics, moderate concentrations of alkyldithiol solutions were introduced to the crystals in the dark in three portions followed by UV irradiation, extensive washing, and

supercritical CO₂ activation to afford insoluble single-crystalline HcOF-2, HcOF-3, and HcOF-4, respectively.³⁰ No residual free thiol signal was recorded in the FT-IR and Raman spectra (Figures S66-71) and carbon resonances attributed to dithioether moieties were identified in the aliphatic region (30-45 ppm) of the solid-state ¹³C NMR spectra of HcOFs-2-4 (Figures S18-20), suggesting a complete dithiol-to-dithioether conversion. Although the alkene carbon resonances of **1** disappeared in HcOF-2, residual alkene carbon resonances recorded in HcOF-3 and HcOF-4 suggest some allyl groups remain unreacted after crosslinking. Averaged monomer-to-crosslinker ratios in the supercritical CO₂-activated HcOFs-2-4 were revealed by elemental analysis (Table S1) as $(3.98 \pm 0.13)EDT:\mathbf{1}$ in HcOF-2, $(2.97 \pm 0.03)PDT:\mathbf{1}$ in HcOF-3, and $(2.53 \pm 0.03)BDT:\mathbf{1}$ in HcOF-4, suggesting approximately two and three out of eight allyl groups in **1** remain unreacted in HcOF-3 and HcOF-4, respectively.

When alkyldithiols were diffused into **2**_{crystal}, the soluble crystal samples **P5-P8** were obtained after photo-irradiation.³¹ Dynamic light scattering (DLS, Figure S36) analysis suggests **P5-P8** are composed of polymeric and oligomeric species, in which polymers are the major components of **P5**

($r \sim 50$ nm) and **P6** ($r \sim 70$ nm) whereas oligomers are the dominate species in **P7** ($r \sim 3$ -4 nm) and **P8** ($r \sim 3$ -4 nm). In the NMR analysis (Figures S30-35), dithioethers ($-\text{S}(\text{CH}_2)_n\text{CH}_2\text{S}-$, $n = 1, 2$) are found as the major species in **P5** and **P6**, while mono-hydrothiolated motifs ($-\text{S}(\text{CH}_2)_4\text{SH}$) were abundant in **P7** and **P8**. These results indicate that dithioether crosslinking motifs were consistently formed between periodically aligned pairs of alkenes in the solid-state in **P5** and **P6**. In contrast, a mismatched alkene-to-dithiol distance will lead to a rapid decrease in the degree of polymerization and generate oligomeric species, as was observed in **P7** and **P8**.

Transparent light-yellow crystals of HcOF-2-4 and **P5-P7** were found to retain the orthorhombic space group $Fddd$ by SCXRD analysis (**P8** did not diffract well). Compared to **1_{crystal}**, HcOFs-2-4 demonstrate 2.7 %, -1.9 %, and 1.5 % changes in unit cell volume, respectively (Tables S5). Compared to **2_{crystal}**, **P5-P7** demonstrate -2.3 %, -0.4 %, and -1.3 % changes in unit cell volume, respectively (Tables S4). Refinement of the TPE-melamine entity for HcOFs-2-4 and **P5-P7** is straight forward, which allows for conformational comparisons to those of the monomers by calculation of the root mean square deviation (RMSD) of atomic positions. RMSD analysis (Table S8) reveals that the melamine-TPE scaffold of the HcOFs and **P5-P7** possess nearly identical conformations to their corresponding monomer crystals in the solid state. Among them, the melamine-TPE scaffolds of HcOF-3 and **P6** showed the lowest deviation, suggesting PDT crosslinking can be facilitated with the least perturbation of the framework structure.

In HcOF-2-3 and **P5-P6**, electron density consistent with the presence of dithioethers are observed between the allyl groups (C_{25} , C_{28} , and C_{25}) in the hexagonal pores described in **1_{crystal}** and **2_{crystal}**. Refinement of the electron density revealed one conformer of the dithioether moieties in HcOF-2 and HcOF-3, **P5** and **P6**, while residual electron density in close proximity suggests the existence of multiple conformers. The disorder of dithioether moieties in HcOF-4 and **P7** is too severe for reliable structural analysis. Since **1_{crystal}** and **2_{crystal}** are isomorphous, comparisons between the X-ray structures of the HcOFs and their analogous polymers, i.e. HcOF-2 with **P5**, etc., could confirm the accuracy of the refined cross-linking chains. As shown in Figure 3a, an S-shaped linear polymer **P5** was found with ethanedithioethers bridging the C_{25} -allyl groups of two monomer **2** that are packed in two neighboring layers angled at 120° relative to each other along the [010] direction. These S-shaped polymers then weave into each other along the [001] direction (Figure S42). In contrast, a Z-shaped linear polymer **P6** was formed (Figure 3b) with propanedithioethers connecting the C_{25} -allyl groups of two different monomer **2** that are packed in parallel in the same layer along the [010] direction. The C_{22} -allyl groups of **P5** and **P6** remained unreacted and their atomic positions are found nearly identical to that of **2_{crystal}** in RMSD analysis (Table S8). The conclusion from the SCXRD analysis of **P5** and **P6** is consistent with the formation of linear 1D polymers, as was suggested by NMR and DLS analysis (Figures S30-36).

In HcOF-2, a set of S-shaped dithioether linkages in similar positions to those of **P5** were first identified (red, Figure 3c) between two C_{25} -allyl groups in **1_{crystal}**. The second set of Ω -

shaped dithioether linkages were identified between C_{28} -allyl groups of monomer **1** packed in non-adjacent layers along the *c*-axis (yellow, Figure 3c, right). In HcOF-3 (Figure 3e), a set of Z-shaped dithioether linkages (red) that are similar to those of **P6** were identified between C_{25} -allyl groups of monomer **1**. Another set of S-shaped dithioether linkages (yellow, Figure 3e) were identified between C_{28} -allyl groups of monomer **1** that are packed in two neighboring layers also angled at 120° relative to each other. C_{22} -allyl groups of **1** remained unreacted in HcOF-3 (green, spacefill, Figure 3e), which are consistent with our elemental analysis and solid-state NMR results. Although identifying the diethioether moieties between the symmetry-related C_{22} and C_{31} atoms of HcOF-2 and HcOF-3 (formerly C_{22} and C_{31} allyl groups of **1**) could reveal the crosslinking topologies of HcOF-2 and HcOF-3, refinement of the existing residual electron density around the clustered junction cannot provide unambiguous atomic positions of these linking motifs and, therefore a complete topological assignment is yet to be determined.

Due to the ambiguity of the residual electron density between the C_{22} and C_{31} atoms, SCXRD analysis suggests there may exist seven and three crosslinking possibilities (Scheme S5) for HcOF-2 and HcOF-3, respectively. ET-substituted **1_{crystal}**, composed of largely [**1**•(3-4)ET] with blocked C_{25} - and C_{28} -allyl sites, was employed to reveal the crosslinking taking place at the clustered junction. When [**1**•(3-4)ET] crystals were soaked in neat EDT followed by photo-irradiation, surprisingly, a majority of the EDT-reacted crystal samples were soluble in $\text{DMSO-}d_6$. The ^1H NMR spectrum (Figure S44a) suggests that approximately two allyl groups out of the four residual ones were reacted. Neither polymeric nor oligomeric species were detected in DLS, and monomeric species of [**1**•(3-4)ET•2EDT] were observed in mass spectrum analysis (Figure S45-46). These crystals were washed extensively to remove EDT and were re-subjected to UV-irradiation. This time, the product became insoluble crystals, indicating the formation of a cross-linked polymer network and an extensive allyl-to-thioether conversion. Among the seven linking possibilities, only the connection between non-adjacent C_{22} - and C_{31} -allyl groups (Figure 3c, inset) matches the experimental outcomes (see the SI for detailed analysis, page S55-59). That is, the C_{31} -allyl groups of **1** were substituted with two equivalent EDT in the junction area first, forming soluble monomeric species. The remaining thiol groups subsequently reacted with the non-adjacent C_{22} -allyl groups to form insoluble interlocked 2D networks. When PDT was diffused into [**1**•(3-4)ET] crystals, the photo-irradiated crystals were completely soluble in $\text{DMSO-}d_6$, leaving two unreacted allyl groups and forming linear polymeric species revealed by NMR and DLS analysis (Figures S47-48). The experimental evidence and SCXRD analysis suggest that a homotopic-crosslinking had taken place between C_{31} -allyl groups (Figure 3e, inset, see the SI for detailed analysis, page S61-62), connecting monomers packed in parallel in the same layer. These analyses complement the SCXRD data and allow for topological determination of the complete HcOF-2 and HcOF-3 structures. The fully resolved crosslinking topology of HcOF-2 is revealed to be a 3D interconnected polymer network (Figure 3d), and HcOF-3 as a 2D polymer network (Figure 3f). When the four-connected (4-*c*) TPE core, three-

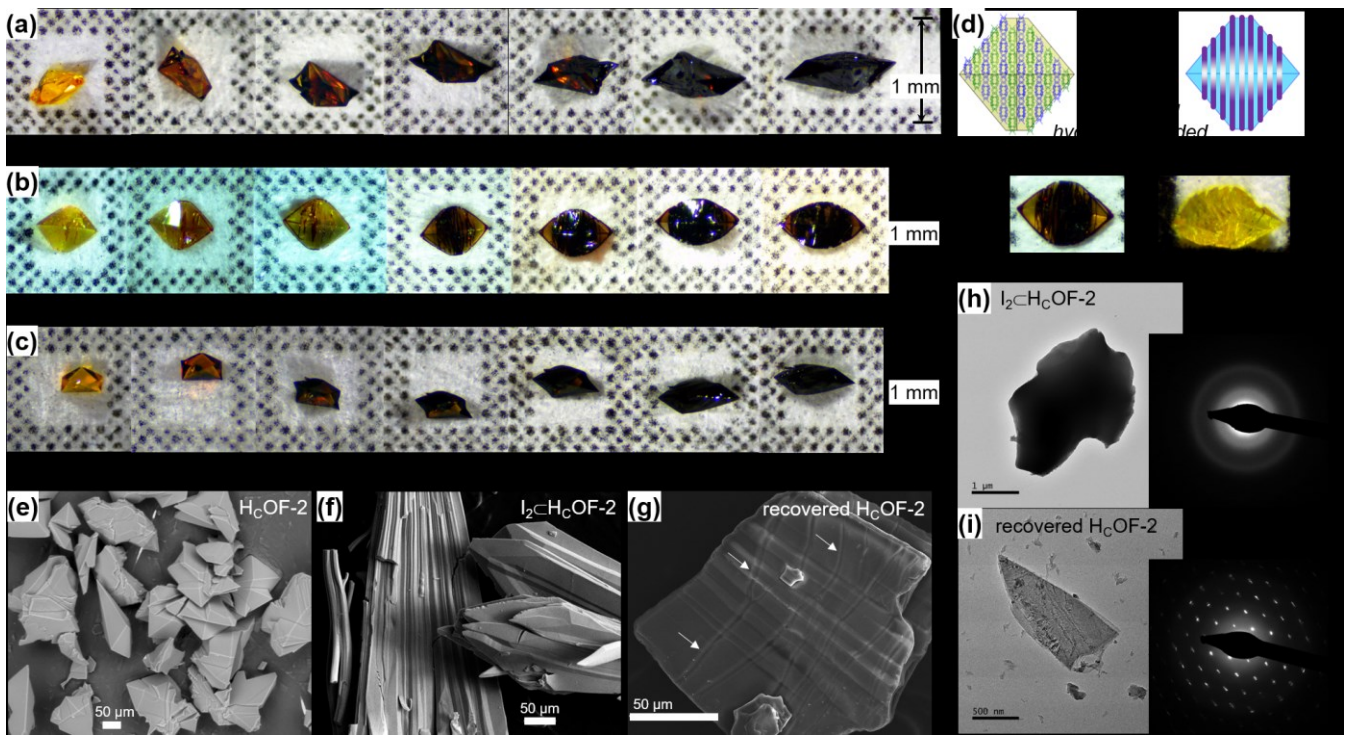


Figure 4. Anisotropic expansion and contraction of the HcOFs crystal sizes upon guest sorption and structural elucidation of the dynamic behavior of HcOFs. (a-c) Images of crystal samples of (a) HcOF-2, (b) HcOF-3, and (c) HcOF-4 immersed in methanolic solution of iodine (50 mM) for different times. The crystals were placed in a 1×1 mm square box for size comparison. (d) Images of crystals of $I_2 \subset H_cOF-3$ and $DMSO \subset H_cOF-3$, and the simulated crystal morphology of an HcOF-3 crystal (void spaces are highlighted in yellow). (e-g) SEM images of (e) HcOF-2, (f) $I_2 \subset H_cOF-2$, and (g) recovered HcOF-2 after iodine desorption. (h-i) TEM images and electron diffraction patterns of (h) $I_2 \subset H_cOF-2$ and (i) HcOF-2 after I_2 desorption.

connected (3-c) melamine moieties, and the dithioether moieties are defined as nodes, topological analysis suggests that (i) HcOF-2 is a 3-nodal 3,3,4-connected self-entangled 3D network with stoichiometry $(3\text{-}c)_2(3\text{-}c)_2(4\text{-}c)$ and point symbol of $(6.10.12)_2(6.12^2)_2(6^2.10.12^3)$; and (ii) HcOF-3 is a 2D 3,4-connected net with stoichiometry $(3\text{-}c)_2(4\text{-}c)$ 2-nodal $(7.8^2)_2(7^3.8^3)$ network, in which two neighboring packed layers of **1** are covalently crosslinked and extended along the [010] plane.

Due to their narrow pore openings and small voids, HcOFs-2-4 demonstrate no N_2 uptake at 77 K and only adsorb small amounts of CO_2 at 273 K (Figure S98). Remarkably, record-breaking aqueous iodine uptake capacities of 3.23 ± 0.18 g, 3.00 ± 0.09 g, and 3.57 ± 0.10 g of I_2 per gram of HcOF were recorded³² for HcOFs-2-4 (Table S9), respectively, which exceed the previous record²⁴ set by HcOF-1. Based on the adsorption values, the calculated van der Waals volumes^{33,34} of adsorbed I_2 (26633 \AA^3 for HcOF-2 and 23878 \AA^3 for HcOF-3) exceed the available void space of HcOF-2 and HcOF-3 by 473 %, and 668 %, respectively, requiring drastic volumetric expansion of these HcOF crystals to accommodate I_2 molecules as guests! When an HcOF-2 crystal ($\sim 0.50 \times 0.25$ mm, Figure 4a) was immersed in a methanolic I_2 solution,³⁵ gradual anisotropic expansion of the crystal and a directional I_2 diffusion path were observed. After 8 h, the sample had expanded to a much larger size ($\sim 0.50 \times 1.20$ mm) while maintaining a single cohesive shape and developing a visibly darker color. Similar anisotropic I_2 -induced expansion was also recorded in HcOF-4 (Figure 4c).

The I_2 -induced anisotropic size expansion of HcOF-3 is noticeably slower, and visible parallel linear I_2 diffusion paths were observed orthogonal to the axis of greatest expansion (Figure 4b and 4d). Since the morphology of this HcOF-3 crystal matches well to its calculated Bravais-Friedel-Donnay-Harker (BFDH) morphology (Figure 4d), the experimental observation suggests that I_2 diffuses between the crosslinked 2D bilayers resulting in de-lamination of the crystal.

Scanning electron microscopy studies (SEM, Figure 4e-g) revealed that diamond-shaped crystals of HcOF-2 were transformed into bundles of $I_2 \subset H_cOF-2$ as a result of the size expansion, and energy dispersive X-ray (EDX) spectra (Figure S56) confirmed the presence of large quantities of iodine. The dark crystal-like $I_2 \subset H_cOF-2$ exhibited poorly defined diffraction patterns by transmission electron microscopy (TEM, Figure 4h) and powder X-ray diffraction (PXRD, Figure 5c), suggesting a crystalline-to-amorphous phase transition. The adsorbed iodine could be released by washing with DMSO, ethanol, and water, resulting in gradual size shrinking as the dark crystals turned to a light-red (Figure S74). The recovered HcOF-2 samples possess wrinkles (Figure 4g) generated by the expansion/contraction process and some of the recovered HcOF-2 samples in TEM analysis (Figure 4i) are found to recover their periodic alignment as shown in the electron diffraction pattern. Similar phenomena were also observed in HcOF-3 (Figure S57-61).

When I_2 molecules were adsorbed in HcOFs-2-4, the carbon resonances assigned to the triazine ring and aliphatic

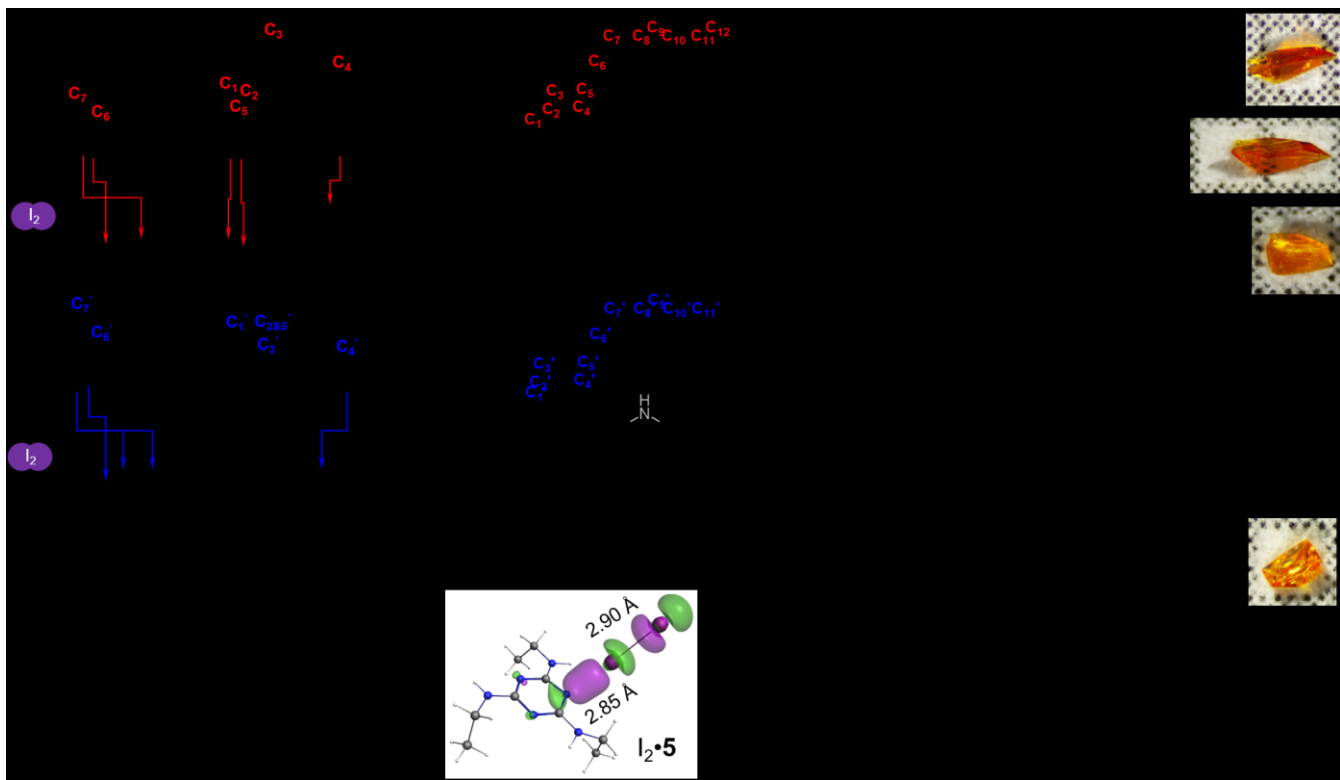


Figure 5. (a) Solution-phase ¹³C NMR spectra of model compound **3** (top, 6.53 mM) and solid-state ¹³C NMR spectra of HcOF-2 (bottom) in the absence and presence of I₂ (14.0 equivalent to **3**). (b) Chemical structures of **4-5**, binding affinity of 3I₂•**4**, and DFT calculated I₂•**5** with calculated N-I and I-I bond distances of 2.85 Å and 2.90 Å, respectively. (c) PXRD profiles of (from bottom up) **1**crystal, supercritical CO₂ activated HcOF-2, I₂ adsorbed HcOF-2 with an uptake of 0.2 g/g I₂, I₂ adsorbed HcOF-2 with an uptake of 3.2 g/g of I₂, DMSO-soaked HcOF-2 at 50 °C, DMSO-soaked HcOF-2 at 70 °C, and air cooled DMSO-soaked HcOF-2 at 25 °C after heating.

crosslinkages shifted upfield (up to 15 ppm) in the solid-state ¹³C NMR spectra (Figure 5a), suggesting that the I₂-induced crystal size expansion is attributed to a change of the local environment of the melamine motifs. The detailed I₂-framework interaction is revealed through the investigation of model compounds **3-5** with I₂ (Figure 5a and 5b, Figure S80-83). In DMSO-*d*₆, **3** binds multiple I₂ molecules, the -NH- proton resonances of the melamine moieties of **3** shift downfield, and triazine carbon resonances shift upfield, similar to those phenomena in HcOF-2. A strong binding affinity of $K_a = 5.7 (\pm 2.7) \times 10^3 \text{ M}^{-1}$ was measured between **4** and I₂ (in 1:3 stoichiometry, Figure 5b) in DMSO-*d*₆ at 298 K, despite the large excess of DMSO competing with I₂. Density functional theory (DFT) calculations (Scheme S9, Table S10, Figures S78-79) revealed that strong halogen bonds are formed between the lone pair of triazine nitrogen atoms and σ^* orbital of I₂ in the I₂•**5** complex, with a high binding enthalpy (~12 kcal/mol per N-I bond). The DFT and NMR studies suggest that, when I₂ is adsorbed into HcOFs, the strong N-I halogen bonding interactions outcompete the original hydrogen-bonded network. The flexible crosslinkers in HcOFs are thus stretched and -NH- moieties are rotated to expand the polymer network and accommodate more I₂ molecules without breaking any covalent bonds.

When DMSO is employed as a weaker hydrogen bonding network disrupter, less extensive framework expansion and contraction of HcOFs was observed (Figure 5c). Interestingly a temperature-gated crystal size expansion and

contraction was observed when HcOF crystals were soaked in DMSO, and the crystalline-to-amorphous and amorphous-to-crystalline phase transitions were recorded by PXRD (Figures 5c, S90-92). After heating HcOF-2 at 50 °C, HcOF-3 at 70 °C, and HcOF-4 at 70 °C in DMSO, crystal samples of HcOF-2 expanded anisotropically (Figure 5c, inset), while the crystal samples of HcOF-3 de-laminated into sheets (Figure 4d). After cooling these crystal samples to room temperature, the long-range order of the molecular entities of HcOFs-2-4 recovered as shown in their PXRD profiles (Figure 5c, S90-92). The different I₂ uptake capacities and phase-change temperature in DMSO between HcOF-2 and HcOF-3 also highlights the significant contribution from the different crosslinking topologies, despite the common monomer crystal and identical hydrogen bonding network from which these HcOFs are derived.

From a thermodynamic perspective, when the guest molecules are desorbed from the framework, the reformation of the hydrogen bonds between the melamine groups directs the framework recovery. Depending on the extensiveness of the framework expansion, the melamine groups may form kinetically stable, randomly oriented hydrogen bonding networks to maximize the network entropy rather than recovering to the highly ordered original form. This analysis highlights the important contribution from the flexible (disordered) crosslinkers. During the elastic expansion process, the flexible crosslinkers are stretched to entropically disfavored conformations. During the framework recovery

process, the stretched flexible crosslinkers return to their disordered state and contribute positive entropy for the reformation of the highly ordered framework. Similar structures are also found in many proteins³⁶⁻³⁹, where the amorphous domain of the protein enables the rigid domain to dynamically alter its functions. Hence, although the co-existence of multiple conformations of the dithioether motifs in H_cOFs increases the complexity for SCXRD analysis, the integration of disordered linkages as molecular level 'amorphous domains' enables the elastic expansion and contraction of the framework materials. In addition, the guest-activated dormant voids in H_cOFs are inversely related to the crosslinking density of H_cOFs, similar to the swelling behavior of traditional crosslinked polymer networks.⁴⁰ For example, H_cOF-2 possesses a higher I₂ adsorption capacity compared to the more densely crosslinked H_cOF-1, despite the permanent voids of H_cOF-2 being significantly smaller than those of H_cOF-1.

Conclusions

In summary, we demonstrated a topochemical strategy to synthesize single crystalline porous organic frameworks H_cOFs-2-4 with divergent crosslinking topologies through a single-crystal to single-crystal transformation. Although the tetraphenylethylene and melamine moieties of these H_cOFs remain largely undisturbed compared to their parent molecular crystal, the disordered dithioether linkages generated after photo-crosslinking obstruct the molecular-level structural analysis of current, and likely future, H_cOFs consisting of flexible crosslinkers. We overcome this obstacle by (1) synthesizing two isostructural single-crystalline polymer analogues to the H_cOFs (**P5** and **P6**), followed by (2) investigating the crosslinking outcomes using crystal samples with partially blocked allyl sites. The unveiled topological assignments of H_cOF-2 and H_cOF-3 suggest them to be a self-entangled 3D network and a 2D-bilayer crosslinked polymer, respectively. H_cOFs-2-4 demonstrated superior iodine uptake capacities (up to 3.5 g/g) and significant visible crystal size expansions (> 200 %) in an aqueous environment, and the different guest sorption capacities and morphology changes of the H_cOFs comes from the different crosslinking topologies. These results suggest that the flexible dithioether linkages in H_cOFs can give rise to a much larger dormant void activated by the presence of guest molecules. The binding energy difference between the substrate-framework (I₂ with melamine moieties) and the hydrogen-bonded framework of H_cOFs determines the energy barrier to activate the guest-induced voids, as observed in their temperature-gated DMSO sorption behaviors. Our investigations not only point out that incorporating flexible crosslinking moieties in H_cOFs is critical to facilitate large dynamic structural variation without sacrificing the highly ordered structure of H_cOFs, it also illustrated a clear structure property relationship in H_cOFs, revealing that the crosslinking topology plays a key role in determining the guest-induced void space. We believe that these research outcomes not only demonstrate a topologically-controlled synthesis of H_cOF materials with integrated high structural order and flexibility, but, more importantly, reveal a molecular-level understanding of the roles of 'ordered' and 'disordered' segments of H_cOFs in relation to their dynamic guest sorption behaviors. Such insights lay the foundation for

future H_cOF designs and, more broadly, the design of crystalline porous organic framework materials with dynamic sorption behaviors for both high uptake capacity and selectivity.

ASSOCIATED CONTENT

Supporting Information. The Supporting Information is available free of charge via the Internet at <http://pubs.acs.org>. Syntheses and characterizations of monomers, H_cOFs and polymers, SCSC transformations, single-crystal and powder X-ray diffraction analysis, guest sorption investigations, DFT calculations, and NMR studies (PDF)
Crystal structures (CIF)

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(32) The iodine uptake capacities of H_cOFs-2-4 were measured in aqueous KI₃ solutions at room temperature, which reversibly release I₂ upon H_cOFs' adsorption. These iodine uptake capacities have been cross verified by gravimetric, elemental, and thermogravimetric analysis, which holds the record of iodine adsorption in the aqueous environment. A higher iodine uptake capacity of 6.2 g/g has been reported in iodine gas vapor, see Wang, P.; Xu, Q.; Li, Z.; Jiang, W.; Jiang, Q.; Jiang, D. Exceptional iodine capture in 2D covalent organic frameworks. *Adv. Mater.* **2018**, *30*, 1801991. Related iodine adsorption values obtained using other porous materials are summarized in Table S11 in the Supporting Information.

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