Amine–Boranes as Transfer Hydrogenation and Hydrogenation Reagents: A Mechanistic Perspective
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Transfer hydrogenation (TH) has historically been dominated by Meerwein–Ponnord–Verley (MPV) reactions. However, with growing interest in amine–boranes, not least ammonia–borane (H$_3$N·BH$_3$), as potential hydrogen storage materials, these compounds have also started to emerge as an alternative reagent in TH reactions. In this Review we discuss TH chemistry using H$_3$N·BH$_3$ and their analogues (amine–boranes and metal amidoboranes) as sacrificial hydrogen donors. Three distinct pathways were considered: 1) classical TH, 2) nonclassical TH, and 3) hydrogenation. Simple experimental mechanistic probes can be employed to distinguish which pathway is operating and computational analysis can corroborate or discount mechanisms. We find that the pathway in operation can be perturbed by changing the temperature, solvent, amine–borane, or even the substrate used in the system, and subsequently assignment of the mechanism can become nontrivial.

1. Introduction

Hydrogenation is one of the most important and fundamental transformations used in chemistry. The direct addition of dihydrogen gas (H$_2$) across an unsaturated moiety is a well-developed area of reduction chemistry which has resulted in H$_2$ as the preferred hydrogen source in many of these transformations.[1] However, transfer hydrogenation (TH) offers an alternative pathway that avoids the use of highly pressurized gas and potentially proffers greater control in the level of reduction. Here, a sacrificial TH agent is used to donate hydrogen, whereby the TH agent is usually cheap, abundant, and easily manipulated. Early examples include Meerwein–Ponnord–Verley (MPV) reductions using secondary alcohols as the TH agent to reduce aldehydes and ketones to their respective alcohols.[2] Furthermore, progression into asymmetric transfer hydrogenation (ATH) reactions, mediated by ruthenium catalysts, were pioneered in the 1980s using isopropanol or an azetropic mixture of formic acid and triethylamine as the TH agent; this first report has since spawned a great interest in this area alone.[3]

Beyond isopropanol and formic acid, where the by-products formed, acetone and CO$_2$, respectively, are usually trivial to separate from the reaction mixture, additional TH agents reported in literature includes Hantzch esters,[4] dimethylformamide with a base as additive,[5] sodium hypophosphite,[6] benzothiazoline,[7] and hydrazine[8] although more commonly in the form of hydrazine hydrate.[9] More recently, ammonia–borane (H$_3$N·BH$_3$) has been studied and shown to be a promising addition to the numerous TH agents reported to date. The low molecular weight (30.87 g mol$^{-1}$), high hydrogen content (19.6 % wt %), and ease of handling as a bench-stable crystalline solid makes H$_3$N·BH$_3$ an attractive TH agent. Furthermore, derivatives of H$_3$N·BH$_3$, amine–boranes (R$_n$·H$_3$N·BH$_3$·R$_n$) and metal amidoboranes (MAB)[10] have also been studied as hydrogen donors but so far are less developed in the area of TH chemistry. The advantages of using amine–boranes over H$_3$N·BH$_3$ can be found in their: 1) greater solubility in common analytical solvents such as benzene and chloroform, 2) ease of identification of by-products in TH reactions as some amine–boranes are less likely to form insoluble polymeric substances, and 3) greater control of reduction by altering R groups on both the nitrogen and boron atom and by virtue of fewer hydrogen atoms available to transfer to the acceptor molecule.

In this Review we will present recent publications that use H$_3$N·BH$_3$ and amine–borane TH agents with an emphasis on understanding the mechanism operating in these reactions.[11] Pertinently, this Review is not an evaluation of the chemistry of dehydrogenation/dehydrocoupling (DHC) of amine–boranes, for which there are numerous reviews,[12] but instead focuses exclusively on the related tandem dehydrogenation TH process. Therefore, two fundamental questions present themselves in order to clarify the following discussion: What is the origin of the hydrogen atoms and how are they transferred to the substrate? We find that the literature studies reviewed here can be classed as: 1) classical TH processes whereby the double hydrogen transfer comes from both the amine and borane (Sections 2 and 3), 2) nonclassical TH processes whereby hydroboration from the amine–borane is followed by solvolysis (Section 4), and 3) hydrogenation via H$_2$ released from dehydrogenation of amine–borane (Sec-
2. Catalyst-Free Classical TH of Preactivated Substrates

The TH of substrates without a catalyst has been achieved for molecules containing a polarized unsaturated bond. These reactions therefore are not applicable to a great range of substrates but still provide vital mechanistic understanding into this elementary reaction that can be informative for catalyzed processes (Section 3). In addition to lowering the activation barrier using preactivated substrates, the formation of by-products from dimerization and cyclisation of amine–boranes could provide the entropic and enthalpic driving forces of the forward reactions. In this section we will review notable examples from the literature that have pioneered uncatalyzed TH using amine–boranes and confirm whether classical TH, nonclassical TH, or hydrogenation mechanism is in operation.

In 2010, Berke and co-workers reported the TH of imines with 1–2 equiv of H₃N-BH₃ in tetrahydrofuran (THF) to generate the corresponding amines in good to excellent yields with concomitant formation of borazine (BZ) or polyborazine (PBZ) as by-product (Scheme 1).[14] The working hypothesis for this reaction was that the polarity match between the protic H₂ and hydridic H₂ of H₃N-BH₃ with the polarized N≡C⁻ moiety of the substrate would allow for spontaneous double H transfer. To probe the mechanism of this reaction and confirm this hypothesis, the reaction temperature was kept below 60°C ensuring that no thermal decomposition of H₃N-BH₃ occurred and avoiding H₂ release, therefore simple hydrogenation was omitted as a reaction pathway.[15] Additionally, heating a mixture of H₃N-BH₃ and D₂N-BD₃ at 60°C in THF resulted in no deuterium scrambling, indicating that the adduct does not dissociate and therefore Lewis acid (BH₃) or base mediated (NH₃) transfer hydrogenation could also be discounted.

Deuterium labeling experiments were performed using benzyldiene aniline as the model substrate. Using H₃N-BD₃, deuterium incorporation was found solely at the C atom of benzylidene aniline as the model substrate. Using H₃N-BD₃, deuterium incorporation was found solely at the C atom of benzylidene aniline as the model substrate. Using H₃N-BD₃, deuterium incorporation was found solely at the C atom of benzylidene aniline as the model substrate.

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the imine moiety; conversely, using D$_3$N-BH$_4$ resulted in deuterium incorporation exclusively at the N atom. Using D$_3$N-BD$_3$ gave deuterium incorporation both at the C and N atoms. These experiments corroborated the polarity match mechanism, which was found to be feasible based on quantum mechanics calculations performed. A six-membered transition state (TS) whereby the double H transfer of N–H–N and the B–H–C was found to be 23.5 kcal mol$^{-1}$ more favorable than the polarity mismatch configuration. In order to decipher whether the mechanism was via a concerted or stepwise process, primary deuterium kinetic isotope effect (DKIE) experiments were undertaken, revealing: 1) inverse DKIE (0.87) when H$_2$N-BD$_3$ was used, 2) a normal DKIE (1.93) when D$_3$N-BH$_4$ was used, and 3) a small positive DKIE (1.39) when D$_3$N-BD$_3$ was used. Hammett correlations also revealed positive values of the sensitivity constants ($\rho$) for para-substituted benzylidene anilines (substitution at aniline side, $\rho = 1.61$ and substitution at benzylidene side, $\rho = 0.69$). All these results indicate an asynchronous concerted double H transfer, whereby the breaking of the N–H bond was the rate-determining step (RDS) of the transformation.

In related studies, Berke and co-workers expanded this methodology for polarized olefins (Scheme 2).[16] The reactions were too quick to be monitored by NMR spectroscopy in THF, so acetonitrile was used as slower reactivity was observed in this solvent. Deuterium labeling experiments confirmed the polarity match of the hydridic H$_a$ and protic H$_b$ transfer to the C atom with aryl/alkyl groups and C atom with electron-withdrawing substituents, respectively. However, the measured DKIE using 2-cyclohexylidenemalononitrile as the model substrate revealed 1) no DKIE (1.00) when H$_2$N-BD$_3$ was used, 2) a normal DKIE (1.55) when D$_3$N-BH$_4$ was used, and 3) a normal DKIE (1.61) when D$_3$N-BD$_3$ was used. This indicated that this was a stepwise process whereby the RDS involved cleavage of the N–H bond. Further experimentation of a 1:3 mixture of H$_2$N-BH$_4$ with 2-cyclohexylidenemalononitrile at $-40^\circ$C in [D$_6$]THF (and also in CD$_3$CN) allowed the authors to observe the hydroborate intermediate INTI by in situ multinuclear NMR spectroscopy, suggesting the mechanism involved a facile hydroboration step preceding the RDS. This mechanism was distinctly different to that observed with imines (vide supra). Continuing on from this work, Berke and co-workers were also able to effect the TH of polarized olefins using MeH$_2$N-BH$_4$, BuH$_2$N-BH$_4$, and Me$_3$HN-BH$_4$ as well as H$_2$N-BH$_4$ as the hydrogen donor.[17]

Mechanistic investigation showed similar results of a stepwise double H transfer.

In 2011, Manners and co-workers reported the metal-free TH between several amine–boranes and the aminoborane $i$Pr$_3$N=BH$_2$ in THF at 20$^\circ$C (Scheme 3a).[18] Experimental and computational investigation into this reaction followed.[19] Me$_3$HN-BH$_4$ was chosen as the model substrate in these studies. Experimentally the reaction proceeded more cleanly than the reaction with Me$_3$N-BH$_4$ and H$_2$N-BH$_4$, with the only side-product being cyclodiborazane [Me$_3$N–BH$_3$]. However, the reaction reached equilibrium at $\approx$50% conversion.

![Scheme 2. TH of polarized olefins via stepwise double H transfer.](image)

Overall, a mechanism analogous to that reported by Berke for the TH of imines[15] was proposed, involving an asynchronous concerted double H transfer (Scheme 3b). However, the thermodynamic accessibility of the two reactions was vastly different. When the TH of $i$Pr$_3$N=BH$_2$ with Me$_3$N-BH$_4$ was monitored by multinuclear NMR spectroscopy at varying temperatures, the calculated thermodynamic parameters showed the TH from Me$_3$HN-BH$_4$ to $i$Pr$_3$N=BH$_2$ was endergonic ($\Delta G^{\circ}_{(298)} = 10 \pm 7$ kJ mol$^{-1}$) but the dimerization of the transient [Me$_3$N=BH$_3$] species was more exergonic ($\Delta G^{\circ}_{(298)} = -28 \pm 14$ kJ mol$^{-1}$) and therefore driving the reaction in the forward direction. The measured large entropy and small enthalpy of activation for the forward TH reaction ($\Delta S^{\circ}_{(298)} = -210 \pm 11$ kJ mol$^{-1}$ and $\Delta H^{\circ}_{(298)} = 29 \pm 5$ kJ mol$^{-1}$) were consistent with a highly ordered bimolecular TS, suggesting a concerted TS with values similar to those previously reported for Diels–Alder reactions.[20] DKIE experiments with $i$Pr$_3$N=BH$_2$ showed a large positive DKIE ($k_{\text{DKIE}} = 6.7 \pm 0.9$) when Me$_3$DN=BH$_4$ was used, but a small inverse DKIE ($k_{\text{DKIE}} = 0.7 \pm 0.1$) with Me$_3$HN-BD$_3$ and a large positive DKIE ($k_{\text{DKIE}} = 3.2 \pm 0.8$) with Me$_3$DN-BD$_3$. Manners and co-workers rationalized the

![Scheme 3. a) Reversible TH of $i$Pr$_3$N=BH$_2$ with Me$_3$HN-BH$_4$ and b) simplified reaction profile of the forward reaction.](image)
small inverse DKIE obtained for the hydride transfer as the result of a secondary kinetic isotope effect and the change in the geometry around the boron atom at the TS. Density functional theory (DFT) calculations of these thermodynamic parameters gave a good match to the experimental values. Furthermore, DFT calculations showed that alternative pathways, such as stepwise addition with B–H→B transfer first, stepwise addition with N→H→N transfer first, or a dissociative process were energetically unfeasible and did not align with the experimental evidence.

It is worth noting that this study focused on Me$_2$HN-BH$_3$ as the TH partner. However, when RR'HN-BH$_3$ was used (where R = H and R' = Me or H), an additional by-product was observed: [H$_2$B(μ-H)(μ-NRR')BH$_3$]. This would suggest that under these reaction conditions a stepwise or dissociative pathway could be operating, and highlights the sensitivity of these reaction pathways and how they could be perturbed by simply changing the substituents on the amine–boranes used.

More recently, Braunschweig and co-workers reported the TH of three iminoboranes with bulky R substituents (R-N=B-R', where R = Bu and R' = Bu, mesityl, or 2,3,5,6-tetramethylphenyl) with H$_2$N-BH$_3$.[21] They calculated that the formation of two aminoboranes as the products (more accurately the cyclization products from H$_2$N-BH$_3$) would be thermodynamically favorable to drive the reaction forward. Placing Bu-N=B-Bu under high H$_2$ pressure led to no hydrogenation even in the presence of Pd/C catalyst, indicating that a classical TH process was occurring. Although the multinuclear NMR and FTIR spectroscopic data supported the formation of BuHN=BBuH as the product, the cis/trans stereochemistry of this aminoborane was not clear. Isolation of the products for X-ray diffraction analysis from the subsequent reaction of BuHN=BBuH with HCl or the N-heterocyclic carbene 1,3-disopropylimidazol-2-ylidene (IPr) suggested the aminoborane carried trans stereochemistry.

Probing the mechanism further, deuterium labeling experiments using D$_2$N-BH$_3$ and H$_2$N-BD$_3$ confirmed the polarity matching of the substrates. However, no DKIE experiments were reported to substantiate the DFT calculations, which supported a concerted double H transfer through a very low-energy TS (5.4 kcal mol$^{-1}$) (Scheme 4). This concerted addition would lead to the cis-aminoborane, which was 8.4 kcal mol$^{-1}$ higher in energy than the trans-aminoborane. Observation of the cis conformation would align with the proposed mechanism, but the trans-aminoborane as the final product was inferred experimentally (vide supra) and could indicate a stepwise pathway instead. However, a rotation around the N–B bond to allow the isomerization from cis to trans was found through a relatively high barrier of 17.8 kcal mol$^{-1}$ at room temperature. This isomerization step would therefore be the RDS and in theory the cis-aminoborane should be observed by in situ multinuclear NMR spectroscopy prior to isomerization, but this was not reported by the authors. This could indicate that an alternative isomerization pathway with a much lower barrier could be involved than the one calculated and reported; low-temperature studies could give insight.

In 2012, Chen and co-workers reported a number of studies using lithium amidoborane (LiH$_2$N-BH$_3$) and calcium amidoborane (Ca(H$_2$N-BH$_3$)) to chemoselectively reduce α,β-unsaturated ketones to allylic alcohols under ambient temperature (Scheme 5a,b).[19,23] Using MABs circumvented the use of conventional reducing agents such as NaBH$_4$, which often has poor selectivity from over-reduction of the substrate, or using lithium aminoborohydrides (LiR-N-BH$_3$, where R = H), which requires a subsequent hydrolysis step. Optimization of the reaction found THF to be the best solvent, as MeOH resulted in solvolysis of the MAB. Keeping the reactions at ambient temperature negated dehydrogenation of the MAB, with these processes occurring at elevated temperatures (LiH$_2$N-BH$_3$, ≈40°C[21]; Ca(H$_2$N-BH$_3$), ≈80°C[24]). Deuterium labeling experiments using [M''-](D$_2$N-BH$_3$)$_{n-}$ (M = Li or Ca) showed deuterium incorporation only at the oxygen, and when [M''-](H$_2$N-BD$_3$)$_{n-}$ was used only at the C atom of the carbonyl moiety, confirming the double H transfer process and that the hydrogen came from the respective MAB. A further comparison of the two different MABs revealed that Ca(H$_2$N-BH$_3$)$_2$ was more competent at the TH of α,β-unsaturated aldehydes to the allylic alcohol (Scheme 5c). When the group reacted LiH$_2$N-BH$_3$ with the substrate,
they observed full conversion of the starting material but only \( \approx 50\% \) of the allylic product was formed according to \( ^1H \) NMR spectroscopy. A white precipitate observed in the reaction mixture was assigned to a lithium aminoboranylborane species, which upon hydrolysis with aqueous HCl gave the desired alcohol. This lower reactivity was postulated as a result of poor solubility of the intermediate in THF and the potentially higher enthalpic penalty of losing a second (Li)N–H bond versus (Ca)N–H bond.\(^{[10b,c]}\)

Chen and co-workers followed up this work by reporting the TH of ketones and imines with LiH,N-BH\(_3\), Ca-(H,N-BH\(_3\)), and also sodium amidoboranes (Na(H,N-BH\(_3\))) with high conversion to secondary alcohols and amines, respectively, across all MABs used.\(^{[25]}\) Higher reactivities were displayed by the MABs in comparison to H,N-BH\(_3\) in these TH reactions. This was attributed to the weaker B–H bond of the former due to a more electron-rich B center\(^{[10b]}\) and also M–...–B interactions.\(^{[26]}\) The reaction mechanism of these TH reaction kinetic studies was probed using LiH,N-BH\(_3\) with benzophenone and N-benzylideneaniline, and a first order dependence with respect to LiAB and a zeroth order dependence on the substrate was found. Additional DKIE experiments carried out using benzophene revealed a small positive DKIE (1.26) with LiD,N-BH\(_3\) and a larger positive DKIE (1.89) with LiH,N-BD\(_3\). Similar values were obtained when N-benzylideneaniline was used (LiD,N-BH\(_3\); 1.26; LiH,N-BD\(_3\); 2.12). From these experiments they proposed that the B–H bond breaking is involved in the RDS. Although MABs were reported to be superior TH agents than H,N-BH\(_3\) in these studies, the addition of the alkali and alkaline earth metals complicates the mechanism operating in these reactions. Chen et al. reported a complementary DFT investigation of the TH of N-benzylideneaniline with LiH,N-BH\(_3\) (Scheme 6).\(^{[25]}\) A calculated pathway was found which involved the initial elimination of LiH from LiH,N-BH\(_3\), representing the RDS of the reaction at \( \Delta G^\ddagger = 17.2 \text{kcalmol}^{-1} \). This RDS agreed with the kinetic and DKIE experiment showing zeroth order dependence on the substrate and the breaking of the B–H bond in this step. Chen et al. also attributed the small DKIE observed when LiD,N-BH\(_3\) was used as a consequence of the small difference in energy between the two TSs, \( \Delta G^\ddagger = 3.2 \text{kcalmol}^{-1} \), involving both the N–H and B–H bond-breaking steps. Furthermore, they found a higher RDS (\( \Delta G^\ddagger = 28.0 \text{kcalmol}^{-1} \)) when H,N-BH\(_3\) was used as the TH agent which matched the higher reactivity displayed by LiH,N-BH\(_3\) in the reduction reactions.

3. Catalyzed Classical TH Reactions

In this section we highlight the chronological development of catalyzed, along with some stoichiometric, TH reactions where amine–boranes are required as hydrogen source and precatalyst activator; it is vital to comprehend that in a classical TH the amine–borane assumes this double role, allowing the formation of an active species/catalyst and also furnishing the H’/H’ critical to reduction. Classifying the following reactions as classical TH therefore makes it possible to distinguish them from nonclassical TH (Section 4) and hydrogenation reactions (Section 5). Following the aim of this Review, the focus will be given to studies where the mechanistic investigations are detailed. It is worth noticing that some reactions cannot be classified exactly in the three main categories that we have chosen to investigate, and that grey areas exist with mechanistic changes occurring with varying substrates and/or reaction conditions. Therefore, we carefully comment and propose a rationale for these unclear points to allow a complete description of the topic and provide a thoughtful analysis of classical TH with amine–boranes.

3.1. Metal Catalysis

One of the first examples of metal-catalyzed transfer hydrogenation of olefins with amine–boranes was reported by Berke and co-workers (Scheme 7).\(^{[27]}\) The authors used 1 mol% of [Re(Br)\(_2\)(NO)(PCy\(_3\))(η\(^3\)-H\(_2\))] \( \text{I} \) to convert octene into octane with Me,HN-BH\(_3\). The reaction allowed quantitative conversion in 1 h, irrespective of whether the reaction was carried out in an open or closed vessel, suggesting that H\(_2\) was not responsible for the reduction. Initial stoichiometric studies hinted to the importance of transient phosphine dissociation from the metal precursor to allow oxidative addition of the amine–borane to Re\(^{\text{II}}\), with an excess of phosphine found to decrease reactivity drastically. The reaction mechanism proceeded stepwise, with initial B–H o-bond activation and oxidative addition to form a Re\(^{\text{III}}\) species, followed by hydride insertion and Re–alkyl bond formation. After a \( \beta \)-hydride shift to liberate cycloborazine, the reductive elimination step ensured product formation and catalyst regeneration.
The authors followed up on the TH with Me₂HN-BH₃ by using the five-coordinate [Re(Br)(NO)(PCy₃)(H)(L)] (L = 1,3-dimesitylimidazol-2-ylidine IMes or PCy₃) precatalyst, which allowed a slight increase in the reaction efficiency (Scheme 13b). Subsequently the Re-catalyzed TH reaction was applied to the reduction of terminal olefins, relying, however, on the ethanalysis of amine–boranes (Section 4).

In 2013, Lin and Peters explored the reduction of olefins by Co–boryle complex 2 (Scheme 8a). The formation of a Co–hydridoborane complex 4 was found when 2 was subjected to an excess of Me₂HN-BH₃ (Scheme 8b), and the structure was confirmed by X-ray analysis and NMR spectroscopy. It was noted that the reduction of styrene was dramatically faster under an atmosphere of H₂ with full conversion to ethylbenzene in 1 h versus the 24 h needed when Me₂HN-BH₃ was used. Paul and co-workers analyzed the computational details of this reduction, and the authors calculated that in the presence of amine–borane, complex 2 can form 3 via an associative mechanism with an activation energy of 24.7 kcalmol⁻¹. However, the generation of active catalyst 3 was found to be more energetically demanding with Me₂HN-BH₃ than its formation in the presence of molecular H₂ (17.3 kcalmol⁻¹), which confirms the results observed experimentally. Co–hydridodiborane 4, which formed only in the presence of amine–borane, was found to be off-cycle and was described as resulting from the decomposition of 3. Further experimental and computational details from Paul and co-workers showed that an excess of base NEt₃ can convert 4 back to 2.

Studies from Cazin and co-workers showed an efficient [Pd(NHC)(PR₃)]-catalyzed TH of alkenes and alkynes with H₂N-BH₃ (Scheme 9). The active intermediate [Pd(H)₃(NHC)] 5 was isolated, and its formation and role in the hydrogenation was computationally clarified by Yi and co-workers. Intermediate 5 formed via a sequential ligand-assisted N–H followed by B–H activation of H₂N-BH₃ (∆G* = 23.8 kcalmol⁻¹). Stepwise TH from 5, instead of molecular H₂, was found to be kinetically and thermodynamically favorable with an energy barrier of 22.3 kcalmol⁻¹, thus highlighting TH not hydrogenation is in place in this Pd-catalyzed system. The role of iPrOH in the reaction mechanism was not analyzed in detail, thus the possibility of solvolysis for this TH process cannot be ruled out (Section 4).

A nickel version of alkyne TH was performed by Garcia and Barrios-Francisco using [Ni(dppe)(η²-dpa)] (dpa = diphenylacetylene); the authors showed the selective semi-TH of alkynes with H₂N-BH₃. The observed stereoselective divergency to cis or trans olefin was dependent on the solvent, with THF favoring the former while MeOH furnished trans selection. It is worth highlighting that a nonclassical TH might be in action when polar protic solvent is used, following our considerations on solvolysis-mediated TH reactions (Section 4). No trace of H₂ was evident by GC-MS in the catalytic tests, which highlights that the TH mechanism might be the preferred pathway for this transformation.

The first example of semi-TH of alkynes with H₂N-BH₃ in the presence of copper was reported in 2017, the authors used air-stable [Cu(IPr)(OH)] and an excess of H₂N-BH₃ to selectively reduce alkynes to (Z)-alkynes in THF at 50°C, and further expanded the substrate scope to the full reduction of propiolates. A blank reaction under 1 bar of H₂ with a catalytic amount of H₂N-BH₃ (20 mol%) allowed only 20% conversion into product, which suggested a direct hydride transfer mechanism to be in place.

Wang, Liao, and co-workers reported an efficient sequential dimerization/semihydrogenation reaction of alkynes into (E,Z)-1,3-dienes with Co⁹ complex 6 and H₂N-BH₃ (Scheme 10a). The authors distinguished the two sequential catalytic cycles and studied the full reaction profile both experimentally and by DFT analysis. Initial dimerization of the alkyne to a 1,3-enzyme proceeded through metal–ligand cooperative

Scheme 7. a) General TH of olefins with Me₂HN-BH₃ and with Re¹ precatalyst 1; b) range of Re¹ precatalysts and TOF results for the TH of octene after 1 h reaction at 85°C.

Scheme 8. a) General scheme of the Co¹-catalyzed hydrogenation of styrene with Me₂HN-BH₃; b) major reaction intermediates with details on the RDS for active catalyst formation (kcal mol⁻¹).
activation of the alkyne by 6 to form an alkylidene–Co complex 7 (Scheme 10b); the latter could further react through an anti-Markovnikov addition to a second equivalent of alkyne, releasing the 1,3-enyne, and reform precatalyst 7. The subsequent addition of H₃N·BH₃ to the reaction mixture allowed the pyridonate ligand mediated formation of a [Co–H] intermediate (Scheme 10c); this step was proposed to proceed via borane activation by nucleophilic attack of the pyridonate ligand with ammonia release (ΔG° = 22.1 kcal mol⁻¹), which further re-enters the cycle and attacks the new borate formed to allow Co–H bond formation (RDS with ΔG° = 23.9 kcal mol⁻¹). The latter hydride species further reacts with the enyne for the transfer reduction; the hydride transfer to the α-carbon versus the β-carbon atoms of the enyne differed by 0.5 kcal mol⁻¹, which did not allow to distinguish the fate of H⁻ incorporation. This hypothesis was further corroborated by deuterium labeling studies, which showed that there was no distinction between deuterium incorporation in the 1,3-diene with H₃N·BH₃ or D₃N·BH₃. Final facile intramolecular protonolysis by the amino group releases the product with a cis configuration of the reduced triple bond.

Another example of TH and hydrogenation of olefins was reported by Wolf and co-workers. The authors used catalytic amount of a reduced cobaltate anion [K(thf)₁.₅{(IPr(BIAN))Co(COD)}] (BIAN = bis(iminoacenaphthene)dia mine) to perform dehydrogenation of H₃N·BH₃, TH of disubstituted olefins and imines with H₃N·BH₃ and hydrogenation of tri- and tetrabuckled alkynes (Scheme 11).

The kinetic profiles for the dehydrogenation reaction showed a second order rate in catalyst 8 (Scheme 12), suggesting the formation of a dinuclear [Co–H] active species, with catalyst deactivation observed at higher conversions; Hg drop test and P(OMe)$_3$ poisoning experiment did not affect conversions. However, when the strongly coordinating ligand dibenz[a,e]cyclooctatetraene (dct) was used, the reaction slowed down and traces of dct partial hydrogenation were found, which suggested the process to be homogeneous. Moreover, an induction period was observed at low catalyst loading that was found to be related to partial hydrogenation of 1,5-cyclooctadiene through poisoning experiments. The mechanism for TH was derived from these initial findings, with an observed reaction rate similar to that of the dehydrogenation reaction, which highlights that the dinuclear [Co–H] species is the common species formed in both the TH and dehydrogenation reactions. When TH of α-methylstyrene was performed in a D₂ atmosphere (1.1 bar), 20% of deuterium incorporation into the cumene product was found, which supported a H₃N·BH₃-mediated reduction. In contrast, when bulky olefins were subjected to the optimized reaction conditions, 10 bar of H₂ was necessary to ensure product formation. This highlights that hydrogenation is in action with hindered substrates.

Liu et al. presented a catalyzed TH of nitriles by H₃N·BH₃ (Scheme 12). The NNP-type cobalt pincer complexes 9 and 10 were active for a chemodivergent nitrile hydrogenation to primary, secondary, or tertiary amines, depending on the solvent. When hexane was used and benzonitrile was subjected to the reaction conditions, benzylamine was found to be the predominant product, while switching to hexafluoro-2-propanol (HFIP) led to the selective formation of secondary dibenzylamine. The reaction worked well with N–H or
N–Me complexes, which highlights that an outer-sphere mechanism with ligand cooperativity might not be involved in this reaction, while an inner-sphere mechanism should be considered. Moreover, the involvement of HFIP should not be discarded, as the authors reported when studying olefin TH (Section 4.2).[39]

Webster and co-workers presented a rare example of Fe$^{II}$-catalyzed TH and semi-TH of olefins.[40] The β-diketiminate iron alkyl precursor 11 in the presence of sacrificial amines and borane selectively reduced alkenes and alkynes (Scheme 13a). The catalytic system was found to be extremely robust when amino substituents were included in the substrate, by eliminating the use of a sacrificial external amine. Labeling experiments showed that selective anti-Markovnikov mono-deuteration occurred with deuterated aniline, while incorporation of deuterium was found to be predominant in the internal positions (Markovnikov product) when DBpin was used (Scheme 13b).

Gelation was observed when the TH of olefins was performed, which, together with the lack of competitive hydroboration of substrates, prompted the authors to analyze the involvement of oligomeric species in the reaction mechanism. The formation of dimeric and tetrameric species of [nBuH,N-BHPin]$_x$ ($x=1$: AB2; $x=2$: AB4) was proven based on the evident shift of the B–H signal versus free pinacol borane in the $^1$B NMR spectrum; these species were found to be entropically favored compared to DHC adducts. Subsequent DFT calculation of the catalytic cycle highlighted the formation of an [Fe–H] active species and 1,2-alkene insertion followed by rate-limiting protonolysis with a $\Delta G^\ddagger = 17.3$ kcal mol$^{-1}$ (Scheme 13c). Therefore, the formation of oligomers in this transformation was found to be crucial to decrease the concentration of free borane and amine in solution and to allow the TH process to be the preferred pathway.

Punji and Sharma studied the TH of nitriles to secondary amines using [Co(Xantphos)Cl$_2$].[41] The transformation was found to be dependent on the amine–borane used; with H$_3$N·BH$_3$, the selective formation of symmetrical secondary amine was observed. When Me$_2$HN·BH$_3$ was used and the reaction was performed in the presence of a second equivalent of amine, the synthesis of unsymmetrical amines resulted instead.

Wang, Liao, and co-workers also described a catalytic TH of nitriles using a molybdenum–thiolate complex to synthesize primary amines in the presence of H$_3$N·BH$_3$ (Scheme 14a).[42] The authors used [Cp*Mo(1,2-Ph$_2$PC$_6$H$_4$S)(h$_2$-NCMe)] which promptly reacts with H$_3$N·BH$_3$, activating the B–H bond via a metal–ligand cooperative action and forming a neutral Mo$^{II}$-H/borohydride species (13). The mixed species 13 was analyzed by variable-temperature $^1$B NMR spectroscopy, which showed that a fast interchange was in effect between the Mo–H–B and the B–H bond at room temperature.

\[\text{Scheme 13. a) Fe}^{\text{II}}\text{-catalyzed TH of alkenes and alkynes; b) selective deuterium incorporation with labeled aniline or HBpin; c) details of the computed free energy profiles (kcal mol}^{-1}\text{) for the competing DHC of AB2, preferred quintet spin state shown.}\]

\[\text{Scheme 14. a) Molybdenum–thiolate catalyzed TH of nitriles; b) formation of the [Mo–H]/borohydride mixed species 13 and details regarding its }^{11}\text{B NMR chemical shifts; c) details of the Gibbs free energy diagram (kcal mol}^{-1}\text{) of the catalytic cycle.}\]
temperature (Scheme 15b). The latter species 13 could catalyze the TH of nitriles but only in the presence of added H,N-BH₃. Kinetic studies showed a first order dependence in H,N-BH₃ and precatalyst 12, while the reaction was zeroth order in substrates, which ruled out nitrile activation as the RDS. Further DKIE values were determined, (kₒ,D,N-BH₃ = 1.4, kₒ,H,N-BH₃ = 2.6, and kₒ,D,N-BD₃ = 3.3) which could suggest that both N–H and B–H bond activation are involved in the RDS. However, further computational calculations to describe the catalytic cycle indicate that, instead, Mo–H insertion into the C= N bond with a ΔG° of 22 kcal mol⁻¹ is the RDS (Scheme 14c). Intermediate 13 was found to be the resting state of the catalytic cycle. Protonolysis via H,N-BH₃ releases the product, while protonation by free NH₃ was discarded.

![Scheme 15. a) Initial finding for the catalyzed reduction of azoarenes with 14 using H,N-BH₃ as the hydrogen source and the proposed formation of P₃–hydride 16, b) revised mechanistic study highlighting P–ONO ligand cooperativity.](image)

3.2. Metal-Free TH

In the metal-free reduction reactions with amine–boranes, most of the examples follow concerted TH pathways with the formation of a six-membered-ring transition state analogous to the work published by Berke’s group on the TH of imines with H,N-BH₃.[12] with the exception of a reported stepwise pathway for CO₂ reduction (vide infra).

The development of pinoctig-catalyzed TH of azoarenes with H,N-BH₃ was initiated by the Radosevich group (Scheme 15a).[13] The authors reported the synthesis of the T-shaped strained P₃III compound 14, which was able to oxidatively add H₂ from H,N-BH₃ and form the active P₃V compound 16. The latter was isolated and proposed to be the active species which enabled this catalytic transformation. However, later computational studies on the reaction mechanism suggested that a P₃V–ligand cooperativity might be in action instead of the initially hypothesized active P₃III/P₃V redox cycle (Scheme 15b).[14] Concerted activation of H,N-BH₃ with a barrier of ΔG° = 27.1 kcal mol⁻¹ allowed formation of species 15 (ΔG° = 13.5 kcal mol⁻¹), the active species for the TH of azobenzene to 1,2-diphenylhydrazine through a concerted six-membered-ring TS (ΔG° = 28.1 kcal mol⁻¹). This step can therefore be denoted as the RDS of the reaction.

Dimerization of 15 resulting in formation of the P₃–hydride species 16, originally isolated by Radosevich’s group, was found to be energetically feasible (ΔG° = 8.9 kcal mol⁻¹). However, it was determined that 16 was an off-cycle species for the TH of azobenzene, and other potential mechanistic pathways, for example, insertion of the N=N bond into P–H bond (43.2 kcal mol⁻¹) or reduction of azobenzene through ion-pair interaction with P–H (29.4 kcal mol⁻¹), were discarded because of the high energy requirements.

Following these findings, Hirao and Kinjo reported the catalytic reduction of azoarenes with H,N-BH₃ and 5 mol % 1,3,2-diazaphospholenes (DAPs) (Scheme 16a).[15] This process was more efficient than Radosevich’s procedure, where catalysis was generally faster. A proposed catalytic cycle was presented which starts with insertion of the P–H bond into N=N, followed by hydrogenolysis of the exocyclic P–N bond via H,N-BH₃. The latter hydrogen transfer was found to proceed through a concerted six-membered-ring TS with protic and hydridic hydrogen transfer to N and P atoms, respectively. This pathway was energetically feasible (ΔG° = 25.2 ± 4.2 kcal mol⁻¹, ΔH° = ± 2.2 kcal mol⁻¹, and ΔS° = ± 11.6 ± 6.8 e.u.), with oligomerization of H,N-BH₃ accounting for the slightly endergonic nature of the reaction. Additional stepwise pathways were analyzed but discarded as they were found to be more energetically demanding. DKIE analysis for the reduction of azobenzene to 1,2-diphenylhydrazine was carried out using isotopically labeled H,N-BH₃. These experiments revealed normal KIEs with D,N-BD₃ (3.05), H,N-BD₃ (1.44), and D,N-BD₃ (4.67). These values demonstrated that B–H and N–H activation of H,N-BH₃ are

![Scheme 16. a) Proposed catalytic cycle for the TH of azoarenes by DAPs with H,N-BH₃; b) phosphite reduction of azoarenes with H,N-BH₃.](image)
involved in the RDS. When H$_2$N-BD$_3$ was used, incorporation of deuterium was found to be selective for 1,3,2-diazaphospholene recovered at the end of the reaction, with traces of PH$_3$ formation. The role of phosphane in the reaction was not investigated further.

Recent findings of Landaeta and co-workers on a similar reduction reaction with an acyclic phosphite precatalyst (Scheme 16b) substantiated that the reduction of azoarenes with pnictogenate precatalysts is prone to an associative mechanism. The authors presented a detailed mechanistic study of the latter confirming, through DKIE values following the same trend as that in Hirao and Kinjo’s work ($k_{\text{H}_2\text{N-BH}_3} < k_{\text{D,N-BH}_3} < k_{\text{D,N-BH}_3}$), that concerted B–H and N–H bond breaking was involved in the slowest reaction step.[47]

With the rise of frustrated Lewis pair (FLP) chemistry and following the finding that H$_2$ is released from H$_2$N-BH$_3$ initiated by Bi(C$_5$H$_5$)$_3$ (BCF),[46] TH reactions were performed using Lewis acid activation. In one of the first examples, Du and co-workers[49] reported the activation of H$_2$N-BH$_3$ with BCF (10 mol%) to obtain a stereoselective reduction of pyridines to piperidines (Scheme 17). The authors proposed the formation of a zwitterion species of type 17 resulting from hydride abstraction of H$_2$N-BH$_3$ from the Lewis base/Lewis acid adduct. A similar hypothesis was proposed by Shi and co-workers in the development of N-heteroarene reduction,[50] by Xiao and co-workers for the deoxygenation of amides and lactams,[51] and by Zhong and co-workers for the reductive amination of ketones.[52]

Du, Meng, and co-workers reported an asymmetric and stereoselective TH of imines using H$_2$N-BH$_3$ (Scheme 18a).[53] The authors exploited the idea of zwitterion pair ion by using chiral tert-butyl sulfonamide and Piers’ borane HB(C$_5$H$_5$)$_3$, originally finding success in the stoichiometric reaction and then transposing it into a catalytic process (10 mol %). It is worth noting that H$_2$ (20 bar) was not an efficient reductant, with low conversion found compared to the reaction performed with H$_2$N-BH$_3$ (10% versus 99% conversion after 20 h). The O-isomer 18 was proposed to be the active intermediate and $^1$B NMR analysis of the catalytic reaction allowed identification of the species, which showed a broad signal at $-4.8$ ppm (Scheme 18b). Formation of this isomer, from addition of Piers’ borane and the sulfonamide, was also found to be the most likely by DFT calculations. From this active species TH occurred via the eight-membered TS1-(S), which is responsible for the enantioinductive step. Release of chiral amine and formation of dehydrated species 19 were confirmed by NMR spectroscopy, with 19 having a characteristic $^1$B NMR signal at 1.3 ppm. Stoichiometric experiments showed that 19 could quickly regenerate active catalyst 18 in the presence of H$_2$N-BH$_3$ via an energetically viable concerted pathway ($\Delta G^+ = 14.4$ kcalmol$^{-1}$). The authors hypothesized the role of 19 as a Brønsted acid initiator for the reaction, although the barrier found for this process ($\Delta G^+ = 29.0$ kcalmol$^{-1}$) was not comparable to that of the FLP mechanism. Moreover, 20, a dimer of 19, was isolated and proven to be unable to perform catalysis and described as an off-cycle species for the TH of imines.

Further advancement of the asymmetric TH of imines and ketones with H$_2$N-BH$_3$ was described by Du and co-workers when they used enantioenriched phosphoric acid (Scheme 19a,b).[54] The chiral ammonia–borane complex 21 was isolated but, while it was found to be off-cycle for ketone reduction (Scheme 19b), it was proven to be an active intermediate in the TH of imines by stoichiometric studies and DFT calculations (Scheme 19a). Interestingly for both processes, DFT calculations supported the formation of a six-membered-ring TS which accounted for substrate activation by amine–borane chiral complex 21 for imine reduction, while it involves substrate activation by phosphoric acid to form species 22, followed by TH from H$_2$N-BH$_3$ for ketone reduction.

A recent development of catalysis by pnictogens was reported by Cornella and co-workers who used a well-defined Bi$^3$ compound to deliver TH of azo- and nitroarenes (Scheme 20a,b).[55] The authors suggested an elusive bismuthine(III) hydride species might form by oxidative addition to Bi$^3$ (Scheme 20c); the evidence for such species was given by analyzing the catalytic reaction by high-resolution mass spectrometry (HRMS), which showed an adduct at 453.1738 g mol$^{-1}$ assigned to a cationic Bi$^{10}$ monohydride complex 23. Release of H$_2$ was evident when the dehydrogenation of H$_2$N-BH$_3$ was performed with bismuthine.
kinetic isotope effect, with $k_{D,N,BH_3} = 1.63$, $k_{H,N,BD_3} = 3.94$ and $k_{D,N,BD_3} = 7.05$ which indicated a concerted TS as RDS, reminiscent of the results found for DPAs and phosphite-catalyzed TH (Scheme 16).

A computational exploration of the potential of SCS-Ni pincer complexes in the TH of acetone, acetonaphone, and methanamine with H$_2$N-BH$_3$ has also been reported.[50] The calculations reveal that a proton-coupled hydride transfer is the more energetically demanding step for the reduction of ketones, while a stepwise hydride and proton transfer might occur in the TH of imines (Scheme 21). The key to this transformation was the imidazolium substituent on the SCS ligand that acted as proton shuttle. The reactivity of these complexes was compared to the reactivity of lactate racemates,[55] where the potential role of the metal center might be solely to stabilize the molecular entity which needs to perform the transformation, as observed by others.[56] Future experimental and mechanistic details would be of great interest to clarify and test the potential of this theoretical exploration.

The final example of TH is that of CO$_2$ and it differs slightly from the non-metal-catalyzed TH examples reported so far. Initially reported by Ménard and Stephan in 2010 (Scheme 22a),[59] the stoichiometric reduction of CO$_2$ to MeOH was performed by tris(2,4,6-trimethylphenyl)phosphine (PMes$_3$) and an excess of AlX$_3$ (X = Cl, Br) FLP species; the formation of FLP-CO$_2$ adducts 26 and 27 was confirmed by NMR spectroscopy and X-ray analysis, and they could be further reduced with H$_2$N-BH$_3$ and quenched with H$_2$O. The reduction reaction was found to be facile allowing a moderate yield (37–51%) of MeOH after 15 min at room temperature. The mechanism of this TH was further studied by computational methods by Paul and co-workers (Scheme 22b).[60] The reduction of CO$_2$ to liquid fuel was found to be initiated by interaction of the hydridic B-H with the C1 atom of FLP-CO$_2$ and PMes$_3$ displacement. The energy barrier to TS$_{1}$ of 15.1 kca mol$^{-1}$ was in line with the mild experimental conditions. Subsequent reduction steps were calculated to be driven by B–H activation, with the exception of the hydrolysis step which allows final C–O bond cleavage, which was postulated to be facilitated by H$_2$N-BH$_3$ or dehydrated oligomers, as found by Webster and co-workers (Scheme 13, Section 3.1). Interestingly, the authors compared this mechanism to the uncatalyzed reduction of CO$_2$ to formic acid,[61] for which they could locate a six-membered-ring TS, which favors the hypothesis of a concerted mechanism.
In a creative theoretical experiment, Maeda, Sakaki, and co-workers described the use of computationally designed ONO and NNN pincer PIII compounds for the activation of CO₂ (Scheme 23). The author’s calculations were found to be in line with literature findings, with ligand-PIII dehydration of H₃N·BH₃ being the RDS (19.7 kcal mol⁻¹), followed by CO₂ reduction. The latter proceeded via a concerted pathway when optimizations were performed with ONO-P pincer ligands. When calculations were focused on NNN-P pincer ligands, a stepwise coordination of formate to P-H followed by reduction to formic acid was most likely to be in action.

### 3.3. Supramolecular and Heterogeneous Examples

Reaction mechanisms for TH reactions with supramolecular and heterogeneous systems are somewhat less studied and the lack of mechanistic investigation does not allow discrimination between these systems as being either TH or standard hydrogenation reactions. Therefore, we will highlight in this section only the reactions where the mechanism has been proven to be a classical TH by amine–boranes.

Initial reports of supramolecular systems of amine–boranes to perform reduction reactions were published in 1984. Allwood and co-workers explored the formation of a supramolecular adduct formed between substituted chiral 18-crown-6-ethers and H₃N·BH₃; the adducts were isolated and characterized by X-ray diffraction, and were found to be active in the enantioselective reduction of ketones with selectivities up to 67% ee.

A rare example of heterogeneous TH was reported by Li and co-workers in 2015 (Scheme 24a). The authors built cobalt nanoparticles on graphitic carbon nitride dyad (Co/CN or Co/g-C₃N₄) on a mesoporous carbon nitride as the catalyst support which resulted in a hybrid structure of amorphous shells (Co²⁺) and metallic core (Co⁰) as confirmed by X-ray photoelectron spectroscopy (XPS). The Co/CN material was highly active in the TH of nitroarenes, olefins, ketones, and aldehydes with H₃N·BH₃ as the hydrogen transfer agent. The TH was efficiently performed at room temperature in less than 1 h. The reaction supported small-scale application with 20 mg of catalyst per 0.5 mmol of substrate, but could also be scaled up by a factor of 10. It is important to note that when the reaction was performed in an atmosphere of H₂ (1 bar), there was no conversion after 12 h. A potential Co²⁺/Co⁰ redox pair was postulated to be involved in the reaction mechanism (Scheme 24b); however, the formation of CoH₃ and/or Co–amidoborane intermediates could not be excluded.

Non-supported commercially available CuO was used for the TH of nitro compounds (Scheme 25). H₃N·BH₃ was the only reducing agent capable of performing the reaction, while NaBH₄, hydrazine, acetic acid, and H₂ showed limited conversion (<10%). The reaction was run in alcoholic solvents, with MeOH being optimum; when CD₃OD was used, no deuterium incorporation was found in the final product, which discounted the solvolysis of H₃N·BH₃ in the reaction mixture. Reduced intermediates, such as azoxybenzene and diazobenzene, which were found by ¹H NMR analysis, favored a stepwise TH mechanism.
In a detailed computational exploration, Ankan and co-workers described the potential fixation of $\text{N}_2$ onto tantalum atoms supported on a silica surface, and its further reduction using $\text{Pt}_3\text{HN-BH}_3$ (Scheme 26). The authors analyzed the reduction in the presence of $\text{Pt}_3\text{HN-BH}_3$ as the reducing agent, and noticed that the latter, in contrast to $\text{H}_2\text{N-BH}_3$, does not oligomerize once dehydrogenated; this could allow the aminoborane product $\text{Pt}_3\text{N}=\text{BH}_3$ to be rehydrogenated to the amine–borane equivalent. Concerning the fixation/reduction of nitrogen on tantalum, the author proposed the formation of a tantalum amid imido intermediate $[≡\text{SiO}_2\text{Ta}(=\text{NH})(\text{NH}_3)]$ (28) as the end-product of the reaction, as found by others when analyzing the fate of $\text{N}_2$ reduction with molecular $\text{H}_2$. $\text{N}_2$ could approach the supported Ta atoms and be activated through a relatively low activation barrier of 14.5 kcal mol$^{-1}$, resulting in elongation of the $\text{N}–\text{N}$ bond length (1.20 Å) compared to free $\text{N}_2$ (1.09 Å), indicating activation. Further stepwise proton and hydride transfer to the $\text{N}–\text{Ta}$ bond allowed formation of a diazenido species, which was further reduced to $[≡\text{SiO}_2\text{TaH}(\text{NHNH})]$ by hydride migration from Ta to N. A second equivalent of amine–borane could further activate species 28 and form $[≡\text{SiO}_2\text{TaH}_3(\text{NH}_3\text{NH})]$: the following second hydride migration from the Ta centre was predicted to be RDS with an activation energy $\Delta G^*=33.8$ kcal mol$^{-1}$ to form $[≡\text{SiO}_2\text{Ta}(=\text{NH})(\text{NH}_3)]$ (28). This event was found to be energetically favored at 13.5 kcal mol$^{-1}$ compared to other models using molecular $\text{H}_2$ as reductant with an activation barrier of 43 kcal mol$^{-1}$. Furthermore, the author predicted that the reaction could be implemented experimentally at low temperature (160–170°C) and suggested a way to circumvent catalyst decomposition via exposing the surface to $\text{N}_2$ at high pressure in order to maximize fixation and subsequently allow more facile hydrogenation to occur. However, even though the results looked promising, no further experimental evidence to disprove the authors’ findings have been reported yet; this is certainly an encouragement to expand on the topic.

Recently Jiang and co-workers synthesized a core–shell CuPd@ZIF-8 composite, with a cubic CuPd core and a MOF shell, and applied the system to the selective semi-TH of alkynes with $\text{H}_2\text{N-BH}_3$. This system is notable because of the synergistic behavior of the Cu and Pd centres, which allows selective absorption of $\text{H}_2\text{N-BH}_3$ ($–1.38$ eV on Cu versus $–1.49$ eV on Pd) and phenylacetylene ($–2.43$ on Pd versus $–0.61$ eV on Cu), respectively. The MOF shell protects the core, decreasing potential chemical etching of Cu nanocubes, even after five consecutive runs. Deuterium labeling experiments allowed assessment of the role of $\text{H}_2\text{N-BH}_3$ in the system; a first order rate dependence on the reductant was observed under catalytic conditions for the reduction of phenylacetylene, with a KIE of 4.08 using $\text{H}_2\text{N-BD}_3$, indicating that the B–H activation is rate determining. Incorporation of deuterium occurs also in the presence of $\text{D}_2\text{N-BH}_3$, while solvolysis was not in action with no deuterium incorporation into styrene when MeOD was used instead. The negligible capacity of $\text{H}_2$ to perform the reduction was also described and further DFT calculations on the catalytic system could define clearly that a classical TH reaction is in effect.

### 4. Solvolysis of Amine–Boranes in Nonclassical TH Reactions

In this section, we are concerned with the parallel/alternative route that can occur once $\text{H}_2\text{N-BH}_3$ dissociates into free $\text{NH}_3$ and the solvent adduct of $\text{BH}_3$—that is the reduction of an unsaturated bond by an initial hydroboration step followed by protic solvent work-up. This route should be categorized as nonclassical TH as the protic hydrogen is donated from the solvent and not the amine counterpart, but importantly, nor are the hydrogens due to $\text{H}_2$ released from the solvolysis of $\text{H}_2\text{N-BH}_3$. Similar to Section 3, the literature reviewed here, opens up some ambiguity into the precise mechanism operating. We have inferred a solvolysis pathway occurring where the authors themselves have not classified whether classical or nonclassical TH is undergoing in their systems.

Early examples of solvolysis of amine–boranes in reduction reactions have been reported by Jones using $\text{Me}_2\text{N-BH}_3$ to reduce 4-tert-butylicyclohexanone in benzene followed by aqueous work up. In 1971, Borsch and Levitan investigated the asymmetric reduction of ketones with optically active phenethylamine–borane with high conversion but very poor optical purity of the final product. These early examples, although they did not provide a great deal of mechanistic insight, showed the potential of amine–boranes as reducing agents in combination with aqueous work-up or action of solvolysis.
4.1. Uncatalyzed Solvolysis

Continuing on their previous work of uncatalyzed TH of polarized bonds (Section 2), and on metal catalyzed TH of olefin (Section 3.1) Berke and co-workers explored the reactivity of aldehydes and ketones with H$_2$N-BH$_3$ (Scheme 27). Rather than observing analogous TH reactions in THF they found only hydroboration of the C=O moiety to form borate esters. The presence of free NH$_3$ was also observed in situ by NMR spectroscopy ($\delta_{NH} = 0.4$ ppm), suggesting an alternative mechanism was operating in contrast to those previously reported in Section 2. When benzophenone was used as the model substrate, deuterium labeling experiments showed only deuterium incorporation at the carbon position of the C=O unit when H$_2$N-BD$_3$ or D$_2$N-BD$_3$ was used, confirming the „spectator“ role of NH$_3$ (e.g. D$_2$N-BH$_3$ led to no deuterium incorporation into the product). Furthermore, similar reactivities were observed with H$_3$B-THF as the hydrogen source. DKIE experiments showed normal DKIEs ($k_{D,N-BH}/k_{H,N-BH} = 1.74$ and $k_{D,N-BD}/k_{H,N-BD} = 1.10$) with D$_2$N-BH$_3$ and normal DKIEs ($k_{D,N,BD}/k_{H,N,BD} = 1.28$ and $k_{H,N,BD}/k_{D,N,BD} = 1.49$) with H$_2$N-BD$_3$. These experiments suggest the dissociation of H$_2$N-BH$_3$ is the RDS and the values are indicative of a secondary KIE due to changing geometry at the N and B atoms. The H$_3$B-THF species can then undergo standard hydroboration reactions with aldehydes and ketones to form borate esters.

Performing the reaction in MeOH, Berke found formation of the desired primary and secondary alcohols along with B(O Me)$_3$ and free NH$_3$ as the by-products. It was postulated that dissociation of H$_2$N-BH$_3$ would be the RDS to form free NH$_3$ as a spectator molecule and BH$_3$ as the reagent. BH$_3$ could immediately form an adduct with the C=O moiety of the substrate and hydroboration would form the borate ester intermediate which then undergoes methanolysis to give the products. Alternatively, a MeOH-BH$_3$ adduct could form after dissociation and undergo direct hydrogenation via a double H transfer with the protic hydrogen coming from the alcohol. Deuterium labeling experiments were unable to distinguish between these two pathways. However, using MeOD confirmed that the deuterium incorporation at the O atom of the carbonyl moiety was solely from the solvent and again indicating that this reaction does not undergo a classical TH process.

It is worth noting that in countering studies, Chen and co-workers published a DFT study on the reduction of benzaldehyde with H$_2$N-BH$_3$.[72] The authors first examined S$_0$-versus S$_2$-type processes for the dissociation of H$_2$N-BH$_3$, for a reaction involving THF, MeOH, and benzaldehyde. The S$_0$ pathway was the most favorable route to the common adduct, PhCHO-BH$_3$, with minor differences in the energies in THF ($\Delta G^* = 23.5$ kcal mol$^{-1}$) and MeOH ($\Delta G^* = 24.5$ kcal mol$^{-1}$). Furthermore, the RDS in all the S$_0$ routes was after the initial H$_2$N-BH$_3$ dissociation and involved a second dissociation of the BH$_3$-solvent adduct to form PhCHO-BH$_3$. Therefore, only the boron counterpart, and not NH$_3$, appears to be involved, which would contradict the observed normal DKIE effects with D$_2$N-BH$_3$ or D$_3$N-BD$_3$ reported by Berke and co-workers.[72]

From the PhCHO-BH$_3$ species, a hydroboration step (THF, $\Delta G^* = 31.8$ kcal mol$^{-1}$; MeOH, $\Delta G^* = 32.8$ kcal mol$^{-1}$) was found representing the RDS of the pathway (Scheme 30a). In comparison, the direct TH route from H$_2$N-BH$_3$ was found to be more kinetically favorable, with the RDS involving the concerted double H transfer ($\Delta G^* = 27.1$ kcal
The small difference of 4.7 kcal mol\(^{-1}\) (Scheme 29b). The small difference of 4.7 kcal mol\(^{-1}\) between the TH route and hydroboration may suggest that there is some interchangeability between the two pathways depending on the product-determining step, and potentially could explain the difference in reactivity observed by Berke's group and Chen's group for different aldehydes used in their investigations (Scheme 29).

### 4.2. Homogeneous Mediated Solvolysis

In 2016 Liu, Luo, and co-workers published the TH of alkynes to cis- and trans-alkenes selectively using PNP- and NNP-type Co-pincer complexes\(^{[39]}\). Controlling the steric profile around the cobalt center by altering the groups on the pincer ligands allowed them to access good chemo- and stereoselective transformation of numerous alkenes (Scheme 30). The role of H\(_3\)N·BH\(_3\) was seemingly just as the borohydride source rather than other conventional borohydrides (NaBH\(_{4}\), NaBH\(_4\)CN, Me\(_2\)S·BH\(_3\), NaBH\(_2\)(OAc), Me\(_2\)HN·BH\(_3\)); 4) The reaction in alcohols had the higher activity than that in THF or toluene, with MeOH chosen as the preferred solvent.

In order to identify the hydrogen source, deuterium labeling experiments were performed. When diphenylacetylene was used as the model substrate, CD\(_3\)OH showed no deuterium incorporation into the product, but CD\(_3\)OD allowed the isolation of the monodeuterated trans-1,2-diphenylethene. The reaction of H\(_3\)N·BH\(_3\) with 1 mol% under standard conditions with and without 1 equiv of diphenylacetylene always gave B(OMe)\(_3\), as the product with formation of H\(_3\)O. Without any catalyst or substrate, formation of B(OMe)\(_3\) in only 5% yield was observed after 16 h at 50°C, suggesting the methanolysis of H\(_3\)N·BH\(_3\) is a catalytic process in this system. The group also demonstrated that without H\(_3\)N·BH\(_3\) no product was observed, so MeOH alone could not act as the hydrogen source.

A plausible mechanism was proposed (Scheme 31) based on all the experimental evidence, suggesting the role of H\(_3\)N·BH\(_3\) was to generate the active [Co–H] species, which hydrometalates the alkyne across the triple bond to generate an alkenyl cobalt complex. Methanolysis of the Co–Cbond releases the cis-alkene product and forms a [Co–OMe] complex, observed by NMR spectroscopy. Regeneration of the active [Co–H] species is enabled by H\(_3\)N·BH\(_3\) and after 3 turnovers can give B(OMe)\(_3\) as the by-product. The authors also propose the competitive isomerization cycle to give the trans-alkene product from the common [Co–H] species. It is worth noting that the use of NaBH\(_4\) also gave successful results in the optimization reaction and would further corroborate the spectator role of the amine counterpart in H\(_3\)N·BH\(_3\). However, the lack of success when using NaBH\(_{4}\), a stronger hydride donor, to generate the [Co–H] species is somewhat surprising given the precedence\(^{[74]}\) and may allude to a more complex process or alternative process operating to form the active [Co–H] species. Using D\(_3\)N·BD\(_3\) may help to confirm the formation of a [Co–D] species, adding more weight to the mechanism. Furthermore, a control experiment under an atmosphere of H\(_2\) would be informative and allow further scrutiny whether hydrogenation from H\(_2\) participates in the mechanism.

**Scheme 29.** Simplified reaction profile of a) hydroboration of benzaldehyde and b) TH of benzaldehyde with H\(_3\)N·BH\(_3\).
The solvolysis of H₂N-BH₃ to effect reduction reviewed in this section can therefore be viewed as nonclassical TH reactions. The role of the H₂N-BH₃ is akin to that of borohydride reagents to reduce and activate the catalyst and to regenerate the active metal hydride complex during the cycle, with the NH₃ component not partaking in the active cycle. Instead, proton transfer is from the protic solvent, namely MeOH, and formation of B(OMe)₃ or H₃N·B(OMe)₃, is observed as the by-product. Deuterium labeling experiments, using alternative borohydride sources and using tertiary ammonia boranes (R₂N·BH₃, R ≠ H), are simple methods in the chemist’s toolbox that could be used to determine whether classical TH is taking place or whether hydride transfer and solvolysis is occurring instead.

What is interesting and less understood is the mechanism of activation of the precatalyst by H₂N-BH₃. These transition metal hydride species are often invoked based on the precedence of related hydrogenation reactions but not further scrutinized within these systems. Parallel DKIE experiments, kinetic experiments, and initial rates would have provided additional invaluable data towards understanding this pre-activation step. Inference from transition metal mediated dehydrogenation/dehydrocoupling of H₂N-BH₃ may be pertinent in this instance.⁷⁰⁻⁷², ¹⁵c, ⁷⁵

### 4.3. Heterogeneous Mediated Solvolysis

The literature around the reduction of unsaturated substrates by methanolysis of amine–boranes under heterogeneous conditions is scarce. This may be due in part to the dearth of mechanistic data available in order to determine whether the reactions are simply dehydrogenation of amine–boranes with molecular H₂ transferred to a surface to participate in subsequent hydrogenolysis. However, in 2001 Couturier and co-workers reported the methanolysis of primary, secondary, tertiary, and aromatic amine–boranes with Pd/C and Raney Ni at room temperature in MeOH.⁷⁶ The absence of protic hydrogens on the tertiary and aromatic amines would confirm that the hydrogen release is due to methanolysis of the amine borane. In follow-up studies they envisioned the reduction of nitroareyls using amine—boranes, provided the rate of reduction was faster than the rate of H₂ release.⁷⁷ Me₃N-BH₃ was chosen in this study. Reaction times for the reduction of nitroarens varied from 0.7–22 h for room temperature reactions (Scheme 32). A control reaction with Me₃N-BH₃ and 10 mol% Pd(OH)₂/C showed a reaction time of 20 h for complete methanolysis, which was determined by monitoring the amount of H₂ released. This provided good evidence that reduction was occurring faster than H₂ release.

\[
R\text{NO}_2 + \text{Me}_3\text{N}-\text{BH}_3 + 1.2 \text{ equiv } \text{Pd(OH)}_2/\text{C} \rightarrow \text{Me}_3\text{N}·\text{BH}_3 \rightarrow \text{R·NH}_2 \text{BH}_3 \rightarrow \text{R·NH}_2 + \text{H}_2 \text{BH}_3 \rightarrow \text{R·NH}_2 + \text{MeOH} + \text{H}_2
\]

Scheme 32. Reduction of nitroarens with Me₃N-BH₃ mediated by Pd(OH)₂/C.

In 2013, Stratakis and co-workers reported the reduction of nitroarens and nitroalkanes into amines and alkylhydroxylamines, respectively, using H₂N-BH₃ as the reductant and Au NPs supported on TiO₂ as the catalyst (Scheme 33).⁷⁸ Optimization reactions using p-nitrotoluene showed the reaction performed best in EtOH and H₂O as the solvent with less than 5% conversion observed with polar aprotic and nonpolar solvents. A control reaction without any Au NPs in EtOH showed no conversion to product. Stratakis et al. noted that the reaction was unlikely to involve H₂ gas as in related studies by Corma and co-workers, where high temperatures (100–140°C) and high pressures of H₂ (9–25 bar) were required to mediate the chemoselective reduction of nitroarens by the same catalyst system.⁷⁹ Instead, the authors suggested involvement of Au–H species without further scrutiny of the mechanism and they were unable to identify the fate of H₂N-BH₃ after the reaction. However, based on the solvent optimization reactions, the effect of EtOH indicates that solvolysis pathway might be in operation, but without further experimental evidence, this cannot be substantiated.

Following up on this study, Stratakis and co-workers expanded the scope of their reaction to report the stereoselective cis-semihydrogenation of alkynes to alkenes.
Solvent optimization of their system found again that aprotic polar solvents and nonpolar solvents resulted in poor conversion, with EtOH again showing the best results. Interestingly adding 5% v/v H₂O in THF improved reduction from 11 to > 99% conversion when compared to just using THF. Investigating the redundant source, they found using H₂N·BH₃, Me₃HN·BH₃, and MeH·N·BH₃ gave full conversion, but BuH·N·BH₃ and Me₂N·BH₃ resulted in 15% and no conversion, respectively. Furthermore, using H₂B·SMe₂ or HBpin also gave no product, which suggested the amine counterpart is important but also both the B–H and N–H are involved with the reduction. Reaction of 0.5 equiv of H₂N·BH₃ with deuterium-labeled p-methoxyphenylacetylene to afford the stereoselective cis-addition product indicated a potential concerted addition of the hydrogens. In further studies of the reaction mechanism, ¹¹B NMR spectroscopy provided information on the destination of the H₂N·BH₃ after the reaction. Analysis of the liquid phase using CD₃OD showed a peak at δ₂=9.0 ppm, which was assigned as NH₂B(OCD₃)₂; this was the only additional peak observed in the ¹¹B NMR spectra at the end of the reaction. Based on their experimental data, they proposed involvement of Au–H species generated from insertion of the borane moiety of the complex and proton from the solvent, to finally give the borate salt NH₂B(OR), as the by-product in the reaction. When the reaction was performed using CD₃OD or THF/MeOD and p-methoxyphenylacetylene as the substrate, there was 60–65% deuterium incorporation on both carbon atoms of the styrene moiety, corroborating with the proposed mechanism. This study represents involvement of both classical TH and nonclassical TH (solvolyis) processes at different stages of the reaction with the choice of starting amine–borane salient to the success of the reduction reactions. This further highlights the difficulty distinguishing the “true” mechanism in operation of these reactions with amine–boranes as the reductant, as the easy interchangeability of pathways that can be undertaken by the amine–borane based on reaction conditions could cloud the mechanism.

In 2017 Fu and co-workers reported the reduction of nitrile and nitro groups to primary amines using Ni₃P NPs with H₂N·BH₃ in a mixed ethanol/water solvent system (1/4, v/v) (Scheme 35a). Control reactions using H₂ (1 atm) as the hydrogen source showed no formation of product, suggesting that the dehydrogenation of H₂N·BH₃ is not operating in this system. The reaction mechanism was further probed by DFT calculations using 4-methoxybenzonitrile as the model substrate with Ni₃P NPs as the catalyst in water (Scheme 35b). The initial hydrolysis of H₂N·BH₃ mediated by Ni₃P NPs was previously reported by the group and was shown to be exothermic to give INT1. Subsequent steps involve the interaction between INT1 with another H₂O molecule and the substrate at the Ni₃P₂ surface. This orientation allowed the transfer of two H atoms from the H₂O molecule and the BH₃ moiety in INT1 to the C=N group, respectively, to form benzylicam. A second transfer of two H atoms from the BH₃(OH) moiety and another H₂O molecule to the C=N moiety represented the kinetic key step (3.17 eV) and resulted in the formation of benzylicamide as the product.

Very recently, Glorius and co-workers reported the TH of benzene derivatives and heteroarenes using H₂N·BH₃ mediated by [Rh(cod)(μ-Cl)]₂. Moderate to excellent yields were achieved along with good diastereomeric ratio for numerous substrates (Scheme 36). Optimization found that the reaction performed best in fluorinated alcohols, with TFE (2,2,2-trifluoroethanol) giving the best yield and d.r. values; reactions performed in hexane, THF and EtOH resulting in
no yield of product. The formation of a black suspension over
the course of the reaction indicated the involvement of
heterogeneous complexes, which was supported by Hg drop
test experiment resulting in no product formation. In
addition, Rh nanoparticles (60–100 nm), boron clusters, and
aluminum impurities were observed by SEM analysis of
the black suspension, supporting a heterogeneous mediated
catalysis. Probing the reaction further, the average deuterium
incorporation into the model substrate (tert-butylidimethyl(3-
tolyloxy)silane) using different deuterated H2(D)3BH(D)3
and TFE indicated that the protic hydrogen was from the
solvent and not the NH3 counterpart. Furthermore, the
reaction was also successful using Me2N-BH3 or HBpin as
the boron source. To test whether hydrogenation played a role
in the mechanism, the reaction using the model substrate was
performed under 1 bar H2 without any H2N-BH3 and gave no
conversion to the desired product even at 2 bar H2. Moreover,
letting [(Rh(cod)(µ-Cl))3] react with H2N-BH3 for ≈3 h then
adding the substrate under 1 bar H2 resulted in only 8% conversion,
suggesting that H2 is deleterious to the reaction. Cumulatively,
these experiments indicated a nonclassical TH mechanism in operation mediated by Rh nanoparticles.

5. Hydrogenation Reactions

To conclude, in this section we examine examples of
hydrogenation using amine–boranes. This route differs from
the classical and nonclassical TH reactions presented so far,
because the real reducing agent is the H2 released in situ.
When screening the literature, we observed a paucity of
hydrogenation reaction using amine–boranes in homogene-
cous catalytic systems (Section 3.1).[77] We rationalize this
finding, as differentiating whether a homogeneous system is
undergoing classical TH or hydrogenation is not trivial.
However, we cannot be certain that there have not been
examples of the use of alkene traps to monitor gas release in
investigations into the DHC of amine–boranes—most litera-
ture on this area of chemistry has monitored the direct
release of H2.[72] What has been reported is the use of
cyclohexene to trap H2N=BH3, with the formation of
Cy2BNH2 as the product but no mention of the formation of
cyclohexane.[84] Experimental control reactions can help elucidate
whether TH or hydrogenation is occurring. Primarily if no
reduction occurs in a homogeneous system when the reaction
is performed with H2 instead of amine–boranes—this indi-
cates classical TH. If reduction is observed but at a different
rate to that observed using amine–borane, then it would also
indicate a classical TH process. However, if reduction occurs
in the system with H2 at the same rate as that using amine-
boranes, then the identity of the mechanism is ambiguous and
computational insight could be helpful.

The key question in Section 3 is whether the amine–
borane’s role is specific to forming the active catalytic species
to mediate the reduction process as well as providing the
hydrogen source? The complication in answering this ques-
tion is that a common catalytic species is often associated with
both classical TH and hydrogenation pathways. However, if
the direct release of H2 from the amine–borane results in the
formation of the active species, then the role of amine–borane
is no different to just using H2 in the reaction and therefore we
classify this as standard hydrogenation.

In contrast, we find that a plethora of examples using
heterogeneous catalysts have been reported,[85] and we high-
light only those which present productive mechanistic studies
for the understanding of the reaction.

5.1. Heterogeneous Hydrogenation Reactions with Amine–
Boranes

A notable example of a hydrogenation reaction per-
formed with amine–boranes was reported by Manners and co-
workers, who analyzed the formation of catalytically active
Rh colloids when reacting [[Rh(cod)(µ-Cl)]3] with
H2N-BH3.[86] The new system was able to dehydrogenate
H2N-BH3 and sequentially hydrogenate cyclohexene with
molecular H2 in a closed vessel. When the reaction was
performed in an open vessel, no alkene reduction was
observed, clearly demonstrating the direct hydrogen addition
was taking place in this transformation. Further studies from
the same research group on heterogeneous hydrogenation
showed that the air-stable Rh/Al2O3 system in the presence
of Me2HN-BH3 could perform the reduction of alkenes without
external H2.[87] However, these reactions were still performed
in closed vessels and no further evidence of indirect hydrogen
transfer was furnished.

Nanoparticulate systems (NPs) based on different metals,
alloys, and sizes have been developed and tested in the
catalytic reduction of nitriles and nitroarenes, for example,
Pd@MIL-101, Pd NPs enclosed in an mesoporous MOF,[88]
and g-Cu20NiNP, CuNi NPs grafted on graphite.[89] When H2N-BH3
was simply replaced with H2, comparable yields could be
found, indicating a clear involvement of the gaseous source.
Moreover, tests conducted with open vessels gave lower
conversion (<20%) than experiments with higher pressur-
ized closed vessels, also highlighting that gas evolution and
solubility is paramount for the reduction to occur.

Xu and co-workers elegantly described a tandem dehy-
drogenation/hydrogenation of alkenes by H2N-BH3 using
Pickering emulsions,[90] which are emulsions stabilized by
solid particles instead of surfactants. The authors chose Pd
NPs coated onto g-C3N4 and carefully analyzed the efficiency
and behavior of these microreactors (Figure 1). Importantly,
when H2N-BH3 was replaced with gaseous H2, low reactivity
was observed mainly due to mass transfer effects from the gas
to the liquid phase. Moreover, limited emulsification could
decrease the reaction efficiency and stirring was found to be
beneficial to increase interface area (H₂ interaction with Pd NPs). These control reactions showed that the Pickering emulsions also function as transient H₂ storage materials, with potential chemisorbed gas onto Pd and/or formation of gas microbubbles. More importantly, these results highlight that simple interchange between H₂,N-BH₃ and H₂ might not be sufficient to define the nature of the reduction mechanism, while gas solubility and mass transfer effects need to be considered and tested.

Figure 1. a) Light microscopy image of Pickering emulsions; b) schematic representation of H₂,N-BH₃, DHC/styrene hydrogenation by the Pickering emulsions. Reproduced with permission.

6. Summary and Outlook

In this Review we have examined the use of amine–boranes as TH agents. We have carefully analyzed and classified the reduction reactions following three major mechanistic pathways: 1) classical TH, 2) nonclassical TH or solvolysis, and (3) hydrogenation with amine–boranes. In each of these contexts we have defined the role of the amine–borane species as the reducing agent and/or precatalyst activator. We have further highlighted the major types of amine–borane derivatives, such as H₂,N-BH₃, more elaborate amine–boranes, and MAB, and categorized these according to their role in reduction reactions.

We have examined the experimental and theoretical characterization techniques which have been used to allow such a mechanistic portrayal. The leading actor is a simple "H₂ test", which allows the initial classification of TH versus hydrogenation. However, this experiment is underrated, and it is apparent that this simple test is not used routinely. Moreover, it is crucial to study the role of the solvent to identify nonclassical TH, which is still ill-defined. The lack of differentiation between solvent-mediated protonolysis and amine-mediated protonolysis thereby places some reactions along the continuum between classical and solvolysis-mediated TH. The indisputable leading techniques are isotope labeling experiments and kinetic analysis for uncatalyzed and catalyzed TH reactions that can allow differentiation between stepwise versus concerted routes.

Each class of TH presented carries its own merits. There is not a single best route but a plethora of different options that can be selected in order to meet the user’s requirements. As described, classical TH reactions have proven to be an efficient method to access selectively deuterate substrates with relatively cheap and easy-to-handle amine–boranes. Solvolysis TH reactions further allow the introduction of benign and green solvents, whilst classical hydrogenation reactions using amine–boranes allow the use of a very precise quantity of an easy-to-handle „drop-in“ source of H₂ gas and are still the best way to reduce challenging substrates, for example, hindered alkenes. As stated, the area of amine–borane DHC is a buoyant one, but there are virtually no papers which clearly set out to undertake the type of consecutive dual catalysis that is necessary to dehydrogenate an amine–borane then use the H₂ released to reduce an unsaturated bond, particularly in the field of homogeneous catalysis. This presents a unique opportunity: With a well-defined catalyst that can undertake these dual roles and with full mechanistic understanding, researchers could be in a strong position to develop further divergent, asynchronous reactions.

We have highlighted that the stepwise transfer hydrogenation mechanism is kinetically and thermodynamically more favorable than dehydrogenation of amine–boranes when homogeneous systems are used. Vice versa, a predominant hydrogenation mechanism is in place when reduction reactions are performed with heterogeneous catalysts. It would be useful to be able to pinpoint why these differences exist: Does it depend on the rate of dehydrogenation versus the rate of stepwise B-H/N-H activation? Does this effect depend on the substrate affinity of the catalyst (e.g. better substrate affinity for styrene) versus amine–borane dehydrogenated side-products? In highlighting this, we hope to fuel discussion and research into this area.

As discussed, most TH substrates are azaones and unsaturated hydrocarbons; strikingly few examples of small-molecule activation have been reported. Rauchfuss et al. have undertaken a highly relevant study on the reduction of O₂ to H₂O₂, a key transformation in fuel cell research. Although not the focus of their study, the authors do report on the ability of H₂,N-BH₃, along with other hydrogen donors, to undertake the reaction in the presence of their Ir catalyst. Beyond this, it appears that the literature is very limited, with only one example of CO₂ reduction reported by Stephan (Section 3.2) along with a theoretical study of N₂ reduction on a Ta surface from Paul. It is evident that the TH of small molecules is an area of unmet need: can small molecules such as N₂, NO₂, and N₂O undergo activation and reduction using an amine borane in a laboratory setting and what is the mechanism of these processes? Can we expand on the chemistry of TH of CO₂ and O₂, develop new catalysts, and obtain deeper mechanistic understanding?

Another area that is underdeveloped is that of enzymatic TH. Many of the examples reported are deraezimations in the presence of H₂,N-BH₃ and an oxidase. However, it could be argued that the substrates presented are more challenging, or at least more complex, than those tackled using transition metal catalysis. Although mechanistic elucidation is likely to be demanding, a certain level of detail is needed to clearly understand the precise nature of the bond-breaking and bond-making process in biological media; the latter aspect is a highly attractive feature of enzymatic chemistry, harmonizing with the benign nature of a TH agent such as H₂,N-BH₃.

Finally, although we have covered several different amine–boranes and their role in TH, great opportunities...
must exist beyond these standard hydrogen sources. Indeed, elegant studies into the hydrogen-release properties of amino complex borane encapsulated in metal organic frameworks (MOFs),[90] ethyldiaminoboranes (EDABs),[91] and metal ethyldiaminoboranes (MEDABs)[92] and theoretical studies into boron nitride nanotubes (BNNTs)[93] have been undertaken, but detailed synthetic investigations into their ability to reduce classes of substrate are lacking. Although the TH reagents may not be suitable over a broad spectrum of substrates, they do present an opportunity to probe the extent of reactivity and functional group tolerance, which may allow key or unique targets to be met. As an example, the structures of these unusual amine boranes may lend themselves well to selective reduction of targeted sets of double bonds in a multiply double-bonded system, for example, in terpenoid feedstocks.

In conclusion, TH presents great opportunities in mechanistic studies, organic synthesis, and catalyst design. We have identified a plethora of variables associated with TH reactions from: 1) complex substrates through to traditionally inert small molecules, 2) catalyst design ranging from homogeneous, heterogeneous, enzymatic, or even catalyst-free transformations, 3) the variability of the TH reagent itself, with many possible reagents still to undergo comprehensive testing and opportunities for regio- and stereoselectivity, and 4) the nuanced means by which TH takes place that fall into three broad categories. All factors combined indicate that amine–borane-mediated TH is an area set for growth and likely to be a rich topic of research for years to come.

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Conflict of interest

The authors declare no conflict of interest.

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