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Structures and coalescence behavior of size-selected silicon nanoclusters studied by surface-plasmon-polariton enhanced Raman spectroscopy

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The structures and coalescence behavior of size-selected, matrix-isolated silicon clusters have been studied using surface-plasmon-polariton (SPP) enhanced Raman spectroscopy. The cluster ions were produced in a laser vaporization source, mass selected then deposited into a co-condensed matrix of Ar, Kr or N_2 on a liquid He cooled substrate. Raman spectra from monodisperse samples of the smaller clusters studied, Si₄, Si₆ and Si₇, show sharp, well-resolved, vibrations which are in good agreement with predictions based on ab initio calculations. From these comparisons we confirm that Si_4 is a planar rhombus, and assign Si_6 as a distorted octahedron and Si_7 as a pentagonal bypyramid. Si₅ depositions down to 5 eV did not reveal a measurable Raman spectrum under our experimental conditions. Evidence for cluster-cluster aggregation (or fragmentation) was observed under some conditions, even for a "magic number" cluster such as Si₆. The spectra of the aggregated small clusters were identical to those observed for directly deposited larger cluster "bands," such as Si_{25-35} . The Raman spectra of the aggregated clusters bear some similarity to those of bulk amorphous silicon. Cluster-deposited thin films were prepared by sublimating the matrix material. Even under these "soft landing" conditions, changes in the Raman spectrum are observed with the thin films showing even greater similarity to amorphous silicon. © 1999 American Institute of Physics. [S0021-9606(99)70521-0]

I. INTRODUCTION

In recent years, metal and semiconductor clusters made up of 2-100 atoms have been the topics of intense activity and interest.¹ Silicon clusters, in particular, have demonstrated many fascinating properties.²⁻⁴ While impressive progress has been made in the understanding of this range of matter between atoms and bulk materials, structural information has been more difficult to obtain. High level calculations⁵⁻⁷ and studies using empirical potentials^{8,9} predict that silicon clusters differ significantly from the bulk crystal lattice and surfaces, i.e., they are extensively reconstructed compared to microcrystal fragments. Up to this point, experimental information about the structures was either indirect¹⁰ or limited to very small clusters of up to four atoms.11-13 This lack of knowledge about structures has limited our understanding of the physical, electronic and chemical properties of these clusters.

Many of the observed properties of silicon clusters have been interpreted as evidence for three dimensional reconstruction,^{2,3} even for clusters containing up to 70 atoms $(\sim 10 \text{ \AA} \text{ diameter})$. Since small particles of silicon of 20–30 Å diameter have been observed to have crystalline bulklike lattices,^{14,15} obtaining information about the structures of small clusters and the transition to bulklike crystalline or amorphous structure would be extremely important in the study of basic material properties and growth processes.

The paucity of experimental structural information is largely due to the difficulty of preparing monodispersed samples of sufficient quantity and density, and to the difficulty of applying traditional spectroscopy techniques to gas phase clusters. The application of mass spectrometry to molecular and ion beam studies of metal and semiconductor clusters has made it possible to experimentally determine size-dependent behavior on an atom-by-atom scale. In addition, a great deal of information about very small clusters, mostly metal dimers and trimers, has been obtained by matrix isolation experiments.¹⁶ And many properties of small particles have been elucidated by studies on substrate supported particles, albeit with wide size distributions.^{17,18}

Combining mass spectrometry with cluster deposition is technically very difficult, but makes new experiments on size-selected clusters possible. The first experiments of this kind were done by Wöste and co-workers, who studied the

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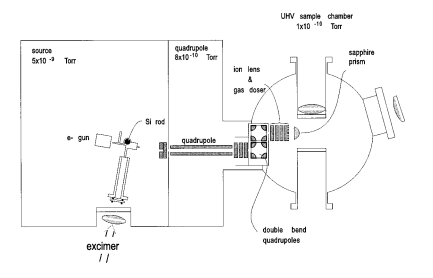


FIG. 1. Schematic diagram of silicon cluster deposition apparatus.

photographic properties of small Ag_n clusters.¹⁹ Subsequently, DiCenzo, Berry and Hartford recorded photoemission spectra for size-selected gold clusters deposited on amorphous carbon.²⁰ Further photoemission studies on size-selected clusters were performed by Eberhardt *et al.*²¹ Harbich, Lindsay and co-workers have done a number of studies on size-selected, matrix-isolated clusters,^{22,23} while Jarrold and co-workers have done several experiments on size-selected and deposited silicon clusters.^{24–26} In a recent paper, Moskovits and co-workers identified the structure of Ag₅ deposited in an argon matrix using Raman spectroscopy.²⁷

The relatively low flux of cluster ions in these experiments results in small quantities of deposited material within a reasonable amount of time. In the experiments reported here, a typical deposition rate for Si_6^+ was 1.5 Å per hour. We have used surface plasmon-polariton (SPP) enhanced Raman spectroscopy to study the clusters,^{28,29} a technique which is capable of detecting molecular species at monolayer surface coverages. Here we report on Raman spectroscopy studies of several small cluster sizes: Si₄, Si₆ and Si₇. As described in an initial communication,³⁰ the sharp, well-resolved vibrational spectra were found to be in good agreement with predicted Raman frequencies and intensities from *ab initio* structure calculations, making it possible to assign the structures of these small clusters.

In addition to the properties of the isolated clusters, we describe some preliminary observations of interactions between silicon clusters. Since most conceivable technological applications of clusters would involve clusters deposited on a support or assembled in some way to form new materials, cluster-substrate and cluster-cluster interactions are important areas of study. Cluster deposition or aggregation has been used to achieve different growth kinetics and hence different material properties.^{31,32} Using Raman spectroscopy, we have examined the aggregation properties of small silicon clusters under two sets of conditions: warming the matrix to allow the clusters to diffuse in the matrix, and sublimating the matrix completely to "soft land" the clusters onto a substrate. We find that small silicon clusters aggregate under these very gentle conditions, even the relatively stable, highly symmetric ones. The resulting Raman spectra of the aggregates are independent of initial cluster size and for both methods of preparation bear some similarities to those of amorphous silicon.

II. EXPERIMENT

A. Production and size-selection of silicon cluster ions

Figure 1 shows a diagram of the experimental apparatus for depositing size-selected clusters. The apparatus is essentially the same as that used for previous studies of sizeselected deposited clusters^{25,26} with modifications to do the in-situ SPP enhanced Raman measurements and increase the stability of the ion beam. A pulsed laser-ablation source produced the silicon cluster ions (see Fig. 2). A Lambda Physik excimer laser (EMG103MSC) operating at 308 nm (xenon chloride) between 100 and 200 Hz ablates silicon from a rotating and translating polycrystalline target rod (99.999%) purity) into a continuous flow ($\sim 25 \text{ cm}^3/\text{s}$) of He buffer gas (99.9999%) at 13–20 Torr. The production of positive cluster ions was enhanced by preionizing the He buffer gas with a 1 keV electron beam (emission current 5 mA) fired into the He gas through a pinhole upstream of the silicon target rod. The injected electrons are also crucial for effective ion transport because the electrons neutralize space-charge effects and reduce the expansion of the ion beam. After a mild ex-

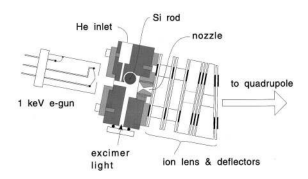


FIG. 2. Laser vaporization source used to produce the silicon clusters. An electron gun at 1 keV produces a He plasma and the clusters are ionized by charge exchange. See text for details.

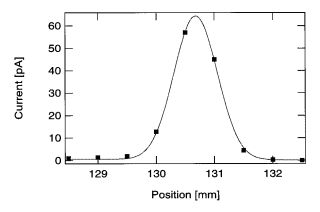


FIG. 3. Current profile measured on a 0.5 mm wire moved perpendicularly to the beam. The continuous line is a fit to the data taking into account the thickness of the wire and assuming a Gaussian beam profile. This yields a diameter for the cluster beam of 0.85 mm full width at half maximum (FWHM).

pansion into a low-vacuum chamber (base pressure 5×10^{-9} Torr, operating pressure 6×10^{-4} Torr), the cluster ions are focused into a low energy ion beam, pass into a differentially pumped chamber and are injected into a quadrupole mass spectrometer in a medium vacuum environment (base pressure 8×10^{-10} Torr, operating pressure 6×10^{-7} Torr). The cluster ions are mass filtered, focused through an aperture, and then undergo a double right-angle bend into the ultrahigh vacuum (UHV) deposition chamber which is cryo- and turbo-pumped to a base pressure of 7×10^{-11} Torr (operating pressure 5×10^{-8} Torr of He). The double right-angle bend deflects the clusters out of the path of any residual effusive flow from the low and medium vacuum chambers that might contaminate the sample during deposition.

Immediately before the deposition substrate, the cluster ions are slowed to their final deposition energy while being focused with a three element zoom lens. Typically, 27 eV was used as the final deposition energy, with some depositions done at 63 eV. Typical cluster ion currents measured at the substrate holder ranged from 120 up to 800 pA for Si_6^+ in a spot varying between 0.8 to 3.5 mm in diameter. Deposition times of 1–3 hours yielded the equivalent of 0.5–1 monolayers of clusters distributed in the matrix. In early experiments, Rutherford backscattering (RBS) was used to calibrate cluster deposition rates and measure the spot size. In later experiments, the spot could be directly observed by imaging the shift in the surface plasmon polariton resonance angle as well as measuring the current profile directly, as shown in Fig. 3.

B. Deposition and matrix isolation

The matrix gas to be co-condensed with the clusters was directed through an element which served as both the final ion optic and a gas diffuser. The substrate was a hemicylindrical sapphire prism, cooled by a continuous flow liquid He cryostat to between 8 and 15 K. The prism was positioned 5 mm in front of the final ion optic/gas diffuser as a compromise between maintaining a small cluster beam spot size and ensuring a uniform thickness of the matrix. First, a buffer layer of 50 Å was deposited to ensure that no cluster came into contact with the silver film, then the cluster-containing matrix was built to a thickness of typically 500–1500 Å. After the clusters were deposited, a layer of pure matrix material was deposited to form a "cap" several microns thick, providing the necessary optical conditions for the plasmon polariton, and isolating the clusters from subsequent contamination. Optical interferometry was used to calibrate the deposition rate of the matrix, while the position of the SPP dip was used to calibrate its index of refraction.

The cluster concentration in the matrix was controlled by monitoring the cluster current and regulating the deposition rate to produce the desired concentration. The cluster ions were neutralized in the matrix by electrons generated from a shielded hot filament (+4 eV to ground, 3, μ A emission current). Solid N₂ was used as the matrix material for most of the studies. Surface chemistry and gas phase cluster reactivity experiments have shown that N₂ is unreactive with bulk silicon surfaces and silicon clusters at low temperature.³³ Matrices of Kr with 1% N₂, or Ar with 1% SF₆, were used for some of the experiments. Spectra recorded in the different matrices were not significantly different.

C. SPP enhanced Raman spectroscopy of matrixisolated silicon clusters

SPP enhanced Raman spectroscopy^{19,20,34} was used to characterize the vibrational properties of the clusters. The sample geometry and Raman optics configuration are shown in Fig. 4. The excitation laser was an argon ion laser operating at between 514.5 and 457.94 nm (10-200 mW, *p*-polarized) with spatial and plasma line filters. The beam was focused by a cylindrical lens to $100 \,\mu\text{m} \times 2 \,\text{mm}$. This leads to $\sim 10^{11}$ clusters in the laser spot. A re-entrant quartz viewport provided access for the F2 collection optics at \sim 55 deg from normal on the Ag side of the prism. The scattered light collimated by the collection lens was focused by an *f*-matching lens into a triple grating spectrometer (SPEX Triplemate) with an 1800 gr/mm grating final stage, providing approximately 1 cm⁻¹/pixel dispersion onto a backthinned, LN₂ cooled, Tektronix 512×400 charge coupled device (CCD) detector (Princeton Instruments). The resolution of the spectrometer was approximately 3 cm^{-1} . Exposure times were typically 1200 sec, with multiple spectra averaged together. A 3×3 median filter was used to remove cosmic rays from unbinned images. The images were then summed along the nondispersive CCD columns to obtain a 2-D spectrum of intensity versus Raman energy loss. Coarse alignment of the substrate and optics was done with two intersecting laser beams through the chamber. The Raman signal from the N₂ matrix or N₂ or SF₆ molecular species, seeded in the Kr or Ar matrices, respectively, was used to optimize the alignment and as an *in situ* calibration of the Raman cross sections of the clusters. The unenhanced Raman lines from the Al_2O_3 substrate were attenuated by the Ag film (by $\sim 1/200$), but were still detected and used to calibrate the vibrational energy scale.³⁵

In our experiments, the surface-plasmon-polariton (SPP) was excited using the attenuated total reflection (ATR), or

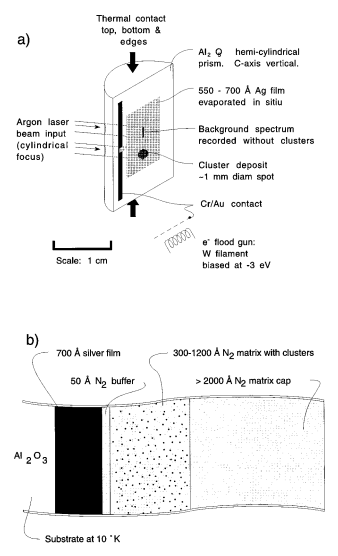


FIG. 4. Schematic of substrate configuration, showing prism, Ag, cluster/ matrix and capping layers, and incident and collected light angles.

Kretschmann configuration. In this arrangement, SPP modes of a metal film deposited on a dielectric prism are excited by adjusting the incident angle of the laser beam. This matches the laser wave vector to that of the nonradiative branch of the dispersion curve of the surface-plasmon-polariton at the metal/prism interface. Approximately 550 Å of Ag was deposited in-situ on the sapphire prism, with the clustercontaining matrix deposited on the Ag film. (Later this was increased to 700-750 Å in order to further attenuate the sapphire substrate Raman lines.) The lower refractive index of the matrix relative to the prism forms an asymmetric slab configuration that causes the nonradiative SPP field to decay exponentially from the Ag/matrix interface. The resonant nature of the SPP excitation causes the field intensity at the Ag/matrix interface to be a factor of $\sim 50-100$ times the incident laser intensity. This enhancement and exponential localization of the incident optical field to a volume (by definition) comparable to the sample size makes possible the large and selective enhancement of the Raman signal of our samples.

The enhanced field strength decreases exponentially with distance from the Ag/matrix interface with a calculated 1/*e*

distance of \sim 700 Å. The calculated enhancement was found to depend critically on the thicknesses and optical constants of the matrix and Ag film. We found that the details of preparing the Ag film produced observable differences in the resonance condition and enhancement. The largest SPP enhancement and narrowest dip (see below) were found for Ag films deposited at 200 K and annealed for several hours at room temperature.³⁸

Experimentally, we can determine when the conditions for SPP excitation are met by observing the attenuated total reflection of the laser beam from the Ag film, a small temperature rise of the substrate, and a large increase in the Raman signal of the matrix materials. Figure 5(a) is an image of the laser light reflected from the Ag film when the N₂ matrix on the film contained 0.8% Si₄. For this image, the incident laser beam was expanded to a diameter of 3 cm so that the entire length of the sapphire prism could be illuminated at once. The vertical axis of the image is the spatial position along the vertical axis of the sapphire prism. The horizontal axis of the image is the angle at which the laser light was reflected from the Ag film. The dark line in the image is the angle at which the plasmon polariton is resonantly excited in the Ag film. Over most of the prism length, the resonant condition is at a constant angle (49.5 deg). In the region where the clusters were deposited in the matrix, however, there is a large shift in the resonance angle, indicating a change in the dielectric constant of the matrix, resulting in a visible dip in Fig. 5. Figure 5(b) shows line sections taken from the image which show the angle, width and depth of the resonance condition both on and off the spot containing the clusters. The line sections show the broadening and reduction in peak intensity of the resonance produced by the absorption from the Si clusters. Most importantly, the resonance condition results in an increase in the observed Raman intensities. Figure 5(c) shows the 2328 cm⁻¹ N₂ matrix³⁶ Raman signal versus incidence angle of the laser beam for a pure matrix of N2 and a matrix with 0.2% concentration by volume Si₄. The solid line through the experimental points is from calculations of the integrated field intensity using the Fresnel equations for a multilayer structure with complex dielectric constants. The optical constants of Johnson and Christy were used for Ag.³⁷ The calculated angle of 51 deg and field enhancement of ~ 90 for the pure N_2 matrix matches the experimental results using an index of refraction of 1.34 for the N₂. Similarly, a complex index of 1.2+i0.003 for the cluster-containing matrix was found to match the experimentally determined enhancement for this set of deposition conditions. From the cluster concentration and effective medium theory, we can, in principle, extract a cross section for the Raman process for the clusters. In practice, however, it turns out that we would have to take into account many parameters in this estimation, such as the spatial distribution of the clusters, the spatial variation in the intensity of the laser spot, and the shift in the resonance with the concentration of clusters.

Since the clusters were deposited in a small spot $\sim 1-3$ mm in diameter, it was possible to take a background spectrum a few mm away under virtually identical conditions. This background can then be subtracted from the spec-

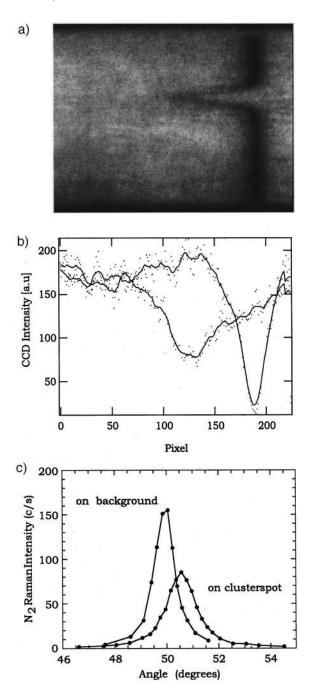


FIG. 5. SPP enhancement: (a) Laser light reflected from the silver film. At the SPP resonance angle the light is not reflected and a dark line appears. The observed dip is the signature of the presence of clusters in the matrix, which slightly shifts the SPP resonance angle. (b) Line section taken from the image above. (c) N_2 signal versus angle for both neat matrix and matrix with clusters. The solid lines through the data points were calculated with different matrix indices of refraction, including a small imaginary part for cluster containing matrix.

trum taken at the cluster position. In addition to the Raman lines from the Al_2O_3 substrate, the background consisted of broadband surface plasmon-polariton emission, stray light, fluorescence and possibly Raman scattering from contaminants on the Ag film or in the matrix. The latter were typically derived from hydrocarbons (as identified by their characteristic Raman bands) and were minimized by baking the vacuum system and depositing a fresh Ag film prior to each cluster deposition run. In later experiments, the hydrocarbon contamination and the accompanying Raman and luminescence background were virtually eliminated by a thorough cleaning of the chambers, the cryotrapping of the diffusion pumps on the low and medium vacuum chambers, and the installation of a gate valve which allowed the low vacuum chamber to be isolated from the other chambers except during cluster deposition.

III. RESULTS AND DISCUSSION

A. Raman spectra of small clusters

Figures 6(a)-6(c) show SPP enhanced Raman spectra from 180 to 560 cm^{-1} for Si₄, Si₆ and Si₇, respectively. The predicted geometries, frequencies and Raman intensities from ab initio calculations are shown in the figure for comparison. For each cluster size, the experimental spectrum shown is an average of several exposures and cluster depositions. Typical conditions for the spectra shown were $\sim 0.1\%$ cluster concentration by volume and an N₂ Raman intensity of 200-300 cps; aggregation begins to occur for concentrations higher than 0.1%. The higher Raman intensities measured for Si₄, compared to Si₆ and Si₇, may suggest an additional enhancement arising from electronic resonance effects. Numerous attempts to measure the Raman signature of Si₅ and Si₁₀ were unsuccessful, presumably because of low Raman cross sections and a limited signal-to-noise ratio. A background spectrum and slowly varying component have been subtracted to increase the visibility of the sharp cluster lines. No lines attributed to the clusters were observed at higher frequencies and the experimental conditions did not permit measurements at significantly lower frequencies.

For Si₄, Si₆ and Si₇ the observed lines are all less than $600-650 \text{ cm}^{-1}$. The Raman spectra of bulk crystalline silicon consist of a single line around 520 cm^{-1} , while the spectrum for amorphous silicon is broad and reflects the density of states.³⁹ The multiple, sharp lines observed for the small clusters are indicative of their molecular nature and demonstrate that these clusters cannot be understood in terms of bulk silicon properties.

B. Effect on Raman intensities from proximity to metal surface

The intensities of the scattered Raman signal are significantly affected by the clusters' proximity to the metal film. These "surface selection rules" can result in large modedependent differences in the observed intensities compared to the standard 90 deg scattering geometry. The intensity versus angle of a radiating dipole close to the surface of a good conductor can be qualitatively understood in terms of image dipoles [Fig. 7(a)]. For dipoles oriented parallel to the surface of the metal, the radiation from the image is out of phase with that of the real dipole, resulting in little or no observed radiation. Alternately, for a dipole oriented perpendicular to the Ag surface, the fields from the real and image dipoles add constructively, although the radiation pattern of the dipoles results in a strong angular dependence of the observed radiation.

For the case of a molecule directly on a metal/vacuum interface, intensities and angular dependencies have been ex-

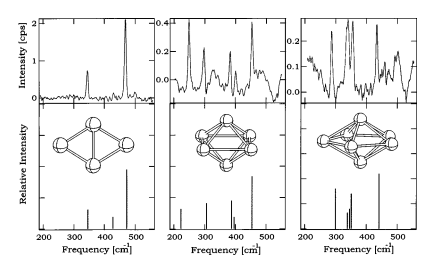


FIG. 6. Measured spectra of Si₄, Si₆, and Si₇, predicted structures, and predicted spectra.

plicitly calculated by several researchers, including Greenler and Slager,⁴⁰ Moskovits⁴¹ and Campion.⁴² With some modifications, we can use these methods to compare Raman intensities for a standard 90 deg scattering geometry to the case of clusters distributed in a matrix at a finite distance from an Ag film. The geometry for calculating the radiated intensity from a dipole *P*, in a matrix of index n_1 , a distance *z* above an Ag surface, is shown schematically in Fig. 7(b). The calculation relies on finding the combined response of the wave directly radiated from the dipole and the wave reflected from the metal surface. The dipole radiation pattern and the polarization and angle-dependent phase shift of the reflected wave give rise to variations in intensity for different angles of reflection and dipole orientations. For dipoles radiating visible wavelength light above an Ag surface, the greatest intensity

> a) f = 0f = 0

FIG. 7. Surface selection rules: (a) Dipoles above a metal surface. (b) Geometry for calculating the radiated intensity from a dipole near a metal surface (see text).

comes from z oriented dipoles, with the maximum integrated intensity coming from around 60 deg from normal.^{41–43}

Since these effects have been worked out in detail by others, we limit our discussion to the consequences of the clusters being distributed in a matrix rather than located exactly at a metal/vacuum interface. This results in an additional phase shift for the reflected wave because of the longer path,

$$\Delta s = \frac{z(1+\cos 2\,\theta)}{\cos\,\theta}.\tag{1}$$

Since the SPP field falls off exponentially over a distance which is less than a wavelength, the radiated intensity will be dominated by those dipoles which are closest to the Ag surface. Compared to the case of the molecule directly on the metal/vacuum interface, this additional phase shift has the effect of slightly increasing the relative intensity of the dipoles which are surface-parallel while slightly decreasing the intensity of those that are surface-normal.

Refraction of the radiated light at the matrix/vacuum interface must also be included in our calculation. Assuming this interface is parallel to the Ag film and the matrix layer is of sufficiently high quality that there is negligible scattering, the external angle ϕ at which the radiated light is observed is related to the internal angle θ by Snell's Law, $\sin \phi$ $=n_1 \sin \theta$. For a matrix of index $n_1=1.25$, the radiated light at an internal angle larger than ~53 deg is totally internally reflected at the matrix/vacuum interface.

With these effects included, we have calculated the relative intensities versus external angle for x, y and z oriented dipoles distributed in the matrix. The intensities were integrated over the z direction with the contributions for the dipoles within a thickness dz weighted by an exponential factor reflecting the decay of the SPP excitation field with distance away from the Ag/matrix interface. Since the SPP electric field is oriented in the z direction, the Raman intensities are related to the α_{xz} , α_{yz} and α_{zz} components of the polarizability tensor. Averaging over all orientations of the clusters yields the intensities in terms of the invariants of the polarizability tensor, $\bar{\alpha}$ and γ^2 , the isotropic and anisotropic parts respectively.^{43,44} Integrating over the range of angles

TABLE I. Vibrational parameters for silicon clusters.

| Cluster | Experimental frequency cm ⁻¹ | Calculated ^a frequency cm ⁻¹ | Absolute deviation cm ⁻¹ | Symmetry assignment | Calculated ^b intensity (90°) | Corrected ^c intensity |
|-----------------|---|--|---|------------------------|---|-------------------------------------|
| Si ₄ | 345 | 346 | 1 | a_g | 1.0 | 3.1 |
| | | 428 | | b_{3g} | 0.6 | 1.9 |
| | 470 | 473 | 3 | a_g | 7.0 | 9.5 |
| Si ₆ | 252 | 223 | 29 | b_{2g} | 1.0 | 3.2 |
| | 300 | 306 | 6 | a_{1g} | 1.7 | 4.1 |
| | 386 | 388 | 2 | b_{1g} | 1.4 | 4.5 |
| | 404 | 396 | 8 | e_g | 0.6 | 1.9 |
| | 458 | 454 | 4 | a_{1g} | 8.2 | 8.4 |
| Si ₇ | 289 | 300 | 11 | e'2 | 1.0 | 3.2 |
| | 340 | 339 | 1 | $e_2^{\tilde{\prime}}$ | 0.4 | 1.3 |
| | 340 | 346 | 6 | $e_1^{\tilde{n}}$ | 0.5 | 1.6 |
| | 358 | 352 | 6 | a_1' | 1.4 | 2.8 |
| | 435 | 441 | 6 | a_1' | 4.3 | 4.4 |

^aQCISD/6-31G^{*} frequencies for Si₄ and Si₆, and MP2/6-31G^{*} frequencies for Si₇, all scaled down uniformly by 5%. See text.

^bRelative intensities at the HF/(7*s*,6*p*,2*d*) level of theory (see text) for a 90 deg scattering geometry assuming no electronic resonance effects.

^cThe calculated intensities corrected for the experimental geometry effects (see text).

subtended by the light collection lens, and using appropriate values of the dielectric constants for the experimental conditions⁴⁵ of the spectra shown in Fig. 6, the calculated intensity is

$$I \sim \bar{\alpha}^2 + 0.5 \gamma^2. \tag{2}$$

Therefore, the intensity observed for each Raman active vibrational mode may be enhanced or suppressed relative to the standard 90 deg scattering geometry where $I \sim \bar{\alpha}^2 + (7/45) \gamma^2$.

C. Theory

The ground state structures of Si3-Si7 have been carefully investigated by ab initio quantum chemical techniques as well as by density functional methods.⁵⁻⁷ In a previous work,³⁰ we reported the vibrational frequencies of Si₄, Si₆, and Si7 at the MP2 level (second-order Møller-Plesset perturbation theory) with the polarized 6-31G* basis set.⁴⁶ We have now refined the the structure and vibrational frequencies of $Si_4 - Si_6$ with the more accurate QCISD (quadratic configuration interaction)⁴⁶ method using the same 6-31G* basis set. The Raman-active QCISD/6-31G* frequencies for Si₄-Si₆, along with the MP2/6-31G* values for Si₇, have been uniformly scaled down by 5% and listed in Table I. The scaling takes into account the systematic overestimations present in the theoretical calculations due to the neglect of anharmonicity effects. The nature of the vibrations can be seen from the associated normal modes which are shown in Figs. 8-10.

The evaluation of Raman intensities requires the use of much larger basis sets. We have evaluated the Raman intensities using Hartree–Fock calculations with a large contracted basis set containing 7s,6p,2d functions on each silicon atom (derived from McLean and Chandler's contracted 6s,5p basis set by addition of a set of diffuse sp functions and two sets of d functions).⁴⁷ The theoretical intensities which correspond to a 90 deg scattering geometry are then

corrected for the experimental geometry effects and the resulting relative intensities are shown along with the experimental spectra in Fig. 6.

The predicted ground state structure of Si₄ is a planar rhombus with D_{2h} symmetry. The perimeter bond length at the QCISD/6-31G* level is 2.31 Å, while the distance along the short diagonal is slightly larger (2.40 Å). The rhombus has three allowed Raman lines at 346(a_g), 428(b_{3g}) and 473(a_g) cm⁻¹. The frequencies and relative intensities of the

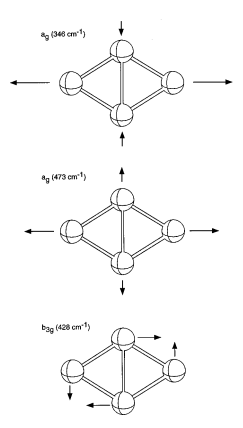


FIG. 8. Normal modes for the three Raman-active vibrations in Si₄ (D_{2h}) .

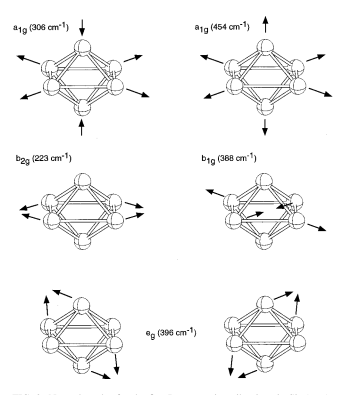


FIG. 9. Normal modes for the five Raman-active vibrations in Si₆ (D_{4h}) .

two a_g modes are in essentially perfect agreement with the experimentally observed modes at 345 and 470 cm^{-1} . Inspection of the normal modes (Fig. 8) shows that the higher frequency a_g mode (473 cm⁻¹) corresponds to the symmetric breathing vibration for the molecule. This has the highest Raman intensity. The lower frequency $a_g \mod (346 \text{ cm}^{-1})$ corresponds to stretching along one axis and compression along a perpendicular axis, and has a lower Raman intensity. The b_{3g} vibration predicted at 428 cm^{-1} has not been observed experimentally, perhaps due to its weak intensity, which makes detection difficult with the existing signal-tonoise. If there is a resonant Raman contribution to the spectrum of Si₄, the two a_g modes may be enhanced further than the b_{3g} mode. The b_{3g} mode has a larger component along the perimeter bonds than the a_g modes. The other three vibrational modes of Si₄ are not allowed in Raman spectroscopy, though some of them have been seen in matrix infrared spectra.48

The ground state structure predicted for Si₆ is a distorted octahedron, although the nature of the distortion is difficult to determine due to the shallow nature of the potential energy surface. At the MP2/6-31G* level of theory, which includes electron correlation effects, Si₆ has a tetragonal bipyramidal structure with D_{4h} symmetry. At other levels of theory (such as Hartree–Fock or density functional theory), additional distortions leading to lower symmetries have been found.^{5–7} However, such distortions, which lead to structures such as a bicapped tetrahedron or a capped trigonal bipyramid, involve shallow degrees of freedom and very little energy. We find that the experimental spectra can be conveniently interpreted in terms of the D_{4h} structure. It can be considered as a strongly compressed octahedron where the two axial atoms are about 2.7 Å apart. Much of the bonding

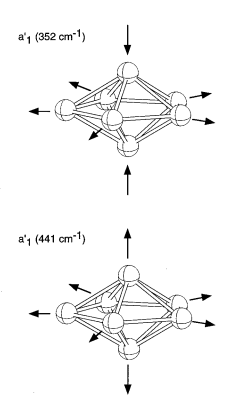


FIG. 10. Normal modes for the two Raman-active a_1' vibrations in Si₇ (D_{5h}) .

strength in this structure comes from the eight equivalent axial-equatorial bonds (2.36 Å). The four "bonds" between the equatorial atoms are about 2.7 Å and do not contribute much to the binding energy. In fact, the distortions involving the weak equatorial bonds are responsible for the shallow energy surface.

At the QCISD/6-31G^{*} level, the D_{4h} form has five allowed Raman lines at 223 (b_{2g}) , 306 (a_{1g}) , 388 (b_{1g}) , 396 (e_g) , and 454 (a_{1g}) cm⁻¹. With the exception of the b_{2g} low frequency mode, all other frequencies are within $10 \,\mathrm{cm}^{-1}$ of the experimental values. The normal modes corresponding to these frequencies are shown in Fig. 9. As in the case of Si_4 , there are two a_{1g} modes corresponding to the symmetric breathing vibration (454 cm⁻¹) and the vibration involving elongation and contraction along two perpendicular directions (306 cm^{-1}) . Again, the symmetric breathing mode has a higher frequency as well as a higher intensity. The b_{2g} mode predicted at 223 cm⁻¹ has a low frequency since it involves only the weak equatorial bonds. The degenerate e_{o} mode has a similar normal mode character, though it has a higher frequency (396 cm^{-1}) since it involves the stronger axial-equatorial bonds. It is interesting to note that the b_{2g} and e_{g} modes result from the splitting of a triply degenerate mode of an octahedron as it distorts to D_{4h} symmetry. The largest deviation from experiment (29 cm^{-1}) for the b_{2g} mode may be related to the difficulty in describing the relatively weak equatorial bonds. Finally, the b_{1g} mode at $388 \,\mathrm{cm}^{-1}$ is in excellent agreement with the corresponding experimental value (386 cm^{-1}) .

Si₇ is predicted to have a pentagonal bipyramidal geom-

etry with D_{5h} symmetry. As in the case of Si₆, it is highly compressed so that the two axial atoms are only 2.51 Å apart (MP2/6-31G*). The equatorial bonds (2.48 Å) are shorter than the corresponding bonds in Si₆. The ten equivalent axial-equatorial bonds (2.46 Å) are somewhat longer than in Si₆. Overall, the D_{5h} form of Si₇ is a compact stable cluster. At the MP2/6-31G* level, it has five allowed Raman lines at 300 (e_2') , 339 (e_2') , 346 (e_1'') , 352 (a_1') , and 441 (a_1') cm⁻¹. Figure 10 shows the normal modes of the two a_1' vibrations. As in the case of Si₄ and Si₆, the symmetric breathing mode is highest in frequency. However, the two modes in this case appear to be roughly equal in intensity. The normal modes for the other degenerate vibrations of Si₇ are more complicated and hence not shown. Experimentally, four lines at 289, 340, 358, and 435 cm^{-1} have been observed, although the width of the 340 cm^{-1} mode suggests the possibility of two overlapping lines. The agreement between theory and experiment is clearly excellent.

Overall, the agreement between the theoretical frequencies of the calculated ground state structures and the experimental Raman spectra is impressive. Among all the Raman active modes predicted for Si₄, Si₆ and Si₇, only the relatively weak b_{3g} mode of Si₄ has not been observed. Sharp vibrational lines have been experimentally seen for all other modes with a mean absolute deviation of only 7 cm⁻¹ between theory and experiment. Consideration of other possible structural isomers^{5–7,30} yields qualitative and quantitative results which are completely inconsistent with the experimental spectra. Thus, our results provide convincing evidence for the structural assignments of small silicon clusters as very different from microcrystalline forms.

D. Fragmentation and/or aggregation of deposited clusters in the matrix

An important issue in experiments on size-selected clusters is that the deposition process does not result in dissociation of individual clusters and/or aggregation into larger particles. Some of the uncertainties in the deposition process include the redistribution of energy in the deposition and neutralization processes and the exact kinetic energy of the cluster ions prior to deposition due to charging of the matrix. One concern was the possibility that the incident cluster ions would sputter or evaporate matrix material during the deposition or deionization process, since this could dramatically effect the concentration of the clusters in the matrix. However, from optical interference measurements, the thickness of the region of the matrix which contains the clusters is within 1% of that of the surrounding matrix.

Although many of the details of the deposition process are not completely understood, there is little question that monodisperse cluster samples can be prepared by sizeselected cluster ion deposition. Harbich, Lindsay and coworkers have reported that up to 75% of Ag_2^+ survive deposition at 30 eV.²² As demonstrated in the cluster–surface collision experiments of St. John and Whetten,⁴⁹ fragmentation is even less likely for silicon clusters deposited at 25 eV, since they have higher dissociation energies and the larger clusters have correspondingly larger heat capacities.³ In our

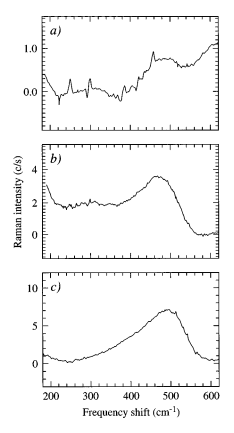


FIG. 11. Cluster aggregation: (a) Raman spectra for Si₆ clusters as deposited and showing some aggregation. (b) Spectrum for Si₆ matrix aggregated at $T_{\text{max}} \sim 28$ K for several hours. (c) Soft-landed Si₆ cluster-deposited film after sublimating the matrix.

experiments, evidence of fragmentation was only observed at the higher deposition energies. For example, Si_{10} clusters show the Raman signature of Si_4 and Si_6 clusters at high deposition energies.

E. Matrix aggregation of clusters and clusterdeposited thin films

While cluster fragmentation can be avoided in the deposition process by using a sufficiently low deposition energy, there is still the possibility of clusters aggregating during the deposition process or once they are in the matrix. We found that it was important to work at low cluster concentrations (around 0.1% by volume) and low temperatures to avoid coalescence. Aggregation of the clusters was easily identified, because it results in Raman spectra that are independent of the deposited cluster size and very similar to spectra from deposition of much larger cluster "bands," such as Si_{25–35}.

Cluster aggregation was found for relatively high cluster concentrations or if the matrix was annealed for several hours at a temperature just below the point where rapid sublimation occurs. Figure 11 shows several spectra from the same Si₆ deposition after raising the temperature, and then recooling to measure the sample under similar conditions. Figure 11(a) shows the as-deposited signal where the prism holder temperature was less than 15 K for both the deposition and measurements. This deposition was done at relatively high concentration, and as discussed below, the broad peak at 470 cm⁻¹ indicates that some cluster aggregation has already occurred. In Fig. 11(b), the spectrum was taken after the temperature was raised to ~28 K for several hours.⁵⁰ The sharp peaks of the isolated Si₆ clusters in Fig. 11(a) are barely visible, replaced with a much broader spectrum due to aggregated clusters. Diffusion of the clusters in the matrix has resulted in the clusters coalescing into larger silicon particles. The Raman cross section for the large silicon particles has increased relative to the isolated clusters because the integrated intensity is much larger. Similar behavior was observed for Si₇: the sharp lines from the isolated clusters evolved into the broad, higher intensity spectrum after warming the matrix to ~29 K overnight. Some of this increase in intensity could be due to part of the material moving closer to the silver surface.

In the spectra showing coalescence of the small clusters, we were unable to find any evidence of intermediate size aggregates. For instance, when Si₄ was deposited at high concentration, only the two sharp lines of the Si₄ clusters and the broad spectrum of the aggregate material were observed, with no evidence of Si₈, Si₁₂ or some other relatively small cluster with sharp Raman spectra. Because of its larger cross section compared to the small clusters, we had much greater sensitivity for detection of the aggregated material. The inability to observe the intermediate sizes would suggest that either those clusters have small Raman scattering cross sections, or the aggregation process is unstable, triggering further coalescence once the process has started. For instance, fusion of two Si_6 clusters to form Si_{12} would release roughly 3 eV of energy.³ A small fraction of this energy could melt a large volume of matrix, further increasing diffusion and subsequent aggregation of clusters.

By warming the substrate and sublimating the cryogenic matrix, we can "soft land" the clusters on to the Ag surface.³¹ Figure 11(c) shows the Raman spectrum of the matrix-aggregated Si₆ clusters deposited as a thin film. This was done by warming the substrate to 60 K to evaporate the matrix and then redepositing a matrix "cap" to achieve similar enhancement conditions as in Fig. 11(a). Raman spectra from samples prepared in this way are reproducible and independent of the initial cluster size, and the degree of aggregation that has already occurred in the matrix. The Raman intensity has increased further, compared to the isolated small clusters. Part of this enhancement is likely due to the material now being closer to the Ag/matrix interface and experiencing a larger excitation field strength.

To try to understand the material formed by the aggregation of the clusters, we can compare the spectra to that of bulk amorphous^{51–53} and microcrystalline⁵⁴ silicon and small silicon particles.⁵⁵ Raman spectra of small silicon particles less than 100 Å diameter are similar to those of amorphous silicon, even when the particles are crystalline as determined by transmission electron microscopy (TEM) or x-ray scattering.^{54,55} This has been interpreted in terms of the relaxation of the k=0 selection rule due to finite size effects⁵⁶ or to arise from field enhancements of an amorphous surface layer on the microcrystals.^{57,58} Although this means that Raman spectroscopy is not sensitive to the degree of crystalline order of small particles, differences in short-range order of the cluster-assembled materials compared to amorphous silicon can be identified. Indeed, evidence for thickness dependent Raman spectra of ultrathin films and clusters of Ge has been observed.¹⁸

The spectra in Figs. 11(b) and 11(c) show some similarities to those of amorphous silicon. The broad peak at $470 \,\mathrm{cm}^{-1}$ in the matrix aggregated material and at $490 \,\mathrm{cm}^{-1}$ in the cluster-deposited material is similar to the transverseoptic-like (TO) $480 \,\mathrm{cm}^{-1}$ peak in amorphous silicon, with the rising edge below $200 \,\mathrm{cm}^{-1}$ corresponding to the transverse-acoustic (TA) band. The longitudinal-acoustic (LA) and -optic (LO) region of the spectrum around $300 \,\mathrm{cm}^{-1}$ is different for the two cluster-derived materials even though the thickness of the cluster-deposited thin film is only ~ 1 cluster diameter. For the matrix-aggregated material, there is more intensity in this region compared to either the cluster-deposited material or bulk amorphous silicon. Given the low temperatures during aggregation and the limited mobility of cluster units compared to individual atoms, it is likely that the material structure is "frozen" into a metastable form.

The matrix aggregated and soft-landed cluster materials appear to have some differences in short-range-order compared to each other and to amorphous silicon. This may be due to finite size effects or the degree of relaxation in the materials. In the work of Lannin and co-workers on Ge clusters and ultrathin films, the TO-like peak shifts and increases in intensity relative to the LA- and LO-like regions as the film thickness increases.¹⁸ Studies of the width and position of the TO-like peak in amorphous silicon have shown a correspondence with the degree of relaxation of the material.^{54,59} The intensity around $300 \,\mathrm{cm}^{-1}$ and the especially broad TO-like peak observed for the matrixaggregated silicon suggest that this material is less relaxed than amorphous silicon prepared by ion implantation or lowtemperature growth. Further characterization of these clusterassembled materials is necessary to elucidate the details of their structures.

IV. CONCLUSIONS

By depositing size-selected silicon clusters into an inert matrix, we have measured Raman spectra of Si₄, Si₆ and Si₇, obtaining the first experimental evidence of the detailed structures of silicon clusters containing more than four atoms. We find excellent agreement with predictions from ab initio theory, confirming that these small clusters have compact, close-packed structures which are completely different from bulk microcrystal fragments. Warming the matrix to allow diffusion of the clusters, or complete sublimation of the matrix to "soft land" the clusters, results in irreversible changes in the spectra. Even magic number sizes such as Si₆ and Si_7 aggregate under these very gentle conditions. The spectra of the resulting cluster-assembled material differ from those of amorphous silicon prepared by conventional methods, but other methods are required to fully characterize the material.

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