

Cite this: *Chem. Sci.*, 2025, 16, 13221

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Structural control of dynamic covalent cages: kinetic vs. thermodynamic assembly and PFAS removal from water†

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Dynamic covalent chemistry is a powerful tool to synthesise complex structures from simple building blocks. However, even minor variations in the numerous parameters governing self-assembly can drastically influence the size and structure of the resulting assemblies. Herein, we report the selective formation of three cages belonging to the low-symmetry $\text{Tri}_2^2\text{Tri}^2$ cage topology for the first time, using highly symmetric tritopic building blocks, confirmed by single-crystal X-ray (SC-XRD) analysis. Fluorinated and non-fluorinated aldehydes were combined with two amines differing in their degree of structural flexibility. Applying either kinetic or thermodynamic control through solvent selection allowed for the selective synthesis of either the low-symmetry $\text{Tri}_2^2\text{Tri}^2$ or the larger, highly symmetric Tri^4Tri^4 assemblies. While the fluorinated linker strongly preferred the formation of the $\text{Tri}_2^2\text{Tri}^2$ cage topology under thermodynamic control, the non-fluorinated linker selectively formed the Tri^4Tri^4 species. Kinetic control, using methanol as a poor solvent, allowed for the selective precipitation of the $\text{Tri}_2^2\text{Tri}^2$ intermediate. Reduction of the Janus-like fluorinated $\text{Tri}_2^2\text{Tri}^2$ cages yielded the cages $\text{Et}^2\text{F}_{\text{red}}^2$ and $\text{TREN}^2\text{F}_{\text{red}}^2$, which showed high potential for removing perfluorooctanoic acid (PFOA) from water, with $\text{Et}^2\text{F}_{\text{red}}^2$ exhibiting structural rearrangements in organic solvents to accommodate PFOA, as observed by ^1H and ^{19}F NMR titrations in combination with ^{19}F DOSY measurements.

Received 24th March 2025

Accepted 18th June 2025

DOI: 10.1039/d5sc02247a

rsc.li/chemical-science

Introduction

Molecular self-assembly using organic building blocks, in combination with metals or in their absence, gives access to a variety of nanostructures through the organisation of tailor-made molecular building blocks that exhibit suitable intra- and intermolecular interactions.¹ Organic chemistry-rooted dynamic covalent chemistry (DCC) is therefore an efficient synthetic strategy that employs multitopic precursors to form reversible covalent bonds, combining the advantages of error correction during self-assembly with stability, allowing for in-solution analysis and application of the assembled molecule.² It has enabled the synthesis of a variety of supramolecular architectures, from macrocycles³ to cages,⁴ polymers,⁵ and covalent organic frameworks (COFs),⁶ with potential energy surfaces being the central concept in understanding the assembly outcome.^{4a,7}

Building blocks of similar topology, through changes in geometry,^{1c,8} size,^{1,4,8} rigidity of linkers,^{1,9} or even changes in substituents¹⁰ can build a multitude of accessible cage topologies.^{4,11} Recently, Jelfs and co-workers rationalised expected topologies (connection patterns) by analysing the geometry and topology of building blocks through calculations combined with experimentally observed structures.^{11b} They also introduced a systematic nomenclature for describing cage topologies, denoted as X^m_pY^n , providing clarity and avoiding confusion with bracket notation, such as $[4 + 2]$, which is more commonly associated with pericyclic reactions in organic chemistry.

By combining ditopic (Di), tritopic (Tri), or tetratopic (Tet) building blocks, a diverse library of cage geometries can be envisioned. Although several gaps have been filled in recent years, some predicted geometries remain unobserved, with no corresponding crystal structures reported. Within the Tri^mTri^n family (Fig. 1), Tri^4Tri^4 is the only geometry for which crystal structures have been commonly obtained.¹² In contrast, Tri^1Tri^1 has mostly only been observed in solution,¹³ while the lower-symmetry $\text{Tri}_2^2\text{Tri}^2$ topology has yet to be reported. The formation of such complex, lower-symmetry structures is relatively rare compared to highly symmetric Platonic or Archimedean solid-based topologies, such as cubes¹⁴ or tetrahedra.¹⁵ Computational studies have shown that the higher symmetry of such assemblies is generally preferred from an entropic point of view,^{16a} however, it cannot be fully disregarded that the formation

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† Electronic supplementary information (ESI) available. CCDC 2430764. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d5sc02247a>

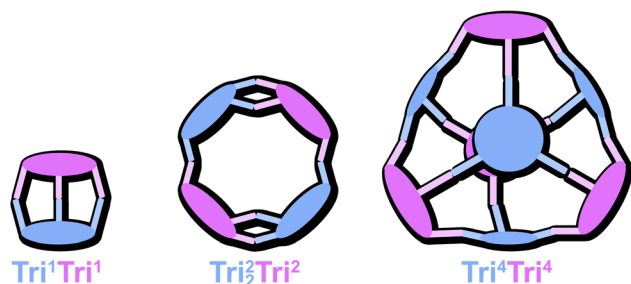


Fig. 1 Schematic representation of the cage geometries obtained when reacting two tritopic building blocks with each other, giving different topologies.

of multiple smaller structures, disregarding their symmetry, should be overall more beneficial for the entropic term.^{11b,16b}

Especially in recent years, the interest in these intricate assemblies has grown due to their unique and selective host-guest chemistry.^{17,18} For metal-organic cages (MOCs), various strategies have been explored to achieve complex assemblies.¹⁷ One common approach involves employing multiple linkers to form heteroleptic cages, effectively disrupting the symmetry of the final structure.¹⁹ In DCC-based systems this approach is of great interest, however, the desired social self-sorting^{4c,20,21} is rare, and narcissistic self-sorting^{10,22} or statistical mixtures²³ are dominating this space. Thus, lower-symmetry assemblies are typically achieved by employing less symmetric building blocks, often exhibiting an inherent chirality.^{18,22a,b,24} For example, He and Zhang *et al.* demonstrated that the use of C_2 and C_{2v} building blocks leads to the formation of a C_2 -symmetric imine cage of the unusual Tet⁴Di⁸ topology.^{24b}

Beyond linker design, reaction conditions can significantly influence the assembly process and resulting topology. Even if using the same starting materials, solvent choice²⁵ and/or the concentration²⁶ used can drastically shift the equilibrium towards different topologies by either enhancing or suppressing inter-/intramolecular interactions, respectively.^{26a}

These examples, however, predominantly focus on the formation of the thermodynamic product. While many examples support the thermodynamically controlled formation of imine cages, some observations suggest that they may instead be kinetically controlled products, especially when precipitating from the reaction mixture.^{12d,25b,27,28} This can be rationalised by considering the potential energy surface of the system, where the cage structure may correspond to a kinetically trapped state rather than the global thermodynamic minimum, and cage formation can be driven by precipitation, preventing further equilibration towards more stable assemblies, highlighting the complexity of controlling self-assembly.

Results and discussion

Synthesis and characterisation of the cages

Herein, we report the synthesis of three novel cages of the unique Tri₂Tri₂ geometry, including topological assembly control and the complexation of per- and polyfluoroalkyl

substances (PFAS) in organic media and their removal from aqueous solutions. For that purpose, we chose the flexible aldehyde **F**, offering the combination of an electron-rich core and electron-deficient fluorinated panels, aiming for a Janus-type nanocavity to facilitate the intermolecular host-guest interactions.^{12c} Stirring (2,4,6-triethylbenzene-1,3,5-triyl) trimethanamine (**Et**) with the flexible **F** in an equimolar ratio for 3 days in methanol at room temperature resulted in the precipitation of a colourless solid. To our surprise, both the ¹H and ¹⁹F NMR of a redissolved sample showed a complex set of signals (Fig. 2c), whereas, on the contrary, MALDI-MS only showed a singular signal belonging to a condensation of two **F** with two **Et** building blocks. The lower C_{2h} symmetry of the obtained cage **Et**²**F**² results in a splitting of the observed resonance signals, showing a 2 : 1 ratio for all signals, which is in good accordance with the ratio expected for a cage of Tri₂Tri₂ geometry. This unrepresented cage topology distinguishes itself through its lowered symmetry as a result of the two double connections found within its structure (Fig. 2a).^{11b} The resulting inherent strain and rigidity of the assembly cause a diastereotopic splitting of the protons H_E and H_F belonging to the double-connected amine motif (Fig. 2c).

Intrigued by these results, we investigated different conditions for the cage formation, aiming for either thermodynamic or kinetic control over the assemblies. For that purpose, we chose chloroform as solvent, heating to 60 °C to allow all intermediates to remain in solution, over time reaching the thermodynamic equilibrium. For experiments under kinetic conditions, we investigated the two most common “poor solvents” (lower solubility of intermediates and products), acetonitrile and methanol, to induce precipitation of possible intermediate structures. The respective mixtures were stirred at room temperature upon dropwise addition of a solution of the respective amine.²⁷

Using acetonitrile as the solvent, stirring at room temperature for 3 days again selectively led to the formation of the Tri₂Tri₂ species **Et**²**F**² precipitating from the reaction solution. Stirring **Et** and **F** in chloroform at 60 °C for three days resulted in the formation of a second, more symmetric species, without any precipitation occurring. MALDI-MS of the obtained mixture revealed the formation of the highly symmetric (T_d) Tri⁴Tri⁴ topology alongside **Et**²**F**² (see Fig. S1 and S2†). In an attempt to isolate the observed different cage topologies, the dynamic covalent imines were reduced by *in situ* reaction with sodium borohydride.²⁹ The respective amine cage **Et**²**F**²_{red} could be isolated in 37% yield from the one-pot two-step reaction of **Et** and **F** or in 94% yield from **Et**²**F**². Again, HRMS (ESI⁺) and NMR analysis confirmed the formation of **Et**²**F**²_{red}, whereas the larger **Et**⁴**F**⁴_{red} structure could not be isolated. **Et**²**F**²_{red} readily crystallised from a chloroform solution, and the obtained crystals were subjected to single-crystal X-ray analysis (SC-XRD), unambiguously confirming the anticipated Tri₂Tri₂ structure (Fig. 2b). As a result of the two double connections between two singular **F** and **Et** motifs, the amine cage is flattened overall, resembling a double-walled macrocycle of C_{2h} symmetry, with two different angles for Ar–O–Ar_F bonds to accommodate the inherent strain expected with less flexible building blocks.^{11b} One ethyl group of



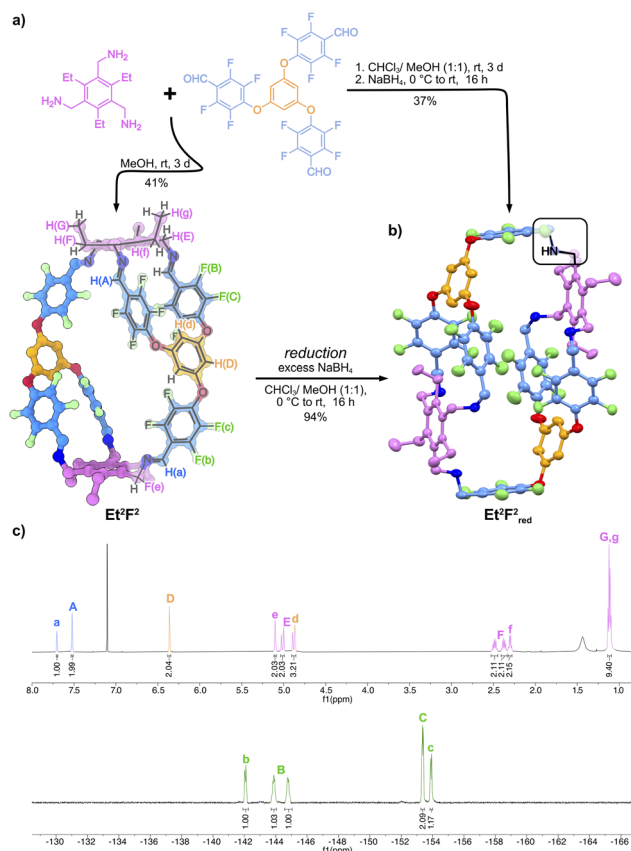


Fig. 2 (a) Synthetic route towards imine cage **Et²F²** and the respective amine cage **Et²F²_red**; (b) the SC-XRD structure of **Et²F²_red** with thermal ellipsoids set at 50% probability, hydrogens and solvents were omitted for clarity; (c) ^1H and ^{19}F NMR (600 MHz/282 MHz, CDCl_3 , 298 K) spectra of the imine cage **Et²F²** showing the splitting of all signals due to the reduced symmetry.

the **Et** motif is located between the fluorinated panels of each double connection. The same is true for the analogous hydrogen of the phloroglucinol motif, explaining the strong upfield shifts observed in ^1H NMR (Fig. S123,† and 4b, H_f , H_g). Meanwhile, all free electron pairs of the amine nitrogen are pointing inwards, forming two main cavities where residual chloroform solvent molecules are located. Overall, even with the increased flexibility of the amine bonds compared to the more rigid imine bonds, the structure still appears potentially strained.

Thus, in addition to **Et**, the flexible tris(2-aminoethyl)amine (**TREN**) was investigated in combination with aldehyde **F**, anticipating that under thermodynamic control, enrichment of the geometry encoded in the linker would be observed, as the rigidity is reduced.

Interestingly, again the $\text{Tri}_2^2\text{Tri}_2$ topology was favoured over the larger $\text{Tri}_4^4\text{Tri}_4$ species (see Table S1†). Heating the reaction mixture in chloroform led to the almost exclusive formation of **TREN²F²** over **TREN⁴F⁴** (96 : 4). In contrast, precipitation from methanol or acetonitrile reaction mixtures favoured the formation of the $\text{Tri}_4^4\text{Tri}_4$ species, likely due to the increased solubility of the cages and their intermediates compared to the **Et**-based counterparts.²⁷ This is supported by the observation

that in acetonitrile solution, only **TREN⁴F⁴** was found in the precipitate, whereas the filtrate contains a mixture of cage topologies similar to those observed in chloroform (acetonitrile filtrate **TREN²F²** : **TREN⁴F⁴** ratio 94 : 6). Overall, these findings indicate that aldehyde **F** preferentially directs the formation of the $\text{Tri}_2^2\text{Tri}_2$ cage topology. Under thermodynamic control, this is less pronounced when employing the more rigid amine **Et** in comparison to **TREN**, with kinetic conditions exclusively leading to the formation of **Et²F²** alongside insoluble oligomeric species typical for these conditions.^{12d} The inverse behaviour of the **TREN**-based cages is likely a result of two factors; (a) their increased solubility, allowing for a higher proportion of $\text{Tri}_4^4\text{Tri}_4$ species to be formed before precipitation occurs, and (b) the distance between the fluorobenzene motifs, which is influenced by the rigidity of the amine building block.

We rationalised that the $\text{Tri}_2^2\text{Tri}_2$ topology is directed through small intramolecular interactions between the fluorinated benzene motifs leading to a preorganisation,³⁰ where two panels are near to each other in solution leading to a preferred formation of a double connection between one **Et** molecule and two aldehyde groups of a singular **F** molecule. This is in line with previous observations where we found that using fluorinated aldehydes can result in the formation of the unusual $\text{Tri}_6^6\text{Di}_9$ topology alongside the otherwise strongly favoured, well-known $\text{Tri}_4^4\text{Di}_6$ topology.³¹ Additionally, the preferred formation of **TREN²F²** over **TREN⁴F⁴** suggests that these interactions are (a) lessened with the more preorganised and less flexible amine **Et** and (b) play a significant role in the stabilisation of the newly obtained $\text{Tri}_2^2\text{Tri}_2$ cage topology.

Therefore, we also investigated the non-fluorinated aldehyde derivative **H**, expecting no significant interactions between its panels. Aldehyde **H** was readily prepared in a two-step procedure starting from phloroglucinol (see the ESI†). To our delight, the initial screenings under either thermodynamic (in chloroform at 60°C) or kinetic control (acetonitrile or methanol at room temperature) revealed that the highly soluble and flexible amine **TREN** exclusively formed the $\text{Tri}_4^4\text{Tri}_4$ geometry (Fig. 3). The respective imine cage **TREN⁴H⁴** could be isolated from chloroform in quantitative yield (Fig. 3a). As expected for reactions under kinetic control, **TREN⁴H⁴** precipitated from the reaction mixture along with insoluble by-products. Similarly, the **Et**-based cage (**Et⁴H⁴**) formed quantitatively under thermodynamic control. In contrast, NMR and MALDI-MS analyses of the precipitate from methanol revealed the clean formation of the lower symmetry cage **Et²H²** of $\text{Tri}_2^2\text{Tri}_2$ topology (Fig. 3c). Upon extraction from the insoluble by-products, the lower symmetry cage could be isolated in 38% yield.

Monitoring a redissolved sample of **Et²H²** in CDCl_3 (2.3 mM, consistent with the concentration used for cage synthesis) at either 60°C or at room temperature resulted in the appearance of a new set of signals. After one day, noticeable amounts of **Et⁴H⁴** had formed, whereas at room temperature only marginal amounts of the $\text{Tri}_4^4\text{Tri}_4$ species **Et⁴H⁴** were observed. Over the course of seven days, the $\text{Tri}_4^4\text{Tri}_4$ species became increasingly enriched, and after 24 days, the complete conversion to **Et⁴H⁴** was observed under both conditions. At 60°C , this initial transformation proceeded notably quicker, reaching a 1 : 1 cage

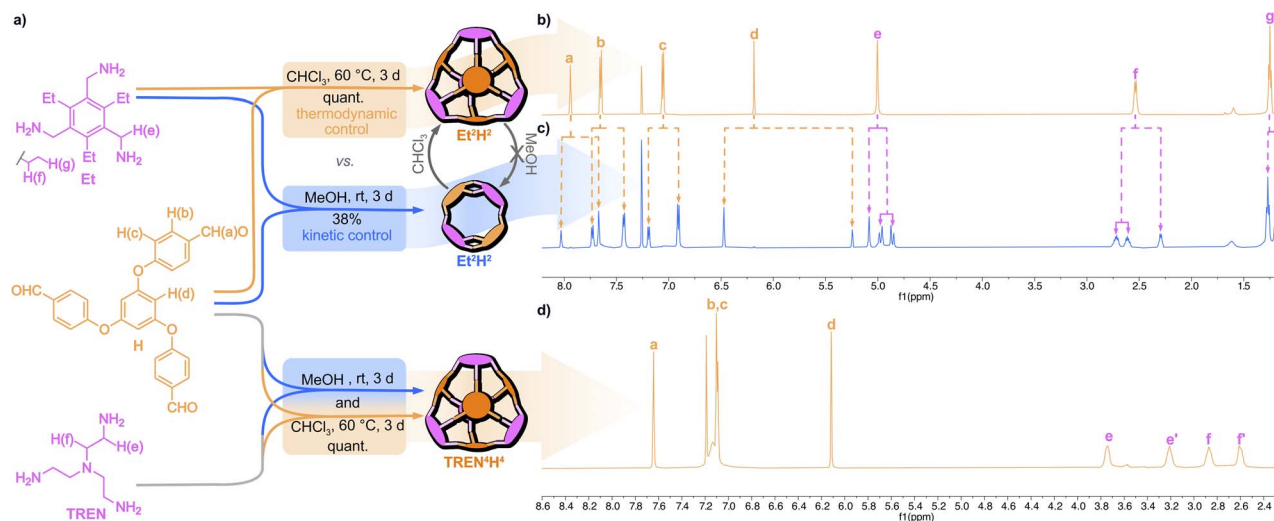


Fig. 3 (a) Synthesis of the non-fluorinated cages under either thermodynamic or kinetic control with a schematic representation of the Tri₂²Tri₂ and Tri⁴Tri⁴ cage geometries and the respective ¹H NMR (600 MHz, CDCl₃, 298 K) spectra of (b) **Et⁴H⁴**, (c) **Et²H²**, and (d) **TREN⁴H⁴**. The signal assignments are shown in (a).

ratio already after five days, while at room temperature this was reached only after about nine days (Fig. S14–S16†). Similarly, when stirring **Et²H²** in CDCl₃ for 3 days at 60 °C, an almost complete cage-to-cage transformation towards **Et⁴H⁴** was observed. At room temperature, only small amounts (~34%) of **Et⁴H⁴** were formed. Suspending **Et⁴H⁴** in methanol and stirring for three days, even at 60 °C, did not result in any observable interconversion (Fig. S13†).

This underlines the bias towards the larger, highly symmetric, and less internally strained Tri⁴Tri⁴ topology over the low-symmetry Tri₂²Tri₂ topology in solution,^{11b} strongly suggesting that **Et²H²** is an intermediate structure formed on the pathway towards **Et⁴H⁴**. To rule out that reaction temperature plays a role in this observation, we reacted **Et** with **H** in chloroform at room temperature, again selectively forming **Et⁴H⁴**, while stirring in methanol, even at 60 °C, leading to **Et²H²** as the singular species, strongly suggesting that the solubility of the intermediates is the discriminating factor.

To test this, during cage formation studies, the chloroform content was fixed at 10% to fully dissolve the starting materials, while the acetonitrile concentration was varied. For a methanol/chloroform mixture (90 : 10), **Et²H²** almost exclusively precipitated from the solution. As the acetonitrile content increased, the proportion of **Et⁴H⁴** in the precipitate also increased, until at 40% acetonitrile content, only **Et⁴H⁴** precipitated from the reaction solution (see Fig. S10 and S11†). This trend aligns with the observed solubility differences of aldehyde **H**, which is poorly soluble in methanol but highly soluble in acetonitrile. Additionally, the faster precipitation observed in mixtures with lower acetonitrile content (Fig. S12†) suggests that (a) the solubility of intermediates in the respective solvents can be roughly estimated from the solubility of the employed building blocks and (b) the primary factor driving the formation of **Et²H²** is precipitation.^{25b} These findings highlight that, under kinetic control, a mixture of species might precipitate, but adjusting

the solvent composition can dramatically shift the equilibrium. Thus, to achieve a desired outcome, the solubility of the building blocks should be carefully considered when selecting the solvent for self-assembly reactions.

PFOA removal

With the understanding of the dynamic covalent chemistry behind the formation of the Tri₂²Tri₂ cage topology, we shifted our focus back to the fluorinated Tri₂²Tri₂ cages. As previously shown, the imine cages could be easily reduced *in situ* to the respective amine cages **Et²F₂^{red}** and **TREN²F₂^{red}** with sodium borohydride. Additionally, **Et²F₂^{red}** with the electron-rich cores with fluorinated, electron-deficient panels, featuring exposed amine groups, was investigated for the uptake of perfluorinated octanoic acid (PFOA) and trifluoroacetic acid (TFA). Per- and polyfluoroalkyl substances (PFAS), or “forever chemicals,” are synthetic compounds used in industry for their thermal stability and resistance to degradation. Found in coatings, foams, and textiles, they persist in the environment, accumulate in organisms, and pose serious health risks.³² Despite restrictions, PFAS contamination remains a major global challenge due to their removal complexity.³³ Supramolecular chemistry approaches have been employed to tackle the complex challenge of sieving PFAS molecules, which exhibit both hydrophilic and lipophilic behaviour, such as leveraging the rich host–guest chemistry of supramolecular compounds³⁴ including cyclodextrin derivatives,^{35d} water-soluble iron-^{35e} or palladium-based,^{35a} and insoluble Zr-based^{35b} MOCs. Recently, Sessler and Chi *et al.* used a water-insoluble fluorinated amine-based cage feasible for the removal of PFOA from aqueous solutions through the reduction of the respective dynamic covalent imine cage.^{35c} Contrary to the binding of neutral compounds, *e.g.*, perfluorocarbons (PFCs), which is primarily governed by hydrophobic effects and π -stacking within nonpolar, shape-complementary cavities,³⁶ the binding of PFAS typically relies



on an interplay of electrostatic and fluorophilic interactions by using the polar headgroup as an anchor.^{35c-e}

First, we performed NMR titrations in organic media (chloroform/methanol mixture 95:5) at a concentration of 2.5 mM, investigating the cage's ability to capture PFOA and to study present interactions. Additionally, the symmetric "open-cavity" model compound **Bn³F¹_{red}** was prepared for the comparison with the "closed-cavity" cages. **Bn³F¹_{red}** was synthesised by condensation of aldehyde **F** with benzylamine, followed by *in situ* reduction with sodium borohydride.

Upon addition of the cages to a PFOA solution, shifts for all PFOA signals could be observed in ¹⁹F NMR spectra. The CF₂-group (F_a) neighbouring the carboxylic acid exhibited the most significant shift of 1.6–1.8 ppm, while the other signals, even including the terminal CF₃-group (F_g), were roughly shifted by −0.2 ppm (see ESI†). For the symmetric model compound **Bn³F¹_{red}**, however, only marginal shifts of (<0.02 ppm) for all signals besides F_a were noticeable. The addition of PFOA to the respective hosts led to a shifting of all host signals in ¹H and ¹⁹F NMR, with the CH₂–NH–CH₂ motif expressing the strongest shifts in ¹H NMR, while other signals were only slightly shifted (<0.05 ppm) in the case of **TREN²F²_{red}** and **Bn³F¹_{red}**. **Et²F²_{red}** showed noticeable downfield shifts for almost all signals. Most interestingly, the protons H_f and H_g were strongly downfield shifted by 0.78 ppm and 0.48 ppm, respectively, until almost overlaying with the other ethyl groups (H_f and H_g) of the **Et** motif. Simultaneously, both phloroglucinol protons H_D and H_d shifted upfield, indicating that **Et²F²_{red}** undergoes a structural rearrangement. Upon addition of PFOA, the singular ethyl group of the **Et** located between the double-connected fluorinated benzenes of **F** is supposedly pushed outwards, away from the fluorinated panels, being deshielded in the process. This assumption is supported by ¹H–¹H NOESY NMR data, which reveal that upon the addition of PFOA, the previously isolated ethyl group exhibits a new correlation pattern similar to that observed for the other ethyl groups (Fig. S75†). A similar structural change can be observed in the solid-state. When comparing the crystal structures of **PFOA@Et²F²_{red}** (grown from a dichloromethane/acetonitrile solution) and **Et²F²_{red}** (Fig. S17–S19†). **PFOA@Et²F²_{red}** shows a more elongated cage structure, where the ethyl groups are all pointing away from the fluorobenzenes and two PFOA molecules are located outside of the cage (Fig. 4a). The addition of a strong acid like trifluoroacetic acid did not result in a significant shifting of these signals, instead, a broadening of the signals is observed (Fig. S50 and S51†). When titrating octanoic acid, of comparable structure and with a pK_a of 3.8 ± 0.1 (vs. 2.2 ± 0.2)³⁷ no shifting of the cage signals can be observed (Fig. S45 and S46†), rendering the observed structural rearrangement of **Et²F²_{red}** to be selective towards PFOA, not being a result of simple protonation. ¹⁹F DOSY experiments unambiguously confirmed the formation of a complex between PFOA and **Et²F²_{red}** in organic media, showing almost identical diffusion coefficients (3.68 × 10^{−10} m² s^{−1} for PFOA and 3.81 × 10^{−10} m² s^{−1} for **Et²F²_{red}**) corresponding well to the one observed for pure cage (3.64 × 10^{−10} m² s^{−1}, see Table S3†).

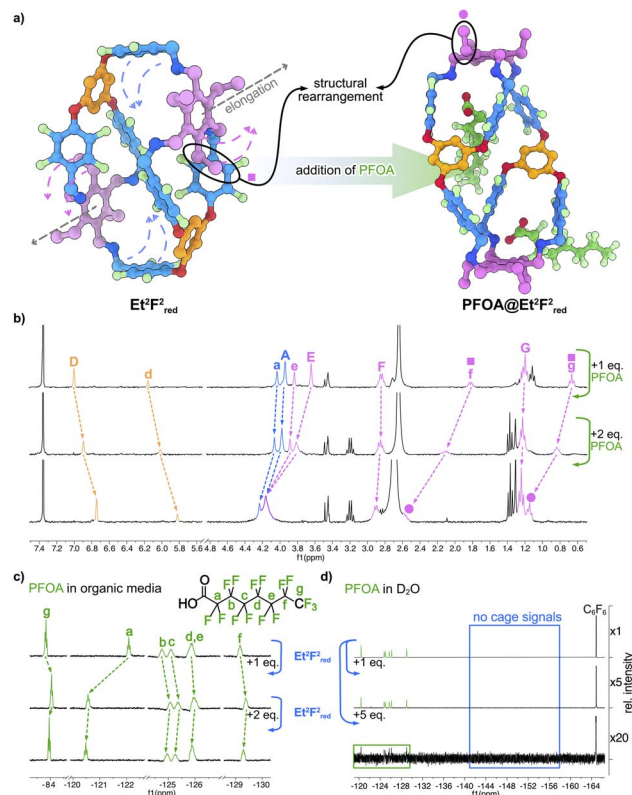


Fig. 4 (a) Schematic illustration of the PFOA-induced structural rearrangement in **Et²F²_{red}**, where one ethyl group (pink square) of the **Et** motif is pushed away from the two neighbouring fluorobenzenes of the **F** motif; (b) stacked ¹H NMR spectra (300 MHz, CDCl₃/MeOD, 95 : 5, 2.5 mM, 298 K), highlighting the chemical shifts in **Et²F²_{red}** upon the addition of PFOA, signal assignment analogous to Fig. 2a; (c) stacked ¹⁹F NMR spectra (282 MHz, CDCl₃/MeOD, 95 : 5, 2.5 mM, 298 K), highlighting the chemical shifts of PFOA upon the addition of **Et²F²_{red}**; (d) stacked ¹⁹F NMR spectra (565 MHz, 128 scans, D₂O, 298 K) showing the removal of PFOA from an aqueous solution using **Et²F²_{red}** as a heterogeneous adsorbent.

The observed ¹⁹F NMR shifts suggest that PFOA binding in organic media arises from a combination of electrostatic and fluorophilic interactions. Binding is primarily driven by electrostatic attraction between the carboxylic acid group of PFOA and the protonated, quaternary amines of the hosts. However, only the cages show additional interactions, as indicated by pronounced shifts of the PFOA signals, particularly of the terminal CF₃ group. This points to a binding mode in which the perfluoroalkyl chain remains largely outside the cage, while the CF₃ group partially inserts into the cage windows and interacts with the fluorinated panels. This interpretation is in good accordance with the preliminary SC-XRD data. Overall, this fluorophilic interaction appears to be weak and of dynamic nature, with the apparent encapsulation resulting mainly from spatial proximity rather than strong inclusion (see the ESI† for a detailed discussion). These findings are consistent with studies on COFs,³⁸ porous polymers³⁹ and macrocycle-based hydrogels,⁴⁰ where quaternisation of amine groups enhances PFAS uptake from aqueous solutions through an interplay of electrostatic and hydrophobic effects.

Encouraged by these promising observations, we investigated the ability of these highly hydrophobic cages to remove PFOA from aqueous solution. A 1 mg mL⁻¹ solution of PFOA in deionised water was prepared, to which 1 equivalent of the completely insoluble cages was added. After stirring the colourless suspension for one hour, the mixture was filtered through a syringe filter, and the clear filtrate was analysed by ¹⁹F NMR. To our delight, 1 equivalent of the cages already removed approximately 80% of the initial PFOA amount. Upon addition of 5 equivalents, no clear signals corresponding to PFOA or the cage were detected, indicating an almost complete removal (Fig. 4d). These results highlight the potential of these cages as heterogeneous, low-molecular-weight adsorbent materials for PFOA removal.

Conclusions

In conclusion, we were able to gain access to an unrepresented cage geometry of the Tri₂Tri² topology using only highly symmetric building blocks. This was accomplished by leveraging either thermodynamic or kinetic control over the self-assembly process. As building blocks, a fluorinated and a non-fluorinated aldehyde alongside one of two amines with differing degrees of preorganisation, flexibility, and solubility were used.

Investigations revealed that the fluorinated linker favoured the Tri₂Tri² cage topology under thermodynamic control, whereas the non-fluorinated linker **H** exclusively formed the larger symmetric Tri₄Tri⁴ derivatives under these conditions. Applying kinetic control allowed for the selective formation of the low-symmetry Tri₂Tri² cages that precipitated from the reaction mixtures. Studies conducted strongly suggest that the flexibility of the building blocks plays a crucial role, enabling the formation of the Tri₂Tri² species as an intermediate towards larger structures. This understanding allowed us to use solvent selection to direct the assembly pathway, enabling the formation of either the intermediate low-symmetry Tri₂Tri² cages under kinetic control or the larger high-symmetry Tri₄Tri⁴ structures under thermodynamic control.

Additionally, the reduction of the fluorinated Tri₂Tri² cages led to the low-symmetry Janus-like cages **Et²F²_{red}** and **TREN²F²_{red}**, which demonstrated promising potential for the removal of PFOA from aqueous solutions. Even in organic solvents, **Et²F²_{red}** indicated interactions selectively with PFOA, undergoing a structural rearrangement to accommodate PFOA.

Our findings highlight the delicate aspects of self-assembly pathways in directing cage assembly and provide new insights into the use of fluorinated and non-fluorinated linkers to tailor structural outcomes. This approach expands the toolbox of supramolecular chemists, offering a new route to design cage-like compounds with enhanced structural complexity, paving the way for enzyme-like complex host-guest structures and advanced lightweight functional materials.

Data availability

The data supporting this article have been included as part of the ESI.† Crystallographic data for **Et²F²_{red}** (2430764) has been

deposited in the joint Cambridge Crystallographic Data Centre (CCDC) and Fachinformationszentrum Karlsruhe Access Structures service, available free of charge.

Author contributions

The study was conceptualised by T. P. and B. M. S. The majority of experiments were conducted by T. P., including the synthesis and characterisation, assisted by P. M. M. P. M. M. and F. Z. performed the titration experiments. A. M. was responsible for ¹⁹F DOSY NMR measurements and data analysis, B. M. S. conducted SC-XRD data acquisition and refinement. The first draft of the manuscript, review, editing, and data presentation was executed by T. P. and B. M. S. All authors approved the final submission.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank Jona Voss, MSc, for help preparing ¹⁹F DOSY NMR samples. T. P. is grateful for a PhD fellowship from the Evangelisches Studienwerk Villigst. B. M. S. acknowledges the support from the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) SCHM 3101/6. F. Z. and P. M. M. are thankful towards the Ministerio de Ciencia e Innovación of Spain, FEDER project PID2020-113483GB-I00 and the Fundación Séneca Región de Murcia (CARM) Project 21956/PI/22. P. M. M. is very thankful for a mobility aid for beneficiaries of the pre-doctoral contract programme of the Plan de Fomento de la Investigación, according to resolution R-606/2021, financed by the University of Murcia.

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