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The non-bonding carbone lone pair in geometrically-constrained antimony and bismuth carbodiphosphorane complexes readily complexed AuCl to afford rare examples of geminal bimetallic carbone coordination featuring a main-group metal.

Since the introduction of hexaphenylcarbodiphosphorane (C(PPh₃)₂ or CDP) by Ramirez *et al.* in 1961, the bonding situation of the divalent carbon center has been the subject of much debate.¹ However, the turn of the century witnessed remarkable advances in the experimental and theoretical elucidation of carbones (CL₂), which are now widely recognized as double Lewis bases comprised of two orthogonal (σ - and π -symmetric) lone pairs within a formally C(0) atom.² This facilitated their increased adoption as electronically-flexible ligands, which can serve as two- or four-electron donors depending on their coordination environment.³ While the stabilization of main-group Lewis acids through only one carbone lone pair (type I) is now ubiquitous in literature, complexes involving the contribution of both lone pairs (types II and III) are sparse (Fig. 1).



Fig. 1 Examples of carbone interactions with main-group elements (NHC = N-heterocyclic carbone).

Geminal bimetallic coordination of a carbone to main-group and transition metals[†]

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Coordination type II to double Lewis acids (mononuclear centers possessing acceptor σ and π orbitals) are of fundamental importance as they represent a non-reductive approach to carbon-element multiple bonding through the formation of novel double dative $C \rightrightarrows E$ bonding.⁴ Achieving this bonding situation depends on both metal Lewis acidity and orbital geometry. Carbone bonding to the hardest double Lewis acid Be²⁺ forms predominantly $C \rightarrow Be \sigma$ -donation (type I) in monodentate (carbone)BeCl₂ complexes,⁵ where further interactions (types II or III) are inhibited by unfavorable stereoelectronics. Examples of $C \rightrightarrows E$ multiple bonding in comparative *p*-block element complexes (E = B, Ge, Sb, Bi) were typically promoted by cationization to enhance the electrophilicity of the Lewis acid.⁶ The elusive $C \rightrightarrows$ Be bond was recently achieved within an *ortho*beryllated carbodiphosphorane (Be(CDP)), wherein the overlap of beryllium and carbone frontier orbitals were enhanced by encapsulating the metal within the tridentate ligand.⁷

Heteronuclear coordination (type III) establishes the intrinsic nature of carbones as containing stereochemically active lone pairs,⁸ but remains underdeveloped and extremely rare among main-group elements (MGEs).9 Indeed, this contrasts the affinity of main-group Lewis acids to form bimetallic complexes with double Lewis bases such as chalcogenides via non-covalent interactions.¹⁰ Petz, Neumüller, Frenking, and coworkers reported a gem-diborylated carbone complex in 2009 in their reaction of CDP and BH₃, which afforded [(CDP)(B₂H₅)][BH₄], presumably via (CDP)(BH₃)₂.¹¹ A recently isolated Li₂(CDP) etherate features a bimetallic MGE carbone complex stabilized by a kinetic chelating effect,12 although quantum chemical studies were not performed for comparisons with (CDP)(BH₃)₂. As these instances feature small 2nd row MGEs (Li, B), we sought to investigate this bonding scenario for larger main-group Lewis acids and its implications for their reactivity profiles.

We recently discovered that *ortho*-bismuthination of CDP yields predominantly $C \rightarrow Bi \sigma$ -donor interactions,¹³ which contrasts $C \rightrightarrows Bi$ interactions in the simple coordination adducts.^{6c,d,14} The constrained carbone interaction in the former is robust and yielded air-stable complexes with enhanced activity for Bi(n)/Bi(m)

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redox catalysis.^{13,15} However, the see-saw geometry of the fourcoordinate Bi(m) center is not symmetry correct for proper overlap with the carbone π lone pair. Encouraged by the robust nature of this framework, we anticipated that the non-bonding lone pair may readily complex metallic electrophiles to achieve novel bimetallic carbone coordination to heavy MGEs.¹¹ We now report the isolation and bonding analyses of carbodiphosphorane antimony and bismuth complexes involving bis(metalated) carbone centers, obtained by reactions of geometrically constrained carbonepnictogen complexes and AuCl.

For comparisons with their bismuth analogues, the antimony complexes 1-3 were prepared in likewise manner,¹³ and isolated as analytically pure white solids in 83 – 90% yields (Scheme 1). Characterization by ³¹P{¹H} and ¹³C{¹H} NMR spectroscopy reveal upfield resonances for the phosphorous and ylidic carbon atoms, which may be diagnostic of metal Lewis acidity.¹⁶ However, 3⁺, like the bismuth analogue, is a weak Lewis acid and does not complex triethylphosphine oxide for quantitative analysis *via* the Gutmann-Beckett method. The fluoride ion affinity (FIA) values in CH₂Cl₂ (calculated at PW6B95-(D3BJ)/def2-TZVPP level of theory) are comparable for 3⁺ (156 kJ mol⁻¹) and the Bi analogue (146 kJ mol⁻¹), and contrasts the superacidic nature of the monodentate carbodicarbene adducts.^{6c}

In their solid-state structures (Fig. 2), **1** and **2** feature the anticipated *trans* carbone-metal-halide ligation observed at bismuth, resulting in highly elongated Sb–Cl (2.8382(10) Å vs 2.39 for $\sum R_{cov}$) and Sb–Br (3.1063(10) Å vs 2.54 for $\sum R_{cov}$) bonds, which are comparable to the bismuth analogues.¹³ Solvent interactions through halide hydrogen bonding also contribute to the elongated metal-halide bonds. Charge separation in **3**[**SbF**₆] resulted in a downfield ³¹P NMR shift and shortened ^{carbone}C–Sb bond length compared to **1** and **2** (Fig. 2 caption) due to enhanced metal Lewis acidity.

Compound **3**[**SbF**₆] is further stabilized by weak cation-anion contacts. Attempts to isolate a fully charge separated complex without cation-anion contacts by the reaction of **2** and Na[BAr^F₄] (Ar^F = 3,5-(CF₃)₂C₆H₃) led to the protonated adduct [H(CDP)SbBr][BAr^F₄] (**4**[**BAr**^F₄]) due to hydrolysis by adventitious moisture. Because the anticipated [(CDP)M][BAr^F₄] complex was successfully isolated for bismuth under the same reaction and solvent conditions,¹³ it is suspected that the cationic unit in this framework is more reactive for antimony than bismuth, which is consistent with reports by Gabbai,¹⁷ Venugopal,¹⁸ and our laboratory,^{6c} who observed higher Lewis acidity for Sb *versus* Bi cations with the same ligand environments. The phosphorus resonance for **4**[**BAr**^F₄] ($\delta_P = 28.0$ ppm) is slightly upfield from **2**

 $sbx_{3} \xrightarrow{\text{Li}_{2}(\text{CDP})} \xrightarrow{Ph} \xrightarrow{P} \stackrel{Ph}{\longrightarrow} \stackrel{Ph}{\rightarrow} \stackrel{Ph}{\longrightarrow} \stackrel{Ph}{\rightarrow} \stackrel{Ph}{\longrightarrow} \stackrel{Ph}{\longrightarrow} \stackrel{Ph}{\longrightarrow} \stackrel{Ph}{\rightarrow} \stackrel{Ph}{\rightarrow} \stackrel{Ph}{\rightarrow} \stackrel{Ph}{$

Scheme 1 Synthesis of carbodiphosphoranyl antimony complexes 1-3.



Fig. 2 Molecular structures of **1–4**. Thermal ellipsoids set at 50% probability. Aromatic hydrogens, co-crystallized solvent molecules, and a non-coordinating BAr_4^F anion in **4[BAr_4]** are omitted for clarity. Selected bond distances (Å) and angles (°): **1** [and **2**]: Sb1–C1, 2.247(3) [2.228(4)]; Sb1–Cl1, 2.8382(10) [Sb1–Br1, 3.1063(10) Å]; C1–P1, 1.682(4) [1.692(5)]; C1–P2, 1.682(4) [1.678(5)]; C2–Sb1–C20, 102.40(13) [104.41(17)]; P1–C1–P2, 135.1(2) [134.2(3)]. **3**: Sb1–C1, 2.158(7); C1–P1, 1.708(7); C1–P2, 1.697(7); Sb1…F1, 3.061(9); C2–Sb1–C20, 99.7(2); P1–C1–P2, 130.6 (4).**4**: Sb1–C1, 2.426(3); Sb1–Br1, 2.8045(4); C1–P1, 1.761(3); C1–P2, 1.773(3); C2–Sb1–C20, 97.01(10); P1–C1–P2, 120.67(15).

 $(\delta_{\rm P} = 31.7 \text{ ppm})$, but the ¹³C{¹H} NMR spectrum reveals more dramatic differences whereby the ylidic carbon resonance for **4[BAr^F₄]** ($\delta_{\rm C} = 16.3 \text{ ppm}$, ¹*J*_{CP} = 62.5 Hz) is similarly upfield from 2 ($\delta_{\rm C} = 6.4 \text{ ppm}$, ¹*J*_{CP} = 99.1 Hz) but extends a much smaller ¹*J*(¹³C-³¹P) coupling constant.

Having established structural and spectroscopic similarities between these antimony complexes and their bismuth analogues, we probed the potential for geminal bimetallic coordination at the carbone center. The reaction of **1** or (CDP)BiCl with (Me₂S)AuCl readily affords the anticipated coordination adducts **5** and **6** as colorless solids (Scheme 2). Upon isolation, **5** and **6** are stable as solids for several weeks under ambient conditions, but rapidly decompose in solution (CH₂Cl₂) within 30–45 minutes by gradually depositing a gold film on the walls of the reaction flask. Despite their extreme sensitivity, compounds **5** and **6** were isolated in 69–73% yields as single crystals, which were grown by combining concentrated, cold DCM solutions of their respective reagents and immediately storing at -37 °C in the dark for two (**6**) or four (**5**) hours.

The phosphorus resonances of 5 ($\delta_P = 30.1$ ppm) and 6 ($\delta_P = 32.8$ ppm) suggest electronic similarities between these compounds in solution. Notably, the phosphorus resonance of 5 is comparable to that of 1 ($\delta_P = 29.5$ ppm), but the formation



of a new product was confirmed by unique ¹H and ¹³C{¹H} NMR resonances, combustion microanalysis, and X-ray crystallography (Fig. 3). The ¹*J*(¹³C-³¹P) coupling constant for the ylidic carbon in **6** (50.9 Hz) is smaller than those of (CDP)BiCl (98 Hz) and [H(CDP)BiCl][BPh₄] (70.7 Hz), which further highlights the diverse nature of carbone coordination in these bismuth complexes. Similar comparisons were not possible for **5**, as the ylidic carbon was not observed in the ¹³C{¹H} NMR spectrum due to its poor solubility and propensity to rapidly decompose in solution.

The carbone-pnictogen bonds of compounds 5 (2.453(6) Å) and 6 (2.574(4) Å) are elongated from their starting materials, and expectedly shorter for 5 compared to 6. Indeed, they are comparable to the carbone-pnictogen bonds of the protonated adducts (*e.g.*, 4^+ and [H(CDP)BiCl]⁺), suggesting similarities in the coordination mode of the carbone. The geometry about the carbone is a distorted tetrahedron, and the P1–C1–P2 bond angles (Fig. 3 caption) are comparable to 4^+ , as well as the *gem*-diborylated¹¹ and -diaurated^{*8b*} complexes. The ^{carbone}C1–Au and Au–Cl bond distances are each identical in 5 and 6, and their Au–pnictogen interactions are comparable and within the anticipated van der Waals radii for Sb–Au ($r_W = 3.72$ Å) and Bi–Au ($r_W = 3.73$ Å) aurophilic interactions.

To gain more insight into the bonding nature of the bimetallic complexes 5 and 6, DFT calculations at the BP86-D3(BJ)/def2-TZVP level of theory were performed. When the implicit solvation model (PCM, DCM) was included, the optimized geometrical parameters



Fig. 3 Molecular structures of **5** and **6**. Thermal ellipsoids set at 50% probability. Co-crystallized dichloromethane molecules and all H atoms are omitted for clarity. Selected bond distances (Å) and angles (°): **5**: Sb1–C1, 2.453(6); Sb1–C1, 2.6686(16); Sb1–Au1: 3.4133(5); Au1–C1, 2.038(7); Au1–C12, 2.2916(17); C1–P1, 1.748(7); C1–P2, 1.765(6); C3–Sb1–C21, 102.8(2); P1–C1–P2, 119.2(4). **6**: Bi1–C1, 2.574(4); Bi1–C11, 2.7311(11); Bi1–Au1, 3.4094(3); Au1–C1, 2.035(4); Au1–C12, 2.3068(10); C1–P1, 1.751(5); C1–P2, 1.743(4); C3–Bi1–C21, 100.97(16); P1–C1–P2, 119.8(3).



Fig. 4 (a) Frontier molecular orbitals (FMOs) and (b) Mayer bond indices for **5** (left) and **6** (right), calculated at the BP86-D3(BJ)/def2-TZVPP level of theory. (c) Selected Intrinsic Bond Orbitals (IBOs) of **5** enclosing 80% of the density of the orbital electron.

agree with those observed for the crystal structures, as discussed in our previous report.¹³ The HOMOs of **5** and **6** are located on the *gem*-dimetalated structural units with contributions from the carbone π electron and Au *d* orbitals (Fig. 4a). The lone pair from the Sb(m) atom in **5** features the largest HOMO orbital coefficient (28.3%), while the lone pair of the Bi(m) atom in **6** only has a small contribution (8.6%) to its HOMO. The two-armed PPh₃ in the CDP unit contribute predominantly to the energetically close-lying LUMO and LUMO+1 of **5** and **6** (Fig. S21 and S22, ESI[†]).

The Mayer bond indices are comparable for the C1-Au bonds in 5 (0.62) and 6 (0.64), but significantly higher for

Table 1 Calculated data for the C1-Au interactions in 5 and 6

| | 5 | 6 |
|--|-------|-------|
| Bond length $(C1-Au)^a$ | 2.05 | 2.05 |
| Bond order (C1–Au) | 0.62 | 0.64 |
| BDE in CH_2Cl_2 (C1–Au) ^b | 55.6 | 57.2 |
| NPA $(Au)^c$ | +1.24 | +1.19 |
| NPA (C1) | -1.02 | -1.00 |

 a Bond length (in Å). b bond dissociation energy (BDE, in kcal mol $^{-1}$). c Natural population analysis (NPA).

C1–Sb (0.54) than C1–Bi (0.46). These values are indicative of single dative bond characters (Fig. 4b). Intrinsic bond orbital (IBO) analyses for 5 and 6 reveal that there are one polarized 2c–2e σ -bonding feature for C1–Au, and three σ -bonds between the CDP moiety and the Sb or Bi atom (Fig. 4c and Fig. S20, ESI†). Specifically, the carbone π and σ electrons interact with the σ^* orbitals of Au–Cl and Sb–Cl or Bi–Cl, respectively, which are consistent with the frontier orbital analyses (Fig. 4a). Quantum theory of atoms-in-molecules (QTAIM) analyses corroborate these observations (Tables S2 and S3, ESI†). Thus, 5 and 6 are the first examples of geminal bimetallic CDP coordination to both main-group and transition metals.

Table 1 highlights some electronic features of the carbone-gold coordination. The calculated C1–Au bond dissociation energies (in CH_2Cl_2) in 5 (55.6 kcal mol⁻¹) and 6 (57.2 kcal mol⁻¹) suggest that AuCl coordination is strongly favored and deauration is not facile. Therefore, the rapid decomposition of 5 and 6 in CH_2Cl_2 likely does not result from initial deauration, but may proceed through alternative pathways. The stronger C1–Au interaction in 6 is likely due to a weaker Bi–C1 interaction compared to Sb–C1.

In conclusion, we have isolated and characterized rare examples of *gem*-dimetalated main-group carbone complexes. This coordination mode is a fundamental feature of double Lewis bases in bonding, widely observed in chalcogenide maingroup metal complexes, but hitherto unobserved in likewise manner for carbones. Therefore, these complexes further highlight the electronic flexibility of carbones at main-group metal centers, and may have significant implications for carbonestabilized heterobimetallic reactivities.

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Conflicts of interest

There are no conflicts to declare.

Notes and references

 (a) F. Ramirez, N. B. Desai, B. Hansen and N. McKelvie, J. Am. Chem. Soc., 1961, 83, 3539–3540; (b) N. D. Jones and R. G. Cavell, J. Organomet. Chem., 2005, 690, 5485–5496; (c) H. Schmidbaur, Angew. Chem., Int. Ed., 2007, **46**, 2984–2985; (d) G. Frenking, B. Neumüller, W. Petz, R. Tonner and F. Öxler, *Angew. Chem., Int. Ed.*, 2007, **46**, 2986–2987.

- 2 (a) R. Tonner, F. Öxler, B. Neumüller, W. Petz and G. Frenking, Angew. Chem., Int. Ed., 2006, 45, 8038–8042; (b) R. Tonner and G. Frenking, Chem. – Eur. J., 2008, 14, 3260–3272; (c) V. H. Gessner, Modern ylide chemistry: applications in ligand design, organic and catalytic transformations, Springer, 2018; (d) A. L. Liberman-Martin, Cell Rep. Phys. Sci., 2023, 4, 101519; (e) R. W. A. Havenith, A. V. Cunha, J. E. M. N. Klein, F. Perolari and X. Feng, Phys. Chem. Chem. Phys., 2021, 23, 3327–3334.
- 3 (a) S.-k Liu, W.-C. Chen, G. P. A. Yap and T.-G. Ong, Organometallics, 2020, 39, 4395–4401; (b) B. S. Aweke, C.-H. Yu, M. Zhi, W.-C. Chen, G. P. A. Yap, L. Zhao and T.-G. Ong, Angew. Chem., Int. Ed., 2022, 61, e202201884; (c) L. Zhao, C. Chai, W. Petz and G. Frenking, Molecules, 2020, 25, 4943–4990; (d) W.-C. Chen, J.-S. Shen, T. Jurca, C.-J. Peng, Y.-H. Lin, Y.-P. Wang, W.-C. Shih, G. P. A. Yap and T.-G. Ong, Angew. Chem., Int. Ed., 2015, 54, 15207–15212.
- 4 (a) M. Hermann and G. Frenking, *Chem. Eur. J.*, 2017, 23, 3347–3356;
 (b) M. A. L. Johansen and A. Ghosh, *Nat. Chem.*, 2023, 15, 1042.
- J. E. Walley, G. Breiner, G. Wang, D. A. Dickie, A. Molino, J. L. Dutton, D. J. D. Wilson and J. R. J. Gilliard, *Chem. Commun.*, 2019, 55, 1967–1970; (b) W. Petz, K. Dehnicke, N. Holzmann, G. Frenking and B. Neumüller, *Z. Anorg. Allg. Chem.*, 2011, 637, 1702–1710.
- 6 (a) S. Khan, G. Gopakumar, W. Thiel and M. Alcarazo, Angew. Chem., Int. Ed., 2013, 52, 5644–5647; (b) B. Inés, M. Patil, J. Carreras, R. Goddard, W. Thiel and M. Alcarazo, Angew. Chem., Int. Ed., 2011, 50, 8400–8403; (c) L. S. Warring, J. E. Walley, D. A. Dickie, W. Tiznado, S. Pan and R. J. Gilliard Jr, Inorg. Chem., 2022, 61, 18640–18652; (d) J. Walley, L. Warring, G. Wang, D. A. Dickie, S. Pan, G. Frenking and R. J. Gilliard, Angew. Chem., Int. Ed., 2021, 60, 6682–6690; (e) K. K. Hollister, A. Molino, G. Breiner, J. E. Walley, K. E. Wentz, A. M. Conley, D. A. Dickie, D. J. D. Wilson and R. J. Gilliard, J. Am. Chem. Soc., 2022, 144, 590–598; (f) C.-L. Deng, A. D. Obi, B. Y. E. Tra, S. K. Sarkar, D. A. Dickie and R. J. Gilliard, Nat. Chem., 2023, DOI: 10.1038/s41557-023-01381-0.
- 7 M. R. Buchner, S. Pan, C. Poggel, N. Spang, M. Müller, G. Frenking and J. Sundermeyer, *Organometallics*, 2020, **39**, 3224–3231.
- 8 (a) C. Esterhuysen and G. Frenking, *Chem. Eur. J.*, 2011, 17, 9944–9956; (b) J. Vicente, A. R. Singhal and P. G. Jones, *Organometallics*, 2002, 21, 5887–5900; (c) M. Alcarazo, C. W. Lehmann, A. Anoop, W. Thiel and A. Fürstner, *Nat. Chem.*, 2009, 1, 295–301.
- 9 M. Fustier-Boutignon, N. Nebra and N. Mézailles, *Chem. Rev.*, 2019, 119, 8555–8700.
- 10 (a) K. T. Mahmudov, A. V. Gurbanov, V. A. Aliyeva, M. F. C. Guedes da Silva, G. Resnati and A. J. L. Pombeiro, *Coord. Chem. Rev.*, 2022, 464, 214556; (b) F. A. Devillanova and W.-W. Du Mont, *Handbook of chalcogen chemistry: new perspectives in sulfur, selenium and tellurium*, Royal Society of Chemistry, 2013.
- 11 W. Petz, F. Öxler, B. Neumüller, R. Tonner and G. Frenking, *Eur. J. Inorg. Chem.*, 2009, 4507–4517.
- 12 S. C. Böttger, C. Poggel and J. Sundermeyer, *Organometallics*, 2020, **39**, 3789–3793.
- 13 A. D. Obi, D. A. Dickie, W. Tiznado, G. Frenking, S. Pan and R. J. Gilliard Jr, *Inorg. Chem.*, 2022, **61**, 19452–19462.
- 14 J. E. Münzer, N.-J. H. Kneusels, B. Weinert, B. Neumüller and I. Kuzu, *Dalton Trans.*, 2019, **48**, 11076–11085.
- 15 The isostructural phosphorus species was recently isolated, and enabled metallomimetic P(iii)/P(v) redox catalysis. See K. Chulsky, I. Malahov, D. Bawari and R. Dobrovetsky, Metallomimetic Chemistry of a Cationic, Geometrically Constrained Phosphine in the Catalytic Hydrodefluorination and Amination of Ar–F Bonds, *J. Am. Chem. Soc.*, 2023, **145**, 3786–3794.
- 16 (a) P. Erdmann and L. Greb, Angew. Chem., Int. Ed., 2022,
 61, e202114550; (b) Q. Teng and H. V. Huynh, Dalton Trans., 2017,
 46, 614-627.
- 17 G. Park, D. J. Brock, J.-P. Pellois and F. P. Gabbaï, *Chemistry*, 2019, 5, 2215–2227.
- 18 D. Sharma, S. Balasubramaniam, S. Kumar, E. D. Jemmis and A. Venugopal, *Chem. Commun.*, 2021, 57, 8889–8892.