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Introduction

It is now 50 years since Schrock reported on the synthesis and structural characterization of tantalum methylidene Cp₂Ta (CH₂)Me as the first isolable molecular metal-methylidene complex.¹ Soon after in 1978, Tebbe described the heterobimetallic titanium methylidene Cp₂Ti(CH₂)(Cl)AlMe₂,² which ever since advanced organic synthesis and, more specifically, excelled as Tebbe's reagent in functional group tolerant methylenation reactions.³ Early mechanistic and structural investigations have also contributed to an in-depth understanding of the role of the organoaluminum component.⁴ For example, while Schrock pointed out that Cp₂Ta(CH₂)Me does reversibly form an adduct with AlMe₃,¹ ideal performance of Tebbe's compound is achieved in the presence of a Lewis base like NEt₃, resulting in the displacement of "stabilizing" Me₂AlCl and the formation of transient [Cp₂Ti=CH₂].⁵ Not surprisingly, research into Tebbe-like nucleophilic carbene

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Half-sandwich scandium methylidenes†

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AlMe₃ sticks, GaMe₃ quits. A series of Lewis acid stabilized scandium methylidenes with commercially available cyclopentadienyl ligands Cp^R (Cp^R = C₅Me₅ (Cp*), C₅Me₄SiMe₃ (Cp')) were synthesized. The-salt metathesis reaction of new half-sandwich dichloride precursors Cp^RScCl₂(µ-Cl)Li(thf)₃ with LiAlMe₄ and AIMe₃ at ambient temperature yielded [Cp^RSc(AIMe₄)Cl]₂. [Cp'Sc(AIMe₄)Cl]₂ was further methylated at ambient temperature to yield Cp'Sc(AlMe₄)Me. At 70 °C, the reaction of Cp^RScCl₂(µ-Cl)Li(thf)₃ with LiAlMe₄ and AlMe₃ led to the formation of Lewis acid stabilized Sc/Al₂ methylidenes $Cp^{R}Sc(CH_{2})(AlMe_{3})_{2}$. The new mixed Sc/Al/Ga methylidene Cp'Sc(CH₂)(AlMe₃)(GaMe₃) was obtained from the reaction of Cp'Sc (AlMe₄)Me with GaMe₃. When heated, complex Cp'Sc(CH₂)(AlMe₃)(GaMe₃) converted into the Sc/Al methylidene [Cp'Sc(CH₂)₂AlMe]₃ via release of the comparatively weak Lewis acid GaMe₃ and methane. The core of trimeric [Cp'Sc(CH₂)₂AlMe]₃ can be described as a triscandacyclohexane {Sc(CH₂)₃ stabilized by a trialacyclohexane {Al(CH₂)}₃ via Pearson hard/hard matching. Complexes Cp^RSc(CH₂)(AlMe₂)₂ and [Cp'Sc(CH₂)₂AIMe]₃ differ in rigidity, thermal stability and reactivities toward ketones and Lewis bases. The isolated methylidenes were analyzed by ¹H, ¹³C(¹H), ⁴⁵Sc, and variable temperature ¹H NMR spectroscopy, SC-XRD, IR spectroscopy, and elemental analysis. Complexes Cp^RSc(CH₂)(AlMe₃)₂ feature pronounced Sc···HC α -agostic interactions. The reaction of Cp^RScCl₂(μ -Cl)Li(thf)₃ with LiAlMe₄ and AlMe₃ was investigated via in situ ¹H and ⁴⁵Sc NMR spectroscopy.

> chemistry also triggered feasibility studies on discrete methylidene complexes of metals adjacent to titanium. More recent studies include the structural characterization of the first monometallic, terminal methylidenes of zirconium(IV) and vanadium(v), (PNP)Zr(OAr)(CH₂) and (PNP)V(NAr)(CH₂) (PNP = $N[2-P(iPr)_2-4-methylphenyl]_2$, Ar = 2,6- $iPr_2C_6H_3$) by the group of Mindiola.^{6,7} The latter two synthesis protocols do not involve any organoaluminum components. In contrast, the first discrete scandium methylidene complex (PNP)Sc(CH₂) $(AlMe_3)_{21}^{8}$ also reported by the Mindiola group in 2008 draws not only on kinetic stabilization of the [Sc=CH₂] moiety via a sterically demanding PNP pincer but also Lewis acid/base interactions with AlMe₃ (Chart 1, I).⁸⁻¹⁰ We have previously shown that such multifaceted stabilization is also applicable for the larger rare-earth metals (Ln) when employing bulky hydrotris(3-R'-5-R-pyrazolyl)borato scorpionate ligands Tp^{R',R} and either AlMe₃ or GaMe₃, e.g., $(Tp^{tBu,Me})La(CH_2)(EMe_3)_2$ (E = Al, Ga) and (Tp^{Me,Me})Sm(CH₂)(AlMe₃)₂.⁹ For these larger metals, the ubiquitous cyclopentadienyl ancillary ligand only led to trimetallic clusters of the type $(C_5Me_5)_3Ln_3(\mu-CH_2)X_4$ (X = Cl, Br).11,12 Compared to d-transition-metal methylidenes, the bonding in rare-earth-metal methylidenes is increasingly ionic and stabilization of the highly polarized Ln-C(methylidene) bond is achieved via cluster formation involving bridging CH_2^{2-} moieties. The cluster size is greatly impacted by the steric shielding of the ancillary ligand. It is also noteworthy



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that except for the cuboidal cluster $[(C_5Me_5)Sc(\mu-CH_2)]_4$ (Chart 1, **II**, Cheng, 2021)¹³ all previously reported methylidene complexes of the smallest rare-earth metal scandium feature noncyclopentadienyl ancillary ligands (Chart 1, **I** and **III-VI**).¹³⁻¹⁸ The formation of tetrametallic **II** *via* thermal treatment (90 °C) of $[(C_5Me_5)Sc(CH_3)(\mu-CH_3)]_2$ clearly parallels the transformation of the Petasis reagent Cp₂Ti(CH₃)₂, which generates transient [Cp₂Ti=CH₂] upon heating at 60–80 °C.¹⁹

Many of the isolated nucleophilic metal methylidenes were probed in the methylenation of ketones and compared to Tebbe's reagent.

Given the feasibility of the half-sandwich complex (C_5Me_5) Sc(AlMe₄)₂,¹⁸ although decomposing at temperatures >–20 °C, we wondered about the existence of discrete scandium methylidene complexes with the commercially available ligands Cp* (C_5Me_5) and Cp' ($C_5Me_4SiMe_3$). This might bring us a step closer to a scandium analog of the cyclopentadienyl-supported Tebbe reagent. Further, fathoming any group 13 effect (AlMe₃ *versus* GaMe₃) might unveil distinct methylidene formations.^{9b,20}

Here, we present discrete AlMe₃-stabilized half-sandwich scandium methylidenes of the known motif **I**. Additionally, we present a novel mixed scandium aluminum gallium methylidene complex of motif **I**. This trimetallic complex decomposes under the release of GaMe₃ and CH₄ granting access to a hexametallic $\{Sc_3Al_3\}$ cluster motif which is new in rare-earth-metal chemistry.

Results and discussion

Half-sandwich scandium chloride precursors

Initial investigations bore on the known metathesis reaction of $ScCl_3(thf)_3$ and Cp'Li ($Cp' = C_5Me_4SiMe_3$) to yield Cp'

 $ScCl_2(thf)_2^{21}$ as well on Okuda's work on half-sandwich methyl scandium complexes (*vide infra*).²² In our hands, the metathesis reactions of $ScCl_3(thf)_3$ and Cp^RLi ($Cp^R = C_5Me_5$, $C_5Me_4SiMe_3$) yielded repeatedly crystals of ate complexes $Cp^RScCl_2(\mu$ -Cl)Li(thf)₃ ($\mathbf{1}^R$) (Scheme 1). While 1' was obtained in 74% yield, the initial low yield of $\mathbf{1}^*$ could be increased from 31% to 55% by raising the reaction temperature to 40 °C and additional toluene extraction. Complexes 1' and $\mathbf{1}^*$ are isostructural exhibiting a piano stool configuration, and were further analyzed by ¹H, ¹³C{¹H}, ⁷Li, and ⁴⁵Sc NMR spectroscopy in C_6D_6 , as well as elemental analysis. Compound $\mathbf{1}^*$ loses THF at prolonged exposure to vacuum.

Targeting a potential Tebbe-like reactivity, the dichlorido precursors were treated with 5.2 (1') or 5.5 (1*) equivalents of AlMe₃. Not unexpectedly, addition of AlMe₃ to 1^R did not afford alkylation but displacement/precipitation of LiCl only,²³ along with the formation of white $[Cp^RScCl_2]_4$ (2^R). The reaction mixture was heated to 130 °C for one day, yielding a palevellow solution. Upon cooling to ambient temperature, complexes 2^R crystallized and were analyzed by single-crystal X-ray diffraction (SC-XRD). Tetrameric 2^R are isostructural to other rare-earth-metal half-sandwich dihalides including [Cp^RScI₂]₄.^{21b,24} Although LiCl is not included in the crystal structure, 2^R showed a ⁷Li NMR signal even after crystalline isolation. Treatment of 2^{R} with THF yielded 1^{R} (Scheme 1).

Metathesis reactions of Cp^RScCl₂(µ-Cl)Li(thf)₃ with LiAlMe₄

In 2008, the group of Okuda reported on metathesis reactions of $Cp'ScCl_2(thf)_2$ with two equivalents of $LiAlMe_4$. This led to the successful isolation of $[Cp'ScMe_2]_2$.²² When $AlMe_3$ was added in the metathesis reaction, an "orange oil" was obtained at ambient temperature, then depicted as $[Cp'Sc(AlMe_4)_2]$. The orange oil could not be identified unequivocally. The actual congener $Cp*Sc(AlMe_4)_2$ was isolated by our group in 2019,



 $\label{eq:scheme1} \begin{array}{l} \mbox{Synthesis of scandium chloride precursors } Cp^RScCl_2(\mu-Cl)Li $$ (thf)_3 (1^R) and their reactions with AlMe_3 to yield [Cp^RScCl_2]_4 (2^R). $$ \label{eq:scheme1} \end{array}$

showing high sensitivity toward temperatures above -20 °C.18 This sparked our interest in re-investigating Okuda's orange oil. Accordingly, complexes 1^{R} were reacted with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ in toluene at ambient temperature overnight. This resulted in the isolation of mixed chlorido tetramethylaluminato complexes $[Cp^{R}Sc(AlMe_{4})Cl]_{2}$ (3^R), which were analyzed by ¹H, ¹³C{¹H}, and ⁴⁵Sc NMR spectroscopy in C₆D₆ and toluene-d₈, SC-XRD, as well as elemental analysis. Complexes 3^R crystallized as chlorido-bridged dimers.^{11b,25} The ⁴⁵Sc NMR spectra displayed three signals (cf. ESI[†]). The ¹H NMR spectra, however, did not indicate multiple species. Therefore, we assumed that 3^R are subject to monomer-dimer equilibria or to the formation of even larger aggregates in solution which is only seen in the ⁴⁵Sc NMR spectra. A variable temperature (VT) NMR study in toluene- d_8 lent support to this hypothesis (cf. ESI[†]). The spectrum at 60 °C revealed only one broad peak in the methyl region. At -80 °C, four sharp peaks appeared, assignable to the monomer and the dimer. The VT 45Sc NMR spectra show that the peak at about 210 ppm increases with lower temperatures, whereas the peak at about 400 ppm gets bigger with higher temperatures. We were hoping that these mixed chlorido/tetramethylaluminato complexes 3^R may directly lead to methylidene complexes, since the congeners of the larger rare-earth metals form trinuclear methylidenes when reacted with THF.¹¹ In the case of the small scandium however, treatment of 3^{R} with THF did not form such methylidenes. Instead, undefined decomposition occurred, and, on one occasion, the extremely sensitive aluminate cleavage product [Cp*ScMeCl(thf)]₂ could be identified via a crystal snapshot.

Prolonging the reaction of 1' with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ in toluene for at least two days led to the isolation of [Cp'Sc(AlMe₄)Me] (4') (Scheme 2). Compound 4' features the orange oil reported by Okuda (vide supra),²² also in our hands opposing successful crystallization. To prove its identity, 4' was analyzed by ¹H, ¹³C{¹H}, and ⁴⁵Sc NMR spectroscopy in C_6D_6 . The ¹H NMR signal of the Sc-CH₃ protons of 4' appeared as one broad peak, suggesting a high fluxionality, as known for tetramethylaluminato and methyl ligands. Moreover, this broad peak showed integrals between 15 and 22 which is less than the expected 24 for putative $[Cp'Sc(AlMe_4)_2]$. The higher integral numbers (compared to 15 in 4') originate from residual AlMe₃(thf) as the oil had to be evacuated and co-evaporated with toluene extensively in order to get rid of all AlMe₃(thf). The ⁴⁵Sc NMR resonance of 4' was detected at 404.2 ppm, shifted considerably to lower field compared to Cp*Sc(AlMe₄)₂ (245.1 ppm).¹⁸ This downfield shift of 4' suggests that it indeed is not putative $[Cp'Sc(AlMe_4)_2]$. To strengthen our hypothesis and without a crystal structure on hand, we envisaged derivatization reactions to further prove the identity of 4' (Scheme 2). First, we reacted $[Cp'ScMe_2]_2$ with 1 equiv. AlMe₃ which led to the sole quantitative formation of 4'. Compound 4' reacted with TMEDA according to a donorinduced aluminate cleavage yielding single crystals of (tmeda) (AlMe₃)₂,²⁶ verified by a unit cell check. The ¹H and ⁴⁵Sc NMR spectra of this reaction identified [Cp'ScMe₂]₂ as the coproduct



Scheme 2 Syntheses of Cp'Sc(AlMe₄)Me (4') using various precursors, and reactivities of Cp'Sc(AlMe₄)Me (4').

(*cf.*, ESI[†]).²² Reacting 4' with the donor DMAP afforded the monomeric adduct Cp'ScMe₂(dmap) (5'), which was analyzed by ¹H, ¹³C{¹H}, and ⁴⁵Sc NMR spectroscopy, SC-XRD, and elemental analysis. The ¹H NMR spectrum of 5' in C₆D₆ revealed that coproduct (dmap)AlMe₃ could not be removed entirely by crystallization. (dmap)AlMe₃ itself was crystallized from the supernatant and identified by a unit cell check.²⁷ To rule out that the orange oil itself is a methylidene complex, 4' was treated with cyclohexanone. This led to the identification of dimeric cyclohexenolate complex [Cp'ScMe(OC₆H₉)]₂ (**6**') *via* a crystal snapshot (*cf.*, ESI[†]). The small quantity of **6**' was also analyzed by ¹H and ⁴⁵Sc NMR spectroscopy in C₆D₆.

In order to skip the isolation of 1^{R} , we probed a one-pot reaction of ScCl₃(thf)₃, 1.1 equiv. Cp'H and 5 equiv. MeLi in THF. This led to the new half-sandwich scandium trimethyl ate complex [Li(thf)₄][Cp'ScMe₃] (7'), which was analyzed by ¹H, ¹³C{¹H}, ⁷Li, and ⁴⁵Sc NMR spectroscopy in C₆D₆, SC-XRD, and elemental analysis. Complex 7' is isostructural with [Li (tmeda)₂][Cp*LuMe₃], which was obtained from a one-pot reaction of LuCl₃, NaCp*, and MeLi.²⁸ When treated with 6 equiv. AlMe₃, complex 7' gave 4' in essentially quantitative yield. Compound 4' was also obtained from the reaction of halfsandwich dibenzyl complex Cp'Sc(CH₂C₆H₄NMe₂-*o*)₂ with trimethylaluminum, further proving the orange oil 4' to exhibit the connectivity [Cp'Sc(AlMe₄)Me]. Compound 4' could not be isolated in pure form. It contained at least 5% impurities such as 8' (*vide infra*) or Me₂Al(CH₂C₆H₄NMe₂-*o*).

Monomeric methylidenes Cp^RSc(CH₂)(AlMe₃)₂

The thermal behavior of 4' at temperatures >70 °C indicated release of methane but yielded a red oil containing an undefined mixture of products. Similarly, performing the metathesis reaction of $1^{\mathbb{R}}$ with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ at an increased temperature of 70 °C led to methane formation, yielding a red oil as well. However, now repeated recrystallization of that red oil from an n-hexane solution gave methylidenes Cp^RSc(CH₂)(AlMe₃)₂ (8^R) as colorless crystals in moderate yields of 66% ($R = Cp^*$) and 47% (R = Cp') (Scheme 3, Fig. 1 and Fig. S74[†]). Complexes 8^R were analyzed by ¹H, ¹³C ^{1}H , and ⁴⁵Sc NMR spectroscopy in C₆D₆ and toluene- d_8 , SC-XRD, DRIFTS, and elemental analysis. Compounds 8^R feature Lewis acid stabilized methylidenes with the recurring structural motif $[Ln(CH_2)(EMe_3)_2]^+$ (Ln = rareearth metal, E = Al, Ga) as detected for (PNP)Sc(CH₂)(AlMe₃)₂ (I) and $Tp^{tBu,Me}Ln(CH_2)(EMe_3)_2$ (Ln = La, Ce, Nd; E = Al, Ga; Ln = Sm; E = Ga).⁸⁻¹⁰ The ¹H NMR spectrum of 8^{R} at ambient temperature displayed the methylidene protons as two broad signals (Table 1: δ = 0.95, -0.01 ppm for 8', δ = 0.84, -0.13 ppm for 8*). The slight upfield shift of one methylidene proton points to Sc…HC α-agostic interactions. The methyl groups appeared as one singlet ($\delta = -0.33$ ppm for 8', $\delta =$ -0.43 ppm for 8*), accounting for a rapid exchange of bridging and terminal methyl groups at ambient temperature. To further elucidate this fluxional behavior in solution, a variable temperature ¹H NMR study was conducted between -80 and +80 °C (cf., ESI[†]). At low temperatures, the methylidene signals get sharper and geminal coupling of the two inequivalent protons is detected. The bridging and the two terminal methyl groups split into three signals at -80 °C. At 80 °C, the methylidene signals coalesce into one very broad peak. Crucially, the



Scheme 3 Synthesis of AlMe₃-stabilized methylidenes $Cp^{R}Sc(CH_{2})$ (AlMe₃)₂ (8^R) via tandem metathesis/thermally induced methyl deprotonation.



Fig. 1 Crystal structure of **8***. Atomic displacement ellipsoids were set at 50% probability. Hydrogen atoms except methylidene hydrogen atoms omitted for clarity. Selected interatomic distances [Å] and angles [°]: Sc1-C11 2.1624(17), Al1-C11 2.0914(18), Al2-C11 2.0647(17), Sc1-H11A 2.17(2), Sc1-C11-H11A 77.2(13) (ESI†). Complex **8**' is isostructural (Fig. S74†).

spectroscopic signature of the methylidene moiety is distinct from that reported for (PNP)Sc(CH₂)(AlMe₃)₂ (**I**), which showed a singlet at 0.69 ppm at ambient temperature shifting to 0.75 ppm at -50 °C.⁸

The ${}^{13}C{}^{1}H$ NMR spectra of 8^R at ambient temperature show peaks at 53.2 (8^*) and 53.0 ppm (8') for the methylidene carbon atoms (Table 1), rather comparable to the respective shift of $Tp^{tBu,Me}La(CH_2)(AlMe_3)_2$ ($\delta = 50.9 \text{ ppm}$)^{9a} than of (PNP) $Sc(CH_2)(AlMe_3)_2$ (I, 28.8 ppm). The signals of the methylidene carbon atoms are broadened due to ${}^{1}J$ coupling to 7/2 and 5/2 nuclei of ⁴⁵Sc and ²⁷Al, respectively. The peaks were assigned to the methylidene carbon atom by ¹H-¹³C HSQC NMR coupling with the methylidene protons. The Sc…HC α-agostic interaction is also seen in the SC-XRD analysis with short Sc-H (agostic) distances of 2.17(2) Å (8*) and 2.14(3) Å (8'). The Sc-C (methylidene)-H(agostic) angles are 77.2(13)° (8*) and 75.1 (16)° (8'). The Sc–C(methylidene) distances are 2.1624(17) Å (8*) and 2.1735(18) Å (8') (Table 1 and Fig. 1). To the best of our knowledge, these are the shortest distances of scandium to a methylidene carbon atom detected so far. In accordance with the NMR spectroscopic signatures, the Sc-C(methylidene) distance is markedly shorter than the 2.3167(17) Å detected for (PNP)Sc(CH₂)(AlMe₃)₂ (I).⁸ The short Sc-C(methylidene) distance and the ¹J coupling hint to comparatively strong Scmethylidene interactions in 8^R. The Al-C(methylidene) distances of **8*** are 2.0647(17) Å and 2.0914(18) Å, and of **8**' are 2.0898(18) Å and 2.0879(18) Å. Compared to (PNP)Sc(CH₂) (AlMe₃)₂ (I, 2.0387(18) Å and 2.0369(18) Å), the longer Al-C (methylidene) distances in 8^{R} support the assumption of a stronger Sc-C(methylidene) interaction in 8^R compared to $(PNP)Sc(CH_2)(AlMe_3)_2.$

The complexes $\mathbf{8}^{R}$ and $Tp^{tBu,Me}La(CH_2)(AlMe_3)_2$ show α -agostic interactions, fluxionality at ambient temperature, geminal coupling of the methylidene protons at -80 °C and a similar ${}^{13}C{}^{1}H{}$ NMR shift of the methylidene carbon atom (*vide supra*). Thus, one could draw the comparison of the frag-

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Table 1 Scandium–C(methylidene) distances and selected NMR spectroscopic signatures (δ)

Compound	CN	Sc–C [Å]	$^{13}\mathrm{C}\{^{1}\mathrm{H}\}[\mathrm{ppm}]$	¹ H [ppm]	Ref.
$Cp*Sc(\mu_3-CH_2)(AlMe_3)_2$ (8*)	6	2.1624(17)	53.2	0.84(br)/-0.13(br)	This work
$Cp'Sc(\mu_3-CH_2)(AlMe_3)_2(8')$	6	2.1735(18)	53.0	0.95(br)/-0.01(br)	This work
$Cp'Sc(\mu_3-CH_2)(AlMe_3)(GaMe_3)$ (10')	6	2.1724(15)	53.1	0.94(br)/0.10(br)	This work
$[Cp'Sc(\mu_3-CH_2)(Al(\mu_3-CH_2)Me)]_3$ (11')	6	2.245(3) - 2.3338(18)	81.2/35.0	0.73(d)/-0.13(s)/-1.37(d)	This work
$(PNP)Sc(\mu_3-CH_2)(AlMe_3)_2$ (I)	6	2.3167(17)	28.8	0.69(s)	8
$\left[Cp^*Sc(\mu_3-CH_2) \right]_4 (II)$	6	2.210(7) - 2.303(7)	_	1.71(s)	13
$L_{3}^{1}Sc_{3}(\mu_{3}-CH_{2})(CH_{3})_{4}$ (III)	6	2.367	123.9	3.32(s)	15
$Sc_3(\mu_4-CH_2)_2(\mu_2-CH_3)_3(AlMe_4)_2(AlMe_3)_2$ (IV)	6	2.298(3) - 2.442(2)	78.9	2.06(s)	18
$L_{2}^{2}Sc_{2}(\mu_{2}-CH_{2})(\mu_{2}-PC_{6}H_{3}iPr_{2}-2,6)$ (V)	5	2.193(3)/2.232(3)	_	$5.62(d, {}^{3}J_{\rm PH})$	17
$[L^{3}Sc(\mu_{2}-CH_{2})]_{2}$ (VI)	5	_	_	1.51/1.48/1.13/1.09	16
$(NN^{FC})Sc(\mu_3-CH_2)(AlMe_3)_2$	5	2.6921(18)	-4.7	-1.42(d)	35

 $L^{1} = PhC(NC_{6}H_{4}iPr_{2}-2,6)$. $L^{2} = N_{2}C_{3}HAr^{iPr}_{2}-1,5-Me_{2}-2,4$. $L^{3} = NC_{4}H_{2}(CH_{2}PCy_{2})_{2}-2,5$. $NN^{Fc} = 1,1'-fc(NSitBuMe_{2})_{2}$. Structures of I-VI are depicted in Chart 1.

ments $[Cp^RSc]^{2+}$ and $[Tp^{tBu,Me}La]^{2+}$. The congener with the smaller yttrium, $[Tp^{tBu,Me}Y(CH_2)(AlMe_3)_2]$, is elusive since the reaction comes to a halt at $Tp^{tBu,Me}YMe(AlMe_4)$.^{9*a*} It seems that the steric bulk of the ancillary ligand (L) has to match the size of the rare-earth-metal cation (Ln) in order to obtain methylidene complexes from the intermediary formed $[LLn(AlMe_4)_2]$. Such size match is the case for the fragments $[Cp^RSc]^{2+}$ and $[Tp^{tBu,Me}La]^{2+}$. In **8**^R, the steric requirement is achieved with the commercially available ligands C_5Me_5 and $C_5Me_4SiMe_3$.

To gain more insight into the formation mechanism of 8', the reaction of 1' with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ was monitored by ¹H and ⁴⁵Sc NMR spectroscopy without agitation of the reaction mixture (Fig. 2 and Scheme 4). At initial ambient temperature, the chlorido-bridged dimer 3' is detected as the main species along with different unidentified species, maybe differently sized clusters of [Cp'ScCl₂] (2'). Upon heating to 70 °C, full conversion into the mixed methyl/ tetramethylaluminato complex 4' evolved within 5 minutes. Over the course of two hours at 70 °C, methylidene complex 8' emerged as the main product. Relating to the 45Sc NMR spectra, at the beginning, the peak pattern of 3' appeared in the range between 200 and 420 ppm. Then, after 5 minutes at 70 °C, complex 4' (δ = 404.2 ppm) emerged as the main intermediate (Scheme 4). During prolonged heating at 70 °C, 8' became the main product already after 2 h (δ = 272.6 ppm). Consequently, 8' could be obtained according to another reaction path by treating isolated 4' in *n*-hexane with excess AlMe₃ at 70 °C.

On one occasion, the red supernatant of precipitated **8'** yielded very few colorless crystals of another methylidene complex, $[(Cp'ScCH_2)_3LiAlMeCl_3]_n$ (**9'**). The obtained small quantity of **9'** was analyzed by ¹H and ⁴⁵Sc NMR spectroscopy in C₆D₆, and SC-XRD. Compound **9'** consists of three [Cp' ScCH₂] units which are linked to a one-dimensional coordination polymer *via* LiAlMeCl₃ moieties (*cf.*, ESI[†]). The ¹H NMR spectrum of **9'** shows two very broad signals at 0.97 ppm and 1.36 ppm attributable to the methylidene protons. Compound **9'** revealed a very broad ⁴⁵Sc NMR signal at 429.2 ppm being in stark contrast to the chemical shift of **8'** ($\delta = 272.6$ ppm), and more in the range of cluster Sc₃(µ₄-CH₂)₂(µ₂-



Fig. 2 Monitoring of the reaction of **1**' with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ in a J. Young-valved NMR tube without agitation *via in situ* ¹H (top) and ⁴⁵Sc (bottom) NMR spectroscopy in C₆D₆, beginning with **3**' (blue), then intermediate **4**' (red), and final product **8**' (yellow). Unmarked spectra were measured (after cooling) at 26 °C. Spectra marked with an asterisk (*) were measured at 70 °C.

 $CH_3)_3(AlMe_4)_2(AlMe_3)_2$ (Chart 1, **IV**, $\delta = 388.3/331(2:1)$ ppm). However, the amount of pure isolated **9**' was not sufficient for further analytics and various attempts failed to reproduce **9**'.



Scheme 4 Mechanism of reaction of 1' with $LiAlMe_4$ and $AlMe_3$ with intermediates 3' and 4' to yield the final product 8'.

The Cp*-analog compound $[Cp*Sc(AlMe_4)Me]$ (4*) was not encountered. Prolonged reaction times of 1* with 3 equiv. LiAlMe₄ and 4 equiv. AlMe₃ at ambient temperature only led to the isolation of $[Cp*Sc(AlMe_4)Cl]_2$ (3*), alongside small amounts of 8*. A reaction of $Cp*Sc(CH_2C_6H_4NMe_2-o)_2$ with 3 equiv. AlMe₃ at ambient temperature did not yield a distinct product nor ¹H and ⁴⁵Sc NMR spectra similar to 4'.

GaMe₃ in place of AlMe₃: formation of a trimeric Lewis acid stabilized scandium methylidene

GaMe₃ is a weaker Lewis acid than AlMe₃, according to Pearson's HSAB concept.²⁹ This was key to the isolation of otherwise elusive compounds by our group. For example, GaMe₃ gets easily displaced in imido complexes Tp^{tBu,Me}Ln $(NAr)(GaMe_3)$ (Ln = Nd, Sm; Ar = C₆H₃iPr₂-2,6) via treatment with THF to yield the terminal imides.^{9b} The AlMe₃-containing congener did not show the same reactivity. In another case, GaMe₃ was released under reduced pressure from Tp^{tBu,Me}Ln (Me)(GaMe₄) to yield the terminal methyl complex Tp^{tBu,Me}LnMe₂.³⁰ Thus, we envisaged organogallium instead of organoaluminum-stabilized scandium methylidenes. The initial reactions of $Cp'Sc(CH_2C_6H_4NMe_2-o)_2$ with 3 or 4 equiv. GaMe₃ at ambient temperature yielded a mixture of ill-defined products. Since the instability of elusive $[Cp'Sc(GaMe_4)_2]$ was safe to assume or any subsequently evolved methylidene complex might also be unstable, we targeted a mixed aluminum/gallium compound.

Given the formation of Sc/Al_2 methylidene 8' in a reaction of 4' with AlMe₃ at 70 °C, 4' was treated with GaMe₃ in a

J. Young-valved NMR tube and heated to 70 °C (Scheme 5). After 6.5 h, the heterotrimetallic methylidene $Cp'Sc(CH_2)$ (AlMe₃)(GaMe₃) (10') was evidenced as the main product through a 45 Sc NMR chemical shift similar to 8' (275.3 vs. 272.6 ppm). Compound 10' was (re-)crystallized from a saturated *n*-hexane solution several times, and analyzed by ${}^{1}H$, ${}^{13}C$ $\{^{1}H\}$, and ^{45}Sc NMR spectroscopy in toluene- d_{8} , SC-XRD, and elemental analysis. Compound 10' is isotypic to 8' and features a very similar Sc-C(methylidene) distance (2.1724(15) vs. 2.1735(18) Å). The ¹H NMR spectrum of **10**' shows a broad peak for each AlMe₃ and GaMe₃ moiety. While the shift of the AlMe₃ moiety equals that of 8^{R} , the GaMe₃ signal appeared at lower field (-0.17 ppm). The ¹H NMR spectroscopic monitoring also revealed that 10' decomposes slowly at ambient temperature via methane evolution. This lack of stability may be due to GaMe₃ being a softer Lewis acid compared to AlMe₃²⁹ once more underpinning the limited coordination capability of GaMe₃ in Pearson-type hard/hard metal-ligand pairs. Following this trend, AlMe₃ instantly replaces GaMe₃ when AlMe₃ was added to 10' to form 8'.

The decomposition of **10**' is complete after 3 days at 80 °C. During this time, **10**' releases GaMe₃ and another equivalent of CH₄. Afterwards, crystals of the new Sc/Al methylidene [Cp' Sc(CH₂)₂AlMe]₃ (**11**') could be isolated, devoid of any gallium component (Scheme 5). Thus, the decomposition pathway of **10**' favors the formation of a second methylidene over the formation of a methylidyne.^{13,31} Compound **11**' was analyzed by ¹H, ¹³C{¹H}, and ⁴⁵Sc NMR spectroscopy in C₆D₆, SC-XRD, DRIFTS, and elemental analysis. The molecular structure of **11**' can be described as a trimer of a half-sandwich scandium methylidene stabilized by a trimeric aluminum methyl methylidene (Fig. 3).



Scheme 5 Reactions of Cp'Sc(AlMe₄)Me (4') with AlMe₃ or GaMe₃ to form Sc/Al₂ methylidene **8**' or mixed Sc/Al/Ga methylidene **10**' and its reaction to [Cp'Sc(CH₂)₂AlMe]₃ (**11**').



Fig. 3 Crystal structure of 11'. Atomic displacement ellipsoids were set at 50% probability. Hydrogen atoms except methylidene hydrogen atoms omitted for clarity. Selected interatomic distances [Å] and angles [°]: Sc1-C21 2.246(3), Sc1-C24 2.322(3), Al1-C21 2.054(3), Al1-C24 2.015(3), Sc1-H21A 2.29(3), Sc1-H24B 2.41(3), Sc1-C21-H21A 79.6(17) (ESI†).

Overall, **11**' features a $Sc_3Al_3(CH_2)_6$ distorted hexagonal prismatic core, or in other words a triscandacyclohexane ring $\{Sc_3(CH_2)_3\}$ faced with a trialacyclohexane ring $\{Al_3(CH_2)_3\}$ through Pearson-type hard/hard matching. In terms of nuclearity, **11**' ranges between the homometallic parent methylidene complex $[Cp'Sc(CH_2)]_4$ (**II**)¹³ and dodecameric $[MeAl(CH_2)]_{12}$.³² The core structure of **11**' with two chair-like six-membered rings is a common motif for aluminum imides.³³ For rareearth-metal chemistry, it has been rarely observed. A crystal snapshot of homometallic $Cp^*{}_6Y_6(CH_2)_4Me_4$, which was obtained *via* thermolysis of $[Cp^*YMe_2]_3$, comes close,³⁴ but organo-heterobimetallic Ln/Al clusters exhibiting this structural motif have not been encountered to this date.

The ¹H NMR spectrum of **11**' shows one signal set for the three Cp' ligands and one signal set for the three methyl groups, accounting for a C₃ symmetry in solution. The methylidene moieties show three signals. The methylidene protons of the $\{Sc_3(CH_2)_3\}$ ring appeared as a singlet at -0.13 ppm at ambient temperature. In contrast, the ¹H NMR signal of the methylidene protons of the $\{Al_3(CH_2)_3\}$ ring is split at 0.73 and -1.37 ppm, mainly due to the distinct axial and equatorial ring positions. This splitting might also involve weak α-agostic interactions of one methylidene proton with scandium. In contrast to the rather fluxional methylidene moieties in 8^{R} and 10', the ¹H NMR methylidene signals in 11' do not coalesce into one signal even at 120 °C. Instead, the methylidene protons of the $\{Al_3(CH_2)_3\}$ ring show geminal coupling at ambient temperature and at low temperatures, arising from the rigid chair conformation.

Considering the ⁴⁵Sc NMR chemical shifts, that of **11**' (δ = 396.7 ppm) is shifted downfield compared to **8**' (δ = 272.6 ppm), and hence similar to **9**' (δ = 429.2 ppm). Like observed for **9**', the ⁴⁵Sc NMR signal of **11**' is much broader

than that of 8' (FWHM: 5566 Hz vs. 2715 Hz, cf., ESI[†]). The 13 C {¹H} NMR spectrum of **11**' at ambient temperature shows two broadened signals for the methylidene carbon atoms of each ${M_3(CH_2)_3}$ ring (M = Sc: δ = 81.2 ppm, Al: δ = 35.0 ppm). Similarly, compound $Sc_3(\mu_4-CH_2)_2(\mu_2-CH_3)_3(AlMe_4)_2(AlMe_3)_2$ (Chart 1, IV) featuring two $Sc_3Al(\mu_4-CH_2)$ coordination environments revealed a ${}^{13}C{}^{1}H$ (methylidene) resonance at 78.9 ppm.¹⁸ Apparently, the ${}^{13}C{}^{1}H$ NMR chemical shifts of the methylidene carbon atoms are strongly influenced by the number of adjacent rare-earth-metal and aluminum centers. For example, $[Ln_3(\mu_3-CH_2)]$ moieties revealed the largest down- $\label{eq:2.1} field\ shifts\ (\textit{e.g.}\ (NSiMe_{3}Ar^{iPr})_{3}Y_{3}(\mu_{2}\text{-}Me)_{3}(\mu_{3}\text{-}Me)(\mu_{3}\text{-}CH_{2})(thf)_{3},$ δ = 100.2 ppm).^{12a} Changing the ratio of coordinating metal centers from Sc₂Al in **11**' (δ = 81.2 ppm) to ScAl₂ in **8**' with a very short Sc–C(methylidene) distance (2.1735(18) Å) causes a significant shift to higher field (δ = 53.0 ppm). The signal of the methylidene carbon atom $[ScAl_2(\mu_3-CH_2)]$ in 11' (δ = 35.0 ppm) featuring a longer Sc-C(methylidene) distance (2.246(3) Å) is similar to the methylidene shifts in (PNP)Sc $(CH_2)(AlMe_3)_2$ (I, $\delta = 28.77$ ppm)⁸ and (TiPTAC)Y $(Me_3AlCH_2AlMe_3)(\mu$ -MeAlMe₃) ($\delta = 34.2$ ppm).^{10c} An even higher upfield chemical shift was observed by the group of Diaconescu for the moiety $[ScAl_2(\mu_3-CH_2)]$ in $(NN^{Fc})Sc(CH_2)$ $(AlMe_3)_2$ ($\delta = -4.7$ ppm; NN^{Fc} = 1,1'-fc(NSitBuMe_2)_2), in accordance with a very long Sc-C(methylidene) distance of 2.6921(18) Å.³⁵ The latter chemical shift is very close to the ${}^{13}C{}^{1}H$ NMR shift of the methylidene carbon atoms in $[MeAl(CH_2)]_{12}$ ($\delta =$ -5.1 ppm).³²

The Sc-C(methylidene) distances in 11' (2.245(3)-2.3338(15) Å) are elongated compared to the monomeric structures of 8^{R} (2.1624(17)/2.1735(18) Å) and 10' (2.1724(15) Å) (Fig. 3). Consequently, the Sc-H(agostic) distances in 11' (2.24(4) Å/ 2.29(3) Å) are longer than those in 8^R (2.17(2) Å/2.14(3) Å), but the Sc-C(methylidene)-H(agostic) angles (79.6(17)°/76(2)°) are similar to those in 8^R (77.2(13)° for 8*, 75.1(16)° for 8'). The {Al₃(CH₂)₃} ring shows a chair-like conformation with methylidene protons in distinct axial or equatorial positions, whereas the $\{Sc_3(CH_2)_3\}$ ring is distorted due to the steric pressure of the Cp' ligands. Here, the methylidene protons don't assume the classical axial or equatorial positions. This may go handin-hand with a high mobility of the increasingly ionic methylidene moiety, resulting in only one signal for the two methylidene protons in the ¹H NMR spectrum (*vide supra*). A strategy for reducing the ratio of stabilizing aluminum moieties per rare-earth-metal center to 1:1 was described for the Sc/Al silycomplex (PNP)Sc(µ2-CHSiMe3)(µ2-Me)[Al(Me) lalkylidene (CH₂SiMe₃)] by the group of Mindiola.³⁶ There, the chemical shift of the methylidene carbon atom was noted as a broad signal at 137 ppm, corresponding to a very short Sc-CHSiMe₃ distance of 2.0845(14) Å. The heterobimetallic complex Cp*Ir $(\mu_2$ -CHSiMe₃) $(\mu_2$ -NtBu)Sc(Cp')(py)_x displayed similar spectroscopic and structural signatures (x = 0: ¹³C{¹H}, 114.4 ppm; x =1: Sc-C, 2.139(9) Å).³⁷ For further comparison, scandium complexes bearing dianionic [C(PPh₂S)₂] carbene pincer-type ligands revealed Sc-C distances as short as 2.200(3) Å,³⁸ and non-chelated phosphoniomethylidene complex (PNP)Sc(μ_2 - CPPh₃)(AlMe₃) feature the shortest Sc–C distance (2.036(5) Å) recorded to date.^{36,39}

Reactivity studies of methylidenes Cp^RSc(CH₂)(AlMe₃)₂ and [Cp'Sc(CH₂)₂AlMe]₃

Monomeric methylidenes 8^R eliminate methane at 80 °C, vielding unidentified products. Trimeric 11', in contrast, is stable at 100 °C. Compounds 8^R methylenate fluorenone instantly at ambient temperature, whereas the reaction of 11' with fluorenone takes 3 h at 80 °C until completion (cf., ESI[†]). Full conversion of methylidene complexes 8^{R} is indicated by ⁴⁵Sc NMR spectroscopy, revealing chemical shifts to higher field by ca. 50 ppm, likely ascribed to the symmetric oxo species $Cp^{R}Sc(O)(AlMe_{3})_{2}$ (cf., ESI[†]). Since the reactions were conducted with two equivalents of fluorenone, methylenation is competing with or followed by methylation.9c,12c Hence, comparatively low yields of dibenzofulvene were obtained (8*, 45%; 8', 46%; 11', 19%). It should be noted that DFT calculations on the methylenation of ketones by the trinuclear cluster $[PhC(NC_6H_3iPr_2-2,6)_2]_3Sc_3(\mu_3-CH_2)(\mu_3-Me)(\mu_2-Me)_3$ showed that methylene transfer rather than methyl transfer applies.40 Reactivities of the methylidenes toward multiple donors were tested as well. Compound 8* was treated with pyridine, leading to an immediate color change from colorless to vellow. After two hours at ambient temperature, a dark-brown solution had evolved. From this dark-brown solution, colorless crystals of AlMe₃(pyr) could be obtained at $-40 \, ^{\circ}\text{C}$ (cf., ESI[†]). Compound 11' does not react with pyridine, THF nor diethyl ether even at 100 °C. Compounds 8^R and 11' oligomerize δ-valerolactone instead of promoting methylenation (cf., ESI[†]). This suits the known reactivities of rare-earth-metal methylidenes in contrast to Tebbe's reagent.¹¹ A preliminary evaluation of the oligomerization/polymerization performance of complexes 8^{R} reveal an activity comparable to $[Cp'ScMe_{2}]_{2}$ (ref. 22) but significantly higher than Cp*₂YMe(thf) or commercially available yttrium isopropoxide (conditions: toluene as a solvent, ambient temperature, 1 h; Table S5[†]).

Conclusions

Cyclopentadienyl-supported discrete methylidenes $Cp^{R}Sc(CH_{2})$ (AlMe₃)₂ ($Cp^{R} = C_{5}Me_{5}$, $C_{5}Me_{4}SiMe_{3}$) are accessible directly by a tandem metathesis reaction/thermally induced methane elimination by treatment of $Cp^{R}ScCl_{2}(\mu-Cl)Li(thf)_{3}$ with the alkylating mix LiAlMe₄/AlMe₃. These reactions proceed *via* isolable intermediates as evidenced for $[Cp^{R}Sc(AlMe_{4})Cl]_{2}$ and $[Cp'Sc(AlMe_{4})Me]$. Here, *in situ* ⁴⁵Sc NMR spectroscopy provides a valuable tool to assess the progress of such one-pot reactions. The mixed methyl/tetramethylaluminato complex $[Cp'Sc(AlMe_{4})Me]$ reacts with GaMe₃ to yield the first example of a mixed Sc/Al/Ga methylidene, $Cp'Sc(CH_{2})(AlMe_{3})(GaMe_{3})$. Thermal treatment of $Cp'Sc(CH_{2})(AlMe_{3})(GaMe_{3})$ generates the new trimeric scandium aluminum bis(methylidene) cluster $[Cp'Sc(CH_{2})_{2}AlMe]_{3}$, under release of GaMe₃ and CH₄. The $\{Sc_{3}Al_{3}\}$ hexametallic cluster is closing the gap between methylidenes solely bound to rare-earth-metal centers of the $\{Sc_4\}$ cubane-type, $[Cp^*Sc(CH_2)]_4$ (ref. 13), and $\{ScAl_2\}$ -type methylidenes bound to only one rare-earth-metal center and two stabilizing Lewis acids, like $Cp^RSc(CH_2)(AlMe_3)_2$. X-ray diffraction and NMR analyses revealed enhanced rigidity of the thermally quite stable $\{Sc_3Al_3\}$ cluster in contrast to the fluxional methylidenes $Cp^RSc(CH_2)(AlMe_3)_2$. Distinct reactivity of the $\{ScAl_2\}$ and $\{Sc_3Al_3\}$ methylidenes is found toward donors and for ketone methylenation. The $\{ScAl_2\}$ -type methylidene converts fluorenone immediately at ambient temperature, whereas the $\{Sc_3Al_3\}$ cluster needs prolonged reaction time at elevated temperatures. Finally, taking the earlier reported orange oil, putative $[Cp'Sc(AlMe_4)_2]$, under close scrutiny led to the development of this new line of methylidenes with commercially available ligands Cp^* and Cp'.

Author contributions

GTLZ, synthesis and characterization of compounds, writing and editing original draft; SAZ, synthesis and characterization of compound **11**'; JR, synthesis and characterization of compound **8***; HS, crystallography of compound **8**'; CM-M, crystallography, editing original draft; RA, conceptualization, supervision, writing and editing original draft, project administration, funding acquisition.

Data availability

Experimental, spectroscopic and structural data supporting this article have been uploaded as part of the ESI.† Crystallographic data for all compounds have been deposited at the CCDC under 2410874–2410889,† and can be obtained from https://www.ccdc.cam.ac.uk/structures/.

Conflicts of interest

There are no conflicts to declare.

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