Bromide substitution in lithium borohydride, LiBH₄–LiBr

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ABSTRACT

By means of in situ synchrotron radiation powder X-ray diffraction, powder neutron diffraction, attenuated total reflectance infrared spectroscopy, differential scanning calorimetry and the Sieverts techniques we have investigated how anion substitution in the LiBH₄–LiBr system leads to changes in the structural, physical, chemical and hydrogen storage properties of this material. Mechano-chemical treatment facilitates formation of a hexagonal solid solution h-Li(BH₄)₁₋ₓBrₓ whereas heating at elevated temperatures, i.e. T > 112 °C appears to allow full solubility in the system LiBH₄–LiBr. The first step in the anion substitution process may be dissolution of small amounts of LiBH₄ in α-LiBr deduced from observation of a hexagonal solid solution with unit cell volume similar to β-LiBr for a hand-mixed sample. The solid solution, Li[(BH₄)₁₋ₓBrₓ], is isostructural to the hexagonal high temperature polymorph of LiBH₄. This solid solution melts at a significantly higher temperature depending on the composition as compared to h-LiBH₄. Furthermore, a new hexalithium borate tribromide, Li₆(BO₃)Br₃ was discovered and structurally characterized.

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1. Introduction

The renewable energy sources, e.g. solar cells, wind turbines or wave energy, are sustainable, environmentally friendly, inexhaustible alternatives to fossil fuels. However, the widespread use of these energy sources is in general hampered by their fluctuation in time and their non-uniform distribution geographically [1–3]. One solution is the development of a safe, efficient and inexpensive energy storage system, and hydrogen is world-wide considered as an alternative energy carrier. Hydrogen is conveniently produced by electrolytic water splitting, but storage of hydrogen remains a significant unsolved problem [4–6]. A wide range of materials has been investigated during the past decade initially with a focus on magnesium or alanate based hydrides [7–11]. Sodium alanate is an excellent candidate for reversible storage of hydrogen, however, the capacity remains too low when measured on a system basis [12]. This has prompted intense research in metal borohydrides, which often have higher hydrogen contents [13–15]. Metal borohydride materials are interesting solid state hydrogen storage materials due to the generally high hydrogen content [14,15]. Unfortunately, the borohydrides...
usually have insufficient kinetic and thermodynamic properties, e.g. lithium borohydride, LiBH₄, which contains 18.5 wt% H₂ and 122.5 g H₂/L, releases hydrogen at fairly high temperatures of 410 °C (p(H₂) = 1 bar) [16]. The decomposition mechanism is generally described as shown in reaction scheme (1) [17]:

$$2\text{LiBH}_4 \rightarrow 2\text{LiH} + 2\text{B} + 3\text{H}_2 \quad \rho_{\text{m}}(\text{H}_2) = 13.97\text{wt%}$$ (1)

Although other decomposition products of LiBH₄ have been observed, e.g. Li₄B₂H₆, none of them are well characterized and the rehydrogenation has been reported at harsh conditions, T = 600 °C and p(H₂) = 155 bar [18–21]. LiBH₄ is known to exist in four polymorphic forms [22]. At ambient pressure and temperature, LiBH₄ exhibits an orthorhombic structure (α-LiBH₄) with the space group Pnma (no. 62) and unit cell parameters a = 7.141(5), b = 4.431(3), c = 6.748(4) Å [22]. Upon heating to ~112 °C, this structure transforms into a hexagonal polymorph (h-LiBH₄), a = 4.3228(10) and c = 7.0368(10) Å, with space group P6₃mc (no. 186) [23–25]. At high pressure, two structurally different polymorphs are identified with space group symmetries Am2 (no. 40) and Fm-3m (no. 225) [23,26].

The properties of LiBH₄ can be modified using appropriate additives, e.g. SiO₂, TiF₃ or TiC₁₆ [16,27–30]. Reactive hydride composites (RHC) may significantly improve the thermodynamic properties of LiBH₄, e.g. as observed in the 2LiBH₄–MgH₂ system [31–38]. Anion substitution is another interesting strategy, which may stabilize the too unstable borohydrides or destabilize the too stable borohydrides [28,39–49]. Substitution of BH₄⁻ with Cl⁻, Br⁻ and I⁻ anions in LiBH₄ is reported to significantly improve the ion conductivity of LiBH₄ [50–52].

This has prompted the present study of the mechanism for bromide, Br⁻, substitution in LiBH₄ and the effect on hydrogen release in the system LiBH₄–LiBr. Lithium bromide, α-LiBr, has a cubic NaCl-type structure (Fm-3m, no. 225) and a hexagonal polymorph, β-LiBr (P6₃mc, no. 186) which can be synthesized at T ≤ 0 °C using a substrate to initiate crystal growth [53–55]. We have used in situ synchrotron radiation powder X-ray diffraction (SR-PXD), powder neutron scattering (PND), attenuated total reflection infrared spectroscopy (ATR-IR), differential scanning calorimetry (DSC), and the Sieverts method, to study the effect of anion substitution on the structural, physical, chemical and hydrogen storage properties of LiBH₄.

2. Experimental

Samples were prepared from mixtures of lithium borohydride, LiBH₄, (95.0%, Aldrich) and lithium bromide, α-LiBr (98.0%, Fluka) in molar ratios of 1:0.25 (denoted S1), 1:0.5 (S2), 1:1 (S3) and 1:2 (S4) using mechano-chemical synthesis (ball milling). A sample for PND was prepared mixing Li₁₁BD₄ (98%, KatChem) and α-LiBr (98%, Fluka) in the molar ratio 1:1 (denoted S5). All samples were ball milled under inert conditions (argon atmosphere) using a Fritsch Pulverisette no. 4. The ball milling consists of 2 min milling intervened by 2 min breaks to avoid heating of the sample. This was repeated 6 times. The sample to ball ratio was 1:40 and tungsten carbide (WC) vial (80 mL) and balls (Diameter: 10 mm) was used.

A fraction (~0.5 g) of the samples prepared same way as S1–S5 was transferred to corundum crucibles placed in a sealed argon filled quartz tube and annealed in a furnace kept at a fixed temperature of 280 °C for 96 h (S1A, S2A, S3A and S4A) or 245 °C for 72 h (S3B, S5B). Furthermore, a sample of LiBH₄–LiBr in molar ratio 1:1 was prepared by hand milling (HM) in an agate mortar for 10 min (denoted S6). All investigated samples are listed in Table 1.

Laboratory Powder X-ray Diffraction (PXD) were performed in Debye-Scherrer transmission geometry using a Stoe diffractometer equipped with a curved Ge(111) monochromator (Cu Kα₁ radiation, λ = 1.54060 Å) and a curved position-sensitive detector. Data were collected at RT between 4 and 12° 20 with counting times of ~280 s per step. The samples were mounted in a glovebox in 0.5 mm (outer diameter, o.d.) glass capillaries sealed with glue.

In situ SR-PXD data for sample S2 and S6 were measured at beamline BM01A at the Swiss-Norwegian Beam Lines (SNBL), European Synchrotron Radiation Facility (ESRF), Grenoble, France, using a MAR345 image plate detector. The samples were mounted in glass capillaries (0.5 mm o.d.) sealed with a composite adhesive to prevent contact with air. The data were collected with a sample to detector distance of 20 cm.

### Table 1 – List of investigated samples. The composition of the samples is given as the molar ratios and the molar fractions, and the calculated hydrogen content is denoted $\rho_{\text{m}}(\text{H}_2)$. The preparation methods are either ball milling (BM) or hand-mixing in a mortar (HM) and in some cases combined with annealing in argon atmosphere.

<table>
<thead>
<tr>
<th>Notation</th>
<th>Materials</th>
<th>Molar ratio</th>
<th>n(LiBH₄)/n(total)</th>
<th>Preparation</th>
<th>$\rho_{\text{m}}(\text{H}_2)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1A</td>
<td>LiBH₄–LiBr</td>
<td>1:0.25</td>
<td>0.200</td>
<td>BM, A&lt;sup&gt;a&lt;/sup&gt;</td>
<td>9.3</td>
</tr>
<tr>
<td>S2</td>
<td>LiBH₄–LiBr</td>
<td>1:0.5</td>
<td>0.333</td>
<td>BM</td>
<td>6.2</td>
</tr>
<tr>
<td>S2A</td>
<td>LiBH₄–LiBr</td>
<td>1:0.5</td>
<td>0.334</td>
<td>BM, A&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6.2</td>
</tr>
<tr>
<td>S2C</td>
<td>LiBH₄–LiBr</td>
<td>1:1</td>
<td>0.334</td>
<td>BM</td>
<td>6.2</td>
</tr>
<tr>
<td>S3</td>
<td>LiBH₄–LiBr</td>
<td>1:1</td>
<td>0.498</td>
<td>BM</td>
<td>3.7</td>
</tr>
<tr>
<td>S3A</td>
<td>LiBH₄–LiBr</td>
<td>1:1</td>
<td>0.500</td>
<td>BM, A&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.7</td>
</tr>
<tr>
<td>S3B</td>
<td>LiBH₄–LiBr</td>
<td>1:1</td>
<td>0.500</td>
<td>BM, A&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.7</td>
</tr>
<tr>
<td>S4A</td>
<td>LiBH₄–LiBr</td>
<td>1:2</td>
<td>0.665</td>
<td>BM, A&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.1</td>
</tr>
<tr>
<td>S5B</td>
<td>Li₁₁BD₄–LiBr</td>
<td>1:1</td>
<td>0.500</td>
<td>BM, A&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.7</td>
</tr>
<tr>
<td>S6</td>
<td>LiBH₄–LiBr</td>
<td>1:1</td>
<td>0.500</td>
<td>HM</td>
<td>3.7</td>
</tr>
</tbody>
</table>

Conditions for annealing. <sup>a</sup> 280 °C/96 h, <sup>b</sup> 245 °C/72 h and <sup>c</sup> 250 °C/5 min.

Li₆(BO₃)Br₃ was indexed from the HR-PXD and PND data using the Fullprof suite [60], see further details in the sample holder. The high-resolution powder X-ray diffraction (HR-PXD) measurement (λ = 0.500860 Å) for SSB was performed at the beamline BM01B at the SNBL, ESRF. A channel-cut monochromator Si(111) was used for wavelength selection. The diffractometer was equipped with six analyzer crystals, meaning that six complete patterns were collected simultaneously, with an offset of 1.1° between the detectors. The sample was mounted in a glass capillary (0.5 mm o.d.) sealed with a composite adhesive to prevent contact with air.

SR-PXD data for sample S1A, S2A, S3A and S4A were measured at the beamline I711 at the MAX-II synchrotron in the research laboratory MAX-lab, Lund, Sweden (λ = 0.955 Å) with a MAR165 CCD detector system [56]. The samples were mounted in sapphire (Al₂O₃) single crystal tubes (1.09 mm o.d., 0.79 mm i.d.) in argon filled glovebox (p(O₂), H₂O < 0.5 ppm) [57]. The SR-PXD experiments were conducted in argon atmosphere (p(Ar) = 1 bar) at RT.

Powder neutron diffraction (PND) data were collected at RT on sample SSB with the PUS instrument at the JEEP II reactor at Kjeller, Norway [58]. Neutrons with wavelength 1.5553 Å were obtained from a Ge(Si11) monochromator. The detector unit consists of two banks of seven position-sensitive ³He detectors, each covering 20° in 2θ (binned in steps of 0.05°). Data were collected from 10 to 130° in 2θ. The sample was contained in a rotating cylindrical 6 mm diameter vanadium sample holder.

All SR-PXD data were integrated using the Fit2D program [59]. The SR-PXD data were analyzed by Rietveld refinement using the Fullprof suite [60], see further details in the supplementary information.

The novel compound hexalithium borate tribromide, Li₆(BO₃)Br₃ was indexed from the HR-PXD and PND data collected on SSB using DICVOL2004 [61]. A hexagonal structural model was obtained using direct space algorithms implemented in the program FOX, a = 8.94084(6) and c = 5.77783(6) Å with the space group symmetry P6₃/mmc [62]. The structural model was finally refined using the Rietveld method in the Fullprof suite [60]. The Rietveld refinement converged at R_B = 4.18%, R_F = 3.84%, R_p = 11.3% and R_wP = 12.3%.

Attenuated total reflectance infrared spectroscopy (ATR-IR) measurements were performed for sample S2 and S2C heated to 250 °C (heating rate 50 °C/min, p(H₂) = 250 mbar) using an ATR-IR spectrophotometer (Bruker Alpha equipped with the ATR single-reflection accessory with a diamond crystal), placed in a nitrogen filled glovebox (gas level: H₂O < 0.1 ppm, O₂ < 0.5 ppm). All spectra were measured at RT in the spectral-range 4000 to 375 cm⁻¹ with 2 cm⁻¹ resolution. Sixty four scans were averaged for each spectrum.

Differential scanning calorimetry (DSC) was performed with a Netzsch STA449C Jupiter instrument on sample S3 at a heating rate of 1.5 °C/min from RT to 410 °C and on sample R1 at a heating rate of 1.5 °C/min from RT to 340 °C, both in a flow of He (50 mL/min). The samples were placed in Al₂O₃ crucibles with a small hole in the lid to prevent increase in pressure during desorption of gases.

The cyclic stability of the substituted compounds was characterized by Sieverts measurements recorded on sample S2 with a Gas Reaction Controller from Advanced Materials Corporation [63]. The sample was loaded in a stainless steel autoclave in a glovebox with nitrogen atmosphere (gas level: H₂O < 0.1 ppm, O₂ < 0.5 ppm). Temperature-pressure desorption (TPD) experiments were performed in the temperature range RT to 500 °C with a heating rate of 4.5 °C/min with a backpressure of 0.2–1 bar of H₂. Hydrogen absorption was measured at a fixed temperature of 415 °C for 50 h, applying an initial hydrogen pressure of ca. 100 bar. Three hydrogen desorption/absorption cycles were subsequently measured. In parallel, three desorption/absorption cycles on pure LiBH₄ were measured at experimental conditions identical to those for sample S2.

3. Results and discussion

3.1. Investigation of the mechanism for bromide substitution

3.1.1. Substitution by mechnano-chemical treatment, ball milling

Anion substitution is studied using in situ SR-PXD data measured for a ball milled sample of LiBH₄−LiBr (1:0.5, S2) heated and cooled three times in the temperature range 26–300 °C (5 °C/min), see Fig. 1. The first diffractogram contains Bragg reflections from a hexagonal solid solution h-Li(BH₄)₀.₄Br₀.₃Brₓ LiBr and weak reflections from o-LiBH₄.

![Fig. 1 – In situ SR-PXD data for LiBH₄−LiBr (1:0.5, S2) heated from RT to 300 °C and cooled to RT three times (heating/cooling rate 4 °C/min, λ = 0.709595 Å, ESRF, BM01B). Symbols: ○ o-LiBH₄ × α-LiBr ● h-Li(BH₄)₀.₄Br₀.₃Brₓ ◊ Li₆(BO₃)Br₃.](image)
(determined by Rietveld refinements). This reveals that a significant degree of anion substitution takes place during ball milling, which produces a solid solution. During the first heating the intensity of the reflections from α-LiBH4 and α-LiBr gradually decreases, suggesting that α-LiBH4 and α-LiBr gradually dissolve in the solid solution, h-Li(BH4)1−xBrx. A single phase solid solution with composition h-Li(BH4)0.75Br0.25 is obtained during the first heating at 280 °C. However, during the second in situ heating of a novel compound, hexahaloromatic tribromide, Li4[BO3]Br5, is observed, representing ~10 wt% of the sample after the third heating. This compound may form due to the hygroscopic nature of lithium bromide, which may contain water.

3.1.2. Anion substitution facilitated by hand mixing in a mortar

A hand mixed (HM) sample of LiBH4–LiBr (1:1, S6) was investigated by in situ SR-PXD in the temperature range RT to 300 °C and then cooled to RT to follow the substitution process. The first PXD pattern, measured at 25 °C, contains Bragg reflections from α-LiBr and α-LiBH4, but also from the substituted solid solution, h-Li(BH4)1−xBrx and a small amount of another compound, possibly LiBr xH2O, which disappear at ~60 °C, see Fig. 2. Five Bragg diffraction peaks were identified and indexed with a cubic unit cell, a = 4.0213 Å, which corresponds with that for LiBr xH2O, a = 4.027 Å [64]. The weight fraction of h-Li(BH4)1−xBrx (3.8 wt%) is too small to allow determination of the bromide content in the solid solution prepared with the hand mixing procedure, i.e. compared to 67.9 wt% α-LiBH4, 26.9 wt% α-LiBr, and 1.5 wt% LiBr xH2O. The small unit cell volume per formula unit V/Z = 50.9 Å3 (T = 25 °C) for the solid solution h-Li(BH4)1−xBrx resembles the β-LiBr structure (V/Z = 49.6 Å3 at T = −50 °C [55]) indicating that the substitution process possibly initiates with dissolution of LiBH4 in α-LiBr, which stabilizes the β-LiBr structure due to a small degree of BH3 substitution during the grinding. The notation ‘y’ is used for the solid solution in this case to distinguish from the solid solutions with larger unit cells, which resemble the structure of LiBH4.

Upon heating at T > 40 °C, additional anion substitution is observed as decreasing diffracted intensity from α-LiBr. The polymorphic phase transition from α-LiBH4 to a solid solution h-Li(BH4)1−xBrx with low bromide content is observed at 112 °C, where the sample composition is h-Li(BH4)1−xBrx (~70.2 wt%), h-Li(BH4)1−xBrx (~8.2 wt%) and α-LiBr (21.6 wt%). The simultaneous existence of two hexagonal solid solutions with different composition, y > x, and a small difference in unit cell volumes per formula unit of ~2.5 Å2 are observed for the PXD patterns obtained in the temperature range 112–285 °C, see Fig. 3 and the Rietveld refinement shown in Fig. s1 in the supplementary information. Fig. 3 shows the unit cell volume per formula unit (V/Z) as a function of temperature extracted by sequential Rietveld refinement of the in situ SR-PXD data shown in Fig. 2. The change in the unit cell parameters a and c of h-Li(BH4)1−xBrx and h-Li(BH4)1−xBrx during the experiment is compared in the supplementary information in Fig. s2 showing a large impact on the unit cell parameter c upon substitution. Furthermore, Fig. 2 reveals that the diffracted intensity from α-LiBr and h-Li(BH4)1−xBrx decrease abruptly in the temperature range 249–296 °C, simultaneously with an increase in intensity from h-Li(BH4)1−xBrx. At 296 °C, full anion substitution is obtained resulting in a single phase solid solution of Li(BH4)1−xBrx and a significant decrease in the unit cell volume in the temperature range 281–300 °C despite the expected thermal expansion due to continued heating clearly illustrated in Fig. 3. The solid solution is observed to be stable upon the cooling to 40 °C where a fraction transforms from hexagonal to orthorhombic structure, i.e. formation of α-Li(BH4)1−xBrx with V/Z = 53.5 Å3, slightly smaller than that reported for α-LiBH4 (V/Z = 54.3 Å3) [65]. Formation of α-Li(BH4)1−xBrx is observed as relatively broad Bragg peaks, which may indicate a less well defined composition of the individual crystal grains.
3.1.3. Substitution by annealing
Anion substitution facilitated by thermal treatment is investigated for different compositions of LiBH$_4$–LiBr prepared by annealing at 280 °C for 96 h, i.e. 1:0.25 (S1A), 1:0.5 (S2A), 1:1 (S3A) and 1:2 (S4A). SR-PXD data for the four samples measured at RT two weeks after annealing are compared in Fig. 4. For sample S2A and S3A, a full substitution was obtained, i.e. less than 1 wt% excess x-LiBr is observed. For sample S1A, a full substitution was also obtained, however, excess lithium borohydride is observed as o-LiBH$_4$ (which may contain bromide) accounting for ~18 wt% of the sample. For sample S4A, excess of x-LiBr is observed, accounting for ~21 wt% of the sample. These results reveal that the hexagonal structure of LiBH$_4$ can be stabilized at RT for extended periods of time and suggest the existence of a lower and an upper limit for the bromide substitution in h-LiBH$_4$. The limits for anion substitution in the system LiBH$_4$–LiBr appear to be ~30–60 mol% after two weeks at RT, i.e. h-Li(BH$_4$)$_{1-x}$Br$_x$, 0.3 < x < 0.6. Full solubility in the system LiBH$_4$–LiBr is expected from the fact that both LiBr and LiBH$_4$ are prone to adopt hexagonal structures with space group P6$_3$/m and that the ionic radii for Br$^-$ and BH$_4^-$ are very similar 1.96 and 2.03 Å, respectively [55,66]. The fact that stability limits to the degree of anion substitution in the system LiBH$_4$–LiBr observed may originate from differences in coordination properties of the Br$^-$ and BH$_4^-$ ions. Limited solubility is also observed for LiBr–LiI, i.e. LiBr$_{1-x}$I$_x$ with 0.25 ≤ x ≤ 0.8 [53].

3.1.4. Analysis of the stability of Li(BH$_4$)$_{1-x}$Br$_x$
Further investigation of the stability of anion substitution in the system LiBH$_4$–LiBr was performed using an annealed (245 °C for 72 h) sample (1:1, S3B) characterized by PXD after a few days and after 8 and 14 months and the sample composition was extracted by Rietveld refinement. A few days after the annealing a single phase solid solution and no x-LiBr was observed. However after 8 and 14 months ca. 22 and 38 wt % x-LiBr, respectively, was observed in the sample. These experiments reveal a slow segregation of x-LiBr from the solid solution h-Li(BH$_4$)$_{1-x}$Br$_x$ at RT. Interestingly, the composition of the hexagonal solid solution remains approximately constant at, h-Li(BH$_4$)$_{0.5}$Br$_0.5$ over extended periods of time suggesting segregation of crystalline x-LiBr and amorphous o-LiBH$_4$ at RT. The experimental PXD data presented in Figs. 2 and 4 suggest that crystalline o-LiBH$_4$ may be segregated faster from a solid solution upon cooling. This shows that the hexagonal structure of LiBH$_4$ is efficiently stabilized at RT by bromide substitution over extended periods of time. A slow segregation of LiCl at RT was also observed for solid solutions of Li(BH$_4$)$_{1-x}$Cl$_x$ but in this case the stabilization of the hexagonal structure was less pronounced [39].

The results presented in this paper on anion substitution in the system LiBH$_4$–LiBr were reproduced several times using different samples with similar compositions. The formation of a metastable solid solution in the LiBH$_4$–LiBr system suggests a slightly positive enthalpy of mixing, which can be easily overcome by mechanical and thermal treatments combined with the ease of forming hexagonal structures. Entropic contributions make the solid solution stable at high temperatures. The comparable ionic radii of BH$_4^-$ and Br$^-$ lead to similar values of the charge-to-volume ratio, which are important for interactions in ionic compounds.

3.2. Structural investigation of hexalithium borate tribromide, Li$_6$(BO$_3$)Br$_3$
The new compound hexalithium borate tribromide, Li$_6$(BO$_3$)Br$_3$ has a hexagonal structure a = 8.94084(6), c = 5.77783(6) Å and space group P6$_3$/mmc. The structure was solved and refined from SR-PXD and PND data shown in the supplementary information as Figs. s3 and s4, respectively. Atomic coordinates and selected bond lengths are given in the supplementary information, see Tables s1 and s2. The structural model contains two independent lithium positions and one position for boron, oxygen and bromide. The bromide anions coordinate to eight lithium atoms in a bicapped trigonal prismatic manner, whereas boron forms the trigonal planar composite anion BO$_3^-$.

Each oxygen coordinate to one boron, two Li1 and two Li2. The closest neighbours of lithium Li1 is oxygen and at a greater distance (>2.85 Å) are the bromide atoms found. Lithium Li2 coordinates to two oxygen and two bromide atoms with a bond length of 2.750(9) Å additionally two bromide atoms are found at a distance of 2.8982(7). The new compound Li$_6$(BO$_3$)Br$_3$ has a three dimensional framework structure illustrated in Fig. 5. Visual inspection of the structure reveals open hexagonal ‘LiBr’ channels running along the c-axis and 5.54 Å wide measured as the Li–Br distance across the channel. Hexalithium borate tribromide formed in several samples after prolonged heating possibly due to the hygroscopic nature of lithium bromide, which may contain water, i.e. as LiBr·H$_2$O [64]. Recently, borohydride borates were discovered during release of hydrogen from Ca(BH$_4$)$_2$ and LiBH$_4$–Ca(BH$_4$)$_2$, i.e. Ca$_3$(BH$_4$)$_3$(BO$_3$) and LiCa$_3$(BH$_4$)$_3$(BO$_3$)$_2$ [31,67,68].

3.3. Infrared spectroscopy of Li(BH$_4$)$_{1-x}$Br$_x$
In order to study the changes in the vibrational properties of bromide substituted lithium borohydride, IR-ATR spectroscopic
measurements were performed on a sample of LiBH₄−LiBr (1:0.5, S₂), see Fig. 6. The IR-ATR spectrum of o-LiBH₄ (R₁), shown in Fig. 6 (a), reveal two main sets of peaks due to B–H stretching (2400–2000 cm⁻¹ region) and H–B–H bending (1300–800 cm⁻¹) vibrational modes, as already reported [69–71].

The isolated BH₄⁻ anion has an ideal tetrahedral symmetry, Td, however, the vibrational modes are split in the crystalline state due to lowering of the site symmetry from Td to Cs, i.e. the degenerate fundamental modes ν₂, ν₃, and ν₄ split into several components. In o-LiBH₄ (R₁), Fig. 6 (a), the broad peak at ~1420 cm⁻¹ may be a combination band of BH₄⁻ librational movements in the crystal found at ~400 cm⁻¹ (see [72]) and the fundamental mode ν₂ (A₁) at ~1050 cm⁻¹. The spectrum of as-prepared LiBH₄−LiBr (1:0.5, S₂) exhibits similar vibrational features as o-LiBH₄, see Fig. 6 (b) and (a), respectively, suggesting that o-LiBH₄ is still present and predominant. However, a decrease in the intensity of the peak at ca. 1420 cm⁻¹ and a small blueshift (3–4 cm⁻¹) of all the absorption bands indicates the presence of a small amount of another compound. PXD performed prior to the IR-ATR experiment showed o-LiBH₄, LiBr (which is not visible in the investigated spectral range) and h-Li(BH₄)ₓ LiBrₓ, see the Rietveld refinement profile in Fig. 5-s in the supplementary information. In order to obtain increased substitution, the sample LiBH₄−LiBr (1:0.5, S₂) was heated to 250 °C and cooled to RT. This resulted in differences in the BH₄⁻ stretching and bending regions compared to the spectrum of o-LiBH₄, see Fig. 6 (c), e.g. a decrease of the number of the components related to ν₂ and ν₃ fundamental modes. In the structure of h-LiBH₄, the BH₄⁻ ions have Cᵥ site symmetry and the ν₂ mode is doubly degenerate, while ν₃ has only two components. Therefore, the changes in the spectrum of Fig. 6 (c) with respect to Fig. 6 (a) may be related to the Cs to Cᵥ BH₄⁻ site symmetry change, confirming the phase transition from o- to h-LiBH₄ due to Br⁻ substitution. Similar changes upon heating have been reported for the Raman spectra of LiBH₄ due to the polymorphic phase transition [70].

Furthermore, several new peaks can be observed in Fig. 6 (c): at ~1170 cm⁻¹ (marked with an arrow), 1020 and 980 cm⁻¹, respectively. This indicates the presence of another compound. PXD performed prior to the IR-ATR experiment showed o-LiBH₄, LiBr (which is not visible in the investigated spectral range) and h-Li(BH₄)ₓ LiBrₓ, see the Rietveld refinement profile in Fig. 5-s in the supplementary information. In order to obtain increased substitution, the sample LiBH₄−LiBr (1:0.5, S₂) was heated to 250 °C and cooled to RT. This resulted in differences in the BH₄⁻ stretching and bending regions compared to the spectrum of o-LiBH₄, see Fig. 6 (c), e.g. a decrease of the number of the components related to ν₂ and ν₃ fundamental modes. In the structure of h-LiBH₄, the BH₄⁻ ions have Cᵥ site symmetry and the ν₂ mode is doubly degenerate, while ν₃ has only two components. Therefore, the changes in the spectrum of Fig. 6 (c) with respect to Fig. 6 (a) may be related to the Cs to Cᵥ BH₄⁻ site symmetry change, confirming the phase transition from o- to h-LiBH₄ due to Br⁻ substitution. Similar changes upon heating have been reported for the Raman spectra of LiBH₄ due to the polymorphic phase transition [70].

Furthermore, several new peaks can be observed in Fig. 6 (c): at ~1170 cm⁻¹ (marked with an arrow), 1020 and 980 cm⁻¹, respectively. This indicates the presence of another compound.
958 cm$^{-1}$. These peaks may be assigned to the fundamental modes of BH$_4^-$, slightly shifted owing to the presence of Br anions in the lattice of a substituted compound, even if the straightforward assignment is rather difficult. The peak at 1420 cm$^{-1}$ evidently decreases in intensity after Br substitution: A decrease in intensity of this peak can be related to alteration in the surroundings of BH$_4^-$, i.e. a change of neighbouring atoms and/or distances between the ions in the lattice. This peak may also originate from oxide species, however, no crystalline oxides are observed in the sample during the experiment.

All these changes indicate the formation of the bromide substituted hexagonal compound Li[BH$_4$]$_{1-x}$Br$_x$ formed due to the annealing. PXD performed on LiBH$_4$–LiBr (1:0.5, S2C, heated to 250 °C and cooled to RT) showed h-Li[BH$_4$]$_{1-x}$Br$_x$ and a small amount of residual α-LiBr, see the Rietveld refinement profile in the supplementary information Fig. s5. The spectrum (d) in Fig. 6 measured one month after the annealing confirm the stability of the Li[BH$_4$]$_{1-x}$Br$_x$ found by PXD.

3.4. Investigation of the bromide substitution by differential scanning calorimetry

Differential scanning calorimetry (DSC) was conducted in the temperature range RT to 400 °C with a heating rate of 1.5 °C/min for LiBH$_4$–LiBr (1:1, S3) and compared with α-LiBH$_4$ (R1), see Fig. 7. The polymorphic transition from α-LiBH$_4$ to h-LiBH$_4$ for the pure LiBH$_4$ (R1) is observed at 115.2 °C and the melting point at 286.3 °C. For sample LiBH$_4$–LiBr (1:1, S3) the signal from the polymorphic transition from α- to h-LiBH$_4$ is found to be significantly weaker and the melting point has increased to 377.9 °C, i.e. a stabilization of Li[BH$_4$]$_{1-x}$Br$_x$ compared to LiBH$_4$. The DSC experiment was continued using the same sample, i.e. performing an additional heating. No DSC signal for the polymorphic transition and the melting point was observed within the measured temperature range. These results confirm that a large degree of Br anion substitution occurs during ball milling and that additional anion substitution occurs during thermal annealing, in agreement with the SR-PXD results.

3.5. Rehydrogenation properties of Li[BH$_4$]$_{1-x}$Br$_x$

The rehydrogenation properties of the LiBH$_4$–LiBr system were investigated using the Sieverts method. Three hydrogen release and uptake cycles for LiBH$_4$–LiBr (1:0.5, S2) and for LiBH$_4$ are shown in Fig. 8 and hydrogen absorption measurements are shown in supplementary information Fig. s6.

A total hydrogen release of 12.4 wt% is observed for LiBH$_4$, during the first dehydrogenation (89% of the theoretical capacity: $\rho_0$($H_2$) = 13.88 wt%). A second and third dehydrogenation was measured for LiBH$_4$ after 50 h of hydrogenation, showing release of 6.1 and 5.3 wt% (44 and 38% of theoretical capacity). The dehydrogenation profile of LiBH$_4$–LiBr (1:0.5, S2) is similar to the second dehydrogenation from LiBH$_4$, see Fig. 8. A total hydrogen release of 5.6 wt% is observed for LiBH$_4$–LiBr (1:0.5, S2), i.e. 90% of the theoretical capacity, $\rho_0$($H_2$) = 6.2 wt%. A second and a third desorption was also measured showing hydrogen release of 2.8 and 2.3 wt% $H_2$ (45% and 37% of the theoretical capacity). The observed decrease in hydrogen storage capacity may be due to incomplete hydrogen absorption, see Fig. s6 in the supplementary information. Thus, the reversible hydrogen storage capacity of the system LiBH$_4$–LiBr is found to be 37% after 3 cycles, which is approximately the same as for LiBH$_4$ showing 38% reversibility after 3 cycles under the same conditions, i.e. the thermodynamics of the hydrogen uptake and release seems to be unchanged.

4. Conclusion

The mechanism for anion substitution in the system the LiBH$_4$–LiBr facilitated by mechano-chemical or thermal treatment is investigated by in situ synchrotron radiation powder X-ray diffraction and the data is analyzed by Rietveld refinement. Mechano-chemical treatment facilitates formation of a hexagonal solid solution h-Li[BH$_4$]$_{1-x}$Br$_x$ whereas heating at elevated temperatures $T > 112$ °C appears to allow full solubility in the system LiBH$_4$–LiBr. Hand mixing a sample of LiBH$_4$–LiBr leads to formation of a hexagonal solid solution h-Li[BH$_4$]$_{1-x}$Br$_x$ with small unit cell volume similar to α-LiBr, which suggest that the first step in the anion substitution process may be dissolution of small amounts of LiBH$_4$ in α-LiBr. Two solid solutions can be observed upon heating and the dissolution process is further accelerated at $T > 112$ °C resulting in a single phase solid solution at elevated temperatures. Cooling a solid solution h-Li[BH$_4$]$_{1-x}$Br$_x$ may lead to crystallisation of α-LiBr, α-Li[BH$_4$]$_{1-x}$Br$_x$ and/or a hexagonal solid solution h-Li[BH$_4$]$_{1-x}$Br$_x$. The composition of the solid solution h-Li[BH$_4$]$_{1-x}$Br$_x$ appears to remain at constant $x \sim 0.5$ upon storage at RT for several months, but α-LiBr and presumably amorphous α-LiBH$_4$ are slowly segregated. Bromide substitution
clearly stabilizes the hexagonal structure of LiBH₄ to RT as a solid solution with composition h-Li(BH₄)₂₋ₓBrₓ. However, the Sieverts measurements revealed that the hydrogen uptake and release properties of h-Li(BH₄)₁₋ₓBrₓ are similar to those of LiBH₄. A new hexalithium borate tribromide, Li₆(BO₃)Br₃ with a hexagonal structure was also discovered in this work. This compound may form upon heating LiBH₄-LiBr mixtures to temperatures above 280 °C and reveal the hygroscopic nature of lithium bromide, which may contain water, e.g. in the form LiBr·H₂O.

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Appendix. Supplementary material

Supplementary data related to this article can be found online at doi:10.1016/j.ijhydene.2011.08.087.

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