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# Enhanced Hydrogen Storage on Li Functionalized BC<sub>3</sub> Nanotube

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**ABSTRACT:** First principles calculations based on density functional theory are carried out to study the effect of lithium functionalization of experimentally synthesized BC<sub>3</sub> nanotube on charge transfer, electrostatic potential, and hydrogen storage. Electron-deficient BC<sub>3</sub> nanotube is found to promote charge transfer from Li to the substrate when lithiated. The resulting Li ions on the tube surface can effectively polarize hydrogen molecules and improve their binding energy and storage capacity. While each Li site on BC<sub>3</sub> nanotube is found to adsorb up to two H<sub>2</sub> molecules, zigzag nanotube shows better performance as a hydrogen storage material with good adsorption energy (0)



a hydrogen storage material with good adsorption energy  $(0.11 \text{ eV/H}_2)$  and high gravimetric density (6.9%). These data are consistent with the system target set by the DOE for 2010.

Currently, the biggest challenge in a new hydrogen economy is finding materials that can store hydrogen with high gravimetric and volumetric density under favorable thermodynamic conditions and possess fast kinetics. A considerable amount of experimental and theoretical research effort has been devoted to studying storage of hydrogen in various materials. Due to the high surface to volume ratio and the well-known tunable size-dependent properties of fullerene and carbon nanotubes (NTs), carbon based nanomaterials have received a great deal of attention.<sup>1-5</sup> It has been demonstrated<sup>3-5</sup> that pure carbon materials cannot be used for hydrogen storage because they interact weakly with hydrogen.

In order to improve the binding energy of hydrogen while maintaining the light weight of the storage material, doping of carbon with light metal atoms is desirable. Among these, Li doping seems to be promising since not only is it the lightest metal in the periodic table but also a positively charged Li<sup>+</sup> ion in free space has been shown to store six hydrogen molecules with desirable adsorption energy.<sup>6</sup> Li functionalized C<sub>60</sub> where 12 Li atoms are capped on the 12 pentagons was found to store hydrogen with large gravimetric density.<sup>7</sup> Here, six electrons from the Li atoms are transferred to the threefold degenerate LUMO of  $C_{60}$ , and each Li atom carries a charge of 0.5 electrons. This partial charge transfer resulted in a hydrogen adsorption energy of 0.075 eV/H<sub>2</sub>.<sup>7</sup> This binding is weak for  $Li_{12}C_{60}$  to serve as a hydrogen storage material for room temperature and ambient pressure<sup>8</sup> applications. To further improve this adsorption energy, it was shown that substitutional doping of B at the C site in C<sub>60</sub> may be effective.<sup>9</sup> When some of the C atoms are replaced by B atoms, the system becomes electron-deficient. Consequently, more electrons can be transferred from the Li atoms to the C<sub>60</sub> substrate leaving Li ions in a more positively charged state. Thus, the enhanced local electric field produced

by these ions can significantly improve the adsorption. Indeed, the hydrogen adsorption energy in  $\text{Li}_{12}\text{C}_{48}\text{B}_{12}$  can reach 0.135  $\text{eV/H}_{2,^9}$  which lies in the desirable energy window.<sup>10</sup> However,  $\text{C}_{48}\text{B}_{12}$  has not yet been synthesized, and up to six C atoms in  $\text{C}_{60}$ have been replaced by B. From a practical point of view, it will be ideal to have a substrate which already has been synthesized and can be doped by Li. In this regard BC<sub>3</sub> nanotube (NT) is a good candidate which has been synthesized experimentally<sup>11</sup> through chemical reactions. Further, it has been found that Li absorption can be enhanced with B doping on carbon NT.<sup>12</sup>

In this paper we report first principles calculations, based on density functional theory, of hydrogen adsorption in Li decorated BC<sub>3</sub> NT. We focus on the following questions: (1) Where do Li atoms bind and distribute on BC<sub>3</sub> NT and how does this depend upon different chiralities of the substrate? (2) How strong is the binding of Li and how does it change with tube radius? (3) How different are the hydrogen gravimetric densities for armchair (A-) and zigzag (Z-) BC<sub>3</sub> NT? We show that the distribution and binding of Li as well as the hydrogen gravimetric density depend on the chirality of BC<sub>3</sub> NT and that Z-BC<sub>3</sub> NT is better than A-BC<sub>3</sub> NT for hydrogen storage.

Calculations were carried out using density functional theory (DFT) and the generalized gradient approximation  $(GGA)^{13}$  for the exchange and correlation functional. We used the Perdew–Wang (PW91)<sup>14</sup> form for the GGA. All computations were carried out using the Vienna Ab Initio Simulation Package (VASP),<sup>15</sup> the ultrasoft pseudopotentials, and a plane-wave basis set with the projector augmented plane-wave method (PAW).<sup>16</sup> Periodic boundary condition and vacuum space of ~15 Å along

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*x*- and *y*-directions were applied in order to avoid interactions between two images of BC<sub>3</sub> NTs. The structures were relaxed using the gradient conjugated method with no symmetric constraints. To represent reciprocal space, Monkhorst–Pack k points<sup>17</sup> of  $1 \times 1 \times 7$  mesh are used. Denser k point meshes will affect the total energy within 0.002 eV. The energy cutoff, convergence criteria of energy, and force are set to be 400 eV,  $1 \times 10^{-4}$  eV, and 0.02 eV/Å, respectively. The accuracy of our calculation was tested by computing the H<sub>2</sub> bond length and dissociation energy, which are, respectively, 0.749 Å and 4.50 eV and agree well with the experimental value, namely, 0.741 Å and 4.53 eV. We also calculated geometric and electronic structures of the BC<sub>3</sub> sheet. Calculated B–C and C–C bond lengths of 1.564 and 1.421 Å, respectively, agree with previous results.<sup>18</sup>

First, we study the site preference of Li on BC<sub>3</sub> NT with varying chiralities. Calculations are carried out for the armchair type (n, n) (n = 4, 6, 8) and the zigzag type (m, 0) (m = 6, 8, 10) NTs. The indices *m* and *n* are even since BC<sub>3</sub> sheet has a 2 × 2 reconstruction. Structural optimizations show that nanotubes



**Figure 1.** Structures of (a) (4, 4)  $BC_3 NT$  and (b) (6, 0)  $BC_3 NT$ , (c) 1 Li on (4, 4)  $BC_3 NT$  and (d) 1 Li on (6, 0)  $BC_3 NT$ , (e) 2 Li atoms on (4, 4)  $BC_3 NT$  and (f) 2 Li atoms on (6, 0)  $BC_3 NT$ , (g) 4 Li atoms on (4, 4) and (h) 12 Li atoms on (6, 0)  $BC_3 NT$ .  $R_1$  and  $R_2$  are distances between Li atoms. Simulated unit cells are represented by dashed rectangles in (a), (b), (g), and (h). For (c) and (e) the simulated supercell is a triple unit cell; for (d) and (f) the supercell is a double unit cell.

can still retain their circular cross section after B doping, and the relaxed B–C and C–C bond lengths are  $\sim$ 1.56 and  $\sim$ 1.42 Å, respectively, similar to those of BC3 sheet. It has been reported that Li atom prefers to reside on the hollow site over C hexagonal ring<sup>19-21</sup> in pure C nanotube, and its binding can be further enhanced when B atoms are doped in the hexagonal ring.<sup>12,22</sup> Note that there are three different kinds of hollow sites on BC<sub>3</sub> NT where Li atoms can reside, namely,  $H_A$ ,  $H_B$ , and  $H_C$ , as shown in Figure 1. There are two B atoms in each of the hexagonal H<sub>A</sub> and H<sub>B</sub> sites, while the H<sub>C</sub> hexagonal site is composed of C atoms only. Therefore, we just consider the adsorption sites marked  $H_{\text{A}}$  and  $H_{\text{B}}$  sites in our calculations. When one Li atom is introduced to the H<sub>A</sub> or H<sub>B</sub> site and the geometries are allowed to relax, we find that the H<sub>A</sub> site is more favorable than the  $H_B$  site by 0.12 eV for the (4, 4) NT and 0.4 eV for the (6, 0) NT. The corresponding structures for Li doped BC<sub>3</sub> NT are shown in Figure 1c and d. Different energies are due to different chiralites and curvatures of the NTs. The absorption energy of Li atom on the  $H_A$  site is found to be 2.53 eV for (4, 4) and 2.42 eV for (6, 0) BC<sub>3</sub> NT.

Now we introduce a second Li atom to neighboring sites of H<sub>A</sub> in order to investigate how successive Li atoms will be distributed on the BC<sub>3</sub> NT. We found that the second Li atom still prefers a H<sub>A</sub>-type site for both A-BC<sub>3</sub> NT and Z-BC<sub>3</sub> NT over the H<sub>B</sub>-type site by 0.20 and 0.45 eV, respectively (Figure 1e and f). Based on these findings, we introduce Li atoms to all H<sub>A</sub>-type sites of A-BC3 NT and Z-BC3 NT (Figure 1g and h).<sup>23</sup> The average Li-B and Li-C bond lengths on (4, 4)  $BC_3$  NT are calculated to be 2.31 and 2.14 Å, respectively, while for (6, 0) BC<sub>3</sub> NT, the corresponding values are 2.27 and 2.12 Å. In Table 1 we list the calculated distances between Li-Li and Li-B, average adsorption energy per Li atom, charge on the Li atom, and average binding energy per H<sub>2</sub>. From these data the following conclusions can be drawn: (1) Concentration of Li on  $BC_3$  NT depends on the tube's chirality. The atomic ratios of Li to B and C are 1/8 and 1/4 for A- and Z-BC<sub>3</sub> NT, respectively. This is larger than previous studies of Li dispersed on pure CNT.<sup>21</sup> (2) Li atoms on Z-BC<sub>3</sub> NT are distributed more densely than those on A-BC<sub>3</sub> NT. For example, in the x- and y-direction Li–Li distances  $R_1$  are 3.95 and 3.59 Å for (6, 0) and (10, 0) BC<sub>3</sub> NTs, respectively. The distance in the *z*-direction,  $R_2$ , on the other hand, is 4.42 Å for all three Z-BC<sub>3</sub> NTs. In A-BC<sub>3</sub> NT,  $R_1$  ranges from 6.11 to 6.61 Å, while  $R_2$  is 5.12 Å. Both these distances are larger than those in Z-BC<sub>3</sub> NT. (3) The distance between Li and B as well as the charge transfer from Li decrease slightly with increasing diameter. (4) The average adsorption energy of Li increases with tube diameters of Z-BC<sub>3</sub>NT. This is due to the reduction in the Li–B distance.

In the following we discuss the charge transfer from the Li atom to  $BC_3 NT$  as the positive Li ion is expected to polarize and bind the  $H_2$  molecule. Charge analysis by integrating electrons

Table 1. Atomic Ratio of Li to B and C, Average Distance Li–Li  $(R_1)$  and Li–B (R) (in Å), Average Adsorption Energy per Li on BC<sub>3</sub> NT  $(E_b/Li, \text{ in eV})$ , Charge on Li  $(Q_{Li}, \text{ in electrons})$ , and Average Binding Energy per H<sub>2</sub> on Li  $(E_b/H_2, \text{ in eV})$ 

	4Li-(4, 4)BC <sub>3</sub> NT	6Li-(6, 6)BC <sub>3</sub> NT	8Li-(8, 8)BC <sub>3</sub> NT	12Li-(6, 0)BC <sub>3</sub> NT	16Li-(8, 0)BC <sub>3</sub> NT	20Li-(10, 0)BC <sub>3</sub> NT
ratio	1/8	1/8	1/8	1/4	1/4	1/4
$R_1$	6.61	6.33	6.11	3.95	3.69	3.59
R	2.34	2.30	2.28	2.30	2.27	2.25
$E_{\rm b}/{\rm Li}$	2.42	2.42	2.42	2.24	2.28	2.30
$Q_{\rm Li}$	+0.63	+0.62	+0.61	+0.60	+0.59	+0.58
$E_{\rm b}/{\rm H_2}$	0.10	0.09	0.09	0.11	0.11	0.11



**Figure 2.** (a) Isosurface with value of 0.012 electron/Å<sup>3</sup> and (b) 2D contour of transferred electron density ( $\rho_{Li-BC3NT} - \rho_{Li} - \rho_{BC3NT}$ ), (c) isosurface with value of 0.2 electron/Å<sup>3</sup> and (d) 2D contour of electrostatic potential of 6Li-(6, 6) BC<sub>3</sub> NT. Blue and yellow colors correspond to positive and negative values.

inside the Wigner-Seitz cell of each Li atom showed that charges are transferred from Li to the tube substrate, and transferred charge is mainly located on the hexagon just below Li. Charge on A-BC<sub>3</sub> NT is slightly larger than those of Z-BC<sub>3</sub> NT. To visualize it more clearly, we calculated the transferred electron density  $(\rho_{\text{Li-BC3NT}} - \rho_{\text{Li}} - \rho_{\text{BC3NT}})$  in Li doped on (6, 6) BC<sub>3</sub> NT. The resulting isosurface is plotted in Figure 2a and b in the form of a 2D contour. The blue and yellow curves indicate positive and negative charge density corresponding to electron accumulation and depletion, respectively. To further understand the effect of B doping of the carbon NT, some comparisons should be made between Li decorated on BC3 NT and pure CNT. Recall that the average adsorption energy per Li on pure CNT was calculated to be 1.8 eV using LDA<sup>20</sup> and 1.5 eV using GGA.<sup>21</sup> Our results (2.2-2.4 eV) indicate that B doping in CNT enhances the adsorption of Li and hence makes Li more positively charged (+0.6 e) than on pure CNT (+0.35 e) using  $LDA^{20}$  and +0.45 eusing GGA<sup>21</sup>). Enhanced charge transfer due to B doping can lead to a stronger local electric field produced by the Li anion and hence can lead to stronger binding of the H<sub>2</sub> molecule. Since the electric field is the gradient of electrostatic potential and vertical to the isosurface, it is helpful to analyze the electrostatic potential. This is plotted in Figure 2 c and d for Li-(6, 6) BC<sub>3</sub> NT. We see that electrostatic potential is positive with large gradient near the outer tube while it is negative with small gradient in the inner tube. Therefore, the electric field in the outer tube would be larger than that of the inner tube, and H<sub>2</sub> molecules would preferably bind to the outer surface.

We also calculated the partial density of states (PDOS) in fully decorated Li on (4, 4) and (6, 0) BC<sub>3</sub> NTs (Figure 3). The systems are metallic-like with p electrons across the Fermi level. This is consistent with previous results.<sup>24</sup> It is known that pure



**Figure 3.** Calculated PDOS of (a) 4Li-(4, 4)  $BC_3 NT$  and (b) 12Li-(6, 0)  $BC_3 NT$ .

(4, 4) and (6, 0) carbon NTs are metallic-like. They become semiconducting when doped with B but become metallic again when Li atoms are introduced.

Finally we discuss hydrogen adsorption on Li functionalized  $BC_3$  NT. Although previous studies<sup>12,19–22</sup> show that each Li can



Figure 4. Relaxed structure of (a) 4Li-(4, 4) BC<sub>3</sub> NT-8H<sub>2</sub>, (b) 6Li-(6, 6) BC<sub>3</sub> NT-12H<sub>2</sub>, (c) 8Li-(8, 8) BC<sub>3</sub> NT-16H<sub>2</sub>, (d) 12Li-(6, 0) BC<sub>3</sub> NT-24H<sub>2</sub>, (e) 16Li-(8, 0) BC<sub>3</sub> NT-32H<sub>2</sub>, and (f) 20Li-(10, 0) BC<sub>3</sub> NT-40H<sub>2</sub>.

adsorb up to three or four H<sub>2</sub> molecules, we find that in the present system, due to steric hindrance, it is impossible to store more than two H<sub>2</sub> molecules on each Li site . The average binding energies per  $H_2$  are shown in Table 1. These lie in the range of  $0.09-0.11 \text{ eV/H}_2$ . The binding energy of H<sub>2</sub> on Z-BC<sub>3</sub> NT is slightly larger than that in A-BC3 NT although charges of Li site on Z-BC<sub>3</sub> NT are slightly smaller. This can be understood from the different distribution of Li on BC3 NT. Compared to A-BC<sub>3</sub> NT, the Li-Li distance in Z-BC<sub>3</sub> NT is shorter, and accordingly the total electric field in the outer tube space would be stronger due to superposition. The geometric structures of H<sub>2</sub> adsorbed on Li functionalized A-BC<sub>3</sub> NT and Z-BC<sub>3</sub> NT are plotted in Figure 4. The relaxed average  $H_2$  bond length is  $\sim$ 0.76 Å, which is only marginally larger than that in its free state, namely, 0.749 Å. This elongation results from electrostatic polarization of H2. The distance between Li and  $H_2$  is around 2.0–2.2 Å, which is consistent with the previously reported Li– $H_2$  distance.<sup>12,19–22</sup> Since van der Waals interactions are not included, the GGA exchange correlation usually underestimates the binding energy  $^{25}$  by  $\sim 0.08$  eV/H<sub>2</sub>. Thus, our predicted binding energy  $(0.09-0.11 \text{ eV/H}_2)$  can actually be around 0.17-0.19 eV/H2. Bhatia and Myers studied the optimum thermodynamic conditions for hydrogen adsorption employing the Langmuir equation and then derived relationships between the operating pressure of a storage tank and the enthalpy of adsorption required for storage near room temperature.<sup>26</sup> They have found that the average optimal adsorption enthalpy should be 15.1 kJ/mol, if operated between 1.5 and 30 bar at 298 K. When the pressure is increased to 100 bar, the optimal value becomes 13.6 kJ/mol. Therefore, for the systems studied here, the adsorption energy of H<sub>2</sub> falls in the required window of hydrogen storage under ambient environment. Furthermore, the gravimetric densities of hydrogen are 3.8 and 6.9 wt %, for A-BC<sub>3</sub> NT and Z-BC<sub>3</sub> NT, respectively, and result from different concentrations of Li on BC<sub>3</sub> NT as pointed above. The latter satisfies the target set by the DOE for 2010.

In summary, we have studied the hydrogen storage properties of Li functionalized BC3 NTs using density functional theory. From a detailed analysis of charge density distribution, electrostatic potential, and hydrogen adsorption, we have arrived at the following conclusions: (1) Li atoms bind to the hexagonal rings containing B atoms. (2) The binding energy of Li to B doped carbon nanotubes is larger than that in pure carbon nanotube. (3)The charge transfer from Li to the nanotube surface is enhanced once the carbon nanotube is doped with B. (4) The electric field strength along the outer tube surface of BC<sub>3</sub> NT is larger than that along the inner surface and hence becomes the preferred space for hydrogen molecules to bind. (5) Li functionalized zigzag BC<sub>3</sub> NT shows better hydrogen adsorption energy and higher gravimetric density than Li functionalized armchair BC<sub>3</sub> NT. (6) Lithiation not only improves the performance of hydrogen storage of BC3 NT but also changes its electronic structure resulting in metallic behavior. Since BC3 NTs have been experimentally synthesized, our theoretical prediction is expected to stimulate experimental studies of hydrogen adsorption by functionalizing the nanotube with lithium.

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to too high concentration of Li decoration.

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