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## Twisted diphenoquinones fused with thiophene rings: thiophene analogs of bianthrone†

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The twisted conformer of bistricyclic aromatic enes (BAEs) has a small HOMO-LUMO gap owing to the twisted double bond. In this study, we synthesized diphenoquinones fused with thiophene rings as a new twisted conformer-predominant BAE. They exhibited deep LUMO energy levels and apparent n-type semiconductor properties.

 $\pi$ -Conjugated materials are widely used in various applications such as organic electronics and imaging materials thanks to their distinctive characteristics such as photoabsorption in the UV-vis-NIR region and semiconducting properties. In order to induce a bathochromic shift in the photoabsorption, the  $\pi$ - $\pi$ \* energy gap must be decreased. This can be achieved by extending the  $\pi$ -conjugation,  $^{1a-c}$  utilizing intramolecular donoracceptor interactions <sup>1a-c</sup> and increasing quinoidal character, <sup>1d</sup> which, unfortunately, results in large molecular structures or requires multistep synthesis. Therefore, new molecular designs that realize long-wavelength absorption in small and simple structures are desired. On the other hand, the twisting of the double bond decreases the overlap between the two p-orbitals, leading to increased reactivity, a biradical character, and a lowenergy triplet state.<sup>2</sup> In addition to these characteristics, the twisting of the double bond leads to a decrease in the transition energy by decreasing the energy gap between the  $\pi/\pi^*$  orbitals (Fig. 1a), which is an important consideration in the design of molecules for the development of  $\pi$ -conjugated materials with a small HOMO-LUMO gap. For example, 9,9'-bifluorenylidene, a typical example of bistricyclic aromatic enes (BAEs),3 is known

(a) Schematic diagram of the energy changes in  $\pi/\pi^*$  orbitals due to the twisting of a double bond. Structures of BAEs based on benzene rings (b) and thiophene rings (c).

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to have a twisted conformer to avoid steric hindrance in the fjord region (Fig. 1b). Its central double bond has a dihedral angle of 42°, and its maximum absorption wavelength (458 nm in  $CH_2Cl_2$ )<sup>4</sup> is much longer than that of dibenzo[g,p]chrysene (351 nm in CH<sub>2</sub>Cl<sub>2</sub>),<sup>5</sup> a planar structural isomer of 9, 9'-bifluorenylidene. In BAEs, which does not contain fluorenylidene for the tricyclic structures, a conformational isomer called the folded conformer, in which the tricyclic structures are folded and there is no twisting in the double bond, is generally more thermodynamically stable than its twisted counterpart.<sup>3</sup> For example, bianthrone, which is a diphenoquinone fused with benzene rings, shows thermochromism,6 predominantly exists as a folded conformer, and is light yellowish at room temperature owing to the large HOMO-LUMO gap (Fig. 1b). Whereas various homomerous<sup>7</sup> and heteromerous BAEs<sup>8</sup> have been reported, twisted-conformer-predominant BAEs are limited to those containing a five-membered ring structure such as fluorenylidene, 4,9 and their structures have

Bifluorenvlidene Bianthrone Twisted-predominant Folded-predominant Diphenoquinone  $\lambda_{\text{max}}$  = 392 nm (in CH<sub>2</sub>Cl<sub>2</sub>)  $\lambda_{max} = 458 \text{ nm (in CH<sub>2</sub>Cl<sub>2</sub>)}$ FMes = 2,4,6-tris(trifluoromethyl)phenyl This work Thiophene dDTCB dDTCO  $\lambda_{max}$  = 661 nm (in toluene)

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limited diversity. As a new strategy to control the energy of BAE conformers, we have focused on the use of heterocycles such as thiophene. The more compact structure of thiophene than benzene is expected to markedly influence the electronic structures and the steric hindrance in BAEs. Indeed, we previously reported that the twisted conformer of dDTCB (Fig. 1c), a BAE comprised of thiophene rings and boron, was thermodynamically stable. 10 However, the synthesis of dDTCB was complicated owing to the presence of tricoordinate boron in the structure. In this study, we designed dDTCO, a simple structured diphenoquinone fused with thiophene rings, by substituting sp<sup>2</sup> carbonyl units for the sp<sup>2</sup> boron units of **dDTCB**, as a new twisted-predominant BAE (Fig. 1c). All benzene rings in bianthrone were replaced by thiophene rings in dDTCO. The first synthesis of **dDTCO** was reported by the Mazaki group, <sup>11</sup> but as far as we know, the detailed synthesis methods and properties of the compound have not been reported.

The synthesis of **dDTCO** was accomplished similarly to that of dDTCB (Scheme 1). 10a The precursor of dDTCO, DTCO, was more stable in its enol form than its keto form as only the phenol form was observed in the <sup>1</sup>H NMR spectrum. <sup>12</sup> We found that DTCO was unstable in air; it gradually converted into a quinone upon standing in air both in solution and in the solid state (Fig. S1, ESI†). Therefore, DTCO was used in the subsequent dimerization reaction immediately after synthesis. DTCO quickly dimerized at low temperatures in the presence of iodine and 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), yielding the desired thiophene-fused diphenoquinone (dDTCO) as a dark purple solid. dDTCO was stable in air, showing no decomposition even after storage for more than one year in air in the solid state. In contrast to bianthrone that readily undergoes photocyclization, 13 dDTCO showed very high photostability (Fig. S2, ESI†). dDTCO exhibited poor solubility, being soluble in chloroform but scarcely soluble in toluene or acetone, whereas alkylated derivatives dDTCO-Me and dDTCO-Bu exhibited better solubility. In particular, dDTCO-Bu dissolved completely in toluene. In the <sup>1</sup>H NMR spectrum of **dDTCO**, only two sharp doublet signals corresponding to the protons on the thiophene rings were observed (Fig. S3, ESI†), suggesting that only one isomer is predominantly present in solution.

Next, single-crystal X-ray diffraction analyses were performed. As expected, all the X-ray structures of the thiophenefused diphenoquinones showed twisted conformers similar to dDTCB (Fig. 2 and Fig. S4, ESI†). 10a This contrasted with the folded X-ray structure of bianthrone, 7c suggesting that the substitution of thiophene rings for the benzene rings in bianthrone thermodynamically stabilized the twisted conformer

Scheme 1 Synthesis of dDTCO and its alkylated derivatives.

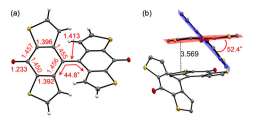


Fig. 2 Monomeric crystal structure with bond lengths in Å (a) and dimeric crystal structure with dihedral angle between the mean planes of the tricyclic structures (b) of dDTCO obtained at 100 K. Thermal ellipsoids are at 50% probability level

relative to the folded conformer. In our previous work, the twist angle of the central double bond was defined by the dihedral angle of the four carbon atoms around the double bond (Fig. 2a). 10a However, for comparison with other studies, here we focused on the dihedral angle between the mean planes of the tricyclic structures (Fig. 2b). The dihedral angle of dDTCO was 52.4°, which was much larger than that of 9,9'bifluorenylidene (42°).4 Although this dihedral angle was slightly smaller than that of dDTCB (53.4°, Fig. S5, ESI†), 10a the difference was only 1.0°. The central C=C double bond of dDTCO was 1.41 Å, much longer than that of bianthrone (1.36 Å), 7c indicating the decreased double bond character in **dDTCO**. Bond alternation in the diphenoquinone structure of dDTCO was smaller than that of bianthrone (Table S1, ESI†), suggesting the highly conjugated structure of dDTCO within the diphenoquinone structure. The monomeric X-ray structures of dDTCO-Me and dDTCO-Bu were similar to that of dDTCO (Fig. S4 and S5, ESI†). Interestingly, despite the twisted structure, **dDTCO** exhibited a dense packing structure with a  $\pi$ - $\pi$ distance of 3.57 Å between two molecules (Fig. 2b), indicating the presence of weak  $\pi$ - $\pi$  interactions. In alkylated derivatives dDTCO-Me and dDTCO-Bu, the intermolecular distances were larger than that of **dDTCO** (Fig. S4, ESI†). The powder X-ray diffraction pattern of dDTCO well matched the pattern simulated from the single-crystal X-ray data (Fig. S6, ESI†).

The UV-vis absorption spectra of the thiophene-fused diphenoquinones in CH<sub>2</sub>Cl<sub>2</sub> are shown in Fig. 3. The strong absorption of dDTCO in the long-wavelength region with a maximum at 608 nm is a typical characteristic of twisted BAEs (Table 1). 4,9,10a Generally, the absorption band shifts to the longer wavelength region as the twist angle of the central double bond

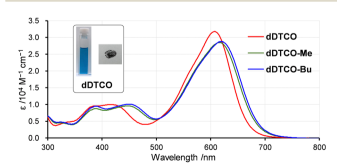


Fig. 3 UV-vis absorption spectra of thiophene-fused diphenoquinones in CH<sub>2</sub>Cl<sub>2</sub> at the concentration of 0.01 mM.

 $\mbox{\bf Table 1} \quad \mbox{Optical and electrochemical data of thiophene-fused diphenoquinones in $CH_2Cl_2$ }$ 

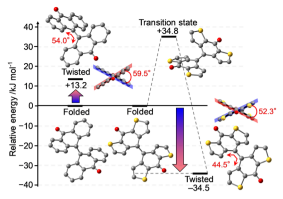
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Compound	$\lambda_{\max}^a/nm$	$E_{\rm ox}^{b}/{\rm V}$	$E_{\mathrm{red}}{}^c/V$	$\mathrm{HOMO}^d/\mathrm{eV}$	LUMO <sup>e</sup> /eV
dDTCO	608	1.01	-0.655	-5.81	-4.15
dDTCO-Me	613	0.895	-0.680	-5.70	-4.12
dDTCO-Bu	617	0.902	-0.698	-5.70	-4.10

<sup>&</sup>lt;sup>a</sup> Absorption maximum. <sup>b</sup> Onset potential of the first oxidation wave in CV ( $\nu$ s. Fc/Fc<sup>+</sup>). <sup>c</sup> Half-wave potential of the first reduction wave in CV ( $\nu$ s. Fc/Fc<sup>+</sup>). <sup>d</sup> HOMO =  $-4.8 - V_{\rm ox}$ . <sup>e</sup> LUMO =  $-4.8 - V_{\rm red}$ .

increases. The dihedral angles in the X-ray structures are ordered as follows: 9,9'-bifluorenylidene < dDTCO < dDTCB,4,10a and accordingly, the absorption maxima are red-shifted. In addition, 1,1',4, 4'-tetramethyl-9,9'-bifluorenylidene, which has methyl groups in the fjord region of 9,9'-bifluorenylidene to increase the steric hindrance,<sup>4</sup> has a dihedral angle of 56°, larger than that of **dDTCO**. Nonetheless, its absorption maximum in CH<sub>2</sub>Cl<sub>2</sub> is 485 nm, much shorter than that of dDTCO. One reason for this is that the aromaticity of thiophene is lower than that of benzene, which effectively extended the conjugation within the molecule for dDTCO.14 The molar absorption coefficient of dDTCO at 608 nm was 31 800 M<sup>-1</sup> cm<sup>-1</sup>, which is similar to that of twisted-conformerdominant **dDTCB** ( $\varepsilon = 32\,200 \text{ M}^{-1} \text{ cm}^{-1}$  at 655 nm in CH<sub>2</sub>Cl<sub>2</sub>). <sup>10a</sup> The absorption maximum of the folded-predominant BAE of bianthrone is around 392 nm in the UV region, 7c revealing a significant impact of the aromatic ring substitution on the electronic structures of BAEs. The absorption maxima of dDTCO-Me and dDTCO-Bu were slightly red-shifted from that of **dDTCO** owing to the inductive effect of the alkyl groups (Fig. 3). Solvent dependence was not observed in the absorption spectra of dDTCO (Fig. S7, ESI†). In the solid-state UV-vis absorption spectrum of dDTCO, the absorption band shifted to longer wavelengths than in solution, probably due to  $\pi$ - $\pi$  interactions in the solid state (Fig. S8, ESI†).

To investigate the influence of each structure on the energy of the conformers, the twisted and folded conformers of dDTCO and bianthrone were optimized in DFT calculations (Fig. 4 and Fig. S9, S10, ESI†). The optimized structures of dDTCO and bianthrone were in good agreement with the single-crystal X-ray structures (Fig. 4 and Fig. S9, ESI†).7c Whereas the folded conformer was 13.2 kJ mol<sup>-1</sup> more stable than the twisted conformer in bianthrone, in the case of **dDTCO**, the twisted conformer was 34.5 kJ mol<sup>-1</sup> more stable than the folded conformer (Fig. 4). The five-membered thiophene rings in dDTCO may reduce the steric hindrance in the fjord region, resulting in a small torsion angle for the twisted conformer and thereby suppressing destabilization in the twisted conformer of dDTCO. Indeed, as shown in Fig. 4, the dihedral angle of the twisted conformer of bianthrone is 59.5°, much larger than that of **dDTCO** (52.3°). We also compared the energies of the corresponding singlet carbenes and dimers (Fig. S11, ESI†), and found a small difference in the formation energies of the folded conformers between bianthrone and dDTCO. However, the formation energy of the twisted conformers is more negative for dDTCO than for bianthrone. This suggest that the thermodynamic stability of the conformers of BAEs is more strongly influenced by the structure of the twisted



**Fig. 4** Gibbs free energy diagram of folded and twisted conformers of bianthrone and **dDTCO** in the closed-shell singlet state at the B3LYP-D3BJ/6-31G(d) level at 298 K. The Gibbs free energy of the folded conformer is set at zero. Hydrogen atoms are omitted for clarity.

conformer than by that of the folded conformer. The energy barrier between the twisted and folded conformers of dDTCO was calculated to be 69.3 kJ mol-1 relative to the twisted conformer (Fig. 4), indicating that the two conformers are in thermal equilibrium at room temperature, like typical BAEs. However, because of the large energy difference between the twisted and folded conformers in dDTCO, the twisted conformer is predominant, as observed in the <sup>1</sup>H NMR and the UV-vis absorption spectra. The triplet state of dDTCO was calculated to be 25.6 kJ mol<sup>-1</sup> higher in energy than the singlet state twisted conformer at 298 K (Fig. S12, ESI†). In the ESR spectrum, the triplet signal was not observed (Fig. S13, ESI†), confirming that the ground state is singlet. From TD-DFT calculations, the  $S_0 \rightarrow S_1$  transition at 599 nm and the  $S_0 \rightarrow S_6$  transition at 420 nm were determined for the twisted conformer of dDTCO (Table S2, ESI†) and found to agree with the experimental absorption spectrum (Fig. S14, ESI†). In the folded conformer, the lowest energy-allowed transition was the  $S_0 \rightarrow S_2$  transition in the UV region at 438 nm (Table S3, ESI†).

Finally, the electrochemical and semiconductor properties of the thiophene-fused diphenoquinones were investigated. dDTCO, dDTCO-Me, and dDTCO-Bu showed irreversible oxidation and reversible reduction waves in the cyclic voltammograms (Fig. 5a and Fig. S15, ESI†). Interestingly, two reversible peaks were observed in the cathodic CV, which may correspond to the formation of radical anion and dianion species with aromatic resonance structures (Fig. 5a).15 The first reduction potentials of dDTCO, dDTCO-Me, and dDTCO-Bu were -0.655, -0.680, and -0.698 V, respectively, from which the LUMO energy levels were calculated to be -4.15, -4.12, and -4.10 eV (Table 1). These LUMO energy levels are deeper than that of **dDTCB**  $(-3.99 \text{ eV})^{10a}$  and comparable to those of typical n-type semiconductors. 16 Therefore, we fabricated organic field-effect transistors (OFETs) using these compounds and evaluated their semiconductor properties. To evaluate the charge carrier transport properties, we fabricated OFETs with top-gate bottomcontact (TGBC) structures<sup>17</sup> using dDTCO-Bu, which has the highest solubility and good film-forming properties. Uniform thin films could not be obtained from dDTCO and dDTCO-Me, and they did Communication ChemComm

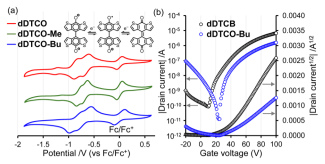


Fig. 5 (a) Cathodic cyclic voltammetry data of thiophene-fused diphenoquinones in dichloromethane containing 0.1 M  $\mathrm{Bu_4NPF_6}$  at the scan rate of 100 mV s $^{-1}$ . (b) Transfer curves of **dDTCB** and **dDTCO-Bu** of OFETs.

not show FET characteristics. Owing to its deep LUMO energy level, dDTCO-Bu exhibited an n-type behavior with a moderate threshold voltage of 39.0 V and a high current on/off ratio of  $1.3 \times 10^5$  in n-channel operation (Fig. 5b), suggesting that the twisted diphenoquinones are a promising building block for n-type semiconductor materials. The maximum electron mobility  $(\mu_e)$  of **dDTCO-Bu** in five cells was  $5.0\,\times\,10^{-4}~\text{cm}^2~\text{V}^{-1}~\text{s}^{-1}\text{,}$  which was lower than that of **dDTCB** (Table S4,  $2.0 \times 10^{-3} \, \text{cm}^2 \, \text{V}^{-1} \, \text{s}^{-1}$ , ESI†). This may be due to the large steric hindrance of the four butyl groups with different tilt angles. Because dDTCO does not contain alkyl groups, it may exhibit superior n-type semiconductor properties. However, owing to its poor film-forming properties in wet processes, OFET evaluation was not possible. By forming films through vapor deposition, it may be possible to create superior OFET devices with dDTCO.

In summary, we performed an investigation of the photophysical properties of diphenoquinones fused with thiophene rings. By substituting thiophene rings for the benzene rings in bianthrone, the twisted conformer became more thermodynamically stable than the folded conformer, and the thiophenefused BAEs were found to exist in the twisted conformer both in the solid state and in solution. The thiophene-fused diphenoquinones possess deep LUMO energy levels, and thin films of **dDTCO-Bu** exhibit clear n-type semiconductor characteristics. The highly twisted BAE structure effectively reduces the HOMO-LUMO gap and is therefore a new molecular design for the development of conjugated molecules with strong absorption in the visible region or organic semiconductors.

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### Data availability

The data supporting this article have been included as part of the ESI.† Crystallographic data for dDTCO, dDTCO-Me, and dDTCO-Bu has been deposited at the CCDC under 2363404-2363406.†

#### Conflicts of interest

There are no conflicts to declare.

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