Conference Paper

Study of Structural and Electronic Behavior of BeH$_2$ as Hydrogen Storage Compound: An Ab Initio Approach

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The quantum mechanical calculations based on density functional theory (DFT) have been performed to study ground state structural and electronic properties of BeH$_2$ and along with doping of two (BeH$_2$ + 2H) and four (BeH$_2$ + 4H) hydrogen atoms. The generalized gradient approximation (GGA) has been employed for the exchange correlation energy. The most stable space group of BeH$_2$ is Ibam. Its optimized equilibrium unit cell volume, bulk modulus and its first-order pressure derivative, and electronic properties have been obtained. Our predicted unit cell parameters for BeH$_2$ $a = 9.2463$ Å, $b = 4.2352$ Å, and $c = 7.8464$ Å are in very good agreement with the earlier reported experimental and theoretical results. The electronic band structure of BeH$_2$ shows its behavior as an insulator. The stability of BeH$_2$ along with doped hydrogen atoms increases, while the energy band gap decreases with the increase in number of doped hydrogen atoms. On these bases, we predict that BeH$_2$ is a promising material for hydrogen storage.

1. Introduction

Energy storage for the future is a great concern for the researchers and scientists. We need materials like metal hydrides for various potential applications, that is, for hydrogen storage, in fuel cells and internal combustion engines, as electrodes for rechargeable batteries, and in energy conversion devices. The metal hydrides for hydrogen storage need to be able to form hydrides with a high hydrogen-to-metal mass ratio, but they should not be too stable, so that the hydrogen can easily be released without excessive heating. Beryllium and magnesium and beryllium-magnesium-based hydrides contain a relatively high fraction of hydrogen by weight but need to be heated $\sim$250 to 300°C in order to release the hydrogen. The alkali-metal and alkaline-earth-metal hydrides represent series with largely ionic bonding. The high pressure behavior of the alkali-metal monohydrides is expected to parallel the alkali-metal halides [1]. However, there is no systematic high-pressure study on the alkaline-earth-metal hydrides. If BeH$_2$ becomes metallic when subjected to high pressures, one can entertain the possibility that its properties could resemble those of metallic hydrogen. BeH$_2$ is commonly considered as a covalent hydride with a postulated polymeric crystal structure made up of H-bridged chains. However, mainly owing to experimental difficulties in the synthesis of the material, the structure has long remained unknown [2]. Crystalline BeH$_2$ has been synthesized and the structure has been established as body centered orthorhombic (Ibam) structure of BeH$_2$ has the lowest total energy.

In this paper, we have carried out the first-principles calculations of the ground-state behavior of BeH$_2$ in Ibam
Table 1: Calculated structural parameters, bulk modulus $B$ and its pressure derivative $B'_0$, and energy for orthorhombic BeH$_2$ are listed with available references.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Lattice parameter (in Å)</th>
<th>$B$ (in Gpa)</th>
<th>$B'_0$</th>
<th>$E$ (in Ry)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$a$</td>
<td>$b$</td>
<td>$c$</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>9.082</td>
<td>4.160</td>
<td>7.077</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>8.982</td>
<td>4.156</td>
<td>7.645</td>
<td>23.79</td>
<td></td>
</tr>
<tr>
<td>BeH$_2$ + 4H</td>
<td>10.078</td>
<td>4.616</td>
<td>8.553</td>
<td>40.04</td>
<td>2.9</td>
</tr>
</tbody>
</table>

Figure 1: Crystal structures of (a) BeH$_2$, (b) BeH$_2$ + 2H, and (c) BeH$_2$ + 4H in $Ibam$ phase.

The obtained unit cell parameters for this structure of BeH$_2$ are listed in Table 1. In order to obtain the equilibrium structural parameters, we carried out optimization of the total energy as a function of unit cell volume. Obtained sets of the unit cell volume and corresponding total energy have been fitted to the Murnaghan’s equation of state [11]. The bulk BeH$_2$ has also been studied with added number of hydrogen for the purpose of comparison between the ground state electronic properties of BeH$_2$ and with the doped hydrogen BeH$_2$. The studied variants are BeH$_2$ + 2H and BeH$_2$ + 4H. The obtained structural parameters for all the variants have been listed in Table 1.

2. Computational Details

The structural and electronic properties of the BeH$_2$ are obtained using density functional theory (DFT) [7, 8]. The full potential-linearized augmented plane wave (FP-LAPW) method has been employed for the calculation as implemented in the Wien2k code [9]. Generalized gradient approximation (GGA) is used to calculate the exchange-correlation functional. We expand the basis function up to $R_{MT} * k_{max} = 7$, where $R_{MT}$ is the smallest radius of the muffin-tin (MT) spheres and $k_{max}$ is the maximum value of the reciprocal lattice vectors. The maximum $l$ value for the wave function expansion inside the atomic spheres was confined to $l_{max} = 10$. The $k$-points used in the calculations were based on $5 \times 5 \times 5$ Monkhorst-Pack scheme [10]. The iteration process was repeated until the calculated total energy and charge of the crystal converge to less than 0.0001 Ry and 0.001 e$, respectively. In order to avoid ambiguities regarding the calculated results, we have used the same value of the energy cutoff as well as $k$-grid density for convergence for all the structural variants studied. The present theoretical approach has been successfully applied to study ambient- and high-pressure phases computationally.

The bulk BeH$_2$ exhibits $Ibam$ structure in ground state. The obtained unit cell parameters for this structure of BeH$_2$ are listed in Table 1. In order to obtain the equilibrium structural parameters, we carried out optimization of the total energy as a function of unit cell volume. Obtained sets of the unit cell volume and corresponding total energy have been fitted to the Murnaghan’s equation of state [11]. The bulk BeH$_2$ has also been studied with added number of hydrogen for the purpose of comparison between the ground state electronic properties of BeH$_2$ and with the doped hydrogen BeH$_2$. The studied variants are BeH$_2$ + 2H and BeH$_2$ + 4H. The obtained structural parameters for all the variants have been listed in Table 1.

3. Results and Discussion

3.1. Structural Properties

The ground state of BeH$_2$ has been synthesized experimentally [3] and studied theoretically [5]. Its orthorhombic unit cell (space group $Ibam$) is composed of twelve BeH$_2$ formula units as shown in Figure 1(a).

The primitive cell contains six formula units. The Wyckoff positions of different atoms are two Be1 (4a), four Be2 (8j), eight H1 (16k), and four H2 (8i). Each Be1 is surrounded by four H1 atoms to build the tetrahedral structure and each Be2 connects with two H1 atoms and two H2 atoms. Calculated atomic structural parameters for orthorhombic BeH$_2$ are included in Table 1. The obtained results of BeH$_2$ are in excellent agreement with the previous theoretical results [5, 6] and available experimental data [3, 4].

BeH$_2$ also studied with two and four extra added hydrogen atoms. Optimized values for unit cell volume and corresponding energy are fitted to Murnaghan’s equation of state [11]. The optimization curve is shown in Figure 2. From
the optimized unit cell volume, we have calculated the lattice parameters of BeH$_2$ + 2H and BeH$_2$ + 4H. The obtained unit cell parameters and other constants are listed in Table 1.

It is evident from Table 1 and plots in Figure 2 that the optimized energy of BeH$_2$ is less than that of BeH$_2$ + 2H and the energy of BeH$_2$ + 2H is less than that of BeH$_2$ + 4H. This indicates clearly that the stability of structures increases with the increase in the number of hydrogen atoms. A plot of total energy as a function of the doped hydrogen atoms is shown in Figure 3(a) and plot of unit cell volume versus number of doped hydrogen atoms is shown in Figure 3(b). The plot in Figure 3(a) shows linear behavior, while plot in Figure 3(b) is nonlinear.

The band structure of BeH$_2$ shows insulator behavior as in this case direct energy band gap is $\sim 5.8$ eV, while band structure plots for BeH$_2$ + 2H and BeH$_2$ + 4H show semiconducting behavior because in these cases indirect energy band gaps are, respectively, $\sim 2.2$ eV and $\sim 2.5$ eV (Figure 4). It is very interesting to note that below Fermi energy level width of a group of valence-band is $\sim 6.5$ eV which is in excellent agreement with earlier reported data [5]. The width of the group of valence-bands increases with
Figure 3: A plot between optimized energy and added extra number of hydrogen.

Figure 4: Electronic band structures for (a) BeH$_2$, (b) BeH$_2$ + 2H, and (c) BeH$_2$ + 4H.

the increase in number of doped hydrogen atoms. For BeH$_2$ + 2H, this width is $\sim$6.5 eV and for BeH$_2$ + 4H it is $\sim$7.5 eV. The behavior of conduction bands is somewhat different. Corresponding to two doped hydrogen atoms conduction bands near Fermi level are separated, while for 6 doped hydrogen atoms they are further compressed. This shows that number of added hydrogen atoms has significant effect on energy levels.

4. Conclusions

In this work, the structural stability of the Beryllium Hydride, BeH$_2$, and doped hydrogen compounds BeH$_2$ + 2H and BeH$_2$ + 4H have been studied by the first-principles calculations. The equilibrium structures of these compounds are obtained from DFT/GGA total energy minimization. The obtained results for BeH$_2$ are in excellent agreement with
the reported experimental and theoretical results. BeH₂ is an insulator and BeH₂ + 2H and BeH₂ + 4H are semiconductor. The effect of added hydrogen atoms is very clear. The stability of BeH₂ + 4H > BeH₂ + 2H > BeH₂. Lower value of total energy in BeH₂ + 4H shows that hydrogen storage in BeH₂ is one of very good option (Figure 2).

**Conflict of Interests**

The authors declare that there is no conflict of interests regarding the publication of this paper.

**References**


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