Calcium formamidinate derivatives by hydride insertion of carbodiimides†

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The carbodiimides, Ar/RN==C==NR (Ar = 2,6-di-isopropylphenyl (Dipp), 4-methylphenyl; R = tert-butyl, cyclohexyl, iso-propyl), react with the polar Ca—H bonds of the dimeric β-diketiminato calcium hydride, [(BDI)CaH]2 (BDI = HC(MeCN-2,6-i-Pr2CaH3)2), to provide a series of heteroleptic calcium formamidinate derivatives. While reactions with carbodiimides bearing more sterically demanding substituents (Dipp, t-Bu, Cy) are straightforward, and provide four-coordinate compounds, less bulky iso-propyl or 4-methylphenyl substitution results in the sequestration of a further equivalent of the carbodiimide and the isolation of a heteroleptic hydride-bridged dinuclear species. This latter observation is suggested to be a reflection of the robust dimeric constitution of the calcium hydride reagent and the reduced steric demands of the di-para-tolyl carbodiimide reagent.

Introduction

Amidinate anions are among the most widely applied ligands in contemporary coordination and organometallic chemistry.1,2 Organic amidines are readily available by a variety of routes and are easily metallated by organometallic or amide bases derived from more electropositive elements. These species may, in turn, be applied in onward metathesis reactions with metal salts to provide a panoply of derivatives that encompasses the entirety of the periodic table. A further reliable route to C-alkyl amidinates involves the insertion of a carbodiimide, RN==C==NR (R = alkyl or aryl), into the M–C bond of a polar organometallic derivative (Scheme 1(a)). In cases where the parent formamidinate species are required, analogous access necessitates the availability of a suitably reactive, molecular metal hydride. Documented examples of this latter process include rare earth metal (Y,3–5 La6 and Lu7), transition metal (Fe8–9, Zr,9, Nb,10 and Re11), p-block (B,12–15 Al,16,17 Zn,18–20 Ga16 and Sn21) and s-block metal (Li22 and Mg23–24) hydrides (Scheme 1(b)).

A variety of calcium amidinate species have been synthesised either by salt metathesis of a pre-formed anion with CaI2, by deprotonation of an amidine by [Ca[N(SiMe3)2]2]2, or by carbodiimide insertion into the polar Ca–C bonds of a terminal acetylide.25–34 Of most relevance to the current work, four heteroleptic calcium formamidinate derivatives, [Ca[Ar/RN]2CH2][THF] (Ar = 2,6-di-isopropylphenyl (Dipp), o-methylphenyl, 2,6-dimethylphenyl; R = i-Pr) have previously been reported. These latter compounds, however, were synthesised by a redox transmetallation process between the neutral amidine and elemental calcium, either in the presence of Hg(C6F5)2 or in liquid ammonia.35–37 Insertion reactions of carbodiimides with calcium hydride bonds are yet to be reported, a lacuna which may be traced to the relative dearth and only recent provenance of appropriately stable and soluble molecular calcium hydrides.32,38–48

A seminal report in s-block chemistry was provided by Harder’s molecular β-diketiminato calcium hydride complex, [[(BDI)CaH][THF]] (BDI = HC(MeCN-2,6-i-Pr2CaH3)2; 1).40 Although compound 1 and related species have been shown to react directly with the polar multiple bonds of small molecules containing C==O, C==O, C==N and C==N functionalities to provide the corresponding alkoxide and amide derivatives,41,49–53 its reactivity with carbodiimides appears to have escaped attention. This observation is particularly surprising given the ability of sterically demanding C-alkyl amidinate ligands to impart sufficient kinetic stability in their own right to stabilise hydrido derivatives of the heavier alkaline earth elements.32,54 We have recently reported that a non-THF solvated analogue of compound 1, compound 2 (see Scheme 2), reacts readily with the C==C bonds of terminal n-alkenes to provide calcium alkyl species which display sufficient nucleophilicity to allow the direct alkylation of benzene and the heterolytic dissociation of dihydrogen under very mild conditions.55,56 In this contribution, we continue to assay the chemistry of compound 2 and report the outcome of its reactions with a representative palette of N,N-dialkyl- and N,N-dialkylcarbodiimides.

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Results and discussion

An initial NMR scale reaction between compound 2 and DippN=CN=NDipp afforded clean access to the expected formamidinate, [(BDI)Ca(DippNC(H)NDipp)] (3, Scheme 2). The formation of compound 3 was most clearly evidenced by the appearance of a diagnostic formamidinate methine singlet resonance at δ 8.07 ppm that integrated 1H:1H with a BDI γ-CH methine signal at δ 4.83 ppm in the resultant 1H NMR spectrum. Cooling a saturated pentane solution of 3 to −35 °C afforded single crystals from which its solid state structure was determined by X-ray diffraction analysis (Fig. 1).

The identity of the formamidinate anion in 3 is clearly indicated by the elongation of the C30–N3 and C30–N4 bonds [1.3250(16), 1.3217(16) Å] and the narrowing of the N4–C30–N3 angle [120.51(11)°] in comparison to the respective measurements in the carbodiimide starting material [1.213(2), 1.221(2) Å and 169.3(2)°]. Despite the steric constraints imposed by the flanking di-isopropylphenyl groups in both the formamidinate and BDI anions, Ca1 lies in a distorted pseudo-tetrahedral environment provided solely by the four nitrogen donors and is effectively coplanar with both of the ligands [Ca1 is displaced from the mean plane defined by N3, N4, C30 by 0.075 Å and from N1, N2, C1, C2, C3, C4, C5 by 0.06 Å]. The planes defined by the chelated ligands are not orthogonal, however, and adopt a mutual orientation which is 49.49° away from coplanarity. The Ca–N distances to the formamidinate ligand [Ca1–N3 2.3877(10), Ca1–N4 2.3875(10) Å] fall at the short end of the range reported for terminal calcium amidinates, 2.306(5)–2.525(3) Å.

The reactivity of compound 2 was then extended to the N,N-dialkylcarbodiimides, t-BuN=CN=t-Bu and CyN=CN=Cy, to afford the corresponding formamidinate derivatives, [(BDI)Ca(t-BuNC(H)Nt-Bu)] and [(BDI)Ca(CyNC(H)NCy)], compounds 4 and 5, respectively (Scheme 2). The formation of both 4 and 5 was again confirmed by the appearance of distinctive formamidinate methine resonances at δ 8.51 and 8.08 ppm in the resultant...
N3
6
observation was resolved through the crystallisation of com-

equivalents of the carbodiimide reagent. The origin of this

from a saturated hexane solution at

reported
tert

dence with the analogous measurements of the previously

N4 bonds [1.321(2), 1.322(2) Å], which are in close correspon-
diagram environment of the calcium centre in the structure of

spectra. We are, thus, satisfied to ascribe the unusual coordi-
nation analysis revealed that the formamidinate complex crystal-
lises with the coordination of an equivalent of i-PrN=C=Ni-Pr
to afford a 5-coordinate calcium complex, [(BDI)Ca(iPrNC(H)

NiPr)(i-PrNCNi-Pr)] (Fig. 3).

The C30–N3 and C30–N4 bonds [1.324(3), 1.317(3) Å] and

the N4–C30–N3 angle [121.3(2)°] in compound 6 are compar-
able with the corresponding measurements within Gordon’s

homoleptic calcium N,N-di-isopropyl formamidinate complex,

[(iPrNCH(NiPr)2)Ca]2, [1.28(3)–1.328(6) Å and 118.6(4)–

122.3(4)°].37 The calcium centre of 6 is five-coordinate with four nitrogen donor interactions provided by the formally

1H NMR spectra. The structure of compound 4 was also deter-


dined by single crystal X-ray diffraction analysis of crystals grown

from a saturated hexane solution at ~35 °C (Fig. 2a).

In a similar manner to compound 3, the formation of a for-

mamidinate anion in 4 is indicated by the C34–N3 and C34–

N4 bonds [1.321(2), 1.322(2) Å], which are in close correspond-

dence with the analogous measurements of the previously

reported tert-butyl formamidinate, [(t-BuNC(H)Ni-Bu)2(B(C6F5)2)]

[1.321(2), 1.314(2) Å].13 While the calcium centre lies only

0.048 Å out of the mean plane defined by N3, N4 and C30 and

is, thus, effectively coplanar with the formamidinate anion, in

contrast to compound 3 and presumably as a consequence of the
differing steric demands of the N-alkyl substituents, the caleium centre resides some 1.37 Å out of the N1, N2, C1, C2,

C3, C4, C5 plane of the BDI ligand. As a result, albeit the four

Ca–N interactions [2.3462(11), 2.3420(11), 2.3242(13) and

2.3867(11) Å] lie within the normal expected range, the two

chelated ligands adopt an unusual relative disposition. Most

notably, the angle between the mean planes containing

[Ca1, N1, N2] and [N1, N2, C1, C2, C3, C4, C5] define an angle

of 127.4°, which leaves the calcium centre exposed with what

appears to be an otherwise vacant coordination site (Fig. 2b).

No similar asymmetry of the formamidinate or BDI environ-

ments, however, could be discerned in the solution NMR

spectra. We are, thus, satisfied to ascribe the unusual coordi-
nation environment of the calcium centre in the structure of 4
to a solid state packing effect.

Reduction of the steric bulk of the carbodiimide reaction

tartner to i-PrN=C=Ni-Pr, also afforded the corresponding

formamidinate, compound 6, as evidenced by the appearance

of the formamidinate methine signal at δ 8.27 ppm in its

1H NMR spectrum. In this case, however, complete conversion

of the hydrogen attached to C34, have been removed for clarity. (b) Space filling representation of compound 4 in the same orientation highlighting the exposed environment of the Ca1 centre. Selected bond distances (Å) and angles (°): Ca1–N1 2.3462(11), Ca1–N2 2.3420(11), Ca1–N3 2.3242(13), Ca1–N4 2.3867(11), N3–C34 1.321(2), N4–C34 1.322(2), N2–Ca1–N1 82.56(4), N3–Ca1–N4 58.57(4), N1–Ca1–N3 116.83(4), N3–Ca1–N2 112.61(4), N3–C34–N4 112.42(13).

Fig. 2 (a) ORTEP representation of compound 4 with thermal ellipsoids at 25%. Hexane solvent, disordered carbon atoms and H atoms, except for the hydrogen attached to C34, have been removed for clarity. (b) Space filling representation of compound 4 in the same orientation highlighting the exposed environment of the Ca1 centre. Selected bond distances (Å) and angles (°): Ca1–N1 2.3462(11), Ca1–N2 2.3420(11), Ca1–N3 2.3242(13), Ca1–N4 2.3867(11), N3–C34 1.321(2), N4–C34 1.322(2), N2–Ca1–N1 82.56(4), N3–Ca1–N4 58.57(4), N1–Ca1–N3 116.83(4), N3–Ca1–N2 112.61(4), N3–C34–N4 121.42(13).

Fig. 3 ORTEP representation of compound 6 with thermal ellipsoids at 25%. H atoms, except for the hydrogen attached to C30, have been removed for clarity. Selected bond lengths (Å) and bond angles (°). Ca1–N1 2.352(2), Ca1–N2 2.3691(19), Ca1–N3 2.407(2), Ca1–N4 2.380(2), Ca1–N5 2.538(2), N3–C30 1.324(3), N4–C30 1.317(3), N5–C37 1.235(4), N6–C37 1.219(5), N1–Ca1–N2 81.17(7), N4–Ca1–N3 57.50(7), N1–Ca1–N3 116.05(8), N1–Ca1–N4 107.04(7), N1–Ca1–N5 95.01(7), N2–Ca1–N3 141.17(8), N2–Ca1–N4 108.32(7), N2–Ca1–N5 98.19(7), N3–Ca1–N5 90.09(7), N4–C30–N3 121.3(2), N6–C37–N5 168.8(6).
anionic formamidinate and BDI ligands [2.352(2)−2.407(2) Å]. The coordination sphere of the calcium is completed by a longer nitrogen interaction [2.538(2) Å] from the molecule of i-PrN=C≡Ni-Pr, that effectively occupies the ‘open’ coordination site of the calcium atom that was a notable feature of the structure of compound 4. This observation is underscored by a calculated τ value of 0.09, which is indicative of a coordination geometry approaching square pyramidal with the N1 atom of the BDI ligand providing the apex of the pyramid.59

A further reaction at room temperature with the N-aryl carbodiimide, p-tolN=C≡Np-tol, afforded a mixed formamidinate-hydride complex, [[BDI][H][p-tolNC(H)Np-tol]] (7, Scheme 2), irrespective of the reaction stoichiometry. The formation of compound 7 was clearly apparent from the appearance of unique formamidinate and BDI methine signals (δ 8.04, 4.95 ppm), which were observed to emerge alongside a further singlet resonance that was assigned to the maintenance of a Ca–H environment at δ 4.41 ppm and which integrated in a respective 1H: 2H: 1H ratio in the resultant 1H NMR spectrum. The structure of compound 7 was confirmed by a further crystal X-ray diffraction analysis of crystals formed by cooling a saturated hexane solution to −35 °C (Fig. 4).

The asymmetric unit of compound 7 comprises two similar di-calcium molecules in which dimerization is propagated through a single amidinate anion, which spans both Ca centres, and a residual μ2-Ca−H−Ca bridging interaction. The C–N bonds [C37–N3 1.321(2), C37–N4 1.321(2), N9–C110 1.321(2), N10–C110 1.324(2) Å] of the formamidinate anions within both entities of the asymmetric unit are closely comparable with the analogous measurements within the previously reported magnesium complex of the same ligand, [[p-tolNC(H)Np-tol)2Mg(THF)2] [1.312(3)−1.323(3) Å].64 In contrast, the chelating disposition adopted by the ligands about magnesium in this previously described complex subtend significantly narrower N–C–N angles [116.0(3), 117.9(3)°] than either of the bridging formamidinate anions in the structure of 7 [N4–C37–N3, 124.82(13); N9–C110–N10 125.05(13)°]. The bridging mode adopted also contrasts significantly with that observed in the only other amidinate-bridged calcium complexes, the aforementioned [[iPrNC(H)iPr]Ca]2 and a similarly homoleptic species comprising a propargyl-substituted anion.37,61 Whereas the bridging amidinate nitrogen atoms in both of these previously reported species are 4-coordinate, engaging with both calcium centres in a μ2-Ca−N−Ca fashion, N3 and N4 within the structure of compound 7 interact solely with Ca1 and Ca2, respectively. Despite this reduction in the number of nitrogen-to-element bonds from 4 to 3, the resultant Ca1–N3 and Ca2–N4 distances [2.4055(12), 2.4207(12) Å] in 7 are not notably shorter than the ranges established previously [2.400(3)−2.650(3) Å]. Both calcium centres, thus, reside in four-coordinate environments, which are completed by two Ca–N interactions from the BDI ligands and the bridging hydride. Attempts to react the remaining hydride of compound 7 with a further equivalent of p-tolN=C≡Np-tol at elevated temperatures were unsuccessful and merely resulted in ligand redistribution to various species, including the known bis-β-diketiminate derivative, [Ca(BDI)]2.62,63 We have previously noted that the reactivity of compound 2 with the C=C bonds of terminal alkynes occurs with the retention of the relatively robust hydrido-bridged dimer.55,56 The formation of compounds 3–6 necessarily ensues with the generation of mononuclear calcium species, which is most likely a reflection of the significant steric bulk of the nitrogen-bound hydrocarbon residues. Notably, inspection of the resultant 1H NMR spectrum immediately after addition of a single further equivalent of i-PrN=C≡Ni-Pr to an isolated sample of compound 7 demonstrated the disappearance of the residual hydride resonance at δ 4.41 ppm and the generation of an equimolar mixture of compound 6 and a new amidinate species, which we assign as the further amidinate complex [[BDI][Ca(p-tolNC(H)Np-tol)]] (Fig. S11†). Although not pursued further, we, thus, tentatively suggest that the dinuclear constitution of compound 7 is a consequence of both the reduced steric demands of the N-p-tolyl substituents and the lower basicity of p-tolN=C≡Np-tol. The consequent inability of a single equivalent of p-tolyl carbodiimide to induce dimer fragmentation during the initial insertion process, thus, restricts access to the remaining Ca–H bond.

In conclusion, reactions between the dimeric calcium hydride, [[BDI]CaH]2 and representative carbodiimide small molecules are facile and result in the generation of well-defined formamidinate products. X-ray crystallographic studies reveal that the resultant calcium coordination environments and coordination numbers are strongly dependent upon the relative steric demands of the N-aryl or N-alkyl substituents.
Experimental details

General considerations and starting materials

All manipulations were carried out using standard Schlenk line and glovebox techniques under an inert atmosphere of argon. NMR experiments were conducted in J Young tap NMR tubes prepared and sealed in a Glovebox. NMR spectra were collected on a Bruker AV300 spectrometer operating at 300.2 MHz (1H), 75.5 MHz (13C) or an Agilent ProPulse spectrometer operating at 500 MHz (1H), 126 MHz (13C). The spectra were referenced relative to residual protio solvent resonances. Solvents (toluene, hexane, pentane) were dried by passage through a commercially available (MBrayn) solvent purification system, under nitrogen and stored over molecular sieves. Calcium iodide molecular sieves. C6D6 was purchased from Sigma-Aldrich Corporation and used without further purification.

[(BDI)CaH]2 (2) was synthesised by a literature procedure.55

[(BDI)Ca(DippNC(H)NDipp)] (3)

Toluene (10 ml) was added to 2 (0.92 g, 1.00 mmol) and DippNC≡C=NDipp (0.73 g, 2.01 mmol) then stirred overnight at room temperature. The resulting solution was evaporated to dryness, redissolved in hexane (10 ml), cannula filtered and concentrated. Colourless crystals were deposited overnight at −35 °C and collected via cannula filtration to yield compound 3. (0.94 g, 57%).1H NMR (500 MHz, benzene-d6) δ 8.07 (s, 1H, NC(CH3)H), 7.06–7.11 (m, 6H, Ar-H, C(CH3)3), 2.30 (hept, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.32 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.10 (br d, JHH = 6.8 Hz, 24H, CH(CH3)2) ppm. Elemental analysis for C54H76CaN4: C, 78.97; H, 9.92; N, 6.82%. Found: C, 78.97; H, 9.92; N, 6.82%.

[(BDI)Ca(t-BuNC(H)t-Bu)] (4)

Toluene (10 ml) was added to 2 (0.92 g, 1.00 mmol) and t-BuNC≡C≡Nt-Bu (386 µl, 2.00 mmol) then stirred at room temperature for ca. 1 hour. The resulting solution was evaporated to dryness, redissolved in hexane (10 ml), cannula filtered and concentrated. Colourless crystals were deposited overnight at −35 °C and collected via cannula filtration to yield compound 4. (0.55 g, 45%).1H NMR (500 MHz, benzene-d6) δ 8.51 (s, 1H, NC(CH3)H), 7.21–7.11 (m, 6H, Ar-H), 4.87 (s, 1H, NC(CH3)H), 3.26 (hept, JHH = 6.8 Hz, 4H, CHF(CH3)2), 1.74 (s, 6H, NC(CH3)H), 1.29 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.27 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 0.99 (s, 18H, NC(CH3)3) ppm. 13C{1H} NMR (126 MHz, benzene-d6) δ 166.1 (NC(CH3)2), 164.3 (NC(CH3)N), 146.8 (Cpreno), 140.4 (Cortho), 124.6 (Cmeta), 123.9 (Cmeta), 91.4 (NC(CH3)CH), 51.3 (NC(CH3)3), 32.5 (NC(CH3)3), 28.8 (CH(CH3)2), 25.8 (CH(CH3)2), 24.8 (NC(CH3)CH), 24.3 (CH(CH3)2) ppm. Elemental analysis for C43H64CaN4: C, 72.62; H, 9.92%; N, 6.82%. Found: C, 72.62; H, 9.92; N, 6.82%.

[(BDI)Ca(CyNC(H)NCyl)] (5)

Toluene (10 ml) was added to 2 (0.92 g, 1.00 mmol) and CyNC≡C=NCy (0.41 g, 1.99 mmol) then stirred at room temperature for ca. 1 hour. The resulting solution was evaporated to dryness, redissolved in hexane (10 ml), cannula filtered and concentrated. Colourless crystals were deposited overnight at −35 °C and collected via cannula filtration to yield compound 5. (0.62 g, 47%).1H NMR (500 MHz, benzene-d6) δ 8.08 (s, 1H, NC(CH3)H), 7.18–7.11 (m, 6H, Ar-H), 4.94 (s, 1H, NC(CH3)CH), 3.29 (hept, JHH = 6.8 Hz, 4H, CH(CH3)2), 2.57–2.51 (m, 2H, NC(CH3)H), 1.77 (s, 6H, NC(CH3)CH), 1.72–1.64 (m, 4H, NHC(CH3)2), 1.61–1.54 (m, 4H, NHC(CH3)2), 1.51–1.43 (m, 4H, NHC(CH3)2), 1.29 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.26 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.14–1.03 (m, 4H, NHC(CH3)2), 0.92–0.81 (m, 4H, NHC(CH3)2) ppm. 13C{1H} NMR (126 MHz, benzene-d6) δ 168.7 (NC(CH3)N), 165.9 (NC(CH3)CH), 146.4 (Cpreno), 141.5 (Cortho), 124.6 (Cmeta), 93.0 (NC(CH3)CH), 61.7 (NC(CH3)CH), 37.2 (NC(CH3)CH), 28.7 (CH(CH3)2), 26.4, 26.3 (NC(CH3)2), 25.6 (CH(CH3)2), 24.5 (NC(CH3)CH), 24.3 (CH(CH3)2) ppm. Elemental analysis for C42H58CaN2: C, 75.85; H, 9.70; N, 8.42%. Found: C, 75.71; H, 9.87, N, 8.37%.

[(BDI)Ca(iprNC(H)NiPr)(iprNCNiPr)] (6)

Toluene (10 ml) was added to 2 (0.92 g, 1.00 mmol) and iprNCNiPr (621 µl, 4.01 mmol) then stirred at room temperature for ca. 1 hour. The resulting solution was evaporated to dryness, redissolved in hexane (10 ml), cannula filtered and concentrated. Colourless crystals were deposited overnight at −35 °C and collected via cannula filtration to yield compound 6. (0.56 g, 39%).1H NMR (500 MHz, benzene-d6) δ 8.27 (s, 1H, NC(CH3)H), 7.18–7.07 (m, 6H, Ar-H), 4.92 (s, 1H, NC(CH3)CH), 3.44 (hept, JHH = 6.8 Hz, 4H, CH(CH3)2), 3.14 (hept, JHH = 6.4 Hz, 2H, 2H, NC(CH3)H), 3.09 (hept, JHH = 6.4 Hz, 2H, NC(CH3)H), 1.76 (s, 6H, NC(CH3)CH), 1.32 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.29 (d, JHH = 6.8 Hz, 12H, CH(CH3)2), 1.09 (d, JHH = 6.8 Hz, 12H, NC(CH3)H), 0.86 (d, JHH = 6.4 Hz, 12H, NC(CH3)H) ppm. 13C{1H} NMR (126 MHz, benzene-d6) δ 169.0 (NC(CH3)N), 165.4 (NC(CH3)CH), 148.0 (Cpreno), 141.8 (Cortho), 124.1 (Cmeta), 93.6 (NC(CH3)CH), 53.9 (NC(CH3)H), 49.5 (NC(CH3)H), 28.3 (CH(CH3)2), 26.8 (NC(CH3)H), 25.8 (CH(CH3)2), 24.9 (NC(CH3)CH), 24.6 (CH(CH3)2), 24.2 (NC(CH3)H), 24.3 (CH(CH3)2) ppm. Elemental analysis for C42H58CaN2: C, 72.62; H, 9.92; N, 11.82%. Found: C, 72.43; H, 9.95, N, 11.69%.
Conflicts of interest

There are no conflicts of interest to declare.

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