Synthesis and reduction chemistry of mixed-Lewis-base-stabilised chloroborylenes†

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The one-electron reduction of (CAACMe)BCl3 (CAACMe = 1-(2,6-diisopropylphenyl)-3,3,5,5-tetramethylpyrrolidin-2-ylidene) yields the dichloroboryl radical [(CAACMe)BCl2]. Furthermore, the twofold reduction of (CAACMe)BCl3 in the presence of a range of Lewis bases (L = CAACMe, N-heterocyclic carbene, phosphine) yields a series of doubly base-supported (CAACMe)LBCl chloroborylenes, all of which were structurally characterised. NMR and UV-vis spectroscopic and electrochemical data for (CAACMe)LBCl show that the boron centre becomes more electron-rich and the HOMO–LUMO gap widens as L becomes less π-accepting. A [(CAACMe)BCl2]− boryl anion coordination polymer was isolated as a potential intermediate in these reductions. In most cases the reduction of the chloroborylenes resulted in the formation of the corresponding hydroborylenes or derivatives thereof, as well as ligand C–H activation products.

Introduction

While the field of boron chemistry has long been dominated by the naturally occurring +3 oxidation state of this electron-deficient element, the last decade has seen tremendous advances in the accessibility of lower oxidation state boron species. Among these, borylenes (:BR), which feature a highly reactive boron(i) centre with two empty p orbitals, have attracted a lot of attention owing to their similarities with organic carbene.1 The parent borylene, :BH, was first generated in the 1930s by exposure of a BCl3/H2 gas stream to an electrical discharge,2 but is so short-lived that it can only be studied spectroscopically. In contrast, boron fluoride (:BF), which calculations have shown to be the most stable diatomic boron fluoride (:BF), which have been studied spectroscopically.

Like carbenes, borylenes may be stabilised by adduct formation with electron-rich metal centres. Since the report by our group of the first stable metal borylene in 1995,4 this field has rapidly expanded and a number of bridging haloborylene complexes have been structurally characterised,10 while terminal fluoroborylene complexes have only been observed in the matrix at 6 K.11

It was not until 2011 that the first stable metal-free borylene, (CAAC)2BH (I, CAAC = 3,3-dimethyl-2-(2,6-diisopropylphenyl)-2-azaspiro[4,5]dec-1-ylidene) was isolated by Bertrand...
and co-workers. Its stability is owed to the two strongly σ-donating/π-acidic cyclic (alkyl)(amino)carbene (CAAC) ligands, which compensate for the build-up of electron density at the boron(I) centre by efficient π delocalisation to form two partial B–CAAC π bonds. In recent years the field of metal-free borylenes has greatly expanded, recently culminating in the synthesis of the first bis(borylene)-stabilised borylene, (TMP) \( B(\text{TMP})_2 \) (TMP = 2,2,6,6-tetramethylpiperidinyl). In order to vary the electronic properties at sp\(^2\)-hybridised borylene centres and tune their reactivity, various synthetic routes toward mixed-base-stabilised L\(_1\)L\(_2\)BR borylenes have been developed, in which L\(_1\) is typically a CAAC ligand and L\(_2\) is a less π-accepting donor ligand. The simplest approach, which consists in adding a suitably small Lewis base (CO, CN\(_t\)Bu, IMeMe\(_{1,3,4,5}\)-tetramethylimidazol-2-ylidene) to an isolated phosphine-supported borylene, compound \( \text{II-PEt}_3 \) (Fig. 1), from the tetrameric precursor \([\text{CAACMe}_2\text{B}]_2\). While facile, this method remains limited by the dearth of isolable dicoordinate borylenes. A number of L(CO)\(_2\)BR borylenes (L = CAAC, CNR, NHC) have also been obtained by the addition of strongly σ-donating ligands to metal-carbonyl-bound borylenes. This is the case of \( \text{III-CO} \), generated by the addition of CAACMe to \([\text{Me}_3\text{P}][\text{CO})_4\text{Fe}]\) (Dur = 2,3,5,6-tetramethylphenyl), and which may itself undergo photolytic CO–L ligand exchange to yield \( \text{III-L} \) (L = CNBu, pyridine, IMe = 1,3-dimethylimidazol-2-ylidene). A more modular route to L\(_1\)L\(_2\)BR borylenes involves the introduction of both donors at the borane stage, prior to reduction. Thus compounds \( \text{IV-L} \) were obtained via multi-step syntheses from a simple (CAAC)\(_2\)BH\(_3\) precursor undergoing (a) replacement of one hydride by a trflate group, (b) displacement of trflate by L, (c) replacement of a second hydride by trflate, and finally, (d) reduction to \( \text{IV-L} \). A much simpler approach, involving the twofold reduction of a (CAAC)\(_2\)BR\(_2\) precursor in the presence of a second donor was used for the synthesis of the first phosphine-supported borylene, compound \( \text{II-PEt}_3 \).

Despite all these advances, isolable metal-free haloborylenes, which are of particular interest because of their potentially reactive B–X bond, remain rather elusive. Only one bromoborylene, compound \( \text{V} \), supported by a complex, chelating carborane-bridged bis(silylene) ligand, has been reported thus far and has already shown promising reduction and ligand exchange reactivity, providing access to the first borylene cation, \( \text{VI} \) (Scheme 2).

In this contribution we present a facile route towards chloroborylenes of the form (CAAC)L\(_2\)B\(_2\)Cl (L = CAAC, NH\(_2\), phosphine) and examine the steric and electronic effects of L on the structural and spectroscopic properties of these borylenes. We also present the first stable dichloroboryl radical and a dichloroboryl anion, both likely intermediates in the formation of these borylenes. Finally, we show that further reduction of these species leads to hydrogen abstraction from the solvent, ligand metalation and C–H activation products.

**Results and discussion**

The reduction of \([\text{CAACMe}_2]\text{BCl}_2\) (1) with one molar equivalent of Li in THF at room temperature yielded a pale green solution, which displayed no \(^{11}\text{B}\) NMR resonance (Scheme 3). Solvent removal and crystallisation of the hexane extract at \(-25^\circ\text{C}\) provided a small crop of orange crystals, which X-ray crystallographic analysis showed to be the \([\text{CAACMe}_2]\text{BCl}_2\) radical, compound 2 (Fig. 2a).

The geometry around the boron atom is trigonal planar (\( \Sigma \angle \text{B} 359.96(13)^\circ \), while the B1–C1 distance (1.498(3) Å) is indicative of a double bond and significantly shorter than in precursor 1 (1.644(2) Å, see Fig. S41 in the ESI† for the solid-state structure of 1). This has been observed previously for other CAAC-supported boryl radicals and is indicative of the delocalisation of the unpaired electron over the B–C–N π framework.\(^{21}\) The EPR spectrum of 2 shows a very broad singlet centred at g\(_{\text{iso}}\) = 2.003 (Fig. 2c), unlike its relative \([\text{CAACMe}_2]\text{B(Dur)Cl}_2\), which displays a triplet from the hyperfine interaction with the \(^{14}\text{N}\) nucleus (\( a(14\text{N}) = 19 \text{MHz} \)). A simulation of

![Scheme 2](image-url)  
**Scheme 2** Halide abstraction at bromoborylene V.

![Scheme 3](image-url)  
**Scheme 3** One-electron reduction of 1.
the EPR spectrum of 2 provided the following hyperfine coupling parameters: $a^{\text{14N}} = 17.3 \text{ MHz}$ and $a^{\text{11B}} = 6.70 \text{ MHz}$, the latter being significantly higher than that in the simulated EPR spectrum of $[(\text{CAACMe})\text{B(Dur)}]^{+}$ (ca. 2.7 MHz),

\[ a^{(\text{14N})} = 17.3 \text{ MHz} \]

\[ a^{(\text{11B})} = 6.70 \text{ MHz} \]

which contributes to the strong signal broadening and loss of resolution. The spin density distribution in 2 was further analysed by density functional theory (DFT) calculations. The unpaired electron is delocalised over the B–C–N π system with atomic spin densities of 0.39, 0.33, and 0.24 on C1, B1, and N1, respectively (Fig. 2d). The lower hyperfine coupling with $^{11}$B, as well as the lower spin density at boron for $[(\text{CAACMe})\text{B(Dur)}]^{+}$ (0.277)\(^{111}\) versus 2 are due to the delocalisation of spin density to the duryl group in the former, whereas no such delocalisation to the chloride ligands is observed in 2. The B–C π-bonding character of the SOMO of 2 (Fig. S43) results in a partial double bond, which is reflected in a B–C Wiberg bond index of 1.16 (Fig. 2b).

Cyclic voltammetry performed on 2 showed a single irreversible reduction wave at $E_{pc} = -2.35 \text{ V}$ (against the ferrocene (Fc/Fc\(^+\)) couple) and no oxidation event, unlike $[(\text{CAACMe})\text{B(Dur)}]^{+}$, for which a first reversible reduction wave (\(E_{pc} = -2.03 \text{ V}\)) and an irreversible oxidation wave (\(E_{oc} = -0.53 \text{ V}\)) were observed.\(^{22b}\) This implies that a $^+$ cation is unlikely to be chemically accessible, whereas further chemical reduction of 2 should be achievable with a suitable reducing agent.

The room-temperature reduction of 1 in the presence of 1 equivalent of CAAC\(^{\text{Me}}\) with 2.5 molar equiv. of KC\(_8\) in benzene yielded a strikingly purple-blue reaction mixture, the colour of which intensified over a period of five hours (Scheme 4).

Removal of volatiles, extraction with hexanes and subsequent solvent removal yielded a crude purple solid displaying a broad $^{11}$B NMR resonance at 18.7 ppm and a single set of $^1$H NMR CAAC\(^{\text{Me}}\) resonances, indicating a symmetrical compound. Crystallisation from pentane at $-25 \degree\text{C}$ over a period of one week yielded large purple crystals suitable for X-ray crystallographic analysis, which provided the structure of a doubly CAAC\(^{\text{Me}}\)-stabilised chloroborylene, $3\text{-CAAC}\text{Me}^{2}$ presenting a trigonal planar boron centre ($\Sigma_{\text{vB}} 359.97(15)$, Fig. 3). The molecule is C\(_2\) symmetric and the two B1–C1 and C1–N1 bonds (1.530(2) and 1.389(2) \(\AA\), respectively) are intermediate between typical lengths of sp\(^2\)-sp\(^3\) single (B–C 1.56; C–N 1.47 \(\AA\))\(^{22}\) and double bonds (B–C 1.44; C–N 1.35 \(\AA\))\(^{24,25}\) suggesting delocalisation of the borylene lone pair over the entire N–C–B–N π framework. The compound is thereby reminiscent of Bertrand’s parent borylene 1, the B–C bonds of which are slightly shorter (1.5175(15) and 1.5165(15) \(\AA\))\(^{13}\) presumably because of the smaller size and lower electronegativity of the hydride compared to that of the chloride ligand.

The analogous reductions of 1 in the presence of one molar equivalent of IMes (1,3-bis(2,4,6-trimethylphenyl)imidazol-2-ylidene) or SIMes (1,3-bis(2,4,6-trimethylphenyl)-4,5-dihydroimidazol-2-ylidene) with 2.5 molar equivalents of KC\(_8\) in benzene similarly yielded deep-pink reaction mixtures from which the corresponding mixed-base chloroborylenes 3-L (L = IMes, SIMes) were extracted in moderate to good yields (58–76%). For smaller L ligands, such as IMe\(^{\text{Me}}\), PEt\(_3\) or PMe\(_3\) using a 1 : 1 1-to-1 L ratio for the reduction resulted in mixtures of the desired chloroborylene 3-L and (CAAC\(^{\text{Me}}\))\(_2\)BH ($\delta_{1\text{H}} 12.5 \text{ ppm}$, \textit{vide infra} compound 5-CAAC\(^{\text{Me}}\)). For PEt\(_3\) or PMe\(_3\), a fivefold excess of the phosphine could be used to obtain the mixed CAAC-phosphate-stabilised chloroborylenes in moderate isolated yields (ca. 50%), with the excess phosphine simply being removed \textit{in vacuo} prior to extraction of the product. Unlike the bis(CAAC)- or CAAC–NHC-stabilised chloroborylenes, these compounds crystallise as pale yellow solids. Finally, 3-IMe\(^{\text{Me}}\) was obtained as a bright red compound by quantitative NHC/phosphine ligand exchange with 3-PMe\(_3\) at room temperature (Scheme 5). Ligand exchange with the more sterically demanding carbene IMes, SIMes and CAAC\(^{\text{Me}}\) was also attempted but remained limited to 2–5% conversion.\(^{8}\)

From the attempted synthesis of the dimethylsulfide-stabilised borylene (CAAC\(^{\text{Me}}\))(SMe\(_2\))BCl using 20 equivalents SMe\(_2\), recrystallisation of the toluene–hexane extract at $-25 \degree\text{C}$ yielded a few isolated crystals of a linear coordination polymer of the dichloroboryl anion 4, $[((\text{CAAC}\text{Me})\text{BCl})_2\text{K}_2\text{(SMe}_2\text{)}_3\text{]}_n$.
The boron atoms in this structure are trigonal planar with respect to their attached chloride and CAACMe ligands \((\Sigma(\angle B1) 359.95(14)°)\) and present a relatively short \(\text{B–CCAAC}\) double bond \((1.427(3) \text{ Å})\) compared to other known \([\text{CAAC})\text{BXY}]^\text{−}\) boryl anions \((\text{XY} = \text{CN}, \text{H(CN)}, \text{HCl}, \text{H}_{2}\text{B}); \text{B–CCAAC} 1.432(6)\text{ Å} \text{ to } 1.473(2) \text{ Å}).\)

Each potassium cation lies in the plane of the sp²-boron atom, coordinating to both chloride ligands and bridging between three adjacent boryl anion units, once via a \(\text{Cl1–K1–Cl1'}\) bridge and once via a \(\text{Cl2–K1–Cl2'}\) bridge. The adjacent \((\text{N1,C1,B1,Cl1,Cl2,K1})\) planes form an angle of 75° between each other. The two potassium cations bridging the Cl and Cl' atoms, K1 and K1', are further bridged by one SMe₂ ligand and their coordination sphere is completed by \(\pi\)-interactions with the adjacent Dip residue. Unfortunately, due to its virtual insolubility in hydrocarbon solvents and extreme air- and moisture-sensitivity, no NMR spectroscopic or elemental analysis data could be obtained for 4. It is likely, however, that similar \([[\text{CAACMe})\text{BCl}_2]_{\text{K}}\text{L}\) coordination oligomers or polymers could be intermediates in the synthesis of the 3-L chloroborylenes shown in Scheme 4.

A comparison of all six 3-L chloroborylenes shows an increasing upfield shift in the \(^{11}\text{B}\) NMR resonances in the following order (Table 2): \(\text{CAACMe} (18.7 \text{ ppm}) > \text{SIMes} (11.9 \text{ ppm}) > \text{IMes} (8.4 \text{ ppm}) > \text{PET}_{3} (5.6 \text{ ppm}) = \text{IME}\text{Me} (5.5 \text{ ppm}) > \text{PMe}_{3} (2.8 \text{ ppm}).\) This correlates well with the trend of the overall electron-donating nature of the ligands \(\text{L}, i.e. \) their combined \(\sigma\) donor and \(\pi\) acceptor abilities, usually determined using the Tolman electronic parameter (TEP),\(^{27}\) which is defined as the \(\nu(\text{CO})\) stretching frequency of \(\text{Ni(}\text{CO})_{3}\text{L}\) complexes: the lower the TEP, the less electron-donating \(\text{L}\) is overall. While the TEP of IMeMe is yet to be determined experimentally,\(^{27}\) the trend in this table suggests a value close to 2062 cm\(^{-1}\) and similar electronic properties to PET\(_{3}\).

Moreover, for \(\text{L} = \text{carbene}\), the \(^{11}\text{B}\) NMR shifts of 3-L decrease with the \(^{77}\text{Se}\) NMR shifts for the corresponding \(\text{L} = \text{Se}\) adducts, which are a measure of the \(\pi\) acidity of the corresponding carbene: the more downfield the \(^{77}\text{Se}\) NMR chemical shift, the more \(\pi\)-accepting the carbene.\(^{28}\) Consequently, the \(^{11}\text{B}\) NMR shift of 3-L represents a reliable measure of the relative electron-donating ability of \(\text{L}\).

Another measure of the relative \(\sigma\) donor and \(\pi\) acceptor abilities of \(\text{L}\) is provided by a comparison of the X-ray structural data of 3-L (Fig. 3, Table 1). For the carbene ligands, this is linked to the degree in which the CAACMe and \(\text{L}\) frameworks, respectively, are rotated out of the plane of the borylene core, represented by the torsion angles \((\text{N1,C1,B1,Cl1})\) and \((\text{N2,C21,B1,Cl1})\), respectively, as well as the relative \(\text{B–CAAC}\) and \(\text{B–}\text{Cl}\) bond lengths. In the \(\text{C}_{2}\)-symmetric 3-CAACMe the \((\text{N1,C1,B1,Cl1})\) torsion angles are only 14.51(16)°, which enables
a large overlap between the borylene lone pair and the empty p₂ orbitals on both carbene carbon atoms. As a result, the borylene lone pair is fully delocalised over the C–C–C π orbital, which is further confirmed by the B–C bond lengths of 1.530(2) Å, which are intermediate between a single and double bond. While the (N₁,C₁,B₁,C₁₁) torsion angle remains very small for all the other derivatives (≤5°), allowing for excellent π overlap of CAACMe with the borylene lone pair, the (N₂,C₂₁,B₁,C₁₁) torsion angle increases in the following order: CAACMe (14.5(2)°) < IMes (44.4(2)°) < IMes (53.5(6)° = avg. of the two molecules present in the asymmetric unit) < IMeMe (79.3(2)°). This is in agreement with the decrease in π acidity in this ligand series (see Table 2). As the (N₂,C₂₁,B₁,C₁₁) torsion angle grows closer to orthogonality, π overlap decreases until it becomes negligible for L = IMeMe. This is also apparent in the lengthening of the B–C₁ bond from 1.530(2) Å in 3-CAACMe to 1.579(3) Å in 3-IMeMe, which suggests a typical dative bond, concomitant with a shortening of the B–Cbond to 1.440(3) Å in the IMeMe analogue, indicating a typical B=C double bond, as the entire π electron density from the borylene lone pair is employed in π backbonding from boron to CAACMe. A similar trend can also be observed in the structural data from Bertrand’s (CAAC)LBH hydroborylenes,13 and IV-L (Fig. 1)13 for which the torsion angles between the borylene plane and π framework of L increase in the order of CAACCl < BIPr < BACIPr, which fits with the decreasing π acidity of L, concomitant with a shortening of the B–C bond lengths of 1.456(2) Å (R = Et) and 1.449(2) Å (R = Me) a indicate a B=C double bond, similarly to the 3-IMeMe derivative. In the only other structurally characterised CAAC-phosphine-stabilised borylene, compound II-PMe₃ (Fig. 1), the B–C bond is slightly longer (1.484(6) Å) due to a small amount of π backbonding to the π-acidic cyano ligand.18

DFT calculations by the group of Bertrand have shown that the π-type HOMO of the (CAAC)LBH hydroborylenes I and IV-L is delocalised over the CCAAC–B=C moiety, with the extent of delocalisation increasing with the π acidity of L.13,23 Given the structural and electronic similarities of these compounds to our 3-CAACMe and 3-NHC derivatives, we limited our DFT analysis to the bonding and electronic structure of 3-PMe₃, in which the phosphine ligand acts as a pure σ donor. As expected, the HOMO of 3-PMe₃ corresponds largely to the B–C π bond, with minor C1–N1 and B–C1 π antibonding interactions (Fig. 5b). The calculated B=C Wiberg bond index of 1.53 indicates a full B=C double bond (Fig. 5a). This is also borne out by natural resonance theory (NRT), which gives the phosphonium allyldene borate as the major resonance form (38.3%, see Fig. S45 in the ESI†), as well as natural population analysis (NPA), which places a large positive charge of +1.25 on P1 and small negative charges of –0.07 and –0.19 on B1 and C1, respectively (Fig. 5a).

### Table 1 Relevant bond lengths (Å) and angles (°) for 3-L, L = CAACMe, IMes, SIMes, IMeMe, PEt₃, and PMe₃

<table>
<thead>
<tr>
<th>3-L, L</th>
<th>CAACMe</th>
<th>IMes</th>
<th>SIMes</th>
<th>IMeMe</th>
<th>PEt₃</th>
<th>PMe₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1-C1</td>
<td>1.530(2)</td>
<td>1.437(7), 1.450(7)</td>
<td>1.481(3)</td>
<td>1.440(3)</td>
<td>1.456(3)</td>
<td>1.449(2)</td>
</tr>
<tr>
<td>B1-C11</td>
<td>1.843(3)</td>
<td>1.884(5), 1.879(6)</td>
<td>1.854(2)</td>
<td>1.855(2)</td>
<td>1.855(2)</td>
<td>1.8516(17)</td>
</tr>
<tr>
<td>B1-C21</td>
<td>—</td>
<td>1.575(7), 1.570(7)</td>
<td>1.550(3)</td>
<td>1.578(3)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>B1-P1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1.912(2)</td>
<td>1.9114(17)</td>
</tr>
<tr>
<td>Σ(B1)</td>
<td>359.97(15)</td>
<td>360(4)</td>
<td>360(4)</td>
<td>360(4)</td>
<td>360(4)</td>
<td>360(4)</td>
</tr>
<tr>
<td>C1–B1–C21</td>
<td>133.9(2)</td>
<td>132.1(4)</td>
<td>128.49(17)</td>
<td>122.96(18)</td>
<td>129.06(17)</td>
<td>131.54(12)[c]</td>
</tr>
<tr>
<td>(N1,C1,B1,C11)</td>
<td>14.51(16)</td>
<td>4.9(8), 1.3(7)</td>
<td>3.2(3)</td>
<td>2.8(3)</td>
<td>3.0(3)</td>
<td>0.6(2)</td>
</tr>
<tr>
<td>(N2,C21,B1,C11)</td>
<td>44.4(2)</td>
<td>57.5(8), 49.2(6)</td>
<td>79.3(2)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

a Torsion angles. b C1–B1–C1. c The asymmetric unit contains two structurally distinct borylene molecules. d Torsion angle (N3,C21,B1,C11). e C1–B1–P1.
With the exception of the two phosphine derivatives, compounds 3-L were intensely coloured (Fig. 6, top). Furthermore, whereas the naphthyl and phosphine derivatives were highly air-sensitive, solutions of 3-CAACMe exposed to air retained their colour for several hours at room temperature, thus demonstrating the unusual stability of this species. An overlay of the UV-vis spectra of all 3-L compounds (Fig. 6, bottom) shows that the wavelength of the most red-shifted absorption band maximum decreases in the order of L = CAACMe (556 nm, purple-blue) > IMes (550 nm, purple-pink) > SIMes (538 nm, pink) > IMeMe (470 nm, orange-red) > PMe3 (350 nm, pale yellow) > PET3 (336 nm, pale yellow). A TD-DFT study of our tetrameric cyanoborylene, [(CAACMe)B(CN)]4, had shown that its tetrameric cyanoborylene, [(CAACMe)B(CN)]4, had shown that its,

### Table 2

<table>
<thead>
<tr>
<th>L</th>
<th>¹¹B NMR shift (ppm) of 3-L</th>
<th>TEP of L⁻²⁻ (cm⁻¹)</th>
<th>⁷⁷Se NMR shift (ppm) of L=Se</th>
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<tbody>
<tr>
<td>CAACMe</td>
<td>18.7</td>
<td>2046.0</td>
<td>492</td>
</tr>
<tr>
<td>SIMes</td>
<td>11.9</td>
<td>2051.5</td>
<td>116</td>
</tr>
<tr>
<td>IMes</td>
<td>8.4</td>
<td>2050.7</td>
<td>35</td>
</tr>
<tr>
<td>PET3</td>
<td>5.6</td>
<td>2061.7</td>
<td>n.a.</td>
</tr>
<tr>
<td>IMeMe</td>
<td>5.5</td>
<td>—</td>
<td>3</td>
</tr>
<tr>
<td>PMe3</td>
<td>2.8</td>
<td>2064.1</td>
<td>n.a.</td>
</tr>
</tbody>
</table>

*Only a calculated TEP of 2051.7 cm⁻¹ has been reported for IMeMe.²⁷a*
identity of 5-CAAC\textsuperscript{Me} was further confirmed by X-ray crystallographic analysis (see Fig. S42 in the ESI). The reduction of 3-IMes with excess Li sand in THF also proceeded cleanly to a red-orange compound presenting a broad \(^{11}\text{B}\) NMR resonance at 2.8 ppm (compound 6, Scheme 6b). X-ray crystallographic analysis of 6 revealed a planar hydroborylene (\(\Sigma(\angle B1) 359.8(8)\)) stabilised on the one side by a strongly \(\pi\)-accepting CAAC\textsuperscript{Me} ligand (B1–C1 1.454(3) Å) and on the other side by a \(\sigma\)-donating IMes ligand (B1–C21 1.583(4) Å) lithiated at the backbone C4 position, the coordination sphere of the lithium atom being completed by three THF residues (Fig. 7, top). Such metatation of unsaturated NHC backbones in the presence of strong, non-nucleophilic bases or reducing agents has been well documented over the last decade.\(^{29}\) Brief exposure of a \(\text{C}_{6}\text{D}_{6}\) solution of 6 to air led to an immediate change from red to orange and NMR spectra showed clean conversion to 5-IMes (Scheme 6c). Interestingly, this route is more reliable and selective than the reduction of 3-IMes with \(\text{KC}_{8}\) (Scheme 6b) or the reduction of (CAAC\textsuperscript{Me})BHX\(_2\) (X = Cl, Br) with a Li(dme)\(_3\) countercation (Scheme 6d, Fig. 7, bottom). The boron centre is planar (\(\Sigma(\angle B1) 359.8(8)\)) and displays a B1–C1 single bond to the now sp\(^2\)-hybridised C1 atom (B1–C1 1.615(3); C1–N1 1.500(3) Å) and a formal B1–C21 double bond (1.435(3) Å) to the former IMes ligand. Alkyldene borates are typically generated by \(\pi\)-deprotonation of a suitable sp\(^3\)-borane precursor with a strong, non-nucleophilic base,\(^{30}\) by nucleophilic quaternisation of a neutral alkylidene borane,\(^{25,3a}\) or by tautomerisation of intramolecular frustrated Lewis pair systems involving an \(\pi\)-proton transfer from a highly electron-withdrawing \(\text{CH}_{3}\text{B}(\text{C}_{6}\text{F}_{5})\text{Cl}\) moiety to the Lewis basic site.\(^{25d}\) In contrast, the formation of 7 presumably results from the 2e\(^-\) reduction of 3-SIMes to a highly reactive dicordinate [CAAC-Me\textsuperscript{Me}](SIMes)B\(^-\) anion (proposed intermediate in Scheme 6), which undergoes intramolecular ortho-methyl C–H activation of the mesityl substituent, followed by a 1,2-hydride migration from the boron centre to the adjacent CAAC carbene carbon atom. Similar mesityl-CH\(_3\)-activations have been observed previously in the reduction of IMes-stabilised or mesityl-substituted boranes,\(^{30}\) as has the 1,2-hydride migration in
CAAC-supported hydroboranes.\textsuperscript{31} Unfortunately, the other products of this reduction could not be cleanly isolated. No tractable products were obtained from various reduction attempts of the phosphine derivatives \textit{3-PMe\textsubscript{3}} and \textit{3-PEt\textsubscript{3}}.

Conclusions
To conclude, we have described a facile synthetic route towards a series of metal-free haloborylenes, (CAAC)\textit{L}BCl\textsubscript{2}, by twofold reduction of a (CAAC)\textit{L}BCl\textsubscript{3} precursor with KC\textsubscript{8} in the presence of a donor ligand \textit{L}. Two likely intermediates in these reduction reactions were isolated in the form of a [(CAAC)BCl\textsubscript{2}]\textsuperscript{—} radical and a [(CAAC\textit{M}e\textsubscript{3})BCl\textsubscript{2}]\textsuperscript{—} anion, the potassium counterion of which is coordinated by \textit{L}. Variation of \textit{L} from a highly \textit{π}-accepting CAAC ligand to purely \textit{π}-accepting NHCs to moderately \textit{π}-accepting phosphines provides a series of borylenes displaying more or less sterically shielded and electron-rich boron centres, as determined by X-ray crystallographic analyses, NMR and UV-vis spectroscopy and cyclic voltammetry. The \textit{PMe\textsubscript{3}} derivative undergoes facile phosphine–NHC ligand exchange with a small NHC but not with the more sterically demanding ones. DFT calculations show that in the absence of possible \textit{π} backbonding to \textit{L}, the borylene lone pair is stabilised solely by \textit{π} backbonding to the CAAC ligand. Finally, the room-temperature reduction of the CAAC and NHC derivatives led to the isolation of several hydroborylenes and the product of an intramolecular C–H activation, suggesting the formation of radical and anionic dicoordinate “[[(CAAC)[NHC]B]]\textsuperscript{—}” reduction intermediates. We are currently studying the reactivity of these compounds, in particular \textit{3-PMe\textsubscript{3}}, towards Lewis acids and bases, unsaturated small molecules and electrophilic substitution, and will be reporting our results in a follow-up study.

Conflicts of interest
The authors declare no conflict of interest.

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Notes and references
\textsuperscript{†} Geometry optimisations were performed at the \textit{w}b97X-D/SD\textsubscript{D} level of density functional theory; bonding analyses reported are based on wave functions obtained from subsequent M06-2X/6-311G(d,p) single-point calculations. See ESI\textsuperscript{‡} for full details.
\textsuperscript{‡} These attempted ligand exchange reactions had to be carried out at room temperature as \textit{3-PMe\textsubscript{3}} started decomposing in solution at 60 °C.
\textsuperscript{§} \textit{3-PMe\textsubscript{3}} and \textit{3-PEt\textsubscript{3}} display near identical solution colours at 0.12 mM.
\textsuperscript{∥} The cyclovoltammograms of \textit{3-PMe\textsubscript{3}} and \textit{3-PEt\textsubscript{3}} are virtually identical (see Fig. S39 and S40 in the ESI) and show multiple irreversible redox events, presumably due to rapid phosphine dissociation upon oxidation or reduction.

** NMR-spectroscopic monitoring of the reduction of \textit{3-BMes} with Li in THF showed that \textit{5-BMes} is an intermediate in the formation of \textit{6}. Since the latter is quantitative, hydrogen abstraction from the ligand backbone can be excluded as source of the BH functionality.

2 W. Loehte-Holtgreven and E. S. van der Vleugel, \textit{Z. Phys.}, 1931, \textit{70}, 188.


