Theoretical Investigation of Paramagnetic Diazabutadiene Gallium(III)−Pnictogen Complexes: Insights into the Interpretation and Simulation of Electron Paramagnetic Resonance Spectra

Tuononen, Heikki; Armstrong, Andrea


All material supplied via JYX is protected by copyright and other intellectual property rights, and duplication or sale of all or part of any of the repository collections is not permitted, except that material may be duplicated by you for your research use or educational purposes in electronic or print form. You must obtain permission for any other use. Electronic or print copies may not be offered, whether for sale or otherwise to anyone who is not an authorised user.
Theoretical Investigation of Paramagnetic Diazabutadiene Gallium(III)-Pnictogen Complexes: Insights into the Interpretation and Simulation of EPR Spectra

Heikki M. Tuononen†* and Andrea F. Armstrong‡

Departments of Chemistry, University of Jyväskylä, P.O. Box 35, FIN-40014 Jyväskylä, Finland and University of Calgary, Calgary, Alberta, Canada T2N 1N4

† University of Jyväskylä
‡ University of Calgary

* To whom correspondence should be addressed.
Telephone: +358-14-260-2618
Fax: +358-14-260-2501
E-mail: hetuonon@cc.jyu.fi
Abstract

The electronic structures and the spin density distributions of the paramagnetic gallium1,4-diaza(1,3)butadiene (DAB) model systems \{('Bu-DAB)Ga(I)[Pn(SiH\textsubscript{3})\textsubscript{2}])\} and the related dipnictogen species \{(‘Bu-DAB)Ga[Pn(SiH\textsubscript{3})\textsubscript{2}]_2\} (Pn = N, P, As) were studied using density functional theory. The calculations demonstrate that all systems share a qualitatively similar electronic structure and are primarily ligand centered $\pi$-radicals. The calculated EPR hyperfine coupling constants (HFCCs) for these model systems were optimized using iterative methods and used to create accurate spectral simulations of the parent radicals \{(‘Bu-DAB)Ga(I)[Pn(SiMe\textsubscript{3})\textsubscript{2}])\} (Pn = N, P, As) and \{(‘Bu-DAB)Ga[Pn(SiMe\textsubscript{3})\textsubscript{2}]_2\} (Pn = P, As), the EPR spectra of which had not been simulated previously due to their complexity. Excellent agreement was observed between the calculated HFCCs and the optimum values, which can be considered the actual HFCCs for these systems. The computational results also revealed inconsistencies in the published EPR data of some related paramagnetic group 13-DAB complexes.
Introduction

Research into the chemistry of inorganic radicals of the main group elements has undergone a renaissance in recent years.\(^1\) Since many of the p-block elements have at least one naturally occurring spin-active isotope, the EPR spectra of these radicals are often quite complex due to hyperfine interactions of the unpaired electron with several magnetically active nuclei. The resultant poor resolution of these EPR spectra renders it difficult to extract accurate values of the hyperfine coupling constants (HFCCs) to the various spin-active nuclei. Because HFCCs are used to glean information regarding the spin density and spin distribution of the unpaired electron throughout a paramagnetic molecule,\(^2\) this is quite problematic, as it can impede researchers from gaining a thorough understanding of a particular radical system. One approach to overcoming this difficulty is to use DFT calculations to predict the magnitudes of the HFCCs. The calculated values can then be incorporated into a simulation of the experimental EPR spectrum, which, once optimized, yields accurate HFCCs. We have recently shown this methodology to be successful in assigning and interpreting the complex EPR spectra of both tetraisimidophosphate\(^3\) and boramidinate\(^4\) radicals.

The 1,4-diaza(1,3-)butadiene ligand (R-DAB) (1) readily undergoes a one-electron reduction to form the monoanionic diazabutadiene radical (2). By incorporating bulky substituents such as tert-butyl or 2,6-di\(\text{iso}\)-propylphenyl on the nitrogen atoms and chelating this bidentate anion radical to a metal center, it has proven possible to produce a series of stable paramagnetic compounds. Complexes of aluminum\(^5\) and gallium\(^6\)–\(^8\) have been reported where the metal is bis-chelated by one DAB anion radical 2 and a neutral
DAB ligand 1 forming a spirocyclic ligand-centered monoradical; isostructural radicals of the s-block metals lithium and magnesium are also known.  

\[
\begin{align*}
1 & \quad 2 \\
\text{H} & \quad \text{H} \\
\text{R} & \quad \text{R} \\
\text{N} & \quad \text{N} \\
\text{C} & \quad \text{C} \\
\text{C} & \quad \text{C} \\
\end{align*}
\]

More recent developments in this area have resulted in the isolation and characterization of a wide range of paramagnetic DAB complexes of the group 13 metals, particularly gallium.  

\[
\begin{align*}
3a, \text{Pn} = \text{N} & \quad 4a, \text{Pn} = \text{P} & \quad 5a, \text{Pn} = \text{As} \\
3a & \quad 4a & \quad 5a \\
\end{align*}
\]

Among these are the monocyclic pnictogen compounds \{(′Bu-DAB)Ga(I)[Pn(SiMe₃)₂]\}⁺ (3a, Pn = N; 4a, Pn = P; 5a, Pn = As) and the related dipnictogen species \{(′Bu-DAB)Ga[Pn(SiMe₃)₂]₂\}⁺ (7a, Pn = P; 8a, Pn = As), though the nitrogen analog of the latter radicals “{(′Bu-DAB)Ga[N(SiMe₃)₂]₂}” (6a) has not been isolated.  

The EPR spectra of these novel radicals are quite complex due to hyperfine couplings to the \(^1\text{H}\) and \(^{14}\text{N}\) nuclei of the DAB backbone, as well as to \(^{69,71}\text{Ga}\) \((^{69}\text{Ga}, 60 \% \text{, } I = 3/2; ^{71}\text{Ga}, 40 \% \text{, } I = 3/2)\), the pnictogen atom(s) \((^{14}\text{N}, 99 \% \text{, } I = 1; ^{31}\text{P}, 100 \% \text{, } I = 1/2; ^{75}\text{As}, 100 \% \text{, } I = 3/2)\), and the iodine nucleus \((^{127}\text{I}, 100 \% \text{, } I = 5/2)\) in the case of 3–5.

\[
\begin{align*}
3–5 & \quad 6–8 \\
\end{align*}
\]

At present, accurate simulations of the EPR spectra of 3a–5a, 7a, and 8a have not been obtained due to their complexity: though the magnitudes of some of the HFCCs have been estimated, these values alone do not accurately reproduce the experimental spectra. Additional complications in simulating these spectra have been attributed to “slight differences in \(^{69,71}\text{Ga}\) hyperfine couplings and isotropic g-values”.  

However, the electronic Zeeman interaction does not depend on nuclear isotopes.  

In addition, the
HFCCs of $^{69}$Ga and $^{71}$Ga isotopes in a given magnetic environment are inter-related by the ratio of their nuclear g-factors provided that the EPR spectrum shows no significant anisotropy. The near-perfect inversion symmetry with respect to peak positions exhibited by the experimental EPR spectra of these radical indicates the presence of negligible amounts of anisotropy.

A recent theoretical investigation$^{14}$ of a series of paramagnetic group 13-DAB complexes has indicated that the singly occupied molecular orbitals (SOMOs) of these systems are compositionally equivalent, and, like their spin density distributions, exhibit only a minor dependence upon the identity of the central metal and the substituents which complete its coordination sphere. The magnitudes of the HFCCs for these ligand-centered $\pi$-radicals were calculated and found to be in very good agreement with values extracted from experimental EPR spectra, in cases where the identity of the paramagnetic species has been definitively established. In light of these results, we have now undertaken a DFT study of the pnictogen-containing radicals $3$–$8$ in order to determine their electronic structures and spin density distributions. The complex EPR spectra of these radicals will then be interpreted in terms of their electronic structures and predicted HFCCs. Some implications of these results regarding the characterization of other know group 13-DAB complexes are also discussed.

Computational Details

The geometries of the radical systems \{('Bu-DAB)Ga(I)[Pn(SiH$_3$)$_2$] $^\prime$ ($3b$, Pn = N; $4b$, Pn = P; $5b$, Pn = As) and \{('Bu-DAB)Ga[Pn(SiH$_3$)$_2$] $^\prime$ ($6b$, Pn = N; $7b$, Pn = P; $8b$,}
Pn = As) were optimized in their doublet ground states using DFT. The calculations utilized the hybrid PBE0 exchange-correlation functional\(^\text{15}\) and Ahlrichs’ triple-zeta valence basis sets augmented by one set of polarization functions (TZVP).\(^\text{16}\) For iodine, the corresponding large-core ECP basis sets (46 electrons frozen) were used.\(^\text{16}\) The PBE0 hybrid density functional was chosen on the grounds of several published benchmarks that have shown it to perform well in calculating molecular properties for a wide variety of chemical systems.\(^\text{17}\) The way in which the PBE0 functional is constructed and the lack of empirical parameters fitted to specific physical properties also renders it appealing from a purely theoretical standpoint. Appropriate molecular point groups were used to improve the efficiency of the calculations.

Hyperfine coupling constants were calculated for 3b–5b in their geometry optimized structures using both quasi-relativistic and fully relativistic methods, and the unrestricted Kohn-Sham formalism. For compounds 6b–8b, only non-relativistic calculations were done. The non-relativistic calculations utilized the same TZVP basis sets and density functional as the geometry optimizations; several benchmarks have recently been published which demonstrate the suitability of TZVP basis set for the calculation of EPR parameters using DFT.\(^\text{18}\) For 3b–5b, the use of a quasi-relativistic ECP for the heavier nucleus iodine prevents the direct determination of its HFCCs using the same method. Thus, all-electron relativistic calculations were carried out for systems with iodine atoms. The calculations utilized the large QZ4P STO basis set, the PBEPBE GGA functional\(^\text{15a–c}\) as well as the scalar-relativistic ZORA formalism. In quasi-relativistic calculations, large-core ECPs were used on iodine; test calculations showed that a virtually same set of HFCCs was obtained when small-core ECPs were utilized.
All geometry optimizations were done with the Turbomole 5.71\textsuperscript{19} and Gaussian 03\textsuperscript{20} program packages. The hyperfine coupling constant calculations were done with the Gaussian 03\textsuperscript{20} (non-relativistic and quasi-relativistic) and ADF 2004.01\textsuperscript{21} program packages (fully relativistic). Population analyses were done using the NBO 3.1 program as implemented in Gaussian03.\textsuperscript{22} Spectral simulations were carried out using XEMR\textsuperscript{23} and WINEPR SimFonia.\textsuperscript{24}

Results and Discussion

**Gallium Monopnictogen Diazabutadiene Radicals.** Calculations were carried out on the gallium-pnictogen diazabutadiene model systems \{(t-Bu-DAB)Ga(I)[Pn(SiH\textsubscript{3})\textsubscript{2}]\}• 3b–5b. As a whole, the calculated bond angles and bond lengths are in good agreement with the values determined by X-ray crystallography for the parent systems \{(t-Bu-DAB)Ga(I)[Pn(SiMe\textsubscript{3})\textsubscript{2}]\}• (3a–5a),\textsuperscript{13} taking into account the different electronic and steric effects of the SiH\textsubscript{3} groups compared to SiMe\textsubscript{3}. Pertinent calculated and experimental metrical parameters are compiled in Table 1. The geometry-optimized structures of 3b–5b contain a planar DAB ligand chelated to a gallium atom which displays a tetrahedral coordination environment. In the case of 3b, the Pn(SiH\textsubscript{3})\textsubscript{2} nitrogen atom is trigonal planar; for the heavier-pnictogen analogs 4b and 5c, pyramidal geometries are observed about the group 15 atoms, with the sum of the bond angles totaling 307.1° and 297.9° for phosphorus and arsenic, respectively. This result is consistent with the experimentally determined solid-state structures of 3a–5a.\textsuperscript{13} The predicted Ga–Pn (Pn = P, As) bond lengths are consistently slightly longer (0.05 Å) than
the experimentally determined distances; conversely, the calculated Ga–I bonds in 4b and 5b are shorter (0.03 Å) than the actual bond lengths of 4a–5a. These can be attributed to the different steric and electronic effects of the SiMe$_3$ substituents as compared to the SiH$_3$ units of the model systems; the discrepancies in Ga–I bond lengths arise also from the quasi-relativistic nature of the calculations. The lesser steric bulk of the SiH$_3$ groups (cf. SiMe$_3$) is also the reason for the large variance between the calculated and experimental sums of bond angles for the pnictogen atoms (see Table 1). For 3a/3b, where the pnictogen atom (N) is trigonal planar, better agreement between the experimental and calculated structural parameters is observed, as expected.

Despite the difference between the geometry of 3 and 4/5, the SOMOs of 3b–5b were found to be essentially identical which in agreement with the $\pi$-radical nature of the systems. In all three instances, the SOMO is localized on the DAB ligand and is composed primarily of nitrogen and carbon $p$-orbitals. The SOMOs have no $s$-contribution from the central metal or DAB hydrogen atoms, and only very small contributions from the pnictogen and iodine nuclei; the SOMO of 5b is depicted in Figure 1. Mulliken population analyses of 3b–5b were consistent with the appearance of the SOMOs, indicating that the unpaired electron in each molecule is equally delocalized over the nitrogen and carbon atoms of the DAB backbone (a list of spin densities is included as Supporting Information).

Since all complexes 3–5 contain a DAB anion radical 2, the formal oxidation state on the electropositive gallium centers is $+\text{III}$ in all cases. Further analysis using the natural population analysis (NPA) method provides a deeper understanding of the bonding between the gallium and pnictogen/iodine nuclei, and facilitates a comparison
with the data reported for the related spirocyclic monoradical \{Ga(R-DAB)\}_2^\cdot\textsuperscript{25} the numerical results of these are summarized in Table 2.

As expected, the Ga atom in \textit{3b–5b} bears a large positive charge which decreases as the Ga–Pn bond becomes more covalent \textit{i.e.} upon descending group 15. The DAB anion radical contains a near-constant negative charge of 0.50 electrons; the negative charge is more strongly concentrated on the pnictogen atom on the Pn(SiH\textsubscript{3}) group. This is reminiscent of what has been observed for the spirocyclic radical \{Ga(R-DAB)\}_2^\cdot which displays a strong concentration of negative charge on the DAB ring system that has no spin density.\textsuperscript{25} The calculated Wiberg bond indices (see Table 2) for \textit{3b} show only a partial bonding character (0.35–0.45) between gallium and the surrounding nitrogen atoms, indicating a strong donor-acceptor character in these bonds. For the heavier-pnictogen compounds \textit{4b} and \textit{5b}, the Wiberg index for the Ga–Pn bond approaches unity as the covalent contribution within the bond increases; the Ga–I bond is calculated to be almost purely covalent (Wiberg index 0.90) for all systems \textit{3b–5b}.

The hyperfine coupling constants (HFCCs) for \textit{3b–5b} were calculated using both fully relativistic and quasi-relativistic methods and are compiled in Table 3. The HFCCs do not vary significantly from \textit{3b} to \textit{4b} to \textit{5b}, other than that the higher symmetry of \textit{3b} due to the trigonal planar geometry of the pnictogen atom renders the \textsuperscript{1}H and \textsuperscript{14}N atoms of the DAB ligand symmetry-equivalent. In consequence, HFCCs to two equivalent \textsuperscript{1}H and two equivalent \textsuperscript{14}N atoms are predicted for \textit{3b}, while slight inequivalences are expected for the two \textsuperscript{1}H and two \textsuperscript{14}N nuclei of \textit{4b} and \textit{5b}. Relatively small HFCCs have been calculated to the pnictogen atom of \textit{3b} (0.5 G), \textit{4b} (6.4 G), and \textit{5b} (3.5 G); the
values of these couplings vary with the relative magnitudes of the gyromagnetic ratios of $^{14}\text{N}$, $^{31}\text{P}$, and $^{75}\text{As}$, as is expected.\textsuperscript{2}

The average absolute value of the $^{69}\text{Ga}$ HFCC is 19.9 G using quasi-relativistic methods; this decreases to 14.4 G when a full relativistic ZORA method is applied. Using the quasi-relativistic method, it is not possible to calculate the $^{127}\text{I}$ HFCCs since an effective core potential is used to model the inner electrons of the iodine atom, which naturally results in near-zero values for this coupling constant. However, fully relativistic calculations have indicated that a significant $^{127}\text{I}$ HFCC should be observed with an average expected magnitude of 17.6 G in $\text{3b--5b}$; in contrast, the HFCC to iodine nuclei was assumed to be negligible in the published analysis.\textsuperscript{13}

The $^1\text{H}$ HFCCs are not expected to change significantly from $\text{3b--5b}$, nor is there a marked difference between the two inequivalent $^1\text{H}$ couplings in either $\text{4b}$ or $\text{5b}$. The range of predicted $^1\text{H}$ HFCCs (absolute values vary between 5.4–6.0 G) is consistent with those observed in other DAB radicals,\textsuperscript{9,26} and with those calculated for the related group 13-DAB radicals described elsewhere.\textsuperscript{14} Both computational methods yielded similar values of these $^1\text{H}$ hyperfine coupling constants, with differences of no more than 0.3 G observed. The $^{14}\text{N}$ HFCCs are similar in magnitude to the $^1\text{H}$ couplings but show even less variation throughout systems $\text{3b--5b}$, with values of either 4.6 G or 4.7 G predicted in all instances using quasi-relativistic calculations. These values are in agreement with the $^{14}\text{N}$ HFCCs determined for the DAB nitrogen atoms of a variety of other paramagnetic DAB complexes.\textsuperscript{9,26} The implementation of fully-relativistic method appears to result in a significant (1 G) decrease in the $^{14}\text{N}$ HFCCs; however, this is caused by the use of
different functionals (hybrid PBE0 vs. GGA PBEPBE) and basis sets (TZVP vs. QZ4P) for these calculations.  

Though precise values of the HFCCs have not been extracted from the experimental EPR spectra of 3a–5a, some conclusions can be drawn regarding the accuracy of the trends in the calculated HFCCs. First, the calculated HFCCs are quite similar for 3b–5b; in consequence, the width of their EPR spectra should be very similar as well. This is borne out in the experimental spectra, where a deviation in width of no more than 10 G is observed from 3a–5a.  

Second, the experimental spectra are not dominated by the (1:1:1:1) four-line pattern that would be expected if the only large coupling present were to the gallium atom, as had been suggested previously. The appearance of the experimental spectra is thus believed to be influenced equally by the hyperfine couplings to 127I and 69,71Ga, all of which are high-spin nuclei and, thus, have a significant effect on the overall shape of the spectra. In fact, the EPR spectra show a partially concealed sextet pattern due to a large, approximately 20 G, coupling to the spin-5/2 nucleus, 127I.

In light of our recent success in accurately calculating EPR HFCCs for a series of related DAB complexes, attempts were made to simulate the experimental EPR spectra of 3a–5a using the HFCCs that had been calculated for the model systems 3b–5b. Though the calculated values do not accurately reproduce the experimental spectra, optimization of the HFCCs using iterative methods yielded excellent simulations. The experimental and simulated EPR spectra of 3a and 4a are shown in Figures 2 and 3, respectively; optimized HFCCs are included in Table 3. The spectral simulation for 3a was created using identical isotropic g-values for both the 69Ga and 71Ga isotopomers, and is a near-perfect match with the experimental spectrum. The slight differences
between the left- and right-hand sides of the spectrum (i.e. the lack of perfect inversion symmetry in the experimental spectrum with respect to signal intensities) is believed to be caused by higher-order hyperfine interactions due to the presence of large HFCCs to the heavy nuclei $^{69,71}$Ga and $^{127}$I. To accurately model the experimental spectrum, a third-order perturbation theory based Hamiltonian was utilized when creating the spectral simulation. The HFCCs used to create the simulation are very close to the calculated values for the model system 3b, with the optimized $^1$H, $^{14}$N, and $^{127}$I HFCCs slightly larger than the calculated values, while excellent agreement between the optimized (14.9 G) and calculated (−15.4 G) $^{69}$Ga HFCCs is observed.

Similarly, the simulated EPR spectrum of 4a is in good agreement with the experimental spectrum; this simulation was also created using equivalent isotropic g-values for the $^{69,71}$Ga isotopomers and a third-order perturbation theory based Hamiltonian. As was seen for 3a, the optimized $^1$H, $^{14}$N and $^{127}$I HFCCs used to create the spectral simulation of 4a are slightly larger than the predicted HFCCs to the model system 4b; conversely, the optimum $^{69}$Ga HFCC is 2.6 G smaller than the calculated value. While perfect agreement between the optimized and calculated $^{31}$P HFCCs is observed, it must be noted that the assignment of this HFCC to the pnictogen atom phosphorus rather than to one of the DAB hydrogen atoms is somewhat arbitrary since both $^1$H and $^{31}$P are spin-1/2 nuclei with a relative natural abundance of 100%. However, a $^1$H HFCC of 7 G seems somewhat improbable when this value is compared to the $^1$H couplings present in other related paramagnetic group 13-DAB complexes, lending support to the assignment of this coupling to the phosphorus atom. Though intuitively the EPR spectrum of 3a would be expected to consist of more lines than that of 4a due to the
presence of the higher-spin group 15 element nitrogen ($^{14}\text{N}, I = 1$; c.f. $^{31}\text{P}, I = 1/2$), the $^{14}\text{N}$ HFCC is so small (1.03 G) compared to the experimental linewidth (2 G) that it remains unresolved and does not produce additional spectral lines. In consequence, the EPR spectrum of 3a actually contains fewer lines (32) than does that of 4a (36). It is also noted that the optimized HFCCs for both 3a and 4a include couplings which are approximately multiples of each other (see Table 3): this accidental degeneracy of lines makes the spectra deceptively simple and complicates extraction of accurate HFCCs solely by means of spectral simulations.²⁹

The experimental EPR spectrum of the arsenic analogue 5a is not as well resolved as those of 3a and 4a due to hyperfine coupling of the unpaired electron with the spin-3/2 nucleus $^{75}\text{As}$. Because of the resultant line-broadening, it is more difficult to ascertain whether a given spectral simulation is a definitive match. However, a reasonable simulation of this spectrum (not shown) can be obtained by including similar HFCCs to those used to create the simulation of 4a, substituting a $^{75}\text{As}$ HFCC of 3.0 G in place of the $^{31}\text{P}$ HFCC of 4a.

Compounds 3a–5a have also been characterized by ENDOR spectroscopy,¹³ which has indicated the presence of $^1\text{H}$ HFCCs of approximately 1.4 G in all cases: no larger $^1\text{H}$ couplings were detected. No explanation for this observation can be made at this time, as both the calculated HFCCs and spectral simulations support the presence of much larger $^1\text{H}$ HFCCs to the hydrogen atoms of the DAB backbone, as does previous work in this area.⁹¹⁴²⁶ In a study of some related DAB-containing radicals,⁸ it has proven difficult to detect continuous wave ENDOR signals which result from the DAB-backbone hydrogen atoms due to the broad nature of the resonances: it is possible that a similar effect occurs
for 3a–5a. However, the origin of the 1.4 G ¹H couplings remains uncertain: they are not thought to arise due to the tert-butyl substituents of the DAB ligands, as the HFCCs to those methyl protons were calculated to be an order of magnitude smaller. Further research is needed in order to explain the conflicting results.²⁰

In the original spectral analyses,¹³ the possibility for a significant iodine coupling in 3a–5a was discounted based primarily on results obtained for the related systems {(R-DAB)AlI₂}⁺ and {(R-DAB)GaI₂}⁺. For these species, experimental evidence – or rather lack thereof – has suggested that only minor (< 1 G) ¹²₇I couplings are present.⁸,¹² However, an alternative interpretation of the poorly resolved EPR spectrum of {(‘Bu-DAB)GaI₂}⁺ has also been expounded which employs an approximately 8 G HFCC to iodine.¹⁰ In addition, EPR studies on the isoelectronic zinc systems {[{(‘Bu-DAB)ZnX₂]−}} (X = Cl, Br) which produce well-resolved spectra, have demonstrated that the bromine derivative displays an approximately 9 G coupling to the heavy halogen ⁷⁹,⁸¹Br.³¹ In light of these experimental data, recent computational results¹⁴, and the present analysis, it appears highly unlikely that the group 13 systems {(R-DAB)MI₂}⁺ (M = Al, Ga) would exhibit such small HFCCs to iodine as was suggested previously.⁸,¹²

**Gallium Dipnictogen Diazabutadiene Radicals.** The model systems {(‘Bu-DAB)Ga[PN(SiH₃)₂]₂}⁺ 6b–8b were geometry-optimized and found to be closely structurally related to one another. Each gallium atom displays a tetrahedral coordination sphere; in the case of 6b, a planar geometry is seen at the pnictogen (N) centers, while both the phosphorus and arsenic atoms of 7b and 8b, respectively, are pyramidal. This is reminiscent of the structural trends described above for 3–5. Both {(‘Bu-
DAB)Ga[Pn(SiMe$_3$)$_2$]$_2$} (7a, Pn = P; 8a, Pn = As) have been characterized in the solid state by X-ray crystallography, and found to be isostructural with each other.$^{13}$ The observed pyramidal geometries of the P and As atoms are consistent with the calculated structures of the model systems. Key metrical parameters for the calculated model systems are compiled in Table 4 alongside the corresponding empirical data for 7a and 8a. As a whole, the calculated and experimental bond lengths and bond angles are in good agreement with each other; the biggest discrepancies are again observed for the bond angles involving the pnictogen atoms (see Table 4) which arise from the different steric effects of the SiH$_3$ units in 7b and 8b compared to the SiMe$_3$ groups of 7a and 8a.

The singly occupied molecular orbitals of 6b–8b were calculated to be essentially identical to those of 3b–5b; the SOMO of 8b is shown in Figure 4. Mulliken population analyses indicated that the spin density in 6b–8b is equally distributed over the carbon and nitrogen atoms of the DAB ligand backbone, and only a small contribution from the pnictogen atoms as well as from the gallium nuclei was observed. NAO population analyses displayed similar trends that were observed also for 3b–5b (see Table 2); the Ga–N bonds in 6b are more ionic in nature whereas the Ga–P and Ga–As bonds in 7b and 8b, respectively, are close to purely covalent.

Hyperfine coupling constants were calculated for 6b–8b, though the parent nitrogen analogue 6a has not yet been isolated, and are compiled in Table 5. Since these dipnictogen species do not contain the heavy element iodine, only non-relativistic calculations were carried out. The higher (C$_2$) symmetry of 6–8 as compared to 4 and 5 renders the hydrogen and nitrogen nuclei on the DAB backbone equivalent by symmetry: in consequence, couplings to two equivalent $^1$H and two equivalent $^{14}$N HFCC are
anticipated. The calculated values of the $^1$H HFCC for $6b$–$8b$ were found to not differ significantly from the predicted values for $3b$–$5b$ or experimental values obtained for related DAB-centered radicals. However, they differ considerably from the values determined in the previously published simulation (vide supra). Similarly, little variation is observed between the calculated $^{14}$N HFCCs which range from 4.4 G ($6b$) to 4.8 G ($8b$); these values are again consistent with previous experimental and calculated results. The predicted $^{69}$Ga coupling constants for $6b$–$8b$ are consistently larger than those calculated for the monopnictogen systems $3b$–$5b$ due to minor changes in spin density which result from the substitution of a second Pn(SiH$_3$)$_2$ (Pn = N, P, As) unit in place of the iodine atoms of $3b$–$5b$. The only significant difference between the calculated HFCCs for $6b$, $7b$, and $8b$ are the values of the pnictogen couplings which are predicted to be largest for phosphorus (9.2 G) and smallest for $^{14}$N (1.9 G) which is consistent with the relative magnitudes of the gyromagnetic ratios of the nuclei. In consequence, any major difference in the appearance of the experimental EPR spectra of $7a$ and $8a$ is expected to result from the differing HFCCs to the phosphorus ($7a$) and arsenic atoms ($8a$), and the different nuclear spins of these two isotopes.

Experimental EPR spectra of both $7a$ and $8a$ have been reported and display near-perfect inversion symmetry, indicating that higher-order hyperfine interactions in these systems are negligible. The EPR spectrum of $8a$ is ca. 30 G wider than that of $7a$, consistent with the smaller calculated HFCC to the gallium atom of $7a$ and the lesser nuclear spin of $^{31}$P compared to $^{75}$As. An attempt to simulate the experimental EPR spectrum of $7a$ using the HFCCs that were calculated for $7b$ yielded a spectrum with the correct general features; further optimization of the HFCCs gave an excellent simulation
which is shown in Figure 5. The HFCCs used to create this simulation (Table 5) are very similar to the calculated HFCCs with the most significant deviation observed for the $^{31}$P HFCC, which was adjusted by 2.5 G in order to optimize the spectral simulation. Curiously, the calculated $^1$H HFCC is nearly identical to the optimized $^{14}$N HFCC, and vice versa. Though this suggests that these HFCCs have been assigned incorrectly, the agreement between the simulated and experimental EPR spectra is poor when the optimized $^1$H and $^{14}$N HFCCs are interchanged. The apparently too-low and too-high values of the calculated $^{14}$N and $^1$H HFCCs, respectively, has been observed in previous work with related paramagnetic group 13-DAB complexes. This systematic error arises from the apparent tendency of this functional basis set combination to underestimate couplings which arise due to the spin density of the unpaired electron, such as the DAB-nitrogen atom couplings, while overestimating those which arise due to spin polarization effects, as do the $^1$H couplings.

An excellent simulation (Figure 6) was also obtained for the arsenic-containing radical $8a$ by making slight adjustments to the calculated HFCCs of the model system $8b$. As was seen for $7a/7b$, the largest difference between the calculated and optimized couplings occurs for the pnictogen ($^{75}$As) HFCC; in the case of $8a/8b$, the calculated $^{69}$Ga HFCC also deviates slightly (2.9 G) from the optimal value. Considering the near-perfect agreement between the simulated and experimental EPR spectra of $7a$ and $8a$, the optimized HFCCs derived from these DFT calculations can be regarded as the actual HFCCs present in these systems. Given that both gallium and arsenic are forth-period elements, the fact that their calculated and actual HFCCs differ by less than 3 G is considered an excellent agreement.
Conclusions

The electronic structures and spin densities of the paramagnetic gallium(III)-

pnictogen complexes \{('Bu-DAB)Ga(I)[Pn(SiH₃)₂]\}' and \{('Bu-DAB)Ga[Pn(SiH₃)₂]₂\}'

(Pn = N, P, As) were studied using density functional methods. The calculations
demonstrate that these \(\pi\)-radicals have qualitatively similar SOMOs with the unpaired
electron residing on the DAB backbone. The HFCCs calculated for the model systems
were used to simulate the EPR spectra of the closely related radicals \{('Bu-

DAB)Ga(I)[Pn(SiMe₃)₂]\}' (Pn = N, P, As) and \{('Bu-DAB)Ga[Pn(SiMe₃)₂]₂\}' (Pn = P,

As) with excellent accuracy. As shown, the complexity in these EPR spectra does not
arise from different isotropic g-values for the \(^{69}\text{Ga}\) and \(^{71}\text{Ga}\) isotopomers as previously
suggested, but from the presence of higher-order splitting effects and HFCCs which are
approximately multiples of each other. The results also demonstrate the occurrence of
significant hyperfine coupling to the \(^{127}\text{I}\) nucleus in \{('Bu-DAB)Ga(I)[Pn(SiMe₃)₂]\}'
systems as well as confirm the presence of large HFCCs to the \(^1\text{H}\) atoms in the DAB
backbones of both mono- and dipnictogen species. Taken in concert with some previous
experimental work, the spectral analyses described herein suggest that the published
experimental EPR results of some related paramagnetic group 13-DAB complexes might
have been misinterpreted. Detailed theoretical investigations of these systems are
currently being pursued in order to resolve this controversy.

Acknowledgement. The authors gratefully acknowledge Helsingin Sanomain 100-
vuotissäätiö (H. M. T.) and the University of Calgary (A. F. A.) for financial support.
Supporting Information Available: XYZ-coordinates and spin densities of compounds 3b–8b.

This material is available free of charge via the Internet at http://pubs.acs.org.

References and Notes

1. For a recent review, see: Power, P. P. Chem. Rev. 2003, 103, 789.


    2002, 1196.
    3844.
    2005, 44, 2098.
    Perdew, J. P.; Burke, K.; Ernzerhof, M., Phys. Rev. Lett. 1997, 78, 1396. (c) Perdew, 
16. The basis sets were used as they are referenced in the Turbomole 5.7.1 internal basis 
    set library. These basis sets can be downloaded from: ftp://ftp.chemie.uni- 
17. (a) Adamo, C.; Barone, V. J. Chem. Phys. 1999, 110, 6158. (b) C Adamo, C.; Barone, 
    (Theochem) 1999, 493, 145. (d) Adamo, C.; Barone, V. Theor. Chem. Acc. 2000, 105, 
    169.
18. (a) Hermosilla, L.; Calle, P.; Garcia de la Vega, J. M.; Sieiro, C. J. Phys. Chem. A 
22. The population analyses have been done according to the Weinhold-Reed partitioning scheme. For a review, see: Reed, A. F.; Curtiss, L. A.; Weinhold, F. Chem. Rev. 1988, 88, 899.

23. Eloranta, J. XEMR version 0.7, University of Jyväskylä, Finland.


27. This has been confirmed by carrying out non-relativistic calculations for 3b–5b using the PBEPBE GGA functional and STO basis sets.

28. It should be noted that very minor distortions of the line-shape of the experimental EPR spectra occurred through digitization of the spectra for compatibility with the selected EPR software. For high quality versions of the experimental spectra, see Reference 13.

29. The same is also true for the HFCCs in radicals 6a–8a.

30. As originally suggested by our referee, it would be highly enlightening to synthesize DAB analogs of 3–5 in which the backbone hydrogen atoms have been replaced by deuterium, as this would unambiguously resolve the question of whether there is a large HFCC to the backbone ¹H nuclei.

### Table 1. Experimental and Calculated Metrical Parameters for 3–5

<table>
<thead>
<tr>
<th>Parameter</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Calc.(^a)</td>
<td>Exptl.(^b)</td>
<td>Calc.(^a)</td>
</tr>
<tr>
<td>r GaN</td>
<td>1.982</td>
<td>1.956</td>
<td>1.981</td>
</tr>
<tr>
<td>r GaI</td>
<td>2.587</td>
<td>2.590</td>
<td>2.563</td>
</tr>
<tr>
<td>r GaPn</td>
<td>1.881</td>
<td>1.868</td>
<td>2.341</td>
</tr>
<tr>
<td>r NC</td>
<td>1.327</td>
<td>1.329</td>
<td>1.328</td>
</tr>
<tr>
<td>r CC</td>
<td>1.398</td>
<td>1.406</td>
<td>1.397</td>
</tr>
<tr>
<td>(\angle) NGaN</td>
<td>84.9</td>
<td>85.9</td>
<td>85.0</td>
</tr>
<tr>
<td>(\angle) IGaPn</td>
<td>119.3</td>
<td>118.0</td>
<td>113.6</td>
</tr>
<tr>
<td>(\angle) GaNC</td>
<td>108.6</td>
<td>108.6</td>
<td>108.6</td>
</tr>
<tr>
<td>(\angle) NCC</td>
<td>118.8</td>
<td>118.3</td>
<td>118.7</td>
</tr>
<tr>
<td>(\Sigma\ \angle) Pn</td>
<td>360.0</td>
<td>360.0</td>
<td>307.1</td>
</tr>
</tbody>
</table>

\(^a\) Values calculated for 3b–5b.

\(^b\) Averaged experimental values of 3a–5a taken from reference 13.
<table>
<thead>
<tr>
<th>Compound</th>
<th>DAB</th>
<th>Ga</th>
<th>I</th>
<th>Pn(SiH₃)₂</th>
<th>WBI</th>
</tr>
</thead>
<tbody>
<tr>
<td>3b Pn = N</td>
<td>−0.67</td>
<td>−0.53</td>
<td>1.56</td>
<td>−0.37</td>
<td>−1.83</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.68</td>
<td>0.33</td>
</tr>
<tr>
<td>4b Pn = P</td>
<td>−0.65</td>
<td>−0.50</td>
<td>1.21</td>
<td>−0.34</td>
<td>−0.69</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.37</td>
<td>0.35</td>
</tr>
<tr>
<td>5b Pn = As</td>
<td>−0.65</td>
<td>−0.50</td>
<td>1.16</td>
<td>−0.35</td>
<td>−0.57</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.32</td>
<td>0.35</td>
</tr>
<tr>
<td>6b Pn = N</td>
<td>−0.67</td>
<td>−0.51</td>
<td>1.98</td>
<td>-</td>
<td>−1.80</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.73</td>
<td>0.32</td>
</tr>
<tr>
<td>7b Pn = P</td>
<td>−0.66</td>
<td>−0.50</td>
<td>1.28</td>
<td>-</td>
<td>−0.70</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.39</td>
<td>0.34</td>
</tr>
<tr>
<td>8b Pn = As</td>
<td>−0.66</td>
<td>−0.51</td>
<td>1.18</td>
<td>-</td>
<td>−0.57</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>−0.34</td>
<td>0.35</td>
</tr>
</tbody>
</table>
Table 3. Experimental and Calculated Hyperfine Coupling Constants [G] for 3–5

<table>
<thead>
<tr>
<th>Nucleus</th>
<th>3 (Pn = $^{14}$N)</th>
<th></th>
<th></th>
<th></th>
<th>Exptl. $^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Calcd. $^a$</td>
<td>Optim. $^b$</td>
<td>Calcd. $^a$</td>
<td>Optim. $^b$</td>
<td>Calcd. $^a$</td>
</tr>
<tr>
<td>$^{69}$Ga</td>
<td>−19.8 (−15.4)</td>
<td>14.90</td>
<td>−19.6 (−13.5)</td>
<td>10.88</td>
<td>−20.4 (−14.2)</td>
</tr>
<tr>
<td>$^{71}$Ga</td>
<td>−25.3 (−19.7)</td>
<td>18.80</td>
<td>−25.1 (−17.3)</td>
<td>13.81</td>
<td>−26.1 (−18.2)</td>
</tr>
<tr>
<td>Pn</td>
<td>0.5 (0.5)</td>
<td>1.03</td>
<td>6.4 (7.0)</td>
<td>7.00</td>
<td>3.5 (4.1)</td>
</tr>
<tr>
<td>$^1$H</td>
<td>−5.7 (−5.6)</td>
<td>6.61</td>
<td>−6.0 (−5.7)</td>
<td>6.50</td>
<td>−6.0 (−5.8)</td>
</tr>
<tr>
<td>$^1$H</td>
<td>−5.7 (−5.6)</td>
<td>6.61</td>
<td>−5.5 (−5.5)</td>
<td>6.52</td>
<td>−5.4 (−5.4)</td>
</tr>
<tr>
<td>$^{14}$N</td>
<td>4.6 (3.5)</td>
<td>5.48</td>
<td>4.7 (3.5)</td>
<td>5.85</td>
<td>4.7 (3.6)</td>
</tr>
<tr>
<td>$^{14}$N</td>
<td>4.6 (3.5)</td>
<td>5.48</td>
<td>4.6 (3.6)</td>
<td>5.24</td>
<td>4.6 (3.6)</td>
</tr>
<tr>
<td>$^{127}$I</td>
<td>0 (16.9)</td>
<td>20.01</td>
<td>0 (17.8)</td>
<td>23.08</td>
<td>0 (18.2)</td>
</tr>
</tbody>
</table>

$^a$ Values calculated for 3b–5b using quasi-relativistic and (relativistic) methods.

$^b$ Optimized values used to create spectral simulation of 3a–5a.

$^c$ Values previously estimated from experimental spectra (taken from Reference 13).
Table 4. Experimental and Calculated Metrical Parameters for 6–8

<table>
<thead>
<tr>
<th>Parameter</th>
<th>6b</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>r GaN</td>
<td>1.989</td>
<td>1.994</td>
<td>1.986</td>
</tr>
<tr>
<td>r GaPn</td>
<td>1.889</td>
<td>2.363</td>
<td>2.343</td>
</tr>
<tr>
<td>r NC</td>
<td>1.326</td>
<td>1.328</td>
<td>1.330</td>
</tr>
<tr>
<td>r CC</td>
<td>1.399</td>
<td>1.396</td>
<td>1.394</td>
</tr>
<tr>
<td>∠ NGaN</td>
<td>84.5</td>
<td>84.3</td>
<td>84.7</td>
</tr>
<tr>
<td>∠ PnGaE</td>
<td>117.9</td>
<td>115.8</td>
<td>115.2</td>
</tr>
<tr>
<td>∠ GaNC</td>
<td>109.0</td>
<td>109.1</td>
<td>108.6</td>
</tr>
<tr>
<td>∠ NCC</td>
<td>118.7</td>
<td>118.8</td>
<td>119.5</td>
</tr>
<tr>
<td>Σ ∠ Pn</td>
<td>359.8</td>
<td>305.0</td>
<td>330.4</td>
</tr>
</tbody>
</table>

*a* Values calculated for 7b–8b.

*b* Averaged experimental values of 7a–8a taken from reference 13.
Table 5. Experimental and Calculated Hyperfine Coupling Constants [G] for 6–8

<table>
<thead>
<tr>
<th>Nucleus</th>
<th>6 (Pn = ^14N)</th>
<th>7 (Pn = ^31P)</th>
<th>8 (Pn = ^75As)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Calcd.(^a)</td>
<td>Calcd.(^a)</td>
<td>Optm.(^b)</td>
</tr>
<tr>
<td>^69Ga</td>
<td>−22.0</td>
<td>−22.6</td>
<td>−22.20</td>
</tr>
<tr>
<td>^71Ga</td>
<td>−28.2</td>
<td>−28.9</td>
<td>−28.20</td>
</tr>
<tr>
<td>Pn</td>
<td>1.9</td>
<td>9.2</td>
<td>11.70</td>
</tr>
<tr>
<td>^1H</td>
<td>−5.9</td>
<td>−5.6</td>
<td>−4.80</td>
</tr>
<tr>
<td>^14N</td>
<td>4.4</td>
<td>4.7</td>
<td>5.80</td>
</tr>
</tbody>
</table>

\(^a\) Values calculated for model systems 6b–8b.

\(^b\) Optimized values used to create spectral simulations of 7a and 8a.
Figure Captions

Figure 1. SOMO of \{('Bu-DAB)Ga(I)[As(SiH₃)₂]⁻\}’ (5b) (isosurface value = ±0.05).

Figure 2. Experimental (top) and simulated (bottom, 1w. 2.6 G) EPR spectrum of 3a. (Experimental spectrum reproduced with permission from Reference 13. Copyright ACS 2005.)

Figure 3. Experimental (top) and simulated (bottom, 1w. 2.0 G) EPR spectrum of 4a. (Experimental spectrum reproduced with permission from Reference 13. Copyright ACS 2005.)

Figure 4. SOMO of \{('Bu-DAB)Ga[As(SiH₃)₂]₂\}’ (8b) (isosurface value = ±0.05).

Figure 5. Experimental (top) and simulated (bottom, 1w. 2.4 G) EPR spectrum of 7a. (Experimental spectrum reproduced with permission from Reference 13. Copyright ACS 2005.)

Figure 6. Experimental (top) and simulated (bottom, 1w. 1.9 G) EPR spectrum of 8a. (Experimental spectrum reproduced with permission from Reference 13. Copyright ACS 2005.)
Figure 2
Figure 3
Figure 5
Figure 6