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Introduction

Metallo-ligands have found wide scientific interest due to their electronic flexibility and ligand-cooperativity.¹ Here, silylenes have recently developed into a highly versatile and widely applied ligand class. $2,3$ In contrast, examples of ligands with aluminium – the second most abundant metal in the earth crust after silicon – as donor atom are scarce.^{3,4} Here, the discovery of $[(\text{AlCp*})_4]$ as first molecular Al(r) compound represented a milestone for inorganic coordination and cluster chemistry.⁵ Although being tetrameric in the solid state, the roomtemperature stable, air-sensitive $[(\text{AlCp*})_4]$ can reversibly dissociate into its monomers in solution to undergo bond formations with various small molecules or transition metals.⁶ Hence, a large variety of AlCp*-coordination complexes with electron-rich, late transition metals have been reported $(Cr,^7 W,^8)$ Fe,^{9,10} Ru,^{9,11} Co,¹² Rh,¹³ Ni,^{14,15} Pd,^{16,17} Pt,^{17,18} Cu¹⁹). Due to the diverse reactivity of AlCp*, which is isolobal to CO, complexes with terminal AlCp*-units, bridging AlCp*-ligands in binuclear complexes as well as large clusters could be prepared. $[M(AICp^*)_x]$ formation was often challenged by competing C–H activation, most likely due to the highly electrophilic nature of the coordinated Al atom in AlCp*.^{8,9} Complexes of AlCp* with

From mixed group 13 cations $[M(AICp^*)_3]^+ (M = Ga/H)(m^2)$ In/Tl) to an \tilde{Al}_4^+ cluster†

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AlCp*-complexes with transition metals have shown to be highly reactive and enable C–H or Si–H bond activation. Yet, complexes of AlCp* with low-valent main-group metals are scarce. Here, we report the syntheses of $[M(AICp^*)]_3[A(CR^F)_4]$ ($R^F = C(CF_3)_3$) with M = Ga, In, Tl, which include the first covalent Al– In and Al-Tl bonds. For $M = Ga$, AlCp*-coordination induced the formation of the dication $[Ga_2(AICp^*)_6]^{2+}$ in the solid state, which exhibits a solvent and temperature dependent monomer–dimer equilibrium in solution. By contrast, the In and Tl complexes are monomeric and prone to reduction to the metal by the electron-rich AlCp*-moieties. The QTAIM analysis suggests that the metal centres are already highly reduced in the complexes, while the positive charge is distributed onto the AlCp* units. Addition of Me₃TACN (1,4,7-trimethyl-1,4,7-triazacyclononane) to the Ga- and Tl-complex salts resulted in an isomerization to the novel low-valent $Al_4{}^+$ cation [(Me₃TACN)Al(AlCp*)₃][Al(OR^F)₄]. Intermittently formed tetrahedral GaAl₃⁺ clusters could be structurally characterized. From a detailed mechanistic study of this isomerization, the very high yield and clean preparation of [(Me₃TACN)Al(AlCp*)₃][Al(OR^F)₄] was devised from $[M(Me_3TACN)][Al(OR^F)₄]$ (M = Ga, Tl) and $[(AlCp[*])₄]$. **EDGE ARTICLE**
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electron-poor metals are scarce and limited to the f-block metals (Eu,²⁰ Yb,²⁰ U²¹). Quantum-chemical analysis of the Al– M bonding in heterobimetallic complexes revealed a dominant electrostatic character for the bonding interactions, where the covalent part largely constitutes to a ligand-to-metal σ -donation from the lone-pair at AlCp^{*} and diminished π -backbonding.10,15,20,22

However, no bimetallic cluster with $[M_xA]_{4-x}$ core, mimicking the tetrahedral structure of $[(\text{AlCp*})_4]$, is known to date. In addition, and despite the year-long research on the coordination chemistry of AlCp*, complexes to electron-rich, low-valent main-group metals have only been reported with heavy group 15 metals.²³ Hence, we were interested in bonding interactions between AlCp* and its heavier homologues in lowoxidation states. Examples for complexes with Al–Ga or Al–In bonds are scarce. The first characterized Al-Ga bond was reported by Cowley in 2005 with the donor–acceptor complex $[CP^*Ga-Al(C_6F_5)_3]$ I.²⁴ One year later, Schulz described the synthesis of the mixed group 13 donor–acceptor complexes of type $[Cp*M1-M2(^tBu)_3]$ (I-III: M1/M2 = Ga/Al, Al/Ga, In/Al).²⁵ Here, the Cp*In–Al($\rm ^fBu)_{3}$ molecule features the only structurally characterized In–Al bond known in literature. Intriguingly, Schulz reported the synthesis of the group 13 Lewis acid–base adduct $[(BDI)Ga \rightarrow Al(C_6F_5)]$ (V, BDI = β -diketiminiate), which readily inserted in benzaldehyde.²⁶ Recently, the observation of the first non-dative bonding interaction between gallium and aluminium was described by Okuda in the complex salt [(BDI) Ga(H)-Al(H)(tmeda)][B(3,5-Me₂-C₆H₃)₄] **VI.**²⁷ With the high reactivity of low-valent transition metal–AlCp* complexes as

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well as the absence of covalent Al–In and Al–Tl bonds in literature, we were interested to study the interaction of the strongly electron-donating AlCp* moiety with the M^+ cations of lowvalent gallium, indium and thallium salts of the weakly coordinating anion $\left[\text{Al}(\text{OR}^{\text{F}})_{4}\right]^{-}$ $\left(\text{OR}^{\text{F}} = \text{C}(\text{CF}_{3})_{3}\right)^{28-31}$ Very recently, we isolated the first accessible low-valent aluminium complex salt $\text{[Al}(\text{AlCp*})_3\text{][Al}(\text{OR}^{\text{F}})_4\text{]}$ VII prepared via removal of LiCp* from $\rm (AICp^*)_{4}$ with $\rm Li^+.32$ Here, we report independent syntheses and characterizations of the novel salts $[\rm M(\rm AICp^*)\rm_{3}][\rm Al(\rm OR^F)\rm_{4}]$ $(M = Ga (1), In (2), T1 (3)).$ Unexpectedly, but very useful, all the complex salts isomerize upon addition of 1,4,7-trimethyl-1,4,7 triazacyclononane (Me₃TACN) via unprecedented mixed group 13 clusters to the novel Al_4^+ cluster $[(\text{Me}_3 \text{TACN}) \text{Al} (\text{AlCp*})_3]$ $[Al(OR^F)₄]$ in high yields (Fig. 1).

Results and discussion

Syntheses of $[\mathbf{M}(\mathbf{AICp^*}_3)][\mathbf{Al}(\mathbf{OR^F})_4]$

The complex salts 1–3 were synthesized starting from $[(\text{AlCp*})_4]$ and the readily accessible fluorobenzene complexes of gallium and indium $[M(\text{PhF})_{2-3}][Al(\text{OR}^F)_4] (M = Ga^{31} \text{ In}^{28,29})$ as well as Tl $\left[\text{Al}(\text{OR}^{\text{F}})_{4}\right]$ (Scheme 1).³⁰

In a first experiment, $[\mathrm{Ga}(\mathrm{PhF})_{2\text{--}3}][\mathrm{Al}(\mathrm{OR}^{\mathrm{F}})_{4}]$ was reacted with $[(\text{AlCp*})_4]$ in 1,2-DFB (DFB = difluorobenzene) at room temperature to yield an orange solution from which orange crystals were isolated upon layering with *n*-heptane at -30 °C . Single crystal Xray structure determination (sc-XRD) revealed the formation of the dicationic cluster $[\text{Ga}_{2}(\text{AlCp*})_{6}][[\text{Al}(\text{OR}^{\text{F}})_{4}])_{2}$ $(\mathbf{1}_{\text{A}}, \textit{vide infra}).$ Yet, formation of a metallic precipitate was observed at room temperature already after a few minutes. This was prevented by conducting and keeping the reaction at -30 °C. Interestingly, switching to the less-polar, but more strongly-coordinating solvent fluorobenzene, a yellow solution was obtained from which no metal precipitation occurred at room temperature. Crystallization at room temperature by layering with n -pentane yielded orange crystals of $[\text{Ga}_{2}(\text{AlCp*})_{6}][[\text{Al}(\text{OR}^{\text{F}})_{4}])_{2}$, however,

Scheme 1 Synthesis of the complex salts $[M(AICp*)₃][AI(OR^F)₄].$

now in a second modification with a slipped Ga^+ - Ga^+ interaction $(1_B,$ Fig. 2a, vide infra).

The reactions of $[\rm{In}(\rm{PhF})_{2{\rm -}3}][\rm{Al}(\rm{OR}^{\rm{F}})_{4}]$ and $\rm{Tl}[\rm{Al}(\rm{OR}^{\rm{F}})_{4}]$ with $[(\text{AlCp*})_4]$ in fluorinated arenes yielded yellow solutions from which yellow crystals of monomeric $[\text{In}(\text{AlCp*})_3][\text{Al}(\text{OR}^{\text{F}})_4]$ 2 (Fig. 2d) and $[Tl(AlCp*)_3][Al(OR^F)_4]$ 3 separated. However, crystallization was always accompanied with decomposition and formation of some metallic indium and thallium. Hence, the respective complexes are best prepared in situ. Nevertheless, 3 could be isolated in high purity by precipitation via cannulation of the PhF-solutions onto n -heptane at room temperature.

Molecular structures

Related to the independently prepared complex salt $[AI(ALCp^*)_3]$ $[Al(OR^F)₄]₂³²$ scXRD analyses on the obtained crystals yield a trigonal-pyramidal coordination of the formal cationic gallium or indium atom by three AlCp* groups (Fig. 2b and c). In the monomeric units $[M(AICp*)_3]^+$, the average Al-Al distances are long (3.074 \pm 0.009) Å in 1_A, 3.09 \pm 0.04 Å in 1_B, 3.12 ± 0.06 Å in 2^{33} which precludes a strong covalent bonding interaction. By contrast, the Ga1–Al bond lengths in both molecular structures are similarly short and average to 2.520 \pm 0.006 A in 1_A and 2.532 \pm 0.007 A in 1_B . Hence, the Ga–Al bond lengths are close to the bond length reported by Okuda for complex VI (2.5238(9) \AA ²⁷ as well as in the formal Lewis-adducts between $Al^{III}(C_6F_5)_3$ and Ga^I -based Lewis bases $Ga^I(BDI)$ $(2.5482(4)$ $\AA)^{26}$ or $Ga^{I}Cp^{*}$ $(2.515(11)$ $\AA)^{25}$ As expected, the average In–Al bond lengths are longer at 2.75 \pm 0.01 Å. Still, the observed In–Al distances are signicantly shorter than the Edge Article

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Fig. 1 Known compounds with Al–Ga or Al–In bonds as well as novel complex salts of type [M(AlCp*)₃][Al(OR^F)₄] (OR^F = C(CF₃)₃).

Fig. 2 $\,$ (a) Orange crystals of [Ga₂(AlCp*)₆]([Al(OR^F)₄])₂ 1_A obtained by crystallization of 1 from PhF/n-pentane at room temperature. (b) Molecular structure of a [Ga(AlCp*)₃] + unit in the scXRD structure of 1_A . Hydrogen atoms and [Al(OR^F)₄] – anions omitted for clarity. Thermal displacement of the ellipsoids was set at 50% probability. (c) Molecular structure of a [In(AlCp*)₃]⁺ unit in <mark>2</mark>. Hydrogen atoms and [Al(OR^F)₄][–] anions omitted for clarity. Thermal displacement of the ellipsoids was set at 50% probability. (d) Yellow crystals of [In(AlCp*)₃][Al(OR^F)₄] **2** along with indium metal mirror. (e and f) Visualization of the two independent dicationic $[Ga_2(AICp^*)_6]^{2+}$ units obtained at -30 °C from 1,2-DFB/n-heptane (1_A) and at room temperature from PhF/n-pentane (1_B). Hydrogen atoms and [Al(OR^F)₄][—] anions omitted for clarity. Thermal displacement of the ellipsoids was set at 50% probability. (g) Visualization of the intermolecular \ln^+ –Cp* interaction in the molecular structure of 2. Hydrogen atoms and $[Al(OR^F)₄]$ ⁻ anions omitted for clarity. Thermal displacement of the ellipsoids was set at 50% probability.

previously reported dative In $\text{I} \!\to\! \text{Al}^{\text{III}}$ bond length of 2.843(2) $\rm \AA$ in III.²⁵ Interestingly, molecular structures of the indium and gallium complex salts reveal decisive differences in the intermolecular interaction between the $[M(AICp^*)_3]^+$ cations in the solid state. For 1, the dimerization to $[Ga_2(AICp^*)_6]^{2+}$ $\text{Al}(\text{OR}^{\text{F}})_4])_2$ followed in the molecular structures $\textbf{1}_{\text{A}}$ and $\textbf{1}_{\text{B}},$ with a shorter Ga–Ga distance of 2.860(1) \AA in $\mathbf{1}_A$ (Fig. 2e). By contrast, 1_B displays a "slipped" interaction between the [GaAl $_3$] ^\dagger tetrahedra with a longer Ga–Ga distance of 3.108 \pm 0.006 Å (avg., Fig. 2f). The Ga-Ga distances in 1 are significantly longer compared to literature known covalent "Ga $^+$ –Ga $^+$ " bonds as in $[Ga_5(dmap)_5]^{5+}$ (2.495 Å (avg.), dmap = dimethylaminopyridine), $^{34}\,[\mathrm{Ga}_4({}^{t}\mathrm{BuNC})_8]^{4+}\,(2.495\,\mathrm{\AA\, (avg.)})^{35}$ and $[\mathrm{Ga}_4(\mathrm{dmpe})_4]^{4+}$ $(2.485 \text{ Å} (avg.))$.³⁶ Notably, in the latter complexes the gallium cations are only coordinated by two strongly electron donating ligands that induce cluster formation via delocalization of the cationic charge onto the ligand. In contrast to 1, the $\left[{\rm In}({\rm AICp^*})_3\right]^+$ cations in 2 form coordination oligomers with the indium atom loosely coordinating to the Cp*-ligand of a neighbouring cation. Related coordination environments were reported for the solid-state structures of $M(C_5H_5)(M = In,$ Tl) and TlCp*.³⁷ The molecular structure of $[T1(AICp^*)_3]$ $\left[\text{Al}(\text{OR}^{\text{F}})_{4} \right]$ 3 shows similar interionic interactions as the indium cluster. Yet, extensive disorder of the cations and anions precludes a detailed discussion of the bond lengths.

Monomer–dimer equilibria

Interestingly, the dimerization of the gallium complex 1 in 1,2- DFB solution is an equilibrium reaction. Orange solutions matching the colour of the crystals prevail at higher

concentrations and/or low temperatures. In the UV/VIS spectrum at -40 °C (Fig. 3) a broad UV/VIS band was observed at 476 nm, which fits to HOMO/HOMO−2 \rightarrow LUMO excitations of the $\text{[Ga}_2(\text{AlCp*})_6]^{\text{2+}}$ dimer-dication computed by TD-DFT at 486 and 493 nm. No excitations in this spectral region were computed for the monomer. Warming the solution to room temperature is accompanied by a disappearance of the UV/VIS band at 476 nm and colour change to yellow, which suggests a break-up of the dimers into monomeric $[\mathsf{Ga}(\mathrm{AlCp^*})_3]^+$ cations. This monomer–dimer equilibrium is also dependent on the polarity of the solvent: in contrast to the equilibria noted in polar 1,2-DFB $(\varepsilon_{r}$ (295 K) = 13.8), dimer formation is supressed in the less polar fluorobenzene (ε_{r} (295 K) = 5.7) by its Coulomb explosion into the monocations (see ESI, Fig. S64†). Hence, suppression of dimerization in fluorobenzene allows to handle solutions of 1 at room temperature. By contrast and as expected from the distinct differences of their solid-state structures, no colour changes that would indicate a dimerization were observed for solutions of the (monomeric) indium or thallium complex 2 and 3.

NMR-spectroscopy

NMR spectra of the crystals of 1 as well as of the precipitated powders of 2 and 3 in PhF indicate a high purity, since only one Cp*-resonance is detected and no NMR signals of the starting materials were observed. Moreover, the 27 Al NMR spectra of the $[M(AICp*)_3]^+$ complex show distinct resonances of the $(AICp^*)_3$ units (Fig. 3): they shift to higher field following 1 (δ^{27} Al = -43) > 2 (δ^{27} Al = −60) > 3 (δ^{27} Al = −70). These observed ²⁷Al NMR shifts lie in between the shifts observed for the $(AICp^*)_3$ units in

Fig. 3 (a) UV/VIS spectrum of a solution of 1 in 1,2-DFB (c = 19 mg mL⁻¹) measured at rt and −40 °C as well as the computed spectrum for the dimer with the long Ga–Ga distance (TD-DFT with bp86-d3bj/def2-svp, 20 singlet excitations with Fermi exchange). (b) ²⁷Al NMR spectra of PhF solutions of the complex salts 1, 2 and 3. $[{\sf AICp^*}_2]^+$ represents a common decomposition product of AlCp * .

 $\text{[Al}(\text{Al} \text{C} \text{p}^*)_3$]⁺ ($\delta^{27} \text{Al} = -40$) and free $\text{[}(\text{Al} \text{C} \text{p}^*)_4$] ($\delta^{27} \text{Al} = -79$).³² Since both isotopes, 203 Tl and 205 Tl (29.5 and 70.5% natural abundance) possess a nuclear spin of $\frac{1}{2}$, the ²⁷Al NMR resonance of AlCp* in 3 is observed as a doublet with a coupling constant of 1 J(²⁷Al⁻²⁰⁵Tl) = 3577 Hz. As 3 represents the first reported compound with a Tl-Al bond, no other $J(27\text{Al}-205\text{Tl})$ are literature-known. In addition, only few coupling constants of $Tl(i)$ are known due to the typically highly ionic nature of $Tl(i)$ complexes making 3 rather the exception than the rule.³⁸ The quantitative decomposition of solutions of 2 and 3 with formation of a metallic precipitate took a few days at RT. The spectra of NMR-scale reactions showed a clean conversion to [(AlCp*)₄] and [AlCp*₂][Al(OR^F)₄]. Hence, a disproportionation of $[\mathrm{M}(\mathrm{AICp}^*)_3][\mathrm{Al}(\mathrm{OR}^{\mathrm{F}})_4]$ into elemental indium or thallium and $[\text{Al}^{\text{III}}\text{Cp*}_2]^+$ has occurred. Related observations were made for highly concentrated solutions of $\text{[Al}(\text{AlCp*})_3\text{][} \text{Al}(\text{OR}^{\text{F}})_4\text{].}$ Moreover, characteristic Raman bands corresponding to symmetric and asymmetric stretching vibrations of the M–Al bonds in the MAl_3^{-+} units were detected (see ESI, Fig. S7, S16 and S24†). Here, the trends ($\tilde{\nu}_{\rm{sym}}$: 515 cm $^{-1}$ (1) > 496 cm $^{-1}$ (2) > 495 cm $^{-1}$ (3) ; $\tilde{\nu}_{\rm asym}$: 471 $\rm cm^{-1}$ (1) > 460 $\rm cm^{-1}$ (2) > 451 $\rm cm^{-1}$ $(3))$ reflect the decreasing bond strength of the $M-Al₃$ cluster bonds in the order Ga > In > Tl.

Addition of Me₃TACN

To investigate the effects of cation coordination on the group 13 intermetallic clusters $1-3$, Me₃TACN $(1,4,7$ -trimethyl-1,4,7triazacyclononane) was added to fluorobenzene solutions giving yellow solutions for the gallium and indium complexes, from which yellow crystals were obtained along a few colourless, unidentiable crystals. The reaction with 3 yielded a black, metallic precipitate and yellow crystals. Surprisingly, in all reactions the yellow crystals were identified by scXRD measurements as the novel, cationic $\mathrm{Al_4}^+$ cluster [(Me₃TACN) $\text{Al}(\text{AlCp*})_3][\text{Al}(\text{OR}^F)_4]$ 4 (Fig. 4). Unexpectedly, the molecular structure of 4 includes a symmetrically bound $\mathrm{Al_4}^+$ cluster with

Al–Al bond lengths between 2.736(1)–2.789(1) \AA and very similar average Al $_{\rm Cp^*}$ –Al $_{\rm Cp^*}$ and Al $^+$ –Al $_{\rm Cp^*}$ distances of 2.75 \pm 0.03 and 2.765 \pm 0.009 Å. Apparently, the neutral ligand Me₃TACN mimics almost ideally the electronics of a Cp* ligand and matches in 4 the average Al–Al bond lengths of 2.758 A found in (AlCp^{*})₄. This contrasts with the other known³² coordinated Al_4^+ clusters that for cdp- and tmeda-ligands exhibit shorter Al⁺- Al_{Cp} * bonds of 2.662(1) A (cdp = C(PPh₃)₂) and 2.695(1) A (tmeda = $Me₂NC₂H₄NMe₂$) with slightly longer $Al_{CP} A_{CP} A_{CP}$ separations of 2.773(1) and 2.782(2) Å. For the $(dmap)_{3}$ -coordinated Al_4^+ tetrahedron, the situation is inverted: $d(\text{Al}^+ - \text{Al}_{\text{CP}}*) =$ 2.802(1) \mathring{A} , $d(\mathring{Al}_{\mathring{Ch}^*} - \mathring{Al}_{\mathring{Ch}^*}) = 2.671(1) \mathring{A}$. Hence, the electronics of the Al_4^+ -cluster bonding is highly flexible.

While the proton resonances at the Cp^* and $Me₃TACN$ residues in 4 are visible in the ${}^{1}\mathrm{H}$ NMR spectrum, the ${}^{27}\mathrm{Al}$ NMR spectrum only shows a broad signal attributed to the $(AICp^*)_3$ units at $\delta = -77$. Yet, in contrast to the highly labile (dmap)₃ complex, 4 is stable in solution over weeks and hence will be an interesting starting compound to explore the potential of these clusters as source of a cationic low-valent aluminium.

Aiming for the isolation of a mixed $GaAl_3^+$ cluster, the reaction mixture was layered directly after addition of $Me₃TACN$ with pentane. Along with crystals of 4, some yellow crystals of the mixed cluster $[(Me_{3}TACN)Al(AlCp^{*})_{2}(GaCp^{*})][Al(OR^{F})_{4}]$ 5 were obtained (Fig. 4). Here, an isomerization occurred, yielding a formally cationic Al^+ coordinated by $Me₃TACN$ along with a GaCp^{*} unit in the tetrahedral base. In 5, the Cp^{*} ring is bound very loosely, almost η^1 to the gallium atom with a short Ga–C distance of $d(Ga-C1) = 2.440(4)$ A. Interestingly, the shortest M–M bonds in the tetrahedron are the Ga–AlCp* bonds of on avg. 2.726(2) \AA , whereas all Al–Al and the Ga–Al_{TACN} distances are longer: AlTACN–ACp* $(d(A11–A12) = 2.800(2)$ Å, $d(A11-A13) = 2.837(2)$ Å), Al_{CP} * Al_{CP} * (2.826 (2) Å) and Ga- Al_{TACN} $(2.832 \, (2)$ $\AA)$. Performing the reaction and crystallization at low temperatures yielded yellow crystals of the cluster $[(Me₃TACN)$ $Ga(AlCp^*)$ ₃][Al $(OR^F)_4$] 6 before isomerization and with

Fig. 4 Molecular structures of cationic clusters in [(Me₃TACN)Al(AlCp*)₃][Al(OR^F)₄] **4**, [(Me₃TACN)Al(AlCp*)₂(GaCp*)][Al(OR^F)₄] **5** and [(Me₃TACN) Ga(AlCp*)₃][Al(OR^F)₄] **6**. Hydrogen atoms and [Al(OR^F)₄] – anions were omitted for clarity. Thermal displacement of the ellipsoids was set at 50% probability.

coordination of $Me₃TACN$ to the formally cationic gallium atom. Here, the Ga–Al distances ranging from 2.726(2) to 2.759(2) A with an average value of 2.75 \pm 0.01 A reflect the symmetric bonding of the Ga(Me3TACN) unit to the three AlCp* moieties. Whereas the average AlCp*–AlCp* bond lengths of 2.79 ± 0.06 Å (range 2.765(4)–2.833(4) Å) are similar compared to the bond lengths observed in the Al_4^+ cluster 4, more significant differences are observed upon comparison of the average Ga–N bonds in 6 and Al–N bonds in 4 with average lengths of 2.33 \pm 0.02 Å and 2.153 \pm 0.009 Å.

In situ NMR study

To investigate the mechanism of the formation of 4, the reaction of 1 with excess $Me₃TACN$ was studied by in situ NMR spectroscopy (Fig. 5). Here, already one minute after addition of Me3TACN to a solution of 1 in PhF, a quantitative consumption of the starting material 1 and formation of $[(\text{AlCp*})_4]$ was observed in the ²⁷Al NMR spectrum. Hence, the Me₃TACN ligand displaces the three AlCp*-ligands at $Ga⁺$ to form the respective $\left[{\rm Ga(Me_3TACN)}\right]^+$ complex. This is supported by the $^{1}{\rm H}$ NMR spectrum, where proton resonances fitting to the signals detected for individually prepared $\left[{\rm Ga(Me_3TACN)}\right]\hspace{-0.1cm}\left[{\rm Al(OR}^{\rm F})_4\right]$ 7 can be observed (see ESI† for characterization). After 2 h, NMR resonances attributed to the product 4 were observed: a broad shoulder in the $[(\text{AlCp*})_4]$ signal and the resonances of the $Me_{3}TACN-$ and $Cp*-$ methyl-groups of $[(Me_{3}TACN)Al(AlCp*)_{3}]^{+}$ were detected as singlets at δ (Me₃TACN) = 2.40 and 1.85 (Cp^{*}) in the 1 H NMR spectrum. With appearance of the product peaks of 4, the signals assigned to $[(\text{AlCp*})_4]$ and $[\text{Ga}(\text{Me}_3\text{TACN})]^+$ reduce in intensity. Moreover, the 71 Ga NMR spectra show the formation of monomeric GaCp*, which resonates as a characteristic singlet at $\delta = -647$ (Lit.: $\delta^{71}Ga = -653$, $\delta^{1}H = 1.93$ in C₇D₈;³⁹ $[Ga(PhF)_{2-3}][Al(OR^F)_4]$: $\delta^{71}Ga = -756$ (ref. 31)). At this stage, three overlapping singlets appear at δ^1 H = 1.92, 1.93 and 1.94, which hint to the presence of several GaCp* species. Concomitantly, a novel broad singlet was observed at δ^{27} Al = -65, potentially assigned to the mixed GaAl₃⁺ 5 cluster (δ^{27} Al_{calc.} = -60) structurally characterised independently. After 24 h, the $^1\mathrm{H}$ NMR spectrum was completely depleted from $[Ga(Me₃TACN)]^+$

and only a small peak of free $[(\text{AlCp*})_4]$ was left, whereas the signals assigned to the products GaCp* (δ^1 H = 1.93) and the Al₄⁺ cluster 4 represent the major resonances. In the 27 Al NMR spectrum, signals of 4 and (potentially) 5 became significantly more intense. In the course of the next 11 days, a slow decrease of the signals attributed to $[(\text{AlCp*})_4]$ and 5 was observed. Since no quantitative conversion could be achieved, the NMR tube was warmed to 60 $^{\circ}$ C for one hour. Subsequently, the ²⁷Al NMR spectrum displays only the broad signal of the product 4 at δ = [−]77. In the ¹ H NMR spectrum, the respective proton resonances of 4 are observed next to proton resonances of remaining free ligand and GaCp^{*} as side-product of the isomerization (δ = 1.93). Interestingly, orange crystals formed in the NMR tube upon storage, which were identified by scXRD analysis to be the ${Al_4}^+$ cluster 4, as well as traces of a metallic precipitate.

Mechanism of the formation of 4

The results of these NMR studies together with the molecular structure of the mixed $GaAl_3^+$ cluster allow for the formulation of a mechanism underlying the cluster rearrangement reaction (Fig. 5). At first, Me₃TACN reacts rapidly with $[Ga(AlCp*)_3]^+$ to form $[(\text{AlCp*})_4]$ and $[\text{Ga}(\text{Me}_3 \text{TACN})]^+$. Subsequently, the $[Ga(Me₃TACN)]⁺$ cation replaces an AlCp^{*} unit from the cluster to form the elusive, structurally characterized $[(Me₃TACN)$ Ga(AlCp^{*})₃]⁺ **6** (δ^{27} Al_{calc.} = -76), which rapidly isomerizes to give the structurally characterized $[(Me₃TACN)Al(AlCp[*])₂(GaCp^*$]⁺ in 5. From the latter a slow exchange of $GaCp^*$ for AlCp^{*} occurs, finally yielding the Al_4^+ cluster 4 and GaCp^{*} as products. To further elucidate the reaction, $Me₃TACN$ and the thallium cluster 3 were investigated by in situ NMR spectroscopy (see ESI, Section S2†). Similarly, the quick initial formation of $[(\text{AlCp*})_4]$ and $[\text{TI}(\text{Me}_3 \text{TACN})]^+$ was observed. Subsequently, $[(\text{AlCp*})_4]$ was quantitatively transformed into 4 during only 10 hours. The reaction was accompanied by formation of a black precipitate, most likely elemental Tl. Unfortunately, no intermediates were observed in the 27 Al NMR spectra, hinting to a greater lability of the mixed TIAI_{3}^{+} clusters. The proposed overall mechanism delineated in Fig. 5 is supported by computational thermodynamics.

Fig. 5 $\,$ (a)–(c) Mechanistic studies on the reaction of [Ga(AlCp*)₃] $^+$ [Al(OR F)₄] $^-$ 1 with Me₃TACN by ²⁷Al NMR (a), 1 H NMR (b) spectroscopy and ⁷¹Ga NMR (c) spectroscopy. (d) Postulated mechanism of the formation of 4 on the basis of NMR studies and scXRD structures and computed thermodynamics for the postulated reaction pathway (b3lyp-d3bj/def2-tzvpp//bp86-d3bj/def2-svp gas phase energies with Cosmo RS solvation energies).

Optimized route to 4

Since we identified the initial formation of the $Me₃TACN$ complexes as the first step of the reaction mechanism, we tested the preparation of the cationic Al_4^+ cluster by reaction of the easily accessible complex salts $[M(Me_3TACN)][Al(OR^F)₄]$ $(M =$ Ga (7), Tl (8); $c\!f$. ESI†) with [(AlCp*) $_4$] at 50 °C. Thereby, complex salt 4 could be obtained in only 4 h and in high yields exceeding 87%, significantly improving on the low yield, two-step procedure towards Al_4^+ clusters reported previously.³² Hence, these exchange reactions represent a highly promising route towards isolation of novel cationic low-valent aluminium complexes.

DFT calculations on the bonding in the mixed clusters

To investigate differences in bonding and reactivity of the isolated complex cations, a computational DFT analysis was performed. Here, similar frontier orbitals are computed for the $[M(AICp*)₃]⁺$ cations (*e.g.* for $M = Ga$ in Fig. 6a). The lone-pair at M is represented by the HOMO−2 orbital. The two degenerate HOMO/−1 display the interaction of the AlCp*-lone pairs with the p_x - and p_y -orbitals at the unique metal atom M. The LUMO has a strong p_z -character at the heavier group 13 metals. The calculated HOMO/−1/−2 energies decrease from Ga over In to Tl (HOMO−2/HOMO/−1: In at −8.9/−8.3 eV; Tl at −9.1/−8.2 eV). Due to the increase of the HOMO–LUMO gaps for the In and Tl compounds as well as the more diffuse orbitals, the formation of dimeric structures of type $[M_2(AICp^*)_6]^{2^+}$ is not feasible for In and Tl. For the observed dimer dication $[Ga₂(-)]$ AlCp* ₆]²⁺, an EDA-NOCV (energy decomposition analysis with natural orbitals for chemical valence) revealed a rather weak σ bonding interaction with a significant contribution of dispersion forces on the total attractive interactions ($\Delta E_{\rm Orb.} = -25.6$ kcal mol^{−1}, $\Delta E_{\mathrm{Disp.}} = -18.97$ kcal mol^{−1}, Fig. 6c see ESI, Table S2†). Furthermore, the differences between the Ga and the In/Tl based cations are also reflected in the EDA-NOCV analyses (Fig. 6e, more detail see ESI, Section 4†).

Here, the total interaction energies between the formal M^{\dagger} and (AlCp^{*})₃ fragments are lower for 2^+ and 3^+ compared to 1^+ , which originates from a significant drop of the total orbital interaction energy. Nevertheless, even in the indium and thallium complexes, the orbital interaction energy is greater than the electrostatic contribution to the total interaction energy. This reveals the covalent nature of the M-Al bonds ($M = AI$, Ga, In, Tl), which contrasts the dominant electrostatic character of

Fig. 6 (a) Kohn–Sham orbitals of [Ga(AlCp*)₃]⁺ computed at pbe0-d3bj/def2-tzvpp//bp86-d3bj/def2-svp level of DFT (isovalue 0.05). (b) QTAIM charges computed for the dimeric [M₂(AlCp*)₆]²⁺ cations (scXRD-structures with shortest M⁺—M⁺ distances used). (c) EDA-NOCV results for the interaction of two [M(AlCp*)₃]⁺ fragments (S) in [M₂(AlCp*)₆]²⁺ (scXRD-structures with shortest M⁺–M⁺ distances used) at bp86-d3bj/tz2p// bp86-d3bj/def2-svp. (d) QTAIM charges computed for the [M(AlCp*)₃]⁺ cations (gas-phase optimized structures). (e) EDA-NOCV results for the interaction of M⁺ (S, (s²)(p⁰)) with (AlCp*)₃ (S) computed at bp86-d3bj/tz2p//bp86-d3bj/def2-svp with ZORA.

TM-AlCp* bonds.^{10,15,20} Moreover, the gallium complex 1^+ shows an even greater total orbital interaction than the previously reported $[Al(AlCp*)_3]$ ⁺ complex.³² This observation can be attributed to the similar covalent radius⁴⁰ of aluminium ($r_{\text{cov}} = 1.21$) and gallium ($r_{\text{cov}} = 1.22$) and the even higher electronegativity⁴¹

of gallium ($X_{\rm P} = 1.81$) compared to aluminium ($X_{\rm P} = 1.61$). The EDA-NOCV results agree with the QTAIM analysis, where a negative QTAIM charge of $q_{Ga} = -0.49$ was calculated (Fig. 6b). With the dampened covalent interactions in 2^+ and 3^+ , also greatly reduced negative QTAIM charges of −0.06 and

Scheme 2 Optimized synthesis procedure for the cationic Al₄⁺ cluster [(Me₃TACN)Al(AlCp*)₃][Al(OR^F)₄] 4.

−0.19 follow for M. Hence, a formal reduction of the metal atoms by the AlCp* units occurred. In contrast, the aluminium atoms possess positive QTAIM charges, in particular for the gallium complex. This combination of a reduced metal centre covalently bonded to a positively charged and hence potentially Lewis-acidic atom suggests a promising reactivity.

Conclusion and outlook

Complexes between low-valent Al and Ga/In/Tl have been largely unexplored. Here we report the synthesis and characterization of such elusive complex salts $[M(AICp^*)_3][Al(OR^F)_4]$ $(M = Ga, In,$ Tl). Interestingly, the $\text{GaAl}_3{}^+$ complex dimerises in solution and in the solid state, but no dimerization is observed for the In and Tl complexes. Addition of $Me₃TACN$ to all the mixed group 13 cations resulted in an isomerization to the novel low-valent Al_4^+ cation ${\rm [(Me_3TACN)Al(AlCp^*)_3]}^+.$ Here, first mixed GaAl $_3^+$ clusters were structurally characterized, representing intermediates in the isomerization reaction. The novel Al_4^+ cluster could be isolated in exceptionally high yields by the optimized route starting from $[M(Me_3TACN)]^+$ and $\frac{3}{4}(AICp^*)_4$ delineated in Scheme 2. As already suggested by the decomposition of complexes in solution under formation of metallic Ga/In/Tl, negative QTAIM values indicate a formal reduction of the heavier group 13 metal cations in the complexes. With the adjacent positively charged aluminium atoms, a potentially bimetallic reactivity of the complexes will be studied in future research. Edge Article
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Data availability

Reference numbers CCDC 2193060 (1A), 2193049 (1B), 2194540 (2), 2194543 (3), 2193851 (4), 2194404 (5) and 2194403 (6). All other data supporting the findings are contained in the main text or the ESI.†

Author contributions

PD planned and carried out all experiments and analysed the data (conceptualization/investigation). PD carried out the single-crystal X-ray diffraction measurements and conducted the computational investigations (formal analysis). PD and IK wrote the manuscript (visualization/writing - original draft).

Conflicts of interest

There are no conflicts to declare.

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