Synthesis and Oxidation of Triarylamines Bearing Hydrogen-Bonding Groups

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Received February 15, 2007

Hydrogen-bonding triarylamines, 4-(N,N-bis(4-methoxyphenyl)amino)benzoic acid (TPA1), 5-(N,N-bis(4-methoxyphenyl)amino)isophthalic acid (TPA2), and N-(4-(1H-benzimidazol-2-yl)phenyl)-N,N-bis(4-methoxyphenyl)amine (BImTPA), were synthesized as radical cation precursors. TPA1 and TPA2 are readily p-doped by AgSbF6 to give highly persistent radical cations. Poor solid-state spin yields of the radical cation from BImTPA may be due to spin delocalization.

Cationic doping of triarylamines is an important strategy in organic-based magnetic materials.1 Hydrogen-bonding interactions are useful to help assemble molecules in the solid state with some degree of control, to try to control intermolecular exchange between unpaired electrons.2 However, hydrogen-bonded assembly has not been much used to our knowledge in p-dopable triarylamines, although 4-(N,N-bis(4-methoxyphenyl)-amino)benzoic acid, TPA1, was made as a synthetic intermediate for studies of photoactive and luminescent organometallic complexes.3 To our knowledge, the solid-state hydrogen bonding associated with the corresponding radical cations.

The transient spectroscopy of photooxidization and the chemical oxidation of TPA1 with Br2 show absorption bands at 750 and 590 nm that are attributed to the radical cation; these are in agreement with the present results. Part of the spectrum of BImTPA-rc is significantly perturbed by comparison, with doubling of a peak found around 350–360 nm in the other two systems (new peak at ~400 nm) and a red shift of the band at about 500–600 nm to about 630 nm. However, the lowest energy, strongest peak in all is at 780–800 nm (±1.55 eV).

Table 1 summarizes solution ESR hyperfine couplings for the radical cations generated by AgSbF6 oxidation of TPA solutions in dichloromethane at room temperature. TPA1-rc and TPA2-rc show hyperfine coupling (hfc) from the aryl protons, but any analogous hfc in BImTPA-rc was masked by line broadening, despite attempts in several solvents in the temperature range of 100–300 K. The ESR spectra of samples produced by SbCl5 vapor oxidation were essentially the same. Under no conditions or solvents attempted were ESR peaks observed that were clearly attributable to triplet state radical pairs, either in fluid solution or at 77 K in frozen solution.

Numerous attempts to crystallize BImTPA failed to yield single crystals of sufficient quality for X-ray diffraction (XRD) analysis. Its high melting point suggests strong intermolecular interactions. FTIR shows a broad NH stretching envelope over 3100–3300 cm⁻¹ that suggests NH···N hydrogen bonding (NH···N chain formation is common for 2-substituted benzimidazoles). TPA1 and TPA2 crystallize well from chloroform/methanol and were readily analyzed by single-crystal XRD.

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10.1021/jo070318a CCC: $37.00 © 2007 American Chemical Society
TPA2 also crystallizes well from THF, but this solvent is incorporated into the lattice.

Powdered TPA1, TPA2, or BImTPA exposed to SbCl₅ vapor for 24 h turned blue-black, blue-black, and green-black, respectively. The oxidized samples remained visually unchanged after exposure to room air for weeks. Measurements of the vapor-doped paramagnetic susceptibilities of powdered TPA1 and TPA2 showed 28 and 12% of the theoretical moments expected if all molecules yielded $S^{1/2}$ spin units (Supporting Information). Single crystals of TPA1 and TPA2 that were doped for 24 h showed pale yellow interiors when broken, proving that extensive diffusive doping does not occur readily from the vapor phase.

By comparison, dichloromethane solutions of TPA1, TPA2, or BImTPA treated with AgSbF₆ immediately turned deeply colored and could be precipitated into hexane to give dark colored powders that were readily studied by paramagnetic susceptibility (Figure 2). The data indicate 59, 79, and 4% yields of spin units from this procedure, relative to a theoretical value of 0.375 emu·K/Oe·mol for 100% generation of $S^{1/2}$ spin carriers, and assuming that the basic spin units are monoradical cations with SbF₆⁻ counterions.

Neutral TPA1 forms crystallographic dimers, a typical motif for monocarboxylic acids.⁴ TPA2 forms 1-D ribbons, as is common in isophthalic acids.⁴ In both, the aryl rings are significantly twisted relative to one another. All methoxy and some aryl groups show moderately large thermal motion ellipsoids at room temperature, but the molecules are not otherwise disordered. Although the hydrogen bonds hold the amine molecules closer together than might occur without the interaction, the crystallographic geometries found in the neutral amines appear unlikely to assist strong exchange. The closest N···N distances (Supporting Information) in TPA1 are 5.8–5.9 Å, and in TPA2, they are 6.74 Å, too large to allow strong, direct exchange due to direct overlap of N(2p) orbitals from the amonium radical cation centers. It seems likely that swelling of the lattice to accommodate dopant counterions would move the TPA radical cations further apart, although this was not directly probed. However, $\pi$-orbital contact between aryl units could also give exchange between radical cations, given their significant spin densities delocalized from the amonium centers. In TPA1, the $\pi$-carbon ipso to the carboxylic acid group is only 3.52 Å from a $\pi$-carbon ortho to the acid group on a neighboring molecule. For TPA2, inversion symmetry gives an antiparallel close contact of 3.36 Å between the phenyl–CO₂H bonds (Supporting Information). These contacts do not provide ideal $\pi$-system overlap for inter-radical exchange, but modest realignment during doping could easily improve them.

UB3LYP/6-31G* geometry optimized computations⁵ for the three radical cations were carried out in order to evaluate further their spin delocalization and to compare to the experimental ESR hfc. Solution-phase ESR spectra of TPA1-rc and TPA2-rc show aryl C–H hyperfine coupling consistent with the computational 6–9% and (−)2−4% spin populations on the phenyl $\pi$-carbons ortho and meta to the amonium nitrogen, respectively. These delocalized spin density sites could provide exchange sites where close contacts between doped radical cations occur in the solid state, depending on the intermolecular contacts between molecules in the doped form.
Unresolved benzimidazole hfc in BImTPA-rc probably gives heterogeneous line shape broadening that obscures ary1 CH hyperfine in its ESR spectrum. The UB3LYP/6-31G* computations support this, showing that (−)1.6% of spin density is delocalized onto the benzimidazole 2-C position, 5.4% onto the imine 3-N position, and about (±)1−6% onto the benzimidazole ary1 carbons. The significant shifts in some UV−vis bands of BImTPA-rc by comparison to those of TPA1-rc and TPA2-rc also reflect electronic conjugation between the radical site and the benzimidazole group.

Vapor-phase p-doping of neat powdered TPA1 and TPA2 with SbCl₅ gives a surface radical cation layer that persists for many weeks but does not fuse the surface bulk, as explained above. At low temperatures, the χ(T) data for vapor p-doped samples show weak antiferromagnetic (AFM) exchange (Supporting Information), so the amount of surface doping is sufficient to give exchange interaction between radical cation sites by proximity. The neutral amines TPA1 and TPA2 are held in sufficient proximity in their crystal lattices that their doped forms could show small to modest intermolecular exchange interaction, assuming that the dimer and chain hydrogen-bond motifs are maintained after doping.

Solution-phase doping of TPA1 and TPA2 with AgSbF₅ gives precipitated products with much higher spin yields. The exchange behaviors of these products are not necessarily related to those of the vapor-doped TPA samples, but they do also show χ(T) downturns at lower temperature, indicating AFM exchange interactions, and their 1/g(T) versus T plots give Weiss constants of (−)6.3 and (−)2.6 K, respectively. Solid powder samples and solutions of the radical cations generated this way are highly persistent under ambient conditions and retain their characteristic deep coloration for over a year at −30 °C. The carboxylic acid units at the para (TPA1-rc) and both meta (TPA2-rc) positions of the non-methoxylated phenyl rings are apparently effective steric blockading groups, limiting reactions such as radical dimerization or oxygen trapping. However, these samples were not readily subjected to further purification or recrystallization. Attempts at slow evaporation from dichloromethane and slow diffusion of hexane into dichloromethane did not yield crystals of sufficient quality for single-crystal X-ray diffraction analysis, so detailed magnetocrystalline analysis of the antiferromagnetic interactions in the radical cations is not possible at this time, but hydrogen bonding in these systems is a major part of bringing the molecules closer together in the neutral TPA systems and is likely to be an important contributor to holding the radical cations in proximity in the neat solids. Although BImTPA gives radical cations in amounts readily detected by ESR from both vapor and solution oxidation techniques, magnetic studies of products from both methods showed disappointingly poor spin yields of only 4−5%. Solutions of this radical cation also are not as color-persistent as those from TPA1 and TPA2. The radical cation BImTPA-rc is formed to a significant extent, however, based on the spectroscopic results described above, as well as cyclic voltammetry (CV). CV in CHCl₃ shows roughly quasi-reversible behavior, with E° values for (n,+) of 720, 640, and 560 mV at 10 mV/s sweep rate for TPA1, TPA2, and BImTPA, respectively. In order of ease of oxidation, BImTPA > TPA2 > TPA1. Since BImTPA is the most easily oxidized of the triarylamines in this study, its low solid-state spin yield may be due to increased reactivity in the solid. BImTPA-rc has spin delocalization onto its benzimidazole unit based on its UV and ESR spectroscopy, as well as the computational modeling results. This could cause solid-state dimerization and other reactions with consequent loss of spin moment.

Attachment of one carboxylic acid group to a TPA leads to solid-state dimer formation, while attachment of an isophthalic acid group leads to ribbon formation, both due to directional hydrogen bonding between the carboxylic acid groups. The hydrogen bonding between carboxylic acid groups in TPA1 and TPA2 holds them together in motifs that were expected by analogy to other benzenoid and isophthalic acid functionalized systems. Use of benzimidazole as a hydrogen-bonding substituent did not yield diffraction quality crystals, but hydrogen-bonded chains would be expected by analogy to other benzimidazoles. Hence, hydrogen bonding does serve to bring the triarylamines into close solid-state proximity. This has potential use for the crystal assembly of dopable organic paramagnet precursors and for bringing electron donor triarylamines into proximity with other molecules to act as energy transfer or quenching agents.

BImTPA exhibited only small spin yields in cationic doping experiments, with limited persistence of its radical cation solutions. This was attributed to spin delocalization in this system. Cationic doping of the carboxylic acid functionalized TPA2 gives highly persistent radical cations readily characterized by solution molecular spectroscopy. Solid-state magnetic measurements of the latter radical cations show strong doping levels, with modest antiferromagnetic exchange interactions at low temperature. This high persistence of the carboxylic acid functionalized radical cation systems makes them promising as possible spin-bearing components in solid-state magnetic materials and related electronic materials.

**Experimental Section**

4-(N,N-Bis(4-methoxyphenyl)amino)benzoic acid (TPA1). 4-Lo-benzoic acid methyl ester (3.43 g, 13.1 mmol) and N,N-bis(4-methoxyphenyl)amine (3.00 g, 13.1 mmol) were dissolved in toluene (19.6 mL). Sodium tert-butoxide (1.51 g, 15.7 mmol), tri-tert-butylphosphine (0.318 g, 1.57 μmol), and tris(dibenzylideneacetonedi)palladium (0.300 g, 0.328 μmol) were added, and then the mixture was heated at 100 °C for 7 h. The reaction mixture was neutralized with 1 N aqueous ammonia and extracted with chloroform. The organic layer was washed with brine and evaporated to give the crude product, which was added to 146 mL of ethanol and mixed with 146 mL of 1 M aq NaOH. The mixture was heated to reflux for 1 h and then cooled to room temperature and concentrated. The precipitate that formed was filtered. Then 182 mL of water was added, and 182 mL of 1 M aq HCl was added.

**TABLE 1. Solution ESR Hyperfine Coupling Constants**

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<tr>
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<th>a(N)</th>
<th>a(H)</th>
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<tbody>
<tr>
<td>TPA1-rc</td>
<td>8.58</td>
<td>1.37(6H)</td>
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<tr>
<td></td>
<td>1.00(6H)</td>
<td>0.56(6H)</td>
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<td></td>
<td>8.57</td>
<td>1.38(6H)</td>
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<tr>
<td></td>
<td>1.05(6H)</td>
<td>0.57(6H)</td>
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<tr>
<td></td>
<td>8.33</td>
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<tr>
<td>TPA2-rc</td>
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<tr>
<td>BImTPA-rc</td>
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(5) All computations in this paper were carried out using Spartan 2002 for Irix 6.5 by Wavefunction Inc., Irvine CA.
added dropwise while the reaction was stirred. The resulting yellow precipitate was filtered, washed with water, and dried. Chromatography of the solid (silica gel, ethyl acetate) yielded TPA1 (2.15 g, 70%), which crystallized as yellow rhombi from chloroform/methanol: mp 204–206 °C; 1H NMR (CDCl3) δ 7.85 (d, 2H, J = 9.0 Hz), 7.11 (d, 4H, J = 9.0 Hz), 6.87 (d, 4H, J = 9.0 Hz), 6.81 (d, 2H, J = 9.0 Hz), 3.81 (s, 6H); 13C NMR (CDCl3) δ 171.1, 157.0, 153.4, 139.3, 131.6, 127.9, 119.0, 116.8, 115.0, 55.5; IR (KBr pellet, cm⁻¹) 2958 (OH str), 1669 (C=O str); MS (EI, m/z) calcd for C27H23N3O2: 421, found 421.5. Anal. Calcd for C27H23N3O2: C, 76.94; H, 5.50; N, 9.57. Found: C, 76.49; H, 5.60; N, 9.71.

Oxidation of TPA1 with AgSbF6. A solution of AgSbF6 (53.9 mg, 157 μmol) in dichloromethane (5 mL) was added dropwise to a solution of TPA1 (50.0 mg, 143 μmol) in dichloromethane (2.5 mL); the reaction instantly turned from yellow to deep violet. The solution was then filtered and evaporated to give the crude product, which was dissolved in a small amount of dichloromethane and poured into excess hexane to precipitate the radical cation as a deeply colored violet-black solid (44.2 mg, 53%) that was dried under vacuum for ESR and magnetism studies.

Oxidation of TPA2 with AgSbF6. A solution of AgSbF6 (95.9 mg, 279 μmol) in dichloromethane (9 mL) was added dropwise to a solution of TPA2 (100 mg, 254 μmol) in dichloromethane (2.5 mL); the reaction instantly turned from yellow to red-purple. The solution was then filtered and evaporated to give the crude product, which was dissolved in a small amount of dichloromethane and poured into excess hexane to precipitate the radical cation as a darkly purple colored solid (66.0 mg, 41%) that was dried under vacuum for ESR and magnetism studies.

Oxidation of BImTPA with AgSbF6. A solution of AgSbF6 (17.9 mg, 52.2 μmol) in dichloromethane (2 mL) was added dropwise to a solution of the BImTPA (20 mg, 47.5 μmol) in dichloromethane (5 mL); the reaction instantly turned from yellow to green. The solution was then filtered and evaporated to give the crude product, which was dissolved in a small amount of dichloromethane and poured into excess hexane to precipitate the radical cation as a deeply colored green-to-black solid (17.9 mg, 57%) that was dried under vacuum for ESR and magnetism studies.

Acknowledgment. This work was supported by the National Science Foundation under Grant CHE-0415716. The University of Massachusetts Mass Spectrometry Facility, Nanomagnetic Characterization Facility, X-ray Structural Characterization Laboratory, and EPR Facility are also supported in part by the National Science Foundation.

Supporting Information Available: General methods used, crystallographic summaries for TPA1 and TPA2 and TPA2-THF, including full details in CIF format; computational spin density summaries for radical cations; solution ESR spectra and hyperfine simulations for radical cations, FTIR spectra for neutral amines; characterization NMR spectra, summary of Curie and Weiss constants from doping experiments, cyclic voltammetry for TPA1, TPA2, and BImTPA. This material is available free of charge via the Internet at http://pubs.acs.org.


FIGURE 2. (a) Paramagnetic susceptibility γ versus temperature plots for solution doped samples of TPA1 (●), TPA2 (○), and BImTPA (□) at 1000 Oe; (b) Curie–Weiss plot for TPA1; (c) Curie–Weiss plot for TPA2.