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Assembly of a Planar, Tricyclic B₄N₈ Framework With s-Indacene Structure

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A neutral, formally 16-π-electron, tricyclic tetrahydridotetraborane was obtained in a two-step procedure involving self-assembly of a dilitiodiborate with B₄N₈ framework and subsequent oxidation of the phenylborate moieties to boranes and biphenyl using Fe(II) as an oxidant.

The chemistry and properties of borazines, the “inorganic” analogues of benzene, were intensely investigated in the mid-sixties.¹ Borazine research has experienced a resurgence in the last few years, driven by the applications as precursors to boron nitride fibres and ceramics,² as well as molecular materials.³ Boron derivatives of hydrazines have received considerably less attention. A limited number of molecules with bi and tricyclic BN frameworks have been reported as well,⁴ including analogues of phenalene,⁵ naphthalene⁶ and Dewar benzene.⁷

We were interested in studying the ligand properties of the planar 6-π-electron system 1,⁸ which was obtained through quantitative self-assembly of monomethylhydrazine with PhB(NMe₂)₂, according to a reported procedure.⁹ Deprotonation of 1 with KH followed by methylation with MeI resulted in clean replacement of the more acidic ring proton in 1 with a methyl group. The disappearance of the signal corresponding to the ring proton of 1 at 7.48 ppm in the ¹H NMR spectrum clearly indicated the regiochemistry of the reaction. The lithium amide produced by deprotonation of the exocyclic nitrogen in 2 with LiTMP dimerized through N-B bond formation, producing the dilitium salt of a tricyclic borate dianion 3. The ¹H and ¹³C NMR spectra confirmed the presence of two inequivalent phenyl groups and three inequivalent methyl groups, while the ¹¹B NMR spectrum featured signals corresponding to the borane and the borate moieties at 28.5 and 2.7 ppm, respectively. The chemical shift of the ³Li NMR signal (-2.18 ppm) is typical for Li(thf)₄⁺, suggesting the presence of solvent separated ion pairs in THF solution.⁷

A crystallographic determination for 3 revealed a centrosymmetric structure featuring a tricyclic, annelated B₄N₈ skeleton containing two B₃N₃ rings connected through a B₄N₃ ring (Figure 1). As indicated by the NMR spectra, the skeleton contains two borane and two borate moieties. The tetracoordinated boron, B(2), is situated at 0.6 Å outside the best plane formed by the other skeletal atoms, determining the envelope conformation of the five

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Scheme 1 Synthesis of polycycles 3 and 4.

Fig. 1 Molecular structure of 3 with 50% probability level thermal ellipsoids. Hydrogen atoms on the organic substituents have been omitted for clarity and atoms marked with a prime (') character are at equivalent position (-x, -y, 1-z). Selected bond lengths (Å): B(1)-N(2) 1.441(2), B(1)-N(3) 1.408(2), B(2)-N(3) 1.562(2), B(2)-N(3') 1.556(2), B(2)-N(4') 1.566(2), N(1)-N(2) 1.475(2), N(3)-N(4) 1.445(2), Li(1)-N(1) 2.189(3), Li(1)-N(3) 2.326(3), Li(1)-N(4') 2.035(3), Li(1)-O(1) 1.902(3), N-C 1.566(2), N-N 1.475(2)
membered rings (folding angle along the N(1)–N(3) axis ca. 30°) and the chair conformation of the six-membered ring (folding angle along the N(3)–N(4) axis ca. 55°). The N-N bond lengths are 1.443(2) and 1.475(2) Å, comparable to the N-N bond length in hydrazine (1.45 Å). The B-N bonds involving the borane center

Fig. 2 One of the two independent molecules in the structure of 4 with 50 % probability level thermal ellipsoids. Atoms marked with a prime (') character are at equivalent position (-x, 2-y, -z). Selected bond lengths (Å) and angles (°); calculated values are reported in square brackets: B-N 1.411 – 1.416, N-C 1.446(3) – 1.463(3) [1.440 – 1.447], C-B 1.556(3) [1.565], 1.570(3) [1.565], B(3’)-N(5)-N(6) 113.72(15) [113.5], N(5)-N(6)-B(3) 122.09(15) [122.2], N(6)-B(3)-N(5) 124.18(18) [124.2], B(3)-N(6)-B(4) 108.26(17) [108.0], B(4)-N(8)-N(7) 109.49(15) [109.8], N(8)-N(7)-B(3) 106.70(15) [107.1], N(7)-B(3)-N(6) 108.35(16) [108.1].

B(1) (1.408(2) and 1.441(2) Å) are much shorter than those involving the borate center B(2) (1.556(2) – 1.566(2) Å). The lithium ions are positioned 1.7 Å outside the best plane of the ligand, and are coordinated by the three nitrogen atoms connected to the borate center. The coordination sphere of the lithium ion is completed by an intramolecular contact involving the ipso carbon of the phenylborate moiety, and a THF molecule. Oxidation of 3 with FeCl₄(THF)²⁺ in THF produced the neutral BN tricycle 4 in low (22 %) but reproducible isolated yield. The probable product of this reaction, FePh₃(THF)²⁻, subsequently decomposed to metallic iron and biphenyl, which were identified in the reaction mixture. A similar decomposition pathway was reported for FePh₃(PEt₃)²⁻ at temperatures above 0 °C. The¹H and¹³C NMR spectra of 4 featured the resonances expected for one phenyl group and three inequivalent methyl groups, and the signals corresponding to the two inequivalent boron centers merged into a broad singlet at 25.1 ppm. The use of a stoichiometric quantity of Li₄ as oxidizing agent instead of FeCl₄(THF)²⁺ did not result in the formation of 4. The electrochemical oxidation of tetraphenylborane to boric acid and biphenyl in aqueous conditions at a potential of 0.216 V vs. SCE has been reported, however, the reaction of Li[B(PPh₃)₄] with FeCl₄(THF)²⁺ failed to produce BPh₃. Thus, the formation of the stable polycycle appears to be an essential thermodynamic contributor to the oxidation of the phenylborate 3 to biphenyl and the borane 4.

The X-ray diffraction study of 4 reveals two very similar, centrosymmetric, independent molecules containing planar B₄N₄ frameworks with s-indacene structure (Figure 2). The tricyclic skeleton is regular, with all B-N and N-N bonds having the same length, between 1.423(3) and 1.441(2) Å. This length is characteristic of a partial multiple bond character for N-B, as observed in borazines (1.42 – 1.44 Å), and for a single N-N bond (vide supra). The phenyl substituents show little deviation from the ring plane (5.3°), while the C-N bonds form angles of 22.7 – 31.1° with this plane, in an all-trans arrangement. The all-carbon analog of 4, s-indacene, is an unstable molecule with formally antarimatic character according to the Hückel rule (12-π-electrons). Its more stable substituted derivatives¹ can be reduced to formally aromatic, 14-π-electron diianions. It could therefore be expected that a two electron oxidation or reduction of 4 would yield a stable dication or dianion, respectively, satisfying the Hückel condition of aromaticity. In order to investigate this hypothesis, theoretical investigations were performed for 4 as well as for its doubly reduced and oxidized forms.¹⁵ The geometry optimized structure of 4 is in excellent agreement with the data from the structural determination (Figure 2). The analysis of Kohn-Sham orbitals of 4 confirms that it is only formally a 16-π-electron system: though the tricyclic B₄N₄ framework is essentially planar, the geometry around all methyl substituted nitrogen atoms is pyramidal, giving rise to MOs with only partial π-like character (see Electronic Supplementary Information). The HOMO is π antibonding through all four N-N linkages, which readily explains the observed long bond distances indicative of single bonds.

The B₄N₄ framework is significantly nonplanar in the anion [4]²⁻ due to occupation of a MO with B-N antibonding character. Hence, a two-electron reduction of 4 leads to a structure which does not fulfill the general conditions of aromaticity. On the other hand, the two-electron oxidation of 4 yields a dication with a nearly planar structure because two electrons are removed from the N-N antibonding HOMO (see Electronic Supplementary Information). The minor geometric distortions present in the system arise from steric interactions, as evidenced by the geometry optimized structure of a hydrogen substituted tricycle [B(NH₂)₄]²⁺, displaying a perfectly planar Hückel aromatic geometry. Interestingly, the closed shell singlet state SCF solution of [4]²⁺ has an internal instability indicating that the singlet ground state of this dication has nitrogen-centered diradical character. Hence, the open-shell singlet diradical state of [4]²⁺ was modeled using broken symmetry formalism and the calculated structure was compared to the lowest energy triplet state. The calculations showed that the two spin states are very close in energy, with the formally open-shell singlet state being ca. 5 kJ mol⁻¹ lower than the triplet state.

Unfortunately the experimental search for a stable oxidation product of 4 remained fruitless. Cyclovoltammetric measurements were performed in THF in the range of −2.5 to 1.5 V, and they showed two irreversible oxidation steps at 0.35 and 0.90 V vs. SCE (−0.21 and 0.34 V vs. Fc). Chemical oxidation with [Cp₂Fe]PF₆ produced Cp₂Fe and a solid that was insoluble in organic solvents and was not further characterized. Radical species were not detected by EPR upon in-situ electrochemical oxidation. We note that stable radicals with s-indacene-like frameworks are well-known in the chalcogen-nitrogen chemistry.¹⁹

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Notes and references
All operations were conducted under strict exclusion of air and moisture. Solvents were dried prior to use and methylhydrazine was dried over CaH2. Methylated hydrazines are highly toxic and probable carcinogens, and their handling requires special precautions; all residues were neutralized using commercial bleach solution.

Synthesis of 2: A solution of 1 (0.700 g, 2.65 mmol) in THF (20 mL) was slowly added to a suspension of KH (0.106 g, 2.65 mmol) in THF (15 mL). KH dissolved with evolution of hydrogen producing a yellow solution that was stirred at ambient temperature for another 2 h. Mel (0.165 mL, 2.65 mmol) was added to the mixture and the colourless suspension was stirred for an additional hour and then concentrated under vacuum to ca. 3 mL. Hexane (30 mL) was added and the Kl by-product was filtered off. The solvent was subsequently removed in vacuo, leaving behind the product as a colourless powder (638 mg, 88%). 1H NMR (400 MHz, CD2Cl2, 25 °C): δ 2.32 (3H, JHH = 6.4 Hz, HNCH2), 2.88 (6H, (NCH3)2), 3.76 (qH, JHH = 6.4 Hz, HNCH2), 7.25 - 7.34 (3H, m - p-C6H4), 7.74 (4H, JHH = 6.6 Hz, o-C6H4). 13C NMR (100 MHz, CD2Cl2, 25 °C): δ = 31.7 (NCH3), 43.5 (NCH3), 128.8 (m - s-C6H4), 129.1 (p - s-C6H4).

Synthesis of 3: A solution of lithium 2,2,6,6-tetramethylpiperidide, LiTMP was prepared from 1.6 M n-butyllithium in hexane (1.12 mL, 1.80 mmol) and tetramethylpiperidine, TMP (0.254 g, 1.80 mmol) in THF (3 mL). The solutions of 2 (0.500 g, 1.80 mmol) in THF (3 mL) and LiTMP was pre-cooled to -35 °C and mixed, yielding an orange solution that was stored at -35 °C for a day. Subsequently it was allowed to warm to room temperature and volatiles were removed in vacuo, leaving behind a yellow residue that was washed twice with hexane (30 mL) and dried under vacuum. The product was obtained as a colourless powder (388 mg, 60.5%). 1H NMR (400 MHz, THF-d5, 25 °C): δ 2.01 (6H, NCH3), 2.28 (6H, NCH3), 2.71 (6H, NCH3), 7.12 - 7.31 (12H, m - p-C6H4), 7.53 (4H, JHH = 6.8 Hz, o-C6H4), 7.77 (4H, JHH = 6.8 Hz, o-C6H4). 13C NMR (100 MHz, THF-d5, 25 °C): δ 34.2 (NCH3), 39.1 (NCH3), 41.3 (m - s-C6H4), 126.3 (p - s-C6H4), 127.2 (p - s-C6H4), 127.5 (m - s-C6H4), 128.0 (m - s-C6H4), 134.0 (m - s-C6H4), 135.9 (m - s-C6H4). 11B NMR (128 MHz, THF-d5, 25 °C): δ = 2.7 (s and 28.5 (s, br); Li[N(CH3)2] 155 MHz, THF-d5, 25 °C): δ -2.18 (s). Needle-like X-ray quality crystals were obtained by recrystallization of 3 from THF.

Synthesis of 4: A solution of 3 (80.0 mg, 0.112 mmol) in THF (3 mL) was added to solid FeCl3(H2O)2 (38.0 mg, 0.141 mmol). The dark mixture was allowed to stand at room temperature for a day, and was then concentrated to 1 mL and cooled to -35 °C for several days until colourless crystals of 4 formed and were separated by decantation (10 mg, 22%). 1H NMR (400 MHz, THF-d5, 25 °C): δ 2.66 (6H, NCH3), 2.83 (6H, NCH3), 2.87 (6H, NCH3), 3.10 - 3.33 (6H, m - p-C6H4), 3.55 (4H, JHH = 5.6 Hz, o-C6H4). 13C NMR (100 MHz, THF-d5, 25 °C): δ 32.9 (s, N(CH3)2), 33.7 (s, N(CH3)2), 128.4 (m - s-C6H4), 128.9 (s, NCH3), 134.2 (s, m - s-C6H4). 11B NMR (128 MHz, THF-d5, 25 °C): δ 25.1 (s, br); MS (EI, 70 eV); m/z (%): 400 (51%) [M+2 - Me]-, 385 (13%) [M - Me], 278 (40) [M - Ph - 3Me]+, 154 (41) [B,NH3]-, HRMS for H2C2N5Ba2+ calculated 400.2809, found 400.2838.

Crystal data for 3: C26H22Ba2Li2N10O6, monoclinic, space group P21/n, a = 2.2996 (4), b = 16.4156 (4), c = 16.4156 (4), β = 100.171° (3), V = 1943.13 (13) Å3, Z = 2, D = 1.217 g cm-3, F(000) = 760, μ(Mo-Kα) = 0.074 mm-1, λ = 0.7073 Å, T = 123.2 K. K 8425 reflections measured, 4398 unique data (2θmax = 55.0°, Rint = 0.027). GOFF on F2 = 1.02, wR(F2) = 0.126 for all data, R(F) = 0.047 on reflections having I > 2σ(I), CCDC #649697.

Crystal data for 4: C26H22Ba4N8O4, 39.72% triclinic, space group P-1, a = 17.2400 (3), b = 9.1576 (12), c = 16.9832 (12), α = 78.588 (3), β = 83.194 (4), γ = 76.554° (4), V = 1059.71(11) Å3, Z = 2, D = 1.253 g cm-3, F(000) = 400, μ(Mo-Kα) = 0.077 mm-1, λ = 0.7073 Å, T = 173(2) K. 6791 reflections measured, 3693 unique data (2θmax = 50.0°, Rint = 0.031). GOFF on F2 = 1.01, wR(F2) = 0.146 for all data, R(F) = 0.046 on reflections having I > 2σ(I), CCDC #649698.

Cyclovoltammetry of 4: The measurements were performed in THF at an analyte concentration of 1.0mM, and using 0.1 M [NaBu4N]PF6 as a supporting electrolyte. A PARstat 2273 potentiostat was used at scan rates between 50 to 100 mVs⁻¹. The cell had a platinum disk working electrode, a platinum wire auxiliary electrode, and a silver wire pseudo-reference electrode. [Cu2CoCl4]⁺² with E° = +1.16 V vs. ferrocene and -0.80 V vs. the SCE was used as an internal standard.