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# Isolation and X-ray Structural Characterization of Tetraisopropylpyrene Cation Radical

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### Abstract

A practical synthesis of 1,3,6,8-tetraisopropylpyrene and the isolation and X-ray structural characterization of its monomeric cation radical salt are described.

Pyrene has been extensively utilized as a fluorescent probe for studying the conformational changes in proteins in solution owing to its long excited-state lifetime and characteristic excimer formation due to its self association.<sup>1</sup> The ability of pyrene to self associate into ordered structures has also been exploited in its usage as liquid crystalline materials.<sup>2</sup> In stark contrast, it is this ability of  $\pi$ - $\pi$  stacking and excimer formation in solution and solid state that has limited the usage of pyrene as an emissive material in organic light-emitting diodes (OLED's) and related applications.<sup>3</sup>

Although, pyrene undergoes selective and quantitative bromination at the 1,3,6,8-positions to produce the corresponding tetrabromopyrene,<sup>4</sup> attempted electrophilic substitutions to functionalize the pyrene core at the 1,3,6,8-positions with isopropyl groups, and thereby sterically congesting the flat fluorophore,<sup>5</sup> led to complex mixtures of polyalkylated pyrenes from which the separation of the desired isomer was rather tedious.<sup>6</sup>

Our continuing interest in the design and syntheses of stable organic cation radicals or hole carriers, which are of fundamental importance to organic materials science,<sup>2</sup> led us to obtain an efficient and scalable route for the preparation of 1,3,6,8-tetraisopropylpyrene (3) and demonstrate the efficacy of sterically hindered tetraisopropylpyrene for the isolation and X-ray crystallographic characterization of its monomeric cation-radical salt. The details of these findings are described herein.

After several failed attempts, in our hands, to synthesize **3** by Friedel–Crafts alkylation,<sup>56</sup> we resorted to a different approach as summarized in Scheme 1. Thus, a 4-fold Suzuki coupling between the readily-available 1,3,6,8-tetrabromopyrene,<sup>4</sup> and isopropenylboronic acid,<sup>8</sup> afforded cleanly the 1,3,6,8-tetraisopropenylpyrene (**2**) in >92% isolated yield. A catalytic hydrogenation of **2** over 10% Pd-C in a mixture of ethyl acetate–ethanol furnished **3** in quantitative yield. It was noted that under these hydrogenation conditions, the reduction of the pyrene core was not observed.



Scheme 1 Synthesis of 1,3,6,8-tetraisopropylpyrene (3).

The absorption spectrum of **3** in dichloromethane was characteristically similar to that of the parent pyrene with a modest bathochromic shift (~30 nm) owing to the alkyl substitutions. Furthermore, unlike the parent pyrene<sup>2</sup> which showed a broad featureless excimer emission centered at ~472 nm in the concentration range studied (0.05–0.1 M), the tetraisopropylpyrene **3**showed only the monomeric emission (Fig. 1). As such, the observation of monomeric emission for **3** suggests that the bulky isopropyl groups exert sufficient steric

inhibition to prevent a face to face approach of  $\sim$ 3.5 Å necessary for an efficient  $\pi$ -stacking<sup>10</sup> and consequent excimer formation (Fig. 1, right).



**Fig. 1** Left: selected excitation (black lines) and emission (red lines) of **3** as dichloromethane solution ranging from a concentration of  $1.66 \times 10^{-6}$  M to  $1.0 \times 10^{-5}$  M. Right: calculated structures of dimeric pyrene and **3** using DFT at B3LYP-631G\* level.

Next with sufficient quantities of the tetraisopropylpyrene at our disposal, it was subjected to electrochemical oxidation at a platinum electrode as a  $2 \times 10^{-3}$  M solution in dichloromethane containing 0.2 M *n*-Bu<sub>4</sub>NPF<sub>6</sub> as the supporting electrolyte. The cyclic voltammograms (Fig. 2, left) consistently met the reversibility criteria at various scan rates of 100–500 mV s<sup>-1</sup>, as they all showed cathodic/anodic peak current ratios of  $i_a/i_c = 1.0$  (theoretical) as well as the differences between anodic and cathodic peak potentials of  $E_{pa} - E_{pc} = 70$  mV at 22 °C. The reversible oxidation potential of **3** ( $E_{ox} = 0.98$  V vs. SCE) was calibrated with added ferrocene as an internal standard ( $E_{ox} = 0.45$  V vs. SCE). It is important to note that under similar conditions as above, the parent pyrene undergoes an irreversible electrochemical oxidation at  $E_{ox} = 1.36$  V vs. SCE.



**Fig. 2** Left: cyclic voltammograms of  $2 \times 10^{-3}$  M **3** in CH<sub>2</sub>Cl<sub>2</sub> containing 0.2 M *n*-Bu<sub>4</sub>NPF<sub>6</sub> at scan rates between 100 and 500 mV s<sup>-1</sup>. Right: spectral changes upon the reduction of  $1.33 \times 10^{-5}$  M **MA**<sup>-+</sup> by incremental addition

of **3** in dichloromethane at 22 °C. Inset: a plot of increase in absorbance of **3**<sup>'+</sup> (monitored at 494 nm) against the equivalent of added **3**.

The electrochemical reversibility and relatively low oxidation potential of **3**, prompted us to generate its cation radical by chemical oxidation using a stable aromatic cation radical salt (MA<sup>+</sup> SbCl<sub>6</sub><sup>-</sup>;  $E_{red} = 1.11 V vs.$  SCE) as a one-electron oxidant.<sup>11</sup> Thus Fig. 2 (right) shows the spectral changes attendant upon an incremental addition of sub-stoichiometric amounts of tetraisopropylpyrene to a  $1.3 \times 10^{-5}$  M MA<sup>+</sup> [ $\lambda_{max}$  (log  $\varepsilon$ ) = 518 nm (3.86)] in dichloromethane at 22 °C. Furthermore a plot of formation of the tetraisopropyl pyrene cation radical (*i.e.* increase in the absorbance at 494 nm) against the increments of added neutral **3** (see inset of Fig. 2, right), established that MA<sup>++</sup> was completely consumed after the addition of 1 equiv. of **3**; and the resulting highly structured absorption spectrum of **3**<sup>++</sup> [ $\lambda_{max} = 330$ , 345, 362, 494 (log  $\varepsilon = 4.83$ ), 451 (sh), 590, and 695 nm] remained unchanged upon further addition of neutral **3** (*i.e.* eqn (1)).

$$3 + \underbrace{(1)}_{0} \xrightarrow{0} MA^{+}SbCl_{6} \xrightarrow{1} 3^{+}SbCl_{6} \xrightarrow{1} + MA (1)$$

The green-colored solution of the **3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup> is highly persistent and did not show any decomposition at room temperature during the course of 12 hours. Moreover, a reduction of a dichloromethane solution of **3**<sup>+</sup> with zinc dust regenerated neutral **3** quantitatively, which further lends support to the high stability of the **3**<sup>+</sup>. As in the neutral tetraisopropylpyrene which showed no signs of aggregation (*i.e.*Fig. 1), the formation of the dimeric cation radical resulting from the cofacial stacking of **3**<sup>+</sup> with neutral **3**, was not observed as judged by the singular absence of the charge-resonance transition in the NIR region in the presence of a large excess of **3**. Note that the parent pyrene cation radical readily forms a dimeric cation radical with characteristic absorption bands at 395 and 520 nm together with a broad charge-resonance transition centered at ~1600 nm.<sup>22</sup>

The high stability of the tetraisopropylpyrene cation radical in solution prompted us to isolate its crystalline salt by chemical oxidation using nitrosonium hexachloroantimonate as a  $1-e^-$  oxidant according to the stoichiometry in eqn (2).

$$3 + \text{NO}^+\text{SbCl}_6^- \xrightarrow{\text{DCM}} 3^{\bullet+}\text{SbCl}_6^- + \text{NO} \uparrow (2)$$

Thus, a solution of **3** in anhydrous dichloromethane was added to crystalline NO<sup>+</sup> SbCl<sub>6</sub><sup>-</sup> under an argon atmosphere at ~0 °C. The gaseous nitric oxide produced was entrained by bubbling argon through the solution to yield a green-colored solution, which upon spectrophotometric analysis indicated the formation of **3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup> (see Fig. 2). An excellent crop of dark-colored crystals, suitable for X-ray crystallographic studies, were obtained by a slow diffusion of toluene into the above solution of **3**<sup>+</sup> during a period of 2 days at -20 °C.

The crystal structure of  $3^{+}$  SbCl<sub>6</sub><sup>-</sup> revealed that cationic tetraisopropylpyrene moieties pack in a herringbone arrangement (see Fig. 3, left) with a pair of embedded toluene molecules.<sup>‡</sup> One of the two crystallographically independent toluenemolecules forms an individual 1:1 complex with  $3^{+}$  while the other toluene molecule and the counteranion (SbCl<sub>6</sub><sup>-</sup>) fill the space between the herringbone stacks of  $3^{+}$  (see Fig. 3, left).§



**Fig. 3** The ORTEP diagram of **3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup> cation radical salt (right), with the packing diagram (left) showing that the toluenemolecules are embedded between the herringbone stacks of **3**<sup>+</sup>. (The thermal ellipsoids are drawn with 55% probability.)

A closer look at the bond length changes in the cation radical **3**<sup>+</sup>, together with a comparison with its neutral form, the structure of which was established by X-ray crystallography, points to the following important observations: (i) one electron oxidation causes no perceptible change in the bonds marked **A** (d<sub>A</sub> 139 pm) and **E** (d<sub>B</sub> 142 pm). (ii) The increased aromatization of the two internal rings of the pyrene molecule on oxidation occurs by a simultaneous lengthening of the short external bonds **B**and **D** by 2.2 and 2.9 pm, respectively, and shortening of the adjacent long bonds **C** by 2.1 pm. Interestingly, the bonds which undergo most dramatic lengthening in **3**<sup>+</sup> (*i.e.* bonds **B** and **D**) are the bonds on which the HOMO resides, *i.e.*Fig. 4 (right). (iii) The central bond **F** undergoes a shortening of 1.8 pm in order to accommodate the changes in the bond lengths of various annulenic bonds (*i.e.***B**, **C**, and **D**).



**Fig. 4** Left: lettering scheme for the pyrene skeleton. Right: showing the localization of the HOMO of **3**, obtained by DFT calculations at the B3LYP-631G\*\* level, on bonds labelled **B** and **D**.

The experimental observations of the bond length changes upon 1-electron oxidation of **3** were found to be in reasonable agreement with the calculated values using DFT calculations at the B3LYP-631G<sup>\*\*</sup> level (see Table 1).<sup>33</sup>

**Table 1** Experimental and theoretical bond lengths of the neutral and cation radical of 3 presented in picometres(pm)

	B3LYP/6-31G**			X-Ray data		
Bond type	3	3 *	Δ	3	3.+	Δ
Α	139.6	139.5	-0.1	138.9 (	3) 138.6	(2) –0.3
В	141.7	144.0	+2.3	141.1 (	3) 143.5	(3) +2.4
С	143.5	142.4	-1.1	143.4 (	3) 141.2	(3) -2.2
D	135.9	138.0	+2.1	134.7 (	3) 137.5	(3) +2.8
E	143.4	143.0	-0.4	142.7 (	3) 142.4	(3) –0.3
F	144.0	143.1	-0.9	144.5 (	3) 142.7	(4) -1.8

In summary, a simple and practical synthesis of 1,3,6,8-tetraisopropylpyrene (3) has been accomplished from readily available precursors. The emission and absorption spectroscopy of the neutral and cationic 3 clearly show that the  $\pi$ -stacking is inhibited owing to the presence of bulky isopropyl groups. The isolation and X-ray crystal structure determination of 3'+ SbCl<sub>6</sub>-as well as DFT calculations provide unequivocal evidence that introduction of a cationic charge (or polaron) in 3 largely affects the bonds on which the HOMO resides. Studies are underway for a more comprehensive investigation of the steric modulation of the  $\pi$ -stacking in various polyaromatic hydrocarbons.

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#### Footnotes

<sup>+</sup> Electronic supplementary information (ESI) available: Synthetic details of **3** and procedure for the isolation of its cation radical. Crystallographic data for CCDC 674750 and 674751. See DOI: 10.1039/b800168e

‡ *Crystal structure data for***3**. A suitable crystal (0.20 × 0.18 × 0.06 mm<sup>3</sup>) of **3** was obtained from a mixture of dichloromethane– acetonitrile solution at 22 °C. MW = 370.55, triclinic, space group  $P\bar{1}$ , *a* = 11.3272 (7), *b* = 12.6764(7), *c* = 16.9660(12) Å, *α* = 94.595(4)°, *β* = 92.311(4)°, *γ* = 115.064(3)°, *D*<sub>c</sub> = 1.123 Mg m<sup>-3</sup>, *V* = 2192.2(2) Å<sup>3</sup>, *Z* = 4. The total number of reflections measured were 25 845, of which 7547 reflections were symmetrically non-equivalent. Also note that unit cell contained one molecule in a general position and two half molecules lying about their inversion centres. Final residuals were *R*1 = 0.0644 and w*R*2 = 0.1740 for 7547 reflections with *I* > 2*a*(*I*). *Crystal structure data for* [**3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup>, 2C<sub>7</sub>H<sub>8</sub>]. A suitable crystal (0.51 × 0.23 × 0.14 mm<sup>3</sup>) of **3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup> was obtained from a mixture of dichloromethane–toluene solution at −30 °C. MW = 889.27, orthorhombic, space group *Pnma*, *a* = 13.8054(3), *b* = 12.5572(3), *c* = 23.6942(6) Å, *α* = 90°, *β* = 90°, *P* = 90°, *D*<sub>c</sub> = 1.438 Mg m<sup>-3</sup>, *V* = 4107.56(17) Å<sup>3</sup>, *Z* = 4. The total number of reflections measured were 34 504, of which 3746 reflections were symmetrically non-equivalent. Final residuals were *R*1 = 0.0268 and w*R*2 = 0.659 for 3746 reflections with *I* > 2*a*(*I*). Note that all four components have crystallographically imposed mirror symmetry. CCDC numbers of complexes **3** and [**3**<sup>+</sup> SbCl<sub>6</sub><sup>-</sup>, 2C<sub>7</sub>H<sub>8</sub>] are 674750 and 674751.

§ Note that a cofacial arrangement between the toluene and 3<sup>+</sup> at an inter-planar separation of 3.5 Å may stabilize the cationic tetraisopropylpyrene *via* an electron-donor acceptor interaction. Also note that both the toluene molecules are rotationally disordered (within their respective molecular planes) with the occupations of the minor components being 23 and 37%.