Covalent organic frameworks (COFs) have emerged as a versatile material platform for such applications as chemical separations, chemical reaction engineering, and energy storage. Their inherently low mechanical stability, however, frequently renders existing methods of pelletization ineffective, contributing to pore collapse, pore blockage, or insufficient densification of crystallites. Here, we present a process for the shaping and densifying of COFs into robust centimeter-scale porous monoliths without the need for templates, additives, or binders. This process minimizes mechanical damage from shear-induced plastic deformation and further provides a network of interparticle mesopores that we exploit in accessing analyte capacities above those achievable from the intrinsic COF structure. Using a lattice-gas model, we accurately capture the monolithic structure across the mesoporous range and tie pore architecture to performance in both gas-storage and -separation applications. Collectively, these results represent a substantial step in the practical applicability of COFs and other mechanically weak porous materials.

INTRODUCTION

Porous materials capable of reducing both the cost and energy intensity of industrial chemical processes are critically needed in transitioning to a carbon-neutral energy cycle. Constructed from Earth-abundant elements and affording a combination of chemo-structural diversity, ease of synthetic modification, and relative chemical stability, covalent organic frameworks (COFs) have emerged as attractive alternatives to existing porous materials, including activated carbons, zeolites, and metal-organic frameworks (MOFs). However, a technological limit has been reached where traditional methods of adsorbent post-processing are poorly suited to COFs as a result of the low mechanical stability frequently exhibited by these materials. To date, these mechanical characteristics have been shown to limit the pressures that can be used during pelletization and the selection of fluids available for activation, deviations from which can result in sharp losses in capacity. Although factors such as framework topology and linker length can be synthetically tuned to target more robust architectures, the inverse approach—i.e., whereby a desired COF can be shaped into an industrially relevant form factor without compromising key performance metrics—has not been attempted.
Here, we report a simple and rapid process for the shaping of COFs into macroscopic pellets without the use of binders, templates, or additives and without any further processing steps needed for a final application. For an archetypical two-dimensional (2D) COF, TPB-DMTP-COF, we demonstrate control over the degree of aggregation of crystallites within pellets and systematically identify the presence of a lower limit in intercrystallite pore size for a given activation solvent. We tie this limit to the onset of capillary-action-induced, turbostratic disordering of crystallites and further confirm that mechanical damage can be avoided through the use of an activation fluid with an ultra-low surface tension. COF monoliths thus prepared are mechanically robust and exhibit low-pressure adsorption characteristics identical to those of the best-reported powder analogs. They additionally benefit from a system of interparticle mesopores that push final adsorption capacities above levels expected for single crystals. We capture these structural characteristics in a lattice-gas model, which accurately reproduces experimentally derived isotherms for COF monoliths in silico. The combination of intact crystallites, mechanical robustness, high bulk densities, and regular hierarchical mesopores is unique among COF monoliths demonstrated to date. These properties result in industrially suitable monoliths that afford better gas-adsorption performance characteristics for both pure-component gas storage (CO₂ and CH₄) and mixed-gas chemical separation (CO₂/N₂ and CO₂/CH₄) applications than do unprocessed powder controls. On the basis of these findings, our work provides not only a path forward for the industrial applicability of COFs but also a systematic framework through which COF microstructure and final adsorption properties can be tuned without altering the underlying COF chemistry.

RESULTS AND DISCUSSION

High-throughput calculation of mechanical properties

2D COFs are thought to be unstable to in-plane mechanical shear. To evaluate the mechanical properties of COFs across topologies and linkage chemistries, we first performed a high-throughput screen of all reported COFs as inventoried in the CURATED (Clean, Uniform, Refined with Automatic Tracking from Experimental Database) COF database and compared their bulk moduli, shear moduli, and elastic constants with those of MOFs (Figure 1). Within a largest cavity diameter (LCD) range of 15–40 Å, the bulk and shear moduli of COFs were found to be similar to those of MOFs, although COFs exhibited marginally higher bulk moduli and shear moduli on average. However, at lower LCD ranges characteristic of ultramicroporous (<7 Å) and microporous (<20 Å) materials, the bulk and shear moduli of COFs were found to be substantially lower than those of MOFs, suggesting an inherently greater tendency of COFs to mechanically deform even in the absence of larger (>15 Å) pores. To gain insights into the mechanical stability of these materials, we then analyzed the elastic constants (cij) of a representative COF subset, hexagonal 2D COFs, which currently account for 54% of 2D COFs and 45% of all COFs synthesized to date. Applying the stability criteria, c₁₁ > |c₁₂|, c₂₂(c₁₁ + 2c₁₂) > 2(c₁₁)², c₁₁c₂₂ > (c₁₂)² and c₄₄ > 0, we found a majority (64%) of hexagonal 2D COFs to be unstable, confirming weakness to mechanical shear as a predominating feature of these materials and possibly shedding light on the low degrees of crystallinity frequently exhibited by these materials. Given that conventional methods of powder pelletization routinely employ pressures in the range of 1–3 GPa, which are known to trigger losses in capacity within MOFs, we sought a revised approach for COF processing and pelletization.

Sol-gel processing of COFs

To permit ease of experimental benchmarking and analysis, we identified TPB-DMTP-COF as a representative 2D COF with an LCD of 25 Å and excellent known

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crystallinity. Upon screening a variety of synthesis solvent systems, we identified acetonitrile and a 1:1 (v/v) mixture of 1,3,5-trimethylbenzene (mesitylene) and 1,4-dioxane (dioxane) as two systems capable of both solubilizing the starting materials and producing crystalline samples of TPB-DMTP-COF. However, whereas the 1:1 (v/v) mixture of mesitylene and dioxane produced powder samples consisting of aggregated particles > 500 nm in diameter (Figures 2E and S5A), the acetonitrile system produced dense pellets consistent with those previously described for MOF monoliths and composed of particles of approximately 40 nm in diameter (Figures 2E and S5H)—well within the limits previously established for monolith formation in MOFs17,18 (i.e., <120 nm). Taking these two systems as extremes, we used solvent compositions consisting of different fractions of each to prepare pellets as follows: (1) reaction for a fixed amount of time (typically 30 min), (2) centrifugation, (3) purification and solvent exchange to methanol, and (4) controlled drying and activation (Figure 2).

Upon processing, scanning electron microscopy (SEM) of the finished pellets revealed a gradual progression in microstructure from larger, loosely aggregated particles to densely packed monoliths exhibiting conchoidal fracture and little to no interparticle free volume (Figures 2E and S5). Analysis of the nitrogen adsorption isotherms (Figures 3A and S3) collected for these pellets, however, revealed a striking
 Whereas we observed a monotonic increase in Brunauer-Emmett-Teller (BET) area—calculated by BET surface identification (BETSI)—for pellets synthesized in solvent systems containing acetonitrile fractions ranging from 0.000 to 0.750 (v/v), we observed a sharp decrease in BET area to 4 m$^2$ g$^{-1}$ for samples prepared at higher acetonitrile fractions (Figure 3C). As a result, the highest BET area that could be obtained for TPB-DMTP-COF with methanol as the activation solvent was 1,122 m$^2$ g$^{-1}$, suggesting the presence of a lower limit in intercrystallite pore size beyond which pore disruption takes place. To test whether this pore disruption was being induced by capillary action, we prepared a further sample in a pure acetonitrile solvent system and processed it as before but this time dried and activated it in supercritical carbon dioxide (scCO$_2$) instead of in methanol and air. The finished pellet not only recovered full porosity but also lay on the monotonic trend previously described.
Figure 3. Structural characterization of TPB-DMTP-COF monoliths

(A) N₂ adsorption isotherms at 77 K for the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) TPB-DMTP-COF monolith (turquoise squares) and powder (gray diamonds); the inset shows the respective mercury pore-size distributions (PSDs) of the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) TPB-DMTP-COF monolith (turquoise) and powder (gray).

(B) Indentation load versus penetration depth for 56 indents in the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) TPB-DMTP-COF monolith; insets are optical micrographs showing the array of residual indents taken in different areas of the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) TPB-DMTP-COF monolith. Scale bars represent 100 μm.

(C) BET area versus acetonitrile fraction (v/v) of methanol-activated samples (turquoise squares), samples with mechanical damage (white triangles), and supercritically activated (3 bar h⁻¹) sample (yellow star).

(D) SAXS data of the supercritically activated (3 bar h⁻¹) TPB-DMTP COF monolith (turquoise) and powder (gray). The inset shows the characteristic Bragg peaks in the WAXS region; the monolith curve has been offset along the y-axis for clarity.

(E) Indentation modulus and hardness (inset) versus penetration depth averaged over 56 indents. Error bars represent standard deviations calculated from 56 measurements made for the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) TPB-DMTP-COF monolith.

(F) Comparison of density (g cm⁻³) and BET area (m² g⁻¹) values for different COF aerogels, monoliths, and pellets synthesized in the literature. Filled symbols are for COF aerogels, monoliths, and pellets synthesized without the use of any binders, additives, or high pressure: ◆, 3 bar h⁻¹ scCO₂-activated 1,000 (v/v) TPB-DMTP-COF monolith (this work); ◈, TFPT-HZ-COF; ▲, TPT-HZ-COF; ▼, TAPA-TFPA; □, TAPB-PDA.
in that it exhibited a BET area of 2,125 m² g⁻¹—slightly above those previously described for powder analogs of TPB-DMTP-COF. When a higher rate of scCO₂ pressure release (8 versus 3 bar h⁻¹) was used during the activation of an identically prepared 1.000 acetonitrile pellet, a reduction in BET area to 1,439 m² g⁻¹ was observed, further suggesting that losses in BET area occur as a result of damage induced by capillary action. Collectively, these results suggest that intercrystallite pore size is modulated during synthesis by control over particle size—where interparticle pores are the void space created by the fundamentally imperfect packing of approximate hard spheres. Where characteristic interparticle pore size falls below a certain threshold, capillary action during drying causes damage to crystallites.

To gain deeper insights into the structural changes accompanying these bulk characteristics, we used a combination of pair distribution function (PDF) and X-ray diffraction (XRD) (supplemental information section S6). Non-negative matrix factorization of the PDF-XRD data revealed three independent underlying components that we attribute to non-crystalline-layer COF content, residual starting-material content, and multilayer (i.e., crystalline) COF content: components A, B, and C, respectively (Figure 3G). For methanol-activated pellets below an acetonitrile fraction of 0.75, a respective decrease and increase in components A and C were observed as the acetonitrile fraction was increased, indicating that TPB-DMTP-COF crystallinity gradually improves before the onset of mechanical damage. Above an acetonitrile fraction of 0.75 (v/v), crystallinity sharply declines, resulting in an increased content of non-crystalline-layer TPB-DMTP-COF, as seen from the increasing weighting of component A. When scCO₂ is used during drying and activation, the multilayer content is recovered—an observation consistent with findings from nitrogen adsorption studies and providing clear evidence for a correlation between mechanical disruption of COF crystallites during post-processing and observable gas-uptake capacities as previously noted for powdered COF systems.

To better understand the mechanism of crystallite disordering into non-crystalline layers within COF monoliths, we performed a combination of high-resolution transmission electron microscopy (HR-TEM) and NanoBlitz indentation studies on a methanol-activated 1.000 acetonitrile control sample for which crystallites are sufficiently disrupted to afford a BET area of 4 m² g⁻¹ (Figure 4). Analysis of the microstructure both within the bulk and within a few layers of the monolith indicated a series of small multilayer crystalline domains bridged by less-ordered regions. Fourier transforms of the image (Figure 4C, inset) further revealed that these features result in a single diffuse band corresponding to a real-space length of 0.36 nm—consistent with interlayer spacing values obtained from analysis of components A (0.37 nm) and C (0.35 nm) derived from the PDF-XRD data. These results suggest that a crystalline-to-turbostratic-disordering mechanism, similar to that observed in mechanically milled graphite, might be responsible for losses in observed porosity within monoliths. At the macroscopic level, NanoBlitz indentation mapping revealed heterogeneities in both the indentation modulus and the mechanical hardness across a 200 × 200 μm region of the material. Given that turbostratic disordering is triggered by capillary action, differences in the local structure of the monolith during drying...
could give rise to regions of greater or lesser disruption, resulting in macroscopic domains with slightly differing mechanical properties in the finished pellet. Collectively, these findings both confirm the presence of disrupted crystallites in non-porous monoliths and suggest that a turbostratic disordering mechanism is responsible for such observable losses in porosity.

After establishing post-processing conditions capable of explicitly avoiding crystallite damage, we then used a combination of mercury porosimetry and small-angle X-ray scattering (SAXS) to gain insights into the structure of monolith free-volume elements across the mesoscale. Pore-size distributions derived from mercury intrusion curves for a scCO₂-activated 1.000 acetonitrile monolith revealed the presence of sharp mesoporosity at 18.7 nm attributable to narrow and regular interparticle...
free-volume elements (Figure 3A, inset). Broader macroporosity centered at a pore width of around 3 μm was also observed. By contrast, a non-monolithic powder control prepared via the method of Xu et al. (BET area of 1,985 m² g⁻¹) exhibited no regular meso- or macroporosity. Analysis of the respective mercury areas for accessible pore widths down to 3.9 nm (above that of the intrinsic framework, i.e., 2.5 nm) further showed an area of 504 m² g⁻¹ for the monolith and 196 m² g⁻¹ for the powder. These results are comparable to those for classical mesoporous templated silicas and carbons and are consistent with those derived from SAXS (Figure 3D and supplemental information section S7). The scCO₂-processed monolith was well fit by a spheroidal particle model with two log-normalized-distribution models with mean diameters of 25.8 nm (σ = 0.4) and 99.8 nm (σ = 0.2), indicating the presence of mesoporous interparticle free-volume elements and providing evidence for additional macroporosity. By contrast, the non-monolithic powder control was found to possess an interparticle size distribution beyond the 0.5–100 nm range and, consequently, could not be fitted. These results suggest that COF processing into monoliths can not only be used to avoid pore collapse but also provide additional mesoporosity (inaccessible from powders) that can be used for tuning final uptake performance characteristics—potentially beyond those of purely crystalline systems.

To examine the impact of crystallite disordering on mesoporous free-volume elements, we also analyzed a scCO₂-processed monolith activated at an accelerated depressurization rate of 8 bar h⁻¹ (BET area of 1,439 m² g⁻¹) by using SAXS (Figure S4). The sample was fit by three spheroidal size-distribution models exhibiting mean diameters of 14.7 nm (σ = 0.3), 21.1 nm (σ = 0.6), and 98.5 nm (σ = 0.1). The emergence of a third, narrow free-volume element along with an overall shift in mesopore distribution to smaller values suggests that disruption of crystallites is concomitant with a reduction in interparticle free volume. Because this reduction in interparticle pore size can be controlled by the scCO₂ pressure release rate, future opportunities exist for top-down control over monolith microstructure and gas-adsorption properties.

To assess the extent to which COF monoliths can be used as industrial pellets, we carried out nanoindentation studies, from which we derived mechanical indentation moduli and hardness values. For a 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) pellet, we obtained an indentation modulus of 3.71 ± 0.20 GPa and a hardness of 0.18 ± 0.02 GPa (Figures 3B and 3E). These values are significantly higher than those previously obtained for COF aerogel pellets and slightly above those known for high-molecular-weight polyethylene. A full comparison of the mechanical properties of the COF monolith and those of other COF bodies reported in the literature is included in Table S6. These results suggest mechanical robustness and potential industrial suitability, possibly as a result of weak, non-crystalline-layer interfaces between COF crystallites, which serve to dissipate stress. By contrast, powder controls crumbled readily and could not be mounted in the instrument to yield reliable results. Further densification of monoliths with the use of higher-surface-tension activation solvents yielded slight increases in both indentation modulus and hardness. For a methanol-activated 1.000 (v/v) monolith, where pore collapse is complete, the indentation modulus and hardness values were 4.21 ± 0.37 and 0.34 ± 0.03 GPa, respectively (Figures 4D–4H). However, because these values represent modest improvements over the fully porous analog, potential design trade-offs between mechanical properties and porosity are likely to favor porous monoliths. Collectively, the combination of mechanical robustness, high bulk densities, high surface areas, and regular hierarchical mesopores is unique among COF-shaped
bodies demonstrated to date. This is illustrated in Figure 3F and Table S6, in which the 3 bar h⁻¹ scCO₂-activated 1.000 (v/v) monolith is compared with other COF bodies (including those featuring additives and binders) on the basis of density, BET area, and mechanical figures of merit.

**Molecular simulations and lattice-gas model**

To accurately capture the adsorption characteristics of TPB-DMTP-COF *in silico*, we carried out grand canonical Monte Carlo (GCMC) simulations on TPB-DMTP-COF crystalline fragments exhibiting varying degrees of interlayer slip (supplemental information section S2). Starting from perfect AA stacking (0% slip), one of two sequential layers of the COF was gradually shifted until perfect AB stacking was achieved (100% slip). Using cells derived from 0%, 25%, 40%, 50%, 75%, and 100% slipped starting structures (Figure S1), we then used GCMC simulations to...
generate predicted nitrogen isotherms at 77 K. Upon comparison of the respective low-pressure regions and mesoporous steps of the experimental adsorption isotherms to those derived from theory, a 40% slipped structure was found to provide the best agreement with experiment, giving almost identical low-pressure adsorption characteristics up to the mesoporous step (Figure 5D). Above the mesoporous step, however, whereas experimental isotherms for TPB-DMTP- COF powders maintained a reasonable agreement with those calculated from the 40% slipped structure until saturation, substantial deviations from theory were observed for experimental isotherms derived from TPB-DMTP-COF monoliths as a result of interparticle mesoporosity. Given that these deviations ultimately push total nitrogen uptake within the monolith above levels expected for purely crystalline systems, the ability to accurately capture such deviations computationally is critical in evaluating and subsequently tuning final gas-uptake characteristics for a desired target application.

To model contributions to total gas uptake arising from interparticle mesopores, we moved to a lattice-gas model of the TPB-DMTP-COF monolith (supplemental information section S4). Lattice-gas models have been extensively used in the past for studying the nature of sorption hysteresis for fluids in confined interconnected void spaces of porous glasses.35–37 We numerically reconstructed the structural model of the monolith (Figure 5B) used in the lattice-gas model from the SAXS data for the TPB-DMTP-COF monolith by means of generating a two-point correlation function $S_2(r)$ and using it in the reconstruction algorithm. A 3D reconstructed structure and its 2D slice used in the lattice-gas model for the TPB-DMTP-COF monolith activated by scCO$_2$ are shown in Figures 5A and 5C, respectively. To model the trajectory of the system in the grand canonical ensemble, we subsequently employed kinetic Monte Carlo (kMC) simulations from which nitrogen adsorption isotherms at 77 K could be obtained. The numerically generated isotherms show an excellent agreement with experimental data for the TPB-DMTP- COF monolith within the high-pressure region of the adsorption isotherms, providing complementary data to the GCMC-calculated isotherms and demonstrating the applicability of lattice-gas models in capturing the interparticle mesoporosity of COF monoliths. Collectively, these results suggest that the hierarchical porosity of COF monoliths can be accurately described computationally across the micro- and mesoporous ranges, enabling robust future predictions of adsorption characteristics.

Gas-adsorption characteristics of TPB-DMTP-COF monoliths

To demonstrate the utility of monolithic processing of COFs in gas-storage applications, we performed pure-component adsorption studies on TPB-DMTP-COF powders and monoliths. Low-pressure isotherms collected at 298 K revealed good CO$_2$ (Figures 6A and 6D) uptake for both powders and monoliths with modest to low CH$_4$ (Figure 6A) and N$_2$ (Figure 6D) uptake, respectively, for both systems. However, up to pressures of 1 bar, although higher CO$_2$ uptake was obtained for monoliths than for powders, lower uptake for both CH$_4$ and N$_2$ was obtained for monoliths than for powders. These results suggest that the presence of interparticle mesopores in monoliths can be used not only for improving final storage capacities for a single component but also for favorably or disfavorably influencing final uptake characteristics of various components within a mixed feed.
information section S8). At low pressures, the selectivity for CO₂ relative to other components was substantially improved, providing evidence that monolithic COF structuring can be used to provide separation enhancements relative to unstructured COF powders. To confirm this, we performed dynamic breakthrough studies on TPB-DMTP-COF monoliths and powders by using mixed-gas feeds. For the 15% CO₂/85% N₂ mixture (Figure 6F), although comparable separations were achieved for the monolith and powder (and some additional evidence for axial dispersion was observed), the total CO₂ uptake was found to be 13.4% higher for the monolith. For the 50% CO₂/50% CH₄ mixture (Figure 6C), a markedly sharper separation for the monolith than for the powder was observed, and an additional improvement in CO₂ capacity of 8.6% was achieved. Tables S6 and S7 compare these results against those for similar separations reported in the literature. Collectively, these results not only demonstrate the utility of monolithic processing for adsorbent-based chemical storage and separation but also afford additional
degrees of freedom through which the properties of COFs can be systematically designed and tuned.

**Conclusions**

Using a simple and general processing workflow, we introduce methods for the preparation of hierarchically porous COF monoliths without the need for additional materials or processing components. We show that such processing methods are compatible with mechanically weak materials and further afford degrees of design freedom in the control of both extrinsic and intrinsic porosities. These characteristics endow monolithic COFs with properties that are distinct from both powder and single-crystal analogs, which we accurately capture *in silico* by using a lattice-gas model. We envision that such computational approaches can be used in the future to predict gas-uptake properties for broad classes of monolithic mesoporous materials. The extrinsic porosity present in COF monoliths can further be leveraged for simultaneously increasing and decreasing the final uptake capacities for various gas constituents relative to powder benchmarks, which we make use of in demonstrating improved separation performance for industrially relevant gas compositions. We believe that this study not only opens up new possibilities for the practical applicability of COFs but also provides a pathway forward for tuning sorbent-analyte interactions where changes to the underlying framework chemistry might not be possible or synthetically accessible.

**EXPERIMENTAL PROCEDURES**

**Resource availability**

**Lead contact**

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, David Fairen-Jimenez (df334@cam.ac.uk).

**Materials availability**

The experimental dataset and materials generated and/or analyzed during the current study are available from the lead contact upon reasonable request.

**Data and code availability**

Adsorption information files (AIFs), Materials Studio scripts, mechanical-property data, RASPA input files, SAXS data, and crystallographic information files (CIFs) are included in the supplemental information. Any raw data and code used for analysis not included in the supplemental information are available from the lead contact upon reasonable request.

**Materials**

Scandium(III) trifluoromethanesulfonate (98%) was purchased from Alfa Aesar; 1,3,5-tris(4-aminophenyl)benzene (93%) was purchased from TCI; 2,5-dimethoxybenzene-1,4-dicarboxaldehyde (97%) was purchased from Sigma-Aldrich; and methanol (99.9%), acetonitrile (99.9+%), 1,3,5-trimethylbenzene (99%), and 1,4-dioxane (99.5%) were purchased from Acros Organics. All chemicals were used as received without further purification.

**Synthesis of TPB-DMTP-COFs**

**Monoliths**

To a 50 mL centrifuge tube were added 1,3,5-tris(4-aminophenyl)benzene (140.60 mg, 400 μmol) and 2,5-dimethoxybenzene-1,4-dicarboxaldehyde (29.13 mg, 150 μmol). Solvent (16 mL) was then added, and the mixture was sonicated briefly to a homogeneous suspension. Scandium(III) trifluoromethanesulfonate (12 mg, 24 μmol) was
added, the tube was sealed, and the mixture was sonicated again for approximately 20 s. The mixture was then left to react for 30 min undisturbed. The sample was collected by centrifugation for 50 min, washed with three portions of solvent (40 mL each) and an additional portion of methanol (40 mL), and was solvent exchanged in methanol (40 mL) at 50°C for 48 h; the solvent was then replaced after 24 h. The solvent was then decanted, and the sample was washed with methanol (40 mL) and left to dry at 20°C for a further 24 h or dried with scCO2. The sample was activated overnight at 120°C under vacuum prior to characterization.

**Powders**
The TPB-DMTP-COF powder controls were synthesized according to a previously reported procedure.14

**Characterization of TPB-DMTP-COF**

**Total scattering data**
Total scattering data were collected at beamline 11-ID-B of the Advanced Photon Source of the Argonne National Laboratory.39 Monoliths were segmented into regions (rim, top, and bottom; Figure S6) and lightly ground before being loaded into 1.1 mm (outer diameter) Kapton capillaries. High-energy X-ray scattering data were recorded with a Perkin Elmer amorphous silicon-based area detector with an X-ray wavelength of 0.2115 Å at a sample-to-detector distance of ca. 180 mm—experimental geometry was calibrated with a CeO2 diffraction standard. The images were calibrated and reduced to 1D diffraction data within GSAS-II.40 The X-ray scattering measured for an empty Kapton capillary was used as the sample background. The data were background corrected in xPDFsuite,41 and \( g(r) \) was calculated with data in the range 0.1 Å\(^{-1}\) \( \leq Q \leq 23.1 \) Å\(^{-1}\). Full details and discussion on the total scattering data are included in supplemental information section S6. SEM images were acquired by an FEI XL30 FEGSEM with an accelerating voltage of 5 kV. Samples were sputter coated with gold. TEM was carried out on a FEI Tecnai F20 TEM operated at 200 kV, and images were acquired with a Gatan OneView camera.

**Helium pycnometry**
Helium pycnometry was obtained with an AccuPyc 1330 Pycnometer from Micromeritics. This technique was used to estimate the particle density and the volume of both powders and monoliths by measuring the pressure change of helium in a calibrated volume. Each volume was recorded as an average value of six consecutive runs. Prior to the analysis, all samples were activated overnight at 120°C (vacuum) before the mass was measured.

**Mercury porosimetry**
Mercury porosimetry was obtained up to a final pressure of 2,000 bar with an AutoPore IV 9500 instrument from Micromeritics. This technique was used to estimate the particle density of both powders and monoliths at atmospheric pressure. Prior to the analysis, all samples were activated overnight at 120°C (vacuum) before the mass was measured and then degassed in situ thoroughly before the mercury porosimetry.

**Critical point dryer-CO\(_2\) procedure**
A SPI-DRY critical point dryer (jumbo size) modified with a manometer at the chamber was used to dry and activate the COF monoliths. First, the sample was transferred into a dialysis membrane (Spectra/P.1 MWCO 6–8 kD) and sealed. Then, the membrane was introduced into the critical point drying equipment. Then, it was immersed in
subcritical (l) CO₂ at 283 K and 50 bar for half an hour. Then, the exchanged methanol was removed through a purge valve and then flushed with fresh (l) CO₂. This process was repeated three times. Subsequently, the temperature was raised 5 K min⁻¹ up to 313 K to exceed the scCO₂ point. Finally, under constant temperature (313 K), the chamber was vented at 8 or 3 bar h⁻¹ to atmospheric pressure.

**Gas-adsorption measurements**

Ultra-high-purity-grade CH₄, N₂, and CO₂ were used for gas-sorption experiments. Adsorption experiments (up to 1 bar) for different pure gases were performed on a Micromeritics 3 Flex surface-area and pore-size analyzer. About 200 mg of activated sample was used for the measurements. A temperature-controlled bath was used to maintain a constant temperature in the bath throughout the duration of the experiment. Samples were degassed on a Micromeritics PrepStation instrument prior to the analysis.

**Dynamic mixed-gas breakthrough studies**

In a typical experiment, ca. 0.3 g of pre-activated sample was placed in a quartz tube (Ø = 8 mm) to form a fixed bed held in place by quartz wool. For monolithic samples, we broke and sieved individual monoliths to reduce the particle diameter to ca. 2 mm to ensure good packing within the sample tube. We heated each sample to 353 K under a dry helium flow to remove atmospheric contaminants. Upon cooling, the chosen gas mixture was passed over the packed bed with a total flow rate of 2 cm³ min⁻¹ at 298 K. The outlet gas concentration was continuously monitored with an Agilent 5975 MSD mass spectrometer. Upon complete breakthrough and saturation of the packed bed adsorbent, the gas mixture was switched off, and dry helium was flowed over the solid. Heating was switched on, and samples were heated to 353 K to aid regeneration.

To calculate the CO₂ uptake, we initially passed the gas mixture through an empty reactor containing quartz wool at a flow rate of 2 cm³ min⁻¹ as a blank reference. The gas flow was constantly monitored by the mass spectrometer. We integrated the CO₂ curve to calculate the area of the curve (A_ref). Upon completion of a CO₂ breakthrough experiment with an adsorbent, we also integrated the area of the CO₂ adsorption curve (A_exp). To calculate the total amount of CO₂ adsorbed, we used the following equation:

\[
\text{total CO}_2 \text{ uptake} = (A_{\text{Ref}} - A_{\text{Exp}}) \times \text{CO}_2 \text{ flow (cm}^3 \text{ min}^{-1})
\]

**Molecular simulations**

The adsorption isotherms of N₂ were simulated by the GCMC method as implemented in the RASPA simulation package. The geometric properties were calculated with Poreblazer. A more detailed description of the methodology and model parameters is given in supplemental information section S2.

**Lattice-gas model**

First, the collected SAXS data were converted into a two-point correlation function S₂(r) (defined as the probability of two points separated by distance r, belonging to the pore space of the medium), which was then used as a benchmark for creating a 3D reconstruction of the TPB-DMTP-COF monolith. In order to model the trajectory of the system in the grand canonical ensemble and to obtain the adsorption isotherms, we employed kMC simulations. A more detailed description of the reconstruction procedure and the kMC simulations is given in supplemental information section S4.
**Calculation of mechanical properties**

Mechanical properties for the COFs present in the CURATED COF database were calculated with classical molecular mechanics via the “constant strain approach” as implemented in the Forcite module of Materials Studio. Some CURATED structures for which either the mechanical or the geometric property calculation failed were excluded. The mechanical properties calculated included the shear modulus, bulk modulus, and Young’s modulus. A more detailed description of the methodology used is given in supplemental information section S5.

**Calculation of BET area**

BET areas were calculated with a computational tool called BETSI, which makes an unambiguous calculation of the possible BET area. More details about BETSI can be found in supplemental information section S9.

**Nanoindentation tests**

Nanoindentation tests were carried out for measuring the mechanical properties, namely indentation modulus\(^45\) (E) and hardness (H) of the 3 bar h\(^{-1}\) scCO\(_2\)-activated 1.000 (v/v) TPB-DMTP-COF monolith, the methanol-activated 1.000 (v/v) TPB-DMTP-COF monolith, and the TPB-DMTP-COF powder. In order to perform the tests, we cold mounted the sample in epoxy and polished it with sandpapers and diamond suspension (up to a 0.1 \(\mu\)m grain size). This was not possible for the TPB-DMTP-COF powder because it crumbled into small pieces. All tests were carried out with a KLA iMicro nanoindenter equipped with a 50 mN force actuator. A Berkovich diamond indenter tip was used. Continuous stiffness measurements (CSMs) were performed, allowing measurement of E and H as a function of the indentation depth. The maximum indentation depth was set to 2,000 nm for all tests. Average values of E and H were computed in the range of 500–2,000 nm. For the methanol-activated 1.000 (v/v) TPB-DMTP-COF monolith, the NanoBlitz 3D mode was used to generate indentation maps of nanomechanical properties. Three maps in three different areas of the sample were taken.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at [https://doi.org/10.1016/j.chempr.2022.07.013](https://doi.org/10.1016/j.chempr.2022.07.013).

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**AUTHOR CONTRIBUTIONS**

M.E.C. and D.F.-J. designed the research. M.E.C. synthesized and characterized the materials. M.E.C., D.G.M., and C.C. performed the N₂ gas adsorption at 77 K. D.G.M. and C.C. performed the N₂, CO₂, and CH₄ gas adsorption at 298 K. D.G.M. conducted the dynamic mixed-gas breakthrough analysis with supervision from T.C. N.P.M.C. carried out the SAXS and WAXS data collection. D.O.N. performed the SAXS and WAXS data analysis. D.O.N and K.W.C. collected and analyzed the PDF-XRD data. G.D. carried out the TEM analysis. M.E.C. carried out the SEM analysis. N.R. and R.C. carried out the lattice-gas modeling under the supervision of S.T. N.R. carried out the molecular simulations and mechanical property screens. J.A.M.-I. performed the scCO₂ drying and activation under the supervision of F.Z. J.S.-A. carried out the mercury porosimetry. M.T. performed the nanoindentation experiments and data analysis under the supervision of J.-C.T. M.E.C., N.R., and D.F.-J. wrote the manuscript with contributions from all authors.

**DECLARATION OF INTERESTS**

M.E.C. and D.F.-J. are inventors on international patent application no. WO2021052969A1, which covers COF monoliths, as well as aspects of their use. M.E.C. and D.F.-J. have a financial interest in the start-up company Immaterial Labs, which is seeking to commercialize MOFs.

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