

A Phosphine-Coordinated Boron-Centered Gomberg-Type Radical**

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Abstract: The P-coordinated boryl radical $[\text{Ph}_2\text{P}(\text{naphthyl})\text{BMes}]^\bullet$ (Mes = mesityl) was prepared by (electro)chemical reduction of the corresponding borenium salt or bromoborane. Electron paramagnetic resonance (EPR) analysis in solution and DFT calculations indicate large spin density on boron (60–70 %) and strong P–B interactions (P $\rightarrow$ B σ donation and B $\rightarrow$ P negative hyperconjugation). The radical is persistent in solution and participates in a Gomberg-type dimerization process. The associated quinoid-type dimer has been characterized by single-crystal X-ray diffraction.

Over the last few years, significant progress has been achieved in boron chemistry with the isolation of a variety of compounds featuring original structures, exhibiting interesting properties, and displaying rich reactivity. These developments concern highly reactive diamagnetic species (such as boryl anions,^[1] diborynes and oxoboryls,^[2] 1,3-diboratacyclobutane-1,3-diyls,^[3] boriniums,^[4] and borylene adducts^[5]) as well as radicals for which unusual bonding situations (such as one-electron B–B σ and π bonds)^[6] have been uncovered and important synthetic applications have emerged.^[7] In particular, boryl radicals (L $\rightarrow$ BR₂)^{*}, which can be considered as neutral tricoordinate boron radicals, have attracted considerable attention.

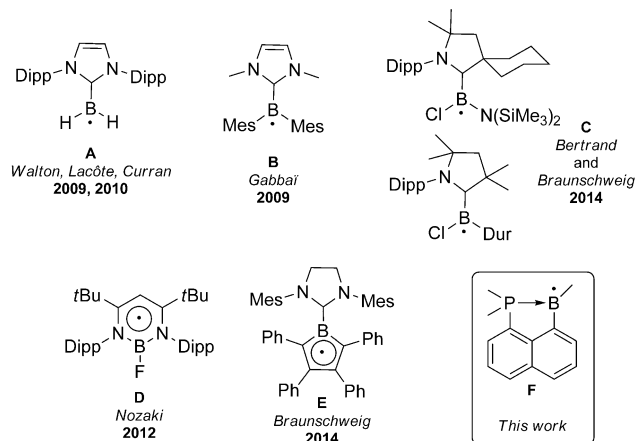


Figure 1. Different types of EPR-characterized/isolated boryl radicals **A–E** and the P-coordinated boryl radical **F** studied herein. Dipp = 2,6-diisopropylphenyl; Dur = 2,3,5,6-tetramethylphenyl.

Some reported examples of boryl radicals are shown in Figure 1. Basically, two different types can be distinguished: 1) acyclic systems, such as **A**,^[8a,b] **B**,^[8c] and **C**,^[5c,8d] in which a five-electron BR₂^{*} radical is stabilized by a strong Lewis base, typically a N-heterocyclic carbene (NHC), and 2) cyclic systems, such as **D** and **E**, which are stabilized by π delocalization.^[9]

In this context, we report herein a persistent P-coordinated boryl radical. A compound of type **F** (Figure 1) has been prepared by (electro)chemical reduction. The structure of the radical has been thoroughly analyzed by electron paramagnetic resonance (EPR) spectroscopy and DFT calculations. Single-crystal X-ray diffraction analysis revealed that the molecule undergoes Gomberg-type dimerization by B–C coupling. This B-centered radical is a new member in the family of naphthyl-bridged P/B compounds,^[10] and is a stable version of the transient phosphine–boryl radicals (R₃PBH₂)^[11]

Taking advantage of the strong P $\rightarrow$ B interaction enforced by the naphthyl bridge,^[10c–e] we have recently isolated the P-coordinated borenium salt **1** (see Figure 2, inset).^[10f] This prompted us to try to generate the corresponding boryl radical. The reduction of **1** was first assessed electrochemically. The cyclic voltammogram showed a quasi-reversible one-electron wave at -0.48 V (versus ferrocene/ferrocenium Fc/Fc⁺) in fluorobenzene (Figure 2).^[12] This reduction potential is significantly more positive than that of other borenium/boryl couples. For example, the NHC-stabilized radical **B** is formed at much lower potential ($E_{1/2} = -1.81$ V)^[8c] and the lowest reported reduction waves occur at $E_{1/2} = -0.86$ V for 9-acridinyl BMes₂ derivatives.^[13] As a result of its cationic

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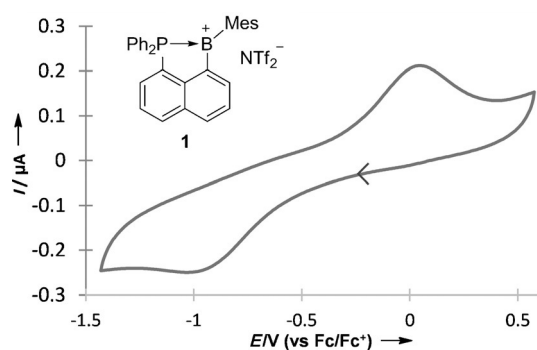
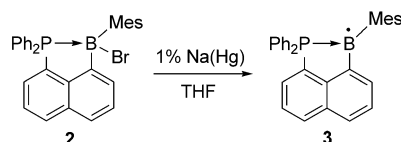


Figure 2. Cyclic voltammogram of **1** in a fluorobenzene solution containing $[n\text{Bu}_4\text{N}][\text{NTf}_2]$ (0.06 M) as the electrolyte. Scan rate: 500 mV s^{-1} ; potential reported against Fc/Fc^+ as the internal standard.

character, **1** can also be reduced much more easily than boranes, including the highly electrophilic tris(pentafluorophenyl)borane ($E_{1/2} = -1.17\text{ V}$).^[14]

Encouraged by the electrochemical reduction of **1**, we then explored the chemical reduction of its precursor, the P-coordinated bromoborane **2** (Scheme 1). Using zinc powder



Scheme 1. Synthesis of radical **3** by reduction of bromoborane **2**.

or 1% Na(Hg) amalgam in THF, a dark-red solution was formed from which compound **3** could be isolated by filtration and precipitation over pentane (83% yield).

The X-band EPR spectrum of **3** in THF/toluene solution at room temperature shows a complex but nicely resolved signal centered at an isotropic g value of $g_{\text{iso}} = 2.0026$ (Figure 3). No change was observed over days at room temperature, indicating the highly persistent character of this boryl radical. The hyperfine structure could be resolved by simulation,^[15] taking into account couplings to the ^{31}P and $^{11}\text{B}/^{10}\text{B}$ nuclei, as well as to two ^1H nuclei. The coupling constants to ^{31}P and ^{11}B nuclei (31 and 11 G, respectively) fall in the same range as those of transient phosphine–boryl radicals (R_3PBH_2).^[11] Delocalization of the unpaired electron over

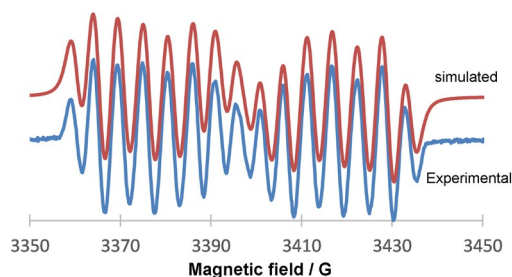


Figure 3. Experimental (gray) and simulated (black) X-band EPR spectra of **3** in toluene/THF solution at room temperature. $G = \text{gauss}$.

the organic substituents (in particular the naphthyl moiety, see DFT calculations described below) is indicated by the necessity to include two ^1H nuclei in the simulation with couplings of 4 and 5 G. However, the large $a(^{11}\text{B})$ coupling indicates that the spin density at boron is unusually high in **3**.^[16] The value measured for **3** falls in the upper range of those reported for persistent/isolable boryl radicals (7.3–13.2 G).^[5c,8] It is also higher than those of the triphenylboron and trimesitylboron anion radicals ($a(^{11}\text{B}) = 7.8$ and 9.9 G , respectively), which have approximately 60% spin density on boron.^[17]

To gain more insight into the structure of **3**, DFT calculations were performed.^[18] The geometry was optimized at the UB3PW91/6-31G** level of theory and the hyperfine coupling constants were then computed at the B3PW91/IGLO-II(P),EPR-II(H,B,C) level (see the Supporting Information, Tables S1 and S2).^[15] The obtained values (35.3 and 10.3 G for ^{31}P and ^{11}B nuclei, respectively) match remarkably well with those determined experimentally. DFT also nicely reproduces the two couplings to ^1H nuclei (5.7 and 5.5 G) and enables us to assign the nuclei as the protons on the naphthyl ring in the *ortho* and *para* positions with respect to boron (suggesting some delocalization of spin density). The optimized structure of **3** (Figure 4a) is also informative. The

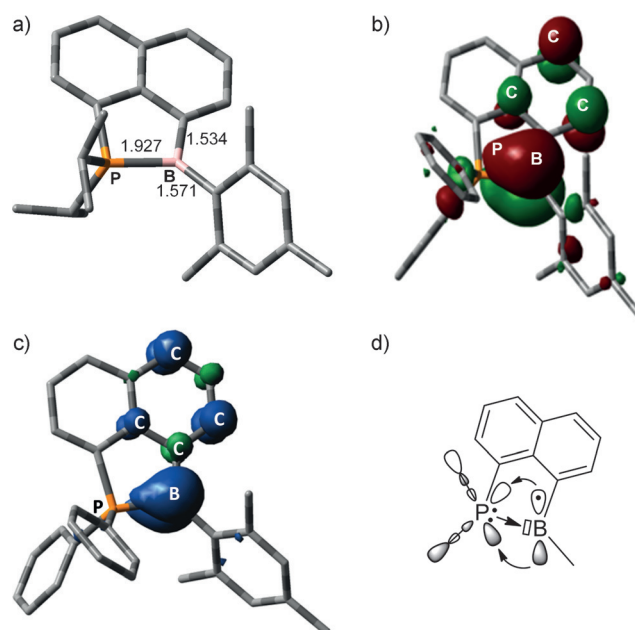


Figure 4. DFT analysis of compound **3**: a) optimized geometry (distances in Å); b) SOMO (cutoff=0.04); c) spin density distribution (isovalue=0.005 au); d) representation of the PB donation/back donation. H atoms are omitted for clarity.

boron center is in a trigonal planar environment (the sum of the bond angles $\Sigma B_\alpha = 359.6^\circ$) and the Mes group is twisted by 68.3° from the (naphthyl,P,B) plane, suggesting very little, if any, delocalization of spin density over the mesityl ring. Consistently, the $\text{B}-\text{C}_{\text{Mes}}$ bond (1.571 Å) is longer than the $\text{B}-\text{C}_{\text{naphthyl}}$ bond (1.534 Å). Bond lengths in pm changed to Å to correspond to Figure 4a and throughout the manuscript for

consistency. The presence of a strong $P \rightarrow B$ interaction is apparent from the short $P-B$ distance (1.927 Å). Comparison with the corresponding borenium salt **1** (2.006 Å) is interesting and reveals some shortening of the bond in **3**. This is rather counterintuitive given that the boron center is less electrophilic in the boryl radical **3** than in the borenium **1**, and thus reduction of **1** to form **3** would be expected to weaken the $P \rightarrow B$ interaction.

The electronic structure of **3** was then carefully analyzed by DFT calculations.^[15] The singly occupied molecular orbital (SOMO; Figure 4b) and the total spin density distribution (Figure 4c) are mainly localized on the B atom (60–70 % spin density) with small contributions of the P atom as well as the C_{ipso} , C_{ortho} , and C_{para} atoms of the naphthyl ring. Most noticeable are the overlaps of the $2p(B)$ orbital with 1) the adjacent $2p(C_{ipso})$ orbital (π bonding) and 2) the neighboring $\sigma^*(PC)$ orbitals (negative hyperconjugation). This picture is corroborated by natural bond orbital analysis (NBO), which gives at the first order a $\pi(BC_{ipso})$ orbital centered on boron (62 %) with an occupancy of $0.86e^-$. Stabilizing interactions with $\pi^*(C=C)_{naphthyl}$ and $\sigma^*(PC)$ orbitals are found at the second-order perturbation level (with NBO delocalization energies $\Delta E_{int} = 15.3$ and $9.5 \text{ kcal mol}^{-1}$, respectively).^[15] Interaction of the singly occupied $2p(B)$ with adjacent vacant $\sigma^*(PC)$ orbitals is reminiscent of the negative hyperconjugation occurring within P ylides (Figure 4d). The effect of the negative hyperconjugation is probably to contribute to the shortening of the PB distance in **3** ($P \rightarrow B$ donation/ $B \rightarrow P$ back donation) compared to borenium **1**.

The stability and high spin density of **3** at the B atom encouraged us to try to crystallize it. Our attempts were finally rewarded and suitable crystals were obtained from a THF solution layered with pentane at room temperature. According to X-ray diffraction, the molecule adopts in fact a dimeric quinoid-type structure $[3]_2$ in the solid state (Figure 5).^[19] The dimerization occurs at the boron atom of one boryl unit and the C_{para} (naphthyl) atom of the other boryl unit. The length of the resulting $B-C$ bond ($B2-C4 = 1.670(7) \text{ Å}$) is typical of a single $B-C$ bond. The two boron atoms are in different environments. The B atom directly involved in the dimerization (B2) becomes tetrahedral and its bond to phosphorus is elongated ($2.115(5) \text{ Å}$) due to steric shielding. The other boron atom (B1) remains trigonal planar. Atom B1 is engaged in a strong $P \rightarrow B$ interaction ($1.933(6) \text{ Å}$)^[20] and the bond towards C_{ipso} (naphthyl) is significantly shortened ($1.463(7) \text{ Å}$), indicating that the bond has double bond character.

The formation of $[3]_2$ is reminiscent of the dimerization of the trityl radical $Ph_3C^\bullet$ as reported by Gomberg (Scheme 2).^[21] Dimerization of triarylmethyl radicals is a particularly fascinating transformation in organic chemistry. The reaction has always attracted significant interest and undoubtedly represents a founding principle of radical chemistry.^[22] Comparatively, very little is known about related transformations with main-group radicals. Monomer/dimer interconversions have been observed with p-block radicals (in particular with $E = Si, Sn, P, Sb, Bi$),^[23] but they usually involve $E-E$ bond formation/cleavage. In a recent study on aryl-substituted silyl radicals, Sekiguchi et al. proposed a $Si-C_{para}$ dimerization process to

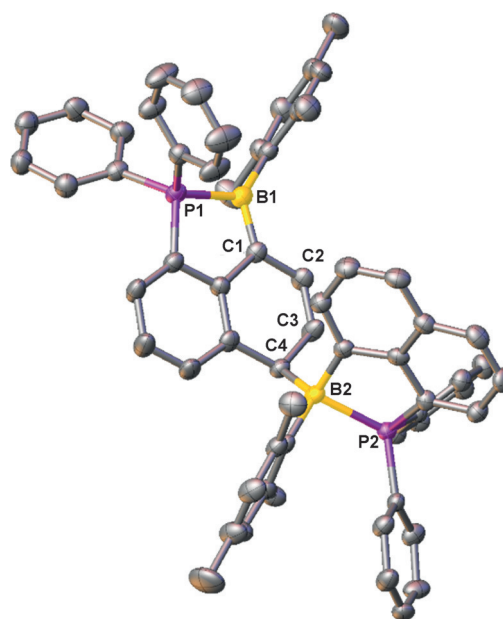
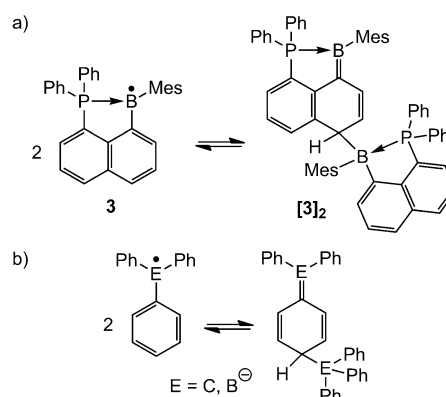


Figure 5. OLEX2 plot of $[3]_2$ with thermal ellipsoids set at 50% probability. For clarity, the hydrogen atoms have been omitted. Selected bond lengths [Å] and angles [°]: $B1-C1$ 1.463(7), $B1-P1$ 1.933(6), $B2-C4$ 1.670(7), $B2-P2$ 2.115(5), $C1-C2$ 1.434(7), $C3-C4$ 1.509(7), $C2-C3$ 1.338(6); $C1-B1-C11$ 131.4(5), $C1-B1-P1$ 101.0(4), $C11-B1-P1$ 127.5(3).



Scheme 2. a) Monomer–dimer equilibrium of **3** and b) the Gomberg-type dimerization of the triphenylmethyl radical ($E=C$) and the triphenylboron anion radical ($E=B^-$).

rationalize the decrease of the EPR signal intensity observed upon cooling.^[24] In the case of boron compounds, Ph_3B was reported early on to dimerize under reductive conditions (Na).^[25] The ionic nature and instability of the transient triphenylboron anion radical prevented detailed analysis of the dimerization process, but a Gomberg-type dimeric structure could be proposed based on NMR spectroscopic data.^[26] The dimerization process observed with the boryl radical **3** is unique among persistent/isolable boryl radicals^[27] and the $B-C_{para}$ coupling is probably favored by the same factors as for the trityl radical: $B-B$ coupling suffers from strong steric congestion whereas $C-C$ coupling is disfavored by a) the low spin density at the C_{para} position (20–25 % versus

60–70% at B according to DFT) and b) high loss of aromatic character (the two naphthyl units would adopt quinoid structures).

Further characterization of **[3]₂** indicated complete dimerization of **3** in the solid state, as demonstrated by the fact that the red powder obtained by crystallization is EPR silent. Additionally, in the cross-polarization magic angle spinning (CP-MAS) NMR spectrum, relatively sharp signals for ³¹P and ¹¹B nuclei were detected at expected chemical shifts ($\delta(^{31}\text{P}) = +11.5$ and -3.9 ppm; $\delta(^{11}\text{B}) = 25.0$ and 5.2 ppm). Finally, the equilibrium between **3** and **[3]₂** was analyzed by variable-temperature EPR spectroscopy in solution. From the variation of the integral of the EPR signal, the Gibbs free energy associated with the dimerization of **3** was estimated to $-15 \text{ kcal mol}^{-1}$.^[15] This value is slightly larger than that of the trityl radical ($\Delta G = -11 \text{ kcal mol}^{-1}$)^[28] which may be attributed, at least in part, to the lower aromaticity of the naphthyl unit (versus phenyl).

In summary, a new P-coordinated boryl radical has been prepared. According to EPR data and DFT calculations, the spin density is mainly localized on the boron center, with small contributions of the P atom and naphthyl unit. The radical is persistent in solution and exists in equilibrium with a head-to-tail B–C dimer, which has been characterized by X-ray crystallography. Gomberg-type dimerizations of this type are extremely rare with main-group radicals and future work will seek to explore further the chemistry of P-coordinated boryl radicals.

Keywords: boron · cyclic voltammetry · EPR spectroscopy · phosphorus · radicals

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