Chemical Science

EDGE ARTICLE

Cite this: Chem. Sci., 2014, 5, 2546

Multiple metal-bound oligomers from Ir-catalysed dehydropolymerisation of H₃B·NH₃ as probed by experiment and computation⁺

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Received 11th March 2014 Accepted 7th April 2014 DOI: 10.1039/c4sc00735b

www.rsc.org/chemicalscience

Multiple metal-bound oligomers in the dehydropolymerisation of $H_3B \cdot NH_3$ have been observed by electrospray-ionisation mass spectrometry and NMR spectroscopy using the catalytic metal fragment $\{Ir(PCy_3)_2(H)_2\}^+$. A computational study suggests that sterics dictate whether multiple dehydrogenation/ B-N coupling of amine-boranes $H_3B \cdot NRR'H$ (R, R' = Me or H) is observed, and also demonstrates the experimentally observed requirement for additional amine-borane to promote dehydrocoupling.

Introduction

The dehydropolymerisation of amine-boranes $H_3B \cdot NRH_2$ (R = H, Me) is a promising methodology for the synthesis of new B–N materials, for example polymeric materials that are isoelectronic with societally ubiquitous polyolefins,^{1,2} or precursors to B–N ceramics such as white graphene.³ Catalysis of these processes by a transition metal fragment offers potential for control of kinetics and final product distributions, and various systems have been shown to promote dehydropolymerisation.^{4–12} Non-metal catalysed processes have also been discussed.^{13,14}

The mechanism of catalytic dehydropolymerisation of $H_3B \cdot NH_3$ or $H_3B \cdot NMeH_2$ has been suggested to be based upon dehydrogenation followed by a second metal-mediated coordination polymerisation step.^{6,9,10,12,15,16} In particular, there is growing evidence to suggest that transient amino-borane (*e.g.* H_2B ==NH₂ or H_2B ==NMeH), that arises from dehydrogenation of the precursor amine-borane, remains associated with the metal.¹⁷ If liberated these unsaturated fragments form the corresponding borazine by oligomerisation (Scheme 1), or can be trapped by hydroboration of exogenous cyclohexene – assuming such reactions are faster than polymerisation (*i.e.* B–N bond formation leading to a growing polymer chain). In addition bulky primary amine-boranes, $H_3B \cdot N^tBuH_2$,¹⁸ or secondary amine-boranes, *e.g.* $H_3B \cdot NMe_2H$,^{19,20} give simple amino-borane

products rather than extensive oligomerisation. Adding to the complexity, different metal/ligand combinations likely lead to subtly different mechanisms.^{8,10,12}

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Direct mechanistic insight into the dehydropolymerisation process through the observation of intermediates has been sparse. Recently we reported the isolation of the product of the first oligomerisation event in such a process by reaction of $[Ir(PCy_3)_2(H)_2(H_2)_2][BAr^F_4]$, 1, with 2 equivalents of $H_3B \cdot NMeH_2$ to form $[Ir(PCy_3)_2(H)_2(\eta^2 - H_3B \cdot NMeHBH_2 \cdot NMeH_2)][BAr^F_4]$, 5b, (Scheme 2).21 This reaction is slow and does not produce higher oligomers, and a tentative mechanism was suggested to account for this selectivity. With bulkier H₃B·NMe₂H only dehydrogenation to form the bound amino-borane (*i.e.* 4a*) is observed.¹⁹ We now report that with $H_3B \cdot NH_3$ dehydropolymerisation can also be promoted by 1 and that, in contrast to $H_3B \cdot NMeH_2$, higher oligomeric products bound to the metal centre (6a-e, Scheme 2) can be observed by electrospray ionisation mass spectrometry (ESI-MS) and NMR spectroscopy. ESI-MS provides the ideal analytical platform to study these processes as it allows for the convenient analysis of mixtures of products under inert conditions.^{22,23} Computational studies^{10,19,24-27} offer a mechanistic rationale for oligomerisation that explains both the difference in the degree of oligomerisation with increasing steric bulk between the amine-boranes H₃B·NH₃, H₃B·NMeH₂



Scheme 1 Dehydrogenation and dehydropolymerisation of amineboranes; full pathways not shown.

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[†] Electronic supplementary information (ESI) available: Full experimental details, ESI-MS, NMR spectra, details of X-ray crystallographic analysis and full computational details. CCDC 971346, 971347 and 988696. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c4sc00735b

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Scheme 2 Selected examples of compounds discussed in this study. $[Ir(PCy_3)_2(H)_2(\eta^2-H_3B\cdot(NRHBH_2)_n\cdot NRH_2)][BAr^F_4]$, R = Me, 5; H, 6; n = 0, a; n = 1, b; n = 2, c; n = 3, d; n = 4, e. $[BAr^F_4]^-$ anions are not shown. * = corresponding amino-borane.

and $H_3B \cdot NMe_2H$ and the previously noted requirement for additional amine-borane to promote this process.²¹

Results and discussion

Addition of one equivalent of H₃B·NH₃ to 1¹⁹ in C₆H₅F solvent results in the immediate formation of the sigma amine-borane complex $[Ir(PCy_3)_2(H)_2(\eta^2 - H_3B \cdot NH_3)][BAr_4^F]$ 6a in quantitative yield by NMR spectroscopy. There is no onward dehydrogenation after 4 hours under these conditions, but addition of further $H_3B \cdot NH_3$ (10 equivalents total) results in the formation of higher oligomers, $[Ir(PCy_3)_2(H)_2\{\eta^2 - H_3B \cdot (NH_2BH_2)_n \cdot NH_3\}][BAr_4] n = 1 -$ 4. This requirement for additional amine-borane to promote dehydrogenation has been noted before in these systems, although its role has only been speculated upon.²¹ Fig. 1 shows the ESI-MS spectra of the reaction of 1 with the amine-boranes $H_3B \cdot NMe_x H_{3-x}$ (x = 0-3) demonstrating the increasing degrees of dehydrogenation and oligomerisation with decreasing steric bulk of the amine-borane. Under these conditions H₃B·NMe₂H undergoes dehydrogenation with no subsequent B-N coupling $(4a^*)$,¹⁹ while H₃B·NMeH₂ gives the product of one dehydrocoupling event (5b).²¹ By contrast for H₃B·NH₃ metal-bound oligomers arising from up to four of these dehydrocoupling events are observed by ESI-MS (6b-e), which all show excellent fits with calculated isotopomer patterns, with 6d/e (n = 4, 5; Scheme 2) observed as $[M-H_2]^+$ cations. In the ${}^{1}H{}^{11}B$ NMR spectrum of this mixture three distinct pairs of Ir…H-B and Ir-H environments are observed in an approximate 1:10:10 ratio (see ESI[†]), which are assigned to 6a, 6b and 6c respectively (vide infra), consistent with the major species observed by ESI-MS (6a–c). The ${}^{11}B{}^{1}H{}$ NMR spectrum of this mixture shows broad, potentially overlapping, signals in the Ir...H₃B and $\{BH_2\}$ regions, and the ³¹P $\{^1H\}$ NMR spectrum shows two tightly-coupled AB doublets in approximately equal ratio, the third species (i.e. 6a) being too low in intensity to be observed. The identity of these complexes has been confirmed by the independent synthesis of 6b and 6c from the preformed



Fig. 1 ESI-MS (positive mode) of 1 (C₆H₅F solution) and 10 equivalents of: (A) H₃B·NMe₃, **3a**; (B) H₃B·NMe₂H, **4a*** (96 h, 3 equiv.); (C) H₃B·NMeH₂, **5a/5b** (D) H₃B·NH₃, **6a–e**; Calculated isotopomer *m/z* given in italics; n = 4 obs. m/z = 871.63 [M – H₂]⁺, calc. 871.62. After 4 hours unless otherwise stated. See Scheme 2 for numbering, and ESI⁺ for an expansion of (D).

borazanes $H_3B \cdot NH_2BH_2 \cdot NH_3^{28}$ and $H_3B \cdot (NH_2BH_2)_2 \cdot NH_3^{29}$ respectively. Scheme 3 shows the solid-state structure (as the $[BAr^{Cl}_4]^-$ salts³⁰ from $[Ir(PCy_3)_2(H)_2(H_2)_2][BAr^{Cl}_4]$, 2) of 6c, alongside that of 6a, which confirm formulation, being closely related to analogous complexes 3a, 4a, 5a and 5b.^{19,21} Over time (24 h) these mixtures of products degrade to give bimetallic products identified by ESI-MS as $[{Ir(PCy_3)_2(H)_2}_2{H_3B(NH_2BH_2)_nH}]^+$ 7a–d (n =0 to 3 respectively), presumably in which the anionic aminoboranes $[H_3B(NH_2BH_2)_nH]^{-29}$ bridge between two cationic metal fragments. Recrystallisation of this mixture afforded small amounts of the borohydride complex³¹ $[{Ir(PCy_3)_2(H)_2}_2(\eta^2,\eta^2 H_2BH_2)][BArF_4]$ 7a (see ESI† for a solid-state structure). We were unable to definitely characterise the other byproducts of this decomposition.

Borazine was also observed during the oligomerisation of $H_3B \cdot NH_3$ (~10% by ¹¹B NMR spectroscopy relative to $[BAr^F_4]^-$), which might suggest free amino-borane is formed as a transient intermediate during the reaction.^{15,18} Addition of excess cyclohexene to the reaction did not result in the observation of any hydroboration product, $Cy_2B=NH_2$, a trapping reaction that



Scheme 3 Synthesis of 6a, 6b and 6c. Solid-state structures (50% displacement ellipsoids) of 6a and 6c. Selected hydrogen atoms are shown and the $[BAr^{Cl}_4]^-$ anions are omitted for clarity. See ESI† for full details.

has previously been suggested to be indicative of free aminoborane in dehydrocoupling reactions.¹⁵ As recently noted, however, this process relies on hydroboration being kinetically competitive with oligomerisation, which might not necessarily be the case.¹⁰

Although these data are consistent with a growing oligomeric chain at the ${Ir(PCy_3)_2(H)_2}^+$ fragment, similar to those observed by ESI-MS for olefin polymerisation,^{32,33} these observations cannot discount a scenario where metal-catalysed dehydrogenation forms the free amino-borane, H₂B=NH₂, which then polymerises off-metal,13 with the most soluble shortchain oligomers then coordinating to the metal fragment. However, as computation suggests (vide infra) that the first dehydrogenation has a significantly higher barrier than subsequent oligomerisation we propose that this scenario is less likely. To probe further the oligomerisation process, three sequential additions of 1.1 equivalents of $H_3B \cdot NH_3$ to 1 gave progressively longer oligomer chains (i.e. 6a-6c) as measured by ESI-MS (see ESI[†]), although this mixture was biased towards 6a and 6b, suggesting that the sigma-bound oligomeric units, e.g. 6b or 6c, are only weakly bound with the metal centre and can be displaced by excess H₃B·NH₃. Confirming this, addition of two equivalents of $H_3B \cdot NH_3$ to 6c immediately results in a mixture of 6a-c and free $H_3B \cdot (NH_2BH_2)_2 \cdot NH_3$, with 6c the major observed product. After 4 hours this has developed into a mixture of 6a-e with 6b and 6c the major products. Addition of 2 equivalents of $H_3B \cdot NH_2BH_2 \cdot NH_3$ to **6a** results in the formation of 6b and relatively smaller amounts of 6c-6e (by ESI-MS), the latter presumably derived from further dehydrocoupling events from **6b** with $H_3B \cdot NH_3$ (Scheme 4). Overall this suggests a mechanism in which the formed sigma-bound oligomer can be displaced by other amine-boranes, *i.e.* reversible chain transfer can occur. At the end of the reaction (24 h) a white solid is recovered that shows an IR spectrum essentially identical to polyaminoborane.34 Use of 5 equivalents each of H3B·NH3 and H₃B·NMeH₂ gave a mixture of metal-bound co-oligomers $[Ir(PCy_3)_2(H)_2{H(H_2BNH_2)_x(H_2BNMeH)_yH}]^+$ (x = 0, 1, y = 1, 2; x x = 1, y = 0; x = 2, y = 1).



Scheme 4 Addition of 2 equivalents of $H_3B \cdot (NH_2BH_2) \cdot NH_3$ to **6a** results in the formation of higher oligomers.

Density functional theory (DFT) calculations³⁵ have been used to study the mechanism of the dehydrocoupling of $H_3B \cdot NH_3$ at **6a** with particular focus on (i) the requirement for additional $H_3B \cdot NH_3$ to induce dehydrogenation, (ii) the mechanism of the B–N coupling step and (iii) the varying affinities of the different amine-boranes toward oligomerisation. These calculations employed PMe₃ ligands, with $[Ir(PMe_3)_2(H)_2(\eta^2-H_3B \cdot NMe_xH_{3-x})]^*$ (denoted **6a**', x = 0, **5a**', x = 1 and **4a**', x = 2) the model initial reactants, and use a BP86-D3(C₆H₅F) protocol. We report free energies derived from gas-phase BP86-optimisations, corrected for dispersion and solvation effects. Each key step in the dehydrocoupling process (B–H/N–H bond activation and B–N bond coupling) presented more than one possible transition state and the most accessible of these are presented here, with alternative structures given in the ESI.[†]

We have previously modelled the dehydrogenation of $H_3B \cdot NMe_2H$ in $[Ir(PMe_3)_2(H)_2(\eta^2-H_3B \cdot NMe_2H)]^+$ (4a') to form the corresponding amino-borane adduct (*i.e.* 4a'*, a model of 4a* in Scheme 2) and defined a mechanism based on sequential B–H activation, H_2 loss and rate-limiting N–H activation.¹⁹ Applying this mechanism to $H_3B \cdot NH_3$ dehydrogenation in 6a' reveals a barrier of 33.8 kcal mol⁻¹ in which the N–H activation step is again rate-limiting (see Fig. S1–3, ESI†). With an added $H_3B \cdot NH_3$ molecule a related mechanism can be characterised but with a significantly reduced barrier of 26.7 kcal mol⁻¹ (Fig. 2). In this process the second $H_3B \cdot NH_3$ molecule first adds to 6a' to give $[Ir(PMe_3)_2(H)_2(\eta^1-H_3B \cdot NH_3)_2]^+$, I6a'₁, with a binding energy of 5.0 kcal mol⁻¹. This stabilisation is in part due to a $BH(\delta^-) \cdots H(\delta^+)N$ dihydrogen interaction between the two $H_3B \cdot NH_3$ ligands.^{36,37} B–H activation in I6a'₁ entails a



Fig. 2 Computed free energy reaction profile (kcal mol⁻¹, BP86-D3(C₆H₅F)) for dehydrogenation of $H_3B \cdot NH_3$ in **6a**' in the presence of added $H_3B \cdot NH_3$.



Fig. 3 Computed structures of the rate-limiting N–H activation transition states of (a) a first and (b) a second $H_3B \cdot NH_3$ molecule at **6a**'. Key distances are in Å and PMe₃ H atoms are omitted for clarity.



TS6a'

Fig. 4 Computed free energy reaction profile (kcal mol $^{-1}$, BP86-D3(C₆H₅F)) for B–N coupling and formation of oligomerisation product **6b**'.

barrier of 16.6 kcal mol⁻¹ *via* **TS6a'**_{BH1} and proceeds with concomitant reductive coupling of the two hydride ligands to give $[Ir(PMe_3)_2(BH_2NH_3)(H)(H_2)(\eta^1-H_3B\cdot NH_3)]^+$, **I6a'**₂ (*G* = +15.9 kcal mol⁻¹). H₂ loss then leads to **I6a'**₃ (*G* = +10.7 kcal mol⁻¹) from which rate-limiting N-H activation occurs *via* **TS6a'**_{NH1} (*G* = +26.7 kcal mol⁻¹) to give **I6a'**₄ in which both an amine- and an amino-borane are bound to the metal centre.

The computed geometry of TS6a'_{NH1} is shown in Fig. 3a and shows transfer of H^{14} from the BH₂NH₃ ligand to Ir (N¹…H¹⁴ = 1.42 Å; $Ir \cdots H^{14} = 1.74$ Å) while a dihydrogen bonding interaction is maintained with the spectator $H_3B \cdot NH_3$ ligand (H^{24} ... $H^{13} = 1.80$ Å). This feature stabilises both TS6a'_{NH1} and its precursor $\mathbf{I6a'}_3$ and so contributes to a reduction in the overall barrier to dehydrogenation of 7.1 kcal mol⁻¹ compared to the reaction direct from 6a' without added amine-borane. An alternative transition state, TS6a'_{NH1}(Alt 1), in which the second $H_3B \cdot NH_3$ ligand adopts an η^2 -(B,H) bonding mode (similar to the amino-borane ligand in TS6a'_{NH2}, see Fig. 3b and below) is comparable in energy ($G = +26.9 \text{ kcal mol}^{-1}$, see Fig. S6(b)†). Both forms of TS6a'_{NH1} are consistent with dehydrogenation being facilitated by the addition of amine-borane to 6a'. Similar reductions in barriers to dehydrogenation have very recently been reported for H₃B·NMe₂H dehydrogenation using {Rh(chelating phosphine)}⁺ fragments.³⁸

For the subsequent B–N coupling step a total seven different pathways have been characterised. Four of these stem from intermediate $I6a'_4$ and entail B–H activation in the $H_3B \cdot NH_3$ ligand to produce a Lewis acidic { H_2BNH_3 } moiety that then couples with H_2B = NH_2 . In most cases these processes occur in one step. Two further pathways have been characterised for the direct reaction of free H_2B = NH_2 with either 6a' or its B–H activated form. All of these pathways, however, have computed barriers in excess of 28 kcal mol⁻¹, and as this is higher than the barrier to dehydrogenation these pathways would be inconsistent with the lack of any bound amino-borane intermediates being observed experimentally. Full details of these alternative pathways are given in the ESI (see Fig. S12†).

A significantly more accessible B–N coupling route was characterised that involved the direct reaction of two $H_2B=NH_2$ units. This process therefore requires the prior dehydrogenation

of a second H₃B·NH₃ molecule and a pathway for this, analogous to that shown in Fig. 2, has been defined starting from I6a'4 and forming $[Ir(PMe_3)_2(H)_2(\eta^2-H_2B=NH_2)]^+$ (I6a'₇) and free $H_2B=NH_2$ (see also Fig. S7-9[†]). I6a'₇ is closely related to that calculated for the product of dehydrogenation of H₃B·NMe₂H by the same fragment.¹⁹ The key N-H activation transition state in this process, TS6a'_{NH2} (Fig. 3b), has a free energy of +24.2 kcal mol⁻¹ and features a spectator $\eta^2(B,H)-H_2B=NH_2$ ligand³⁹ that stabilises the metal centre. Oligomerisation then proceeds through the reaction of I6a'7 with H2B=NH2 and the associated reaction profile (Fig. 4) shows B-N coupling via TS6a'_{BN} at only +17.9 kcal mol⁻¹. The structure of this transition state (Fig. 5) shows that the Ir-bound amino-borane has rearranged to an η^2 -(B,H) mode that exposes the pendant $\{NH_2\}$ moiety to attack by the second, incoming amino-borane ($N^1 \cdots B^2 = 2.37$ Å). As this occurs a hydride transfers from Ir onto N² (Ir-H²⁴ = 1.63 Å; $H^{24} \cdots N^2 = 1.64$ Å) to generate an η^2 -(B,H)– $H_2B \cdot NH_2BH_2 \cdot NH_3$ ligand in the resultant intermediate I6a'₈ ($G = \pm 1.6$ kcal mol⁻¹). Addition of H₂ (I6a'₉, G = +7.3 kcal mol⁻¹) and facile B-H reductive coupling gives the final model product, [Ir(P- $Me_{3}_{2}(H)_{2}(\eta^{2}-H_{3}B\cdot NH_{2}BH_{2}\cdot NH_{3})]^{+}$, **6b**' (G = -10.9 kcal mol⁻¹).⁴⁰ This coupling process is similar to that suggested by



Fig. 5 Computed B–N coupling transition state with key distances in Å and PMe_3 H atoms omitted for clarity.

Schneider and co-workers in bifunctional $Ru(H)_2(PMe_3)(PNP)$ catalysis [PNP = $HN(CH_2CH_2P^tBu_2)_2$], in which an N-H activated $H_3B \cdot NH_3$ group undergoes B-N coupling with H_2B ==NH₂ during dehydropolymerisation.¹⁰

Reaction profiles analogous to those in Fig. 2 and 4 were also computed for the dehydrocoupling of $H_3B \cdot NMeH_2$ at 5a' and H₃B·NMe₂H at 4a'. Similar dehydrogenation barriers are found in each case $(5a'/H_3B\cdot NMeH_2: 25.2 \text{ kcal mol}^{-1}; 4a'/H_3B\cdot NMe_2H:$ 26.2 kcal mol^{-1}) and in the absence of a second amine-borane molecule these barriers increase to above 33 kcal mol⁻¹, reiterating the promotional effect of added amine-borane on this process. In contrast the B-N coupling transition states are more substrate-dependent and increase significantly in energy with the size of the amine-borane $(6a'/H_3B \cdot NH_3: 17.9 \text{ kcal mol}^{-1}; 5a'/$ $H_3B \cdot NMeH_2$: 19.9 kcal mol⁻¹; $4a'/H_3B \cdot NMe_2H$: 26.5 kcal mol^{-1}). This trend is consistent with oligomerisation being accessible for both $H_3B \cdot NH_3$ and $H_3B \cdot NMeH_2$, but this step becoming significantly more difficult for the larger H₃B·NMe₂H. Indeed oligomerisation is not seen experimentally for 4a/H₃B·NMe₂H under the conditions used here.⁴¹

An analogous mechanism based on dehydrocoupling of $H_3B \cdot NH_3$ and $H_3B \cdot NH_2BH_2 \cdot NH_3$ can account for the formation of the $H_3B \cdot (NH_2BH_2)_2 \cdot NH_3$ trimer seen in **6c** (modelled by **6c**'). The key energetics are similar to those computed in the pathway for the formation of 6b': dehydrogenation of $H_3B \cdot NH_3$ (in the presence of H₃B·NH₂BH₂·NH₃) has an overall barrier of 26.3 kcal mol⁻¹, then dehydrogenation of H₃B·NH₂BH₂·NH₃ (now in the presence of $H_2N=BH_2$) has a barrier of 24.3 kcal mol⁻¹. The order of dehydrogenation is important, however, as the alternative initial dehydrogenation of $H_3B \cdot NH_2BH_2 \cdot NH_3$ (in the presence of $H_3B \cdot NH_3$ has a higher barrier of 28.1 kcal mol⁻¹ (see Fig. S15[†]). The subsequent B-N coupling transition state is again more accessible than dehydrogenation, TS6b'_{BNa} (Fig. 6a) having a computed energy of 21.2 kcal mol⁻¹. In this case there are two possible B-N coupling outcomes, depending on whether $H_2B=NH_2$ (as in TS6b'_{BNa}) or $H_2B=NHBH_2 \cdot NH_3$ (TS6b'_{BNb}, Fig. 6b) is bound to Ir in the transition state. The former case leads to a straight chain oligomer product, and is 2.3 kcal mol^{-1} more stable than the alternative that gives a branched chain product. The barrier for this second oligomerisation step is close to that for the B-N coupling of H₃B·NMeH₂ $(19.9 \text{ kcal mol}^{-1})$, highlighting the similar behaviour of these two mono-substituted amine-boranes. This in turn suggests

Fig. 6 Transition states for B–N bond coupling leading to (a) straight chain and (b) branched chain formation in **6***c*' as well as (c) straight chain formation in **5***c*'. Free energies (kcal mol⁻¹, BP86-D3(C₆H₅F)) are quoted relative to [Ir(PMe₃)₂(H)₂(η¹-H₃B·NMe_xH_{3-x})(η¹-H₃B·NMe_xH_{2-x}BH₂·NMe_xH_{3-x})]⁺ (**I6***b*'₁, *x* = 0; **I5***b*'₁, *x* = 1) as appropriate.

(c)

H₂B

Me

-NMeH₂

TS5b'_{BNa}

+24.8 kcal/mo

Me.

PMe₃

TS6b'_{BNb}

+23.5 kcal/mol

(b)

(a)

H₂N

H₂B

BH-

that subsequent chain growth with further $H_3B \cdot NH_3$ may proceed *via* transition states related to $TS6b'_{BNa}$ in which the growing oligomer chain extends away from the metal centre with minimal additional steric impact. By the same token, $H_3B \cdot NMeH_2$ trimerisation is more difficult with the equivalent transition state, $TS5b'_{BNa}$, equating to a higher barrier of +24.8 kcal mol⁻¹ (Fig. 6c). This trend towards higher oligomerisation barriers as the size of the amine-borane increases is consistent with the experimental observations (*i.e.* 5a giving 5b alone whereas 6a can undergo multiple oligomerisation steps to give 6b-e).

A potential side reaction within this mechanistic picture involves the cyclisation of two aminoborane fragments, either directly at the metal (*e.g. via* reaction of H_2B = NH_2 with [Ir(P-Me_3)_2(H)_2(\eta^2-H_2B=NH_2)]^+, **I6a**'₇) or *via* an off-metal process^{13,42} involving two free aminoboranes. In fact for H_3B · NH_3 both these processes are computed to be competitive with B–N coupling *via* **TS6a**'_{BN}, dimerisation at **I6a**'₇ having a transition state energy of +15.4 kcal mol⁻¹ while the off-metal process has a barrier of 16.2 kcal mol⁻¹ (see Fig. S16†). Some dimerisation (and trimerisation) may therefore be anticipated, and indeed evidence of this is seen in the small amount of borazine that is observed as minor products in the oligomerisation processes.

Overall the proposed dehydrogenation/oligomerisation mechanism captures the key trends observed experimentally by ESI-MS and NMR spectroscopy. In particular the promotional effect of added amine-borane on dehydrogenation for all three $H_3B \cdot NMe_x H_{3-x}$ (x = 0-2) species and the decreasing propensity toward oligomerisation as the size of the amine-borane increases are reproduced. However, some issues do remain: (i) the absolute barriers computed for the dehydrogenation are ca. 26 kcal mol^{-1} and so are rather high for a (albeit slow) room temperature process; (ii) once dehydrogenation has occurred, the competing H2B=NH2 dimerisation processes are computed to be slightly more favourable than oligomerisation. One reason for these discrepancies may be the use of a model system in the present study, where PMe₃ is used in place of PCy₃ ligands. However, an additional factor may be that both the key N-H activation (e.g. TS6a'_{NH1}) and B-N coupling (e.g. TS6a'_{BN})



Scheme 5 Key steps in the oligomerisation of $H_3B \cdot NH_3$ at $I6a'_1$ in the presence of a third $H_3B \cdot NH_3$. Free energies (kcal mol⁻¹, BP86-D3(C₆H₅F)) are in kcal mol⁻¹.

PMe-

l₂[−]NH₃ TS6b'_{BNa}

+21.2 kcal/mol



Fig. 7 Computed B–N coupling transition state in the presence of a third $H_3B \cdot NH_3$ molecule. Key distances in Å and PMe₃ H atoms omitted for clarity.

transition states exhibit a vacant site that offers the potential for further stabilisation. Indeed a third $H_3B\cdot NH_3$ molecule was found to promote both of these steps (see Scheme 5 and Fig. 7). Starting from **I6a'_1.AB** dehydrogenation proceeds with a reduced overall barrier of 22.4 kcal mol⁻¹ to give **I6a'_7.AB** at -4.9 kcal mol⁻¹ and from here B–N coupling has a barrier of only 9.4 kcal mol⁻¹. Moreover, B–N coupling (and the completion of the oligomerisation process) are now kinetically preferred over dimer formation. Therefore several substrate molecules may cooperate to promote the oligomerisation process. Alternatively a solvent molecule may interact with the unsaturated metal centre and so promote the oligomerisation step, although we have not attempted to explicitly model this here.

Conclusions

In summary, we report the observation and characterisation of multiple metal-bound oligomers in the dehydrocoupling of H₃B·NH₃. This contrasts with only a single oligomerisation event being observed for H₃B·NMeH₂ and none for H₃B·NMe₂H. Interrogation of the likely mechanism using computational methods reveals that initial dehydrogenation of H₃B·NH₃ is a higher energy process than both the subsequent dehydrogenation of a second amine-borane and metal-promoted B-N bond formation to form an oligomeric borazane bound to the metal centre. Steric factors play an important role in determining the barrier to B-N coupling which increases with x in the $H_3B \cdot NMe_x H_{3-x}$ series (x = 0-2). These studies also suggest a role for additional amine- or amino-borane in promoting dehydrocoupling processes through the formation of adduct species and complementary N-H···H-B interactions, an observation we noted from experimental studies both here and previously²¹ and recently from computational studies on related systems.43

An overall mechanism that captures these observations is shown in Scheme 6. For $H_3B \cdot NH_3$ initial dehydrogenation of the amine-borane (step 1) has the highest barrier (+26.7 kcal mol⁻¹), with the subsequent dehydrogenation of a second amine-borane (step 2) proceeding through a slightly lower energy transition state at +24.2 kcal mol⁻¹. The transition state for the B–N coupling of the resultant amino-boranes (step 3) is then most



Scheme 6 Overall mechanism for the dehydrogenation and B–N bond forming events for $H_3B \cdot NH_3$. R = H (first oligomerisation, *i.e.* to form **6b**); $R = BH_2NH_2$ (second oligomerisation, **6c**). Numbers on parenthesis are calculated barriers for the model system (kcal mol⁻¹). [Ir] = {Ir(PR_3)_2}^+ (R = Cy, experiment; R = Me, computation).

accessible of all (+17.9 kcal mol⁻¹). The rather high barrier to dehydrogenation (step 1) means that these systems turnover rather slowly, especially compared to others that rapidly promote dehydropolymerisation.⁶⁻¹² However, the corollary is that intermediates such as 6a-e can be observed, allowing for direct mechanistic insight. For subsequent oligomerisations (e.g. to form 6c, $R = BH_2NH_3$ Scheme 6) the key transition state energies retain the same pattern, thus promoting formation of a growing oligomeric chain at the metal centre. When the amineborane is changed to H₃B·NMeH₂ the same computed pattern still holds for the initial oligomerisation, but the second B-N coupling transition state (+24.8 kcal mol⁻¹) does becomes very close in energy to those for the two dehydrogenation steps (+25.4 kcal mol⁻¹ and +24.0 kcal mol⁻¹). Clearly B–N coupling is disfavoured by the greater bulk and experimentally only 5b is observed to be formed. For H3B·NMe2H no B-N bond formation to give a linear diborazane is observed under these experimental conditions, with 4a* formed only.

B–N coupling is also calculated to be competitive with amino-borane cyclisation, consistent with the observation of a small amount of borazine. However, coupling must be faster than reaction of exogenous cyclohexene with amino-borane as no hydroborated product is observed under these conditions. Our mechanism therefore has some similarities to those recently proposed for the catalytic dehydropolymerisation of $H_3B\cdot NH_3$ using a bifunctional Ru-based catalyst¹⁰ and of $H_3B\cdot NMeH_2$ using $Ir({}^tBuPOCOP{}^tBu)H_2$.^{6,12} Although the intimate mechanistic details of these two systems likely differ, both propose dehydrogenation to form an amino-borane, that then must undergo fast metal-mediated B–N coupling, as neither system promotes hydroboration when exogenous cyclohexene is added.

Amine-borane dehydrocoupling presents a high degree of mechanistic complexity that is additionally highly catalyst specific. Although the precise mechanism outlined here might be rather system specific, the observations and suggested pathways presented might help guide future work on developing and understanding this challenging transformation. Ultimately the goal is the design of improved catalysts for this important process that have the potential to produce B–N materials "to order".

Acknowledgements

The Rhodes Trust (A.K.), the University of Oxford, EPSRC (EP/J02127X/1) and the Spanish government (A.G.A.) for a Post-doctoral Fellowship (EX2009-0398).

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Grimme's D3 parameter set (*i.e.* BP86-D3) and solvent (C_6H_5F , PCM approach). See ESI[†] for references and full details.

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