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A crystalline sponge based on dispersive forces suitable for X-ray structure determination of included molecular guests†

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A crystalline porous material showing one-dimensional (1-D) rectangular micropores ($12 \times 9 \text{ \AA}^2$) has been assembled from a semirigid macrocyclic tetraimine and EtOAc as the templating agent. The 1-D nature of the material is intrinsic to the conformationally rigid structure of a macrocyclic sub-unit bearing four cyclohexylidene residues. The multiple dispersive forces established among the aliphatic residues glutted the 1-D channels and provided thermal stability to the material at temperatures below 160°C . Upon removal of the template, the structure of the empty solid exhibited permanent microporosity ($S_{\text{BET}} = 342 \text{ m}^2 \text{ g}^{-1}$). Being a true molecular sponge, the channel framework of this material allowed the inclusion of a variety of molecular sample guests without compromising its crystalline nature. Remarkably, this crystalline material enabled the structure determination by X-ray diffraction of the included molecules. Theoretical studies demonstrated the vital role played by the dispersive forces in the overall stabilization of the crystal packing.

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Introduction

Porous organic materials¹ are currently of growing interest owing to their potential applications in diverse fields such as storage and separation of gases,² molecular sorting³ and catalysis.⁴ In this context, the synthesis of porous materials constructed from discrete molecules⁵ is particularly attractive. This approach takes advantage of the structural versatility provided by organic synthesis and the possibility to obtain porous solid materials simply by crystallization.⁶ The molecular components of porous organic molecular crystals (POMC) lacking an extended directional covalent or coordination bonding are held together only by weak intermolecular forces. Hence, molecular materials based on non-directional intermolecular forces are metastable and tend to collapse upon removal of the template agents used to stabilize the voids of their porous crystalline structures.⁷ Related materials exclusively stabilized by weak dispersive interactions are scarce. In this regard, Atwood and Barbour's contributions are particularly significant. These

authors showed that interstitial van der Waals interactions could be employed as the main component for the design of porous host-guest assemblies.⁸ A logical development of this approach that produced more stable porous molecular materials relied on the use of molecules interacting through hydrogen bonding and aromatic interactions, either separately or in combination.^{2d-f,7} Using this strategy, Shimizu and colleagues reported a microporous crystalline bis-urea macrocyclic host that self-assembled by hydrogen bonding and aromatic stacking interactions. In this example, the macrocyclic ureas aligned into columnar 1-D assemblies and maintained its porous structure even upon guest removal.⁹

In this work, we present a new organic material featuring permanent porosity based on the macrocyclic tetraimine **3** (Scheme 1). A key aspect of our design is the use of multiple weak dispersive interactions as the glue element to organize and maintain together the components of the stable porous molecular material. Our report constitutes a paradigmatic example of the so-called Gulliver's principle¹⁰ applied to the design of porous organic materials.

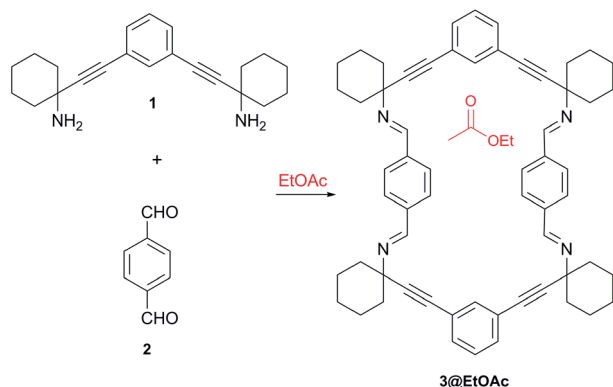
Previously, we have described the synthesis of the 1,3-phenylene-bis-propargylic diamine **1** used in this work as rigid spacer to avoid the collapse of the macrocycle **3**.¹¹ The hierarchical self-assembly of the macrocycle **3** into a porous material arises from the combination of multiple van der Waals and aromatic interactions.¹² These interactions are established between the phenyl and cyclohexylidene residues, the "sticky groups" of **3**. The resulting porous material displays 1-D channels that are large enough for liquid-solid sorption applications.¹³

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† Electronic supplementary information (ESI) available: Synthesis of macrocycle **3** together with characterization data. TGA and DSC traces. CIF files of all structures. Computational details and additional calculations. CCDC 1012389–1012397 and 1063714. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5sc01838b



Scheme 1 Synthesis of macrocyclic tetraimine 3.

More specifically, we were interested in knowing if a crystalline porous material based only on dispersive forces, could be used as the crystalline solid support in X-ray diffraction studies by applying the “crystalline sponge method” developed by Fujita.¹⁴ In this method, accurate molecular structures of samples can be obtained by analyzing the diffraction patterns of single crystals without requiring the crystallization of the samples. However, the crystalline sponges reported so far are based on metal–organic frameworks (MOFs) that include electron-rich atoms, such as iodine, bromine, chlorine as well as transition metals in their structures.¹⁵ These atoms contribute heavily to the overall diffraction pattern masking the information available for the guest. From this point of view, our system based only in light-atoms, could offer an alternative to MOF-based crystalline sponges.

Results and discussion

Synthesis and structure of the microporous material

The macrocyclic tetraimine 3 was prepared by Schiff base condensation of the diamine 1 with terephthalaldehyde 2 in EtOAc solution. On standing at room temperature, single crystals of 3@EtOAc slowly grew from the solution (yield 50%).[‡] The solid material was characterized by single crystal X-ray diffraction,¹⁶ ¹³C CP MAS NMR and thermogravimetric analysis (TGA).

The single crystal X-ray structure showed the macrocycle 3 molecules stacked one on top of another forming a columnar assembly. The solid material displayed one-dimensional channels with distorted rectangular morphology along the *b* axis (Fig. 1). The total solvent accessible void volume within the channels calculated from PLATON/Squeeze, was 1096 Å³ (274 Å³ per each macrocyclic imine) representing 19.4% of the unit cell volume.¹⁷ We observed that the channels were filled with partially disordered EtOAc molecules (Fig. 1a). In the crystal packing, the four aromatic rings of 3 were tilted *ca.* 60° with regard to the plane of the macrocycle. Furthermore, the two phenyl rings located at the wider sides of 3 oscillated between two positions (75 : 25 occupancy) and acted as revolving doors (Fig. 1b).

From this structure, it became clear that the combined action of intermolecular displaced π – π aromatic stacking, as

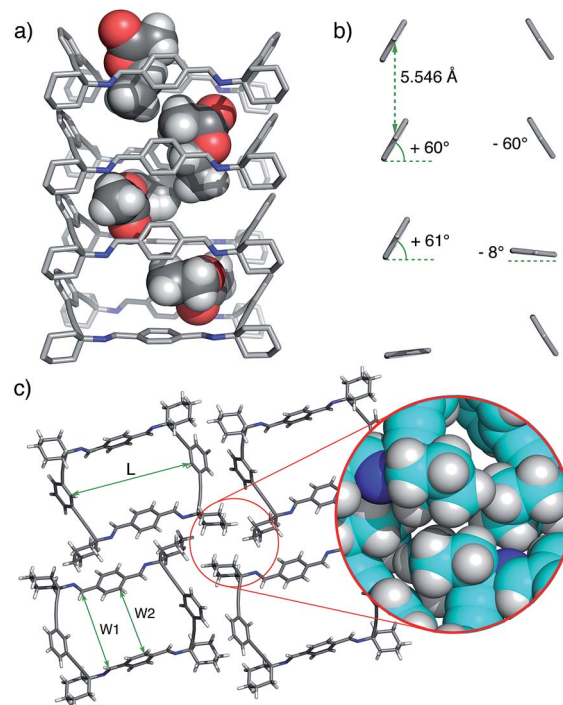


Fig. 1 (a) Representative X-ray perspective side-view in *b* direction of 3@EtOAc crystals displaying disordered EtOAc molecules inside the channel. (b) Partial view showing the relative orientation of the 1,4-phenylene rings (see text). (c) Top view of a molecular cell unit of the monoclinic crystals ($V_c = 5642 \text{ \AA}^3$, $Z = 4$) of 3@EtOAc ($D_c = 1.121 \text{ g cm}^{-3}$; $L \sim 11.98 \text{ \AA}$, $W1 \sim 9.13 \text{ \AA}$; $W2 \sim 6.11\text{--}6.90 \text{ \AA}$) (CCDC no. 1012389). Disorder and EtOAc solvent molecules are removed for clarity. The inset shows multiple van der Waals contacts between cyclohexylidene residues.

well as multiple CH (sp^3)–CH (sp^3) van der Waals interactions extending throughout the channel, provided the necessary stabilization of the assembly.

The results obtained in ¹³C CP-MAS NMR experiments confirmed the role of EtOAc as guest template. The carbon signals of the relatively mobile molecules of EtOAc included in the solid, but not those of the host tetraimine 3 that formed the porous framework were observed in the standard mode ¹³C MAS spectra (Fig. S7 in ESI[†]).

Stability and dynamic behaviour of the porous material

Guest solvent molecules were completely removed by treating the solvate 3@EtOAc under vacuum for several hours at room temperature (12 h, $2 \times 10^{-2} \text{ mm Hg}$). This treatment afforded a new crystalline material, the apohost 3(I). The X-ray structure of 3(I) (Fig. 2) revealed the absence of electron density in the channels previously filled with EtOAc. The lack of EtOAc in the crystalline structure was also confirmed by registering the ¹H NMR spectra of single crystals of 3(I) dissolved in CDCl₃.

More important, the crystal packing of 3(I) was almost identical to that of the 3@EtOAc solvate (see Table S1 in ESI[†]), indicating that the porous structure remained intact after vacuum treatment.



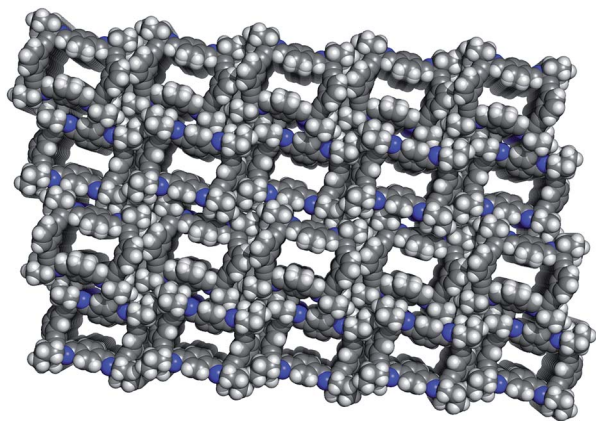


Fig. 2 X-ray perspective top-view of the apohost **3(I)** ($V_c = 5639 \text{ \AA}^3$, $Z = 4$) ($D_c = 0.9859 \text{ g cm}^{-3}$; $L \sim 11.98 \text{ \AA}$, $W1 \sim 9.20 \text{ \AA}$; $W2 \sim 6.90 \text{ \AA}$) (CCDC no. 1012398).

The relatively easy removal of the included EtOAc molecules at room temperature implied that they were weakly retained in the channels.

Remarkably, the **3@EtOAc** solvate could be regenerated by simply soaking crystals of the empty material **3(I)** in EtOAc. These findings were consistent with the absence of strong dipole-dipole or hydrogen bonding interactions between the cycloimine **3** and the included EtOAc molecules.

The dynamic nature of the guest exchange constituted an essential feature of this material. The kinetics of the guest exchange process was monitored by using ^{13}C NMR. Crystals of **3@ ^{13}C -[1,2]-EtOAc** obtained by treating **3(I)** with ^{13}C labelled EtOAc were suspended in unlabelled EtOAc.

We observed that the combined integral values of the isotopic ^{13}C doublet grew steadily with time and reached a plateau after 8–10 h (Fig. 3). This experiment demonstrated that the almost complete exchange of the included ^{13}C labeled EtOAc was slow in the human time-scale. Desolvation of EtOAc was also monitored using thermogravimetry. The TGA trace of **3@EtOAc** revealed an 11.8% weight loss in a temperature range from room temperature to 175°C . This weight loss corresponded to the vaporization of 1.3 molecules of EtOAc per macrocycle, therefore indicating the 3 : 4 stoichiometry found for this complex (see Fig. S9 in ESI †). In complete agreement, the differential scanning calorimetry (DSC) thermogram of **3@EtOAc** exhibited an endothermic peak at around 160°C due to vaporization of the entrapped guest solvent (Fig. 4b).

On the other hand, the porosity of **3(I)** was also confirmed by gas absorption of nitrogen at 77 K (Fig. 4a). The empty solid **3(I)** exhibited a typical type I adsorption isotherm indicative of a microporous ($<2 \text{ nm}$) structure with significant specific surface area ($S_{\text{BET}} = 342 \text{ m}^2 \text{ g}^{-1}$).¹⁸ In addition, the DSC trace of the apohost **3(I)** displayed an exothermic and irreversible process at around 175°C (Fig. 4b and S10 †), which we assigned to a monotropic solid-phase transition of **3(I)**. In both cases, the X-ray structure analysis of the material obtained after thermal treatment above 160°C , revealed the formation of a new non-porous polymorphic form **3(II)**.

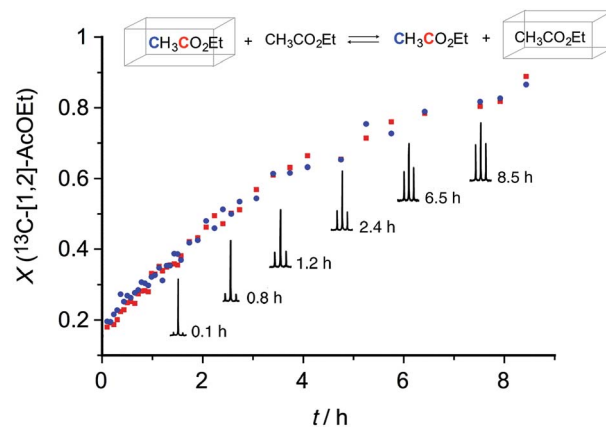


Fig. 3 Plot representing the fraction of ^{13}C -[1,2]-EtOAc in a solution of EtOAc. This plot was obtained from 50 mg of labelled **3@ ^{13}C -[1,2]-EtOAc** suspended in 0.5 mL of EtOAc at 298 K , by integration of the ^{13}C satellites with respect to the ^{12}C signal and assuming the complete replacement of ^{13}C -[1,2]-EtOAc. The inset shows representative increasing carbonyl doublet resonances (169.7 ppm , $J_{\text{C-C}} = 59.5 \text{ Hz}$) used to extract the integral data.

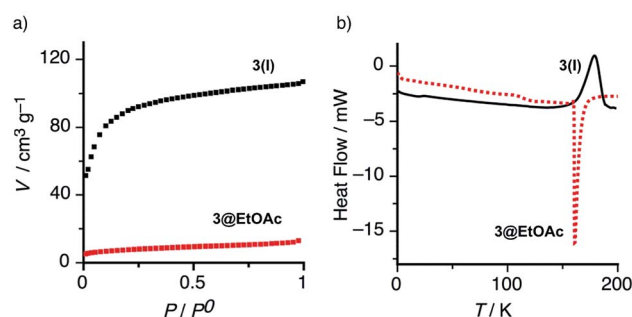


Fig. 4 (a) Adsorption isotherm of N_2 for the solvate **3@EtOAc** compared to the empty solid **3(I)** ($T = 77 \text{ K}$, $P^0 = 1 \text{ atm}$). (b) DSC traces of **3@EtOAc** displaying an endothermic peak at 160°C due to solvent vaporization (red dotted line) and of **3(I)** showing an exothermic peak at 177°C assigned to irreversible phase transition to **3(II)**.

The X-ray structure of **3(II)** (see Fig. S11 in ESI †) showed densely packed layers of displaced macrocyclic imine units **3** ($D_c = 1.127 \text{ g cm}^{-3}$) and the concomitant loss of the columnar assembly characteristic of the **3(I)** polymorphic form.

Liquid–solid sorption experiments

The void volumes of the porous frameworks exhibited by the crystalline materials **3@EtOAc** and **3(I)** are sufficiently large to include a variety of molecular guests. This characteristic, together with the absence of polar groups that can be displayed to the included guests and the high thermal stability of the crystalline materials, suggested us the potential use of **3@EtOAc** and **3(I)** as suitable supports for X-ray diffraction studies by the crystalline sponge method.^{14,15}

To test this functionality, we soaked single crystals of **3@EtOAc** and **3(I)** in different neat liquid samples with molecular volumes ranging from $55\text{--}60 \text{ \AA}^3$ *i.e.* nitromethane



and ethylene glycol, to $>200 \text{ \AA}^3$ of *Z*-stilbene (see, Table S2 in ESI†). In favourable cases, *i.e.* less than 5 min for nitromethane, the initially floating crystals of **3@EtOAc** gradually sank to the bottom of the vial providing a visual indication of the exchange status. During the process, we did not observe any apparent degradation of the crystal integrity. The exchange of the guest samples was confirmed by ^1H NMR of CDCl_3 solutions of the solvates. Also, ^{13}C CP-MAS NMR experiments confirmed the inclusion of the liquid guests within the porous framework. The carbon signals of non-volatile guests (*i.e.* diethyl squarate) appeared slightly upfield shifted (0.5–2 ppm) with respect to those of diethyl squarate in the bulk (Fig. S8 in ESI†).

Owing to the poor solubility of the solvates in CDCl_3 , the molar ratios were better evaluated by digesting the solvates with $\text{DCl}/\text{D}_2\text{O}$ (50 μL) before registering the ^1H NMR spectra in a mixture of $\text{DMSO}-d_6$: CDCl_3 (2.5 : 1 v/v, 600 μL) (see, ESI†).^{15c} In general, the molar ratios of **3** : sample evaluated by ^1H NMR after a common period of 72 h, agree well with the crystallographic stoichiometries (Table S2 in ESI†). In one case, the average experimental molar ratio calculated for *Z*-stilbene of 0.8 was consistently inferior to that expected for a 1 : 1 **3@Z-stilbene** solvate. This result is likely due to the incomplete inclusion of *Z*-stilbene within the channels of the **3(I)**. Therefore, the molecular volume of *Z*-stilbene of $216 \text{ \AA}^3$, approaching the maximum available void volume of $274 \text{ \AA}^3$ per macrocycle calculated using PLATON/Squeeze,¹⁷ suggested us a practical upper limit for the effective inclusion of samples in **3(I)**.

Fig. 5a–h, depict ORTEP views of representative X-ray structures obtained by guest exchange from single crystals of **3@EtOAc** or alternatively, by soaking a single crystal of the empty apohost **3(I)** with *ca.* 1 mL of the liquid sample. Both methods produced identical results in all tested cases (see ESI for X-ray structural data†). Both, the empty material **3(I)** and the solvate **3@EtOAc** allowed the formation of inclusion solid

complexes (solvates) with aromatic, unsaturated and aliphatic neutral substrates bearing a variety of functional groups such as hydroxy, aldehyde, ester and nitro groups, among others.

Since the included guests could, eventually, establish hydrogen bonds or dipole–dipole interactions with **3** that would lead to degradation of the porous structure of the solids, the obtained results in the guest exchange/inclusion experiments confirmed that the interactions established between **3** and the incoming sample guests must be feeble.

Remarkably, the inclusion of the different guests into the crystal lattice did not disrupt the pore size significantly ($\Delta L < 2\%$; $\Delta W_1 < 2.6\%$). However, to obtain a better fit between the size and shape of the pore and those of the included molecules (induce fit), the two opposite phenyl rings located on the wider sides of the macrocycle **3** oscillated with a mechanism that resembles the functioning of the revolving doors. The adaptation process experienced by the pore was revealed by careful analysis of the X-ray structures of **3@EtOAc**, **3@p-anisaldehyde**, and **3(I)@S-(–)-nicotine** ($\Delta W_2 \sim 20\%$).

The induced fit process, in which the width of the channels responded to the size and shape of the included sample guests took place without any loss of crystal integrity (Table S1 in ESI†). In contrast, the X-ray structure of the non-porous polymorphic form **3(II)** displayed a narrowed entrance of the pore that can be referred as “closed” doors. Most likely, this structural change was responsible for inhibiting the guest sorption of liquid guests ($W_1 \sim 9.91 \text{ \AA}$; $W_2 \sim 5.54 \text{ \AA}$).

Theoretical calculations

(a) **Host–host interactions.** In order to rationalize the sorption properties of **3@EtOAc** and **3(I)**, we first studied the non-covalent interactions responsible for the formation of the one-dimensional channels and the columnar assembly in the

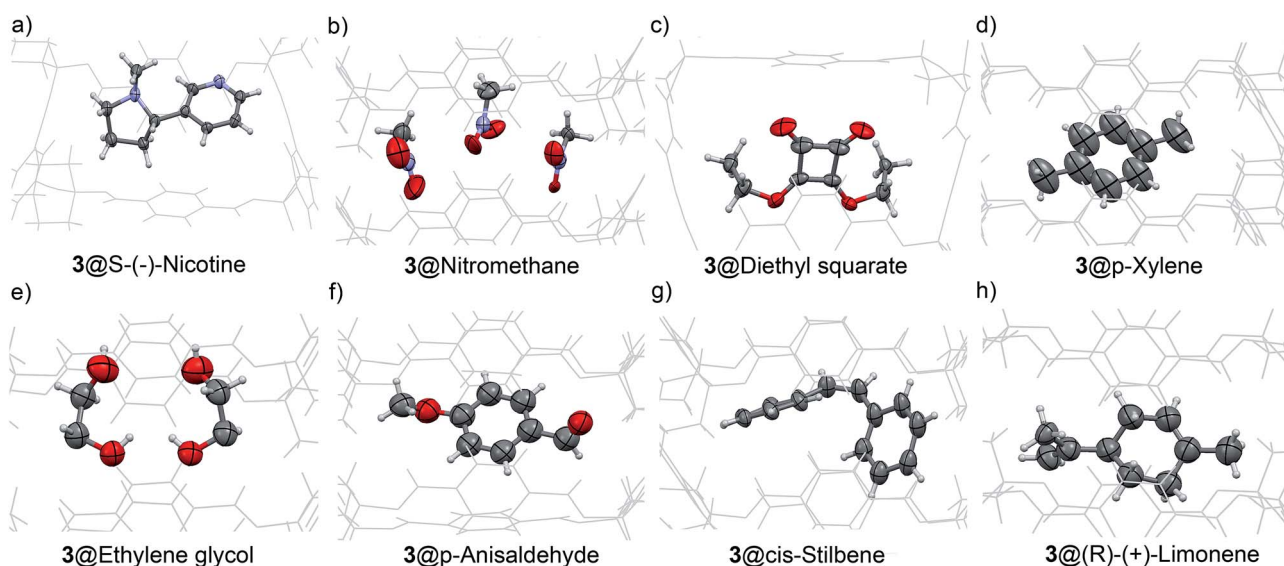
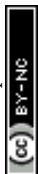


Fig. 5 (a to h) ORTEP views of different liquid molecules diffused into the channel structure of **3(I)** with thermal ellipsoids set at 50% probability. Disorder is removed for clarity. CCDC codes 1012389 to 1012397 and 1063714 contain the structural X-ray data (see ESI for RX structural details†).



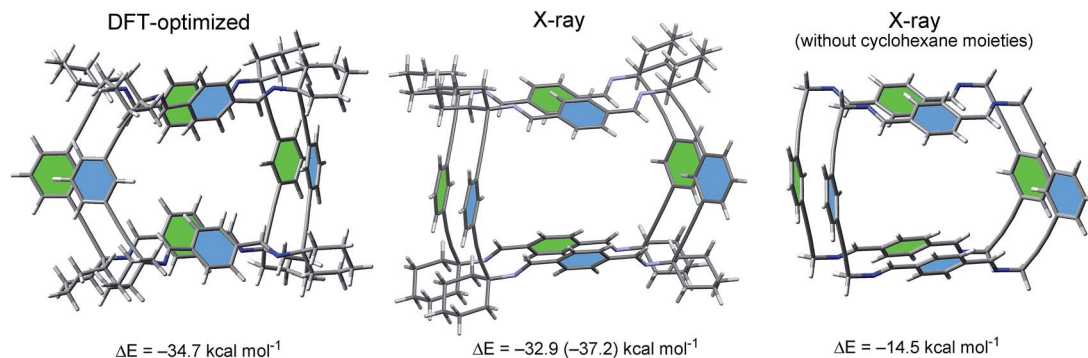


Fig. 6 Comparison of the X-ray and DFT-D3 optimized structures of **3(I)** and interaction energies at the BP86-D3/def2-TZVP level of theory (value in parenthesis corresponds to the BP86-D3/def2-SVP level).

crystal packing. We fully optimized a dimer of the macrocycle **3** starting from the crystallographic coordinates of **3(I)** and evaluated its interaction energy. The dimerization energy was large and negative ($-34.3 \text{ kcal mol}^{-1}$), indicating a strong binding (Fig. 6). The geometry of the energy-optimized dimer **3₂** was very similar to the one found in the solid state.

In addition, the interaction energy computed for the dimer using the crystallographic coordinates (without optimization) gave a very similar value ($-32.9 \text{ kcal mol}^{-1}$). Removal of the cyclohexylidene residues led to a significant reduction of the interaction energy ($-14.5 \text{ kcal mol}^{-1}$). This finding supported the strong influence exerted by the C-H $\cdots$ H-C interactions to the stability of the porous material.¹⁹

The energy difference calculated for the two energy-minimized dimers ($18.4 \text{ kcal mol}^{-1}$) represented an estimation of the contribution of these interactions to the overall dimerization process. Since four cyclohexylidene residues are present in the macrocycle **3**, each intermolecular cyclohexylidene $\cdots$ cyclohexylidene interaction contributes in $4.6 \text{ kcal mol}^{-1}$. This value was in good agreement with previously reported high level *ab initio* results for dodecahedrane.¹⁹

To further analyze the non-covalent interactions in the dimer **3₂** we used the NCI (Non Covalent Interaction) index.²⁰ This visualization index enables the identification and visualization of weak non-covalent interactions efficiently.

The isosurfaces corresponded to both favourable and unfavourable interactions, as differentiated by the sign of the

second density Hessian eigenvalue and defined by the isosurface color. NCI analysis allowed an assessment of host-guest complementarity and the extent to which weak interactions stabilize a complex. Fig. 7, shows the representation of the NCI plot computed for the dimer **3₂**. In addition to a panoply of C-H $\cdots$ H-C interactions, two weak equivalent and oppositely located parallel-displaced π - π interactions and four C H/ π interactions involving the π -system of the triple bond were also observed. The largest isosurface was located around the cyclohexylidene rings that established a multitude of C-H $\cdots$ H-C interactions with another cyclohexylidene and the aromatic ring.

(b) Substitution analysis. We theoretically analyzed the progressive substitution of EtOAc by diethyl squarate (SQA) inside the channel using a minimalist model of **3@EtOAc**. This exchange process was experimentally favourable. In our model system, we started from a minimal representative channel composed of only three macrocyclic imines complexed to three EtOAc molecules and locating one diethyl squarate (SQA) molecule at one end of the channel. We energetically evaluated the process of inserting the SQA molecule through this end and, simultaneously, removing one included EtOAc molecule out.

The process was favourable by $4.1 \text{ kcal mol}^{-1}$ (Fig. 8). Subsequently, we assessed the energy required to exchange the expelled EtOAc by a new SQA molecule. This second process was

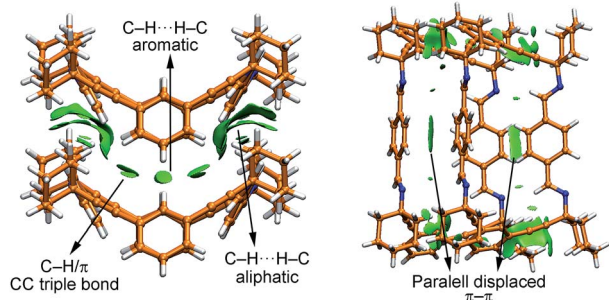


Fig. 7 NCI surfaces of a dimer of **3** used as a minimalist model of the 1-D channel. The isosurfaces (pale and dark green) correspond to favourable.

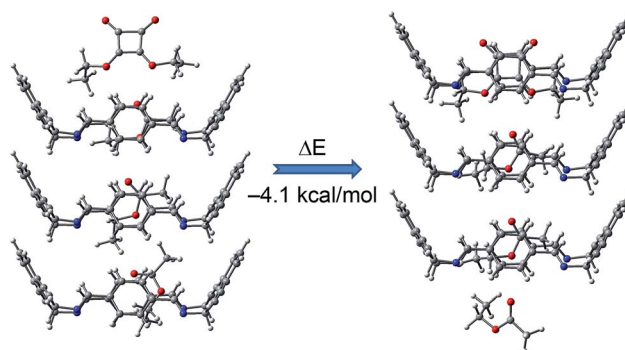


Fig. 8 Favourable energy change due to the substitution of one EtOAc molecule by one diethyl squarate molecule in a minimalist representative channel of **3@EtOAc**.



also favourable by 3 kcal mol⁻¹. This sequence was repeated to evaluate the energy of the exchange processes of the second and third molecules of EtOAc by SQA molecules (see Fig. S15–S17 in ESI†).

These processes were also energetically favourable and showed similar magnitudes for the exchange energies. Hence, the results of this theoretical study agreed well with the aforementioned experimental results and supported the viability of the liquid–solid guest exchange process.

Conclusions

In summary, we have synthesized an unprecedented conformationally semirigid macrocyclic tetraimine **3** that is decorated with four peripheral cyclohexylidene substituents. The tetraimine **3** crystallized from EtOAc solutions affording a porous crystalline material **3@EtOAc** composed of one-dimensional channels filled with solvent molecules. The crystal packing of **3@EtOAc** was stabilized mainly by dispersive interactions. We have demonstrated that the templating EtOAc solvent molecules can be removed *in vacuo* to yield an empty but crystalline and porous solid **3(I)**, which can be considered as a polymorphic form of **3@EtOAc**. Both solid materials **3@EtOAc** and **3(I)**, are rare examples of porous crystalline structures stabilized exclusively by dispersive forces. We confirmed the use of these solid crystalline materials for X-ray structure determination of molecular guests included in their pores by the crystalline sponge method. Finally, using high-level theoretical calculations we evaluated the strength of the cohesive interactions responsible for the porous structure, as well as the favourable energy change produced by guest exchange of molecules in the pores. Because porous materials based on dispersive forces have energetic advantages for the reversible storage of polar molecules in comparison to porous materials containing polar groups that can be presented to the included guests, we anticipate an emergence of materials of the former type in the near future.

Acknowledgements

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Notes and references

† Crystal data for **3@EtOAc** (CCDC no. 1012389): C_{65.36}H_{70.72}N₄O_{2.68}, *M* = 955.18, monoclinic, *a* = 34.794(16) Å, *b* = 5.836(3) Å, *c* = 28.131(14) Å, α = 90.00°, β = 97.862(11)°, γ = 90.00°, *V* = 5659(5) Å³, *T* = 100(2) K, space group *C2/c*, *Z* = 4, 35 400 reflections measured, 10 696 independent reflections (*R*_{int} = 0.0434). The final *R*₁ values were 0.0597 (*I* > 2σ(*I*)). The final *wR*(*F*²) values were 0.1624 (*I* > 2σ(*I*)). The final *R*₁ values were 0.0931 (all data). The final *wR*(*F*²) values were 0.1931 (all data). Crystal data for **3(I)**, (CCDC no. 1012398): C₆₀H₆₀N₄, *M* = 837.12, monoclinic, *a* = 34.8696(19) Å, *b* = 5.8099(3) Å, *c* = 28.1212(16) Å, α = 90°, β = 98.143(2)°, γ = 90°, *V* = 5639.6(5) Å³, *T* = 100(2) K, space group *C2/c*, *Z* = 4, 26 236 reflections measured, 8377 independent reflections (*R*_{int} = 0.0577). The final *R*₁ values were 0.0602 (*I* > 2σ(*I*)). The final *wR*(*F*²) values were 0.1686 (*I* > 2σ(*I*)). The

final *R*₁ values were 0.0825 (all data). The final *wR*(*F*²) values were 0.1883 (all data). Crystal data for **3(II)**, (CCDC no. 1012399): C₃₀H₃₀N₂, *M* = 418.56, triclinic, *a* = 6.5695(4) Å, *b* = 13.4759(8) Å, *c* = 14.1239(9) Å, α = 91.0310(10)°, β = 99.2650(10)°, γ = 91.6850(10)°, *V* = 1233.21(13) Å³, *T* = 100(2) K, space group *P1*, *Z* = 2, 13 275 reflections measured, 13 275 independent reflections (*R*_{int} = 0.0000). The final *R*₁ values were 0.0529 (*I* > 2σ(*I*)). The final *wR*(*F*²) values were 0.1334 (*I* > 2σ(*I*)). The final *R*₁ values were 0.0860 (all data). The final *wR*(*F*²) values were 0.1550 (all data).

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