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# Theoretical and experimental studies on stability, bonding and isolation of elusive bis-(dichloro-aluminium) oxides supported by donor-base ligands†

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Herein, we depict the stability, and bonding studies of bis-(dichloro-aluminium) oxides  $Cl_2Al-O-AlCl_2$  supported by a pair of homo-/hetero-leptic donor base ligands L, L' [L, L' = cyclic alkyl(amino) carbene (cAAC<sup>Me</sup>; 1); N-heterocyclic carbene (NHC<sup>Me</sup>; 2); di-amido carbene (DAC<sup>Me</sup>; 3); L = cAAC<sup>Me</sup>, L' = NHC<sup>Me</sup>; (4)] with a general formula (L)Al(Cl)<sub>2</sub>-O-Al(Cl)<sub>2</sub>(L') (1-4) by NBO, QTAIM and EDA-NOCV analyses. Theoretical calculations suggest that 1-4 possess favorable interaction energies ( $\Delta E_{int}$ ), and bond dissociation energies between L/L', and Al(Cl)<sub>2</sub>-O-Al(Cl)<sub>2</sub> fragments *via* the formation of two dative L  $\rightarrow$  Al bonds [1 > 4 > 2 > 3]. This trend is rationalized by the  $\sigma$ -donor ability of the ligands L/L'. Moreover, we depict the first successful solid-state isolation of the colorless compound (cAAC)Al(Cl)<sub>2</sub>-O-Al(Cl)<sub>2</sub>(cAAC) (1') by reacting cAAC and AlCl<sub>3</sub> in the presence of a controlled amount of H<sub>2</sub>O, where two equiv. of cAAC is being utilized as the base. 1' has been structurally characterized by single-crystal X-ray diffraction, and further studied by NMR spectroscopy.

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### Introduction

The elusive oxo-bridged di-chloro aluminium compounds have been rarely documented in the literature. The low coordinate oxo-bridged Al compounds, known as the aluminoxanes have attracted significant interest among chemists due to their utility as active catalysts in the polymerization processes of epoxides, 1,2 aldehydes,3,4 and olefins.5 The discovery of methyl-aluminoxane (MAO) [(Me)AlO]<sub>n</sub> in 1980 has been marked as the pivotal development in this field introducing a highly active co-catalyst that effectively promotes the polymerization of ethylene, and propylene. 6,7 Aluminoxanes with general formula  $[(R)AIO]_n$  or  $[R_2AlOAlR_2]_n$ , can be synthesized through the controlled hydrolysis of organo-aluminium compounds. This process can be initiated using water in the form of the moisture present in hydrated metal salts8 or by reacting those compounds with oxygen-containing entities.9 In recent years, substantial advancements in the structural characterization of aluminoxanes have been documented, despite the inherent complexity of their synthetic routes.<sup>8,9</sup> In 2003, Roesky and co-workers

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unveiled the isolation of the pioneering bimetallic aluminoxane, featuring the terminal hydroxide groups (Fig. 1, **A**) *via* hydrolysis of an aluminium compound [LAl(I)<sub>2</sub>] (L = HC {MeCNDipp}<sub>2</sub>, Dipp = 2,6- ${}^{i}$ Pr<sub>2</sub>C<sub>6</sub>H<sub>3</sub>) within a biphasic liquid ammonia/toluene system, leading to the isolation of a previously unknown molecule [{LAl(OH)}<sub>2</sub>( $\mu$ -O)]. In 2005, the same group highlighted the reaction between an Al(I) monomer LAl (L

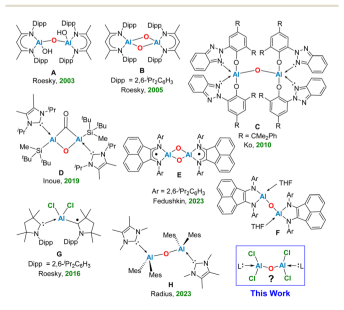


Fig. 1 Representative oxo-/di-chloro oxo-aluminium compounds isolated in the laboratory.

<sup>†</sup> Electronic supplementary information (ESI) available: Computational details, synthesis and single-crystal X-ray diffraction data of 1', NMR spectra, elemental analysis (PDF). CCDC 2374368 (1') and 2376845 ([cAACH]AlCl<sub>4</sub>). For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d5ra00211g

Paper

=  $HC[(CMe)(NAr)]_2$ ,  $Ar = 2,6^{-i}Pr_2C_6H_3$ ), and molecular oxygen, leading to the stabilization of a neutral bis-aluminium oxide  $[LAlO]_2$  (Fig. 1, B).<sup>11</sup>

In 2010, Ko and colleagues discovered the aluminium-methyl derivative with bulky ligands [(CMe<sub>2</sub>PhBTP)<sub>2</sub>AlMe] (CMe<sub>2</sub>PhBTP = 2-(2H-benzotriazol-2-yl)-4,6-bis(1-methyl-1-phenylethyl)-phenolate), which readily reacted with water to produce the dimeric oxo-aluminium species C (Fig. 1).12 In a ground-breaking study, Inoue and team successfully synthesized the first known di-alumene species ( $[((NHC)(Si)Al)_2]$  Si =  $Si^tBu_2Me$ ) stabilized by the donor base ligand NHC (N-heterocyclic carbene) with a formal Al=Al double bond. The reaction of a toluene solution of this dialumene under an atmosphere of CO₂ at −78 °C to 50 °C by a gradual heating, produced the novel oxo-di-aluminacarbonyl compound **D**, showcasing an unusual central Al(μ-CO)(μ-O)Al bridged motif (Fig. 1).13 Very recently, Fedushkin and colleagues have elucidated a fascinating reaction of a dialane [(dpp-bian)Al-Al(dpp-bian)] as a potential four-electron reducing agent with N<sub>2</sub>O in toluene at room temperature, leading to the formation of the corresponding oxides  $[(dpp-bian)Al(\mu_2-O)_2Al(dpp-bian)]$  (E), and  $[(dpp-bian)Al(THF)-(\mu-O)Al(THF)(dpp-bian)]$  (F) (dpp-bian)1,2-bis[(2,6-diisopropylphenyl)imino]-acenaphthene) (Fig. 1).<sup>14</sup> In 2016, Roesky and co-workers reported a neutral radical (Me2cAAC)<sub>2</sub>AlCl<sub>2</sub> (G, Fig. 1) stabilized by cyclic (alkyl)(amino)carbenes (cAACs), synthesized by reduction of the Me<sub>2</sub>-cAAC→AlCl<sub>3</sub> adduct with KC<sub>8</sub> in the presence of another equiv. of Me<sub>2</sub>-cAAC.<sup>15</sup> The EDA-NOCV analysis performed on G suggested an interaction between the neutral AlCl<sub>2</sub> radical and (cAAC)<sub>2</sub> in a singlet electronic state, and (cAAC)2 in a triplet electronic state interacting with AlCl<sub>2</sub>, featuring the equal probability electron sharing and coordinate bond due to negligible energy difference between  $\Delta E_{\rm orb}$  values of the two bonding possibilities. The mono-radical  $(Me_2\text{-cAAC})_2AlCl_2$  (G) is a  $\pi$ -type radical with the electron densities on  $\pi_{C=N}^*$  orbital of cAAC ligand, similar to the diradical (Me<sub>2</sub>cAAC)2SiCl2.15b Radius and colleagues synthesized dialane compounds stabilized by NHC and examined their reactivity with chalcogenides and oxygen. Using pyridine N-oxide as an oxygen donor, they isolated the compound ({(IMe<sup>Me</sup>)·AlMes<sub>2</sub>}<sub>2</sub>-μ-O) (Mes = 2,4,6-trimethylphenyl) (H, Fig. 1), exhibiting a central linear Al-O-Al bridge positioned on a crystallographic inversion center.16 However, no reports have been documented on the computational and experimental studies on the stabilization of the elusive neutral bis-(dichloro-aluminium) oxides [Cl2Al-O-AlCl<sub>2</sub>]. In this report, we shed light on the stability and bonding of five neutral monomeric bis-(dichloro-aluminium) oxides, supported by a pair of donor-base ligands with general formula  $(L)Al(Cl)_2OAl(Cl)_2(L')$   $(L, L' = cAAC^{Me}, 1; L, L' = NHC^{Me}, 2; L' = NHC$ diamidocarbene (DAC<sup>Me</sup>, 3),  $L = cAAC^{Me}$ ,  $L' = NHC^{Me}$ , 4), and the first successful laboratory isolation of the cAAC-stabilized bis-(dichloro-aluminium) oxide (1') by the activation of water molecule in presence of cAAC and AlCl<sub>3</sub>.

# Computational methods

The geometry optimizations, and frequency calculations for the hypothesized compounds (1–4) have been performed at BP86-D3(BJ)/def2-TZVPP<sup>17a-c</sup> level in both singlet and triplet electronic

states, employing Grimme D3 with Becke-Johnson damping dispersion correction. The potential energy surface (PES) was found to have a minimum as there was no imaginary frequency present. The Gaussian 09 program package was utilized to perform all the density functional theory (DFT) calculations.18 The natural bond orbital (NBO)19 analysis has been performed using NBO 6.0 program to evaluate the partial charges, Wiberg bond indices (WBI),20 and the natural bond orbitals. All the predicted compounds (1-4) have been found to possess singlet ground states. To elucidate the bonding characteristics within the (L)Al(Cl)2-O-Al(Cl)2(L') species, Energy Decomposition Analysis (EDA)21 integrated with the Natural Orbital for Chemical Valence (NOCV)22 approach was employed using the ADF 2020.105 software suite. The EDA-NOCV analysis, predicated on geometries optimized at the BP86-D3(BI)/def2-TZVPP level and conducted at the BP86-D3(BJ)/TZ2P level, facilitates the dissection of the intrinsic interaction energy  $(\Delta E_{\rm int})$  between two molecular fragments into four constituent energy components (see ESI†).

### Results and discussion

We initiated our studies with optimizations of the predicted molecules in both singlet and the triplet states. The optimized geometries of the hypothetical neutral monomeric bis-(dichloroaluminium) oxides, (L)Al(Cl)<sub>2</sub>–O–Al(Cl)<sub>2</sub>(L'), where, L, L' = cAAC<sup>Me</sup> (1); L, L' = cAAC<sup>Dipp</sup> (1'); L, L' = NHC<sup>Me</sup> (2); L, L' = NHC<sup>Dipp</sup> (2'); L, L' = DAC<sup>Me</sup> (3) and L = cAAC<sup>Me</sup>, L' = NHC<sup>Me</sup> (4) in their singlet states are represented in Fig. 2.

The computed  $C_L$ -Al distances for these compounds are found to be 2.047, 2.031, 2.097 and 2.045 Å (Table 1), respectively, which are comparable to the previously reported carbene donor  $C \rightarrow Al$  bond lengths in  $({}^{i}Pr_2Im) \cdot AlH_3$  (2.0405(17) Å),<sup>23</sup>

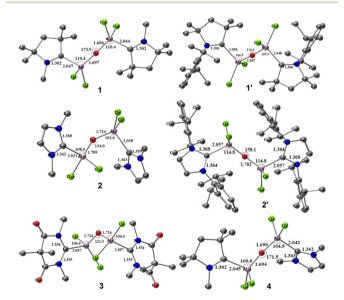


Fig. 2 Optimized geometries (at BP86/def2-TZVPP level of theory) of the species (L)Al(Cl)<sub>2</sub>-O-Al(Cl)<sub>2</sub>(L) (1-4) in the singlet ground states. L,  $L' = CAAC^{Me}$  (1); L,  $L' = NHC^{Me}$  (2), L,  $L' = DAC^{Me}$  (3) and  $L = CAAC^{Me}$ ;  $L' = NHC^{Me}$  (4).

Table 1 Selected bond lengths (Å) of compounds 1-4, calculated at BP86/def2-TZVPP level of theory

Compound	Bond	Bond length	Compound	Bond	Bond length
1	$\mathrm{C_{cAAC}} ext{-Al}$	2.047	<b>2</b> '	C <sub>NHC</sub> -Al	2.057
	Al-O	1.697		Al-O	1.702
1′	$\mathrm{C_{cAAC}} ext{-Al}$	2.050	3	$C_{DAC}$ -Al	2.097
	Al-O	1.697		Al-O	1.724
2	$C_{ m NHC}$ –Al	2.031, 2.038	4	$C_{cAAC}$ – $C_{NHC}$	2.045
					2.042
	Al-O	1.709, 1.724		Al-O	1.694
		•			1.699

(cAAC)<sub>2</sub>Al(Cl)<sub>2</sub> (2.097(2) Å), cAAC: →AlCl<sub>3</sub> (2.037(1) Å),<sup>15</sup> and (NHC)<sub>2</sub>Al<sub>2</sub>H<sub>4</sub> (2.0860(13) Å).<sup>24</sup> However, they are found to be slightly longer than the covalent Al–C single bond lengths observed in LHAlMe<sub>2</sub>Cl (1.950(3) and 1.980(2); L = HC[C(Me) N(Ar)]<sub>2</sub>, Ar = 2,6-<sup>i</sup>Pr<sub>2</sub>C<sub>6</sub>H<sub>3</sub>),<sup>25</sup> and Al<sub>2</sub>[CH(SiMe<sub>3</sub>)<sub>2</sub>]<sub>4</sub> (1.982(3)–1.985(3) Å).<sup>26</sup> In the species 1–4, the Al–O distances remain relatively consistent, hovering around 1.7 Å. In compounds 1 (L, L' = cAAC<sup>Me</sup>) and 4 (L = cAAC<sup>Me</sup>; L' = NHC<sup>Me</sup>), the central Al(Cl)<sub>2</sub>–O–Al(Cl)<sub>2</sub> moiety demonstrates a nearly linear configuration, with an Al–O–Al bond angle of 173.9° and 171.5°. In contrast, compounds 2 and 3 exhibit reduced bond angles of 134.0° and 121.5°, respectively. But when the Me groups of 2 were replaced by the Dipp (2,6-diisopropylphenyl) groups (2'), the Al–O–Al bond angle was increased to 158.1°, which can be attributed to the higher steric effect (see ESI†).

To have a detailed understanding of the distribution of electron densities in the hypothetical species 1-4, we have performed the Natural Bond Orbital (NBO) analyses (Table S9, ESI†). The respective Kohn-Sham orbitals show that the LUMOs for all three species are the  $\pi^*$  orbitals of the C=N bond of the carbenes, which play a crucial role in their reactivity and interaction with other molecules. The lower lying  $\pi_{C=N}^*$ orbital of cAAC, in general, is beneficial for the intra-molecular charge transfer, small molecule activation, enhancement of photoluminescence properties, hosting the extra electron densities from bonded element, and formation of the cAACcentred radical anion intermediate for reduction of cAAC containing compounds leading to the formation of low-valence cAAC-containing compounds. 15b,c For compounds 1, 2, and 4 the HOMO represents the lone pairs of electrons on the central bridging oxygen atom. We observed that the fully occupied frontier orbitals of these molecules do not only represent the  $\pi$ orbitals of the phenyl rings of the Dipp groups, rather, those exhibit through-space interaction of these  $\pi$  orbitals with the non-bonding electrons of the Cl and the O atoms (see ESI† for the respective Hirshfeld plots). Such secondary interactions indirectly play important role in overall stabilization of the respective molecules. For example, the HOMO-1 of 4 represents through space interaction of the lone pair of electrons on bridging O-atom with the  $\pi$  electron cloud of the aromatic ring of the Dipp group. The HOMO of 1' shows through-space interaction between the lone pair of electrons on Cl atoms with the central O atom and the  $\pi$  electron cloud of the aromatic rings. Similarly, the HOMO of 2' shows the through-space

interaction between the lone pair of electrons on Cl-atoms and the  $\pi$  electron cloud of the aromatic ring.

Compound 3 demonstrates σ-donation from the DAC moiety to Al. Moreover, for compounds 1, 2, and 4, the  $\sigma$ -donation from cAAC/NHC is represented in the HOMO-1, and HOMO-8 orbitals, respectively; suggesting variations in the electron donation capabilities among the compounds 1-4 (see ESI†). We could observe comparable results when the methyl groups in 1, and 2 are replaced by sterically bulkier Dipp groups (1'-2'). The NBO analyses performed on bulky analogues 1', and 2' at BP86/ def2-TZVPP level of theory showed the presence of a single occupancy bond between CcAAC/NHC-Al, which is majorly polarised towards C<sub>L</sub> atom (86–88.3%). The WBI for C-Al bond is approximately 0.4, which implies its single bond nature. The HOMO-3, and HOMO-10 of 1', and 2', respectively confirm  $\sigma$ donation from carbene to Al. The bond between Al-O is polarized exclusively towards the O atom due to its higher electronegativity. The LUMO of 1' and 2' corresponds to the  $\pi^*$  of C= N. HOMO of  $\mathbf{1}'$  and  $\mathbf{2}'$  lies on the  $\pi$  orbitals of the phenyl ring. HOMO of  $\mathbf{1}'$  also represents the lone pairs present on the oxygen atom (Fig. 3).

The single bond occupancy was observed along the C<sub>carbene</sub>-Al bond, which is significantly polarized towards the carbene moiety. This polarization is indicative of the unique bonding interactions within 1-4. The Wiberg Bond Indices (WBI) further elaborate on the nature of the C-N bonds within the compounds; for instance, compound 1 possesses a WBI of 1.60, implying a double bond characteristic. Contrastingly, in compound 2, the WBI drops to 1.29, suggesting a diminishing double bond nature, as visualized in the HOMO-4 orbital. The NBO results do not distinguish between a covalent dative bond or a covalent electron-sharing bond, and therefore NBO cannot accurately determine the true nature of the bond. In this regard, the energy decomposition analysis natural orbitals of chemical valence (EDA-NOCV)21,22 approach is the most suitable tool for gaining insight into the nature of the chemical bonds of (L)Al(Cl)2- $OAl(Cl)_2(L')[L, L' = cAAC^{Me}(1), L, L' = NHC^{Me}(2), L, L' = DAC^{Me}$ (3),  $L = cAAC^{Me}$ ,  $L' = NHC^{Me}$  (4)]. The magnitudes of  $|\Delta E_{orb}|$  from different bonding scenarios were compared, and the bonding scenario with the lowest value of  $\Delta E_{\rm orb}$  is considered to be the best (i.e., the most feasible) bonding scenario, since it will require the least change in the electronic structure of the fragments to maintain the electronic structure of the final molecule at its equilibrium geometry.15a In the present study, the best bonding description of 1-4 is illustrated considering different bonding

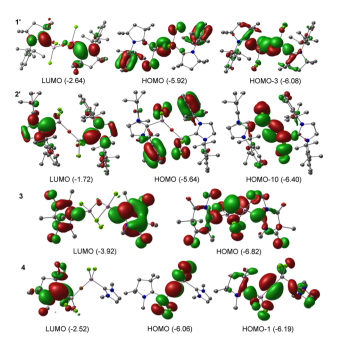
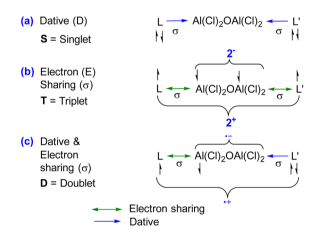


Fig. 3 Selected Kohn-Sham orbitals of compounds 1'-2', and 3 and 4 at BP86/def2-TZVPP level of theory. The values in the parentheses are the energies of the orbitals in eV.



Scheme 1 Possible bonding scenarios between interacting fragments [(L)(L')] and Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub> of 1-4.

possibilities (Scheme 1) by varying the charge, and electronic states of the interacting fragments [(L)(L')], and Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub>, specifically: (a) neutral [(L)(L')] and Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub> fragments in their electronic singlet state forming two dative bonds, (b) doubly charged [(L)(L')]<sup>2+</sup> and [Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub>]<sup>2-</sup> fragments in their electronic triplet state leading to the formation of two  $\sigma$  electron sharing bonds, and (c) singly charged  $[(L)(L')]^+$  and  $[Al(Cl)_2]^+$ OAl(Cl)<sub>2</sub>] fragments in electronic doublet state, which would interact to form an electron sharing and a dative bond. The EDA-NOCV results depicted from Table 2 show that for carbene-containing species 1-4, the fragmentation scheme involving neutral [(L)(L')] and  $[Al(Cl)_2OAl(Cl)_2]$  fragments in the electronic singlet state (Scheme 1(a)) forming dative bonds gives the smallest  $\Delta E_{\rm orb}$ and is therefore the best bonding scenario.

Table 2 The EDA-NOCV analyses of L-Al(Cl)2OAl(Cl)2-L' and LAI(CI)2-O-AI(CI)2L' bonds of LAI(CI)2-O-AI(CI)2L' compounds 1-4 using [ligands] and [Al(Cl)2OAl(Cl)2] in the electronic singlet (S) and  $[(LAl(Cl)_2)(L'Al(Cl)_2)]^{2+}$  and  $[O]^{2-}$  in the electronic singlet (S) states as interacting fragments at the BP86-D3(BJ)/TZ2P level of theory. Energies are in kcal mol<sup>-1</sup>

Compound	Energy	$L\text{-Al}(Cl)_2OAl(Cl)_2\text{-}L$	LAl(Cl) <sub>2</sub> -O-Al(Cl) <sub>2</sub> L
1	$\Delta E_{ m int}$	-157.9	<b>−770.45</b>
	$\Delta E_{\mathrm{Pauli}}$	231.9	405.59
	$\Delta E_{ m ele}$	-233.6 (59.9%)	-899.60 (76.5%)
	$\Delta E_{ m dis}$	-28.6 (7.3%)	-4.33 (0.4%)
	$\Delta E_{ m orb}$	-127.7 (32.8%)	-272.10(23.1%)
2	$\Delta E_{ m int}$	-154.0	-783.38
	$\Delta E_{\mathrm{Pauli}}$	225.6	405.22
	$\Delta E_{ m ele}$	-227.7 (60%)	-905.05 (76.1%)
	$\Delta E_{ m dis}$	-25.6 (6.7%)	-3.91 (0.3%)
	$\Delta E_{ m orb}$	-126.4 (33.3%)	-280.64(23.6%)
3	$\Delta E_{ m int}$	-136.6	-795.98
	$\Delta E_{\mathrm{Pauli}}$	221.3	441.01
	$\Delta E_{ m ele}$	-204.7 (57.2%)	-916.54 (74.1%)
	$\Delta E_{ m dis}$	-30.4 (8.5%)	-4.15 (0.3%)
	$\Delta E_{ m orb}$	-122.8 (34.3%)	-316.31 (25.6%)
4	$\Delta E_{ m int}$	-154.1	-774 <b>.</b> 1
-	$\Delta E_{\mathrm{Pauli}}$	225.3	408.0
	$\Delta E_{ m ele}$	-228.1 (60.1%)	-907.2 (76.7%)
	$\Delta E_{ m dis}$	-25.1 (6.6%)	-4.16 (0.4%)
	$\Delta E_{ m orb}$	-126.2 (33.3%)	-4.10 (0.470) -270.7 (22.9%)
	<b>⊥</b> Lorb	-120.2 (33.370)	-270.7 (22.970)

Table 2 encapsulates the results, revealing that in these compounds, the bonding between the aluminium (Al) and the carbene ligand (L/L') predominantly stems from electrostatic interactions, constituting approximately 57.2-60% of the bond character. The covalent interactions contribute significantly as well, accounting for 32.8-34.3% of the bonding nature, while dispersion forces contribute marginally between 8.5% and 6.6%, which is significant.

A deeper analysis involves dissecting the  $\Delta E_{\rm orb}$ , the energy associated with the orbital interactions, into pairwise contributions to understand the specifics of bonding. The study

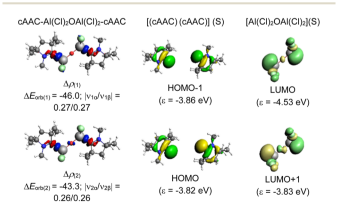


Fig. 4 The shape of the deformation densities  $\Delta \rho_{(1)-(2)}$  that correspond to  $\Delta E_{orb(1)-(2)}$ , and the associated MOs of cAAC-Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub>cAAC (1) and the fragments orbitals of [(cAAC)(cAAC)] and [Al(Cl)2-OAI(CI)<sub>2</sub>] in the singlet state (S) at the BP86-D3(BJ)/TZ2P level. Isosurface values are 0.003 au for  $\Delta \rho_{(1)-(2)}$ . The eigenvalues  $|\nu_n|$  give the size of the charge migration in e.

identifies two significant orbital contributions,  $\Delta E_{\text{orb}(1)}$  (with same phase; ++) and  $\Delta E_{orb(2)}$  (with opposite phase; +-) which mainly correspond to the  $\sigma$  donation from the ligand (L) to the

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aluminium centre (Al) (Fig. 4).

The dative in-phase (++)  $\sigma$  donation from HOMO-1 of the ligands [(L)(L)] into LUMO of the [Al(Cl)2OAl(Cl)2] forms the  $\Delta E_{\text{orb}(2)}$  in 2, and 3. The  $\Delta E_{\text{orb}(2)}$  of 4 features the  $\sigma$  donation from HOMO-1 of the ligands [(cAAC)(NHC)] into LUMO+1 of the [Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub>] in out-of-phase (+-) combination.

Roesky and co-workers studied the bonding of the neutral radical (Me2-cAAC)2AlCl2 (ref. 15) using EDA-NOCV studies, and predicted two bonding scenarios. The first scenario proposed an interaction between an AlCl<sub>2</sub> radical and (cAAC)<sub>2</sub> in a singlet electronic state, whereas the second scenario treated (cAAC)2 in a triplet electronic state interacting with AlCl<sub>2</sub>, leading to the formation of a shared electron bond and a coordination bond. The findings indicated that the energy difference for both scenarios ( $\Delta E_{\rm orb}$ ) was nearly identical, suggesting both are similarly probable. Further analysis of the fragments' orbitals revealed the primary contribution to  $\Delta E_{\text{orb}}$  in the original study was the transfer of the unpaired electron from aluminium to the cAAC ligands' LUMO, amounting to -123.9 kcal mol<sup>-1</sup>. However, in our analysis of compound 1, the primary contribution to  $\Delta E_{\rm orb}$ , totalling -89.3 kcal  ${\rm mol}^{-1}$  or 70% of  $\Delta E_{\rm orb}$ , stemmed from the electron donation from the cAAC2 ligand fragments to the Al(Cl)2OAl(Cl)2 center. Whereas, the stabilization arising from the coordinate bond ( $C_{cAAC} \rightarrow Al$ ) in (Me<sub>2</sub>cAAC)<sub>2</sub>AlCl<sub>2</sub> is only -59.6 kcal mol<sup>-1</sup>, i.e., 27.7% of the  $\Delta E_{\rm orb}$ . The bond dissociation energy for the (cAAC)2 and Al(Cl)2-OAl(Cl)<sub>2</sub> is 118.9 kcal mol<sup>-1</sup> is significantly higher compared to (Me<sub>2</sub>-cAAC)<sub>2</sub>AlCl<sub>2</sub> (-71.0 kcal mol<sup>-1</sup>). The computed interaction energies ( $\Delta E_{\rm int}$ ) and bond dissociation energies (BDEs) establish a distinct trend of the ligand strength in the order of 1 > 4 >2 > 3, which directly correlates with the  $\pi$ -acceptor and  $\sigma$ -donor properties of the respective carbenes. The BDE values follow a similar pattern with 1 showing 118.9 kcal mol<sup>-1</sup>, while 2 and 3 exhibit 113.9 kcal mol<sup>-1</sup> and 89.6 kcal mol<sup>-1</sup>, respectively. The heteroleptic system, 4 falls in between at 113.1 kcal mol<sup>-1</sup> for the L-(Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub>)-L' bonds. The stronger bonding interaction in 1 and 4 is attributed to the enhanced  $\pi$ -acceptor ability of cAAC, which enables greater orbital overlap with the electrondeficient aluminium centre, leading to better charge delocalization and bond stabilization. On the other hand, the DACstabilized compound 3 exhibits the weakest interaction, as reflected in both its lower BDE (89.6 kcal mol<sup>-1</sup>) and interaction energy (-136.6 kcal mol<sup>-1</sup>). This can be attributed to its diminished donor-acceptor interactions and reduced charge transfer efficiency, making it inherently less stable compared to cAAC and NHC. The WBI analysis also supports these findings, with the C-N bond in 1 exhibiting a WBI of 1.60, indicative of substantial double-bond character, while 2 shows a lower value of 1.29, signifying weaker  $\pi$ -conjugation.

To study the nature of the central Al-O bonds, we performed EDA-NOCV calculations, cleaving Al-O bonds in 1-4. Three bonding possibilities were generated by changing the charge and the multiplicity of the fragments. The first possibility considered the interaction of neutral triplet fragments leading

(a) Electron (E)  $L(CI)_2AI \longrightarrow O \longrightarrow AI(CI)_2L'$ Sharing (σ) **T** = Triplet (b) Dative and  $L(CI)_2AI \xrightarrow{+} O \xrightarrow{\uparrow} AI(CI)_2L'$ Electron Sharing (σ) **D** = Doublet  $L(CI)_{2}AI \xrightarrow{+} O \xrightarrow{} AI(CI)_{2}L'$ (c) Dative (D) S = Singlet

Scheme 2 Possible bonding scenarios of the Al-O bonds of 1-4.

to the formation of electron-sharing bonds (Scheme 2(a)). The second bonding possibility considered the interaction of a combination of dative and electron-sharing bonds from the interaction of singly charged doublet fragments (Scheme 2(b)). The third and last considers the interaction of doubly charged singlet fragments [(O<sup>2-</sup>) and ((L)Al(Cl<sub>2</sub>))(Al(Cl)<sub>2</sub>(L'))<sup>2+</sup>] forming dative bonds (O  $\rightarrow$  Al). It was found the  $\Delta E_{\rm orb}$  was found the least for the third possibility and hence considered the best bonding scenario (Scheme 2(c)).

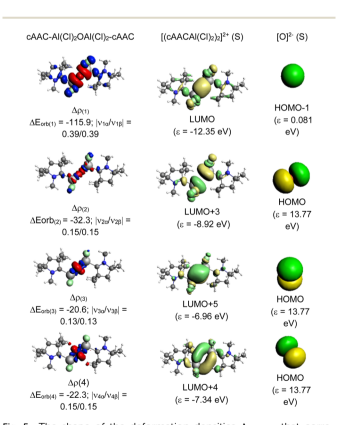


Fig. 5 The shape of the deformation densities  $\Delta 
ho_{(1)-(4)}$  that correspond to  $\Delta E_{\mathrm{orb}(1)-(4)}$ , and the associated MOs of (cAAC)Al(Cl)<sub>2</sub>-O- $Al(Cl)_2(cAAC)$  (1) and the fragments orbitals of  $[(cAACAl(Cl)_2)_2]^{2+}$  and [O]<sup>2-</sup> in the singlet state (S) at the BP86-D3(BJ)/TZ2P level. Isosurface values are 0.001 au for  $\Delta \rho_{(1)-(4)}$ . The eigenvalues  $|\nu_n|$  give the size of the charge migration in e. The direction of the charge flow of the deformation densities is red  $\rightarrow$  blue

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L: 
$$\longrightarrow$$
 AI  $\longrightarrow$  O  $\longrightarrow$  AI  $\longrightarrow$  CI  $=$  CAAC (1)  $=$  NHC (2)  $=$  DAC (3)  $=$  cAAC, L'  $=$  NHC (4)

Scheme 3 Predicted best bonding scenario between O and Al centres

The bonding between the Al and O atoms predominantly stems from electrostatic interactions, constituting approximately 74.1-76.7% of the bond character (Fig. 5, see ESI†). The covalent interactions contribute significantly, accounting for 22.9-25.6% of the bonding nature, while dispersion forces contribute marginally between 0.3 and 0.4%, which is significantly lower than that of C<sub>L</sub>-Al bonds. A deeper bonding analysis involves dissecting the  $\Delta E_{\rm orb}$ , the energy associated with the orbital interactions, into pairwise contributions to understand the specifics of covalent bonding. The study identifies four significant orbital contributions,  $\Delta E_{\mathrm{orb}(1)} - \Delta E_{\mathrm{orb}(4)}$ , which mainly correspond to the  $\sigma$  and  $\pi$  donation from the O-atom to the aluminium centre (Al).  $\Delta E_{\mathrm{orb}(1)}$  and  $\Delta E_{\mathrm{orb}(2)}$  correspond to the  $\sigma$ donation from  $\mathrm{O}^{2-}$  to Al centre in which the major contribution is from  $\Delta E_{\text{orb}(1)}$  (42.6–50.3%).

In 1 and 3 it is the dative out-phase (+-)  $\sigma$  donation from HOMO-1 of O into LUMO of the  $((L)Al(Cl_2))(Al(Cl)_2(L'))^{2+}$ whereas in 2 and 4 it is the dative in-phase (++)  $\sigma$  donation from HOMO-1 of O into LUMO of the  $[((L)Al(Cl_2))(Al(Cl_2)(L'))]^{2+}$ .  $\Delta E_{\text{orb}(3)}$  and  $\Delta E_{\text{orb}(4)}$ , which contribute minorly to the corresponds to the  $\Delta E_{\rm orb}$ , are  $\pi$  donation from O to Al fragments. The dative in-phase (++)  $\pi$  donation from O to Al forms the  $\Delta E_{\text{orb}(3)-(4)}$  in 1 and 4, whereas, dative out-phase (+-)  $\pi$  donation from O to Al form the  $\Delta E_{\text{orb}(3)-(4)}$  in 2 and 3 (Scheme 3).

# **Synthesis**

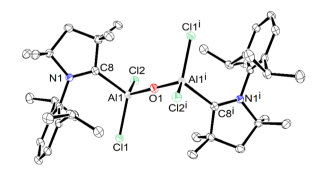
To a 1:1 molar mixture of Me<sub>2</sub>-cAAC: → LiOTf (Me<sub>2</sub> $cAAC:=:C(N-2,6^{-1}Pr_2C_6H_3)(CMe_2)_2(CH_2))$  and  $AlCl_3$ , toluene (10) mL) was added at -5 °C (ice/water mixture) to which 1 equiv. of water was added using a micro-syringe and the reaction mixture was slowly raised to room temperature (rt), The resulting reaction solution was stirred at rt for 12 h. Afterwards, the white precipitate was filtered, and the pale-yellow filtrate was concentrated under reduced pressure, and stored for crystallization at ambient temperature to obtain the colorless cubes of compound 1' after 1-2 days in 42% yield (Scheme 4).

Scheme 4 Synthesis of [(cAAC<sup>Dipp</sup>)Al(Cl<sub>2</sub>)-O-Al(Cl<sub>2</sub>)(cAAC<sup>Dipp</sup>)] (1').

Compound 1' was structurally characterized by single-crystal X-ray diffraction, and further studied by NMR spectroscopy, and elemental analyses. The white residue obtained was also characterized by the NMR spectroscopy, which revealed the formation of the salt [cAACH]AlCl<sub>4</sub> (see ESI†). The powder of 1' was found to be thermally stable up to 200 °C under an inert atmosphere, and then decomposed to a black liquid in the temperature range of 201-204 °C.

The crystals of 1' were found to be soluble in organic solvents, such as, toluene, THF, etc. The toluene solution of 1' was found to be stable for over one month under an argon atmosphere. The <sup>13</sup>C NMR spectrum of a deuterated benzene solution of 1' exhibited the C<sub>carbene</sub> peak at 206.6 ppm at 298 K, which is upfield shifted compared to that of the free carbene,27 and downfield shifted when compared to that of ({(IMe<sup>Me</sup>)· AlMes<sub>2</sub> $_{2}$ - $\mu$ -O)(172.4 ppm). The molecular structure of 1' is shown in Fig. 6.

Compound 1' crystallizes in the  $P\bar{1}$  triclinic space group, and features a linear (Cl)2AlOAl(Cl)2 units with an inversion centre passing through O1. The aluminium atoms in  $\mathbf{1}'$  display a distorted tetrahedral geometry and form bonds with two chlorine atoms, a cAAC ligand, and a µ-O unit. The Al-Cl bond lengths in 1' measure 2.1659(7) and 2.1494(8) Å, which are comparable to those observed in cAAC2AlCl2 (2.1612(7)-2.1638(7) Å), and  $cAAC: \rightarrow AlCl_3$  adduct (2.1315(5)-2.1439(5) Å). The Al-C<sub>cAAC</sub> bond length in 1' is found to be 2.063(2) Å, which lies inbetween to those observed in Me<sub>2</sub>-cAAC: →AlCl<sub>3</sub> adduct (2.037(1) Å), featuring the  $\sigma$ -donating  $C_{cAAC}$ :  $\rightarrow$  Al bond, and in (Me<sub>2</sub>-cAAC)<sub>2</sub>AlCl<sub>2</sub> (2.097(2) Å), featuring one shorter C<sub>cAAC</sub>:-Al electron-sharing covalent bond (1.967(2) Å) and a relatively longer  $C_{cAAC}$ :  $\rightarrow$  Al σ-donating bond (2.097(2) Å). <sup>15</sup> The Al- $C_{cAAC}$ bond in 1' seems to be longer as expected than those present in compounds stabilized by bulky non-carbene ligands, e.g.,  $LHAlMe_2Cl$  (1.967(2) Å) (L =  $HC[C(Me)N(Ar)]_2$ , Ar =  $2,6^{-1}Pr_2C_6H_3$ ) and  $Al_2[CH(SiMe_3)_2]_4$  (1.982(3)–1.985(3) Å), where the Al-C<sub>cAAC</sub> bonds are covalent single bonds.<sup>25,26</sup> The Al-O bond length observed in 1' is 1.6780(6) Å, which is significantly



Molecular structure of isolated (cAAC<sup>Dipp</sup>)Al(Cl<sub>2</sub>)-O-Al(Cl<sub>2</sub>)(cAAC<sup>Dipp</sup>) (1'). The anisotropic displacement parameters are depicted at the 50% probability level. Hydrogen atoms, and solvent (toluene) are omitted for clarity. Selected experimental [calculated at BP86/def2-TZVPP] bond lengths [Å] and bond angles [°]: C8-Al1 2.063(2) [2.050], Al1-Cl2 2.1659(7) [2.168], Al1-Cl1 2.1494(8) [2.175], Al1-O1 1.6780(6) [1.697], C8-N1 1.297(2) [1.306]; N1-C8-Al1 129.73(14) [128.5], Al1-O1-Al1<sup>i</sup> 180.00 [176.5].

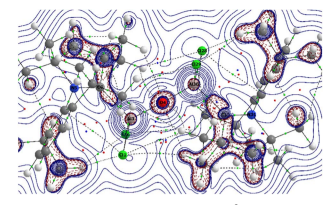


Fig. 7 Contour plot of Laplacian distribution  $[\nabla^2 \rho(r)]$  in the Al3–O4–Al30 plane of  $(cAAC^{Dipp})Al(Cl_2)-O-Al(Cl_2)(cAAC^{Dipp})$  (1'). Solid blue lines indicate the areas of charge concentration  $(\nabla^2 \rho(r) < 0)$  while dotted purple lines denotes charge depletion  $(\nabla^2 \rho(r) > 0)$ . Solid lines connecting atomic nuclei (black) are the bond paths and green spheres on bond path indicates the BCP.

shorter than that (1.7252(4) Å) present in ( $\{(IMe^{Me})\cdot AlMes_2\}_2-\mu$ -O) (H), possibility due to the bulky Mes group on Al.<sup>16</sup>

The QTAIM<sup>28</sup> analyses on 1' at the BP86/def2-TZVPP level of theory depicted the electron densities  $(\rho(r))$  of 0.068–0.109 and positive Laplacian  $(\nabla^2 \rho(r))$  at the bond critical point (BCP) of the C–Al and Al–O bonds in 1' suggested the closed-shell interactions as expected (Fig. 7).

### Conclusions

In summary, we have hypothesized the possible stabilization of the elusive neutral bis-(dichloro-aluminium) oxides in presence of donor-base ligands  $[(L)Al(Cl)_2OAl(Cl)_2(L)][(L, L' = donor base$ ligands; e.g., cAAC, NHC, and DAC); 1-4], and predicted their electron densities distribution and bonding scenarios by quantum chemical calculations. The NBO analysis revealed that the LUMOs comprise of the  $\pi^*$  orbitals of the C=N bond, a finding that portends significant repercussions for their reactivity and potential interaction abilities. The HOMOs in compounds 1, 2, and 4 are the lone pairs on the oxygen atoms.  $\sigma$ -Donation from  $C_{carbene}$  to the aluminium centre in 1, and 4 is represented by the HOMO-1, and in 2 by HOMO-8 orbital. Moreover, the analysis revealed a noteworthy polarization of the Ccarbene-Al bond towards the carbene, thereby introducing an additional layer of complexity to the bonding interactions. The EDA-NOCV analysis revealed that the bond between the ligands (L, L'), and the Al(Cl)<sub>2</sub>OAl(Cl)<sub>2</sub> core is mediated through a purely dative mechanism involving neutral singlet fragments. Intriguingly, the bond between the [((L)Al(Cl<sub>2</sub>))(Al(Cl)<sub>2</sub>(L'))] units and oxygen was identified as a dative bond, albeit with singlet doubly charged fragments, thus delineating a complex interaction framework within these molecular systems. The cAAC<sup>Dipp</sup> analogue (cAAC<sup>Dipp</sup>)Al(Cl<sub>2</sub>)-O-Al(Cl<sub>2</sub>)(cAAC<sup>Dipp</sup>) (1') has been successfully synthesized, isolated, and structurally characterized by single-crystal X-ray diffraction, and studied further by NMR spectroscopy.

# Data availability

Deposition numbers 2374368 (for  $\mathbf{1}'$ ), 2376845 (for [cAACH]AlCl<sub>4</sub>) contain the supplementary crystallographic data for this paper.

## Conflicts of interest

There are no conflicts to declare.

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