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# Self-organization of supramolecular helical dendrimers into complex electronic materials

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The discovery of electrically conducting organic crystals<sup>1</sup> and polymers<sup>1-4</sup> has widened the range of potential optoelectronic materials<sup>5-9</sup>, provided these exhibit sufficiently high charge carrier mobilities<sup>6-10</sup> and are easy to make and process. Organic single crystals have high charge carrier mobilities but are usually impractical<sup>11</sup>, whereas polymers have good processability but low mobilities<sup>1,12</sup>. Liquid crystals exhibit mobilities approaching those of single crystals and are suitable for applications<sup>13-18</sup> but demanding fabrication and processing methods limit their use. Here we show that the self-assembly of fluorinated tapered dendrons can drive the formation of supramolecular liquid crystals with promising optoelectronic properties from a wide range of organic materials. We find that attaching conducting organic donor or acceptor groups to the apex of the dendrons leads to supramolecular nanometre-scale columns that contain in their cores  $\pi$ -stacks of donors, acceptors or donor-acceptor complexes exhibiting high charge carrier mobilities. When we use functionalized dendrons and amorphous polymers carrying compatible side groups, these co-assemble so that the polymer is incorporated in the centre of the columns through donoracceptor interactions and exhibits enhanced charge carrier mobilities. We anticipate that this simple and versatile strategy for producing conductive  $\pi$ -stacks of aromatic groups, surrounded by helical dendrons, will lead to a new class of supramolecular materials suitable for electronic and optoelectronic applications.

We have elaborated a library based on a semifluorinated tapered dendron<sup>19,20</sup> that was functionalized at its apex with a diversity of electroactive donor (D) and acceptor (A) groups, (**3,4,5**)12F8G1-D/A, (Figs 1 and 2). The resulting functional dendrons self-assemble into columns containing the optoelectronic element in their cores. For synthesis and structural analysis see the Supplementary Infor-

mation. The supramolecular columns self-organize into homeotropically aligned liquid crystals with hexagonal columnar  $(\Phi_h)$ , centred and simple rectangular columnar  $(\Phi_{r-c} \text{ and } \Phi_{r-s})$  morphologies (Fig. 2). We name this desirable homeotropic selforganization between electrodes 'self-processing'.

Carbazole (D1), naphthalene (D2) and pyrene (D3 and D4) derivatives have been introduced at the apex as D groups and 4,5,7-trinitrofluorenone-2-carboxylic acid (TNF) as an A group (A1) (Fig. 1). We use a diethylene glycol or tetraethylene glycol spacer between the dendron and the D or A groups to decouple their motion and facilitate fast self-assembly. The approach is simple and allows for diverse synthetic methods, and ensures that the self-assembled D and A groups are protected from moisture by the internal compartmentalization of the column and external jacketing with a fluorinated coat (Fig. 2).

Co-assembling a D-dendron with an A-dendron incorporates an electron donor–acceptor (EDA) complex in the centre of the column (see A1 + D1). Disordered amorphous polymers with A and D side groups form EDA complexes when the D-dendron (D1) is mixed with the A-polymer (AP1) and when the A-dendron (A1) is mixed with a D-polymer (DP1 or DP2). The EDA complexes assemble into columns, which in turn self-organize into  $\Phi_h$  or  $\Phi_r$  liquid crystals. The repulsion between the dendron's fluorine coat and the aromatic groups is so strong that the complexed polymer is forced to reside in the centre of the column (left and right columns in Fig. 2). EDA interactions<sup>21</sup> rather than the previously reported covalent bonding<sup>22</sup> are generating a supramolecular polymer<sup>23</sup> with dendritic side groups leads to a more ordered polymer backbone.

The self-assembly and self-organization of the material into a  $\Phi_h$ (D1, D2, D3, D4, D1 + A1, A1 + DP1, A1 + DP2),  $\Phi_{r-s}$ (D1 + AP1) and  $\Phi_{r-c}$  (A1) liquid crystals was demonstrated by a combination of techniques including differential scanning calorimetry (DSC), X-ray diffraction (XRD) (Fig. 3a), thermal optical polarized microscopy (TOPM) (Fig. 3b, c) and electron diffraction (ED) (Supplementary Information). Calculations based on density and XRD measurements indicate that between 3.8 and 4.6 dendrons self-assemble into a 4.8 Å stratum of the column (Supplementary Information). These self-processed systems have a column density of  $4.5 \times 10^{12}$  to  $5.8 \times 10^{12}$  columns per square centimetre (Supplementary Information), which exceeds the active element density achieved in other systems<sup>10</sup> by two orders of magnitude.

Figure 1 summarizes the structures of the supramolecular assemblies and their electron mobilities ( $\mu_e$ ) and hole mobilities ( $\mu_h$ ). The field- and temperature-independent  $\mu_e$  and  $\mu_h$  values were determined by the time-of-flight (TOF) method (Supplementary Information). Tapered D-dendrons lead to  $\mu_h$  ranging from  $10^{-4}$  to  $10^{-3}$  cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup> in the liquid crystal state whereas the A-dendron (A1) has an  $\mu_e$  of  $10^{-3}$  cm<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>. The mobilities  $\mu$  of the EDA polymer complexes are in the same range. All these values are two to five orders of magnitude higher than those of the related D and A compounds in the amorphous state (Fig. 1), reflecting the  $\pi$ -stacked one-dimensional ordered structure of the D, A or EDA complexes in the centre of the supramolecular column. The charge migration through the columns is not well understood but may involve a polaron hopping process<sup>16,24</sup>. XRD and <sup>1</sup>H-nuclear magnetic resonance (NMR) studies carried out in solution and bulk by using previously developed techniques<sup>25</sup> demonstrate  $\pi$ - $\pi$ -interactions between the D or A groups. The  $\mu$  value of these very simple D-dendrons are similar to those of more complex discotic liquid crystals<sup>13-16</sup> produced from disc-like molecules; reports of discotic liquid crystals exhibiting higher  $\mu$  values<sup>13,15,26,27</sup> refer either to measurements on the  $\Phi_{ho}$  crystal phase, or to measurements of the one-dimensional intracolumnar mobility by contactless pulseradiolysis time-resolved microwave conductivity (PR-TRMC)<sup>13,15,26,27</sup>. TOF data are inherently lower owing to structural imperfections within the inter-electrode gap, but are more relevant

for technical applications than the higher values obtained by PR-TRMC.

Cooling the liquid crystal state leads to crystallization in the case of **D2**, whereas the two-dimensional (2D) order of **D1**, **D3** and **A1** is enhanced in the glassy state showing helical short-range order. For example, the XRD of a fibre of A1 in the glassy ordered state cooled from the liquid crystal phase exhibits, in the small-angle region, the pattern of the  $\Phi_h$  order (Fig. 3a). This demonstrates that the liquid crystal order of the large and uniform homeotropic domain (100  $\mu$ m to a few centimetres in size) obtained by slow cooling on

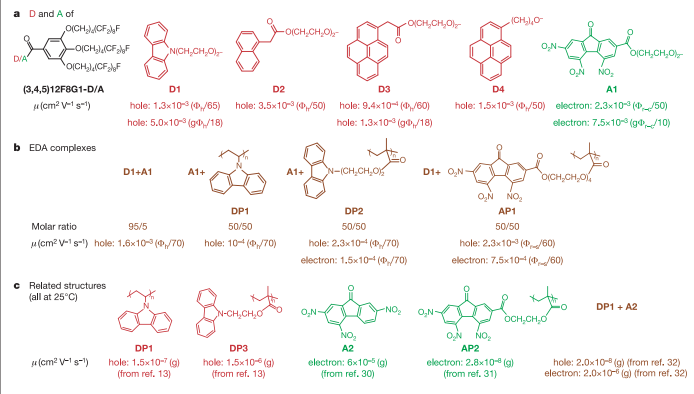


Figure 1 Charge carrier mobility of supramolecular columns and related systems. **a**, Donor (**D1–4**) and acceptor (**A1**) groups of (**3,4,5**) **12F8G1-D/A**. **b**, EDA complexes. **P**, polymer. **c**, Related structures (all at 25 °C). The charge carrier mobility values ( $\mu$ ) are shown for all structures. ( $\Phi_h/65$ ), in hexagonal columnar phase at 65 °C. ( $\Phi_{r,c}/50$ ), in centred rectangular phase at 50 °C. (g $\Phi_h$ /18), in glassy hexagonal columnar phase at 18 °C. (g $\Phi_{r-c}$ /10), in glassy centred rectangular phase at 10 °C. ( $\Phi_{r-s}$ /60), in simple rectangular phase at 60 °C. (g), glassy.

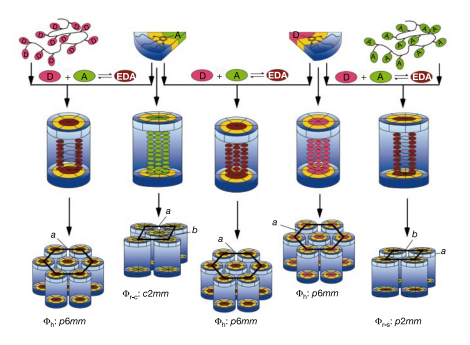
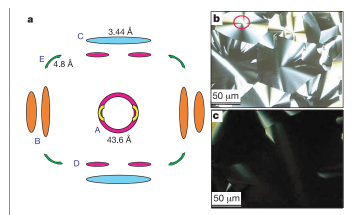


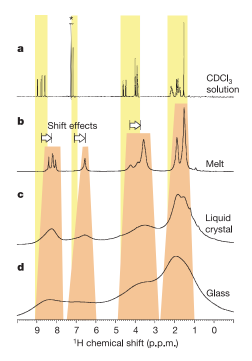
Figure 2 Schematic illustration of the liquid crystal assembly processes. Shown are the self-assembly, co-assembly and self-organization of dendrons containing donor (D) and acceptor (A) groups with each other and with disordered amorphous polymers containing

D and A side groups. The different systems form hexagonal columnar ( $\Phi_{r,b}$ ), centred rectangular columnar ( $\Phi_{r-c}$ ) and simple rectangular columnar ( $\Phi_{r-s}$ ) liquid crystals. *a* and *b* are lattice dimensions; space groups are shown.



**Figure 3** Structural characterization of self-assembled liquid crystals. **a**, Characteristic small- and wide-angle XRD of **A1**. The individual XRD features are identified as (A) supramolecular column to column distance, short-range, (B) order in tail region within a supramolecular cylinder, (C)  $\pi$ -stacks within the column, (D) correlation along a direction at an angle to the supramolecular column axis, (E) helical correlation. **b**, Thermal optical polarized micrograph (TOPM) of **D4** in the  $\Phi_h$  liquid crystal state, obtained by cooling at 20 °C min<sup>-1</sup> from the isotropic phase on a hydrophobic indium—tin oxide coated substrate showing defecting alignment. The circle indicates a highly tilted defect characteristic of the  $\Phi_h$  phase. **c**, TOPM of a homeotropically aligned **D4**, obtained by cooling at 0.1 °C min<sup>-1</sup> on the same substrate. Hydrophilic substrates such as plasma-treated carbon induce planar orientation of the columns.

a hydrophobic substrate from the isotropic state (Fig. 3b, c and Figure S3 of the Supplementary Information) is frozen into a glassy  $\Phi_{\rm h}$  state. In addition, at wide angles the XRD pattern reveals diffuse spots arranged in an X-like fashion that indicates short-range helical correlation along the column axis with an average repeat distance of 4.8  $\pm$  0.3 Å (E, Fig. 3a). This value arises from the packing of the aromatic regions of the dendron near the aliphatic tails. This XRD also shows diffuse spots assigned to disordered  $\pi$ -stacks of TNF units in the core of the column with an average separation of 3.44 Å



**Figure 4** <sup>1</sup>H NMR spectra of **A1**. **a**, In CDCl<sub>3</sub> at 22 °C; **b**, in the isotropic melt at 120 °C; **c**, in the  $\Phi_h$  liquid crystal state at 75 °C; and **d**, in the  $\Phi_h$  glassy state at 25 °C. Spectra **b–d** were recorded under fast magic-angle spinning. The high-field shifts caused by adjacent  $\pi$ -electron systems are indicated by arrows.

(C, Fig. 3a). Consequently,  $\mu$  of this glassy liquid crystal state increases since the motion of the D groups from the core of the supramolecular column is decreased leading to reduced dynamic disorder (Fig. 1). Furthermore, the motion of the ionic impurities is frozen in the ordered state, whereas the photo-generated charges continue to migrate within the system. Therefore, the glassy ordered state is insensitive to ionic impurities.

Figure 4 depicts <sup>1</sup>H NMR spectra of A1 in CDCl<sub>3</sub> solution at 22 °C, in the isotropic melt at 120 °C, in the  $\Phi_{\rm h}$  liquid crystal state at 75 °C and in the  $\Phi_{\rm h}$  glassy state at 25 °C. In this sequence, the loss of molecular mobility is reflected by the increase of the line widths. As fast magic-angle spinning is applied (except for the solution), four groups of <sup>1</sup>H resonances can be distinguished in all spectra: aromatic fluorenone (9-8 p.p.m.), dendron phenyl (7-6 p.p.m.), OCH<sub>2</sub> (4.5-3 p.p.m.) and alkyl (2.5-1 p.p.m.). In the glass, the liquid crystal phase and the melt, the lines are shifted to high field by about 0.5–0.7 p.p.m. relative to the solution-state values except for the alkyl resonances. This effect is due to aromatic  $\pi$ -electrons situated above or below the respective protons<sup>25</sup> and, thus, provides direct evidence for  $\pi - \pi$  packing of the fluorenones and the dendron phenyls. Analogous effects are observed in the <sup>1</sup>H spectra of the other materials discussed here. Moreover, in A1 a distance of 3.5 Å was determined between the protons in adjacent fluorenone moieties by <sup>1</sup>H-<sup>1</sup>H double-quantum NMR spectroscopy<sup>25</sup>, which leads to the assumption of a sandwich-type packing of pairs of nitro-fluorenone groups (Fig. 5a). Combining the NMR information with the XRD data, it can be concluded that, in the glassy  $\Phi_{\rm h}$  phase, the columns adopt a supramolecular structure of the form

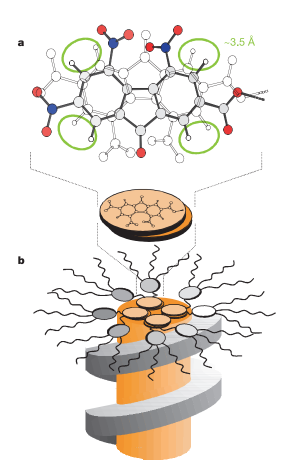


Figure 5 The molecular arrangement in the supramolecular column self-assembled from A1. a, Sandwich-type stacking of the nitro-fluorenone moieties with proton—proton distances of 3.5 Å, as determined by <sup>1</sup>H-<sup>1</sup>H double-quantum NMR spectroscopy.
b, Structure of the supramolecular columns with stacks of fluorenone sandwiches in the centre of the columns, jacketed by helical dendrons.

schematically depicted in Fig. 5b. In the centre, the fluorenone sandwiches are stacked in a column surrounded by dendrons whose phenyl groups are arranged in a helical fashion around the central column. Furthermore, NMR data indicate separation of the fluorenones (centre), dendron phenyls (inner ring) and alkyl chains (outer ring) from each other, because there are no proton–proton distances detectable between the different units on a length scale of <4 Å. This clear separation is not completely achieved when the material is precipitated from solution, but only when the system is allowed to self-organize during slow cooling from the melt into the liquid crystal and glassy  $\Phi_h$  phases. Before this cooling and 'self-repair' process, the supramolecular column contains defects introduced by intramolecular backfolded A1 dendrons.

The supramolecular arrangement shown in Fig. 5b resembles the base-pairing in DNA, which is known to support charge migration<sup>28</sup>, and points to a new structural framework for incorporating dendritic molecules into functional materials<sup>29</sup>.  $\Box$ 

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#### **Competing interests statement**

The authors declare that they have no competing financial interests.

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# Copepod hatching success in marine ecosystems with high diatom concentrations

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Diatoms dominate spring bloom phytoplankton assemblages in temperate waters and coastal upwelling regions of the global ocean. Copepods usually dominate the zooplankton in these regions and are the prey of many larval fish species. Recent laboratory studies suggest that diatoms may have a deleterious effect on the success of copepod egg hatching<sup>1-4</sup>. These findings challenge the classical view of marine food-web energy flow from diatoms to fish by means of copepods<sup>5-7</sup>. Egg mortality is an important factor in copepod population dynamics<sup>8</sup>, thus, if diatoms have a deleterious *in situ* effect, paradoxically, high

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erratum

## Self-organization of supramolecular helical dendrimers into complex electronic materials

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In this Letter, the volume numbers appeared incorrectly in the footer of pages 384, 385 and 386 as 'VOL 417'. It should have read 'VOL 419'.  $\hfill \Box$