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Introduction

The organometallic chemistry of aluminum is strongly dominated by the oxidation state +III, and subvalent aluminum species are quite rare. $1-4$ Even though currently on the transition from chemical curiosities to a rather well-established class of compounds, the synthesis of molecules with aluminum in the low oxidation states +I and +II is still experimentally challenging and consistently hampered by their high reactivity and pronounced tendency to disproportionation.5,6 Thus, only a handful of examples of dialanes with Al(π) centers have been reported so far.⁶⁻¹² Monovalent aluminum was first realized by Schnöckel and coworkers with their seminal discoveries of metastable AlX (X = halide)^{13,14} and $(Cp^*Al)_4$.^{15,16} These studies have inspired many main group element chemists, and several other Al(I) species have been made available,^{17,18} including other Cp-based derivatives,¹⁹⁻²² β -diketiminate-stabilized systems,²³ the recently reported NHC-stabilized dialumenes,²⁴ and aluminyl anions.²⁵ Among these, Schnöckel's $(Cp*Al)_4$ and derivatives derived thereof still represent the best-studied subvalent aluminum systems. It was shown that $(Cp^RA)_{4}$ tend to be tetrameric in

Monomeric Cp^{3t} Al(I): synthesis, reactivity, and the concept of valence isomerism†

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With the isolation of Cp^{3t} Al (1), the first monomeric Cp -based Al(I) species could be realized in a pure form via a three-step reaction sequence (salt elimination/adduct formation/adduct cleavage) starting from readily available AlBr₃. Due to its monomeric structure, reactions involving 1 were found to proceed more selectively, faster, and under milder conditions than for tetrameric (Cp*Al)₄. Thus, 1 readily formed simple Lewis acid–base adducts with t BuAlCl₂ (6) and AlBr₃ (7), reactions that before have always been interfered with by the presence of aluminum halide bonds. In addition, the 2 : 1 reaction of 1 with AlBr₃ enabled the realization of the very rare trialuminum adduct species 8. 1 also reacted rapidly with N_2O and PhN₃ at room temperature to afford Al₃O₃ and Al₂N₂ heterocycles 9 and 10, respectively. With the structural characterization of products 4 and 5, the reaction of monovalent 1 with Cp^{3t} AlBr₂ (2) provided the first experimental evidence for the concept of valence isomerism between dialanes and their Al(I)/ Al(III) Lewis adducts. **EDGE ARTICLE**

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the solid state. In solution, however, an equilibrium with their monomeric form (Cp^RA) is established, which is highly dependent on the sterics of Cp^R and temperature.¹⁹⁻²²

For $(Cp^*Al)_4$, this equilibrium lies almost completely on the side of the tetrameric aggregate at ambient temperatures, which together with the low solubility of the aggregate in common organic solvents efficiently masks the high reactivity of the Al(I) centers. Accordingly, $(Cp^*Al)_4$ appears rather unreactive, and often higher temperatures and/or longer reaction times are required. For this reason, much efforts have been made to develop monomeric Cp^RAl species by introducing bulky Cp^R ligands, which are expected to prevent any aggregation. As shown in Table 1, increasing the steric demand of Cp^R to C₅(CH₂Ph)₅, C₅H₂(SiMe₃)₃, and C₅iPr₄H proved actually successful, and no tetramer $(Cp^RAl)₄$ was detected in solution via ²⁷Al NMR spectroscopy. However, the synthetic approaches proved either tedious (long reaction times) and/or required special equipment (generation of metastable AlX precursors), and did not allow for the isolation of Cp^R Al.²⁰

Recently, we reported an alternative approach towards the generation of such $\text{Cp}^R\text{Al}(\text{I})$ species by Lewis base-induced disproportionation of a Cp*-substituted dialane into $Cp^*Al(i)$ and Cp*Al(m)Br₂ L (L = Lewis base).²⁶ We envisaged that it might be feasible to transfer this strategy to systems with bulkier Cp groups, eventually allowing for a straightforward isolation of pure monomeric Cp^RAI . Thus, we initially set out to study the suitability of 1,3,5-tri-tert-butylcyclopentadienyl (Cp^{3t}) as a stabilizing ligand for subvalent $CP^{R}Al(I)$.

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^{*a*} Decomposes above $-60\degree C$. ^{*b*} Cp^RAl detectable at 60 $\degree C$. ^{*c*} Decomposes above -30 °C.

Results and discussion

The synthetic protocol used for the generation of Cp^{3t} Al (1) is outlined in Scheme 1. Accordingly, we first developed a convenient synthesis for the precursor $Cp^{3t}AlBr_2(2)$. While simple salt elimination reactions of MCp^{3t} (M = Li, Na, and K) with AlBr₃ were found to be inefficient showing only poor conversion and were, in part, accompanied by decomposition processes, 2 was isolated in good yields by applying $(Cp^{3t})_2Mg$ in pentane solution (Scheme 1). 27 Due to the high solubility of the magnesium reagent in hydrocarbon solvents, the reaction proceeded quickly and quantitatively within 30 minutes. Characterization

of 2 in solution provided no difficulties and NMR spectroscopic parameters agreed very well with the anticipated structure. Thus, the chemical shift of the ²⁷Al NMR resonance of 2 (δ –42) strongly resembles that of Cp*AlBr₂ (δ –46).²⁸ Also, the ¹H NMR spectrum features a signal for the aromatic protons at δ 6.64, and two signals for the *t*Bu groups at δ 1.41 and 1.24. Colorless crystals of 2 were obtained by cooling a saturated hexane solution. 2 is highly sensitive towards water and oxygen, which is evident by an immediate color change from colorless to slightly brownish.

The solid state structure of 2 nicely reflects the increased steric requirements of the Cp^{3t} ligand with respect to Cp^* as evidenced by its monomeric structure (Fig. 1).²⁷ By contrast, Cp^*AIBr_2 is a halide-bridged dimer in the solid state. However, both the Al–C bond lengths $(2.175(4)-2.239(4)$ Å), as well as the distance between the aluminum center and the Cp centroid (Al– cent 1.835 \AA) compare very well with the values found in Cp^*AlBr_2 (av. Al–C 2.223 Å; Al–cent 1.865 Å).²⁸

With $Cp^{3t}AlBr_2(2)$ in hand, we next tried to make $Cp^{3t}Al(1)$ accessible via direct reductive dehalogenation, a route that has been successfully applied to the synthesis of $(Cp*Al)₄$, for instance.¹⁶ However, reduction of 2 with alkali metal-based reductants such as KC_8 , Na $K_{2.8}$, Na, Na $[C_{10}H_8]$, and Li under various conditions consistently failed, and no traceable materials could be isolated. Thus, we next attempted the reductive elimination pathway recently introduced by Fischer and Frenking for an alternative synthesis of $(Cp^*Al)_4$.²⁹ This approach requires $(Cp^{3t})_2$ AlH as the reagent, though, which upon heating might be susceptible to Cp^{3t} H elimination to generate the desired monovalent Cp^{3t} Al (1) . Again, all our attempts to selectively prepare $(Cp^{3t})_2$ AlH remained unsuccessful. Heating a mixture of $(Cp^{3t})_2Mg$ and HAlCl₂ always resulted in the formation of elemental Al among several other undefined species. According to 27 Al NMR spectroscopy, only trace amounts of a compound with a chemical shift of δ -161, indicative of the presence of 1, were present in the reaction mixtures. Thus, we were not able to separate and isolate 1. Openical Science

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> After these rather disappointing results we reviewed our most recent findings on the related Cp^* system, *i.e.* the Lewis base-induced cleavage of dialane Cp*(Br)Al–Al(Br)Cp* into

Scheme 1 Experimental approach towards the synthesis of 1.

Fig. 1 Molecular structures of 2 (left) and 3 (right) in the solid state. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å): 2 Al1–C1 2.196(4), Al1–C2 2.214(4), Al1–C3 2.175(4), Al1–C4 2.187(4), and Al1–C5 2.239(4); 3 Al1–Al2 2.533(1), Al1–C1 2.005(3), Al2–C11 2.196(3), Al2–C12 2.180(3), Al2–C13 2.136(3), Al2–C14 2.192(3), and Al2–C15 2.220(3).

 $(Cp^*Al)_4$ and $Cp^*Al(m)Br_2 \tcdot L^{26}$ Unfortunately, no Cp^{3t-1} substituted dialane was accessible by stoichiometric reduction of 2. However, keeping in mind that (i) DFT calculations have established an equilibrium between dialane Cp*(Br)Al–Al(Br) Cp* and the corresponding Lewis acid–base adduct Cp*Al·Al(Br)₂Cp*,²⁶ which is in line with the valence isomerism between these two classes of compounds, as already suggested by Cowley,³⁰ and (ii) Al-Al bond formation has also been accomplished by the reaction of $Al(1)$ species with $H₂AlNacNac$ (NacNac = [ArNC(Me)CHC(Me)NAr]⁻; Ar = 2,6-iPr₂C₆H₃)³¹ and AlX₃ cAAC (X = Cl and I; cAAC = $1-[2,6-iPr_2C_6H_3]-3,3,5,5-1$ tetramethyl-2-pyrrolidinylidene),³² we wondered if an asymmetric dialane is available by the comproportionation of $(Cp^*Al)_4$ and $Cp^{3t}AlBr_2$, or if an initial Lewis acid-base adduct is already susceptible to release the targeted Cp^{3t} Al (1) upon treatment with Lewis bases. To this end, a solution of 2 in toluene was treated with $(Cp^*Al)_4$ under sonication, whereupon the color of the suspension turned from yellow to colorless (Scheme 1).²⁷ The progress of the reaction was easily monitored by the disappearance of the ²⁷Al NMR signals of 2 (δ –46) and $(Cp^*Al)_4$ $(\delta$ –78), while no product resonances were apparent. ¹H NMR spectroscopic studies on the reaction mixture evidenced the selective formation of a single product with Cp^* and Cp^{3t} ligands in a 1 : 1 ratio, thus pointing to the formation of either an asymmetric dialane or the asymmetric $Al(I)-Al(m)$ Lewis acidbase adduct 3. In its ¹H NMR spectrum, the reaction product features one signal at δ 1.90 for the Cp* protons, as well as signals at δ 6.25, δ 1.38, and 1.24 for the Cp^{3t} ligand. In particular, the highfield shift of the aromatic Cp^{3t} protons in comparison to 2 is noteworthy and favors a Cp^{3t} ligand as part of an $Al(r)$ fragment, thus suggesting the presence of 3 (Scheme 1). Edge Article

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X-ray diffraction studies on suitable single crystals of 3 eventually verified the adduct formation.²⁷ As evident from Fig. 1, $(Cp^*Al)_4$ acted as a reductant in its reaction with 2, and the Cp^{*} fragment now features a Lewis-acidic Al (m) center containing the two bromides formerly belonging to 2. Accordingly, it is the Cp^{3t} fragment that plays the Lewis-basic Al(1) part in adduct 3, while the Cp^{3t} ligand adopts a η^5 coordination mode with bonding parameters (Al–C 2.136(3)–2.220(3) \AA , Al– cent 1.815 Å) similar to those in 2 (Al–C 2.175(4)–2.239(4) Å, Al– cent 1.835 Å). By contrast, the Cp^{*} ligand adopts a η^1 coordination mode with the $Al(m)$ center with an Al–C distance $(2.005(3)$ Å) in the same region as observed before. The Al-Al separation distance in 3 (2.533(1) \AA) is somewhat shorter than that in the Al(I)–Al(III) adducts Cp*Al·AltBu₃ (2.689(2) \AA ³³ or $\text{CP*Al}\cdot\text{Al}(C_6F_5)$ ₃ (2.591(2) Å),³⁰ which indicates a quite strong dative Al–Al interaction in 3. The Al–Al distance in 3 is rather reminiscent of the Al–Al bond length found in dialane Cp*(Br) Al–Al(Br)Cp* $(2.530(2)$ A),¹² which might be taken as another piece of evidence for the close relationship between dialanes and the corresponding $Al(I)-Al(m)$ adducts as valence isomers.

Adduct 3 appears to be an ideal precursor for the release of Cp^{3t} Al (1) by the addition of strong Lewis bases, for which reason we reacted solutions of 3 in C_6D_6 with PMe₃, IPr (1,3diisopropyl-imidazole-2-ylidene), and cAAC. In all cases, 27Al NMR spectroscopy showed the formation of a new signal at

 δ –161, indicative of a monomeric Al(1) species. In addition, the expected resonances of $Cp^*AlBr_2 \cdot PMe_3$ (δ 48) and $Cp^*AlBr_2 \cdot IPr$ (δ 106) could be found in the respective ²⁷Al NMR spectra of the reaction mixtures, while the signal of $Cp^*AlBr_2 \cdot cAAC$ was not detected, most likely due to its broadness. The formation of all three adducts was, however, clearly verified by ¹H NMR spectroscopy, which showed the expected signal patterns for one equivalent of Cp*AlBr₂ \cdot L, as well as signals at δ 5.94, 1.36, and 1.20 for monovalent 1. Compound 1 was eventually isolated on a larger scale by the reaction of 3 with cAAC in pentane, making use of the poor solubility of $Cp^*AlBr_2 \cdot cAAC$ under these conditions (Scheme 1).²⁷ After filtration and removal of the solvent under reduced pressure, a yellow oil remained, which contained essentially pure 1. Even though all attempts to obtain suitable single crystals of 1 for X-ray diffraction failed, its 27 Al NMR resonance (cf. monomeric $Cp*Al(r)$ shows a signal at δ -150 at 60 °C),²⁰ the absence of characteristic signals for a tetrameric aggregate (usually between δ -60 and δ -110),²⁰ and the results of ¹H DOSY NMR spectroscopic studies (single species, diffusion coefficient $D = 8.703 \times e^{-10} \text{ m}^2 \text{ s}^{-1}$; see Fig. S32 in the ESI†) leave no doubt on its constitution and its description as a monomeric, monovalent aluminum species.

The isolation of pure 1 also allowed us to evaluate the concept of valence isomerism between dialanes and their respective $Al(I)-Al(m)$ adducts, and to probe the possibility of generating a Cp^{3t} -based dialane by simply reacting 1 with one equivalent of Cp^{3t} AlBr₂ (2; Scheme 2).²⁷ The ²⁷Al NMR spectrum of the reaction mixture features a single broad resonance at δ –64 with a chemical shift intermediate between those of 1 (δ -161) and 2 (δ -46). Also, only one set of signals is evident in the ¹H NMR spectrum, while the chemical shift of the aromatic Cp^{3t} protons (δ 6.39) is found again intermediate between the values of 1 (δ 5.94) and 2 (δ 6.64). Hence, while the observation of a single set of NMR spectroscopic parameters would suggest the formation of a symmetric dialane (5) , the observed ²⁷Al NMR

Scheme 2 Valence isomerism between adduct 4 and dialane 5 (top). The energy profile of the isomerisation from 4 to 5 via transition state TS (bottom; M06L/Def2-SVP; free energies given in kcal mol⁻¹).

chemical shift $(\delta - 64)$ significantly differs from that of dialane Cp*(Br)Al–Al(Br)Cp* (δ –11).¹² On the other hand, for adduct 4, two sets of signals are to be expected with substantially different 27 Al NMR chemical shifts as observed in this case (cf. $\text{Cp*Al}\cdot \text{Al}(C_6F_5)$ ₃ δ -116, 107).³⁰ For a related gallium system comprising Cp*Ga and GaX₂Cp* (X = Cl and I), Jutzi and coworkers similarly reported only one set of NMR signals for the Cp^{*} moieties upon mixing of the components, even at -80° C,³⁴ which indicates a corresponding valence isomerism in the case of gallium. Thus, our NMR spectroscopic studies rather indicate the presence of a rapid equilibrium between adduct 4 and dialane 5, which we tried to verify by VT NMR spectroscopic studies in toluene solution (–90 °C to +100 °C). Unfortunately, we were not able to derive a clear picture of this process, most likely because the exchange process is too fast on the NMR time scale across the entire temperature range. Nevertheless, a few hints can be extracted from the measurements. While low temperature 27 Al NMR spectroscopy was not very helpful in solving this puzzle (the signal only broadened, eventually disappearing in the baseline; see Fig. $S28-S31$ in the ESI[†]), the ²⁷Al NMR resonance at δ –64 is shifted to a higher field $(\delta$ –90) at increased temperatures. This chemical shift is reminiscent for Al(I) centers in Lewis acid–base adducts (cf. Cp*Al·Al(C₆F₅)₃ δ -116).³⁰ In addition, a low intensity signal appears at δ 90, which is found in the same region as the tetracoordinate aluminum atom of Cp*Al·Al(C₆F₅)₃ (δ 106) or adducts 6 (δ 146) and 7 (δ 95; see below). These findings suggest that at higher temperatures, the equilibrium mixture of 4/5 is pushed to the side of adduct 4. By contrast, the 1 H NMR signals of the aromatic $C_5H_2tBu_3$ protons are gradually shifted to a lower field upon cooling the sample ($\delta_{\rm rt}$ 6.39; $\delta_{-60\degree\rm C}$ 6.59) and finally became chemically non-equivalent at -80 °C , as evidenced by two low-field signals ($\delta_{-80^{\circ}\mathrm{C}}$ 6.89, 6.64). It should be noted that this decoalescence is not related to the 4/5 isomerization process, but only to the rotation of the Cp^{3t} rings, which gets frozen at low temperatures. Thus, we still see only one Cp^{3t} ligand environment, which in combination with the low-field chemical shift of these protons (characteristic for three-coordinate aluminum species such as 2) indicates an equilibrium mixture shifted to the side of the symmetric dialane 5. In order to evaluate the thermodynamic stabilities of both species 4 and 5, we performed DFT calculations at the M06L/Def2-SVP level of theory (Scheme 2 (bottom)). Accordingly, dialane 5_{calc} is energetically more stable by 4.0 kcal mol^{-1} than the asymmetric adduct structure 4_{calc}, and both species are separated by a lowlying transition state TS with an activation barrier of only 5.6 kcal mol⁻¹ for the transformation of 4 into 5. These results fit very well to our experimental findings that only mixtures of adduct 4/dialane 5 can be generated in solution, and that the interconversion is possible over the entire temperature range. Eventually, both isomers could be characterized in the solid state by X-ray diffraction (Fig. 2). Consistent with the spectroscopic results and the calculated stabilities, we were able to selectively crystallize dialane 5 at $-30\,^{\circ}\mathrm{C}$ and adduct 4 at room temperature from the same pentane reaction mixture.²⁷ In agreement with the equilibrium shown in Scheme 2, both isolated species showed the same averaged NMR spectra as the Openical Science Works Article. Published on 14 of dialation 4

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Fig. 2 Molecular structures of 4 (left) and 5 (right) in the solid state. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å) of 4 Al1–Al2 2.599(3), Al1–C1 2.209(7), Al1–C2 2.183(7), Al1–C3 2.216(7), Al1–C4 2.192(7), Al1–C5 2.178(7), and Al2–C11 2.063(7); 5 Al1–Al2 2.586(3), Al1–C1 2.231(8), Al1–C2 2.343(8), Al1–C3 2.392(8), Al1–C4 2.247(8), and Al1–C5 2.172(8).

initial reaction mixture. Importantly, we were able to isolate again 4 and 5 from both of these solutions under the same conditions as described above, thus providing a definite experimental proof for this isomerization process. To the best of our knowledge, this represents the first instance, in which the suggested valence isomerism between dialanes and their respective $AI(I) - AI(m)$ adducts could be demonstrated directly within a single molecular system. It should be noted, however, that Nikonov and co-workers had previously observed the reversible disproportionation of a dialane to the corresponding pair of discrete Al(I) and Al(II) species, which, in contrast to 4, did not form an $Al(I)-Al(m)$ Lewis pair.³¹

The structural parameters of adduct 4 are comparable to those of 3, but illustrate the increased steric demand of the Cp^{3t} ligand. Accordingly, the Al–Al separation distance of 4 (2.599(3) Å) is slightly larger than that of the less bulky adduct $3(2.533(1))$ A), but in the same region as in Cp*Al·Al(C₆F₅)₃ (2.591(2) A).³⁰ In analogy to 3, the Cp^{3t} ligand of the monovalent aluminum center of 4 (Al1; Al-C 2.178(7)-2.216(7) Å, Al-cent 1.824 Å) adopts a η^5 coordination mode with essentially the same bonding parameters as in 3 (Al–C 2.136(3)–2.220(3) Å, Al–cent 1.815 Å). Similarly, the Cp^{3t} ligand of the Al(III) fragment of 4 (Al2) is again bound in a η^1 fashion, while the Al2–C11 distance $(2.063(7)$ Å) is slightly elongated with respect to 3 $(2.005(3)$ Å). The bulkier nature of the Cp^{3t} ligand is also evident in the molecular structure of dialane 5. Thus, the torsion angle Br1– Al1–Al2–Br2 $(-152.71(8)^\circ)$ and the Al–cent distances of 5 (1.925 \AA , 1.930 \AA) are significantly larger than in the related dialane Cp*(Br)Al–Al(Br)Cp* (Br1–Al1–Al2–Br2 102.04(5); Al–cent 1.902 Å, 1.904 Å).¹² Other relevant structural features of 5 strongly resemble those found for Cp*(Br)Al–Al(Br)Cp*, i.e. both species show η^5 -coordination modes for the Cp rings and similar Al-Al distances (5: Al–C 2.164(8)–2.392(8) Å, Al–Al 2.586(3) Å; Cp^{*}(Br) Al–Al(Br)Cp*: Al–C 2.169(4)–2.365(4) A, Al–Al 2.530(2) A).¹²

The reaction of isolated 1 with simple haloalanes aimed at elucidating similarities and differences of the reactivity patterns of 1, $(Cp^*Al)_4$, and other monovalent Al(I) species. It has been shown that halide substituents are usually non-innocent in the reaction of Al(I) species with haloalanes, leading to ionic or

comproportionation products. Thus, the reaction of $(Cp^*Al)_4$ with AlCl₃ enabled the isolation of the decamethylaluminocenium cation $[\text{Cp}^*{}_{2}\text{Al}]^{{\scriptscriptstyle +}},^{35}$ while ß-diketiminatestabilized LAl(1) asymmetric dialanes were formed via comproportionation.³² Only when reacted with halide-free alanes such as Al $(\mathrm{C}_6\mathrm{F}_5)_3$ (ref. 30) or Al $t\mathrm{Bu}_3, ^{33}(\mathrm{Cp^*Al})_4$ acts as a Lewis base. In our work, monovalent 1 did not show any evidence for reduction chemistry upon treatment with t BuAlCl₂ and AlBr₃, but rather afforded adducts 6 and 7 quantitatively within seconds (Scheme 3).²⁷ As expected, ¹H NMR spectroscopic parameters of 6 and 7 strongly resemble each other and those of adducts 3 and 4, both featuring the typical set of three signals for the monovalent Cp^{3t}Al part (6: δ 6.13, 1.30, and 1.22; 7: δ 6.10, 1.13, and 0.99). The ¹H NMR resonance for the *t*Bu group of 6 (δ 1.09) integrates very well with 9H, thus confirming the presence of a 1 : 1 ratio of the Al(I) and Al(III) fragments. The ²⁷Al NMR spectra of 6 (δ 146) and 7 (δ 95) showed only a single resonance each for the tetracoordinate $Al(m)$ fragments, while the signals of the $Al(r)$ centers were not detected presumably because of their large full widths at half maximum. The molecular structures of 6 and 7 are unobtrusive and reminiscent of known $Al(i)$ / $Al(m)$ Lewis acid–base adducts such as 3 and 4, or the aforementioned literature examples.²⁷ The Al-Al separation distances (6: 2.621(2) Å; 7: 2.554(1) Å), as well as the Al-C (6: $2.176(4) - 2.203(4)$ Å; 7: $2.142(2) - 2.188(2)$ Å) and Al-cent distances (6: 1.820 Å; 7: 1.796 Å) are found in the expected regions, while the Cp^{3t} ligand in 7 appears to be more tightly bound to the $Al(r)$ center as in all other adducts (Fig. 3).

A closer inspection of the synthesis of 7 revealed another highly interesting reactivity of 1. When a solution of 1 was added dropwise to a solution of $AlBr₃$, 7 was formed quantitatively as a single product. By reverse addition, however, a minor

Scheme 3 Lewis base reactivity of monovalent 1.

Fig. 3 Molecular structures of 6 (left) and 7 (right) in the solid state. Hydrogen atoms are omitted for clarity. The asymmetric unit of 6 contains two independent molecules with similar metrical parameters; only one molecule is shown. Selected bond lengths (A) : 6 Al1-Al2 2.621(2), Al1–C1 2.202(3), Al1–C2 2.174(4), Al1–C3 2.203(4), Al1–C4 2.194(4), Al1–C5 2.176(4), and Al2–C11 1.973(4); 7 Al1–Al2 2.554(1), Al1–C1 2.188(2), Al1–C2 2.177(2), Al1–C3 2.142(2), Al1–C4 2.178(2), and Al1-C5 2.172(2)

amount of a second aluminum-containing species was observed by ²⁷Al NMR spectroscopy (δ -71), which we could eventually isolate as colorless crystals in low yield. This compound was subsequently identified by X-ray diffraction as the trialuminum species 8 (Fig. 4) with a very rare structural motif. In the solid state, 8 features three distinct aluminum centers with aluminum in two different oxidation states, being best described as the Lewis pair of the asymmetric dialane $Br₂Al-$ Al(Br)Cp^{3t} and **1.** Thus, the Lewis-basic Al(I) atom Al3 of **1** forms a dative interaction with the sterically less crowded and more Lewis-acidic Al (II) center Al2 of the dialane fragment $\text{(Br}_2\text{Al} Al(Br)Cp^{3t}$, while the second $Al(n)$ center Al3 remains 'threecoordinate' and covalently bound to Al2 $(Br₂Al–Al(Br)Cp^{3t})$. The observed Al–Al bond lengths are consistent with this picture, and Al1–Al2 $(2.538(1)$ Å) and Al2–Al3 distances $(2.601(1)$ Å) show typical values for covalent and dative Al–Al bonds, respectively. To the best of our knowledge, only one related, even though ionic, non-cluster-type trialuminum species has been mentioned in the literature, $[Cp^*{}_2Al_3I_2]^+ [Cp^*Al_2I_4]^{-.36}$ The selective, large scale synthesis of 8 was accomplished by either reacting adduct 7 with one equivalent of 1 or by dropwise addition of 0.5 equivalents of AlBr₃ to a solution of 1 (Scheme 4). The ²⁷Al NMR spectrum of 8 features only a single resonance for a three-coordinate aluminum atom $(\delta$ -71), which clearly substantiates the formulation of 8 as a dialane adduct. The absence of resonances for the other two aluminum atoms can be rationalized in combination with ¹H NMR data, which both suggest the fluxional behavior of the bromide atoms. Thus, only one set of signals for the Cp^{3t} protons is evident in the ${}^{1}H$ NMR spectrum. These findings strongly indicate scrambling of the bromide substituents between the two Cp^{3t} Al units. While the exact mechanism for the formation of 8 is not known, the fluxional behaviour in solution is consistent with a mechanism proceeding either (i) by reversible oxidative addition of one Al– Br bond of 7 to the Al (i) center of 1, (ii) by reversible coordination of the Lewis-basic $Al(1)$ center of 1 to the hypothetical valence isomer of 7 (*i.e.* Cp^{3t}(Br)Al-AlBr₂), or (iii) by initial formation of the bis(adduct) $Cp^{3t}Al \cdot AlBr_3 \cdot AlCp^{3t}$ and subsequent bromide migration. So far, we do not have any spectroscopic evidence for a favoured reaction pathway. Edge Article

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> Monovalent aluminum compounds have also been used to study Al=E multiple bonding $(E = N, P, As, O, Se, and Te)$. Thus, $(Cp^*Al)_4$ was shown to form tetrameric heterocubane type

Fig. 4 Molecular structures of 8 in the solid state. Hydrogen atoms are omitted for clarity. Selected bond lengths (Å): 8 Al1-Al2 2.538(1), Al2-Al3 2.601(1), Al1–Br1 2.473(1), Al2–Br2 2.319(1), and Al2–Br3 2.319(1).

clusters $(Cp^*A|E)_4$ when reacted with elemental O_2 , $N_2O,^{37}$ Se, and Te,^{16,38} respectively. Similarly, treatment of $(tBu₃SiAl)₄$ with molecular oxygen afforded $(tBu₃SiAlO)₄$ among other products.³⁹ Employing β -diketiminate-stabilized LAl(1) in the reaction with O_2 led to a dimeric product, $(LAIO)_2$.⁴⁰ These results imply a strong impact of the electronic and steric parameters of the $Al(i)$ species on the product structure, while a truly monomeric $Al=O$ compound is still absent in the literature.⁴¹ Accordingly, we became curious about what kind of product 1 might generate in view of its larger steric bulk as compared to $(Cp^*Al)_4$. To this end, a degassed solution of 1 in C_6D_6 was subjected to an atmosphere of N_2O , which instantaneously resulted in the formation of the oxygenated species 9 (Scheme 5).²⁷ NMR spectroscopic studies suggested the presence of a symmetric compound in solution, as evidenced by a single set of signals for the Cp^{3t} ligand in the ¹H NMR spectrum of 9 (δ 6.49, 1.61, and 1.39). However, NMR data were not sufficient to clarify the identity of 9, particularly because no 27 Al NMR resonance was found. Operation Science $\gamma = \frac{1}{G\mu_1} + \frac{G\lambda_0 R_1}{G\mu_2} + \frac{G\lambda_0 R_2}{G\mu_3} + \frac{G\lambda_0 R_1}{G\mu_4} + \frac{G\lambda_0 R_2}{G\mu_5} + \frac{G\lambda_0 R_1}{G\mu_6} + \frac{G\lambda_0 R_2}{G\mu_7} + \frac{G\lambda_0 R_2}{G\mu_8} + \frac{G\lambda_0 R_1}{G\mu_9} + \frac{G\lambda_0 R_2}{G\mu_9} + \frac{G\lambda_0 R_2}{G\mu_$

It was an X-ray diffraction study on suitable single crystals of 9 that eventually helped to assign the exact composition.²⁷ As can be seen from Fig. 5, oxygenation of 1 with N_2O goes along with the formal oligomerization of hypothetical monomeric Cp^{3t} Al=O to afford the six-membered Al₃O₃ heterocycle 9. Thus, a new member of the $[LAO]_n$ family with $n = 3$ is accessible making use of the steric demand of the Cp^{3t} ligand. The central Al_3O_3 ring of 9 is almost planar with all Al-O distances within a narrow range $(1.697(2)-1.705(2)$ Å). The Al-O distances of 9 are thus signicantly shorter than those in dimeric $(LAIO)₂$ ⁴⁰ and tetrameric $(R^*AIE)₄$ $(R = SitBu₃$ 1.836 $A³⁹$;³⁹ R = Cp*_{calc} 1.845 Å),³⁷ which might indicate larger ionic contributions to the Al–O bonds in 9. As a consequence, the Al–C distances $(2.216(3)-2.295(3)$ Å) of all three exocyclic η^5 -bound Cp^{3t} ligands are somewhat elongated in comparison to all other Cp^{3t} based compounds described in this study.

The related $RAI=NR'$ chemistry appears even more unpredictable due to the presence of a substituent at the nitrogen center. Hence, the reaction of $(Cp^*Al)_4$ with different azides $\rm R^{\prime}N_3$ was used to realize a variety of Al–N-based compounds, the formation of which can be rationalized by the stability of the

Scheme 5 Synthesis of Al_3O_3 (9) and Al_2N_2 (10) heterocycles from 1.

Fig. 5 Molecular structures of 9 (left) and 10 (right) in the solid state. Hydrogen atoms are omitted for clarity. Symmetry operation for the conversion of Al1 to Al1' in 10: $-x + 1$, $-y + 1$, $-z + 1$. Selected bond lengths (Å): 9 Al1-O1 1.701(2), Al1-O3 1.703(2), Al2-O1 1.702(2), Al2-O2 1.697(2), Al3–O2 1.705(2), Al3–O3 1.701(4), Al–C 2.216(3)– 2.296(3); 10 Al1-N1 1.809(2), Al1-N1' 1.827(2), Al1-C11 2.295(2), Al1-C12 2.244(2), Al1–C13 2.216(2), Al1–C14 2.198(2), Al1–C15 2.249(2), N1–C1 1.396(3).

initial iminoalane Cp*Al=NR' and the nature and size of R' . While for $R' = SiIPr₃$, SiPh₃, and SitBu₃ symmetric iminoalane dimers with a regular Al_2N_2 core have been obtained,⁴² the reaction with $Me₃SiN₃$ was more complicated affording an irregular iminoalane dimer.⁴³ With the bulky azide $MesN₃$, the monomeric iminoalane did not participate in oligomerization reactions, but was rather stabilized by a C–H bond activation/ proton migration sequence.⁴³ Other $(RAINR')_n$ structures ranging from monomeric iminoalanes $(n = 1)^{44}$ to species with $n = 16$ were also described in the literature.⁴⁵⁻⁴⁷ However, either β -diketiminate-stabilized LAl(1) was used as a reagent in the reaction with azides, or other approaches were applied in their synthesis.

In our work, the reaction of 1 with PhN_3 proceeded smoothly at room temperature to afford dimeric iminoalane 10 in good yields (Scheme 5).²⁷ Again, the degree of Cp^{3t} Al=NPh aggregation is not apparent from NMR spectroscopy in solution, which pointed to either the presence of a monomeric iminoalane, or a symmetric species with a single set of 1 H NMR signals for the Cp^{3t} ligand (δ 6.78, 1.57, and 1.18) and the phenyl group (δ 7.35, 7.11, and 6.97) in a relative ratio of 1 : 1. No ²⁷Al NMR resonance was observed for 10. X-ray diffraction eventually established a dimeric iminoalane structure of 10 in the solid state with the Cp^{3t} ligands in typical η^5 coordination modes (Al–C 2.198(3)– 2.295(2) \AA , Al-cent 1.883 \AA ; Fig. 5). The basic structural parameters of 10 are very similar to those of the iminoalane $(Mes*AlNPh)_2$ described by Power,⁴⁸ while noticeable differences are found in Roeskys' actually more related iminalane $(\mathbb{C}p^*\text{AlNS}it\text{Bu}_3)_2$.⁴² In analogy to $(\text{Mes}^*\text{AlNPh})_2$ (Al–N 1.824(2) Å, $\Sigma_{\rm N}$ 360°), the Al₂N₂ core of 10 is perfectly planar, as expected for a centrosymmetric heterocycle, with Al–N bond lengths of 1.809(2) and 1.827(2) Å, and trigonal planar nitrogen atoms ($\Sigma_{\rm N}$) 359.9°). By contrast, $(Cp^*AlNSitBu_3)_2$ features somewhat elongated Al–N bonds $(1.835(2)-1.842(2)$ Å) and nitrogen atoms that significantly deviate from planarity $(\Sigma_N, 353.4^\circ, 353.7^\circ)$, and most importantly η^1 -coordinated Cp* ligands. These structural distinctions most likely trace back to the steric demand of the extremely bulky $NSitBu₃$ fragments.

Conclusions

In this contribution, we demonstrated that Cp^{3t} Al (1) can be obtained by a three-step protocol involving the Lewis baseinduced release of 1 from an $Al(i)/Al(m)$ adduct as the key step. Thus, it was possible for the first time to isolate an analytically pure monomeric Cp-based Al(I) compound, a task that has consistently been hampered by synthetic difficulties. Detailed reactivity studies clearly furnished evidence that reactions involving 1 proceed more selectively (adducts 6 and 7), faster, and under milder conditions (heterocylces 9 and 10) than for $(Cp^*Al)_4$, which is in line with our expectations that the monomeric nature of 1 might provide some experimental benefits. Thus, monovalent 1 is expected to show a rich chemistry, particularly with respect to the realization of novel structural motifs (dialane adduct 8), and the activation of small molecules, two areas that are part of our current research efforts. One finding that deserves special attention is the experimental verification (reaction of 1 with 2) of the concept of valence isomerism between dialanes and the respective $Al(i)$ / $Al(m)$ Lewis acid–base adducts (structural characterization of 4 and 5), which so far has only been proposed on the basis of computational studies. Edge Article

Conclusions we deconvented dut Cp³M (1) can be 12/2024 2019. Nonethed, 2.5 are the set of the control interviewing the Task has 12 May 2004. Downloaded at the Same May 2002 2019. In Exam and Alexander Chan

Conflicts of interest

The authors declare no conflict of interest.

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