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First-principles calculated decomposition pathways for LiBH₄ nanoclusters

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We analyze thermodynamic stability and decomposition pathways of LiBH₄ nanoclusters using grand-canonical free-energy minimization based on total energies and vibrational frequencies obtained from density-functional theory (DFT) calculations. We consider (LiBH₄)ₙ nanoclusters with n = 2 to 12 as reactants, while the possible products include (Li)ₙ (B)ₙ (LiB)ₙ and LiₙBH₄: off-stoichiometric LiₙBₙH₄ (m ≤ 4n) clusters were considered for n = 2, 3, and 6. Cluster ground-state configurations have been predicted using prototype electrostatic ground-state (PEGS) and genetic algorithm (GA) based structural optimizations. Free-energy calculations show hydrogen release pathways markedly differ from those in bulk LiBH₄. While experiments have found that the bulk material decomposes into LiH and B, with Li₃B₁₂H₄ as a kinetically inhibited intermediate phase, (LiBH₄)ₙ nanoclusters with n ≤ 12 are predicted to decompose into mixed LiₙBₙ clusters via a series of intermediate clusters of LiₙBₙH₄ (m ≤ 4n). The calculated pressure-composition isotherms and temperature-pressure isobars exhibit sloping plateaus due to finite size effects on reaction thermodynamics. Generally, decomposition temperatures of free-standing clusters are found to increase with decreasing cluster size due to thermodynamic destabilization of reaction products.

Hydrogen is a promising next-generation energy carrier due to its high energy density, high energy conversion efficiency, and absence of harmful emissions. However, on-board hydrogen storage is still a serious challenge for developing economically viable hydrogen-powered passenger vehicles.¹ To achieve widespread commercialization, hydrogen storage systems should simultaneously possess several characteristics such as safety, high gravimetric and volumetric densities, fast reversible hydrogen release and uptake under moderate pressures and temperatures matched to the operating conditions of proton exchange membrane (PEM) fuel cells.

Complex metal hydrides emerged as feasible high-density hydrogen storage materials after Bogdanović and Schwickardi demonstrated reversible (de)hydrogenation reactions in transition metal doped sodium alanate (NaAlH₄)³. Spurred by this discovery, many other complex hydrides have been explored as candidate hydrogen storage materials, including alanates, amides, and borohydrides. Lithium borohydride, LiBH₄, has received particular attention owing to its high gravimetric (18.5 wt.% H₂) and volumetric (121 g H₂/L) densities. However, due to its high thermodynamic stability and slow kinetics, hydrogen release from bulk LiBH₄ requires very high temperatures that are incompatible with proton-exchange membrane (PEM) fuel cells³⁻⁵. In addition, the tendency to release diborane during dehydrogenation leads to irreversibility and causes poisoning of the fuel cell⁶⁻¹⁰.

Destabilization has been extensively explored as a means of improving the thermodynamics of hydrogen storage reactions¹¹⁻¹⁴. The main idea of this approach is to add a second reactant, which upon decomposition forms a stable low-energy product phase and lowers the overall reaction enthalpy. The canonical example of a destabilized reaction is LiBH₄ + MgH₂ → LiH + MgB₂ + (5/2)H₂, where the addition of MgH₂ leads to the formation of MgB₂ as a low-energy product phase. This significantly lowers the reaction enthalpy relative to the decomposition reaction of the pure compound, LiBH₄ → LiH + B + (3/2)H₂. Unfortunately, destabilization is not very effective in lowering the kinetic barriers to hydrogen release, leading to only modest improvement in the hydrogen release temperatures.

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Another approach to improving the thermodynamics and kinetics of hydrogen release consists of using LiBH₄ nanoparticles incorporated in support materials with nanoscale pores, such as nanoporous carbon or metal-organic frameworks (MOFs). Besides improving the kinetics and lowering the temperature of hydrogen release, this strategy has also been shown to suppress the formation of diborane (H₂B₂), which is a major byproduct in the dehydrogenation of LiBH₄. First-principles density-functional theory (DFT) based studies have investigated the thermodynamic properties of nano-LiBH₄, suggesting that bonding with the nanoporous carbon support plays a key role in improving the thermodynamic properties of nano-LiBH₄. However, due to the large configurational space, these studies did not perform comprehensive examination of the most stable cluster structures and compositions as functions of cluster size. Hence, a complete picture of the size-dependent thermodynamics properties of LiBH₄ nanoparticles and their decomposition products is not yet available.

Decomposition pathways and pressure–composition isotherms are determined using free energy minimization in the grand canonical ensemble. In addition to clusters with the chemical composition of bulk LiBH₄ and its predicted bulk decomposition products Li₂B₆H₁₂, LiH and B₃0, we considered (Li)ₙ, (LiB)ₙ, and Li₂BH₄ clusters as possible products. To assess the role of other cluster compositions, further detailed studies were performed for the decomposition reactions of (LiBH₄)ₙ, (LiBH₄)₀ and (LiBH₄)₂ including Li₂B₆H₁₂ (n = 1 to 7), Li₂BH₄ (n = 1 to 11) and Li₂B₆H₈ (n = 1 to 23) clusters as possible intermediate products. Our analysis shows that the reaction end products are changed from (LiH)ₙ + Bₖ + (3/2)H₂ to (LiB)ₙ + 2mH₂ when the size of reactants is reduced to the nanoscale. The calculated reaction enthalpies decrease from 238 (n = 2) to 133 kJ/mol H₂ (n = 12) with increasing cluster size, remaining significantly above the bulk decomposition enthalpy of LiBH₄. The hydrogen release temperatures are predicted to behave in a non-trivial way, with multiple intermediates appearing in the simulated dehydrogenation processes. The finite size effects and presence of intermediates manifest in multiple plateaus in the calculated release curves as functions of temperature, while the calculated pressure-composition isotherms and temperature-composition isobars exhibit sloping plateaus. Our study provides an in-depth understanding of the thermodynamics of hydrogen release from nanoscale particles and shows the importance of engineering appropriate support materials that can bind the reaction products and lower the reaction enthalpies to achieve reversible hydrogen storage in a practically viable range of temperatures and pressures.

Results

Structure of reactants and products. We start by briefly describing our findings for the atomic geometries of the reactants and products, which are summarized in Fig. 1. The reactants, LiBH₄ nanoclusters, are composed of the cation, Li⁺, and the anionic group, BH₄⁻. We find that the lowest energy conformations of small (LiBH₄)ₙ clusters are chairlike, forming closed rings. The tetrahedral BH₄⁻ anion is a near perfect tetrahedron in all cases with some elongation in the B–H bond length from 1.22 to 1.24 Å. Note that the length of the B–H bond in bulk LiBH₄ is 1.22 Å. The diameter of LiBH₄ clusters increases gradually from 6.0 to 10.4 Å upon increasing the cluster size from n = 2 to 12. (LiH)ₙ clusters form rock salt-derived structures for sizes n ≥ 4.

For boron clusters, our GA method reproduced the lowest energy structures found in earlier studies. Small boron clusters are found to form 2-dimensional planar structures except for the B₆ cluster which has two boron atoms on two sides of the circular substructure with seven boron atoms forming a ring. Changes in the cluster geometries with increasing number of boron atoms indicate that boron nanoclusters prefer to have a coordination number of six.

For pure lithium clusters, the GA again finds the same structures as those in previous studies. Unlike boron clusters, lithium clusters agglomerate in three-dimensional motifs when the cluster sizes exceed four atoms.

In the mixed lithium–boron clusters (LiB)ₙ, the boron atoms generally tend to gather in the center of the cluster with lithium atoms surrounding them on the periphery. In addition, for n ≤ 5 the arrangement of boron atoms is the same as in the pure boron clusters. When n > 5, boron atoms in the center are distorted due to interactions with the surrounding lithium atoms. We also found that the B–B distances in the center of the (LiB)ₙ clusters are gradually increased from 1.53 Å for n = 2 to 1.85 Å for n = 12, while the Li–B bond lengths stay approximately constant. It appears that interactions with the outer Li ions and charge transfer from Li to B combine to push the B atoms outward.

Cluster energies. To better understand the thermodynamic stability, reaction pathways and reaction enthalpies, we first discuss the cluster-size dependent energetics of reactants and reaction products. Figure 2 shows the relative energies of clusters normalized by their bulk energies without the zero-point energy (ZPE) corrections. This quantity is defined as the ratio of the energy of the cluster per formula unit versus the energy of the bulk phase. Bulk lithium is calculated in the Im3m body-centered cubic lattice, bulk boron in the α-B phase, and bulk LiH is in the rocksalt structure. Bulk LiBH₄ has several near-degenerate ground-state structures, and the Pnma structure is used in our calculations. As for bulk LiB, we used a P6₃/mmc structure reported in a previous experimental study. The calculated reaction enthalpy for the decomposition of bulk LiBH₄ at T = 0 K is ΔH = 60 kJ/(mol H₂) with zero-point energy (ZPE) corrections, while excluding ZPE we obtain 82 kJ/(mol H₂). The predicted decomposition temperature is Tdec = 340 °C at p = 1 bar hydrogen pressure, which is about 100 °C below experimental results. We find that the stabilities of the reaction products are dramatically reduced with decreasing cluster size, with LiH and LiB clusters being more stable than pure Li and B clusters. However, the behavior of the reactant LiBH₄ clusters is very different: the energy per formula unit of (LiBH₄)ₙ is nearly flat from n = 2 to n = 12, which signals that their thermodynamic stability is reduced only slightly upon decreasing cluster size. This suggests that most of the binding energy is stored in the polar covalent B–H bonds, while the electrostatic interactions between lithium and BH₄⁻ complexes are relatively weak. Examination of the size-dependent stabilities of each cluster type shows that clusters with even-numbered formula units are generally more stable than the odd ones. For pure clusters, Liₙ and Bₙ we find that Liₙ, Li₁₄, Li₁₅, Li₁₁, B₇, B₁₅, and B₁₁ possess certain geometric symmetries, but they are unstable with respect to decomposition into even-numbered n + 1 and n − 1
clusters due to unpaired electron filling in the highest occupied molecular orbital. As for the LiH and LiB clusters,
even though the highest orbitals in (LiH)$_n$, (LiH)$_{11}$, and (LiB)$_n$ are fully occupied, these clusters are unstable because of broken geometric symmetry. Moreover, (LiH)$_{10}$ and (LiB)$_5$ are also unstable because they are at the structural transition point between distinct bonding topologies.

Decomposition pathways of LiBH$_4$ clusters. Previous computational work$^{30}$ has shown that, thermodynamically, decomposition of bulk LiBH$_4$ should occur via the formation of an intermediate closobarane Li$_2$B$_{12}$H$_{12}$ compound. However, the formation of closobaranes is kinetically inhibited in experiments and decomposition proceeds directly to a mixture of LiH and B. We begin our discussion of the decomposition pathways of LiBH$_4$ nanoclusters by considering simplified decomposition reactions with closobarane Li$_2$B$_n$H$_n$, elemental (Li)$_n$ and (B)$_n$, and binary (LiH)$_n$ and (LiB)$_n$ clusters as the end products; the full treatment using the ensemble grand canonical formalism is given below. First, we consider the following reaction pathways:

\[
\text{(LiBH}_4\text{)}_n \rightarrow \text{(LiH)}_n + \text{B}_n + \frac{3n}{2}\text{H}_2, \quad (1)
\]

\[
\text{(LiBH}_4\text{)}_n \rightarrow \text{Li}_n + \text{B}_n + 2n\text{H}_2, \quad (2)
\]
Equations 1 and 2 are the nanocluster analogues of the bulk decomposition reactions occurring at $T = 400$ and $900 \, ^\circ\mathrm{C}$, respectively. Reaction 4 is the nanocluster analogue of the bulk decomposition reaction into $\text{Li}_2\text{B}_{12}\text{H}_{12}$, accounting for the formation of various closoborane ($\text{B}_n\text{H}_n$)$_2^-$ species with $n \leq 12$, while the product of Eq. 1 can be obtained by further decomposition of a mixture of $\text{Li}_2\text{B}_n\text{H}_n$ and $(\text{LiH})_{n-2}$. Finally, Eq. 3, is similar to the favored decomposition reaction predicted for nano-$\text{NaAlH}_4$.

The calculated DFT reaction enthalpies of Eqs 1–4 without vibrational free energy contributions are shown in Fig. 3. In all cases, the calculated DFT reaction enthalpies are larger than the decomposition enthalpy of bulk $\text{LiBH}_4$ $[82 \, \text{kJ/(mol H}_2\text{)}]$. This does not mean that $\text{LiBH}_4$ nanoclusters are generally more stable with respect to decomposition than the bulk compound because these data are based on a limited number of possible reactions and reaction products; we will show below that the Gibbs free energy minimization approach and inclusion of product clusters with intermediate hydrogen content can lower the reaction enthalpies significantly. The trend towards higher reaction enthalpies for the hypothesized reactions in Eqs 1–4 can be easily explained from the size-dependence of the cluster binding energies in Fig. 2: since the $(\text{LiBH}_4)_n$ clusters show weak size dependence, while all the product clusters show much stronger finite size energy penalty, the reaction enthalpies are significantly higher than the bulk limit. It is interesting to note that of all four reaction pathways, the bulk-like Eq. 1 is:

$$\text{(LiBH}_4)_n \rightarrow (\text{LiB})_n + 2n\text{H}_2,$$

$$\text{(LiBH}_4)_n \rightarrow \text{Li}_2\text{B}_n\text{H}_n + (\text{LiH})_{n-2} + (n + 1)\text{H}_2.$$

Figure 2. Calculated DFT total energies (without zero-point energy corrections) of $(\text{LiBH}_4)_n$, $(\text{LiH})_n$, $(\text{LiB})_n$, $\text{B}_n$ and $\text{Li}_n$ clusters, normalized to the corresponding bulk values.

Figure 3. Calculated DFT reaction enthalpies (without ZPE correction) of the four decomposition pathways of nano-$\text{LiBH}_4$ given by Eqs 1–4, shown as functions of the cluster size $n$. 

$$\text{(LiBH}_4)_n \rightarrow (\text{LiB})_n + 2n\text{H}_2,$$  

$$\text{(LiBH}_4)_n \rightarrow \text{Li}_2\text{B}_n\text{H}_n + (\text{LiH})_{n-2} + (n + 1)\text{H}_2.$$
the least favorable one in nano-LiBH₄, which shows that the reaction path can change dramatically upon reducing the particle size to the nanometer regime; similar effects have been predicted for sodium alanate²⁹,³⁹. Both pathways in Eqs 2 and 3 liberate all the hydrogen, and our results show that Eq. 3 is more favorable than Eq. 2 because the binary (LiB)ₙ clusters are more stable than the mixture of (Li)ₙ and (B)ₙ clusters. We note that the formation of Li₂BₙHₙ clusters via Eq. 4 is the preferred path for \( n = 6 \) to 12 when closoborane species form as the end product, while the pathway in Eq. 3 is preferred for \( n = 3 \) to 5. In general, we find that the calculated reaction enthalpies decrease with increasing cluster size \( n \), and are significantly higher than the calculated DFT enthalpies for decomposition of bulk-LiBH₄ into Li₂B₁₂H₁₂ ³⁰. Due to the high stability of \( -BH₁₂ \) complex anions, the reaction enthalpy of Eq. 4 drops significantly from \( n = 11 \) to \( n = 12 \), and one may expect that these species will persist as the dominant end product for \( n > 12 \).

To investigate the possibility that intermediate products other than those in Eqs 1–4 may form during decomposition, we have carried out calculations for clusters with the general formula LiₙBₙHₘ, where \( 0 < m < 4n \) and \( n = 2, 3, \) and 6. For this series, we adopted the random search scheme rather than the genetic algorithm. The random search scheme is implemented as follows. New initial structures were randomly generated by removing one or more H atoms from nano-LiBH₄ and by gradually adding H atoms to LiB nanoclusters, breaking symmetries by slightly distorting all atoms away from the original positions. These randomly generated structures were then relaxed using DFT forces to a local minimum configuration, and the energies of the most favorable structures at each composition were used in the Gibbs ensemble formalism given by Eqs S1–S3 (See Supplementary Information). The zero point energy, temperature-dependent vibrational free energy, and the free energy of the diatomic H₂ gas are included in these calculations. Our results for the cluster type fractions \( p_f \) are summarized in Fig. 4 assuming a hydrogen pressure of \( p = 0.01 \) bar, which is representative of experimental measurements.

Figure 4(a) shows the calculated dehydrogenation pathway of (LiBH₄)₂ and the structures of the predicted intermediate products as functions of temperature. (LiBH₄)₂ nanoclusters start to release hydrogen at approximately 450 °C by first forming Li₂B₂H₆. The structure of this cluster resembles diborane, with a significant difference that the donation of two electrons from Li atoms reconfigures the three-center two-electron bonds of diborane into a direct B-B bond. At 500 °C, Li₂B₂H₄ clusters form and exist as the dominant species from 580 to 980 °C. Finally, Li₂B₂H₂ clusters appear at 800 °C and dominate until full dehydrogenation into (LiB)₂ as the end product above 1400 °C (not shown). We note that the intermediate products in the dehydrogenation pathway of (LiBH₄)₂ all contain an even number of hydrogen atoms, and each intermediate reaction step releases one H₂ molecule per cluster.

Figure 4(b) shows the decomposition pathway of (LiBH₄)₃. It is seen that hydrogen release starts at 530 °C where two H₂ molecules per cluster are released to form Li₃B₃H₆, which exists in a narrow temperature interval. The latter cluster is essentially a superposition of Li₃B₃H₄ and (LiBH₄)₂ clusters, which explains its marginal stability. In the next step, two more H₂ molecules are released to form Li₃B₃H₄, bypassing Li₂B₂H₄ entirely. At temperatures of 920 °C and higher, a Li₃B₃H₄ cluster is formed by releasing another H₂ molecule, before full dehydrogenation to the end product Li₃B₃. Small fraction of Li₃B₃H₄ and Li₃B₃H₂ clusters with odd-numbered hydrogen atoms is predicted to appear intermittently.

Finally, Fig. 4(c) shows the dehydrogenation pathway of (LiBH₄)₆. Even-numbered Li₆B₆H₁₂ and Li₆B₆H₁₀ clusters stage a brief appearance at 250 and 300 °C, respectively. The structures of these clusters can be described as derived from the parent borohydride cluster by releasing a hydrogen molecule and forming a B₃H₉ complex anion. It is tempting to hypothesize that \( 2BH_4^2- \rightarrow B_3H_9^2- \) also represents a kinetically viable hydrogen release.
pathway, but this has to be verified by separate studies of the activation energies. In any case, the negative charge carried by the \( \text{BH}_4^- \) anions should present a significant barrier to bringing these complexes together for the reaction. Most of the hydrogen is released around 420 °C with the appearance of \( \text{Li}_6\text{B}_6\text{H}_7 \) and \( \text{Li}_6\text{B}_6\text{H}_6 \) clusters. The latter cluster exhibits a high degree of symmetry and appears to be stable over a wide range of temperatures up to approximately 700 °C. In the final reaction steps, \( \text{Li}_6\text{B}_6\text{H}_3 \), \( \text{Li}_6\text{B}_6\text{H}_2 \) and \( \text{Li}_6\text{B}_6 \) show up one by one as the majority cluster phases. Again, it is seen that most of the stable clusters in the decomposition sequence contain an even number of hydrogen atoms, with the brief appearance of \( \text{Li}_6\text{B}_6\text{H}_7 \) as the only exception.

Based on the data in Fig. 4, we have calculated the cumulative mass fraction of released \( \text{H}_2 \) as a function of temperature at constant pressure (i.e., temperature-composition isobar). Figure 5 shows the results of these calculations. It is seen that the calculated hydrogen release curves exhibit nontrivial behavior as functions of temperature and cluster size.

For instance, the initial onset of hydrogen release occurs at a lower temperature for the \( n = 2 \) cluster than for the \( n = 3 \) cluster. This is due to the fact that the \( n = 2 \) cluster can release one \( \text{H}_2 \) molecule and form \( \text{Li}_2\text{B}_2\text{H}_6 \) (which consists of two \( \text{Li}^+ \) ions and a \( \text{BH}_2^- \) anion), while the corresponding \( \text{Li}_3\text{B}_3\text{H}_{10} \) compound is unfavorable because it involves a mixture of \( \text{BH}_2^- \) and \( \text{BH}_4^- \) anions. Furthermore, all three clusters exhibit two sloping plateaus in their dehydrogenation curves. Focusing on the \( n = 6 \) cluster, most of the hydrogen comes off in a narrow temperature interval before the first plateau, corresponding to successive removal of hydrogen molecules and eventual formation of \( \text{Li}_6\text{B}_6\text{H}_6 \) with an overall formula Eq. 4. In the second stage, a smaller amount of hydrogen is released by decomposing \( \text{Li}_6\text{B}_6\text{H}_6 \) and forming \( \text{Li}_6\text{B}_6\text{H}_3 \), etc. With increasing cluster size of nano-\( \text{LiBH}_4 \), we expected that the amount of hydrogen released in the first stage will increase and that the main hydrogen release step will shift to lower temperatures. According to the reaction enthalpies in Fig. 3, the overall reaction for \( n \leq 12 \) is expected to proceed according to Eq. 4, even though it seems likely that the full pathway will consist of multiple intermediate steps, similar to those discussed above. For \( n > 12 \), we hypothesize that the formation of clusters with \( \text{Li}-\text{compensated} \text{BH}_m^- \) complex anions (6 \( \leq m \leq 12 \)) will be favorable, but the existence of intermediate steps is also likely, in a manner similar to the \( n = 6 \) case. Unfortunately, reliable structure prediction for such large systems is beyond the capabilities of current computational methods.

Due to the increased enthalpy difference between the reactants and products, to achieve dehydrogenation at the same pressure as bulk \( \text{LiBH}_4 \), \( \text{LiBH}_4 \) nanoclusters need temperatures that are approximately 200 °C higher. Since experimental measurements on nanoconfined borohydrides18–20 often find lowering of the desorption temperature, this indicates that the supporting substrate plays an important role by binding the supported nanoclusters and lowering the enthalpy of the reaction products relative to \( (\text{LiBH}_4)_6 \).

While dehydrogenation is usually carried out by increasing temperature at a constant pressure, as shown in Figs 4 and 5, hydrogen absorption measurements are commonly done by increasing pressure at constant temperature. Hence, pressure-composition isotherms (PCT), showing the equilibrium pressure of the \( \text{H}_2 \) gas coexisting with a material with a given the hydrogen content, are of considerable interest. For bulk materials, these curves show flat regions corresponding to equilibria between the hydrogen gas and two solid phases on the opposite sides of the miscibility gap. In nanomaterials, clusters with different hydrogen content play the role of the coexisting solid phases. However, in contrast to bulk systems where the plateaus are perfectly flat in the thermodynamic limit with no kinetic limitations, in finite size clusters the plateaus acquire slope. This is illustrated in Fig. 6, which shows the calculated DFT pressure-composition isotherms for \( (\text{LiBH}_4)_6 \) clusters at three different temperatures. The left-hand side is the dehydrogenated phase, \( \text{Li}_6\text{B}_6\text{H}_6 \), corresponding 4.6 wt.% \( \text{H}_2 \) on the horizontal axis. The fully hydrogenated phase, \( (\text{LiBH}_4)_6 \), is on the right-hand side, at 18.5 wt.% \( \text{H}_2 \). Going right-to-left, the sloping plateaus describe the series of increasingly hydrogen-poor clusters in the decomposition sequence of \( (\text{LiBH}_4)_6 \).
with the slope arising from the intermediate steps and from the fact that the cluster probabilities $p_i$ in Eq. S2 vary continuously with pressure and temperature. We note that sloping plateaus are seen in experimental study of pressure-composition isotherms of bulk-LiBH$_4$, but this is likely due to kinetic limitations, not thermodynamic effects. Additionally, the apparent width of the plateau increases slightly with increasing temperature, which we attribute to the narrowing of the thermodynamic stability windows for the intermediate cluster compositions between (LiBH$_4$)$_5$ and Li$_2$B$_6$H$_6$.

Conclusions
We present a detailed computational study of the decomposition thermodynamics of small LiBH$_4$ nanoclusters using DFT-based ground state structure prediction algorithms and total DFT free energies, including vibrational contributions. With a few exceptions at very small sizes, nanoclusters favor the same decomposition sequence into closoborane Li$_2$B$_n$H$_n$ species as bulk LiBH$_4$. The calculated reaction enthalpies are higher than those for bulk materials because the total formation energies of the reaction product clusters approach their bulk values significantly slower than (LiBH$_4$)$_n$. Hence, reduction of cluster size is not expected to improve the thermodynamic properties of LiBH$_4$ for use in reversible hydrogen storage. Since experimental studies have demonstrated significant reduction of hydrogen release temperatures in supported nanoclusters, this confirms the importance of cluster-substrate binding energies for the thermodynamics of hydrogen release reactions. It also suggests that technologically favorable thermodynamics of LiBH$_4$ nanocluster decomposition could be engineered by varying the microscopic structure and composition of the porous substrate to lower the enthalpies of the reactants and achieve hydrogen release at temperatures compatible with the operating conditions of PEM fuel cells. Furthermore, we find that in all cases the decomposition pathways of (LiBH$_4$)$_2$, (LiBH$_4$)$_3$, and (LiBH$_4$)$_6$ consist of multiple intermediate stages, most of them separated by release of an integer number of H$_2$ molecules. While in bulk systems each reaction occurs along a sharp temperature-pressure curve given by the van't Hoff relation, in nanoclusters the transitions between the cluster phases of different compositions are smooth and characterized by broad coexistence regions on the temperature and pressure; within these regions, clusters of different compositions coexist and their equilibrium fractions vary continuously. As a result, the calculated pressure-composition isotherms exhibit sloping plateaus and smooth variation near the regions of stable bulk compositions.

Computational Methods. The structural and thermodynamic properties of small (LiBH$_4$)$_n$ ($n \leq 12$) clusters are studied using first-principles DFT calculations in conjunction with state-of-the-art genetic algorithm (GA) and prototype electrostatic ground state search (PEGS) for structure predictions. The detailed descriptions of our computational methods can be found in the Supplementary Information. Decomposition pathways and pressure-composition isotherms are determined using free energy minimization in the grand canonical ensemble (see Supplementary Information).

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Figure 6. Calculated pressure-composition isotherms for (LiBH$_4$)$_n$ clusters.
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Author Contributions

F.C.C. and V.O. conceived and initiated the study. Z.Q.H. performed the GA calculations, W.C.C. performed the DFT calculations, and E.H.M. performed the PEGS calculations. W.C.C., F.C.C. and V.O. performed the analysis and contributed the discussions. F.C.C., W.C.C. and V.O. wrote the manuscript. All authors reviewed the manuscript.
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