A Co-conformationally “Topologically” Chiral Catenane

Arnau Rodríguez-Rubio, Andrea Savoini, Florian Modicom, Patrick Butler, and Stephen M. Goldup*

ABSTRACT: Catenanes composed of two achiral rings that are oriented (C_{nh} symmetry) because of the sequence of atoms they contain are referred to as topologically chiral. Here, we present the synthesis of a highly enantioenriched catenane containing a related but overlooked “co-conformationally ‘topologically’ chiral” stereogenic unit, which arises when a bilaterally symmetric C_{nv} ring is desymmetrized by the position of an oriented macrocycle.

Topology is the study of the properties of objects and networks that are preserved under deformations that do not break connections/surfaces or require surfaces/edges to pass through one another. Chemical topology applies these ideas to molecules. At the simplest level, constitutional isomers are topologically distinct, as they differ in the network of atoms. More interesting topological isomerism arises when structures contain identical atomic connections, the most famous examples of which are Möbius ladders (isomers of the untwisted macrocycle), molecular knots (isomers of the unknotted ring), and [2]catenanes (isomers of two non-interlocked rings). These structures have nonplanar graphs in that there is no two-dimensional projection of their structures in which bonds do not cross over one another and this property is topologically invariant in three-dimensional space — no matter how the structure is distorted, even drastically altering the geometry around atoms, a planar graph cannot be achieved.

Such topologically nontrivial structures can display chirality in the absence of covalent stereogenic units. Depending on their topology, Möbius ladders and molecular knots can be chiral as a result of the pattern of bond crossing points. Although [2]catenanes do not display unconditional topological stereochemistry, as recognized by Wasserman and Frisch, they can be chiral as a result of the constitutional symmetry of the rings; rings that are “oriented” (C_{nh} symmetry) due to the sequence of atoms in the cycle give rise to topologically chiral catenanes (Figure 1a). The absolute stereochemistry of topologically chiral objects is invariant under all topologically allowed deformations in three-dimensional space.

We recently identified “missing” stereogenic units that arise in interlocked molecules and give rise to classes of chiral rotaxanes and catenanes that had yet to be discussed or synthesized. An example that presents particular linguistic problems are [2]catenanes in which one ring is oriented (C_{nh}) and the other is bilaterally symmetric (e.g., C_{2v}) (Figure 1b). The time averaged structure of such catenanes is achiral, but any co-conformation in which the oriented ring does not lie on the internal mirror plane of the C_{2v} ring is chiral. If the structure is designed such that the oriented ring is permanently prevented from occupying said mirror plane, the molecule will display kinetically fixed molecular chirality (Figure 1c).

As with related co-conformational-covalent and co-conformational mechanical planar stereochemistry in rotaxanes, this stereogenic unit can be considered to appear due to the oriented ring acting as a substituent of the region of C_{2v} ring that it encircles, effectively reducing its symmetry to C_{1h}. Thus, this stereogenic unit arises because one ring is oriented due to its constitution and the other by the molecular co-conformation and so we have previously provisionally termed such molecules “co-conformationally “topologically” chiral” to clearly make the link with the established stereogenic unit of topologically chiral catenanes while also highlighting that the stereochemistry of the system is clearly not topologically invariant.

Semantic arguments aside, we set out to synthesize an enantioenriched co-conformationally “topologically” chiral
The majority of enantioenriched topologically chiral catenanes, which was then extended to a co-conformationally chiral target.

The stereoselective synthesis of a co-conformationally chiral catenane requires (i) the oriented ring to be incorporated at a defined position around the C2v macrocycle and (ii) the oriented ring to be installed stereoselectively. The first requirement can be met by forming the mechanical bond such that the oriented ring is trapped between bulky groups. The second is the same problem as encountered in the synthesis of any topologically chiral [2]catenane. Although the majority of enantioenriched topologically chiral catenanes in which the mechanical bond is the sole source of stereochemistry have been accessed by chiral stationary phase HPLC (CSP-HPLC) separation, recently developed an auxiliary approach in which a chiral covalent auxiliary directs the stereoselective formation of the mechanical bond. However, in this proof-of-concept synthesis, the stereoselectivity of the mechanical bond formation was low (dr ∼ 2:1), which required the mechanical epimers to be separated prior to removal of the auxiliary, limiting the utility of this methodology for more complicated targets. To overcome this challenge, we set out to extend a phenylalanine-based auxiliary, developed for the synthesis of mechanically planar chiral rotaxanes, to the synthesis of topologically chiral [2]-catenanes.

Tyrosine-derived pre-macrocycle (S)-1a was synthesized (96% ee, Figure S40) and reacted under pseudo high-dilution active template Cu-mediated alkene−azide cycloaddition (AT-CuAAC) conditions with bipyridine macrocycle 2. Catenane 3a was produced with reasonable stereoselectivity (Table 1, entry 1), based on 1H NMR analysis of the crude reaction product; proton Hα of the diastereomers of 3a resonate at 8.98 (major) and 9.07 (minor) ppm, respectively (Figure S111). 1H NMR analysis also suggested the presence of several other interlocked species, characterized by higher ppm (9.51−9.61; Figure S286) triazole resonances. LCMS analysis indicated that these signals were due to [3]catenane 4 (Scheme 1), which can be formed as three diastereomers, and the corresponding [2]catenane (not shown, two diastereomers) containing a single bipyridine ring (Supporting Information (SI) section S10). We were unable to obtain pure samples of these compounds.

Longer addition times (entry 2) resulted in diminished diastereoselectivity, perhaps due to epimerization of the covalent stereogenic center, and lower conversion of macrocycle 2. Lowering the reaction temperature resulted in enhanced diastereoselectivity (74% de) and reduced quantities of oligomeric species, allowing catenane 3b to be isolated in 39% yield and 74% de (entry 3). Although increasing the equivalents of 1a resulted in higher conversion of 2, lower yields of 3a were obtained as the non-interlocked triazole-containing macrocycle was challenging to remove. Varying the solvent did not improve diastereoselectivity or conversion of 2 (SI section S8). Applying the same conditions to (S)-1b, which features a bulkier Pr ester, gave catenane 3b in 82% de, albeit the conversion of macrocycle 2 was diminished and the formation of oligomeric biproducts was increased, resulting in a low isolated yield (26%, 82% de, entry 4). Surprisingly, (S)-1c gave poor stereoselectivity (68% de, entry 5) and low conversion of 2 (~25%). Pleasingly, single crystal X-ray diffraction (SCXRD) analysis of a racemic sample of catenane 3b produced using rac-1b allowed the relative stereochemistry of the major diastereomer to be tentatively assigned as (S*). Thus, the major product of (S)-1b and macrocycle 2 is assigned as (S,S*,S*)-3b (Figure 2a).

We then turned to methods to remove the covalent stereogenic unit from the mixture of catenane 3b diastereomers (Scheme 2). Attempts to ablate the covalent stereocenter of a model compound by radical decarboxylation met with failure due to scission of the triazole N1−C substituent bond (SI section S9). Ultimately, we found that reduction of ester 3b to give alcohol catenane 5 followed by tandem Oppenauer-type oxidation/RhI-mediated decarbonylation gave rise to catenane 6 in reasonable isolated yield.

**Table 1. Effect of Reaction Conditions and Structure of 1 on the AT-CuAAC Synthesis of Topologically Chiral Catenanes**

| entry | R | T (°C) | t (h) | 2:3:oligos | de | yield |
|-------|---|--------|------|------------|----|-------|
| 1     | Et| 60     | 4    | 34:44:22   | 70%| n.d.  |
| 2     | Et| 60     | 8    | 47:37:16   | 62%| n.d.  |
| 3     | Et| 60     | 25   | 15:44:41   | 74%| 39%   |
| 4     | Pr| 60     | 25   | 14:30:56   | 82%| 26%   |
| 5     | Bu| 60     | 25   | 77:11:12   | 68%| n.d.  |

* Determined by 3H NMR analysis of the crude reaction product (SI section S10).
* Not isolated due to low conversion of 2.
a mixture of diastereomers (88% ee). The product, topologically chiral [2]catenane, was subjected to the AT-CuAAC reaction with macrocycle 2b.

Crystals of a rac-6 suitable for SCXRD analysis were obtained, allowing the structure of the product to be confirmed (Figure 2b).

Finally, we turned to the synthesis of a co-conformationally “topologically” chiral target (Scheme 3). Pre-macrocycle (S)-7 was subjected to the AT-CuAAC reaction with macrocycle 2. The product, topologically chiral [2]catenane 8, was isolated as a mixture of diastereomers (88% ee), as judged by 1H NMR (Figure 3a). By analogy with catenane 3b, which seems reasonable given the similarities of the functional groups reacting and the similar stereoselectivity obtained, the major isomer is tentatively assigned as (S,S<sub>mt</sub>-8).

Auxiliary removal from (S,S<sub>mt</sub>-8) (88% ee) yielded [2]-catenane 9, which contains no previously described stereogenic units—it lacks covalent stereogenic units, and the triazole containing ring is not oriented and so the system does not conform to the definition of a topologically chiral catenane. Nevertheless, whereas the compounds produced from 3b and (S,S<sub>mt</sub>-8) produce identical 1H NMR spectra (Figure 3a and 3ai respectively), the latter is clearly highly enantioenriched, whereas the former is racemic as judged by CSP-HPLC analysis (Figure 3b). The major stereoisomer of 8 was assigned as (S<sub>mt</sub>-8) based on the assigned stereochemistry of the major diastereomer of 3b. Crystals of a rac-6 suitable for SCXRD analysis were obtained, allowing the structure of the product to be confirmed (Figure 2b).

In conclusion, we have developed an auxiliary for the synthesis of topologically chiral catenanes in high enantiopurity and applied it to the synthesis of catenane (S,S<sub>mt</sub>-9), a molecule containing a previously unreported co-conformationally “topologically” chiral stereogenic unit, unambiguously demonstrating the chiral nature of this overlooked form of mechanical stereochemistry. However, it poses a problem of nomenclature—how can the topological stereochemistry of a molecule depend on its co-conformation? In short, it cannot, but once the fixed co-conformation is considered, the covalent subcomponents of catenane 9 display the same symmetry properties as those that comprise the established stereogenic unit of topologically chiral catenanes, which leads to our linguistic conundrum. One solution to this would be to rename “topologically chiral” catenanes as “mechanically planar chiral”, to bring them in line with the analogous rotaxanes to which they are strongly related, but this would require further discussion in the field. Linguistic issues aside, chiral interlocked molecules are attracting increasing attention for applications in catalysis, sensing, and as chiroptical or stereodynamic switches. By highlighting their potential to display molecular chirality due to unexplored stereogenic units, we hope to inspire further investigation of their rich stereochemistry.
(b) HPLC analysis of catenane

Figure 3. (a) Partial $^1$H NMR (CDCl$_3$, 298 K) of i. catenane 8, ii. catenane rac-9, and iii. enantioenriched catenane ($S_{enanti}^9$)-9. Atom labels and colors as in Scheme 3, except macrocycle signals (blue). (b) HPLC analysis of catenane rac-9 and ($S_{enanti}^9$)-9. (c) Circular dichroism spectra of catenane rac-9 and ($S_{enanti}^9$)-9. (d) Solid state structure of rac-9 showing a pair of enantiomeric structures related by a point of inversion (orange). Colors as in Scheme 3 except O (gray), N (dark blue), H (white). Majority H atoms omitted for clarity.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c02029.

Procedures and full characterization data (NMR, MS, CD, SCXRD, HPLC as appropriate) for all novel compounds and supplementary discussion. (PDF)

Accession Codes

CCDC 2125552, 2129422, and 2129424 contain 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.

AUTHOR INFORMATION

Corresponding Author

Stephen M. Goldup — Chemistry, University of Southampton, Highfield, Southampton SO17 1BJ, United Kingdom;

orcid.org/0000-0003-3781-0464; Email: s.goldup@soton.ac.uk

Notes

The authors declare no competing financial interest.

Data (characterization data for reported compounds) is available from the University of Southampton data repository (https://doi.org/10.5258/SOTON/D2279).

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(27) Catenanes of the form of 4 contain two covalent stereogenic units and two topological stereogenic units as both the central and peripheral rings are oriented. See SI section S6 for a more detailed discussion.

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(30) That catenane 9 is formed from 8 at high temperature, but the stereopurity of the starting material matches closely with the product is consistent with co-conformational motion being completely blocked. In keeping with this, heating a purified sample of 9 in mesitylene at 170 °C for 24 h did not result in any loss of stereopurity (Figure S260).

(31) We note that that SCXRD data for rac-9 are poor due to it crystallizing as very thin needles (see SI section S7). However, it is sufficient to confirm the connectivity of the product and the lack of any covalent stereogenic unit in the structure.

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