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Lithium nickel borides: evolution of [NiB] layers driven by Li pressure[†]

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Here we show the effect of Li chemical pressure on the structure of layered polymorphs with LiNiB composition: *RT*-LiNiB (room temperature polymorph) and *HT*-LiNiB (high temperature polymorph), resulting in stabilization of the novel *RT*-Li_{1+x}NiB ($x \sim 0.17$) and *HT*-Li_{1+y}NiB ($y \sim 0.06$) phases. Depending on the synthesis temperature and initial Li content, precisely controlled via hydride route synthesis, [NiB] layers undergo structural deformations, allowing for extra Li atoms to be accommodated between the layers. *In situ* variable temperature synchrotron and time-dependent laboratory powder X-ray diffraction studies suggest Li step-wise deintercalation processes: *RT*-Li_{1+x}NiB \rightarrow *RT*-LiNiB (high temp.) \rightarrow LiNi₃B_{1.8} \rightarrow binary Ni borides and *HT*-Li_{1+y}NiB \rightarrow *HT*-LiNiB (high temp.) \rightarrow LiNi₃B_{1.8} \rightarrow binary Ni borides. Quantum chemistry calculations and solid state ⁷Li and ¹¹B NMR spectroscopy shed light on the complexity of real superstructures of these compounds determined from high resolution synchrotron powder diffraction data.

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Introduction

The story of lithium nickel borides began in 1976 when four ternary phases were identified in the Li–Ni–B system.^{1–3} Crystal structures were reported for three of these compounds, LiNi₃B_{1.8} (MgNi₃B₂ structure type, *P*6₂₂ space group), Li₃Ni₂₀B₆ (Cr₂₃C₆, *Fm*3_m), and Li_{2.8}Ni₁₆B₈ (Li₃Ni₁₆B₈, *P*4/*mbm*). Only a tentative composition of 1:1:1 was assigned for the fourth compound, a phase with the richest Li content. To the best of our knowledge, no other ternary compounds in all *A*–T–B systems (*A* = alkali metal, *T* = Ti–Zn) have been reported. The lack of the ternary *A*–T–B compounds can be due to the drastically different reactivity of refractory boron (melting point 2348 K) and alkali metals (boiling point 944–1615 K) making traditional synthesis methods (*e.g.* arc melting) non-applicable.

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The hydride synthesis route, where salt-like alkali metal hydrides are used as a source of alkali metals, has been successfully utilized for the preparation of alkali/alkali-earth antimonides, arsenides, germanides, and silicides.^{4–10} Application of this method for the Li–Ni–B system significantly shortened the synthesis time of ternary phases. For instance, LiNi₃B_{1.8} can be synthesized in a span of \sim 24 hours compared with the previously reported 3-month annealing for the reaction of elements.⁹ The hydride route also permitted the compositional control and comprehensive screening of unknown compounds. This way, two lithium nickel boride polymorphs *RT*-LiNiB (room temperature polymorph) and *HT*-LiNiB (high temperature polymorph), with unique layered crystal structures, were discovered.¹¹ In these polymorphic compounds, Li layers, featuring hexagonal-like or squared motif, alternate with nearly planar [NiB] layers, whose structure differs depending on the polymorph (Fig. 1). A structural relationship between the *RT*-LiNiB and *HT*-LiNiB polymorphs, Li-depleted phase LiNi₃B_{1.8}, Ni₂B, and Li has been found,¹¹ indicating on the ∞ [NiB] framework evolves with the changing of Li content and/or temperature variation (Fig. S1[†]).

Layered lithium nickel borides are of interest from the functional materials perspective. Firstly, \sim 33 at% of Li is stored between [NiB] layers in RT- and HT-LiNiB (9.1 wt% of Li as compared to 8.8 wt% of Li in LiC₆ anode material) gives these compounds potential for applications in Li-ion batteries. Secondly, due to the unique arrangement of Li and [NiB] layers in their structures, exfoliation of these materials can open a



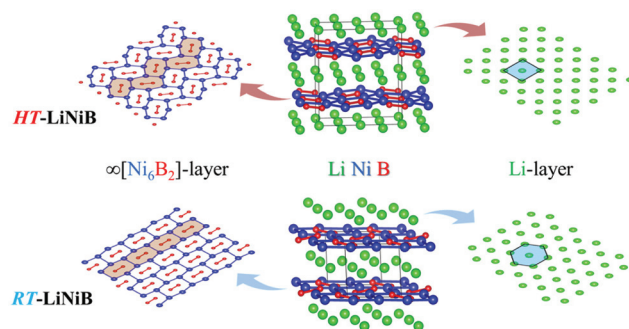


Fig. 1 Crystal structure of *RT*-LiNiB and *HT*-LiNiB polymorphs.

pathway to novel two-dimensional (2D) nickel borides.^{11,12} The search for 2D borides or MBene has been accelerated after successful preparation of the 2D transition metal carbides or MXene.^{13,14} MXenes are extensively studied due to their attractive thermal, magnetic, mechanical and electric properties.^{15–20} MBenes were predicted to have similarly fascinating properties^{21–24} but have not yet been obtained experimentally, despite several prominent attempts.^{24–28}

Recent theoretical investigation of the Li–Ni–B system²⁹ revealed a new Li-rich layered compound Li_2NiB , whose structure is derived from *RT*-LiNiB. According to the calculations, only a little extra energy is needed to accommodate an additional Li layer in the structure of the LiNiB parent phase. Here, we have experimentally explored the Li-rich compositions of the Li–Ni–B phase diagram and discovered two compounds that are Li-enriched versions of *RT*-LiNiB and *HT*-LiNiB polymorphs. Both *RT*- $\text{Li}_{1+x}\text{NiB}$ ($x \sim 0.17$) and *HT*- $\text{Li}_{1+y}\text{NiB}$ ($y \sim 0.06$) phases feature layered crystal structures with the original topology of the [NiB] layers modified due to the incorporation of extra Li between the layers. We use *in situ* synchrotron and laboratory powder X-ray diffraction, solid state NMR spectroscopy, and quantum chemical calculations to study structural relationships, reactivity, and phase transformations upon heating or exposure to air of these lithium-rich layered nickel borides.

Results and discussion

Synthesis and composition

In our recent study,¹¹ the Li–Ni–B ternary system was evaluated using an adaptive genetic algorithm (AGA) search for potentially stable ternary phases. As a result, a *RT*-LiNiB compound with a unique layered structure was computationally predicted and further synthesized experimentally using the hydride route. The structure of *RT*-LiNiB can be represented as almost planar [NiB] layers alternating with Li layers (Fig. 1).

However, as evident from high-resolution synchrotron data, the real structure is more complex and includes intergrowth of two polytypes (Table 1). Above 1173 K *RT*-LiNiB transforms to its high-temperature polymorph *HT*-LiNiB featuring different topology of [NiB] and Li layers (Fig. 1).

Previously we showed^{10,11} the hydride route is feasible for preparation of ternary lithium nickel borides providing fast reactions and accurate compositional control. The experimentally determined, $\text{LiH} : \text{Ni} : \text{B} = 1.3 : 1 : 1.15$, molar ratio results in phase-pure samples of LiNiB phases with minimal excess of LiH and B needed.¹¹ A further increase of the LiH ratio to values of 1.4 to 1.9 leads to stabilization of two new phases *RT*- $\text{Li}_{1+x}\text{NiB}$ and *HT*- $\text{Li}_{1+y}\text{NiB}$ (Table 1, Fig. 2). A slight excess of lithium hydride ($\text{LiH} : \text{Ni} : \text{B} = 1.4 : 1 : 1.15$) is necessary for the synthesis of the *HT*- $\text{Li}_{1+y}\text{NiB}$ compound. For stabilization of the *RT*- $\text{Li}_{1+x}\text{NiB}$ phase, a considerable excess of LiH

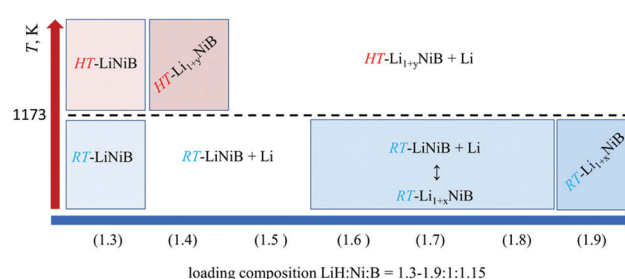


Fig. 2 Temperature-composition diagram for the synthesis of layered lithium nickel borides.

Table 1 Family of layered lithium nickel borides

Phase (space group)	$a, \text{\AA}$	$b, \text{\AA}$	$c, \text{\AA}$	$\alpha, {}^\circ$	$\beta, {}^\circ$	$\gamma, {}^\circ$	$V, \text{\AA}^3$	$V/Z, {}^e \text{\AA}^3$	Ref.
<i>RT</i> (<i>m</i>)-LiNiB ($P2_1/c$) ^a	4.6114	4.8333	6.156	90	109.607	90	129.26	32.32	11
<i>RT</i> *(<i>m</i>)-LiNiB ($P2_1/c$) ^b	18.451	4.8336	6.1573	90	109.636	90	517.21	32.33	11
<i>RT</i> (<i>o</i>)-LiNiB ($Cmc2_1$) ^a	4.8341	6.1573	8.6888	90	90	90	258.62	32.33	11
<i>RT</i> (<i>m</i>)- $\text{Li}_{1+x}\text{NiB}$ ($P2_1/c$) ^c	18.277(1)	4.86606(5)	6.1818(2)	90	107.623(1)	90	524.00(2)	32.75	This work
<i>RT</i> (<i>t</i>)- $\text{Li}_{1+x}\text{NiB}$ ($P1$)	18.2763(4)	4.86589(3)	18.5447(5)	89.9810(6)	107.6202(9)	90.056(2)	1571.82(3)	32.75	This work
<i>HT</i> -LiNiB ($P2_1/m$)	3.9095	8.8087	7.5234	90	90.070	90	259.08	32.39	11
<i>HT</i> (<i>m</i>)- $\text{Li}_{1+y}\text{NiB}$ ($P2_1/c$) ^d	3.92591(7)	7.5593(1)	8.8181(2)	90	92.6245(7)	90	261.42(1)	32.68	This work
<i>HT</i> (<i>t</i>)- $\text{Li}_{1+y}\text{NiB}$ ($P1$)	7.85184(9)	15.1183(2)	8.8179(1)	89.9079(7)	92.6271(5)	90.0008(5)	1045.63(3)	32.68	This work

^a Different polytypes of *RT*-LiNiB compound. ^b Superstructure of two polytypes of *RT*-LiNiB. ^c Idealized structure of *RT*- $\text{Li}_{1+x}\text{NiB}$ compound. ^d Idealized structure of *HT*- $\text{Li}_{1+y}\text{NiB}$ compound. ^e Volume normalized per formula units.

must be used ($\text{LiH} : \text{Ni} : \text{B} = 1.9 : 1 : 1.15$). Moreover, we found that synthesis of $RT\text{-Li}_{1+x}\text{NiB}$ compound requires that the sample be ground and annealed following the initial heating of precursors. After the first annealing, samples always comprise the $RT\text{-LiNiB}$ phase. $RT\text{-LiNiB}$ was the major phase and only a small amount of $RT\text{-Li}_{1+x}\text{NiB}$ compound was detected experimentally in the sample synthesized with loading composition $\text{LiH} : \text{Ni} : \text{B} = 1.6 : 1 : 1.15$. With a further increase in Li content, the fraction of $RT\text{-Li}_{1+x}\text{NiB}$ increases accordingly (Fig. S2†), while loading composition $\text{LiH} : \text{Ni} : \text{B} = 1.9 : 1 : 1.15$ gives phase pure $RT\text{-Li}_{1+x}\text{NiB}$ samples.

As evident from the high-resolution synchrotron powder X-ray diffraction (PXRD) data (Fig. 3), experimental PXRD patterns of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ compounds are similar. Both feature an intensive peak corresponding to the interlayer distance of $d \sim 4.34\text{--}4.41 \text{ \AA}$ between the [NiB] layers (yellow-shaded, Fig. 3). In contrast, the structure of [NiB] layers in two phases is different, resulting in the difference in PXRD patterns (blue-shaded, Fig. 3).

Crystal structures

Determination of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ crystal structures is challenging for several reasons: (1) extreme air-sensitivity (*vide infra*), (2) hydride synthesis did not yield large enough crystals suitable for laboratory single crystal X-ray diffraction; (3) the presence of light elements, Li and B, which makes X-ray data less informative, since heavier Ni atoms dominate X-ray scattering. Therefore, we used a synergistic combination of PXRD data refinement and theoretical calculations to elucidate crystal structures of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$. Solving structures from PXRD data yields the positions of Ni

atoms, thus the [NiB] layers (Tables S1–S6†). The structure of Li layers, including the atomic coordinates of extra-Li atoms in $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$, was determined using theoretical calculations.

Variations of [NiB] layers. Comparison of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ crystal structures with their respective parent phases $RT^*(m)\text{-LiNiB}$ and $HT\text{-LiNiB}$ are shown in Fig. 4 and 5. Additional Li atoms neither change the topology of [NiB] layers in the case of $RT\text{-Li}_{1+x}\text{NiB}$ nor the stacking sequence of [NiB] layers (Tables S1–S3†). Thus, $RT\text{-Li}_{1+x}\text{NiB}$ exhibits the same space group and Wyckoff sequence of the atomic sites as $RT^*(m)\text{-LiNiB}$ (Tables 1, S1–S3†), while variation in the monoclinic β -angle is responsible for variations in the PXRD pattern (Fig. 3 and 4). The stacking sequence of [NiB] layers for both $RT\text{-Li}_{1+x}\text{NiB}$ and $RT\text{-LiNiB}$ is ABCB (Fig. 4).

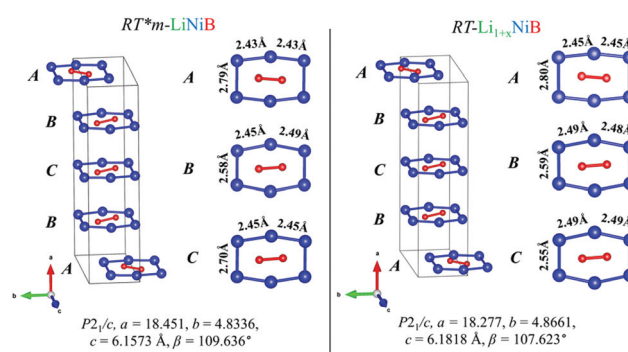


Fig. 4 Comparison between [NiB] layers (ABCB stacking sequence) of $RT\text{-LiNiB}$ and $RT\text{-Li}_{1+x}\text{NiB}$ compounds. Bond distances between Ni atoms in the $[\text{Ni}_6\text{B}_2]$ units are highlighted.

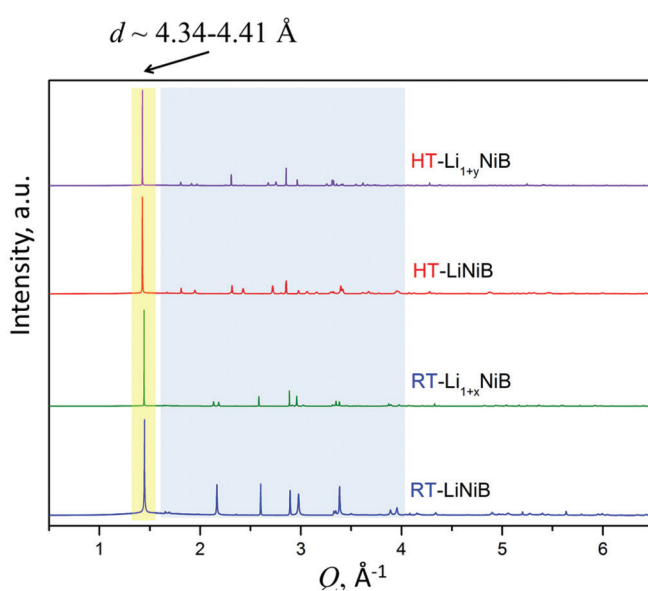
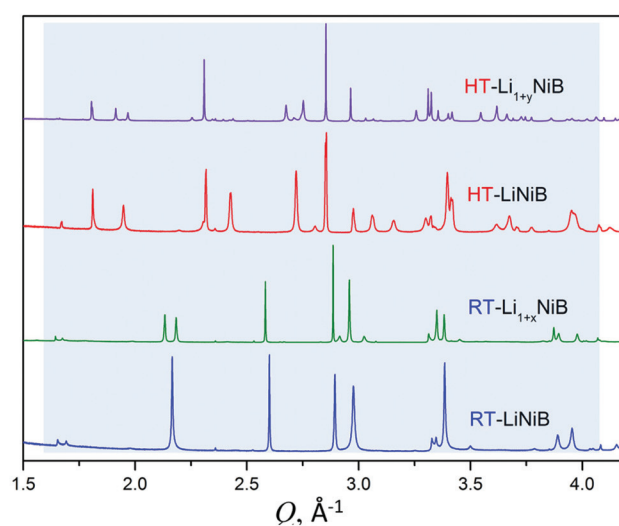


Fig. 3 Experimental synchrotron powder X-ray diffraction patterns of $RT\text{-LiNiB}$ [11], $HT\text{-LiNiB}$ [11], $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$. The most intense characteristic peak $d \sim 4.34\text{--}4.41 \text{ \AA}$ corresponds to the distance between [NiB] layers and is common for all phases. However, compounds have different atomic arrangement within the [NiB] layers, as indicated by the different sequence of the diffraction peaks in the blue-shadowed region, an enlarged view is shown on the right.



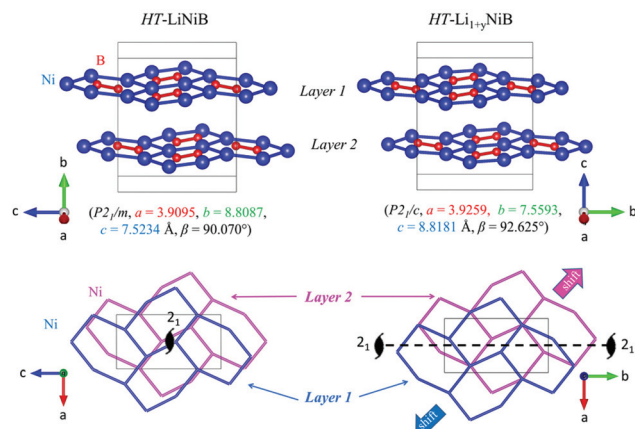


Fig. 5 Comparison between [NiB] layers in the structures of *HT*-LiNiB and *HT*-Li_{1+y}NiB compounds. The lower panel shows a relative shift between two adjacent [NiB] layers in the structures of *HT*-LiNiB and *HT*-Li_{1+y}NiB.

The unit cell volume of *RT*-Li_{1+x}NiB is larger than that of *RT*^{*(*m*)}-LiNiB, consistent with the incorporation of extra Li atoms (Table 1). This increased volume is due to the longer Ni–Ni distances within the Ni₆B₂ units leading to the slightly longer *b* and *c* parameters in the structure of *RT*-Li_{1+x}NiB compared to *RT*^{*(*m*)}-LiNiB.

The topology of the [NiB] layers in *HT*-LiNiB and *HT*-Li_{1+y}NiB compounds is similar. Here the fused Ni₆B₂ units form zig-zag chains within the layers. However, the stacking of [NiB] layers differs between the two compounds, resulting in the respective change of the space group from *P2*₁/*m* to *P2*₁/*c*, as well as in a swap of the *b*- and *c*-axes (Fig. 5 and Tables 1, S4–S6†). The [NiB] planar layers in *HT*-LiNiB are stacked along the two-fold screw axis 2₁, *e.g.* *b* axis is a stacking direction. On the contrary, for the *HT*-Li_{1+y}NiB phase the *c* axis is a stacking direction, thus perpendicular to the axis 2₁. The change in the space group arises from the more pronounced relative shift between the two adjacent [NiB] layers in structure of *HT*-Li_{1+y}NiB as compared to the *HT*-LiNiB structure (Fig. 5).

Li chemical pressure as a driving force for structural variation. The distance between [NiB] layers in the structures of *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB increases only slightly compared to the parent *RT*-LiNiB and *HT*-LiNiB, *d* ~ 4.34–4.41 Å, as evident from the high resolution synchrotron powder X-ray diffraction data (Fig. 3). The normalized volume per formula unit (*V/Z*) increases by 0.43 Å³ for *RT*-Li_{1+x}NiB and by 0.29 Å³ for *HT*-Li_{1+y}NiB compared to the respective parent compounds (Table 1). The increase of only ~1.3% is not sufficient to presume double Li-layers in the structure of *RT*-Li_{1+x}NiB or *HT*-Li_{1+y}NiB, as in the structure of theoretically predicted Li₂NiB compound.²⁹ Since the intensity of the PXRD peaks is dominated by scattering from Ni atoms, we utilized a constrained GA (genetic algorithm) search^{30,31} to shed light on the structure of Li layer in the *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB compounds.

Li layers in *RT*-Li_{1+x}NiB phase. A constrained GA search for the structure of the Li layers was performed while fixing the positions of Ni and B atoms to those in the experimentally determined structure (*P2*₁/*c* space group, Table 1). Since the complex “Li_{1+x}” packing can only be modeled by a superlattice, we performed the search on the supercell of the current lattice. The structure searching is very challenging for *RT*-Li_{1+x}NiB structure, because it already contains 48 atoms for *x* = 0 (*i.e.* Li₁₆Ni₁₆B₁₆). On the other hand, as indicated in Fig. 4, the layered structure of *RT*-Li_{1+y}NiB contains only AB- and BC-types of [NiB] stacking. Therefore, we searched for the Li layers between the AB and BC stacking types of [NiB] layers separately to reduce the computational cost. We built a smaller Li₈Ni₈B₈ lattice with only AB- or BC-type NiB layers. Two different searches with the lattices extended along *b* and *c* direction by 1 × 2 × 2 (*i.e.* Li_{32+m}Ni₃₂B₃₂ with *m* = 1, 2, 3, 4) and 1 × 3 × 3 (*i.e.* Li_{72+n}Ni₇₂B₇₂ with *n* = 2, 4, 6, 8, ..., 18) were performed. With 1 × 2 × 2 supercell, no energetically favored structures were found. Interestingly, as shown in Fig. 6a, the search with 1 × 3 × 3 supercell revealed an energetically stable Li packing with 16.7% excess of Li compared to the *RT*-LiNiB parent phase.

The Li layer configuration with *x* = 0.167 is stable in both AB-type and BC stacking types since they have the same formation energy as *RT*-LiNiB. Incorporation of extra Li atoms into the Li layer is accomplished by shortening of Li–Li distances to 2.6–2.8 Å, resulting in the denser hexagonal lattice (Fig. 6c) compared to *RT*-LiNiB (Li–Li distances ~2.8–3.1 Å). The configurations with intermediate Li excess 0 < *x* < 0.167 form combinations of loose (*x* = 0) and dense (*x* = 0.167) hexagonal lattices, resulting in a locally disordered packing motif and have an excess in the formation energy in a range of 3 meV (Fig. 6a).

Li layers in *HT*-Li_{1+y}NiB phase. We performed a similar GA search of the Li layers by implementing constraints to fix the

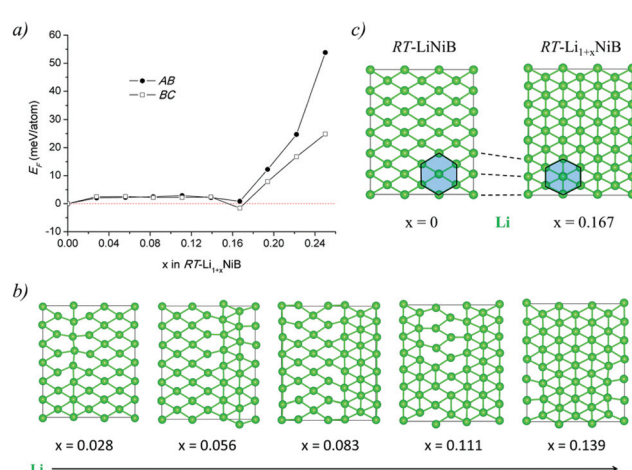


Fig. 6 (a) Energetics of *RT*-Li_{1+y}NiB structure as a function of *x*. Formation energies are calculated as $E_F = [E(\text{Li}_{1+y}\text{NiB}) - x \times E(\text{Li}) - E(\text{LiNiB})]/(3 + x)$, where the *P2*₁/*c* structure is used for *x* = 0. (b) Top views of the Li layers in the selected optimized structures are plotted. (c) Comparison of Li layer in *RT*-LiNiB and *RT*-Li_{1+y}NiB.



positions of Ni and B atoms as those in the experimentally determined structure of $HT\text{-Li}_{1+y}\text{NiB}$ (space group $P2_1/c$). Only Li positions were optimized during the structure searches. The $2 \times 2 \times 1$ supercell was used in the search with 32 Li atoms for $y = 0$. The number of excessive Li atoms ranges from 2 to 10, *i.e.* $\text{Li}_{32+k}\text{Ni}_{32}\text{B}_{32}$ with $k = 0, 2, 4, \dots, 10$. In the constraint searches, EAM (embedded atom method³²) classical potential selected from the AGA searches was used to screen the structures, and the final structures were then relaxed by DFT. For $y = 0$, Li atoms form a square lattice as plotted in the lower panel of the Fig. 7. The Li layer is similar to that experimentally found for $HT\text{-LiNiB}$.¹¹ With the increase of Li content, Li atoms tend to have coordination numbers larger than 4, *e.g.* 5 or 6 (Fig. 7a). Eventually, after adding enough Li atoms, such as in $\text{Li}_{42}\text{Ni}_{32}\text{B}_{32}$, Li atoms form a modulated hexagonal lattice. The addition of a small amounts of Li atoms lowers the formation energy of $HT\text{-Li}_{1+y}\text{NiB}$ at first (Fig. 7a), while for $y > 0.125$, the structure becomes less stable, even though with positive formation energies. The internal pressure induced by incorporating more Li atoms inside the structure as the function of y is plotted in Fig. 7b. If the unit cell is not allowed to expand (solid points), adding more Li atoms significantly increases this pressure for $y > 0.1$, again indicating the possible structural instability of $HT\text{-Li}_{1+y}\text{NiB}$ with large y . Therefore, we estimate the upper limit for y in $HT\text{-Li}_{1+y}\text{NiB}$ to be 0.125.

Overall structure. Careful analysis of synchrotron powder XRD patterns of the $RT\text{-Li}_{1+x}\text{NiB}$ or $HT\text{-Li}_{1+y}\text{NiB}$ compounds reveals the plethora of the low-intensity Bragg peaks (Fig. S3†). Part of these diffraction peaks can be accounted by superstructures, obtained by multiplying of the unit cell parameters by a factor of 2 or 3 and simultaneous lowering of the sym-

metry to triclinic: $a_{\text{superstr}} = a$, $b_{\text{superstr}} = b$, $c_{\text{superstr}} = 3 \times c$, $Z_{\text{superstr}} = 3 \times Z$ for $RT\text{-Li}_{1+x}\text{NiB}$; $a_{\text{superstr}} = 2 \times a$, $b_{\text{superstr}} = 2 \times b$, $c_{\text{superstr}} = c$, $Z_{\text{superstr}} = 4 \times Z$ for $HT\text{-Li}_{1+y}\text{NiB}$ (Tables 1 and S1–S6, cif files are provided in ESI†). We utilized these unit cells and created the structural models with [NiB] layers obtained from the PXRD data and Li layers found computationally, as described above. Rietveld refinement of these models resulted in a satisfactory fit (Tables S1, S4, Fig. S4–S11†), allowing for fitting of low intensity diffraction peaks while accounting for additional Li atoms between [NiB] layers. The obtained structural models are consistent with denser packing of Li atoms within the layer. As it is highlighted in Fig. 8, the denser packing of Li atoms results in 2 or 3 Li atoms per each Ni_6B_2 hexagonal unit, while solely 2 Li atoms per Ni_6B_2 are present in $RT\text{-LiNiB}$ and $HT\text{-LiNiB}$ parent structures (Fig. 8). This is accompanied by shortening of the interatomic distances between Li atoms. We also note that not all low intensity diffraction peaks can be accounted by the triclinic supercell (Fig. S3†), and the real structure of $RT\text{-Li}_{1+x}\text{NiB}$ or $HT\text{-Li}_{1+y}\text{NiB}$ might be even more complex (modulated or even larger cell), however the present model agrees fairly well with PXRD data, has reasonable bond distances and accounts for extra Li atoms in the structures. Most Li–B bonds in the structures of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ compounds have the same length as in the parent compounds ($d_{\text{Li–B}} \sim 2.3\text{–}2.4 \text{ \AA}$) (Fig. S12†). However, due to the distortion of Li-layers in the Li-enriched $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ compounds (Fig. 6 and 7), some of the Li atoms have longer distances to the nearest B atoms (2.4–2.6 \AA).

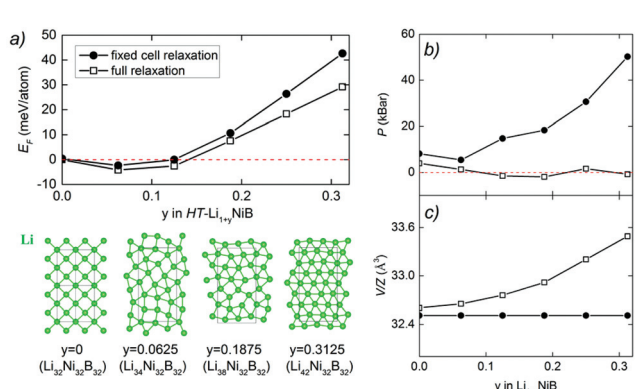


Fig. 7 (a) Energetics of $HT\text{-Li}_{1+y}\text{NiB}$ structure as a function of y . Formation energies are calculated as $E_F = [E(\text{Li}_{1+y}\text{NiB}) - y \times E(\text{Li}) - E(\text{LiNiB})]/(3 + y)$, where the $P2_1/m$ structure is used for $y = 0$. Top views of the Li layers in the optimized structures are plotted in the lower panel of (a). (b) Pressure of the $HT\text{-Li}_{1+y}\text{NiB}$ structure calculated as the function of y . (c) Volume per formula unit of the $HT\text{-Li}_{1+y}\text{NiB}$ generated structure calculated as the function of y . Note that here a $2 \times 2 \times 1$ supercell was used. Structure relaxations were carried out for two scenarios: unit cell fixed to the $2 \times 2 \times 1$ supercell during relaxation (solid circles) and full relaxation (hollow squares).

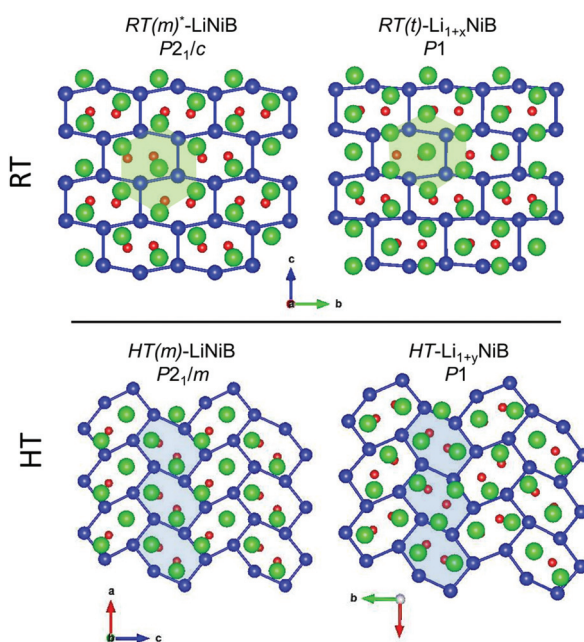


Fig. 8 Comparison of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ with the $RT\text{-LiNiB}$ and $HT\text{-LiNiB}$ parents. Light green and blue highlighted areas emphasize denser packing of Li atoms in the case of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$, leading to 3 Li atoms per each Ni_6B_2 distorted hexagon.

Temperature stability, reactivity and phase transformations

High-temperature synchrotron powder X-ray diffraction (HT-PXRD) was used to study the thermal stability of *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB. Samples were vacuum-sealed into silica capillaries for high temperature PXRD experiments. Upon heating Li is gradually “leached” from the structure, presumably reacting with silica. This leads to *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB transformation into *RT*-LiNiB and *HT*-LiNiB at 461 and 473 K, respectively (Fig. 9a and b). This again confirms the presence of extra Li atoms in *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB compared to their parent compounds. With a further increase in temperature above 684 K, *RT*-LiNiB transforms to Li-depleted LiNi₃B_{1.8} phase¹⁰ and binary Ni₂B. Between 684 K and 885 K, all three phases, *RT*-LiNiB, LiNi₃B_{1.8} and binary Ni₂B, are present. Above ~885 K, binary nickel borides Ni₂B, Ni₄B₃, Ni₃B are the major products.

Interestingly, HT-PXRD data of the *HT*-LiNiB polymorph (Fig. S13b†) shows considerable structural distortion, which was refined in the same structure type (same space group $P2_1/m$ and Wyckoff sequence) (Table S7†). The distorted structure

features substantially deformed Ni₆B₂ units within [NiB] layers and the relative shift of [NiB] layers (Fig. S13c and d†). This distorted structure was only observed in *in situ* PXRD experiments, while attempts to quench it from ~620 K were not successful. With further increase in temperature, the *HT*-LiNiB phase decomposes into Li-depleted LiNi₃B_{1.8}, and finally, binary Ni borides similarly to *RT*-LiNiB polymorph (Fig. S13†).

Air stability of *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB. In addition to the temperature, the transition between *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB compounds and their respective parents *RT*-LiNiB and *HT*-LiNiB can be driven by oxidation with oxygen in air. *HT*-Li_{1+x}NiB, if exposed to air, rapidly transforms to the *HT*-LiNiB phase (Fig. S14†). However, decomposition of the *RT*-Li_{1+x}NiB phase in air is considerably slower (Fig. 10). The higher reactivity of *HT*-Li_{1+x}NiB might originate from the considerably distorted Li-layer in the structure as well longer Li–B bonds, compared to the *RT*-Li_{1+x}NiB (Fig. 6, 7 and Fig. S12†). Powder X-ray diffraction patterns of *RT*-Li_{1+x}NiB show its evolution in air is completed within 160 min (Fig. 10). The process is relatively fast during the first ~1 hour (patterns in blue) and slows down afterwards. The last acquired powder pattern (at 160 min) contains mostly the *RT*-LiNiB compound with traces of the initial *RT*-Li_{1+x}NiB. Also, the fraction of LiOH·H₂O increases with the exposure time, suggesting that the overall process can be described as follows: $\text{Li}_{1+x}\text{NiB} + \text{O}_2 + \text{H}_2\text{O} \rightarrow \text{LiNiB} + x\text{LiOH}\cdot\text{H}_2\text{O}$, which again confirms the presence of extra-Li in *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB compounds.

Solid-state NMR spectroscopy

To further shed light on the evolution of Li and B coordination environment upon incorporation of extra-Li in the structures of *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB compounds, ⁷Li and ¹¹B solid-state magic angle spinning (MAS) NMR spectra were collected (Fig. 11a). The ⁷Li MAS NMR spectra of *RT*-Li_{1+x}NiB compound

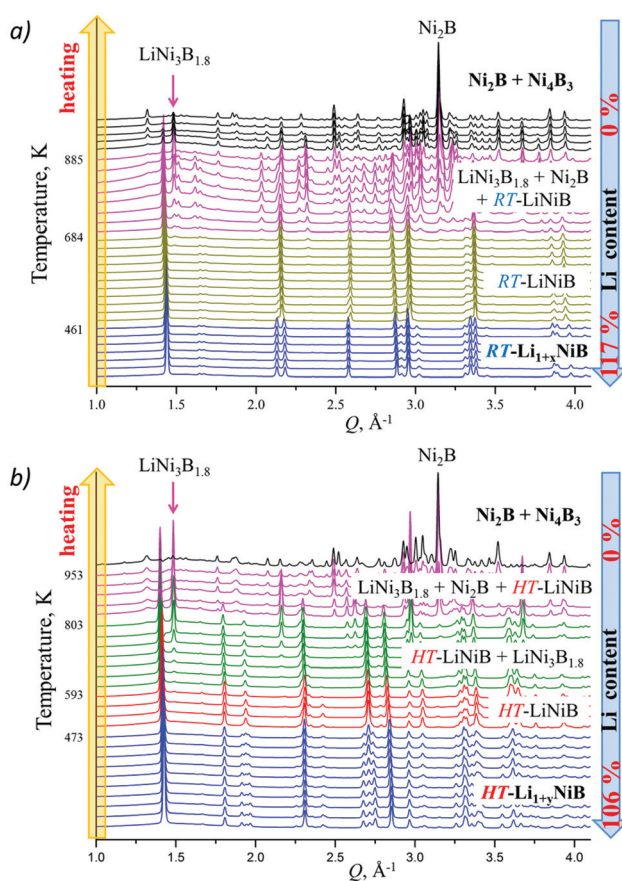


Fig. 9 High-temperature *in situ* powder X-ray diffraction patterns showing the equilibria in the Li–Ni–B system. With the increase of temperature Li is gradually “leached” from the compounds because of reaction with capillary material (SiO₂) involving (a) *RT*-Li_{1+x}NiB → *RT*-LiNiB → LiNi₃B_{1.8} → Ni₂B and Ni₄B₃ transformations; (b) *HT*-Li_{1+y}NiB → *HT*-LiNiB → LiNi₃B_{1.8} → Ni₂B and Ni₄B₃ transformations.

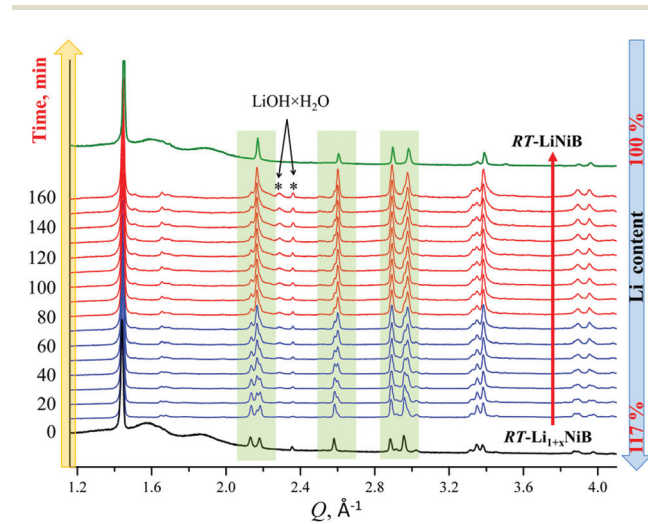


Fig. 10 X-ray diffraction patterns showing evolution of *RT*-Li_{1+x}NiB phase (bottom, black) upon exposure to air in 0–160 min range. Reference powder pattern of *RT*-LiNiB compound (top, green) is provided for comparison.



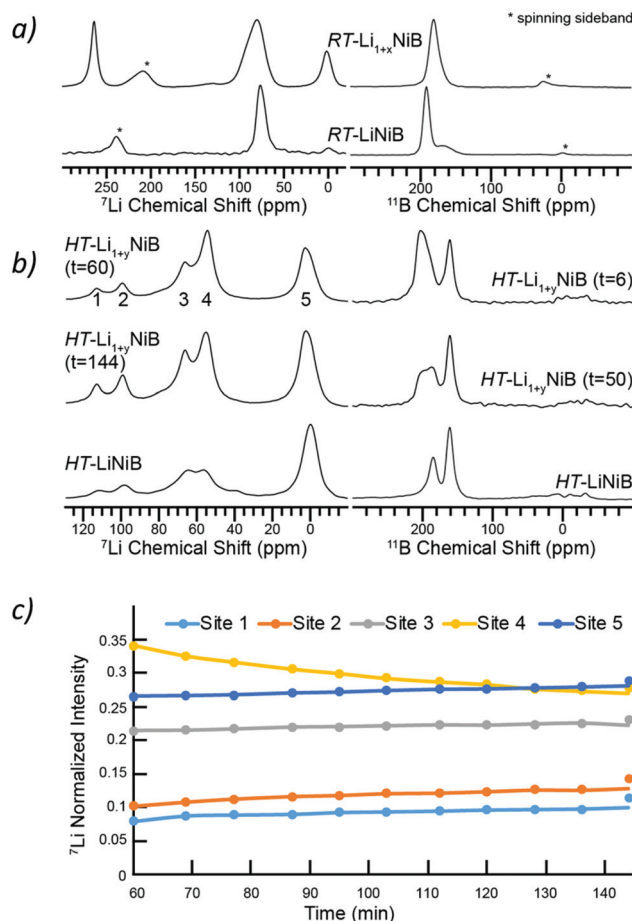


Fig. 11 (a) ⁷Li and ¹¹B MAS solid-state NMR spectra of *RT*-Li_{1+x}NiB in comparison to *RT*-LiNiB phase. (b) ⁷Li and ¹¹B MAS solid-state NMR spectra of *HT*-Li_{1+y}NiB in comparison to *HT*-LiNiB phase. The time in minutes after starting the magic angle spinning is indicated. (c) Variation in intensities of characteristic peaks (1, 2, 3, 4, 5) with time.

have two major peaks at \sim 265 and \sim 80 ppm and a minor one at \sim 0 ppm. The peak at 0 ppm originates from diamagnetic impurity phases, such as Li₂O (2.8 ppm), LiOH-H₂O (0.4 ppm), or LiOH (1.3 ppm), and is also present in ⁷Li MAS NMR spectrum of *RT*-LiNiB phase.¹¹ The peak at 265 ppm is most likely due to a Li metal impurity,³³ as a result of the significant excess of LiH required for synthesis and its further decomposition upon heating. The peak at \sim 80 ppm is attributed to the *RT*-Li_{1+x}NiB phase and is consistent with a single type of Li site expected in the structure. It is slightly positively shifted compared to the peak at \sim 75 ppm for *RT*-LiNiB phase but within the range of ⁷Li chemical shifts in other Li-containing intermetallics.^{9,11} The slight increase in the ⁷Li chemical shift for *RT*-Li_{1+x}NiB as compared to *RT*-LiNiB can likely be attributed to the subtle differences in the Li coordination environment in the two compounds. The increased width of the ⁷Li peak may be a result of the increased complexity of *RT*-Li_{1+x}NiB structure, where extra Li atoms are incorporated between the [NiB] layers, leading to a variety of local ⁷Li environments and distribution of shifts. The peak at \sim 80 pm

is shifted to higher frequencies compared to the diamagnetic signal, indicative of Knight shift from conduction electrons, and thus metallic behavior is expected for *RT*-Li_{1+x}NiB phase. The ⁷Li MAS NMR spectra of *HT*-Li_{1+y}NiB is akin to *HT*-LiNiB and is distinctive from that of *RT*-Li_{1+x}NiB and *RT*-LiNiB. There are two intense peaks at \sim 55–65 ppm and two peaks with lower intensity at \sim 95 and 110 ppm, in addition to the peak at \sim 0 ppm (diamagnetic impurity, *e.g.* Li₂O). Multiple peaks with positive chemical shifts are expected for *HT*-Li_{1+x}NiB, since it would be consistent with its metallic nature and a structure model with three Li sites (2a, 2c, 4e, *P*₂₁/*c*).

The NMR spectroscopy data also confirms the high reactivity of *HT*-Li_{1+y}NiB compound, which rapidly transforms in the air to *HT*-LiNiB compound (Fig. S14†). Fig. 11b shows the evolution of NMR spectra of a *HT*-Li_{1+x}NiB sample with time while the sample is undergoing magic angle spinning in a 2.5 mm rotor. The nitrogen gas used for spinning is approximately 99.9% pure and the 2.5 mm rotors are not gas-tight. Hence the reaction is likely driven by the slow diffusion of residual oxygen and water in the MAS gas into the sample. There are clear variations in intensities of five observed ⁷Li peaks, likely because of the changes in Li local environment between the layers. Li is also partly deintercalated and forms Li₂O, explaining the increased intensity of the 5th peak. During the course of MAS experiments, the ¹¹B solid-state NMR spectra of *HT*-Li_{1+y}NiB become more similar in appearance to *HT*-LiNiB.

The ¹¹B solid-state NMR spectrum for *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB is consistent with their metallic behavior. The 150–220 ppm range of ¹¹B chemical shifts is similar to the chemical shift observed for *RT*-LiNiB and *HT*-LiNiB compounds¹¹ and other metallic borides and borocarbides.^{9,34,35} Boron atoms in all *RT*-Li_{1+x}NiB, *RT*-LiNiB, *HT*-LiNiB, *HT*-Li_{1+y}NiB layered phases form B–B pairs, which are further surrounded by 6 Ni atoms (Fig. 1, 4 and 5). However, the different topology of the layers, as well as differences in boron coordination by Li, likely explains the observed differences in the ¹¹B solid-state NMR spectra.

Magnetic properties

Temperature dependence of the molar magnetic susceptibility in the range 5–300 K of *RT*-Li_{1+x}NiB and *HT*-Li_{1+y}NiB (Fig. S15†) is consistent with both phases being temperature-independent Pauli paramagnets. No magnetic ordering was observed down to 5 K. The field dependence of the magnetization in the same temperature range (Fig. S15†) at high magnetic field (10 000 Oe–50 000 Oe) is linear with no tendency for saturation, which is again consistent with temperature independent paramagnetism. Small deviation from linearity of the field dependence of the magnetization at low magnetic field (0–10 000 Oe, *see* Fig. S15†) could be attributed to some magnetic moment bearing impurity, concentration of which is under the detection limit of X-ray analysis (below 1%). The intrinsic susceptibility values were estimated to be in the range of $0.7\text{--}1.2 \times 10^{-4}$ emu mol^{−1} (Fig. S15†). The parent compounds *RT*-LiNiB and *HT*-LiNiB also exhibit temperature-independent paramagnetism.¹¹ Thus, incorporation of additional



Li atoms does not affect the magnetic properties. This is in stark contrast to the partial deintercalation of Li from paramagnetic LiNiB, inducing magnetic ordering in $\text{Li}_{1-\delta}\text{NiB}$.¹²

Experimental

Synthesis

Starting materials for the synthesis were powders of lithium hydride (AlfaAesar, 99.4%), nickel (AlfaAesar, 99.996%), and boron (AlfaAesar, amorphous & crystalline, 98%). All manipulations of the starting materials and samples were performed in an argon-filled glovebox with $p(\text{O}_2) < 1 \text{ ppm}$. Powders of the starting materials (total mass = 0.3 g) were weighted in $\text{LiH} : \text{Ni} : \text{B} = 1.9 : 1 : 1.15$ ($RT\text{-Li}_{1+x}\text{NiB}$) and $1.4 : 1 : 1.15$ ($HT\text{-Li}_{1+y}\text{NiB}$) molar ratios and loaded into polystyrene grinding vial set with slip-on cap. These loading compositions were determined empirically to get the highest yield of the target phases. The vials were then sealed into double-sided polypropylene bag under argon atmosphere and brought out for the powder mixing/milling using ball-mill (8000M MIXER/MILL) for 18 min (vials were enclosed in poly-bags during the process to ensure O_2 -free environment). Afterward, freshly prepared fine powders were loaded in Ta tubes inside argon-filled glovebox. The metal containers were sealed by arc-melting and placed into silica reactors equipped with Swagelok safety check valves to prevent over-pressurization of the reactors due to the release of hydrogen gas during the heat treatment. The silica reactors were evacuated down to 4×10^{-5} Bar and slowly (1.25 K min^{-1}) heated from room temperature to 1173 K, held at that temperature for 12 hours and quenched into cold water for the preparation of $HT\text{-Li}_{1+y}\text{NiB}$ phase or cooled to room temperature by switching off furnace in the case of $RT\text{-Li}_{1+x}\text{NiB}$ phase. The second annealing of the manually ground sample loaded to the Ta tube using the same temperature profile was necessary to obtain $RT\text{-Li}_{1+x}\text{NiB}$, otherwise $RT\text{-LiNiB}$ compound forms. The synthesized compounds are air and moisture sensitive.

Powder X-ray diffraction

The purity of polycrystalline samples was checked using Rigaku MiniFlex600 powder diffractometer with $\text{Cu K}\alpha$ radiation ($\lambda = 1.54051 \text{ \AA}$) and $\text{Ni-K}\beta$ filter. Data were collected in zero-background plate holder for air-sensitive samples at room temperature. Phase analysis was performed using the PDF-2 database incorporated into PDXL2 program software.³⁶ Powder diffractograms during controlled samples oxidation in the air were collected using open zero-background plate holder with ~ 10 min time intervals.

Crystal structure determination from synchrotron powder X-ray diffraction data

High-resolution synchrotron powder diffraction data were collected at beamline 11-BM-B Advanced Photon Source (APS), Argonne National Laboratory at 295 K and 100 K and $\lambda = 0.412815 \text{ \AA}$ for a sample of $RT\text{-Li}_{1+x}\text{NiB}$ and $\lambda = 0.412818 \text{ \AA}$ for

a sample of $HT\text{-Li}_{1+y}\text{NiB}$. Data were collected at ambient pressure.

The unit cell symmetry, metrics, and atomic arrangement in the structures of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ were obtained using program FOX for the *ab initio* structure determination from powder diffraction.³⁷ After indexing, positions of heavy Ni atoms were determined and further refined in the triclinic space group *P*1. After the atomic coordinates of heavy Ni atoms were found, the structure was analyzed with the program Platon,³⁸ and the correct space group was determined. The residual electron density peaks were assigned to tentative B and Li atomic positions based on the analysis of interatomic distances.

RT-Li_{1+x}NiB. Indexing of the high-resolution synchrotron powder X-ray diffraction data using program FOX³⁷ resulted in the *P*-monoclinic unit cell (space group $P2_1/c$, $a = 18.277$, $b = 4.8660$, $c = 6.1818 \text{ \AA}$, $\beta = 107.623^\circ$, $V = 524.00 \text{ \AA}^3$ at 295 K, $Z = 16$). Ni atoms occupy four 4e sites. Further refinements showed residual electron density peaks, which were assigned to be B in four 4e sites and Li in four 4e sites based on the analysis of interatomic distances.

HT-Li_{1+y}NiB. Indexing of the high-resolution synchrotron powder X-ray diffraction data using program FOX³⁷ resulted in *P*-monoclinic unit cell (space group $P2_1/c$, $Z = 8$, $a = 3.92591$, $b = 7.5593$, $c = 8.8181 \text{ \AA}$, $\beta = 92.6245^\circ$, $V = 261.42 \text{ \AA}^3$). Ni atoms occupy two 4e sites. Further refinements showed residual electron density peaks assigned to be B in 4e, 4e, and Li occupying 2a, 2c, 4e sites based on the analysis of interatomic distances.

Rietveld refinement with the GSAS II software package³⁹ was further performed using the derived structural models. The profile parameters, background parameters, zero shift, and cell parameters were refined first. The background was fitted using a shifted 24-points linear interpolation function, a pseudo-Voigt function was applied to generate the profile shape. Atomic coordinates of Ni atoms were refined, while atomic coordinates of light B and Li atoms were fixed to the values obtained computationally. Experimental details and crystallographic data are compiled in Tables S1–S3, Fig. S4–S7† ($RT\text{-Li}_{1+x}\text{NiB}$), and Tables S4–S6, Fig. S8–S11† ($HT\text{-Li}_{1+y}\text{NiB}$). The final refinements of the structures were performed with isotropic displacement parameters for all atoms. Crystal structures were visualized using the program VESTA 3.⁴⁰

In situ high-temperature synchrotron powder X-ray diffraction (HT-PXRD). Transformation of the $RT\text{-LiNiB}$, $HT\text{-LiNiB}$, $RT\text{-Li}_{1+x}\text{NiB}$, and $HT\text{-Li}_{1+y}\text{NiB}$ compounds at elevated temperatures was evaluated from *in situ* high-temperature synchrotron X-ray powder diffraction data collected at beamline 17-BM-B APS ($\lambda = 0.24158 \text{ \AA}$). Powdered samples were filled into 0.7 mm outer diameter thick-walled (0.1 mm) silica capillaries and sealed under vacuum. Capillaries were mounted on a standard base equipped with two coiled resistance heaters. The further details of experimental setup can be found elsewhere.⁴¹ High energy synchrotron X-ray diffraction spectra were collected in the temperature range 295–1000 K.

Solid-state NMR spectroscopy. ^7Li and ^{11}B solid state NMR spectra were measured on a Bruker widebore 9.4 T ($\nu_{1\text{H}} =$



400 MHz) NMR magnet equipped with a Bruker Avance III HD spectrometer. All spectra were acquired using a 2.5 mm triple resonance MAS probe. Samples were diluted to approximately 50 weight% in powdered polyfluoroethylene to minimize conductivity effects and packed into MAS rotors while still under inert atmosphere. The MAS rate was 25 kHz in all cases.

Magnetic properties. Magnetic measurements were performed on polycrystalline samples ($m = 50$ mg), which were loaded under an inert atmosphere in EPR quartz tubes and sealed under vacuum. The data were collected using a Quantum Design MPMS XL and MPMS3 SQUID Magnetometer DC. Magnetic susceptibility was measured in an applied field of 1000 Oe, in the 5–300 K range. Field-dependent measurements were carried out at 2 K, 5 K and 300 K.

Computational methods. The crystal structure searches were performed using the adaptive genetic algorithm (AGA).^{30,31} AGA integrates auxiliary interatomic potentials and first-principles calculations in an adaptive manner to ensure high efficiency and accuracy. Interatomic potentials based on the embedded-atom method (EAM)³² were chosen as the auxiliary classical potentials to describe the Li–Ni–B system (see ref. 11 for more details). The packing of Li atoms in $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ were determined using constraint genetic algorithm (GA) searches by fixing the atomic positions on the NiB layers. During the searches, no symmetry constraint was applied. The EAM potential obtained from the AGA search was also used to screen the high-energy structures, the final structures from constraint GA were relaxed and evaluated by first-principles calculations.

First-principles calculations were carried out using the density functional theory (DFT) within a generalized gradient approximation (GGA) of the exchange–correlation functional as implemented in the VASP code.^{42,43} The projector augmented-wave method⁴⁴ was used to describe the valence configuration: $1s^2 2s^1$ for Li, $3d^8 4s^2$ for Ni, and $2s^2 2p^1$ for B. The GGA exchange–correlation energy functional parametrized by Perdew, Burke, and Ernzerhof was used.⁴⁵ A plane-wave basis was used with a kinetic energy cutoff of 650 eV. The Monkhorst–Pack's scheme⁴⁶ was adopted for Brillouin zone sampling using a k -point grid with a spacing of $2\pi \times 0.033 \text{ \AA}^{-1}$, and the ionic relaxations stopped when the forces on every atom became smaller than 0.01 eV \AA^{-1} .

Conclusions

Li chemical pressure is a driving force for the formation of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$, which accommodate up to 16.7% and 12.5% excess Li into their structure compared to their parent compounds $RT\text{-LiNiB}$ and $HT\text{-LiNiB}$. The crystal structures of $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ compounds are similar to that of $RT\text{-LiNiB}$ and $HT\text{-LiNiB}$, built up from alternating Li layers with hexagonal or squared motif respectively, and [NiB] layers. The complex Li environments in layers of these two new compounds were determined using DFT methods, while subtle variations in the [NiB] layers were elucidated from synchrotron

powder X-ray diffraction. Deintercalation of Li from the [NiB] layers was observed at elevated temperatures resulting in the gradual “leaching” of Li from the layers of these Li-enriched compounds and their subsequent transformation into their parent phases: $RT\text{-Li}_{1+x}\text{NiB} \rightarrow RT\text{-LiNiB}$ at 461 K and $HT\text{-Li}_{1+y}\text{NiB} \rightarrow HT\text{-LiNiB}$ at 473 K. Upon further heating decomposition into $\text{LiNi}_3\text{B}_{1.8}$ and finally, into binary nickel borides (~ 900 K) takes place. Upon oxidation in air at room temperature, both $RT\text{-Li}_{1+x}\text{NiB}$ and $HT\text{-Li}_{1+y}\text{NiB}$ phases slowly release Li and transform to $RT\text{-LiNiB}$ and $HT\text{-LiNiB}$ compounds, respectively. Controlled oxidation and deintercalation as a route of further modification of layered LiNiB structures is a subject of future investigation.

Conflicts of interest

There are no conflicts to declare.

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