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Synthesis and structural evaluation of closed-shell folded and open-shell twisted hexabenzo[5.6.7]quinarene

We synthesized hexabenzo[5.6.7]quinarene and demonstrated spin-state switching by a combination of chemical oxidation/reduction and thermal stimulation. This spin-state switching cycles like a planet in orbit around three stars exhibiting chemical oxidation, reduction, and thermal stimulation, as shown in the artwork.



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Synthesis and structural evaluation of closed-shell folded and open-shell twisted hexabenzo[5.6.7]quinarene†‡

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We describe the synthesis and characterization of hexabenzo[5.6.7] quinarene, which is composed of an anthraquinodimethane (AQD) central core that is end-capped with fluorenyl and dibenzosuberenyl groups. Due to strong intramolecular spin-spin interaction through the central AQD unit, this compound is obtained as a stable folded form. Isolation of the stable twisted dication by oxidation and reduction of the dication yields a twisted triplet species, which thermally reverts to the folded form. This is a spin-switching system based on a combination of chemical oxidation/reduction and thermal stimulation.

Thiele's hydrocarbon (TH)¹ is composed of two triphenyl methyl radical units and is sometimes considered as a prebiradical molecule due to the quinoidal π -skeleton. However, the strong intramolecular spin-spin interaction of its two unpaired electrons results in a stable closed-shell species with a large singlet-triplet energy gap (ΔE_{S-T} , Fig. 1a). With the recent research interest in biradicaloid² and multiradicaloid π -systems³ for the development of molecule-based spintronic materials, there is growing attention towards spin-state controllable molecules by external stimuli such as heating,⁴ photoirradiation,⁵ or mechanical stress.⁶ To introduce an open-shell nature to the TH skeleton, an anthraquinodimethane (AQD) unit has been employed as its central core to create steric repulsion between the terminal aromatic rings and the central AQD unit. This facilitates transformation to a biradical form by twisting the ethylene units (Fig. 1b). In fact, several AQD-extended TH derivatives end-capped with tricyclic aromatic units such as fluorenyl (FLU),⁷ 9,10-dihydro-10,10-dimethyl-9-anthryl(DMA),⁸

or dibenzosuberenyl (DBS)⁹ units have been synthesized as systems that can be switched from a stable closed-shell ground state to an open-shell biradical state via heating or photoirradiation. Typically, AQD-extended TH derivatives have symmetric terminal π -skeletons, which leads to homolytic cleavage of the π -bond orbitals on the C=C bonds, resulting in a biradical state.¹⁰ On the other hand, it is also interesting to investigate whether heterolytic π -bond cleavage of C=C bonds, leading to a zwitterionic state, is possible or not in a biradical system. Several zwitterionic biradicaloids with heteroatom(s) embedded in their skeletons have been synthesized.¹¹ Due to the difference in electronegativity between carbon and heteroatoms, electrontransfer between them can occur, changing the electronic state from biradical to zwitterion and vice versa. Moreover, several neutral planar hydrocarbon π -systems including fiveand seven-membered rings have been suggested to exhibit a resonance structure between a biradical state and an electronpolarized state.12



Fig. 1 (a) The structure of Thiele's hydrocarbon. (b) Typical examples of reported π -extended THs, having fluorenyl, 9,10-dihydro-10,10-dimethyl-9-anthryl, or dibenzosuberenyl end-caps. (c) The structure of hexabenzo[5.6.7]quinarene **1** and its structural isomers **1-F** (closed-sell) and **1-T** (open-shell) as well as a formal resonance structure of **1-P** (polarized form).

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 π -extended [5.6.7]quinarene¹³ **1**, composed of an AQD central core end-capped with FLU and DBS groups, is a good candidate for understanding the nature of biradical and polarized character and for investigating whether these states can coexist in a pure hydrocarbon skeleton (Fig. 1c). Previously, we investigated persistent FLU and DBS radicals protected by a 9-anthryl ring and a redox potential difference (ΔE_{redox}) of the DBS radical (E_{ox}) and the FLU radical (E_{re}) was determined to be 0.80 V, which is expected to be a moderate ΔE_{redox} to construct a neutral chargetransfer (CT) complex (Fig. S1, ESI‡).¹⁴ Herein, we report the synthesis of compound **1** and elucidate the structure and properties of folded closed-shell **1-F** and twisted open-shell **1-T** that can be switched by a combination of chemical oxidation/ reduction and thermal stimulation. We also discuss whether twisted polarized species **1-P** can exist.

The synthesis of **1** is shown in Scheme **1**. To introduce a nonsymmetric terminal π -skeleton into the AQM core, key intermediate **3**, possessing an anthrone and DBS unit, was synthesized in two steps from 9,10-dibromoanthracene and dibenzosuberenone. Compound **2** was converted to **3** by treatment with an excess amount of trifluoroacetic acid and water. Although the Barton–Kellogg reaction is a well-known method to construct an overcrowded ethylene¹⁵ to introduce a FLU unit into carbonyl compound **3**, we employed the Peterson-olefination reaction¹⁶ because the stable FLU anion and 9-(trimethylsilyl)fluorenyl anion **4** are readily prepared by *n*-butyllithium in a one-pot reaction. The reaction of **3** and **4** afforded target hexabenzo[5.6.7]quinarene **1** as a pale yellow solid in 74% yield.

To evaluate the structure of **1**, single crystal X-ray diffraction and ¹H NMR measurements were conducted. These measurements revealed that compound **1** possesses a folded structure (**1-F**) rather than a twisted one. **1-F** can exhibit *exo-* and *endo*forms of the DBS unit, and the *exo-*form was identified by X-ray crystallography in the solid state (Fig. 2a and Fig. S3, ESI‡). In addition, the 2D NOESY ¹H NMR spectrum showed that only the *exo-*form is present even in solution, as confirmed by cross peaks between *a-f* and *i-j* protons (Fig. S5, ESI‡). The DFT calculations showed that the *endo-*form is 2.27 kcal mol⁻¹



Fig. 2 (a) X-ray crystallography of **1-F**. Hydrogen atoms are omitted for clarity. (b) Comparison of the energy difference ΔE between *exo-* and *endo-*forms of **1-F** (∞ B97X-D/6-31G**).

higher in energy than the *exo*-form, probably because the *endo*-form of **1-F** has larger steric repulsion between the DBS unit and central AQD core than that of the *exo*-form. From the X-ray analysis, the C=C bond lengths *a*, *b*, and *c* are 1.357 Å, 1.348 Å, and 1.339 Å, respectively. This indicates that, despite the anthracene bridge, the intramolecular spin–spin interaction between the FLU and DBS units is strong, resulting in a folded closed-shell structure.

Next, the structural transformation of 1-F into the twisted structure 1-T was investigated. UV-vis absorption measurements of 1-F in various solvents such as dichloromethane (DCM), toluene, and DMF were attempted to evaluate solventinduced charge transfer, but no significant difference in absorption maximum (λ_{abs} = 354–356 nm) or absorption coefficient ($\varepsilon = ca. 23000 \text{ cm}^{-1} \text{ M}^{-1}$) was observed among them (Fig. S6, ESI[‡]). Mechanical grinding of 1-F to generate 1-T was also attempted,^{6e} but no color change in the solid state was observed. This observation is in accordance with DFT calculations, which revealed that the twisted biradical structure **1-T** is 20 kcal mol⁻¹ higher in energy than **1-F** (Fig. S7, ESI^{\ddagger}). Photoirradiation at 365 nm to 1-F did not induce isomerization to 1-T. Despite these results, the cyclic voltammogram of 1-F showed unique irreversible waves (Fig. 3b). There are three anodic peaks at E = -1.06, +0.24, and +0.92 V and three cathodic peaks at E = +0.41, -0.22, and -2.05 V. This observation indicates that the quinoidal folded form of 1-F undergoes a one-step two-electron oxidation/reduction process, producing the twisted dication $\mathbf{1}^{2+}$ at E = +0.92 V or dianion $\mathbf{1}^{2-}$ at



Scheme 1 Synthetic route to **1** and its dication **1**²⁺. Reagents and conditions: (a) (i) 1.0 eq. of *n*-BuLi, -78 °C, THF, and 30 min; (ii) 0.8 eq. of dibenzosuberenone, -78 °C to RT.; (b) TFA/H₂O, 1,2-dichloroethane; and (c) 4.0 eq. of antimony pentachloride, dichloromethane.



Fig. 3 (a) Cyclic voltammogram of **1-F**. (0.1 M $^{n}Bu_{4}NPF_{6}$ in CH₂Cl₂, scan rate = 0.1 V s⁻¹). Black arrows indicate the scan direction. (b) Conformational change of **1** during the oxidation/reduction processes.

E = -2.05 V, respectively (Fig. 3c). Due to the different redox potentials of charged FLU ($E^{Ox}_{1/2} = +0.28$ V, $E_{1/2}^{Re} = -1.11$ V) and DBS ($E_{1/2}^{Ox} = -0.30$ V, $E_{pc} = -1.78$ V) units (Fig. S1, ESI‡),¹⁴ the twisted dication 1^{2+} or dianion 1^{2-} showed a stepwise reduction or oxidation, respectively, with redox potentials corresponding to those of FLU and DBS radicals attached with a 9-anthryl unit, affording the twisted biradical form 1-T. It is considered that the generated 1-T thermally reverted to 1-F due to the irreversible nature of the waves. Thus, it is expected that the twisted biradical form 1-T can be generated by reduction of dication 1^{2+} at low temperatures.

The key compound of dication 1^{2+} was prepared using antimony pentachloride as the oxidative reagent in DCM, and it was obtained in a good yield (96%, Scheme 1). Single crystals of 1²⁺were obtained, which were suitable for X-ray crystallography and stable under ambient conditions even though it has an antiaromatic fluorenyl cation unit with large positive NICS(0) =+25.7 ppm (Fig. S8, ESI \ddagger). The X-ray structure of $\mathbf{1}^{2+}$ is shown in Fig. 4 and Fig. S4 (ESI[‡]). Unlike the structure of 1-F, the central core of 1^{2+} changed from AQD form to anthryl (Ant) form, which is confirmed by the C-C bond length of the central Ant core (Fig. S3 and S4, ESI‡) and C-C bond length between FLU and the Ant unit (a = 1.436 Å) as well as DBS and the Ant unit (b = 1.516 Å). In addition, the twist angles between FLU and Ant (\angle FLU-Ant) and DBS and Ant (\angle DBS-Ant) are 57.5° and 82.9°, respectively. The smaller twist angle of \angle FLU-Ant is due to lesser steric hindrance of the FLU unit than that of the DBS unit. Therefore, the positive charges are mainly localized on both the DBS and FLU units, but slight charge delocalization on the central Ant unit from the FLU unit is considered, judging from the C-C bond length and the twist angle. This was also suggested by natural population analysis (Fig. S9, ESI‡). The UV-vis spectrum of 1²⁺ exhibited a weak broad band at 1040 nm (Fig. S10, ESI[‡]). The TD-DFT calculations revealed that the broad band is derived from intramolecular CT from the central Ant core to the electron-deficient FLU unit (Fig. S11 and Table S1, ESI‡).

With the dication 1^{2+} in hand, the reduction to afford the twisted biradical **1-T** was conducted by using Zn powder in MeCN solution at -40 °C, affording a dark red powder (Scheme 2). The ESR measurement of biradical **1-T** in toluene showed a characteristic pattern for $\Delta M_{\rm S} = \pm 1$ peaks associated with triplet species (Fig. 5a). Using the point dipole approximation,



Scheme 2 Generation of twisted biradical 1-T from its dication 1^{2+} .

the average spin-spin distance was estimated to be 6.98 Å, which is the midpoint distance between the ring centers of FLU and DBS units (8.67 Å), and between the 9-position of FLU and 5-position of DBS units (5.82 Å, Fig. 5b). These findings provide unambiguous evidence for the conclusion that the triplet state of twisted 1-T can be generated. In addition, DFT calculations revealed that twisted 1-T has an open-shell ground state singlet with large biradical character $y_0 = 0.99$ estimated from the natural orbital occupation number¹⁷ and ΔE_{S-T} is only -0.06 kcal mol⁻¹, indicating that the singlet and triplet energy levels are almost degenerated. The recrystallization of 1-T was performed at room temperature, but the obtained crystal structure was 1-F due to the isomerization occurring in solution even though the color of the solution remained slightly dark red. Thus, through a combination of chemical oxidation/reduction and thermal stimulation, compound 1 exhibits spin-switching behavior between 1-F and 1-T via 1^{2+} . Although the experimental determination of the equilibrium kinetics of 1-T to 1-F was attempted, it was not possible to exclude an equilibrium of intermolecular σ -dimerization between DBU units,^{12c} presumably leading to two FLU radical units in the σ-dimer, similar to the equilibrium of the Chichibabin's hydrocarbon (Scheme S1, ESI[‡]).¹⁸

The optical property of **1-T** was evaluated to determine whether it can exist in a polarized form **1-P**. Unlike dication 1^{2+} , **1-T** did not show near-infrared absorption bands above 1000 nm and the spectrum is similar to that of the 9-Ant FLU radical (Fig. S2, ESI‡). Although several solvents with different polarity such as DCM, toluene, and DMF were used for the measurement of the UV-vis spectra, only minor differences were observed in the longer-wavelength broad weak band over 700 nm (Fig. S12, ESI‡). TD-DFT calculations revealed that the broad weak band originated from a combination of transitions from the FLU unit to the central Ant core and from the FLU-Ant



Fig. 4 X-ray crystallography of $\mathbf{1}^{2+}$. FLU and DBS units are represented by purple and orange colors, respectively. (a) Front view and C–C bond lengths of *a*, *b*, and *c*. (b) Side view and twist angles between FLU and the central anthryl (Ant) unit (\angle FLU-Ant) as well as DBS and the Ant unit (\angle DBS-Ant). Protons and counter anions SbCl₆⁻ are omitted for clarity.



Fig. 5 (a) ESR spectrum of **1-T** ($\Delta M_{\rm S} = \pm 1$) in toluene at -113 °C. (b) Calculated structure of **1-T** and carbon–carbon distances between the ring centers of the FLU and DBS units and between the 9-position of FLU and 5-position of DBS units (U ω B97X-D/6-31G**).

unit to the DBS unit (Fig. S13 and Table S2, ESI[‡]). On the other hand, the electrostatic potential surfaces of **1-T** in both openshell singlet and triplet structures showed no significant charge deflection between the FLU and DBS units (Fig. S14, ESI[‡]). Additionally, negligible difference in absolute spin density of the FLU and DBS units was observed between open-shell singlet and triplet structures (Fig. S15, ESI[‡]). These results indicate that there is no electron transfer between FLU and DBS in the singlet state and that the polarized form of **1-P** cannot be obtained under ambient conditions.

In summary, we synthesized hexabenzo[5.6.7] quinarene 1 and evaluated its properties. Although 1 has a folded form (1-F) as the most stable structure and does not undergo mechanochemical conversion to its twisted form (1-T) in the solid state, dication 1^{2+} could be readily prepared, which has the twisted form, enabling us to obtain twisted 1-T by chemical reduction. The generation of 1-T was unambiguously confirmed from the triplet signals in ESR measurement. Although the polarized form of 1-P could not be observed, a combination of chemical oxidation/reduction and thermal stimulation led to the discovery of spin-switching behaviour between 1-F and 1-T *via* 1^{2+} .

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Conflicts of interest

There are no conflicts to declare.

Notes and references

- (a) J. Thiele and H. Balhorn, *Ber. Dtsch. Chem. Ges.*, 1904, 37, 1463;
 (b) L. K. Montgomery, J. C. Huffman, E. A. Jurczak and M. P. Grendze, *J. Am. Chem. Soc.*, 1986, 108, 6004.
- 2 (a) M. Abe, Chem. Rev., 2013, 113, 7011; (b) Z. Zeng, X. Shi, C. Chi, J. T. Navarrete, J. Casado and J. Wu, Chem. Soc. Rev., 2015, 44, 6578; (c) T. Stuyver, B. Chen, T. Zeng, P. Geerlings, F. D. Proft and R. Hoffmann, Chem. Rev., 2019, 119, 11291; (d) T. Kubo, Bull. Chem. Soc. Jpn., 2021, 94, 2235.
- 3 T. Y. Gopalakrishna, W. Zeng, X. Lu and J. Wu, Chem. Commun., 2018, 54, 2186.

- 4 (a) C. Wentrup, M. J. Regimbald-Krnel, D. Müller and P. Comba, Angew. Chem., Int. Ed., 2016, 55, 14600; (b) S. Nishida, Y. Morita, K. Fukui, S. Maki and K. Nakasuji, Angew. Chem., Int. Ed., 2005, 44, 7277; (c) Y. Hamamoto, Y. Hirao and T. Kubo, J. Phys. Chem. Lett., 2021, 12, 4729.
- 5 (a) K. Fujita, S. Hatano, D. Kato and J. Abe, Org. Lett., 2008, 10, 3105;
 (b) Y. Kishimoto and J. Abe, J. Am. Chem. Soc., 2009, 131, 4227;
 (c) K. Uchida, S. Ito, M. Nakano, M. Abe and T. Kubo, J. Am. Chem. Soc., 2016, 138, 2399.
- 6 (a) T. Nishiuchi, S. Uno, Y. Hirao and T. Kubo, J. Org. Chem., 2016, 81, 2106; (b) T. Nishiuchi, S. Aibara and T. Kubo, Angew. Chem., Int. Ed., 2018, 57, 16516; (c) T. Nishiuchi, K. Kisaka and T. Kubo, Angew. Chem., Int. Ed., 2021, 60, 5400; (d) T. Nishiuchi, S. Aibara, T. Yamakado, R. Kimura, S. Saito, H. Sato and T. Kubo, Chem. – Eur. J., 2022, 28, e202200286; (e) T. Nishiuchi, S. Aibara, H. Sato and T. Kubo, J. Am. Chem. Soc., 2022, 144, 7479; (f) T. Nishiuchi, Y. Makihara, R. Kishi, H. Sato and T. Kubo, J. Phys. Org. Chem., 2023, 36, e4451.
- 7 X. Yin, J. Z. Low, K. J. Fallon, D. W. Paley and L. M. Campos, *Chem. Sci.*, 2019, **10**, 10733.
- 8 T. Nishiuchi, R. Ito, E. Stratmann and T. Kubo, *J. Org. Chem.*, 2020, 85, 179.
- Y. Ishigaki, Y. Hayashi and T. Suzuki, J. Am. Chem. Soc., 2019, 141, 18293; (b) V. G. Jiménez, P. Mayorga-Burrezo, V. Blanco, V. Lloveras, C. J. Gómez-García, T. Šolomek, J. M. Cuerva, J. Veciana and A. G. Campaña, Chem. Commun., 2020, 56, 12813; (c) Y. Hayashi, S. Suzuki, T. Suzuki and Y. Ishigaki, J. Am. Chem. Soc., 2023, 145, 2596.
- 10 (a) Y. Ishigaki, T. Hashimoto, K. Sugawara, S. Suzuki and T. Suzuki, Angew. Chem., Int. Ed., 2020, 59, 6581; (b) K. Li, Z. Xu, J. Zu, T. Weng, Z. Chen, S. Sato, J. Wu and Z. Sun, J. Am. Chem. Soc., 2021, 143, 20419.
- (a) E. Niecke, A. Fuchs, F. Baumeister, M. Nieger and W. W. Schoeller, *Angew. Chem., Int. Ed. Engl.*, 1995, 34, 555; (b) M. Abe, W. Adam and W. M. Nau, *J. Am. Chem. Soc.*, 1998, 120, 11304; (c) Z. Zeng, S. Lee, M. Son, K. Fukuda, P. M. Burrezo, X. Zhu, Q. Qi, R.-W. Li, J. T. L. Navarrete, J. Ding, J. Casado, M. Nakano, D. Kim and J. Wu, *J. Am. Chem. Soc.*, 2015, 137, 8572; (d) S. Arikawa, A. Shimizu and R. Shintani, *Angew. Chem., Int. Ed.*, 2019, 58, 6415.
- 12 (a) X. Yang, X. Shi, N. Aratani, T. P. Gonçalves, K.-W. Huang, H. Yamada, C. Chi and Q. Miao, *Chem. Sci.*, 2016, 7, 6176;
 (b) X. Fu, H. Han, D. Zhang, H. Yu, Q. Hi and D. Zhao, *Chem. Sci.*, 2020, 11, 5565; (c) C. Zhen, S. Lu, M. Lin, J. Wu, I. Chao and C. Lin, *Chem. – Eur. J.*, 2021, 27, 16682; (d) K. Horii, R. Kishi, M. Nakano, D. Shiomi, K. Sato, T. Takui, A. Konishi and M. Yasuda, *J. Am. Chem. Soc.*, 2022, 144, 3370.
- 13 (a) K. Takahashi, S. Takenaka, Y. Kikuchi, K. Takase and T. Nozoe, Bull. Chem. Soc. Jpn., 1974, 47, 2272; (b) K. Takahashi, I. Oikawa and K. Takase, Chem. Lett., 1974, 1215; (c) K. Takahashi and K. Takase, Tetrahedron Lett., 1975, 15, 245.
- 14 T. Nishiuchi, R. Ito, A. Takada, Y. Yasuda, T. Nagata, E. Stratmann and T. Kubo, *Chem. – Asian J.*, 2019, **14**, 1830.
- (a) H. Staudinger and J. Siegwart, *Helv. Chim. Acta*, 1920, 3, 833;
 (b) D. H. R. Barton and B. J. Willis, *J. Chem. Soc. D*, 1970, 1225;
 (c) R. M. Kellogg and S. Wassenaar, *Tetrahedron Lett.*, 1970, 11, 1987;
 (d) A. M. Schoevaars, W. Kruizinga, R. W. J. Zijlstra, N. Veldman, A. L. Spek and B. L. Feringa, *J. Org. Chem.*, 1997, 62, 4943.
- 16 D. J. Peterson, J. Org. Chem., 1968, 33, 780.
- 17 D. Doehnert and J. Koutecky, J. Am. Chem. Soc., 1980, 102, 1789.
- 18 (a) A. E. Tschitschibabin, Chem. Ber., 1907, 40, 1810; (b) W. J. V. Hart and L. J. Oosterhoff, Mol. Phys., 1970, 18, 281; (c) L. K. Montgomery, J. C. Huffman, E. A. Jurczak and M. P. Grendze, J. Am. Chem. Soc., 1986, 108, 6004.