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Isolation of a chiral anthracene cation radical: X-ray crystallography and computational interrogation of its racemization

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Abstract

Chiral cation-radical salts hold significant promise as charge-transfer materials, chiroptical switches, and electron-transfer catalysts for enantioselective synthesis. Herein we demonstrate that the readily-available chiral 9,10-diphenyleanthracene derivative (*i.e.*^s**ANT**) forms a robust cation radical, whose structure was elucidated by X-ray crystallography and DFT calculations. While ^s**ANT** was observed to racemize on a timescale ($t_{1/2}$) of 1.1 hours, a computational conformational search and kinetic analysis of the racemization pathway led

us to identify a simple methyl substituted ^{*s*}**ANT** derivative, which does not racemize (racemization $t_{1/2}$ 10¹³–10¹⁷ years).

Introduction

There is continuing interest in identifying chiral molecules, which form stable cation radicals for applications in electron-transfer catalysed enantioselective synthetic transformations,¹ as chiroptical molecular switches,² in electro-magneto-chiral anisotropy,³ and for enhancing conducting properties under an applied magnetic field,⁴ among others. For example, it was shown for several radical-cation salts that enantiopure materials show enhanced conductivity when compared with the corresponding racemic mixtures owing to the structural disorder.^{5,6}

Unfortunately, examples of stable enantiopure cation radicals are scarce.^{2,5,6} As polycyclic aromatic hydrocarbons (PAHs) are best suited for isolation of stable cation radicals, the axially chiral PAHs^Z are ideal candidates for chiral cation-radical salts. For example, Agranat and coworkers^{8,9} reported that McMurry coupling of the readily-available dibenzo[b,f]suberone unexpectedly afforded a chiral doubly-bridged 9,10-diphenylanthracene in excellent yield (eqn (1)). It was concluded that this reaction produced only chiral 'synanthracene' (^sANT) isomer as a racemic mixture – not the achiral (meso) 'anti-anthracene' (^sANT). Careful chromatographic separation and racemization studies showed that ^sANT racemizes at ambient temperatures despite an activation barrier of ~23 kcal mol⁻¹.^{8.9}



Our continuing interest in design and synthesis of stable cation radicals^{10–15} as charge-transport materials for modern photovoltaic applications prompted us to examine if ^s**ANT** forms a stable cation radical, and whether racemization be prevented by a modification of its structure. In this communication, we will show that this readily available chiral anthracene derivative forms a robust cation-radical salt, which can be isolated and characterized by X-ray crystallography. A detailed conformational analysis of racemization of ^s**ANT** by DFT calculations and kinetic analysis of racemization led us to design an improved ^s**ANT** derivative that will not be subject to racemization under ambient conditions. The details of these preliminary findings are described herein.

Initial indication of the stability of the ^s**ANT** cation radical was apparent from completely reversible cyclic voltammograms at scan rates varying from 50 to 400 mV s⁻¹ in dichloromethane (Fig. 1). The measured electrochemical oxidation potential ($E_{ox} = 0.46$ V vs. Fc/Fc⁺ as an added internal standard) ^s**ANT** was 360 mV lower than 9,10-diphenylanthracene—a non-encumbered analogue of ^s**ANT**.



Fig. 1 Cyclic voltammograms of 1 mM ^S**ANT** at different scan rates (50 to 400 mV, as indicated) in CH₂Cl₂ containing 0.2 M tetra-n-butylammonium hexafluorophosphate at 22 °C.

The cation radical of ^s**ANT** can be generated in solution via quantitative redox titrations using [**THEO**⁺'SbCl₆⁻] (**THEO** = 1,2,3,4,5,6,7,8-octahydro-9,10-dimethoxy-1,4:5,8-dimethano-anthracene, $E_{red1} = 0.67 V vs. Fc/Fc^+$, $\lambda_{max} = 518 nm$, $\varepsilon_{max} = 7300 cm^{-1} M^{-1}$)^{<u>17</u>} and [**NAP**^{+'}SbCl₆⁻] (**NAP** = cycloannulated naphthalene derivative, $E_{red1} = 0.94 V vs. Fc/Fc^+$, $\lambda_{max} = 672 nm$, $\varepsilon_{max} = 9300 cm^{-1} M^{-1}$)^{<u>18,19</sub> as oxidants</u>. For example, Fig. 2A displays the electronic spectra obtained upon an incremental (sub-stoichiometric) addition of a concentrated CH₂Cl₂ solution of ^s**ANT** to a solution of [**THEO**^{+'}SbCl₆⁻] at 22 °C. A complete consumption of the oxidant (i.e. **THEO**^{+'}) and clean formation of ^s**ANT**^{+'}, after the addition of 1 equivalent of neutral ^s**ANT**, was ascertained by a quantitative deconvolution^{<u>20</u>} of the component spectra of various species present at each titration point (Fig. 2B). A plot of the mole fractions of various species, obtained from Fig. 2B, against the added equivalents of ^s**ANT** established a 1 : 1 stoichiometry of the redox reaction, i.e. ^s**ANT** + **THEO**^{+'} \rightarrow ^s**ANT**^{+'} + **THEO** (Fig. 2C). The ^s**ANT**^{+'} spectra were identical when generated using **NAP**^{+'} as oxidant (Fig. S3–S5 in the ESI⁺). Also note that the spectra of ^s**ANT**^{+'} remained unchanged at tenfold higher concentration or at temperatures between 22 °C to $-50^{\circ}C.\frac{21}{2}$}



Fig. 2 (A) Spectral changes observed upon the reduction of 0.049 mM **THEO**⁺⁺ in CH₂Cl₂ (3 mL) by addition of 4 μ L increments of 3.6 mM solution of ^S**ANT** in CH₂Cl₂. (B) Clean deconvolution of each spectrum from figure **A** into its component spectra, i.e.**THEO**⁺⁺ (burgundy), ^S**ANT**⁺⁺ (blue), and ^S**ANT** (grey), with negligible residual (<0.1%). (C) Plot of the mole fractions of **THEO**⁺⁺ (burgundy) and ^S**ANT**⁺⁺ (blue) against the added equivalents of neutral ^S**ANT**.

Symbols represent experimental points, while the solid lines show best-fit to experimental points using $\Delta G_1 = E_{red}(THEO^{+}) - E_{ox}(^{S}ANT) = 0.2 \text{ V}.^{20}$

The stability of $[{}^{s}ANT^{+}SbCl_{6}^{-}]$ was further attested by its quantitative isolation by precipitation using diethyl ether from a reaction of ${}^{s}ANT$ (0.25 mmol) with NO⁺SbCl₆⁻ (0.25 mmol) in CH₂Cl₂ (10 mL) at 0 °C (see Section S3 in the ESI⁺ for details). The X-ray quality single crystals of $[{}^{s}ANT^{+}SbCl_{6}^{-}]$ were obtained by a careful layering of its CH₂Cl₂ solution with toluene and storage in a refrigerator (~-10 °C) for 48 h. The determination of the precise X-ray structures of $[{}^{s}ANT^{+}SbCl_{6}^{-}]$ as well as of neutral ${}^{s}ANT$ allowed a detailed elucidation of the structural changes induced by 1–e⁻ oxidation, which was compared with theory.

The ORTEP diagrams of racemic ^SANT and ^SANT⁺⁺ as well as their crystal packing diagrams are compiled in Fig. 3A–D.[‡] The enantiomers of both neutral and cationic ^SANT pack in separate stacks; and the stacks of individual enantiomers of ^SANT⁺⁺ and SbCl₆⁻ counter ions form mixed layers along the ab plane. No contacts less than the sum of van der Waals radii amongst the ^SANT⁺⁺ moieties in its crystals were observed; and the anthracene fragment of both ^SANT and ^SANT⁺⁺ were significantly distorted from planarity (see Fig. S16 in ESI⁺). The dihedral angle between phenyl and anthracene core showed a negligible change (~0.5°) upon 1–e⁻ oxidation due to the rigidity of bimethylene bridges.¹⁶



Fig. 3 The ORTEP diagrams (50% probability) of ^sANT (A) and [^sANT⁺SbCl₆⁻] (B), and the packing diagrams of neutral (C) and cation-radical salt (D) of ^sANT.

A comparison of the bond length changes in the ^sANT \rightarrow ^sANT^{+*} transformation showed that the bond contractions and elongations were mostly confined to the anthracene core, and the 9,10-phenyl groups did not undergo significant changes, suggesting that the spin/charge is largely confined to the anthracene core (Fig. 4). Furthermore, a plot of the bond length changes in ANT \rightarrow ANT^{+*} transformation obtained by X-ray crystallography against those obtained by DFT calculations clearly shows a linear correspondence (Fig. 4C). Indeed, the oxidation-induced bond length changes in ^sANT^{+*} closely followed the disposition of the bonding/antibonding lobes of HOMO, i.e. bonds with bonding HOMO lobes undergo elongations, whereas the bonds with antibonding lobes undergo contractions (compare Fig. 4A and C).²²



Fig. 4 Isovalue plots of HOMO of neutral **ANT** (A) and spin density of **ANT**⁺⁺ (B) [B1LYP-40/6-31G(d)+PCM(CH₂Cl₂)]. Comparison of the oxidation-induced C–C bond length changes in ^S**ANT** obtained by X-ray crystallography and DFT calculations with a linear trend line y = 0.79x + 0.03, R² = 0.71. Error bars correspond to the standard deviation of the bond length changes ($\sigma = 0.8$ pm). (C) The bond length changes are based on the averages of equivalent bonds.

Building upon this success, we then undertook a detailed conformational analysis at the B1LYP-40/6- $31G(d)+PCM(CH_2CI_2)$ level of theory to examine why the racemization of ^sANT occurs at ambient temperatures, despite a relatively high energy barrier. We also examined whether this racemization barrier is altered upon oxidation to cation radical.

A careful search in the conformational space showed that enantiomer of ^s**ANT** first converts to ^A**ANT** by flipping one of its phenyl rings, while the second remains fixed (Fig. 5, above). This interconversion proceeds through a highly twisted transition state ($\Delta G^{\ddagger} = 27.6 \text{ kcal mol}^{-1}$, **TS1**), in which the rotation of a single phenyl group accompanies a significant puckering of anthracene core and prompts an inversion at both bridging methylenes. As the inversion of methylenes and rotation of the phenyl group progresses, the puckering in the anthracene reduces (see superimposed structures in Fig. 5), producing a high-energy intermediate ($\Delta G = 12.3 \text{ kcal mol}^{-1}$, **I**), which then transforms to achiral ^A**ANT** through an almost barrierless process ($\Delta G^{\ddagger} = 2.0 \text{ kcal mol}^{-1}$, **TS2**). The calculations predict that ^A**ANT** lies 2.8 kcal mol⁻¹ higher than ^s**ANT** (Fig. 5) and the racemization barrier of 27.6 kcal mol⁻¹ is in fair agreement with the barrier of 22.6 kcal mol⁻¹ obtained by experiment.^{8.9}



Fig. 5 IRC (Intrinsic reaction coordinate) pathway of the ^SANT(R,R) \rightleftharpoons ^SANT(S,S) racemization and the structures and Gibbs free energies (in kcal mol⁻¹) of transition states (**TS1**, **TS2**) and intermediates (**I**) along the pathway. Superimposed structures (^SANT, **TS1**, **I**, **TS2** and ^AANT) showing the rotation of single phenyl group during ^SANT \rightarrow ^AANT transformation.

Interestingly, racemization of a model compound (**M**) containing only one bridged phenyl group follows an uncannily similar pathway with a similar activation barrier of 24.5 kcal mol⁻¹ (Fig. S6 in the ESI⁺), signifying that the second phenyl group is not involved in the ^s**ANT** \Rightarrow ^A**ANT** interconversion. However, once the meso^A**ANT** is formed, the flipping of the second phenyl ring via an identical mirror-imaged pathway completes the conversion of original ^s**ANT** into its enantiomer. Thus, the racemization of ^s**ANT** proceeds via achiral ^A**ANT** as an intermediate, i.e.eqn (2).

Racemization of neutral and cation radical of ^s**ANT** follows identical pathways, and their activation barriers differ only by 1.2 kcal mol⁻¹ (see Fig. S7 in the ESI⁺), suggesting that the structural changes in ^s**ANT**^{+*} induced by 1–e⁻ oxidation have a little impact on the conformational switching. Also, the energy difference between ^s**ANT**^{+*} and ^A**ANT**^{+*} of 2.2 kcal mol⁻¹ is similar to that observed in the neutral analogues (i.e. 2.8 kcal mol⁻¹).

Despite a high estimated racemization barrier (22.6 kcal mol⁻¹), the enantiopure ^S**ANT** undergoes racemization with a half-life (i.e. $t_{1/2}$) of 1.1 hours at ambient temperatures, as determined using the Eyring equation (see Section S5 in the ESI⁺).⁹ As the reaction half-life exponentially depends on the activation barrier, the rate of racemization can be controlled or even halted by increasing the activation barrier. In order to assess the necessary increase of the activation barrier required to prevent racemization, we performed a kinetic analysis of ^S**ANT**(S,S) interconversion process.

Simulated time-evolutions for the formation of racemic mixture of ^sANT(R,R)/^sANT(S,S) from pure (meso)^AANT or enantiopure ^sANT was obtained from numerical solution of the system of the first-order reactions from eqn (2) (see Section S5 in the ESI⁺). The simulations predict that ^AANT quickly converts into equal amounts of ^sANT(R,R) and ^sANT(S,S) with the reaction half-life of ~20 seconds (Fig. 6A), while the enantiopure ^sANT has a reaction half-life of 1.1 hours (Fig. 6B).



Fig. 6 Simulated mole fraction plots of ^S**ANT**(R,R), ^S**ANT**(S,S) and ^A**ANT** with respect to time with initially (A) only (meso)^A**ANT** and (B) enantiopure ^S**ANT** using $\Delta G^{\ddagger} = 22.6$ kcal mol⁻¹. (C) Plot of the racemization reaction half-lives (t_{1/2}) against the activation barrier (ΔG^{\ddagger}) from Table 1. Note that kinetic simulations in (A) and (B) show little difference on the racemization half-lives when the high-energy intermediate (I) is included in the kinetic analysis and can therefore be excluded for sake of simplicity (see Fig. S15 in the ESI⁺).

The two-order of magnitude difference in half-lives (compare Fig. 6A and B), dependent on the starting reactant, cannot be simply accounted for by the presence of two reaction channels in case of ^AANT, i.e.^sANT(R,R) \leftarrow ^AANT \rightarrow ^sANT(S,S), as it accounts only for an increase by a factor of 2. Rather, the energy difference of 2.8 kcal mol⁻¹ between ^sANT and ^AANT contributes to lowering of the activation barrier for ^AANT \rightarrow ^sANT transformation and is largely responsible for the reduced half-life (Fig. 6C). Interestingly, a simple inspection of Fig. 6C suggests that the doubling of the activation barrier for ^sANT \rightarrow ^sANT transformation (22.6 kcal mol⁻¹) would increase the racemization half-life by a factor of 10¹⁶ (Fig. 6C). As the ^sANT(R,R) \rightleftharpoons ^sANT(S,S) interconversion requires a rotation of the phenyl rings, this can be most effectively impeded by placing a substituent at the ortho position of phenyl groups, i.e. the boxed structure in Fig. 7.



Fig. 7 Structure of four derivatives of ^SANT.

Consistent with this expectation, calculated racemization barriers for various methyl substituted (**B**, **C**, **D**) ^s**ANT** derivatives (Table 1) shows that the substitution on the phenyl rings (i.e.**C**) produces a much higher activation barrier (51 kcal mol⁻¹) than when the substitution is on anthracene core (**B**, 34 kcal mol⁻¹).

Table 1 Computed activation energies of racemization (ΔG^{\ddagger} , kcal mol⁻¹), energy difference between ^sANT and ^AANT conformations (ΔG_{anti} , kcal mol⁻¹) and racemization half-lives ($t_{1/2}$) of ^sANT derivatives. All ^sANT derivatives

	Neutral			Cation radical		
	∆G⁺	ΔG_{anti}	t 1/2	ΔG	ΔG_{anti}	t 1/2
s ANT (exp) ⁹	22.6	n/a	1.1 h	n/a	n/a	n/a
^s ANT	27.6	2.8	5.3 × 10³ h	26.4	2.4	696 h
Μ	24.5	0.0	28 h	22.8	0.0	1.6 h
А	26.9	2.5	1.6 × 10³ h	26	2.1	354 h
В	34.3	3.1	4.9 × 10⁴ year	34	2.4	3.0 × 10⁴ year
С	51.2	4.1	1.2 × 10 ¹⁷ year	42.2	3.0	3.0 × 10 ¹⁰ year
D	56.7	2.6	1.3 × 10 ²¹ year	50.9	1.0	7.2 × 10 ¹⁶ year

follow a racemization pathway similar to that of parent ^sANT (Fig. S6–S11 in the ESI). See Table S16 in ESI for racemization half-lives scaled to reproduce experimentally observed half-live of ^sANT

Examination of the **B** and **C** transition state structures reveals the considerable structural distortion required for the methyl substituted rings (i.e. anthracene core in case of **B** and phenyl group in case of **C**) during the rotation of the phenyl group (see Fig. S12 in the ESI⁺). It is well known that the energy needed to distort PAHs such as anthracene is rather modest as compared to the distortion of a single aromatic ring.²³ Indeed, methyl substitutions both at phenyl groups and anthracene (i.e.**D**) increases the barrier minimally (57 kcal mol⁻¹) as compared to **C** (51 kcal mol⁻¹). Note also that substitution of ^s**ANT** by remote t-butyl substituents (**A**) does not affect the activation barrier (27 kcal mol⁻¹).

Although the ^sANT derivatives **B** or **C** or **D** have sufficiently large racemization barriers to halt racemization on a practical timescale (Table 1), structure **C** is the most viable candidate for synthetic pursuit rather than **B** and **D** because the peri substitution of anthracene core raises the energies of **B** and **D** by 11 kcal mol⁻¹ as compared to **C** owing to the steric crowding (see Fig. S13 in the ESI⁺).

In conclusion, we have shown that the ^s**ANT** scaffold is well suited for the formation of a stable cation radical, by isolation and structural characterization via X-ray crystallography. Unfortunately, ^s**ANT** racemizes with the half life of ~1 hour. A systematic conformational and kinetic analysis of the racemization pathway led us to identify a ^s**ANT** derivative, containing a methyl group on each phenyl group, which is not subject to racemization on a practical timescale. Efforts are underway for the preparation of this minimally altered ^s**ANT** derivative for the isolation and study of a new class of enantiopure cation radicals.

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⁺ Electronic supplementary information (ESI) available: Full experimental details for ^sANT and [^sANT^{+*}SbCl₆⁻⁻] and computational details. CCDC 1525472 and 1525473. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6cc10307c

‡ Crystal structure data for SANT [C30H22] (raj25q): FW = 382.48, P21/c, a = 7.64722(7) Å, b = 10.95523(14) Å, c = 22.9991(3) Å, β = 94.4658(10)°, Z = 4, V = 1920.95(4) Å3, D = 1.323 g cm-3, T = 100 K, 3895 reflections measured, 3466 unique reflections, Rint = 0.0259, 271 parameters refined, R(all) = 0.0370, wR(all) = 0.0975, S = 1.045 (CCDC 1525472). Crystal structure data for [SANT+'SbCl6-] [C30H22 SbCl6] (raj25s): FW = 716.93, Fdd2, a = 29.7613(10) Å, b = 18.2178(6) Å, c = 10.1451(3) Å, α = 90°, β = 90°, γ = 90°, Z = 8, V = 5500.5(3) Å3, d = 1.731 g cm-3, T = 100 K, 2540 reflections measured, 2368 unique reflections, Rint = 0.0794, 168 parameters refined, R(all) = 0.0389, wR(all) = 0.0966, S = 1.046 (CCDC 1525473).