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Reversible Reductive Elimination in Aluminum(II) Dihydrides

Rosalyn L. Falconer, Gary S. Nichol, Ivan V. Smolyar, Scott L. Cockroft, and Michael J. Cowley*

Abstract: Oxidative addition and reductive elimination are defining reactions of transition-metal organometallic chemistry. In main-group chemistry, oxidative addition is now wellestablished but reductive elimination reactions are not yet general in the same way. Herein, we report dihydrodialanes supported by amidophosphine ligands. The ligand serves as a stereochemical reporter for reversible reductive elimination/ oxidative addition chemistry involving Al^I and Al^{III} intermediates.

Oxidative addition to aluminum(I) compounds is now well established, but examples of reductive elimination from Al^{III} to Al^I—and strategies to engineer it—are much scarcer.

Al^I compounds can be broadly divided into neutral and anionic classes. Of the neutral compounds, oxidative addition of E-H and E-X bonds has been reported for (Cp*Al)₄ (I, Figure 1),^[1] bulkier derivatives,^[2] and for NacNacAl(I) (II).^[3] More recently, several anionic Al^I compounds have been reported. Starting with the dimeric precursor to **IV**^[4] several Al^I compounds supported by diamido ligands have been prepared.^[5] Diverging from diamides, mixed amido/alkyl^[6] and dialkyl^[7] Al^I anions are now also known. Unsurprisingly, the anionic systems are even more reactive than the neutral compounds and also exhibit oxidative addition chemistry.

Reductive elimination chemistry at aluminum is much less developed. This is unsurprising considering the lower stability of the Al^I oxidation state compared to Al^{III}. Nevertheless, there are notable examples of reductive elimination from Al^{III}. Cp*₂AlH reversibly reductively eliminates Cp*H to form Cp*Al, I.^{[1b,8] Dipp}NacNacAlH₂ undergoes reversible oxidative addition to ^{Dipp}NacNacAl(I), **II**, though the product III has not been isolated $(^{Dipp}NacNac = (DippNCMe)_2CH).^{[9]}$ Most remarkably, the monomeric Al^I anion **IV** reversibly inserts into the C-C bond of benzene (by oxidative addition).^[10]

A notable characteristic of main-group systems is that base-coordination can induce reductive elimination. For

[*] Dr. R. L. Falconer, Dr. G. S. Nichol, I. V. Smolyar, Prof. S. L. Cockroft, Dr. M. I. Cowley

School of Chemistry, University of Edinburgh

Joseph Black Building, David Brewster Road, Edinburgh, EH9 3FJ (UK)

E-mail: michael.cowley@ed.ac.uk



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example, treating Si₂Cl₆ with Lewis bases induces reductive elimination, to form SiCl₄ and base-coordinated SiCl₂.^[11] Similar reactivity was recently reported for the dialane V, which disproportionates to AI^I and Al^{III} fragments upon treatment with Lewis bases.^[12]

We have reported Al^{III} dihydrides **1a** and **1b** supported by mixed-donor N/P ligands that exhibit flexible coordination behavior.^[13] Notably, the same ligand class was used by Kato to mediate reversible oxidative addition to Si^{II} compounds.^[14] We thus wondered whether the amidophosphine ligands of 1a and **1b** might prime Al^{II} systems for reductive elimination.

The study of dialanes-Al^{II} compounds-predates most work on Al^I. Dialanes are prone to disproportionation to Al⁰ and Al^{III}, but this can be avoided using bulky substituents.^[15]

Despite the useful and extensive reactivity of aluminum-(III) hydrides,^[16] there are scant examples of Al^{II} hydrides.



Figure 1. Selected neutral and anionic aluminum(I) and aluminum(II) compounds exhibiting (reversible) reductive elimination/oxidative addition behaviour. Dipp = 2,6-diisopropylphenyl. (2,2,2)cryptand = 4,7,13,16,21,24-hexaoxa-1,10-diazabicyclo[8.8.8]hexacosane.

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Jones has reported the NHC-coordinated parent dialane Al_2H_4 as well as amidinato- and guanadinato-coordinated dihydrodialanes.^[17] The reactivity of these compounds has not been explored.

Here, we report the reversible reductive elimination reactivity of dihydrodialanes supported by amidophosphine ligands. The complex stereochemistry of the N/P ligand enables the detection of "hidden" reductive elimination processes by revealing the interconversion of multiple diastereomers by reversible reductive elimination.

We initially prepared the dihydrodialanes **3a** and **3b** by reduction of Al^{III} dihydrides **1a** and **1b** (Scheme 1).^[13] Jones *et al.* used Mg^I reducing agents to access hydride-substituted dialanes.^[17] Accordingly, treatment of **1a** or **1b** with [^{Mes}NacNacMg]₂^[18] forms the Al^{II} dihydrido dialanes **3a** and **3b**, alongside the expected Mg^{II} hydride. The formation of **3a** and **3b** was slow (2–4 days at 70 °C) and limited quantities of starting material were converted (**1a**: 22%, **1b**: 12%). In practice, the dialanes **3a** and **3b** are accessed in better yields from reduction of the OTf substituted Al^{III} hydrides **2a** and **2b**, which leads to good conversion to **3a** and **3b** (**3a** 43%, **3b** 78%) and acceptable isolated yields (**3a** 20%, **3b** 43%).

Reducing instead Al^{III} dihydrides with decreased steric bulk at the nitrogen center—**4a** and **4b**—further improves the reaction. The reduction to the dialanes **5a** and **5b** proceeds smoothly with quantitative conversion and in good yields (**5a** 81%; **5b** 37%).

The Al^{II} dihydrodialanes **3** and **5** are structurally very similar. X-ray crystallography reveals the expected fourcoordinate tetrahedral geometry at the aluminum centers, which are κ^2 -coordinated by the amidophosphine ligands. Although the parent dialane Al₂H₄ is predicted to favor hydride-bridged structures,^[19] base coordination induces H₂Al-AlH₂ connectivity and terminal hydrides.^[17] Thus,





Scheme 1. Reductions of Al^{III} hydrides **1**, **2** and **4** to form Al^{III} dihydrodialanes **3** and **5**. Dipp=2,6-diisopropylphenyl. Mes=2,4,6-trimethylphenyl.

crystallography reveals terminal hydride substituents; this is consistent with the observed infrared terminal Al-H stretches which, as would be expected, are at lower wavenumber than the dihydride precursors (e.g. **5**: 1688–1725 cm⁻¹; **4**: 1795– 1802 cm⁻¹).^[17,19] In all of the structures, stereocenters at aluminum and in the ligand lead to crystallographic disorder in the norbornene backbone due to co-crystallization of multiple diastereomers (see later). We also note that in solving the structure of **5a** (Figure 2) we identified a cocrystallized minor component with a substantially shorter Al-Al distance than **5a**.^[20]



Figure 2. X-ray crystal structure of **5** a (ligand hydrogen atoms omitted). Thermal ellipsoids at 50% probability. Only the major component of the disordered ligand is shown (**5** a-**B**).^[24]

Unsurprisingly, increasing the steric bulk at the nitrogen or phosphorus centers increases N-Al and P-Al distances (Table 1). The Al–Al bond distances follow the same general trend, though the Al–Al bond distance in the *N*-mesityl dialane **5b** is longer than in its Dipp counterpart **3b** (2.886(2) vs. 2.8386(18) Å).

The Al-Al bond distances in **3** and **5** are remarkably long. The shortest distance is found in **5a**, which at 2.6585(16) Å is close to reported amidinato aluminum(II) hydride dimers (2.57-2.67 Å).^[17] The Al-Al distances in **3b** and **5b** are much longer than any previously reported for 3- or 4-coordinate dialanes, including that of the dialane(4) [('Bu₃Si)₂Al]₂ (2.751 Å).^[15d] Since dialane(6) compounds on average have longer Al-Al distances than dialane(4)s (Figure S1), the long Al-Al distances in **3** and **5** are consistent with the large steric bulk of the ligands.

To understand why the Dipp-substituted **3b** has a shorter Al–Al bond than its Mes analogue **5b**, we used an energy decomposition analysis. Multiple density functional theory

Table 1: Selected bond distances [Å] and angles [°] from the X-ray structures of 3a, 3b, 5a and 5b.

	Al-Al′	N-Al	P-Al	N-Al-P
3a N(Dipp)/P ^t Bu ₂	2.7323(19)	1.927(2)	2.4977(11)	85.12(7)
3b N(Dipp)/PN ₂ Si	2.8386(18)	1.939(2)	2.4982(10)	85.06(7)
5 a N(Mes)/P ^t Bu ₂	2.6586(16)	1.918(2)	2.4677(9)	85.86(6)
5 b N(Mes)/PN ₂ Si	2.886(2)	1.911(3)	2.4671 (11)	86.51 (8)

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methods, both with and without dispersion correction, underestimate the Al–Al bond in **5b** by ≈ 0.3 Å (e.g. M062X-GD3/ def2SVPP vs. X-Ray: 2.581 vs. 2.886(2) Å, Table S6). We ascribe the difference between predicted (gas phase) and experimental Al-Al distances to intermolecular dispersion interactions in the crystal structures. Indeed, a relaxed potential energy scan of the Al-Al distance in 5b (M062-X-GD3/def2SVP) reveals a minimal cost of 15 kJ mol⁻¹ to increase the distance from the DFT-optimized value to the experimentally observed one. Consistent with this, SAPT(2)/ 6-311G* computations on models of 3b and 5b extracted from M062X/Def2SVPP optimized geometries (Figure S3) revealed stabilizing intramolecular interactions of near identical magnitude. Although 3b obtains greater stabilization from dispersion, this is offset by larger exchange interactions (Figure S3) leading to near identical stabilization overall (3b, **5b**: -34.4, -32.7 kJ mol⁻¹). Thus, we believe the differing solid-state Al-Al bond distances in 3b/5b arise from intermolecular dispersion in the crystalline material.

³¹P{¹H} NMR spectroscopy of isolated **3** and **5** reveals they exist as mixtures of stereoisomers. Taking the NMes/PtBu₂ coordinated dialane **5a** as an illustrative example,^[21] crystal-

^tBu

^tBu

tBu

5a-D

line samples dissolved in C₆D₆ give rise to a spectrum (Figure 3 a.iii) containing three resonances; two singlets at δ 17.8 and 16.7 and a pair of mutually coupled doublets at δ 17.5 and 16.9 (${}^{3}J_{PP} = 17.8 \text{ Hz}$). These resonances are ca. 10 ppm downfield from Al^{III} dihydride starting material 4a. In the ¹H NMR spectrum, a broad resonance is observed for the Al-H groups at δ 5.1 (shifted downfield relative to Al^{III} dihydride **4a**: δ 4.6). Although the ¹H spectrum is complex, certain distinctive resonances (e.g. mesityl-CH₃ groups and norbornene CH₂ groups) again reveal the presence of three stereoisomers (see SI).

The spectroscopic situation is even more tortuous^[22] when the reduction at room temperature of **4a** by [^{Mes}NacNacMg]₂ is monitored by ³¹P{¹H} NMR spectroscopy. In addition to the three stereoisomers in the crystalline samples of dialane 5a, three other stereoisomers are also present, resulting in the observation of a total of 8 resonances (Figure 3 a.i). To understand these spectra, it is necessary for us to consider the possible stereoisomers of 5a.

The dialanes 3 and 5 have 16 possible stereoisomers (although they contain 6 stereocenters, those on the norbornenyl backbone can be treated as a single center for our





Figure 3. a) Diastereomers A-F of the dihydrodialane 5a with relative energies by DFT (M062X/Def2SVPP); ³¹P{¹H} NMR spectra for 5a i) reaction mixture showing mixture of 5 a diastereomers A-F produced after reduction of 4 a at room temperature, ii) reaction mixture after reduction at 70 °C, iii) isolated crystalline 5a in C₆D₆ solution. b) Proposed mechanism for interconversion of diastereomers by reversible reductive elimination and the Al^{I} compound **7***a*.

ťΒú tBu

7a

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^tBú

4a

. Bu

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tRu

5a-C

Mes

purposes), which on inspection reduce to the six diastereomers A–F (Figure 3a) (see SI). In diastereomers A–C the phosphine ligands are located *anti* across the Al-Al bond, the variation between A–C arising from the relative orientation of the norbornene CH₂ bridges (e.g. "*cis*" or "*trans*"). A–C are the diastereomers present in crystallographically characterized 3 and 5, and those observed spectroscopically in solutions of isolated 5a. Diastereomers A and B are *meso* compounds, due to the inversion center between the two Al atoms, rendering the phosphorus centers equivalent. We assign the singlets observed in the ³¹P{¹H} NMR spectrum at δ 17.8 and 16.7 to these isomers. Diastereomer C has inequivalent phosphorus centers which we therefore assign to the doublets at δ 17.5 and 16.9.

Diastereomers **D**–**F** have the phosphorus ligands located *syn* about the Al–Al bond. These diastereomers are observed in solution alongside **A**–**C** when the room-temperature reduction of the dihydride precursors is monitored by NMR spectroscopy, and the three isomers should result in four ³¹P NMR signals. The first two isomers **D** and **E** have equivalent phosphorus centers; **F** has two inequivalent phosphorus centers are found as singlets at δ 18.1, 17.7, 17.6 and 16.9; no doublets are observed for **F**, which we attribute to a decrease in the magnitude of ³J_{P-P} as the P-Al-Al-P torsion angle is reduced from \approx 180° in **C** to \approx 90° in **F**.

Consistent with the observation of all six diastereomers of **5a** in solution, and the experimentally observed order of stability, DFT calculations (M062X/Def2SVPP) reveal that diastereomers **A**–**F** fall within 23 kJ mol⁻¹ of each other (Figure 3a). Notably, the isomer with the lowest calculated energy, **5a**-**C**, is also found as the major species in solutions of crystalline **5a** ($\approx 60\%$ **5a**-**C**, remainder **5a**-**A** and **5a**-**B**). Isomers **D**–**F** with *syn* phosphorus centers are higher in energy than **A**–**C**.

We carried out an energy decomposition analysis to deconvolute ligand/ligand interactions and reveal the origins of the preference for isomers A-C. SAPT(2)/6-311G* computations were used to decompose the interactions between ligands in models of the 5a-C/5a-F structures (geometries extracted from M062X/Def2SVPP optimized structures, see SI). Substantial stabilizing non-covalent interactions were found to occur between the ligand N and P substituents (Figure S4). Whilst attractive dispersion interactions between the syn NMes groups in 5a-F contribute substantial stabilization ($-20.9 \text{ of } -50.0 \text{ kJ mol}^{-1}$ total), this is opposed by a larger exchange repulsion component in this configuration ($+59.9 \text{ kJ mol}^{-1}$). Indeed, despite the *anti* **5a-C** isomer that places NMes groups close to P^tBu₂ being less stabilized by dispersion $(-34.4 \text{ kJ mol}^{-1})$, it is overall more stable than 5a-F due to weaker exchange repulsion $(+36.2 \text{ kJ mol}^{-1})$. This general picture also holds for **3a** although the bulkier NDipp substituents result in overall lower stabilization from non-covalent interactions (Figure S6).

The six stereoisomers of 5a interconvert in solution. Mixtures of 5aA-F, generated by reduction at room temperature, equilibrate over time. The process requires weeks at room temperature or 24 hours at 80°C. It results in consumption of isomers **D**–**F** and production of thermodynamically favored isomers **A**–**C**. Lower quantities of **D**–**F** are also observed when reduction of **4a** is carried out at 70 °C (Figure 3 a.ii).

The interconversion of diastereomers **A**–**F** requires inversion of stereochemistry at the aluminum centers. We propose that this steroisomerisation proceeds by reductive elimination of an Al–H bond from dihydrodialanes (**3** or **5**) to generate the Al^I intermediates (**6** or **7**) and the Al^{III} dihydrides (**1** or **4**) with destruction of the Al stereocentres (Figures 3b and S8). Subsequent oxidative addition of an Al–H bond of the Al^{III} dihydrides **1** or **4** by Al^I intermediate **6** (NDipp) or **7** (NMes) reforms the dihydrodialanes **3** or **5**, thus regenerating two Albased stereocenters. Over time these processes bring mixtures of **A**–**F** to thermodynamic equilibrium.

Besides reversible reductive elimination, other mechanisms for inversion of stereochemistry at aluminum are conceivable. We can discount a mechanism involving inversion of stereochemistry at aluminum by reversible phosphine dissociation from one Al center on the basis that this pathway is unfavorable. We have already reported that the energetic cost for phosphine dissociation for a κ^2 -(amidophosphine)AlH₂ compound related to **1a/b** and **4a/b** is 83 kcal mol^{-1.[13]} The Al-Al bond in dialanes can dissociate photolytically, generating monomeric Al^{II} radicals,^[7] which could also explain interconversion of isomers A-F. We have no evidence for such a process in 3 or 5. In the case of the 5a, isomers A-F equilibrate no faster when exposed to light than they do in the dark. Also, treatment with *n*Bu₃SnH does not generate the Al^{III} dihydrides 1 or 4 (as could be expected by H-atom abstraction from nBu_3SnH by an Al^{II} radical).

Nikonov *et al.* reported the reductive elimination of Al–H bonds from **III**.^[9] In this case, the equilibrium constant for reductive elimination is large enough to detect the Al¹ and Al¹¹ compounds **II** and **III** alongside the Al¹¹¹ dihydride. In contrast, from **3** or **5** we did not observe Al¹¹¹ or Al¹ species in equilibrium, indicating the equilibrium constant for their formation from **3** or **5** is low. Nevertheless, there is further experimental evidence for reversible Al-H reductive elimination from dihydrodialanes **3** and **5**.

When a mixture of the P'Bu₂ Al^{II} dihydrodialane 5a and the $P(N^tBu)_2SiMe_2 Al^{III}$ dihydride **1b** in C₆D₆ was heated to 65°C, formation of the "crossover" dialane 8 was observed (Scheme 2). Formation of this compound proceeds by reductive elimination of 4a from 5a, generating the Al^I intermediate 7a (not observed) which then reacts with 1b to generate the mixed dialane 8. Both the dialane 8 (75.1-74.1 ppm) and the Al^{III} dihydride by-product **4a** (8.7 ppm) are apparent in the ³¹P{¹H} NMR spectrum. After 20 hours, the reaction generates an equilibrium mixture of the dialanes 5a and 8 together with the Al^{III} hydrides 1b and 4a, which precluded isolation of 8. A quantity of the symmetrical P(N^tBu)₂SiMe₂ dihydrodialane 3b might also be expected, but was not detected, indicating 5a and 8 are more stable. Similar "crossover" behavior was observed in the reactions of the dialane 5a with the hydride triflates 2a and 2b (SI, p40).

We also note here the related observation that reduction of the aluminum(III) hydride triflates 2 leads to formation of, besides dihydrodialanes 3a and 3b, the Al^{III} dihydrides 1a or



Scheme 2. "Crossover" experiment showing reaction between 5a and 1b to generate the "mixed" dihydrodialane 8 by reductive elimination from 5a and oxidative addition to the proposed intermediate 7a.

1b. The formation of the Al^{III} dihydride by-products is explained by reductive elimination of Al–H bonds from the dialane product **3** under reaction conditions.

Very few reactions of dihydrodialanes or dialane(6) compounds have been reported. We thus investigated some rudimentary reactivity of the dihydrodialane **5a**. Reported dialane(4) compounds are oxidized by iodine.^[15d] In contrast, when **5a** is treated with iodine the Al–H bonds are selectively iodinated in preference to the Al–Al bond, and diiododialane **9** is formed instead (Scheme 3). Crystallization of **9** from toluene solution allowed its solid-state structure to be determined, revealing that replacement of the Al–H bond distance or other structural features compared to **5a** (Al-Al **5a**: 2.6586(16) Å; **9**: 2.664(3) Å).

In summary, we have prepared a series of N,P-coordinated dihydrodialanes. Increasing the steric bulk of the N or P substituents lengthens the Al-Al bond distance. The stereocenters of Al^{II} compounds **3** and **5** reveal the interconversion of diastereomers by reversible reductive elimination. This suggests that stereoactive ligands could be useful tools for probing "hidden" reversible reductive elimination processes elsewhere in main-group chemistry, similar to the use of chiral silanes as mechanistic probes in B(C₆F₅)₃ chemistry.^[23] We are now investigating whether greater control of reductive elimination is possible by varying the nature of the donor ligand (phosphine in **3** and **5**).

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Scheme 3. Reaction of 5 a with iodine to form Al^{II} iodide dimer 9.

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Conflict of interest

The authors declare no conflict of interest.

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Communications



Communications



R. L. Falconer, G. S. Nichol, I. V. Smolyar, S. L. Cockroft, M. J. Cowley* _____ ####-#####

Reversible Reductive Elimination in Aluminum(II) Dihydrides



Low-oxidation-state main-group compounds exhibit rich oxidative addition chemistry. The same is not true for the reverse process, reductive elimination. Put together, the processes enable numerous catalytic cycles in transition-

metal chemistry. Here, using a stereoactive ligand as a reporter, it is revealed that AI^{II} dihydrodialanes exhibit transition-metal-like reversible reductive elimination. Mes = 2,4,6-trimethylphenyl

^tBu /^tBu

Mes

Mes

^tBu

AI(II)

^tBu

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