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Lemniscular [16]Cycloparaphenylene: a Radially Conjugated Figure-Eight Aromatic Molecule

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ABSTRACT: A cycloparaphenylene-based molecular lemniscate (CPPL) was obtained in a short synthesis involving masked p-phenylene equivalents. The strained figure-eight geometry of CPPL is sustained by the incorporated 9,9'-bicarbazole subunit, which also acts as a stereogenic element. The shape of the distorted [16]cycloparaphenylene nanohoop embedded in CPPL is accurately approximated with a Booth lemniscate. The structure of CPPL, investigated using NMR and Raman spectroscopic methods, revealed strain-dependent features, consistent with the variable curvature of the ring. The electronic and optical properties of CPPL combine features more characteristic of smaller cycloparaphenylenes, such as a reduced optical bandgap, and red-shifted fluorescence. CPPL was resolved into enantiomers, which are configurationally stable and provide strong chiroptical responses, including circularly polarized luminescence.

INTRODUCTION

The topology and distortion of a cyclic π-electron system have a fundamental influence on its properties. In the most common case, found e.g. in benzene, the π-conjugated framework, consisting of an array of sp²-hybridized centers, has a planar structure with the constituent pₓ orbitals in perfectly parallel alignment (Figure 1A). By gradual distortion of such π systems, it is possible to produce molecules with radial (in-plane) conjugation, in which the pₓ orbitals are perpendicular to a cylindrical surface (Figure 1B). This type of conjugation is found in various tubular aromatics, notably [n]cycloparaphenylenes ([n]CPPs), carbon nanobelts, and nanotube endcaps. Planar and radial aromatics can be represented respectively by an annulus and an open cylinder, which are topologically equivalent. Since these surfaces are orientable, the corresponding π systems follow the Hückel aromaticity rules in the ground state. Radial π conjugation is therefore fundamentally related to planar π conjugation and has indeed been shown to sustain macrocyclic (global) Hückel aromaticity in charged [n]CPP rings. When the π system is twisted along the perimeter, its topology is altered, resulting in the increase of the so-called linking number. The latter parameter, denoted Lk, is expressed in units of π (i.e. 180°) and corresponds to the number of half-twists introduced into the aromatic surface. Such twisted aromatics alternate between ground-state Hückel and Möbius aromaticities, for even and odd Lk values, respectively. Among these systems, molecules with Lk = 1 and Lk = 3 have aroused particular interest since the first example of a Möbius-aromatic hydrocarbon was reported in 2003 by Herges et al.
Surfaces with $Lk = 2$ have two edges that are interlocked in space. Aromatic molecules with this topology adopt figure-eight (lemniscular) shapes, which help maximize $\pi$-orbital overlap and reduce internal strain. Lemniscular aromatics reported to date, mostly porphyrinoids\cite{25,26} and some rare carbocyclic frameworks,\cite{15,18} have been invariably characterized by locally planar in-plane conjugation in lemniscular aromatics (Figure 1D) may provide fundamental insight into the intricate relationship between three-dimensional structure and $\pi$-electron conjugation. This task is however challenging because of the unfavorable strain of such constructs, which requires an effective structural design and needs to be introduced by appropriate synthetic methodology.\cite{27} Here we show that it is possible to combine radial conjugation with a double twist of the $\pi$ surface in an appropriately bridged [16]CPP derivative. The resulting system adopts a chiral lemniscular shape while retaining a contiguously conjugated macrocyclic circuit.

RESULTS AND DISCUSSION

Synthesis. The target cycloparaphenylene lemniscate (CPPL, 1), was obtained from 9,9’-bicarbazole tetraboronate 3 and Jasti’s U-shaped building block 4\textsuperscript{1} in a concise three-step synthesis (Scheme 1). Carbazole-2,7-diyd building blocks have been previously incorporated into nanohoop structures, to produce rigidified yet circularly shaped CPP derivatives.\cite{28} The octamethoxy intermediate 6, obtained using the Yamamoto coupling in a double macrocyclization step, was structurally characterized and found to possess the requisite figure-of-eight structure, albeit with incomplete $\pi$ conjugation (Scheme 1). The macrocyclic loops in 6 are relatively unstrained, as evidenced by the nearly planar geometries of all benzene rings. Reductive aromatization of the four masked $p$-phenylene units in 6 was performed using the method reported by the Yamago group.\cite{29} Under these conditions, 6 was cleanly converted into the CPPL product 1, which was isolated in an 85% yield as a yellowish powder.

Molecular Structure and Dynamics. The structure of CPPL was unambiguously determined using one- and two-dimensional NMR spectroscopy, mass spectrometry, and a range of optical methods, with additional support of theoretical calculations. The lowest-energy $D_{2d}$-symmetrical conformer of 1 found in the gas phase using density functional theory (DFT) is shown in Figure 2. Variable-temperature $^1$H NMR spectra of 1, analyzed with the aid of correlation methods, are consistent with such a figure-eight structure possessing internal degrees of freedom (Figure 3 and Supporting Information). At 300 K in dichloromethane-$d_2$, the spectrum contains a unique ABCD spin system (gg’hh’), which indicates that the macrocycle is indeed twisted and that the corresponding phenylene rings (D, cf. Scheme 1) do not rotate within the oligophenylene loop. Each of the other phenylene groups, B and C, is represented by an AA’BB’ spin system, consistent with fast internal rotations. Ring C produces a
sharp ABCD pattern at 200 K in CDCl$_3$F (ee'ff'), whereas the signals of ring B (cc'dd') reach the coalescence point at ca. 160–170 K. These observations clearly show that the rate of phenylene rotations for rings B–D decreases significantly with the increasing local curvature of the loop. Protons located closer to the concave surface of the loop (b', e', f, g, and h) give resonances shifted to higher field, reflecting the increased magnetic shielding inside the lemniscate.

**Figure 2.** Lowest-energy conformation of \((P)\text{-1}\) determined in DFT calculations (ωB97XD/6-31G(d,p), in vacuo). (A) Side view and key geometrical parameters: mean boat angles, interplanar angles, and mean CCCC torsion angles are shown in blue, black, and red, respectively. (B) Pictorial representation of the π electron system, showing that the blue and red parts are interlocked. (C) Numerical fitting of the carbon framework of CPPL with a Booth lemniscate. Projections of several \([n]\)CPP rings are shown for comparison.

In addition to the differential decoalescence described above, some $^1$H NMR signals of \(\textbf{1}\) revealed temperature-dependent changes of chemical shifts. These variations are relatively small (up to ca. 0.1 ppm in the 160–220 K range in CDCl$_3$F) and may reflect an equilibrium between several conformations that differ in the relative orientations of some phenylene rings, but are nearly degenerate energetically. In such a case, \(\textbf{1}\) would exist as a mixture of rapidly exchanging conformers even at 160 K, while still producing a spectrum with effective $D_s$-symmetry. In spite of this presumed conformational inhomogeneity, almost all of the $^{13}$C NMR signals of \(\textbf{1}\) could be assigned using correlation spectroscopy (CDCl$_3$F, 200 K, Supporting Information). Importantly, the corresponding chemical shifts yielded a satisfactory correlation with the theoretical GIAO shieldings. The latter result provides additional evidence in support of the proposed structure of \(\textbf{1}\).

The curvature of the cycloparaphenylene belt contained in CPPL varies along the periphery. Rings D are most strongly curved, with the mean boat angle of 12.2°, comparable with the value of 12.7° determined for [6]CPP.\(^{30}\) The boat angles decrease for rings C (8.5°) and B (3.3°), and the carbazole units are nearly planar. The dihedral angle between carbazole planes (70.9°) is smaller than observed in the crystal structures of \(\textbf{2}\) (85.6–89.3°), \(\textbf{3}\) (79.2°), and \(\textbf{6}\) (90.4° inside the loop, Scheme 1). Nevertheless, the scissor-like compression of the bicarbazole unit is relatively small in \(\textbf{1}\). Mean torsional angles between consecutive phenylenes show alternating signs in the regions of low curvature (rings A–B–C), a typical feature found in \([n]\)CPP structures. The high-curvature regions (rings C–D–D–C) are characterized by uniform signs of phenylene torsions, corresponding to the helical shape of the loop. In spite of the significant overall twist of the macrocycle in CPPL, the largest interphenylene torsions do not exceed 40°, i.e. are within the range observed experimentally for circular CPP systems. The π surface of \(\textbf{1}\) is therefore characterized by fairly effective and contiguous overlap of $p$ orbitals along the macrocycle. Thus, CPPL and similarly designed systems can be expected to sustain macrocyclic conjugation in their charged states, in analogy to CPP nanohoops.\(^{11–13}\) \(\textbf{1}\) resembles some recently reported multi-loop CPP derivatives,\(^{31–33}\) but it is the first such system containing a contiguous \([n]\)CPP substructure. Because of the large interplanar angle between the carbazoles, the π system is effectively decoupled along the N–N linkage, which thus serves only as a steric restraint necessary to support the lemniscular conformation.

**Figure 3.** $^1$H NMR spectra of \(\textbf{1}\) at two different temperatures (600 MHz). Labeling is defined in Scheme 1. Additional NMR data are given in the Supporting Information.

The lemniscular structure of \(\textbf{1}\) is characterized by a linking number of $|L_k| = 2$, corresponding to a double half-twist of the π system.\(^{14}\) The latter feature can be verified...
by noting that the two sides of the π surface (red and blue in Figure 2B) are interlocked in space, forming the so-called Hopf link.\(^1\) The linking number \(Lk\) can be decomposed into twist (\(Tw\)) and writhe (\(Wr\)) according to the Călugăreanu formula, \(Lk = Tw + Wr\).\(^1\) The twist parameter \(Tw\) was calculated for 1 as the cumulative rotation of POAV\(^3\) vectors along the backbone of quaternary \(p\)-phenylene carbons. The resulting value of \(|Tw| = 0.478\) is lower than typically observed in figure-eight aromatics with planar conjugation,\(^1,15\) and indicates a high level of \(p\)-orbital overlap in the π-conjugated system. Interestingly, when the DFT structure of 1 is projected onto a plane along the N–N bond, the network of carbon atoms can be very accurately approximated by noting that the two sides of the Booth lemniscate.\(^36\) In polar coordinates, the latter curve is defined as \(r^2 = 4b(a - b \sin^2 \theta)\), and the best fit for CPPL is obtained with \(a = 4.07\ \text{Å}\) and \(b = 11.57\ \text{Å}\), which correspond to a lemniscate with lateral dimensions of 27.4 Å × 8.1 Å (Figure 2C).

The solid-state Raman spectrum of 1, in combination with a theoretical normal-mode analysis (Figure 4), provides an important insight into the curvature and strain of CPPL. The CC stretching modes of the benzene rings are observed in the 1629–1576 cm\(^{-1}\) range. The 1576 cm\(^{-1}\) mode correlates with those determined for [6]- and [7]CPP (1567 cm\(^{-1}\) and 1574 cm\(^{-1}\), respectively),\(^37\) indicating that a fragment with a similar degree of quinoidal character may be present in CPPL. The highest frequency band in this range (1629 cm\(^{-1}\)) corresponds mostly to stretches in the carbazole units and agrees well with the corresponding stretch in free carbazole (1625 cm\(^{-1}\)). In the low wavenumber region, the experimental bands at 509 and 476 cm\(^{-1}\) were assigned to C–H out-of-plane deformation vibrations. Four additional bands (404.5, 364, 204, and 169/159 cm\(^{-1}\)), were identified as pseudo radial breathing modes (p-RBMs), i.e., out-of-plane CCC modes with their motion amplitudes along the radial direction of the loop, analogous to those found in [\(n\)]CPPs.\(^37\) The previously established empirical dependence between the p-RBM frequency and internal strain in CPPs produced an estimated strain energy of 122.5 kcal/mol for CPPL. A direct homodesmotic calculation, performed in analogy to a previously reported scheme,\(^39\) yielded a strain enthalpy of \(\Delta H = 102.7\) kcal/mol for CPPL, in very good agreement with the above empirical prediction (Supporting Information).

Electronic Structure. In dichloromethane, 1 exhibits a single absorption peak at 357 nm (Figure 5A, \(\varepsilon = 142\ 000\ \text{M}^{-1}\ \text{cm}^{-1}\)), somewhat red-shifted in comparison with [16]CPP (339 nm).\(^40\) The spectrum of 1 features a long-wavelength shoulder, which extends up to ca. 450 nm and is responsible for the pale-yellow color of the compound. The greenish fluorescence emission of 1 has a maximum at 496 nm and a quantum yield of \(\Phi_F = 0.36\). A similar emission band, with a more clearly resolved vibronic structure, was observed in 2-methyltetrahydrofuran at 80 K (Figure S2). As a fluorophore, CPPL differs markedly from the cycloparaphenylene series, with which it is structurally related. Large \([n]\)CPPs \((n > 9)\) have been found to emit strongly from the \(S_2\) state, whereas smaller nanohoops emit primarily from \(S_1\), albeit with progressively diminishing quantum yields.\(^1,13,41–44\) The large apparent Stokes shift observed for CPPL suggests that its emission is "S\(_2\)" in spite of the large size of the embedded nanohoop. Indeed, photoluminescence properties of 1 differ significantly from those of the parent [16]CPP \((\lambda_{em} = 415\ \text{and} 438\ \text{nm}; \Phi_F = 0.88)\). The position of the emission maximum is closest to that reported for [9]CPP (494 nm), which is however a stronger emitter (\(\Phi_F = 0.73\)).\(^1,40\) The emission lifetime of 1 (2.92 ns) is longer than that of [16]CPP (ca. 0.9 ns) and matches those of smaller CPPs \((n \sim 10–11)\),\(^45\) reflecting the expected influence of increased curvature in CPPL on exciton dynamics.
Figure 5. (A) Electronic absorption spectrum (solid line) and emission spectrum of 1 (dashed line, both in dichloromethane). (B) Electronic circular dichroism spectra of (P)-1 and (M)-1 (298 K, ca. 10^{-6} M in dichloromethane). (C) Circularly polarized luminescence spectra of (P)-1 and (M)-1 (ca. 10^{-6} M in dichloromethane) at 298 K (\lambda_{\text{exc}} = 350 \text{ nm}), \Delta f = l_{n}-k_{n}.

Kohn–Sham HOMO and LUMO levels of CPPL, obtained from DFT calculations, are both non-degenerate and their nodal characteristics are analogous to those of [16]CPP (Figure 6). At the B3LYP/6-31G(d) level of theory, the bandgap of CPPL is 3.35 eV, i.e. it is smaller than the calculated gap of [16]CPP (3.66 eV). In comparison, the dicarbazole nanohoop 1-H_2, which is structurally intermediate between [16]CPP and CPPL, is predicted to have an energy gap essentially identical with that of its CPP parent. In both 1 and 1-H_2, HOMO and LUMO amplitudes at the nitrogen atoms are small, indicating that the perturbation of frontier energy levels caused by the heteroatoms is insignificant. Thus, since the conjugation length is identical in [16]CPP and CPPL, the reduced optical bandgap of the latter species can be mostly attributed to its higher curvature. The lower symmetry of CPPL (D_2) in comparison with the circular CPPs results in a more complex pattern of electronic transitions, as determined in time-dependent (TD) DFT calculations (Figure 6). The lowest-energy transition of 1 is dominated by the HOMO–LUMO excitation and is weakly allowed (f = 0.03). It is followed by three other weak transitions (S_2-S_3, f = 0.01–0.10), and a very strong one (S_3, f = 3.56). This pattern differs from the spectra of [n]CPPs, which feature a dipole-forbidden S_1 transition, followed directly by the very strong S_2. The partially allowed character of the lowest excited states may contribute the relatively strong S_1-type fluorescence emission of 1, which is more typical of smaller CPP rings (n < 10).

Figure 6. Top: Kohn–Sham HOMO and LUMO orbitals for CPPL and its analogues (0.01 a.u. isosurfaces, energies in eV). Bottom right: lowest energy electronic transitions determined using time-dependent DFT. Arrows indicate principal excitations. Dotted arrows correspond to f < 0.10. All calculations were performed at the B3LYP/6-31G(d) level of theory.

The first oxidation of 1 observed in dichloromethane using voltammetric methods has limited chemical reversibility (Supporting Information). The oxidation potential of 0.64 V vs. Fc/Fc+ is substantially lower than the values reported for large CPPs (ca. 0.9 V), and correlates with the shift of the HOMO level discussed above. The first reduction of 1 was observed at –2.36 V, yielding an electrochemical gap of 3.00 eV, in good agreement with the gaps derived from optical absorption and TD-DFT.

Since a 64-electron macrocyclic circuit can formally be distinguished in the structure of 1, the molecule might in principle be expected to exhibit Hückel-antiaromatic (4n-electron) characteristics. However, the purely benzenoid framework of CPPL, which is not cross-conjugated with the central N–N unit, produces a conjugation pattern that is accurately described in terms of 16 Clar sextets, in analogy to other cycloparaphenylene systems. Such a formulation is reflected in the magnetic properties of 1 determined using ^1H NMR, and also in the computationally modeled bond length patterns, obtained at various levels of theory (Supporting Information), which show that \pi bond delocalization is confined to six-membered rings. Quinoidal features, caused by curvature
of individual benzene units, can nevertheless be traced in the Raman spectra of 1, as discussed above.

**Chirality.** CPPL is a chiral molecule, in which the twisted 9,9′-bicarbazole unit acts as the stereogenic element. CPPL is thus different from other chiral CPP-related systems, in which enantiomer differentiation was achieved via locked internal rotation of desymmetrized aromatic subunits, or by introduction of sp³ stereocenters. The oligophenylene loops in 1 are apparently not essential for configurational stability, as suggested by the atropisomerism of the naturally occurring dixiamycins A and B, which also contain the 9,9′-bicarbazole moiety.

Enantiomers of 1 were precisely resolved by means of supercritical fluid chromatography on a chiral stationary phase. According to the convention previously adopted for figure-eight aromatics, the enantiomer of CPPL shown in Scheme 1 and Figure 2 is labeled (P)-1, to reflect the local helicity of the constituent oligophenylene loops. At room temperature, the enantiomers of 1 do not racemize in solution, but their stability upon heating could not be reliably verified because of gradual decomposition. According to DFT calculations, inversion of the lemniscate should occur via a C₃-symmetrical transition state with a predicted in-vacuo barrier of ΔG°,298 = 51.4 kcal/mol. This high value implies that enantiomers of 1 should be configurationally stable even at elevated temperatures.

The figure-eight shape of the CPPL chromophore makes it of interest as a chiroptical material. The electronic circular dichroism (ECD) spectra of pure CPPL enantiomers each contain two major Cotton effects of opposite signs (Figure 5B). The more intense peak (∆ε = 300 M⁻¹·cm⁻¹) is found at 360 nm, whereas the other is located at 390 nm and approximately coincides with the shoulder observed in the absorption spectrum. An additional weaker Cotton effect is found at ca. 250 nm (∆ε = 100 M⁻¹·cm⁻¹). The absolute configuration of P and M enantiomers of 1, indicated in Figure 5B, was established on the basis of TD-DFT calculations employing B3LYP and CAM-B3LYP functionals (Supporting Information). In particular, in the CD spectrum calculated using B3LYP/6-31G(d), the highest rotatory strengths corresponded to the S₁ and S₂ transitions. The two enantiomers yielded mirror-image circularly polarized luminescence (CPL) spectra with dissymmetry factors [gₜₐₜ] of around 3.7·10⁻³ at 498 nm, i.e. in the range typically observed for organic CPL emitters. Interestingly, this gₜₐₜ value is close to the dissymmetry factor g₁₂ observed in the ECD spectra of 1 (g₁₂ = 4·10⁻³ at 400 nm), suggesting that both the ground state and the emitting excited state of 1 have similar chiral geometries.

**CONCLUSIONS**

CPPL, the lemniscal analogue of [16]cycloparaphenylene described in this work is the first example of a radially conjugated π-electron system with a double half-twist. The structural design of CPPL provides a unique opportunity to observe the effect of curvature variations on the electronic structure in isolation from other influences. The electronic properties of this new system differ from those of the parent [16]CPP ring, showing absorption and emission features more characteristic of smaller cycloparaphenylenes. The twisted nanohoop of CPPL can be resolved into configurationally stable enantiomers, which are found to produce circularly polarized luminescence. More generally, the approach to curvature control presented herein enables reduction of electronic bandgaps while retaining large conjugation lengths in nanohoop systems. Additionally, by creating small loops with chiral interiors the new design is of interest for its potential use in supramolecular chemistry. Efforts to explore these new possibilities are currently underway in our laboratories.

**ASSOCIATED CONTENT**

**Supporting Information**

Synthetic and spectroscopic details, computational data. The Supporting Information is available free of charge on the ACS Publications website.

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**REFERENCES**


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