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Experimental and Computational Study of the Formation Mechanism of the Diammoniate of Diborane: The Role of Dihydrogen Bonds

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Supporting Information

ABSTRACT: The mechanism of formation of ammonia borane (NH $_3$ BH $_3$, AB) and the diammoniate of diborane ([H $_2$ B(NH $_3$) $_2$][BH $_4$], DADB) in the reaction between NH $_3$ and THF \cdot BH $_3$ was explored experimentally and computationally. Ammonia diborane (NH $_3$ BH $_2$ (μ -H)BH $_3$, AaDB), a long-sought intermediate proposed for the formation of DADB, was directly observed in the reaction using 11 B NMR spectroscopy. The results indicate that dihydrogen bonds between the initially formed AB and AaDB accelerate the formation of DADB in competition with the formation of AB.

Adihydrogen bond is an attractive interaction between a positively charged H atom (such as N–H, O–H, or other proton donor) and a negatively charged H atom (such as M–H or B–H, where M is usually a transition metal). Such $H\cdots H$ interactions were first noticed by Brown through solution IR spectra in the late 1960s. Although substantial progress has been made over the past two decades in the identification and characterization of the dihydrogen bond, is possible applications in synthetic chemistry have not been widely explored. Recent investigations of the role of the dihydrogen bond in amine—borane systems indicate that $N-H\cdots H-B$ dihydrogen bonds could promote new syntheses/reactions.

A diborane ammonia complex, which was first observed in the 1920s by Stock et al. in a reaction between ammonia (NH₃) and diborane (B₂H₆), was characterized with the empirical formula B₂H₆·2NH₃ and thus was called the diammoniate of diborane (DADB). Over the years, four possible structural formulas were proposed on the basis of the chemical composition: (a) [NH₄]₂ $[B_2H_4]^{,8}$ (b) $[NH_4][H_3BNH_2BH_3]^{,9}$ (c) $[(NH_4)(BH_2NH_2)]^{,9}$ (BH_4)], and (d) [NH₃BH₂NH₃][BH₄]. The structure of DADB was shown to be formula d in a series of papers from Parry's group in 1958.¹¹ Recently, the crystal structure of DADB was also reported.¹² Interestingly, the anion in the structural formula b, [H₃BNH₂BH₃]⁻, was synthesized very recently through the reaction of Na or NaNH₂ with ammonia borane (NH₃BH₃, AB) under reflux in THF. 13 DADB was also found as a side product in a simple displacement reaction between NH3 and tetrahydrofuran borane (THF · BH₃) along with the desirable product AB, ¹⁴ which is a widely studied compound for hydrogen storage. 15

The mechanism of DADB formation has attracted considerable attention. ¹⁶ Factors such as steric effects and solvent properties were considered ¹⁶ but were found to be insufficient to explain the DADB formation. ¹⁷ Ammonia diborane (NH₃BH₂(μ -H)BH₃, ¹⁸ denoted as AaDB in order to distinguish it from aminodiborane (NH₂B₂H₅, ADB^{5,19})), has long been contemplated as an intermediate

Scheme 1. Proposed Mechanism Leading to DADB and AB

associated with the formation of DADB, although it was never identified experimentally. Here we present the first direct ¹¹B NMR evidence for the existence of the AaDB intermediate in the reaction of NH₃ with a THF·BH₃ solution. Moreover, the mechanism leading to DADB formation from the AaDB intermediate is experimentally and computationally explored.

A three-step mechanism for the formation of a mixture of AB and DADB is proposed (Scheme 1). In step 1, AB is produced in a displacement reaction. AB further reacts with THF \cdot BH $_3$ to form the AaDB intermediate in step 2. In step 3, NH $_3$ can react with AaDB in one of two pathways: (I) NH $_3$ attacks B $_a$ to form DADB or (II) NH $_3$ attacks B $_b$ to form two molecules of AB.

To study the mechanism of the DADB formation, ¹¹B NMR spectroscopy, which can easily distinguish AB from DADB, was employed to follow the reaction. Figure 1 shows a stacked plot of ¹¹B NMR spectra recording an 11 min period from the beginning to completion of the reaction of NH₃ gas with a THF · BH₃ solution at -78 °C. As the reaction proceeded, the THF · BH₃ signal (δ -0.9 ppm) decreased, and a new signal characteristic of AB (δ -22.5 ppm) increased after a slight delay. Two additional small signals at δ -13.8 and -25.5 ppm appeared within 3 min and were assigned respectively to B_aH₂ (triplet) and B_bH₃ (quartet) in the AaDB intermediate (Scheme 1). At \sim 7 min, a broad peak emerged at δ -38.0 ppm, which gradually increased in amplitude and changed

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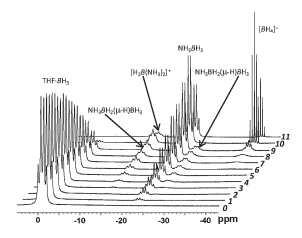


Figure 1. 11 B NMR spectra of the reaction of NH₃ with a THF·BH₃ solution at -78 °C. Samples were extracted at 1 min intervals and stored at -78 °C. After the reaction was complete, spectra were rapidly recorded for each sample as it warmed to ambient temperature.

to a quintet characteristic of $\mathrm{BH_4}^-$. Finally, after 11 min of reaction, only signals due to AB and DADB were present in the spectrum.

The transformation from the AaDB intermediate to DADB product is clearly reflected in the formation of the BH₄⁻ anion in the 11 B NMR spectra. When the quartet signal at δ –25.5 ppm, assigned to B_bH₃ in the AaDB, gradually decreased and finally disappeared, a new BH₄⁻ peak at δ –38.0 ppm appeared and gradually increased at the same time. In contrast, the triplet at δ –13.8 ppm assigned to B_aH₂ increased steadily in the same period of time rather than producing the corresponding chemical shift change. One possible explanation is the coincidence of the chemical shifts of B_aH₂ and $[H_2B(NH_3)_2]^+$ at δ –13.8 ppm.

The observed values of 11 B chemical shifts, $\delta - 13.8$ and -25.5 ppm, for B_aH_2 and B_bH_3 groups in AaDB are not consistent with the results calculated at the B3LYP/6-311++G(d,p) level of theory with GIAO method. This discrepancy is attributed to a dihydrogen bond effect on the 11 B NMR spectra of AaDB. When an AB binds AaDB through $B-H\cdots H-N$ dihydrogen bonds, the value calculated for B_b shifts significantly to $\delta - 19.8$ ppm. When two AB molecules interact with AaDB, the calculated B_b chemical shift is $\delta - 24.4$ ppm, in good agreement with the observed result. For comparison, the solvation effect of THF on the 11 B chemical shifts of AaDB is less pronounced because the $N-H\cdots O$ H-bonds formed are remote from B_b . A calculated chemical shift of $\delta - 17.9$ ppm for B_b was obtained when three THF molecules were H-bonded to AB (see Supporting Information for computational details).

Attempts to identify the AaDB intermediate by 1 H, 1 H{ 11 B}, and two-dimensional NMR spectroscopies were unsuccessful. The weak signals from AaDB were obscured by the large THF \cdot BH₃ and AB signals. However, a control experiment clearly revealed separate resonances for NH₃BH₂NH₂BH₃ and AaDB, proving that the resonances at δ -13.8 and -25.5 ppm are not from NH₃BH₂NH₂BH₃ (Figures S3 and S4).

The proposed mechanism in Scheme 1 is supported by a variety of experimental results. When THF \cdot BH $_3$ reacted with a large excess of liquid NH $_3$ at -78 °C, pure AB (Figure S8) was obtained, indicating that only step 1 had occurred. No THF \cdot BH $_3$ remained for the formation of AaDB in step 2. In contrast, when NH $_3$ gas was passed over a THF \cdot BH $_3$ solution at -78 °C, the reaction proceeded along all three steps to produce a mixture of AB and DADB (Figure 1).

Table 1. Relationship between Dihydrogen Bonds and the Ratio of DADB to AB

	H-bonds	ratio DADB/AB ^a
$THF \cdot BH_3 (1 M) + NH_3$	н…н	1:4.0
$THF \cdot BH_3 (1 M) + ND_3$	$D \cdots H$	1:4.1
$THF \cdot BH_3 (0.5 M) / AB (0.5 M) + NH_3$	$H \cdots H^b$	1:2.6
THF · BH ₃ (0.5 M)/TMAB (0.5 M) c + NH ₃	$H\cdots H^b$	1:15

 a The ratio of DADB to AB is based on the integrated values of the ${\rm [BH_4]}^-$ anion and formed AB in $^{11}{\rm B\{^1H\}}$ NMR spectra. b The intentionally added AB is much more effective in forming dihydrogen bonds than the intentionally added TMAB. c The actual concentration of TMAB is ca. 0.3 M.

Formation of DADB (pathway I) was perhaps enhanced by the dihydrogen bond formation between AB and AaDB. To test this hypothesis, NH₃ gas was passed over an equimolar solution (0.5 M) of THF·BH₃ and AB at -78 °C. The ratio of DADB to AB in the product increased, consistent with increased dihydrogen bonding to AaDB promoted by the high concentration of AB. When NH₃ gas was passed over an equimolar solution (0.5 M) of THF·BH₃ and trimethylamine borane (TMAB) at -78 °C, the ratio of DADB to AB decreased significantly. The presence of methyl groups makes TMAB less effective than AB in forming dihydrogen bonds with AaDB. When ND₃ gas was passed over a THF·BH₃ solution at -78 °C, the deuterium isotope effect was found to be minimal (Table 1).

The proposed mechanism presented in Scheme 1 and the role of dihydrogen bonds between AB and AaDB in the formation of DADB were supported by computational studies carried out at the MP2/6-31++G(d,p) level of theory (see Supporting Information for computational details). Step 1 is recognized as a typical S_N2 reaction. In the transition state, the borane moiety is planar, and breaking of the B–O bond and formation of the N–B bond occur simultaneously. The calculated activation energy barrier is 8.4 kcal/mol, and the reaction is exothermic (–5.6 kcal/mol) (Table S1).

The optimized reactant complex (RC), transition state (TS), and product complex (PC) for step 2 are shown in Figure 2. In the RC, AB interacts with THF \cdot BH $_3$ via a bifurcated dihydrogen bond. The transition state is created by binding a hydridic H of AB (negative charge -0.11 according to natural population analysis (NPA), Figure S26) to the borane moiety on THF \cdot BH $_3$. In the TS, a planar borane moiety of THF \cdot BH $_3$ is identified, suggesting simultaneous cleavage of the B-O bond and formation of an H-B bond. The predicted reaction energy barrier is 15.5 kcal/mol (Table 2). In the PC, a N-H \cdot · · · O hydrogen bond with THF stabilizes AaDB. The formation of AaDB from AB and THF \cdot BH $_3$ is slightly endothermic (1.4 kcal/mol), suggesting that formation of AaDB may be reversible.

In step 3 of Scheme 1, NH₃ attacks either B_a or B_b of the AaDB intermediate (pathway I or II, respectively). The calculated energy barrier for pathway I (30.6 kcal/mol in a vacuum) was significantly higher than the barrier of pathway II (14.3 kcal/mol) (Table 3, Figure S28). When the THF solvation effect was included (a polarizable continuum model (PCM) method), the resulting energy barrier for pathway I was still much higher than that for pathway II (Table 3). NPA charge analysis for the TS of pathway I showed that the BH₄ moiety has a negative charge (-0.79) and the NH₃BH₂ moiety has a positive charge (+0.65) (Figure S29). Their strong attractive interaction may be responsible for the high energy barrier to create the BH₄ anion and the

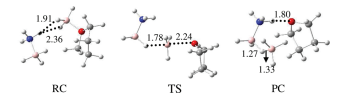


Figure 2. Optimized RC, TS, and PC for the reaction between AB and $THF \cdot BH_3$ to produce the AaDB intermediate. Colors: C, gray; H, white; N, blue; B, pink; and O, red. Bond lengths are shown in Å.

Table 2. Energetics of the Reaction between AB and THF·BH₃ To Produce AaDB (kcal/mol at 0 and 195 K), Calculated at the MP2/6-31++G(d,p) Level of Theory^a

	ΔE_0	ΔH_0	ΔH_{195}	ΔG_{195}	$\Delta H_0(PCM)$	$\Delta G_{195}(PCM)$
RC	0.0	0.0	0.0	0.0	0.0	0.0
TS	16.9	15.5	15.5	15.1	15.2	14.9
PC	1.2	1.3	1.3	1.7	3.4	3.8

^a The THF solvation effect was analyzed using the PCM approach. The single-point energies were calculated using optimized geometries in a vacuum with dielectric constant $\varepsilon = 7.4257$.

NH₃BH₂NH₃⁺ cation for formation of DADB. In contrast, there was no significant charge separation for the TS of pathway II. As a consequence, AB formation has a lower energy barrier.

Computational analysis of the reaction of NH₃ with AaDB in the presence of one, two, three, and four AB molecules provides a possible explanation for DADB formation as the reaction nears completion (Table 3). As the proportion of AB increases, the activation energies for pathways I and II become nearly comparable. Dihydrogen bonds from one AB to AaDB reduced the calculated activation energy barrier leading to DADB by 5.7 kcal/mol. Figure 3 contains the relevant RC, TS, and product structures for pathways I and II of step 3. TS1, a protonic H of AB, forms a dihydrogen bond with the BH₄ group, facilitating the separation of BH₄⁻ to produce DADB. More dihydrogen bonds form when two AB molecules interact with AaDB, providing assistance in the separation of BH₄⁻. The calculated energy barrier of pathway I decreases to 20.7 kcal/mol. When four AB molecules are considered, the predicted activation energy for pathway I is only 15.9 kcal/mol, compared to 11.8 kcal/mol for pathway II.

The catalytic role of AB molecules in the formation of DADB is attributed to the fact that the AB molecule has a partially positively charged protonic end and a partially negatively charged hydridic end. The partially positively charged end favors the BH₄ group, and the partially negatively charged end prefers to interact with the NH₃BH₂ moiety through B-H···H-N dihydrogen-bonding interactions in the TS leading to DADB. As a consequence, the attractive interaction between the BH₄⁻ group and the NH3BH2 group is weakened. Therefore, the reaction energy barrier for the formation of DADB is lowered due to the assistance of AB molecules via the intermolecular dihydrogen bonds, rendering pathway I to form DADB feasible. In comparison with pathway II to form AB, pathway I still has a relatively higher energy barrier. Therefore, one can expect AB to be the major product and DADB a minor product, which is consistent with the experimental observations.

Here we have explicitly analyzed the influence of dihydrogen bonds on the activation energy of pathways I and II. The solvation effect of polar solvents ²¹ or ionic liquids ²² plays a similar role through dipole—dipole or ion—dipole interactions. However, the dihydrogen

Table 3. Energetics of the Reaction between NH₃ and AaDB in the Presence of Zero, One, Two, Three, and Four AB (at the MP2/6-31++G(d,p) Level, in kcal/mol, at 0 and 195 K)^a

1411 2/0-3111	G(u,p)						
	ΔE_0	ΔH_0	ΔH_{195}	ΔG_{195}	$\Delta H_0(PCM)$	$\Delta G_{195}(PCM)$	
Without AB Molecule							
Pathway I: Attack B _a							
RC1	0.0	0.0	0.0	0.0	0.0	0.0	
TS1	32.1	30.6	30.1	31.9	19.9	21.1	
DADB	-3.0	-1.8	-2.6	-0.4	-9.5	-8.0	
Pathway II: Att	ack B _b						
RC2	0.0	0.0	0.0	0.0	0.0	0.0	
TS2	16.1	14.3	14.5	14.0	12.4	12.1	
2AB	-14.6	-13.1	-13.3	-12.8	-15.8	-15.5	
With One AB Molecule							
Pathway I: Atta	ack B _a						
RC1	0.0	0.0	0.0	0.0	0.0	0.0	
TS1	26.5	24.9	24.5	26.0	18.2	19.3	
$DADB \cdots AB$	-10.2	-9.0	-9.7	-7.1	-13.7	-11.8	
Pathway II: Att	ack B _b						
RC2	0.0	0.0	0.0	0.0	0.0	0.0	
TS2	14.2	12.5	12.6	12.4	11.0	10.9	
$2AB\cdots AB$	-14.0	-12.4	-12.6	-12.5	-16.1	-16.2	
		With '	Two AB	Molecu	ıles		
Pathway I: Atta	ack Ba						
RC1	0.0	0.0	0.0	0.0	0.0	0.0	
TS1	22.1	20.7	20.3	21.4	18.2	18.9	
DADB···2AB	-12.7	-11.3	-11.9	-10.1	-14.1	-13.0	
Pathway II: Att	ack B _b						
RC2	0.0	0.0	0.0	0.0	0.0	0.0	
TS2	12.9	11.5	11.4	11.8	10.9	11.2	
2AB · · · 2AB	-21.2	-18.7	-19.4	-17.2	-19.0	-17.5	
		With T	hree Al	3 Molec	rules		
Pathway I: Atta	ack B _a						
RC1	0.0	0.0	0.0	0.0	0.0	0.0	
TS1	19.5	18.0	17.7	18.6	16.8	17.4	
DADB···3AB	-16.0	-14.5	-15.1	-13.6	-15.9	-15.0	
Pathway II: Att	ack B _b						
RC2	0.0	0.0	0.0	0.0	0.0	0.0	
TS2	13.6	12.6	12.3	13.5	11.2	12.1	
2AB···3AB	-21.1	-19.1	-19.7	-17.2	-19.1	-17.3	
With Four AB Molecules							
Pathway I: Attack B _a							
RC1	0.0	0.0	0.0	0.0	0.0	0.0	
TS1	17.5	15.9	16.0	16.0	15.6	15.7	
DADB···4AB	-22.8	-20.7	-21.4	-18.2	-18.6	-16.2	
Pathway II: Attack B _b							
RC2	0.0	0.0	0.0	0.0	0.0	0.0	
TS2	12.4	11.8	11.8	12.1	11.1	11.3	
$2AB \cdots 4AB$	-24.8	-22.6	-22.8	-21.8	-20.6	-19.7	

^a The THF solvation effect was considered by using the PCM approach. The single-point energies were calculated using optimized geometries in a vacuum with dielectric constant $\varepsilon = 7.4257$.

bond may play a dominant role in determining the product types, based on this experimental and computational study. For instance, in

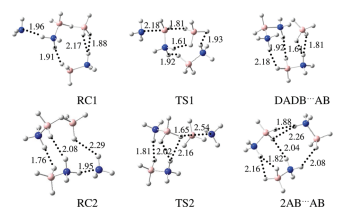


Figure 3. Optimized RC, TS, and products for both reaction pathways of NH_3 to AaDB in the presence of one AB. Colors: H, white; N, blue; B, pink. Bond lengths are shown in Å.

the intentional additions of AB or TMAB discussed above, the solvation effect of the reaction systems should be similar. However, the alteration of dihydrogen bonding between AB/TMAB and AaDB causes a significant difference in the ratio of DADB to AB.

In summary, both experimental and computational investigations have been applied to the classic displacement reaction between NH $_3$ and THF \cdot BH $_3$ to explore the formation mechanism of DADB, a side product. The results indicate that the initially produced AB reacts with THF \cdot BH $_3$ to yield an AaDB intermediate, which has been directly observed in the reaction using 11 B NMR spectroscopy. AB, produced by NH $_3$ reacting with either THF \cdot BH $_3$ or B $_b$ of AaDB, facilitates the separation of the BH $_4$ via dihydrogen bond interactions with AaDB that lower the reaction energy barrier to DADB production. Dihydrogen bonds may play a significant, but previously unrecognized, role in other reactions where reactants, intermediates, or products have both hydridic and protonic hydrogens.

■ ASSOCIATED CONTENT

Supporting Information. Experiments, computations, and characterization. This material is available free of charge via the Internet at http://pubs.acs.org.

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