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Sub-stoichiometric 2D covalent organic frameworks from tri- and tetratopic linkers

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Covalent organic frameworks (COFs) are typically designed by breaking down the desired network into feasible building blocks - either simple and highly symmetric, or more convoluted and thus less symmetric. The linkers are chosen complementary to each other such that an extended, fully condensed network structure can form. We show not only an exception, but a design principle that allows breaking free of such design rules. We show that tri- and tetratopic linkers can be combined to form imine-linked [4+3] sub-stoichiometric 2D COFs featuring an unexpected bex net topology, and with periodic uncondensed amine functionalities which enhance CO_2 adsorption, can be derivatized in a subsequent reaction, and can also act as organocatalysts. We further extend this class of nets by including a ditopic linker to form [4+3+2] COFs. The results open up possibilities towards a new class of sub-valent COFs with unique structural, topological and compositional complexities for diverse applications.

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ovalent organic frameworks (COFs) represent an intriguing class of crystalline porous materials, where appropriately functionalized organic molecules are linked with an atomic level precision to give extended periodic structures^{1–6}. Such materials are synthesized by topologically directed condensation reactions, based on matching the geometry of the building blocks to form the desired net composed of the respective vertex and edge linker units^{3,7-9}. The shapes and dimensions of the vertex and the edge linkers determine the pore shape and size, respectively. Most of the 2D COF structures reported till date are based on four simple topologies for which the vertices are connected by only one kind of edge-the honeycomb, square, kagome, and hexagonal lattice complexes³. In order to make the COFs, these nets are typically deconstructed into highly symmetric building blocks (e.g., Type I COFs, Fig. 1). On the other hand, increased compositional and structural complexity is paramount in order to broaden the scope and applicability of COFs (or any other material). Reports in this regard are limited to either using complex molecules or macromolecules as vertices 10-20, or using combinations of linear linkers of different lengths with appropriate vertex molecules^{21,22}, to arrive at complicated network structures (e.g., Type II COFs, Fig. 1). However, most such efforts produce deformations of the aforementioned simple topologies only. More importantly, most of these approaches involve cumbersome and time consuming multistep organic synthesis before the COF formation reaction which is yet to be understood well and controlled. In all the possibilities discussed thus far, the vertices and the linkers are always complementarily chosen such that formation of a fully condensed network structure is possible. In other words, only fully condensed structures are considered as targets for synthesis of COFs. Synthesis is not attempted if due to geometric constraints or otherwise this rule is deemed not possible to be followed, thus limiting creativity.

Contrary to these typical design principles, in this manuscript, we show that triangular tritopic and rectangular tetratopic linkers can actually be combined to form a crystalline, dual pore 2D COF belonging to a previously unidentified *bex* topology with two distinct vertex molecules (Fig. 1, Type-III COFs). Uniquely, in such COFs the tetratopic linker simultaneously acts as a bi- and tetradentate linker in two different coordination environments and thus inherently results in uncondensed functional groups in the network. This has significant implications with regard to the potential of COFs for applications in diverse

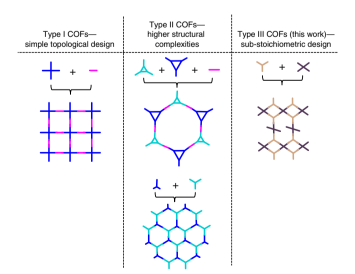


Fig. 1 Hierarchy of the evolution of structural complexity in COFs

fields, including sensing^{23,24}, storage^{25,26}, (photo)catalysis^{27–29}, ion conduction^{30,31}, and optoelectronics^{32,33}. Such diverse applicability demands diversified structural characteristics and importantly, appropriate functionality, which is often difficult to achieve in a single-step and requires post-synthetic modification of the framework. With examples limited to only a few^{5,26,34,35}, it is still challenging to tune the structure and the physical properties of the COF post-synthetically while maintaining its crystallinity and porosity. This is because generating functional groups in the framework by post-synthetic modification requires orthogonal reaction chemistry, which also limits probable options considerably. In this regard, the unconventional combination of building blocks to produce periodic, uncondensed amine functional groups in the fully extended, crystalline and porous imine-linked COFs, is a new approach toward parallel and simultaneous functionalization of 2D COFs. Also, this method of introducing free functional groups by generating defined subvalent COFs is fundamentally different from previous examples that rely on the introduction of defects^{36,37}, or the introduction of periodic linker vacancies into a saturated host lattice to introduce dangling functional groups in the framework as reported by Loh and co-workers³⁸. We further show that the free amine functional groups enhance CO2 adsorption of the framework, can be transformed in a subsequent reaction, and can also act as catalysts in organic transformations.

All three-component COFs reported thus far use building blocks of only two different symmetries ^{13,21,22,39}. In this report, we also demonstrate the combination of tritopic, tetratopic, and linear ditopic linkers to form COFs, which have crystallinity and surface area comparable to the state-of-the art examples.

This class of sub-stoichiometric COFs with a frustrated bonding network prompts us to think outside the set rules of topological deconstruction and paves way for a new class of structures with unique structural, topological, and compositional complexities. Such COFs are inherently linked to free functional groups, and can be exploited in a diverse, application-specific manner.

Results

Synthesis of PT- and PY-COFs. Triazine tribenzaldehyde (T) and benzene tripicolinaldehyde (Y) were chosen as the triangular tritopic linkers for reaction with pyrene tetraaniline (P) as the tetratopic linker. A fully condensed 2D framework cannot be envisaged from a combination of molecules with such symmetries, and thus periodic, crystalline frameworks are typically not expected to form. Nevertheless, when these linkers are reacted in an equimolar ratio in 1:1 mesitylene/dioxane under solvothermal conditions, crystalline PT-, and PY-COFs result (Fig. 2 and Supplementary Methods). Other typical solvent combinations and different ratios of mesitylene and dioxane, as well as different molar ratios of the reactants, either had no noticeable effect or decreased crystallinity while still retaining the powder x-ray diffraction (PXRD) pattern (Supplementary Methods).

IR and NMR characterization of PT- and PY-COFs. Fourier transform infrared spectroscopy (FT-IR) measurements (Fig. 3a) of PT- and PY-COFs show almost complete disappearance of $v_{\rm C=O}$ (stretch) at ~1700 cm $^{-1}$ corresponding to the aldehyde groups in the starting materials, T and Y (Supplementary Fig. 1). The disappearance of the carbonyl stretching signal is accompanied by the appearance of $v_{\rm C=N(stretch)}$ at $1625\,{\rm cm}^{-1}$ corresponding to the imine bonds formed in the framework by condensation of the aldehyde and the amine groups in the starting materials. Interestingly, similar to that in P (Supplementary Fig. 1), $v_{\rm N-H(stretch)}$ in the region $3430\,{\rm cm}^{-1}$ – $3190\,{\rm cm}^{-1}$ is observed for both PT- and

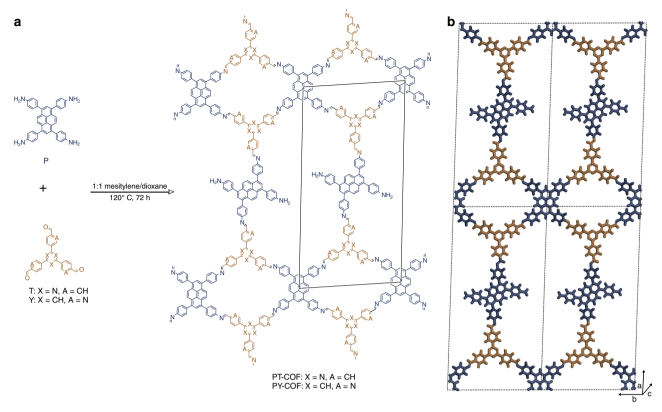


Fig. 2 Structure of PT- and PY-COFs. a Synthesis and molecular structure of PT- and PY-COFs. The in-plane unit cell is outlined in black. b Illustration of the COF structure model with bex toplogy (viewed perpendicular to the ab plane)

PY-COFs. ¹³C cross-polarization magic angle spinning solid-state NMR (ssNMR) spectroscopy (Fig. 3b) shows the disappearance of the aldehyde carbonyl 13 C resonance at ~190 ppm as present in the precursor aldehydes, T, and $Y^{27,40,41}$. This is accompanied by the appearance of the characteristic imine ¹³C signal^{41,42} at 157 ppm (1) for PT-COF and at 159 ppm (1') for PY-COF proving the condensation of the starting materials into the framework. Retention of the molecular structure of the building blocks in the framework is also evident from an analysis of the 13C NMR spectrum. For PT-COF, the signals for carbon 2, 3, and 4 can be identified at 170 ppm, 150 ppm, and 116 ppm, respectively⁴². Likewise, for PY-COF, carbon 3', 4', 5', and 6' appear at 150 ppm, 116 ppm, 155 ppm, and 146 ppm, respectively. This analysis is further corroborated with ¹⁵N-ssNMR measurements (Fig. 3b and Supplementary Figs 2, 3) and corresponding quantum-chemical calculations (Supplementary Fig. 4-9 and Supplementary Tables 1, 2). For PT-COF, the imine nitrogen 7 appears at -46 ppm and the triazine nitrogen 8 appears at -127 ppm^{42} . For PY-COF, the imine (7') and the pyridine (10') nitrogen appear between -40 ppm and -78 ppm. The signal at -262 ppm in the ¹⁵N ssNMR spectrum of PY-COF probably arises from oxidation of a fraction of the imine-linkages to amides in the COF (Supplementary Figs 9, 10, Supplementary Table 1)43-47. Interestingly, both PT- and PY-COFs show the presence of free amine functional groups (9 and 9') at -323 ppm to -330 ppm (Fig. 3b, inset).

PXRD analysis and structural modeling of PT- and PY-COFs. PY- and PT-COFs are crystalline materials (see Supplementary Fig. 11 for chemical stability tests). The experimental PXRD patterns (Fig. 3c) show eight reflections at 3.7, 4.0, 4.9, 6.1, 7.3, 8.7, 10.9, and 15.2° 2θ . In addition, a broad stacking reflection can be seen at $2\theta = 24^{\circ}$ for both COFs hinting towards the

formation of 2D rather than a 3D-COF (vide infra). The PXRD patterns could be simulated by a structure model (Supplementary Methods) as shown in Fig. 2a and b. In this model, the tetratopic P plays a dual role, a behavior that is currently unknown in COF chemistry. The triangular linkers connect two P molecules at two points on both sides; P thus acts as a node and is tetra-coordinated to produce a ribbon-like architecture. The free -CHO termini of the triangular linkers constituting the ribbons are then interconnected by a single P molecule by its diagonal amine groups, which now acts as a linear ditopic linker. As a consequence, two uncondensed -NH2 groups result per unit cell, the presence of which were confirmed with 15N-ssNMR and IR experiments. The unit cell parameters for this structure model with P1 space group and bex crystallographic network topology were obtained by force-field optimizations and were then Pawley refined against the experimental PXRD pattern (Supplementary Table 3). The refined unit cell parameters are observed to be similar to the parameters obtained by structure simulations. The structure model for AA-type stacking yields the best agreement with the experimental pattern (Supplementary Fig. 12). Finally, using this model and the parameters obtained from the Pawley fit, the experimental powder pattern was Rietveld refined, yielding the unit cell parameters of $a = 52.516 \,\text{Å}$, $b = 23.556 \,\text{Å}$, c = 3.681 Å, and $\alpha = \beta = 90^{\circ}$, $\gamma = 90.759^{\circ}$, $R_{wp} = 3.22$ for PT-COF (Supplementary Data 1) and a = 52.718 Å, b = 24.117 Å, c = 3.815 Å, and $\alpha = \beta = 90^{\circ}$, $\gamma = 88.393^{\circ}$, $R_{wp} = 2.0$ for PY-COF (Supplementary Data 2). It is important to note that the bex net in the present case is a more complex net and has a transitivity of 222 with two kinds of vertices from two different molecules as vertices in the network structure and has not been previously identified for COFs⁴⁸. Almost all of the hundreds of 2D COF structures with considerable complexities produced till date belong to only four highly symmetric network topologies with

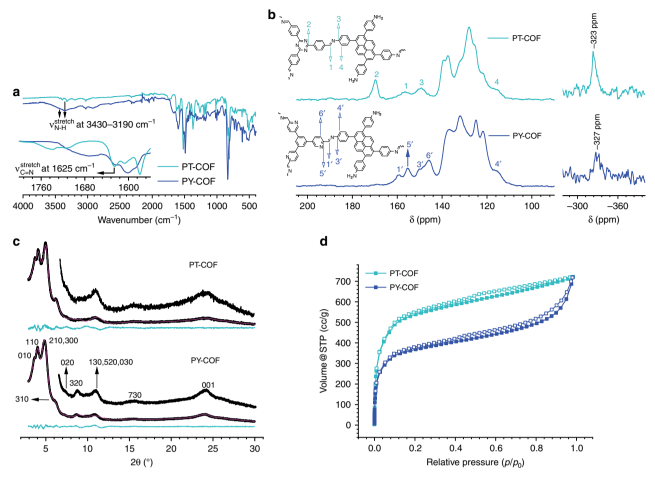


Fig. 3 Characterization of PT- and PY-COFs. **a** FT-IR characterization of PT- and PY-COFs. Free amine functionalities can be identified by the characteristic $\nu_{\text{N-H(stretch)}}$ in the region 3430 cm⁻¹-3190 cm⁻¹. The inset shows the presence of $\nu_{\text{C=N(stretch)}}$ **b** ¹³C ssNMR spectra with corresponding assignments. The unassigned signals correspond to the pyrene moiety and the phenyl rings. ¹⁵N ssNMR depicting the free amine functionalities in PT- and PY-COFs is shown on the right as an inset. **c** PXRD patterns of PT- and PY-COFs with corresponding Rietveld refinements (magenta) showing good fits to the experimental data (black) with minimal differences. The turquoise traces show the difference between the experimental data and the refinement. The corresponding Miller indices of the identifiable reflections are shown as well. **d** Argon sorption isotherms of PT- and PY-COFs at 87 K. Filled and open symbols represent the adsorption and the desorption branches, respectively

minimal transitivities³. A 3D-COF based on the tbo net, as known from the metal organic framework HKUST-1⁴⁹, can also be envisaged by connecting the P and T/Y linkers and was thus modeled (Supplementary Fig. 13, 14). However, the experimental PXRD patterns, with the stacking reflection at $2\theta = 24^{\circ}$, did not match with that expected for this model. It should be noted that the formation of a crystalline 2D COF from P and T/Y linkers is classically inconceivable because these linkers cannot form a fully condensed 2D framework. It is also worth noting the reported formation of 3D-COFs from tetraphenylethylenebased tetratopic and triphenylbenzene/triphenylamine-based tritopic linkers⁵⁰. This points to a probable design principle for the formation of such 2D sub-stoichiometric COFs: the increased planarity of linkers (as with our P and T/Y linkers) possibly increases the predilection towards stacking and thus the consequent formation of such structures over stoichiometric 3D-COFs.

Synthesis of PT₂B- and PY₂B-COFs. In order to further validate the structure model, we decided to decouple the two linking modes of P and check if a COF could be made by replacing one equivalent of this linker with the linear ditopic benzidine (B) linker. In the resulting structure, the ribbons should now be

linked by B molecules instead of P, and thus the COF should not have any free-amine functional groups. Thus, P, T/Y, and B were reacted in a 1:2:1 molar ratio in the same solvent mixture as before under the same conditions to get highly crystalline PT₂B-and PY₂B-COFs (Fig. 4 and Supplementary Methods).

IR and NMR characterization of PT₂B- and PY₂B-COFs. FT-IR measurements (Fig. 5a) again show a near complete loss of $v_{C=0}$ (stretch) signal corresponding to the aldehyde groups in the starting materials and a simultaneous appearance of $v_{C=N(stretch)}$ at 1625 cm⁻¹, validating the formation of the framework. Indeed, as expected, the v_{N-H(stretch)} signal, as observed for PT- and PY-COFs, is absent in PT₂B- and PY₂B-COFs. ¹³C ssNMR spectrum (Fig. 5b) of PT₂B- and PY₂B-COF is very similar to PT- and PY-COF, respectively, and again shows the disappearance of the aldehyde carbonyl ¹³C resonance at ~190 ppm and appearance of an imine ¹³C signal at 157 ppm (11) for PT₂B-COF and at 159 ppm (11') for PY₂B-COF proving the formation of the framework. Retention of the molecular structure of the building blocks in the framework is again evident. 15N ssNMR is identical to PT- and PY-COFs (Supplementary Fig. 15), except for an almost complete absence of nitrogen signals corresponding to free amine functional groups (Fig. 5b, inset).

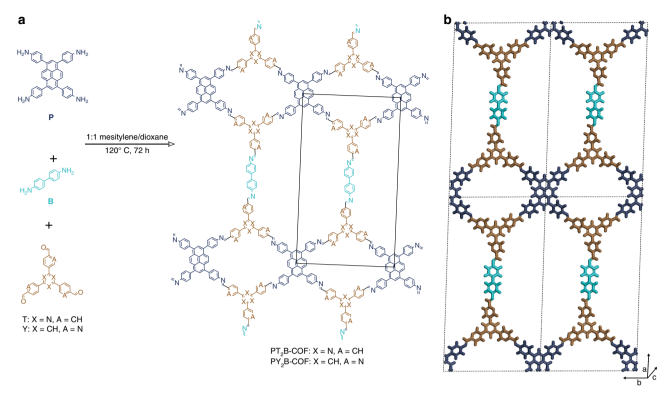


Fig. 4 Structure of PT₂B- and PY₂B-COFs. **a** Synthesis and molecular structure of PT₂B- and PY₂B-COFs. The in-plane unit cell is outlined in black. **b** Illustration of the COF structure model (viewed perpendicular to the *ab* plane)

PXRD analysis and structural modeling of PT₂B- and PY₂B-COFs. PT₂B- and PY₂B-COFs have higher crystallinity as compared with PT- and PY-COFs. The PXRD patterns (Fig. 5c) have 12 and 14 prominent reflections, respectively, for PT₂B- and PY₂B-COF at 1.8, 3.6, 4.1, 5.3, 5.9 (for PY₂B-COF), 6.9, 7.3, 9.4, 10.6, 12.3, 14.7 (for PY₂B-COF), 15.9, 24.2, 25.8–26.5° 2θ . The Pawley refined unit cell parameters (Supplementary Table 3) are again similar to the parameters simulated by modeling the structure (Supplementary Fig. 16). The unit cell parameters as obtained by Rietveld refinement of the experimental powder pattern were a = 46.264 Å, b = 23.872 Å, c = 3.802 Å, and $\alpha = \beta =$ 90°, $\gamma = 90.247$ °, $R_{\rm wp} = 2.64$ for ${\rm PT_2B\text{-}COF}$ (Supplementary Data 3) and a = 45.845 Å, b = 24.352 Å, c = 3.743 Å, and $\alpha = \beta =$ 90°, $\gamma = 90.981$ °, $R_{wp} = 2.59$ for PY_2B -COF (Supplementary Data 4). The replacement of the longer P in PT- and PY-COFs with a shorter B in PT₂B- and PY₂B-COFs to interlink the ribbons is immediately apparent from the crystal parameters with the corresponding unit cell length a decreasing from 52.5-52.7 Å in the former to 46.2-45.8 Å in the latter. In addition, the unit cell length b, corresponding to the separation of the pyrene moieties in the ribbon in both the COF systems stays the same at ~23.5-24.4 Å. This further proves the correctness of our structural model. The lower crystallinity of PT- and PY-COFs compared with PT₂B- and PY₂B-COFs could be argued to be owing to the presence of free amine groups in the P molecule that acts as the linker interconnecting the ribbons. Reaction only along the diagonal amine groups of such linkers will give rise to the described structure; condensation and reaction propagation along other directions will lead to defects and disorder reducing the crystallinity and crystallite size as observed with TEM measurements (vide infra). Making COFs by combining three different linkers has been reported, albeit using intuitive combinations of either two differently sized linear linkers of the same geometry 13,21,22, or using two molecules of similar geometry but having different linking

groups³⁹. PT₂B- and PY₂B-COFs, on the other hand, are the first example of COFs made from a combination of three linkers with different topicity.

TEM analysis. TEM data (Fig. 6 and Supplementary Fig. 17) provide strong support for the structural model (see Supplementary Fig. 18 for SEM measurements). Different d-spacings can be observed in the same sample-and often in the same crystallite-strongly suggesting a structural model based on a unit cell with significantly different a and b parameters describing the in-plane structure. The observed crystallites, in many cases bent, feature a large repeat unit of 3.9-4.1 nm for PT2B- and PY2B-COFs (Fig. 6d, f) corresponding to the edge distance a, which at 4.8-5.0 nm is longer for PT- and PY-COFs (Fig. 6h, i). The other prominently observed d-spacing is much smaller at ~1.9 nm (Fig. 6e) and is similar for all four COFs and corresponds to the unit cell length b. The shorter edge lengths in TEM measurements, as compared with those observed with PXRD measurements, is known to result from shrinkage of the structure owing to damage by the electron beam⁴². In addition, in the [100] direction, while PT2B- and PY2B-COFs show a monomodal intensity variation (Fig. 6d, f, g), PT- and PY-COFs exhibit a bimodal intensity variation (Fig. 6h-j) in each unit cell as a result of the higher electron density corresponding to the pyrene moieties between the ribbons in the latter systems, as also verified by simulated potential maps of the model structures. The COF crystallites also possess a large anisotropy and differ significantly in their extension in [100] and [010] directions (Supplementary Table 4). Elongated strip-like crystallites can be observed with predominant extension along [010] direction, i.e., the direction of b. This demonstrates that the ribbons can act as supramolecular motifs, i.e., secondary building units for hierarchical design of differently expanded frameworks. Along [100] or the direction

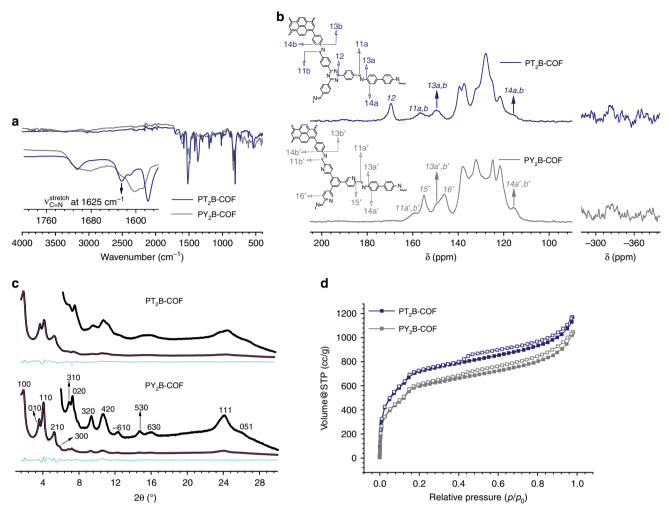


Fig. 5 Characterization of PT_2B - and PY_2B -COFs. **a** FT-IR characterization of PT_2B - and PY_2B -COFs. The inset shows the presence of $\nu_{C=N(stretch)}$, **b** ^{13}C ssNMR spectra with corresponding assignments. The unassigned signals correspond to the pyrene moiety and the phenyl rings. ^{15}N ssNMR depicting the absence of free amine functionalities in PT_2B - and PY_2B -COFs is shown on the right as an inset. **c** PXRD patterns of PT_2B - and PY_2B -COFs with corresponding Rietveld refinements (magenta) showing good fits to the experimental data (black) with minimal differences. The turquiose traces show the difference between the experimental data and the refinement. The corresponding Miller indices of the identifiable reflections are shown as well. **d** Argon sorption isotherms of PT_2B - and PY_2B -COFs at 87 K. Filled and open symbols represent the adsorption and the desorption branches, respectively

of *a*, the crystallites are much shorter. This is especially true for PT- and PY-COFs. It should be noted that such detailed TEM analysis of COFs is rare^{42,51,52} as useful measurements are typically impeded by the low TEM contrast and most importantly, because of an easy degradation of most COFs under the electron beam, for most except highly crystalline samples⁴².

Surface area and porosity measurements. The higher crystal-linity of PT₂B- and PY₂B-COFs is also reflected in their higher Brunauer–Emmett–Teller (BET) surface area as evaluated by measuring Argon sorption isotherms at 87 K (Figs. 3d, 5d). Although the BET surface areas of the PT- and PY-COFs are 1791 m² g⁻¹ and 1220 m² g⁻¹, respectively, those for the PT₂B- and PY₂B-COFs are 2367 m² g⁻¹ and 1984 m² g⁻¹, respectively. The BET surface areas of PT₂B- and PY₂B-COFs are comparable to some of the most porous 2D COFs, such as ILCOF-1 (2723 m² g⁻¹)⁵³, 4PE-1P-COF (2140 m² g⁻¹)⁵⁴ and TPB-DMTP-COF (2015 m² g⁻¹)⁵⁵. The absence of free amine functional groups in PT₂B- and PY₂B-COFs also reduces the possibility of chemisorbed molecules clogging the pores and possibly contributes to the larger surface area. A pore size distribution (PSD) was calculated from the Argon sorption isotherms using a Quenched

Solid DFT adsorption kernel for Argon at 87 K on carbon with cylindrical pores. For the PT- and PY-COFs, only one type of pore centered at 1.77–1.83 nm is observed (Supplementary Fig. 19a). This matches with the theoretical pore sizes predicted from the crystal structures (Supplementary Fig. 20), with the larger pore having pendant amine functionalities behaving as two smaller pores having sizes closer to the small pores in the ribbons. On the other hand, owing to the absence of the pendant amine functionalities and hence a larger second pore, two distinct pores can be seen in the PSD for PT₂B- and PY₂B-COFs at ~1.83 nm and 2.37 nm (Supplementary Fig. 19b), again in agreement with crystal structure predictions. The pore volumes for PT- and PY-COFs are 0.896 cm³ g⁻¹ and 0.837 cm³ g⁻¹, respectively. The pore volumes for PT₂B- and PY₂B-COFs are larger at 1.03 cm³ g⁻¹ and 1.259 cm³ g⁻¹, respectively.

Photoluminescence measurements. The COFs being distinctly emissive when dispersed in acetonitrile (Supplementary Fig. 21 and Supplementary Table 5), the presence of free amine functionalities can be corroborated with photoluminescence measurements. The detailed photophysical characterizations are listed in Supplementary Table 5. For PT- and PY-COFs in

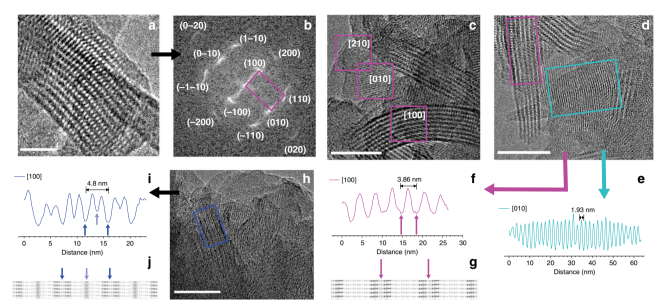


Fig. 6 TEM analysis. **a** A PY₂B-COF crystallite viewed along [001] showing the pores in the structure. **b** Fast Fourier Transform (FFT) filtered image of **a** showing the different planes in the PY₂B-COF crystallite in accordance with the PXRD data. **c** Individual crystallites of PY₂B-COF in different orientations. Multiple lattice spacings can be identified as shown. **d** Individual crystallites of PT₂B-COF in different orientations. **e** Line scan analysis in the [010] direction corresponding to the turquoise outlined region in **d**. A small *d*-spacing of 1.93 nm can be identified and is similar for all four COFs showing a similar constitution of the ribbon structure. A line scan analysis with monomodal intensity variation in the [100] direction (magenta outline) is shown in **f**. The magenta arrows correspond to the electron rich ribbons in PT₂B-COF as shown with magenta arrows in the projected potential map **g** of the COF. Dark contrast corresponds to higher projected potentials. **h** Individual crystallites of PT-COF in different orientations. Multiple lattice spacings can be identified. A line scan analysis in the [100] direction (blue outline) showing bimodal intensity variation is shown in **i**. The deep blue arrows correspond to the electron rich ribbons in PT-COF as shown with similar arrows in the projected potential map **j** of the COF, wherein dark contrast corresponds to higher projected potentials. The light blue arrow in the line scan analysis designates the bi-coordinating P linkers in PT-COF, which are more electron rich than the B linkers in PT₂B- and PY₂B-COFs. Scale bars: **a**: 20 nm; **c**, **d**, **h**: 50 nm

acetonitrile with ~2 equivalents of 2,6-lutidine (per -NH₂ group) as the base for trapping the HI generated, the emission intensity first drops and then increases beyond the initial value on addition of increasing amounts of methyl iodide (MeI) (Supplementary Fig. 22, 23). The drop in emission intensity can be related to the initial formation of -NHMe and -NMe2 species which further increase the photoinduced electron transfer quenching of pyrene-based fluorescence by the donor amine⁵⁶. As the concentration of MeI is increased further, -NMe3+ species form, which cannot participate in electron transfer quenching anymore and thus the emission intensity increases. On the other hand, no such obvious trend was observed for PT2B-COF, which does not have the free amine functional moieties in it. PY₂B-COF was too weakly emissive to be studied. The change in the photoluminescence emission intensity thus substantiates the presence of a COF structure with free amine groups.

 ${
m CO_2}$ adsorption studies. Free amine functional groups are known to enhance ${
m CO_2}$ adsorption capacity of porous frameworks and could thus again corroborate their presence $^{57-60}$. Thus, ${
m CO_2}$ sorption isotherms were recorded at 273 K and the absolute ${
m CO_2}$ uptake capacities of PT-, PY-, PT₂B-, and PY₂B-COFs at 1 atm are found to be 95, 146, 85, and 127 mg g⁻¹, respectively. To put this in perspective, the ${
m CO_2}$ uptake capacity of highly active carboxylic acid functionalized porphyrin based ${
m H_2P-COFs}$ reported by Jiang and co-workers is in the range 96–174 mg g⁻¹ ²⁶. ${
m CO_2}$ uptake capacity normalized to the BET surface area of the individual samples shows a clear trend (Supplementary Fig. 24): PT- and PY-COFs have 1.46 and 1.67 times higher ${
m CO_2}$ adsorption capacity than ${
m PT_2B-}$ and ${
m PY_2B-COFs}$,

respectively, an effect attributed predominantly to the presence of free amine groups.

Heterogeneous organocatalysis. The presence of these free amine functionalities in the COF framework could be further substantiated by employing them as heterogeneous organocatalysts. The aromatic amine-catalyzed cyclization-substitution cascade reaction of 2-hydroxylcinnamaldehyde with trimethylsilyl enol ether to form substituted chromenes was chosen because no radical intermediates are known to be involved and the reaction takes place at room temperature⁶¹. For PT₂B- and PY₂B-COFs, product formation is negligible (1-2% only). On the other hand, with PT- and PY-COFs, the reaction yields of 22% and 8%, respectively, with excellent regioselectivity (Supplementary Table 6, Supplementary Fig. 25-28), show that these COFs are porous heterogeneous amine-group terminated materials indeed. It is important to consider the proposed mechanism for the aforesaid catalytic reaction according to which a Schiff base adduct is initially formed in situ between the pendant amine groups and 2-hydroxylcinnamaldehyde (Supplementary Fig. 29)⁶¹. The fact that the reaction works with COFs as the amine catalysts thus notably shows that the pendant amine moieties in the pores can actually be modified suitably and reversibly in the COF template and can be used to link functional moieties specific to particular applications.

Functional group interconversion. As a proof-of-concept, we explored a functional group interconversion reaction, namely the conversion of amines to isothiocyanates. The COF as the amine substrate was reacted with CS_2 to generate the corresponding dithiocarbamate salt in situ followed by elimination to the

isothiocyanate product using cyanuric chloride as the desulfurylation reagent (Figs 7, 8, and Supplementary Methods)^{62,63}. Fig. 7a compares the FT-IR spectrum of PY-COF and PY-NCS-

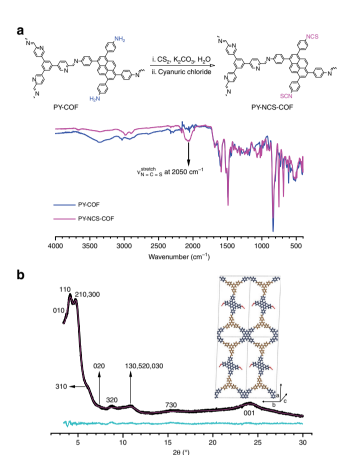


Fig. 7 Post-crystallization amine to isothiocyanate group interconversion reaction with PY-COF. **a** FT-IR spectrum of PY-NCS- and PY-COF showing the appearance of $\nu_{\rm N=C}$ $_{\rm S(stretch)}$ signal post modification. **b** PXRD pattern of PY-NCS-COF with corresponding Rietveld refinement (magenta) showing good fits to the experimental data (black) with minimal differences. The turquoise trace shows the difference between the experimental data and the refinement. The corresponding Miller indices of the identifiable reflections are shown as well. The inset shows an illustration of the COF structure model (viewed along the stacking direction)

COF. The $v_{\text{N-H(stretch)}}$ in the region 3430 cm⁻¹-3190 cm⁻¹ corresponding to the free amines in PY-COF is found to be almost absent in PY-NCS-COF. This is accompanied with the appearance of $v_{N=C=S(stretch)}$ signal at 2050 cm⁻¹. ¹⁵N NMR spectrum (Supplementary Fig. 30) again shows a diminished intensity of the amine signal at -330 ppm together with the concurrent appearance of a weak, but prominent signal at $-274 \,\mathrm{ppm}$ corresponding to the nitrogen atom of the isothiocyanate group. The chemical shift of this signal matches well with the ¹⁵N NMR signal for PhNCS. From the ¹³C spectrum (Supplementary Fig. 31) it is difficult to identify a signal due to the isothiocyanate carbon (see Supplementary Fig. 32 for EDX analysis); it is also known for the isothiocyanate carbon to be near-silent or not observed at all in ¹³C NMR spectra⁶⁴⁻⁶⁷. Interestingly, although there is a slight broadening of reflections, the PXRD pattern of PY-NCS-COF shows that crystallinity is retained in this functional group transformation reaction (Fig. 5b). Similar to PY-COF, the experimental PXRD pattern shows eight peaks at 3.7, 4.1, 4.7, 6.1, 7.3, 8.7, 10.9, and 15.2° 2θ , together with the broad stacking reflection at $2\theta = 24^{\circ}$. The changes in reflection intensities as compared to PY-COF could be successfully modeled by replacing the dangling amine groups with isothiocyanate functional groups during Rietveld refinement, further validating the structural transformation. Rietveld refinement of the experimental powder pattern resulted in unit cell parameters of a = 53.024 Å, b = 23.551 Å, c = 3.794 Å, and $\alpha = \beta =$ 90°, $\gamma = 92.106$ °, $R_{wp} = 1.62$ (Supplementary Data 5). From the Argon sorption isotherm at 87 K (Supplementary Fig. 33), the BET surface area of PY-NCS-COF is measured to be 979 m² g⁻¹. Thus, a complete topochemical functional group interconversion could be achieved with our sub-stoichiometric COFs.

Discussion

We report sub-stoichiometric COFs (Figure 8), by demonstrating an unconventional combination of tritopic and tetratopic linkers to form a hitherto unexpected 2D COF of a previously unidentified network topology where the pyrene-based linker P uniquely assumes a simultaneous dual role of di- and tetratopic linker in two different coordination environments. The presence of periodic uncondensed amine functional groups in the imine-linked substoichiometric PT- and PY-COFs is a principal observation with immediate consequences for both the design principles and functionalization of COFs: this design concept goes beyond conventional post-synthetic functionalization, while still retaining its essence.

Although we have demonstrated this concept for imine-linked COFs, it may be extended to other COF linkages as well. The implication of these results is profound; it shows that complex

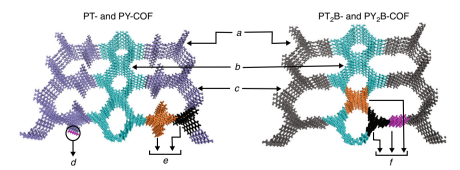


Fig. 8 Summary of the unique features of the sub-stoichiometric and the three-component COFs. a bex network topology. b Supramolecular ribbon-like motifs as secondary building units for hierarchical design. c Highly crystalline and porous, dual pore COFs. d Free functional groups by sub-stoichiometric design—free amines in an imine-linked COF, functionalizable, and catalytically active. e Combination of tetratopic and triangular linkers—dual linking mode of tetratopic molecule. f Combination of di-, tri-, and tetratopic linkers

structures can be achieved by unusual linker combinations, without having to pursue arduous synthesis of sophisticated linkers. For example, in an analogous way, the combination of appropriate hexatopic and tetratopic linkers can be envisaged to produce a sub-stoichiometric COF (Supplementary Fig. 34). Our result shows that by just considering the linker geometry, it may not always be possible to predict the correct COF structure and it prompts us to think beyond conventional retrosynthetic topological deconstruction rules for designing COFs and to create a new class of structures with unique complexities, including possibly quasicrystalline COFs (Supplementary Fig. 35). Most importantly, such sub-stoichiometric COFs are characteristically linked to free functional groups (without the need of separate functionalization of the linkers), which can either be designed for use as such, or can be further derivatized for specific applications with the COF as the template, which again opens up a new direction of research. For example, the free amine moieties can be used to link molecular proton reduction co-catalysts to increase the photocatalytic H₂ evolution efficiency⁶⁸. The possibilities are numerous.

Methods

Synthesis of PT-/PY-COF. P (0.076 mmol, 43.2 mg), T/Y (0.076 mmol, 30 mg), 1,4-dioxane (3 ml), mesitylene (3 ml), and aqueous acetic acid (6 M, 0.6 ml) were added to a Biotage© precision glass vial, sealed, and heated under autogenous pressure at 120 °C for 72 h. After the reaction was allowed to cool down, the reaction mixture was filtered and washed thoroughly with water, tetrahydrofuran, chloroform, and acetone, and then dried in high dynamic vacuum overnight to get PT- and PY-COF in 72% and 68% yield, respectively.

Synthesis of PT₂B-/PY₂B-COF. P (0.064 mmol, 36 mg), T/Y (0.127 mmol, 50 mg), B (0.064 mmol, 12 mg), 1,4-dioxane (5 ml), mesitylene (5 ml), and aqueous acetic acid (6 M, 1 ml) were added to a Biotage© precision glass vial, sealed, and heated under autogenous pressure at 120 °C for 72 h. After the reaction was allowed to cool down, the reaction mixture was filtered and washed thoroughly with water, tetrahydrofuran, chloroform, and acetone and then dried in high dynamic vacuum overnight overnight to get PT₂B- and PY₂B-COF in 75% and 72% yield, respectively.

Synthesis of PY-NCS-COF. To a solution of K_2CO_3 (20 mg, $0.145\,\mathrm{mmol})$ in water (10 ml), 50 mg of PY-COF was added. To the resulting suspension, CS_2 (40 µl, 0.66 mmol) was added dropwise. After the addition was complete, the reaction mixture was stirred overnight at room temperature. The reaction mixture was then cooled to 0 °C and a solution of cyanuric chloride (5.1 mg, 0.0276 mmol) in 5 ml dichloromethane was added dropwise. After complete addition, the reaction mixture was stirred for 3 h and then filtered. The residue was basified with 6 N NaOH solution and then washed with water until the washings were pH neutral followed by washing with acetone, chloroform, and tetrahydrofuran and then dried in high dynamic vacuum overnight to get PY-NCS-COF in 62% yield.

Data availability

All data supporting the findings of this study are available within the Article and its Supplementary Information and/or from the corresponding authors upon reasonable request. Reprints and permissions information is available online at www.nature.com/reprints.

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Author contributions

T.B. and F.H. contributed equally. T.B. initiated the project, planned and performed all experiments and wrote the manuscript with inputs from F.H. F.H. developed the structure model and analyzed PXRD and TEM data. S.T. performed sorption measurements. B.P.B. synthesized linker Y with F.H. and co-performed post modification of COFs. G.S. and C.O. performed quantum-chemical calculations of NMR chemical shifts. V.D. recorded EM images and simulated the projected potential maps. I.M. recorded solid-state NMR spectra. B.V.L. supervised the research. All authors read and commented on the manuscript.

Additional information

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