# Dalton Transactions



**PAPER** 

View Article Online
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**Cite this:** *Dalton Trans.*, 2016, **45**, 6023

# Exploring structural and electronic effects in three isomers of tris{bis(trifluoromethyl)phenyl}borane: towards the combined electrochemical-frustrated Lewis pair activation of $H_2\dagger$

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Three structural isomers of tris{bis(trifluoromethyl)phenyl}borane have been studied as the acidic component of frustrated Lewis pairs. While the 3,5-substituted isomer is already known to heterolytically cleave  $H_2$  to generate a bridging-hydride; *ortho*-substituents in the 2,4- and 2,5-isomers quench such reactivity through electron donation into the vacant boron  $p_z$  orbital and steric blocking of the boron centre; as shown by electrochemical, structural and computational studies. Electrochemical studies of the corresponding borohydrides identify that the two-electron oxidation of terminal-hydrides occurs at more positive potentials than observed for  $[HB(C_6F_5)_3]^-$ , while the bridging-hydride oxidizes at a higher potential still, comparable to that of free  $H_2$ .

Received 21st May 2015, Accepted 22nd July 2015 DOI: 10.1039/c5dt01918d

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## Introduction

Since the pioneering work of Stephan's group  $^1$  the field of frustrated Lewis pair (FLP) chemistry has grown rapidly.  $^{2-7}$  The archetypal FLP system combines the Lewis acid  $B(C_6F_5)_3$  with a sterically demanding Lewis base such as  $P(^tBu)_3$ , to effect the heterolytic cleavage of  $H_2$ , resulting in the formation of the corresponding hydridic and protic products respectively.

FLPs have found applications as catalysts or mediators for a variety of reactions such as the metal-free hydrogenation of imines and nitriles, alkynes, silyl enol ethers, and ketones. The activation of small molecules such as CO<sub>2</sub>, silyl and alkynes by FLPs has also found applications in the synthesis of heterocycles and other aromatic systems.

While much of the literature has focused on tris-(pentafluorophenyl)borane,  $B(C_6F_5)_3$  as the Lewis acidic component in FLPs, other electron-deficient boranes have been used, including a range of halogenated triarylboranes, <sup>14,16–18</sup> and borenium cations, <sup>19,20</sup> Other examples of Lewis acids that have found use in FLPs include: the triaryl aluminium species,  $Al(C_6F_5)_3$ , which generates a bridging-hydride following cleavage of  $H_2$ ;<sup>21</sup> and the carbon based *N*-methylacridinium salts which activate  $H_2$  even in the presence of  $H_2O$ .<sup>22</sup> FLPs are not limited to the main group, with zirconocene–phosphane complexes pioneered by Wass and co-workers shown to act as intramolecular FLPs that exhibit unprecedented reactivity towards small molecules;<sup>23</sup> similar chemistry has also been demonstrated with zirconocene-amines which act as hydrogenation catalysts with a wide range of substrates.<sup>24</sup>

In 2014 we introduced the concept of "combined electrochemical-frustrated Lewis pairs", 20,25,26 that couple the heterolytic cleavage of H2 by a conventional FLP with in situ electrochemical oxidation of the resultant borohydride and subsequent regeneration of the parent borane. The combined electrochemical-FLP systems were shown to be electrocatalytic for the oxidation of H<sub>2</sub> (to form two protons and two electrons - a key reaction in many hydrogen-based energy technologies). These preliminary reports represent the first application of FLP chemistry other than to catalyse the hydrogenation of small molecules. For example, by combining the archetypal  $B(C_6F_5)_3/P(^tBu)_3$  FLP in the presence of  $H_2$  whilst at the same time oxidizing the resultant [HB(C<sub>6</sub>F<sub>5</sub>)<sub>3</sub>]<sup>-</sup> intermediate formed, the potential for oxidation (the energetic driving force) of the borohydride at a glassy carbon electrode was found to be reduced by 0.61 V, in comparison to the direct oxidation of H<sub>2</sub> (observed at ca. +1.49 V vs.  $[FeCp_2]^{0/+}$  under the same conditions), an energy saving equivalent to 117.7 kJ mol<sup>-1</sup>.<sup>25</sup>

Herein we report studies on three isomers of tris{bis(tri-fluoromethyl)phenyl}borane (Fig. 1) and their associated

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 $<sup>\</sup>dagger$  Electronic supplementary information (ESI) available: Comprising: annotated ORTEP plot of the X-ray crystal structure of 3; cyclic voltammograms of  $B(C_6F_5)_3$  under corresponding conditions; further computational data. CCDC 1061234. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5dt01918d

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Fig. 1 Isomers of tris{bis(trifluoromethyl)phenyl}borane

tris{bis(trifluoromethyl)phenyl}borohydrides. Electrochemical and computational investigations explain their varying ability to heterolytically cleave  $H_2$  as part of a FLP and allow us to extend our studies into combined electrochemical-FLPs.

## Results and discussion

### Synthesis and characterization

The synthesis of tris{3,5-bis(trifluoromethyl)phenyl}borane 1 has been previously reported, 18,27 and whilst a synthetic route to tris{2,4-bis(trifluoromethyl)phenyl}borane 2 has been reported by Cornet *et al.*28 their method led to a mixture of products requiring the isolation of 2 in low yield by fractional sublimations. The commercial availability of 2,4-bis(trifluoromethyl)bromobenzene and 2,5-bis(trifluoromethyl)bromobenzene allows us to report an improved synthetic route to 2 and also the synthesis of the novel tris{2,5-bis(trifluoromethyl)phenyl}borane 3.

Boranes 2 and 3 were synthesized by lithium–halogen exchange of  ${\rm BrC_6H_3(CF_3)_2}$  with  ${\rm ^nBuLi}$  at -77 °C generating the aryllithium species  ${\rm LiC_6H_3(CF_3)_2}$ . Treatment of this with a third of an equivalent of  ${\rm BCl_3}$ , followed by warming to room temperature, subsequent removal of the volatiles and purification by sublimation or recrystallization allowed for the isolation of pure 2 and 3, in 90 and 77% yields respectively. It should be noted that applying the same synthetic route for the synthesis of 1, leads to a mixture of products with  $[{\rm Li}({\rm OEt_2})_n]$ - $[{\rm B}\{3,5\text{-}({\rm CF_3})_2{\rm C_6H_3}\}_4]$  as the major component.

The crystal structures of  $1^{29}$  and  $2^{28}$  have been previously reported. Single crystals of 3 were obtained by slow diffusion of a saturated  $\mathrm{CH_2Cl_2}$  solution of 3 into *n*-hexane at -25 °C, from which the crystal structure was obtained in collaboration with the EPSRC UK National Crystallography Service<sup>30</sup> (Fig. 2 and S1,† and Table 2).

The structures of 1–3 all show similar features, with a trigonal-planar boron centre and the three aryl rings twisted with respect to the  $BC_3$  plane to minimize steric interactions between the aryl rings. The degree of twist can be quantified by an appropriate choice of C–B–C–C torsion angles, which are both smaller and more consistent for 1 {mean  $36(2)^\circ$ , range

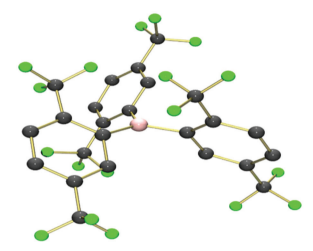


Fig. 2 The crystallographic molecular structure of  $B\{2,5-(CF_3)_2C_6H_3\}_3$  3 (hydrogen atoms removed for clarity).

33.2–38.9°} than for 2 {mean  $54(11)^\circ$ , range 42.3– $68.3^\circ$ } or 3 {mean  $53(9)^\circ$ , range 40.9– $61.2^\circ$ }, due to the steric bulk of *ortho*-trifluoromethyl substituents. For 2 and 3 this results in the *ortho*-CF<sub>3</sub> groups orientated above/below the boron centre resulting in B···F distances of 2.81(1), 2.80(7) Å for 2 and 3 respectively. The orientation of these groups suggests the possibility of electron donation from the fluorine atoms into the formally vacant  $p_z$  orbital at boron; an effect together with the steric influence of the *ortho*-CF<sub>3</sub> groups which would be expected to have a significant influence on the Lewis acidity/reactivity of these compounds. An example of the type of reactivity which could be expected from the *ortho*-CF<sub>3</sub> groups, was observed by Cornet *et al.*, who reported evidence of B–Cl/B–F exchange in mixtures containing the boranes  $B(Ar^F)_2Cl$  where  $Ar^F = 2,4,6-(CF_3)_3C_6H_2$ ,  $2,4-(CF_3)_2C_6H_3$  or  $2,6-(CF_3)_2C_6H_3$ .<sup>28</sup>

The reactivity of 1 as the Lewis acidic component of an FLP has been previously studied, <sup>18</sup> which showed that the reaction of 1 with  $H_2$  in the presence of 2,2,6,6-tetramethylpiperidine (tmp) leads to rapid formation of the bridging hydride species [tmpH][( $\mu$ -H)(1)<sub>2</sub>]. Analogous reactions of 2 and 3 with  $H_2$  in the presence of the Lewis bases tmp or  $P({}^tBu)_3$ , result in neither Lewis acid-base adduct formation nor any evidence of

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H<sub>2</sub> activation observable by NMR spectroscopy over a minimum period of 48 hours.

Direct synthesis of authentic terminal-hydride species [HB- $\{C_6H_3(CF_3)_2\}_3$  proved to be possible for all three isomers 1-3 by direct reaction of the borane with sodium triethylborohydride in toluene solution, resulting in near quantitative conversion to the corresponding borohydrides. NMR spectra of the terminal-hydride species Na[1-H]-Na[3-H] show doublets in the <sup>11</sup>B spectra ( $\delta_B$ : -9.1, -15.3, -14.2 and <sup>1</sup> $J_{BH}$ : 88, 93, 84 Hz respectively) and broad 1:1:1:1 quartets in the proton spectra ( $\delta_{\rm H}$ : +3.66, +4.06, +3.06 respectively). Additionally, for Na[2-H] and Na[3-H] clear evidence of through-space coupling between the hydride and ortho-CF3 groups is observed in the <sup>19</sup>F spectra ( $I_{\rm FH}$  = 6.8, 7.6 Hz respectively); further, in the case of Na[3-H] this coupling is also observable in the proton spectrum with the hydride signal observable as a 1:1:1:1 quartet of broad 1:3:3:1 quartets (Fig. 3).

Synthesis of Na[2-H] and Na[3-H] allows for the confirmation that the inability of the  $2,3/P(^{t}Bu)_{3}$  FLPs to cleave H<sub>2</sub> is not due to unfavourable thermodynamics, but due to the significant kinetic barrier resulting from the steric and electronic effects of the ortho-CF3 groups. Reaction with authentic [(tBu)3PH]Cl results in rapid metathesis (indicated by precipitation of NaCl) and formation of the salts  $[(^tBu)_3PH][2-H]$  and [(<sup>t</sup>Bu)<sub>3</sub>PH][3-H]. NMR spectra of which, show no liberation of H<sub>2</sub>, regeneration of free borane/phosphine, or any other evidence of reaction over a 66 hour period.

## **Electrochemical studies**

Cyclic voltammetric studies of 1-3 were performed in the noncoordinating solvent  $CH_2Cl_2$  using  $[^nBu_4N][B(C_6F_5)_4]$  as the added electrolyte at a glassy carbon electrode (GCE). In all three cases a one-electron, reduction process is observed with quasi-reversible (moderately fast) electron transfer kinetics. However for 1 (Fig. 4a) the process appears to be chemically irreversible (but not *electrochemically* irreversible) at scan rates up to 2.0 V s<sup>-1</sup> due to fast chemical follow-up kinetics cf. the rate of electron transfer, while for both 2 (Fig. 4b) and 3 (Fig. 4c) an associated oxidation wave is observed indicating

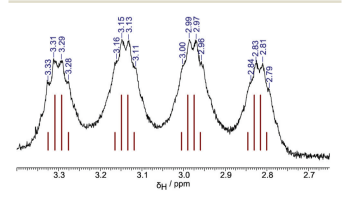
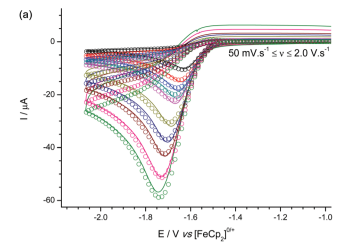
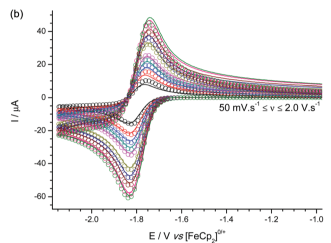


Fig. 3 Hydride resonance in <sup>1</sup>H NMR spectrum of Na[3-H], showing coupling to both <sup>11</sup>B (I = 3/2) and an ortho-CF<sub>3</sub> group  $\{3 \times {}^{19}F, I = 1/2\}$ .





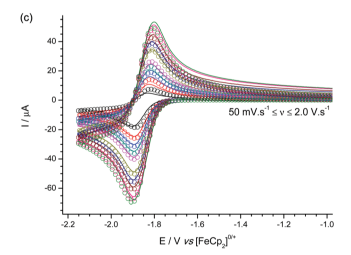


Fig. 4 Experimental (line) and simulated (open circles) cyclic voltammograms for the reduction of (a) 1, (b) 2, and (c) 3.

that the radical anions 2'- and 3'- are sufficiently stable on the electrochemical timescale for their subsequent (re-)oxidation to be observable.

Based on our previous experience from an electrochemical study of B(C<sub>6</sub>F<sub>5</sub>)<sub>3</sub>,<sup>31</sup> we performed digital simulations of the **Paper** 

Table 1 Mechanistic parameters obtained by digital simulation of voltammetric data for the one-electron reductions of 1-3

		1	2	3
$BArF_{18} + e^- = BArF_{18}^{\bullet -}$	$E^{\circ}$ vs. $[\text{FeCp}_2]^{0/+}/\text{V}$ $\alpha$ $k^{\circ}/\text{cm s}^{-1}$	$-1.61 \pm 0.01$ $0.419$ $4.56 \times 10^{-3}$	$-1.79 \pm 0.01$ $0.498$ $2.61 \times 10^{-2}$	$-1.85 \pm 0.01$ $0.468$ $2.22 \times 10^{-2}$
BArF <sub>18</sub> *- $\Rightarrow$ 'decomposition' $D(BArF_{18}) = D(BArF_{18}^{})/cm^2 s^{-1} a$	$k_{\rm f}/{\rm s}^{-1}{}^{b}$ 3.76 × 10 <sup>-5</sup>	$\geq 25$ $1.13 \times 10^{-5}$	$1.92 \times 10^{-2} \\ 1.13 \times 10^{-5}$	$8.08 \times 10^{-2}$

<sup>&</sup>lt;sup>a</sup> Diffusion constants (D) obtained via <sup>1</sup>H and <sup>19</sup>F DOSY NMR spectroscopy. <sup>b</sup> k<sub>f</sub> values are modelled as a pseudo first-order process.

experimental voltammetric data modelled using an EC-mechanism (*i.e.* a reversible, heterogeneous electron transfer step followed by an irreversible, homogeneous chemical step which generates electro-inactive products. Other postulated mechanisms produced a poor fit to the data). These digital simulations allowed us to extract pertinent mechanistic parameters such as the formal redox potentials and charge transfer coefficients ( $E^{\circ}$  and  $\alpha$  respectively) and kinetic parameters for the electron transfer ( $k^{\circ}$ ) and follow-on chemical step ( $k_{\rm f}$ ) as shown in Table 1.

The formal reduction potentials ( $E^{\circ}$ ) suggest that 1 is the most electrophilic of the three boranes, while 3 is the least. While the difference in electrophilicity between 2 and 3 is consistent with a simplified view of the inductive electron withdrawing effects of the different *meta*- and *para*-(CF<sub>3</sub>) group positions based on Hammett parameters { $\sigma_{meta}(\text{CF}_3) = 0.43 \text{ vs.}$   $\sigma_{para}(\text{CF}_3) = 0.54$ }<sup>32</sup> the relative electrophilicity of 1 cannot be

Table 2 Crystallographic data for 3

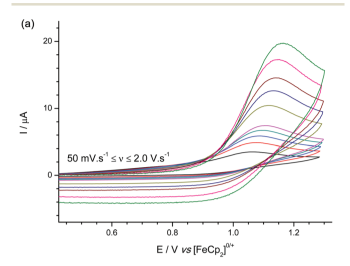
	B{2,5-(CF <sub>3</sub> ) <sub>2</sub> C <sub>6</sub> H <sub>3</sub> } <sub>3</sub> 3		
Empirical formula	$C_{24}H_9BF_{18}$		
Formula weight	650.12		
Temperature/K	100		
Crystal system	Triclinic		
Space group	$Par{1}$		
a/Å	7.2951(5)		
b/Å	10.6358(7)		
c/Å	15.9794(11)		
$\alpha/^{\circ}$	85.950(4)		
β/°	86.582(4)		
γ/°	74.846(4)		
Volume/Å <sup>3</sup>	1192.64(14)		
Z	2		
$ ho_{ m calc}/ m mg~mm^{-3}$	1.810		
$\mu/\text{mm}^{-1}$	0.204		
F(000)	640.0		
Crystal size/mm <sup>3</sup>	$0.30 \times 0.06 \times 0.02$		
Radiation	Mo Kα ( $\lambda = 0.71075 \text{ Å}$ )		
$2\Theta$ range for data collection	3.974 to 54.96°		
Index ranges	$-9 \le h \le 8, -13 \le k \le 13,$		
_	$-20 \le l \le 20$		
Reflections collected	17 457		
Independent reflections	$5459 \left[ R_{\text{int}} = 0.0820, \right]$		
•	$R_{\text{sigma}} = 0.0575$		
Data/restraints/parameters	5459/0/388		
Goodness-of-fit on F <sup>2</sup>	1.055		
Final <i>R</i> indexes $[I \ge 2\sigma(I)]$	$R_1 = 0.0555$ , $wR_2 = 0.1508$		
Final R indexes [all data]	$R_1 = 0.0712, \text{ w} R_2 = 0.1631$		
Largest diff. peak/hole/e A <sup>-3</sup>	0.56/-0.39		

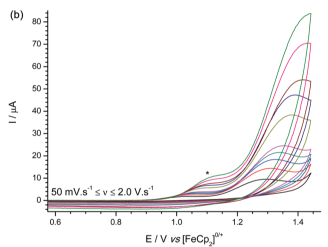
similarly rationalized. This is in part because simple Hammett parameters for ortho-(CF<sub>3</sub>) substituents do not satisfactorily account for any additional steric and/or electronic effects. We can rationalize the reduced electrophilicity of 2 and 3 in comparison to 1 by considering the electronic effect of having ortho-CF<sub>3</sub> groups present in 2 and 3. As noted above, the crystal structures of 2 and 3 show the ortho-(CF<sub>3</sub>) groups are positioned at sufficiently close distances above the central BC<sub>3</sub> plane such that donation from the lone pairs on the fluorine atoms into the vacant boron  $p_z$  orbital on boron could occur; such donation of electron density would be expected to reduce the electrophilicity of the boron centre, as is observed experimentally by our electrochemical measurements and indicated in electronic structure DFT calculations ( $vide\ infra$ ).

Furthermore, the steric shielding of the boron centre by the ortho-(CF<sub>3</sub>) groups in 2 and 3 is qualitatively evident from the observation of more reversible redox processes in the cyclic voltammetric data. Quantitatively this is shown in the values of the rate constants  $(k_f)$  obtained from voltammetric digital simulation for the radical anion decomposition step, which is assumed to proceed in a similar fashion as was previously ascertained for the analogous [B(C<sub>6</sub>F<sub>5</sub>)<sub>3</sub>]\*- intermediate via reaction between solvent molecules and the boron centre in the reduced radical anion intermediates, 1\*-, 2\*-, or 3\*-.31 The value of  $k_{\rm f}$  is at least three orders of magnitude greater for the decomposition of 1'- than for 2'- or 3'-, where, in the latter two cases, the presence of ortho-CF<sub>3</sub> groups provides significant steric shielding to the boron centre. It is worth noting here that these findings demonstrate the ease with which synthetic chemists working in this area can gain powerful insights into the chemistry of Lewis acidic species by the application of simple, rapid electrochemical characterization techniques in addition to the more ubiquitous crystallographic and spectroscopic characterization techniques. Simple examination of the shape and position of the voltammetry of each borane, obtained in a 20 minute experiment using inexpensive equipment can tell us qualitatively that in comparison to 1, boranes 2 and 3 are more sterically hindered, and less electrophilic, and therefore less Lewis acidic and less likely to be active FLP components for H2 activation. The voltammetry even allows us to infer why this is so, given that the only thing boranes 2 and 3 have that 1 does not have and that could simultaneously sterically shield the borane and reduce the electronic demand at the boron centre are the o-CF<sub>3</sub> groups.

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In comparison with the archetypal Lewis acid, B(C<sub>6</sub>F<sub>5</sub>)<sub>3</sub> {which is observed under the same conditions as a quasireversible reduction at  $E^{\circ} = -1.518 \text{ V} \text{ vs. } [\text{FeCp}_2]^{0/+} \text{ (see}$ 





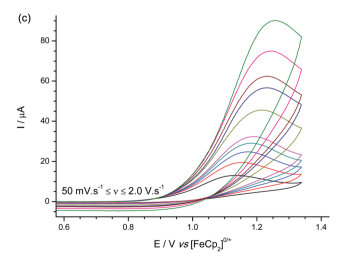


Fig. 5 Experimental cyclic voltammograms for the oxidation of (a) Na-[1-H], (b) Na[2-H], and (c) Na[3-H] shoulder (\*) visible at the higher scan rates due to trace impurity in the solvent/electrolyte.

Fig. S2†)}, boranes 1-3 are all less electrophilic; while  $[B(C_6F_5)_3]^{\bullet-}$  is more stable (smaller  $k_f$ ) with respect to followon decomposition reactions than 1.-, but still considerably less stable than the *ortho*-(CF<sub>3</sub>) stabilized 2<sup>•–</sup> and 3<sup>•–</sup>.

Borohydride oxidation. Our combined electrochemical-FLP concept requires the electrochemical oxidation of borohydrides generated by the heterolytic cleavage of H2. Despite the only such species generated from the B{C<sub>6</sub>H<sub>3</sub>(CF<sub>3</sub>)<sub>2</sub>}<sub>3</sub> isomers, being the previously reported bridging hydride  $[(\mu-H)(1)_2]^{-18}$ the ease of direct synthesis of the terminal hydrides  $[HB\{C_6H_3(CF_3)_2\}_3]^-$  allows for their electrochemical study, to add to our understanding of the electrochemistry of the triarylborohydrides. Therefore, electrochemical studies of Na[1-H]-Na[3-H] and  $[tmpH][(\mu-H)(1)_2]$  were performed under the same conditions as their borane precursors (although Na[1-H] and  $[tmpH][(\mu-H)(1)_2]$  proved only sparingly soluble in  $CH_2Cl_2$ , and hence their concentrations cannot be accurately determined). In all cases a single irreversible oxidation wave is observed at scan rates up to  $2.0 \text{ V s}^{-1}$ .

For the terminal hydrides Na[1-H]-Na[3-H] (Fig. 5a-c) oxidations occurred with peak potentials of +1.08, +1.31, +1.13 V vs. [FeCp2]0/+ at a scan rate of 100 mV s-1 respectively; with no evidence of electroactive product species (such as the parent boranes) being regenerated in sufficient quantities to be observed.

The bridging hydride  $[tmpH][(\mu-H)(1)_2]$  (Fig. 6) oxidation is observed as a shoulder on the edge of the solvent window, at ca. +1.55 V vs.  $[FeCp_2]^{0/+}$  at a scan rate of 100 mV s<sup>-1</sup> (observable distinct to the solvent/electrolyte breakdown at scan rates below 1.0 V s<sup>-1</sup>), trace amounts of the terminal hydride species [1-H] are also observed. Following oxidation of  $[(\mu-H)(1)_2]^-$ , sweeping to negative potentials results in observation (at 100 mV s<sup>-1</sup> scan rate and above) of an irreversible reduction wave at ca.  $-1.7 \text{ V} \text{ vs.} [\text{FeCp}_2]^{0/+}$  characteristic of the reduction wave observed for the parent borane 1.

In comparison to our previous studies on  $[HB(C_6F_5)_3]^{-,25}$  all of the terminal-hydrides oxidize at more positive potentials,

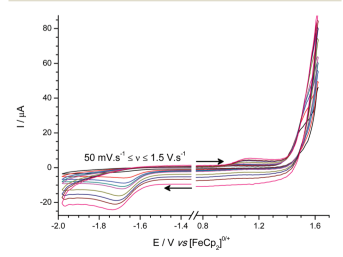


Fig. 6 Experimental cyclic voltammogram(s) for the oxidation of  $[tmpH][(\mu-H)(1)_2].$ 

(a) (c) (c)

Fig. 7 Lowest unoccupied molecular orbitals (LUMOs) for (a) 1, (b) 2, and (c) 3.

whilst the bridging-hydride species,  $[(\mu-H)(1)_2]^-$ , is oxidized at an even more positive potential approaching that of the direct oxidation of H<sub>2</sub> at a GCE (*ca.* +1.5 V vs.  $[FeCp_2]^{0/+}$ ), and is effectively behaving as an electrolyte under these conditions.

## **Computational studies**

**Paper** 

To further our understanding of the tris{bis(trifluoromethyl)-phenyl}borane isomers 1–3 we have investigated them and their associated radical-anions 1<sup>\*-</sup>–3<sup>\*-</sup> using density functional theory (DFT).

For all the neutral boranes 1–3 the LUMOs (Fig. 7) showed a high degree of boron  $p_z$  character (19, 22, 19% respectively, the majority of the remaining contributions being from the phenyl  $\pi$ -systems) as do the SOMOs (Fig. S4†) for the radical-anions 1<sup>-</sup>-3<sup>-</sup>.

Total molecular energy calculations indicate that the ground state of **1** is lower in energy than those of **2** and **3** by *ca.* 43 kJ mol<sup>-1</sup>. Analysis of calculated Mulliken atomic charges for **1**–**3**, show that any stabilisation of **1** compared to **2** and **3** cannot be attributed to electron withdrawing effects of the aryl rings; and therefore may be attributed to the reduced steric hindrance caused by the lack of *ortho*-CF<sub>3</sub> groups in **1**.

As noted previously, the *ortho*-CF $_3$  groups in 2 and 3 are orientated such that there is potential for B···F bonding interactions. Such interactions are clearly identified by the calculated bonding parameters: the B···F bonding parameter for 2 is 0.065, whilst for 3 it is 0.076 (in both cases averaged over all contributing B···F pairs). As expected, in 1, where no interaction occurs, the B···F bonding parameter to the *meta*-CF $_3$  groups is <0.001.

## Conclusions

We have investigated three structural isomers of tris{bis(tri-fluoromethyl)phenyl}borane in terms of both their electrochemical redox chemistries and also as the Lewis acidic components of an FLP for the heterolytic cleavage of  $\rm H_2$ .

The two isomers of tris{bis(trifluoromethyl)phenyl}borane that incorporate *ortho*-CF<sub>3</sub> groups were not found to be active as the Lewis acidic component of FLPs for H<sub>2</sub> cleavage reac-

tions. This lack of reactivity is due to a combination of, kinetic effects resulting from in part steric shielding of the boron centre; but also quenching of the boranes' electrophilicity through B···F bonding interactions, which are quantified by DFT calculations and electrochemical measurements. Whilst electrochemical studies show that all three isomers are less electrophilic than the archetypal Lewis acid  $B(C_6F_5)_3$  the link between Lewis acidity, activity towards  $H_2$  in an FLP, and electrophilicity, as shown in our previous studies  $^{20,25,26}$  and those of other groups  $^{33-35}$  is complex and requires further study.

The direct synthesis of all three tris{bis(trifluoromethyl)-phenyl}borohydride species under mild conditions allows for their reduction potentials to be measured, and their propensity for combined electrochemical-frustrated Lewis pair catalysis to be screened in a straightforward manner. The oxidation potentials of all three terminal borohydrides studied were found to be more positive than that of  $[HB(C_6F_5)_3]^-$  yet are still less than the potential required for the direct oxidation of  $H_2$  at a GCE under identical conditions.

Electrochemical studies of the bridging hydride formed when  $H_2$  is cleaved by the  $B\{3,5\text{-}(CF_3)_2C_6H_3\}_3$ /tmp FLP, show that this species is oxidized at comparable potentials to that of the direct oxidation of  $H_2$ . However, following oxidation, the regeneration of the parent borane species is clearly observed, which is not the case for any of the terminal borohydride Na[1-H]-Na[3-H] species studied.

What this report demonstrates is the utility of electrochemical characterization methods to enable synthetic chemists to rapidly screen prospective new Lewis acids using simple electrochemical techniques to gain insights into the chemical behaviour of new species. It also provides insights to guide the design of new Lewis acids for researchers wishing to employ the combined electrochemical-frustrated Lewis pair activation of  $H_2$ , which is the focus of our ongoing studies.

# Experimental

All reactions and manipulations were performed under an atmosphere of dry, oxygen-free N<sub>2</sub>, using either standard Schlenk techniques or in either a MBraun UNIlab or LABmaster

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glovebox. All solvents were dried prior to use by refluxing over an appropriate drying agent {Na/benzophenone for petroleum ether (b.p 40-60 °C) and diethyl ether; Na for toluene; CaH2 for dichloromethane}, collected by distillation under an inert N<sub>2</sub> atmosphere and stored over 4 Å molecular sieves prior to use. All other reagents were obtained from commercial suppliers and used as received.

NMR Spectra were obtained on either a Bruker Avance III 500 MHz or Bruker AV 400 MHz spectrometer, all deuterated solvents were dried over 4 Å molecular sieves prior to use. For <sup>1</sup>H spectra residual *protio*-solvent was used as an internal standard; for <sup>13</sup>C the solvent resonance(s) were used as an internal standard;36 for 19F spectra CFCl3 was used as an external standard; for 11B spectra BF<sub>3</sub>·Et<sub>2</sub>O was used as an external standard. 1H and 19F DOSY experiments were performed on a Bruker Avance III 500 MHz spectrometer equipped with a broadband multinuclear probe, using a longitudinal eddy current delay incorporating bipolar gradients for diffusion and spoil gradients (ledbpgp2s) pulse sequence.<sup>37</sup>

Mass spectrometry was performed by the EPSRC Mass Spectrometry Service at the University of Swansea for 3, or by Dr L. Haigh at Imperial College using a Micromass Autospec Premier spectrometer for 2. Elemental analyses were performed by Mr S. Boyer of the Elemental Analysis Service at London Metropolitan University.

Single crystals of 3 were grown by slow diffusion of a saturated CH<sub>2</sub>Cl<sub>2</sub> solution of the compound into n-hexane; data collection and processing was performed at the UK National Crystallographic Service at the University of Southampton.<sup>30</sup> Using Olex2,38 the structure was solved and space group assigned with SuperFlip/EDMA<sup>39</sup> using charge flipping, and then refined with the ShelXL version 2014/740 refinement program using least squares minimization.

CCDC 1061234 contains the supplementary crystallographic data for this paper.

Electrochemical studies were carried out using a Metrohm Autolab PGSTAT302N potentiostat linked to a computer running Metrohm Autolab NOVA version 1.11 software, in conjunction with a three electrode cell comprising: a glassy carbon disc working electrode (Bioanalytical Systems Inc., ca.  $7.0~\text{mm}^2$  area calibrated using the  $[\text{FeCp}_2]^{0/\text{+}}$  redox couple), a platinum wire (99.99% purity) counter electrode, and a silver wire (99.99% purity) pseudo-reference electrode; all electrodes were polished with 0.3  $\mu m$   $\alpha$ -alumina and dried prior to use. All electrochemical measurements were performed at ambient temperature under a dry N2 atmosphere, in CH2Cl2 containing  $0.05 \text{ M} \text{ } [^{n}\text{Bu}_{4}\text{N}][\text{B}(\text{C}_{6}\text{F}_{5})_{4}] \text{ as the supporting electrolyte and}$ between 1.0 and 2.0 mM of the analyte species of interest. Cyclic voltammetric measurements were iR-compensated using positive-feedback to within 85  $\pm$  5% of the uncompensated solution resistance. ["Bu<sub>4</sub>N][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>] was synthesized according to published methods. 41 All potentials were referenced to the [FeCp2]0/+ redox couple, which was added as internal standard. Simulations of electrochemical processes were performed using ElchSoft DigiElch version 7.096 software. 42

DFT calculations were performed using the Gaussian 09 computational package. 43 Geometry optimization calculations have been carried out using the three-parameter exchange functional of Becke44 (B3) and the correlation functional of Lee, Yang, and Parr (LYP), B3LYP. 45 The 6-311+G(d,p) basis set has been implemented for all atoms. 46 Structures were geometry optimized in the gas phase with the default convergence criteria and confirmed as minima through frequency calculations. All optimized structures were confirmed as minima by frequency analysis with thermodynamic properties extracted for the gas phase at 298.15 K and 1 atm. Bonding parameters between B and F atoms were calculated as the absolute values of the associated non-diagonal elements of the condensed to atoms electron density matrix. Density matrixes were computed in a separate calculation taking into consideration basis set superposition error (BSSE) correction using an unrestricted Hartree-Fock (H-F) calculation with counterpoise (CP) correction approach as implemented in Gaussian suite.

 $\mathbf{1}^{18,27}$  $B\{3,5-(CF_3)_2C_6H_3\}_3$ and  $[tmpH](\mu-H)(B{3,5 (CF_3)_2C_6H_3\}_3)_2$  [tmpH][( $\mu$ -H)(1)<sub>2</sub>]<sup>18</sup> were synthesized as previously reported.

#### $B\{2,4-(CF_3)_2C_6H_3\}_3$ 2

2,4-Bis(trifluoromethyl)bromobenzene (2.00 g, 1.16 cm<sup>3</sup>, 6.83 mmol) and Et<sub>2</sub>O (100 cm<sup>3</sup>) were combined and the solution cooled to -77 °C. With the aid of rapid stirring, <sup>n</sup>BuLi (2.87 cm<sup>3</sup>, 7.17 mmol, 2.5 M in hexanes) was added slowly by means of a syringe. Following one hour of stirring, BCl<sub>3</sub> (2.28 cm<sup>3</sup>, 2.28 mmol, 1.0 M in heptane) was syringed into the amber solution and the mixture permitted to warm slowly to room temperature. The volatiles were removed in vacuo and the off white residue extracted with  $CH_2Cl_2$  (3 × 25 cm<sup>3</sup>) and filtered through Celite. Volatiles were removed under vacuum, and following a high vacuum sublimation step (10<sup>-6</sup> mbar) at 85 °C, a pure white solid was obtained. Yield 1.33 g (2.04 mmol, 90%).

<sup>1</sup>H NMR (400.4 MHz,  $CD_2Cl_2$ , 25 °C,  $\delta$ ): +8.06 (s, 3H, 3-H), +7.87 (d, 3H,  ${}^{3}J_{HH}$  = 8 Hz, 5-H), +7.46 (d, 3H,  ${}^{3}J_{HH}$  = 8 Hz, 6-H); <sup>11</sup>B NMR (128.4 MHz,  $CD_2Cl_2$ , 25 °C,  $\delta$ ): +74.0 (br.s); <sup>13</sup>C{<sup>1</sup>H} NMR (100.6 MHz,  $CD_2Cl_2$ , 25 °C,  $\delta$ ): +144.2 (br, 1-C), +135.9 (s, 6-CH), +134.2 (q,  ${}^{2}J_{CF}$  = 34 Hz, 2/4-C), +133.7 (q,  ${}^{2}J_{CF}$  = 34 Hz, 2/ 4-C), +127.9 (q,  ${}^{3}J_{CF}$  = 3 Hz, 5-CH), +123.9 (q,  ${}^{1}J_{CF}$  = 273 Hz, 2/ 4-CF<sub>3</sub>), +123.6 (sept.,  ${}^{3}J_{CF} = 3$  Hz, 3-CH), +123.6 (q,  ${}^{1}J_{CF} =$ 273 Hz, 2/4-CF<sub>3</sub>). <sup>19</sup>F NMR (376.8 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): -56.6 (s, 9F, 2-CF<sub>3</sub>), -63.8 s, 9F, 4-CF<sub>3</sub>). HRMS-EI (m/z): [M] calc. for C<sub>24</sub>H<sub>9</sub>BF<sub>18</sub>, 650.0510; found, 650.0491. Elemental analysis (calc. for C<sub>24</sub>H<sub>9</sub>B<sub>1</sub>F<sub>18</sub>): C 44.34 (44.48), H 1.40 (1.47).

## $B\{2,5-(CF_3)_2C_6H_3\}_3$ 3

2,5-Bis(trifluoromethyl)bromobenzene (3.1 cm<sup>3</sup>, 17.9 mmol) and  $Et_2O$  (100 cm<sup>3</sup>) were combined and cooled to -77 °C. <sup>n</sup>BuLi (11 cm<sup>3</sup>, 17.6 mmol, 1.6 M in hexanes) was added to the stirred solution. After one hour BCl<sub>3</sub> (5.8 cm<sup>3</sup>, 5.8 mmol, 1.0 M in heptane) was added to the orange solution and the mixture permitted to slowly warm to room temperature. The volatiles were removed in vacuo and the pale yellow residue extracted

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with  $CH_2Cl_2$  (3 × 25 cm<sup>3</sup>) and filtered (*via* cannula). The product is then recrystallized from  $CH_2Cl_2/n$ -hexane and isolated as a white micro-crystalline solid. Yield 2.89 g (4.44 mmol, 77%).

<sup>1</sup>H NMR (500.21 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): +7.95 (s, 3H, 3/4-H), +7.95 (s, 3H, 3/4-H), +7.47 (s, 3H, 6-H); <sup>11</sup>B NMR (160.49 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): +70.7 (br.s); <sup>13</sup>C{<sup>1</sup>H} NMR (125.78 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): +141.5 (br.s, 1-C), +137.1 (q,  $^2J_{\text{CF}}$  = 33 Hz, 2/5-C), +133.3 (q,  $^2J_{\text{CF}}$  = 33 Hz, 2/5-C), +132.4 (q,  $^3J_{\text{CF}}$  = 3.7 Hz, 3/4/6-C), +129.3 (q,  $^3J_{\text{CF}}$  = 3.7 Hz, 3/4/6-C), +124.1 (q,  $^1J_{\text{CF}}$  = 275 Hz, 2/5-CF<sub>3</sub>), +123.9 (q,  $^1J_{\text{CF}}$  273 Hz, 2/5-CF<sub>3</sub>); <sup>19</sup>F NMR (470.67 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): -56.5 (s, 9F, 2-CF<sub>3</sub>), -63.7 (s, 9F, 5-CF<sub>3</sub>). HRMS-APCI (*m*/*z*): [M - F]<sup>+</sup> calc. for C<sub>24</sub>H<sub>9</sub>BF<sub>17</sub>, 631.0525; found, 631.0519. Elemental analysis (calc. for C<sub>24</sub>H<sub>9</sub>B<sub>1</sub>F<sub>18</sub>): C 44.22 (44.48), H 1.38 (1.47).

## $Na[HB{3,5-(CF_3)_2C_6H_3}_3] Na[1-H]$

To a solution of 1 (0.40 g, 0.62 mmol) in toluene (10 cm<sup>3</sup>) was added Na[HBEt<sub>3</sub>] (0.6 cm<sup>3</sup>, 0.6 mmol, 1.0 M in toluene), the reaction mixture was stirred for 6 hours to give a colourless solution. All volatiles were removed *in vacuo*, to give a white residue which was washed with petroleum ether ( $2 \times 5$  cm<sup>3</sup>) and dried *in vacuo* to give a white solid. Yield 0.40 g (0.59 mmol, 95%).

<sup>1</sup>H NMR (500.21 MHz, CD<sub>3</sub>CN, 25 °C, δ): +7.70 (s, 6H, 2,6-H), +7.58 (s, 3H, 4-H), +3.66 (br.q,  ${}^{1}J_{\rm HB}$  = 84 Hz, 1H, HB); <sup>11</sup>B NMR (160.49 MHz, CD<sub>3</sub>CN, 25 °C, δ): -9.1 (d,  ${}^{1}J_{\rm BH}$  = 88 Hz); <sup>13</sup>C{<sup>1</sup>H} NMR (125.78 MHz, CD<sub>3</sub>CN, 25 °C, δ): +165.0 (q,  ${}^{1}J_{\rm CF}$  = 49 Hz, 1-C), +135.7 (s, 2,6-C), +130.1 (q,  ${}^{2}J_{\rm CF}$  = 32 Hz, 3,5-C), +126.0 (q,  ${}^{1}J_{\rm CF}$  = 272 Hz, CF<sub>3</sub>), +118.5 (s, 4-C); <sup>19</sup>F NMR (470.67 MHz, CD<sub>3</sub>CN, 25 °C, δ): -63.0 (s, 18F, CF<sub>3</sub>). Elemental analysis (calc. for C<sub>24</sub>H<sub>10</sub>B<sub>1</sub>F<sub>18</sub>Na): C 42.95 (42.76), H 1.61 (1.50).

## $Na[HB{2,4-(CF_3)_2C_6H_3}_3]Na[2-H]$

To a solution of 2 (0.275 g, 0.42 mmol) in toluene (8 cm<sup>3</sup>) was added Na[HBEt<sub>3</sub>] (0.43 cm<sup>3</sup>, 0.43 mmol, 1.0 M in toluene), the reaction mixture was stirred for 4 hours to give a cloudy white suspension. The reaction mixture was concentrated *in vacuo* reducing its volume to ca. 2 cm<sup>3</sup>, and the product precipitated by addition of petroleum ether and cooling to -25 °C. The white solid was isolated by filtration and dried *in vacuo*. Yield 0.231 g (0.34 mmol, 81%).

<sup>1</sup>H NMR (500.21 MHz, CD<sub>3</sub>CN, 25 °C, δ): +7.76 (s, 3H, 3-H), +7.47 (d,  ${}^{3}J_{\rm HH}$  = 7.9 Hz, 3H, 5-H), +7.11 (d,  ${}^{3}J_{\rm HH}$  = 7.9 Hz, 3H, 6-H), +4.06 (br.q,  ${}^{1}J_{\rm HB}$  = 93 Hz, 1H, BH); <sup>11</sup>B NMR (160.49 MHz, CD<sub>3</sub>CN, 25 °C, δ): -15.33 (d,  ${}^{1}J_{\rm BH}$  = 93 Hz); <sup>13</sup>C{<sup>1</sup>H} NMR (125.78 MHz, CD<sub>3</sub>CN, 25 °C, δ): +139.1 (s, 6-CH), +135.3 (q,  ${}^{2}J_{\rm CF}$  = 29 Hz, 2/4-C), +126.7 (q,  ${}^{3}J_{\rm CF}$  = 3.7 Hz, 5-CH), +126.7 (q,  ${}^{1}J_{\rm CF}$  = 275 Hz, 2/4-CF<sub>3</sub>), +126.3 (q,  ${}^{2}J_{\rm CF}$  = 32 Hz, 2/4-C), +126.2 (q,  ${}^{1}J_{\rm CF}$  = 272 Hz, 2/4-CF<sub>3</sub>), +122.6 (sept.,  ${}^{3}J_{\rm CF}$  = 3.7 Hz, 3-CH),; <sup>19</sup>F NMR (470.67 MHz, CD<sub>3</sub>CN, 25 °C, δ): -59.5 (d,  $J_{\rm FH}$  = 6.8 Hz, 9F, 2-CF<sub>3</sub>), -62.6 (s, 9F, 4-CF<sub>3</sub>). Elemental analysis (calc. for C<sub>24</sub>H<sub>10</sub>B<sub>1</sub>F<sub>18</sub>Na): C 42.83 (42.76), H 1.57 (1.50).

### $Na[HB{2,5-(CF_3)_2C_6H_3}_3]Na[3-H]$

To a solution of 3 (0.275 g, 0.42 mmol) in toluene (8 cm<sup>3</sup>) was added Na[HBEt<sub>3</sub>] (0.43 cm<sup>3</sup>, 0.43 mmol, 1.0 M in toluene), the reaction mixture was stirred for 4 hours to give a colourless solution. The reaction mixture was concentrated *in vacuo* reducing its volume to ca. 2 cm<sup>3</sup>, and the product crystallized by addition of petroleum ether and cooling to -25 °C. The white crystalline solid was isolated by filtration and dried *in vacuo*. Yield 0.207 g (0.31 mmol, 74%).

<sup>1</sup>H NMR (500.21 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): +7.70 (d,  ${}^{3}J_{\rm HH} = 8.2$  Hz, 3H, 3/4-H), +7.45 (d,  ${}^{3}J_{\rm HH} = 8.5$  Hz, 3H, 3/4-H), +7.23 (br.s, 3H, 6-H), +3.06 (br.qq,  ${}^{1}J_{\rm HB} = 82$  Hz,  $J_{\rm HF} \approx 9.1$  Hz, 1H, BH);  ${}^{11}$ B NMR (160.49 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): -14.23 (d,  ${}^{1}J_{\rm BH} = 84$  Hz);  ${}^{13}$ C{ ${}^{1}$ H} NMR (125.78 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): +134.8 (q,  ${}^{2}J_{\rm CF} = 28.0$  Hz, 2/5-C), +133.7 (q,  ${}^{3}J_{\rm CF} = 3.7$  Hz, 3/4/6-C), +132.3 (q,  ${}^{2}J_{\rm CF} = 31.0$  Hz, 2/5-C), +127.1 (q,  ${}^{1}J_{\rm CF} = 275$  Hz, 2/5-CF<sub>3</sub>), +126.3 (br.m, 3/4/6-C), +124.9 (q,  ${}^{1}J_{\rm CF} = 273$  Hz, 2/5-CF<sub>3</sub>), +121.6 (q,  ${}^{3}J_{\rm CF}$  3.9 Hz, 3/4/6-C);  ${}^{19}$ F NMR (470.67 MHz, CD<sub>2</sub>Cl<sub>2</sub>, 25 °C, δ): -59.3 (d,  $J_{\rm FH} = 7.6$  Hz, 9F, 2-CF<sub>3</sub>), -63.6 (s, 9F, 5-CF<sub>3</sub>). Elemental analysis (calc. for C<sub>24</sub>H<sub>10</sub>B<sub>1</sub>F<sub>18</sub>Na): C 42.93 (42.76), H 1.61 (1.50).

# Acknowledgements

G. G. W. and A. E. A. thank the Royal Society for financial support by University Research Fellowships. E. J. L. thanks the EPSRC for financial support by a DTA studentship under grant number EP/J500409. The research leading to these results has received funding from the European Research Council under the ERC Grant Agreement no. 307061 (PiHOMER).

We thank Dr C. MacDonald (University of East Anglia) for assistance with the <sup>1</sup>H and <sup>19</sup>F DOSY NMR experiments.

We acknowledge the use of the EPSRC funded National Chemical Database Service hosted by the Royal Society of Chemistry, and the EPSRC UK National Mass Spectrometry Facility (NMSF) at the University of Swansea. We thank the EPSRC UK National Crystallography Service at the University of Southampton for the collection of the crystallographic data. We thank the Research Computing Service at the University of East Anglia for access to the high performance computing cluster.

## References

- 1 G. C. Welch and D. W. Stephan, *J. Am. Chem. Soc.*, 2007, 129, 1880–1881.
- 2 D. W. Stephan, Dalton Trans., 2009, 3129-3136.
- 3 D. W. Stephan and G. Erker, *Angew. Chem., Int. Ed.*, 2010, 49, 46–76.
- 4 D. W. Stephan, Comp. Inorg. Chem. II, 2013, 1, 1069-1103.
- 5 D. W. Stephan and G. Erker, *Chem. Sci.*, 2014, 5, 2625-2641.
- 6 D. W. Stephan, Acc. Chem. Res., 2015, 48, 306-316.

7 D. W. Stephan and G. Erker, *Angew. Chem., Int. Ed.*, 2015, 54, 6400–6441.

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- 8 P. A. Chase, G. C. Welch, T. Jurca and D. W. Stephan, *Angew. Chem., Int. Ed.*, 2007, **46**, 8050–8053.
- 9 K. Chernichenko, Á. Madarász, I. Pápai, M. Nieger, M. Leskelä and T. Repo, *Nat. Chem.*, 2013, 5, 718–723.
- 10 H. Wang, R. Fröhlich, G. Kehra and G. Erker, *Chem. Commun.*, 2008, 5966–5968.
- 11 T. Mahdi and D. W. Stephan, *J. Am. Chem. Soc.*, 2012, **136**, 15809–15812.
- 12 D. J. Scott, M. J. Fuchter and A. E. Ashley, *J. Am. Chem. Soc.*, 2014, **136**, 15813–15816.
- 13 G. Ménard, T. M. Gilbert, J. A. Hatnean, A. Kraft, I. Krossing and D. W. Stephan, *Organometallics*, 2013, 32, 4416–4422.
- 14 Z. Lu, Y. Wang, J. Liu, Y. Lin, Z. H. Li and H. Wang, *Organometallics*, 2013, **32**, 6753–6758.
- 15 R. L. Melen, Chem. Commun., 2014, 50, 1161-1174.
- 16 S. C. Binding, H. Zaher, F. M. Chadwicka and D. O'Hare, *Dalton Trans.*, 2012, **41**, 9061–9066.
- 17 D. J. Scott, M. J. Fuchter and A. E. Ashley, *Angew. Chem.*, Int. Ed., 2014, 53, 10218–10222.
- 18 T. J. Herrington, A. J. W. Thom, A. J. P. White and A. E. Ashley, *Dalton Trans.*, 2012, **41**, 9019–9022.
- 19 J. M. Farrell, J. A. Hatnean and D. W. Stephan, J. Am. Chem. Soc., 2012, 134, 15728–15731.
- 20 E. J. Lawrence, T. J. Herrington, A. E. Ashley and G. G. Wildgoose, *Angew. Chem.*, *Int. Ed.*, 2014, 53, 9922– 9925.
- 21 G. Ménard and D. W. Stephan, Angew. Chem., Int. Ed., 2012, 53, 8272–8275.
- 22 E. R. Clark and M. J. Ingleson, Angew. Chem., Int. Ed., 2014, 53, 11306–11309.
- 23 A. M. Chapman, M. F. Haddow and D. F. Wass, *J. Am. Chem. Soc.*, 2011, 133, 18463–18478.
- 24 X. Xu, G. Kehr, C. G. Daniliuc and G. Erker, *J. Am. Chem. Soc.*, 2015, **137**, 4550–4557.
- 25 E. J. Lawrence, V. S. Oganesyan, D. L. Hughes, A. E. Ashley and G. G. Wildgoose, *J. Am. Chem. Soc.*, 2014, 136, 6031– 6036.
- 26 E. J. Lawrence, R. J. Blagg, D. L. Hughes, A. E. Ashley and G. G. Wildgoose, *Chem. – Eur. J.*, 2015, 21, 900– 906.
- 27 E. L. Kolychev, T. Bannenberg, M. Freytag, C. G. Daniliuc, P. G. Jones and P. D. M. Tamm, *Chem. – Eur. J.*, 2012, **18**, 16938–16946.
- 28 S. M. Cornet, K. B. Dillon, C. D. Entwistle, M. A. Fox, A. E. Goeta, H. P. Goodwin, T. B. Marder and A. L. Thompson, *Dalton Trans.*, 2003, 4395–4405.
- 29 W. V. Konze, B. L. Scott and G. J. Kubas, *Chem. Commun.*, 1999, 1807–1808.
- 30 S. J. Coles and P. A. Gale, Chem. Sci., 2012, 3, 683-689.
- 31 E. J. Lawrence, V. S. Oganesyan, G. G. Wildgoose and A. E. Ashley, *Dalton Trans.*, 2013, 42, 782–789.

- 32 D. H. McDaniel and H. C. Brown, *J. Org. Chem.*, 1958, 23, 420-427.
- 33 T. A. Rokob, A. Hamza and I. Pápai, *J. Am. Chem. Soc.*, 2009, **131**, 10701–10710.
- 34 I. B. Sivaev and V. I. Bregadze, *Coord. Chem. Rev.*, 2014, 270-271, 75-88.
- 35 Z. M. Heiden and A. P. Lathem, *Organometallics*, 2015, 34, 1818–1827.
- 36 G. R. Fulmer, A. J. M. Miller, N. H. Sherden, H. E. Gottlieb, A. Nudelman, B. M. Stoltz, J. E. Bercaw and K. I. Goldberg, *Organometallics*, 2010, 29, 2176–2179.
- 37 D. H. Wu, A. D. Chen and C. S. Johnson, J. Magn. Reson., Ser. A, 1995, 115, 260–264.
- 38 L. J. Bourhis, O. V. Dolomanov, R. J. Gildea, J. A. K. Howard and H. Puschmann, *Acta Crystallogr., Sect. A: Fundam. Crystallogr.*, 2015, 71, 59–75; O. V. Dolomanov, L. J. Bourhis, R. J. Gildea, J. A. K. Howard and H. Puschmann, *J. Appl. Crystallogr.*, 2009, 42, 339–341.
- 39 L. Palatinus and G. Chapuis, J. Appl. Crystallogr., 2007, 40, 786–790; L. Palatinus and A. van der Lee, J. Appl. Crystallogr., 2008, 41, 975–984; L. Palatinus, S. J. Prathapab and S. van Smaalen, J. Appl. Crystallogr., 2012, 45, 575–580.
- 40 G. M. Sheldrick, Acta Crystallogr., Sect. C: Cryst. Struct. Commun., 2015, 71, 3–8; G. M. Sheldrick, Acta Crystallogr., Sect. A: Fundam. Crystallogr., 2008, 64, 112–122.
- 41 S. Lancaster, *ChemSpider Synthetic Pages*, 2003, DOI: 10.1039/SP215, http://cssp.chemspider.com/215; T. E. Krattf, Boulder Scientific Company, *Eur. Pat*, 1186608, 2002; R. J. LeSuer, C. Buttolph and W. E. Geiger, *Anal. Chem.*, 2004, 76, 6395–6401.
- 42 DigiElch-Professional, ElechSoft, Kleinromstedt, Germany.
- 43 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, J. E. Peralta Jr., F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, Gaussian 09, revision A.02, Gaussian, Inc., Wallingford, CT, 2009.
- 44 A. D. Becke, J. Chem. Phys., 1993, 98, 5648-5652.
- 45 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter*, 1988, 37, 785–789.
- 46 P. C. Hariharan and J. A. Pople, *Theor. Chim. Acta*, 1973, **28**, 213–222.