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Guanidinium octahydrotriborate: an ionic liquid with high hydrogen storage capacity

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**Publication Details**

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Abstract
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Keywords
capacity, storage, liquid, hydrogen, ionic, high, guanidinium, octahydrotriborate

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Guanidinium octahydrotriborate: An ionic liquid with a high hydrogen capacity

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For chemical hydrogen storage, capacity is one key criterion that has spurred intense efforts to investigate compounds with high hydrogen content. The guanidinium cation and the octahydrotriborate anion possess 6 H⁺ and 8 H⁻, respectively. The combination of these two ions yields guanidinium octahydrotriborate with 13.8 wt.% hydrogen. This paper presents its facile synthesis, as confirmed by ¹¹B and ¹H nuclear magnetic resonance spectroscopy. The results show that guanidinium octahydrotriborate is an ionic liquid with a melting point below –10 °C, which makes it a possible injectable/pumpable hydrogen carrier. It decomposes selectively to hydrogen, in stark contrast to the formation of various boranes from related solid octahydrotriborates. The much improved H₂ purity can be ascribed to the more effective combination of H⁺ and H⁻, and the higher H⁺/H⁻ ratio in the liquid guanidinium octahydrotriborate.

Introduction

Hydrogen has been considered as a clean and efficient energy carrier to store electricity generated by wind and solar farms and to deliver the energy upon demand. The current means of storing hydrogen, however, are either not safe, not compact, or too energy intensive. These disadvantages constitute a big barrier to the wide deployment of devices powered by hydrogen, in particular, where both volumetric and gravimetric densities are critical. Material-based hydrogen storage has thus been intensively studied, since it outperforms traditional storage via high pressure or cryogenic tanks. Among all the materials studied, hydrogen-rich alanates, amides, and borohydrides consisting of light elements have attracted particularly intense attention. Due to the light atomic weight, boron-nitrogen-hydrogen-containing compounds (abbreviated as BNH) could also possess remarkably high hydrogen content. Alkali/alkaline metal borohydrides and ammonia borane have therefore been extensively investigated. Despite their intrinsic high hydrogen content, they suffer from serious problems, such as slow kinetics, problematic by-products, or unfavourable thermodynamics, which make them unsuitable for practical application.

Strategies such as introducing additives and making derivatives have been employed to suppress the formation of impurities and/or enhance the kinetics. For example, replacing H atom on the N by Li/Na or Ca can effectively suppress the formation of by-products from ammonia borane, and introducing MgH₂ can effectively destabilize LiBH₄ and improve its dehydrogenation kinetics. Its performance, however, still falls short of the criteria. New BNH compounds have thus been synthesized and screened. BNH compounds feature protic H (H⁺) on N and hydridic H (H⁻) on B atoms, which normally results in an extensive dihydrogen bond network throughout the crystal structure. The formation of H₂ via the combination of H⁺ and H⁻ features a high enthalpy (H⁺ + H⁻ = H₂, ΔH = −17.37 eV), which is considered as one of the driving forces of dehydrogenation. To take advantage of these H⁺ --- H⁻ interactions, H⁺ containing units have been introduced to destabilize borohydrides that contain four H⁻ atoms. Several borohydride/ammonia and borohydride/amide systems have thus been developed. Other N–H containing units such as guanidinium (C(NH₂)₃) have also been employed. The interaction between H⁺ and H⁻ observed in guanidinium borohydride leads to a low activation energy pathway for H₂ evolution. The excess H⁺ in guanidinium borohydride can be offset by introducing LiBH₄ or Ca( BH₄)₂, and these have been reported to suppress the formation of ammonia from pristine guanidinium borohydride.

Compared with the well-studied borohydrides (also known as tetrahydroborate (with BH₄⁻)), octahydrotriborates (with B₃H₈⁻) have recently attracted considerable interest. They have been proposed as an intermediate during the thermal decomposition of borohydrides. Sodium octahydrotriborate has been reported to outperform the most studied NaBH₄ and NH₃BH₃ for hydrogen storage via hydrolysis. Octahydrotriborate is also used to make ammonia triborane (NH₃B₃H₇), which has shown exceptional hydrolytic properties.

For practical applications, solid hydrogen storage materials are inferior to liquid ones with respect to the control of hydrogen supply upon demand. Liquid compounds are very compati-
ble with the current liquid fuel distribution techniques, which can offer precise control of hydrogen release in terms of rates and quantity. Their sluggish mass transfer and poor heat dissipation make the control of hydrogen release from solid materials very challenging. Ionic liquid based hydrogen carriers have thus attracted the interest of researchers.\textsuperscript{29,35}

In this paper, we report guanidinium octahydrotriborate (Fig. 1) from synthesis to hydrogen release. Theoretically, guanidinium octahydrotriborate has 13.8 wt.% hydrogen (14/101 = 13.8 wt.%), and the 6 \textit{H}\textsuperscript{+} in the (C(NH\textsubscript{2})\textsubscript{3})\textsuperscript{2+} cation and the 8 \textit{H}\textsuperscript{–} in the (B\textsubscript{3}H\textsubscript{8})\textsuperscript{–} anion would contribute to a facile formation of H\textsubscript{2}. Guanidinium octahydrotriborate was once reported as a solid at 20 °C and it is not pure.\textsuperscript{36} We found, however, this compound has a melting point below –10 °C, which makes it interesting for injectable/pumpable liquid fuel for hydrogen supply. The high hydrogen capacity could also make guanidinium octahydrotriborate a green hypergolic fuel.\textsuperscript{37-40}

**Fig. 1** Schematic diagram of the structure of guanidinium octahydrotriborate (grey, C; blue, N; pink, B; white, H).

**Experimental**

**Reagents**

Guanidinium chloride (98%) was purchased from Alfa Aesar and used as received. Tetrahydrofuran (THF) was purchased from Merck (liquid chromatography grade, ≥ 99.9%). Sodium octahydrotriborate (NaB\textsubscript{3}H\textsubscript{8}, > 97%) was prepared following the method reported.\textsuperscript{41}

**Characterization**

Solvent nuclear magnetic resonance spectroscopy (NMR) and solid-state magic-angle-spinning nuclear magnetic resonance spectroscopy (MAS NMR) were performed on a Bruker Avance 500 MHz spectrometer equipped with liquid and solid detectors, respectively. Solution \textsuperscript{1}H NMR was referenced to CD\textsubscript{3}CN and \textsuperscript{11}B NMR to BF\textsubscript{3}Et\textsubscript{2}O. Powder X-ray diffraction (XRD) characterization was performed on a PANalytical X’Pert diffractometer (Cu K\textalpha, 40 kV, 40 mA) with a self-made sample holder that protects the samples from exposure to air. Temperature programmed desorption (TPD) coupled with a mass spectrometer (MS, Hiden Analytical HPR 20) (TPD-MS) was employed to analyse the released gas. Volumetric release measurements (VR) were performed on a Sieverts apparatus.\textsuperscript{32} Decomposition in the solvents was conducted in an autoclave fitted with a pressure detector. Both solution NMR and solid state MAS NMR were used to characterize the intermediates and final products. Fourier-transform infrared (FT-IR) data were collected on a Varian 3100 FTIR spectrophotometer (Varian, USA, Excalibar Series) in the diffuse reflection infrared Fourier transform spectroscopy (DRIFT) mode. Low tempera-

ture differential scanning calorimetry (DSC) was performed on a Netzsch DSC 204 unit (Netzsch) with a heating rate of 2 °C/min. Differential thermal analysis (DTA) measurements were carried out on a Netzsch STA449C thermal analysis system (Netzsch) with a heating rate of 2 °C/min. All the manipulations were carried out in an air-free environment.

**Results and Discussion**

**Synthesis**

For the synthesis of guanidinium octahydrotriborate, NaB\textsubscript{3}H\textsubscript{8} and guanidinium chloride were ball milled together in a 1:1 molar ratio at 150 rpm for 4 h. Guanidinium octahydrotriborate can also be produced by reacting these two starting materials in tetrahydrofuran (THF). The resultant precipitate, i.e., NaCl, was filtered away in THF. Powder X-ray diffraction (XRD) confirmed the identity of the solid powder as NaCl (Supporting Information Fig. S1). Based upon the weight of NaCl and the quantity of the starting materials, the following reaction (Eq. 1) can be derived.

\[
\text{NaB}_3\text{H}_8 + \text{C(NH}_2\text{)}_3\text{Cl} \rightarrow \text{C(NH}_2\text{)}_3\text{B}_3\text{H}_8 + \text{NaCl}
\]

This was corroborated by NMR studies. The \textsuperscript{11}B NMR spectrum (Fig. 2, left) indicates the integrity of the B\textsubscript{3}H\textsubscript{8} unit after the reaction. The \textsuperscript{1}H NMR spectrum (Fig. 2, right) confirms the coexistence of both NH\textsubscript{2} and B\textsubscript{3}H\textsubscript{8} units in a ratio close to 3:1, based upon the integration of the \textsuperscript{1}H NMR resonance.

![Fig. 2](image)

This was confirmed by a low temperature DSC measurement from 40 °C down to –40 °C, which revealed a melting point below –10 °C upon heating (Fig. S2). Guanidinium octahydrotriborate is highly miscible and stable in various solvents such as diethyl ether, acetonitrile, dichloromethane, tetrahydrofuran, 1,2-dimethoxyethane, etc. Clear solutions with concentrations ranging from 0.2 mol/L to 10 mol/L could be prepared in THF (Fig. S3). No obvious change was detected in the NMR characterization after 2 months in storage at 4 °C. Isolation of guanidinium octahydrotriborate from several solvents was unsuccessful, since it tends to decompose during the prolonged pumping to remove solvents. In measuring the ther-
mal decomposition of the ball-milled product, NaCl was not removed after synthesis. Considering the high thermal stability of NaCl, it is believed that NaCl will not affect the decomposition of guanidinium octahydrotriborate.26, 27

**Thermal decomposition of the as-prepared guanidinium octahydrotriborate**

The TPD-MS curve (Fig. 3) shows that only H₂ was detected during the heat treatment to 350 °C, a big improvement in purity over that of guanidinium borohydride, where ammonia was observed as a by-product.25 Guanidinium octahydrotriborate starts to release H₂ at about 70 °C and peaks at about 78 °C, which are noticeably lower than for guanidinium borohydride (dehydrogenation starting at 102 °C and peaking at 110 °C).25 A slight jump in temperature was detected, indicating that the decomposition is exothermic. This was confirmed by DTA measurement (Fig. S4).

![Fig. 3 TPD-MS profiles of guanidinium octahydrotriborate with a ramping rate of 2 °C/min.](image)

To quantify the hydrogen released, the as-prepared guanidinium octahydrotriborate sample from ball milling was heated at 83.5 °C in a closed vessel. The temperature was set to increase from 20 °C to 85 °C at a rate of 2 °C /min and then dwell at 85 °C. Inside the vessel, the temperature was measured to be 83.5 °C. Only H₂ was detectable by MS as the gaseous product. The VR results (Fig. 4) show that about 5.5 equiv. H₂ was released in less than 10 minutes, corresponding to 6.9 wt.% hydrogen, taking into account the NaCl.

![Fig. 4 Dehydrogenation of the as-prepared guanidinium octahydrotriborate at 83.5 °C. Volumetric release measurement (VR) (left) and MS analysis (right).](image)

The thermal decomposition of NH₃B₃H₇, NH₄B₃H₈, and (NH₃)₂BH₂B₃H₈ gives off noticeable amount of B₂H₆, B₃H₈, and B₃N₃H₆.43 The negligible amount of impurities in the case of guanidinium octahydrotriborate may be due to the improved ratio of H⁺ to H⁻, i.e., from 3:6 and 3:5 in NH₄B₃H₈ and (NH₃)₂BH₂B₃H₈, to 3:4 in C(NH₂)₃B₃H₈. In addition, the highly mobile C(NH₂)₃⁺ and B₃H₈⁻ in the liquid state could effectively contribute to the combination of H⁺ with H⁻ upon heating, without the need to overcome a mass transfer barrier as in all the solid crystalline octahydrotriborates. More efficient H₂ evolution via the combination of H⁺ and H⁻ is the result, and at the same time, the formation of volatile species is suppressed.

**Decomposition in solvents**

Being liquid at room temperature gives guanidinium octahydrotriborate a great potential as an injectable hydrogen carrier, which would effectively facilitate the control of hydrogen evolution in terms of both rates and quantity, using the current liquid fuel distribution techniques. To test the feasibility, 0.1 mol/L guanidinium octahydrotriborate in THF solution was heated to different temperatures. THF was used to separate NaCl from the product and to reduce the viscosity of the guanidinium octahydrotriborate. The concentration can be adjusted to maximize the capacity. Other high boiling point solvents such as dimethyl sulfoxide have also been tried with similar results obtained (not shown here). The released gas was analysed by MS (Fig. 5). When heated at 90 °C, H₂ is released with high purity. Note that the peak at 40 amu in Fig. 5 is associated with argon. In total, 3.9 equiv. H₂ was released at 90 °C in 1 hour (see Table 1). When the solution was treated at 100 °C, the dehydrogenation took place in multiple steps (Fig. 6). About 6.5 equiv. or 12.9 wt.% H₂ in all was released after 95 h, with about 4 equiv. H₂ released in the first hour and the other 2.5 equiv. H₂ over the remaining time. Solution ¹¹B NMR spectra were collected during this period (Fig. S5). Most guanidinium octahydrotriborate decomposed in less than 30 min, as the resonance associated with B₃H₈⁻ became very weak, while trace amounts of B₂H₆ (present as THF•BH₃ at –0.4 ppm) and B₃N₃H₆ (27.5 ppm) were produced. Meanwhile, a large amount of precipitate was found. Since mass transport in solvent is faster than in a solid, the subsequent hydrogen evolution is slow. Both B₂H₆ and B₃N₃H₆ participate in the subsequent reaction(s), since they were not observed after 6.5 equiv. H₂ was released (Fig. S5).

![Table 1: H₂ evolution at different temperatures and on different time scales.](image)
Solvent alters the decomposition pathway of guanidinium octahydrotriborate. Without solvent, the compound decomposes at about 70 °C, and there was no further decomposition at higher temperatures. The VR results revealed ~6.9 wt % H₂ released at 83.5 °C. When guanidinium octahydrotriborate is added to THF, however, there are stepwise release of H₂ at 100 °C, and 12.9 wt.% H₂ is ultimately released. The solvent contributes to the interactions between the hydrogen-rich cations and anions. This is in contrast to the slower mass transfer in the solid-state reaction.

Fig. 5 MS spectra of the gaseous products from 0.1 mol/L guanidinium octahydrotriborate in THF at 90 °C (left), and 100 °C (right).

Concentration effects were considered in the first two hours. From the dehydrogenation curve (Fig. 7), the 0.1 mol/L solution features a two-step hydrogen release, while the 0.5 mol/L and 1 mol/L solutions feature only one step. MS analysis (Fig. S6) indicates that the concentration does not affect the purity of the H₂. There is a slight decline in quantity, however, from 0.1 mol/L to 0.5 mol/L, and then to 1 mol/L.

Fig. 6 H₂ release profile of 0.1 mol/L guanidinium octahydrotriborate in THF at 100 °C (inset is the first 2 h).

Cross-polarization and one-pulse ¹¹B MAS NMR were both performed (Fig. 8). Signals at –42 ppm can be assigned to –BH₄. The small peaks at –16 ppm and at –12 ppm are related to tetra-coordinated B species, with the differences in shift arising from the different coordination environments. Other shifts at –2 ppm, 0 ppm, 15 ppm, and 19 ppm correspond to tri-coordinated B species such as –BN₃ and/or –BN₂H. Solid-state cross-polarization ¹³C MAS NMR spectra of these two stages were also collected (Fig. 9). They show nearly identical signals, which means that there are no significant changes in the carbon skeleton between these two stages. The ¹³C MAS NMR spectra both have signals at 159, 69, 63, 35, 26, 20, and 14 ppm. The peak at 159 ppm can be assigned to C=N double bonds and peaks at 69 and 63 ppm to C-N single bonds. From 4 D to 6.5 D, the main changes are the increase in intensity for the peak at 63 ppm and the decrease for peak at 69 ppm. Therefore, the peak at 69 ppm is likely to be associated with an intermediate, which could further dehydrogenate to certain species linked to the peak at 63 ppm. To validate this, the 6.5 D products were further treated at 120 °C for 3 h under dynamic conditions.

Fig. 7 Hydrogen release from guanidinium octahydrotriborate at different concentrations in THF with heating up to 100 °C.

Solid products characterization

Solid-state ¹¹B and ¹³C MAS NMR spectroscopy (Figs. 8 and 9), which give the coordination environments of B and C, were employed to study the solid products. Solid samples were collected at two points for 0.1 mol/L guanidinium octahydrotriborate in THF: after release of about 4 equiv. H₂ (4 D for short) and after release of 6.5 equiv. H₂ (6.5 D for short).

Fig. 8 Solid state MAS ¹¹B NMR of the precipitate: (a) ¹¹B MAS cross-polarization NMR and (b) ¹¹B MAS one-pulse NMR. Blue line: 4 D products; black line: 6.5 D products.
vacuum, after which, the peak at 69 ppm nearly disappeared, and only the peak at 63 ppm remained (Fig. S7). Normally peaks at 15-40 ppm are indicative of C-H single bonds, but there are no C-H bonds in the starting materials. The Fourier transform infrared (FTIR) results confirm the existence of C-H stretching vibrations in 4 D and 6.5 D (Fig. S8). These may come from the residual solvent, since there is a chance that the THF was not removed completely, although the formation of C-H bonds during the dehydrogenation of guanidinium octahydrotriborate is also likely.

![Cross-polarization MAS $^{13}$C NMR spectra of the 4 D and 6 D products.](image)

To elucidate the origins of the C-H bonds, TPD-MS was employed to monitor the thermal decomposition of the 6.5 D products (Fig. S9). THF, diborane, and ammonia were not detected, while only trace amounts of hydrogen were released from 150 - 250 °C. Furthermore, another $^{13}$C NMR spectrum was collected on the 6.5 D products, which were further treated at 120 °C for 3 h under dynamic vacuum. Under these conditions, the THF can be removed nearly completely due to its high volatility. The peaks at 15-40 ppm are still visible (Fig. S7). Based upon these results, it can be said that C-H bonds are formed in the dehydrogenated products. Due to the typical high bonding energy of C-H, this portion of the hydrogen would be difficult to extract under the applied experimental conditions.

After the dehydrogenation at 100 °C, a very weak chemical shift associated with $-\text{B}_3\text{H}_8$ was detected in the solution $^{11}$B NMR spectrum (Fig. S5), and a weak shift correlated to $-\text{BH}_4$ was detected in the solid state $^{11}$B NMR spectrum (Fig. 8). The formation of $-\text{BH}_4$ during the thermal decomposition of $-\text{B}_3\text{H}_8$ has been reported. The observed trace amounts of hydrogen at 150 - 250 °C from the precipitate (Fig. S9) is likely to be due to the decomposition of the $-\text{BH}_4$ species. The appearance of different tetra- and tri-coordinated B environments (Fig. 8) is commonly observed in the decomposition products of BNH compounds. These are similar to the results of thermal decomposition of guanidinium borohydride, where protic and hydric hydrogens combine to form H$_2$ and B-N bonds and polymerize to form a precipitate at the same time. The precipitate has a lower mass transfer rate, so the evolution rate of H$_2$ decreases. Besides the release of hydrogen, there is partial dehydrogenation of carbon. For practical applications, the formation of the precipitate during dehydrogenation would require off-board digestion and regeneration of the spent fuel.

For solutions with different concentrations, the resultant solid products feature slightly different structural features, as evidenced by the solid-state NMR spectra (Figs. S10 and S11). In the solid state MAS cross-polarization $^{11}$B NMR, the broad peak at 19 ppm shifts to 22.2 ppm and 26.7 ppm when the concentration increases. For the MAS cross-polarization $^{13}$C NMR, when the concentration increases, the peaks at 26 ppm and 69 ppm become weak, while the peak at 63 ppm increases in intensity. Since the concentration has a slight impact on the amount of hydrogen evolved (Fig. 7), the remaining hydrogen is responsible for the differences in the C and B coordination environments, but the main framework of the dehydrogenated products is similar in all cases.

The solid product from decomposition at 100 °C for 95 h in THF bears some similarities to the remains of the guanidinium octahydrotriborate decomposed without solvent. The products of these two different dehydrogenation methods are similar in respect to the B environments. (Fig. S12). For the C coordination environments, the main peak at 159 ppm related to C=N appears under both conditions (Fig. S13), but the intensity of this peak for decomposition in THF is much weaker. This indicates that the guanidinium cation structure has largely disappeared when more H is released (6.5 H$_2$ released versus 5.5 H$_2$).

**Conclusions**

Guanidinium octahydrotriborate, with a high hydrogen content of 13.8 wt.%, has been successfully synthesized. It behaves like an ionic liquid with a melting point below –10 °C. Its liquid state could effectively facilitate H$_2$ supply upon demand using the current liquid fuel distribution techniques. High purity H$_2$, as evidenced by MS analysis, was released at a temperature as low as 70 °C. Quantitative measurements demonstrated that about 6.9 wt.% (if excluding NaCl, 11.2 wt.%) hydrogen can be released at 83.5 °C within a short period. When dissolved in THF, about 4 equiv. H$_2$ (7.9 wt.% based on guanidinium octahydrotriborate) was released in less than 1 h, and about 6.5 equiv. H$_2$ (12.9 wt.% based on guanidinium octahydrotriborate) was released after an extended period. High purity H$_2$ with negligible amounts of impurities was confirmed by the MS spectra. During the dehydrogenation, solid spent fuel is formed, which slowly gives off trace amounts of H$_2$ upon further heating. To fully evaluate this compound for hydrogen storage, further tests on the thermal stability, vapour pressure, and viscosity in various solvents, as well as the changes in these properties with temperature and the temperature dependent kinetics, are needed.

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