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# Donor-free 9-aluminafluorenes: molecular structures and reactivity†

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Aluminum-doped polycyclic aromatic hydrocarbons (PAHs) are underexplored despite the broad applications of boron-containing PAHs in areas such as catalysis and optoelectronics. We disclose the donor-free, sterically unprotected 9-methyl-9-aluminafluorene (Me-AlFlu; **2**), synthesized by heating a 9,9-dimethyl-9-stannafluorene and AlMe<sub>3</sub> in hexanes. The compound is a dimer, (**2**)<sub>2</sub>, with *trans*-positioned AlMe substituents in the solid state. In solution, (**2**)<sub>2</sub> shows a dynamic *cis/trans*-interconversion rather than a monomer-dimer equilibrium (Tol- $d_8$ , RT). Lewis bases L cleave (**2**)<sub>2</sub> into monomeric adducts **2**·L (L = OEt<sub>2</sub>, thf, pyridine). Lewis acidic AlBr<sub>3</sub> transforms (**2**)<sub>2</sub> into a 2,2'-(Br<sub>2</sub>Al)<sub>2</sub>-1,1'-biphenyl (**3**), crystallographically characterized as dimeric (**3**)<sub>2</sub>. (**3**)<sub>2</sub> is a synthetic equivalent for the elusive free Br-AlFlu: Treatment with donor molecules furnishes Br-AlFlu·L adducts (L = OEt<sub>2</sub>, pyridine); the three-coordinate, monomeric aluminafluorene Mes\*-AlFlu was prepared from (**3**)<sub>2</sub>, Mes\*Li, and a 2,2'-dilithio-1,1'-biphenyl in quantitative yield (Mes\* = 2,4,6-(tBu)<sub>3</sub>C<sub>6</sub>H<sub>2</sub>).

#### Introduction

Doping organic  $\pi$ -electron systems with other p-block elements is an effective strategy to impart new chemical and physical properties to these species. Specifically, the combination of a polycyclic aromatic hydrocarbon (PAH) such as fluorene with boron as a dopant to generate 9-borafluorenes (BFlus) can have a particularly pronounced effect, as a conjugation barrier (*i.e.*, the CH<sub>2</sub> fragment in the carbonaceous species) is removed and a vacant B(p<sub>z</sub>) orbital is introduced instead, which can now: (i) mediate electron delocalization and bring about novel optoelectronic properties, (ii) facilitate reduction, and (iii) act as a Lewis acid to promote bond-activation reactions or the expansion of the five-membered central borole ring.

Compared to the extensive research on BFlus, their heavier homologues, the 9-aluminafluorenes (AlFlus),  $^{9,10}$  are far less well explored. This is unfortunate, because AlFlus are expected to exhibit a lower degree of aryl-heteroatom double-bond character than BFlus,  $^{11}$  leading to a greater propensity to form structurally intriguing aggregates through  $Al \cdots \pi(Ar)$  complexes

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or Al-C-Al' two-electron-three-center (2e3c) bonds. Relative to open-chain arylaluminum compounds, AlFlus should possess a structurally enforced enhanced Lewis acidity due to their small endohedral C-Al-C angle. This angle (108° in a regular five-membered ring) deviates more from the ideal 120° angle of three-coordinate AlR<sub>3</sub> species than from the corresponding angles of perfectly tetrahedral (109.5°) adducts. By the same token, the behavior of Al-based Lewis acids is more diverse than that of their B-based counterparts, as Al sites, unlike B centers, can readily accommodate coordination numbers larger than four.

In 1962, Eisch et al. reported the formation of Ph-AlFlu through the metalative cyclization of o-biphenylyl(diphenyl) aluminum at 200 °C. Their claim was mainly based on the analysis of hydrolysis and iodinolysis products. 12,13 The topic lay dormant until 2015, when Chujo and Tanaka used salt-metathesis protocols to synthesize AlFlus carrying Al-bonded phenyl rings with one or two chelating (dimethylamino)methyl substituents at their ortho positions (Fig. 1). Their research focused on the emission properties of the obtained four- and five-coordinate AlFlus. 14,15 More recently, Braunschweig et al. disclosed the synthesis of various aluminafluorenes R-AlFlu  $[R = 1,2,4-(tBu)_3C_5H_2 (92\%; Fig. 1), Ph_2(tBu)Si (44\%), 2-C_4H_3S$ (79%), tBu (23%; Fig. 1), Br (53%)]. The compounds were again prepared from 2,2'-dilithio-1,1'-biphenyl by salt-metathesis reactions and isolated and structurally characterized as their ether adducts – with the exception of the  $\eta^5$ -cyclopentadienide derivative, which is monomeric in the solid state, and the tBu derivative, which crystallizes as a dimer. 16

Me<sub>2</sub>N-Al-NMe<sub>2</sub>

tBu
tBu
tBu
tBu
tBu
tBu

Cp3t-AIFI

(tBu-AIFIu)<sub>2</sub>

Chujo & Tanaka 2015 Braunschweig 2021

**Fig. 1** Known mono- and dimeric 9-aluminafluorenes featuring 2,6-bis [(dimethylamino)methyl]phenyl (BDMAPh), 1,2,4-(tBu)<sub>3</sub>C<sub>5</sub>H<sub>2</sub> (Cp<sup>3t</sup>), and tert-butyl (tBu) substituents.

One aim of our study outlined herein was to develop straightforward, high-yield synthesis protocols for base-free R-AlFlus featuring (i) the small substituent R = Me to minimize steric shielding of the Al center, and (ii) the reactive substituent R = Br for late-stage derivatization. Particular emphasis was placed on the molecular structure of Me-AlFlu in nondonor solvents and in the solid state, as well as on the synthesis of the first base-free, three-coordinate, monomeric aluminafluorene, Mes\*-AlFlu (Mes\* = 2,4,6-(tBu) $_3C_6H_2$ ). All our AlFlus were equipped with tBu groups in their 2,7-positions to enhance solubility in non-polar solvents and to facilitate NMR-spectroscopic analysis.

#### Results and discussion

#### **Syntheses**

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The base-free Me-AlFlu (2) was synthesized by heating the 9,9-dimethyl-9-stannafluorene 1 with 1 equiv. of AlMe $_3$  <sup>17</sup> in either hexanes or C<sub>6</sub>H<sub>6</sub>/toluene (Scheme 1). The only by-product formed is the volatile and relatively inert SnMe $_4$ . <sup>18,19</sup> An advantage of using hexanes as the solvent is that the dimer (2) $_2$  precipitates in pure form already upon cooling the reaction mixture to room temperature (yield: 74%); when C<sub>6</sub>H<sub>6</sub>/toluene is employed, the yield of (2) $_2$  is higher (91%), but some further workup is required. In the presence of the donor molecules Et $_2$ O, THF, or pyridine, (2) $_2$  is cleanly split into its constituting monomers to furnish the monoadducts 2·OEt $_2$ , 2·thf, or 2·py (Scheme 1).

Treatment of  $(2)_2$  with 4 equiv. of  $AlBr_3^{17}$  in  $C_6H_6$  results not only in quantitative AlMe/AlBr exchange but also in the incorporation of two  $AlBr_3$  molecules to afford dimeric  $2,2'-(Br_2Al)_2$ -1,1'-biphenyl [(3)<sub>2</sub>, 95%; Scheme 1]. Upon addition of  $Et_2O$  to (3)<sub>2</sub> in  $C_6H_6$ , the donor adduct of Br-AlFlu,  $4\cdot OEt_2$ , precipitates quantitatively as a colorless solid. In terms of yield, our overall synthesis cascade to  $4\cdot OEt_2$  improves upon the published protocol<sup>16</sup> by about 40 percentage points. Although pyridine can also reconstitute the AlFlu scaffold from (3)<sub>2</sub>, it proved challenging to separate the target product  $4\cdot py$  from byproducts such as  $[AlBr_2(py)_4][X]$  ([5][X];  $X = Br^-$ ,  $AlBr_4^-$ ; Fig. S44 and S45†).

A particularly notable application of  $(3)_2$  as a synthetic equivalent of donor-free Br-AlFlu is the preparation of Mes\*-AlFlu

Scheme 1 Synthesis of donor-free (2) $_2$  through Sn/Al exchange between the 9-stannafluorene 1 and AlMe $_3$  (ArH:  $C_6H_6$ /toluene). The addition of AlBr $_3$  to (2) $_2$  furnishes (3) $_2$ . Lewis bases (L: Et $_2$ O, THF, or pyridine), cleave (2) $_2$  or (3) $_2$  into the monomeric adducts 2·L or 4·L. (i) Hexanes, 140 °C, 3 d (74% yield) or  $C_6H_6$ /toluene, 120 °C, 3 d (91% yield); sealed glass ampoule. (ii)  $C_6H_6$ , room temperature, 1 d (95% yield). (iii) 2·OEt $_2$ : in Et $_2$ O, room temperature; 2·thf:  $C_6D_6$ , room temperature; 2·py:  $C_6H_6$ , room temperature (quantitative conversions). (iv) 4·OEt $_2$ :  $C_6H_6$ , room temperature (quantitative conversion); 4·py:  $C_6D_6$ , room temperature (not isolated). Note: in (3) $_2$ , four bonds were arbitrarily chosen as formally intermolecular (highlighted in orange) to facilitate the distinction between the monomers M and M'.

(6): sequential addition of Mes\*Li (4 equiv.) and 2,2'-dilithio-4,4'-di-*tert*-butyl-1,1'-biphenyl (2 equiv.) to  $(3)_2$  in  $C_6H_6$  gave 6 in 97% yield (Scheme 2).

#### Solid-state structures

In the solid state, Me-AlFlu forms centrosymmetric dimers, with the Al-bonded Me substituents adopting a *trans*-configuration (*trans*-(2)<sub>2</sub>; Fig. 2).<sup>20</sup> The individual monomers, M and M', are linked by two Al···C interactions, resulting in two Al(1) ···Al(1)' bridging aryl rings (Ar<sub>b</sub>) and two terminal rings (Ar<sub>t</sub>), with bridging [C(11)] and terminal [C(21)] *ipso*-C atoms. The position of Ar<sub>b</sub> is asymmetric between Al(1) and Al(1)', as indicated by the differing angles Al(1)-C(11)···C(14) = 153.13(17)° and Al(1)'-C(11)···C(14) = 128.38(16)°. The fact that the

Scheme 2 Synthesis of Mes\*-AlFlu (6) using (3) $_2$  as a synthetic equivalent of the elusive Br-AlFlu. (i)  $C_6H_6$ , room temperature, 1 d (97% yield).

C(21) Al(1) C(11) C(21) Al(1) C(11) C(11)

Fig. 2 Top: molecular structure of *trans*-(2)<sub>2</sub> in the solid state; H atoms omitted for clarity (C: black, Al: turquoise). Bottom: structural formulae of *trans*-(2)<sub>2</sub> and of the comparable 9-borafluorene dimer (H-BFlu)<sub>2</sub>.

Al(1)'-C(11)-C(14) angle is significantly closer to 90° than the Al(1)-C(11)-C(14) angle can still be viewed as a remnant of the initial intermolecular Al $\cdots\pi$ (Ar) complex when the two heterofluorene units first encountered each other. Correspondingly, the 'intermonomer' Al(1)'-C(11) bond (2.148(3) Å) is longer by 0.055 Å than the 'intramonomer' Al(1)-C(11) bond (2.093(3) Å; cf. Al(1)-C(21) = 1.971(3) Å). The range of C-C bond lengths in  $Ar_b (1.382(5)-1.427(5) \text{ Å})$  is close to that in  $Ar_t (1.389(5)-1.409(5) \text{ Å})$ , indicating that the bridging mode does not lead to a systematic bond-length alternation. However, the two C-C bonds involving the bridging C(11) atom are slightly longer than the other four (1.409(4) and 1.427(5) Å vs. 1.382(5)-1.401(5) Å). Finally, we note that trans-(2)2 has very similar structural parameters to Braunschweig's (tBu-AlFlu)<sub>2</sub>, 16 while the comparable 9-borafluorene dimer (H-BFlu)<sub>2</sub> shows one B-(μ-H)-B two-electronthree-center bond and one B···B'-bridging aryl ring (the three other rings remain terminally bonded).<sup>21</sup>

X-ray crystallography reveals that the compound  $(3)_2$  no longer contains the 9-aluminafluorene motif but instead forms a centrosymmetric  $2,2'-(Br_2Al)_2-1,1'$ -biphenyl dimer (Fig. 3).

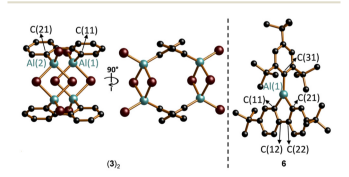


Fig. 3 Left: molecular structure of  $(3)_2$  in the solid state, viewed from two different perspectives; tBu-groups in the 2,7-positions of the biphenyl backbones and H atoms omitted for clarity. Right: molecular structure of 6 in the solid state; H atoms omitted for clarity (C: black, Al: turquoise, Br: brown).

The two Br<sub>2</sub>Al substituents in each monomer adopt an approximate *s-trans* configuration with a torsion angle Al(1)–C(11)–C(21)–Al(2) of 129.55(19)° [Al(1)–C(11) = 1.945(5) Å, Al(2)–C(21) = 1.956(4) Å]. Four Br atoms occupy bridging positions between Al centers of different monomers, assembling the cage-like structure of (3)<sub>2</sub>. The underlying structural feature, a four-membered R(Br)Al–( $\mu$ -Br)<sub>2</sub>–Al(Br)R ring, is common not only for aluminum tribromide (R = Br) but also for numerous dibromo(organo)alanes.<sup>22</sup>

The 2,4,6-(tBu)<sub>3</sub>C<sub>6</sub>H<sub>2</sub>-substituted Mes\*-AlFlu (6) exists as a monomeric species with a three-coordinate Al center in the crystal lattice (Fig. 3). The sum of C-Al-C angles is 360°, confirming a trigonal-planar ligand environment, although the endocyclic C(11)-Al(1)-C(21) bond angle is nearly rectangular (91.79(6)°). All three Al–C bonds are of equal length, regardless of whether they are endo- or exocyclic, or whether the respective ipso-C(pz) orbital is positioned parallel or orthogonal to the vacant  $Al(p_z)$  orbital [Al(1)-C(11)/C(21)/C(31) = 1.9611(14)/C(21)/C(31) = 1.9611(14)/C(21.9516(14)/1.9606(13) Å]. Within the five-membered AlC<sub>4</sub> core, the length of the central C(12)-C(22) bond (1.5024(19) Å) approaches that of a typical C-C single bond (1.54 Å), 23 while benzannulated bonds are significantly shorter [C(11)-C(12)/C(21)-C(22) = 1.4208(18)/1.4134(18) Å]. The other ten C-C bonds within the biphenyl fragment fall within a narrow range of 1.389(2)-1.4041(19) Å, closely matching the corresponding bonds in C<sub>6</sub>H<sub>6</sub> (1.39 Å).<sup>24</sup> Taken together, this analysis of bond lengths suggests that the AlFlu moiety of 6 preserves two largely unperturbed Clar sextets within its two C<sub>6</sub>H<sub>3</sub> fragments, with no indication of a delocalized (antiaromatic)  $\pi$ -system, nor any significant Al(1)=C(11)/C(21) doublebond character in the AlC<sub>4</sub> heterocycle.

The donor adducts  $2 \cdot \mathrm{OEt}_2$ ,  $2 \cdot \mathrm{py}$ , and  $4 \cdot \mathrm{py}$  were subjected to X-ray analysis to confirm that  $(2)_2$  and  $(3)_2$  can indeed serve as precursors of Me-AlFlu and Br-AlFlu, respectively (Fig. S40, S41, S43†). Furthermore, compared to donor-free **6**, the C–C bond lengths within the  $C_6H_3$ – $C_6H_3$  units of  $2 \cdot \mathrm{py}$  and  $4 \cdot \mathrm{py}$  were found to differ by no more than  $3\sigma$  (and much less for most bonds). This observation again suggests that the vacant  $Al(p_z)$  orbital exerts no significant electron-withdrawing mesomeric effect on the  $\pi$ -electron system.

#### NMR analysis

At room temperature, **2** gives severely broadened <sup>1</sup>H NMR signals, providing limited diagnostic value (Tol-*d*<sub>8</sub>; Fig. 4 and S7†). At 70 °C, two sharp resonances are detectable in the aliphatic region of the spectrum (integral ratio 3H:18H); the aromatic region contains one broad feature and two doublets with coupling constants of about 8.2 Hz (Fig. 4 and S6†). At –30 °C, the <sup>1</sup>H NMR spectrum of **2** is characterized by two well-resolved sets of signals attributable to two different but closely similar components (Fig. 4 and S8†); the same is true for the <sup>13</sup>C{<sup>1</sup>H} NMR spectrum (Fig. S9†). The proton-integral values of the two sets indicate a minor-to-major component ratio of approximately 0.12:1 (Fig. S8†). Focusing on the major component, the <sup>1</sup>H NMR spectrum reveals one singlet at –0.67 ppm (6H), and two additional singlets at 1.47 and

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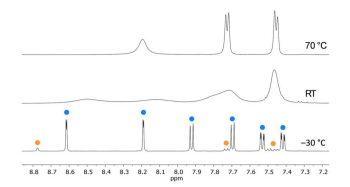


Fig. 4 Aromatic regions of  ${}^{1}H$  NMR spectra of (2)<sub>2</sub> in Tol- $d_8$  (500.2 MHz). Top: 70 °C. Middle: room temperature. Bottom: -30 °C.  $\bullet$ :  $trans-(2)_2$ .  $\bullet$ :  $cis-(2)_2$ .

1.28 ppm (2 × 18H), assignable to two equivalent AlMe substituents and two pairs of non-equivalent tBu groups, respectively. In the aromatic region, four doublets  $(4 \times 2H; 2 \times {}^{3}J_{H,H} =$ 8.2 Hz,  $2 \times {}^4J_{\rm H,H} = 2.2$  Hz) and two doublets of doublets (2 × 2H) are observed, indicative of two pairs of non-equivalent C<sub>6</sub>H<sub>3</sub> fragments. In principle, these NMR features would align with both the molecular structure of the cis- and trans-(2)<sub>2</sub> dimer (as observed in the solid state). Vice versa, the minor signal set likely arises from trans- or cis-(2)2. At low temperatures, both isomers are present in an (essentially) static mixture, while some dynamic rearrangement equilibrium is established at higher temperatures. This preliminary conclusion raises two questions: (i) Does cis- or trans-(2)2 dominate at low temperatures? (ii) Is the dynamic equilibrium at high temperatures due to monomer/dimer association/dissociation, or is it the result of a rapidly interconverting cis/ trans dimeric form of  $(2)_2$ ?

To address question (i), quantum-chemical calculations predict that the crystallographically characterized trans-(2)<sub>2</sub> is 1.6 kcal mol<sup>-1</sup> more favorable in energy than cis-(2)<sub>2</sub> (Scheme S1†; experimental value, determined at -30 °C from the proton-integral values of the minor/major component:  $\Delta G^{\circ} = 0.7$  kcal mol<sup>-1</sup>). Furthermore, the relative proportion of the minor component increases with solvent polarity, consistent with the existing dipole moment of cis-(2)<sub>2</sub> (<sup>1</sup>H NMR spectroscopic control; Table S1 and Fig. S1, S2†). Finally, the computed <sup>13</sup>C chemical shift values for cis/trans-(2)<sub>2</sub> align more closely with the assumption that the major component is trans-(2)<sub>2</sub> rather than  $vice\ versa\$ (Tables S9–S11†). It is therefore safe to assume that the major component in an equilibrating cis/trans-(2)<sub>2</sub> mixture is the trans isomer.

Regarding question (ii), we note that the computed energy required for cleaving trans-(2)<sub>2</sub> into its constituting monomers is 19.4 kcal mol<sup>-1</sup> (in CH<sub>2</sub>Cl<sub>2</sub>). In contrast, the computed energy barrier of the cis/trans interconversion of (2)<sub>2</sub> is only  $\Delta G^{\ddagger} = 14.8$  kcal mol<sup>-1</sup>, which agrees well with the value of  $\approx 14.5$  kcal mol<sup>-1</sup> experimentally determined from the coalescence temperature ( $T_c$ ) in conjunction with the maximum peak separation ( $\Delta \nu$ ) in the slow-exchange limit (CD<sub>2</sub>Cl<sub>2</sub>; see the

ESI† for full details). The observed NMR features are therefore more convincingly attributed to a dynamic *cis/trans* equilibrium rather than to a monomer/dimer association/dissociation equilibrium.

In the temperature range of -30 to 70 °C,  $(3)_2$  exhibited only extremely broadened signals in the <sup>1</sup>H and <sup>13</sup>C{<sup>1</sup>H} NMR spectra, providing no structural information.

The <sup>1</sup>H and <sup>13</sup>C{<sup>1</sup>H} NMR spectra of all adducts formed between our R-AlFlus and Lewis bases are in accord with the proposed molecular structures, as is the case for ligand-free 6 (see the ESI† for the fully assigned spectra). In addition to aiding in structure elucidation, <sup>13</sup>C{<sup>1</sup>H} NMR spectroscopy is also a valuable tool for mapping the  $\pi$ -charge density distribution in conjugated systems, as the shielding of a specific  $C(sp^2)$  atom depends linearly on the corresponding  $\pi$ -electron density at that position.26 Given this background, we compared the <sup>13</sup>C chemical shift values of the C atoms constituting the C<sub>6</sub>H<sub>3</sub>-C<sub>6</sub>H<sub>3</sub> fragment of **6** with those of the equivalent atoms in the corresponding fragments of the adducts 2.OEt2, 2.thf, 2.py, 4.OEt2, and 4.py. Except for the Al-bonded ipso-C atoms, whose shift differences varied from  $\delta(6)$ - $\delta(adduct) = 3.3$  to -4.7 ppm without a systematic trend, the  $\Delta\delta(^{13}\text{C})$  values for all other structurally analogous C atoms were less than ±1.8 ppm. In other words, we found no evidence of an overall <sup>13</sup>Cdeshielding effect or  $\pi$ -electron depletion in 6 that could be attributed to a mesomerically electron-withdrawing Al(sp<sup>2</sup>)

<sup>13</sup>C{¹H} NMR spectroscopy on 2·py and 4·py provides a method to evaluate the relative Lewis acidities of free, monomeric Me-AlFlu and Br-AlFlu: in pyridine complexes of maingroup elements, stronger acids induce increased shielding of the C-2,6 and deshielding of the C-3,4,5 nuclei of the ligand.<sup>27</sup> For 2·py/4·py, our observations consistently indicate that Me-AlFlu is the stronger acid, comparable in this respect to BPh<sub>3</sub>.<sup>28</sup> X-ray crystallography, however, offers a contrasting view: 4·py exhibits a shorter Al−N bond and a more pyramidalized Al center, implying higher Lewis acidity for Br-AlFlu.<sup>28</sup> Given the small differences in the key NMR and structural parameters between 2·py and 4·py, these conflicting observations highlight the limitation of relying on a single method to determine Lewis acidity, emphasizing the need for complementary approaches.

#### Conclusions

We synthesized the donor-free 9-aluminafluorene Me-AlFlu (2), which was characterized as its dimer (2)<sub>2</sub> through X-ray crystallography and VT NMR spectroscopy (Tol- $d_8$ ). The key to this success was the highly selective reaction between the 9,9-dimethyl-9-stannafluorene 1 and AlMe<sub>3</sub>,<sup>17,18</sup> which proceeds in non-donor solvents and releases volatile SnMe<sub>4</sub> as the sole byproduct. Unlike the bulky *tert*-butyl group in *t*Bu-AlFlu,<sup>16</sup> the sterically less demanding methyl substituent in Me-AlFlu allows relatively unhindered access to the electrophilic Al center, as demonstrated by the straightforward formation of

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various base adducts  $2 \cdot L$  ( $L = OEt_2$ , thf, py). In terms of an umpolung of the Al center, exploring the reduction of 2 on a preparative scale could be promising (a non-benzannulated alumole has previously been reduced to its corresponding dianion). The resulting product,  $[2]^{2-}$ , could potentially serve as an Al-centered nucleophile – analogous to the  $[H-BFlu]^{2-}$  dianion, which is a valuable B-centered nucleophile.  $^{3,6,29}$ 

Treatment of  $(2)_2$  with AlBr<sub>3</sub> furnishes the 2,2'-(Br<sub>2</sub>Al)<sub>2</sub>-1,1'-biphenyl  $(3)_2$ . Although this ring-opened product no longer retains the AlFlu motif, it rearranges back to afford Br-AlFlu adducts, such as 4·L (L = OEt<sub>2</sub>, py), in the presence of Lewis bases. The use of Mes\*Li, which provides the extremely bulky, negatively charged Lewis base [Mes\*]<sup>-</sup>, grants unprecedented access to three-coordinate, monomeric aluminafluorenes, specifically Mes\*-AlFlu (6), *via* LiBr elimination. This reaction proves the utility of  $(3)_2$  as a synthetic equivalent for the still-elusive free Br-AlFlu. A comparison of characteristic structural and NMR features of 6 with those of  $2 \cdot \text{L}/4 \cdot \text{L}$  reveals that the three-coordinate Al center exerts only a negligible  $\pi$ -electron withdrawing effect and does not mediate significant  $\pi$ -electron delocalization.

#### Author contributions

P. L. L. performed the experimental studies and characterized all new compounds. P. L. L. and J. G. performed the quantum-chemical calculations. A. V. performed the X-ray crystal structure analyses of all compounds. H.-W. L. and M. W. supervised the project. The manuscript was written by P. L. L. and M. W. and edited by all co-authors.

### Data availability

The datasets supporting this article have been uploaded as part of the ESI.  $\dagger$ 

#### Conflicts of interest

There are no conflicts to declare.

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