Structure and bonding in reduced boron and aluminium complexes with formazanate ligands†

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Group 13 complexes of the type [(PhNNC(p-tol)NNPh)ZPh2]2− (Z = B, Al) containing a highly reduced, tri-anionic formazanate-derived ligand were studied and the differences in the structure, bonding and reactivity between the B and Al compounds were investigated. The increased ionic character in the bonding of the Al complex is evident from the enhanced charge delocalization onto the peripheral ligand substituents (N–Ph) via the π-framework, as shown by the rotation barrier around the N–C(Ph) bond. The electron-rich nature of these compounds allows facile benzylation at the ligand, and the structures of the products were analysed by X-ray crystallography. The products are inorganic analogues of 1-alkylated 1,2,3,4-tetrahydro-1,2,4,5-tetrazines ('leucoverdazyls'). The six-membered heterocyclic cores of the B and Al compounds are shown to be different, having envelope- and boat-type conformations, respectively. Homolysis of the N–C(benzyl) bond in these compounds was studied by NMR spectroscopy under conditions that trap the organic radical as TEMPO-Bn. Analysis of the reaction kinetics affords activation parameters that approximate the N–C(benzyl) bond strength. The ionic Al compound has one of the weakest N–C bonds reported so far in this type of inorganic leucoverdazyl analogues.

Introduction

The direct involvement of ligands in redox reactions by coordination complexes bearing non-innocent ligands is an emerging research area.1–3 In this context, there is growing interest in designing new types of redox-active (non-innocent) ligands. In 2007, Hicks and coworkers observed reversible ligand-based reduction in a formazanate boron acetate compound.4 This demonstrated for the first time that the formazanate ligand can be considered as a redox-active analogue of the well-known β-diketiminate ligands.5–8 Taking inspiration from this work, our group has explored the coordination chemistry, redox behavior and reactivity of complexes with formazanate ligands.9–16 Along with our work, the Gilroy group17–26 and others27–29 have been involved in the synthesis and application of new molecular complexes with formazanate ligands. Previously, we showed that the ligands in boron and aluminium compounds with formazanate ligands could be sequentially reduced by 1- and 2-electrons at moderate reduction potentials (Scheme 1).30,31 The 2-electron reduced formazanate boron compound (12−) subsequently reacted with electrophiles (E+) such as benzyl bromide (BnBr) and water (H2O) to form ligand-benzylated and -protonated products (Bn1− and H1− in Scheme 1).32 In these compounds, the formazanate ligands are modified by the ‘storage’ of [2e−/E+], which could be converted to Bn· and H· radicals by the homolytic cleavage of the N–C (Bn) and N–H bonds, respectively (Scheme 1).32 These reactions occur readily, because the boron-containing radical that

Scheme 1. Ligand-based storage of [2e−/E+], and subsequent conversion to E· (Bn·/H·) radicals that can be trapped with TEMPO.
is generated (1⁻⁻) is relatively stable due to the presence of a low-energy SOMO that is delocalized over all four N-atoms in the ligand backbone. It is anticipated that the basicity (nucleophilicity), radical stability and N-C/N-H bond strength of compounds bearing functionalized formazanate ligands could be tuned via either ligand substituent effects or incorporation of a different central element (main group or transition metal) in the formazanate chelate ring.

In order to investigate the effects of changing the central element (boron) in 1 to the more electropositive aluminum, here we provide a detailed comparison of formazanate B and Al complexes with an identical ligand. The comparison includes an analysis of resonance delocalization in the two-electron reduced formazanate aluminum diphenyl compound (2⁻⁻) via dynamic NMR spectroscopy. The synthesis of ligand-benzylated products Bn₂⁻⁻ is described, and crystallographic and spectroscopic characterization data are provided. Furthermore, the kinetics of homolytic N-C(benzyl) cleavage is studied.

Results and discussion

The two-electron reduced formazanate aluminium diphenyl compound [PhNCC(p-tol)NNPh]AlPh₂⁻⁻ (2⁻⁻) was synthesized as its disodium salt according to a previously published procedure. The product 2⁻⁻, which has an electron-rich, formally trinionic formazanate ligand, is highly air-sensitive, but stable at room temperature under inert conditions. The ¹H NMR spectrum of 2⁻⁻ in THF-d₈ was shown to be temperature-dependent: at 233 K, the spectrum shows 5 inequivalent resonances due to the N-Ph groups. Increasing the temperature results in line broadening and ultimately coalescence of three resonances due to the N=C(Ph) groups. The ¹H NMR spectra of 2⁻⁻ in THF-d₈ was shown to be temperature-dependent: at 233 K, the spectrum shows 5 inequivalent resonances due to the N-Ph groups. Increasing the temperature results in line broadening and ultimately coalescence of three resonances due to the N=C(Ph) groups. The observed down to 233 K. We interpret the difference between the B and Al complexes with an identical formazanate ligand (1⁻⁻ and 2⁻⁻) as a reflection of the difference in bonding within the boron and aluminum heterocycles. In particular, the largely ionic character of the Al–N bonds in comparison with the more covalent B–N interactions is responsible for the increased accumulation of negative charge within the NNCNN framework in the Al compound. An increase in N–C(Ph) π-bonding due to resonance delocalization of the negative charge into the N–Ph group is also observed in the solid state structures obtained by X-ray crystallography: the N(C(Ph)) bonds in 2⁻⁻ are 1.371(2)/1.375(3) Å, whereas those in 1⁻⁻ are marginally larger at 1.379(3)/1.385(3) Å.

To further probe the bonding differences between formazanate boron and aluminium complexes, a DFT computational study was carried out (B3LYP functional and 6-311+G(d,p) basis set). The geometries were optimized in the gas phase starting from the crystallographic coordinates, with the Na(THF), cations removed (the computational results for 1²⁻⁻ were described previously). The metrical parameters of the optimized structures of 1²⁻⁻ and 2²⁻⁻ are in good agreement with the experimental structures, albeit the N–N bonds are somewhat shorter in the DFT models (see Table S2†). Visual inspection of the frontier orbitals shows that the HOMO is primarily π antibonding between the N-atoms in the formazanate ligand, and in addition, evidences the presence of the π bonding character in the N–C(Ph) fragment (Fig. S15†). Although the differences are small, the calculated Wiberg bond index for the N–C(Ph) bonds is larger in the Al compound (2²⁻⁻: 1.24) than in the B compound (1²⁻⁻: 1.22), corroborating the trend for the strength of π-bonding that was obtained from the NMR study. In addition, a higher Wiberg bond index is found for the B–N bonds in 1²⁻⁻ (0.68) in comparison with 2²⁻⁻ (0.64).

†The additional broadening observed above 283 K is attributed to the presence of a small amount of radical species (the paramagnetic monoanion 2⁻) that engages in electron transfer with 2⁻⁻.

§The estimated upper limit of the barrier for N–C(Ph) bond rotation is ca. 40 kJ mol⁻¹. See ESI† for details.
comparison with the Al–N bonds in $2^{2-}$ (0.40) and the natural charges indicate that the formazanate ligand bears a significantly higher negative charge in the Al complex (−2.40e) than in the B analogue (−1.99e).

Computational evaluation of the barrier to rotation around the N–C(Ph) bond in $2^{2-}$ was carried out by scanning the dihedral angle between the Ph ring and the ligand backbone. As expected, a maximum was found at a dihedral angle of ca. 90°, and at this geometry a transition state optimization was carried out to arrive at a saddle point ($2^2$-TS, $N_{\text{imag}} = 1$). The computed barrier for N–C(Ph) bond rotation was found to be 55.9 kJ mol$^{-1}$, which is in good agreement with the experimental value (−54.9 kJ mol$^{-1}$ at 298 K). A comparison between the ground and transition states shows that π-bonding is lost in the N–Ph ring that is rotated (Wilberg bond index = 1.04), whereas a small increase is observed in the other N–C(Ph) bond (1.26). This is also reflected in the different N–C(Ph) bond lengths that are calculated for $2^2$-TS (1.411 Å and 1.365 Å; cf. 1.370 Å in $2^{2-}$).

We subsequently evaluated the reactivity of compound $2^{2-}$ towards the electrophile benzyl bromide. Treatment of an orange THF-$d_6$ solution of $2^{2-}$ with 1 equivalent of Bu$_4$NBr resulted in an immediate colour change from orange to yellowish-green. The $^1$H NMR spectrum of the reaction mixture at room temperature (400 MHz, THF-$d_6$) reveals the diagnostic resonances for the diastereotopic protons of the benzyl-CH$_2$ group at $\delta$ 4.29 and 4.45 ppm with a geminal coupling constant $J_{HH} = 12.5$ Hz. Also, the $^1$H NMR spectrum shows two inequivalent resonances for the para-protons of the N–Ph rings at $\delta$ 6.09 and 6.34 ppm due to a descent in symmetry from the $C_{2v}$-symmetric precursor $2^{2-}$. The NMR data of the reaction product are similar to those of the boron analogue, and suggest that it contains a benzyl group that is attached to a N-atom of the ligand backbone. Consequently, we formulate the product as an anionic, ligand-benzylated complex [PhNN\(\text{(Bz)}\text{C(Ph)}\text{–NNPh}\)]AlPh$_2$$^{-}$ ([$^{\text{[Bn2]}\text{[Na]}$], Scheme 2). On a preparative scale, the reaction between $2^{2-}$ and Bu$_4$NBr allowed isolation of the product as an anionic, ligand-benzylated complex [PhNN\(\text{(Bz)}\text{C(Ph)}\text{–NNPh}\)]AlPh$_2$$^{-}$ ([$^{\text{Bn2]}\text{[Na]}$] and [$^{\text{Bn1]}\text{[Na]}$] in yields of 75%. Gratifyingly, crystals of [$^{\text{Bn2]}\text{[Na]}$] had a much higher melting point, and structure determination using X-ray was successfully completed for both B- and Al-containing products. The molecular structures of [$^{\text{Bn1]}\text{[Na]}$] and [$^{\text{Bn2]}\text{[Na]}$] show that in both of these compounds, the benzyl group is attached to one of the internal N-atoms to retain the 6-membered chelate ring structures of the dianionic precursors (Fig. 2, pertinent metrical data in Table 1). A closer inspection of the central six-membered

![Scheme 2](image)

Table 1: Selected bond lengths (Å) and bond angles (°) of [$^{\text{Bn1]}\text{[NBu}_4\text{]}$] and [$^{\text{Bn2]}\text{[Na]}$].

| Bond Lengths | [$^{\text{Bn1]}\text{[NBu}_4\text{]}$] | [$^{\text{Bn2]}\text{[Na]}$] |
|--------------|---------------------------------|--------------------------|
| N1–N2        | 1.428(2)                        | 1.446(1)                 |
| N3–N4        | 1.393(2)                        | 1.402(1)                 |
| N2–C7        | 1.396(2)                        | 1.437(1)                 |
| C7–N3        | 1.289(2)                        | 1.292(1)                 |
| A1–N1        | 1.867(1)                        | 1.867(1)                 |
| A1–N4        | 1.914(1)                        | 1.914(1)                 |
| B1–N1        | 1.575(2)                        |                          |
| B1–N4        | 1.584(2)                        |                          |
| N1–C1        | 1.414(2)                        | 1.387(1)                 |
| N4–C15       | 1.392(2)                        | 1.399(1)                 |
| N2–C33       | 1.468(2)                        | 1.486(1)                 |
| N1–B1–N4     | 105.78(1)                       |                          |
| N1–A1–N4     | 95.24(4)                        |                          |
| N1–N2–C7     | 115.41(1)                       | 112.27(8)                |
| N1–N2–C33    | 113.94(1)                       | 110.33(8)                |
| C7–N2–C33    | 118.79(1)                       | 112.13(8)                |
heterocyclic rings in these compounds reveals some noteworthy differences. Compound [Bn1][NBu4] adopts an envelope conformation which is similar to the precursor 12−31 with the formazanate backbone atoms (NNCNN) nearly coplanar and the B atom residing 0.568 Å above that plane. In contrast, the six-membered ring in the Al compound [Bn2][Na] is found to be present in a boat conformation in which N1, A1, N3 and C7 are coplanar, whereas N2 and N4-atoms are displaced out from that plane by 0.502 and 0.507 Å, respectively (Fig. 2). This is markedly different from the nearly planar conformation in the precursor 22−31. A Cremer–Pople puckering analysis16 was carried out using PLATON,37 which allowed a more quantitative distinction between the divergent geometrical parameters within the core of [Bn1][NBu4] vs. [Bn2][Na]. In this analysis, a combination of the puckering amplitude (Q) and two polar coordinates (θ and ϕ) are used to map the conformations onto a sphere. The following puckering parameters are obtained for these compounds: for [Bn1][NBu4] (Q = 0.44 Å; θ = 57.0°; ϕ = 332.2°) and [Bn2][Na] (Q = 0.58 Å; θ = 89.8°; ϕ = 300.2°). Of these parameters, the θ angles in six-membered rings distinguish among the chair, envelope and boat conformations, whereas ϕ defines the pseudorotation pathways that transverse twisted conformations as well. The angles θ found for both compounds are indicative of either an envelope (idealized: 54.7°; [Bn1][NBu4]: 57.0°) or boat conformation (idealized: 90°; [Bn2][Na]: 89.8°).

In the B and Al dianions (12− and 22−) the bonding in the six-membered core is delocalized, as shown by the equivalent N−N and C−N bond lengths, whereas ligand benzylation leads to a more localized bonding picture. For example, the C−N bonds in the ligand backbone change from 1.332(2)/1.328(2) Å in 22− to 1.437(1)/1.292(1) Å in [Bn2][Na]. Similar changes in bond lengths occur in the boron analogue (see Tables S2/S3† for a full comparison of metrical data). In both compounds, the benzylated N-atom is pyramidal as indicated by the sum of angles around N(2) being much smaller than 360° (Σξ(N(2))) = 348.1° and 334.7° for [Bn1][NBu4] and [Bn2][Na], respectively. In [Bn2][Na] the Ph-substituted N-atoms are planar (sp2-hybridized) with Σξ(N) = 359.5° for N(1) and 354.5° for N(4). Conversely, in the boron analogue [Bn1][NBu4] the N(4) atom is almost planar (Σξ(N) = 357.5°), while N(1) is pyramidalized (Σξ(N) = 347.8°). It appears that these differences reflect a larger degree of directionality (covalency) in the B−N bonds in comparison with the more ionic Al congener.

The anions Bn1− and Bn2− were also evaluated by DFT calculations. Optimized geometries of Bn1− and Bn2− were obtained starting from the crystallographic coordinates (counter cations were removed) and were shown to be in good agreement with the empirical structures. Importantly, the disimilarities in pyramidalization at the N-atoms of both the B and Al compounds are reproduced in the computational models, suggesting that these are intrinsic and are not due to packing or ion pairing effects. In both the experimental structures and the DFT models, there is a small but noticeable elongation of the N−C(Bn) bond in the Al compound Bn2− (X-ray: 1.486(1) Å; DFT: 1.492 Å) compared to that in the B congener Bn1− (X-ray: 1.468(2) Å; DFT: 1.472 Å). An NBO analysis indicated that the bonding within the NNCNN ligand backbone is similar in both B and Al compounds and can be explained by a normal σ-bonding framework with an additional localized N=−C σ-bond (Fig. 3A). The hybridization of the benzyl-substituted nitrogen atom is intermediate between sp2 and sp3, with a lone pair that has a small but non-negligible amount of the s-character (10% for Bn1−: 16% for Bn2−). Importantly, the NBO analysis shows that the N−B and N−Al interactions in both the compounds are quite different. For the boron compound Bn1−, we find two natural bond orbitals that represent the relatively covalent B−N σ-bonds (ca. 77% contribution from N; 23% from B). For the Al analogue Bn2−, on the other hand, no 2-center Al−N bonds are obtained and, instead, the NBO analysis indicates that the N-based lone pairs interact with an empty Al orbital (Fig. 3). Similarly, the natural charges indicate that the Al complex is much more ionic (NPA charge for Al in Bn2−: 1.82; B in Bn1−: 0.74).

A comparison of the UV/Vis spectra of the Na+ and Bu4N+ salts of Bn1− and Bn2− shows that the absorption maxima in dilute THF solution are independent of the nature of the cation. The boron compounds Bn1− and the aluminium compounds Bn2− absorb at 395 and 389 nm, respectively (Fig. S1†). These absorbance maxima are most likely due to a π−π* transition of the localized N=−C bonds present in these compounds, as shown by the crystallographic data (vide supra), and are blue-shifted compared to the delocalized precursors 12− and 22− (λmax = 486 nm).31

†The UV/Vis spectrum of 12− reported in the ESI† of ref. 30 is incorrect, likely due to the decomposition of this highly sensitive compound.
The compounds $^{\text{Bn}}1^-$ and $^{\text{Bn}}2^-$ are inorganic, anionic analogues of purely organic 1-alkylated tetrahydro-1,2,4,5-tetrazines (‘leucoverdazyls’). Leucoverdazyls (with N–H bonds) and the alkylated (N–C) derivatives are known to have weak N–H/N–C bonds because homolytic cleavage generates a stable verdazyl radical. The stability of this type of radical extends to inorganic systems. In order to examine the effect on the N–C(Bn) bond dissociation energy (BDE) due to the replacement of B to the more electropositive Al in $^{\text{Bn}}2^-[\text{Na}]$, the cleavage of the N–C bond in $^{\text{Bn}}2^-[\text{Na}]$ was investigated. The N–C(Bn) bond dissociation enthalpy in $^{\text{Bn}}2^-[\text{Na}]$ was obtained experimentally by NMR spectroscopic monitoring of the kinetics of benzyl radical transfer from $^{\text{Bn}}2^-[\text{Na}]$ to TEMPO (present in excess) in the temperature range of 65–85 °C (Scheme 3).

Clean exponential decay of the starting material and concomitant appearance of TEMPO-Bn were observed, which allowed the rate constants to be determined (Fig. S8†). An Eyring analysis provided the activation parameters $\Delta H^\ddagger$ and $\Delta S^\ddagger$ of 107 ± 4 kJ mol$^{-1}$ and 17 ± 11 J mol$^{-1}$ K$^{-1}$, respectively, for the benzyl transfer (Fig. 4; see the ESI† for details). Both these values are somewhat smaller than those found for the boron analogue $^{\text{Bn}}1^-[\text{Na}]$ ($\Delta H^\ddagger = 121 ± 5$ kJ mol$^{-1}$; $\Delta S^\ddagger = 77 ± 14$ J mol$^{-1}$ K$^{-1}$). A likely explanation for these differences is the ground-state destabilization of the N–C(Bn) bond which is indicated by the somewhat larger N–C(Bn) distance found by both experiment and theory (vide supra).

To investigate a possible influence of ion-pairing on the N–C(Bn) bond dissociation energy, the kinetics of benzyl transfer were also measured for the tetrabutyl ammonium salts $^{\text{Bn}}1^-[\text{NBu}_4]$ and $^{\text{Bn}}2^-[\text{NBu}_4]$ using the same methodology. The resulting rate constants are in good agreement with those of the sodium salts (see the ESI† for details). Thus, even though in the solid state the sodium cation is bound to the ligand backbone, this weak interaction is likely broken in the solution. This is further supported by the observation that the UV/Vis spectra of the solution do not depend on the nature of the cation.

These data indicate that N–C(Bn) bond homolysis is modulated by the central element in these heterocyclic leucoverdazyl analogues: on going from relatively covalent, C-based parent structures (i.e., 1-benzyl-substituted 1,2,3,4-tetrahydro-1,2,4,5-tetrazines), the N–C(Bn) bond strength progressively decreases with an increase in the electropositive nature of the central element (i.e., C > B > Al).

Conclusions

In conclusion, this work addresses the differences in the structure and bonding between diatomic formazanate boron ($1^-$) and aluminum ($2^-$) complexes. Experimental (NMR, UV/Vis spectroscopy, and X-ray crystallography) and computational studies (Wiberg bond indices and NBO analysis) reveal that the increased ionic character of the Al compounds results in a higher degree of resonance delocalization of the ligand negative charge into the periphery of the ligand (i.e., the N–Ph substituents), which is reflected in the rotation barrier around the N–C(Ph) bond. For both the compounds, facile ligand benzyl transfer occurs upon reaction with benzyl bromide to form $^{\text{Bn}}1^-$ and $^{\text{Bn}}2^-$ as anionic analogues of carbon-based leucoverdazyls. X-ray diffraction studies of $^{\text{Bn}}2^-$ and the boron congener $^{\text{Bn}}1^-$ are reported and a comparison shows distinct differences in the solid state structures between these complexes which can be related to a different degree of the ionic character in the bonding. The kinetics of benzyl transfer show that the N–C (Bn) bond homolysis is modulated by the nature of the central element present in the six-membered heterocyclic rings of $^{\text{Bn}}1^-$ and $^{\text{Bn}}2^-$, with the ionic Al-based compound $^{\text{Bn}}2^-$ having the weakest N–benzyl bond.

Experimental section

General considerations

All manipulations were carried out under a nitrogen atmosphere using standard glovebox, Schlenk, and vacuum-line techniques. THF and hexane (Aldrich, anhydrous, 99.8%) were passed over columns of Al$_2$O$_3$ (Fluka). The compounds $^{\text{Bn}}1^-[\text{NBu}_4]$, $^{\text{Bn}}2^-[\text{Na}]$ and $^{\text{Bn}}2^-[\text{NBu}_4]$ are highly air-sensitive, and the solvents (THF and hexane) used for their preparation and characterization were additionally dried on a Na/K alloy and subsequently vacuum transferred and stored under nitrogen. All solvents were degassed prior to use and stored under nitrogen. THF-$d_8$ (Sigma-Aldrich) was vacuum transferred from the Na/K alloy and stored under nitrogen. The compounds $^{\text{Bn}}1^-[\text{Na}]$ and $2^-$ (as its disodium salt, [PhNNC(p-tol)NNPh])

Scheme 3 Benzyl group transfer from $^{\text{Bn}}2^-[\text{Na}]$ to TEMPO.

Fig. 4 Eyring analysis for benzyl transfer from $^{\text{Bn}}2^-[\text{Na}]$ to TEMPO.
AlPh$_3$][Na$_2$(DME)$_3$]$^{31}$ were synthesized according to published procedures. NMR spectra were recorded on a Varian Mercury 400 or Inova 500 spectrometer. The $^1$H and $^{13}$C NMR spectra were referenced internally using the residual solvent resonances and reported in ppm relative to TMS (0 ppm); $J$ is reported in Hz. Assignments of NMR resonances were aided by COSY, NOESY, HSQC and HMBC experiments using standard pulse sequences. UV-Vis spectra were recorded in THF solution ($\sim 10^{-4}$ M) in a quartz cuvette using an AVANTEA AvaSpec-2048. Samples for elemental analyses were sent to Kolbe Microanalytical Laboratory (Mülheim an der Ruhr, Germany). However, despite our best efforts, no satisfactory analysis data could be obtained for these compounds, which is likely due to their highly air-sensitive nature.

### Synthesis of $[^{18}$Na$]_{[BnLAlPh_2][NBu_4]}$ ($[^{18}$Na$]_{Na_2}$)$^{31}$

Compound $[^{18}$Na$]_{[BnLAlPh_2][NBu_4]}$ ($[^{18}$Na$]_{Na_2}$) was synthesized as follows: 20.71 mg ($0.072$ mmol, 78%) $^1$H NMR (400 MHz, THF-$d_8$) 7.70 (d, $J = 7.1$ Hz, 2H, B-Ph(1) o-H), 7.11–7.06 (m, overlapped, 4H, B-Ph(1) (m-)H and p-tol-CH$_3$), 1.29 (s, 8H, NBu$_4$), 1.58 (s, 8H, NBu$_4$), 1.34–1.30 (m, 3H, NBu$_4$), 0.95 (t, $J = 6.8$ Hz, 12H, NBu$_4$).

### Synthesis of $[^{18}$LAlPh$_3$][Na$_2$(THF)$_3$]$^{32}$

Compound $[^{18}$LAlPh$_3$][Na$_2$(THF)$_3$]$^{32}$ was synthesized as follows: 20.71 mg ($0.072$ mmol, 78%) $^1$H NMR (400 MHz, THF-$d_8$) 7.70 (d, $J = 7.1$ Hz, 2H, p-tol-CH$_3$), 7.11–7.06 (m, overlapped, 4H, o-H and Al-Ph(1) o-H), 6.86–6.79 (m, 3H, benzyl), 6.64–6.53 (m, 5H, overlapped, 5H, N(1) Ph m-H and B-Ph(2) (m+p)-H), 6.46 (d, $J = 4.2$ Hz, 4H, N(2)Ph (o+m)-H), 6.17–6.13 (m, 1H, N(2)Ph p-H), 6.09 (t, $J = 7.1$ Hz, 1H, N(1)Ph p-H), 3.78 (d, $J = 15.3$ Hz, 1H, benzyl-CH$_3$), 3.67–3.59 (m, 4H, THF), 3.44 (d, $J = 15.3$ Hz, 1H, benzyl-CH$_3$), 3.12 (s, 8H, NBu$_4$), 1.82 (s, 8H, NBu$_4$), 1.58 (s, 8H, NBu$_4$), 1.34–1.30 (m, 3H, NBu$_4$), 0.95 (t, $J = 6.8$ Hz, 12H, NBu$_4$).

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Compound $[^{18}$LAlPh$_3$][Na$_2$(THF)$_3$]$^{32}$ was synthesized as follows: 20.71 mg ($0.072$ mmol, 78%) $^1$H NMR (400 MHz, THF-$d_8$) 7.70 (d, $J = 7.1$ Hz, 2H, p-tol-CH$_3$), 7.11–7.06 (m, overlapped, 4H, B-Ph(1) o-H), 1.58 (s, 8H, NBu$_4$), 1.34–1.30 (m, 3H, NBu$_4$), 0.95 (t, $J = 6.8$ Hz, 12H, NBu$_4$).

In 30 minutes, all the volatiles were removed under reduced pressure and the crude product was washed with hexane (3 × 2 ml). Subsequently, drying under vacuum gave the compound $[^{18}$Na$]_{[BnLAlPh_2][NBu_4]}$ as an oily green material. The oil was dissolved in a minimal amount of THF, and a layer of hexane (3 ml) was added on top of the THF solution. The layers were allowed to diffuse slowly at −30 °C. After three days, the light green crystals that had precipitated were isolated and washed with hexane (3 × 2 ml). The crystals were dried under vacuum to give the compound $[^{18}$Na$]_{[BnLAlPh_2][NBu_4]}$.
Suitable crystals of compounds \([\text{Bn}_1][\text{NBu}_4]\) and \([\text{Bn}_2][\text{Na}]\) were mounted on top of a cryoloop and transferred into the cold (100 K) nitrogen stream of a Bruker D8 Venture diffractometer. Data collection and reduction were performed using the Bruker software suite APEX2. Data collection was carried out at 100 K using Cu radiation (1.54178 Å) (for \([\text{Bn}_1][\text{NBu}_4]\)) and Mo radiation (0.71073 Å) (for \([\text{Bn}_2][\text{Na}]\)). The final unit cell was obtained from the xyz centroids of 9929 (all other degrees of freedom were optimized). Starting from the highest energy points along the scan coordinate, the transition state was optimized and confirmed by frequency analysis (number of imaginary frequencies = 0). A SAME instruction was applied for the disordered THF molecule without disorder. Some atoms in the disordered THF molecule showed non-positive definite displacement parameters when refined freely, and RIGU/DELU constraints were applied. One of the carbon atoms of the remaining Na-THF fragment was split into two disorder components, such that the two disordered THF molecules were restrained to have a similar geometry to that found for the THF molecule without disorder. Some atoms in the disordered THF molecule showed non-positive definite displacement parameters when refined freely, and RIGU/DELU instructions were applied. All of the carbon atoms of the remaining Na-THF fragment was split into two disorder components, which refined to an s.o.f. of 0.73 for the major component. The hydrogen atoms were generated by geometrical considerations, constrained to idealized geometries and allowed to ride on their carrier atoms with an isotropic displacement parameter related to the equivalent displacement parameter of their carrier atoms. Crystal data and details on data collection and refinement are presented in Table S1.†
