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Isolation of Elusive Phosphinidene-Chlorotetrylenes: The Heavier Cyanogen Chloride Analogues

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Abstract: The elusive phosphinidene-chlorotetrylenes, [PGeCI] and [PSiCI] have been stabilized by the hetero-bileptic cyclic alkyl(amino) carbene (cAAC), N-heterocyclic carbene (NHC) ligands, and isolated in the solid state at room temperature as the first neutral monomeric species of this class with the general formulae (L)P-ECI(L') (E = Ge, 3a-3c; E = Si, 6; L = cAAC; L' = NHC). Compounds 3a-3c have been synthesized by the reaction of cAAC-supported potassium phosphinidenides $[cAAC=PK(THF)_x]_n$ (1a-1c) with the adduct NHC:→GeCl₂ (2). Similarly, compound 6 has been synthesized via reaction of 1a with NHC:→SiCl₂ adduct (4). Compounds 3a-3c, and 6 have been structurally characterized by single-crystal X-ray diffraction, NMR spectroscopy and mass spectrometric analysis. DFT calculations revealed that the heteroatom P in 3 bears two lone pairs; the non-bonding pair with 67.8% of s- and 32% of p character, whereas, the other lone pair is involved in π backdonation to the C_{cAAC} -N π^* of cAAC. The Ge atom in 3 contains a lone pair with 80% of s character, and slightly involved in the π backdonation to C_{NHC}. EDA-NOCV analyses showed that two charged doublet fragments {(cAAC)(NHC)}⁺, and {PGeCl}⁻ prefer to form one covalent electron-sharing σ bond, one dative σ bond, one dative π bond, and a charge polarized weak π bond. The covalent electronsharing σ bond contributes to the major stabilization energy to the total orbital interaction energy of 3, enabling the first successful isolations of this class of compounds (3, 6) in the laboratory.

Introduction

Cyanogen chloride [N=C-CI] is a stable volatile compound, utilized in various organic synthesis.^[1] In contrast, chloroand chloro-phosphasilyne phosphagermyne [P≡Ge−Cl], [P=Si-Cl], the elusive heavier analogues of cyanogen chloride are expected to be highly unstable due to the presence of significantly weaker PEE triple bond (E = Ge, Si) because of the participation of significantly diffused 3s/3p and 4s/4p orbitals with internal nodes and Pauli repulsion energy, leading to the huge deviation from compact overlap of these orbitals of P and E atoms.^[2] The predicted lower thermodynamic stability of the heavier analogues of cyanogen halides can be attributed to the dramatically decreased ability of these heavier group 15/14 elements to form the multiple bonds,[3] which in turn potentially increases the contribution of the phosphinidene-tetrylene^[4] type isomer B to the ground state of the molecule (Figure 1). Such species has been theoretically predicted to be stabilized by using significantly bulky alkyl or aryl substituents, where the stability of the multiple bonded linear form of $[:P \equiv ER]$ (A, E = heavier group 14 elements, e.g., Si, Ge, etc.) is much higher compared to the corresponding bent structure **C** [RP=E:] with an electronegative R group (R = Cl, F) at the group 14 atom (E).^[5] However, none of the experimental attempts to synthesize these molecules have been successful so far, except the approach, where the sterically demanding amido ligands have been employed to stabilize the phosphinidene-tetrylene dimer **D**,^[6d] and the monomer **G**^[6f] (Figure 1).^[6]



Figure 1. Representative heavier analogues of cyanide (A-H) and carbenestabilized low coordinate germanium halide (I).

Although, the syntheses and reactivity of main group higher analogues of alkenes, alkynes and nitriles have attracted huge attention for past few decades,^[7] the stabilization of the higher analogues of cyanogen chloride as the neutral, monomeric, low coordinate phosphorus-(chloro)-germanium/silicon compounds has been rarely explored due to the high synthetic challenges caused by the lower thermodynamic stability and tendency to oligomerise. Although, a few groups have reported the isolation of the corresponding non-halogenated analogues in the dimeric forms, e.g., the phosphagermylyne dimers D [LGeP]₂ (Figure 1),^[8] synthesis of the donor base-stabilized corresponding chloro substituted species, phosphinidene-chlorotetrylenes [(L)P \equiv ECI(L'); E = Ge, Si; L = L' = donor base ligands] could not be achieved so far. In recent past, a phosphorus containing digermanium species^[2b] and a germylene-P=C=O^[2c] have been reported. A decade ago, Ge-P containing clusters have been isolated by reaction of BuLi with GeCl₂(dioxane).^[2d] The interconversion of donor base ligand free P=Si-Cl and Si=P-Cl species have been theoretically studied by Chu et al.[5a] The first phosphasilene R-P=Si(R₂) was characterized in 1984 by Bickelhaupt et al.[2e] and such class of compound was first structurally characterized by Niecke et al.^[2f] Since then, a large

number of phosphasilenes have been synthesized, isolated and characterized by different techniques. $^{[2g\text{-}h]}$

Since the discovery of N-heterocyclic carbenes (NHCs) in 1991^[9] by the group of Arduengo, and cyclic alkyl(amino) carbenes (cAACs) in 2005^[10] by the group of Bertrand, NHCs and cAACs have been hugely employed as Lewis base donor ligands to stabilize a variety of otherwise unstable exotic main group compounds with low coordinate main-group elements.[11] However, the chemistry of low coordinate exotic germaniumbased compounds are still scarce in the literature. Recently, the group of Roesky reported the cAAC-stabilized neutral germanium(I) radicals H (Figure 1).[12] The stabilization of heterodiatomic group 14/15 compounds with low coordinate group 14/15 elements stabilized by donor base ligands has been of significant interest during the past few decades.^[13] NHCs have been employed for the stabilization of low coordinate, cationic phosphorus-germanium compound **E**,^[14] and the chlorogermyliumylidene chloride I.^[15] Recently, phosphinidenes. the reactive group 15 analogues of carbenes with two lone pairs of electrons on phosphorus have gained considerable attention for stabilizing main group exotic species. One such example is the bis-NHC-phosphinidenide-stabilized tetrylenes F (Figure 1).^[16] Lappert et al. reported the possible stabilization of such species by utilizing a variety of anionic ligands, e.g., the bulky amido ligands (-NR₂) due to the effective p_{π} - p_{π} interaction between the lone pair of electrons at N atom and the vacant p orbital at the heavier group 14 elements.^[17] However, the synthetic viability of phosphinidene stabilized chlorotetrylenes, more precisely the phosphinidene-chlorogermylenes, and phosphinidene-chlorosilylenes are not explored till date. In this context, it is worth mentioning that recently, we have theoretically predicted that the laboratory isolation of the elusive homo- and hetero-bileptic chloro-phosphasilynes or the positional isomers, the phosphinidenecorresponding chlorosilylenes would be feasible by using homo- or heterobileptic donor base ligands, such as, carbenes (NHCs or cAACs).^[18] Inspired by these predictions, we attempted the syntheses of such species along with the corresponding germanium analogues, i.e., the elusive chlorophosphagermylynes or the phosphinidene-chlorogermylenes. Herein, we report on the successful syntheses of room temperature stable, neutral, monomeric, hetero-bileptic phosphinidene-chlorogermylenes 3a-3c with general formula (L)P-GeCI(L') (L' = NHC (1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene); Cy-cAAC (:C(N-2.6-1 = = ${}^{i}Pr_{2}C_{6}H_{3})(C_{6}H_{10})(CMe_{2})(CH_{2}))$ (3a), Et₂-cAAC = (:C(N-2,6- ${}^{i}Pr_{2}C_{6}H_{3})(CEt_{2})(CMe_{2})_{2}(CH_{2}))$ (3b), Me₂-cAAC = (:C(N-2,6-ⁱPr₂C₆H₃)(CMe₂)₂(CH₂)) (**3c**)), and phosphinidene-chlorosilylene 6 with general formula (L)P-SiCl(L') (L' = NHC; L = Me₂-cAAC) using cAACs and NHC as stabilizing ligands. The compounds 3a-3c, and 6 have been further structurally characterized in the solid state by single-crystal X-ray diffraction, NMR spectroscopy and mass spectrometric analysis.

Results and Discussion

The reaction of 1:1 molar ratio of cAAC-stabilized potassium phosphinedenides $[cAAC=PK(THF)_x]_n^{[19]}$ (1a-1c) ((Cv $cAAC=PK(THF)_x)_n$, **1a**; $(Et_2-cAAC=PK(THF)_x)_n$, **1b**; (Me₂cAAC=PK(THF)_x)_n, 1c), and NHC-stabilized dichlorogermanium NHC:→GeCl₂ (2) (NHC = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene) in anhydrous THF at -78 °C resulted in a dark brownish-red solution. The temperature of the reaction mixture was then slowly raised to room temperature over 1 h and further kept at that temperature with constant stirring for 2 h until the color of the reaction mixture changes to orange-red. THF was then completely removed under reduced pressure and the orange-red crystalline solid was extracted in anhydrous hexane. The hexane solution was then filtered to remove the salt (KCI), and the dark red filtrate was concentrated under reduced pressure to 1-2 mL, and kept for crsytallization. The orange-red block or rod shaped crystals of compounds 3a-3c $[cAAC: \Rightarrow: P-Ge(CI): \leftarrow: NHC]$ (cAAC = Cy-cAAC (3a), Et₂-cAAC (3b), and Me₂-cAAC (3c)) were obtained in 40-48% vields from the concentrated hexane solutions kept at -40 °C (Scheme 1).



Scheme 1. Syntheses of cAAC and NHC-stabilized elusive phosphinidenechlorogermylenes (cAAC)P-GeCl(NHC) (3a-3c).

A longer reaction time led to the dissociation of the free NHC along with the formation of the NHC salt (confirmed by ¹H, and ¹³C NMR spectra). To study the reaction progress with time, we have recorded the ³¹P NMR spectra of the reaction mixture at room temperature in a regular time interval. The ³¹P spectrum recorded after 5 min of the addition of NHC:->GeCl₂ (2) to a precooled THF-D₈ solution of (Me₂-cAAC=PK(THF)_x)_n (1c) at -110 ^oC exhibited a broad singlet at 207.5 ppm corresponding to the potassium phosphinidenide 1c.^[19] The ³¹P spectrum of the same reaction mixture recorded after 30 min showed the appearance of a sharp singlet at the upfield region of 90.5 ppm with the disappearance of the resonance, characteristic of 1c. No further changes were observed when the ³¹P spectrum was recorded after 1 h.

The controle of the reaction temperature was found to be critical for the increased yields of the products **3a-3c**. When the reaction was carried out at room temperature or there was a sudden increase in the reaction temperature, the formation of a tetraphosphinidenide-anchored Ge(IV) compound (cAACP)₄Ge (4) was isolated as a light yellow crystalline by-product, which was characterized by single-crystal X-ray diffraction, and NMR spectroscopic analysis (see SI). The formation of compound 4 ((Me₂-cAACP)₄Ge) was found to be more significant in case of the less bulkier Me₂-cAAC ligand. A concentrated hexane solution kept at -40 °C afforded the light yellow block shaped crystals of **4** after 2-3 days. Several attempts to grow single crystals of **4**, suitable for X-ray diffraction always produced very weakly diffracting crystals for which the data quality was poor

(see SI for the molecular structure of **4**, given only for the atom connectivity and not for comparing the experimental bond parameters). The ³¹P NMR spectrum of **4** in C_6D_6 exhibited a singlet at 37.7 ppm.

A similar synthetic approach to isolate the corresponding phosphinidene-chlorosilylene 6 became successful when the cAAC-stabilized potassium phosphinedenide ((Me2 $cAAC=PK(THF)_x)_n$ 1c and NHC-stabilized dichlorosilylene NHC: \rightarrow SiCl₂ (5) reacted in 1:1 molar ratio in THF at -110 °C. The temperature of the reaction mixture was slowly raised to 0 °C over 30 min, and further stirred at that temperature for additional 30 min to obtain a dark brownish-red solution. The dark-red crystalline solid obtained upon evaporation of the THF was extracted in hexane and filtered. The dark red filtrate was concentrated under reduced pressure to 1-2 mL, and kept for crsytallization. The bright red plates of compound 6 [Me2cAAC: =: P-: Si(CI) ←: NHC] was obtained in 23% yield from the concentrated hexane solution, kept at -40 °C after three weeks (Scheme 2). Recently, the bonding and stability of the donor base stabilized (L)PSi-Cl(L') (6) have been theoretically predicted by the groups of Roy and Mondal.^[18]



Scheme 2. Synthesis of cAAC and NHC-stabilized elusive phosphinidenechlorosilylene (cAAC)P-SiCI(NHC) (6).

Compounds 3a-3c, 6 were found to be extremely air and mosture sensitive. Exposure of the orange-red crystals of 3a-3c under air for 5 min led to the decomposition of the compounds to the colorless solid of the carbene salts, while the dark red crystals of 6 decomposed immediately after exposure to air. The solutions of 3a-3c, 6 in hexane were stable at -40 to 0 °C for several months (3) under an inert atmosphere. However, an exposure to air, turned the solution to light yellow to colorless within 1-2 min. The crystals of 3a-3c are stable at room temperature for more than 6 months and those of 6 for several days under an inert atmosphere. 3a-3c melt above 200 (3a), 182-184 (3b), 163-165 °C (3c). The ³¹P NMR spectra of the respective C₆D₆ solutions of 3a-3c, and THF-D₈ solution of 6 showed sharp singlets at 87.8, 79.8, 86.7, and 75.7 ppm, respectively, which are much upfield shifted when compared with those of the corresponding cAAC=PK (1a-1c) (200-210 ppm, broad singlets).^[19] The ²⁹Si NMR spectrum of 6 showed a doublet (due to coupling with ³¹P nucleus with $I = \frac{1}{2}$) at -20.7 ppm, which is slightly upfield shifted when compared to that of the compound NHC \rightarrow SiCl₂ \rightarrow P-Tip (-19.12 ppm).^[20] The J_{Si-P} for compound 6 was found to be 219.2 Hz, which is higher than that of NHC \rightarrow SiCl₂ \rightarrow P-Tip (J_{Si-P} = 197.7 Hz).^[20] This observation is in accordance with the previous prediction of the slight π electron delocalization between the silicon and phosphorus atoms in 6, resulting in a stronger coupling between ²⁹Si and ³¹P nuclei.^[18] The ¹³C resonances of **3a-3c** for C_{cAAC} were observed as doublets in the range of 213.8-212.5 ppm with

 J_{C-P} in the range of 83.0-86.0 Hz, which are slightly downfield shifted when compared with those of the corresponding [cAAC=PK(THF)_x]_n (**1a-1c**) (202-208 ppm),^[19] but upfield shifted when compared with those of the corresponding free carbenes (309.4 and 304.2 ppm for Cy-cAAC and Me₂-cAAC, respectively).^[21] The ¹³C NMR spectrum of **6** showed a comparatively downfield shifted doublet at 221.9 ppm ($J_{C-P} = 112.1$ Hz). The UV-vis spectra of **3a-3c** were recorded in hexane under an inert atmosphere, which showed broad (300-500 nm) with an absorption maxima (λ_{max}) at 428 nm for **3a**, 369 nm for **3b**, and 360 nm for **3c** (see SI for details).

Single crystal X-ray structure determinations of compounds **3a**-**3c**, and **6** revealed that all of the compounds have a similar molecular structure (Figures 2-3), and therefore, the structural aspects of only compound **3a** are described herein (for structural details of **3b-3c**, and **6** see SI).

Compound **3a** crystallizes in the triclinic space group *P*-1 (Figure 2). The molecular structure of (Cy-cAAC)P-GeCl(NHC) (**3a**) exhibits bent geometry at phosphorus atom which is bound to one carbene (Cy-cAAC) and one chlorogermanium (GeCl) unit, which is further bonded to the NHC.



Figure 2. Molecular structure of compound (Cy-cAAC)P-GeCl(NHC) (3a). H atoms are omitted for clarity. Selected bond lengths [Å] and bond angles (°): N3-C1 [1.371(2)], C1-P1 [1.7240(17)], P1-Ge1 [2.3872(5), Ge1-Cl1 [2.3222(5), Ge1-C2 [2.0857(16)], C2-N1 [1.348(2)], C2-N2 [1.359(2)], N3-C1-P1 (119.95(13)), C1-P1-Ge1 (105.479(6)), P1-Ge1-Cl1 (93.886(18)), C1-Ge1-C2 (99.002(4)), Ge1-C2-N1 (137.438(9)), Ge1-C2-N2 (118.132(9)).

The germanium atom adopts a trigonal pyramidal geometry (sum of the bond angles at Ge1 is 279.67°), indicating the presence of a lone pair of electrons. Ge/Si atom is 1.11 Å above the plane containing PCIC atoms. The Cy-cAAC and the NHC units of **3a** are oriented in a *trans*-position (*E*) (C1–P1–Ge1–C2 torsion angle 151.52°) with respect to the central P–GeCI moiety. The phosphorus atom in **3a** is bound to one germanium atom with the Ge1–P1 bond length of 2.3872(5) Å, which is comparable with the the Ge–P electron sharing single bond (2.3823(12) Å) in the diphosphidogermylene [(Ar₂P)₂Ge:],^[22] but much shorter than the P–Ge donor-acceptor type interaction (2.525(3), 2.612(3) Å) in the dimeric phosphinoamido-chlorogermylene [Ph₂PN(Ar)GeCI]₂.^[23] The Ge1-C2 bond length in **3a** is 2.0857(16) Å, which is shorter than that in NHC:–GeCl₂

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(2.112(2) Å),^[24] but comparable with those (Ge1–C1: 2.058(3) Å, and Ge1–C2: 2.057(3) Å) of the Ge(II) salt in [bis(NHC)–GeCI]*CI⁻, referring to a stroger chelating interaction of the C_{NHC} to the Ge atom.^[15] The Ge1–Cl1 bond distance in **3a** is 2.3222(5) Å, which is much longer than that in the adduct NHC:->GeCl₂ (**2**) (average 2.277 Å),^[24] and in the related fourcoordinate Ge(II) species 1,2,3,4-tetrakis(di-*tert*-butylmethylsilyI)-3,4-dichlorotetragermetene (2.248(1) Å),^[25] but comparable (2.310(1) Å) with the Ge(II) salt in [bis(NHC)–GeCI]*CI⁻.^[15] The P1–C1 bond length in **3a** is 1.7240(17) Å, which is longer than that in cAAC=PK (**1a**) (1.702(3) Å)^[19] and in between those in Cy-cAAC₂P₂ (1.719(7) Å)^[26] and NHC₂P₂ (1.750(2) Å),^[27] indicating a donor-acceptor type bond and the phosphinidene nature [cAAC:=:::P:].

Several attempts to reduce compound **3** in the presence of various reducing agents, e.g., $K/Na/Na-C_{10}H_8$ could not be successful, instead the corresponding cAAC-stabilized bisphosphinidnene cAAC₂P₂^[28] was isolated each time. However, the *in situ* generation of the corresponding radical (Et₂-cAAC)P–Ge(NHC) (**3b-CI**) was confirmed by the EPR spectroscopic analysis of the reaction mixture containing (Et₂-cAAC)P–GeCl(NHC) (**3b**) and finely cut pieces of potassium metal in 1:1 molar ratio in THF at 293 K to *in situ* generate **3b-CI** radical. The X-band EPR spectrum exhibited five hyperfine lines due to the coupling of the unpaired electron with two neighbouring ¹⁴N nuclei (*I* = 1), which have been further replicated two times due to the coupling of the unpaired electron with neighbouring ³¹P nucleus (*I* = 1/2) (see SI).

The molecular structure of compound **6** is shown in Figure 3 (see SI for details), the structural features are very much similar to those of **3**, and hence not described here.



Figure 3. Molecular structure of compound (Me₂-cAAC)P–SiCl(NHC) (6). H atoms are omitted for clarity. Selected experimental bond lengths [Å] and bond angles [^{*}]: N3-C3 1.379(6), C3-P1 1.713(5), P1-Si1 2.2951(18), Si1-Cl1 2.1857(17), C1-Si1 1.968(4), C1-N1 1.372(6), C1-N2 1.349(6), N3-C3-P1 122.0(3), C3-P1-Si1 107.32(16), P1-Si1-C1 87.02(14), P1-Si1-Cl1 98.84(7).

We have carried out density functional theory (DFT) calculations on a model compound **3a'**, where the Dip (2,6-diisopropylphenyl) substituents at the cAAC and NHC donor moieties are replaced by the methyl groups (to save the computational time) in order to

elucidate the electron densities distribution and bonding situation in compound 3a. We have optimized the geometry of 3a' at the BP86/def2-TZVPP level of theory and then performed the natural bond orbital (NBO) analysis (see SI). The optimized bond lengths and bond angles of 3a' are in good agreement with the experimental values obtained for 3a (see SI). The optimized geometry of the model compound 3a' was found to be comparable to that of 3a (see SI). The natural bond orbital (NBO) analysis of 3a' supports the bonding picture of the molecule in terms of the donor-acceptor interactions, depicted as cAAC: ⇒: P-Ge(CI): ⇒: NHC. Figure 4 shows the relevant NBOs of 3a' participating in bonding. The HOMO and HOMO-2 are the σ lone pair orbitals at P and Ge centres, respectively; whereas, the HOMO-3 is solely the σ lone pair orbital at P centre. HOMO-1 is the π bonding orbital of the C_{cAAC}=P bond, which is majorly polarised towards the P centre (58.9%). The lone pair on P has 67.8% of s character, and 32.0% of p character. Another lone pair on P is involved in π backdonation to the vacant orbital of cAAC (LUMO), which can be very well visualised in HOMO-1.



Figure 4. Representative NBOs of model compound 3' at BP86/def2-TZVPP level of theory. Hydrogen atoms are omitted for clarity. Energy values are in eV.

The Wiberg bond index (WBI = 1.51) and two bonding occupancies have been computed for the C_{cAAC} -P bond in **3a'** imply the presence of a partial double bond between C_{cAAC} and P atoms of **3a'**. The NBO analyses reveal the presence of a lone pair of electrons on Ge, which is 80.8 and 23.8% of s p characters, respectively. HOMO of **3a'** suggests that there is a secondary bonding interaction between P-Ge single bond with $C_{cAAC/NHC}$. HOMO-2 shows the diffusion of electron densities on Ge towards P and C-P bond (Figure 4).

To study the topological features of the model compound **3a'**, we have performed Bader's quantum theory of atoms in molecules (QTAIM).^[29] The strength of a bond is echoed in the parameter, called the electron density at (3,-1) the bond critical point (BCP). The electron density, $\mathbf{p}(\mathbf{r})$ of C_{cAAC}-P bond (0.16 a.u.) suggests a closed shell interaction, and more polarised towards the P centre (Figure 5; see SI). $\mathbf{p}(\mathbf{r})$ for **3a'** was found to be comparatively lower for the bonds P-Ge and Ge-C_{NHC}, indicating a weaker closed-shell interaction. The positive values of $\nabla^2 \mathbf{p}(\mathbf{r})^{[29d]}$ for the bonds C_{cAAC}-P (0.16) and Ge-C_{NHC} (0.11) reveal the closed-shell interactions between them. A small negative value (-0.07) of $\nabla^2 \mathbf{p}(\mathbf{r})$ for the P-Ge bond implies that it may be a covalent electron sharing bond. The ellipticity (ϵ)

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parameter of C_{cAAC} -P bond is 0.05 suggesting single or multiple bonding interactions between them. However, QTAIM cannot unambiguously predict the nature of chemical bond especially when heavier elements are involved.



Figure 5. Contour plot of Laplacian distribution $[\nabla^2 \mathbf{p}(r)]$ in the P3–Ge1–C12 plane of **3a'**. Solid blue lines indicate the areas of charge concentration $(\nabla^2 \mathbf{p}(r) < 0)$, while dotted purple lines denote charge depletion $(\nabla^2 \mathbf{p}(r) > 0)$. Solid lines connecting atomic nuclei (black) are the bond paths.

To get a deeper insight into the bonding scenario between the ligand fragments (cAAC NHC) and PGeCI moiety, and the nature of the P-GeCl bond, we performed the energy decomposition analysis coupled with natural orbital for chemical valence (EDA-NOCV).^[30] For the study of the bonding nature of (cAAC, NHC) and PGeCI fragments, we considered four different bonding possibilities by changing the charge and the multiplicity of the interacting fragments: (a) neutral (cAAC NHC) and PGeCI fragments in singlet state, interacting to form the dative bonds; (b) neutral (cAAC NHC) and PGeCI fragments in quintet state, implying the interaction to form electron sharing covalent bond; (c) doubly charged fragments of [(cAAC NHC)]²⁺ and [PGeCI]²⁻ in triplet state, interacting to form σ electron sharing and π dative bonds; and finally (d) singly charged fragments of [(cAAC NHC)]⁺ and [PGeCI]⁻ in doublet state, which would interact to form both electron sharing and dative bonds (Scheme S5).

Table 1. The EDA-NOCV results at the BP86-D3(BJ)/TZ2P level of theory forcAAC-PGe(CI)-NHCbondsofcAAC-P-Ge(CI)-NHC(3a') using $[(cAAC)(NHC)]^+$ and $[P-Ge(CI)]^-$ in the electronic doublet (D) states asinteracting fragments. Energies are in kcal/mol.

Energy (kcal/mol)	Type of bonding interaction in 3'	[(cAAC NHC)] ⁺ (D) +
		[P-Ge(Cl)] ⁻ (D)
ΔE_{int}		-239.5
ΔE_{Pauli}		488.1
$\Delta E_{disp}^{[a]}$		-23.4 (3.2%)
$\Delta E_{\text{elstat}}^{[a]}$		-374.9 (51.5%)
$\Delta E_{\rm orb}^{[a]}$		-329.4 (45.2%)
$\Delta E_{\text{orb(1)}}^{[b]}$	cAAC-PGe(CI)-NHC	-161.7 (49.1%)
	σ electron sharing	
$\Delta E_{orb(2)}^{[b]}$	cAAC→PGe(Cl)←NHC σ dative	-60.9 (18.5%)
$\Delta E_{\text{orb(3)}}^{[b]}$	cAAC←PGe(Cl)	-65.8 (20.0%)
	π dative backdonation	
$\Delta E_{\text{orb}(4)}^{[b]}$	cAAC←PGe(Cl)→NHC	-11.7 (3.5%)

	charge polarized	
	π backdonation	
$\Delta E_{\text{orb(rest)}}$		-29.3 (8.9%)

Table 1 summarizes the EDA-NOCV results from which it is evident that the central PGeCI unit in **3a'** prefers to be anionic (PGeCI⁻) and the ligand-pair [(cAAC^{Me})(NHC^{Me})] prefers to be cationic [(cAAC^{Me})(NHC^{Me})]⁺ in their doublet states due to the electron deficient nature of the central PGeCI unit and the electron rich nature of the ligand pair (the best bonding scenario; Scheme 3). The contribution due to the electrostatic interaction energy between (cAAC NHC)⁺ and PGeCI fragments is signifiantly higher (55.5%) that of orbital interaction energy (($\Delta E_{orb} = 45.2\%$). The stabilization due to the favorable dispersion energy (3.2%) is small. The total orbital interaction energy (ΔE_{orb}) can be further divided into four pairwise contributions ($\Delta E_{orb(1)} - \Delta E_{orb(4)}$) (Figure 6), where four different types of bonding interactions have been observed.



Figure 6. The shape of the deformation densities $\Delta \rho_{(1)-(4)}$ that correspond to $\Delta E_{orb(1)-(4)}$, and the associated MOs of cAAC-PGe(CI)-cAAC (**3**') and the fragments orbitals of [(cAAC) (NHC)] and [PGeCI] in the doublet state (D) at the BP86-D3(BJ)/TZ2P level. Isosurface values are 0.002 au for $\Delta \rho_{(1-3)}$ and 0.0005 au for $\Delta \rho_{(4)}$. The eigenvalues $|v_n|$ give the size of the charge migration in e. The direction of the charge flow of the deformation densities is red-blue.

The $\Delta E_{orb(1)}$ represents an electron-sharing C_{cAAC/NHC}-PGe σ bond via the interaction of SUMO of (cAAC NHC)⁺ having σ symmetry with $\pi\text{-type}$ SUMO of PGeCl- in doublet states. It contributes nearly 50% of the total orbital interaction energy (ΔE_{orb}) . The $\Delta E_{orb(2)}$ shows the formation of one σ -dative $C_{\text{NHC/cAAC}} \rightarrow PGe$ bond due to the overlap of HOMO-1 of (cAAC NHC)⁺ with π^* -orbital of P=Ge bond contributing only 18.5% to $\Delta \textit{E}_{\text{orb}}.$ The formation of a $C_{\text{cAAC}}\text{=P}$ $\pi\text{-bond}$ has been in the deformation densities of $\Delta E_{orb(3)}$ contributing 20% to ΔE_{orb} . The $\Delta E_{orb(3)}$ is slightly higher than $\Delta E_{orb(2)}$. $\Delta E_{orb(3)}$ involves the overlap of lone pair of electrons on P-atom of PGeCl with $\pi^*_{\text{C-N}}$ orbital of cAAC. The $\Delta E_{\text{orb}(1,3)}$ show the loss of double bond character of P-GeCl bond in their deformation densities (charge flow red→blue). $\Delta E_{orb(4)}$ represents the weak π -back donations from P and Ge atoms of PGeCl fragment to the π^* -orbitals on cAAC/NHC. $\Delta E_{orb(4)}$ is six times smaller in magnitude than $\Delta E_{orb(2)}/\Delta E_{orb(3)}$ (Table 1). The magnitude of C_{cAAC/NHC}-PGe bond is significantly higher than other interactions due to the covalent electron-sharing nature of the bond (see SI for detailed EDA-NOCV analyses). EDA-NOCV analyses of the P-Ge bond reveals that it is an electron sharing σ -bond. It contributes 81.8% of the ΔE_{orb} . The stabilization due to the favourable dispersion energy is found to be 3.2%. The best bonding scenario of compound 3' has been depicted in Scheme 3.



Scheme 3. Best bonding scenario in compound 3'. The cAAC \leftarrow P π -bond [$\Delta E_{orb(3)}$] is six time stronger than that (Ge \rightarrow NHC)) Ge-C_{NHC} bond [$\Delta E_{orb(4)}$, Figure 4].

Conclusion

In conclusion, we have demonstrated the first efficient synthetic route for the elusive, neutral, monomeric hetero-bileptic ligandstabilized singlet phosphinidiene-chlorogermylenes (3a-3c), and phosphinidiene-chlorosilylene (6) under the elimination of potassium chloride salt. 3, 6 are new class of main group compounds containing low coordinate germanium/silicon and phosphorus atoms in their lower oxidation states. 3a-3c, and 6 have been isolated in moderate yields and characterized by Xray single-crystal diffraction, NMR spectroscopy amd mass spectrometry. The stability, electron densities distributions and the chemical bonding of the model compound 3' has been studied by quantum chemical calculations, such as, DFT, NBO, QTAIM and EDA-NOCV analyses. The HOMO, HOMO-1, and HOMO-2 of 3' are lone pair of electrons on Ge/P-atoms, C_{cAAC}=P π bond, P=Ge π -bonds, respectively. EDA-NOCV calculations of 3' showed that the central unit PGeCI prefers to be anionic (PGeCl⁻) and the ligand-pair [(cAAC^{Me})(NHC^{Me})] prefers to be

cationic [(cAAC^{Me})(NHC^{Me})]+ in their doublet states to give rise to the best bonding scenario due to the electron deficient nature of the central PGeCI unit and the electron rich nature of the ligand pair. There are electron-sharing $C_{CAAC/NHC}$ -PGe σ bond, contributing –161.7 kcal/mol, one coordinate $C_{\text{NHC/cAAC}} \rightarrow PGe \sigma$ bond contributing –60.9 kcal/mol, and dative $C_{cAAC} \leftarrow P \pi$ bond contributing -65.8 kcal/mol between these two charged fragmments. The fourth minor bonding interaction is due to charge flow from lone pair of electrons on Ge and P-Ge σ-bond to the cAAC ligand contributing -11.7 kcal/mol as weak π stronger than the $C_{\text{NHC/cAAC}}{\rightarrow}\text{PGe}\ \sigma$ donation. The magnitude of C_{cAAC/NHC}-PGe bond is significantly higher than other interactions due to the covalent electron-sharing nature of the bond. This is a new addition to the low coordinate Ge-P chemistry utilizing the bonding diversities of two dffierent carbene ligands.

Experimental Section

Synthesis of compound 3a: A THF solution of NHC:→GeCl₂ (2, 269 mg, 0.5 mmol, 1 equiv) cooled to -78 °C (acetone/liquid nitrogen bath) was slowly added through a cannula to the Schlenk flask containing a THF solution of Cy-cAAC=PK (1a, 200 mg, 0.5 mmol, 1 equiv) at -78 °C with constant stirring and maintaining the same reaction temperature for 15 min. Afterwards, the temperature of the reaction mixture was slowly raised to room temperature and stirred for 2 h, resulting in a color change from brownish-red to orange-red. The reaction mixture was then filtered, and the filtrate was concentrated under vacuum up to complete dryness. resulting in a reddish-brown crystalline solid, which was further extracted with anhydrous hexane (15-20 mL), filtered and the filtrate was concentrated under vacuum to 1-2 mL and kept at -40 °C to obtain the orange-red rod-shaped single crystals (suitable for X-ray single-crystal diffraction) of 3a (170 mg, 40% yield) after 3-4 days. ¹H NMR (400 MHz, C₆D₆, 298 K) δ: 7.31 (d, J = 7.4 Hz, 3H, -ArH), 7.18 (d, J = 7.8 Hz, 6H, -ArH), 6.50 (s, 2H, -CH), 2.96-2.84 (m, J = 6.8 Hz, 6H, -CH(CH₃)₂), 1.60 (s, 3H, -CH₃), 1.55 (d, J = 6.7 Hz, 2H, -CH₂), 1.52 (d, J = 6.8 Hz, 16H, -CH(CH₃)₂), 1.45 (d, J = 6.8 Hz, 2H, -(cyclohexyl)), 1.40 (d, J = 6.7 Hz, 3H, -(cyclohexyl)), 1.33-1.23 (m, 4H, -(cyclohexyl)), 1.12 (dd, J = 15.7, 7.7 Hz, 24H, -CH(CH₃)₂) ppm; ¹³C NMR (101 MHz, C₆D₆, 298 K) δ: 212.9 (d, J_{C-P} = 84.8 Hz) (Ccarbene), 175.0, 146.1, 145.8, 137.9, 134.7, 131.2, 130.4, 129.3, 125.7, 124.5, 124.3, 124.0, 67.3, 51.0, 29.3, 29.1, 29.0, 25.9, 25.8, 25.7, 23.3, 23.1, 21.4 ppm; ³¹P NMR (162 MHz, C₆D₆, 298 K) δ: 87.8 ppm; HRMS (ESI): m/z calculated for C₅₀H₇₁ClGeN₃P as [M+NH₄]⁺ 871.4624; found: 871.4490 [M+NH4]+. MP: 198-200 °C.

https://www.ccdc.cam.ac.uk/services/structures?id=doi:10.1002/chem.20 2201242 Deposition Numbers 2167245 (for **3a**), 2167246 (for **3b**), 2167248 (for **3c**), and 2180182 (for **6**) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service https://www.ccdc.cam.ac.uk/structures/?.

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Elusive hetero-bileptic carbenes-stabilized singlet phosphinidiene-chlorosilylene, and chlorogermylenes with majorly contributing covalent electron-sharing σ bonds are isolated and structurally characterized in the solid state.