# Dalton Transactions



PAPER View Article Online
View Journal | View Issue

hydrogenations†



Cite this: Dalton Trans., 2017, 46,

K. Chernichenko,<sup>a</sup> B. Kótai,<sup>b</sup> M. Nieger,<sup>a</sup> S. Heikkinen,<sup>a</sup> I. Pápai\*<sup>b</sup> and T. Repo\*<sup>a</sup>

Replacing C<sub>6</sub>F<sub>5</sub> groups with Cl and H atoms in

frustrated Lewis pairs: H<sub>2</sub> additions and catalytic

a few synthetic steps and demonstrated the cleavage of  $H_2$  under mild conditions. Depending on the nature of the dialkylamino group, X, and Z, the stability of the produced zwitterionic  $H_2$  adducts varies from isolated solids indefinitely stable in an inert atmosphere to those quickly equilibrating with the initial aminoborane and  $H_2$ . Using a combined experimental/computational approach on a series of isostructural aminoboranes (dialkylamino = 2,2,6,6-tetramethylpiperid-1-yl), it was demonstrated that the electronegativity and the steric effect of the substituents generally follow the trend  $C_6F_5 \sim Cl \gg H$ . This observation is useful for designing new FLPs for practical applications. As an example, we demonstrated the hydrogenation of alkynes to *cis*-alkenes under mild conditions that was catalyzed by a chloro-analogue of the  $C_6F_5$ -substituted aminoborane developed previously. The presence of a BHCl group in the aminochloroboranes or in their  $H_2$  adducts features facile redistribution of the H and Cl atoms and the formation of polychloro and polyhydrido species.

2-(Dialkylamino)phenylboranes containing the BXZ group, where X,  $Z = C_6F_5$ , Cl, and H, were prepared in

Received 8th December 2016, Accepted 10th January 2017 DOI: 10.1039/c6dt04649e

rsc li/dalton

## Introduction

High Lewis acidity and hydrolytic stability of (perfluoroaryl)-boranes have uniquely positioned these compounds as catalysts in organic synthesis and  $\alpha$ -olefin polymerization. Recently, such boranes in combination with sterically demanding amines and phosphines have shown unprecedented reactivities as components of frustrated Lewis pairs (FLPs). Particularly, metal-free heterolytic  $H_2$  splitting and its transfer to other organic molecules in a catalytic fashion have been fruitfully explored.

Motivated by the development of cost-efficient and light weight FLPs for catalytic applications, we have been studying ansa-aminoboranes (where "ansa" refers to the close vicinity of amino and boryl groups), in which the  $C_6F_5$  groups of the borane moiety are replaced with elemental substituents X

<sup>†</sup>Electronic supplementary information (ESI) available: Experimental procedures, NMR spectra, crystallographic data, and detailed computational analysis. Crystallographic data (excluding structure factors) for the structures reported in this work. CCDC 1511243 (2c), 912583 (4c), 912582 (4e), 912585 (5c), and 912584 (5e). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6dt04649e

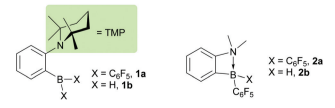


Fig. 1 Previously reported 2-(dialkylamino)phenylborane FLPs.

<sup>(</sup>where X = H, halogens). Recently, we have reported two archetypical C<sub>6</sub>F<sub>5</sub>-substituted ortho-aminophenylboranes, 1a and 2a differing in the Lewis basic amino component (Fig. 1).5 The presence of a highly sterically demanding 2,2,6,6-tetramethylpiperid-1-yl amino group (TMP) and a sterically accessible dimethylamino (Me2N) group substantially affected the thermodynamics and the reactivity of H2. Whereas 1a produced an extremely thermally stable H2 adduct, 2a reacted with H2 reversibly, showing smooth intramolecular protonation<sup>6</sup> and other unexpected behaviour. The replacement of a single C<sub>6</sub>F<sub>5</sub> group with H in 2a provided 2b serving as a catalyst in an unprecedented metal-free selective hydrogenation of alkynes into cis-alkenes. Aminoborane 2b has also been shown to insert readily into sp<sup>2</sup>-C-H bonds of simple arenes and alkenes.<sup>7</sup> On the other hand, the complete replacement of the C<sub>6</sub>F<sub>5</sub> groups in 1a with hydrogens gave aminoborane 1b that activates H<sub>2</sub> reversibly<sup>8</sup> and efficiently catalyses the C-H borylation of

<sup>&</sup>lt;sup>a</sup>Department of Chemistry, University of Helsinki, P.O. Box 55, FIN-00014, Finland. E-mail: timo.repo@helsinki.fi

<sup>&</sup>lt;sup>b</sup>Research Centre for Natural Sciences, Hungarian Academy of Sciences, Magyar tudósok körútja 2, H-1117 Budapest, Budapest, Hungary. E-mail: papai.imre@ttk.mta.hu

**Paper** 

hetarenes with pinacolborane.9 In continuation of our efforts,

we report herein new ansa-aminoboranes, the derivatives of 1 and 2, in which the C<sub>6</sub>F<sub>5</sub> groups are partially or completely replaced with Cl or H atoms. 10 We studied H2 addition to these aminoboranes following the established dichotomy between ortho-TMP- and ortho-Me2N-phenylboranes such that the former defined general reactivity patterns, whereas the more labile and reactive Me<sub>2</sub>N compounds were used for catalytic implementations.

According to spectroscopic Lewis acidity scales, inorganic boranes BX<sub>3</sub> (X = H or halogen) have similar acidities to  $B(C_6F_5)_3$ . These data are supported by experimental results on the H2 splitting by FLPs comprising chloroboranes as the Lewis acidic component. 12 At the same time, comparative reactivity studies of isostructural FLPs with systematic  $C_6F_5 \rightarrow Cl$ replacement at the Lewis acidic site and motivated by the development of catalytic applications have never been addressed previously and, therefore, are of particular interest.

## Results and discussion

### Synthesis and characterization of new ansa-aminoboranes

Chloroboranes 1c and 1e were prepared in one step starting from a readily available lithium compound 3 5 and BCl3 or C<sub>6</sub>F<sub>5</sub>BCl<sub>2</sub>, <sup>13</sup> respectively (Scheme 1). Both aminoboranes were isolated in close to quantitative yields, similar to the previously reported 1a. Apparently, high steric bulkiness of the TMP group suppressed the double addition of 3 to the starting boranes. Reduction of dichloroborane 1c with 2 eq. of Me<sub>3</sub>SnH<sup>14</sup> provides an alternative approach to a dimeric ansaaminodihydroborane 1b (Scheme 1) that was previously reported by us.9 With smaller amounts of Me<sub>3</sub>SnH, ansaaminochloroborane 1d is formed. In solution, it does not exist individually, but it forms an equilibrium with 1c and 1b. The equilibrium is instantly established at room temperature and even at -15 °C due to the rapid B-H/B-Cl exchange. The equilibrium state is slightly shifted to 1d in aromatic hydrocarbons and strongly in more polar dichloromethane- $d_2$  and 1,2dichloroethane (see the highlighted part of Scheme 1). 15

Scheme 1 Synthesis of aminoboranes 1b-1e

Frustrated aminoboranes can exist in several forms as illustrated in Scheme 2. The intramolecular N-B dative adducts and the µ-H-bridged dimeric species possess a reduced reactivity potential in comparison to the unquenched open structures. The aminoboranes 1a, 1e, and 1c exist in their open forms as evident by the 11B NMR shifts typical of noncoordinated boranes: 55.8, 62.2 and 62.3 ppm, respectively. 16 A combination of highly sterically demanding TMP and  $B(C_6F_5)_2$  moieties in 1a prevents the formation of an intramolecular N  $\rightarrow$  B dative bond. Despite the smaller size of a chlorine atom as compared to the C<sub>6</sub>F<sub>5</sub> group, both 1c and 1e have unquenched acid/base sites. In line with the experimental findings, DFT calculations predict open equilibrium structures for 1a, 1e, and 1c.

The closed forms (i.e. four-membered ring structures with internal B-N dative bonds) could not be identified as energy minima on the potential energy surfaces. Computations point to the coexistence of two conformers for these aminoboranes with the phenylene bridge occupying either the equatorial (structure A) or the axial position (structure B, Scheme 2b) of the piperidine ring.<sup>17</sup> The former structure is predicted to be slightly more favoured for all aminoboranes 1a, 1e, and 1c (for details, see the ESI†). Monochloroborane 1d appears as a doublet in the 11B NMR spectrum evidencing its monomeric form. Variable temperature (-12-90 °C, in toluene- $d_8$ ) <sup>11</sup>B NMR spectroscopy revealed a strong drift in the chemical shift of 1d ( $\delta$  = 20-42 ppm) attributed to a very rapid equilibrium between its open and dative forms, which is supported by calculations as well (see the ESI†). We showed previously that the trans-dimeric form of dihydroborane 1b dominates in solutions whereas in the solid state it is the exclusive form as evident from X-ray diffraction analysis.8

a) TMP 
$$\frac{1}{2} \sum_{X} \sum$$

b) 
$$X = C_6F_5, 1a$$

$$X = C_1, 1c$$

$$X = C_1, 1c$$

$$X = H, open-1b$$

Scheme 2 (a) Appearance of ansa-TMP-phenylboranes as the open and the quenched forms; (b) conformational variation in compounds

**Dalton Transactions** Paper

#### Addition of H2 to the ansa-aminoboranes

As solutions in hydrocarbons or in chlorinated hydrocarbons, aminochloroboranes 1c, 1e and 1d react with H<sub>2</sub> (2 bar) within the first few minutes at room temperature, producing the respective ammonium chloroborohydrides 4c, 4e and 4d. Compounds 4c and 4e were isolated almost quantitatively as white crystalline powders indefinitely stable under an inert atmosphere.

Owing to the existing equilibrium between 1d, 1c, and 1b in solutions, the reaction with H2 "freezes" it to some extent, producing mixtures of chloroborodihydride 4d contaminated with varying amounts of 4c and 1b (Scheme 3). Dichloromethane and 1,2-dichloroethane are advantageous solvents for producing mixtures rich in 4d owing to the higher content of 1d in these solvents. Previously, we reported that the addition of H<sub>2</sub> to ortho-TMP-dihydroborane 1b is a rapid and thermodynamically nearly neutral process. The equilibrium can thus be shifted towards the H<sub>2</sub> adduct 4b by using a more polar solvent, higher H2 pressure and low temperatures (72% conversion in  $CD_2Cl_2$ , 10 bar  $H_2$ , -15 °C).

The solid state structures of H2 adducts 4c and 4e were determined using single crystal X-ray diffraction (Fig. 2). The structure of 4c displays the proximity of the NH and BH hydrogens pointing to the existence of a dihydrogen bond similarly to that observed for analogous ansa-aminoborane-H2 adducts. 5,18 Interestingly, the X-ray structure of 4e does not involve this type of interaction, but instead, H...Cl bond formation is apparent. To characterize the structure of dihydrogen adducts 4a, 4c-4e in dichloromethane solution, the

Scheme 3 Addition of H<sub>2</sub> to aminoboranes 1c, 1d and 1e

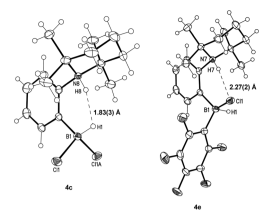


Fig. 2 Structures of chloroborohydrides 4c and 4e in a solid state (displacement parameters are drawn at the 50% probability level).

H<sub>N</sub>-H<sub>B</sub> bond lengths were studied by 1D NOE <sup>1</sup>H NMR spectroscopy and they were compared to data from DFT calculations (see the ESI† for details). Similarly to the solid state, a pronounced preference for the dihydrogen-bonded isomer in solution was established for 4c by both methods. Adduct 4d could not be isolated in the pure form, therefore, only solution-phase computational and NOE data are available, which indicate that dihydrogen-bonded species are clearly favoured in DCM solutions.

#### Computational study of H<sub>2</sub> addition to ansa-aminoboranes 1a-c

The results reported above point to the similar reactivities of C<sub>6</sub>F<sub>5</sub>- and chloro-substituted ansa-aminoboranes, but also to a somewhat different behaviour of 1b. To rationalize the observed reactivities, hydrogen addition to compounds 1a-1c was studied by DFT calculations. The results are summarized in Fig. 3.

The structures of the transition states located along the H<sub>2</sub> splitting pathway (TS<sub>1a</sub>, TS<sub>1b</sub>, TS<sub>1c</sub> in Fig. 3) share common features with those of the previously investigated FLP systems.<sup>19</sup> The slightly elongated H-H bond, the pyramidalization of the borane unit, and the typical end-on N···H<sub>2</sub> and side-on H<sub>2</sub>···B arrangements of the reacting partners are all in line with the electron transfer reactivity model.<sup>20</sup> In the case of **1a** and **1c**, the activation barriers are fairly low ( $\Delta G^{\ddagger} = 17.7$  and 16.1 kcal mol<sup>-1</sup>, respectively),<sup>21</sup> which is consistent with the observed reaction rates. Likewise, the thermodynamics of H2 additions to 1a and 1c, resulting in 4a and 4c, are substantially exergonic and the computed reaction free energies are similar ( $\Delta G_{\rm r}$  = -12.0 and -11.1 kcal mol<sup>-1</sup>). Although the open form of aminoborane **1b** is still rather reactive with an unprecedentedly low barrier  $(TS_{1b} \text{ is only } 11.8 \text{ kcal mol}^{-1} \text{ above } open-1b + H_2), \text{ the overall}$ barrier is predicted to be slightly higher (20.5 kcal mol<sup>-1</sup>) than those with 1a and 1c, which is clearly due to the reactant state stabilization arising from dimerization. For the same reason, the reaction with 1b becomes thermodynamically less favoured as well (slightly endergonic in toluene).

Naturally, the trend obtained for the Gibbs free energies of the reaction is closely related to the variation of the Lewis

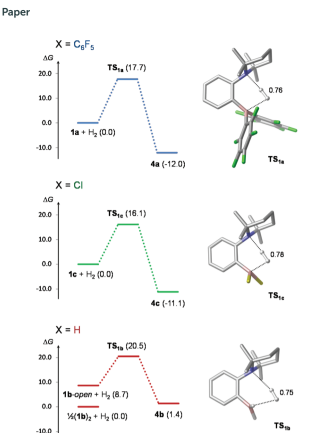
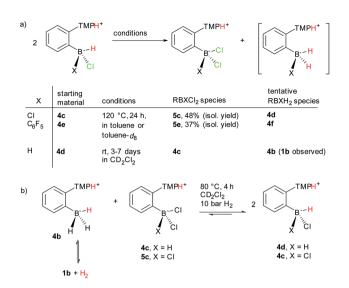


Fig. 3 Computed Gibbs free energy profiles for dihydrogen activation by 1a, 1b and 1c. Relative stabilities are given in parenthesis (in kcal  $\text{mol}^{-1}$ ; with respect to separated reactants; solvent = toluene). H–H bond distances are in Å (the bond length of free  $H_2$  is 0.74 Å). In TS structures, CH hydrogens are omitted for clarity.

acidity of boryl units in the **1a–1c** series. In light of the hydride affinities of  $B(C_6F_5)_3$ ,  $BCl_3$  and  $BH_3$  boranes ( $\Delta G_{ha} = -72.5$ , -64.2 and -46.3 kcal  $mol^{-1}$ , respectively),<sup>22</sup> one expects somewhat larger differences between the thermodynamics of  $H_2$  addition to the corresponding aminoboranes **1a**, **1c** and **1b**. However, our energy decomposition analysis reveals that the proton affinity of the TMP group is notably influenced by the nature of the boryl substituent, and also that the acid–base cooperativity taking place through the *ortho*-phenylene linker in these aminoboranes is an important factor.<sup>23</sup> This self-compensatory reactivity potential mechanism operating *via* a conjugated phenylene linker is a remarkable feature of the *ortho*-aminophenylborane FLPs.

## Thermal behaviour of H<sub>2</sub> adducts

Unlike **4b**,  $H_2$  adducts **4c–4e** do not demonstrate reverse hydrogen release, but instead they tend to decompose under certain conditions (Scheme 4). Compound **4d** has limited stability in  $CD_2Cl_2$  solution dismutating to **4c** and presumably **4b** upon standing at room temperature for several days. Upon heating of **4c** or **4e** for 24 h at 120 °C in toluene, tri- **5c** and dichloroborate **5e** are isolated in 48% and 37% yields, respectively (Scheme 4a), as crystalline solids precipitating



Scheme 4 (a) Decomposition of 4c, 4e and 4d with the formation of chloroborates 5c, 5e and 4c; (b) formation of 4c and 4d via "retrodismutation".

from the solution upon cooling (for X-ray structures, see the ESI†). The filtrate solution is a complex mixture of unidentified products, except for  $C_6F_5H$ , that is formed in an equimolar amount to  $\bf 4e$ , as evident from  $^{19}F$  and  $^{1}H$  NMR spectroscopies. We suggest that the B-H/B-Cl exchanging dismutation of  $\bf 4c$  and  $\bf 4d$  takes place at elevated temperatures and progresses until reaching the ultimate trichloro-  $\bf 5c$  and trihydroborate species  $\bf 4b$ , whereas  $\bf 4b$  decomposes into  $\bf 1b$  and  $\bf H_2$ . Since  $\bf 1b$  is not detected among the products, we presume that it is unstable under harsh reaction conditions.

Additional evidence for such a decomposition pathway is provided by demonstration of a "retrodismutation" reaction: dichloroborohydride  $\bf 4c$ , aminoborane  $\bf 1b$  and  $\bf H_2$  produced  $\bf 4d$  upon heating for 4 h at 10 bar  $\bf H_2$  pressure and 80 °C. Similarly, the reaction between trichloroborate  $\bf 5c$ ,  $\bf 1b$  and  $\bf H_2$  results in the formation of varying amounts of  $\bf 4c$  and  $\bf 4d$  with their ratio depending on the ratio of the starting materials. Trichloroborane  $\bf 5c$  can be completely converted into  $\bf 4c$  and  $\bf 4b$  provided  $\bf 1b$  is present in sufficient amounts (Scheme  $\bf 4b$ ).

The formation of the B-H/B-Cl exchange products during the addition of  $H_2$  to the  $ClB(C_6F_5)_2/2,2,6,6$ -tetramethylpiperidine and  $BCl_3/2,6$ -dimethylpyridine FLPs was reported previously. In the absence of the stabilizing factors, the easy redistribution of Cl and H atoms between chloro- and hydroborates seems to be a common reactivity pattern for these species. To gain deeper insight into the thermally-promoted transformations of 2-(TMP)-phenyl-chloroboranes and their adducts, we examined a series of reactions involving various  $H_2$  and HCl addition/elimination steps computationally as shown in Scheme 5. The results are summarized in Fig. 4 in the form of a free energy profile.

It is apparent from this profile that the adduct 4c lies in a free energy minimum with respect to  $H_2$  and HCl elimination. The barrier towards  $H_2$  elimination is notably lower, therefore

**Dalton Transactions** 

Scheme 5 Series of reactions investigated computationally

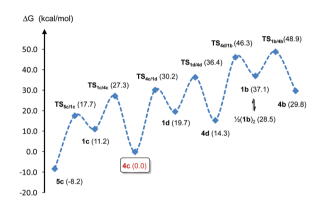


Fig. 4 Computed Gibbs free energy profile for the series of reactions shown in Scheme 5. The zero level of the diagram is arbitrarily chosen at 4c.

 $4c \rightarrow 1c + H_2$  might be the first step of the thermally induced transformation and decomposition. Although H2 elimination from 4c is unfavoured thermodynamically, this reaction may shift towards the formation of 1c as H2 is continuously discharged from the solution in these experiments.

The reaction between 1c and 4c to produce 5c and 4d is thermodynamically feasible as calculations predict  $\Delta G_r$  = 0.3 kcal mol<sup>-1</sup> in toluene and 1.4 kcal mol<sup>-1</sup> in DCM for this process. We found that this transformation can occur in a single step via a concerted H-/Cl- exchange (for the identified transition states, see the ESI†). The related activation barrier is fairly high  $(\Delta G^{\ddagger} = 30.2 \text{ kcal mol}^{-1} \text{ in toluene})$ and 26.9 kcal mol<sup>-1</sup> in DCM), but it is consistent with the experimental conditions (120 °C, 24 h).

As for the destiny of the tentative 4f formed via the H/Cl redistribution at the initial stage of 4e thermolysis (Scheme 4a), we suggest that it decomposes by the intramolecular protonative splitting of the B-C<sub>6</sub>F<sub>5</sub> bond that produces 1b and C<sub>6</sub>F<sub>5</sub>H as detected experimentally. Such a reaction was previously shown to proceed surprisingly easily in the ortho-aminophenylborane core. Besides, we revised the thermal behaviour of compound 4a and found that its decomposition via a similar protonative pathway becomes apparent at 150 °C (see the ESI† for details).

#### Catalytic hydrogenations

Recently, we have reported the highly cis-selective semihydrogenation of internal alkynes catalysed by ansa-aminohydroborane **2b** generated *in situ* from aminoborane **2a** (Scheme 6). The ansa-phenylene junction of the active B and N centres in 2b proved to be essential for such a catalytic activity based on the well-established reaction mechanism. Herein we report the similar catalytic activity of aminoborane 2c (Table 1), a light weight chloro analogue of 2a, prepared in 40% yield via a simple three-step protocol from inexpensive starting materials: N,N-dimethylaniline, butyllithium and boron trichloride (Scheme 6).

Internal alkynes were converted into respective cis-alkenes within 24 h or less at 100 °C and 2.2 bar H<sub>2</sub> using 2c as a catalyst. Remarkably, sterically hindered amine 1,2,2,6,6-pentamethylpiperidine (6) serves as an efficient promoter enhancing

2a: 
$$X$$
,  $Z = C_0F_5$   
2b:  $X = C_0F_5$ ,  $Z = H$   
2c:  $X$ ,  $Z = CI$   
1) BuLi, 5 mol. % TMEDA, hexane or cyclohexane, 70 °C, 18h  
2) 2 eq. BCl<sub>3</sub>, toluene, -80 °C  
3) 80 °C, 90 min, vacuum

Scheme 6 ansa-Aminoboranes 2a and 2b reported recently to catalyse the hydrogenation of alkynes and the synthesis of the isostructural chloro-analogue 2c. X-ray diffraction structure of 2c (displacement parameters are drawn at the 50% probability level).

Table 1 cis-Selective semi-hydrogenation of internal alkynes catalysed by 2ca

$$R^{1} = -R^{2} \qquad \begin{array}{c} 2.2 \text{ bar } H_{2}, \text{ CI}(\text{CH}_{2})_{2}\text{CI} \\ \\ \underline{5 \text{ mol.}\% 2c, 5 \text{ mol.}\% 6} \\ \\ 100 \text{ °C} \end{array} \qquad \begin{array}{c} H \\ R^{1} \\ \\ R^{2} \end{array}$$

Substrate	<b>2c</b> , mol%	Time, h	Conversion <sup>b</sup> (Isol. yield), %
_=/	5	3	100
	5	24	100
SiEt <sub>3</sub>	5	24	56
	5	24	100 (90)
	10	24	92

<sup>a</sup> 125 ml Schlenk tube was charged with 0.5 mmol of alkyne, a catalytic amount of 2c and 6 and 0.35 ml of 1,2-dichloroethane, pressurized with  $H_2$  (2.2 bar) and stirred at respective temperatures.  $^{\dot{b}}$  Conversions were determined by the <sup>1</sup>H NMR analysis of crude reaction mixtures.

**Paper** 

the catalytic activity approximately two fold. Under standard conditions, only 5 mol% of both 2c and 6 loadings are sufficient for reaching complete conversions of acetylenes. At the same time 2a appears to be more catalytically active than 2c, because the majority of substrates are completely hydrogenated with the aid of 2a in 3 h at 80 °C. Regarding the feasibility of catalysis and high cis-stereoselectivity during hydrogenations, we suggest that the mechanism of catalysis by 2c is very similar to the one previously reported for 2a/2b though the details are yet to be established in the ongoing studies.

# Conclusions

In our present work, we studied structural analogues of previously reported frustrated 2-aminophenylboranes 2-(Alk<sub>2</sub>N)-C<sub>6</sub>H<sub>4</sub>-B(C<sub>6</sub>F<sub>5</sub>)<sub>2</sub>, in which C<sub>6</sub>F<sub>5</sub> groups were partially or completely replaced with H or Cl atoms. With the Alk2N group represented by 2,2,6,6-tetramethylpiperid-1-yl, all the considered aminoboranes react with H<sub>2</sub> within minutes at room temperature. We found strong similarities between C<sub>6</sub>F<sub>5</sub>-substituted and chloro-substituted boranes in their reactivities as well as the energetic and kinetic parameters of H2 addition. At the same time, the replacement of C<sub>6</sub>F<sub>5</sub> or Cl with H atoms leads to a significant drop in the reactivity potential, mainly due to the formation of the quenched forms of the starting B-Hsubstituted aminoboranes. This is consistent with the FLP concept as the compact size of the H atom cannot provide sufficient steric separation of the Lewis acidic and basic centres in the aminoboranes. On the other hand, our computations revealed a self-compensatory mechanism for this class of FLPs: more Lewis acidic boryl units diminish the basicity of the TMP group via the phenylene ring. Consequently, the energetics of H2 addition to the aminoboranes that vary in the boryl part  $(B(C_6F_5)_2, BCl_2, BH_2)$  differs less than one expects from the comparison of the Lewis acidities of the corresponding parental boranes alone.

The attempted thermally promoted dehydrogenation of ammonium chloroborohydrides (H2 adducts) leads to the redistribution of B-H and B-Cl substituents resulting in the isolation of polychloroborates 5c and 5e. These processes are feasible only under conditions when H<sub>2</sub> is discharged from the reaction as shown by the reversible formation of chloroborohydrides in "retrodismutation" experiments. For C<sub>6</sub>F<sub>5</sub>-substituted borates the decomposition involves protonative cleavage of the B-C<sub>6</sub>F<sub>5</sub> bond yielding C<sub>6</sub>F<sub>5</sub>H. In the molecules of the studied ammonium chloroborohydrides, a protic hydrogen atom can be connected to either a Cl or H atom of the BH(Cl)X unit through intramolecular Cl···H or dihydrogen bonds. We found that these forms are usually nearly equal in energy and can be easily interconverted *via* rotation around the B-C bond.

Experimental and computational comparisons between isostructural chloro- and C<sub>6</sub>F<sub>5</sub>-substituted aminoboranes revealed a high degree of similarity in reactivities to H<sub>2</sub>, which is reflected by the energetics of the overall reactions and transition states as well as by the stability of H2 adducts. This similarity was pronouncedly demonstrated by the similar catalytic abilities of chloro- and C<sub>6</sub>F<sub>5</sub>-substituted aminoboranes 2c and 2a in the hydrogenation of alkynes. Simple and lightweight FLPs derived from boranes with elementary substituents are promising catalysts for hydrogenation and C-H borylation reactions and studies of their catalytic properties are currently in progress in our groups.

# Acknowledgements

This work was supported by the Academy of Finland (139550, 276586), COST action CM0905, and the Hungarian NKFI Grant (K-115660).

## Notes and references

- 1 (a) W. E. Piers, Adv. Organomet. Chem., 2004, 52, 1-76; (b) M. Rubin, V. Gevorgyan, S. Chandrasekhar, B. Nagendra Babu and G. Chandrashekar, Tris(pentafluorophenyl)borane in Encyclopedia of Reagents for Organic Synthesis, John Wiley & Sons, 2009, DOI: 10.1002/047084289X.rn00259.pub2.
- 2 E. Y.-X. Chen and T. J. Marks, Chem. Rev., 2000, 100, 1391-1434.
- 3 (a) Frustrated Lewis Pairs I: Uncovering Understanding, in *Topics in Current Chemistry*, ed. G. Erker and D. W. Stephan, Springer, Berlin, Heidelberg, 2013, vol. 332; (b) Frustrated Lewis Pairs II: Expanding the Scope, in Topics in Current Chemistry, ed. G. Erker and D. W. Stephan, Springer, Berlin, Heidelberg, 2013, vol. 334; (c) D. W. Stephan and G. Erker, Angew. Chem., Int. Ed., 2015, 54, 6400-6441; (d) D. W. Stephan, J. Am. Chem. Soc., 2015, 137, 10018-10032; (e) D. W. Stephan, Acc. Chem. Res., 2015, 48, 306-316.
- 4 For reviews on FLP-catalysed hydrogenations, see: (a) D. W. Stephan, S. Greenberg, T. W. Graham, P. Chase, J. J. Hastie, S. J. Geier, J. M. Farrell, C. C. Brown, Z. M. Heiden, G. C. Welch and M. Ullrich, Inorg. Chem., 2011, **50**, 12338–12348; (b) D. W. Stephan, Org. Biomol. Chem., 2012, 10, 5740-5746; (c) J. Paradies, Angew. Chem., Int. Ed., 2014, 53, 3552-3557; (d) L. J. Hounjet and D. W. Stephan, Org. Process Res. Dev., 2014, 18, 385-391. For a selection of the most recent achievements, see: (e) S. Tussing, L. Greb, S. Tamke, B. Schirmer, C. Muhle-Goll, B. Luy and J. Paradies, Chem. - Eur. J., 2015, 21, 8056-8059; (f) M. Lindqvist, K. Borre, K. Axenov, B. Kótai, M. Nieger, M. Leskelä, I. Pápai and T. Repo, J. Am. Chem. 2015, **137**, 4038-4041; (g) P. Eisenberger, B. P. Bestvater, E. C. Keske and C. M. Crudden, Angew. Chem., Int. Ed., 2015, 54, 2467-2471; (h) I. Chatterjee and M. Oestreich, Angew. Chem., Int. Ed., 2015, 54, 1965-1968; (i) Z. Zhang and H. Du, Angew. Chem., Int. Ed., 2015, 54, 623-626; (j) Á. Gyömöre, M. Bakos, T. Földes, I. Pápai, A. Domján and T. Soós, ACS Catal., 2015, 5, 5366-5372;

(k) D. J. Scott, T. R. Simmons, E. J. Lawrence, G. G. Wildgoose, M. J. Fuchter and A. E. Ashley, ACS Catal., 2015, 5, 5540–5544; (l) Z. Zhang and H. Du, Org. Lett., 2015, 17, 6266–6269; (m) S. Tussing, K. Kaupmees and J. Paradies, Chem. – Eur. J., 2016, 22, 7422–7426; (n) D. J. Scott, N. A. Phillips, J. S. Sapsford, A. C. Deacy, M. J. Fuchter and A. E. Ashley, Angew. Chem., Int. Ed., 2016, 55, 14738–14742.

**Dalton Transactions** 

- 5 K. Chernichenko, M. Nieger, M. Leskelä and T. Repo, *Dalton Trans.*, 2012, 41, 9029–9032.
- 6 K. Chernichenko, Á. Madarász, I. Pápai, M. Nieger, M. Leskelä and T. Repo, Nat. Chem., 2013, 5, 718–723.
- 7 K. Chernichenko, M. Lindqvist, B. Kótai, M. Nieger, K. Sorochkina, I. Pápai and T. Repo, J. Am. Chem. Soc., 2016, 138, 4860–4868.
- 8 K. Chernichenko, B. Kótai, I. Pápai, V. Zhivonitko, M. Nieger, M. Leskelä and T. Repo, *Angew. Chem., Int. Ed.*, 2015, **54**, 1749–1753.
- 9 (a) M. A. Légaré, M. A. Courtemanche, É. Rochette and F. G. Fontaine, *Science*, 2015, **349**, 513–516; (b) M. A. Légaré, É. Rochette, J. L. Lavergne, N. Bouchard and F. G. Fontaine, *Chem. Commun.*, 2016, **52**, 5387–5390. Very recently, aminoborane Me<sub>2</sub>N-C<sub>6</sub>H<sub>4</sub>-BH<sub>2</sub> was also synthesized by Fontaine *et al.*, and it was shown to undergo a dehydrogenative B-B homocoupling reaction. See: (c) É. Rochette, N. Bouchard, J. L. Lavergne, C. F. Matta and F. G. Fontaine, *Angew. Chem., Int. Ed.*, 2016, **55**, 12722–12726.
- 10 Preliminary results of this work have been included in a PhD thesis. See: K. Chernichenko, Ph.D. Thesis, University of Helsinki, Helsinki, Finland, 2013, http://urn.fi/URN: ISBN:978-952-10-9388-3.
- 11 For review publications on the Lewis acidity of boranes, see ref. 10, pp. 21–28, and I. B. Sivaev and V. I. Bregadze, *Coord. Chem. Rev.*, 2014, 270–271, 75–88.
- 12 (a) C. Jiang, O. Blacque and H. Berke, *Organometallics*, 2009, 28, 5233–5239; (b) B. Ginovska, T. Autrey, K. Parab, M. E. Bowden, R. G. Potter and D. M. Camaioni, *Chem. Eur. J.*, 2015, 44, 15713–15719; (c) E. R. Clark, D. A. Grosso and M. J. Ingleson, *Chem. Eur. J.*, 2013, 19, 2462–2466; (d) M.-A. Courtemanche, E. Rochette, M.-A. Legare, W. Bi and F.-G. Fontaine, *Dalton Trans.*, 2016, 45, 6129–6135.

- 13 R. D. Chambers and T. Chivers, J. Chem. Soc., 1965, 3933–3939.
- 14 We found Me<sub>3</sub>SnH to be extremely efficient for the conversion of chloroboranes to hydroboranes. A by-product, Me<sub>3</sub>SnCl, can be easily removed in a vacuum unlike when a more common tin hydride, Bu<sub>3</sub>SnH, is used. For preparation of Me<sub>3</sub>SnH, see: R. H. Fish, H. G. Kuivila and I. J. Tyminski, *J. Am. Chem. Soc.*, 1967, 89, 5861–5868.
- 15 Aminoborane 1c has been recently used by us as an intermediate for the preparation of its diarylboryl analogues: V. V. Zhivonitko, K. Sorochkina, K. Chernichenko, B. Kótai, T. Földes, I. Pápai, V.-V. Telkki, T. Repo and I. Koptyug, *Phys. Chem. Chem. Phys.*, 2016, 18, 27784–27795.
- 16 B. Wrackmeyer, Ann. R. NMR S., 1988, 20, 61-203.
- 17 DFT calculations were carried out using the dispersion-corrected range-separated hybrid ωB97X-D functional along with the 6-311G(d,p) basis set as implemented in Gaussian 09. The electronic energies were refined by single-point energy calculations using a larger basis set (6-311++G (3df,3pd)). The SMD continuum model was employed to describe solvation. The reported energies refer to solvent-phase Gibbs free energies. For further details, see the ESI.†
- (a) F. Schulz, V. Sumerin, S. Heikkinen, B. Pedersen,
   C. Wang, M. Atsumi, M. Leskelä, T. Repo, P. Pyykkö,
   W. Petry and B. Rieger, J. Am. Chem. Soc., 2011, 133, 20245–20257; (b) V. Sumerin, K. Chernichenko, M. Nieger,
   M. Leskelä, B. Rieger and T. Repo, Adv. Synth. Catal., 2011, 353, 2093–2110.
- 19 For a review of previous theoretical mechanistic studies on FLP-mediated H<sub>2</sub> activation, see: T. A. Rokob and I. Pápai in *Topics in current chemistry*, Springer, 2013, vol. 332, pp. 157–212.
- 20 T. A. Rokob, I. Bakó, A. Stirling, A. Hamza and I. Pápai, J. Am. Chem. Soc., 2013, 135, 4425–4437.
- 21 Solution-phase Gibbs free energies reported in Fig. 2 and 3 refer to toluene as a solvent. Results obtained for DCM are provided in the ESI.†
- 22 The hydride affinity values were obtained from M05-2X calculations as described in our previous work. See: T. A. Rokob, A. Hamza and I. Pápai, *J. Am. Chem. Soc.*, 2009, **131**, 10701–10710.
- 23 For a detailed energy decomposition analysis, see the ESI.†