Modeling and stabilities of Mg/MgH₂ interfaces: A first-principles investigation

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We have theoretically investigated the modeling and the structural stabilities of various Mg/MgH₂ interfaces, i.e. Mg(10 ¯10)/MgH₂(210), Mg(0001)/MgH₂(101) and Mg(10¯10)/MgH₂(101), and provided illuminating insights into Mg/MgH₂ interface. Specifically, the main factors, which impact the interfacial energies, are fully considered, including surface energies of two phases, mutual lattice constants of interface model, and relative position of two phases. The surface energies of Mg and MgH₂, on the one hand, are found to be greatly impacting the interfacial energies, reflected by the lowest interfacial energy of Mg(0001)/MgH₂(101) which is comprised of two lowest energy surfaces. On the other hand, it is demonstrated that the mutual lattice constants and the relative position of two phases lead to variations of interfacial energies, thus influencing the interface stabilities dramatically. Moreover, the Mg-H bonding at interface is found to be the determinant of Mg/MgH₂ interface stability. Lastly, interfacial and strain effects on defect formations are also studied, both of which are highly facilitating the defect formations. Our results provide a detailed insight into Mg/MgH₂ interface structures and the corresponding stabilities. © 2014 Author(s). All article content, except where otherwise noted, is licensed under a Creative Commons Attribution 3.0 Unported License.

I. INTRODUCTION

Magnesium (Mg) is a promising candidate¹,² for application of mobile hydrogen storage material due to the high gravimetric hydrogen content³ (7.6% in MgH₂), high volumetric density³ (6.7 × 10¹⁹ H atoms/L), and high energy density⁴ (9MJ/kg) of its hydride MgH₂. To make Mg applicable, there are still two technical obstacles¹,² to be solved: (1) Slow hydrogenation/dehydrogenation kinetics; (2) A too high working temperature of 350 - 400°C. In order to improve the material hydrogen sorption performance, extensive theoretical and experimental researches including strain engineering,⁵,⁶ material surface/bulk ratio increasing,⁷–⁹ transition metal catalyzing¹⁰–¹² and mixing¹³,¹⁴ with other compounds have been done to mitigate the thermodynamic and kinetic drawbacks of Mg.

Hydrogen sorption in Mg concerns to the Mg/MgH₂ or other related interfaces. Recently, some experimental and theoretical efforts have been made to study the Mg/MgH₂ interface structure and improve MgH₂ dehydrogenation properties. Epitaxial growth of MgH₂(110) on Mg(0001) is experimentally observed.¹⁵ The significant increase of equilibrium hydrogen pressure in Pd/Mg systems was first believed to be caused by the clamping effect¹⁶ but later found to be induced by the interfacial alloy hydride.¹⁷ Mooij et al.⁹ have found that the nanoconfined MgH₂ is more easily destabilized compared to its bulk counterpart because of an interface energy effect. Moreover, decreased dehydrogenation enthalpies could be also observed in Mg/Mg₂Ni interface system¹⁸ or on

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MgH₂ clusters⁷ with smaller size. Noticeably, the MgH₂ dehydrogenation mechanism was found¹⁹ not to be simply a reverse of the adsorption one. Although the barrier for Mg nucleation is small¹⁹ in dehydrogenation process, the interfacial void formations¹⁹, ²⁰ accompany the Mg nucleation thus leading to a difficult determination of Mg/MgH₂ interface structures. Theoretically, it is found²¹ that the dehydrogenation facilitation of (100) MgH₂/TiH₂ multilayers attributes to structural deformations and interfacial energy variations. Moreover, by means of Car-Parrinello molecular dynamics method (CPMD), it is found that the atomic hydrogen diffusivity at MgH₂(110)/Mg(010) interface steeply increase²² at 700 K, and is facilitated²³ when Fe is doped. Though these experimental and theoretical studies provide interesting and illuminating knowledge of Mg/MgH₂ interface properties, there is still a lack of systematic study of relative stabilities of various Mg/MgH₂ interfaces, i.e. various combinations of different Mg and MgH₂ surfaces. Additionally, the theoretical studies have not fully taken into account the influences of relative position,²² interlayer spacing²², ²⁴, ²⁵ and lattice mismatch²⁴, ²⁶–²⁹ at the first-principles level, which may be determining the Mg/MgH₂ interface stabilities.

In this paper, we firstly investigate the matching between Mg and MgH₂ surfaces by considering their surface energies, lattice constants and cell shapes. Then we consider the interfacial elastic effect by determining the mutual lattice constant which leads to minimized energy cost on elastic effect. Furthermore, we indicate that the most stable interfacial atomic structures would be found according to their lowest interfacial energies and the interfacial geometric characters would be discussed in detail. Lastly, the interfacial properties and strain effects on defect formations are also provided. The methods are specified in Section II, the results and discussions are in Section III, while conclusions are in Section IV.

II. METHODOLOGY

The incoherent interface between two phases, namely magnesium and magnesium hydride, is chosen to be modeled in this paper. The two phases are modeled to be thick enough so that they could be seen as the atom reservoirs, so as to stimulate the hydrogen sorption process. During the establishment of interface model, we do not consider the diffusion of H atoms at the interface of Mg/MgH₂, though it does not reflect the practical scenario as the hydrogenation/dehydrogenation often occurs above 600 K for Mg-H system. The trend of the H diffusion at the interface will be discussed from the formation energies of interstitial and vacancy H defects around the interface.

We start with the investigation of surface energies. Afterwards, some combinations of two phases could be found. Of these the chosen are determined according to the proposed rules, where the relative positions of two phases and the biaxial strains along each axis are considered. Finally, the comparison between the stabilities of interface models is made to discuss what the factors that dominate the interface stabilities are.

A. Computational details

Our calculations are performed using Vienna ab initio Simulation Package.³⁰–³² We use the exchange-correlation with generalized-gradient approximation (GGA)³³ and projector-augmented wave (PAW) potentials.³⁴ In all the calculations, adequate tests with k-points meshes³⁵ have been done to show excellent converged results and the plane-wave cutoff energy was set to be 310 eV. Monkhorst-Pack³⁵ k-point samplings of 6 × 2 × 1, 5 × 2 × 1 and 5 × 2 × 1 are used in the Mg(1010)/MgH₂(210), Mg(0001)/MgH₂(101) and Mg(1010)/MgH₂(101) interfaces, respectively, for the Brillouin-zone integration. The internal optimization is converged with the force on each atom to be less than 0.01 eV/Å. The energy convergence with respect to the k-points and cutoff energy has also been tested to be in the precision of ~0.01 eV per cell.

B. The calculations of surface energy, interfacial energy and H defect formation energy

1. The surface energy

Surface energies are seen as the energy cost of creating a surface by dividing a crystal into two parts. Here, the surface energies of magnesium (Mg) and magnesium hydride (MgH₂) are calculated
by the following formula

\[ E_{\text{surf}} = \lim_{n \to \infty} \left( \frac{1}{2A} \left( E_{\text{tot}}(n) - E_{\text{bulk}}(n) \right) \right), \]  

(1)

where \( E_{\text{tot}}(n) \) and \( E_{\text{bulk}} \) represent the total energies of the slab model and the bulk with \( n \) formula units, \( A \) is the area of the surface slab and the factor 2 denotes two surfaces being created. Of note, the Fiorentini approach\textsuperscript{36} is adopted to get a reasonable convergence of surface energies.\textsuperscript{37}

2. The interfacial energy

The interfacial energy \( E_i \) at the grain boundary between two phases is defined as the formation energy of the interface system in a particular context, where the chemical potentials at which two phases are at equilibrium could be determined. The calculated formula of interfacial energy proposed by Arras \textit{et al.}\textsuperscript{38} is given by

\[ E_i = E_{\text{tot}} - N_{\text{Mg}} \mu^0_{\text{Mg}} - N_{\text{H}} \mu^0_{\text{H}}, \]  

(2)

where \( E_{\text{tot}} \) is the total energy of interface model, \( N_{\text{Mg}} \) and \( N_{\text{H}} \) are the total number of atoms of magnesium and hydrogen respectively in the interface system, \( \mu^0_{\text{Mg}} \) and \( \mu^0_{\text{H}} \) are the chemical potentials of magnesium and hydrogen when two phases, namely Mg and MgH\textsubscript{2}, are at equilibrium. In our model in which the two phases represent the atom reservoirs,\textsuperscript{39} \( \mu^0_{\text{Mg}} \) and \( \mu^0_{\text{H}} \) are calculated by

\[ \mu^0_{\text{Mg}} = \frac{(x_{\text{Mg}} \varepsilon_{\text{MgH}_2} - x_{\text{MgH}_2} \varepsilon_{\text{Mg}})}{(x_{\text{Mg}} - x_{\text{MgH}_2})}, \]  

(3)

and

\[ \mu^0_{\text{H}} = \frac{(x_{\text{Mg}} \varepsilon_{\text{MgH}_2} - x_{\text{MgH}_2} \varepsilon_{\text{Mg}})}{(x_{\text{Mg}} - x_{\text{MgH}_2})}, \]  

(4)

where \( \varepsilon \) is the total energy per formula unit and \( x \) is the atomic concentration of an element (subscript) in a phase (superscript). Specifically, \( x_{\text{Mg}}^{\text{Mg}}, x_{\text{Mg}}^{\text{MgH}_2}, x_{\text{H}}^{\text{Mg}}, x_{\text{H}}^{\text{MgH}_2} \) are equal to 1, 0, 1/3 and 2/3, respectively.

3. The H defect formation energy

The H defect formation energy \( E_f \) is defined as the formation energy of the H vacancy in the MgH\textsubscript{2} phase, or the H interstitial in Mg phase of the interface model. The calculated formula of the defect formation energy is given by

\[ E_f = E_{\text{tot}} - E_{\text{perfect}} \pm N_{\text{defect}} \cdot \mu_{\text{H}}, \]  

(5)

where \( E_{\text{tot}} \) and \( E_{\text{perfect}} \) are the total energies of interface models with and without H defect, \( N_{\text{defect}} \) indicates the number of H defects, and the signs + and - represent H vacancy and interstitial, respectively. Here, in order to stimulate the H-rich context, \( \mu_{\text{H}} \) is equal to 1/2\( E_{\text{H}_2} \).

C. The criterion of choosing surface combinations

1. Surface energies

The surface energies strongly reflect the possibilities of formation of a particular surface. It should be noted that our first-principles calculation of surface energies is conducted at 0K, where the bond cutting and the surface reconstruction are the two major factors that dictate the surface energies. Previous papers\textsuperscript{37,40,41} showed that both of these two effects played an important role on surface energies. Moreover, the contribution of vibration entropy, another factor in determining the surface energies, goes up when the temperature rises so that the two major factors may not be the only determining ones and the surfaces with high surface energies at 0 K may appear.\textsuperscript{42} As a matter of fact, the hydrogen desorption in magnesium hydride requires a temperature much higher than the room temperature. Therefore, the surfaces with high surface energies at 0 K also have to be considered in dictating the combination of two surfaces of two phases.
TABLE I. The information, namely the lattice constants, the included angle and the surface energy, of Mg and MgH2 surface cells. The Mg surface energies are sourced in Ref. 33. The stoichiometric MgH2 surface energies are calculated in this paper.

| Surface index | a (Å) | b (Å) | Included angles γ (°) | Calculated surface energies (meV/Å²) | (J/m²) |
|---------------|-------|-------|------------------------|--------------------------------------|--------|
| Mg            |       |       |                        |                                      |        |
| 0001          | 3.19  | 3.19  | 120.00                 | 39a                                  | 0.624  |
| 1010          | 3.19  | 5.21  | 90.00                  | 60.6a                                | 0.970  |
| 1011          | 3.19  | 6.10  | 105.15                 | 45.9a                                | 0.734  |
| 1012          | 3.19  | 7.59  | 90.00                  | 54.6a                                | 0.874  |
| 1013          | 3.19  | 9.92  | 99.26                  | 49.1a                                | 0.786  |
| 1014          | 3.19  | 12.22 | 90.00                  | 48.7a                                | 0.779  |
| 1015          | 3.19  | 14.85 | 96.17                  | 49.5a                                | 0.792  |
| 1016          | 3.19  | 17.37 | 90.00                  | 45.7a                                | 0.731  |
| 1017          | 3.19  | 20.09 | 94.55                  | 49.5a                                | 0.792  |
| 1018          | 3.19  | 22.71 | 90.00                  | 48.3a                                | 0.773  |
| 1019          | 3.19  | 25.45 | 93.59                  | 46.5a                                | 0.744  |
| 1020          | 5.21  | 5.53  | 90.00                  | 51.4a                                | 0.822  |
| 2130          | 5.21  | 8.44  | 90.00                  | 56.1a                                | 0.898  |
| MgH2          |       |       |                        |                                      |        |
| 001           | 4.45  | 4.45  | 90.00                  | 31.3b                                | 0.501  |
| 100           | 4.45  | 2.99  | 90.00                  | 23.6b                                | 0.378  |
| 101           | 5.36  | 4.45  | 90.00                  | 24.2b                                | 0.387  |
| 110           | 2.99  | 6.30  | 90.00                  | 21.3b                                | 0.341  |
| 210           | 2.99  | 9.95  | 90.00                  | 56.1b                                | 0.898  |

* Reference .33
† This work.

2. The matching of shape and lattice constants of the cells

The periodic interface model which consists of two phases requires that the two phases share the same shape and close lattice constants. The perfectness of matching leads incoherent interfaces to coherent or semi-coherent interfaces.

The matching of shape takes priority over the matching of lattice constants. Specifically, the cells of rectangle shape (NB: the hexagonal shape can be transformed into rectangle shape) could always be matched easily after some unit repeating operations, which is in pursuit of existence of a coincidence site lattices43,44 (CSL). Comparatively, the cells of parallelogram with unequal neighboring sides and a non-right angle are difficult to be matched with the others. For instance, it is found that the Mg(1011) surface cell is in its difficulty to be matched due to its shape of parallelogram where \( a_{\text{Mg}(1011)} = 6.11 \, \text{Å}, \ b_{\text{Mg}(1011)} = 6.38 \, \text{Å} \) and the included angle \( \gamma = 74.87^\circ \). The cell of this shape, unlike the rhombus, is also impossible to be transformed into the periodic cell of rectangle shape due to its specified atomic arrangement.

The unit repeating operations follow two geometrical criteria45,46: (1) The lattice misfit strain between two phases should be as small as possible. (2) The size of repeated cells, manifested by the period of the CSL, is as small as possible. The first criterion is easily understandable. The second one is based on the observation3,33 that the grain boundaries between two identical materials are often found to have a CSL with small unit cell volume. The information about the size of our repeated cells is specified in Table II.

In summary, the rectangle cell (original or transformed) is always preferred in the process of choosing the surface of each phase. In practice, we choose the Mg(0001), Mg(1010), MgH2(101) and MgH2(210) surfaces whose information is specifically listed in Table I, which will be further discussed in following section. Fig. 1 provides an intuitive illustration of interface modeling and interface pairs in which we are interested.
FIG. 1. (a) Cartoon for our interface model whose \(xy\) plane shape of cell is rectangle. \(\Delta x\), \(\Delta y\) and \(\Delta z\) denote the variation of relative position of two phases, which is detailed in the next chapter. Front and side views of supercells of interfaces of (b) \(\text{Mg}(10\overline{1}0)/\text{MgH}_2(210)\), (c) \(\text{Mg}(0001)/\text{MgH}_2(101)\) and (d) \(\text{Mg}(10\overline{1}0)/\text{MgH}_2(101)\). Magnesium and hydrogen atoms are in green and white respectively. The dashed lines in (b), (c) and (d) denote the size of repeated cells as listed in Table II.

3. Determination of the mutual lattice constants

For an interface system with lattice mismatch, both of the two phases would be possibly under elastically tensile or compressive effect. Therefore, the elastic cost subject to the mutual lattice constants is required when the interface is created. Here, the mutual constants are defined as the lattice constants of the interface model where two phases coexist with minimized elastic energy cost. The minimum of this elastic energy cost is expected to be found in our interface models. Basing on the assumption that both of two phases would compromise energetically to reach the mutual constant, we propose that the energy cost on elastic effect is calculated by

\[
E_{\text{EEC}} = \Delta E_{\text{Phase.1}} + \Delta E_{\text{Phase.2}},
\]

where \(E_{\text{EEC}}\) represents the total elastic energy cost of two phases, \(\Delta E_{\text{Phase.1}}\) and \(\Delta E_{\text{Phase.2}}\) denote the elastic energy costs of two phases with respect to the most energetically favored structure.

The search of the mutual lattice constants is required to determine the mutual constants. In practice, the calculations of \(\Delta E_{\text{Phase.1}}\) and \(\Delta E_{\text{Phase.2}}\) are calculated respectively in bulk forms of two phases by fitting the \(E-c\) curve at given \(a\) and \(b\). Thus, the mutual constants could be determined by the minimized elastic energy cost. It is in line with widely adopted theoretical models, although it
is relatively simple to the real interface where the lattice constants of both phases will be restored beyond corresponding critical thickness.\textsuperscript{47}

4. Determination of relative position between two phases

The stability of interface, after the determination of the mutual lattice constants, is highly influenced by the bonding nature in interfacial region, namely the interfacial structure. Naturally, the factors that lead to a change of interfacial structure should be focused on. One may easily come up with several factors including the relative position of two phases, the concentration gradient between two phases and the effect of defect or dopants in interfacial region.

The authentic hydrogen sorption process always exhibits migration of grain boundaries, which is affected not only by the thermodynamic stability but also by the kinetic one. From the point of view of theoretical calculation, it is not easy to describe all these factors simultaneously. However, when it comes to the discussion of the interfacial bonding, we simply assume that the relative position of two phases should be the primary factor in affecting the interfacial structure in our calculations since the variation of the relative position would lead to a dramatic change of interfacial bonding, whose effect is similar as the sliding shear acted on the system.

III. RESULTS AND DISCUSSIONS

A. The bulk properties of magnesium and magnesium hydride

The crystal structure of magnesium is hexagonal close packing and the calculated Mg bulk lattice constants are $a_{Mg} = b_{Mg} = 3.190\,\text{Å}$ with the $c/a$ value of 1.623, which are in good agreement with the experimental values\textsuperscript{48,49} and previous theoretical results.\textsuperscript{50–52}

Magnesium hydride, whose crystal structure is simple tetragonal, is a highly stable hydride with strong Mg-H bond, which is a mixture of ionic and covalent bonding.\textsuperscript{53} The calculated bulk lattice constants are $a_{MgH_2} = b_{MgH_2} = 4.452\,\text{Å}$, $c_{MgH_2} = 2.992\,\text{Å}$ with the $c/a$ value of 0.672, which agree well with the previous experimental\textsuperscript{54–56} and theoretical\textsuperscript{57–59} results.

B. The surface energies of Mg and MgH$_2$

Here, surface energies of two phases are calculated and compared to find the relative stabilities of surfaces. It is known that the low-index surfaces are often more stable than the high-index ones since the number of breaking bonds\textsuperscript{60} that are required to create a low-index surface is relatively less. However, it has been reported\textsuperscript{37} that the low-index Mg(1010) has a high surface energy thanks to its massive basal bond cuttings. MgH$_2$(210) is of the highest surface energy in Table I, while the other MgH$_2$ surfaces share low surface energies.

In a theoretical calculation, the adopted model should be periodic. Consequently, the shapes and lattice constants of two phases are identical, which are widely accepted in previous literatures\textsuperscript{24,26,38,61–63} concerning interface. It means that the shapes and the lattice constants should be taken into account in combining the two surfaces of two phases.

Considering these factors mentioned in the above two paragraphs, i.e. the surface energies, the shape and lattice constants of a surface-slab cell, we choose the combinations, as listed in Table I, of which two surfaces of two phases are identical in shape and close in lattice constants (repeated along the axes if necessary). In practice, the low-energy Mg(0001) and MgH$_2$(101) are chosen while the high-energy Mg(1010) and MgH$_2$(210) are selected to exemplify our approaches.

C. The search of mutual constants

1. Elastic effect

Formation of interfaces always exhibits energy compromise of two phases, which can be reflected by elastic effect. We define here, for an ideal condition, the mutual constant as a constant which leads to minimized energy cost on elastic effect. Therefore, the search of mutual constants

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for interface models could be interpreted as searching the minimum $E_{ECC}$ among a wide range of lattice constants.

In practice, the search of mutual lattice constant for interface could be divided into two steps: (1) The elastic costs on each phase are estimated. (2) Take into account the whole contribution from two phases caused by elastic effect. It should be pointed out here that the interface-induced strain effect on both phases may lead to different strains along axes $a$ and $b$. Therefore the consideration of nonequivalent strain on axes $a$ and $b$ makes sense. Since the total energies of Mg and MgH$_2$ per formula unit are $-1.56$ eV and $-8.91$ eV respectively, the contributions from two phases, caused by elastic effect, also differ. It is clear that the thickness of the two phases significantly affects the minimization of the elastic energy in our model. Practically, the critical thickness, including the strain release effect of the layers within the critical thickness, should be included in the minimization of the elastic energy in our model. Practically, the thickness of two phases is significant in affecting the stability of the two phases significantly affects the minimization of the elastic energy in our model. Practically, the critical thickness, including the strain release effect of the layers within the critical thickness, should be included in the minimization of the elastic energy in our model.

Moreover, the number of Mg layers is nearly twice as that of MgH$_2$ layers so that the contributions from two phases could be competing. Here, we assume that our choice of thicknesses, made by the two above mentioned strategies, is capable of investigating the elastic effect, i.e. determining the mutual constants. Similar ratio between numbers of two phases, i.e. Ge and GeO$_2$, is used in previous literature.

Fig. 2 shows the case of Mg(1010)/MgH$_2$(210) interface. We apply various structures, which are distinguishable from their lattice constants, to two phases in their bulk form cell. The elastic effects are found to be obviously influencing the phase stabilities. The relative stabilities of varying lattice constants could be reflected by the energy difference of the respective structures with respect to the most energetically favored one (as shown in Figs. 2(a) and 2(b)). Considering the Possion effect, we fixed $a$ and $b$ but varied $c$, and then fitted the $E$-c curve to determine the energy for each structure. Afterwards, the whole elastic contribution from two phases are estimated by simply calculating $E_{ECC} = E_{Phase,1} + E_{Phase,2}$ [see Fig. 2(c)]. The most energetically favored structure could be found among many structures that we considered. In this Mg(1010)/MgH$_2$(210) interface case, the mutual constants are found to be $a = 3$ Å and $b = 10.2$ Å. Here, we have made a cell repeating operation for Mg(1010) in order to match the interface (see Table II).

Similarly, the mutual constants searching proceeds on the cases of Mg(0001)/MgH$_2$(101) and Mg(1010)/MgH$_2$(101) interfaces as shown in Fig. 3. The details of three interface combinations are specifically listed in Table II. With the obtained lattice constants, the interfaces could be modeled.

### 2. The search of relative position of two phases

The determination of mutual constants mainly focuses on elastic effect but does not involve the interfacial bonding, which primarily affects the interface stabilities. Recent papers have shown that interface properties highly depend on the lattice matching or interface chemistry. At Mg/MgH$_2$ interfaces, the Mg-H bonding is the most important bonding that leads to interface stability.

Authentically, there are interfaces consisted of two phases with distinctive symmetries in many cases, especially for the cases with high-index surfaces. For example, correlation between interface trap densities and interface anisotropy are found on the interfaces consisted of high-index surfaces. Another case is that the interface of Ag/high-index substrate enhances resonance by enhancing the scattering cross-sections. From a point of view of theoretical calculation, it is rather difficult to find the most energetically-favored structure for interfaces where two phases are with different symmetries. Therefore, the full consideration of relative position is essential.

We propose that the relative position of two phases could be seen as the vital reason causing change of interfacial bonding in theoretical calculations. For theoretical modeling, considerable interfacial structures could be obtained by adjusting the relative position of two phases. Therefore, it is possible to compare the stabilities of relaxed interfaces with various structures.

We have calculated the effect of relative position in Fig. 4. It could be obviously observed the varying of relative position could cause a dramatic change in interfacial energies (the largest difference is 73.2 meV/Å$^2$ (1.171 J/m$^2$), up to 22.73% of the maximum). In every structure (at particular $x$ and $y$), an energy dependence on $d$ could be seen. Here, $d$ is defined as the distance...
FIG. 2. Contour maps showing the energy preference for the lattice constants reflecting the elastic effect on two phases of Mg(10\bar{1}0)/MgH2(210) interface. The \( \Delta E_{\text{Mg}(10\bar{1}0)} \) and \( \Delta E_{\text{MgH}_2(210)} \) are calculated by the difference between the structure energies and the most stable one among them. The structures considered are within the range of constant \( a \) and \( b \): \( a \) ranges from 2.8–3.6 Å with an interval of 0.1 Å while \( b \) from 9.8–10.6 Å with an interval of 0.1 Å. The energy unit of interfacial energies is J/m².
FIG. 3. Elastic effects on two interfaces including Mg(0001)/MgH$_2$(101) and Mg(10$\bar{1}$0)/MgH$_2$(101). The calculation method is the same as the Fig. 2. The structures considered are within the range of constant $a$ and $b$: $a$ ranges from 5–5.8 Å with an interval of 0.1 Å while $b$ from 9.8–10.6 Å with an interval of 0.1 Å. Each point denotes the lowest energy among structures with $a$ fixed and $b$ varied from 12.1–13.7 Å with an interval of 0.2 Å, as shown in the inset. The mutual constants of Mg(0001)/MgH$_2$(101) and Mg(10$\bar{1}$0)/MgH$_2$(101) are found to be $a = 5.4$ Å, $b = 12.2$ Å and $a = 5.2$ Å, $b = 12.2$ Å.

TABLE II. Specific data and information of search of the mutual constants and interface modeling.

| Interface                  | Mg(10$\bar{1}$0)/MgH$_2$(210) | Mg(0001)/MgH$_2$(101) | Mg(10$\bar{1}$0)/MgH$_2$(101) |
|----------------------------|---------------------------------|-----------------------|---------------------------------|
| Shape change of unit cell (Mg/MgH$_2$) | No/No                           | Yes/No                | No/No                           |
| Size of repeated cell (Mg/MgH$_2$)     | $1 \times 2/1 \times 1$       | $1 \times 4/1 \times 3$ | $1 \times 4/1 \times 3$       |
| No. of layers (Fixed) (Mg/MgH$_2$)     | 10(4) / 5(2)                   | 5(2) / 3(1)           | 5(2) / 3(1)                     |
| No. of Mg atoms (Mg/MgH$_2$)           | 20 / 10                        | 40 / 18                | 40 / 18                         |
| Mutual Constant (MC) (Å)               | $a = 3$                        | $a = 5.4$             | $a = 5.2$                       |
|                                           | $b = 10.2$                     | $b = 12.9$            | $b = 12.9$                      |
| Searching range (Å)                   | $a$: 2.8–3.6                   | $a$: 5–5.8            | $a$: 5–5.8                      |
|                                           | $b$: 9.8–10.6                  | $b$: 12.1–13.7        | $b$: 12.1–13.7                  |
| Interval in MC searching (Å)           | $a$: 0.1                       | $a$: 0.1              | $a$: 0.1                        |
|                                           | $b$: 0.1                       | $b$: 0.2              | $b$: 0.2                        |

between two phases of initial configuration along z axis, as shown in the inset of Fig. 4(c). This is easy to understand since the increasing of $d$ lengthens the distance of two phases and thus greatly alters the Mg-H bond lengths at interface. It would lead to the dramatic change of Mg-H bond energies and thus influence the interfacial stabilities. We find there are local energy minimaums in every structure within the proposed $d$ range. It means that there is a preference for $d$, which dovetails with strengthening of interfacial bonds, in interface models. Logically, it could be interpreted that the optimization of $d$ is always required for interface models.

Moreover, it is found the varying of relative position on x-y plane (at fixed $d$) also induce a change of interfacial energies although not as dramatic as that induced by $d$ varying. Specifically, the varying of x-y relative position would cause a change of local structure. But this effect would be obviously offset by relaxations of interfacial atoms and thus just induce a slight effect on the change of interfacial energies. Comparably, the varying of $d$ would be more effective on tuning
FIG. 4. (a) The contour map showing the lowest interfacial energies of structures with varying relative positions of two phases of Mg(10\overline{1}0)/MgH\textsubscript{2}(210) interface (\(a = 3\) Å, \(b = 10.2\) Å). The lowest interfacial energies at grid line crossings, like the red up-triangle, are determined by the method described in part (c). Each structure at grid line crossings represents a specific \(\Delta x\) and \(\Delta y\): \(\Delta x\) ranges from 0 to 3 Å with an interval of 0.6 Å (20% of 3 Å) while \(\Delta y\) ranges from 0 to 10.2 Å with an interval of 1.02 Å (10% of 10.2 Å). (b) The visualization of tuning the relative position of two phases of Mg(10\overline{1}0)/MgH\textsubscript{2}(210) interface (\(a = 3\) Å, \(b = 10.2\) Å) in the \(x\)-\(y\) plane. The symbols □ and ◦ represent the initial \(x\)-\(y\) plane position of the first and second phases respectively. (c) The figure showing the determination of the lowest interfacial energies in the range of varying distance between two phases along \(z\) direction, namely \(d\), at a fixed relative position. The inset visualizes the varying distance between two phases along \(z\) direction.

interfacial bonding. Higher effectiveness is mainly attributed to the fact that all interfacial bonds have to respond as \(d\) varies. Compared with \(x\)-\(y\) relative position varying, it is more global, which relaxations of interfacial atoms are not easy to offset.

Similar findings could be observed at interfaces of Mg(0001)/MgH\textsubscript{2}(101) and Mg(10\overline{1}0)/MgH\textsubscript{2}(101) interfaces (see Fig. 5). In details, every structure experiences quadratic curves. Thus, the relative position of two phases, in theoretical interface models and calculations, should be always taken into account, especially for the influence of \(d\) varying on interfaces where no obvious symmetries are adopted.

Notably, it is also observed that the interfacial energies of three interface pairs in Fig. 4 and Fig. 5 are quantitatively distinct from each other. This draws our attention to the descriptions of
FIG. 5. The contour maps showing the lowest interfacial energies of structures with varying relative positions of two phases of (a) Mg(0001)/MgH$_2$(101) and (d) Mg(1010)/MgH$_2$(101) interfaces ($a = 5.5$ Å, $b = 12.9$ Å). Each structure at grid line crossings represents a specific $\Delta x$ and $\Delta y$: $\Delta x$ ranges from 0 to 5.5 Å with an interval of 1.1 Å (20% of 5.5 Å) while $\Delta y$ ranges from 0 to 12.9 Å with an interval of 1.075 Å (8.33% of 12.9 Å). The method where the data is obtained is the same as the part (c) of Fig. 4. The varying distance between two phases of (b) Mg(0001)/MgH$_2$(101) and (c) Mg(1010)/MgH$_2$(101) interfaces along $z$ direction could also be visualized.
TABLE III. The percentage of bond lengths in specified range. The standard Mg-Mg and Mg-H bond lengths are 3.19 Å and 1.96 Å extracted from calculated lattice constants of Mg and MgH₂ bulks.

| Interface Pair          | Structure                   | Percentage in specified range (%) |
|-------------------------|-----------------------------|----------------------------------|
|                         |                             | Mg-Mg Bond | Mg-H Bond |
|                         |                             | ±5%         | ±3%       | ±5%         | ±3%       |
| Mg(10̅10)/MgH₂(210)     | The most stable configuration | 82.28       | 46.84     | 88.24       | 64.71     |
|                         | The least stable configuration | 78.89       | 39.89     | 73.53       | 52.9      |
| Mg(0001)/MgH₂(101)     | The most stable configuration | 76.86       | 62.78     | 92.68       | 45.12     |
|                         | The least stable configuration | 87.28       | 81.58     | 82.9        | 44.74     |
| Mg(10̅10)/MgH₂(101)     | The most stable configuration | 22.94       | 13.76     | 90.36       | 62.79     |
|                         | The least stable configuration | 31.28       | 22.58     | 85          | 65        |

interfacial bonding. Table III lists the figures of interfacial bonding of various structures. Observations show obviously that the most stable interfacial structures exhibit a higher percentage of Mg-H bonds in specified range and a lower one of Mg-Mg bonds. Specifically, the interfacial atom relaxations allow Mg-H bonds to be in their most energetically favored length. Thus it indicates that the Mg-H bonding leads to higher interfacial stability. Comparatively, the Mg-Mg bonding is much less contributive to interfacial stability. Therefore we argue that the Mg-H bonding is the primary cause leading to higher interfacial stabilities. The most energetically favored interfacial bonding structure could be obtained by varying the relative position, i.e. the search of relative position of two phases.

Interestingly, we found that the interfacial stabilities are influenced by surface energies of two phases. The Mg(10̅10)/MgH₂(210), the interfacial pair of surfaces with high surface energy, are expected to have low interfacial energy due to great number of dangling bonds that each phase has. However, the results show that Mg(10̅10)/MgH₂(210) are owning a high interfacial energy around 4 meV/Å² (0.064 J/m²) while the Mg(0001)/MgH₂(101) pair consisted of two low energy surfaces has a much lower one of nearly 2.1 meV/Å² (0.034 J/m²). Meanwhile, the interfacial energy of Mg(10̅10)/MgH₂(101) pair, which is with one high energy and one low energy surfaces, is about 2.8 meV/Å² (0.045 J/m²). This difference of interfacial energies could be explained by the fact that the partial saturation of interfacial dangling bonds is not capable enough to offset the instabilities of high energy surfaces. Experimentally, it is observed⁴⁴ that the grain boundaries comprised of low energy surfaces have relatively low energies. Our finding is in line with the experimental ones, and indicates that consideration of surface energies should be taken into when choosing interface combinations.

D. Defect formation at interface

The properties of Mg/MgH₂ interface, especially the hydrogen sorption property, are intriguing. As hydrogen sorption dovetails with the migration of Mg/MgH₂ interface, both of the hydrogen adsorption or desorption are considered. We here report comparative formation energies of hydrogen vacancies and interstitials, representing interfacial effect on hydrogen desorption and adsorption respectively.

Fig. 6(a) shows the interfacial effects on hydrogen vacancies. Clearly, it could be observed that hydrogen vacancy formations are highly facilitated in near-interface region. Overall, the formation of hydrogen vacancies at interface is much easier than that in MgH₂ bulk. It could be well understood as the formation of interface dovetails with some dangling (under-coordinated atoms) or floating (over-coordinated atoms) bonds. The existence of dangling or floating bonds means the interfacial H atoms are less bound by Mg atoms and weaker stabilities than those in MgH₂ bulk and thus the interface dramatically lowers the formation energies of hydrogen vacancies. Tensile strain monotonically increases the formation energies of $V_{1st}^H$ and $V_{2nd}^H$ while the curves of far-interface $V_{3rd}^H$, $V_{4th}^H$, $V_{5th}^H$, $V_{6th}^H$ and $V_{7th}^H$ are slightly parabolic. When hydrogen vacancy is more distanced from interface, the Mg-H bonding approaches to that in MgH₂ bulk. Thus similar trends induced by interfacial effect on hydrogen vacancy formation could be clearly observed under various strain states. Bi axial strain
FIG. 6. Formation energies of (a) single hydrogen vacancies and (b) single hydrogen interstitials at (c) different sites of Mg(10\textbar 10)/MgH\textsubscript{2}(210) interface. Biaxial strains with magnitude $m$ ($m = -6\%$, $-3\%$, $0\%$, $+3\%$ and $6\%$, five points from left to right) are applied on the interface. The numbered white hydrogen atoms denote vacancies while those grey ones with underlined numbers represent interstitials. The superscripted ordinal numbers (in (a) and (b)) indicate the relative distances, from the nearest to the furthest, between the defect site and the interface. The same concentration data in Mg and MgH\textsubscript{2} bulks are also shown with corresponding strains applied on Mg(10\textbar 10) and MgH\textsubscript{2}(210). Figure (c) is shown in size of 1 \times 2 supercell. The dash and dot vertical lines in (a) and (b) correspond to the areas of unstrained Mg(10\textbar 10) and MgH\textsubscript{2}(210) surfaces, respectively.

effect affects the H vacancy formation energy as expected. As the interface lattice constants (tuned by strain) are close to those of MgH\textsubscript{2} bulk, the $V\text{H}_\text{bulk}$ formation energies go up due to increased stability of MgH\textsubscript{2} phase of interface thus parabolic curves appear. Interestingly, the MgH\textsubscript{2} bulk data is found to be monotonic, which is not parabolic. Actually, it is raised from difference between interface and bulk data. Specifically, the $C$ axis of bulk is allowed to relax when strains are applied. It means all the atoms are able to relax in response to the strain. Two MgH\textsubscript{2} atom layers in the interface model are fixed in order to stimulate the stable MgH\textsubscript{2} substrate therefore the atom relaxations in far-interface regions are not enough so that similar parabolic curves could be observed. In authentic interface environments, the latter situation is more likely to happen since inadequate responses happen to a substrate that are ‘thick’ enough, which we usually deem to be strains. Based on the above analyses, we conclude that hydrogen vacancies are more energetically preferred in interface regions, especially the near-interface one, compared with the bulk region.

The interface also facilitates hydrogen interstitials. It is extremely easy for H\textsubscript{inte} \textsuperscript{1st} to form, which is reflected by its negative value of formation energy. H interstitial formations are much more approachable to the bulk region situation as it is more distance from interface. The reason for this phenomenon is that Mg bulk context would facilitate hydrogen uptake thanks to the strong Mg-H bonding. The strong bonding strongly increases stability of system. When a tensile strain is applied, the space for H interstitials is enlarged and thus resulting in a monotonic trend of formation energies. It is observed that the lengths of Mg-H bond under strains of $-6\%$ and $+6\%$ are 1.92 Å and 1.96 Å for H\textsubscript{inte} \textsuperscript{1st}. The altering of Mg-H bonding rationalizes the monotonic trend of formation energies.

IV. CONCLUSIONS

Three combinations of Mg/MgH\textsubscript{2} interface models are built to investigate the interface stabilities. The criterions including surface energies, interface matching, elastic effects and relative positions are discussed for Mg/MgH\textsubscript{2} interface modeling. It is found that the elastic effects and relative positions
strongly affect the incoherent interface stabilities by affecting incoherent interfacial structure. Thus the searches of mutual constants and relative positions provide feasible solutions and make prediction of interfacial structure possible. Lastly, the defect formations are highly facilitated at Mg/MgH2 interfaces.

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