Conjugated Polycycles

Heptagon-Embedded Pentacene: Synthesis, Structures, and Thin-Film Transistors of Dibenzo[*d*,*d*']benzo[1,2-*a*:4,5-*a*']dicycloheptenes**

Xuejin Yang, Danqing Liu, and Qian Miao*

Abstract: This study presents a new class of conjugated polycyclic molecules that contain seven-membered rings, detailing their synthesis, crystal structures and semiconductor properties. These molecules have a nearly flat C_6 - C_7 - C_6 - C_7 - C_6 polycyclic framework with a p-quinodimethane core. With field-effect mobilities of up to 0.76 cm²V⁻¹s⁻¹ as measured from solution-processed thin-film transistors, these molecules are alternatives to the well-studied pentacene analogues for applications in organic electronic devices.

Conjugated polycyclic hydrocarbons containing non-hexagonal carbocycles^[1] have received considerable attention in organic chemistry and materials science because of their novel properties and potential applications. The most widely studied subgroup of these conjugated polycyclic hydrocarbons contains five-membered rings. This group of molecules include curved π -conjugated molecules known as aromatic bowls^[2,3] (or buckybowls^[4]) and electron acceptors^[5] as applied in organic electronics, since the five-membered rings can not only introduce positive curvature but also stabilize injected electrons by forming the aromatic cyclopentadienyl anion. Unlike five-membered rings, seven-membered rings can give rise to saddle-shaped π -molecules with negative curvature^[6] and stabilize holes injected into p-type organic semiconductors by forming the aromatic cycloheptatrienyl cation.^[7] However, conjugated polycyclic hydrocarbons containing seven-membered rings have been rarely explored. Herein, we report dibenzo [d,d'] benzo [1,2-a:4,5a']dicycloheptene derivatives (**1a–c**, Figure 1),^[8] which have an unprecedented pentacyclic conjugated backbone containing two seven-membered carbocycles.^[9] Unlike the recently reported indenofluorenes (**3a**,**b** and **4**, Figure 1),^[10] **1a**–**c** have seven-membered rings rather than five-membered rings replacing two benzene rings in pentacene, a benchmark

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Figure 1. Structures of ethynylated dibenzo[*d*,*d*']benzo[1,2-*a*:4,5-*a*']-dicycloheptenes (**1 a**-**c**) and related molecules.

organic semiconductor for applications in organic thin-film transistors (OTFTs). The synthesis, X-ray crystal structures and OTFTs containing compounds **1a–c** are described.

Scheme 1 details the synthesis of molecules **1a-c** starting from compound 5,^[11] which was prepared from *p*-xylene in four steps by modification of reported methods.^[12] Cyanation of compound 5 was achieved by heating with CuCN to reflux in DMF to form 6 as a mixture of isomers, which were used in the subsequent catalytic hydrogenation without separation. In contrast, lithiation of molecule 5 followed by reaction with CO2 afforded the corresponding diacid in very poor yield, and subsequent catalytic hydrogenation of 5 led to debromination. Hydrolysis of dicyanide 7 resulted in the diacid 8. The total yield of 8 from p-xylene in seven steps was 49%. In comparison to this synthesis, the earlier reported preparation of 8 from phenylacetic acid and pyromellitic dianhydride required fewer steps but had a much lower total yield (14%), and involved a byproduct that could lead to explosion.^[13] Double cyclization of diacid 8 with polyphosphonic acid at elevated temperature and subsequent dehydrogenation yielded dione **10**, which was first reported 40 years ago^[13] but almost neglected afterwards. Addition of various acetylide compounds to 10 produced diols 11a-c. Subsequent reduction of the intermediate diol 11a in THF with a solution of 37% HCl that was saturated with SnCl₂ led to the formation of a deep purple solution, from which 1a was isolated as a stable compound in good yield. The same

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Scheme 1. Synthesis of 1 a-c. NBS = N-bromosuccinimide. DBN = 1,5diazabicyclo[4.3.0]non-5-ene.

method was used to prepare **1b** and **1c** from diols **11b** and **11c**, respectively, with lower yields.

Two findings related to the stability of **1a-c** should be noted. First, 1a-c appeared unstable on common silica gel or neutral alumina for unknown reasons. However, these compounds were successfully purified with column chromatography using silica gel or neutral alumina that were deactivated with 6% of water (detailed in the Supporting Information). Second, it was necessary to keep the concentration of 11b,c at 2 mm or lower to achieve a good yield of 1b,c. When the concentration of **11b**,**c** in THF was higher, the yield of **1b**,**c** decreased dramatically and the ¹H NMR spectra of the crude products exhibited broad signals, which suggested polymerization of 1b,c under the acidic conditions. To further investigate this observation, the stability of **1a-c** in solution (concentration 0.02 M) was monitored with ¹H NMR spectroscopy. It was found that after storage of the solutions under ambient conditions for six days, no changes were apparent in the ¹H NMR spectra of the solutions of **1a** and **1b** in CDCl₃ (Supporting Information). In contrast, the solution of 1c in CDCl₃ turned dark red and exhibited a broad signal in the aromatic region of the ¹H NMR spectra after storage under the same conditions for one day. The instability of 1c may be related to polymerization of the reactive C-C double bonds in the π -conjugated backbone. In contrast, compounds **1a** and 1b are more stable, likely because their larger substituents can hinder polymerization by blocking overlap of the reactive C-C double bonds.



Figure 2. a) The pentacyclic backbone of **1a** in the crystal structure with partial carbon-atom labeling and some highlighted bond lengths; b) π -stacking between **1a** molecules in the single crystal; c) molecular packing of **1b** as viewed along the *c*-axis of the unit cell. (Hydrogen atoms are removed for clarity. Thermal ellipsoids set at 50% probability, silylethynyl substituents are shown as capped sticks, and disordered triethyl groups of **1b** are shown as dots.)

Crystals of **1a** and **1b** suitable for single-crystal X-ray analysis were grown from solutions in hexane and ethyl acetate.^[14] Figure 2a shows the pentacyclic backbone of **1a**, which is essentially flat. Examination of the bond lengths indicates that the central six-membered ring has two short C-C bonds (C6–C6a and C14–C14a: 1.37 Å) and four long C–C bonds (C5a-C6 and C13a-C14: 1.43 Å; C5a-C14a and C6a-C13a: 1.45 Å) and is bonded to C5 and C13 by relatively short bonds (C5–C5a and C13–C13a: 1.40 Å). These bond lengths are similar to the corresponding bond lengths in the crystal structure of 3b,^[10a] and are in agreement with a *p*-quinodimethane structure. The seven-membered ring contains a C-C double bond (C15-C16) which measures 1.33 Å, a typical bond length for alkenes. In comparison, the C5-C4a, C14a-C15, and C16-C16a bonds in the seven-membered ring measure 1.44–1.48 Å and resemble single bonds between two sp²-hybridized carbon atoms.^[15] Unlike its pentacene analogue 2 (Figure 1),^[16] which stacks in a two-dimensional faceto-face π -stacking arrangement, compound **1a** forms one-

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dimensional offset π -stacks (Figure 2b). In each stack, two parallel neighbors have roughly two rings overlapped with a π -to- π distance of 3.47 Å. The non-overlapped space above and below the pentacyclic backbone is occupied by the triisopropylsilyl groups of neighboring stacks. The fact that **1a** forms one-dimensional π -stacks rather than two-dimensional π -stacks is likely related to the substitution position of bulky triisopropylsilyl groups. With a smaller substituting group, **1b** exhibits a special herringbone-type arrangement (shown in Figure 2c), where one molecule of **1b** has the edge of its termini contacted with the π -face of a neighboring molecule. This packing motif involves much less intermolecular overlap of π -orbitals than the established herringbone packing of pentacene.

Molecules 1a-c are deep purple and nonfluorescent in solution when excited with UV light, and exhibit almost identical UV/Vis absorption spectra. The UV/Vis absorption spectra of compounds 1a and 2 are shown in Figure 3 for comparison. In the visible-light region, 1a exhibits more



Figure 3. UV/Vis absorption spectra of 0.05 mm solutions of 1a and 2 in dichloromethane.

intense absorption than 2. The longest wavelength absorption maxima of **1a** occurs at $\lambda = 587$ nm, which is blue-shifted by 55 nm relative to that of 2. To determine the energy levels of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), the redox behavior of compound 1a in solution in dichloromethane was investigated with cyclic voltammetry. The cyclic voltammogram of **1a** (Supporting Information) exhibited a reversible reduction wave and a reversible oxidation wave with halfwave potentials of -1.66 V and 0.12 V versus ferrocene/ ferrocenium, respectively. Based on these electrochemical potentials, the HOMO and LUMO energy levels of 1a are estimated as -4.92 eV and -3.14 eV, respectively.^[17] These energy levels lead to a HOMO-LUMO energy gap of 1.78 eV, which is in good agreement with the absorption edge at approximately $\lambda = 649$ nm (1.91 eV). In comparison, the HOMO and LUMO energy levels of 2, also estimated from electrochemical potentials, are -5.17 eV and -3.30 eV, respectively,^[18] and the HOMO and LUMO energy levels of **3a** are -5.88 eV and -4.00 eV, respectively.^[10b] The higher HOMO and LUMO energy levels of 1a correspond to the fact that the seven-membered ring in 1a is more easily oxidized than the benzene ring in 2 and the five-membered ring in 3a by forming an aromatic cycloheptatrienyl cation.

To test the semiconductor properties of **1a–c**, thin-film transistors of **1a–c** were fabricated by dip-coating or dropcasting a solution onto an oxidized silicon substrate. Highquality films of **1a** were formed by immersing a SiO₂/Si substrate in a solution of **1a** (2 mg mL⁻¹) in a solvent mixture of dichloromethane and acetone (1:1 by volume) and then pulling it up with a constant speed of 5.4 μ ms⁻¹. The films of **1a** deposited on the SiO₂ surface were composed of aligned crystalline fibers, as shown in the polarized-light micrograph of Figure 4a. X-ray diffraction patterns from the films of **1a** (Supporting Information) exhibited one peak at a *d*-spacing of 11.19 Å (2 θ =7.90°) and three higher-order peaks at *d*spacings of 5.59 Å, 3.72 Å, and 2.79 Å. These peaks do not correspond to any diffractions derived from the single-crystal structure of **1a**, and thus indicate a polymorph different from



Figure 4. a) Reflection polarized-light micrograph of a film of **1a** as dip-coated on SiO₂ at a substrate temperature of room temperature. b) Drain current (I_{DS}) versus gate voltage (V_{C}) with drain voltage (V_{DS}) at -50 V for the best-performing OTFT of **1a** measured in air with an active channel of W=1 mm and L=50 µm.

its bulk crystals. The device fabrication was completed by depositing a layer of gold on the film of **1a** through a shadow mask to form top-contact source and drain electrodes. As measured in air from these devices. **1a** functioned as a p-type semiconductor with a field-effect mobility in the range of 0.19–0.76 $\text{cm}^2 \text{V}^{-1} \text{s}^{-1}$. The highest mobility was extracted from the transfer I-V curves shown in Figure 4b using the equation: $I_{\rm DS} = (\mu W C_i/2L)(V_G - V_T)^2$, where $I_{\rm DS}$ is the drain current, μ is field-effect mobility, C_i is the capacitance per unit area (11 nF cm^{-2}) for the 300 nm-thick dielectric layer of SiO₂, W is the channel width, L is the channel length, and V_G and V_T are the gate and threshold voltage, respectively. It was found that the mobility of $0.76 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ decreased to $0.57 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ and $0.39 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ after the device was stored in air for one week and one month, respectively. Such degradation of mobility can be attributed to the fact that the thin-film phase gradually changes to the more stable bulkcrystal phase. This is shown by the X-ray diffractions from the film stored for one week (Supporting Information), which indicate the existence of both the thin-film phase and the bulk-crystal phase of 1a. The thin-film transistors of 1b and 1c exhibited field-effect mobilities of 6×10^{-4} and $5 \times$ $10^{-5} \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$, respectively, for holes. The low mobility of

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1b is in agreement with its molecular packing, which only has poor edge-to-face interactions between the π -backbone of neighboring molecules.

The device exhibits a positive threshold voltage of approximately 10 V and a large drain current of circa $1 \times$ 10^{-5} A at zero gate bias (Figure 4b). The drain current is suppressed by a positive gate voltage of 12 V to approximately 1×10^{-9} A, which corresponds to an on/off ratio of greater than 105 when compared with the on-current obtained at a gate voltage of -50 V. The large drain current at zero gate bias and the positive threshold voltage of 1a are similar to those of the solution-processed OTFTs of 2 reported earlier,^[19] and indicate that the film of **1a** is already p-doped at zero gate bias. This observation is in agreement with the fact that molecule 1a has an even higher HOMO energy level than 2. To identify the possible dopants, the thin-film transistors of 1a were then tested under vacuum. It was found that under vacuum the transistor of 1a exhibited essentially the same drain current of approximately 1×10^{-5} A at zero gate bias, suggesting that the film is doped not by oxygen from the air but likely by oxygen species on the SiO₂ dielectric surface. To test this hypothesis, the SiO₂ surface was pretreated with a self-assembled monolayer (SAM) of organosilane. As shown in the Supporting Information, when the SiO₂ surface was modified with phenyltrichlorosilane,^[20] the drain current at zero gate bias was reduced to circa 1×10^{-8} A and the threshold voltage shifted to approximately 3 V. This change is probably because the oxygen species on the SiO₂ were passivated by the SAM of phenyltrichlorosilane. When the SiO₂ was modified with 3-(aminopropyl)triethoxysilane,^[21] the drain current at zero gate bias was further reduced to approximately 1×10^{-10} A and the threshold voltage shifted to -26 V. This change occurs because the surface amino groups are known to cause a shift in threshold voltage to more negative values, presumably by electron transfer into organic semiconductors.

In summary, this study puts forth a new class of conjugated polycyclic molecules that contain a linearly annulated C_6 - C_7 - C_6 polycyclic framework. The crystal structures of **1a** and **1b** indicate that their pentacyclic π -backbone is nearly flat with a *p*-quinodimethane core. With bulky substituents, **1a** and **1b** exhibit robust environmental stability in solution. It is found that **1a–c** function as p-type organic semiconductors in solution-processed OTFTs with field-effect mobilities of up to $0.76 \text{ cm}^2 \text{V}^{-1} \text{s}^-$. These results suggest that heptagonembedded conjugated polycyclic hydrocarbons are promising candidates for organic electronic materials.

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Communications

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Heptagon-Embedded Pentacene: Synthesis, Structures, and Thin-Film Transistors of Dibenzo[*d*,*d*']benzo[1,2*a*:4,5-*a*']dicycloheptenes



All sixes and sevens: A new class of conjugated polycyclic molecules have a nearly flat C_6 - C_7 - C_6 - C_7 - C_6 polycyclic framework with a *p*-quinodimethane core. With a field-effect mobility of up to 0.76 cm²V⁻¹s⁻¹ as measured from solution-processed thin-film transistors, these molecules are alternatives to the pentacene analogues for application in organic electronic devices.

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