Dynamic host–guest behavior in halogen-bonded two-dimensional molecular networks investigated by scanning tunneling microscopy at the solid/liquid interface†

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The fabrication of supramolecularly engineered two-dimensional (2D) networks using simple molecular building blocks is an effective means for studying host–guest chemistry at surfaces toward the potential application of such systems in nanoelectronics and molecular devices. In this study, halogen-bonded molecular networks were constructed by the combination of linear halogen-bond donor and acceptor ligands, and their 2D structures at the highly oriented pyrolytic graphite/1-phenyloctane interface were studied by scanning tunneling microscopy. The bi-component blend of the molecular building blocks possessing tetracycloxy chains formed a lozenge structure via halogen bonding. Upon the introduction of an appropriate guest molecule (e.g., coronene) into the system, the 2D structure transformed into a hexagonal array, and the central pore of this array was occupied by the guest molecules. Remarkably, the halogen bonding of the original structure was maintained after the introduction of the guest molecule. Thus, the halogen-bonded molecular networks are applicable for assembling guest species on the substrate without the requirement of the conventional rigid molecular building blocks with C3 symmetry.

Introduction

Supramolecular chemistry of well-ordered two-dimensional (2D) networks at surfaces is of great importance owing to the potential application of such systems in nanoelectronics, molecular devices, and lithographic techniques.1–6 The fine-tuning of the molecular organizations at surfaces has been attempted by the design and preparation of molecular building blocks with functional groups for enabling supramolecular interactions such as hydrogen bonding, metal coordination, dispersion force, and halogen bonding. The halogen bonding is a highly directional interaction of halogen atoms with other nucleophilic atoms carrying lone electron pairs, and its strength is often comparable to that of hydrogen bonding.7–10 Halogen bonds have attracted interest because they can play important roles in the molecular recognition processes, and the formation of supramolecular structures in natural as well as artificial systems.11–15

Previously, supramolecularly engineered 2D structures in physiosorbed monolayers at solid/liquid interfaces have been investigated by scanning tunneling microscopy (STM) with submolecular resolution.16–23 The formation of 2D structures at the solid/liquid interface is affected by several parameters such as the solvent type, temperature, concentration, alkyl chain length, intermolecular interactions, and electric field. By tuning these parameters, a wide variety of 2D network structures have been constructed.16–23

The 2D porous networks of supramolecular assemblies have attracted much attention because they can capture and immobilize guest molecules in their pores, and they are potentially applicable in molecular devices, nanoelectronics, sensors, catalysis, etc.24–34 There have been many remarkable studies on the host–guest chemistry at surfaces toward the potential application of such systems in nanoelectronics and molecular devices.24–34 However, in most of these studies, the 2D porous networks were created via hydrogen bonding and dispersion interactions of alkyl chains using rigid molecular building blocks with C3 symmetry.27–34

In a previous study,25 we prepared simple and linear molecular building blocks with halogen bonding acceptor and donor units (Py-Cn and FI-Cn (n = 18), respectively, in Chart 1). The binary blends of Py-C18 and FI-C18 formed hexagonal and porous networks, as observed by STM, at the highly oriented pyrolytic graphite (HOPG)/1-phenyloctane interface. This result
prompted us to apply the halogen-bonded structures to study host–guest chemistry at surfaces. To the best of our knowledge, there are few reports on the host–guest chemistry in halogen-bonded 2D molecular networks.14

In this contribution, we present the formation of halogen-bonded molecular networks, which adaptively capture the guest molecule, coronene, resulting in the transformation of the 2D structure. In this study, we prepared halogen-bond donor and acceptor ligands (Py-Cn and FI-Cn; n = 14 and 15; Chart 1) and studied the 2D structures of single and bicomponent systems at STM at the HOPG/1-phenyloctane interface. In addition, at this interface, we investigated guest inclusion into the 2D molecular networks.

**Experimental**

**Preparation and characterization of molecular building blocks**

The chemicals were purchased from Sigma-Aldrich, Kanto Chemical, Kishida Chemical, or Tokyo Chemical Industry, and they were used without further purification. Except for the change in the length of the alkyl chains, the building blocks of Py-Cn and FI-Cn (n = 14 and 15) were prepared as reported previously.25

Briefly, for the preparation of Py-Cn, the alkylation of catechol with an alkyl bromide followed by iodonation was performed to yield 1,2-bis(alkyloxy)-4-iodobenzene. Then, the Sonogashira coupling of 3-ethylnypridine with 1,2-bis(alkyloxy)-4-iodobenzene was performed in dry tetrahydrofuran with a Pd[PPh3]4 catalyst, Cul co-catalyst, and triethylamine. The flask was cooled, degassed, charged with N2 gas, and stirred at 60 °C for 7 days. The crude product of Py-Cn was purified by silica gel column chromatography with CHCl3 as the eluent.

For the preparation of FI-Cn, methyl 3,5-dihydroxybenzoate was alkylated, and the resultant compound was reduced with LiAlH4 to obtain 3,5-bis(alkyloxy)benzylalcohol. Then, 3,5-bis(alkyloxy)benzylalcohol in a small amount of dry N,N-dimethylformamide was reacted with pentafluoriodobenzene in the presence of Cs2CO3 for 15 h. The crude product of FI-Cn was purified by silica gel column chromatography using CHCl3/n-hexane (1 : 1) as the eluent.

**Chart 1** Chemical structures of the molecular building blocks used in this study: pyridine (Py-Cn), tetrafluoriodobenzene-based molecules (FI-Cn), and coronene (COR) (n = 14 or 15).

The prepared compounds were characterized by 1H and 13C NMR spectroscopy (Bruker Avance 400 or 500 NMR spectrometer) using tetramethylsilane as the internal standard and Fourier-transform infrared spectroscopy (FTIR; JASCO FT/IR 420) by the KBr pellet method. The PhCF3 signal at −62.61 ppm (ref. 36) was used as the internal standard for 19F NMR measurements. Matrix-assisted laser desorption/ ionization high-resolution mass spectrometry (MALDI-HRMS) was conducted on a JMS-S3000 spiralTOF mass spectrometer (JEOL, Tokyo, Japan). Trans-2-[3-(4-tert-Butylphenyl)-2-methyl-2-propenylidene]malononitrile was used as the matrix and silver trifluoroacetate was used as the cationizing agent. The prepared compounds were mixed with the matrix and the cationizing agent in a solvent-free sample preparation (the solid chemicals were ground using a mortar and pestle for a few minutes) and the powdery mixture with an approximate matrix : cation : sample molar ratio of 50 : 1 : 1 was pressed onto a MALDI target using a spatula to form a thin film.

**Py-C14.** 1H NMR (400 MHz, CDCl3): δ 0.88 (t, J = 6.74 Hz, 6H), 1.26 (br, 40H), 1.46 (m, 4H), 1.83 (m, 4H), 4.02 (t, J = 4.98 Hz, 4H), 6.84 (d, J = 8.32 Hz, 1H), 7.04 (d, J = 1.88 Hz, 1H), 7.11 (dd, J1 = 8.28 Hz, J2 = 1.92 Hz, 1H), 7.28 (m, overlapped with CHCl3, 1H), 7.79 (d, J = 7.88 Hz, 1H), 8.54 (s, 1H), 8.76 (s, 1H). 13C NMR (100 MHz, CDCl3): δ 14.1, 22.7, 26.0, 29.18, 29.22, 29.23, 29.39, 29.43, 29.65, 29.69, 29.70, 32.0, 69.1, 69.3, 84.4, 93.1, 113.2, 114.5, 116.6, 120.9, 123.0, 125.2, 138.2, 148.2, 148.8, 150.2, 152.2. FTIR (KBr): 2918, 2849, 1514, 1470, 1254, 1220, 1125 cm−1. MALDI-HRMS: theoretical m/z for silverated C41H63F4IO3 (isotopes 107Ag/109Ag): 710.4061/712.4065; measured: 710.4065/712.4066.

**Py-C15.** 1H NMR (500 MHz, CDCl3): δ 0.88 (t, J = 6.90 Hz, 6H), 1.26 (br, 44H), 1.46 (m, 4H), 1.82 (m, 4H), 4.01 (t, J = 6.60 Hz, 4H), 6.84 (d, J = 8.30 Hz, 1H), 7.04 (d, J = 1.85 Hz, 1H), 7.11 (dd, J1 = 8.25 Hz, J2 = 1.90 Hz, 1H), 7.28 (m, overlapped with CHCl3, 1H), 7.78 (d, J = 7.85 Hz, 1H), 8.53 (d, J = 4.85 Hz, 1H), 8.75 (s, 1H). 13C NMR (125 MHz, CDCl3): δ 14.1, 22.7, 26.0, 29.18, 29.22, 29.38, 29.42, 29.44, 29.68, 29.72, 31.9, 69.1, 69.3, 84.4, 93.1, 113.2, 114.5, 116.6, 120.8, 123.0, 125.2, 138.2, 148.2, 148.8, 150.2, 152.2. FTIR (KBr): 2918, 2849, 1514, 1464, 1254, 1221, 1126 cm−1. MALDI-HRMS: theoretical m/z for silverated C41H65NO2 (isotopes 107Ag/109Ag): 738.4374/740.4378; measured: 738.4376/740.4339.

**FI-C14.** 1H NMR (500 MHz, CDCl3): δ 0.88 (t, J = 6.98 Hz, 6H), 1.26 (br, 40H), 1.43 (m, 4H), 1.76 (m, 4H), 3.92 (t, J = 6.58 Hz, 4H), 5.18 (s, 1H), 6.41 (dd, J = 2.25, 2.25 Hz, 1H), 6.53 (d, J = 2.20 Hz, 2H). 13C NMR (100 MHz, CDCl3): δ 14.2, 22.8, 26.1, 29.26, 29.44, 29.66, 29.68, 29.74, 29.76, 32.0, 64.2 (t, J(C,F) = 28 Hz), 68.1, 76.3, 101.8, 106.3, 106.8, 137.3, 140.9 (d, J(C,F) = 249 Hz), 147.3 (d, J(C,F) = 240 Hz), 160.5. 19F NMR (376 MHz, CDCl3): δ −121.18 to −121.28 (m, 2F), −153.06 to −153.15 (m, 2F). FTIR (KBr): 2920, 2852, 1610, 1480, 1397, 1319, 1159, 1093 cm−1. MALDI-HRMS: theoretical m/z for silverated C41H63F4O3 (isotopes 107Ag/109Ag): 913.2804/915.2807; measured: 913.2809/915.2809.

**FI-C15.** 1H NMR (500 MHz, CDCl3): δ 0.88 (t, J = 6.98 Hz, 6H), 1.26 (br, 44H), 1.43 (m, 4H), 1.76 (m, 4H), 3.92 (t, J = 6.58 Hz, 4H), 5.18 (s, 2H), 6.41 (dd, J = 2.25, 2.25 Hz, 1H), 6.53 (d, J = 2.20 Hz, 2H). 13C NMR (100 MHz, CDCl3): δ 14.1, 22.7, 26.0,
29.23, 29.41, 29.43, 29.62, 29.64, 29.71, 32.0, 64.2 (t, J(C,F) = 28 Hz), 68.1, 76.3, 101.8, 106.3, 106.8, 137.3, 140.9 (d, J(C,F) = 249 Hz), 147.3 (d, J(C,F) = 242 Hz), 160.5. 19F NMR (376 MHz, CDCl3): δ = −121.18 to −121.28 (m, 2F), −153.05 to −153.14 (m, 2F). IR (KBr): 2919, 2850, 1609, 1479, 1394, 1320, 1159, 1097 cm⁻¹. MALDI-HRMS: theoretical m/z for silverated \( \text{C}_{43}\text{H}_{67}\text{F}_{4}\text{I} \) 941.3117/943.2121; measured: 941.3109/943.2121.

**STM observations**

STM was performed with a Nanoscope IIIa system (Digital Instruments). The compounds were dissolved in 1-phenyloctane at different concentrations, and the solutions were deposited on freshly cleaved HOPG substrates (ZYB grade, Momentive Performance Materials Quartz, Inc.) to obtain physisorbed monolayers. A Pt-Ir wire (90/10; 0.25 mm φ) was mechanically cut to prepare STM tips, and STM was carried out at the 1-phenyloctane/HOPG interface. The captured STM images were calibrated and corrected with reference to the HOPG lattice underneath the physisorbed monolayer using the SPIP software.

The solutions of Py-CN and FI-CN (n = 14 and 15) in 1-phenyloctane were prepared, and STM was carried out before and after adding them at a molar ratio of 1 : 1. The guest, corone (COR), was introduced with the coadsorption of molecular building blocks: a saturated 1-phenyloctane solution of COR was mixed with that of the bi-component blend (Py-CN/FI-CN) at a 1 : 1 ratio, and the relative ratio between COR and Py-CN/FI-CN was tuned. The pre-mixed solution of Py-CN/FI-CN/COR deposited on the HOPG surface was stored for 15 h before performing STM on the host–guest system.

**DFT calculations**

The Gaussian 16 program was employed for density functional theory (DFT) calculations using the 6-311G** basis set. The DGDZVP basis set was used for iodides. The B3LYP functional was used with Grimme’s D3 dispersion correction. The geometry of the complex was optimized under imposing \( C_{2v} \) symmetry. BSSE was corrected by the counterpoise method in the calculations of the intermolecular interaction energies.

**Results and discussion**

**Single component system**

The 2D structures of the single-component system at the HOPG/1-phenyloctane interface were studied by STM. Fig. 1A and S1A (ESI†) show the 2D structure of Py-C14. As shown in Fig. 1A, the π-conjugated moieties (bright contrast in the image) are oriented in a face-to-face manner to form a double-columnar structure. The bright column is periodically separated by the dark region with a width of \( L_1 = 1.77 ± 0.10 \) nm, which is identical to the length of the tetradeoxy chain (1.77 nm). The molecular model in Fig. 1C is proposed based on the STM image in Fig. 1A.

In contrast, FI-C14 showed a single-lined columnar structure (Fig. 1B and S1B (ESI†)), where the 3,5-tetradeoxy chains with

**Bi-component system**

Next, the binary blend of Py-C14 and FI-C14 was studied, expecting the formation of a halogen-bonded 2D molecular network. Fig. 2A and B show the STM images of a binary blend of Py-C14/FI-C14 (1 : 1 ratio) at the HOPG/1-phenyloctane interface. Bright spots with cross-shaped assemblies are periodically arranged to form a lozenge-shaped lattice, and the

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L_2 = 1.75 ± 0.09 \text{ nm and } L_3 = 1.78 ± 0.10 \text{ nm exist in an expanded conformation (with an angle of } 121 ± 2^\circ \text{) to follow the HOPG lattice with a herringbone structure; the direction of the iodine atom alternated in neighboring columns, as shown in the proposed molecular model in Fig. 1D. The 2D molecular arrangements in Py-C14 and FI-C14 are almost identical to those of the previously reported Py-C18 and FI-C18 building blocks, respectively, except for the column intervals. A higher concentration of Py-C14 and FI-C14 in 1-phenyloctane was necessary for stable STM investigations as compared to that used for studying Py-C18 and FI-C18. A possible reason for this is that the molecules with shorter alkyl chains (n = 14) have less intermolecular and molecule–substrate interactions compared to those with longer chains (n = 18). The lattice constants of the 2D structures of Py-C14 and FI-C14 are summarized in Table 1.

### Table 1

| Compounds | a (nm) ± std | b (nm) ± std | γ (°) ± std |
|-----------|-------------|-------------|-------------|
| Py-C14    | 1.41 ± 0.02 | 3.43 ± 0.04 | 89 ± 1      |
| FI-C14    | 1.41 ± 0.11 | 4.37 ± 0.06 | 88 ± 1      |
Fig. 2 (A, B) STM images of the bi-component blends of Py-C14/FI-C14 (25 mM) physisorbed at the HOPG/1-phenyloctane interface. A set of arrows indicates the HOPG lattice directions. (C) Proposed molecular model of the lozenge structure is depicted based on the STM image in (B). The building blocks in a part of the repeating units for Py-C14 and FI-C14 are highlighted in green and blue, respectively in (B) and (C). The lengths of \( L_x, L_y, \) and \( L_z \) were measured to be 1.76 ± 0.08 nm; they are almost identical to that of the tetradecyloxy chain (1.77 nm). (D) Optimized structures of the complex for the head groups of arrows indicates the HOPG lattice directions. The lengths of \( Py-C14 \) and \( FI-C14 \) in (1.77 nm). (D) Optimized structures of the complex for the head groups Py-C14 (pyridine) and FI-C14 (2,3,5,6-tetrafluoro-1-iodo-4-methoxybenzene) obtained by DFT calculation. I–N halogen bonding and Ar–F···H–Py interactions are represented as red and cyan dotted lines, respectively. Tunneling conditions: (A) \( I = 25 \) pA, \( V = -805 \) mV; (B) \( I = 25 \) pA, \( V = -813 \) mV.

**Effect of the alkyl chain length on the 2D structure formation in the bi-component blend.**

In our previous study, we observed the formation of halogen-bonded hexagonal networks in a binary blend of Py-C18/FI-C18. In this context, we examined the effect of the alkyl chain length on the 2D structure formation in the bi-component blend. Fig. 3A and B show the STM images of Py-C15/FI-C15 at the HOPG/1-phenyloctane interface. Within the concentration range of 10–25 mM, Py-C15/FI-C15 dominantly formed hexagonal arrays throughout the physisorbed monolayer, although there were significant deviations from those of the single component systems (Py-C14 and FI-C14).

The total interaction energy of the 4 head groups (Fig. 2D) was calculated to be \(-20.2 \) kcal mol\(^{-1}\) (Fig. S3, ESI†), indicating that the complex formation even at the head groups contributes significantly to the stabilization of the 2D structure. In addition, alkyl chain moieties assisted the 2D arrangements in the lozenge structure: one of the two alkyl chains derived from FI-C14 in the complex was aligned along the HOPG lattice, accompanied by two alkyl chains of Py-C14.

Even when the Py-C14 : FI-C14 blend ratio was changed between 1 : 3 and 3 : 1, the 2D structure remained unchanged and showed the same lozenge-shaped lattice (data not shown), suggesting that the lozenge structure is thermodynamically favored in the binary blend of Py-C14/FI-C14. Note that the lozenge structure was observed within the concentration range of 10–25 mM, and no change in the 2D structure was found even after 15 h of storage at room temperature (25 °C).

**Table 2** Lattice constants of Py-CN/FI-CN \((n = 14 \) and \( 15) \) and Py-C14/FI-C14/COR measured from the STM images in Fig. 2–5.

|          | \( a \) (nm)  | \( b \) (nm)  | \( \gamma \) (°) |
|----------|---------------|---------------|-----------------|
| Py-C14/FI-C14 | 3.16 ± 0.07   | 3.73 ± 0.07   | 70 ± 3          |
| Py-C15/FI-C15 | 6.93 ± 0.09   | 6.93 ± 0.09   | 60 ± 1          |
| Py-C14/FI-C14/COR \( a \) | 6.69 ± 0.03   | 6.69 ± 0.03   | 61 ± 2          |
| 25 mM      | 6.67 ± 0.06   | 6.67 ± 0.06   | 60 ± 1          |
| 17 mM      | 6.67 ± 0.06   | 6.67 ± 0.06   | 60 ± 1          |

\( a \) The solution of Py-C14/FI-C14 at the concentration noted in the table was mixed with a saturated COR solution at 1 : 1 ratio.

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the individual components (Py-C15 and FI-C15) displayed linear structures (Fig. S4, ESI†). The triangular helical assembly due to the I···N halogen bonding and Ar–F···H–Py interactions (Fig. 3C) allowed the formation of a honeycomb structure with hexagonal pores. The lattice constants of the honeycomb structure in Py-C15/FI-C15 are listed in Table 2. The results in Fig. 2 and 3 suggest that there is a specific alkyl chain length at which the blend forms either a honeycomb or lozenge structure: hexagonal arrays are constructed by the building blocks of Py-CN/FI-CN (n = 15 and 18), whereas a lozenge structure is formed by those of Py-C14/FI-C14.

We propose that the alkyl chain length-dependence of the 2D structure formation can be explained in terms of both solvent co-adsorption and packing density. The pores of the hexagonal arrays cannot be completely vacant, and they can be dynamically filled with solvent molecules that aid the stabilization of the pore structure.14,24 The longer the alkyl chain length is, the larger the pore size is. This implies that building blocks with long alkyl chains are more advantageous compared to those with short chains in terms of the number of solvent molecules that can be co-adsorbed. When the alkyl chain is short, a sufficient number of solvent molecules cannot be introduced into the pore to maintain the honeycomb structure.

In general, building blocks physisorbed on the surface tend to cover the entire surface and increase the packing density to a large extent, in order to maximize the intermolecular interactions. The surface coverage calculated from the number of building blocks and lattice constant indicates that the lozenge structure (0.356 molecule per nm²) has ca. 16% larger packing density than that of the hexagonal structure (0.307 molecule per nm²), if we assume that Py-C14/FI-C14 forms a hexagonal structure.26 Taking the solvent co-adsorption and packing density into account, it can be concluded that the lozenge structure is energetically favored in comparison to the porous hexagonal structure in the case of Py-C14/FI-C14.

Host–guest co-adsorption

Although Py-C14/FI-C14 formed a lozenge structure without pores, we studied the host–guest behavior of the 2D networks. COR was selected as a flat guest molecule. Fig. 4A–C and S5 (ESI†) show the STM images of the Py-C14/FI-C14/COR mixture physisorbed at the HOPG/1-phenyloctane interface. Interestingly, hexagonal arrays were formed throughout the monolayer, where the lengths of L₉ and L₁₀ were measured to be 1.77 ± 0.07 nm. The lattice constants of the hexagonal structure are listed in Table 2. In some pores, close-packed COR with seven individual molecules was found with clear contrast, as indicated by the arrows in Fig. 4A, whereas the other pores showed fuzzy disk-like spots, suggesting the dynamic adsorption and mobility of COR in the pore.24–26 Note that STM allowed the capture of snapshots indicating the dynamic behavior of guest inclusion. Therefore, variable numbers of COR were observed in the pore, that changed during the STM imaging (Fig. S6 and S7, ESI†). According to the STM image in Fig. 4B, a molecular model with the maximum number of adsorbed COR molecules is proposed, as shown in Fig. 4D. The I···N halogen bonding and Ar–F···H–Py interactions resulted in triangular assemblies (see Fig. 3C), which organized into hexagonal arrays through the interdigitation of alkyl chains. Remarkably, the I···N halogen bonding was found in both 2D structures before (Fig. 2) and after guest inclusion (Fig. 4), that is, the halogen bonding remained substantially unchanged. Therefore, the arrangements of the halogen-bonded supramolecular complex and the alkyl chains in Py-C14/FI-C14 adaptively changed to form hexagonal arrays to accommodate the guest molecules in the 2D structure. Although there are numerous reports on 2D structural change accompanying guest inclusion in molecular assemblies composed of C₅-symmetric rigid molecules,24–26 the present study demonstrates the dynamic host–guest chemistry in halogen-bonded 2D molecular networks formed with linear molecular building blocks.

We further studied the effect of the blend ratio on the guest inclusion. Fig. 5 shows the STM images of Py-C14/FI-C14/COR obtained by tuning the relative ratio of Py-C14/FI-C14 and COR. When the concentration of the 25 mM solution of Py-C14/FI-C14 was decreased to 17 mM and then mixed with the saturated COR solution, the hexagonal structure and its lattice constants remained unchanged (Table 2). However, a larger number of the hexagonal pores were filled with close-packed COR molecules (seven in total) (Fig. 5A–C) than that observed in Fig. 4. This result suggests that increasing the guest concentration relative to that of the host network allows more COR molecules to reside in the pores without the alteration of the hexagonal structure.

Considering that the pore is surrounded by 1.77 nm long tetradecyloxy chains, the diagonal length of the hexagonal pore (pore size) could be 1.77 × 2 = 3.44 nm. The reported lattice constants of the close-packed structure of COR on HOPG are a =
possibly because of the solvent co-adsorption and the molecular system, the 2D structure of \( \text{Py-C14/FI-C14} \) is hexagonal. Once the guest molecule, \( \text{COR} \), is introduced into the halogen-bonded molecular networks composed of simple, linear building blocks. This work could inspire the controlled organization of functional molecules for constructing multi-component 2D nanoarchitectures via supramolecular chemistry at surfaces.

Conflicts of interest

There are no conflicts to declare.

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