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Interaction of $C_{59}$Si with Si based clusters: a study of Janus nanostructures

Miao Miao Wu$^{1,2}$, Xiao Zhou$^{1,2}$, Jian Zhou$^{1,2}$, Qiang Sun$^{1,2}$, Qian Wang$^3$ and Puru Jena$^3$

$^1$ Department of Advanced Materials and Nanotechnology, Peking University, Beijing 100871, People’s Republic of China
$^2$ Center for Applied Physics and Technology, Peking University, Beijing 100871, People’s Republic of China
$^3$ Department of Physics, Virginia Commonwealth University, Richmond, VA 23284, USA

E-mail: sunqiang@pku.edu.cn

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Abstract

Using density functional theory with the generalized gradient approximation (GGA), we show that carbon–silicon Janus anisotropic nanostructures can be synthesized by using $C_{59}$Si heterofullerene as a seed where the doped Si atom preferentially attaches to some well-known silicon and silicon based clusters such as $Si_{10}$, $WSi_{12}$, $TiSi_{16}$, and $BaSi_{20}$. The interaction energy of these clusters with $C_{59}$Si varies from 0.9 to 1.9 eV. The anisotropy of the resulting carbon–silicon Janus structures produces large dipole moments (4–9 D), anisotropic distributions of electronic orbitals, and the anisotropic reactivity.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

Nanoparticles, due to their small size and low dimensionality possess unique properties that differ from their bulk form. The large surface area of these particles and quantum confinement are mainly responsible for these properties. The minimization of surface energies usually leads nanoparticles to have spherical symmetry and hence these particles are not suitable as multifunctional materials. Consequently there has been a great deal of effort in recent years to synthesize anisotropic nanoparticles where different ends of the particle may have different properties. For example, one end may be hydrophilic while the other end is hydrophobic. These anisotropic nanoparticles have been termed by de Gennes [1] as Janus particles and the two hemispheres of these particles have different chemical character. However, synthesizing these Janus nanostructures has been a challenge. In this paper we study the interaction of Si-doped fullerene as a seed to promote anisotropic growth. We have concentrated on carbon–silicon Janus nanostructures for the following four reasons:

(1) Although both Si and C belong to the same group of elements in the periodic table, their chemistry is very different. For example, oxides of carbon, namely CO and CO$_2$, form gases, while oxides of silicon constitute common sand. Similarly the sp$^2$ and sp$^3$ bonding characteristics of carbon give rise to novel forms starting from the well-known planar graphite and tetrahedral diamond to ‘spherical’ fullerenes and cylindrical nanotubes. On the other hand, silicon prefers only tetrahedral bonding due to its larger core. Both carbon based nanostructures such as fullerenes and nanotubes [2, 3] as well as Si based nanostructures [4–8] have potential technological applications. This has led to recent interest in carbon–silicon composite nanostructures [6, 9, 10] which have great potential for applications in quantum dots, quantum wires, nonlinear electronic components and memory devices.

(2) Bulk silicon carbide is a well-known material with exceptional physical and mechanical properties, including low density, high strength, high thermal conductivity, stability at high temperature, high resistance to shock, low thermal expansion, high refractive index, wide (tunable) band gap and chemical inertness. We can expect that nanoscale carbon–silicon composite structures may exhibit novel properties due to the large surface area and quantum size effect. One of the well-known carbon–silicon nanostructures is $C_{60}$@$Si_{60}$, which was suggested by Harada and co-workers for stabilizing the $Si_{60}$ cage with a $C_{60}$ core. This complex has attracted extensive interest theoretically [11–16] and experimentally [17–24]. Unfortunately, the observation of $C_{60}$@$Si_{60}$ has remained
elusive [17–24]. In contrast, it has been found experimentally that Si60 is unlikely to wet the surface of C60 [22], and it has been theoretically confirmed that C60@Si60 is energetically unstable [7]. Due to the stability of the carbon fullerene cage, it is difficult to grow a silicon nanostructure directly on the C60 surface. In this study we show that fullerene substitutionally doped with Si provides us with an embryo for Si growth, with the Si site acting as the growth seed.

(3) Recently, a great deal of attention has been paid to the shape-controlled synthesis [25–30] of nanostructures with anisotropies that possess certain advantages over their spherical counterparts, as they can be dually functionalized for potential applications. Since C and Si have different properties, it is expected that carbon–silicon Janus nanostructures may have interesting properties.

(4) Silicon nanostructures such as Si10, Si12, M@Siₙ (M = Ti, W, Cr, Mo, Ba, n = 8–20) have been extensively studied both theoretically and experimentally, but one of the challenges for their application is how to assemble these structural units to make functional materials [8–10]. Here we show that Si-doped heterofullerene may provide a novel pathway for assembling carbon–silicon Janus nanostructures.

2. Computational methods

The calculations were carried out within the framework of density functional theory (DFT) and the generalized gradient approximation (GGA) for exchange–correlation energy for which the Perdew–Wang 91 (PW91) functional form [31] was used. In order to optimize the geometry effectively, a plane-wave basis set is adopted with the projector-augmented-wave (PAW) method, originally developed byBlochl [32] and recently adapted by Kresse and Joubert, as implemented in the Vienna ab initio simulation package (VASP) [33]. The particular advantage of the PAW method over the ultrasoft pseudopotentials is that the pseudization of the augmentation charge can be avoided. The structure optimization is symmetry unrestricted and carried out using the conjugate gradient algorithm. The convergence and force were set to 0.0005 eV Å⁻¹ and 0.01 eV Å⁻¹, respectively. The kinetic energy cutoff used in our calculations is 300 eV. A supercell approach was used, where the clusters under investigation were surrounded by 15 Å of vacuum space along the x, y, and z directions, which is large enough to avoid the interactions with images. Due to the large supercell, the Γ point was used to represent the Brillouin zone. The accuracy of calculation for fullerene and Si based nanostructures has been extensively tested in our previous studies [6–8, 10]. For example, although GGA usually underestimates the HOMO–LUMO gap, our calculated value for C60 is 1.61 eV, which is comparable to the experimental value of 1.68 eV. GGA + U is better, especially for systems containing d- or f-electrons; for our studied systems C59Si + Si-clusters, the corrections of GGA + U would be small.

3. Results and discussion

We begin with the structure of C59Si where a C atom in the C60 fullerene has been substituted with a Si atom. This heterofullerene was synthesized in 1996 using the laser vaporization cluster beam technique. Our optimization of the C59Si geometry yielded a structure with C2 symmetry (figure 1), the average Si–C bond distance of 1.822 Å. The energy cost to replace one C atom with Si, calculated by taking the total energy difference, namely ΔE = E(C59Si) + E(C) - E(C60) - E(Si), is 5.080 eV. These are in agreement with previous calculations and experiments [34–37].

For C59Si the energy gap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) is 1.17 eV, lower than that (1.61 eV) of C60. Furthermore, the HOMO and LUMO were mainly concentrated on the Si site and its neighboring C sites. Because of the difference in electronegativity between C and Si, charge is transferred from Si to C allowing C59Si to have a dipole moment of 1.42 D. From the geometry, charge distribution and frontier orbitals, we can clearly see the anisotropy induced by Si doping. Thus, the Si site can thus be used as a ‘seed’ for anisotropic growth when ligand molecules are attached. This is demonstrated in the following by using Si10, Si12W, Si16Ti and Si20Ba as ligands.

3.1. Si10 interacting with C59Si

Silicon clusters in the size ranges of 3–12 were initially synthesized in a supersonic beam in 1987 [38]. The ultraviolet photoelectron spectra (UPS) data suggested that among these clusters, those containing 4, 6, 7 and 10 Si atoms are magic. Since then, many experimental and theoretical studies have concentrated on the Si10 cluster [39–41] which has a tetracapped trigonal prism structure with C3v symmetry. The relaxed structure is plotted in figure 2, together with its HOMO and LUMO images. We see that the HOMO is mainly concentrated on the bottom triangular face, while the LUMO originates from both the bottom and the top triangular face. The HOMO–LUMO gap is 2.09 eV.

Based on the distribution of orbitals in C59Si, we can expect that the Si site should be the most likely absorption site [42]. For Si10 we consider two possible configurations interacting with C59Si: configuration I where Si10 binds to the Si atom in C59Si through its bottom triangular face or configuration II where Si10 binds through its top triangular face. For configuration I, the optimized geometry and electronic structure are shown in figure 3. The interaction
energy, defined as, $\Delta E = -[E(C_{59}Si-Si_{10}) - E(C_{59}Si) - E(Si_{10})]$, is calculated to be 1.15 eV with the linking Si–Si distance of 2.441 Å and the HOMO–LUMO gap of 0.65 eV. For configuration II, the adsorption energy is 0.86 eV, weaker than that in configuration I, while the bond length of the Si–Si is enlarged to 2.475 Å. In figure 3, we also see that both the HOMO and LUMO orbitals of the C_{59}Si–Si_{10} complex are mainly concentrated on the Si part and the atoms near the Si site. Therefore the introduction of Si_{10} into C_{59}Si has enhanced the anisotropy of the structure. Furthermore, because of the difference in electronegativity between C and Si, charge is likely to transfer from Si to C. In fact Si in C_{59}Si carries a charge of $+0.852$, which is increased to $+0.996$ when interacting with Si_{10}. The charges are redistributed in the system, in particular the linking Si site in the Si_{10} unit carries a charge of $-0.236$, and the three C atoms linking with Si in C_{59}Si become more negatively charged, as shown in figure 4. The charge redistribution in C_{59}Si–Si_{10} results in a large dipole moment of 7.11 D. The frontier orbital distributions clearly indicate that the Si sites and their neighboring C sites are much more active than other sites.

3.2. W@Si_{12} interacting with C_{59}Si

W@Si_{12} clusters were synthesized using an ion trap by reacting the transition metal ion, W^{+} with silane (SiH_{4}). The W@Si_{12} cluster was found experimentally to be magic and stable, which was also confirmed by ab initio calculation. In W@Si_{12}, the W atom is encapsulated inside a Si_{12} cage, having a hexagonal-shaped cage with D_{6h} symmetry. The stability of this structure is due to both the electronic and the geometrical shell closures [8, 43].

We began the calculations by optimization on the W@Si_{12} cage, plotted in figure 5 (the bonds connecting the Si atom with W in the center are omitted for clarity). The relaxed structure is a hexagonal cage as expected. There are two types of Si–Si bond lengths: 2.381 Å on the top and bottom faces and 2.402 Å in the lateral edges. The bond length between Si and W is 2.667 Å. Its HOMO–LUMO gap W@Si_{12} is calculated to be 1.37 eV. These data agree well with the results in [41]. Note that the HOMO of W@Si_{12} mainly reside on two hexagon rings, and the LUMO arises mainly from $dz^2-W$.

To optimize the geometry of Si_{12}W interacting with C_{59}Si, we began with an initial geometry where the Si site in C_{59}Si is connected to the hexagonal ring of Si_{12}W. The optimized structure is shown in figure 6, together with the HOMO and LUMO plots. We find that the W@Si_{12} is slightly distorted, and tilted at an angle of 55° with respect to the initial orientation to make the two linking Si atoms four-fold coordinated as required by sp^{3} hybridization. The bond length between the two linking Si atoms is 2.434 Å. The corresponding adsorption energy is 1.06 eV and the HOMO–LUMO gap is reduced to 0.43 eV. This is much lower than those of the individual units (1.17 eV for C_{59}Si, and 1.37 eV for W@Si_{12}). Unlike C_{59}Si–Si_{10}, there is only a small amount of charge transfer between these two units (see figure 7). However, the frontier orbitals clearly show the anisotropy of the C_{59}Si–Si_{12}W complex, where the Si and the neighboring C sites are much more reactive. What is more important is that this complex has a large dipole moment of 8.04 D. This character may be very useful for materials self-assembly under an applied electric field.
3.3. Ti@Si\textsubscript{16} interacting with C\textsubscript{59}Si

The Ti@Si\textsubscript{16} cage has been synthesized by a combination of a laser vaporization method and a flow-tube reactor \cite{44}. Extensive investigations \cite{45, 46} have confirmed that this cluster possesses a truncated tetrahedral structure, called the Frank–Kasper (FK) polyhedron, with T\textsubscript{d} symmetry. The optimized structure is shown in figure 8 along with its HOMO and LUMO orbitals. Among the 28 triangles, there are four truncated triangles with bond lengths of 2.437 Å, with the remaining 24 being of one type with bond lengths of about 2.658 Å. We point out that an isolated Ti atom has an electronic configuration of 3d\textsuperscript{2}4s\textsuperscript{2} and a magnetic moment of 2μ\textsubscript{B}, which is quenched when encapsulated in Si\textsubscript{16} cage. Is it possible to recover this magnetic moment when this cluster is attached to C\textsubscript{59}Si? Actually if the attachment can expand the Si cage, the increased Si–W bond length will lead to less orbital hybridization between W and its neighboring Si atoms. Hence, the magnetic moment of Ti can be retained.

Motivated by this expectation, we studied the interaction between C\textsubscript{59}Si and Si\textsubscript{16}Ti. As discussed above, Si\textsubscript{16}Ti has two inequivalent types of triangles which were respectively attached to the Si site in C\textsubscript{59}Si in optimizing the geometry. The preferred structure is shown in figure 9. The two cages are connected with each other through a single Si–Si bond (2.496 Å). The corresponding binding energy and HOMO–LUMO gaps are respectively 0.93 eV, and 0.56 eV. The Si site in C\textsubscript{59}Si interacts with the Si in Ti@Si\textsubscript{16} along the on-top direction. Because of this interaction, the structure of the Ti@Si\textsubscript{16} cage is distorted and the symmetry is changed to C\textsubscript{3v}. The cage is only slightly expanded, but not enough to significantly reduce orbital hybridization. Hence, the Ti@Si\textsubscript{16} complex maintains its non-magnetic character, but displays a huge dipole moment of 9.18 D. The attachment also changes the orbital distributions. In contrast to figure 8(c) where the LUMO of Si\textsubscript{12}W mainly derives from Si sites on the cage surface and W site in the center, in the Ti@Si\textsubscript{16} complex the contribution from the Si cage surface is reduced (see figure 9(c)). The top part of Si cage and the Si site in C\textsubscript{59}Si together with its neighboring C sites become reactive.

3.4. Ba@Si\textsubscript{20} interacting with C\textsubscript{59}Si

The structure of Si\textsubscript{20} is of special interest since the smallest fullerene cage is C\textsubscript{20}. However, Si\textsubscript{20} consists of two Si\textsubscript{10} forming a compact structure. The cage structure can be stabilized by encapsulating one Ba atom \cite{5}, as shown in figure 10(a). The bond length of Si–Si is 2.357 Å and the distance of the center Ba atom to Si is 3.301 Å. The HOMO–LUMO gap of the cage is 0.58 eV. Different from the case of Si\textsubscript{16}Ti, the HOMO and LUMO arise mainly from Si atoms but not from the metal atom (see figures 10(b) and (c)). Due to the cage’s highest symmetry, every atom is equivalent. Thus, to optimize the geometry we initially put the pentagonal
Table 1. Bond lengths between linking Si atoms, binding energy, HOMO–LUMO gap, and dipole moment of the various complexes studied.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Linking bond length (Å)</th>
<th>Binding energy (eV)</th>
<th>HOMO–LUMO gap (eV)</th>
<th>Dipole moment (D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C59Si–Si10</td>
<td>2.441</td>
<td>1.15</td>
<td>0.65</td>
<td>7.11</td>
</tr>
<tr>
<td>C59Si–W@Si12</td>
<td>2.434</td>
<td>1.06</td>
<td>0.44</td>
<td>8.04</td>
</tr>
<tr>
<td>C59Si–Ti@Si16</td>
<td>2.496</td>
<td>0.93</td>
<td>0.56</td>
<td>9.18</td>
</tr>
<tr>
<td>C59Si–Ba@Si20</td>
<td>2.354</td>
<td>1.89</td>
<td>0.58</td>
<td>3.96</td>
</tr>
</tbody>
</table>

Figure 10. (a) Geometry, (b) HOMO and (c) LUMO of the Ba@Si20 cluster.

face of the Ba@Si20 above the Si top site in C59Si. After optimization, the Ba@Si20 cage prefers to connect with the C59Si heterofullerene through a single Si–Si bond with a bond length of 2.354 Å and a tilt angle of about 30° with respect to the initial orientation, see figure 11. The absorption energy of this system is 1.88 eV, which is larger than the others. The reason is the following: in Ba@Si20 the Si sites are negatively charged due to the charge transfer from Ba [5], while the Si site in C59Si is positively charged, which results in a large binding energy and a small linking distance when these two units bind together. Its HOMO–LUMO gap is 0.58 eV, and the dipole moment is found to be 3.96 D. It is interesting to note that the interactions between these two units change the orbital distributions. For example, the HOMO in the complex structure is mainly concentrated on the Ba@Si20 cage, while the LUMO is predominantly on C sites nearest to the Si site in C59Si. This suggests that if the LUMO of some species is close to the HOMO of C59Si–Si20Ba in energy, then these species would preferably attach to the Si cage. On the other hand, if the HOMO of some species is close to the LUMO of C59Si–Si20Ba in energy, then these species will preferably attach to the C cage. Therefore, selective functionalization can be realized in this system.

4. Conclusions

The similarities and differences between Si and C, as well as their potential applications, make studies of the Si–C Janus structures very attractive. In this paper, we have systematically studied the interactions between Si-doped fullerene C59Si and experimentally synthesized and well studied Si clusters including Si10, Si12W, Si16Ti, and Si20Ba. In table 1 we have summarized the main results of our work and arrived at the following conclusions:

1. The Si atom in the heterofullerene provided the attachment site for the ligands during the anisotropic growth of Janus nanostructures.

2. The Si site in C59Si is more reactive than the C atoms.

3. When a Si cluster is attached to C59Si, it preferentially interacts with the Si atom of the ligand cluster along the on-top direction.

4. Si–C Janus nanostructures thus formed exhibit enhanced anisotropic properties. They display large dipole moments and hence can form self-assembled materials under the influence of an external electric field.

5. The attachment of Si clusters to C59Si provides a simple pathway to fabricate silicon–carbon Janus anisotropic nanostructures.

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