

A Neutral Beryllium(I) Radical

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Abstract: The reduction of a cyclic alkyl(amino)carbene (CAAC)-stabilized organoberyllium chloride yields the first neutral beryllium radical, which was characterized by EPR, IR, and UV/Vis spectroscopy, X-ray crystallography, and DFT calculations.

While group 2 chemistry is mainly dictated by the naturally occurring +2 oxidation state of its elements, the last two decades have seen the emergence of a growing number of low-oxidation-state alkaline earth metal compounds. Since the landmark synthesis of the first dinuclear Mg^I complexes, including I (Figure 1),^[1] these compounds have been successfully applied as highly selective reducing agents for the activation of small molecules^[2] and the synthesis of new



Figure 1. Selection of low-oxidation-state molecular group 2 complexes.



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homo- and heteronuclear metal-metal bonds,^[3] culminating most recently in the isolation of the first molecular Mg⁰ species, complex II.^[4]

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Low-valent beryllium complexes long remained confined to the computational realm due to their very high toxicity. Recent years, however, have seen a renewed interest in beryllium coordination chemistry in the areas of organometallic, pure inorganic, and bioinorganic chemistry.^[5] Although the viability of Be^I-Be^I-bonded species has been predicted,^[6] and the Be₂ dimer has been observed spectroscopically,^[7] the low Be-Be bond enthalpy makes mononuclear Be⁰ compounds more accessible targets.^[8] In 2016 our group reported the first Be⁰ compound, complex **III**, which owes its stability to strong three-center-two-electron π backbonding from the Be^0 atom in its $2s^02p^2$ electronic configuration to the neutral cyclic alkyl(amino)carbene (CAAC) ligands,^[9] and this has since been used as a reducing agent to synthesize the first carbene bismuthinidene complex.^[10] Although beryllium radicals have been postulated as intermediates in reduction reactions resulting in ligand activation,^[11] the first isolable Be^I radical cation, IV, was only reported in 2020 from the oneelectron oxidation of an analogue of III with 2,2,6,6-tetramethylpiperidin-1-oxyl.^[12] Calculations showed that the bonding in IV is similar to that in III, with two neutral CAAC ligands stabilizing a Be^I cation in its excited 2s⁰2p¹ electronic configuration through donor-acceptor interactions, and that the spin density is delocalized over the entire N-C-Be-C-N framework, with 38% located at the beryllium center. Furthermore, Paparo and Jones succeeded in isolating the first neutral Be^I complexes, such as V, which present covalent Be^I-Al^I bonding.^[13] We now report the synthesis and computational analysis of the first structurally characterized neutral beryllium radical, stabilized by both a neutral and a C1-protonated CAAC ligand.

The organoberyllium halide precursors (CAAC)-(CAACH)BeX (CAAC = 1-(2,6-diisopropylphenyl)-3,3,5,5tetramethylpyrrolidin-2-ylidene; CAACH = 1-(2,6-diisopropylphenyl)-3,3,5,5-tetramethylpyrrolidin-2-yl; X = Cl, Br) were synthesized by the addition of L-selectride (Li-[HBsBu₃]) to a 1:1 mixture of CAAC and (CAAC)BeX₂^[9] in toluene at -78°C and isolated as pale orange solids in excellent yields (\geq 82%, Scheme 1a). The reaction likely proceeds via formation of a (CAAC)Be(H)Cl intermediate, followed by coordination of the second CAAC ligand to form (CAAC)₂Be(H)Cl, and finally a 1,2-hydride shift from the beryllium center to the CAAC carbene carbon atom. Such 1,2-hydrogen shifts are common in CAAC-stabilized main group hydrides upon coordination of an additional Lewis base.^[14] A characteristic ¹H singlet around 3.0 ppm and the complex ligand resonance patterns of the ¹H NMR spectra confirmed the protonation of one of the CAAC ligands. The **Communications**



Scheme 1. Synthesis and crystallographically derived solid-state structure of [(CAAC)(CAACH)Be][•] (only the major part of the two flipdisordered Be(CAACH) units shown). Atomic displacement ellipsoids represented at 50% and omitted for the ligand periphery. Hydrogen atoms omitted, except for H21. Selected bond lengths (Å) and angles (°): N1-C1 1.321(2), C1-Be1 1.661(7), Be1-C21 1.730(7), C21-N1' 1.548(4); C1-Be1-C21 170.7(3).^[25]

⁹Be NMR shifts of the two complexes appear at 19 and 20 ppm for X = Cl and Br, respectively. In order to place the ⁹Be NMR shifts of (CAAC)(CAACH)BeX in context with comparable CAAC-stabilized tricoordinate beryllium complexes, they are downfield-shifted from those of their (CAAC)BeX₂ precursors at 12.9 and 14.0 ppm, respectively,^[9] slightly upfield-shifted from that of the beryllole (CAAC)- BeC_4Ph_4 at 22.9 ppm,^[15] and similar to that of the diazaborolyl beryllium chloride (CAAC)BeCl(B(NDipCH)₂) at 20 ppm.^[16] Additionally, the complexes were characterized by X-ray crystallographic analyses (see Figures S19 and S21 in the Supporting Information).^[25] Cyclic voltammetry (CV) experiments carried out in difluorobenzene showed a single irreversible reduction wave at $E_{\rm pc} = -1.83$ V versus the ferro- (Fc/Fc^+) cene/ferrocenium couple for (CAAC)-(CAACH)BeCl, whereas (CAAC)(CAACH)BeBr showed two irreversible reduction waves at $E_{\rm pc} = -1.85 \, {\rm V}$ and -2.46 V, which hint at the potential for chemical reduction of both species.

Whereas the reduction of (CAAC)(CAACH)BeBr with a wide range of reducing agents in various solvents at best resulted in partial reduction and the formation of [CAACH]Br as the sole isolable product, the room-temperature reduction of (CAAC)(CAACH)BeCl with lithium sand in diethyl ether over a period of 10 minutes resulted in the formation of the radical species [(CAAC)(CAACH)Be]; which was isolated as a brown-orange crystalline solid in 70% yield (Scheme 1b). While the radical proved stable in the solid state at -30 °C under an argon atmosphere for several weeks, it decomposed within minutes in polar solvents, such as THF and 1,2-difluorobenzene, and within two days in diethyl ether at -30 °C. In aromatic hydrocarbon solvents, such as benzene and toluene, the compound was less soluble but remained stable at room temperature, provided silanized glassware or polyethylene vials were used to avoid its reaction with glassware surface OH groups. Under these conditions [(CAAC)(CAACH)Be] could be heated up to 60 °C before significant decomposition set in.

As expected, [(CAAC)(CAACH)Be]• was NMR-silent but displayed a complex EPR signal centered at $g_{iso} = 2.003$. Simulation provided a hyperfine coupling constant to ⁹Be of 11.6 MHz (4.1 G, Figure 2 a), significantly larger than for **IV** (0.32 G).^[12] Calculations at the UBP86-D3(BJ)/def2SVP level of theory, performed using Gaussian16,^[17] show that the SOMO is mainly delocalized over the Be–C_{CAAC} π bond, with some π -antibonding character on the C–N bond of the CAAC ligand (Figure 2 b), as is the case for most CAAC-stabilized main group radicals.^[18] The calculated spin density at beryllium (23 %) is significantly lower than that calculated for the radical cation **IV** (38 %).^[12,19]



Figure 2. a) Experimental (black) and simulated (red) continuous-wave (CW) X-band EPR spectra of [(CAAC) (CAACH)Be][•] in benzene at room temperature. The simulation parameters are $g_{iso} = 2.003$, $a({}^9Be) = 11.6$ MHz, $a({}^{14}N) = 3.7$ MHz, and $a({}^{1}H) = 6.3$ MHz. b) Plot of spin density of [(CAAC) (CAACH)Be][•] calculated at the UBP86-D3(BJ)/ def2SVP level of theory. Mulliken spin densities: 0.23 (Be1); 0.54 (C1); 0.19 (N1).

The solid-state IR spectrum of $[(CAAC)(CAACH)Be]^{*}$ shows a characteristic band at 2693 cm⁻¹ which calculations attribute to the C–H stretching frequency of the protonated beryllium-bound carbon atom ($\tilde{v}_{calcd} = 2725 \text{ cm}^{-1}$). The UV/ Vis spectrum of $[(CAAC)(CAACH)Be]^{*}$, which had to be recorded in Et₂O in a silanized cuvette to avoid decomposition, shows a broad absorption centered at $\lambda_{max} = 350 \text{ nm}$, spanning over 100 nm at mid-height and extending into the 400–500 nm range, thus accounting for the brown-orange coloration of the radical.^[20] Accordingly, TD-DFT calculations indicate the presence of charge-transfer transitions from the SOMO to low-lying LUMOs in this wavelength window, the one with the largest oscillator strength appearing at 330 nm (UCAM-B3LYP/6-31 ++ G**, see Supporting Information).

Independent of the crystallization conditions [(CAAC)-(CAACH)Be] crystallized in the $P\bar{1}$ space group (see solidstate structure in Scheme 1), with one fully centrosymmetrically disordered molecule per asymmetric unit, which further presents a twofold disorder in the relative *R/S* configuration of the CAACH ligand backbone in a ca. 4:1 ratio.^[21] While the structural data may not, therefore, be discussed in detail, it confirms that the beryllium center displays a near-linear geometry (C1-B1-C21 ca. 171°) and that the Be–C1 bond to the neutral CAAC ligand (ca. 1.66 Å) is significantly shorter than the Be1–C21 single bond to the protonated CAACH ligand (ca. 1.73 Å). The partial double bond character of C1–Be1 is also supported by Mayer bond order calculations, the value of which (1.10) is significantly larger than that of Be1–C21 (0.79).

The energy decomposition analysis in combination with the natural orbitals for chemical valence method (EDA-NOCV), as implemented in ADF 2019,^[22] was applied to [(CAAC')(CAAC'H)Be][•] (truncated model with Me and *i*Pr groups replaced by hydrogen atoms) in order to investigate its bonding situation. The results were obtained at the BP86-D3(BJ)/TZV2P level of theory. The quantitative results for three distinct decomposition schemes are shown in Table S1 in the Supporting Information. These were based on [(CAAC'H)Be] and CAAC' as interacting fragments and varied depending on the electronic configuration and multiplicity of the fragments. The interaction between [(CAAC'H)-Be] in its first excited doublet configuration, where the radical occupies a p₁ orbital of Be, and a ground-state singlet CAAC' resulted in the lowest orbital interaction term ΔE_{orb} . As this is a useful criterion for discerning the best bonding description in terms of interacting fragments,^[15,23] we conclude that donor-acceptor interactions are at play in the stabilization of the [(CAAC')(CAAC'H)Be] neutral radical.

As shown in Table S1 in the Supporting Information, essentially half of the attraction (50.3%) between the [(CAAC'H)Be] and CAAC' fragments is due to the covalent contribution ΔE_{orb} . The dispersion contribution (ΔE_{disp}) accounts for merely 2.6% and the electrostatic attraction ΔE_{elstat} is responsible for the remaining 47.1%. These results are comparable to those observed for the paramagnetic beryllium radical cation **IV**.^[12] The breakdown of ΔE_{orb} into pairwise orbital interactions (Table SX and Figure 3) shows that the strongest contribution comes from the (CAAC'H)Be \rightarrow CAAC' π backdonation from the Be radical



Figure 3. Plots of the deformation densities $\Delta \rho_{1\alpha\beta}$ and $\Delta \rho_{2\alpha}$ of the main pairwise contributions associated with the orbital interaction term ΔE_{orb} in the truncated model [(CAAC')(CAAC'H)Be][•]. Interacting fragments: [(CAAC'H)Be][•] (radical in the p_⊥ orbital of Be) and CAAC' (closed-shell singlet ground state). Values in parentheses are the percentage of the pairwise orbital interaction with respect to the total ΔE_{orb} contribution. The $\nu_{\pm k}$ values correspond to the eigenvalues of the complementary eigenfunctions (ψ_{-k} , ψ_{+k}) in the NOCV representation. Isovalues: 0.003. Charge flows from red to blue.

into the vacant π orbital of the CAAC' ligand ($\Delta E_{orb(2\alpha)} = -56.5 \text{ kcal mol}^{-1}$, see Figure 3 for the corresponding deformation density). This contribution is slightly stronger than that obtained for the radical cation $\mathbf{IV}^{[12]}$ and significantly weaker than that of the neutral Be⁰ species \mathbf{III} , ^[9] the latter on account of the half-empty Be p_{\perp} orbital of [(CAAC')-(CAAC'H)Be], which is a weaker donor in comparison to the doubly occupied Be p_{\perp} orbital of \mathbf{III} . In contrast, the (CAAC'H)Be \leftarrow CAAC' σ donation (see Figure 3 for the corresponding deformation density) is $\Delta E_{orb(1\alpha\beta)} = -43.4 \text{ kcal mol}^{-1}$, weaker than the CAAC \rightarrow Be \leftarrow CAAC σ donation contributions in **III** and **IV**. This is explained by the fact that in this case only one CAAC ligand contributes to the σ donation, while in the previous cases both ligands donate to the central Be atom.

In order to assess the possibility of fluxional hydrogen shifting from CAACH to CAAC in [(CAAC)(CAACH)Be][•] via an intermediate tricoordinate tautomer [(CAAC)₂BeH][•], we also examined the latter computationally (Figure 4). At



Figure 4. Canonical Kohn–Sham molecular orbitals of [(CAAC)-(CAACH)Be][•] and its putative tautomer [(CAAC)₂BeH][•] at the UBP86-D3(BJ)/def2-SVP level of theory.

the UBP86-D3(BJ)/def2SVP level of theory $[(CAAC)_2BeH]^{\cdot}$ lies 10.5 kcalmol⁻¹ higher in energy than $[(CAAC)_{\cdot}(CAACH)Be]^{\cdot}$. Its SOMO is π -delocalized along the two CAAC ligands and the central Be atom and features two nodal planes at the C–N bonds. In contrast, the LUMO of $[(CAAC)_2BeH]^{\cdot}$, which is mostly located at the endocyclic C– N bonds, is stabilized by 0.56 eV compared to that of $[(CAAC)(CAACH)Be]^{\cdot}$. This leads to a decrease in the SOMO–LUMO (SL) gap of $[(CAAC)_2BeH]^{\cdot}$ to merely 0.75 eV, less than half the SL gap of $[(CAAC)(CAACH)Be]^{\cdot}$ (1.61 eV). These results show that a doubly CAAC-stabilized BeH radical is not energetically accessible. Since the nature of the Lewis base (L) strongly influences the electronic and structural features of main group compounds,^[24] a theoretical investigation of various $[L_2BeH]^{\cdot}$ radicals, aiming at the identification of potential synthetic targets, is currently under investigation in our group.

To summarize, we have synthesized and structurally characterized a stable neutral Be^I radical, the first example of an isolable neutral s-block radical with significant spin density located at the metal center. We are currently investigating the reactivity of this species and will report our findings in due course.

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Conflict of Interest

The authors declare no conflict of interest.

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