



Cite this: *Chem. Commun.*, 2016, 52, 8389

Received 31st March 2016,
Accepted 3rd June 2016

DOI: 10.1039/c6cc02720b

www.rsc.org/chemcomm

Catalytic dehydrocoupling of amine-boranes and amines into diaminoboranes: isolation of a Pt(II), Shimoi-type, η^1 -BH complex†

Marta Roselló-Merino,^a Raquel J. Rama,^a Josefina Díez^b and Salvador Conejero*^a

The platinum complex [Pt(I^tBu')(I^tBu)][BAR^F] is a very efficient catalyst in the synthesis of diaminoboranes through dehydrocoupling of amine-boranes and amines. Shimoi-type, η^1 -BH complexes are key intermediates in the process.

In the last few years amine-boranes and related base-stabilised borane adducts have been shown to produce rich chemistry in which metal catalysed dehydrocoupling processes are involved.¹ During some of these reactions the B–H bonds of the amine borane establish an initial interaction with the metal centre to form complexes exhibiting η^1 or η^2 coordination modes, undergoing thereafter subsequent reactivities.^{1a,2}

Dehydrogenation of amine-boranes has been mainly focused on the production and release of dihydrogen and on the generation of amino-boranes that can undergo further dimerization, oligomerization or polymerization processes.^{1,2,3} Less attention has been paid to the production of other boranes, through boron–boron coupling⁴ or the formation of diaminoboranes, (NR₂)₂BH.⁵ With regard to the latter process, only a couple of catalytic processes have been reported by Alcaraz and Sabo-Etienne, using ruthenium^{5a,c} and by Hill,^{5b} with calcium and magnesium based catalysts to produce the corresponding diaminoboranes. Catalyst loadings of 2.5% (Ru, Ca, Mg), together with long reaction times (Ru), and even heating (Ca, Mg) were required.

In this communication we wish to report a very efficient platinum(II) complex that promotes diaminoborane formation at a catalyst loading of 0.5% in a few minutes for most of the substrates tested. In addition, we have been able to characterise

by means of X-ray diffraction studies the first Shimoi-type η^1 -BH complex of a Lewis base-borane adduct and platinum, which provides structural insights into a key intermediate in the dehydrogenation process of amine-boranes and amines.

Previously, we had reported that the [Pt(I^tBu')(I^tBu)][BAR^F] complex (where I^tBu stands for 1,3-di-*tert*-butylimidazolylidene and I^tBu' its cyclometalated form), **1**,⁶ is able to promote the dehydrocoupling of dimethylamineborane (NMe₂H·BH₃) into cyclic [NMe₂BH₂]₂ (Scheme 1).⁷ During NMR mechanistic studies, we observed that bis(dimethylamino)borane (NMe₂)₂BH is also formed in small amounts and its yield increases as the concentration of free NMe₂H in the reaction media increases. Therefore, we analysed the ability of complex **1** to act as a catalyst for the formation of diaminoboranes.

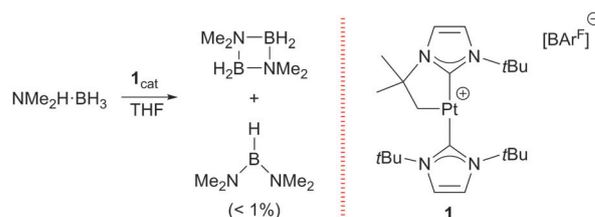
We have first studied the benchmark reaction of *tert*-butylamineborane (^tBuNH₂·BH₃) and 1 equiv. of *tert*-butylamine (^tBuNH₂) in CD₂Cl₂, using a catalyst loading of 0.5% (Scheme 2). The reaction was monitored both by NMR spectroscopy and by measuring the increase of gas pressure in a closed system due to the generated H₂ (see the ESI†). The ¹¹B NMR spectrum revealed the formation of a single species in nearly quantitative yield after 11.5 min showing a doublet signal at δ 25.8 ppm (¹J_{BH} = 127 Hz). In addition, the ¹H NMR spectrum exhibits a broad quartet signal centred at 4.12 ppm that sharpens upon ¹¹B decoupling into a triplet (*J*_{HH} = 8.2 Hz). These values are in agreement with the formation of diaminoborane (^tBuNH)₂BH,⁸ **s1** (Scheme 2). The calculated TON and TOF values for this process are 400 and 2087 h⁻¹, the highest reported to the best of our knowledge (Table 1).

^a Instituto de Investigaciones Químicas (IIQ), Departamento de Química Inorgánica, Centro de Innovación en Química Avanzada (ORFEO-CINCA), CSIC and Universidad de Sevilla, Avda. Américo Vespucio 49, 41092 Sevilla, Spain. E-mail: sconejero@iiq.csic.es; Tel: +34 954489563

^b Laboratorio de Compuestos Organometálicos y Catálisis

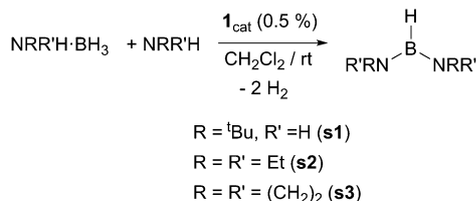
(Unidad asociada al CSIC), Departamento de Química Orgánica e Inorgánica, Universidad de Oviedo, C/Julián Clavería 8, 33006, Oviedo, Spain

† Electronic supplementary information (ESI) available: Experimental section, H₂ evolution graphics and X-ray crystallographic data. CCDC 1468913. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6cc02720b



Scheme 1 Dehydrocoupling of NMe₂H·BH₃ catalysed by complex **1**.





Scheme 2 Catalytic synthesis of symmetric diaminoboranes.

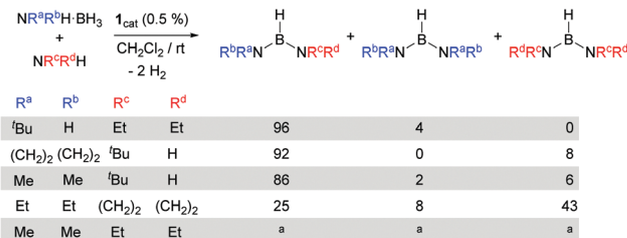
Table 1 Catalytic dehydrocoupling of amine-boranes and amines

Entry ^a	Amineborane	Amine	% Yield ^b (isolated)	TON	TOF (h ⁻¹)
1	^t BuNH ₂ ·BH ₃	^t BuNH ₂	> 99 (99)	400	2087
2	NET ₂ H·BH ₃	NET ₂ H	> 99 (88)	400	2400
3	(CH ₂) ₄ NH·BH ₃	(CH ₂) ₄ NH	95 ^c (81)	380	193
4	^t BuNH ₂ ·BH ₃	NET ₂ H	> 99 (99)	400	3692
5	^t BuNH ₂ ·BH ₃	(CH ₂) ₄ NH	94 (68)	376	376
6	NET ₂ H·BH ₃	(CH ₂) ₄ NH	93	372	248
7	NMe ₂ H·BH ₃	^t BuNH ₂	78 ^c (71) ^d	312	2023
8	NMe ₂ H·BH ₃	NET ₂ H	76 ^c	316	2257

^a Reaction conditions: CH₂Cl₂, rt. ^b Yields determined by ¹¹B NMR spectroscopy with respect to all diaminoboranes formed. ^c Cyclic dimers [NR₂BH₂]₂ and other unidentified species constitute the remaining reaction products (see the ESI). ^d Isolated yield corresponds to all the possible products formed.

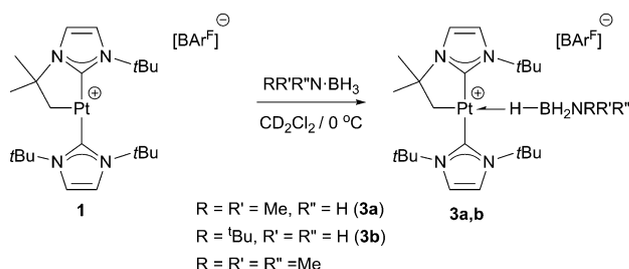
At the very end of the reaction, the catalyst remained the cyclo-metallated species **1**, but is slowly hydrogenated into complex [PtH(^tBu)₂][BAR^f], **2**. Other amine-boranes were tested under identical reaction conditions to form symmetrical diaminoboranes. Diethylamine-borane was also shown to be efficiently dehydrogenated in *ca.* 10 min in the presence of 1 equiv. of NET₂H and catalyst **1** (TON = 400, TOF = 2400 h⁻¹). On the other hand, pyrrolidine-borane (CH₂)₄NH·BH₃ required longer reaction times (2 h) to be converted into [(CH₂)₄N]₂BH (TON = 380, TOF = 193 h⁻¹). In the latter case, 5% of cyclic dimer [(CH₂)₄NBH₂]₂ is observed in the ¹¹B NMR spectra. Bulkier amine-boranes, such as ⁱPr₂NH·BH₃, do not undergo dehydrocoupling under similar reaction conditions, while heating at 60 °C for 12 h results in the formation of amino-borane NⁱPr₂BH₂, with no evidence of the corresponding diaminoborane.

Asymmetrical diaminoboranes are challenging substrates to be produced catalytically in a selective manner. Hill *et al.* have recently reported a method for the preparation of this type of substance using group 2 metal catalysts.^{5b} Although in most of the cases the reaction proceeds with excellent selectivities, long reaction times (24–92 h) and mild heating are usually necessary. Complex **1** has shown very good catalytic behaviour in the formation of unsymmetrical diaminoboranes (Scheme 3). The reaction of ^tBuNH₂·BH₃ with NET₂H takes place very fast (6.5 min, TON = 400, TOF = 3692 h⁻¹). According to NMR spectroscopy, borane (^tBuNH)(NET₂)BH is formed almost exclusively (*ca.* 4% of (^tBuNH)₂BH is observed by ¹H NMR). If the reaction is carried out in the inverse way, starting from NET₂H·BH₃ and ^tBuNH₂, the process is slower (*ca.* 20 min), but selectivities are comparable. The same behaviour is observed in the catalytic dehydrocoupling of (CH₂)₄NH·BH₃ and ^tBuNH₂. In this case, the reaction

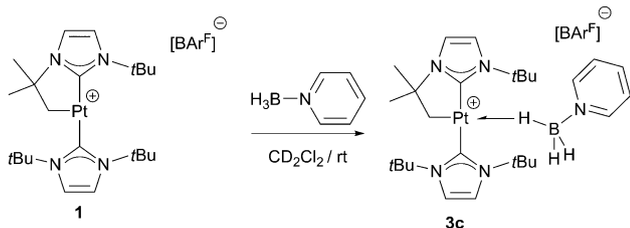
Scheme 3 Catalytic synthesis of asymmetric diaminoboranes. ^a Signal overlapping in the ¹H and ¹¹B(¹H) NMR spectra precluded establishing the exact amount of each diaminoborane (see the ESI[†]).

exhibits good selectivities towards the asymmetric diaminoborane (^tBuNH)[(CH₂)₄N]BH (92%), whereas the only symmetric borane detected is [(CH₂)₄N]₂BH (8%), with complete conversion after 2 h at rt. When the reaction is carried out the other way around, that is, starting from ^tBuNH₂·BH₃ and (CH₂)₄NH, the selectivities are only slightly different (85% of (^tBuNH)[(CH₂)₄N]BH, 15% of [(CH₂)₄N]₂BH), although the reaction proceeds in 1 h. The combination of pyrrolidine and diethylamine substrates indicates that the reaction is not selective yielding all possible reaction products (NET₂)₂BH, [(CH₂)₄N]₂BH and (NET₂)[(CH₂)₄N]BH in a 1 : 5.4 : 3.1 ratio, respectively, independent of the two possible combinations utilised. A similar behaviour was observed in the reaction of NMe₂H·BH₃ with NET₂H, although we could not establish the exact ratio due to signal overlapping in the ¹H NMR (see the ESI[†]). Nevertheless, good selectivities are produced in the reaction of NMe₂H·BH₃ and ^tBuNH₂ (Scheme 3). Very interestingly, the reaction of ^tBuNH₂·BH₃ with ⁱPr₂NH generates diaminoborane (^tBuNH)₂BH (**s1**) and amino-borane ⁱPr₂N-BH₂ in *ca.* 5 min at rt. Since **1** does not catalyse the dehydrocoupling of ⁱPr₂NH·BH₃ at rt, ⁱPr₂N-BH₂ is probably formed through the reorganization of amino-borane ^tBuNH-BH₂ and ⁱPr₂NH.⁹

With regard to the mechanism by which these transformations occur, it has been previously shown that dehydrogenation of amine-boranes NR₂H·BH₃ leading to the corresponding aminoboranes NR₂BH₂ is a key step.⁵ Previously, we have reported that complex **1** is able to dehydrogenate NMe₂H·BH₃ into NMe₂BH₂ (that then dimerises) through a distinct mechanism that involves a first step in which the amine-borane interacts with the platinum atom through the BH₃ protons (complex **3a** in Scheme 4) followed by nucleophilic addition of free amine present in solution to the activated boron atom.⁷ According to DFT calculations the most stable coordination mode of the BH₃ moiety is η¹-BH (Shimoi type complex) but, unfortunately, no experimental evidence for this

Scheme 4 Formation of complexes **3a,b**.

type of coordination could be provided. Here we further investigate these adducts by carrying out stoichiometric reactions between complex **1** and amine-boranes at 0 °C in CD₂Cl₂. ^tBuNH₂·BH₃ reacts with **1** leading to a new species that has been postulated as the η¹-BH derivative **3b** (Scheme 4). The BH₃ group resonates as a broad signal centred at 0.12 ppm that sharpens upon ¹¹B broadband decoupling. This signal is shifted upfield from the free ^tBuNH₂·BH₃ (δ 1.36). In addition, *J*_{PtH} of the CH₂-Pt fragment has a value of 90 Hz, that is, 30 Hz smaller than in complex **1** (120 Hz), consistent with the coordination of a ligand *trans* to the CH₂ moiety.¹⁰ This value is smaller than that observed in dimethylamine-borane **3a** (103 Hz) suggesting that the interaction with the platinum atom is stronger. The ¹¹B NMR spectrum shows a signal at -18.2 ppm that is shifted with respect to free amine-borane (-21.1 ppm). All these data agree well with the formulation of **3b** as depicted in Scheme 4.¹¹ Unfortunately, the instability of **3a** and **3b** due to their propensity to undergo dehydrocoupling precluded their isolation as pure compounds and further characterization. To avoid this problem, tertiary amine-borane NMe₃·BH₃ was used. However, no interaction with **1** was observed, probably due to steric constraints, thus allowing the establishment of a correlation between the bulkiness of the amine-borane and its interaction with **1** (stronger interaction: ^tBuNH₂·BH₃ > NMe₂H·BH₃ > NMe₃·BH₃). Consequently, the less hindered Lewis base stabilised borane C₆H₅N·BH₃ was a judicious choice. This borane reacts with **1** to yield **3c** (Scheme 5) quantitatively by NMR spectroscopy. The main feature of **3c** is the broad NMR signal centred at 1.25 ppm attributed to the BH₃ protons (2.55 ppm in free C₆H₅N·BH₃). The ¹¹B{¹H} spectrum exhibits a signal at -8.9 ppm, nearly 3 ppm down-field shifted with respect to C₆H₅N·BH₃. The apparent coupling constant ¹*J*_{B,H} is *ca.* 85 Hz, that is, 12 Hz smaller than in free C₆H₅N·BH₃, and compares well with previously described Shimoi-type complexes.^{11,12} The coupling constant of the CH₂-Pt protons with ¹⁹⁵Pt of 93 Hz is similar to that of complex **3b**. A definite proof of the nature of this compound came from the solid-state structure obtained by X-ray diffraction studies. Colourless crystals suitable for this analysis were obtained by slow diffusion of a concentrated solution of **3c** in CH₂Cl₂ into pentane at 0 °C. Fig. 1 depicts an ORTEP-type view of the cation of complex **3c**. The complex is four-coordinated, with two N-heterocyclic carbene units in the expected *trans* arrangement (C(1)-Pt(1)-C(12): 168.38(19)°) one of which is cyclometalated. The fourth coordination site is occupied by the C₆H₅N·BH₃ ligand in which one of the hydride atoms of the BH₃ unit bridges the platinum nucleus. The Pt(1)-H(1B) and H(1B)-B(1) bond distances are 1.96(5) and 1.03(2) Å, respectively.



Scheme 5 Synthesis of complex **3c**.

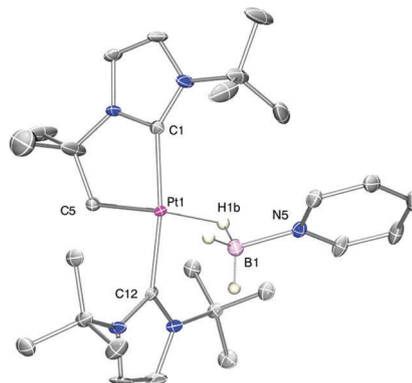


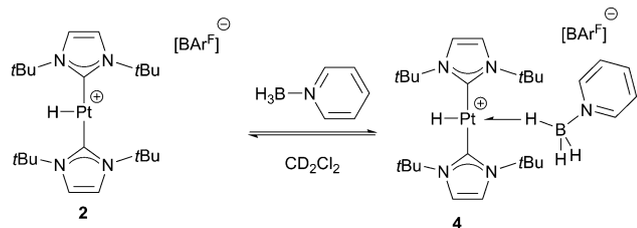
Fig. 1 ORTEP-type view of the cation of complex **3c**. Hydrogen atoms, except those of the BH₃ unit, have been omitted for clarity (ellipsoids are drawn at 30% probability). Selected bonds (Å) and angles (°): Pt1-H1b 1.96(5), Pt1-C5 2.045(5), B1-H1b 1.03(2); Pt1-H1b-B1 147.48(485), C1-Pt1-C12 168.38(19).

The Pt(1)-H(1B)-B(1) angle of 147.48(485)° together with the long Pt(1)··B(1) bond distance (2.8436(5) Å) indicates a negligible interaction between platinum and boron, and therefore this complex is best described as a η¹-BH or Shimoi-type complex.^{12,13}

Variable temperature NMR spectroscopy was used to study the dynamic behaviour due to exchange between bridging and terminal BH protons. The resonance for the BH₃ protons collapses into the baseline at *ca.* 223 K. Upon cooling to 203 K, a new signal of relative integral 2 appears at 2.47 ppm, in the expected region for terminal BH protons. Nevertheless, at this temperature, the signal corresponding to the Pt-H-B proton is not discernible.¹⁴ Further cooling to 183 K provided evidence for a very broad resonance at *ca.* -4 ppm (approx. integral value of 1), with the terminal BH protons resonating at 2.44 ppm. With respect to the ¹¹B NMR spectra, a very broad signal centred at δ -12.2 is detected at this temperature.

For comparison purposes, we have analysed the interaction of C₆H₅N·BH₃ with the hydrogenated form of complex **1**, the hydride derivative [PtH(^tBu)₂][BARF], **2**.¹⁵ No evidence of interaction between **2** and C₆H₅N·BH₃ is observed at 298 K. At this temperature, the BH₃ protons resonate at δ 2.49, whereas the platinum-hydride signal appears at δ -25.47 exhibiting a *J*_{Pt,H} of 2550 Hz, only marginally different from that of complex **2** in the absence of C₆H₅N·BH₃ (*J*_{Pt,H} = 2564 Hz).¹⁵ However, the ¹H NMR spectra at temperatures below 233 K indicate the presence of a new species, **4** (Scheme 6), coexisting with complex **2**, in a 0.3 to 1 ratio (**4**:**2**). This compound is characterised by a signal in the hydride region at -21.85 ppm showing a reduced coupling constant with ¹⁹⁵Pt (*J*_{Pt,H} = 1920 Hz) and another broad signal at 0.28 ppm (3H relative integral) that sharpens upon ¹¹B decoupling. As the temperature decreases, the concentration of **4** increases at the expense of **2**, reaching a maximum at 188 K (**4**:**2** ratio, *ca.* 4:1). The ¹¹B{¹H} NMR spectra at all temperatures exhibit a single very broad signal at -12.3 ppm that does not show splitting upon coupling to ¹H below 208 K. Interestingly, even at 188 K the BH₃ group still shows fast exchange between terminal and bridging hydrogens. The different behaviour of compounds **1** and **2** toward the interaction





Scheme 6 Equilibrium between complexes **2** and **4** in the presence of $C_6H_5N-BH_3$.

with $C_6H_5N-BH_3$ can be easily rationalised in terms of the different steric protection that the cyclometalated tBu ligand exerts on the platinum atom compared to its non-cyclometalated form. When the tBu ligand is cyclometalated the tBu group is tilted¹⁶ away from the metal centre favouring the interaction with amine-boranes.

Once the amine-borane binds the metal in a Shimoï-type fashion, dehydrogenation leading to amino-boranes, NR_2-BH_2 , takes place through a mechanism that involves the intermediacy of boronium cations $(NHR_2)_2BH_2^+$ and the neutral platinum hydride $[PtH(tBu)(tBu)]$, **5**.⁷ It must be noted that we have not observed an interaction between complex **1** and amino-borane NMe_2BH_2 (either using cyclic dimer $[NMe_2BH_2]_2$ as a precursor¹⁷ or during monitoring the dehydrogenation reaction of **1** and NMe_2H-BH_3 at 0 °C).⁷ At this point, we do not have further information on the mechanism by which the amino-borane is converted into diamino-boranes and the role of complex **1** in the process.

In summary, the coordinatively unsaturated Pt(II) complex **1** is a very efficient catalyst for the synthesis of diamino-boranes achieving TON and TOF values of 400 and $3692 h^{-1}$, respectively, the highest reported to date. The process takes place through the initial coordination of the BH protons of amine-boranes to the platinum centre in an end-on mode (Shimoï type) that was demonstrated crystallographically in the pyridine- BH_3 adduct **3c**. Ongoing efforts are geared towards unveiling the mechanism by which amino-boranes are transformed into diamino-boranes.

Financial support (FEDER contribution) from the MINECO (Projects CTQ2013-45011-P and CTQ2014-51912-REDC) and the Junta de Andalucía (Project FQM-2126) is gratefully acknowledged.

Notes and references

- (a) H. C. Johnson, T. N. Hooper and A. S. Weller, *Top. Organomet. Chem.*, 2015, **49**, 153; (b) E. M. Leitao, T. Jurka and I. Manners, *Nat. Chem.*, 2013, **5**, 817; (c) N. E. Stubbs, A. P. M. Robertson, E. M. Leitao and I. Manners, *J. Organomet. Chem.*, 2013, **730**, 84; (d) A. S. John, K. I. Goldberg and D. M. Heinekey, *Top. Organomet. Chem.*, 2013, **40**, 271.
- (a) H. C. Johnson and A. S. Weller, *Angew. Chem., Int. Ed.*, 2015, **54**, 10173; (b) M. A. Esteruelas, A. M. López, M. Mora and E. Oñate, *ACS Catal.*, 2015, **5**, 187; (c) A. G. Algarra, L. J. Sewell, H. C. Johnson, S. A. Macgregor and A. S. Weller, *Dalton Trans.*, 2014, **43**, 11118; (d) S. Muhammad, S. Moncho, E. N. Brothers and A. A. Bengali, *Chem. Commun.*, 2014, **50**, 5874; (e) L. J. Sewell, G. C. Lloyd-Jones and A. S. Weller, *J. Am. Chem. Soc.*, 2012, **134**, 3598; (f) C. Y. Tang, N. Phillips, M. J. Kelly and S. Aldridge, *Chem. Commun.*, 2012, **48**, 11999; (g) R. Kumar and B. R. Jagirdar, *Inorg. Chem.*, 2013, **22**, 28; (h) H. C. Johnson, A. P. M. Robertson, A. B. Chaplin, L. J. Sewell, A. L. Thompson, M. F. Haddow, I. Manners and A. S. Weller, *J. Am. Chem. Soc.*, 2011, **133**, 11076; (i) R. Dallanegra, A. P. M. Robertson, A. B. Chaplin, I. Manners and A. S. Weller, *Chem. Commun.*, 2011, **47**, 3763; (j) G. Alcaraz and S. Sabo-Etienne, *Angew. Chem., Int. Ed.*, 2010, **49**, 7170; (k) C. Y. Tang, A. L. Thompson and S. Aldridge, *J. Am. Chem. Soc.*, 2010, **132**, 10578; (l) T. M. Douglas, A. B. Chaplin, A. S. Weller, X. Yang and M. B. Hall, *J. Am. Chem. Soc.*, 2009, **131**, 15440.
- (a) A. Staubitz, A. P. M. Robertson and I. Manners, *Chem. Rev.*, 2010, **110**, 4079; (b) N. C. Smythe and J. C. Gordon, *Eur. J. Inorg. Chem.*, 2010, 509; (c) C. W. Hamilton, R. T. Baker, A. Staubitz and I. Manners, *Chem. Soc. Rev.*, 2009, **38**, 279.
- H. C. Johnson, C. L. McMullin, S. D. Pike, S. A. Macgregor and A. S. Weller, *Angew. Chem., Int. Ed.*, 2013, **52**, 9776.
- (a) C. J. Wallis, G. Alcaraz, A. S. Petit, A. I. Poblador-Bahamonde, E. Clot, C. Bijani, L. Vendier and S. Sabo-Etienne, *Chem. – Eur. J.*, 2015, **21**, 13080; (b) P. Bellham, M. S. Hill, G. Kociok-Köhn and D. J. Liprott, *Chem. Commun.*, 2013, **49**, 1960; (c) C. J. Wallis, H. Dyer, L. Vendier, G. Alcaraz and S. Sabo-Etienne, *Angew. Chem., Int. Ed.*, 2012, **51**, 3646; (d) It has been reported that aromatic amine-boranes can produce diamino-boranes without a catalyst: H. Helten, A. P. M. Robertson, A. Staubitz, J. R. Vance, M. F. Haddow and I. Manners, *Chem. – Eur. J.*, 2012, **18**, 4665.
- O. Rivada-Wheelaghan, B. Donnadieu, C. Maya and S. Conejero, *Chem. – Eur. J.*, 2010, **16**, 10323.
- M. Roselló-Merino, J. López-Serrano and S. Conejero, *J. Am. Chem. Soc.*, 2013, **135**, 10910.
- (a) P. Bellham, M. S. Hill and G. Kociok-Köhn, *Organometallics*, 2014, **33**, 5716; (b) Y. Kawano, M. Uruichi, M. Shimoï, S. Taki, T. Kawaguchi, T. Kakizawa and H. Ogino, *J. Am. Chem. Soc.*, 2009, **131**, 14946.
- A. P. M. Robertson, E. M. Leitao and I. Manners, *J. Am. Chem. Soc.*, 2011, **133**, 19322.
- M. A. Ortuño, S. Conejero and A. Lledós, *Beilstein J. Org. Chem.*, 2013, **9**, 1352.
- G. Alcaraz and S. Sabo-Etienne, *Coord. Chem. Rev.*, 2008, **252**, 2395.
- (a) R. Dallanegra, A. B. Chaplin, J. Tsim and A. S. Weller, *Chem. Commun.*, 2010, **46**, 3092; (b) M. Shimoï, S. Nagai, M. Ichikawa, Y. Kawano, K. Katoh, M. Uruichi and H. Ogino, *J. Am. Chem. Soc.*, 1999, **121**, 11704.
- (a) A. E. W. Ledger, C. E. Ellul, M. F. Mahon, J. M. J. Williams and M. K. Whittlesey, *Chem. – Eur. J.*, 2011, **17**, 8704; (b) R. Dallanegra, A. B. Chaplin and A. S. Weller, *Angew. Chem., Int. Ed.*, 2009, **48**, 6875; (c) Y. Kawano, K. Yamaguchi, S. Miyake, T. Kakizawa and M. Shimoï, *Chem. – Eur. J.*, 2007, **13**, 6920; (d) Y. Kawano, M. Hashiva and M. Shimoï, *Organometallics*, 2006, **25**, 4420; (e) N. Merle, G. Kociok-Köhn, M. F. Mahon, C. G. Frost, G. D. Ruggerio, A. S. Weller and M. C. Willis, *Dalton Trans.*, 2004, 3883; (f) T. Yasue, Y. Kawano and M. Shimoï, *Angew. Chem., Int. Ed.*, 2003, **42**, 1727.
- A similar behavior has been observed in a ruthenium complex. See ref. 13a.
- (a) M. A. Ortuño, P. Vidossich, S. Conejero and A. Lledós, *Angew. Chem., Int. Ed.*, 2014, **53**, 14158; (b) O. Rivada-Wheelaghan, M. Roselló-Merino, M. A. Ortuño, P. Vidossich, E. Gutiérrez-Puebla, A. Lledós and S. Conejero, *Inorg. Chem.*, 2014, **53**, 4257.
- J.-N. Luy, S. A. Hauser, A. B. Chaplin and R. Tonner, *Organometallics*, 2015, **34**, 5099.
- G. Bénac-Lestrille, U. Helmstedt, L. Vendier, G. Alcaraz, E. Clot and S. Sabo-Etienne, *Inorg. Chem.*, 2011, **50**, 11039.

