Synthesis, Crystal Structure, Thermal Decomposition, and $^{11}$B MAS NMR Characterization of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$

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ABSTRACT: A metal borohydride–ammonia borane complex, Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$, is synthesized via a solid-state reaction between Mg(BH$_4$)$_2$ and NH$_3$BH$_3$. Different mechanochemical reaction mechanisms are observed, since Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ is obtained from $\alpha$-Mg(BH$_4$)$_2$, whereas a mixture of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$, NH$_3$BH$_3$, and amorphous Mg(BH$_4$)$_2$ is obtained from $\gamma$-Mg(BH$_4$)$_2$. The crystal structure of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ has been determined by powder X-ray diffraction and optimized by first-principles calculations. The borohydride groups act as terminal ligands, and molecular complexes are linked via strong dihydrogen bonds (<2.0 Å), which may contribute to the high melting point of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ found to be $\sim$48 °C in contrast to those for other molecular metal borohydrides. Precise values for the $^{11}$B quadrupole coupling parameters and isotropic chemical shifts are reported for the two NH$_3$BH$_3$ sites and two BH$_4^−$ sites in Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ from $^{11}$B MAS NMR spectra of the central and satellite transitions and MQMAS NMR. The $^{11}$B quadrupole coupling parameters agree excellently with the electric field gradients for the $^{11}$B sites from the DFT calculations and suggest that a more detailed structural model is obtained by DFT optimization, which allows evaluation of the dihydrogen bonding scheme.

1. INTRODUCTION

The increasing demand for energy and the environmental problems associated with the use of fossil fuels are the major driving forces for utilization of renewable energy. Hydrogen is recognized as a possible renewable energy carrier since it is widely abundant and environmentally friendly.$^{1,2}$ Complex metal hydrides such as borohydrides and ammidoboranes are considered as possible solid-state hydrogen storage materials owing to their high volumetric and gravimetric capacities.$^{3−8}$ However, they typically suffer from poor thermodynamic and/or kinetic properties.

Magnesium borohydride, Mg(BH$_4$)$_2$, is among the most studied borohydrides as a result of its high hydrogen content (14.8 wt % H$_2$) and complex chemistry. Mg(BH$_3$)$_2$ has been known since the 1950s,$^9$ but the crystal structure of the solvent-free polymorph $\alpha$-Mg(BH$_4$)$_2$ was first reported in 2009.$^{10,11}$ More recently, a new porous polymorph denoted $\gamma$-Mg(BH$_4$)$_2$ has been discovered, which can reversibly adsorb small molecules such as H$_2$, N$_2$, and CH$_4$Cl.$^{12}$ The thermal decomposition of both $\alpha$- and $\gamma$-Mg(BH$_4$)$_2$ has been intensively studied and turns out to be highly complex.$^{13−15}$ It is reported that $\alpha$-Mg(BH$_4$)$_2$ transforms into a high-temperature polymorph, $\beta$-Mg(BH$_4$)$_2$, at $T \approx 185$ °C, which becomes amorphous at $T \approx 300$ °C before it decomposes.

Another potential hydrogen storage material is ammonia borane, NH$_3$BH$_3$ (19.6 wt % H$_2$, 146 g H$_2$/L), which is stable in air.$^{16}$ NH$_3$BH$_3$ releases one equivalent of hydrogen in each of the three decomposition steps forming polyaminoborane, [NH$_2$BH$_2$]$_n$ (90−120 °C), polyaminoborane, [NHBH]$_n$ (120−200 °C), and finally boron nitride, BN (>500 °C).$^{17−20}$ Recently, the regeneration of ammonia borane has been reported to take place from polyaminoborane by reaction with hydrazine in liquid ammonia at 40 °C within 24 h.$^{21}$ However, the hydrogen release is accompanied by volatile and...
toxic byproducts, such as ammonia, diborane, and borazine. Metal amidoboranes are a class of materials that prevent the release of the diborane and borazine. They are synthesized by reacting metal hydrides with ammonia borane and typically decompose with fast kinetics and a release of hydrogen below 100 °C. This has prompted further modifications of ammonia borane with use of other complex anions such as amides and borohydrides. Recently, metal borohydrides and ammonia borane were reacted by mechanochemical treatment providing new compounds, i.e., LiBH$_4$(Am(BH$_3$)$_3$)$_{0.5}$, Mg(BH$_4$)$_2$(Am(BH$_3$)$_3$)$_2$, and Ca(BH$_4$)$_2$(Am(BH$_3$)$_3$)$_2$. However, the thermal decomposition of these compounds still involves release of diborane and borazine, like neat NH$_3$BH$_3$, reflecting a weak interaction between the borohydride and the ammonia borane. This is also observed in a recent study of NaBH$_4$–2NH$_3$BH$_3$ and KBH$_4$–2NH$_3$BH$_3$ mixtures which are not forming new compounds.

Physosorption of a small molecular hydride, i.e., ammonia borane, in a porous hydride is a tempting approach, which has motivated the present investigation. Here, we present new synthesis strategies and further structural investigations of the recently described Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ complex, using solid-state $^{11}$B MAS NMR, synchrotron radiation powder X-ray diffraction data (SR-PXD), and first-principles DFT calculations, which provide new interesting evidence for significant dihydrogen bonding in the borohydride–ammonia borane complex. The thermal decomposition is studied by variable-temperature $^{11}$B MAS NMR, in situ SR-PXD, a new photographic setup, and simultaneously thermogravimetric analysis, differential scanning calorimetry, and mass spectroscopy.

2. EXPERIMENTAL SECTION

2.1. Sample Preparation. Ammonia borane, NH$_3$BH$_3$ (Sigma-Aldrich, 97%), was used as received. $\alpha$-Mg(BH$_4$)$_2$ was synthesized in-house by reacting dibutyl magnesium, Mg-(nBu)$_2$, with (CH$_3$)$_2$S-BH$_3$ at room temperature, which forms the solvate Mg(BH$_4$)$_2$·$^1/2$(CH$_3$)$_2$S. Solvent-free $\alpha$-Mg(BH$_4$)$_2$ was obtained by heating the solvate to 140 °C for 3 h in argon atmosphere and followed by vacuum pumping for 1 h. The porous polymorph, $\gamma$-Mg(BH$_4$)$_2$, was prepared by heating the solvate to 80 °C under vacuum for 12 h. The synthesis products were examined by Fourier transform infrared spectroscopy (FTIR) and PXD.

Mixtures of Mg(BH$_4$)$_2$–NH$_3$BH$_3$ were prepared by ball milling (BM) in the ratios $\alpha$-Mg(BH$_4$)$_2$–NH$_3$BH$_3$ (1:1), (1:2), and (1:3) and $\gamma$-Mg(BH$_4$)$_2$–NH$_3$BH$_3$ (1:2), using a Frisch Pulverisette no. 4. These samples are denoted s1, s2, s3, and s4, respectively, see Table 1. The powders were packed in an inert atmosphere in a tungsten carbide vial (80 mL) together with tungsten carbide balls (diameter: 10 mm) in the w/w ratio 1:30. The powders were ball milled for 3 min followed by a 3 min break at 400 rpm and this sequence was repeated 135 times (405 min). Fractions of $\alpha$-Mg(BH$_4$)$_2$–NH$_3$BH$_3$ (1:2) were collected after 100 or 325 min characterized by PXD.

A mixture of $\alpha$-Mg(BH$_4$)$_2$ and NH$_3$BH$_3$ was also prepared in the molar ratio $\alpha$-Mg(BH$_4$)$_2$–NH$_3$BH$_3$ (1:2) by grinding by hand for 3 min in a mortar (sample denoted s5). Moreover, mixtures of $\gamma$-Mg(BH$_4$)$_2$–NH$_3$BH$_3$ (1:0.66) and (1:2) were ground in a similar manner for 10 min, and these samples are denoted s6 and s7, respectively. Samples s6 and s7 were loaded to a pellet die (diameter = 4 mm) followed by compression of the mixtures two times at 0.4 GPa with use of a PerkinElmer Press. The same procedure was repeated for amorphous Mg(BH$_4$)$_2$ reacted with NH$_3$BH$_3$ in the ratio 1:2 (sample denoted s8). The amorphous Mg(BH$_4$)$_2$ was obtained from $\gamma$-Mg(BH$_4$)$_2$ which becomes amorphous over time.

Fractions of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ (s2), $\alpha$-Mg(BH$_4$)$_2$, and NH$_3$BH$_3$ were heated from room temperature to 220 °C and then cooled to room temperature in a steel autoclave in an argon atmosphere. These samples are denoted s2–220, Mg$_{220}$, and NH$_3$BH$_3$–220, respectively.

Finally, RbBH$_4$ (Katchem, Prague, Czech Republic, 98%) and CsBH$_4$ (Katchem, Prague, Czech Republic, 98%) were mixed with NH$_3$BH$_3$ in the molar ratio of MBH$_4$–NH$_3$BH$_3$ (1:2) (M = Rb, Cs) and exposed to mechanochemical treatment similarly to the procedure described above for the Mg(BH$_4$)$_2$–NH$_3$BH$_3$ mixtures. The samples were ball milled for 2 min followed by 2 min break and this sequence was repeated 80 times.

2.2. In-house Powder X-ray Diffraction (PXD). PXD patterns of the as-prepared samples were measured in-house on a Rigaku Smart Lab diffractometer, using a Cu source and a parallel beam multilayer mirror (Cu K$_\alpha_1$ radiation, $\lambda = 1.540593$ Å, Cu K$_\alpha_2$ radiation, $\lambda = 1.544414$ Å). Data were collected in the 2θ range 8–60° at 5 deg/min, using a Rigaku D/tex detector. All air-sensitive samples were mounted in a glovebox in 0.5 mm glass capillaries sealed with glue.

2.3. Synchrotron Radiation Powder X-ray Diffraction (SR-PXD). In situ SR-PXD data were collected for s1 at the Swiss-Norwegian-Beamline (SNBL) at ESRF, Grenoble, France with a Pilatus area detector, $\lambda = 0.22568$ Å. The sample was packed in a glass capillary (i.d. 0.5 mm) and heated from room temperature to 100 °C (5 deg/min). Additionally, a PXD pattern was obtained at room temperature for s2 with use of the same setup. This data set was used for initial structural analysis. Subsequently, in situ SR-PXD data with higher resolution were collected for s6 at 111 at Diamond, Oxford, England with a MythenII detector, $\lambda = 0.82712$ Å. The sample was packed in a
glass capillary (i.d. 0.5 mm) and was heated from room temperature to 88 °C (10 deg/min). Sample compositions obtained by Rietveld refinement of PXD data only included the crystalline fraction of the sample, since an internal X-ray standard was not used.

2.4. Structure Solution and Refinement. The structure of the complex Mg(BH4)2(NH3BH3)2 was solved and refined from SR-PXD data measured at Diamond, UK (a detailed description is given in the Supporting Information). The final Rietveld refinement (Figure 1) indicated that the sample contains Mg(BH4)2(NH3BH3)2 (86 wt %), α-Mg(BH4)2 (10 wt %), and 4 wt % of NH3BH3.

![Figure 1. Rietveld refinement of SR-PXD data for Mg(BH4)2(NH3BH3)2 measured at room temperature (j = 0.822568 Å, sample s6). Tic marks α-Mg(BH4)2 (top), Mg(BH4)2(NH3BH3)2 and NH3BH3.](image)

The structure data from Mg(BH4)2(NH3BH3)2 were indexed with an orthorhombic unit cell, a = 14.41633(7) Å, b = 13.21283(7) Å, c = 5.11512(2) Å, and V = 974.331(8) Å³, using EXPO 2011.39 The structure was solved in the space group P212121 using global optimization in direct space implemented in the program FOX.40 The structural model (Table S1, Supporting Information) was refined by the Rietveld method, using the program Fullprof, and was checked for higher symmetry by using the ADDSYM routine in Platon.41,42 The final discrepancy factors are R_p = 0.66%, R_wp = 0.97% (not corrected for background), R_p = 14.1%, R_wp = 8.4% (conventional Rietveld R-factors), R_pseudo = 6.1%, and global χ² = 28.9.

2.5. Density Functional Theory Calculation. DFT calculations were further optimized to refine the structure from the SR-PXD data. The calculations were carried out by using the Vienna Ab-initio Simulation Package (VASP).43 For the exchange-correlation functional, the generalized-gradient approximation by Perdew, Burke, and Ernzerhof was adopted.44 Structural optimization was performed by using VASP projector augmented wave potentials45 Mg_pv (with semicore p states as valence electrons), B, N, and H until the force on each atom became smaller than 0.005 eV/Å. The plane-wave cutoff energy was set to 500 eV, and a Γ-centered k-point mesh of 2 × 2 × 6 was used. The lattice parameters were fixed to the experimental values since the optimization of the lattice parameters resulted in only 1% increase in volume. For the calculation of the electric field gradient potentials,46 harder potentials Mg_sv (with s and p semicore states), B_h, N_h, and H_h, and a higher cutoff energy of 750 eV were used. The atomic positions were not reoptimized.

2.6. Thermal Analysis and Mass Spectroscopy. Thermogravimetric analysis (TGA) was performed for s2 by using a PerkinElmer STA 6000 apparatus simultaneously with mass spectrometry (MS) analysis of the residual gas with use of a Hiden Analytical HPR-20 QMS sampling system. The samples (approximately 2 mg) were placed in an Al2O3 crucible and heated from 40 to 500 °C (2 deg/min) in an argon flow of 65 mL/min. The released gas was analyzed for hydrogen, ammonia, diborane, and borazine. Additionally, thermogravimetric analysis/differential scanning calorimetry (DSC) measurements were performed on s7 with Mettler Toledo instruments, TGA/sDTA 851e and DSC 821, and the heating rate of 5 deg/min from 25 to 500 °C. The samples for the TGA and DSC analysis were loaded in an argon glovebox into alumina crucibles with caps or sealed into aluminum pans, respectively. The experiments were performed with 10 mL/min nitrogen flow to prevent the oxidizing reactions.

2.7. Fourier Transform Infrared Spectroscopy (FTIR). α-Mg(BH4)2, NH3BH3, s2, and the samples heat treated to 220 °C were characterized by infrared absorption spectroscopy by using a Nicolet 380 FT-IR from Thermo Electron Corporation. The samples were exposed to air for approximately 15 s when transferring from the sample vial to the instrument.

2.8. Temperature-Programmed Photographic Analysis. Approximately 10 mg of Mg(BH4)2(NH3BH3)2 (s2) and NH3BH3 were sealed under argon in a glass tube placed in a home-built aluminum heating block as described recently.47 The samples were heated from 25 to 200 °C (heating rate 3 deg/min), while photos of the sample were collected every fifth second.

2.9. 11B NMR Measurements. Solid-state 11B MAS and multiple-quantum (MQ) MAS NMR experiments were performed on a Varian INOVA-300 spectrometer (7.05 T) with use of a home-built variable-temperature (VT) CP/MAS probe for 7 mm o.d. rotors and a home-built CP/MAS NMR probe for 5 mm o.d. rotors. The sample temperature in the NMR rotors was lowered by regulating the temperature of the air-bearing gas with an XRII851D00 Air-Jet sample cooling/heating system (FTS Systems, Stone Ridge, NY). The temperature gradient across the sample is less than 2 and 1 °C for the 7 and 5 mm PSZ rotors, respectively, and the actual sample temperatures were calibrated by using 27Pb MAS NMR of an external sample of Pb(NO3)2 under identical experimental conditions. Dry air (dew point below −60 °C) was used for the air-bearing and drive gases in all experiments. The VT 11B MAS NMR spectra following the decomposition of Mg(BH4)2(NH3BH3)2 were performed in situ, employing the 7 mm VT CP/MAS probe and a torlon end-cap with a small cylindrical hole (0.8 mm) throughout its center to allow release of gases during the gradual heating of the sample. The experiments used a 11B rf field strength of γ_B/2π = 40 kHz, 1H TPPM decoupling (γ_B/2π = 50 kHz), a short excitation pulse (τ_p = 0.5 μs), a relaxation delay of 4.0 s, and 256 scans for each spectrum. The 11B MAS NMR spectra of the central and satellite transitions and the 1H MQMAS NMR spectra were acquired at a sample temperature of roughly 31 °C, using the 5 mm CP/MAS probe with the stronger rf field strengths of γ_B/2π = 62 kHz and γ_B/2π = 70 kHz for 11B and 1H, respectively.
The MQMAS spectrum was obtained with the three-pulse, z-filter sequence, employing TPPM $^1$H decoupling during both the $t_1$ evolution and $t_2$ detection periods. A few $^{11}$B MAS NMR spectra were acquired at 14.09 T, using a home-built CP/MAS probe for 4 mm rotors, spinning speeds of $\nu_R = 1.00 - 12.0$ kHz, and a 0.5 μs excitation pulse for an rf field strength of $\nu_B / 2\pi = 60$ kHz. The determination of the $^{11}$B quadrupole coupling parameters ($\varepsilon_Q$ and $\eta_Q$ ) and isotropic chemical shifts ($\delta_{iso}$) was performed by least-squares fitting of simulated to experimental spectra for either the central transition or satellite transitions, using the STARS software package. The $^{11}$B isotropic chemical shifts are in ppm relative to neat $\text{F}_3\text{B-O(CH}_2\text{CH}_3)_2$.

3. RESULTS AND DISCUSSION

3.1. Synthesis of $\text{Mg(BH}_4)_2(\text{NH}_3\text{BH}_3)_2$ from $\alpha$-Mg(BH$_4$)$_2$; PXD patterns (Figure 2) have been collected for $\alpha$-Mg(BH$_4$)$_2$-

NH$_3$BH$_3$ (1:2) after 3 min of grinding (s5) and after various ball milling times (s2) showing the progress of the synthesis of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ ($d = 1.54056$ Å). Symbols: ▲, NH$_3$BH$_3$; ▲, α-Mg(BH$_4$)$_2$; (filled gray square), Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$; ?, 1.

NH$_3$BH$_3$ (1:2, s5) after 3 min of manual grinding and for $\alpha$-Mg(BH$_4$)$_2$—NH$_3$BH$_3$ (1:2, s2) after different BM times. The PXD patterns show reflections from the newly produced compound, Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ complex, and the unreacted starting materials, $\alpha$-Mg(BH$_4$)$_2$ and NH$_3$BH$_3$ for the s5 mixture after 3 min of manual grinding, demonstrating that a mechanochemical reaction readily takes place. After ball milling for 100 min the intensity of the reflections from NH$_3$BH$_3$ decreases, and after 325 min all reflections from Mg(BH$_4$)$_2$ and NH$_3$BH$_3$ have disappeared. Thus, only reflections from Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ and a new set of unidentified reflections, denoted 1 ($d = 6.890, 6.595, 5.075, 4.793, 4.712, 4.393, 4.352, 4.059, 3.678, 3.622, 3.889, 3.000, 2.553$, and 2.538 Å) are observed after mechanochemical treatment for 325 min. The reflections listed above may originate from another new compound, 1, in the system Mg(BH$_4$)$_2$—NH$_3$BH$_3$ which may be formed in a side reaction. The sample was stored at room temperature for 5 days before a SR-PXD pattern was collected (Figure S1, Supporting Information), which also includes a few new unidentified reflections, denoted 2 ($d = 6.617, 5.734, 5.093, 4.479, 4.332$, and 3.180 Å). The change in $d$-values for the unidentified reflections may reflect transformation from one unstable compound to another in the Mg(BH$_4$)$_2$—NH$_3$BH$_3$ system. A Rietveld refinement has been performed for compound 2 with the crystal structure of $\gamma$-Mg(BH$_4$)$_2$. There are significant shifts in the cell parameters for 2 ($a = 16.2234$ Å) as compared to $\gamma$-Mg(BH$_4$)$_2$ ($a = 15.7575$ Å), which may reflect that NH$_3$BH$_3$ is physisorbed in the porous structures of $\gamma$-Mg(BH$_4$)$_2$. However, when a guest molecule is adsorbed in the pores, the relative intensity between the peaks for $\gamma$-Mg(BH$_4$)$_2$ normally changes, which is not observed for the data obtained for 2. After 405 min of BM all reflections can be assigned to Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ reflecting a completed reaction according to reaction 1.

$$\text{Mg(BH}_4)_2(s) + 2\text{NH}_3\text{BH}_3(s) \rightarrow \text{Mg(BH}_4)_2(\text{NH}_3\text{BH}_3)_2(s)$$

Ball milling of the Mg(BH$_4$)$_2$—NH$_3$BH$_3$ (1:1) and (1:3) mixtures also produces Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ however, with an excess of either Mg(BH$_4$)$_2$ or NH$_3$BH$_3$. This is in contrast to the LiBH$_4$—NH$_3$BH$_3$ system, where either LiBH$_4$(NH$_3$BH$_3$) or LiBH$_4$(NH$_3$BH$_3$)$_0.5$ has been synthesized by ball milling depending on the ratio of LiBH$_4$ and NH$_3$BH$_3$. Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ transforms into a foam when it is stored at room temperature for several weeks (Figure S2, Supporting Information). Therefore, all samples containing Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ have been stored at $-35$ °C.

3.2. Synthesis of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ from Porous $\gamma$-Mg(BH$_4$)$_2$. The synthesis of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ by using the porous $\gamma$-Mg(BH$_4$)$_2$ polymorph rather than $\alpha$-Mg(BH$_4$)$_2$ as starting material has also been investigated. This work was prompted by the idea of adsorption of NH$_3$BH$_3$ in the porous structure. After manual grinding of $\gamma$-Mg(BH$_4$)$_2$—NH$_3$BH$_3$ (1:0.66, s6) for 10 min, 50 wt % conversion into the complex Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ is obtained (Figure S3, Supporting Information). After compressing the ground sample two times at 0.4 GPa, the resulting pellet contained more than 85 wt % of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ and ~12 wt % of NH$_3$BH$_3$ based on the detected content of crystalline components only (Figure S4, Supporting Information). Compression (0.4 GPa) of $\gamma$-Mg(BH$_4$)$_2$—NH$_3$BH$_3$ (1:2, s7) also produces Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ but with a larger excess of NH$_3$BH$_3$. The compressed samples also contain varying amounts of amorphous $\gamma$-Mg(BH$_4$)$_2$, observed as a broad hump in the PXD background centered at ~8°–12°. A mechanochemically treated sample of $\gamma$-Mg(BH$_4$)$_2$—NH$_3$BH$_3$ (1:2, s4) reveals crystalline Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ (95 wt %) and unreacted NH$_3$BH$_3$ (5 wt %), as obtained by Rietveld refinement (Figure S5, Supporting Information). The solid-state $^{11}$B MAS NMR spectrum of this sample (Figure S6, Supporting Information) is dominated by the resonances from Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$; however, resonances from unreacted NH$_3$BH$_3$ and amorphous Mg(BH$_4$)$_2$ have also been identified although they overlap with the peaks from the BH$_4$$^-$ sites of the Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ complex, respectively. This indicates that amorphous $\gamma$-Mg(BH$_4$)$_2$ does not react with NH$_3$BH$_3$. This hypothesis is verified by reacting amorphous Mg(BH$_4$)$_2$ with NH$_3$BH$_3$ (s8), which surprisingly does not result in any formation of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ as seen by Figure S7 (Supporting Information). This is a rare observation of different reactivity for the two polymorphs, $\gamma$-Mg(BH$_4$)$_2$ and $\alpha$-Mg(BH$_4$)$_2$. Thus, synthesis of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ from $\gamma$-Mg(BH$_4$)$_2$ appears not to be well reproducible as a result of a partial amorphization of the porous reactant.

To further investigate the unidentified compounds, 1 and 2, observed during synthesis from $\alpha$-Mg(BH$_4$)$_2$, and in an attempt...
Angles are more open for the shorter H···H distances involving the borohydride groups are slightly shorter than 2.0 Å. As expected, the N···B bonds are listed in Table S2 (Supporting Information). Only Mg···H bonds that coordinate Mg are more stretched by 0.02 Å on average than the other B···H bonds for both BH4 anions (corresponds to [110] and [110] directions).

The Mg2+ cation has a distorted tetrahedral coordination, with B–Mg–B angles ranging from 97.0(2)° to 134.1(2)°, similar to the range 92.1(1)–129.0(1)° found for α-Mg(BH4)2 and 100.4–129.6° found in γ-Mg(BH4)2. The MgH4 coordination polyhedron corresponds to a J84 Johnson solid, also known as a snub disphenoid. The same geometry of this coordination polyhedron has been observed in α- and γ-Mg(BH4)2 and is assumed to be important for the stability of these most common polymorphs. The Mg···H distances are not symmetrical, showing a span from 1.75 to 2.33 Å; however, this may be due to the low X-ray scattering power of H. The Mg···B distances involving the borohydride groups are slightly shorter than those for the BH3 groups in NH3BH3 and are equal to 2.41 and 2.48 Å. The Mg···N angles are 120.1(4)° and 114.0(4)°. The molecular complexes are linked in the crystal structure via dihydrogen bonds of the type N–H···B. The main geometrical parameters for the four strongest H···H bonds are listed in Table S2 (Supporting Information). Only one contact is shorter than 2.0 Å. As expected, the N···H···H angles are more open for the shorter H···H bonds, while the B···H···B angles show a large spread.

The DFT optimization starting from the refined PXD structure suggests only minor changes to the structure, confirming the structural model. The optimized Mg···H distances become more homogeneous, distributed between 1.971 and 2.074 Å. In accordance with the experimental structure, borohydride groups approach closer to Mg than the NH3BH3 groups. Thus, average Mg···B and Mg···H distances are 2.394 and 1.996 Å for BH4, and 2.469 and 2.064 Å for NH3BH3. An overview with selected bond lengths and fractional atomic coordinates from the DFT-optimized structure are given in Tables S3 and S4 (Supporting Information), respectively. The B···H bonds that coordinate Mg are more stretched by 0.02 Å on average than the other B···H bonds for both BH4 and NH3BH3.

The DFT-optimized H positions reveal a stronger dihydrogen bonding network than what is expected from the experimental structure. We note that the short dihydrogen bondings always appear intermolecular and form across different molecular complexes, and not between BH4 and NH3BH3 within the same complex. Each neutral molecular complex [Mg(BH4)2(NH3BH3)]2− is linked to the neighboring ones via four strong H···H bonds shorter than 2.0 Å (symmetrically equivalent pairs with distances of 1.957 and 1.995 Å). When the cutoff distance is increased to 2.1 Å, the number of short bondings increases from four to ten. This is in contrast to a previous study where the stronger dihydrogen bonds were observed within one complex between adjacent BH4 and NH3BH3 (2.21 to 2.27 Å) and the weaker between neighboring complexes (2.24 to 2.39 Å). This is the main difference between the structure solutions presented here and in a previous study. In the earlier investigation the unit cell parameters are reported to be a = 14.4135(2) Å, b = 13.2084(2) Å, and c = 5.1118(1) Å, while a = 14.41633(7) Å, b = 13.21283(7) Å, and c = 5.11512(2) Å are determined in this work.

The mononuclear molecular structure of Mg(BH4)2(NH3BH3)2 resembles those of Al(BH4)3 and Zr(BH4)3, where the borohydride groups act as terminal ligands. However, the borohydride groups share a face (η3-coordination) for Zr(BH4)3 and an edge with Mg (η6) in Mg(BH4)2(NH3BH3)2. The three other metal borohydride complexes with NH3BH3, known so far, show strikingly different connectivities, where the BH4− groups and the NH3BH3 molecules act as bridging ligands. In Ca(BH4)2(NH3BH3)2 the borohydride groups are bridging two metal atoms, forming a 2D square net. The NH3BH3 molecules behave as terminal ligands, completing the coordination sphere of Ca to octahedral.
LiBH₄(NH₃BH₃)₀.₅, the borohydride groups are bridging three metal atoms and the NH₃BH₃ molecules are coordinated by two metals atoms, making a 3D polymeric framework. The LiBH₄(NH₃BH₃)₂ structure appears to be very complex, containing four independent borohydride anions and four NH₃BH₃ molecules. While the borohydride anions are coordinated to 2–4 metal atoms, the NH₃BH₃ molecules act as a terminal ligand or even stay isolated from metal atoms, forming a cocrystal. Thus, Mg(BH₄)₂(NH₃BH₃)₂ is the only known molecular complex of ammonia borane, having a potential to show a relatively high vapor pressure at temperatures close to ambient.

3.4. ¹¹B MAS and MQMAS NMR of Mg(BH₄)₂(NH₃BH₃)₂. The ¹¹B MAS NMR spectrum of the central transition region for Mg(BH₄)₂(NH₃BH₃)₂ (Figure 4a, sample s2) clearly reveals the presence of two distinct BH₄⁻ sites, as observed by the narrow resonances at −40 and −42 ppm. Moreover, centerband resonances exhibiting second-order quadrupolar line shapes are observed in the range −24 to −34 ppm. These line shapes are removed in the isotropic (F1) dimension of the MQMAS NMR spectrum (Figure 5), which unambiguously shows the presence of two distinct sites in the −24 to −34 ppm spectral region. Thus, this region is analyzed by spectral simulations/optimizations by using two overlapping quadrupolar line shapes, assigned to the two distinct NH₃BH₃ sites. This results in the optimized ¹¹B isotropic chemical shifts (δiso) and quadrupole coupling parameters (CQ and ηQ) listed in Table 2 and illustrated by the simulations in Figure 4c.d.

Figure 4. (a) ¹¹B MAS NMR spectrum (7.05 T, νR = 9.0 kHz) of the central transitions for Mg(BH₄)₂(NH₃BH₃)₂ (s2), obtained at a sample temperature of 31 °C. (b) Optimized simulated spectrum for the central transitions, employing the δiso, CQ, and ηQ parameters listed in Table 2. Parts c and d illustrate simulated second-order quadrupolar line shapes for the B(13)–NH₃BH₃ and B(21)–NH₃BH₃ sites, respectively, as obtained from least-squares fitting to the spectrum in part a.

Figure 5. ¹¹B MQMAS NMR spectrum (7.05 T, νR = 9.0 kHz) of Mg(BH₄)₂(NH₃BH₃)₂ (s2), obtained at a sample temperature of 31 °C using the three-pulse z-filter sequence, 192 t₁ increments, a 3 s relaxation delay, 72 scans for each increment, and ¹H decoupling during both the t₁ and t₂ periods. The projections onto the F1 and F2 axes represent summations over the 2D spectrum. The asterisks indicate spinning sidebands for the two BH₄⁻ sites and the diamonds artifacts arising around the transmitter frequency in the F1 dimension.

Accurate values for δiso, CQ, and ηQ cannot be obtained by line shape analysis of the central transitions for the two BH₄⁻ sites since these display Gaussian-like line shapes. However, two distinct manifolds of spinning sidebands (ssb’s) from the satellite transitions are observed for the two BH₄⁻ sites, as illustrated in Figure 6a. Least-squares analyses to the intensities and partly overlapping line shapes for the two manifolds allow determination of the quadrupole interaction parameters with high precision (Table 2). This is also apparent from the simulation spectrum of the ssb manifolds for the two BH₄⁻ sites only in Figure 6b, which matches very well the experimental intensities and line shapes. The high precision of the δiso, CQ, and ηQ parameters determined for the two NH₃BH₃ sites and two BH₄⁻ sites are further supported by the isotropic triple-quantum shifts (δ3Qiso, Table 2) observed in the ¹¹B MQMAS NMR spectrum (Figure 5). These shifts depend on the δiso, CQ, and ηQ parameters, following the relationship

$$δ_{3Q}^{calc} = \frac{17}{8} δ_{iso} + \frac{1}{32} \frac{C_Q^2(1 + \eta_Q^2/3)}{ν_L^2} × 10^6$$

(2)

The calculated values (δ3Qcalc), using the parameters determined from the ¹¹B MAS NMR spectra (Figures 4a and 6a) and listed in Table 2, match fully the experimental values within their error limits. Finally, Figure 4b illustrates a full simulation of the central transition region, using the ¹¹B NMR parameters in Table 2 and relative intensities for the individual sites in the range 0.90–1.05. In the ¹¹B MQMAS spectrum, the intensities for the NH₃BH₃ sites are significantly smaller than those for the two BH₄⁻ sites, which reflect the reduced efficiency of triple-quantum (3Q) excitation and triple-quantum to single-quantum conversion for ¹¹B sites with large quadrupole couplings.
Table 2. $^{11}$B Quadrupole Coupling Parameters ($C_Q, \eta_Q$), Isotropic Chemical Shifts ($\delta_{iso}$), and Triple-Quantum Chemical Shifts ($\delta_{3Q}$) Determined for Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ at 31 °C.

<table>
<thead>
<tr>
<th>site $^{b}$</th>
<th>$\delta_{aa}$ (ppm)</th>
<th>$C_Q$ (MHz)</th>
<th>$\eta_Q$</th>
<th>$\delta_{3Q}^{aa}$ (ppm)</th>
<th>$\delta_{3Q}^{ab}$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B(2)−BH$_4^-$</td>
<td>−41.8 ± 0.1</td>
<td>0.409 ± 0.020</td>
<td>0.71 ± 0.03</td>
<td>−88.1 ± 0.2</td>
<td>−88.1</td>
</tr>
<tr>
<td>B(7)−BH$_4^-$</td>
<td>−39.5 ± 0.2</td>
<td>0.568 ± 0.020</td>
<td>0.73 ± 0.03</td>
<td>−82.7 ± 0.2</td>
<td>−82.6</td>
</tr>
<tr>
<td>B(13)−NH$_3$BH$_3$</td>
<td>−23.4 ± 0.3</td>
<td>1.419 ± 0.030</td>
<td>0.35 ± 0.05</td>
<td>−42.5 ± 0.2</td>
<td>−42.6</td>
</tr>
<tr>
<td>B(21)−NH$_3$BH$_3$</td>
<td>−24.7 ± 0.3</td>
<td>1.221 ± 0.025</td>
<td>0.38 ± 0.02</td>
<td>−47.1 ± 0.2</td>
<td>−47.3</td>
</tr>
</tbody>
</table>

The quadrupole coupling and chemical shift parameters are determined from optimization to spectra of the central transition and the satellite transitions for the NH$_3$BH$_3$ and BH$_4^-$ sites, respectively. The quadrupole coupling parameters are $C_Q = \epsilon Q V_{zz}/h$, $\eta_Q = (V_{yy} - V_{xx})/(|V_{zz}|)$, where $V_i$ are the principal elements of the electric field gradient tensor, following the convention: $V_{xx} \geq V_{yy} \geq V_{zz}$. Assignment to the different crystallographic B sites from the SR-PXD structure determination (Table S1, Supporting Information) based on the calculated $^{11}$B electric field gradient tensors by DFT in Table 3. The triple-quantum shifts are calculated from $\delta_{iso}$, $C_Q$, and $\eta_Q$ using the expression in eq 2.

\[
Q_{zz}^{exp} = C_Q,
\]

\[
Q_{yy}^{exp} = -\frac{1}{2}(1 - \eta_Q)C_Q,
\]

\[
Q_{zz}^{exp} = -\frac{1}{2}(1 + \eta_Q)C_Q.
\]

The sign of $C_Q$ cannot be determined from single-pulse NMR experiments and thus it is assumed that the experimental $C_Q$ values exhibit the same sign as the corresponding $V_{zz}^{calc}$ elements from the DFT calculations. The calculated $^{11}$B EFG's and quadrupole coupling parameters from the DFT calculations before and after a DFT optimization of the structure determined from SR-PXD (Table S1, Supporting Information) are listed in Table 3. Both of these sets of calculated $^{11}$B EFG tensor elements result in the same assignment of the observed $^{11}$B quadrupole coupling parameters (Table 2) to the B(2), B(7), B(13), and B(21) sites in the crystal structure for Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ (Table S1, Supporting Information).

Using this assignment, and assuming that the experimental quadrupole tensor elements ($Q^{exp}$) exhibit the same sign as the calculated $V_{zz}^{calc}$ elements, results in the plot of $Q_{zz}^{exp}$ as a function of the $V_{zz}^{calc}$ elements after the DFT optimization of the Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ structure shown in Figure 7. This plot shows a convincing correlation between the experimental and calculated tensor elements and linear regression analysis of the data gives the equation

\[
Q_{zz}^{exp} = C_Q^{calc},
\]

\[
Q_{yy}^{exp} = -\frac{1}{2}(1 - \eta_Q^{calc})C_Q^{calc},
\]

\[
Q_{zz}^{exp} = -\frac{1}{2}(1 + \eta_Q^{calc})C_Q^{calc}.
\]
borane sites, in particular B(13), as evidenced by the calculated $C_Q$ and $\eta_Q$ values in Table 3. For the B(13) site, the asymmetry parameter changes from $\eta_Q^{\calc} = 0.72$ to $\eta_Q^{\calc} = 0.50$ after the DFT optimization, where the latter value is much closer to the experimental value of $\eta_Q = 0.35$. For the B(21) site the two calculated asymmetry parameters ($\eta_Q^{\calc} = 0.34$ and 0.40) agree well with the experimental value (0.38) whereas $C_Q$ changes from 1.34 to 1.24 MHz upon the DFT optimization, where the latter value is in full agreement with the experimental value, $C_Q = 1.221 \pm 0.025$ MHz. Thus, the comparison of the $^{11}$B NMR data with the EFG tensor elements (Tables 2 and 3) indicates that crystal structure data may give a slightly improved representation of the crystal structure as compared to the SR-PXD structure data (Table S1, Supporting Information), principally for the local environments of the two ammonia borane sites. Finally, it is noted that the slopes of the linear correlations in eqs 4 and 5 correspond to $^{11}$B nuclear quadrupole moments of $Q = 0.0361$ barn (after DFT optimization) and $Q = 0.0344$ barn (SR-PXD structure). These values are slightly smaller than the value $Q = 0.0409$ barn, reported in an earlier $^{11}$B NMR–DFT study of inorganic borates, and the value $Q = 0.0406$ barn, reported in recent compilations of nuclear quadrupole coupling moments.

3.5. Thermal Analysis. TGA-MS data are obtained for $\text{Mg(BH}_4\text{)}_2(\text{NH}_3\text{BH}_3)_2$ (s2) from room temperature to 400 °C (2 deg/min) and presented in Figure 8. TGA reveals a mass loss of 27 wt % between room temperature and 125 °C and an additional mass loss of 10 wt % in the temperature range from 125 to 400 °C. The differentiated TGA data (dTGA) indicate that a two-step reaction is taking place between room temperature and 125 °C. From the MS signals a mixture of hydrogen, ammonia, and a small amount of diborane and borazine are observed below 150 °C, while only hydrogen is observed above 150 °C. The TGA signal is remarkably different from those observed for both pristine NH$_3$BH$_3$ and Mg(BH$_4$)$_2$. NH$_3$BH$_3$ decomposes in two distinct steps between 100 and 175 °C, while pristine Mg(BH$_4$)$_2$ decomposes at ~365 °C. The TGA/DSC data measured for γ-$\text{Mg(BH}_4\text{)}_2$–

![Figure 7. Correlation between the principal $^{11}$B quadrupole coupling tensor elements ($Q^{\exp}$) and the corresponding calculated EFG tensor elements ($V^{\calc}$) from the DFT optimization of the SR-PXD structure of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$. The calculated EFG tensor elements are summarized in Table 3. Linear regression analysis of the of the data is given in eq 4](image)

$$Q^{\exp}(\text{MHz}) = 0.0872(\text{MHz}/(\text{V }\text{Å}^2))V^{\calc}(\text{V }\text{Å}^2) - 7.3 \times 10^{-6}(\text{MHz}) \quad (4)$$

with the correlation coefficient $R = 0.991$. A similar analysis of the $Q^{\exp}$ data as a function of $V^{\calc}$ elements before the DFT optimization (i.e., the PXD data in Table 3) results in the equation

$$Q^{\exp}(\text{MHz}) = 0.0831(\text{MHz}/(\text{V }\text{Å}^2))V^{\calc}(\text{V }\text{Å}^2) - 4.7 \times 10^{-9}(\text{MHz}) \quad (5)$$

and the correlation coefficient $R = 0.987$. Thus, a slightly improved agreement is observed between the experimental quadrupole tensor elements and those calculated after the DFT optimization of the structure. The largest difference between the two sets of calculated $^{11}$B EFG data and their correlation with the experimental data are observed for the two ammonia borane sites, in particular B(13), as evidenced by the calculated $C_Q$ and $\eta_Q$ values in Table 3. For the B(13) site, the asymmetry parameter changes from $\eta_Q^{\calc} = 0.72$ to $\eta_Q^{\calc} = 0.50$ after the DFT optimization, where the latter value is much closer to the experimental value of $\eta_Q = 0.35$. For the B(21) site the two calculated asymmetry parameters ($\eta_Q^{\calc} = 0.34$ and 0.40) agree well with the experimental value (0.38) whereas $C_Q$ changes from 1.34 to 1.24 MHz upon the DFT optimization, where the latter value is in full agreement with the experimental value, $C_Q = 1.221 \pm 0.025$ MHz. Thus, the comparison of the $^{11}$B NMR data with the EFG tensor elements (Tables 2 and 3) indicates that crystal structure data may give a slightly improved representation of the crystal structure as compared to the SR-PXD structure data (Table S1, Supporting Information), principally for the local environments of the two ammonia borane sites. Finally, it is noted that the slopes of the linear correlations in eqs 4 and 5 correspond to $^{11}$B nuclear quadrupole moments of $Q = 0.0361$ barn (after DFT optimization) and $Q = 0.0344$ barn (SR-PXD structure). These values are slightly smaller than the value $Q = 0.0409$ barn, reported in an earlier $^{11}$B NMR–DFT study of inorganic borates, and the value $Q = 0.0406$ barn, reported in recent compilations of nuclear quadrupole coupling moments.

3.5. Thermal Analysis. TGA-MS data are obtained for $\text{Mg(BH}_4\text{)}_2(\text{NH}_3\text{BH}_3)_2$ (s2) from room temperature to 400 °C (2 deg/min) and presented in Figure 8. TGA reveals a mass loss of 27 wt % between room temperature and 125 °C and an additional mass loss of 10 wt % in the temperature range from 125 to 400 °C. The differentiated TGA data (dTGA) indicate that a two-step reaction is taking place between room temperature and 125 °C. From the MS signals a mixture of hydrogen, ammonia, and a small amount of diborane and borazine are observed below 150 °C, while only hydrogen is observed above 150 °C. The TGA signal is remarkably different from those observed for both pristine NH$_3$BH$_3$ and Mg(BH$_4$)$_2$. NH$_3$BH$_3$ decomposes in two distinct steps between 100 and 175 °C, while pristine Mg(BH$_4$)$_2$ decomposes at ~365 °C. The TGA/DSC data measured for γ-$\text{Mg(BH}_4\text{)}_2$–

![Figure 8. TGA-MS curves for Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ (s2) heated from room temperature to 400 °C (2 °C/min). Upper part: The TGA data and dTGA are presented by a solid and dashed black line, respectively. Lower part: Corresponding MS signals shown for hydrogen, ammonia, borazine, and diborane presented by green, purple, red, and blue lines, respectively.](image)
NH₃BH₃ (1:2, s7) from room temperature to 500 °C (5 deg/min) are shown in Figure S16 (Supporting Information). This sample contains some unreacted NH₃BH₃ and amorphous Mg(BH₄)₂, as described previously, and the TGA data (Figure S16, Supporting Information) reveal a two- or three-step decomposition from 80 to 170 °C as a result of the presence of NH₃BH₃. The DSC data reveal endothermic peaks at 48 and 85 °C and a strong exothermic peak at 100 °C. The two events at 85 and 100 °C cannot be clearly examined due to the presence of both NH₃BH₃ and Mg(BH₄)₂(NH₃BH₃)₂. However, the endothermic peak at 48 °C is described as the melting point of Mg(BH₄)₂(NH₃BH₃)₂ which is significantly lower than the melting point for NH₃BH₃ (100–114 °C).¹⁹

To confirm the melting point of Mg(BH₄)₂(NH₃BH₃)₂ a series of photographs of the powder of Mg(BH₄)₂(NH₃BH₃)₂ were collected between room temperature and 200 °C and compared with those of NH₃BH₃ (Figure 9). As expected, no changes are observed for the powder of NH₃BH₃ between room temperature and 100 °C, while the powder expands significantly in the temperature range from 108 to 112 °C, as previously reported.¹⁹ In contrast, Mg(BH₄)₂(NH₃BH₃)₂ melts and at 53 °C the powder is liquefied. Above the melting point the sample starts to foam and at 70 °C it starts to expand slowly. When temperatures around 110 °C are reached, it expands significantly faster similar to the observations for pristine NH₃BH₃. A fraction of Mg(BH₄)₂(NH₃BH₃)₂ was heated to 55 °C in argon atmosphere, cooled to room temperature, and characerized by PXD (Figure S17, Supporting Information). It is clearly revealed that the crystal structure of Mg(BH₄)₂(NH₃BH₃)₂ is remained, confirming that it is a melt.

In situ SR-PXD diffractograms (Figure S18, Supporting Information) have been obtained for the sample γ-Mg(BH₄)₂−NH₃BH₃ (1:0.66, s6) heated from room temperature to 88 °C (10 deg/min). These data reveal that all diffraction peaks from Mg(BH₄)₂(NH₃BH₃)₂ disappear at T = 45 °C and no new peaks appear, confirming the melting state. The activation energy for hydrogen release from Mg(BH₄)₂(NH₃BH₃)₂ was calculated in a previous study.²⁰ However, in that work they are not aware of the fact that Mg(BH₄)₂(NH₃BH₃)₂ decomposes from a melt, which then may lead to an inaccurate value for the activation energy.

3.6. Variable-Temperature ¹¹B MAS NMR. Variable-temperature (VT) ¹¹B MAS NMR spectra of the central transition region for Mg(BH₄)₂(NH₃BH₃)₂ following a stepwise heating of the sample from 3.9 to 73.1 °C is shown in Figure 10. The spectra acquired at 3.9 and 33.2 °C resemble closely the spectrum in Figure 4a, thereby reflecting the presence of two well-defined NH₃BH₃ and two BH₄⁻ sites. At 40 °C the overlapping quadrupolar line shapes from the NH₃BH₃ sites become broader and intensity from the low-frequency BH₄⁻ site converts partly into a centerband at ~39.5 ppm. The broadening of the quadrupolar line shapes reflects the formation of a new NH₃BH₃ site with a chemical shift shifted to higher frequency, as apparent from the spectrum recorded at 45.6 °C. At 48.2 °C the quadrupolar line shapes from the two distinct NH₃BH₃ sites have transformed into this new NH₃BH₃ environment, corresponding to the almost symmetric and rather narrow centerband with a center of gravity at δg = −22.7 ppm. The decrease in line width of this centerband may reflect that the new NH₃BH₃ site is influenced by motional processes that partially average out the quadrupole interaction. A similar resonance at δ = −22.5 ppm has been observed in VT-¹¹B MAS NMR spectra following the dehydrogenation of NH₃BH₃ at 88 °C and assigned to a new NH₃BH₃ phase.¹⁸ A structural change is also observed for the two distinct centerbands from the BH₄⁻ sites which at 48.2 °C have merged into a narrow resonance at δg = −38.9 ppm. These changes for the NH₃BH₃ and BH₄⁻ sites may reflect that the increase in temperature reduces the hydrogen bonding network, thereby allowing a higher degree of motion of the structural units, resulting in averaging of the quadrupole interactions for these sites. The rather narrow resonances and symmetric line shapes observed at 62.4 °C indicate a nearly complete averaging of the ¹¹B quadrupole interactions in accordance with the presence of a melt phase with a high viscosity. A further increase in temperature leads to a decrease in the intensity for the NH₃BH₃ site that is almost absent at 73.1 °C. Moreover, a minor fraction of a new type of boron species is observed at 62.4 and 68.6 °C by the presence of a centerband at δg = −11.7 ppm. This resonance may arise from new BH₄⁻ sites,²⁰ and it broadens significantly at 73.1 °C where the spectrum otherwise only includes the resonance from the BH₄⁻ units. The centerband from the latter species shifts slightly to higher frequency on going from 68.6 to 73.1 °C (δg = −38.2 ppm). The main change observed in the VT-¹¹B MAS NMR spectra from 40.3 to 48.2 °C is in accord with the endothermic peak at 48 °C in the DSC experiment and reflects the breakdown of the Mg(BH₄)₂(NH₃BH₃)₂ structure. Moreover, the overall decrease in the ¹¹B centerband intensity on going from 48.2 to 73.1 °C, in particular for the (NH₃BH₃)₂ sites,

Figure 9. Temperature-programmed photographic analysis of NH₃BH₃ (upper row) and Mg(BH₄)₂(NH₃BH₃)₂ (lower row) heated from room temperature to 200 °C with a heating rate 4 deg/min in argon atmosphere.
subsequently characterized by PXD (Figure S19, Supporting Information) of Mg-BH$_4$$_2$ (NH$_3$BH$_3$)$_2$. FTIR reveals one stretch in the N–H region at 3440 cm$^{-1}$ for s$_2$-220 and two stretches in the B–H region at 2494 and 2274 cm$^{-1}$. The Mg-220 sample shows bands similar to α-Mg(BH$_4$)$_2$, while NH$_3$BH$_3$-220 reveals the B–H (2494 cm$^{-1}$) and N–H (3440 cm$^{-1}$) stretches, which are the characteristic bands from polyimidoborane, [NHBBH]$_n$. It is clearly seen that the data for s$_2$-220 can be considered as a sum of s$_2$-220 and NH$_3$BH$_3$-220, revealing that the products present at 220 °C are Mg(BH$_4$)$_2$ and [NHBBH]$_n$. This is confirmed by a $^{11}$B MAS NMR spectrum of Mg-220, (Figure 12) that includes a partly resolved second-order quadrupolar line shape from [NHBBH]$_n$ ($\delta_q = 22.8$ ppm) and a narrow resonance at $-40$ ppm from the BH$_4^-$ sites in Mg(BH$_4$)$_2$. The quadrupolar line shape is in accordance with the parameters $\delta_{iso} = 30.1$ ppm, $\eta_Q = 0.17$, determined for [NHBBH]$_n$ from a $^{11}$B MAS NMR spectrum obtained at lower magnetic field (7.05 T). An evaluation of the centerband intensities gives a molar ratio of 0.44:1.0 between the $^{11}$B sites in [NHBBH] and Mg(BH$_4$)$_2$.

### 3.8. Discussion

To summarize the thermal decomposition, Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ melts at ~48 °C followed by decomposition in the temperature range 75 to 125 °C releasing a gas mixture of H$_2$, NH$_3$, B$_2$H$_6$, and B$_3$N$_3$H$_6$. Hydrogen (10 wt %) is slowly released in the temperature range from 125 to 400 °C, and α- and β'-Mg(BH$_4$)$_2$ as well as [NHBBH]$_n$ are present in the sample at 220 °C. The melting point at ~48 °C is on the one hand lower than the melting point for NH$_3$BH$_3$ (~112 °C), but is on the other hand remarkably higher than melting points for molecular metal borohydrides such as Al(BH$_4$)$_3$ (<0 °C). This is explained by the strong dihydrogen bonds revealed by the DFT-optimized structural model. The thermal decomposition of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ is different from the previously reported M(BH$_4$)$_2$–NH$_3$BH$_3$ (M = Li, Na, K, or Ca) systems. For the system MBH$_4$–2NH$_3$BH$_3$ (M = Na, K), NH$_3$BH$_3$ decomposes independently of the presence of the metal borohydride, while LiBH$_4$(NH$_3$BH$_3$)$_0.5$ and Ca(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ decompose in steps similar to those for the individual components, i.e., first decomposition of NH$_3$BH$_3$ and at higher temperatures decomposition of the metal borohydrides. The difference is explained by the different crystal structures for LiBH$_4$NH$_3$BH$_3$, LiBH$_4$(NH$_3$BH$_3$)$_0.5$, Ca(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$, and Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$. Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ is the only compound that melts, most likely as a result of the presence of molecular complexes of Mg(BH$_4$)$_2$(NH$_3$BH$_3$)$_2$ formed by Mg coordination to both BH$_4^-$ and NH$_3$BH$_3$ groups as terminal ligands.

For the systems M(BH$_4$)$_2$–NH$_3$BH$_3$ (M = Li, Na, K, Mg or Ca), NaBH$_4$ and KBH$_4$ are the only metal borohydrides that do not react with NH$_3$BH$_3$ forming borohydride–ammonia borane complexes. This may be correlated with the decreased polarization power of the metal cations for MBH$_4$ (M = Na, K) as compared with those for M(BH$_4$)$_2$ (M = Li, Mg, Ca). To investigate this further, RbBH$_4$–NH$_3$BH$_3$ (1:2) and CsBH$_4$–NH$_3$BH$_3$ (1:2) were treated mecanochemically and subsequently characterized by PXD (Figure S21, Supporting Information). The PXD data reveal for both samples that all diffraction peaks can be assigned to the starting materials, NH$_3$BH$_3$ and RbBH$_4$ or CsBH$_4$. This shows that the RbBH$_4$–NH$_3$BH$_3$ (1:2) and CsBH$_4$–NH$_3$BH$_3$ (1:2) systems do not react during mecanochemical treatment similar to NaBH$_4$–NH$_3$BH$_3$ (1:2) or KBH$_4$–NH$_3$BH$_3$ (1:2), as expected.
according to the decreased polarization power. A similar trend has recently been observed for the dehydrogenation effect of M(BH$_4$)$_n$ (M = Li, Na, Mg, Ca) on Al(BH$_4$)$_3$·6NH$_3$. In that study, an increased hydrogen purity and lowered decomposition temperature was found for Al(BH$_4$)$_3$·6NH$_3$ as a function of increasing polarization power for the added metal borohydride.

The systems M(BH$_4$)$_n$·NH$_3$BH$_3$ (M = Li, Na, K, Mg, Ca) have in common that they do not suppress the release of the toxic gases, e.g., B$_2$H$_6$ and B$_3$N$_3$H$_6$, as observed for metal amidoboranes. Thus, it may be concluded that the bonding of NH$_3$BH$_3$ is insufficiently modified for MBH$_4$·NH$_3$BH$_3$ (M = Li, Na, K, Mg, Ca), because NH$_3$BH$_3$ maintains its molecular form. This is in contrast to metal amidoboranes where a metal is substituted for a hydrogen atom. Thus, the lack of pure hydrogen release from MBH$_4$·NH$_3$BH$_3$ systems must be addressed to utilize borohydride–ammonia borane systems for hydrogen storage in the future. Several attempts to achieve this goal may be possible, e.g., nanoconfinement, the addition of catalysts or by development of composite systems including metal hydrides such as Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$→LiH or Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$→MgH$_2$, which all have been successful treatments for NH$_3$BH$_3$.

4. CONCLUSION

In summary, Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$ has been mechanochemically synthesized from α- or γ-Mg(BH$_4$)$_2$·NH$_3$BH$_3$ (1:2). The crystal structure has been solved from SR-PXD data and subsequently confirmed by DFT calculations. $^{11}$B quadrupole coupling parameters and isotropic chemical shifts have been determined with high precision for the two ammonia borane sites and the two borohydride sites from $^{11}$B MAS NMR of the central and satellite transitions, respectively, and from $^{11}$B MQMAS NMR. The quadrupole coupling parameters have been compared with $^{11}$B electric field gradient tensor elements, obtained by DFT calculations, and thereby used to validate the crystal structures from SR-PXD and the subsequent DFT optimization. This comparison reveals that the structure gives the best agreement with the $^{11}$B NMR data, most likely reflecting an improved description of the local environments for the two ammonia borane sites. The strong intermolecular dihydrogen bonds revealed by the optimized structural model may explain the relatively high melting point, 48 °C, for Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$, as compared to those for other molecular metal borohydrides. Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$ has a high gravimetric and volumetric H$_2$ content ($\rho_m = 17.4$ wt % H$_2$, $\rho_V = 137$ g H$_2$/L), which makes it a good candidate for a new hydrogen storage material. Furthermore, Mg(BH$_4$)$_2$·(NH$_3$BH$_3$)$_2$ has an attractive decomposition temperature for practical use, but suffers from the lack of pure...
hydrogen release, which needs to be addressed in future work. Metal hydrides with low melting points are of interest for safe, fast, and efficient handling, e.g., refueling.

ASSOCIATED CONTENT
2 Supporting Information
Atomic coordinates and bond lengths in Mg-(BH4)2(NH2BH3)2. Rietveld refinement plots, and supporting PXD, FTIR and 11B NMR data. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION
The authors declare no competing financial interest.

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