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Spontaneous dissociation of a conjugated molecule on the Si(100) surface

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The adsorption mechanism of α-sexithiophene (α-6T) on the clean Si(100)-(2 × 1) surface has been investigated using scanning tunneling microscopy (STM) and first principles electronic structure calculations. We find that at submonolayer coverage, the α-6T molecules are not stable and dissociate into monomers. We observe two different configurations of the monomers and have discussed the corresponding adsorption geometries based on theoretical calculations. The calculations elucidate how the fragments are absorbed on the surface, giving rise to the observed STM images. With increasing coverage, the STM images show the existence of complete α-6T molecules. In addition, results of the adsorption behavior of α-6T molecules on the H-passivated Si(100)-(2 × 1) surface are reported. On this surface the molecules are highly mobile at room temperature due to the weak molecule-substrate interaction. The STM results also indicate that they can easily be anchored at the defect sites. © 2002 American Institute of Physics. [DOI: 10.1063/1.1480857]

I. INTRODUCTION

In recent years studies of organic molecular adsorption on silicon surfaces have been largely stimulated by the scientific interest in organic modifications of silicon surfaces.²⁻⁴ By tapping into the vast resources of organic molecules, a wide variety of functionalities can be designed and incorporated with the existing silicon-based technology for new and improved semiconductor devices, e.g., molecular electronics, chemical sensors and “DNA chips.” Prior to realization of hybrid organic molecule-silicon devices, a detailed understanding of adsorption behavior of various organic molecules on silicon surfaces becomes essential for controllably incorporating the organic functionalities with the silicon micro-electronics technology.³

In the area of organic molecular adsorption on silicon, a lot of work has been focused on alkenes.⁴ Among the alkenes ethylene was the first molecule for which the adsorption structure on the Si(100)-(2 × 1) surface was well established by experimental methods such as scanning tunneling microscope (STM)⁵,⁶ and traditional surface science techniques⁷⁻¹⁰ as well as by theoretical simulations.¹¹,¹² It has been demonstrated that ethylene reacts with the silicon surface via the interaction between the C–C double bond and silicon dimers, producing two Si–C bonds (the so-called di–σ bonding). Other alkenes are found to react in a similar fashion with the Si(100) surface,¹³⁻¹⁷ i.e., they bind to the surface also via the di–σ bonding configuration. Besides simple alkenes, conjugated dienes such as 1,3-butadiene and 1,3-cyclohexadiene have been investigated.¹⁸⁻²⁰ It is found that these conjugated dienes can react with Si–Si dimers not only via one of the two C–C double bonds, but also through both C–C double bonds, resulting in multiple bonding structures on the surface. The reaction between a Si–Si dimer and two C–C double bonds is referred as the Diels-Alder reaction. The interesting aspect of this reaction is that it produces a six membered ring containing a residual C–C double bond which may prove useful in further controlled chemical reactions at the interface. Furthermore, interaction between aromatic compounds, e.g., benzene,²¹⁻²⁵ toluene²⁶ and thiophene,²⁷ and the Si(100) surface has also been investigated. All these molecules show multiple bonding geometries on the surface. For instance, room-temperature STM images²²,²⁵ clearly illustrate that three different images of the benzene molecule can be observed on the Si(100)-(2 × 1) surface and these images correspond to three different adsorption geometries.

In our experiment, we focus on a large rodlike conjugated molecule, α-sexithiophene (α-6T), which is a prototype molecule for devices such as field effect transistors (FETs),²⁸⁻³¹ light emitting diodes (LEDs)²²,²³ and transistors (LETs).³² Besides the technological aspect of this molecule, the other reason for studying this molecule is that growth of oligothiophene thin films onto dielectric (oxidized silicon,³³ mica,²⁶⁻³⁸), semimetal (HOPG)³⁹ and metal surfaces (Au, Ag, Al)⁴⁰⁻⁴³ has been thoroughly studied. In comparison, the growth of large conjugated oligothiophene molecules onto...
clean semiconductor surfaces is relatively unexplored. In order to gain the control needed to form highly perfect thin films, it is of considerable importance to study the adsorption behavior of α-6T on silicon surfaces. In this work, by combining STM experiments with the first principles studies we elucidate the reactive processes occurring upon adsorption of α-6T onto the clean Si(100)-(2×1) surface and compare with the adsorption process of α-6T on the corresponding H-passivated Si surface. The difference of the two Si surfaces lies in the chemical reactivity. Due to the existence of dangling bonds on the clean Si surface, this surface is more reactive than the H-passivated surface where the dangling bonds are passivated by hydrogen atoms. In this study we will show that the high reactivity of the Si~100~ surface leads to fragmentation of the α-6T molecules upon adsorption while the molecules adsorb intactly on the H-passivated surface.

II. EXPERIMENT

The experiments were performed in an ultrahigh vacuum (UHV) chamber pumped by an ion pump and a sublimation pump. The base pressures can be kept below \(1 \times 10^{-10}\) Torr. A load lock system enables efficient transfer of samples and tips without disrupting the vacuum system. The main chamber is equipped with a commercial STM (Danish Micro Engineering A/S). The molecular deposition system is attached to the load lock and can therefore be pumped separately without disrupting the vacuum condition in the main chamber. The deposition system consists of an evaporator, a shutter and a quartz crystal microbalance.

In the experiment commercial n-type Si(100) wafers with a resistivity of 1 \(\Omega\cdot\text{cm}\) were used as substrates. The clean reconstructed Si(100)-(2×1) surfaces were prepared inside the main chamber by flashing the sample up to 1473 K for a few seconds, and cooling the sample with a rate around 5 K/s to room temperature. Prior to the flashing process the sample was degassed at 923 K for at least 12 h. The base pressure during annealing was kept below \(5 \times 10^{-10}\) Torr. The 2×1 reconstruction was verified by STM imaging. H-passivated Si(100)-(2×1) surfaces were prepared by exposing clean Si(100)-(2×1) surfaces to atomic hydrogen for 6 min with a chamber pressure of about \(1 \times 10^{-6}\) Torr. During exposure the temperature of the silicon substrate was kept at 623±25 K. A hot W filament was used for cracking the hydrogen gas.

The molecular structure of α-sexithiophene (hereafter referred to as 6T) is depicted in Fig. 1. The mass spectroscopy measurement showed that except for the initial ionization of the 6T molecules to 6T\(^{+}\) ions, no fragmentation was observed during sublimation. The melting point of 6T is 576 ± 1 K. Prior to the deposition, the molecules were degassed, and then evaporated on the sample surfaces via sublimation at a rate of 0.01–0.02 Å/s, which was monitored with a quartz crystal microbalance. The deposition time for the clean silicon surfaces ranged from 5 to 10 s, while for the H-passivated surface the time ranged from 30 to 60 s with a rate of 0.05–0.10 Å/s. The coverage of the molecules was estimated from the STM images. Tungsten tips were prepared by electrochemical etching and were subsequently cleaned inside the chamber by Ar\(^+\) ion bombardment. Constant current STM images were acquired with the sample bias ranging from −2.0 to −2.5 V.

![FIG. 1. Molecular structure of α-sexithiophene. The α and α’ mark the C atoms adjacent to the sulfur atoms.](image1)

![FIG. 2. (a) An STM image of the Si(100)-(2×1) surface after deposition of a submonolayer coverage of 6T molecules \(I_{\text{sample}} = 0.15\) nA; \(V_{\text{sample}} = -2.5\) V. (b) The magnified STM image of the highlighted part of (a). A and B mark the two different types of the ball-like protrusions. (c) The cross-section profile along the line in (b).](image2)
III. RESULTS

A. Submonolayer coverage

Figure 2 shows an STM image of the Si(100)-(2×1) surface after deposition of 6T molecules. The underlying Si(100) surface structure is visible in the image. On the Si(100) surface, adjacent silicon atoms on the topmost layer pair into dimers via a strong $\sigma$-bond and a weak $\pi$-bond. In the filled state image, Si dimers are imaged as bean-shaped protrusions arranged in rows with an interrow distance of 7.68 Å. At the step edge, with an atomic step height of 1.36 Å the dimer rows on the upper terrace are orthogonal to those on the lower terrace due to the directional covalent bond. Beside the dimer row structure, the image also clearly reveals many bright ball-like protrusions on the surface. They are distributed across the surface as either isolated spots or small clusters. There is no indication of preferential adsorption towards the step edges. To identify these bright ball-like protrusions on the surface, a cross-section profile analysis was performed for 17 isolated bright spots. The size is given by the value of the full width at half-maximum (FWHM) of cross-section line profile. A line profile for one selected ball-like protrusion in the magnified STM image [Fig. 2(b)] is displayed in Fig. 2(c). The sizes of the spots measured in three directions are depicted in Fig. 3. The sizes measured in two directions, perpendicular to and along the dimer rows, are referred as the length and the width, respectively. The third one corresponds to the height of the protrusions. The average value of length, width and height of these spots are found to be $6.5 \pm 0.6$ Å, $5.8 \pm 0.6$ Å, and $1.2 \pm 0.2$ Å, respectively. This result was further confirmed by many other STM images recorded for different samples and tips. Although the values for the length and the width are scattered, they are significantly smaller than the length of a 6T molecule, which is about 24 Å. Thus, no 6T molecules are observed after deposition of 6T molecules on the Si(100)-(2×1) surface at low coverage.

Figure 4 displays the molecular structure of thiophene and its oligomers (2≤$n$≤6) obtained by optimizing the molecular structure using the semi-empirical PM3 method. And we report the lengths as measured by the distance between the two terminal H atoms bound to the C$_a$ atoms at the end rings. By comparing the size of the protrusions obtained in Fig. 2 with that of oligothiophenes, we find that these bright protrusions have sizes most close to 1T. The average ratio of the length to the width of these spots is found to be 1.1, in good agreement with a 1T molecule whose ratio is 1.3. It is also well known that the size of adsorbates estimated by the cross-section profiles normally does not reflect the true geometric size of the adsorbates due to the complex interplay between the STM tip and the electronic structure of the substrate and adsorbate. Thus, the discrepancy between the measured and expected size of 1T could be caused by this kind of the tip-sample convolution. Our observation suggests that upon adsorption on the Si(100)-(2×1) surface at submonolayer coverage, due to the reactive Si dangling bonds, 6T molecules are not stable and dissociate into 1T fragments by breaking the C$_a$–C$_b$ bonds.

The position of these protrusions with respect to the underlying Si dimer rows was further investigated by a careful inspection of the images. It was found that there are two different types of ball-like protrusions, marked by A and B in Fig. 2(b). For type A the ball-like protrusion is located exactly in the middle of the silicon dimer row, so referring to the dimer row it is symmetric. For type B, the bright spot is closer to one of the silicon dimer atoms, and therefore they are asymmetric with respect to the dimer row. Since many of...
these bright spots are agglomerated in clusters, it is difficult to make a reliable estimation of the relative percentage of types A and B.

B. High coverage

Figure 5 shows a sequence of STM images of the clean Si surface upon increasing the dose of 6T. With increasing coverage, the concentration of bright ball-like protrusions is also increasing, as shown in Fig. 5(a). Most protrusions are in clusters and located on top of the dimer rows. Further deposition leads to new features, bar-shaped (BS) protrusions, appearing in the image [Fig. 5(b)]. It is found that all the BS protrusions with different lengths have the long axes oriented in the same direction as the underlying dimer rows. Moreover, the BS protrusions appear brighter than most of the ball-like protrusions. The height profiles reveal that BS protrusions are about 0.4 Å higher than the slightly dim ball-like spots. This gives us an indication that these BS protrusions are not directly lying on top of the silicon dimers. Instead they may locate on top of the ball-like protrusions. By further increasing the coverage, BS protrusions with different sizes become dominant on the surface, as shown in Fig. 5(c). We do not observe any ordering of the BS protrusions.

Seven different BS protrusions, labeled in Fig. 5(b), were selected, and the measured lengths of these seven protrusions are plotted in Fig. 6. Due to the tip-sample convolution, we expect that the apparent size of these BS protrusions will be larger than their real size. Therefore, we also estimated the size broadening by calculating the size difference between the average size of the ball-like protrusions in the image and the size of 1T. The average length and width of the ball-like protrusion are found to be 7.1 and 6.1 Å, respectively, which are in accordance with the values obtained for the protrusions in Fig. 2. Thus the size broadening in this image is estimated to be about 2.5 Å. In order to get a quick impression what these BS protrusions correspond to, in the same figure the “lengths” of thiophene and oligomers up to 6T, including the tip broadening effect, are plotted as the dashed lines for the eye guiding. It is seen that the sizes of the different BS protrusions are close to the sizes of oligothiophene for \(2 \leq n \leq 6\). We admit that there are some uncertainties in the method used to estimate the tip broadening effect. However, we are sure that longer oligomeric fragments (ring number > 1) start to appear on the surface after

![FIG. 5. A sequence of STM images [from (a) to (c)] of the clean silicon surface upon increasing doses of 6T. In (b) seven bar-shaped protrusions of different lengths are labeled and the measured lengths are plotted in Fig. 6.](image)

![FIG. 6. The lengths of the seven bar-shaped protrusions labeled in Fig. 5. The molecular lengths of thiophene and oligomers up to 6T, including the tip broadening effect, are plotted as the dashed lines.](image)

![FIG. 7. Side view (a) and top view (b) of the cluster used to describe the Si(100) surface. There are a total of 56 Si and 48 H atoms. The solid rectangle in (b) shows the irreducible part of the surface.](image)
the dangling bonds are passivated with the monomeric fragments.

IV. FIRST PRINCIPLES MODELING

Though the experimental data indicate that, due to strong interaction with the dangling bonds, 6T breaks into monomers at submonolayer coverage, they are not able to give details on the position of the underlying atoms. An accurate modeling of these fragments on the surface, therefore, helps to elucidate where they are preferentially absorbed on the surface and the main interactions among the atoms, which give rise to the observed STM images. From the molecular structure of 6T, we see that there are two different monomers upon C<sub>a</sub>–C<sub>a</sub>' bond breaking. The four middle monomers (SC<sub>4</sub>H<sub>2</sub>) will be missing two H atoms, while the two outermost monomers (SC<sub>4</sub>H<sub>3</sub>) will be missing only one H atom, relative to a thiophene molecule (1T). For this reason, we have investigated theoretically the adsorption geometry of the two fragments on the Si(100) surface with the DACAPO<sup>45</sup> electronic structure code. These calculations were based on density functional theory (DFT), and the generalized gradient approximation of Ref. 46 (PW91) was used for the exchange correlation energy. The wavefunctions of the valence electrons were expanded in a plane-wave basis set with an energy cutoff of 20 Rydberg, and the core electrons of Si, C, H, and S were described by ultra-soft pseudo potentials.<sup>47</sup> Within this approach the Si lattice constant was determined to be \( a_0 = 5.46 \) Å. The Si(100)-p(2×2) surface was modeled by a c(4×4) slab with four layers, thus a total of 32 Si atoms. The last two layers were fixed in bulk positions and dangling bonds on the lower surface were passivated with H atoms. A Gaussian broadening of 0.1 eV and a (221) Monkhorst-Pack mesh<sup>48</sup> for the Brillouin zone sampling (two k-points in the irreducible zone) were used. The atomic positions of the Si atoms in the first two layers were relaxed until the forces on the atoms were smaller than 0.2 eV/Å.

A systematic search for SC<sub>4</sub>H<sub>2</sub> adsorption positions is too computationally demanding using this approach, and instead the semi-empirical Austin model (AM1)<sup>49</sup> was applied to locate all the potential adsorption positions. These calculations were performed with the GAUSSIAN 98 package,<sup>50</sup> which can only describe cluster geometries. For this purpose we extracted a four-layer (4×3) silicon cluster from the slab coordinates, and passivated all nonsurface dangling bonds with H atoms (see Fig. 7). The solid rectangle in Fig. 7(b)
Two layers were allowed to relax to a second set of calculations where also Si atoms in the first lowest energy obtained from this search were used as input few degrees relative to the surface. The geometries with the surface within a dimer row. In class ~

Different initial positions, and in each geometry all SC4 H2 were tilted a fixed positions. In order to facilitate rotations perpendicular the molecule was turned 360 degrees parallel to the long axis and three along the short axis. In each class the mass of the molecular fragment is positioned within this area that can represent the entire surface. If the center of this area is placed in eight different points, and in each geometry all SC4 H2 are relaxed. Bond lengths are in Ångström and angles in degrees.

<table>
<thead>
<tr>
<th>Energy</th>
<th>SC4H2-al</th>
<th>SC4H2-all</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM1</td>
<td>-8.73 eV</td>
<td>-7.91 eV</td>
<td>-4.81</td>
</tr>
<tr>
<td>DFT(1)</td>
<td>-4.44 eV</td>
<td>-3.65 eV</td>
<td></td>
</tr>
<tr>
<td>DFT(2)</td>
<td>-4.88 eV</td>
<td>-4.28 eV</td>
<td></td>
</tr>
</tbody>
</table>

Structural parameters

C1–C2 1.524 1.513 1.521
C2–C1 1.468 1.633 1.476
C1–C3 1.368 1.517 1.347
S–C1 1.759 1.722 1.881
S–C2 1.855 1.717 1.784
Si1–C1 1.897 1.931 1.908
Si2–C2 2.058 2.006 1.933
Si3–C3 1.993 1.936 2.539
Si4–C4 2.539 2.347
Si3–S 2.347 109.5 106.3
\(\angle SC1C2\) 106.5 109.0 106.5
\(\angle C1C3C2\) 111.1 109.0 110.8
\(\angle C2C1C3\) 117.3 109.3 117.1
\(\angle C1C2S\) 105.9 109.3 112.5
\(\angle C2SC1\) 94.4 99.7 92.5
\(\angle Si2C2C3\) 119.2 104.1 118.8
\(\angle Si2C3C2\) 101.8 94.4 106.4
\(\angle Si3SC1\) 115.1

*Data from Reference 27.

TABLE I. Adsorption energy and structural parameters for the adsorption geometries of SC4H2 shown in Fig. 9. For comparison we show the AM1 energies that are more than 1 eV lower than the adsorption energies calculated using the AM1 method by Jeong, Lee and Kim.27 Comparison of the geometrical parameters shows that the most favorable adsorption geometry al is very similar to the most favorable adsorption geometry of thiophene. In this geometry there is a double bond between C3 and C4 atoms. The main differences observed for SC4H2 are due to the larger reactivity of the C1 and C4 atoms of the diradical fragment, with respect to

Within class (a) there are two local minima, illustrated in Figs. 9(a) and 9(d), which we denote geometry al and alI. In Table I we show the adsorption energies of these two geometries as obtained by subtracting the total energy of the isolated molecule and the energy of the clean Si surface from the total energy of each geometry. These two AM1 geometries were used as starting points for supercell DFT calculations. In Table I we show the calculated adsorption energies when the coordinates of the molecule and the first layer Si atoms were relaxed within the DFT approach (DFT1) and when coordinates of three layers of Si atoms were relaxed (DFT2). Compared to the AM1 results, the absolute value of the adsorption energy changes significantly, while relative differences are quite similar. Thus, in this case, the AM1 results seem to be quite accurate in predicting the most stable geometry. It is interesting to compare the obtained geometries with the adsorption geometries of thiophene on Si(100) which have been calculated with the AM1 method by Jeong, Lee and Kim.27 Comparison of the geometrical parameters shows that the most favorable adsorption geometry al is very similar to the most favorable adsorption geometry of thiophene. In this geometry there is a double bond between C3 and C4 atoms. The main differences observed for SC4H2 are due to the larger reactivity of the C1 and C4 atoms of the diradical fragment, with respect to
thiophene. In fact, though the C3–C4 bond preserves its character of a double bond, as the case of thiophene, the C4 atom has a larger interaction with the neighbor Si atom, indicated by the shorter Si1–C4 bond length. Besides, the Si1–C1 bond length is shortened, while that of Si2–C2 is lengthened. Moreover, as shown for the thiophene molecule, the S atom has a significant bonding interaction with a Si atom. The second stable chemisorbed state of thiophene found by Jeong, Lee and Kim is the one where the C1 atom is not bound to a Si atom. We do not find such a chemisorbed state for the SC4H2 molecule, presumably because in our case C1 and C4 atoms are the most reactive atoms. In the second stable chemisorbed state of thiophene, the C1 atom is not bound to a Si atom, while that calculated for the SC4H2 molecule shows larger interactions, especially among Si1 and C1 atoms as well as Si3 and C4 atoms. In this case, the diradical lies on the surface with C1 and C4 atoms, and C2 and C3 atoms almost equidistant from the closer Si atoms, while the S atom is not bound to a Si atom. In this chemisorbed state all four C atoms interact with the underlying silicon atoms, thus the C–C bonds have a length of a single C–C bond or of a broken bond (C2–C3).

To compare the calculated adsorption geometry with the experimental data, we have made STM simulations using the Tersoff-Hamann model.51 Figures 9(c) and 9(f) show the calculated constant current filled state STM images of aI and aII corresponding to a sample bias of −2.5 V and a local density of states (LDOS) of 10−5 Å−3. This value of the LDOS corresponds to a current around 104 nA.52 Due to the limited numerical accuracy of the plane wave expansion of the electronic states in the vacuum region, this is the lowest LDOS (current) value we can use. We have found that the corrugation in the calculated STM image only depends weakly on the LDOS value, and we expect only minor changes in the STM image for the LDOS value corresponding to the experimental current of 1 nA as used for recording the experimental image. Figure 10 shows the line profiles of the calculated images for both geometries aI and aII together with those for the experimental images of types A and B (Fig. 2) in two directions along the silicon dimer rows and in the perpendicular direction. Line profiles measured along a silicon dimer row and across dimer rows are used for aligning the profiles. The profiles of type B were taken by choosing a type B molecule locating on the left side of the silicon dimer row, while the line profiles along the dimer row were taken along the center of a dimer row. Hence an apparent height difference is visible between the two profiles taken along the dimer row [Fig. 10(a)] for types A and B. Besides, from the line profiles taken across the dimer rows it is also clearly seen that the type B molecule is located on the left side of the dimer. The peak at the position of 3.5 Å in Fig. 10(b) reveals the corrugation of the dangling bond.52 Comparison of the molecular profiles reveals that the width and the height of the profiles in the calculated images are qualitatively consistent with those in the experimentally recorded images. This result

<table>
<thead>
<tr>
<th>Energy</th>
<th>SC4H3–a (eV)</th>
<th>SC4H3–b (eV)</th>
</tr>
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<tbody>
<tr>
<td>AM1</td>
<td>−11.28</td>
<td>−11.16</td>
</tr>
<tr>
<td>DFT(1)</td>
<td>−3.08</td>
<td>−4.07</td>
</tr>
<tr>
<td>DFT(2)</td>
<td>−3.70</td>
<td>−4.49</td>
</tr>
<tr>
<td>DFT(3)</td>
<td>−3.70</td>
<td>−4.49</td>
</tr>
</tbody>
</table>
ments SC$_4$H$_3$ with respect to the surface was investigated. To interact with the Si atoms, the behavior of the terminal fragments SC$_4$H$_2$ monomers upon adsorption on the clean Si surface. The line profile of the aI geometry appears symmetric with respect to the silicon dimer row, and we therefore link type A to the aI geometry. The line profile of geometry aII is very similar to the one of geometry aI, and it would be very hard to differentiate these two geometries experimentally. However, since the adsorption energy of geometry aII is much lower than that of geometry aI, we do not expect geometry aII to be present on the surface.

Having shown how the four middle monomers (SC$_4$H$_2$) interact with the Si atoms, the behavior of the terminal fragments SC$_4$H$_3$ with respect to the surface was investigated. To investigate the adsorption of SC$_4$H$_3$ on the surface, we took as a starting point a selection of geometries that represented the local minima structures of category (a), (b) and (c) in Fig. 8. For each geometry we added one H atom to the SC$_4$H$_2$ molecule, and relaxed the coordinates. Figure 11 shows the four geometries, which have the lowest energy. To check the reliability of the AM1 results we used geometries (a) and (b) as starting points for the DFT calculations. The result is summarized in Table II. We see that there is a large difference between the AM1 results and the DFT results both in the absolute and relative energies. Thus it seems that in this case the AM1 results are not very reliable, which might be related to the fact that in this case the cluster has an odd number of electrons. The DFT(3) result in Table II is for a spin-polarized DFT calculation, and we see that spin polarization has a negligible effect on the DFT results. Since the AM1 results are not very reliable (relative energies have errors of 1 eV), we can not trust that the AM1 geometries are a good starting point for the DFT calculations. Thus, to find the adsorption geometry of SC$_4$H$_3$ with the lowest energy we would probably need to make a full search within the DFT approximation which is not feasible.

Figure 12 shows the DFT optimized structure and corresponding theoretical STM images of the two AM1 geometries of SC$_4$H$_3$ which have the lowest energy. Note that within DFT it is geometry SC$_4$H$_3$-b that has the lowest energy while within AM1 it is SC$_4$H$_3$-a. We see that the SC$_4$H$_3$-b geometry gives rise to an asymmetric image, however, with a dip in the middle which is not seen in the experimental images. In Fig. 13 we plot the line profiles of the theoretical images of the SC$_4$H$_3$-a and -b geometries (Fig. 12) together with those of the STM images of types A and B (Fig. 2). The discrepancies in corrugation and shape between the theoretical and experimental images indicate that the two geometries do not correspond to the experimental images. This can have two reasons, either we have not found the global minimum because we used AM1 geometries as starting points, and this method is not very reliable in this case, or it is favorable to break off an additional H atom from SC$_4$H$_3$, and SC$_4$H$_3$ is therefore not observed on the surface. To investigate the latter we compare the adsorption energy of SC$_4$H$_2$ with the energy when SC$_4$H$_3$ and a single H atom are adsorbed on the surface. The adsorption energy of H is 3.19 eV, while the energy cost of breaking a H atom from the C$_a$ atom of SC$_4$H$_3$ is 4.45 eV. Thus, from the adsorption energies in Tables I and II, we see that it is more favorable to keep the H atom attached to SC$_4$H$_3$. Hence, the asymmetric spots most likely correspond to SC$_4$H$_3$, but the geometry we have found in the theoretical calculation might not be the one with the lowest energy.

Thus the theoretical calculations support the experimental observation that upon adsorption on the silicon surface, 6T molecules dissociate into monomeric fragments by breaking the C$_a$–C’$_a$ bonds. The STM images reveal two different configurations: symmetric and asymmetric with respect to the underlying Si dimer rows and we postulate that the symmetric one could link to the SC$_4$H$_2$ monomer and the asymmetric one to the SC$_4$H$_3$ monomer. Since the ratio of SC$_4$H$_2$ to SC$_4$H$_3$ is 4:2, we would expect double as many symmetric bright spots compared to the number of asymmetric bright spots in the images. However, since many of these bright spots are in clusters, it is difficult to obtain the exact ratio of these two different types by performing a statistical counting.

To understand the dissociation mechanism of 6T, we estimated the total energy involved in the dissociation from bond energies. Albeit we can not find the exact energy for cleaving the C$_a$–C’$_a$ bonds, the mean dissociation energies of C–C single (3.61 eV) and double (6.34 eV) bonds could set
the lower and upper limits of the \( \text{C}_a - \text{C}' \) bond energy, respectively. For each broken C–C bond two Si–C bonds are formed. The calculated Si–C bond lengths are 1.897 and 1.925 Å, in accordance with the 1.89 Å bond length of the Si–C bond in organosilicon compounds.\(^{53}\) In organosilicon chemistry the typical bond energy for Si–C is 3.30 eV.\(^{54}\) Thus by breaking 6T into six monomers and forming ten new Si–C bonds the system will gain an energy, which will fall into the interval between \( 10 \times 3.30 \text{ eV} - 5 \times 6.34 \text{ eV} = 1.3 \text{ eV} \) and \( 10 \times 3.30 \text{ eV} - 5 \times 3.61 \text{ eV} = 14.95 \text{ eV} \). Hence, the dissociation of 6T is energetically favorable.

In general, the same argument applies to all hydrocarbons having single or double C–C bonds, i.e., upon adsorption it is energetically favorable to break single or double C–C bonds and form two new Si–C bonds for every broken C–C bond. However, for most studied hydrocarbons-silicon systems the C–C bond cleavage is not observed. For instance, unsaturated hydrocarbons, such as acetylene, ethylene or dienes,\(^{4}\) nondissociatively chemisorb on the Si(100)–(2 \times 1) surface while molecules having only single C–C bonds, e.g., methane or propane,\(^{13}\) do not adsorb at all. The reason for this lies on the fact the C–C bond cleavage may be prohibited by a high kinetic barrier. In comparison, it seems that the kinetic barrier is much lower for the dissociation of 6T on the clean silicon surface. We speculate that this is due to the following; when the \( \text{C}_a \) atoms react with the Si atoms, they will lose their \( \sigma \) bond character and instead become \( sp^3 \) hybridized. This can, for instance, be seen from the fact that the \( \text{C}_1 - \text{C}_2 \) bond length of 1.52 Å, as listed in Table I, is close to the C–C bond of 1.54 Å in alkanes.\(^{53}\) Since C atoms in a \( sp^3 \) hybrid state will tetrahedrally bind to other C atoms, such a rehybridization would cause a gas phase 6T molecule to twist from the planar conformation. However, when 6T is adsorbed on the surface, all monomeric rings are anchored to the Si atoms and the molecule cannot twist. Thus, the rehybridization will induce a large strain in the molecule that effectively lowers the kinetic barrier for the \( \text{C}_a - \text{C}' \) bond breakage.

**V. ADSORPTION OF 6T ON THE H-PASSIVATED SILICON SURFACE**

In Sec. III we have shown that 6T molecules dissociate on the clean Si(100)-2 \times 1 surface. After passivating the dangling bonds with these monomers, the surface becomes chemically inert and consequently the dissociation process slows down and longer oligomers start to appear on the surface. Therefore, we would expect that by terminating these highly reactive sites with atomic hydrogen, the adsorption behavior of 6T molecules on the H-passivated silicon surface will be different compared to the clean silicon surface. In addition, recently Lopinski et al.\(^{55}\) have demonstrated an approach for fabricating nanoscale organic structures on silicon surfaces, by employing minimal intervention by an STM tip and a spontaneous self-directed chemical growth process. Styrene molecule with C–C double bonds could form straight molecular lines on the H-passivated silicon surface. This self-directed chain growth provides a relatively fast way of forming and/or connecting nanostructures on silicon surfaces, in comparison with constructing nanostructures via an STM tip. Their work also motivates us to further investigate the adsorption behavior of 6T molecules on the H-passivated Si(100) surfaces.

Figure 14(a) is an STM image of the H-passivated silicon surface before 6T deposition. The reconstructed dimer row structure is still maintained after H-passivation. The bright spots on the surface are dangling bonds due to missing hydrogen atoms. After exposing this surface to 6T molecules, the recorded images show bright and oval shaped features having about the same size, but fuzzy. Figure 14(b) shows an example of an STM image recorded for the H-passivated Si(100) surface after deposition of 6T molecules. The average length of the features is found to be about 29 Å, close to the molecular length of 6T. Since these features were not observed before the 6T exposure, we believe that they are 6T molecules which are weakly adsorbed on the H-passivated silicon surface. During scanning, it is observed that bright features may follow the tip movement until they bind to dangling bonds. We believe that the bright noisy spots represent 6T molecules bound to dangling bonds at a single point around which it rotates randomly. Our measurements show no indication of chain reactions as in the case for styrene,\(^{55}\) or breakage of 6T as for the clean silicon surface mentioned above. Based on our results, we propose that 6T molecules probably interact with individual dangling bonds through the sulfur atoms. A similar mechanism was also proposed for thiophene adsorbed on Si(111)-(7 \times 7) at room temperature.\(^{56,57}\)

**VI. CONCLUSIONS**

We have investigated the adsorption mechanism of \( \alpha \)-sexithiophene on the clean and the H-passivated Si(100)-2 \times 1 surface by combining STM measurements with first principles electronic structure calculations. On the clean silicon surface the STM images reveal that at submonolayer coverage the molecules dissociate into monomeric fragments, which gives rise to two different STM spots, that are symmetric and asymmetric, with respect to the underlying silicon dimer rows. The dissociation arises from the strong interaction between the molecules and the dangling bonds. The theoretical calculations support the experimental results and link the STM images to the monomeric
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45 The Dacapo DFT code is publicly available at http://www.fysik.dtu.dk.