The Varied Frustrated Lewis Pair Reactivity of the Germylene Phosphaketene (CH{(CMe)(2,6-Pr2C6H3N)}2)GePCO

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Abstract: The germylene species (CH{(CMe)(2,6-Pr2C6H3N)}2)GePCO 1 is shown to react with the Lewis acids (E(C6F5)3 E = B, Al). Nonetheless, 1 participates in FLP chemistry with electron deficient alkynes or olefins, acting as an intramolecular FLP. In contrast, in the presence of B(C6D6)3 and an electron rich alkyne, 1 behaves as Ge-based carbene donors have been employed. On the other hand, the range of Lewis acids has mostly been limited to electron-deficient boranes or alanes, although carbon-based cationic Lewis acids have received lesser attention. Among these latter cases, trityl[2], benzhydrylium (I),[3] N-methylacridinium (II),[3–4] trioxatriangulenium (III)[5] and a Ru-η6-arene salt (IV)[6] have been served as the Lewis acid in FLP reactivity. Meanwhile, Bazan et al.[7] and Alcarazo[8] have demonstrated that fullerene acts as a neutral carbon-based Lewis acid while Alcarazo et al.[9] have also exploited electron-deficient allenes (V) as carbon-based Lewis acid in FLP chemistry (Figure 1).

The ambiphilic nature of carbenes has also been exploited for FLP chemistry. The report of the activation of H2 by cyclo-aaminoalkyl carbenes (CAACs) by Bertrand[10] can be viewed as FLP chemistry at a single site as the carbene carbon is both an electron donor and an acceptor. Among the heavier analogs, Müller and coworkers[11] have shown that the combination of silylenes (VI) (Figure 1) with either Lewis acids or bases results in intermolecular FLPs capable of activating H2, further emphasizing the ambiphilic character. On the other hand, while the use of Ge(II), Sn(II) and Pb(II) centers have been exploited as

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Example of group-14-element-based Lewis acids for FLP chemistry.

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Lewis acids in intramolecular FLP systems with P, N donors (VII) (Figure 1) to activate alkenes and alkenes 12,13 the use of such heavier cognates as Lewis bases has drawn lesser attention.14

In the last decade, Grützmacher and Goicoechea15 have pioneered the chemistry of 2-phosphaethynolate anion (PCO−), the phosphorus analogue of the cyanate anion. This anion has proved to be very useful building block for phosphorus-containing heterocycles.16 In addition, Grützmacher, Li and co-workers demonstrated the electrophilic character of the carbon center in (R,Ni)P−PCO and P3Si−PCO, as these species react with Lewis basic N-heterocyclic carbenes affording carbene–phosphinidene adducts17 and carbene–phosphakene adducts.18 Moreover, phosphaketene phosphine (R,Ni)P−PCO and phosphakete gemynylenes Mg−PCO have been shown to form phosphanyl supported phosphinidene19 and 1,3-digerma-2,4-diphosphacyclobutadiene20 via the elimination of CO under UV photolysis, respectively. Grützmacher et al. also reported the construction of optoelectronic material precursors 1,3,4-azadiphospholides from NaPCO, in which the nitrogen-containing substrate acting as a nucleophile to attack the electrophilic carbon center of PCO.21 Recently, we have exploited P3SiGePCO as a versatile precursor for the facile synthesis of the phosphorus analogue of DMF, Me2PC(O)H,22 diphospha-ureas,23 phosphakelenes,24 and acylophosphines.25

Herein, we explore the chemistry of the related germylene species L3−GePCO (1, L3= CH[(CMe)-(2,6-Pr2C6H4)N]). We demonstrate that this species exhibits varying FLP reactivity in combinations with Lewis acids and/or alkyne and olefin substrates. These findings are illuminated via state-of-the-art DFT calculations.

The phosphaketene substituted gemylene 1 was readily prepared from the corresponding [L3−GeCl] compound and NaPCO.20,26 This neutral derivative of the Ge(II) cation [Ge(Dipp,nacnac)]+ reported by Power and co-workers in 200127 incorporates nucleophilic P and O centers, an electrophilic C center and an ambiphilic Ge. Thus, a variety of chemo- and regioselectivity is anticipated in the reactivity of 1. DFT calculations at the TPSS-D3/def2-TZVP + COSMO(toluene) level showed that the highest occupied (HOMO) and the lowest unoccupied (LUMO) molecular orbitals of 1 mainly consist of the Ge and P electron lone pairs and the delocalized π* antibond of the Nacnac ligand backbone ring, respectively, with the HOMO-LUMO gap being 2.3 eV (see Supporting Information). Since the HOMO is 0.55 eV above other occupied molecular orbitals, the Ge and P centers are expected to dominate the nucleophilic reactivity of 1.

To probe this basicity of 1, we began examining reactions with Lewis acids. The reaction of 1 with B(C6F5)3 was performed in toluene at −30°C (Scheme 1). Upon mixing, the solution became violet and afforded violet crystals of 2 (Scheme 2). While the cations are planar L3−Ge(II) units, the anion is derived from two PCOB fragments forming a four-membered C3P2 ring, with two exocyclic OB and one exocyclic BP linkages. The observed C−P (1.8013(18), 1.7923(18), 1.8400(18), and 1.8315(18) Å), P−B (2.078(2) Å) and B−O (1.541(2) and 1.555(2) Å) bond lengths are similar to those previously seen in the anion derived from the reaction of NaPCO and B(C6F5)3.20

To probe the mechanism of these reactions, accurate dispersion-corrected DFT calculations at the PW6B95-D3/def2-QZVP + COSMO/R5/TPSS-D3/def2-TZVP + COSMO level of theory in toluene solution were performed (see Supporting Information).28 DFT calculations considering the interactions of 1 with B(C6F5)3 show that the binding free energies of B(C6F5)3 to the Ge, P, γ-C and O sites of 1 are 5.2, −0.2, 19.5 and −1.7 kcal/mol, respectively. These data are consistent with reversible B(C6F5)3 binding to the phosphaketene oxygen as interactions with Ge or P centers are somewhat sterically hindered. B(C6F5)3 binding is expected to enhance the electrophilicity of the phosphaketene carbon and decrease the nucleophilicity of the Ge center thus weakening the Ge–P bond. Indeed, the dissociation of the PCOB adduct is only 7.4 kcal/mol endergonic generating the electrophilic [L3−Ge]1+ and nucleophilic [PCOB(C6F5)3]+ ion in solution. Further B(C6F5)3 binding to the P-site of [PCOB(C6F5)3]+ is 3.5 kcal/mol endergonic, followed by nucleophilic addition of another.
[POB(C₆F₅)₃]⁻ that is 21.6 kcal/mol more exergonic to afford the experimentally observed salt 2.

In an analogous fashion, reaction of 1, with 1 equivalent of Al(C₆F₅)₃ in toluene at room temperature yielded colorless crystals of 3 after storage at −30°C overnight. The molecular structure of 3 reveals that the γ-carbon of the ligand backbone added to the electrophilic C of phosphaketene moiety directly upon coordination of the Al to oxygen atom (Scheme 1, Figure 3). The resulting Ge–P distance is 2.418(1) Å while the C–P distance is 1.707(6) Å suggestive of a C–P double bond. DFT calculations also show the transfer of Al(C₆F₅)₃ to the phosphaketene oxygen site of 1 from Al(C₆F₅)₃·(toluene) is −12.5 kcal/mol exergonic affording the adduct 3a (see Supporting Information). Further ring-closing to give a C–C bond to the γ-carbon is 5.3 kcal/mol endergonic thus not favored in solution but likely favored in the solid state. The greater Lewis acidity of Al(C₆F₅)₃ enhances the Lewis acidity of the CO fragment, prompting reaction with the basic γ-carbon of the Nacnac ligand yielding the formation of 3. In this case, the reaction can be viewed as a C/Al FLP addition to the CO fragment in 1. Related 1,4-additions involving both the γ-carbon and the metal center has been seen in other systems containing magnesium, aluminium, germanium, and platinum.

Compound 1 was also shown to react with the electron-deficient alkyne diisopropyl but-2-yne diolate in toluene (Scheme 2), as slow addition led to an immediate color change from light yellow to dark purple. Workup and recrystallization from a hexane/toluene afforded purple crystals of 4. The ³¹P NMR spectrum of 4 showed a high-field singlet at −119.8 ppm (DFT ³¹P: −107.6 ppm). The absorption bands at 1519 cm⁻¹ in the IR spectrum can be assigned to the PCO moiety (DFT 1550 cm⁻¹). Single crystal XRD analysis revealed 4 (Figure 4a) to be a planar five-membered C₃PGe heterocycle in which the alkyne has added to the Ge and the carbonyl carbon of the PCO fragment. The newly formed Ge–C and C–C bond lengths are 1.946(5) and 1.343(6) Å, respectively. The formation of 4 amounts to the addition of the basic Ge and Lewis acidic carbonyl carbon to the alkyne, suggesting that 1 behaves as an intramolecular FLP in this case.

Subsequent treatment of 4 with 1 equivalent of B(C₆F₅)₃ in C₆D₆ at room temperature led to an immediate color change of

Figure 3. POV-ray depiction of the molecular structure of 3. Hydrogen atoms are omitted for clarity. C: black, F: hot pink, P: orange, N: blue, O: red, Ge: pale pink, Al: plum.

Scheme 2. Reactions of 1, diisopropyl but-2-yne diolate with/without B(C₆F₅)₃.

Figure 4. POV-ray depiction of the molecular structure of (a) 4 (b) 5 and (c) 7. Hydrogen atoms are omitted for clarity. C: black, P: orange, N: blue, O: red, Ge: pale pink.
the resulting solution from dark violet to bright red. With the addition of n-hexane, red crystals of 5 were obtained from the solution in 71% yield. In an analogous fashion, the adduct of 4 with Al(CF$_3$)$_3$ was also prepared, isolated and spectroscopically characterized (Scheme 2). Both products 5 (Figure 4b) and 6 (See Supporting Information) were also crystallographically characterized, revealing simple coordination of the carbonyl oxygen to B and Al, yielding the B–O and Al–O bond lengths of 1.5157(16) and 1.790(3) Å, respectively.

Interestingly, simultaneously mixing of 1, B(C$_6$F$_5$)$_3$ and disisopropyl-but-2-yneidoate (1:1:1) in toluene proceeded at ambient temperature affording colorless crystals of 7 as the main product in 71% yield (Scheme 2). The molecular structure of 7 was established by NMR analysis and a single-crystal X-ray diffraction study. The $^{31}$P and $^{11}$B NMR spectra of 7 displayed singlets at 33.4 and $–$1.7 ppm, respectively, (DFT $^{31}$P: 51.8 $^{11}$B: $–$1.8 ppm) indicating the presence of a tetracoordinated boron anionic center. In the molecular structure of 7 (Figure 4c), the γ-carbon of the Nacnac ligand backbone is linked to the alkyne carbon of the dialkyl but-2-yneidoate. The Ge atom is coordinated by two imine-nitrogen atoms and one phosphorus atom and adopts a trigonal pyramidal geometry. The planar four-membered C$_3$P heterocycle of 7 features a C–C double bond (1.370(3) Å) and a C–C single bond (1.513(3) Å), respectively. The two C–P bond lengths of 1.847(2) Å and 1.907(2) Å in 7 are closed to that of primary acylphosphine Ph$_2$P(CHC(O)PH$_2$ (1.858(3) Å).$^{22}$ In addition to 7, crystals of 5 were also isolated from this reaction albeit in 2% yield.

The formation of 5 and 7 demonstrates the impact of order of reagent addition and suggests that the reaction pathway is altered by the interaction of alkyne with B(C$_6$F$_5$)$_3$. Probing these reactions computationally revealed that in the absence of borane, nucleophilic attack of the electron-poor alkyne iPrO$_2$CC=CCO$_2$Pr by the Ge center of 1 is 8.3 kcal/mol endergonic over a low barrier of 11.1 kcal/mol (via TS1) generating the zwitterion A (Scheme 3). This is followed by a barrierless ring-closing (via TS2) between the anionic alkyne carbon and the electrophilic phosphaketene carbon affording five-membered-ring product 4. The subsequent addition of B(C$_6$F$_5$)$_3$ to the phosphaketene oxygen is barrierless and $–$19.4 kcal/mol exergonic to form the product 5. In contrast, for the simultaneous mixture of 1, B(C$_6$F$_5$)$_3$ and the alkyne iPrO$_2$CC=CCO$_2$Pr, the anion [PCOB(C$_6$F$_5$)$_3$]$^–$ may act as a nucleophile to attack the alkyne yielding a PC$_3$-ring anion which adds further via the C–P bond to the electrophilic Ge$^+$ and the nucleophilic γ-carbon centers of the FLP-like [L$^3$Ge]$^+$ affording 7.

The corresponding reaction of 1, B(C$_6$F$_5$)$_3$, and phenyl-acetylene (ratio 1:1:1) in toluene, gave rise to new singlets at $–$304.2 ppm and $–$16.4 ppm in the $^{31}$P and $^{11}$B NMR spectra, respectively (DFT $^{31}$P: $–$326.7 $^{11}$B: $–$16.0 ppm). Crystallization from toluene solution afforded colorless blocks of 8 in 51% yield. The structure of 8 was confirmed by single crystal X-ray diffraction (Scheme 4, Figure 5a), revealing the regiospecific intermolecular trans-FLP addition of germylene and B(C$_6$F$_5$)$_3$ to the alkyne with the borane adding to the less hindered carbon. The new Ge–C and B–C bond lengths are 1.918(5) and 1.645(7) Å, respectively, yielding an olefinic linkage. At the same time, the PCO group is unchanged.

The simultaneous mixture of 1 with B(C$_6$F$_5$)$_3$ and dimethyl maleate (ratio 1:1:1) in toluene also gave two new species as evidenced by $^{31}$P resonances at $–$9.5 and $–$31.4 ppm and the broad $^{11}$B NMR signals at $–$1.7 ppm. While these two species proved challenging to separate, crystals of 9 that gives rise to the former $^{31}$P signal were isolated and an X-ray crystallographic study (Figure 5b) affirmed the FLP addition of the nucleophilic Ge and the electrophilic C to the olefinic double bond. The diastereomeric, five-membered germaphosphacycles L$^3$GePC(O)(B(C$_6$F$_5$)$_3$)(Me$_2$C=CHCHC=O)Me exhibits a Ge–P double bond of 2.2518(9) Å, similar to that in 4 and to those previously reported.$^{14}$ The C–C bond within the heterocycle is a typical single bond of 1.556(6) Å. Interestingly, efforts to prepare the analogous compound from more stable trans-dimethyl fumarate led to no reaction even after heating at 100°C for 24 h.

These latter two reactions were also considered computationally. For the electron-rich alkyne PhCCH, computations revealed a transient adduct B with Lewis-acidic B(C$_6$F$_5$)$_3$ that is 8.8 kcal/mol endergonic (Scheme 5a). Rapid nucleophilic attack at the Ge center of 1 yields 8 in an overall reaction that is $–$14.0 kcal/mol exergonic over a low free energy barrier of only 12.0 kcal/mol (via TS3). An alternative reaction pathway afford-
ing a 5-membered ring analogous to 5 derived from direct intramolecular FLP addition of PhCCH to 1 was found to be 20.6 kcal/mol exergonic but is prevented by a sizable barrier of 27.6 kcal/mol.

Finally, binding of B(C₆F₅)₃ to one ester group of the electron-deficient olefin MeO₂CC=CCO₂Me to give C is 1.9 kcal/mol endergonic over a low barrier of 12.9 kcal/mol (via TS₄) generating the zwitterion D. Subsequent rapid ring-closing at the electrophilic phosphaketene carbon (via TS₅) affords E and a subsequent shift of the B(C₆F₅)₃ to the phosphaketene oxygen, affords the product 9. For 9, the respective DFT-computed ³¹P and ¹¹B signals at −14.9 ppm and −0.3 ppm agree well with experiment. The additional product observed experimentally is very likely the species E, as supported by the computed ³¹P and ¹¹B signals at −27.6 ppm and −1.8 ppm, respectively. Consistent with experiment, the direct reaction of 1 and MeO₂CC=CCO₂Me alone encounters a sizable barrier of 23.8 kcal/mol, thus is slow even under moderate heating conditions. These results indicated that even in a molecule with multiple reactive sites, computation of reaction profiles provides insights into the subtleties of Lewis acidic/basic chemoselectivity.

In conclusion, despite the observation of direct reactivity of 1 with the Lewis acids (E(C₆F₅)₃ E = B, Al), 1 participates in FLP chemistry with alkylne and olefin substrates. Specifically, 1 is shown to react with electron deficient alkynes or olefins, acting as an intramolecular FLP. In contrast, in the presence of B(C₆F₅)₃ and an electron rich alkyne, 1 behaves as Ge-based nucleophile to effect intermolecular FLP addition reactions. This range of reactivity demonstrates that the precise reaction pathway is controlled by the precise nature of the electrophile and nucleophile generated in solution. We are continuing to examine related systems containing Ge-based nucleophiles and to explore the utility and generality of FLPs in inorganic and organic chemistry.

Crystallographic details
Deposition Number(s) 2095456, 2125894, 21004686, 2095279, 2095451, 2125902, 2095286 and 2095291 contain(s) 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.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.

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