

Electronic properties of Zintl phase hydride for hydrogen storage

K. Khodja and Y. Bouhadda

Unité de Recherche Appliquée en Energies Renouvelables, URAER
Centre de Développement des Energies Renouvelables, CDER
47378, Ghardaïa, Algeria

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Abstract - *In this paper we report the SrAl₂H₂ electronic properties which is a Zintl phase hydride in frame of the Density Functional Theory 'DFT' using the plane wave and pseudo potential method. We discuss the chemical bond nature using total and partial density of states 'DOS and PDOS', also we calculated the bonding distance of hydride compound and its precursor SrAl₂ and the enthalpy formation of the SrAl₂H₂ for hydrogen storage.*

Résumé - *Dans ce travail, nous avons étudié les propriétés électroniques du SrAl₂H₂ qui est un composé hydride de phase Zintl dans le cadre de la Théorie de la Fonctionnelle de la Densité (DFT) en utilisant la méthode des ondes planes et du pseudo potentiel. Grâce à la densité d'état totale et partielle 'DOS et PDOS', nous avons pu discuter la nature des liaisons chimiques. Nous avons aussi calculé les distances des liaisons du composé hydride et de son prédécesseur le SrAl₂ et son enthalpie de formation pour une application dans le domaine du stockage de l'hydrogène.*

Keywords: Hydrogen Storage - Zintl Phase - Metal Hydride - Electronic Properties - Density Functional Theory.

1. INTRODUCTION

Aluminum hydrides have been a strong candidate for hydrogen storage. These compounds have satisfactory hydrogen content for storage but complications with kinetics and reversibility have made them slow to reach practical storage applications.

When hydrogen is absorbed in a metal hydride, the bond between the atoms in the H₂ molecule is broken as electrons from the metal lattice is filled into the anti bonding orbitals of the H₂ molecule and the H atoms are intercalated into the interstitial sites between the metal atoms.

The bond between the hydrogen and the metal atoms should be too strong to make it possible for hydrogen to be stored and released at temperatures and pressures close to ambient.

SrAl₂H₂ which is a Zintl phase compound was synthesized by hydrogenating of its precursor SrAl₂ at about 463 K under pressure of 50 bars [1]. Increasing the hydrogenation temperature to 513 K, SrAl₂H₂ further absorbs hydrogen to form Sr₂AlH₇ and Al.

In order to optimize the hydrogen storage in SrAl₂H₂ compound many experimental and computational works have been realized. Gingl *et al.* [1] have studied the hydrogenation of SrAl₂ by X-ray powder diffraction and found that the reaction proceeds in three steps. SrAl₂H₂ is formed in the first step starting at about 190 °C and

as the temperature is raised; the compound is further hydrogenated to a second hydride phase in different structure with higher hydrogen content.

Orgaz *et al.* [2] investigated the electronic structure of SrAl_2H_2 , Ca_3SnH_2 , and $\text{Ca}_5\text{Sn}_3\text{H}$ by means of the full-potential linearized augmented-plane-wave method. They found the SrAl_2H_2 and $\text{Ca}_5\text{Sn}_3\text{H}$ hydrides are metallic and Ca_3SnH_2 is a small-gap semiconductor. Lee *et al.* [3, 4] have investigated the vibrational properties of SrAl_2H_2 and SrAlSiH by means of inelastic neutron scattering ‘INS’ and first-principles calculations.

The results showed that SrAlSiH is characterized by very weakly dispersed Al-H modes with well-resolved overtones, while SrAl_2H_2 yields a solid-state dispersed phonon spectrum. The bonding characteristics study of SrAlSiH , SrAl_2H_2 , SrGa_2H_2 and BaGa_2H_2 using DFT calculations were done by Subedi *et al.* [5]. Their results indicate that in SrAl_2H_2 the Al layers are nominally neutral, i.e not polyanionic.

The Zintl phase alloy SrAl_2 can absorb hydrogen to form SrAl_2H_2 or Sr_2AlH_7 , the hydrogenation kinetics is quite slow especially for the latter one. Zhu *et al.* [6, 7] studied experimentally the structural and hydrogenation properties of the Zintl phase alloy SrAl_2 .

Zhang *et al.* [8] synthesized the Sr_2AlH_7 hydride by ball milling of a Sr_2Al alloy under hydrogen but it was followed by further hydrogenation of different types of samples.

The aim of our work is to investigate the electronic properties of SrAl_2H_2 and its precursor SrAl_2 by calculating the heat formation and the bonding distances and also discussing the chemical bond nature using total and partial density of states ‘DOS’ of each compound.

2. METHODOLOGY

In this study, all the computations have been done using the Abinit code [9] based on pseudo potentials and plane waves in density functional theory ‘DFT’ [10]. We have used the local density approximation ‘LDA’ of Troullier *et al.* [11] to approximate the exchange-correlation energy, where the Sr (3d, 4p, 5s), Al (3s, 3p) and H (1s) orbitals are treated as valence states.

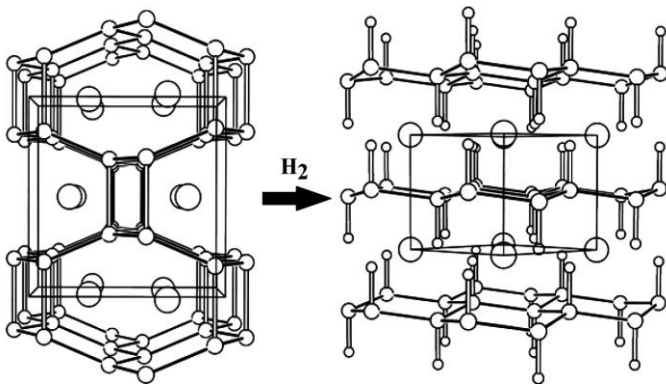


Fig. 1: Crystal structures of SrAl_2 (left) and SrAl_2H_2 (right), big, medium and small circles denote Sr, Al, and H atoms, respectively [1]

The electronic wave functions were expanded in plane waves up to a kinetic energy cutoff of 40 Hartree. Integrals over the Brillouin zone were approximated by sums of $6 \times 6 \times 6$ mesh of special k-point.

3. RESULTS AND DISCUSSIONS

3.1 Crystal structure

The crystal structure of SrAl_2H_2 is known and occurs in a trigonal space group $\bar{P}3m1$ (164). Al atoms are arranged as a slightly puckered graphitic layer. Additionally each Al atom is coordinated to one hydrogen atom. In SrAl_2 the puckered Al hexagonal layers are connected by a long Al-Al interlayer bonds which are cut in SrAl_2H_2 and terminated by hydrogen atoms (Fig. 1).

The relaxed structural parameters agreed very well with the experimentally and computationally determined ones (**Table 1**).

Table 1: Calculated relaxed structural parameters for SrAl_2H_2 compared with computational and experimental parameters

SrAl_2H_2	Our work		Compt		Exptl [2]
a (Å°)	4.6868	4.5267 [4]	4.5280 [6]	4.5289 [12]	4.5283
c (Å°)	4.8867	4.7195 [4]	4.7220 [6]	4.7248 [12]	4.7215
Al (1/3, 2/3, z)	0.4642	0.4613 [4]	0.4608 [6]	0.4614 [12]	0.4589
H (1/3, 2/3, z)	0.0952	0.0978 [4]	0.0964 [6]	0.0988 [12]	0.0976

In **Table 2**, we reported the bonding distances of each element that composes the SrAl_2H_2 compound and its precursor. As already mentioned in Gingl *et al.* [1] work, our Al-Al bond length of SrAl_2H_2 (2.72 Å°) is considerably shorter than the ones in SrAl_2 (2.84 Å°) and the distances between Sr and Al are slightly longer (SrAl_2 : 3.42 Å°, SrAl_2H_2 : 3.53 Å°).

Table 2: Bonding distances

Bond (Å°)	Our work		Exptl [2]	
	SrAl_2H_2	SrAl_2	SrAl_2H_2	SrAl_2
Sr-Al	3.53096	3.42357	3.654	3.59
Sr-H	2.74562	-	2.653	-
Al-H	1.80319	-	1.706	-
Al-Al	2.72845	2.84744	2.641	2.79

3.2 The formation energy (enthalpy formation)

We have take into account two reactions related to the formation of the Zintl phase hydride SrAl_2H_2 :



To calculate the formation heat of the reaction (1) we subtracted the total energies of the pure elements Sr, Al and the hydrogen molecule from their hydride SrAl_2H_2 , and we have done the same with the second one.

$$\Delta H (\text{SrAl}_2\text{H}_2) = \Delta E (\text{SrAl}_2\text{H}_2) - \Delta E (\text{Sr}) - 2\Delta E (\text{Al}) - \Delta E (\text{H}_2) \quad (3)$$

Table 3 contains the total energy and the formation enthalpy computed of SrAl_2H_2 for two different reactions. The total energy of the hydrogen molecule is -30.8229 eV and has been also calculated. We noted that the heat formation of the second reaction is much less than the first one, making the reaction (2) more favorable for the formation of SrAl_2H_2 compound.

Table 3: Calculated heat of formation of SrAl_2H_2

Elément	Total energy (eV)	Enthalpie formation (kJ/mol)
Sr	-576.4821	-
Al	-64.0722	-
H ₂	-30.8229	-
SRAl ₂	-712.5062	-
SrAl ₂ H ₂	-743.7879	804.4945 44.2674

3.3 Electronic structure

In this section the lattice parameters were calculated and reported in **Table 1**. The total and the partial densities of state for SrAl_2H_2 are plotted in Fig. 2 and Fig. 3 which are similar to that obtained previously by Orgaz *et al.* [2], Subedi *et al.* [5]. The electronic structure is metallic without the energy gap. This is in agreement with the computational and experimental work of Lee *et al.* [3]. We can also say that SrAl_2H_2 is weakly metallic because the DOS reach a value of 0.0354 (states/eV/cell) at the Fermi level [2].

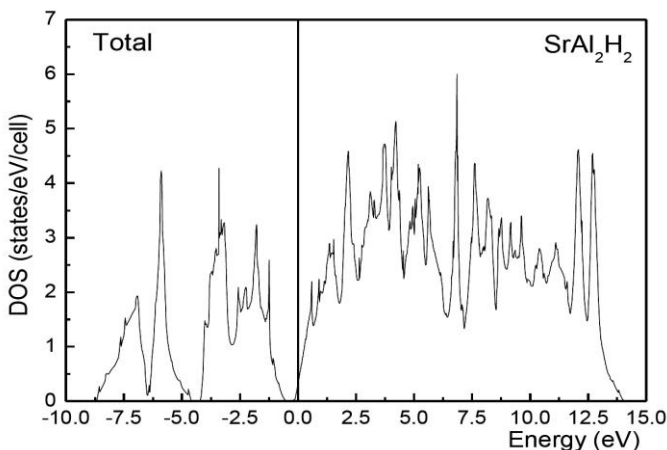


Fig. 2: The total density of state of SrAl_2H_2

In Fig. 2 at the bottom of energy scale, we can see two peaks; the first concerns the H-s/Al-s bonding interaction. This is followed by a second peak mainly produced by the H-s states, where a non-negligible and small Al-s,p orbital contribution is present.

This can be seen in the PDOS plots of Fig. 3. The second part of the DOS plot is well separated from the first. Two main contributions appear in increasing order of energy. First, there is a complex H-s/Al-s orbital interaction including small Sr-d and Al-p contributions. This is completed by the Al-p states up to the Fermi energy. The PDOS of Sr has a very small contribution to the valence band from the region -10 (eV) to 0.

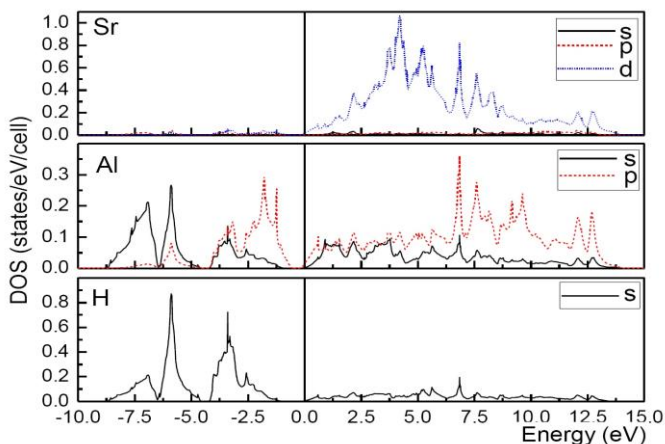


Fig. 3: Partial density of state for SrAl_2H_2

4. CONCLUSION

In this work we report the electronic structure calculations for the ternary hydride SrAl_2H_2 using the ABINIT code. The electronic structure is obtained from LDA total-energy minimizations. Formation energy for SrAl_2H_2 is calculated for two different possible reaction pathways.

This allowed us to know that is better to synthesize our compound from its precursor SrAl_2 than from each element that composes the Zintl phase hydride. Also from the plot of the DOS, we noted that SrAl_2H_2 has no gap which means that our compound is a metallic and the valence band is dominated by hydrogen atoms.

NOMENCLATURE

Sr: Strontium	Al: Aluminum
H: Hydrogen	a , b : Cell parameters
ΔH : Enthalpy formation	Å° : Interatomic distance unit
DOS : Total density of charge	PDOS : Partial density of charge
eV : Energy unit	state/eV / cell : Density of state unit

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