Crystalline boron-linked tetraaminoethylene radical cations†

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Single-electron oxidation of neutral boryl-linked tetraaminoethylene derivatives 4 led to the formation of radical cations 4c+, which have been isolated and fully characterized. X-ray diffraction analysis, EPR spectroscopy, and computational studies revealed that the unpaired electron is delocalized over the B2N4C2 skeleton and the spin density mainly resides on the carbon and boron atoms.

Introduction

Boron-containing radicals have been extensively studied due to their importance in fundamental chemistry and promising applications in various organic syntheses involving radical reactions.1 Thus far, a number of stable anionic2 and neutral3 boron radicals have been reported. By stark contrast, isolable boron radical cations are extremely rare probably due to the intrinsic electron deficient nature of boron. Indeed, only a handful of boron radical cations supported by the strong electron donors L have been isolated and structurally characterized by the Bertrand ([I]4, Braunschweig ([II]5, Xie ([III]6), Himmel ([IV]7, and Harman ([V]8) groups (Fig. 1a). While two boron-containing radical cations ([VI] and [VII]9) without Lewis base stabilization have also been reported, the unpaired electron was most likely localized in the substituents rather than the boron atoms.

Owing to their peculiar ability to serve as powerful organic reducing reagents, electron-rich olefins (EROs) have received a surging interest more than two decades.11,12 During the redox reactions with EROs, the corresponding oxidized forms of EROs, especially the radical cations are concomitantly generated and therefore have been considered as key species (Fig. 1b). Among them, the radical cation of tetrathiafulvalene (TTF) [VIII]13 has been extensively studied owing to its diverse utilities in various applications.14–17 Compared to TTF, tetraazafulvalenes (TAFs) [IX] feature more negative potentials18 which makes them attractive organic super-electron donors14–17 and organocatalysts.12 However, their corresponding radical cations [IX] have never been isolated thus far, mainly due to the facility of two-electron oxidation process attributed to the small electronic coupling. Recently, the Bertrand group reported two isolable triazaolefin radical cations ([X] and [XI]c+) and demonstrated the methylene-tethered [XI]c+ exhibits a larger electronic coupling.20 Similarly, it has been reported that aliphatic tetraaminoethylene (TAEs) [XII] linked by (poly)methylene chains increases

![Fig. 1](image-url) (a) Structurally characterized boron-containing radical cations. (b) Examples of radical cations of electron-rich olefins (Dipp = 2,6-diisopropylphenyl). (c) Present work.
Results and discussion

Treatment of tetraaryloxalamidine 1 (ref. 23) with one equivalent of dibromophenylborane in the presence of two equivalents of diisopropylmethylene in toluene afforded 2a as a yellow powder in 78% yield. A subsequent reaction between 2a and a stoichiometric amount of dibromophenylborane in toluene immediately afforded a brown precipitate, which was collected by filtration and then washed with hexane to give 3a as a white solid in 75% yield. A toluene solution of 3a with two equivalents of potassium graphite (KC₈) was stirred overnight under ambient condition, and after work up, boryl-linked tetraazalefin derivative 4a was obtained as a yellowish green powder in 54% yield (Scheme 1). The ¹¹B NMR spectrum of 4a displays a broad singlet at 25.4 ppm, which is shifted downfield with respect to that (9.8 ppm) of 3a. Compound 4a is thermally stable both in the solid state and in solutions, and it melts at 339 °C without decomposition. When para-substituted dibromophenylboranes with F and Bu groups were employed under the same reaction procedures, the corresponding derivatives 4b and 4c were obtained in moderate yields. In the ¹¹B NMR spectra, a broad singlet appears at δ = 26.4 ppm (4b) and δ = 26.1 ppm (4c), respectively. Compounds 4a-c can be deemed inorganic/organic hybrid versions of pentalene dianion,²⁴ which has been widely utilized as an ancillary ligand of metal complexes.²⁵

X-ray diffraction analysis of 4a confirmed the essentially planar B₂N₂C₂ framework with a propeller-like orientation of the six ary rings (Fig. 2a). Two boron atoms adopt perfectly trigonal-planar geometry (Σ₁ = 360°) with the N1–B1–N2 bond angle of 105.8(4)°. The equal B–N (1.460(5) Å) bond distances are significantly shorter than the corresponding bonds (1.581(4) Å and 1.583(4) Å) in 3a (Fig. S71f) and in the range between typical B–N single bond (1.50 Å) and B≡N double bond (1.31 Å), indicating the partial multiple bond property. Compared to those of 3a, there are lengthening of the C16–N1 bond (1.396(3) Å) and the C17–N2 bond (1.397(4) Å) and a markedly shortening of the C16–C17 bond (1.323(7) Å), which is close to those found in tetrakis(dimethylamino)ethylene (TDAE) (1.350(2) Å), X (1.3459(16) Å) and XI (1.344(2) Å).²⁵ Compound 4b (Fig. S72f) exhibits metric parameters similar to those of 4a. DFT calculation performed on 4a at the M062X/6-31G(d,p) level of theory shows that the HOMO of 4a corresponds to the π-system over the B₂C₂N₄ skeleton featuring a node between each NBN and the central CC π-unit (Fig. 2b). Natural bond orbital (NBO) analysis gave Wiberg bond index values of the C–C bond (1.51), the B–N bonds (0.94 and 0.93) and the C–N bonds (1.05 and 1.05) of the B₂C₂N₄ skeleton.

Cyclic voltammetry of 4a in CH₂Cl₂ (0.1 M Bu₄NPF₆ as the electrolyte) showed two well-separated oxidation waves (Fig. S79f), which is in sharp contrast to TAFs IX exhibiting very close redox waves (ΔE₁/₂ < 0.3 V).²⁴ In the first oxidation of 4a at E₁/₂ = −0.44 V (referenced against the ferrocene/ferroenium (Fc/Fc⁺) couple) was found to be a reversible process associated with the formation of the radical cation species whereas the second oxidation centered at E₁/₂ = 0.43 V was irreversible. Cyclic voltammograms of 4b and 4c show similar oxidation events with the first reversible (−0.44 V for 4b and −0.56 V for 4c) and second irreversible (0.41 V for 4b and 0.33 V for 4c) waves, respectively (Fig. S80 and 81f).

Scheme 1 Synthesis of compound 4 (Mes = 2,4,6-trimethylphenyl).
The first oxidation potentials of 4a-c are less negative than those of tetraazafuvalenes IX (around \(-1.00\) V vs. SCE, Fe/Fe\(^{3+}\): \(E_{1/2} = 0.46\) V vs. SCE). Thus, upon incorporation of the boryl groups, the reducing power of compounds 4a-c becomes weaker than that of tetraazafuvalenes IX.

It has been reported that tetraaminothienylene derivative typically undergoes the addition reaction with an electrophile at the central C=C moiety.\(^2\) To investigate the chemical behavior of 4, we performed their reactions with electrophiles. Treatment of a toluene solution of 4a with an equimolar amount of trifluoromethane sulfonic acid (HOTf) at room temperature afforded the corresponding conjugated acid 5a as a white powder in 85% yield (Scheme 2). The \(^{11}\)B NMR spectrum displays a broad singlet at 38.1 ppm which is shifted downfield with respect to that (25.4 ppm) of 4a. In the \(^1\)H and \(^{13}\)C NMR spectra, a singlet at 7.71 ppm for NC and reddish brown powder (5b) were observed, respectively. Under the same conditions, 4b and 4c underwent protonation with HOTf to afford the respective products 5b and 5c in good yields. An X-ray diffraction studies of 5a revealed that both boron centers feature the trignon-planar geometry. One of the central carbon atoms in the \(\text{B}_2\text{C}_2\text{N}_4\) ring is in the tetragonal configuration with a hydrogen atom whereas its adjacent catonic carbon exhibits the trignon-planar geometry with a formal charge of \(+1\) (Fig. 3a). The distance of the C25–C26 (1.490(5) Å) bond is significantly longer than the corresponding bond (1.323(7) Å) in 4a, and comparable to typical single C–C bond length. These results demonstrate the tetraiminothienylene-type nature of 4.

In line with the oxidation potentials, chemical single-electron-electron oxidations of 4a and 4b were readily accomplished by using of [Ph\(_3\)C][B(C\(_6\)F\(_5\))\(_4\)] as an oxidizing agent. Reactions of 4a and 4b with [Ph\(_3\)C][B(C\(_6\)F\(_5\))\(_4\)] in \(\text{CH}_2\text{Cl}_2\) at ambient temperature smoothly generated 4a\(^{++}\) and 4b\(^{++}\) as NMR-silent red (4a\(^{++}\): 84%) and reddish brown powder (4b\(^{++}\): 87%), respectively (Scheme 2). Crystals suitable for X-ray crystallographic studies were obtained by recrystallization from a mixture of \(\text{CH}_2\text{Cl}_2\) and toluene solution. Like the neutral 4a and 4b, 4a\(^{++}\) and 4b\(^{++}\) bearing the 9π-system retain the planar \(\text{B}_2\text{N}_4\text{C}_2\) skeletons with trigonal planar sp\(^3\) boron atoms (Fig. 3b and S73\(^a\)). Upon oxidation of 4a to 4a\(^{++}\), the B–N bonds (1.458(3)–1.474(3) Å) and the C=C (1.414(3) Å) bond are very slightly lengthened, while the endocyclic C–N bonds (1.352(2)–1.365(2) Å) are shortened, which is in good agreement with the electronic structure of 4a. Thus, upon removal of one electron from the HOMO of 4 (Fig. 2b), the electron density in the C=C and N–B–N π-bonding orbitals decreases, leading to the elongation of the C=C and B–N distances, while the repulsion between the bonding (N–B–N and C=C) electron pairs decreases resulting in the shortening of the C–N bond distances. Note that 4\(^{++}\) represent the first example of radical cations derived from BN heterocycles featuring some aromatic nature (Fig. S84, Table S2\(^b\)\(^,\)\(^c\)\(^,\)\(^d\)\(^,\)\(^e\)\(^,\)\(^f\)\(^,\)\(^g\)\(^,\)\(^h\)\(^,\)\(^i\)\(^,\)\(^j\)\(^,\)\(^k\)\(^,\)\(^l\)\(^,\)\(^m\)\(^,\)\(^n\)\(^,\)\(^o\)\(^,\)\(^p\)\(^,\)\(^q\)\(^,\)\(^r\)\(^,\)\(^s\)\(^,\)\(^t\)\(^,\)\(^u\)\(^,\)\(^v\)\(^,\)\(^w\)\(^,\)\(^x\)\(^,\)\(^y\)\(^,\)\(^z\)\(^,\)\(^{+}\))\(^,\)\(^{++}\)). Moreover, 4\(^{++}\) correspond to the anion radical of organic pentalene, among which only the derivatives featuring thermodynamically highly stabilizing benzo substituents have been structurally characterized.\(^2\)\(^,\)\(^3\)

The EPR spectrum of 4a\(^{++}\) measured in fluorobenzene at room temperature displays a complex system (\(g = 2.0050\)) arising from couplings with two boron (\(a(11\text{B}) = 1.16\) G, \(a(13\text{B}) = 0.63\) G), four nitrogen (\(a(14\text{N}) = 1.53\) G), and four hydrogen atoms (\(a(1\text{H}) = 1.32\) G) at the ortho-positions of phenyl groups (Fig. 4a). The \(^{11}\)B hyperfine coupling constant is smaller than those of I (6.43 G)\(^,\)\(^2\) and isolable boron radical anions,\(^4\) but comparable to those of II \((\leq 1.18\) G)\(^,\)\(^2\) and reported stable neutral boron radicals \((0.96–8.5\) G)\(^,\)\(^4\). The moderate \(a(11\text{B})\) coupling constant together with relative small \(a(14\text{N})\) values suggests delocalization of the unpaired electron over the \(\text{B}_2\text{N}_4\text{C}_2\) core. To gain more insight into the electronic structure of 4a\(^{++}\), unrestricted DFT calculations were performed at the UM062X/6-31G(d,p) level. NBO method confirmed that the spin density is entirely delocalized over the \(\text{B}_2\text{N}_4\text{C}_2\) framework with some extension to the two Ph groups on the B atoms (Fig. 4b). The spin density is estimated to be present mainly on the carbon (0.26e \(\times\) 2) and the boron (0.19e \(\times\) 2) atoms with a relatively small contribution of the nitrogen atoms (0.02e \(\times\) 4). While the several resonance forms including 4\(^{++}\), 4\(^{++}\), 4\(^{++}\) can be drawn for 4\(^{++}\) (Fig. 4c), the X-ray diffraction analysis, ESR spectra in

Scheme 2 Synthesis of 4\(^{++}\) and 5.

Fig. 3 (a) Solid-state structures of 5a (hydrogen atoms are omitted for clarity). Thermal ellipsoids are set at the 50% probability level. Selected bond lengths [Å] and angles [°]: B1–N1 1.514(6), B1–N2 1.422(6), B1–C1 1.546(7), B2–N2 1.430(6), B2–N4 1.489(6), B2–C26 1.542(7), N1–C26 1.330(5), N2–C25 1.469(5), N3–C25 1.484(5), N4–C24 1.365(5), C25–C26 1.490(5), N1–B1–N2 107.3(4), N1–B1–C1 123.8(4), N2–B1–C1 129.2(4), N3–B2–N4 107.0(4), N3–B2–C36 126.7(4), N4–B2–C36 127.6(4). (b) Solid-state structures of 4a\(^{++}\) (hydrogen atoms are omitted for clarity). Thermal ellipsoids are set at the 50% probability level. Selected bond lengths [Å] and angles [°]: B1–C3 1.545(3), B1–N1 1.458(3), B1–N2 1.474(3), B2–N2 1.464(2), B2–N4 1.472(2), C1–N1 1.356(5), C1–N3 1.365(2), C2–N2 1.356(2), C2–N4 1.472(1), C1–C2 1.414(3), C3–B1–N1 126.64(17), C3–B1–N2 126.69 (17), N1–B1–N2 106.66(16), C27–B2–N3 125.82(16), C27–B2–N4 127.73(17), N3–B2–N4 106.40(15).
combination with DFT results indicate that compared with the N-centered radical form $4_{\text{N}}^+$, contribution of others including the tetraaminoalkene radical cation $4_{\text{C}}^+$ and boron radical cation $4_{\text{B}}^+$ is more significant to the actual electronic structure of $4^+$. 

Conclusions

In summary, we have shown that boryl-tethered tetraaminoalkene derivatives 4 can be readily synthesized and they undergo protonation reaction and one-electron oxidation reaction. The former afforded the corresponding salts 5 whereas the latter led to the formation of isolable radical cations $4^+$. X-ray diffraction analysis, EPR spectra as well as computational studies of $4^+$ revealed the unpaired electron delocalized over the $\text{B}_2\text{N}_4\text{C}_2$ framework mainly resides on the carbon and boron atoms.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


