Ab Initio Hydrogen Dynamics and the Morphology of Voids in Amorphous Silicon

Parthapratim Biswas* and Dil Limbu

Dedicated to Professor David A. Drabold on the occasion of his 60th birthday

This article presents an ab initio study of hydrogen dynamics inside nanometer-size voids in amorphous silicon (α-Si) within the framework of the density-functional theory for a varying hydrogen load of 10–30 atoms per void at the low and high temperature of 400 and 700 K, respectively. Using the local density approximation (LDA) and its generalized-gradient counterpart (GGA), the dynamics of hydrogen atoms inside the voids are examined with an emphasis on the diffusion of H atoms/molecules, and the resulting nanostructural changes of the void surfaces. The results from simulations suggest that the microstructure of the hydrogen distribution on the void surfaces and the morphology of the voids are characterized by the presence of a significant number of monohydride Si–H bonds, along with a few dihydride Si–H₂ configurations. The study also reveals that a considerable number (about 10–45 at%) of total H atoms inside a void can appear as H₂ molecules for a hydrogen load of 10–30 H atoms per void. The approximate shape of the voids is addressed from a knowledge of the positions of the void-surface atoms using the convex-hull approximation and the Gaussian broadening of the pseudoatomic surfaces of Si and H atoms.

1. Introduction

Hydrogen plays an important role in the electronic and optical properties of amorphous silicon (α-Si). Although the presence of a small amount (about 6–12 at%) of hydrogen in α-Si is particularly beneficial for producing device-grade samples of α-Si:H, via passivation of coordinating defects (e.g., threefold-coordinated Si atoms or dangling bonds), the presence of too much hydrogen in α-Si can be detrimental to the electronic and optical properties of α-Si:H-based devices. Thin films of α-Si:H are often used for surface passivation of crystalline silicon, which is useful for the generation of high open-circuit voltages in silicon-heterojunction solar cells. A relatively high hydrogen content, in void-rich environment of α-Si:H, is preferred for this purpose to achieve high-quality α-Si:H/α-Si interfaces, indicating the benevolent role of H in α-Si. By contrast, the structure of α-Si:H is altered by the light-induced creation of metastable defect states, known as the Staebler-Wronski effect (SWE), which adversely affects the performance of α-Si:H-based solar cells. Nuclear magnetic resonance (NMR) studies on α-Si:H samples have indicated that the motion of H atoms, which are produced by the light-induced breaking of Si–H bonds, plays an important part in recovering the photovoltaic stability of the affected α-Si:H samples upon annealing at 350–400 °C. Thus, an understanding of the motion of H atoms in α-Si in the presence of inhomogeneities is of crucial importance to address the photovoltaic stability of α-Si:H upon light irradiation.

Although a number of earlier theoretical/computational studies on the microstructure of the hydrogen distribution in α-Si:H have focused on the silicon–hydrogen bonding configurations in the bulk environment of α-Si and their effects on structural, electronic, and optical properties of the material, there exist only a few computational studies that directly address the role of non-bonded hydrogen (NBH) and voids on the network structure of α-Si:H. Sekimoto et al. have recently shown experimentally that the presence of large amount of NBH (e.g., H₂ molecules inside voids) can broaden the vacancy-size distribution and enhance the size of the optical gap in α-Si:H. The mass density of α-Si:H has been also found to be somewhat dependent on the number of NBH, and therefore on the distribution of hydrogen inside nanometer-size voids. Following our recent study on the temperature-induced nanostructural evolution of voids in α-Si and its effect on the intensity of small-angle X-ray scattering (SAXS), the present study focuses on accurate calculations of the atomic dynamics of hydrogen inside voids obtained from the density-functional theory (DFT). The emphasis here is on the dynamical aspects of hydrogen motion inside voids on the timescale of several picoseconds at low and high temperatures in the environment of a varying hydrogen concentration. The formation and dissociation of Si–H bonds on the surface of voids are discussed from a kinetic point of view. The movement of H atoms inside nanometer-size voids and its resulting effects on the hydrogen microstructure and the reconstruction of the void surfaces are also addressed here. The presence of voids in the amorphous matrix suggests that, for an accurate...
determination of the motion of H atoms within a void, one must take into account the inhomogeneities in the electronic charge distribution in the vicinity of the void surface. The standard protocol in DFT calculations advises us to address the problem by computing the self-consistent-field (SCF) solution of the Kohn–Sham (KS) equation, and the presence of voids suggests that the generalized-gradient approximation (GGA) should be used to deal with the density fluctuations near the void surface. We shall therefore address the problem using the GGA and compare the results with those from the local density approximation (LDA). The presence of weak dispersion forces, which are usually included via the van der Waals corrections, is not taken into account in this article due to the computational complexity of the problem and the somewhat limited accuracy of local basis functions used in our work.

The rest of the article is organized as follows. Section 2 provides a short description of the method for generating a-Si models with hydrogenated voids, using the Wooten–Winer–Weaire (WWW) method and ab initio molecular-dynamics (AIMD) simulations. The results are discussed in Section 3, with an emphasis on the dynamics of H atoms inside voids and the resulting microstructure of the hydrogen distribution on the void surface for a varying number of hydrogen atoms. A discussion on the kinetics of Si–H bond formation and dissociation and the shape of the voids is also provided in this section. This is followed by the conclusions of the study in Section 4.

2. Computational Method

For AIMD simulations of hydrogen dynamics inside voids in a-Si, we started with four independent 1000-atom models of a-Si, obtained from the WWW method. A spherical void of radius 5 Å was created at the center of each model and the geometry of the resulting structure was thoroughly relaxed, using the first-principles density-functional code SIESTA. A number of models with a hydrogenated void were then produced by adding 10, 20, and 30 H atoms inside the central cavity so that the mass density of the final models was about 2.22 g cm⁻³. A single void of radius 5 Å corresponded to a number density of void of 4.82 × 10⁻⁹ cm⁻³ and a void-volume fraction of 2.52%, which were close to the values reported in experiments. The H atoms were initially distributed in such a way that they were at a distance of at least 2 Å from Si atoms and 1 Å from each other. The silicon atoms within the spherical region of radius between 5 and 8 Å from the center of the voids were labeled as the void-surface atoms for the analysis of the void surface upon annealing and total-energy optimization of the systems.

To study hydrogen diffusion and the microstructure of hydrogen distribution on the void surfaces, AIMD simulations were performed at 400 and 700 K to examine the temperature dependence of H dynamics and the resulting effects on the morphological structure of the void surfaces, using the density-functional code SIESTA. SIESTA employs local basis functions, based on numerical pseudoatomic orbitals and the norm-conserving Troullier–Martins pseudopotentials to obtain the SCF solutions of the KS equation in the density-functional theory. Electronic and exchange correlations between electrons were taken into account via the LDA and the GGA, by using the Perdew–Zunger and the Perdew–Burke–Ernzerhof (PBE) parameterizations for the LDA and the GGA, respectively. Due to the computational complexity of the problem involving a large number of atoms and the necessity for simulating a reasonably long-time dynamics for studying the diffusive behavior of H atoms, using the SCF solution of the KS equation, the AIMD simulations were conducted using the single-zeta (SZ) basis functions for Si atoms and the double-zeta-polarized (DZP) basis functions for H atoms. The high computational cost associated with the calculation of total energy and forces restricted us to use SZ basis functions for Si atoms during the course of MD simulations. The subsequent total-energy optimizations, however, were conducted using the double-zeta (DZ) basis functions for Si atoms until the total force on each atom was less than or equal to 5 × 10⁻³ eV Å⁻¹. During AIMD simulations, the temperature of the system was controlled by using the Nose thermostat, and a time step of 0.8 fs was used to ensure that the movement of light H atoms can be described accurately at a high temperature of 700 K. The evolution of the system was monitored and recorded for a total time of 20 ps. The simulation procedure was repeated for each of the four independent models, indicated by M1 to M4, with a central void for a hydrogen load of 10, 20, and 30 H atoms per void. In each case, we have used a different random distribution of H atoms within the void to gather as much as statistics as possible. The final results were obtained by averaging over the four independent configurations.

3. Results and Discussion

3.1. Hydrogen Dynamics Inside Voids in a-Si

In this section, we discuss results from the density-functional calculations for studying the dynamics of H atoms/molecules inside voids in a-Si. To this end, we mostly focus on those H atoms/molecules which are within the cavity during the entire course of simulations at a low and high temperature of 400 and 700 K, respectively. Figure 1 shows the time evolution of the average number of H atoms (molecules included) within a void-surface region of radius 8 Å at 400 and 700 K. The results correspond to two different values of the hydrogen load, 10 and 30 H atoms per void, and are obtained by averaging over four independent voids/models using the LDA and GGA. At 400 K, the great majority of H atoms stayed within the void, with the exception of one or two H atoms that left the void-surface region of radius 8 Å. The LDA and GGA dynamics exhibit a more or less similar behavior at a given temperature, although the GGA tends to knock out one or two more H atoms outside the cavity for high H loads at 700 K. Despite limited statistics, it would not be inappropriate to conclude that, on average, the LDA and GGA do not differentiate much as far as the (average) number of H atoms leaving the void at 400 K is concerned.

To study the effect of the hydrogen load on the mean-square displacement (MSD) of H atoms at low and high temperature, of 400 and 700 K, respectively, we have examined the variation of the MSD with time for 10, 20, and 30 H atoms per void using the GGA. Once again, the results were obtained by averaging over four independent voids and are shown in Figure 2. The results (in Figure 2) lead to the following observations. First,
the MSD of H atoms decreases with an increasing presence of H atoms within the voids. This is particularly so as the hydrogen load increases from 10 to 30 H atoms per void. This reduction in the MSD is due to the decrease in the average distance between any two H atoms within the cavity for an increasing presence of hydrogen within the same volume. Second, the sharp rise of the MSD for a hydrogen load of 10 H atoms per void near 12 ps at 400 K can be attributed to a hydrogen atom leaving the void. For a hydrogen load of 10 H atoms per void, H atoms can move a bit more freely than those with a load of 20 or 30 H atoms per void. This makes it possible for few H atoms to diffuse rapidly in the vicinity of the void surface, and eventually to escape the void region—and the simulation cell in one or two cases—depending upon the temperature of the system. In contrast, the presence of too many H atoms for a high value of H load can impede the diffusion process and thus reduces the MSD at a given temperature. Third, the motion of H atoms is affected by the temperature of the system as well. This is evident in Figure 2b, where the MSD for a load of 20 H atoms per void was observed to increase at 700 K during 15–16 ps and it continued to remain so until the end of the simulation at 20 ps. This observation contrasts with the case of 30 H atoms per void at the same temperature, where none of the H atoms was found to leave the void but remained within a root-mean-square (RMS) distance of 5 Å from the center of the void. Finally, one may note that the available statistics do not permit us to comment on the dynamical behavior of few H atoms that diffuse out of the cavity at an earlier time (e.g., the H atom in Figure 2a for a hydrogen load of ten atoms per void), based on the results from four configurations. The movement of H atoms is driven by a combination of factors, such as the concentration of H atoms in a void, temperature, and the disorder in the atomic distribution on the void surface. Since we are not particularly interested in the dynamical behavior of few H atoms that escape the void-surface region of 8 Å, we will not discuss this further.

Having discussed the variation of the MSD with respect to the hydrogen load, we now address to what extent the exchange-correlation (XC) approximation may affect the motion of H atoms inside the voids as far as the MSD is concerned. Figure 3 shows the evolution of the MSD with time for a hydrogen load of 30 H atoms per void at 400 and 700 K. Since we are interested in the dynamics of H atoms inside the void, and the resulting hydrogen microstructure of the void surface, the MSD was computed using only those H atoms that were inside the void region of radius 8 Å for the entire duration of simulation. The plots in Figure 3 suggest that the LDA slightly overestimates the MSD of H atoms compared to the value obtained from its generalized-gradient counterpart. The RMS values of the displacement of H atoms inside the void in the LDA and the GGA at 400 K have been found to be about 4.5 and 4.15 Å, respectively. A more or less similar observation applies to the high-temperature dynamics at 700 K, as shown in Figure 3b. It thus appears that the MSD of the H atoms within the void is not particularly strongly affected by the XC approximation at 400 and 700 K.
This is due to the fact that the majority of H atoms within a void reside as bonded hydrogen (BH) (to Si atoms) on the void surface, mostly in the form of Si—H configurations, and few Si—H₂ configurations, along with a few H₂ molecules within the cavity. Since the calculation of the MSD excludes a few mobile H atoms that have already left the void region, the remaining bonded H atoms show a more or less converged value of the MSD during the course of simulation. Thus, in a sense, a more or less converged value of the MSD (in Figure 3) is reflective of the distribution of bonded H atoms on the walls of the void. We shall see later that a slightly larger value of the MSD of H atoms that we observe here for the LDA case (see Figure 3) results from a minor expansion of the void surface in the LDA calculations. Following Sekimoto et al., one may conclude that the expansion originates from the stress on the void surface due to the presence of a significant number of Si—H bonds in the LDA calculations. Finally, it may be mentioned that the dihydride Si—H₂ configurations are found on the void surface at a distance, which is closer to the center of the void than their monohydride (Si—H) counterpart. Figure 4 shows the average distances of the Si—H₂ and Si—H bonding configurations, which are located on the walls of the voids, from the center of the voids for the GGA at 400 K. This observation is consistent with the results reported by Kageyama et al. from dielectric measurements.

We now briefly discuss the diffusion of a few highly mobile H atoms. Earlier in this section, we have seen that the MSD of H atoms can increase occasionally very rapidly for a hydrogen load of 10 H atoms per void. This behavior of the MSD has been attributed to the movement of few H atoms out of the void region, defined by a sphere of radius 8 Å. It has been observed that such a steep rise of the MSD (see Figure 2a at 12 ps) originates from the high mobility of few H atoms inside the hydrogenated voids. Figure 5 shows the plot of the instantaneous kinetic energy (KE) of H atoms at 400 K, obtained from using the GGA for a period 20 ps. The distribution of the KE values...
along the Y axis for a given H atom indicates the range and the frequency of the KE of the atom during the course of AIMD simulations. An analysis of the real-space trajectory of the H atoms in the vicinity of the void region in models M2 and M3, and the results from Figure 5 revealed that a few highly mobile H atoms, such as H7 in Figure 5a and H17 in Figure 5b, left the void region sometime during the course of simulations. This observation was found to be true for other voids as well, where H atoms were observed to leave the void region due to their high KEs. Conversely, we have noted that all the H atoms in model M4 remained inside the void throughout the course of simulation even at the high temperature of 700 K. The results obtained from such a void is shown in Figure 6, where the KE values of the H atoms are found to be considerably lower than those shown in Figure 5.

3.2. Kinetics of Si–H Bond Formation and Dissociation on Void Surfaces of α-Si

We now address a question of considerable importance concerning the kinetics of Si–H bond formation and dissociation on the surface of voids in α-Si. Although a complete understanding of these events requires the knowledge of bond formation and dissociation energies, the problem can be addressed approximately by considering the KE of H atoms in AIMD simulations.

Earlier, in Figure 3, we have seen that the MSD of H atoms inside the voids fluctuates within the range of 17–22 Å², depending upon the XC approximation. These values are indicative of the fact that the great majority of H atoms stay bonded to Si atoms on the void surface as the time evolution of the system continues. However, it has been observed that a few H atoms in Si–H bonds can break free of the surface and move through the cavity to form new Si–H bonds at nearby active Si sites. This behavior is particularly pronounced at 700 K for a hydrogen load of 30 H atoms per void, where the presence of few H atoms with high KE makes it possible to dissociate an existing Si–H bond and form a new Si–H bond in the vicinity of the void surface.

This is shown in Figure 7, which shows the snapshots of two silicon–hydrogen bonding configurations obtained from the GGA at 3 and 18 ps. Figure 7a shows a set of four Si–H bonds, involving H12, H16, H24, and H30, indicated as green–red pairs, at 3 ps, which are found to dissociate later during simulation. The breaking of Si–H bonds is reflected in Figure 7b, which shows that the H atoms (red) resulted from the dissociation of SiH bonds diffused through the void-surface region to form new Si–H bonds, which are indicated as yellow–red pairs at 18 ps. The presence of a green–red pair suggests that the Si–H broke and formed again, but with another (dissociated) H atom.

Figure 7. The kinetics of Si–H bond formation and dissociation at 700 K for a hydrogen load of 30 H atoms per void obtained from the GGA in model M4. a) The Si–H bonds that dissociated during MD simulations are shown as green–red pairs at 3 ps. b) The H atoms (red) resulted from the dissociation of SiH bonds diffused through the void-surface region to form new Si–H bonds, which are indicated as yellow–red pairs at 18 ps. The presence of a green–red pair suggests that the Si–H broke and formed again, but with another (dissociated) H atom.

Figure 6. The distribution of KE along the Y axis for H atoms inside a void (in model M4) at 700 K. The range and the frequency of KE values are indicated by the numbers and shading (of red circles) along the Y axis, respectively.
3.3. Morphology of Hydrogenated Voids in a-Si

The discussion in the preceding subsections so far is mostly confined to the dynamics of H atoms inside the voids and to what extent the motion of H atoms is affected by the XC approximation and the hydrogen load inside the voids, as far as the MSD of H atoms is concerned. We now examine the microstructure of hydrogen distributions on void surfaces and the morphological character of the voids, which result from the movement of hydrogen and silicon atoms in the vicinity of the voids for a varying hydrogen load. In particular, we discuss the formation of various BHs and NBHs, as well as the restructuring of the void surfaces during annealing, with an increasing concentration of H atoms within voids, and the dependence of hydrogen microstructure on the LDA and GGA. The BH and NBH play an important role in characterizing the structural and optical properties of a-Si: H\cite{9,15,16} These properties can be studied experimentally using an array of experiments, such as positron-annihilation lifetime (PAL) spectroscopy\cite{15}, Rutherford backscattering spectrometry (RBS)\cite{16}, hydrogen-effusion measurements\cite{29,30} and Fourier-transform infrared spectroscopy-attenuated total reflections (FTIR–ATRs)\cite{31}

Table 1 shows the statistics of various silicon–hydrogen bonding configurations and NBHs, e.g., H2 molecules and one or two isolated H atoms, near the void, defined by a spherical region of radius 8 Å. The presence of isolated H atoms in the network is an artifact (of simulations), which arises from our choice of the cutoff value of 1.65 Å for Si–H bonds. It has been observed that the isolated H atoms at 700 K, listed in Table 1, are at a distance of 1.66 Å from the nearest Si atom (for the GGA) and at distances of 1.67 and 1.7 Å from the neighboring Si atoms (for the LDA). Thus, these H atoms are not truly isolated and can be viewed as somewhat stretched Si–H bonds at 700 K. The results from Table 1 and an analysis of the void surfaces for the hydrogen load of 10, 20, and 30 H atoms per void show that the surface of the

![Figure 8](image1.png)

Figure 8. a) The KE of two H atoms, H2 and H30, in monohydride Si–H bonds at 700 K. a) The KE of H2 atom is found to stay below the average KE at 700 K. b) The Si–H bond involving the high-energy H30 atom dissociates near 5.5 ps and forms a new Si–H bond. The average KE of the H atoms is indicated as a dashed horizontal line (green). The KE values correspond to a central moving average over 100 fs.

![Figure 9](image2.png)

Figure 9. a) The variation of the bond length of a stable monohydride pair (Si40, H2) with time. b) The dissociation of a monohydride Si–H bond during MD simulations. The Si–H bond associated with the pair (Si48, H30) breaks at 5.3 ps, which is followed by the formation of a new pair (Si65, H30) at 5.6 ps. The isolated state of H30 atom is indicated by red squares, where the “bond length” refers to the distance between H30 and the nearest Si atom. The cutoff value of the Si–H bond length, 1.65 Å, is indicated by a dashed horizontal line (red).
bonds are also spotted for a hydrogen load of 20 \( \text{H}_2 \) molecules, and SiH/\( \text{SiH}_2 \) bonds. Hydrogen atoms outside the void region are listed as Ex. Hydrogen-bonding configurations with respect to the XC approximation from Table 1 that, in comparison with the GGA, the LDA overestimates the number of SiH bonds but underestimates the count of \( \text{H}_2 \) molecules. The presence of few SiH bonds begins very rapidly within the first few picoseconds and then it gradually converges by 18 ps. The LDA seems to overestimate the number of SiH bonds by about 32\%, compared with the same from the GGA (see, for example, Table 1). This noted difference between the LDA and the GGA results indicates the need for choosing an accurate XC functional and a long simulation time for studying the microstructure of SiH and SiH\(_2\) on the walls of the voids. Likewise, the formation of H–H pairs, or H2 molecules, during annealing is shown in Figure 12b, by plotting the evolution of the RMS distance between two pairs of H atoms at 400 K in the GGA for a hydrogen load of 20 H atoms per void. The first H2 molecule, consists of (H1, H12), was formed within the first 4 ps, whereas the second one, (H5, H13), was formed near 14 ps. The RMS distances between the pair of H atoms and the corresponding standard deviation are obtained by averaging over a moving time window of width 200 fs.

We conclude this section by making a few comments on the linear size and the shape of the voids obtained from annealing and total-energy relaxations in the presence of H atoms. Although the linear size of a void can be estimated from the radius of gyration of a set of atoms, which de

| H load | XC   | H\( \text{iso} \) | H\(_2\) | SiH  | SiH\(_2\) | Ex\( \text{H} \) |
|--------|------|-----------------|--------|------|-----------|---------------|
| 400 K  |      |                 |        |      |           |               |
| 10     | GGA  | 0               | 15.0   | 67.50| 0         | 17.50         |
|        | LDA  | 0               | 10.0   | 82.50| 0         | 7.50          |
| 20     | GGA  | 0               | 32.50  | 47.50| 10.0     | 10.0          |
|        | LDA  | 0               | 25.0   | 62.75| 5.0      | 7.50          |
| 30     | GGA  | 0               | 45.0   | 40.0 | 11.67    | 3.33          |
|        | LDA  | 0               | 28.33  | 47.50| 18.33    | 5.83          |
| 700 K  |      |                 |        |      |           |               |
| 10     | GGA  | 0               | 25.0   | 55.0 | 0        | 20.0          |
|        | LDA  | 0               | 0      | 77.50| 5.0      | 17.50         |
| 20     | GGA  | 0               | 27.50  | 45.0 | 12.50    | 15.0          |
|        | LDA  | 0               | 17.50  | 66.25| 2.50     | 13.75         |
| 30     | GGA  | 0.83 (1\*)      | 35.56  | 41.11| 8.89     | 13.33         |
|        | LDA  | 1.65 (2\*)      | 16.67  | 56.67| 11.67    | 13.33         |

Table 1. Statistics of bonded and NBHs inside voids for varying hydrogen loads. The tabulated values indicate the amount of H atoms (in percent of total H) that resides as isolated hydrogen (H\( \text{iso} \)), H\(_2\) molecules, and SiH/SiH\(_2\) bonds. Hydrogen atoms outside the void region are listed as Ex. Asterisks denote the actual number of isolated H atoms.
by constructing the minimal convex polyhedron, or a convex hull, formed by the set of void-surface atoms. The approach assumes that any restructuring of the void surface—caused by the movement of the void-surface atoms—would be reflected in the shape of the convex hull, which is associated with the void region. However, it has been observed that the actual 3D shape of voids can be rather complex and nonconvex in nature, and it may not be represented accurately by a convex polyhedron. An approximate nonconvex surface can be constructed via convolution of the position of the void-surface atoms using the 3D Gaussian basis functions and choosing a suitable value of the isosurface parameter for the atomic pseudosurface of Si and H atoms. The Xcrysden package can generate such pseudosurfaces. A more general discussion on the reconstruction of such nonconvex void shapes can be found in the studies by Biswas et al. and Biswas and Elliott.

Figure 13 shows the approximate shape of a void with a hydrogen load of 30 H atoms per void (in model M3) obtained from annealing at 400 K, followed by a total-energy optimization. The convex polyhedra shown in Figure 13a,b correspond to the set of void-surface atoms obtained from the GGA and the LDA, respectively. The respective nonconvex shapes of the void, obtained via the convolution of the same set of void-surface atoms using the Gaussian functions centered at the atomic sites, are shown in Figure 13c,d. Although the polyhedra in Figure 13a,b appear different, the difference is not particularly noteworthy.
as far as the hull radius, the convex-hull volume, and the sphericity of the polyhedra are concerned. These values are listed in Table 2. The sphericity, $S$, of an object is defined as the ratio of the surface area of a sphere, $A_s$, to that of the object, $A$, both having an identical volume $V$. This definition leads to \[^{[15]}\]

\[ S = \frac{A_s}{A} = \frac{4\pi(6V)^{2/3}}{A} \] (1)

Here, we have used the volume and the corresponding surface area of the convex hull of a void surface to estimate $V$ and $A$, respectively. A review of $S$ and other values in Table 2 suggests that the LDA marginally overestimates the volume of the void, which is evident from the gyrational and hull radii, and the hull volume of the voids for all hydrogen loads and temperatures. This observation is also consistent with the somewhat smaller number of monohydride Si–H bonds but underestimates the presence of H$_2$ molecules inside the cavities, irrespective of the annealing temperature and the concentration of hydrogen. The surfaces of the voids are found to be primarily passivated with monohydride Si–H bonds and a few dihydride SiH$_2$ bonds at high concentration of hydrogen. Neither the LDA nor the GGA shows any presence of SiH$_3$ configurations even for a high concentration/load of 30 H atoms per void. The densities of the bonded and NBHs observed in this work are found to be consistent with those from the infrared and RBS measurements. The study reveals that the kinetics of Si–H bond formation and dissociation during AIMD simulations can be approximately described and understood by considering the (translational) KE of H atoms inside the voids. Hydrogen atoms with a KE value significantly higher than the average KE of the system at a given temperature are found to dissociate from Si–H bonds on the surface of the voids. The resulting isolated H atoms then diffuse through the void region to form new bonds with nearby active Si atoms within a fraction of a picosecond in our simulations. Finally, a somewhat higher value of the MSD of the H atoms within voids in the LDA can be attributed to the reconstruction of the void surface through the formation of Si–H bonds. This has been found to be reflected in linear size of the voids obtained from a convex-hull approximation.

4. Conclusion

In this article, we have studied the ab initio dynamics of hydrogen atoms inside voids in a-Si with an emphasis on hydrogen diffusion and the resulting structure of the void surfaces with respect to a varying concentration of hydrogen at 400 and 700 K. A comparison of the results obtained from the LDA and the GGA reveals that the former considerably overestimates the number of monohydride Si–H bonds but underestimates the presence of H$_2$ molecules inside the cavities, irrespective of the annealing temperature and the concentration of hydrogen. The surfaces of the voids are found to be primarily passivated with monohydride Si–H bonds and a few dihydride SiH$_2$ bonds at high concentration of hydrogen. Neither the LDA nor the GGA shows any presence of SiH$_3$ configurations even for a high concentration/load of 30 H atoms per void. The densities of the bonded and NBHs observed in this work are found to be consistent with those from the infrared and RBS measurements. The study reveals that the kinetics of Si–H bond formation and dissociation during AIMD simulations can be approximately described and understood by considering the (translational) KE of H atoms inside the voids. Hydrogen atoms with a KE value significantly higher than the average KE of the system at a given temperature are found to dissociate from Si–H bonds on the surface of the voids. The resulting isolated H atoms then diffuse through the void region to form new bonds with nearby active Si atoms within a fraction of a picosecond in our simulations. Finally, a somewhat higher value of the MSD of the H atoms within voids in the LDA can be attributed to the reconstruction of the void surface through the formation of Si–H bonds. This has been found to be reflected in linear size of the voids obtained from a convex-hull approximation.

Acknowledgements

The work was partially supported by the USA National Science Foundation (NSF) under Grant No. DMR 1833035. The authors acknowledge the use of computing resources at the University of Southern Mississippi, supported by the NSF under the Major Research Instrumentation (MRI) program via Grant No. ACI 1626217.

Conflict of Interest

The authors declare no conflict of interest.

Keywords

amorphous silicon, hydrogen dynamics, voids

Received: September 13, 2020
Revised: November 17, 2020
Published online: January 7, 2021

Table 2. Linear [Å] and volumetric [Å$^3$] measures of the voids, reconstructed from the convex-hull approximation. $R_c$, $R_h$, and $V_h$ indicate the radius of gyration, the convex-hull radius, and the volume of the hull, respectively. $S$ and $N_h$ represent the sphericity and the number of atoms on the convex hull, respectively.

| H load | XC    | $R_c$ | $R_h$ | $V_h$ | $S$   | $N_h$ |
|--------|-------|-------|-------|-------|-------|-------|
|        |       |       |       |       |       |       |
| 400 K  |       |       |       |       |       |       |
| 10     | GGA   | 6.392 | 7.162 | 1275.04 | 0.884 | 41    |
|        | LDA   | 6.517 | 7.205 | 1305.45 | 0.888 | 41    |
| 20     | GGA   | 6.416 | 7.204 | 1295.95 | 0.883 | 41    |
|        | LDA   | 6.566 | 7.218 | 1316.80 | 0.890 | 42    |
| 30     | GGA   | 6.489 | 7.231 | 1318.42 | 0.885 | 42    |
|        | LDA   | 6.541 | 7.263 | 1332.45 | 0.887 | 42    |
| 700 K  |       |       |       |       |       |       |
| 10     | GGA   | 6.192 | 7.102 | 1242.36 | 0.883 | 40    |
|        | LDA   | 6.347 | 7.175 | 1287.78 | 0.887 | 41    |
| 20     | GGA   | 6.365 | 7.242 | 1299.63 | 0.880 | 39    |
|        | LDA   | 6.461 | 7.274 | 1323.15 | 0.879 | 39    |
| 30     | GGA   | 6.430 | 7.202 | 1319.43 | 0.892 | 42    |
|        | LDA   | 6.475 | 7.238 | 1317.39 | 0.882 | 41    |
[14] E. Guerrero, D. A. Strubbe, "Phys. Rev. Mater." 2020, 4, 025601.
[15] T. Sekimoto, M. Matsumoto, A. Sagara, M. Hishida, A. Terakawa, "J. Non-Cryst. Solids" 2016, 447, 207.
[16] T. Sekimoto, M. Matsumoto, A. Terakawa, "Jpn. J. Appl. Phys." 2018, 57, 08RB07.
[17] F. Wooten, K. Winer, D. Weaire, "Phys. Rev. Lett." 1985, 54, 1392.
[18] G. T. Barkema, N. Mousseau, "Phys. Rev. B" 2000, 62, 4985.
[19] J. M. Soler, E. Artacho, J. D. Gale, A. García, J. Junquera, P. Ordejón, D. Sánchez-Portal, "J. Phys.: Condens. Matter" 2002, 14, 2745.
[20] A. H. Mahan, Y. Xu, D. L. Williamson, W. Beyer, J. D. Perkins, M. Vanecel, L. M. Gedvilas, B. P. Nelson, "J. Appl. Phys." 2001, 90, 5038.
[21] D. L. Young, P. Stradins, Y. Xu, L. M. Gedvilas, E. Iwaniczko, Y. Yan, H. M. Branz, Q. D. L. Wang, "Appl. Phys. Lett." 2007, 90, 081923.
[22] The effective width of the void surface is chosen to be about 3 Å. This is based on the results from the test calculations at 700 K, which show that the majority of silicon atoms near the void surface can move up to a distance of 2.5–3.0 Å. This distance is roughly equal to the first nearest-neighbor distance between Si atoms in the amorphous environment. For structural analysis of voids, we thus use a void-surface region of 8 Å.
[23] J. Junquera, O. Paz, D. Sánchez-Portal, E. Artacho, "Phys. Rev. B" 2001, 64, 235111.
[24] N. Troullier, J. Martins, "Phys. Rev. B" 1991, 43, 1993.
[25] J. P. Perdew, A. Zunger, "Phys. Rev. B" 1981, 23, 5048.
[26] J. P. Perdew, K. Burke, M. Ernzerhof, "Phys. Rev. Lett." 1996, 77, 3865.
[27] S. Nosé, "J. Chem. Phys." 1984, 81, 511.
[28] S. Kageyama, M. Akagawa, H. Fujiwara, "Phys. Rev. B" 2011, 83, 195205.
[29] W. Beyer, "Sol. Energy Mater. Sol. Cells" 2003, 78, 235.
[30] W. Beyer, H. Wagner, "J. Appl. Phys." 1982, 53, 8745.
[31] T. Sekimoto, M. Matsumoto, M. Hishida, A. Terakawa, "Jpn. J. Appl. Phys." 2014, 53, 095501.
[32] Y. J. Chabal, C. K. N. Patel, "Phys. Rev. Lett." 1984, 53, 210.
[33] P. Biswas, S. R. Elliott, "J. Phys.: Condens. Matter" 2015, 27, 435201.
[34] A. Kokalj, "J. Mol. Graph. Model." 1999, 17, 176.
[35] H. Wadell, "J. Geol." 1935, 43, 250.