

**Review** Article

Chemical Review and Letters journal homepage: www.chemrevlett.com ISSN (online): 2645-4947 (print) 2676-7279



# A Short Chronological Review on the Syntheses of Amine-Boranes

Daniel O. Reddy<sup>a,</sup>\*

<sup>a</sup>Department of Chemistry, Purdue University, West Lafayette, IN USA

#### ARTICLE INFO

ABSTRACT

Article history: Received 13 July 2020 Received in revised form 27 August 2020 Accepted 31 August 2020 Available online 31 August 2020

*Keywords:* Amine-boranes Borane complexes Synthetic history Since their discovery by Burg and Schlesinger in 1937, amine-boranes have enjoyed a rich preparative history and have experienced reinvigorated interest as valuable compounds. These borane complexes have been implemented in a variety of applications, spanning from reagents in organic syntheses to hydrogen storage materials. The importance of amine-boranes derives especially from their reductive abilities. Given the utility of amine-boranes and their current resurgence, a mini review on their general properties and notable preparations is both timely and potentially of interest to organic and materials chemists alike.

## 1. Introduction

Formally, amine-boranes can be described as complexes of borane, which acts as a Lewis acid, and an amine, which acts as a Lewis base. In this complex, the nitrogen atom's lone pair of electrons forms a coordinate covalent, also known as a dative, bond via donation into the vacant borane 2*p* orbital (Figure 1). [1] Such borane adducts can typically be formed with any Lewis base (molecules containing nitrogen, oxygen, phosphorous, or sulfur, wherein the hetero-atom possesses an available pair of non-bonding electrons). Common, simplistic examples of these classical adducts include ammoniaborane (NH<sub>3</sub>-BH<sub>3</sub>, AB), borane-tetrahydrofuran (BH<sub>3</sub>-THF, BTHF), phosphine-borane (PH<sub>3</sub>-BH<sub>3</sub>, PB), boranedimethylsulfide [BH<sub>3</sub>-S(CH<sub>3</sub>)<sub>2</sub>, BDMS, DMSB], etc.



Figure 1. sp $3\sigma$ -p $\sigma$  dative bond formation in an amine-borane The discovery of a boron-nitrogen dative bond is

attributable to Gay-Lussac in 1809 [2], and in 1937, Burg and Schlesinger were credited with the first report of an amine-borane, trimethylamine-borane (Figure 2). [3] With regards to Gay-Lussac, Burg, and Schlesinger, their initial discoveries have ushered more than three-quarters of a century (and beyond) of research into amine-boranes and their derivatives, with diverse applications ranging from reagents in organic syntheses [4] to hydrogen 

The first historical application of amine-boranes is taken to be as reducing agents, as noted in Hutchins et al.'s seminal 1984 review. [7] A full treatment of the numerous applications of amine-boranes and their derivatives, as well as their relatives like phosphine- (or phosphane-) and sulfide-boranes, is not within the scope of this introduction, though such literature serves as the basis for several fine investigations [6c, 8] and reviews (Figure 3 [9]). [1, 2b, 7, 10].



**Figure 3.** Amine-borane applications (Adapted from Kulkarni & Ramachandran, 2017)

Amine-boranes are perhaps most commonly recognized as valuable reagents due to the hydridic nature of the hydrogen atoms on the borane functionality; it is this hydridicity that enables the diverse and unique chemistry of amine-boranes (To illustrate this principle, some older representations of amine-boranes, historically called amine-"borines" or "-borazanes," denote positive and negative charges on the N- and B-atoms, respectively, while maintaining the overall neutrality of the complex) (Figure 4). [2b, 10d, 11] It is evident that AB is the structurally simplest of the amine-boranes, regarding the substitution on the N-atom.



Figure 4. General hydridicity of ammonia-borane (left) & historical representation, intentionally antiquated [10d] (right) By extension, the generally broad classifications of amine-boranes include: ammonia-borane (AB), primary  $(1^{\circ})$ , secondary  $(2^{\circ})$ , tertiary  $(3^{\circ})$ , and heteroaryl (HA) amine-boranes, wherein the N-atom is involved in the aromaticity of the ring; these classifications describe the amount of non-hydrogen substitution on the N-atom. Within the  $2^{\circ}$  and  $3^{\circ}$  classifications, there exists the subset of heterocyclic (HC) amine-boranes, e.g., aziridine-borane, wherein the N-atom is part of the closed ring. Though not as prominent, there are also aminewith hydrazine-bisborane bisboranes (Bis), [12]  $(H_3BNH_2NH_2BH_3)$ and ethylenediamine-bisborane [H<sub>3</sub>BNH<sub>2</sub>(CH<sub>2</sub>)<sub>2</sub>NH<sub>2</sub>BH<sub>3</sub>] as the most recognized examples; there appears to be scant reporting of aminetris-, tetrakis-, or pentakis-boranes, especially no higher than five -BH<sub>3</sub> groups. [13] Interestingly, a recent (2019) patent describes the preparation of a trisborane complex [N,N-bis[2-(dicyclohexylphosphino)ethylamine-

trisborane], wherein two of the three -BH<sub>3</sub> groups are coordinated to P-atoms and the third -BH<sub>3</sub> group is coordinated to an N-atom, representing a hybrid (Hyb) amine/phosphine-borane. [14] Moreover, there exists a subset within each of the aforementioned categories, barring AB, that includes functionalized (Func) amineboranes, for instance methan*ol*amine-borane, 2diethylaminoethanethiol-borane, or allylamine-borane. In some instances, these accompanying functional groups are sensitive to boranation, i.e., installment of the -BH<sub>3</sub> group, so care must be taken when choosing a preparative method, as will later be discussed. In terms of the hydridicity of an amine-borane, this particular property is dependent upon both the N-atom's and the B-atom's substituent(s), though in the case of non-substitution on the B-atom (BH<sub>3</sub>), hydride-donating tunability lies with the N-atom. More generally, the overall stability of an amine-borane is contingent on the substitution(s) of the N- and B-atoms. As an extreme example, aniline-borane is stable at -30 °C, the minimum temperature required for its preparation, but can begin to dehydrocouple as ambient temperatures are approached, assuming standard pressure. [2b, 10d, 15] The most notorious factor influencing amine-borane stability is the steric bulk of the group(s) on both the B- and N-atoms. The steric bulk phenomenon has been quantitatively demonstrated by assessing the molar enthalpies of formation  $(\Delta H_f)$ between certain amines and BTHF; the reported  $\Delta H_f$ trends are: 1)  $nBuNH_2-BH_3 > nBu_2NH-BH_3 > nBu_3N-$ BH<sub>3</sub> and 2) Et<sub>2</sub>NH-BH<sub>3</sub> >  $nPr_2NH$ -BH<sub>3</sub> >  $nBu_2NH$ -BH<sub>3</sub>. [2b, 16] Understandably, the resultant instability from

increasingly bulky groups on the B- and/or N-atoms is due to poor orbital overlap (Figure 1). There have also been suggestions that increased alkyl bulk on the N-atom entails reduced Lewis basicity by increasing the bond angle around the N-atom, thereby reducing the pcharacter of the interacting lone-pair and increasing its s character. [2b, 17] As a general rule of thumb, Hutchins et al. posit that a stable adduct can be formed so long as the p $K_a$  of the amine is greater than 5.0 – 5.5. [7] It is also noted that those amines whose  $pK_a$  values lie within that range can form the dative bond with borane but that this N-B bond is weaker compared to those amines with  $pK_a$ values greater than the specified range. Invariably, some of the same principles governing the stability of an amine-borane adduct also apply to the hydride-donating capability of the amine-borane; it is understood that the amine-borane adduct must itself be stable in order to react. Most prominently, when describing amine-boranes simply bearing the -BH<sub>3</sub> moiety, the identity of the amine governs reductive capabilities of the hydrides. For example, it has been shown that increasing alkyl substitution on the amine decreases reducing ability, illustrated by the trend  $AB > RNH_2-BH_3 > R_2NH-BH_3 >$ R<sub>3</sub>N-BH<sub>3</sub>. [2b, 7] Regarding the sister compounds, heteroaryl- and N-arylamine-boranes, the trend is that those amines with lower  $pK_a$  values are better reducing agents. In an application-related example, several different amine-boranes were tested in their reductive abilities towards a gold salt, AuPPh<sub>3</sub>Cl, for the preparation of nanoparticles, and their capacities to reduce the salt were: AB > t-butylamine-borane  $\approx$ triethylamine-borane > morpholine-borane, illustrating the general effect of the N-atom's substituents on hydridicity. [10b, 18] Another interesting facet of hydride-donating governability is the report that acidic, aqueous/mixed aqueous solvents can enhance the reducing ability of some amine-boranes; this has been demonstrated using morpholine-borane and several of its derivatives and relatives. [2b, 10h, 19] In a specific example, without acidification, N,N,N-trimethylamineborane is unable to reduce cyclohexanone to the corresponding alcohol within 38 hours reaction time. With acidification, the ketone-alcohol transformation is affected within 8 minutes with 80% conversion. It has also been noted that solvent selection and increased temperatures, up to a certain extent (~70 < T < 100 °C), can improve amine-borane hydridicity.<sup>7</sup> Finally, the addition of a Lewis acid catalyst, such as AlCl<sub>3</sub> or BF<sub>3</sub>, markedly improves the reductive effect of certain amineboranes due to a complexing action of the Lewis acid with the carbonyl oxygen, which in turn facilitates the intermolecular hydride transfer from the amine-borane to the  $\delta^+$  carbon. [7, 20] Within the framework of organic chemistry, amine-boranes are most prominently viewed as reducing agents; Hutchins et al. regard amine-boranes as essential "in the arsenal of reductive weapons available to chemists." [7] Though not definitive, it is within reason

to suggest that of the available reducing agents, including amine-boranes, sodium borohydride (NaBH<sub>4</sub>), also known as sodium tetrahydridoborate, is the most widely used and commonly recognized. [21] In fact, it has been boldly asserted that "every beginning organic text mentions the use of sodium borohydride as a reducing agent." [22] Moreover, Ullmann's Encyclopedia of Industrial Chemistry indicates that thousands of metric tons of NaBH<sub>4</sub> are annually manufactured and utilized worldwide, representing a multimillion dollar business; such usage warrants the recognition of NaBH<sub>4</sub> as "by far the most important commercially available complex hydride." [21b, 23] Other popular, often-employed reducing agents include: lithium aluminum hydride (LAH), [24] lithium borohydride (LBH), [25] lithium triethylborohydride [Superhydride®, LiTEBH], [26] sodium cyanoborohydride (NaBH<sub>3</sub>CN), [27] lithium aminoborohydrides (LABs), [28] diisobutylaluminum hvdride (DIBAL, DIBAL-H), [29] sodium-bis(2methoxyethoxy)aluminum hydride (SMEAH, Red-Al®), [30] and triethylsilane (TES). [31] A computational study ranks AB, a somewhat representative amine-borane, as roughly in between TES, a weak reducing agent, and BH4<sup>-</sup> , an intermediate reducing agent, in terms of reducing strength, as measured by  $\Delta G$  values. [21c]

Given such an abundance of reducing agents, one might be prompted to consider how amine-boranes are set apart from their counterparts. With reference to the previous discussion, reducing strength is certainly the most obvious distinction that can be made between amine-boranes and other reducing agents. However, tunability of amine-borane reducing strength is made possible by 1) manipulating the N-atom's substituent(s), 2) acidifying the solution, 3) varying the solvent, 4) adding a Lewis acid catalyst, and/or 5) adjusting the reaction temperature. It is worthwhile however to note that Brown and coworkers experimented with the tunability of sodium borohydride's (SBH) reducing strength by changing the cation and substituting the H atom(s) with alkyl or alkoxy groups. [32] Though Heiden and Latham cite AB as a representative amine-borane in their study on "[e]stablishing hydride donor abilities," [21c] it is difficult to make a generalization about the hydridicity of amine-boranes due to the vastness of this class of compounds. Aside from customizable reducing abilities, amine-boranes are oftentimes far more soluble than their fellow hydride donors in common solvents, both aprotic and protic; some of these solvents include: benzene, dichloromethane, ether, hexane, methanol, tetrahydrofuran, and toluene. Notably, amine-boranes are mostly unreactive towards water and other protic solvents. [2b, 7, 10h] Finally, amine-boranes offer tantalizing advantages, especially over certain reducing agents like LAH, LBH, and Superhydride® as well as boranating agents like diborane (B<sub>2</sub>H<sub>6</sub>, DB), BTHF, and BDMS, in terms of air- and moisture-sensitivity, pyrophoricity, and toxicity. [2b, 33] The current

discussion would be remiss if one of the more exotic applications of amine-boranes was not discussed prior to their synthetic history. By complexing the -BH<sub>3</sub> moiety with an asymmetric amine  $[H_{(3-n)}R_nN]$ , such that  $n = \{2,$ 3} and  $R_1 \neq R_2 \neq R_3$ ], it is possible to impart stereogenicity to the resultant amine-borane and perform an enantioselective reduction (Figure 1). [10d, 34] Similarly, chiral molecules possessing an amine functional group can be boranated and converted to chiral reducing agents. [7, 35] Between those two categories of chiral amine-borane reducing agents, it could be argued that the most infamous example is the Corey-Itsuno, or Corey-Bakshi-Shibata (CBS), reduction. The reaction development was first begun by Itsuno and coworkers in 1981 by reducing prochiral aromatic ketones with chiral alkoxy-amine-borane complexes. [36] Itsuno et al.'s work was extended by Corey, Bakshi, and Shibata in 1987, whereby Corey et al. demonstrated excellent enantioselectivity for the reduction of ketones using chiral oxazaborolidines (Figure 5). [37]



**Figure 5.** Itsuno's alkoxy-amine-borane (left) & Corey's oxazaborolidine-borane (right)

#### 2. Chronological Syntheses of Amine-Boranes

Having provided a cursory overview of amineboranes and their properties, a chronological history of their most notable preparations can be presented. The emphasis of this timeline is to denote the first report of a particular synthetic style; once these syntheses are initially reported, there are seemingly innumerable accounts wherein most of the methods are employed and occasionally developed further. For clarification, the aim of this chronology is to establish a history of synthetic methods for amine-boranes, of which ammonia-borane is both a prime and unique example. Though some of the following synthetic methods are applicable to the synthesis of AB, not all of the methods can accommodate this parent amine-borane. Likewise, there are synthetic methods for attaining AB that are not amenable to other amine-boranes. As such, the current discussion will not emphasize those preparative methods that are exclusive to AB; there are several reviews that seek to address the vast preparative methods for AB alone. [10j, 33b, 38]

As mentioned earlier, the first report of an amineborane, trimethylamine-"borine," was contributed by Burg and Schlesinger in 1937 (Scheme 1). [3] Brown later described the diborane-based process for preparing "borine trialkyl amines" in 1958. [39]

 $3 \text{ N}(\text{CH}_3)_3 + \text{B}_2\text{H}_6 \longrightarrow 2 \text{ N}(\text{CH}_3)_3 \cdot \text{BH}_3 + \text{N}(\text{CH}_3)_3$ Scheme 1. Synthesis of trimethylamine-borine via diborane & free amine

In 1942, Brown, Schlesinger, and Cardon used the diborane protocol to develop a transamination method for

synthesizing amine-boranes, wherein the free amine displaces the amine that is already complexed with borane (Scheme 2). [40] The transamination method was later expounded upon by Baldwin and Washburn in 1961, [41] as well as several others, including the Ramachandran group. [42]

 $C_5H_5N\bullet BH_3 + (CH_3)_3N \xrightarrow{} (CH_3)_3N\bullet BH_3 + C_5H_5N$ Scheme 2. Synthesis of trimethylamine-borane via transamination

More than a decade passed before another novel synthetic procedure for amine-boranes was presented, when Schaeffer and Anderson prepared trimethylamine-borine via LBH in 1949 (Scheme 3). [43]

 $(CH_3)_3NHCl + LiBH_4 \xrightarrow{Et_2O} N(CH_3)_3 \cdot BH_3 + H_2 + LiCl$ Scheme 3. Synthesis of trimethylamine-borine via LBH & amine•hydrochloride

In 1952, Banus, Gibb, Jr., and Bragdon posited a thermaldecomposition method for making amine-boranes (Scheme 4). [44] The pyrolysis method was further explored by Safronov, Jalisatgi, and Hawthorne in their 2019 patent investigating the decomposition of organoammonium tetrahydroborates. [45]

 $(CH_3)_4NBH_4 \xrightarrow{\Delta} (CH_3)_3NHBH_3 + CH_4$ Scheme 4. Synthesis of trimethylamine-borine via tetramethylammonium borohydride

In 1953, Schechter presented a unique synthesis of amine-boranes via electrolysis between an anode and cathode of an ionic borohydride in a non-aqueous solvent like an amine (Scheme 5). [46]

 $NaBH_4 + NH(CH_3)_2 \longrightarrow (CH_3)_2HN \cdot BH_3 + H_2 + Na$ Scheme 5. Synthesis of dimethylamine-borane via electrolysis with Hg cathode and Pt anode

In a similar fashion to Schaeffer and Anderson's LBH preparative method, Taylor, Grant, and Sands synthesized pyridine-borane using SBH with the amine as the solvent in 1955 (Scheme 6). [47] It is likely that previous authors did not report a synthesis of amineboranes via SBH due to difficulties associated with SBH's preparation, though a suitable preparation had in fact been known for some time. [48] However, due to World War II national security concerns, Schlesinger, Brown, and Finholt were unable to publish their findings. [21b, 21e, 32b, 49] The chemical and patent literature is rife with adjustments and alleged improvements to this particular process. [50]

 $C_5H_5N\bullet HCl + NaBH_4 \longrightarrow C_5H_5N\bullet BH_3 + H_2 + NaCl$ Scheme 6. Synthesis of pyridine-borane via SBH & amine•hydrochloride

Shortly after Taylor, Grant, and Sands's synthesis of pyridine-borane, several patents and publications from both academia and the chemical industry began describing the preparation of amine-boranes. First among these descriptions was from Köster in February 1957, wherein he developed a route to amine-boranes using high pressure hydrogenolysis of trialkylboranes (Scheme 7). [51]

 $B(C_2H_5)_3 + N(C_2H_5)_3 + 3H_2 \longrightarrow N(C_2H_5)_3 \cdot BH_3 + 3C_2H_6$ Scheme 7. Synthesis of triethylamine-borane via TEB & free amine

Soon after Köster's work, "Preparation of Amine-Borines" was proposed by Jenkner in March 1957, wherein he prepared various amine-boranes with an emphasis on the addition of boron trichloride to the reaction mixture (Scheme 8); [52] other ensuing patents have extended the scope of Jenkner's work. [53]

 $BCl_3 + N(CH_3)_3 + 3 \text{ NaH} \xrightarrow{\text{Activator}} N(CH_3)_3 \cdot BH_3 + 3 \text{ NaCl}$ 

Scheme 8. Synthesis of trimethylamine-borine via BCl<sub>3</sub>/NaH & free amine

Concluding 1957 as a popular year for amine-borane related patents, Bragdon described a three-step, aminecarbamate-based preparation in August 1957 (Scheme 9). [54]

 $CO_2 + 2C_3H_7NH_2 \longrightarrow C_3H_7NHCOOC_3H_7NH_3$  (1)

 $NaBH_4 + C_3H_7NHCOOC_3H_7NH_3 \longrightarrow C_3H_7NHCOONa + C_3H_7NH_3BH_4 (2)$ 

$$C_{3}H_{7}NH_{3}BH_{4} \longrightarrow C_{3}H_{7}NH_{2}BH_{3} + H_{2} (3)$$

Scheme 9. Synthesis of isopropylamine-borane via SBH & an amine-carbamate

Following Jenkner's patent, as a corporate entity, Farbenfabriken Bayer Aktiengesellschaft formulated a British patent for an amine-borane synthetic process in May 1959, wherein the amine-borane is obtained from a tertiary amine salt, an aqueous solution of metal borohydride, and an inert solvent (Scheme 10). [33b, 55] However, Haberland and Stroh filed an identical U.S. patent earlier in 1958 for the same process. [56] Notably, many other patents have followed suit with "similar yet different" acid-mediated proposals. [57]

 $(CH_3CH_2)_3N + KBH_4 + B(OH)_3 + H_2O \longrightarrow (CH_3CH_2)_3N \cdot BH_3 + KB(OH)_4 + H_2$ Scheme 10. Synthesis of triethylamine-borazane via KBH/boric acid & free amine

Later that same year in July 1959, Lang and Schubert patented an amine-borane synthesis using a metal borohydride and a boron trihalide or boron trihalide etherate (Scheme 11). [58] In 1964, Snover extended Lang and Schubert's work through his patent regarding in-situ preparation of diborane, subsequent reaction of diborane with a free amine, and isolation of the resultant amine-borane by crystallization from water. [59]

 $4 (CH_3)_3N + 4 BF_3 \bullet O(C_2H_5)_2 + 3 NaBH_4 \longrightarrow 4 (CH_3)_3N \bullet BH_3 + 4 (C_2H_5)O + 3 NaBF_4$ Scheme 11. Synthesis of trimethylamine-borane via boron trifluoride-diethyletherate & SBH

Following Lang and Schubert's work, Ashby in August 1959 defined a synthetic process for amine-boranes characterized by reacting a fully-esterified aryl ester of an oxyacid of boron with a metal borohydride (Scheme 12). [60]

 $C_{18}H_{15}BO_3 + NaBH_4 + C_6H_5N(CH_3)_2 \xrightarrow{\text{cat. AlCl}_3} C_6H_5N(CH_3)_2 \bullet BH_3 + NaB(OC_6H_5)_4$ Scheme 12. Synthesis of dimethyl aniline borane via phenyl borate/SBH & free amine

In February 1960, arguably the most commonly-used method for obtaining amine-boranes was established by Kelly and Edwards, wherein BTHF was used for

boranation (Scheme 13). [61] There seems to be agreement that this is the first reported use of BTHF as a boranating agent for amines; [33b] this assertion also agrees with the timeline of chemical history since THF was not commercially available until 1956, [62] so applications involving BTHF as a -BH<sub>3</sub> carrier would understandably follow shortly thereafter.

 $(CH_2NH_2)_2 + 2 C_4H_8O:BH_3 \longrightarrow (CH_2NH_2BH_3)_2 + 2 C_4H_8O$ Scheme 13. Synthesis of ethane 1,2-diamineborane via BTHF After Kelly and Edwards' notable procedure, Marshall prepared amine-boranes via a dialkoxyborane and the free amine in March 1960 (Scheme 14). [63]

3 BH(OCH<sub>3</sub>)<sub>2</sub> + C<sub>5</sub>H<sub>5</sub>N $\xrightarrow{B(OCH_3)_3}$  C<sub>5</sub>H<sub>5</sub>N•BH<sub>3</sub> + B(OCH<sub>3</sub>)<sub>3</sub> Scheme 14. Synthesis of pyridine-borane via dimethoxyborane & free amine

In 1962, Ashby and Foster proposed "A New and Convenient Route to the Amine-Boranes" (Scheme 15). [64] Their protocol uses a borate ester with an amine solvent to obtain the amine-borane.

 $B(OC_6H_5)_3 + Al + 3/2 H_2 \xrightarrow{N(CH_2CH_3)_3} N(CH_2CH_3)_3 BH_3 + Al(OC_6H_5)_3$ Scheme 15. Synthesis of triethylamine-borane via borate ester & free amine as solvent

Following Ashby and Foster's synthesis, Matsumara and Tokura in 1968 published a "superior" preparation using liquid  $SO_2$ , with the superiority purportedly owing to the weakly acidic nature of  $SO_2$  relative to other previously used acids. [65] However, the authors fail to fully characterize the reaction, referring to the products as the desired amine-borane and an unidentified "inorganic product," presumed to be sodium sulfoxylate (Scheme 16).

 $(CH_3CH_2)_3N + NaBH_4 \xrightarrow{liq. SO_2} (CH_3CH_2)_3N \cdot BH_3 + inorganic product Scheme 16. Synthesis of triethylamine-borane via liquid SO_2/SBH & free amine$ 

Later that decade in 1969, Nainan and Ryschkewitsch put forth an amine-borane synthetic procedure involving iodine for the generation of diborane from SBH (Scheme 17). [66]

 $2 \text{ NaBH}_4 + 2 (\text{CH}_3)_2\text{NH} + \text{I}_2 \longrightarrow 2 \text{ NaI} + 2 (\text{CH}_3)_2\text{NH} \cdot \text{BH}_3 + \text{H}_2$ Scheme 17. Synthesis of dimethylamine-borane via SBH/iodine & free amine

Rivalling BTHF as another popular agent for boranation of amines, BDMS appears to have first been used in a similar manner by Burke and Hough in 1976 as part of their patent, "Water Soluble Tertiary Amine Boranes (Scheme 18)." [33b, 67]

In 1984, a Czech patent was prepared by Plesek, Stibr, Drdakova, and Jelinek that describes a method similar to Matsumara and Tokura's method, wherein  $CO_2$  acts as a Lewis acid for the amine-borane formation (Scheme 19). [68] In 1991, Arduengo developed a method similar to the Czech patent, though Arduengo sought to avoid the water-washing step. [33b, 69] Later authors (Cao et al., 2012) suggest that the inorganic product is Na<sub>2</sub>CO<sub>3</sub> as opposed to NaHCO<sub>2</sub>, and that H<sub>2(g)</sub> is in fact a product of the reaction. [70]

 $N_2(C_2H_4)_3 + NaBH_4 + CO_2 \xrightarrow{CH_3CN} N_2(C_2H_4)_3 \cdot BH_3 + NaHCO_2$ Scheme 19. Synthesis of 1,4-diazabicyclo[2.2.2]octanemonoborane via SBH/CO<sub>2</sub>

As a unique example, Kampel and Warshawsky in 1994 published an 18-crown-6-mediated synthesis of various amine-boranes in ether (Scheme 20). [71]

 $(CH_3)_3N$ •HCl + NaBH<sub>4</sub>–18-crown-6 N(CH<sub>3</sub>)<sub>3</sub>•BH<sub>3</sub> + H<sub>2</sub> + NaCl Scheme 20. Synthesis of trimethylamine-borane via 18-crown-6 & amine•hydrochloride

Following Kampel and Warshawsky's work, in 1999, Collins, Lanz, Goralski, and Singaram formulated a lithium *N*,*N*-dialkylaminoborohydride-mediated route towards amine-boranes (Scheme 21). [72]

NC(C<sub>6</sub>H<sub>4</sub>)CH<sub>2</sub>Br  $\xrightarrow{\text{LiH}_3\text{BN}(\text{CH}_3)_2}$  NC(C<sub>6</sub>H<sub>4</sub>)CH<sub>2</sub>N(CH<sub>3</sub>)<sub>2</sub>:BH<sub>3</sub> Scheme 21. Synthesis of *N*,*N*-dimethylcyanobenzylamine-borane via lithium aminoborohydride

There are also several examples of amine-borane synthetic procedures developed in the Brown Center for Borane Research. The first of these described by Ramachandran and Gagare in 2007 focused exclusively on high-purity (>98%) AB synthesis (Scheme 22) by fine-tuning a metathesis procedure developed by Geanangel in 1977. [38, 73] The concentration of the reaction medium was critical in obtaining high-purity AB. This method was further modified by the same group for a large-scale preparation of AB involving ammonia as an additive. [74] Ramachandran and Kulkarni then described that the role of ammonia is that of a reagent (Scheme 22). [75] Water can also be used as a promoting additive, as described by them in 2017 (Scheme 23). [76]

 $2 \text{ NaBH}_4 + (\text{NH}_4)_2 \text{SO}_4 \xrightarrow{5\% \text{ NH}_3} 2 \text{ NH}_3 \cdot \text{BH}_3 + \text{Na}_2 \text{SO}_4 + \text{H}_2$ Scheme 22. Ammonia-promoted synthesis of ammonia borane

 $2 \text{ NaBH}_4 + (\text{NH}_4)_2 \text{SO}_4 \xrightarrow{\text{H}_2\text{O}} 2 \text{ NH}_3 \cdot \text{BH}_3 + \text{Na}_2 \text{SO}_4 + \text{H}_2$ Scheme 23. Water-promoted synthesis of ammonia-borane

In 2012, Ramachandran, Raju, and Gagare proposed a one-pot preparation of AB and several trialkylamineboranes from trimethyl borate, LiH, and AlCl<sub>3</sub> in the presence of the desired amine (Scheme 24). [77]

 $B(OCH_3)_3 + LiH + N(CH_2CH_3)_3 \xrightarrow{AlCl_3} (CH_3CH_2)_3N^{\bullet}BH_3 + LiAl(OMe)_4 + LiCl$ Scheme 24. Synthesis of triethylamine-borane via  $B(OMe)_3/LiH/AlCl_3$  and free amine In 2015, the salt metathesis method was expanded to include other amine-boranes aside from AB (Scheme 25). [78]

 $NaBH_4 + (NH_4)_2SO_4 + 2 (CH_3CH_2)_3N \longrightarrow (CH_3CH_2)_3N \cdot BH_3 + 2 NH_3 + Na_2SO_4 + H_2$ Scheme 25. Synthesis of triethylamine-borane via amine-ammonium salt equilibration

Lastly, in 2016, a bicarbonate-mediated process was optimized and developed that accommodates amines of varying degrees as well as functionalized amines (Scheme 1.24). [76]

NaBH<sub>4</sub> + NaHCO<sub>3</sub> + (CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>)H<sub>2</sub>N + H<sub>2</sub>O $\longrightarrow$  (CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>)H<sub>2</sub>N•BH<sub>3</sub> + H<sub>2</sub> Scheme 26. Synthesis of 1-propanamine-borane via SBH/NaHCO<sub>3</sub> & free amine



**Figure 6.** Web of Science<sup>TM</sup> citation report with "amineboranes" as the Topic from 1937-2020 (per 19-May-2020) **3. Conclusion** 

Simply by looking at the numerous synthetic strategies, it is evident that amine-boranes have maintained their relevance since their discovery. However, a quick Web of Science<sup>TM</sup> search using the "Topic" "amine-boranes" reveals an exponentially growing interest in these salient compounds (Figure 6). In light of the renewed attention received by amine-boranes and their plentiful applications, it is imperative that new methods are optimized and developed to allow access to as many forms of these compounds as possible.

## Acknowledgements

Nil.

#### References

- Burnham, B.S., Amine-Boranes, in Encyclopedia of Metalloproteins, R.H. Kretsinger, V.N. Uversky, and E.A. Permyakov, Editors. 2013, Springer New York: New York, NY. pp. 58-62.
- [2] A. Staubitz, A. P. M. Robertson, M. E. Sloan, and I. Manners, Amine- and Phosphine-Borane Adducts: New Interest in Old Molecules. *Chem. Rev.*, 110 (2010) 4023-4078.
- [3] A. B. Burg, H. I. Schlesinger, Hydrides of Boron. VII. Evidence of the Transitory Existence of Borine (BH3): Borine Carbonyl and Borine Trimethylammine. J. Am. Chem. Soc., 59 (1937) 780-787.
- [4] M. Couturier, B. M. Andresen, J. L. Tucker, P. Dubé, S. J. Brenek, J. T. Negri, The use of borane–amine adducts as versatile palladium-catalyzed hydrogen-transfer reagents in methanol. *Tetrahedron. Lett.*, 42 (2001) 2763-2766.
- [5] (a) A. E. Carre-Burritt, B. L. Davis, B. D. Rekken, N. Mack, T. A. Semelsberger, Enabling ammonia-borane: co-

oligomerizaiton of ammonia-borane and amine-boranes yield liquid products. *Energ. Environ. Sci.*, 7 (2014) 1653-1656. (b) B. L. Dietrich, K. I. Goldberg, D. M. Heinekey, T. Autrey, J. C. Linehan, Iridium-Catalyzed Dehydrogenation of Substituted Amine Boranes: Kinetics, Thermodynamics, and Implications for Hydrogen Storage. *Inorg. Chem.*, 47 (2008) 8583-8585.

[6] (a) S. B. Bruce, Synthesis and pharmacological activities of amine-boranes. *Curr. Med. Chem.*, 12 (2005) 1995-2010.
(b) I. H. Hall, C. J. Gilbert, A. T. McPhail, K. W. Morse, K. Hassett, B. F. Spielvogel, Antineoplastic activity of a series of boron analogues of α-amino acids. *J. Pharm. Sci.*, 74 (1985) 755-758.

(c) A. Sood, C. K. Sood, B. F. Spielvogel, I. H. Hall, O. T. Wong, Synthesis, cytotoxicity, hypolipidemic and antiinflammatory activities of amine—boranes and esters of boron analogues of choline and thiocholine. *J. Pharm. Sci.*, 81 (**1992**) 458-462.

- [7] R. O. Hutchins, K. Learn, B. Nazer, D. Pytlewski, A. Pelter, Amine boranes as selective reducing and hydroborating agents. A review. Org. Prep. Proced. Int., 16 (1984) 335-372.
- [8] (a) M. A. Gamal-Eldin, D. H. Macartney, Host-Guest Complexations of Amine Boranes and Isoelectronic/Isostructural Quaternary Alkylammonium Cations by Cucurbit[7]uril in Aqueous Solution. *Heteroat. Chem.*, 2019 (2019) 8124696.
- [9] A. S. Kulkarni, P. V. Ramachandran, Water-Promoted, Open-Flask Synthesis of Amine-Boranes: 2-Methylpyridine-Borane (2-Picoline-Borane). Org. Synth., 94 (2017) 332-345.
- [10] (a) G. Alcaraz, S. Sabo-Etienne, Coordination and dehydrogenation of amine-boranes at metal centers. *Angew. Chem. Int. Ed.*, 49 (2010) 7170-7179.
  (b) S. Babu Kalidindi, U. Sanyal, B. R. Jagirdar, *Chem. Sus. Chem.*, 4 (2011) 317-324.
  (c) J. M. Brunel, B. Faure, M. Maffei, Phosphane-boranes: Synthesis, characterization and synthetic applications. *Coord. Chem. Rev.*, 187-180 (1998) 665-698.
- [11] A. S. Kulkarni, Amine-Boranes: Novel Syntheses and Application as Green Hypergolic Propellants (Doctoral Dissertation, Purdue University, West Lafayette, USA (2017). Retrieved from https://docs.lib.purdue.edu/dissertations/AAI10281230/.
- [12] (a) H. C. Brown, B. Singaram, Molecular addition compounds. 7. Synthesis of addition compounds of boron trifluoride, borane, and alane with N,N,N'N'tetramethylethylenediamine and triethylenediamine by precipitation from ether solvents. *Inorg. Chem.*, 19 (1980) 455-457.

(b) A. R. Gatti, T. Wartik, Preparation and properties of the mono-and bisborane adducts of N, N'-dimethylpiperazine and triethylenediamine. *Inorg. Chem.*, 5 (1966) 2075-2076.

- [13] L. Zhang, S. Li, Y. Tan, Z. Tang, Z. Guo, X. Yu, Synthesis and hydrogen release properties of alkyl-substituted amineboranes. J. Mater. Chem. A, 2 (2014) 10682-10687.
- [14] Y. Nakayama, H. Nakajima, U.S. Patent No. 10,407,448. Washington, DC: U.S. Patent and Trademark Office (2019).
- [15] (a) H. Helten, A. Robertson, A. Staubitz, J. R. Vance, M. F. Haddow, I. Manners, Ambient Temperature Dehydrocoupling of Aromatic Amine-Boranes. *Chem. Eur. J.*, 18 (2012) 4665-4680.

(b) C. A. Jaska, K. Temple, A. J. Lough, I. Manners, Transition metal-catalyzed formation of boron– nitrogen bonds: catalytic

dehydrocoupling of amine-borane adducts to form aminoboranes and borazines. *J. Am. Chem. Soc.*, 125 (2003) 9424-9434.

- [16] H. Flores-Segura, L. A. Torres, Enthalpies of formation of primary, secondary, and tertiary amineborane adducts in tetrahydrofuran solution. *Struct. Chem.*, 8 (1997) 227-232.
- [17] H. W. Roesky, D. A. Atwood, *Group 13 Chemistry I:* Fundamental New Developments. 2002, Springer: New York.
- [18] N. Zheng, J. Fan, G. D. Stucky, One-step one-phase synthesis of monodisperse noble-metallic nanoparticles and their colloidal crystals. J. Am. Chem. Soc., 128 (2006) 6550-6551.
- [19] H. C. Kelly, M. B. Giusto, F. R. Marchelli, Amineborane reductions in aqueous acid media. J. Am. Chem. Soc., 86 (1964) 3882-3884.
- [20] W. M. Jones, Amine Borane Reductions. The Stereochemistry of the Reduction of 4-t-Butylcyclohexanone with Trimethylamine Borane in the Presence and Absence of Boron Fluoride. J. Am. Chem. Soc., 82 (1960) 2528-2532.
- [21] (a) H. C. Brown, B. C. S. Rao, A New Powerful Reducing Agent—Sodium Borohydride in the Presence of Aluminum Chloride and Other Polyvalent Metal Halides1,2. J. Am. Chem. Soc., 78 (1956) 2582-2588.

(b) W. Büchner, H. Niederprüm, *Sodium Borohydride And Amine-Boranes, Commercially Important Reducing Agents*, In *Boron Chemistry–3*, H. Nöth, Editor. 1977, Pergamon. pp. 733-743.

(c) Z. M. Heiden, A. P. Lathem, Establishing the hydride donor abilities of main group hydrides. *Organometallics*, 34 (2015) 1818-1827.

- [22] Todd, D., J. Chem. Educ., 1979, 56(8): 540.
- [23] Wietelmann, U., Felderhoff, M., and Rittmeyer, P., *Hydrides*, in *Ullmann's Encyclopedia of Industrial Chemistry*. 2016, Wiley-VCH Verlag GmbH & Co. KGaA: Weinheim, Germany. pp. 1-39.
- [24] A. E. Finholt, A. C. Bond, H. I. Schlesinger, Lithium Aluminum Hydride, Aluminum Hydride and Lithium Gallium Hydride, and Some of their Applications in Organic and Inorganic Chemistry. J. Am. Chem. Soc., 69 (1947) 1199-1203.
- [25] H. I. Schlesinger, H. C. Brown, Metallo borohydrides. III. Lithium borohydride. J. Am. Chem. Soc., 62 (1940) 3429-3435.
- [26] H. C. Brown, S. Krishnamurthy, Lithium triethylborohydride. Exceptionally powerful nucleophile in displacement reactions with organic halides. J. Am. Chem. Soc., 95 (1973) 1669-1671.
- [27] C. F. Lane, Sodium cyanoborohydride—a highly selective reducing agent for organic functional groups. *Synthesis*, 3 (1975) 135-146.
- [28] G. B. Fisher, J. C. Fuller, J. Harrison, S. G. Alvarez, E. R. Burkhardt, C. T. Goralski, B. Singaram, Aminoborohydrides.
  4. The Synthesis and Characterization of Lithium Aminoborohydrides: A New Class of Powerful, Selective, Air-Stable Reducing Agents. J. Org. Chem., 59 (1994) 6378-6385.
- [29] T. L. Ho, M. Fieser, L. F. Fieser, R. Danheiser, W. Roush, J. Smith, *Diisobutylaluminum Hydride*, (*DIBAL-H*), in *Fieser and Fieser's Reagents for Organic Synthesis*, L.F. Fieser, M. Fieser, and T.-L. Ho, Editors. (2013) 253-253.
- [30] M. Gugelchuk, L. F. Silva III, R. S. Vasconcelos, S. A. P. Quintiliano, Sodium Bis(2-methoxyethoxy)aluminum Hydride, in Encyclopedia of Reagents for Organic Synthesis (2007).
- [31] J. L. Fry, J. R. Rahaim, R. E. Maleczka, *Triethylsilane*, in *Encyclopedia of Reagents for Organic Synthesis* (2007).
- [32] (a) H. C. Brown, S. Krishnamurthy, Forty years of hydride

reductions. Tetrahedron, 35 (1979) 567-607.

(b) H. C. Brown, P. V. Ramachandran, *Sixty Years of Hydride Reductions*, in *Reductions in Organic Synthesis*. American Chemical Society. (1996) 1-30.

- [33] (a) W. J. Atkins, E. R. Burkhardt, K. Matos, Safe handling of boranes at scale. Org. Process Res. Dev., 10 (2006) 1292-1295.
  (b) Lane, C.F. Ammonia-Borane and Related N-B-H Compounds and Materials: Safety Aspects, Properties and Applications, DOE Chemical Hydrogen Storage Center of Excellence: Northern Arizona University. (2006) 1-33.
- [34] (a) M. B. Eleveld, H. Hogeveen, Enantioselective reduction of acetophenone by borane. chiral amine complexes. *Tetrahedron. Lett.*, 27 (1986) 635-638.
  (b) J. B. Le Toumelin, M. Baboulène, Chiral intramolecular amine-borane complexes as reducing agents for prochiral ketones. *Tetrahedron: Asymmetry*, 8 (1997) 1259-1265.
- [35] (a) M. Periasamy, J. V. B. Kanth, C. K. Reddy, New routes for the synthesis of organometallic reagents. *J. Chem. Soc.*, *Perkin Trans.* 1 (1995) 427-430.
  (b) T. Yamashita, H. Mitsui, H. Watanabe, N. Nakamura, Preparations and asymmetric, reducing abilities of chiral polymeric amine-boranes. *Die Makromolekulare Chemie*, 181 (1980) 2563-2569.
- [36] A. Hirao, S. Itsuno, S. Nakahama, N. Yamazaki, Asymmetric reduction of aromatic ketones with chiral alkoxy-amineborane complexes. J. Chem. Soc., Chem. Commun., 7 (1981) 315-317.
- [37] (a) E. J. Corey, R. K. Bakshi, S. Shibata, Highly enantioselective borane reduction of ketones catalyzed by chiral oxazaborolidines. Mechanism and synthetic implications. *J. Am. Chem. Soc.*, 109 (1987) 5551-5553.
  (b) E. J. Corey, R. K. Bakshi, S. Shibata, C. P. Chen, V. K. Singh, A stable and easily prepared catalyst for the enantioselective reduction of ketones. Applications to multistep syntheses. *J. Am. Chem. Soc.*, 109 (1987) 7925-7926.
- [38] M. G. Hu, J. M. Van Paasschen, R. A. Geanangel, New synthetic approaches to ammonia-borane and its deuterated derivatives. *J. Inorg. Nucl. Chem.*, 39 (1977) 2147-2150.
- [39] H. C. Brown, U.S. Patent No. 2,860,167. Washington, DC: U.S. Patent and Trademark Office (1958).
- [40] H. C. Brown, H. I. Schlesinger, S. Z. Cardon, Studies in stereochemistry. I. Steric strains as a factor in the relative stability of some coördination compounds of boron. J. Am. Chem. Soc., 64 (1942) 325-329.
- [41] R. Baldwin, R. Washburn, Borane Exchange Reactions. J. Org. Chem., 26 (1961) 3549-3550.
- [42] (a) S. G. Shore, X. Chen, U.S. Patent No. 8,637,703 B2. Washington, DC: U.S. Patent and Trademark Office (2014).
  (b) P. V. Ramachandran, A. S. Kulkarni, Nucleophilic displacement of ammonia from ammonia borane for the preparation of alkylamine-, pyridine-and phosphine-boranes. *RSC Adv.*, 4 (2014) 26207-26210.
- [43] G. W. Schaeffer, E. R. Anderson, The Preparation of Trimethylamine-borine, N-Trimethylborazole and N-Dimethylaminoborine. J. Am. Chem. Soc., 71 (1949) 2143-2145.
- [44] M. D. Banus, T. R. P. Gibb, U.S. Patent No. 2,678,949. Washington, DC: U.S. Patent and Trademark Office (1954).
- [45] A. V. Safronov, S. S. Jalisatgi, M. F. Hawthorne, U.S. Patent No. 10,179,795 B2. Washington, DC: U.S. Patent and Trademark Office (2019).
- [46] (a) W. H. Schecter, U.S. Patent No. 3,033,766. Washington,

DC: U.S. Patent and Trademark Office (1962).

(b) W. H. Schecter, R. M. Adams, G. F. Huff, *U.S. Patent No.* 2,938,923. Washington, DC: U.S. Patent and Trademark Office (1960).

- [47] M. D. Taylor, L. R. Grant, C. A. Sands, A convenient preparation of pyridine-borane. J. Am. Chem. Soc., 77 (1955) 1506-1507.
- [48] H. I. Schlesinger, H. C. Brown, A. E. Finholt, The Preparation of Sodium Borohydride by the High Temperature Reaction of Sodium Hydride with Borate Esters1. J. Am. Chem. Soc., 75 (1953) 205-209.
- [49] (a) H. C. Brown, Magnesium amide bases and amido-Grignards. 1. Ortho magnesiation. J. Organomet. Chem., 100 (1975) 3-15.

(b) H. C. Brown, From little acorns to tall oaks-from boranes through organoboranes. *Science*, 210 (1980) 485-492.

- [50] (a) Japanese Patent Laid-open Publication No. 56(1981)-158792. Tokyo, Japan: Japan Patent Office.
  (b) Japanese Patent Laid-open Publication No. 10(1998)-109991. Tokyo, Japan: Japan Patent Office.
  (c) A. R. Cartolano, S. V. Ivanov, C. I. Teich, J. H. Yamamoto, U.S. Patent No. 8,039,666 B2. Washington, DC: U.S. Patent and Trademark Office (2011).
- [51] R. Köster, Some aspects of the surface chemistry of carbon blacks and other carbons. *Angew. Chem.*, 69 (1957) 94-97.
- [52] H. Jenkner, U.S. Patent No. 3,051,754. Washington, DC: U.S. Patent and Trademark Office (1962).
- [53] (a) S. V. Ivanov, B. Casas, U.S. Patent No. 7,718,154 B2. Washington, DC: U.S. Patent and Trademark Office (2010).
  (b) H. Jenkner, U.S. Patent No. 3,103,416. Washington, DC: U.S. Patent and Trademark Office (1963).
- [54] R. W. Bragdon, *U.S. Patent No. 2,927,133*. Washington, DC: U.S. Patent and Trademark Office (1960).
- [55] (a) Farbenfabriken-Bayer-Aktiengesellschaft, *British Patent No.* 822,229. London: The Patent Office (1959).
  (b) A. A. Hinckley, *U.S. Patent No.* 3,127,448. Washington, DC: U.S. Patent and Trademark Office (1964).
- [56] H. Haberland, R. Stroh, U.S. Patent No. 3,013,016.Washington, DC: U.S. Patent and Trademark Office (1961).
- [57] (a) J. A. Marsella, U.S. Statutory Invention Reg. Number H919. Washington, DC: U.S. Patent and Trademark Office (1991).
  - (b) J. M. Sullivan, *U.S. Patent No.* 5,516,909. Washington, DC: U.S. Patent and Trademark Office (1996).
  - (c) J. M. Sullivan, U.S. Patent No. Re. 36,115. Washington, DC: U.S. Patent and Trademark Office (1999).
- [58] K. Lang, F. Schubert, U.S. Patent No. 3,037,985. Washington, DC: U.S. Patent and Trademark Office (1962).
- [59] J. A. Snover, U.S. Patent No. 3,317,525. Washington, DC: U.S. Patent and Trademark Office (1967).
- [60] (a) E. C. Ashby, U.S. Patent No. 3,257,455. Washington, DC: U.S. Patent and Trademark Office (1966).
  (b) E. C. Ashby, U.S. Patent No. 3,153,671. Washington, DC: U.S. Patent and Trademark Office (1964).

- [61] H. C. Kelly, J. O. Edwards, Ethane 1, 2-Diamineborane. J. Am. Chem. Soc., 82 (1960) 4842-4846.
- [62] Archive, A.C.S.-M.o.t.W. *Tetrahydrofuran*. 2015 [cited 2020 06-Apr]; Available from: https://www.acs.org/content/acs/en/molecule-of-theweek/archive/t/tetrahydrofuran.html.
- [63] M. D. Marshall, U.S. Patent No. 3,192,217. Washington, DC: U.S. Patent and Trademark Office (1965).
- [64] E. C. Ashby, W. E. Foster, A new and Convenient Route to the Amine-Boranes. J. Am. Chem. Soc., 84 (1962) 3407-3408.
- [65] S. Matsumura, N. Tokura, Preparation of triethylamine-borane or pyridine-borane in liquid sulfur dioxide and reduction of aralkyl halides with base-borane complex in liquid sulfur dioxide or in nitromethane. *Tetrahedron Lett.*, 9 (1968) 4703-4705.
- [66] K. C. Nainan, G. E. Ryschkewitsch, New synthesis of amineand phosphine-boranes. *Inorg. Chem.*, 8 (1969) 2671-2674.
- [67] A. R. Burke, W. V. Hough, U.S. Patent No. 4,080,381.Washington, DC: U.S. Patent and Trademark Office (1978).
- [68] [68] J. Plesek, B. Stibr, E. Drdakova, T. Jelinek, *Czech Patent* 242,064. Prague, Czech Republic: Industrial Property Office (1986).
- [69] A. J. Arduengo, U.S. Patent No. 5,144,032. Washington DC: U.S. Patent and Trademark Office (1992).
- [70] F. Cao, Z. Y. Fang, F. Chen, Q. Shen, S. Q. Wang, B. Li, Low Cost Chemical Synthesis of Ammonia Borane Complex for Hydrogen Storage. *Key Eng. Mater.*, 519 (2012) 92-95.
- [71] V. Kampel, A. Warshawsky, Organotin compounds and their therapeutic potential: a report from the Organometallic Chemistry Department of the Free University of Brussels. J. Organomet. Chem., 469 (1994) 15-17.
- [72] C. J. Collins, M. Lanz, C. T. Goralski, B. Singaram, Aminoborohydrides. 10. The Synthesis of Tertiary Amine–Boranes from Various Benzyl Halides and Lithium N,N-Dialkylaminoborohydrides. J. Org. Chem., 64 (1999) 2574-2576.
- [73] P. V. Ramachandran, P. D. Gagare, Preparation of ammonia borane in high yield and purity, methanolysis, and regeneration. *Inorg. Chem.*, 2007, 46(19): 7810-7817.
- [74] P. V. Ramachandran, H. Mistry, A. S. Kulkarni, P. D. Gagare, Ammonia-mediated, large-scale synthesis of ammonia borane. *Dalton Trans.*, 2014, 43(44): 16580-16583.
- [75] P. V. Ramachandran, A. S. Kulkarni, The role of ammonia in promoting ammonia borane synthesis. *Dalton Trans.*, 45 (2016) 16433-16440.
- [76] V. Ramachandran, A. S. Kulkarni, Y. Zhao, J. G. Mei, Amine–boranes bearing borane-incompatible functionalities: application to selective amine protection and surface functionalization. *Chem. Commun.*, 52 (2016) 11885-11888.
- [77] P. V. Ramachandran, B. C. Raju, P. D. Gagare, One-Pot Synthesis of Ammonia–Borane and Trialkylamine–Boranes from Trimethyl Borate. *Org. Lett.*, 2012, *14*(24): 6119-6121.
- [78] P. V. Ramachandran, A. S. Kulkarni, Open-Flask Synthesis of Amine–Boranes via Tandem Amine–Ammonium Salt Equilibration–Metathesis. *Inorg. Chem.*, 54 (2015) 5618-5620.

## How to Cite This Article

Daniel O Reddy. "A Short Chronological Review on the Syntheses of Amine-Boranes". Chemical Review and Letters, 3, 4, 2020, 184-191. doi: 10.22034/crl.2020.238849.1070