Electronic Structures and Induced-Hole Carriers of Covalent Semiconductors in External Electric Field

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To investigate electric-field-induced superconductivity in covalent semiconductors, band structures and induced-hole carrier density of hydrogenated Si(110) surface in an external electric field were calculated by using first principles full-potential linearized augmented plane-wave method. Results predict that the induced-hole carriers, which form a Fermi surface with a hole pocket centered at Γ point, are less than the critical density of superconductivity for boron-doped Si. Thus, electric-field-induced superconductivity at Si(110) surface may be difficult to achieve, which is in contrast to the case of diamond(110) surface. [DOI: 10.1380/ejssnt.2014.109]

Keywords: Electric field; Hole carriers; Silicon; Density functional calculations; Superconductivity

I. INTRODUCTION

Controlling carrier density in covalent semiconductors is one of the key issues in the fields of semiconductor physics and engineering. In diamond, carrier doping up to \(10^{21}\) cm\(^{-3}\) was succeeded by boron incorporation. It turns the system to be metallic and to excite superconductivity with superconducting transition temperature \(T_c\) of 11.4 K [1, 2]. Superconductivity in the other covalent semiconductors, e.g., boron-doped Si and SiC with \(T_c\) of 0.35 K [3] and 1.4 K [4], respectively, was further reported experimentally. In order to increase \(T_c\), more efforts to introduce high carrier doping will be practically demanded. In the chemical doping, however, structural disorders always cause an unnecessary complexity in physical properties and may suppress the \(T_c\) [5–7].

Alternately, electric-field-driven superconductivity at insulator surfaces recently has the subject of considerable attention [8], where an application of an external electric field that induces high carrier density opens an advanced way for searching and investigating superconductivity in insulating materials without the chemical doping. Previously [9], we investigated the electric-field-driven superconductivity in hydrogenated diamond(110) surface from first principles calculations, and found that an introduction of a negative electric field that induces hole carriers results in a metallic surface with accumulated hole carriers exceeding the critical carrier density responsible for superconductivity of boron-doped diamond [1].

Here, we extend first principles investigations for treating covalent semiconductors including hydrogenated Si(110) surface by means of first principles full-potential linearized augmented plane-wave (FLAPW) method [10–12]. Results predict that when an external electric field is introduced, the system turns to be metallic, where Fermi level \((E_F)\) locates the top of valence band around the Γ point and the Fermi surface with a hole pocket is formed. In contrast to the diamond case, however, the induced hole carriers are found to be less than the critical density for superconductivity of boron-doped Si.

II. METHOD AND MODEL

We consider the hydrogenated Si(110) surface, modeled as a single slab consisted of 13 atomic-layers, which is terminated by hydrogens on both sides of the slab, as shown in Fig. 1. In-plane lattice constants are assumed to match to those of bulk Si, and all atomic positions are fully optimized by the atomic force calculations in an electric field.

Calculations were carried out by using FLAPW method based on the local density approximation [13]. LAPW basis with a cut-off of \(|k + G| \leq 3.9\) a.u.\(^{-1}\) and muffin-tin (MT) sphere radii of 1.40 a.u. for Si and 0.65 a.u. for H atoms are used. Lattice harmonics with angular momenta up to \(l = 8\) for Si and 6 for H are employed to expand the charge density, potential, and wave functions.

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FIG. 2: Calculated band structure for Si (110) surface in zero external electric field. The energy gap is estimated to 0.61 eV.

FIG. 3: Calculated band structure for Si (110) surface in an external electric field of $-1.0 \text{ V/Å}$. Fermi energy, $E_F$, locates in the valence bands at Γ point. The reference energy ($E = 0$) places $E_F$. In inset, Fermi surface in the external electric field is presented, where the Fermi surface with a hole pocket centered at Γ point appears.

24×16 special $k$-points in two-dimensional Brillouin zone (BZ) for self-consistent calculations were employed. To introduce an external electric field along perpendicular to the surface, we place an electrode (a sheet of charge) far enough outside surface so that electrons have negligible overlap with the sheet [14, 15].

III. RESULTS AND DISCUSSION

We first present atomic and electronic structures of hydrogenated Si(110) surface in zero field. The hydrogen terminations on the Si(110) surface, with the H-Si(1) bond length of 1.52 Å, remove dangling bonds, which recovers almost bulk like $sp^3$ bonding geometry at the Si(110) surface. Also, as seen in the band structure in Fig. 2, the hydrogen terminations remove surface states arising from the dangling bonds that appear in the band gap in the clean surface. The system shows a semiconducting feature with an indirect band gap of 0.61 eV.

When an external electric field of $-1.0 \text{ V/Å}$ is introduced, the bond lengths are slightly compressed. The surface Si-H bond length decreases by 0.007 Å from that in zero field. For the layers below the surface, very small shortening in the bond lengths was observed, where the maximum change appears between the second and third Si layers (Si(2)-Si(3) in Fig. 1) by 0.008 Å. In contrast, as shown in Fig. 3, the surface becomes metallic where the $E_F$ locates at the top of valence bands at the Γ point, and the Fermi surface with a hole pocket centered at the Γ point appears. We confirmed that the area of the Fermi surface increases when the external electric field increases.

Calculated hole carrier density as a function of the $z$-position (along normal to surface plane) is shown in Fig. 4. This is demonstrated by the planar-averaged hole density along the $z$ axis, corresponding to the states in energy range from the $E_F$ to the top of valence band. Although the hole carriers are mainly induced in a few atomic-layers below the surface, they are rather dispersed inside the film, compared to that in diamond [7]. The highest value of the hole density results in $1.1 \times 10^{21} \text{ cm}^{-3}$ at the Si(2) layer, which is less than that in boron-doped Si in experiments, $4.0 \times 10^{21} \text{ cm}^{-3}$ [3].

The mechanism of superconductivity for boron-doped Si may be likely to be electron-phonon coupling [3, 16]. According to ab initio calculations with both virtual crystal and direct supercell calculations, the superconductivity of boron-doped Si is due to the softened optical modes near the zone center caused by the boron doping [16]. The electron phonon coupling constant, $\lambda$, was estimated to be 0.2-0.3, where the superconductivity above 1 K requires the hole doping over 5%, which corresponds to the hole density of $2.8 \times 10^{21} \text{ cm}^{-3}$ [16, 17]. Thus, the electric-field-induced hole density of the Si(110) surface is not enough to provide the hole density to excite superconductivity.
IV. CONCLUSION

We investigated the electronic structures and the induced hole carriers of the hydrogenated Si(110) surface in an electric field by means of first-principles calculations. The electric field introduces the metallic character at the surface where the Fermi surface with the hole pocket appears at the center of the Γ point. The hole carriers are induced up to $1.1 \times 10^{21} \text{cm}^{-3}$ at the Si(2) layer, which are less than the critical hole density of the boron-doped Si superconductivity, and the Si(110) surface may be difficult to excite superconductivity by the electric field.

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