

Intermolecular Hydrogen Bonding Directed by Aryl–Perfluoroaryl π – π Stacking Interactions

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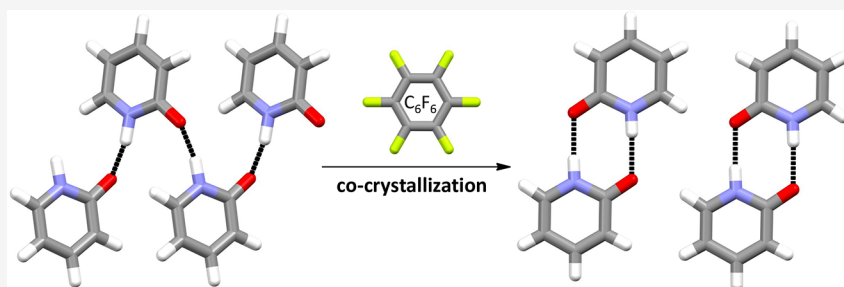
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ABSTRACT: The crystal structures of five compounds capable of forming self-complementary hydrogen bonds but crystallizing as catemers or creating more complex crystal structures were compared with those of their complexes prepared by cocrystallization with perfluoroaryl compounds. The results of X-ray diffraction revealed that in all the cases, the π – π stacking interactions caused the reorganization of hydrogen bonds and induced the creation of the expected self-complementary hydrogen bond dimeric motifs. The results point to the potential of aryl–perfluoroaryl interactions for the control of molecular self-assembly of aromatic compounds able to form hydrogen bonds. Also, the electrostatic potential (ESP) surfaces were investigated in terms of intermolecular interactions in the studied cocrystals. The values of the locally most negative and most positive ESP confirmed the best donor and acceptor sites for hydrogen bonding and other contacts.

INTRODUCTION

The rational design and preparation of crystalline molecular solids rely on control of the molecular arrangement using secondary bonding interactions rather than covalent bonding. Hydrogen bonds (HBs) and π – π stacking interactions are crucial in the construction of materials with the desired structure and function.^{1–6} Because HBs are generally stronger and more directional than other types of molecular interactions, they are vital for the control of self-assembly, orientation, and aggregation processes in the solid state.^{7–13} Therefore, identification and understanding of the preferential formation of specific types of HB arrays are indispensable to predict and rationalize the structural properties of organic molecular crystals.^{14–16} On the other hand, weaker non-covalent interactions, like π – π stacking between aromatic units, also play a significant role in crystal engineering because of their ability to control the relative orientation of molecules toward each other in the solid state.^{17–22} The principal modes of the arene–arene interaction are T-shaped edge-to-face and slipped face-to-face contacts. However, it is well-known that perfluoro-substitution at one aromatic ring causes domination of the face-to-face interaction mode^{23–25} due to the dispersion and quadrupolar interactions between electron-rich and electron-deficient aromatic rings.^{26–28} The electron-deficient regions of perfluorinated rings are now commonly referred to

as π -holes.^{29,30} Consequently, the stacking interactions can also be viewed as an interaction of such a π -hole with aromatic rings. The arene–perfluoroarene stacking interaction emerges as a reliable and robust synthon with potential utility in supramolecular chemistry,^{31–33} crystal engineering,^{34–38} discotic liquid crystals,³⁹ solid-state reactions,⁴⁰ and biomolecular interactions in solution.⁴¹

The majority of the reported crystallographic studies have dealt with cocrystals of aromatic hydrocarbons and perfluorocarbons.^{35–38} Much less research has been done on cocrystals of compounds bearing functional groups.^{42–47} Moreover, the competition or cooperation between hydrogen bonding and π – π stacking interactions in molecular self-assembly has been identified as a useful effect in crystal engineering.^{48–52} For example, we have recently shown that cocrystallization of star-shaped triaryl compounds with pentafluorophenol induces the formation of hydrogen-bonded cyclic trimers due to aryl–perfluoroaryl stacking interactions.⁵³

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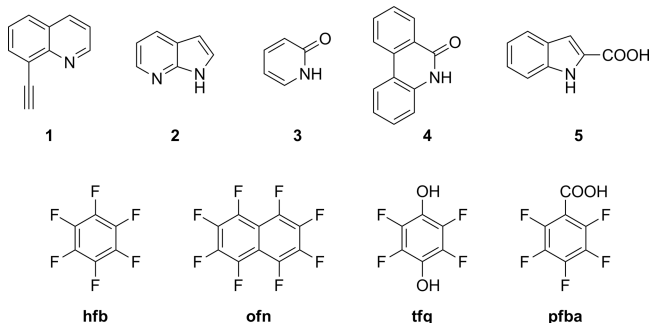
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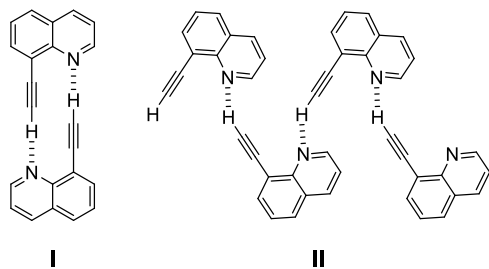
In this context, we were curious about whether $\pi-\pi$ stacking interactions could help to control the supramolecular assembly via HBs. As the first example, we chose 8-ethynylquinoline (**1**, Scheme 1), which was expected to form dimers due to self-

Scheme 1. Chemical Structures of Compounds 1–5 and the Perfluoroarenes Used in the Study



complementary $\equiv\text{C}-\text{H}\cdots\text{N}$ interactions in the form of a $R_2^2(12)$ motif (**I**, Scheme 2). However, the crystal structure of **1**

Scheme 2. Expected (I) and Observed (II) Supramolecular Motif in the Crystal Structure of 1



revealed the catemer C(6) motif (**II**) instead of the cyclic motif **I**. Much to our delight, the situation was reversed upon cocrystallization of **1** with hexafluorobenzene (**hfb**) that caused the self-assembly of molecules of **1** into dimers **I**. Thus, we searched for more examples of compounds crystallizing in rather uncommon packing modes instead of creating self-complementary HBs using popular supramolecular synthons. In this account, we investigated the crystal structures of compounds **1–5** before and after their complexation with perfluoroaryl derivatives such as hexafluorobenzene (**hfb**), octafluoronaphthalene (**ofn**), tetrafluorohydroquinone (**tfq**), or pentafluorobenzoic acid (**pfba**) (Scheme 1). In all of the above cases, aryl–perfluoroaryl stacking interactions led to a significant reorganization of the intermolecular HBs in the cocrystals containing **1–5**.

■ EXPERIMENTAL METHODS

Synthesis and Crystallization. Compounds **1**⁵⁴ and **3**⁵⁵ were prepared according to the literature procedures. The remaining substrates were purchased from Sigma-Aldrich, Alfa Aesar, or Apollo Scientific and used without further purification. The cocrystals **1₂·hfb**, **2₂·hfb**, and **3₂·hfb** were prepared by crystallization from hot hexafluorobenzene (**hfb**). Colorless crystals of the complexes are not air-stable and quickly decompose because of the volatility of **hfb**. However, they can be stored for a long time in the mother liquor or in the presence of **hfb** vapor. The cocrystal **4₂·ofn** was obtained by slow evaporation of the equimolar solution of the components in THF. The cocrystals with tetrafluorohydroquinone (**tfq**) or pentafluoro-

benzoic acid (**pfba**) were obtained by the same method from DCM–heptane. The details and characterization of the cocrystals are listed in the ESI.

Theoretical Calculations. To obtain molecular electrostatic potential maps of the molecules of the compounds studied, the geometries of the isolated molecules were first optimized using the Gaussian 16 package⁵⁶ at the B3LYP/6-31G** level of theory, and then their ESP maps were visualized with GaussView 6.1.1.⁵⁷ Frequency calculations were also included to determine whether the true energy minima were obtained. Finally, the maximum and minimum values of the electrostatic potential ($V_{S,\max}$ and $V_{S,\min}$) on the molecules were calculated.

Crystal Structure Determination. The X-ray diffraction data were collected on an IPDS 2T dual-beam diffractometer (STOE & Cie GmbH, Darmstadt, Germany) at 120.0(2) K with Mo $\text{K}\alpha$ radiation of a microfocus X-ray source (GeniX 3D Mo HighFlux, Xenocs, Sassenage, France, 50 kV, 1.0 mA, $\lambda = 0.71069 \text{ \AA}$) for all the structures. Every crystal was thermostated in a nitrogen stream at 120 K using a CryoStream-800 device (Oxford CryoSystems, UK) during the entire experiment. Data collection and data reduction were controlled by the X-Area 1.75 program.⁵⁸ The structure was solved by the SHELXT method^{59,60} and refined using Olex2^{61,62} and SHELX-2015^{59,60} program packages. Mercury⁶³ was used to prepare Figures 1–7. All the nonhydrogen atoms were modeled as anisotropic, whereas the H atoms were refined as isotropic. Hydrogen atoms were placed in idealized positions and refined with the usual restraints of the riding model. In **3₂·tfq**, the amide moiety (H1–N1–C13–O1) was modeled as disordered (*s.o.f.* 0.815(6)/0.185(6)). The crystallographic data and some details of the structural refinement are summarized in Table S1 in the ESI. Structural details of the hydrogen bonding are listed in Table 1.

Table 1. Parameters of the Selected Hydrogen Bonds in **1 and Cocrystals of **1–5**^a**

	D–H \cdots A	H \cdots A(Å)	D \cdots A(Å)	D–H \cdots A(°)
1	C8–H8 \cdots N1 ⁱ	2.35	3.199(2)	148
1₂·hfb	C11–H11 \cdots N1 ⁱⁱ	2.42	3.357(5)	167
2₂·hfb	N2–H2A \cdots N1 ⁱⁱⁱ	2.04	2.911(2)	168
3₂·hfb	N1–H1 \cdots O1 ^{iv}	1.91	2.773(1)	176
	CS–HS \cdots O1 ^v	2.53	3.399(2)	147
4₂·tfq	N1–H1 \cdots O1 ^{vi}	2.00	2.833(9)	158
	O2–H2A \cdots O1	2.01	2.829(3)	166
4₂·ofn	N1–H1 \cdots O1 ^{vii}	1.92	2.800(5)	177
5₂·pfba	O1–H1 \cdots O4	1.81	2.638(2)	170
	O3–H3A \cdots O2	1.78	2.607(2)	169
	O5–H5A \cdots O6 ^{viii}	1.80	2.627(3)	171

^aSymmetry codes: (i) $1.5 - x, -0.5 + y, 1.5 - z$; (ii) $2 - x, 1 - y, 1 - z$; (iii) $-x, 2 - y, 1 - z$; (iv) $1 - x, 1 - y, 1 - z$; (v) $-1 + x, y, z$; (vi) $1 - x, 1 - y, 1 - z$; (vii) $2 - x, 2 - y, -z$; (viii) $1 - x, -y, 2 - z$.

Crystallographic data for all of the structures reported in this paper have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication no. CCDC 2263874, 2263877, 2263878, 2263881, 2263886, 2263888, 2263893, and 2263894. The data can be obtained free of charge from The Cambridge Crystallographic Data Centre (CCDC) via <http://www.ccdc.cam.ac.uk/structures>.

■ RESULTS AND DISCUSSION

8-Ethynylquinoline (1). Numerous examples of the successful creation of discrete molecular complexes as well as 1D or 2D hydrogen bond networks from alkynyl C–H donors and azaheterocycles have been reported.^{64–69} These findings prompted us to investigate the possibility of constructing dimeric structures assembled by self-complementary $\equiv\text{C}-\text{H}\cdots\text{N}$ interactions. Thus, we obtained compound **1** and crystal-

lized it from AcOEt. Its crystal structure (monoclinic space group $P2_1/n$) revealed infinite antiparallel polymeric tapes spreading along the direction of [010]. These structures are created with the use of $\equiv\text{C}-\text{H}\cdots\text{N}$ bonds and connected via face-to-face π -stacking interactions. The distance between the planes of two adjacent molecules of **1** is 3.51 Å. Instead of the expected cyclic synthon **I**, the molecules are connected by linear $\equiv\text{C}-\text{H}\cdots\text{N}$ HBs creating a catemer structure **II** (Figure 1a).

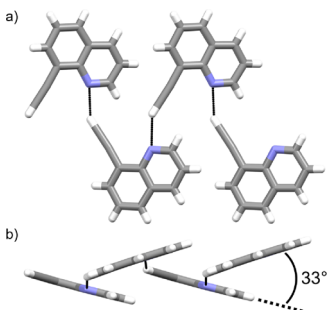


Figure 1. Crystal structure of **1**: (a) top view of a fragment of the tape stabilized by $\equiv\text{C}-\text{H}\cdots\text{N}$ HBs and (b) side view of the fragment showing the dihedral angle between the molecules of **1**.

The $\text{N}\cdots\text{H}$ distance of 2.35 Å indicates a relatively strong $\equiv\text{C}-\text{H}\cdots\text{N}$ interaction, even though the $\text{C}-\text{H}\cdots\text{N}$ bond angle of 148° is quite far from the linear orientation (Table 1).⁶⁸ Furthermore, the dihedral angle between two HB molecules generating the catemer is 33° (Figure 1b), whereas the cyclic synthon **I** requires coplanar orientation of two participating molecules of **1**. Therefore, we came up with the idea that aryl–perfluoroaryl interactions may force two neighboring molecules of **1** into a coplanar orientation and induce the formation of the dimer **I**. Cocrystallization of **1** with hexafluorobenzene (**hfb**) readily afforded 2:1 complex **1₂·hfb**.

It crystallizes in the monoclinic space group $P2_1/c$, and the asymmetric unit contains one molecule of **1** and half of a molecule of **hfb**. The crystal structure of **1₂·hfb** revealed that the molecules of **1** indeed aggregate into dimers in the form of cyclic motif **I**. The dimers are centrosymmetric and essentially planar. The $\text{N}\cdots\text{H}$ distances are 2.42 Å, slightly longer than those in the uncomplexed **1**, but the $\text{C}-\text{H}\cdots\text{N}$ angles are this time closer to 180° (Figure 2a, Table 1). Both factors indicate again relatively strong $\equiv\text{C}-\text{H}\cdots\text{N}$ interactions. The crystals of

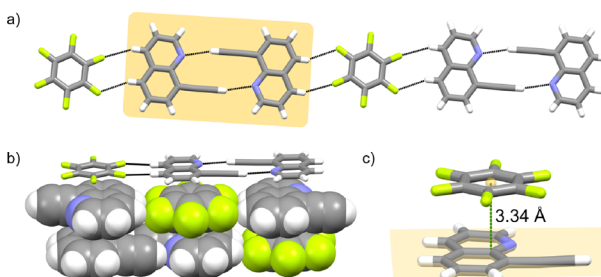
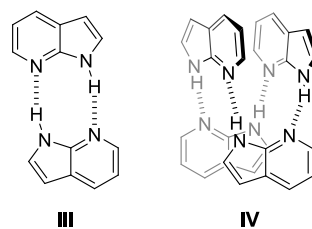


Figure 2. Crystal structure of **1₂·hfb**: (a) top view of a fragment of the tape composed of alternating dimers **I** and **hfb** molecules (one of the dimers is highlighted in light orange), (b) side view of the molecules stacked in a brick-wall motif, and (c) view of the relative orientation of the **hfb** and **1** molecules, with the distance between the centroid of C_6F_6 and the plane of **1**. HBs indicated by black lines.

the complex are composed of parallel tapes of alternating dimers **I** and **hfb** molecules connected by pairs of $\text{C}-\text{H}\cdots\text{F}$ interactions between the aryl and perfluoroaryl units (Figure 2a). It is known that the $\text{C}-\text{H}\cdots\text{F}$ interactions are common in crystals and cocrystals of fluoroaromatic compounds.⁷⁰ The $\text{H}\cdots\text{F}$ distances in **1₂·hfb** are 2.61 and 2.65 Å, so they are slightly shorter than the sum of the van der Waals radii of H and F (2.67 Å),⁷¹ and the $\text{C}-\text{H}\cdots\text{F}$ angles in both cases are approximately 154°. The structure of **1₂·hfb** is further stabilized by π -stacking interactions between the aryl and perfluoroaryl rings of the parallel tapes, which aggregate in a simple brick-wall motif (Figure 2b). The distance between the centroid of an **hfb** molecule and the plane of a quinoline moiety of **1** is 3.34 Å. Moreover, the molecules of each component are almost parallel, as the dihedral angle between the planes of the molecules is ca. 1.4° (Figure 2c).

7-Azaindole (2). Compound **2** (*1H*-pyrrolo[2,3-*b*]-pyridine, also known as 7-azaindole) attracts considerable interest in material chemistry as a blue emitter⁷² and in the field of drug chemistry.⁷³ Because of the close relationship with the nucleic purine bases, i.e., adenine and thymine, the self-association of **2** has been the object of numerous experimental and theoretical studies.^{74–76} In solution, this aza-aromatic compound forms cyclic dimers **III** joined by a pair of complementary N–H···N hydrogen bonds (Scheme 3). In addition to this, in the excited state, dimer **III** undergoes a photoinduced double proton transfer as evidenced by fluorescence spectra.⁷⁶

Scheme 3. Expected (III) and Observed (IV) Supramolecular Motif in the Crystal Structure of 2



Surprisingly, determination of the crystal structure has revealed that **2** crystallizes as tetramers **IV** with the use of four N–H···N HBs (Scheme 3).⁷⁷ One of the possible explanations of this behavior is the van der Waals interaction between roughly parallel heteroaromatic rings within tetramer **IV**. Also, in this case, it seemed reasonable to us to assume that a complexation of **2** with a perfluoroaryl compound, leading to stabilization gained by π – π stacking interactions, should favor the creation of the planar dimer **III** in the solid state. Indeed, cocrystallization of **2** with **hfb** resulted in the formation of the crystalline 2:1 complex **2₂·hfb**. Its crystal structure (the monoclinic space group $P2_1/c$) revealed centrosymmetric dimers **III**, where the molecules of **2** are held together by two complementary N–H···N HBs with an $\text{N}\cdots\text{H}$ distance of 2.04 Å and an N–H···N angle of 168°. The structure of **2₂·hfb** is stabilized by face-to-face stacking interactions between the aryl and perfluoroaryl rings, and the adjacent dimers interact via face-to-edge stacking interactions (Figure 3a). The distance between the centroid of the **hfb** molecule and the plane of the 7-azaindole molecule is 3.36 Å, and the angle between the planes of the molecules is ca. 1.8°. In the cocrystal, sheets composed of molecules of **2** (A) and **hfb** (B) can be

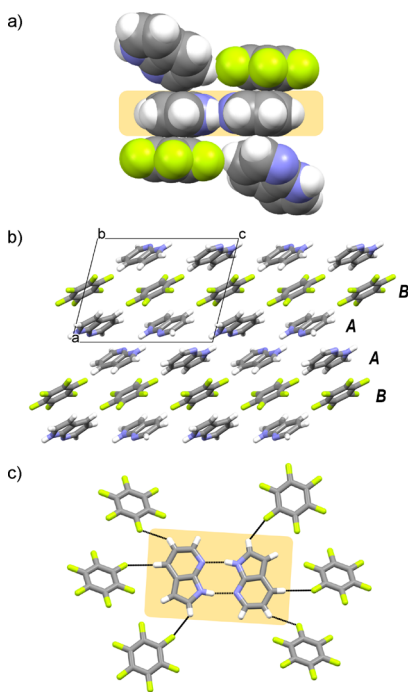
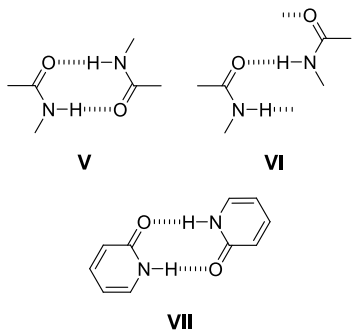


Figure 3. Crystal structure of $2 \cdot hfb$: (a) side view of a fragment of the structure indicating the interactions of 2_2 dimer (highlighted in light orange) with the neighboring molecules, (b) the sheets of molecules of 2 (A) and hfb (B) (view along the b -axis), and (c) view of the C–H···F bonds (black lines) between the dimer and the adjacent hfb molecules.

distinguished, and they are parallel to (100) and oriented in a $(BAAn)_n$ pattern (Figure 3b). Each dimer interacts with as many as six hfb molecules through C–H···F HBs (Figure 3c). The average H···F bond length is 2.62 Å.

2-Pyridone (3). Secondary amides assuming *s-cis* conformation (particularly lactams) most often assemble into dimers with the use of self-complementary $R_2^2(8)$ motif V (Scheme 4).⁸ In fact, self-complementary 2-pyridone (3)

Scheme 4. Supramolecular Motifs V–VI Observed in the Crystal Structures of Secondary Amides and the Dimeric Supramolecular Motif VII Characteristic for 2-Pyridone 3



moieties form one of the most reliable and robust hydrogen-bonded dimeric motifs, VII, which has found broad application in crystal engineering.^{9,78} Surprisingly, the crystal structure of the orthorhombic form of uncomplexed 3 reveals the catemer motif VI (puckered chains linked by linear N–H···O=C HBs) instead of the dimers.^{79,80} Nevertheless, a recently isolated monoclinic polymorph displayed the dimeric structure VII.⁸¹

As in the previous cases, it seemed reasonable to expect that the addition of a perfluoroaryl component should facilitate the formation of 3_2 self-complementary dimers. The cocrystallization of 3 with hfb indeed resulted in the formation of a cocrystal with a 2:1 ratio of the two components ($3_2 \cdot hfb$). Its crystal structure (monoclinic space group $P2_1/n$) contains the expected amide dimers stabilized by strong and complementary N–H···O=C HBs (Table 1). The arrangement of the amide molecules is similar to that of the monoclinic polymorph of 3; the dimers are almost planar and connected via weaker C–H···O HBs into undulated ribbons spreading along the a -axis (Figure 4a). This time, however, they are

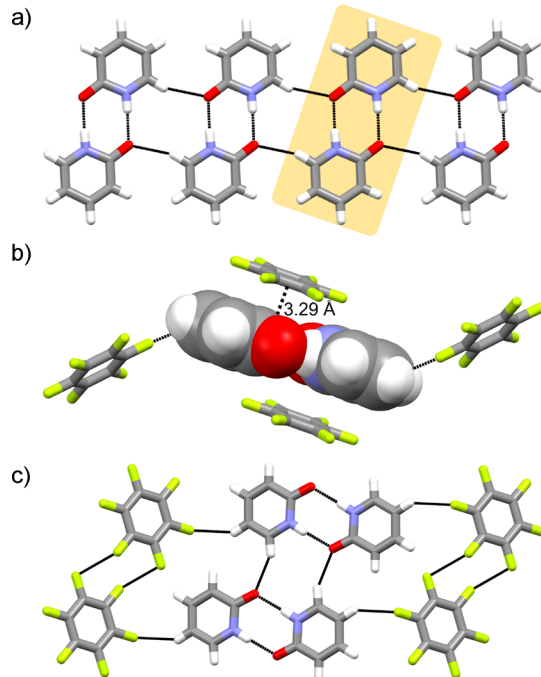


Figure 4. Crystal structure of $3_2 \cdot hfb$: (a) top view of a fragment of the undulated ribbon of molecules of 3 (one of the lactam dimers is highlighted in light orange), (b) side view of the dimer with four neighboring hfb molecules interacting with it and the distance between the centroid of hfb and the plane of 3, and (c) fragment of the structure depicting the network of HBs and F···F contacts (black lines).

centrosymmetric and separated by hfb molecules. In fact, the hfb molecules are located not only above and below the ribbons but also on their sides (Figure 4b). The polymeric structures interact with the hfb molecules through π – π stacking interactions and C–H···F HBs. The distance between the centroid of an hfb molecule and the plane of a molecule of 3 is 3.29 Å, but the molecules are slightly offset from each other. Also, the two molecules are not cofacial; the dihedral angle between the planes of the molecules is ca. 8°. The observed H···F bond lengths are equal to 2.61 Å and are shorter than the sum of the van der Waals radii of H and F by only 0.06 Å. Additional F···F contacts between the neighboring hfb molecules are also present (Figure 4c).

Phenanthridone (4). The aromatic lactam 4 (phenanthridin-6(SH)-one) is another example of a secondary *s-cis* amide whose molecules do not form dimers V in the solid state. Instead, they aggregate into a polymeric helical structure with the use of linear N–H···O=C HBs (motif VI, Scheme

4).⁸² Thus, we attempted to control the self-assembly of 4 by crystallizing it with perfluoroaryl compounds.

Because lactam 4 did not form cocrystals with hfb, we tried two different methods to generate the dimeric self-assembly of this compound, first by cocrystallization of 4 with tetrafluoro-*hydroquinone* (tfq). The 2:1 cocrystal of 4 with tfq (the monoclinic space group $P2_1/c$) is indeed composed of hydrogen-bonded lactam dimers (motif V). The length of both the N–H···O=C HBs is 2.00 Å, and the valence angle is equal to 158° (Table 1). The centrosymmetric dimers are not planar; the two molecules of 4 are located on parallel planes separated by ca. 0.72 Å (Figure 5). The amide moiety (H1–

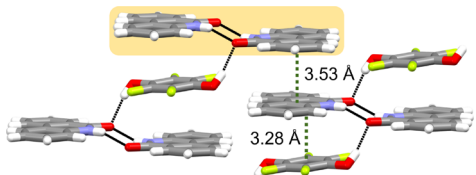


Figure 5. Crystal structure of $4_2\cdot$ tfq: side view of a fragment of the structure with the distances between π -stacked molecules. One of the lactam dimers is colored light orange. HBs are indicated by black lines.

N1–C13–O1) is disordered with a site occupation factor (*s.o.f.*) of 0.815(6)/0.185(6). The dimers are connected by quite unusual O–H···O=C HBs and π – π stacking interactions formed by the tfq molecules. The O···H distances are 2.01 Å, and the valence angle of the O–H···O bond is equal to 166°. Offset stacks of the molecules of 4 (C) and tfq (D) forming a (DCC)_n pattern can be distinguished (Figure 5). The distance between the centroid of a tfq molecule and the plane of the lactam molecule interacting with it is 3.28 Å, whereas the distance between two molecules of 4 is 3.53 Å. The molecules of both components are not parallel; the dihedral angle between the planes constructed for 4 and tfq is approximately 5°. We observed additional stabilization via C–H···F and C–H···O HBs as well as face-to-edge π – π stacking interactions between the lactam molecules.

The control of the self-assembly by cocrystallization with tfq cannot be recognized as fully satisfying because the tfq molecules are engaged in HBs with substrate 4. Thus, we tried cocrystallization with ofn, which afforded cocrystal $4_2\cdot$ ofn (triclinic space group $P\bar{1}$) again with a 2:1 stoichiometry. Its crystal structure revealed the anticipated lactam dimers V that, in contrast to $4_2\cdot$ tfq, are essentially planar. The 4_2 dimers and ofn molecules are coplanar and connected by means of C–H···F interactions forming flat ribbons spreading along [301] (Figure 6a). The polymeric structures aggregate on top of each other into infinite columns stabilized by almost perfectly cofacial π – π stacking interactions (the dihedral angle between the planes of the molecules of both components is equal to 0.4°). The interplanar distances are 3.34 and 3.42 Å for aryl···perfluoroaryl and aryl···aryl interaction, respectively (Figure 6b,c). The adjacent columns are connected via additional C–H···F HBs creating a stairlike 3D architecture.

Indole-2-carboxylic Acid (5). Carboxylic acids generally crystallize as cyclic $R_2^2(8)$ hydrogen-bonded dimers VIII, whereas the catemer C(4) motif IX (Scheme 5), where each carboxyl group is linked to two neighbors via single O–H···O=C HBs, is rarely encountered in the solid state.^{83,84}

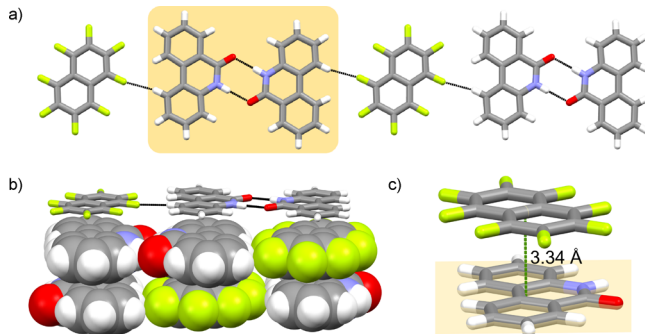
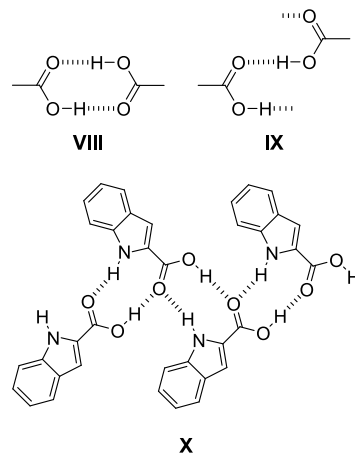


Figure 6. Crystal structure of $4_2\cdot$ ofn: (a) top view of a fragment of the ribbon composed of alternating lactam dimers and ofn molecules (one of the dimers is highlighted in light orange), (b) side view of the molecules stacked in a brick-wall motif, and (c) view of the relative orientation of the molecules of ofn and 4 with the distance between the centroid of ofn and the plane of 4. HBs are indicated by black lines.

Scheme 5. Most Frequent Supramolecular Motifs VIII–IX Found in Crystal Structures of Carboxylic Acids and the Schematic Representation of the Crystal Structure of Uncomplexed 5 (Motif X)



Usually, weaker C–H···O or C–H···N interactions are necessary to stabilize the catemer motif.^{85–88} A prominent example is indole-2-carboxylic acid (5), which attracts increasing interest in various biological activities.^{89,90} The crystal structure of 5 exhibits parallel layers composed of catemers X stabilized by additional N–H···O HBs (Scheme 5).⁸⁹ Because the acid does not crystallize with hfb, we tried cocrystallization with pentafluorobenzoic acid (pfba) in the expectation that stacking interactions with the cyclic homodimers of pfba may induce dimeric self-assembly of 5. Obviously, the success of this experiment relied on the formation of homodimers. However, benzoic acids generally exhibit the opposite tendency and prefer creation of heterodimers by 1 kcal·mol^{−1} for a pair of aromatic carboxylic acids with substituents of different electronegativities.^{42,91} Cocrystallization of the components from CH₂Cl₂ furnished a 2:1 cocrystal, $5_2\cdot$ pfba, and fortunately, the X-ray analysis revealed that the crystals (the triclinic space group $P\bar{1}$) are composed of centrosymmetric homodimers of pfba and noncentrosymmetric dimers of 5. The dimers of 5 are essentially planar, whereas in pfba dimers, because of the steric hindrance, the carboxyl groups are twisted by ca. 43°

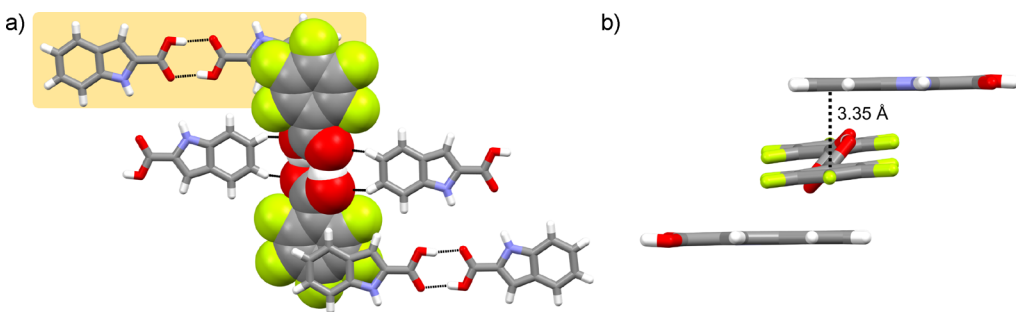


Figure 7. Crystal structure of $S_2\cdot pfba$: (a) top view of a fragment of the structure showing the relative orientation of the homodimers of **S** and **pfba** (one of the S_2 dimers is highlighted in light orange) and (b) π -stacked homodimers with the distance between the centroid of the pentafluorophenyl unit and the plane of **S** (view along the plane of a molecule of **S**). HBs are indicated by black lines.

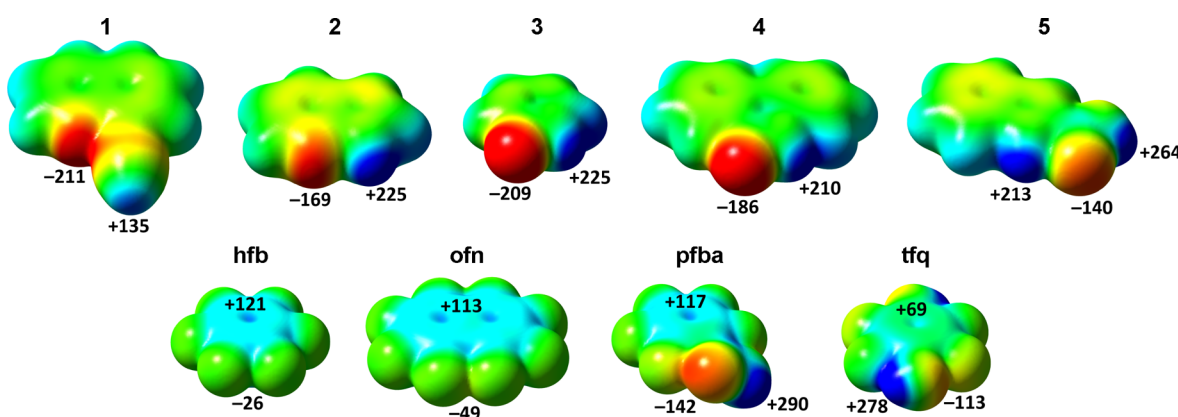


Figure 8. Electrostatic potential (ESP) surfaces of compounds **1–5** and the perfluoroaryls **hfb**, **ofn**, **pfba**, and **tfq** mapped on the molecular $\rho = 0.002$ au isosurface of electron density. Color ranges from -0.06 (red) to $+0.06$ au (blue). Selected most positive ($V_{S,\max}$) and most negative ($V_{S,\min}$) ESP values (in kJ/mol) are indicated.

from the plane of the pentafluorophenyl rings. The homodimers of these components are perpendicularly oriented in such a way that the pentafluorophenyl rings of each **pfba** dimer stack onto the phenyl rings of two different dimers of **S** and *vice versa*. They are further aggregated through $C_{Ar}-H\cdots O$ HBs between the aryl moiety of one of the symmetrically inequivalent molecules of **S** and two oxygen atoms of the **pfba** dimer (Figure 7a, Table 1). As in $3_2\cdot tfq$, the molecules of both components are stacked in an offset (DCC_n) pattern. The perfluorinated rings of the **pfba** (*D*) molecules and the molecules of **S** (*C*) are not parallel to each other; the dihedral angle between the planes of the aryl and perfluoroaryl moieties is about 6.5° . The distance between the centroid of the pentafluorophenyl ring and the plane of **S** is equal to 3.35 \AA . Furthermore, because of the aryl–perfluoroaryl interactions, the strong $N-H\cdots O$ HBs (present in the crystal structure of the uncomplexed **S**) were broken, and the $N-H$ HB donor in the cocrystal does not form any contacts with the neighboring molecules (even though it remains in close proximity to a carbonyl oxygen atom).

Electrostatic Potential Maps. The formation of hydrogen bonds and other noncovalent interactions is mainly the result of electrostatic forces operating between electron-rich and electron-deficient regions of a molecule that, in turn, are associated with the presence of a negative or positive electrostatic potential around a particular atom, respectively.⁹² Therefore, DFT calculations of the electrostatic potential (ESP) have been performed for all nine compounds investigated in this study. Figure 8 represents the ESP maps, where the red color depicts the most electron-rich regions of a

molecule (negative electrostatic potential), whereas the blue color indicates the most electron-deficient regions of a molecule (positive electrostatic potentials). The molecular regions of low electronic density and positive electrostatic potential perpendicular to the plane of the aromatic π -regions can be treated as π -holes that are responsible for intermolecular interactions with negative sites like π -electrons or lone pairs.^{29,30}

The ESP maps indicate that the sp^2 pyridine nitrogen atom is the most electron-rich region in molecules **1** and **2**. The magnitude of the local minimum of the electrostatic potential at the pyridine nitrogen atom in **1** ($V_{S,\min} = -211 \text{ kJ/mol}$) is about 40 kJ/mol lower compared to **2** (-169 kJ/mol), suggesting that the nitrogen atom in 8-ethynylquinoline is a significantly better acceptor of the hydrogen bond than the pyridine nitrogen atom in 7-azaindole. In turn, the most positive electrostatic potential region in **1** is localized at the acetylene hydrogen atom ($V_{S,\max} = +135 \text{ kJ/mol}$) and at the $N-H$ proton ($V_{S,\max} = +225 \text{ kJ/mol}$) in **2**, clearly showing the better HB donor ability of the $N-H$ fragment. Moreover, the area of the repulsive potential extends to the $C-H$ proton adjacent to the $N-H$ moiety in the pyrrole ring of **2**. The presence of such areas is strictly in line with the geometric preferences of $N-H\cdots N$ and $C-H\cdots F$ HBs in the $2_2\cdot hfb$ cocrystal. Electrostatic potential maps of lactams **3** and **4** leave no doubt that the attractive potential is concentrated on the carbonyl oxygen atom (-209 and -186 kJ/mol respectively), whereas the repulsive potential is mainly localized on the hydrogen atom of the amide moiety. Such a distribution of the potential successfully explains the presence of self-complementary

lactam dimers in **3₂**-hfb**** and **4₂**-ofn**** cocrystals. The N—H···O=C HBs that connect the lactam molecules are the result of the interaction between the regions with the minimum and maximum ESP value. It was also noted that the most intense blue color on the ESP map of **3** present at the amide hydrogen atom extended toward the adjacent hydrogen atom (C—H), predicting that it too may be involved in an electrostatic interaction. And, indeed, there are short C—H···O contacts between 2-pyridone dimers in **3₂**-hfb****. Somewhat surprisingly, on the ESP map of **4**, there is also a weak positive local maximum located between the two bay hydrogen atoms (opposite to the amide group; see Figure S2 in the ESI). This is due to the flat geometry of **4** causing the two hydrogens to overlap, which increases the local ESP. This region of the molecule is indeed involved in the C—H···F interaction in the **4₂**-ofn**** cocrystal. Similarly, the ESP map of indole-2-carboxylic acid **5** clearly establishes the presence of a negative electrostatic potential area around the carbonyl oxygen atom and two positive regions of the O—H and N—H protons. Predictably, the higher potential (about 50 kJ/mol) is located on the carboxylic hydrogen atom.

In the case of the perfluoroaryls **hfb** and **ofn** that were used as components to facilitate the dimer formation, the differences between the $V_{S,\max}$ and $V_{S,\min}$ values are not as great as those in compounds **1–5**. As expected, the electrostatic potential below and above the aromatic rings of **hfb** and **ofn**, loaded with electron-withdrawing fluorine atoms, is positive (in contrast with most aromatic systems). These centers of positive potential (π -holes) serve as partners in attractive π – π interactions between the aryl and perfluoroaryl units. The ESP on the periphery of the molecules is only slightly negative (−26 and −49 kJ/mol for **hfb** and **ofn**, respectively). The ESP values are comparable to those reported earlier.^{93,94} Two local minima of the electrostatic potential are present on the ESP map of **pfba**. As expected, the most negative value is present on the carbonyl oxygen atom. The most intense blue color (corresponding to $V_{S,\max} = 290$ kJ/mol) is present on the hydrogen atom of the carboxylic group, indicating the area most prone to HB formation (as in the **5₂**-pfba**** cocrystal). Tetrafluorohydroquinone **tfq** is the only molecule whose potential above and below the aromatic ring is not as high as those in the other perfluorinated compounds, which can be attributed to the presence of strong electron-donating groups. On the other hand, the $V_{S,\max}$ value clearly indicates that the O—H proton is a very strong hydrogen bond donor, which promotes its cocrystallization with (strong or moderate) hydrogen bond acceptors. That is why the strong O—H···O=C HBs are present in the crystal structure of **4₂**-tfq****.

CONCLUSIONS

We described a selection of organic compounds that are capable of forming self-complementary hydrogen bonds but, due to the packing forces, crystallizing as catemers or creating more complex crystal structures and then compared them with the results obtained from their complexation with perfluoroaryl compounds. In all the described cases, the π – π stacking interactions resulted in reorganization of hydrogen bonds and induced the creation of the expected self-complementary synthons. This is apparently due to the fact that most of the catemer structures are not planar, whereas self-complementary synthons expected for the compounds studied require a coplanar orientation of the contributing molecules, which is, in turn, forced by the effective face-to-face aryl–perfluoroaryl

stacking interaction between the components constituting the complex. In summary, our results show that the aryl–perfluoroaryl π – π stacking interactions show significant potential for the control of molecular self-assembly of aromatic compounds able to form hydrogen bonds. The results also demonstrate that the addition of a perfluorinated component often induces the formation of a two-component system with a layered structure. In addition, the calculated electrostatic potential surfaces correlate with the intermolecular interactions within the crystals and thus shed some light on their nature.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.cgd.3c00676>.

Details on the synthesis and characterization of the cocrystals, the crystallographic data and details of the structural refinement, and description of the crystal structure of **4** (PDF)

Accession Codes

CCDC 2263874, 2263877–2263878, 2263881, 2263886, 2263888, and 2263893–2263894 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033. Crystallographic data reported in this paper have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication no. CCDC 2263874 (1), 2263877 (3₂**-hfb**), 2263878 (5₂**-pfba**), 2263881 (4), 2263886 (4₂**-tfq**), 2263888 (1₂**-hfb**), 2263893 (4₂**-ofn**), and 2263894 (2₂**-hfb**). The data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/structures.

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Notes

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