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The unique β-diketiminate ligand in aluminum(I) and gallium(I) chemistry

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Over the past few decades, β-diketiminate ligands have been widely used in coordination chemistry and are capable of stabilizing various metal complexes in multiple oxidation states. Recently, the chemistry of aluminum and gallium in their +1 oxidation state has rapidly emerged. NacNacM(I) (M = Al, Ga; NacNac = β-diketiminate ligand) shows a two coordinate metal center comparable with singlet carbene-like species. The metal center also possesses a formally vacant p-orbital. In this article we present an overview of the last 10 years for aluminum(i) and gallium(i) stabilized by β-diketiminate ligands that have been widely explored in bond breaking and forming species.

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1. Introduction

Complexes containing a low-oxidation metal center is a key topic in modern organometallic chemistry as it leads to the development of new systems. A number of transition metal complexes were synthesized in a broad range of oxidation states to activate small molecules or used as catalysts in organic reactions.¹ It was not until 1991 that the first structurally characterized molecular aluminum(I) compound was reported. 2 Since then, the chemistry of Group 13 metals in the +1 oxidation state have played a great part in the development of p-block chemistry. The synthesis of compounds with aluminum and gallium in the +1 oxidation state is experimentally challenging and consistently hampered by their high reactivity and pronounced tendency towards disproportionation.³ The lone pair of electrons in MR (R = β-diketiminate) (M = Al, Ga) compounds can act as a basic site towards a range of Lewis acids, forming a σ donating bond.⁴ Donor-acceptor bonds have been observed in a range of main group elements, transition metals, and lanthanide and actinide metal complexes. The general M^IR unit (M = Al, Ga) can be considered isolobal with singlet carbenes, CO, and CNR.

β-Diketiminate ligands have found widespread application as supporting ligands in metal-mediated catalysis.⁵ The stoichiometric transformations of NacNacAl(I) and NacNacGa(I) have also been explored widely owing to the lone pair of electrons and a formally vacant p-orbital on aluminum affording high electrophilic and nucleophilic reactivity.⁶ A comparison between main group elements and transition metals was drawn when main group species were found to have reactivity

towards small molecules under ambient conditions. This was rationalized by main group species possessing donor/acceptor frontier orbitals which are separated by modest energy gaps, thus drawing comparisons with open-shell transition metal species.⁷ In 2000, Roesky *et al.* chose the β-diketiminate ligand to synthesize a more kinetically stable monomeric aluminum(1) compound Al ${H C [C (Me) N Dip]_2}$ (Dipp = 2,6-ⁱPr₂C₆H₃) (1).^{8a} This was the first stable dicoordinate aluminum (i) compound to be prepared and structurally characterized in the solidstate. Later, Cui et al. also reported a β-diketiminate ligand stabilized aluminum(1) compound $Al{HC} [C(^tBu)NDipp]_2$ ⁸ Computational studies of β-diketiminate stabilized heavier group metal complexes have shown that their metal lone pairs are associated with the HOMO−2.⁹ As a result, they are good σ donor ligands and poor π acceptors like N-heterocyclic carbenes (NHCs), and thus have the potential to display carbenelike chemistry. Inoue *et al.* reported the first neutral $Al(i)$ compound containing an Al=Al double bond, which was achieved through the reductive dimerization of the corresponding N-heterocyclic carbene (NHC)-stabilized silyl substituted aluminum(m) dihalide (I, Fig. 1), and its reactivity toward the fixation and selective reduction of $CO₂$, both of which can be accessed in a stoichiometric and catalytic fashion.¹⁰ In 2018, Aldridge and Goicoechea et al. reported the first isolation of a nucleophilic aluminyl anion [(NON)Al][−] by employing a chelating ligand (NON = 4,5-bis(2,6-diisopropylanilido)-2,7-di-tertbutyl-9,9-dimethylxanthene) (II, Fig. 1), which acts as an unprecedented aluminum (i) nucleophile $(e.g.,$ in reactions with ${}^{t}Bu_{3}PAuI$), and which effects the formal oxidative addition of the C–C bond in benzene. 11 After that, Coles and co-workers also synthesized a two-coordinate N-heterocyclic aluminyl anion K[Al(NON^{Dipp})] (NON^{Dipp} = $[O(SiMe₂NDipp)₂]^{2-}$, Dipp = 2.65 -ⁱPr₂C₆H₃, III, Fig. 1), which is able to undergo further reactions such as activation of elemental selenium to form an **PERSPECTIVE**
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Fig. 1 Di-tert-butyl(methyl)silyl-substituted dialumene (I), anionic group 13 analogues of N-heterocyclic carbenes (II–III), monomeric 1,2,4-tri-tert-butylcyclopentadienylaluminum(I) (IV), Lewis base stabilized aluminum(I) hydride (V) and an alkyl-substituted aluminum anion (VI).

aluminum complex containing an aluminum–selenium multiple bond and with 1,3,5,7-cyclooctatetraene (COT) to give the first aluminum complex containing a reduced COT-ligand with a strong aromatic character, respectively.¹² Both aluminyl anion complexes reacted with two abundant greenhouse gases $(CO₂$ and N₂O) *via* cycloaddition to generate a monoalumoxane anion. 13 The first isolable example of a room temperature stable monomeric cyclopentadienylaluminum(I) derivative was reported by Braunschweig and co-workers, which was supported by a bulky 1,2,4-tri-tert-butylcyclopentadienyl (Cp^{3t}) ligand (IV, Fig. 1).^{14a} The same group also reported the first example of a monomeric Lewis base stabilized $Al(i)$ hydride that can be isolated and handled under ambient conditions (V, Fig. 1).^{14b} Very recently, Yamashita's group reported an alkylsubstituted aluminum anion that exhibits very strong basicity and nucleophilicity (VI, Fig. 1).^{14c} These species have been observed to form both covalent and donor–acceptor bonds, revealing both the reducing and nucleophilic properties of these novel complexes.

 $Gallium(i)$ compounds are often driven by the thermodynamic preference for the metal center to exist in the +3 oxi-

dation state. Having said this, much of the reported chemistry of monomeric gallium (i) compounds is derived from the significant basicity of the metal through its lone pair of electrons. $Gallium(i)$ compounds are generally more stable towards disproportionation than the corresponding aluminum (i) compounds. So far, several gallium(1) N-heterocycles have been reported.^{15,16} Bi- and tridentate ligand systems have been used in the preparation of a variety of neutral and anionic gallium (i) heterocycles (e.g., five-membered anionic complexes VII-VIII, a guanidinate complex IX and a monomeric tris(pyrazolylborate) complex X, see Fig. 2).¹⁶ These compounds have been prepared either by salt-metathesis reactions between alkali metal salts of the ligands and "GaI" or by alkali metal reduction of Ga^H or Ga^{III} precursors. The monomeric example of Ga^I amide (XI, Fig. 2) can be considered to be having a quasi one-coordinate metal center, which also exhibits weak intramolecular arene interactions in the solid state.^{16m} Recently, a pincer-type gallylene ligand has been successfully synthesized utilizing bis (phosphino)-terpyridine as an efficient scaffold for the Ir–Ga¹ bond, which enabled various reactions at the Ir center by keeping the gallylene ligand intact. 17

The β-diketiminate ligands typically provide monoanionic, bidentate support for metal complexes and offer a much higher degree of steric control through the choice of N-substituents. By tuning the steric and electronic properties of the supporting β-diketiminate ligands, the reactivity of the compounds can be significantly improved.¹⁸ Thus, complexes with a low-valent metal could be stabilized by the employment of sterically encumbering β-diketiminate ligands. It is interesting to note that with a redox-inactive metal bound and appropriate substituents, β-diketiminate ligands become redox-active ligands.¹⁹ Herein we present an overview that is of relevance to the corresponding bond

Fig. 2 Reported examples of gallium(I) N-heterocycles.

activation by aluminum(I) and gallium(I) with β-diketiminate ligands.

2. Chemistry of ^{Dipp}NacNacAl(I) and DippNacNacGa(I)

The monomeric aluminum carbenoid $\text{DippNacNacAl}(I)$ (1) was prepared through reduction of the corresponding $Dipp$ NacNacAlI₂ with potassium (Scheme 1).^{8a} At the same time, Power and co-workers reported a β-diketiminate stabilized Ga(I) monomer $\frac{DippNacNacGa(I)}{2}$ (Scheme 1).²⁰ It was obtained by the reaction of $[Li\{HC(CMeNDipp)_2\}]$ with "GaI". The remarkable thermal stability of the compounds toward disproportionation reaction (decomp. >150 °C) can be attributed to the steric bulk of the β-diketiminate ligand, which provides kinetic protection to the metal center. X-ray crystal structure analysis showed that compound 2 is monomeric and isostructural with its aluminum counterpart. With a singlet lone pair and formally empty p-orbital on the metal, the neutral heterocycles Dipp NacNacAl(I) and Dipp NacNacGa(I) have the potential to exhibit both nucleophilic and electrophilic characteristics. **Column Tens articles**

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2.1. Small-molecule activation

CO and $CO₂$ activation by transition-metal complexes has been studied extensively for many years. However, the activation of CO and $CO₂$ with Group 13 metal elements and their compounds has been explored scarcely.²¹ In 2018, Crimmin et al. reported carbon chain growth from C1 to C3 and to C4 by sequential reactions of CO and $CO₂$ with a transition metal carbonyl complex in the presence of an aluminum(I) complex (Scheme 2).²² Warming a frozen suspension of $[W(CO)_6]$ with 2 equiv. of 1 under 1 atm of CO from -78 °C to r.t. in a benzene-d₆ solvent results in the formation of the C3 homologated product 3. Heating the isolated and purified sample of 3 under one atmosphere of CO leads to the formation of the chain growth product 4. Further chain growth of C3 to a C4 fragment could be achieved upon the reaction of 3 with one atmosphere of $CO₂$. Although the reaction of $CO₂$ with 3 at 25 °C initially produces 5, when the sample 3 is heated at 100 °C for 48 h, it completely converts to 6. No reaction occurs between 1 and CO in the absence of $[W(CO)_6]$. The gallium products of these reactions are not reported.

Scheme 1 Syntheses of ^{Dipp}NacNacAl(I) and ^{Dipp}NacNacGa(I)

Scheme 2 Carbon chain growth by sequential reactions of CO and $CO₂$ with [W(CO)₆] and 1.

Scheme 3 Syntheses of [2.2.1] metallobicycle 7 and reversible CO insertion.

The [2.2.1] metallobicyclic compound 7 was synthesized by the cycloaddition of complex (1) with low-valent aluminum and 1,3-cyclohexadiene. The exposure of a C_6D_6 solution of 7 to one atmosphere of CO generated the insertion product 8. As shown in Scheme 3 the reaction mixture is reversible, when 8 is heated for longer time. Compound 9 was also studied in the reaction with CO, and the insertion of CO into the Al–C bond was observed. However, compound 10 decomposes at 25 °C within 12 h into an intractable mixture of products. 23

Crimmin and co-workers documented the first reversible addition of ethylene to aluminum (i) 1. The monomeric molecular aluminum (i) complex reacted with a series of terminal and strained alkenes including norbornene, $24a$ ethylene, propylene, hex-1-ene, 3,3-dimethyl-1-butene, allylbenzene and 4-allylanisole. Remarkably all these reactions are reversible under mild conditions (Scheme 4). $24b$

2.2. Cleavage of the M–X single bond

Aluminum (i) 1 has been developed to act as a synthon for the preparation of aluminum–metal bonded compounds via oxidative insertion of the Al center into metal-halogen linkages. Jones et al. reported the first example of molecular complexes

Fig. 3 X-ray single crystal structure of compound 20.

containing an unsupported Be–Al bond. The Be–Al bonded complexes 17 and 18 were obtained as yellow crystalline solids from the reaction of $\text{DippNacNacAl}(I)$ (1) with $\text{[BeX}_2(\text{tmeda)}]$ $(X = Br \text{ or } I, \text{ tuneda} = tetramethylethylene diamine) in 1:1 stoi$ chiometry (Scheme 5). The Be–Al bond distances in 17 and 18 are 2.474(1) \AA and 2.432(6) \AA , respectively. They are significantly longer than the sum of single bond covalent radii of the elements (2.28 Å). DFT calculations reveal that the compounds with metal–metal bonds have a high s-character. This is consistent with similar Pauling electronegativities between Al and Be. The isostructural Mg–Al (19) and Zn–Al (20) analogues of these complexes have been isolated in the 1:1 reaction of $\text{DippNacNacAl}(I)$ (1) with $[\text{MgI}_2(\text{tmeda})]$ and $[\text{ZnBr}_2(\text{tmeda})]$, respectively (Scheme 5). 25 The composition of compound 20 was confirmed by means of single-crystal X-ray structural analysis (Fig. 3). Compound 20 has distorted tetrahedral Al and Zn centers. The Zn–Al bond distance is $2.471(1)$ Å, which is longer than the sum of covalent radii of the elements (2.44 Å). DippNacNacGa(1) (2) shows no reactivity towards [BeX₂(tmeda)], even at elevated temperature. Roesky et al. reported unsymmetrical dialumanes by the disproportionation of Dipp NacNacAl(1) (1) with $\int^{\text{Me2}} c \text{AAC} \cdot \text{AIX}_3$ (X = Cl, I) adducts (Scheme 5). The Al–Al bond lengths in compound 21 $(2.6327(11)$ Å) and compound 22 $(2.5953(16)$ Å) are slightly shorter than those of symmetric Al–Al bond lengths, owing to the relaxation of the electrostatic repulsion between the Al atoms.²⁶ The reactions Perspective Data control of the comparison of the comparison of the comparison of the comparison $\frac{1}{2}$ are $\frac{1}{2}$ communities and are particle. The method of the first of the common and the set of the comparison $\$

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Scheme 5 Synthesis of compounds 17–24.

of Dipp NacNacAl(I) (1) with AgX (X = OCN, SCN) resulted in compounds 23 and 24 containing two pseudohalide groups coordinated to the aluminum(III) center.²⁷ The reactions proceed via oxidative addition of the pseudohalides and elimination of the silver metal.

Harder et al. reported a combined attack of \int ^{Dipp}NacNac) $Ca^+(C_6H_6)][B(C_6F_5)_4]$ and ^{Dipp}NacNacAl(1) (1), which led to the complete dearomatization of benzene to give $C_6H_6^2$ that chelates to the $Al(m)$ center (Scheme 6).

The molecular structure of 25, however, showed a heterobimetallic complex in which the $C_6H_6^2$ ⁻ fragment is bridging to Ca and to the Al center.^{28a} Crimmin et al. reported the first catalytic method for the transformation of C–H bonds in unactivated arenes (benzene, toluene and xylene) into C–Al bonds and proposed a mechanism by which the C–H bond is activated by an unusual Al–Pd intermetallic complex.^{28b} Very recently, Harder et al. have described the stoichiometric reactions of $\text{DippNacNacAl}(I)$ (1) with $[(\text{DippNacNac})MgH]_2$ and (^{Dipp}NacNac)ZnH, respectively in benzene, where the Al center inserts into the metal hydride bond, which results in (^{Dipp}NacNac)Al(H)M(^{Dipp}NacNac) complexes (M = Zn 26, Mg 27) (Scheme 7).^{28c} However, the reaction of the calcium hydride complex with 1 in benzene followed a different course leading to benzene C–H alumination. The cleavage of the sp^2 C–H bond in unactivated arenes (benzene, toluene and xylene) by the low valent $Al(i)$ complex 1 at room temperature has been achieved by the catalytic presence of $[(^{Dipp}NacNac)CaH]_2$ (Scheme 8). The possible pathway is the combined action of nucleophilic Al and arene activation by π-coordination to a Lewis acidic Ca center, which is parallel to the first aluminyl anions. $11c$

Schulz et al. reported that a solution of $\text{DippNacNacAl}(i)$ (1) reacts with E_2Et_4 (E = Sb, Bi) in toluene to insert ^{Dipp}NacNacAl(1) (1) into the E–E bond with the formation of $\text{DippNacNacAl}(EEt_2)_2$

Scheme 6 Activation of benzene using Al(i) and Ca^{2+} .

Scheme 7 Synthesis of (DippNacNac)Al(H)M(DippNacNac) complexes (M = Zn 26 and Mg 27).

Scheme 8 Calcium hydride catalyzed C–H bond activation of unactivated arenes by oxidative addition of $\text{DippNacNacAl}(I)$ (1).

 $(E = Sb 32, Bi 33)$ (Scheme 9). Orange crystals of compound 33 were isolated from hexane at −30 °C, and slowly decompose in solution with the formation of $BiEt₃$ and elemental Bi. The analogous reaction of DippNacNacGa(I) (2) with E₂Et₄ is fully reversible and temperature dependent. Analytically pure compounds 34 and 35 were isolated from the 1 : 1 mixture of DippNacNacGa(I) (2) and E₂Et₄, respectively (Scheme 9). By changing the molar ratio of $\text{DippNacNacGa(I)} (2)$ and Bi_2Et_4 into a 1 : 2 ratio, 35 can be isolated in good yield.²⁹ Fischer et al. have achieved the reaction of \overline{D} ^{Dipp}NacNacGa(I) (2) with Bi $\left[OSO_2CF_3\right]_3$ and $\left[\left\{\text{Bi-}(\text{OC}_6\text{F}_5)_3\text{(toluene)}\right\}_2\right]$ to yield $\left[\text{(R}^fO\right)$ $\rm \left(\begin{array}{cc} \text{Dipp} \end{array} \right)$ GaBi=BiGa $\rm \left(\begin{array}{cc} \text{Dipp} \end{array} \right)$ acNac $\rm \left(\text{OTf} \right)$] $\rm \left(\text{Tf = SO}_2\text{CF}_3 \right)$ 36, C_6F_5 37). The dibismuthenes show short Bi–Bi bond lengths of 2.8111(2) and 2.8182(4) Å, respectively (Fig. 4). The reaction proceeds via oxidative addition of the Bi–O bond to $DippNacNacGa(i)$ (2) and concomitant elimination of $DippNacNacGa(OTT)_2$. Finally, Bi(OTf)₃ adds to another equiv. of 2 with subsequent dimerization to form a new type of dibismuthene stabilized by NHC-related ligands. 30 Schulz et al. described a similar insertion reaction of ^{Dipp}NacNacM with BiEt₃ that leads to ^{Dipp}NacNacMEt(BiEt₂) (M = Al 38 and Ga 39)

Scheme 9 Cleavage of Sb–Sb and Bi–Bi bonds by $\frac{Dipp}NacNacAl(1)$ and DippNacNacGa(I), respectively.

Fig. 4 X-ray single crystal structure of compound 36.

(Scheme 9). The consecutive second activation proceeds at a higher temperature through the reductive elimination of $^{\rm Dipp}$ NacNacMEt $_2$, elemental Bi and BiEt $_3$. $^{\rm 31}$

Heavy-metal complexes containing gallium-lead and gallium-mercury bonds were derived from the oxidative addition of DippNacNacGa(i) (2) with the corresponding metal precursors. The reaction of Me₃PbCl with $\frac{Dipp}NacNacGa(I)$ (2) in THF at ambient temperatures afforded compound $[\{({\rm DippNacNac})Ga(Cl)\}PbMe₃]$ (40) in high yield. In addition, the reaction between $[Pb(OSO_2CF_3)_2]$ and $DipPNacNacGa(i)$ (2) (two equiv.) leads to the complex 41 containing a $Ga-Pb^H-Ga$ linkage (Fig. 5). When two equiv. of DippNacNacGa(i) (2) were treated with $[Pb(OSO_2CF_3)_2·2H_2O]$ in THF, deep red crystals of 42 were formed in very poor yield (Scheme 10). The structure of the compound consists of a bent Ga–Pb–Ga backbone with a bridging triflate group between the Ga–Pb bond and a

Fig. 5 X-ray single crystal structure of compound 41; the THF molecule attached to lead is omitted for clarity.

weakly interacting water molecule at the gallium center. Similarly, the reaction of mercury thiolate $Hg(SC_6F_5)_2$ with Dipp NacNacGa(I) (2) (two equiv.) produced the bimetallic homo-

leptic compound 43 (Scheme 10).³² The linear cationic complex $\left[\left\{ \binom{\text{Dipp}}{\text{NacNac}}\right\} _{2}Cu\right]\left[\right.0Tf\right]\cdot 2C_{6}H_{5}F$ (44, Tf = SO₂CF₃) was obtained from the reaction of $DippNacNacGa(i)$ (2) (two equiv.) with $Cu(OTf)$ ·4CH₃CN.³³

In addition, Schulz et al. showed that DippNacNacGa(i) (2) inserted into the In-C bond of InEt₃ resulted in the formation of Dipp NacNacGaEt(InEt₂) (45). The solution of 45 in benzene and toluene tends to decompose slowly at ambient temperature. But the reaction of ^{Dipp}NacNacGa(I) (2) with I^tBu \rightarrow InEt₃ ${I^tBu = [C(Nt-Bu₂CH)₂]$ resulted in a stable compound 46, where the carbene coordinates to the indium atom. The reaction of InEt₃ with 2 equiv. of $\frac{\text{Dipp}}{\text{NacNacGa(i)}}$ (2) resulted in a double insertion product $\begin{bmatrix} \text{Dipp} \text{NacNacGa(Et)} \end{bmatrix}$ ₂InEt (47) (Scheme 11), while further insertion of Dipp NacNacGa(I) (2) into the remaining In–Et bond does not occur. Complex 57 gradually decomposes at 80 \degree C in C₆D₆, resulting in DippNacNacGa (2) and DippNacNacGaEt_2 (48). A similar reaction between DippNacNacGa(i) (2) and Cp*InEt_2 ($\text{Cp*} = \text{C}_5\text{Me}_5$) leads to the production of DippNacNacGaEt_2 (48). These reactions proceeded via the oxidation of DippNacNacGa(i) (2) with the In–Et bond and subsequent reductive elimination of Dipp NacNacGaEt₂ from the indium(III) center.³⁵

Schulz and co-workers have also reported a stibinyl radical [^{Dipp}NacNac(Cl)Ga]₂Sb[•] (49) by the reaction of two equiv. of DiappNacNacGa(i) (2) with Cp*SbCl₂ (Scheme 12), and the traces of $\text{DippNacNac}(Cl)$ GaSb (H) Cp^{*} (50) were formed as a byproduct. Similarly, the equimolar reaction yielded Dipp Nacnac(Cl)GaSb (Cl)Cp* (51), which slowly decomposed to DippNacNacGaCl_2 , decamethylfulvalen (Cp^*_{2}) and the antimony metal. The analogous reaction of $\frac{\text{Dipp}}{\text{NacNacGa(i)}}$ (2) with Cp*BiI₂ yielded a bismuthinyl radical $\binom{\text{Dipp}}{\text{NacNac}}$ ($\text{Gal}_2\text{Bi}^{\dagger}$ (52) (Scheme 12). Their formation illustrates the stepwise insertion of DippNacNacGa(i) (2) into the E–X bond of $Cp*EX_2$ followed by the homolytic bond cleavage of the E–Cp^{*} bond and elimination of Cp^{*}₂. Theoretical calculations showed the significant electron delocalization of the Sb and Bi unpaired radicals onto the Ga ligands. Compounds 49 and 52 adopt V-shaped geometries with Ga–E–Ga bond angles of $104.89(1)°$ (49) and $106.68(3)°$ (52), respectively. Compound 53 was isolated by the direct reaction of 49 with KC_8 in benzene, which gives the first struc-

Scheme 11 Syntheses and thermal stability of compounds 45-48

Scheme 12 Synthesis of compounds 49–52.

Scheme 13 Single-electron oxidation and reduction reactions of compound 49.

turally characterized compound containing a $Ga=Sb$ double bond (Scheme 13). In contrast, the equimolar redox reaction of 2 with the single-electron oxidant $[NO][BF₄]$ occurred with the formation of Dipp NacNacGaClF (54).³⁶

The Cp*AsCl₂ (Cp* = C₅Me₅) reacted with one equiv. of Dipp NacNacGa(I) (2) with the insertion of Dipp NacNacGa(I) (2) into one As–Cl bond and the subsequent formation of $DippNacNac(Cl)GaAs(Cl)Cp*$ (55) (Scheme 14). In contrast, the reaction of two equiv. of DippNacNacGa(i) (2) with Cp*AsCl₂ proceeds with the formation of gallaarsene $\frac{Dipp}{NacNacGaAsCp*}$ (56) with a Ga=As double bond and (DippNacNacGaCp*) (DippNacNacGa) $(n^1, n^2, \mu$ -As₃) (57), containing a central GaAs₃ butterfly type core. The central structural motif of 57 is the bridging As₃ three-membered ring, which coordinates in an n^1 and n^2 fashion to two Ga atoms, respectively (Fig. 6).³⁷

To evaluate the reduction potential of DippNacNacGa(i) (2) in detail, Schulz et al. studied the reactions of $\frac{DippNacNacGa(i)}{DipP}$ (2) with other Sb^{III} reagents. ^{Dipp}NacNacGa(i) (2) reacted with SbX_3 (X = NMe₂, Cl) in a 2:1 molar ratio to form Ga-substituted distibenes $[(\text{DippNacNacGaX})_2\text{Sb}_2]$ (X = NMe₂ 58, Cl 59)

Scheme 14 The reaction of $\frac{Dipp}NacNacGa(I)$ (2) with Cp*AsCl₂.

Fig. 6 X-ray single crystal structure of compound 57.

Scheme 15 Syntheses of compounds 58–61.

(Scheme 15). Heating a toluene solution of 58 at 120 °C for 24 h yielded $[(\text{DippNacNacGaNMe}_2)_{2}(\mu, \eta^{1:1} - Sb_4)]$ (60). But complex 59 required 7 days of heating at 130 \degree C to give $[(\text{Dipp}NacNacGaCl)₂(μ,η^{1:1}-Sb₄)]$ (61). Compound 58 can react with GaCl₃ to form 59; in addition, 59 reacts with Li amide to form 58. The central Sb_4 unit in 60 and 61 adopts a butterfly type conformation (Fig. 7). Interestingly, compound 60 reacted with GaCl₃ in an amide/Cl exchange reaction with the subsequent formation of 61.³⁸

To better understand the reaction mechanism leading to compounds containing $Sb = Sb$ and $Ga = Sb$ double bonds, the reactions of SbCl₃ with different equiv. of $\frac{\text{Dipp}}{\text{NacNacGa(i)}}$ (2) at different temperatures were investigated. The ^{Dipp}NacNacGaSbGa $(X)^{\text{Dipp}}$ NacNac $(X = F 62, Cl 63, Br 64, and I 65)$ compounds were formed by a twofold insertion of $\frac{Dipp}{NacNacGal(I)}$ (2) into two Sb–X bonds, followed by an intramolecular elimination of $DippNacNacGaX₂$ (Scheme 16). The reactions of two equiv. of Dipp NacNacGa(I) (2) with SbCl₃ at 8 °C yielded cyclotristibine 66. The Sb–Sb bond $(2.8205(3)-2.8437(3)$ Å) in cyclotristibine 66 is typical of the Ga–Sb single bond.³⁹

Fig. 7 X-ray single crystal structure of compound 61.

Scheme 16 Syntheses of gallastibenes containing Ga=Sb double bonds and cyclotristibine.

The insertion reactions of two equiv. of $\text{DippNacNacGa(I)} (2)$ with Sbx_3 (X = Cl, Br) at low temperature produced a doubleinserted product $\binom{\text{Dipp}}{\text{NacNacGaX}}_2$ SbX (X = Cl 67, Br 68). In addition, the reaction of isolated compounds 67 and 68 with strong σ-donating carbenes (IDipp(1,3-bis-(2,6-diisopropylphenyl)imidazol-2-ylidene), Me2cAAC) yielded four carbenestabilized stibinidenes (69–72) (Scheme 17). The studies reveal that IDipp-stabilized stibinidenes 69 and 70 show Sb-C_{carbene} single bonds, whereas the Me²cAAC-stabilized derivatives 71 and 72 exhibit Sb– C_{carbene} π-backbonding character.⁴⁰

Scheme 17 Syntheses of IDipp- (69 and 70) and ^{Me2}cAAC-stabilized stibinidenes (71 and 72).

Scheme 18 Syntheses of complexes 73 and 74.

A red-brown complex 73 was obtained by the reaction of the half-sandwich complex $[CP^{R'}Ni(\mu-Br)]_2$ $(Cp^{R'} = C_5(C_6H_4 - 4-Et)_5)$ with DippNacNacGa(i) (2) in THF (Scheme 18).⁴¹ The Cp^{R'}(centroid)-Ni-Ga linkage is bent, and the nickel atom is surrounded by an η^5 -coordinated Cp^{R'} ligand and a σ-coordinated Ga(DippNacNac) ligand, while the bromide bridges the Ni–Ga bond. The reduction of 73 with $KC₈$ afforded compound 74 where the "Cp^{R'}Ni(1)" fragment is trapped by ^{Dipp}NacNacGa. The reduction of $[\text{Cp}^{\text{R}'}\text{Ni}(\mu\text{-Br})]_2$ with KC₈ and the subsequent addition of DippNacNacGa(i) (2) also result in compound 74 albeit in smaller yield (Scheme 18).

2.3. Cleavage of E' –E single and E' =E double bonds

To stabilize Ga-coordinated dipnictenes of the type [DippNacNac $(X)Ga]_2E_2$ (E = P–Bi), the reactions of ^{Dipp}NacNacGa(I) (2) with phosphorus, arsenic, and bismuth halides and amides were studied.⁴² Two equiv. of ^{Dipp}NacNacGa(I) (2) reacted with PX_3 $(X = CI, Br)$ in toluene at ambient temperature with the insertion of ^{Dipp}NacNacGa(1) (2) into two P-X bonds, which resulted in \lceil ^{Dipp}NacNac(X)Ga]₂PX (X = Cl 75, Br 76) (Scheme 19). Similar twofold insertion reactions into $AsCl₃$ and the subsequent elimination of DippNacNacGaCl_2 resulted in the formation of the stable Ga-coordinated diarsene species, which was isolated as a green crystalline solid 77 (Scheme 20). The analogous reaction with $Me₂NASCl₂$ yielded unsymmetrically-substituted diarsene $\int^{\text{Dipp}} \text{NacNac}(Cl)Ga|As=As[Ga]$ $(NMe₂)^{Dipp}NacNac]$ (78) (Scheme 20). In contrast, the reaction of DippNacNacGa(i) (2) with As(NMe₂)₃ required much harsher reaction conditions. A mixture of $\frac{Dipp}NacNacGa(i)$ (2) and As (NMe₂)₃ heated at 165 °C for 5 days resulted in compound 79 (Scheme 19). Its analogous reaction with $\frac{Dipp}NacNacAl(i)$ (1) yielded $\binom{\text{Dipp}}{\text{NacNac}}$ (Me₂N)Al]₂As₂ (80) after heating at 80 °C for one day (Scheme 20). Finally, the reaction of Dipp NacNacGa(I) (2) with $Bi(NEt₂)₃$ also occurred with the insertion and elimination of $\text{DippNacNacGa(NEt}_2)$ and resulted in the corresponding Ga-substituted dibismuthene $\int^{\text{Dipp}} \text{NacNac}(Et_2N)Ga]_2Bi_2$ (81) (Scheme 19). The reaction of 2 with elemental tellurium yielded the Te-bridged compound $\begin{bmatrix} \text{Dipp} \text{NacNacGa-}\mu-\text{Te} \end{bmatrix}$ (82).

Scheme 19 Syntheses of complexes 75–81.

Scheme 20 Te–Te and Te–C bond cleavage reactions using DippNacNacGa(I) (2).

Moreover, the cleavage of the Te–Te and Te–C bonds upon reactions of 2 with Ph_2Te_2 and ⁱPr₂Te resulted in the formation of $^{\rm Dipp}$ NacNacGa $({\rm TePh})_2$ $({\rm 83})$ and $^{\rm Dipp}$ NacNacGa $({\rm ^ipr})$ Te $({\rm ^ipr})$ $({\rm 84}),$ respectively (Scheme 20).³⁴

Nikonov and Crimmin groups separately reported the reactions of the monomeric $Al(r)$ complex with various fluoroalkenes and fluoroarenes, resulting in the breaking of strong $sp²$ and $sp³$ C–F bonds. Aluminum(i) compound 1 undergoes a facile oxidative addition with aryl C–F bonds. 43 The reaction of 1 with an excess of hexafluorobenzene or pentafluorobenzene resulted in compounds 85 and 86, respectively (Scheme 21). A further decrease in the number of fluorine atoms in the starting arene necessitates an increase in the reaction temperature to cleave the C–F bond. The cleaving ability decreases in the order $o \rightarrow p \rightarrow m$. The addition of 1-fluorohexane or fluorocyclohexane to 1 at room temperature yielded the corresponding aluminum alkyl 92 and 93, respectively (Scheme 22).^{43a} The reaction with (E) -1,3,3,3-tetrafluoro-propene(HFO-1234ze) resulted in the immediate formation of a 4 : 1 mixture of 94-E and 94-Z (Scheme 23). The addition of hexafluoropropene to 1 gave two products which were separated by fractional crystallization from hexane. 95 is formed from the internal $sp²$ C–F bond cleavage, while 96 is the result of breaking the terminal $sp²$ C–F bond *trans* to the CF₃ group. The reaction of 1 with 3,3,3-trifluoropropene yielded 97 by the formation of a metallocyclopropane intermediate followed by β-fluoride elimination (Scheme 23). $24a$

Streubel and co-workers described the reaction of monovalent compounds $\frac{Dipp}{NacNacM}$ (M = Al, Ga) with imidazole-2thione based tricyclic 1,4-diphospinine, which produced the corresponding 7-metalla-1,4-diphosphanorbornadiene (98, 99) (Scheme 24).^{44a} Previously Nikonov et al. described the oxi-

Scheme 21 Oxidation addition of $C(sp^2) - F$ bonds by 1.

Scheme 22 Oxidation addition of $C(sp^3)$ –F bonds to 1.

Scheme 23 Reactions of 1 with (E)-1,3,3,3-tetrafluoro-propene, hexafluoropropene and trifluoropropene, respectively.

Scheme 24 Reaction of tricyclic 1,4-diphosphinine with $\frac{Dipp}NacNacAl(1)$ (1) and Dipp NacNacGa $($ |) (2) to get 7-metalla-1,4-diphosphanorbornadienes 98 and 99.

dative cleavage of the C=S bond at the metal center, $44b$ while 98 and 99 undergo the $[4 + 1]$ cycloaddition reaction, which is both kinetically and thermodynamically favorable.

The aluminum(I) compound Dipp NacNacAl(I) (1) reacted with diethyl sulfide at 50 °C, which resulted in the oxidative addition of the $C(sp^3)$ -S bond. This is the first example of $C(sp³)-S$ bond activation by a main-group element.⁴⁵ The groups of Nikonov, Crimmin, and Kinjo independently reported the reactions of monomeric Al(I) compound 1 towards C–O bonds. The oxidative addition reaction of tetrahydrofuran with $\frac{\text{Dipp}}{\text{NacNaCAl(i)}}$ (1) smoothly occurred at room temperature to give complex $101, ^{43b}$ while the reaction between 1 and benzofuran upon heating at 80 °C slowly converted it to product $102.^{43a}$ The reaction of 1 with an equiv. amount of L'_2 PhB (103) ($L' =$ oxazol-2-ylidene) in toluene instantly occurred with the insertion of $\text{DippNacNacAl}(I)$ (1) into the C–O bond, affording complex 104 involving an Al, N, and O mixed heterocyclic carbene or anionic (amino)(boryl) carbene derivative (Scheme 25).46

Treatment of $\text{DippNacNacAl}(I)$ (1) with thiourea resulted in the first carbene-stabilized terminal aluminum sulfide complexes 105 and 106 by the oxidative cleavage of the $C=$ S bond. In contrast, the mixing of compound 1 and triphenyl-

Scheme 25 C–O and C–S bond activation, respectively, by the aluminum(i) compound.

phosphine sulfide in a 1 : 1 ratio afforded a mixture of terminal sulfide $\frac{DippNacNacAl=S(S=PPh_3)}{DipPNacNacAl}$, unreacted 1, and free triphenylphosphine. The existence of the Al–S double bond in 105 and 106 was supported by DFT calculations. Complex 105 undergoes facile cycloaddition with phenyl isothiocyanate to form compound 107 along with zwitterion 108 obtained from the coupling between the liberated carbene and $PhN=C=S$ (Scheme 26).^{44b} To investigate the oxidative cleavage of the unsaturated bond of the $C=N$ unit, the reaction of $DippNacNacAl(r)$ (1) with cyclic guanidine was accomplished and showed the unprecedented cleavage of the C–N multiple bond to give the carbene-ligated amido complex ^{Dipp}NacNacAl (NHTol)(SIMe) (SIMe = $C(N(Me)CH₂)₂$) (109). The splitting of the $C=N$ bond in 109 is the first example of the oxidative addition of the C=N double bond to any metal center. 47 The DFT study supported that the production of 109 occurs via an intermediate of aluminum imide as a result of the oxidative cleavage of TolN=SIMe (SIMe = $C(N(\text{Me})CH₂)₂$) by 1. The reactions of phosphine oxides with 1 occurred readily with the formation of hydroxyl derivatives $\frac{\text{Dipp}}{\text{NacNacAl(OH)}(O=PR_3)}$ (R = Ph 110, Et 111). The C=O bond (179 kcal mol⁻¹) is much stronger when compared with the P=O bond (110 kcal mol⁻¹). Therefore, the reaction of cyclic urea 1,3-dimethyl-2-imidazolidinone with 1 resulted in an unexpected aluminum hydride $\text{DiappNacNacAlH(O=SIMe) (SIMe = C{N(Me)CH₂}₂)$ 112, with the deprotonation of the weakly acidic methyl group in the backbone of the Dipp NacNac ligand.^{48a} In contrast, the reactivity of 1 towards benzophenone afforded a ketylate species NacNacAl $(\eta^2(C,O)-OCPh_2)$ (113) (Scheme 27). The latter compound undergoes easy cyclization reaction with an unsaturated substrate.^{48b}

Recently Stephan et al. have reported the first heteroaluminirenes ^{Dipp}NacNacAl[C(R)=P] (R = ^tBu or adamantyl) 114 and 115 *via* the $\begin{bmatrix} 1 & 2 \end{bmatrix}$ cycloaddition reaction of the aluminum(i) complex $\text{DippNacNacAl}(I)$ (1) with phosphaalkynes, which feature moderate three-centered 2π-electron aromaticity

Scheme 26 Reactivity of 1 toward compounds with $C=$ S and $C=N$ unsaturated bonds.

Scheme 27 Reactivity of 1 toward compounds with $P=O$ and $C=O$ unsaturated bonds.

(Scheme 28). The compounds containing the AlCP ring can be used as synthons to prepare a series of unprecedented Al- and P-containing heterocyclic frameworks.⁴⁹

The Ga(I) compound easily undergoes cyclization with methacrolein at room temperature within 10 min to give gallium enolate 116. Unlike the aluminum congener 1, the gallium compound 2 does not cleave the $P=$ S bond of

Scheme 28 Synthesis of phosphaaluminirenes 114 and 115

Et₃P=S, even upon heating to 80 °C. With Ph₃P=S, however, a slow reaction occurs upon heating to 80 °C to obtain the sulfide $(^{Dipp}NacNacGa-S)_2$ (117). Nevertheless, compound 2 readily reacts with two equiv. of PhNCS to give product 118 via $C = S$ bond cleavage and cyclization, and also the dimer $\binom{\text{Dipp}}{\text{NacNacGa}}_2(\mu\text{-S})(\mu\text{-CNPh})$ (119) at a ratio of 5:1 (3,5- $Me₂C₆H₃$ NCO and PhNCO, respectively, reacts with 2, readily to form the coupling products 120 and 121. 1,3-Di-p-tolylcarbodiimide and DippNacNacGa(i) formed the coupling product 122 (Scheme 29).⁵⁰

 $DippNacNacAl(i)$ (1) reacted with diphenyl disulfide to afford the symmetrically substituted bis(phenyl sulfide) aluminum complex 123 via the cleavage of the S–S bond in diphenyl disulfide. In a similar fashion, the reaction of 1 with bulky tetraphenyl diphosphine resulted in the cleavage of the P–P bond over the course of 3 d at 70 °C to produce the novel aluminum bis(diphenyl phosphido) complex (124) (Scheme 30).⁴⁵

2.4 H–X bond activation

In 2014, Nikonov et al. reported the first examples of oxidative addition of a series of robust H–X bonds $(X = H, B, C, Si, N, P, O)$ to a single $Al(i)$ center. The addition of H_3SiPh , HBPin, HPPh₂,

Scheme 29 Reactivity of 2 toward compounds with unsaturated bonds.

Scheme 30 P–P and S–S bond cleavage by 1.

 HO^i Pr, H_2NtBu and H_2NPh with 1 proceeded smoothly at ambient temperature, while the reaction of bulkier silane $(H₂SiMePh)$, $H₂$, Cp^{*}H with compound 1 required heating at 70 °C. The Al (I) compound reactivity toward the oxidative addition of Dipp NacNacAlH₂ is reversible, proving the possibility of reductive elimination from the species ^{Dipp}NacNacAlH (X) (X = H, B, C, Si, N, P, O).⁵¹ Linti *et al.* described that D ipp_{NacNacGa(1)} (2) undergoes facile oxidative addition reactions towards H_2 , HSnPh₃, HNEt₂, HPPh₂, HOEt and H₂O, leading to a series of gallium hydrides, $\frac{Dipp}NacNacGaH(X)$ (X = H, Sn, O, N, P), substituted by hydride, tin, alkoxy, amido and phosphido groups.⁵² The oxidative addition of DippNacNacGa(i) (2) with HCCCH₂OH, $Ph_2Si(OH)_2$, $(nBuO)_2P(O)(OH)$ and (4-Me- C_6H_4)S(O)₂(OH) resulted in the formation of compounds ρ_{Dipp} NacNacGa(H)(μ-O)CH₂CCH (138), Dipp_{NacNacGa}(H)(μ-O) SiPh₂(OH) (139), ^{Dipp}NacNacGa(H)(µ-O)P(O)(OⁿBu)₂ (140) and $DippNacNacGa(H)(\mu-O)S(O)₂(C₆H₄-4-Me)$ (141), respectively at very low temperature (Scheme 31).⁵³ Very recently, Nikonov *et al.* have reported the *in situ* oxidation of $\text{DippNacNacGa(i)} (2)$ by N_2O or pyridine oxide which results in the generation of NacNacGa(O) as a monomeric oxide intermediate leading to the C–H bond activation (Scheme 32). 54 The oxidation of Dipp NacNacGa(I) (2) by using pyridine oxide led to the C-H bond activation of pyridine oxide, yielding ^{Dipp}NacNacGa(OH) $(\eta^1(C), \kappa^1(O) \cdot o \cdot C_5H_4N-O)$ (142), and pyridine. A similar reaction between $\frac{\text{Dipp}}{\text{NacNacGa(i)}}$ (2) and N₂O in the presence of pyridine and cyclohexanone leads to $\frac{\text{Dipp}}{\text{NacNacGa(OH)}(o-C_5H_4N)}$ (143) and $\frac{\text{Dipp}}{\text{NacNacGa(OH)}(\text{OC}_6\text{H}_9)}$ (144), respectively. The oxidation of DippNacNacGa(i) (2) in the presence of $Ph₂C=O$ resulted in the isolation of compound $D^{1}PP$ NacNacGa $(\kappa^2$ -O₂CPh₂) (145), formed from the sequential oxidation and cyclization. The in situ oxidation of the mixtures of $D^{1}P$ NacNacGa(1) (2) with O=SMe₂ and O=PEt₃ resulted in the sp³ C-H bond cleavage yielding compounds ^{Dipp}NacNacGa Perspective Dation Transcheiment Common Access Articles. Published on 23 2019. The second of the second of the second of the second of the second under a [Creative Commons Attribution 3.0 Unported Licence.](http://creativecommons.org/licenses/by/3.0/) The second under

Scheme 31 Oxidative addition of σ bonds to Al(i) and Ga(i) compounds, respectively.

Scheme 32 Sequential oxidation/C–H activation of Ga(I)

 $(CH_2S(=O)Me)OH$ (146) and ^{Dipp}NacNacGa(CH(Me)P(=O)Et₂) OH (147), respectively.

3. Conclusions and perspectives

In conclusion, we report monomeric aluminum and gallium carbenoid complexes supported by β-diketiminate ligands which possess a lone pair of electrons and a formally vacant p orbital. These features afford high electrophilic and nucleophilic reactivity that could be used in bond activation and cleavage upon reactions with small molecules. The aluminum and gallium carbenoid complexes undergo a series of oxidative addition reactions with σ H–X and E'–E bonds where E is an element from groups 13 to 16. These compounds also demonstrate the oxidative cleavage of multiple bonds and enthalpically strong bonds (M–X, $E'=E$). An extension of the synthetic approach is presented in this review for the synthesis of Group 13 metalloid complexes that consist of an unsupported M′–M $(M' = Al, Ga)$ bond. The oxidative cleavage of multiple bonds shows new and unusual reactivities. These reactivities have opened up a new realm in aluminum and gallium chemistry that could lead to many new products. Nevertheless, the compounds with low valent aluminum and gallium are still showing some limitations in catalytic applications.

However, dialumene (I) promoted both the catalytic and stoichiometric reduction of $CO₂$ to value added C1 products, which is the first example of catalysis using a homonuclear main-group multiple bond. It is important to explore reductive elimination of C–C bonds, which leads to reversible bond activation paving the way towards catalytic applications. Recent developments in the isolation of aluminyl complexes (II) will likely extend the low oxidation state Al chemistry that is used to activate the C–C bonds. We look forward to conducting further studies of the $Al(r)$ and $Ga(r)$ complexes which will result in more unprecedented bond cleavage reactions and important future applications in catalysis.

Conflicts of interest

There are no conflicts to declare.

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