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A Functionalized Ge\(_3\)-compound with a Dual Character of the Central Germanium Atom

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(cAAC)\(\text{Ge(GeL)}\)\(_2\) (1) (cAAC = cyclic alkyl(amino) carbene; \(L = \text{PhC}(\text{tBuN})\)\(_2\)), a functionalized Ge\(_3\)-compound was prepared. Quantum mechanical studies on 1 show a reciprocal relationship between the electronic state of the central Ge atom and its reactivity towards proton, viz. tetravalent Ge(II) in terms of bonding and divalent Ge(0) in terms of reactivity. Thus the central Ge atom can be considered having a hidden but highly reactive lone pair of electrons. However, the terminal Ge atoms can be considered as divalent Ge(I) with an active lone pair of electrons.

The formation of multiple bonds between carbon and heavier group 14 elements (Si, Ge, and Sn) has been an attractive research topic for decades.\(^1\) Stable germenes (>C=Ge<) can be accessed by protecting both carbon and germanium centers using sterically demanding ligands (Scheme 1). The approaches to germenes A\(^2\) and B\(^3\) are assumed via transient carbenes, while B is best described as an adduct of cyclopentadienylidene and germylene. Compound C\(^4\) is prepared by treating Mes\(_2\)Ge(F)\(_4\)C(H)\(_2\) with tBuLi, and D\(^5\) is probably produced under insertion of CS\(_2\) into a Ge=Ge double bond. A germylone\(^6\),\(^a,b\) with composition (cAAC)\(_2\)Ge(0) shows an interesting biradicaloid character.\(^6\) The NBO study demonstrates that the principal orbitals of (cAAC)\(_2\)Ge are a lone pair on Ge and a three-center C-Ge-C \(\pi\)-type orbital where 43% is at Ge atom and 28.5% at each carbene carbon, different from the C-\(\rightarrow\)Ge donor-acceptor bond present in the cyclic germylone.\(^7\) This is mainly attributed to the singlet spin ground state and the smaller HOMO-LUMO energy gap of cAAC, when compared with that of NHIC (N-heterocyclic carbene).\(^8\) This finding enriches the C-Ge species, and a similar bonding situation is theoretically reported for the Si analogues.\(^9\) Although (cAAC)\(_2\)Ge can be prepared easily by using GeCl\(_2\)(dioxane) as the Ge source,\(^6\) we are curious to study the reaction behavior when a substituted germanium precursor is employed. Herein, we report a novel Ge\(_3\)-compound (1) which shows for the first time the direct formation of a Ge=C bond using a stable cAAC. Moreover, theoretical studies predict unusual dual character of the central Ge atom.

The reaction of LGeCl (L= PhC(\text{tBuN})\(_2\))\(^10\) cAAC and K\(_8\) in a molar ratio of 1:0.5:1.2 in THF at -78 \(^\circ\)C for 1 h gives compound (cAAC)\(\text{Ge(GeL)}\)\(_2\) (1) as red crystals in 35 % yield (based on Ge). (Scheme 2) The molar ratio used above is tested to be necessary for the formation of 1. Interestingly, stirring the reaction overnight at room temperature afforded a mixture of 1, dark green plates of (cAAC)\(_2\)Ge (2) and a small amount of colorless crystals of (cAAC)\(_3\)O.\(^11\) 2 is also slowly generated from the mother liquid of 1 (Scheme 2). However, when NHIC (1,3-bis(isopropyl)imidazol-2-ylidene) is used instead of cAAC, only LGe-Gel\(^10\) is produced.

The solution of 1 is highly air sensitive, but it is stable in the solid state, even when exposed to air for two days. 1 decomposes

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above 183°C. I is soluble in THF and toluene, but sparingly soluble in n-hexane. The 1H NMR spectrum shows the resonances of the protons for iPr-H (septet at 3.41 ppm) and CH2 (singlet at 1.42 ppm) of cAAC. While the amide ligand exhibits two tBu-H resonances (1.34 and 1.32 ppm), indicating the asymmetric structure of the molecule. In the 13C NMR spectrum the resonance for the carbene carbon (C-Ge) is observed at 219.4 ppm which shifts upfield when compared with the free cAAC (304.2 ppm), and the conjugated Ar,Ge=C(R)C≡C(R)C=GeAr (Ar = 2,3,4-trimethyl-6-tBu-phenyl; R = nBu and C8H5).12 UV-visible spectrum of I recorded in C6D6 shows a strong absorption band at 490 nm.

**Fig. 1** Molecular structure of I. Selected bond lengths (Å) and angles (°): Ge1-C1 1.881(3), Ge1-Ge2 2.4746(7), Ge1-Ge3 2.4929(6), C1-N1 1.375(5), Ge2-N2 2.015(3), Ge2-N3 2.029(3); Ge2-Ge1-Ge3 107.65(2), C1-Ge1-Ge2 128.95(11), C1-Ge1-Ge3 122.68(11), N1-C1-C2 107.3(3), N1-C1-Ge1 127.5(2), C2-C1-Ge1 124.9(3). Sum of angles around C1 and Ge1 are ca. 359.7 and 359.3°, respectively.

The structure of I (Fig. 1) exhibits an asymmetric Ge3 backbone, in which the central Ge1 atom adopts a three coordinate geometry with one N and two Ge atoms. The geometry around the Ge1 atom is nearly planar, which indicates the absence of a stereo active lone pair at Ge1. On the contrary the Ge2 and Ge3 atoms are both pyramidalized due to the presence of a lone pair of electrons at the apex of a distorted trigonal pyramid. The N2-Ge2-N3 and N4-Ge3-N5 planes are nearly perpendicular (ca. 88°). The Ge2-Ge1-Ge3 bond angle is 107.65(2)° which is comparable to those reported for the stable germathene A (115 and 93 ppm), and the conjugated Ar,Ge=C(R)C≡C(R)C=GeAr (Ar = 2,3,4-trimethyl-6-tBu-phenyl; R = nBu and C8H5).12 UV-visible spectrum of I recorded in C6D6 shows a strong absorption band at 490 nm.

The group charge of cAAC is 40.32 e which indicates a net charge flow from the GeL2 fragment to the carbene ligand. This signifies that the π-back donation from Ge1 to C1-N1 π*-MO of cAAC is much stronger than the σ-donation from cAAC to Ge1 of the GeL2 fragment. The Wiberg bond index of the Ge1-C1 bond (1.35) also indicates the partial double bond character.18 Interestingly, the NBO calculations suggest the presence of p-type lone pair (100% p-character) on Ge1 and a sp-hybrid type lone pair on Ge2 (81.5% s-character and 18.5% p-character) and Ge3 (80.0% s-character and 20.0% p-character). Moreover, the occupancy of the lone pair on Ge1 is only 1.02 e whereas on Ge2 and Ge3 are 1.93 e and 1.94 e, respectively. The molecular orbitals and the NBO charge and population analysis suggest that Ge2 and Ge3 can be considered as divalent Ge(I), where each Ge atom utilizes one electron for forming an electron sharing bond with the terminal Ge atoms and one electron for the electron sharing bond with one N-atom of the amidinate ligand. The second N-atom of amidinate ligand donates two electrons to Ge2/Ge3 atom. Thus two electrons of the terminal Ge atoms are retained as a lone pair. On the contrary, the central Ge1 atom utilizes two electrons for electron sharing bonds with the terminal Ge atoms. The vacant in-plane σ-orbital accepts electron from cAAC and the remaining two electrons on the p-orbital of Ge1 is back donated to the empty C1-N1 π*-MO of cAAC. Hence Ge1 atom can be considered as tetravalent Ge(II).
σ-type and π-type fragment orbitals on the Ge14C and cAAC. The NOCV pairs of orbitals do not have proper σ- or π-symmetry.17 The corresponding deformation density plots Δρ1 (ΔE1 = -62.1 kcal/mol) and Δρ2 (ΔE2 = -77.2 kcal/mol) do not show any significant variation of electron density along Ge1-CAAC bond. However, significant accumulation and depletion of electron density above and below the plane of Ge1-C bond is observed. Hence, the bonding situation in the Ge1-C can be considered as formed by two bent donor-acceptor bonds.17 This is also reflected in the low bond dissociation energy (D∞ = 26.9 kcal/mol).

We have also calculated the proton affinity at each Ge center to understand the reactivity of the lone pairs. The proton affinities of Ge1 (267.6 kcal/mol), Ge2 (264.8 kcal/mol) and Ge3 (266.2 kcal/mol) are similar, which are quite higher than that of the calculated value for five-membered N-heterocyclic germylene (196.0 kcal/mol) at the same level of theory and close to that of germylene (266.1 kcal/mol).6c Even though the lone pair on Ge1 is utilized for the π-bond formation with carbene carbon atom, it is as equally available as those on Ge2 and Ge3 towards protonation. This can be attributed to the compensative π back donation of N1→C1 within the cAAC ring and the more accessible p-type lone pair on Ge1 as compared to the less accessible sp-hybrid type lone pair (approximately 80% s-character) on Ge2 and Ge3 atoms. Thus, as per the structure and bonding analysis, Ge1 can be considered as tetravalent Ge(II) while the high proton affinity indicates its divalent Ge(0) nature. Hence, Ge1 shows a dual character for the electronic state and the lone pair on Ge1 can be considered as hidden-type. We have recently reported a similar type of bonding and reactivity pattern for tri-coordinate beryllium complexes.19 A similar type of bonding and reactivity pattern of divalent carbon(0) is also reported by Frenking and co-workers.20

Scheme 3. Proposed mechanism for the formation of 1 and 2.

Proposed reaction mechanism for the formation of 1-2 (Scheme 3) was theoretically studied (See ESI† for detailed discussion).

In summary we have synthesized and characterized a novel Ge3-com pound (cAAC)GeGeL2 (1) in singlet state. Compound 1 can slowly convert to germylene 2 in solution. Notably, cAAC exclusively favors the generation of 1, while NHC does not. Compound 1 is the first example of direct formation of a Ge=C bond by using a stable cAAC. Quantum mechanical studies show a reciprocal relationship between the bonding and reactivity of the central Ge atom. The bonding pattern of the central Ge atom is substantially different as compared to the terminal Ge atoms. However, the reactivity of all Ge atoms towards protonation is similar. The EDA-NOCV analysis suggests two bent bonds for Ge1-CAAC bond.

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Notes and references

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† Electronic Supplementary Information (ESI) available: Experimental procedures, X-ray crystallographic, and computational information of 1, CCDC 967854 (1). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c000000x/


12 The source of H2O in (cAAC)2O is possibly THF. For the synthesis and characterization of (cAAC)2O, see ref. 6(c).

13 Although no accurate value is reported, it must be smaller than 160 ppm. See reference: F. Meiners, W. Saak and M. Weidenbruch, Pure Appl. Chem., 2009, 81, 1099-1104.


18 (a) See ESI† for the details of computational methodology and its related citations; (b) See ESI† for the detailed bonding description by EDA-NOCV analysis.


A novel Ge$_2$-compound with dual character of the central Ge atom has been synthesized via direct formation of a Ge=C bond by using a stable cyclic alkyl(amo) carbene.