Infinitene: A Helically Twisted Figure-Eight [12]Circulene Topoisomer

Maciej Krzeszewski,† Hideto Ito,*,‡ and Kenichiro Itami*,†⊥

† Department of Chemistry, Graduate School of Science, Nagoya University, Nagoya 464-8602, Japan. † Institute of Transformative Bio-Molecules (WPI-ITbM), Nagoya University, Nagoya 464-8602, Japan

This paper is dedicated to the memory of Prof. François Diederich.

ABSTRACT: New forms of molecular nanocarbons particularly looped polyarenes adopting various topologies contribute to the fundamental science and practical applications. Here we report on the synthesis of an infinity-shaped polycyclic aromatic hydrocarbon, infinitene 1 (cycl[ccc...ccc]dodecakisbenzene) comprising consecutively fused 12-benzene rings forming an enclosed loop with a strain energy of 60.2 kcal·mol⁻¹. Infinitene 1 represents a topoisomer of still-hypothetical [12]circulene, and its scaffold can be formally visualized as the outcome of the “stitching” of two homochiral [6]helicene subunits by their both ends. The synthetic strategy encompasses transformation of a rationally designed dithiacyclophepane to cyclophadiene through the Stevens rearrangement and pyrolysis of the corresponding S,S'-bis(oxide) followed by the UV-light mediated twofold photocyclization. The structure of infinitene 1 is a unique hybrid of helicene and circulene with a molecular formula C₉₈H₅₂, which can be regarded as an isomer for kekulene, [6,6]carbon nanobelt ([6,6]CNB), [12]cycloacene, and tetrabenzo[8]circulene as well. Infinitene 1 is a bench-stable yellow solid with green fluorescence, and soluble to common organic solvents. The figure-eight molecular structure of 1 was unambiguously confirmed by X-ray crystallography. The scaffold of 1, reminiscent of a squeezed spring, stem from its enclosed, fully-fused architecture, is significantly compressed as manifested by a remarkably shortened distance (3.152–3.192 Å) between the centroids of two π-π stacked central benzene rings and the closest C–C distance of 2.920 Å. Combined lamellar and herringbone-like crystal packing suggested three-dimensional electronic interactions. Fundamental photophysical properties of infinitene 1 were thoroughly elucidated by means of UV-vis absorption and fluorescence spectroscopic studies as well as density functional theory (DFT) calculations. Its configurational stability enabled separation of the corresponding enantiomers (P, P) and (M, M) by a chiral HPLC. Circular dichroism (CD) and circularly polarized luminescence (CPL) measurements revealed that 1 has moderate [g₁₂] and [g₂₃] values.

INTRODUCTION

Looped polyarenes adopting various topologies such as cycloarenes,† circulenes,§ carbon nanobelt (CNB),∥ and cyclopaphenynes† have drawn considerable attention in the scientific community. Not surprisingly, many of these synthetically challenging polycyclic aromatic hydrocarbons (PAHs) have been elusive for a long time. However, with the ongoing development of synthetic chemistry at large, many captivating scaffolds were obtained recently, enabling not only theoretical, but also practical investigations. These fascinating molecules contribute to the fundamental science, addressing long-standing questions regarding nature of aromaticity, providing experimental evidences for the distribution of π-electrons in large polyaromatic systems.⊥

The prominent representatives of the cyclic polyarene family are kekulene (cyclo[de...de]dodecakisbenzene), C₉₈H₅₂, and its structural isomers such as [6,6]CNB (cyclo[cccc...cccc]dodecakisbenzene), and tetra-cata-tetrabenzo[8]circulene, as well as unexplored entities such as [12]cycloacene (cyclo[dddd...dddd]dodecakisbenzene) and [12]circulene (cyclo[ee...ee]dodecakisbenzene) (Figure 1a). Very interestingly, these “C₉₈H₅₂ family” include other structurally unique nanographenes such as hexa-peri-hexabenzocoronene, fullerene/higher order bucky bowl, porous graphene sheet, carbon nanotubes and Mackay crystal. These multidimensional carbon architectures were deemed to be potential game-changing materials displaying outstanding physical and mechanical properties. Undoubtedly, controlling size, shape and topology of the...
nanocarbon structures is of utmost importance for fine tuning the features of final materials.\textsuperscript{10}

unique, hybrid of helicene and circulene comprising twelve benzene rings (\textit{C}_{12}H_{16} \textit{1}), dubbed \textit{infinitene} due to its shape reminiscent of the infinity symbol.

**RESULTS AND DISCUSSION**

**Design and Synthesis**

Our molecular design of infinitene \textit{1} was inspired by the historical synthesis of kekulene by Staab and Diederich that utilized a sequence of dihydroxylation formation, sulfur extraction, cyclization, and endgame aromatization.\textsuperscript{20} As a bridge contraction through the Stevens rearrangement was well established and powerful strategy in the chemistry of cyclophanes,\textsuperscript{20,21} we envisioned that sulfur-based, macrocyclization-annulation approach would lead to the first synthesis of infinity-shaped cycloarene \textit{1} (Figure 1). In designing retro-synthesis of \textit{1}, we became aware of the work of Matusbara and Yamamoto who have attempted to synthesize an analogous figure-eight cycloarene nearly twenty years ago.\textsuperscript{24} Although they did not succeed in obtaining their target compound, they did report a successful synthesis of a strained macrocyclic dithioether through a double S\textsubscript{2}′ reaction between 4,4′-bis(bromomethyl)-1,1′-biphenyl and dibenzo[\textit{c}′\textit{f}]chrysene-3,11-diylmethanethiol. As the first step toward the first synthesis of \textit{1}, we succeeded in the crucial macrocyclization of 3,9-bis(bromomethyl)chrysene \textit{2} and dibenzo[\textit{c}′\textit{f}]chrysene-3,11-diylmethanethiol \textit{3} under the influence of Cs\textsubscript{2}CO\textsubscript{3}, furnishing dihydroxylation macrocycle \textit{4} in 25% yield (see SI for details). In this reaction, its rotamer \textit{4}′ and cyclic tetrathiole 4\textsuperscript{′} (dimerization product of \textit{3}) were also formed and isolated in 5% and 16% yield, respectively. Subsequently, we subjected \textit{4} to the methylation reaction with dimethoxycarbenium tetrafluoroborate (the Borch reagent),\textsuperscript{22} followed by NaH-mediated Stevens rearrangement of the thus-obtained salt. As a result, we obtained a complex mixture of stereoisomers of \textit{5} in 57% yield. The resultant mixture was then subjected to the selective oxidation with 2.0 equivalents of m-CPBA to afford the corresponding S,S\textsuperscript{′}-bis(oxide) \textit{6} in 99% yield as a mixture of isomers. Its pyrolysis at 500 °C under vacuum of 0.04–0.05 Torr gave cyclodihene \textit{7} in 46% yield. Finally, cyclodihene \textit{7} underwent smooth twofold photocyclization under UV-light giving the desired target molecule \textit{1} in 89% yield as a bench-stable yellow solid (Figure 2). Unlike barely soluble kekulene, infinitene \textit{1} shows good solubility toward common organic solvents such as CHCl\textsubscript{3}, CH\textsubscript{2}Cl\textsubscript{2}, benzene and ethyl acetate, most likely due to its nonplanar structure.
X-ray Crystallographic Analysis

The structure of infinitene 1 was unambiguously confirmed and analyzed by X-ray crystallography (Figure 3, Table S1 and Figures S2–S6). A suitable crystal was grown by recrystallization of a racemic mixture rac-1 from a chloroform solution via slow vapor diffusion of methanol at room temperature. Infinitene 1 crystallized with the monoclinic space group $P2_1/n$ in the unit cell: $a = 11.2861(11) Å$, $b = 13.3592(10) Å$, $c = 19.508(2) Å$, $\alpha = 90^\circ$, $\beta = 102.261(11)^\circ$, $\gamma = 90^\circ$. The scaffold of 1, reminiscent of a squeezed spring, stem from its enclosed, fully-fused architecture, is significantly compressed contrasting with unsubstituted helicenes. This is manifested by a remarkably shortened distance (3.152–3.192 Å) between the centroids of two π-π stacked central benzene rings and the closest C–C distance 2.920 Å (Figure 3a), compared to the analogous distance between the centroids of two terminal rings of unsubstituted [7]helicene, which reaches 3.804 Å (CCDC 852537). The recent paper on similar figure-eight larger CNB by Wu also reported no interaction between intersected benzene rings. Consequently, the splay angle of 1, determined as the mean torsional angle of the corresponding inner [5]helicene subunit, was much shallower $\phi_{\text{mean}} = 20.34^\circ$ than an analogous splay angle of [7]helicenes $\phi_{\text{mean}} = 24.35^\circ$. The end-to-end twist angle $\phi_{\text{hot}}$ of the junction naphthalene unit reached 22.03°.

The bond lengths of all convex-armchair edges (K-regions) vary only in the range of 1.334–1.359 Å suggesting their significant double bond character. The crystal packing structure indicates the strong three-dimensional intermolecular electronic interactions due to the fact that the $(P,P)$- and $(M,M)$-isomers of 1 tightly arranged themselves in a stacking array lacking any solvent molecules inside the crystal. Concurrently, the crystal packing reveals the presence of stacked lamellar homochiral molecular layers of each enantiomer lying on the ab plane interlacing each other along the c-axis (Figure 3c). The least-squares based mean planes of two intermolecular planes were separated by the length of 9.585 Å (Figure 3d) and the shortest contact between two heterochiral molecules was found between C(5) and H(16) atoms with a gap of 2.873 Å. At the same time, within the homochiral layer, tight herringbone arrangement along the b-axis can be observed (Figure 3e) with the closest intermolecular contact found between the C(1) and H(11) atoms of two different homochiral species with a separation of 2.819 Å (see SI for the packing and short contacts details).

Figure 2. The synthetic pathway towards infinitene 1. Reagents and conditions: (a) CaCO$_3$ (1.1 equiv), DMF, 55 °C, 68 h, 4 (25%), 4* (5%), 4' (16%); (b) i) (MeO)$_2$CHBF$_3$ (6.0 equiv), CH$_2$Cl$_2$, 0 °C to r.t., 6 h, 95%; ii) NaH (10 equiv), THF, 55 °C, 48 h, 61% (overall 57% from 4 in two steps); (c) m-CPBA (2.0 equiv), CHCl$_3$, 0 °C to r.t., 13 h, 99%; (d) 500 °C, 0.04–0.05 Torr, 1 h, 46%; (e) Li, $\delta$-propylene oxide, C$_6$H$_{14}$, r.t., 6 h, 89%.

Figure 3. The X-ray crystal structure of 1. (a) ORTEP drawing of the side view set at 50% probability level. (b) ORTEP drawing of the top view. (c–e) The crystal packing as a space-filling model, along a, b and c axes, respectively. (P,P)-1 enantiomer colored in gray and (M,M)-1 enantiomer colored in red.
UV-vis Absorption and Fluorescence Spectroscopy

UV-vis absorption and fluorescence spectra were measured in order to investigate the effect of the molecular configuration of 12 consecutively fused benzene rings of 1 on its electronic transitions (Figures 4 and S7–S9). The steady-state absorption and emission spectra revealed the basic optical properties of the final product possessing [8]helicene subunit in its structure. Infinitene 1 exhibits three main absorption bands with palatable, fine vibronic features. The first major band is located between 260 and 320 nm with a shoulder extending to 370 nm, showing two distinctive peaks at \( \lambda_{\text{abs}} = 294 \) and 311 nm with the molar extinction coefficients \( (\varepsilon) \) of 95,500 M\(^{-1}\)cm\(^{-1}\) and 59,000 M\(^{-1}\)cm\(^{-1}\), respectively. The second band corresponding to the lower-energy transitions, found between 380 and 420 nm, also shows two peaks at \( \lambda_{\text{abs}} = 389 \) nm \( (\varepsilon = 14,800 \text{ M}^{-1} \text{cm}^{-1}) \) and 412 nm \( (\varepsilon = 15,500 \text{ M}^{-1} \text{cm}^{-1}) \). Finally, the third weak band, which is located to the longest wavelength absorption is located between 440 and 490 nm with the lowest-energy transition at \( \lambda_{\text{abs}} = 481 \) nm \( (\varepsilon = 1,350 \text{ M}^{-1} \text{cm}^{-1}) \).

Infinitene 1 is a weak green light emitter with the maximum of emission located at \( \lambda_{\text{em}} = 505 \) nm (Figures 4 and S9). The Stokes shift was determined as 24 nm (988 cm\(^{-1}\)). Infinitene 1 displays bathochromic shift of the major absorption band (ca. 30 nm) with respect to both [8]helicene\(^{25}\) and [12]helicene.\(^{26,27}\) Emission curve of 1 is red-shifted by ca. 50 nm when compared to that of [8]helicene \( \lambda_{\text{em}} = 454 \) nm (22,050 cm\(^{-1}\))\(^{28}\), and its emission maximum very closely matches with that of [12]helicene \( \lambda_{\text{em}} = 510 \) nm (19,600 cm\(^{-1}\))\(^{28}\). Finally, infinitene 1 exhibits slightly higher fluorescence quantum yield \( (\Phi_f) \) of 0.038 than those of [8]helicene (0.014) and [12]helicene (0.0061).

![Figure 4](image)

**Figure 4.** Absorption (blue line) and emission (red line) spectra of 1 measured in CH\(_2\)Cl\(_2\). Absolute fluorescence quantum yield \( (\Phi_f) \) determined with an integrating sphere.

Computational Study

To gain insight into the structure, \( \pi \)-conjugation, photophysical properties and strain energy of 1, we performed computational studies by the density functional theory (DFT) method with a Gaussian 16 program.\(^{29}\) Among various basis sets, the PBE0\(^{11}\) functional well reproduced the structure obtained in X-ray crystallographic analysis (see Figure S14), and the structure optimization by PBE0/6-311+G\( (d,p) \) gave the most reasonable structure having a \( \pi \)-\( \pi \) stacking distance (3.192 Å) and the closest C–C distance (2.954 Å) (Figure 3a). Using this optimized structure, a NMR spectrum was calculated at the GIAO-DFT\(^{29/30}\)/B3LYP\(^{30}\)/6-311+G\( (2d,p) \) level of theory in CHCl\(_3\) with a SMD\(^{31}\) solvent model. In the experimental \(^1\)H-NMR spectrum of infinitene 1 (Figure 5a), there are six doublet peaks at \( \sigma \) (ppm) = 6.43, 6.99, 7.60, 8.04, 8.16, 8.18, whose values match well to the calculated chemical shifts (ppm) 6.48 \((\text{H}_d)\), 7.04 \((\text{H}_e)\), 7.65 \((\text{H}_f)\), 8.05 \((\text{H}_g)\), 8.22 \((\text{H}_h)\) and 8.23 \((\text{H}_i)\) (Figure 5b). It is reasonable that the only hydrogen atoms H\(_d\) and H\(_e\) were shielded due to an effect from ring current on a lower benzene ring. The calculated NICS\(^{32}(0)\) and NICS\(^{1}(1)\) values of representative three benzene rings are shown in Figure 5c, and a terminal benzene ring A indicated a slightly weakened aromatic nature compared to rings B and C. Considering the bond alternation found in the crystallographic structure and the computational studies, infinitene 1 has two equivalent Clar structures with six aromatic sextets and six double bonds (Figure 5d). Thus, all peripheral K-regions on infinitene 1 seem to have double bond nature like that on phenanthrene, implicating a potential of post-functionalization by annulative \( \pi \)-extension (APEX\(^{30}\)) reactions or direct arylation\(^{31}\) for further extended nanocarbon structures. Through a calculation of hypothetical homodesmotic reaction of 1 with benzene affording non-strained phenanthrene, the value of 60.2 kcal-mol\(^{-1}\) was estimated as a strain energy of 1 (Figure 5e).

![Figure 5](image)

**Figure 5.** (a) Experimental \(^1\)H NMR spectrum of infinitene 1 in the aromatic region (from 6.0 to 8.5 ppm) recorded in CDCl\(_3\). (b) \(^1\)H NMR chemical shift calculations. (c) NICS\(^{0}\) values of 1. NICS\(^{1}\) values are shown in parentheses. (d) Possible Clar structures for 1. (e) Determined strain energy of 1.

We next performed time-dependent (TD)-DFT calculations at the B3LYP/6-311++G\( (d,p) \) level of theory for infinitene 1 geometry-optimized at the PBE0/6-311+G\( (d,p) \) level to elucidate six frontier molecular orbitals and absorptions (Figure 6). Each molecular orbital (MO) is found to be delocalized on entire molecule. While HOMO (\(-5.41 \text{ eV}\)) and HOMO–1 (\(-5.76 \text{ eV}\)) are delocalized only on benzene surface, the orbitals on intersectional center
naphthalene rings in LUMO (−2.21 eV) and LUMO+1 (−2.04 eV) are clearly overlapped. Existence of these through-space interactions is in good agreement with relatively low-lying energy levels of LUMO and LUMO+1. TD-DFT calculation revealed possible energy absorptions at a longer wavelength region: transitions from HOMO−1 to LUMO (2.57 eV, λ = 481 nm, oscillation strength (σ) = 0.0001), HOMO to LUMO (2.60 eV, λ = 477 nm, f = 0.0113), HOMO to LUMO+2 (2.76 eV, λ = 450 nm, f = 0.0024) and HOMO−1 to LUMO (2.89 eV, λ = 430 nm, f = 0.1226) were elucidated. The observed longest-wavelength absorption at 481 nm (Figure 4) can be assigned as forbidden HOMO−1 to LUMO and HOMO to LUMO+1 transitions, whereas weak absorption at 449 nm can be contributed by HOMO to LUMO+2 and HOMO−1 to LUMO transitions.

Chiral Separation and Chiroptical Properties

Intrinsic chirality of the helically twisted 1 prompted us to separate its respective enantiomers by a chiral HPLC. After extensive investigation, we found the optimal conditions for efficient separation of (P,P)-1 and (M,M)-1 enantiomers. The racemic mixture rac-1 was successfully separated by a chiral HPLC (DAICEL CHIRALPAK IE column; eluent: n-hexane/CH2Cl2 = 35:65) (Figure 7 inset and S10). The circular dichroism (CD) spectra of both enantiomers were recorded in CH2Cl2. The CD spectrum measured for the first fraction showed positive Cotton effect peaks at 239, 339 and 375 nm, whereas negative Cotton effect peaks were apparent in the range of 250–300 nm and at 415 nm (Figures 7, S11 and S12). The CD spectrum of the second fraction displayed perfect mirror symmetry, corroborating its opposite configuration. The intensities of the main bands of the molar circular dichroism (Δε) with the local maxima located at 265, 339, 375 and 415 nm, as well as the anisotropy factors (gCD) of the corresponding transitions were distinctly smaller than values reported for unsubstituted [n]helicenes, e.g. (P)-[6]helicene shows (Δε) values of −267 M−1 cm−1 at 246 nm and +259 M−1 cm−1 at 324 nm.69 The aforementioned molar circular dichroism values ([Δε]) and anisotropy factors ([gCD]) measured for 1 reached accordingly: 265 nm: [Δε] = 17.8 M−1 cm−1, [gCD] = 3.5 × 10−1; 339 nm: [Δε] = 52.5 M−1 cm−1, [gCD] = 1.8 × 10−3; 375 nm: [Δε] = 69.0 M−1 cm−1, [gCD] = 4.8 × 10−1; 415 nm: [Δε] = 43.8 M−1 cm−1, [gCD] = 2.9 × 10−1; 457 nm: [Δε] = 10.4 M−1 cm−1, [gCD] = 3.4 × 10−3 and 484 nm: [Δε] = 3.0 M−1 cm−1, [gCD] = 2.3 × 10−3. The lower values of [Δε] and [gCD] are in agreement with the general trend set by the substitution effects on the chiroptical properties of helicenes, which are affected by a sophisticated interplay between geometric and electronic factors.60

Concurrently, unlike large values of optical rotation reported for pristine [n]helicenes, for example (P)-[6]helicene: [α]20D = +3707° (CHCl3)61 and (P)-[7]helicene: [α]20D = +5577° (CHCl3),62 infinitene 1 shows significantly smaller values of [α]20D = +1130° (ε = 8.9 × 104 M−1 cm−1 in CHCl3) in the case of the first fraction (99.98% ee) and [α]20D = −1112° (ε = 8.6 × 104 M−1 cmCHCl3) for the second fraction (99.90% ee). As a rule of thumb in the chemistry of carbohelicenes, levorotary (−) enantiomers tend to adopt left-handed (M) configuration, whereas dextrorotary (+) compounds are mainly, right-handed (P).63 In addition, TD-DFT calculation of both enantiomers also supported that the first fraction (blue line) and the second fraction (red line) were expected to be be (P,P)-1 and (M,M)-1, respectively (Figure S15).

Recently, circularly polarized luminescence (CPL) of helicene derivatives were reported with rational designs for higher dissymmetry factor (gCPL),41 and some of related figure-eight-shaped molecules show 102 to 103 order values of gCPL, whose values are relatively higher than simple carbohelicenes.41 Stimulated by these examples, CPL of both enantiomers of 1 was also measured (Figures 8 and S13). (P,P)-1 and (M,M)-1 displayed −6.2 × 103 and 6.4 × 103 of gCPL, which are reasonable values when considering the same order value of [gCD]. Although infinitene 1 did not show extremely high values of [gCD] and the structure-CPL ability relationship is still unclear, we believe that our new example of shape-persistent, symmetric and fully-conjugated, infinity-shaped molecule will help further molecular design to increase the gCPL value.
CONCLUSIONS

In summary, we have accomplished the synthesis of infinitene 1 (cyclo[cccccccc]dodecakisbenzene), a unique, fully fused, helically twisted [12]circulene topoisomer. We have studied its fundamental properties experimentally as well as computationally. The intrinsic chirality of 1 enabled successful separation of the enantiomers by a chiral HPLC resulting in an investigation of their chiroptical properties. The infinity-shaped geometry of 1 was unequivocally confirmed by X-ray diffraction analysis. Its crystal structure revealed significant compression of the central [8]helicene subunit, manifested by a close distance of 3.152-3.192 Å between two π-π stacked benzene rings. Combined lamellar and herringbone-like crystal packing suggested three-dimensional electronic interactions. CD and CPL measurements revealed that 1 has moderate $g_{\pi \pi}$ and $g_{\sigma \pi}$ values as same as other figure-eight-shaped molecules. These results open the new chapter in the chemistry of fully-fused, looped polyarenes. The established successful synthetic strategy would be an incentive to envision and create new amazing forms of nanocarbons whose architectures are limited only by our imagination.

ASSOCIATED CONTENT

Supporting Information
Experimental procedures, 1H and 13C NMR spectra, characterization of data for all new compounds, optical properties, computational data and crystallographic data of 1.

Accession codes
CCDC 2113525 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.


In this paper, we use the Staub-Vögtle-Diederich nomenclature for cycloarones. See ref 1, 2, and 2d for the definition.


Infinitene

A helically twisted figure-eight [12]circulene topoisomer (C_{48}H_{24})