Atomic-Scale Patterning of Arsenic in Silicon by Scanning Tunneling Microscopy

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ABSTRACT: Over the past two decades, prototype devices for future classical and quantum computing technologies have been fabricated by using scanning tunneling microscopy and hydrogen resist lithography to position phosphorus atoms in silicon with atomic-scale precision. Despite these successes, phosphine remains the only donor precursor molecule to have been demonstrated as compatible with the hydrogen resist lithography technique. The potential benefits of atomic-scale placement of alternative dopant species have, until now, remained unexplored. In this work, we demonstrate the successful fabrication of atomic-scale structures of arsenic-in-silicon. Using a scanning tunneling microscope tip, we pattern a monolayer hydrogen mask to selectively place arsenic atoms on the Si(001) surface using arsine as the precursor molecule. We fully elucidate the surface chemistry and reaction pathways of arsine on Si(001), revealing significant differences to phosphine. We explain how these differences result in enhanced surface immobilization and in-plane confinement of arsenic compared to phosphorus, and a dose-rate independent arsenic saturation density of 0.24 ± 0.04 monolayers. We demonstrate the successful encapsulation of arsenic delta-layers using silicon molecular beam epitaxy, and find electrical characteristics that are competitive with equivalent structures fabricated with phosphorus. Arsenic delta-layers are also found to offer confinement as good as similarly prepared phosphorus layers, while still retaining >80% carrier activation and sheet resistances of <2 kΩ/square. These excellent characteristics of arsenic represent opportunities to enhance existing capabilities of atomic-scale fabrication of dopant structures in silicon, and may be important for three-dimensional devices, where vertical control of the position of device components is critical.

KEYWORDS: scanning tunneling microscopy, density functional theory, atomic fabrication, silicon (001), arsenic, arsine, dopant

Challenges associated with the continued miniaturization of electronic devices toward the atomic-scale limit,¹ and proposals for atomic-scale device architectures, e.g., to enable quantum information processing,² have stimulated renewed interest in the investigation of dopant species in silicon. Of particular interest are methods for controlling the placement of dopant atoms in silicon with atomic-scale precision,³⁴ and understanding the electronic⁵−⁹ and optical¹⁰−¹² properties of single-atom and few-atom clusters of dopants. The atomic-scale control of dopant atoms in silicon using scanning tunneling microscopy (STM) has to date focused on phosphorus donors, which can be introduced into the silicon matrix using the precursor gas, phosphine (PH₃). The chemical reaction of phosphine with the silicon (001) surface has been extensively researched and is now understood in exquisite detail.¹³ This chemical process can be spatially controlled at the atomic scale via the use of STM hydrogen-desorption lithography¹⁴ to produce atomically defined patterns of phosphorus donors in silicon.¹⁵ This has led to numerous exciting advances in our understanding of silicon-based electronic devices at the atomic scale,¹⁶−¹⁹ and is a promising avenue for the use of individual donor spins as qubits in a solid state quantum computer.²⁰−²²

Arsenic donors in silicon are particularly interesting, since they have a lower diffusivity and a higher solid solubility in bulk silicon than phosphorus,²³ as well as a higher ionization...
energy in silicon (53.76 meV, compared to 45.59 meV for phosphorus), a larger atomic radius ($r_{As} = 115$ pm, $r_{P} = 100$ pm), larger atomic spin–orbit interaction ($Z_{As} = 33$, $Z_{P} = 15$), and a higher nuclear spin value ($I_{As} = 3/2$, $I_{P} = 1/2$) than phosphorus. These differing properties present opportunities for atomic-scale device designs with advanced functionality, including quantum computation schemes based on silicon photonic crystal cavities, which would exploit the larger spin–orbit interaction of arsenic, and schemes employing qudits (generalized $d$-dimensional quantum information units) where the 4-state Zeeman splitting of the arsenic 3/2 nuclear spin could be utilized as a $d = 4$ qudit. In this latter case, while arsenic nuclear spins can in principle also be operated as 2-state qubits with some added complexities, accessing the higher dimensionality provided by the 3/2 spin could offer advantages over qubit based quantum computation, including simplifications in physical implementations of quantum gate structures, and greater efficiency and breadth of quantum simulations. Furthermore, the ability to position multiple dopant species in silicon with atomic-scale precision should allow independent addressing of each donor species by exploiting the different orbital excitation energies, and could thus enable principles of device operation such as optically driven silicon-based quantum gates. 

Kipp et al. have studied the adsorption of arsine on Si(001) using STM and X-ray photoemission spectroscopy and suggest that AsH$_3$, adsorbed at room temperature, decomposes into AsH + 2H and saturates at an As concentration of ~20% of a silicon monolayer (ML). Several other reports on the surface chemistry and growth kinetics of arsenic layers on Si(001) from a gas phase arsine precursor have been primarily concerned with rapid growth rates and elevated growth temperatures (>600 °C), which are not compatible with the thermal requirements of STM hydrogen-desorption lithography.
lithography. In general, however, these reports have established that arsenic gas phase dosing of Si(001) is self-limiting at an arsenic coverage of ∼0.25−1 ML, dependent on the growth temperature, and therefore on the degree of H₂ desorption. Previous density functional theory (DFT) studies have considered likely dissociation configurations of AsH₃, kinetic activation barriers for the removal of the first H atom, and comparative thermodynamic stabilities of dissociated AsH₃ and PII. Here, we present a combined atomic-scale STM and DFT investigation of the adsorption and thermal decomposition of arsenic on Si(001). We establish the full chemical pathway of arsenic decomposition and incorporation of arsenic into the Si(001) surface, and also the spatial control of this reaction via STM hydrogen-desorption lithography. In particular, we demonstrate the following: (1) Individual AsH₃ molecules rapidly and fully dissociate on Si(001) at room temperature. (2) Arsenic atoms from dissociated AsH₃ molecules incorporate substitutionally into the silicon lattice at temperatures below 350 °C, and at all surface coverages. (3) Arsenic adsorption on Si(001), at room temperature, is self-terminating at a saturation coverage of 0.24 ± 0.04 molecules/surface silicon atom, independent of gas exposure rate, providing an arsenic coverage well above the metal–insulator transition, allowing for STM fabrication of metallic interconnects. (4) AsH₃ molecules selectively adsorb and dissociate in a STM depassivated region of the Si(001)-H surface, while leaving the surrounding H-resist intact; subsequent annealing incorporates the arsenic atoms into the top layer of the silicon surface, without disturbing the H-resist. (5) Arsenic delta-doped layers can be grown by confining the arsenic incorporated surface below an epitaxial silicon capping layer, grown via temperature-programmed, low temperature epitaxy. (6) Ohmic contacts can be made to the buried, and electrically activated, arsenic-doped layer by aluminum deposition. Fulfilment of these six criteria satisfies the necessary requirements for successful STM fabrication of atomic-scale, electrically contacted, arsenic in silicon electronic devices. While very recent progress has been made using STM patterning of boron in silicon, the present work represents the development of the capability to pattern a dopant species, other than phosphorus, in silicon, at the atomic scale, using hydrogen-desorption lithography. Moreover, this work represents a first step in the development of a materials toolbox, where atomic-scale structures can be constructed using multiple dopant elements: an important step closer toward the emerging field of atomic-scale fabrication and manufacturing.

RESULTS AND DISCUSSION

Dissociation of Isolated AsH₃ Molecules. To ascertain the atomic-scale behavior of the AsH₃/Si(001) adsorption system, we exposed atomically clean Si(001)2 × 1 to a low dose of arsenic (0.015 L at 5 × 10⁻¹¹ mbar) at room temperature. An example of the surface observed by STM, immediately after dosing, is shown in the filled and empty-state STM images of Figure 1a and b. Of the features observed, several are readily identified as native Si(001) surface defects, such as dimer vacancies (labeled DV in Figure 1a and b) and C-defects (labeled Cd). The new features in these images are attributed to the products of AsH₃ dissociation reacting with the surface. These are labeled “type-1” and “type-2”. The type-1 features are by far the most common, accounting for more than 75% of all AsH₃-associated features detected at these coverages, and these are the principal focus here. The minority type-2 features arise due to steric effects, which we briefly discuss below.

High-resolution, filled- and empty-state images of the type-1 feature are shown in Figure 1c and e, respectively. The type-1 feature is characterized by two bright protrusions on two adjacent dimer rows. These protrusions appear in both filled and empty states, but differ in their relative brightness. Across the dimer rows, i.e., along the vertical axis of the images in Figure 1c and e, both protrusions are shifted up from the dimer centers, toward an upper dimer end. The short vertical lines of the overlaid white reticles on the STM images indicate the silicon dimer center positions in the horizontal direction. By reference to these reticles, we see that the upper row protrusion aligns with the third vertical line in from the left, and is thus positioned on top of a single dimer end, while the lower-row protrusion is located between the third and fourth vertical lines, and thus is positioned between two dimer ends. The upper protrusion induces strong dimer pinning, which is apparent in the filled state image (Figure 1c) as the characteristic zigzag appearance along the row.

We assign the type-1 STM feature to the structure shown in Figure 1d, which we call the inter-row end-bridge As + 3H configuration. Gray highlighting in the schematic indicates sites that appear bright in the corresponding filled-state image. These bright sites correspond to the bare Si−Si dimers that surround the adsorbate, and to the two prominent protrusions of the type-1 feature. The two protrusions in the STM images are assigned to the dangling bond end of a Si−Si−H hemihydride dimer on the upper row, and a dimer end-bridge arsenic atom on the lower row (where the remaining two hydrogen atoms are attached to the lower row, directly across from the end-bridge arsenic atom). These two sites are both characterized by a single unpaired electron, i.e., there are half-filled molecular orbitals at these sites that would nominally image brightly in both filled and empty state. The two dimer ends opposite the bridging arsenic atom appear dark in both the filled and empty state image, which is consistent with the electronically saturated, hydrogen-termination at these sites.

Figure 1g shows the proposed reaction pathway giving rise to the inter-row end-bridge As + 3H configuration observed as the type-1 feature. This kinetically preferred pathway is based on DFT calculations of adsorption energies of the various intermediate structures involved and the activation energies for each step along the path. Each successive step leads to a structure of lower energy, providing the thermodynamic driving force toward the final formation of the inter-row end-bridge As + 3H configuration. The first reaction step, labeled (i) in Figure 1g, is the dissociation of a dative adsorbed AsH₃ molecule into an AsH₂ + H species by means of an inter-row hydrogen shift. This reaction transfers one hydrogen atom to the adjacent dimer row, producing a hemihydride dimer. The calculated activation barrier for this reaction step is only 0.23 eV, which corresponds to a reaction that is completed on a time scale of nanoseconds at room temperature. We note that the inter-row hydrogen shift is kinetically preferable to the alternative, (and perhaps more intuitive) on-dimer and interdimer hydrogen shifts (not shown in Figure 1g), which have calculated activation energies of 0.53 and 0.29 eV, respectively. The fact that the inter-row H-shift is preferred, correlates well with the fact that the type-1 STM feature, contains a hemihydride on an adjacent row, and that a majority of adsorbed AsH₃ molecules dissociate by this route.
The second reaction step, labeled (ii) in Figure 1g, is also a hydrogen-shift reaction, which preferentially occurs in the on-dimer direction to produce a structure that we refer to as the dimer-end AsH + 2H structure. The calculated activation energy for this reaction is 0.63 eV, corresponding to a reaction time scale of milliseconds at room temperature. The alternative reaction, in which the hydrogen atom shifts in the interdimer direction (not shown in Figure 1g), has a slightly larger activation barrier of 0.72 eV, and is thus kinetically disfavored.

The dimer-end AsH + 2H configuration containing a single-valent AsH fragment is a highly transient intermediate corresponding to an extremely shallow minimum on the potential energy surface. This species near-instantly stabilizes into the end-bridge AsH + 2H species, in which the AsH fragment bridges between two dimer-ends, labeled (iii) in Figure 1g.

The final reaction step, labeled (iv) in Figure 1g, transfers a hydrogen atom from the AsH fragment to the silicon dangling bond at the opposite dimer end. This reaction produces the inter-row end-bridge As + 3H structure that corresponds to the type-1 STM feature. The reaction barrier for this final hydrogen-shift is calculated to be 0.89 eV; this is the rate-determining activation energy along the full dissociation path. This barrier is still low enough such that full dissociation of an adsorbing AsH₃ molecule into inter-row end-bridge As + 3H will be completed on a time scale of tens of seconds. Overall, the calculated reaction pathway is thus highly consistent with observation of the type-1 STM feature as the majority species following AsH₃ chemisorption on Si(001).

Despite the kinetic preference for this dissociation pathway, STM images also reveal small numbers of a few alternative dissociation structures, e.g., the type-2 feature seen in Figure 1a,b. These arrangements, which are kinetically disfavored on the bare Si(001) surface, are thought to result from steric hindrance provided by neighboring AsHₓ adsorbates, or native Si(001) defects. A single example of the type-2 feature is shown in Figure 1, in the bottom right of panels a and b, and notably occurs immediately adjacent a type-1 feature. At this low coverage, the type-2 feature comprises roughly 10−15% of the total AsHₓ associated STM features, and corresponds to the intrarow end bridge structure, known to be the preferred dissociation configuration for PH₃/Si(001).13

It is instructive here to briefly compare the dissociation of AsH₃ on silicon to the analogous PH₃/Si(001) system. The primary PH₃ dissociation process at low coverage is a progressive sequence of three principal species, namely, PH₂ + H, PH + 2H, and P + 3H.37 Transitions along this sequence occur on a time scale of minutes, making these transitions directly observable in room-temperature STM experiments (see, e.g., Figure 6 in ref 47). In contrast, full AsH₃ dissociation into the As + 3H species is complete before the surface can be imaged by STM after arsine exposure, leaving the type-1 STM feature as the single dominant species at low coverage. Comparison of our DFT calculations for AsH₃/Si(001) with those reported for phosphine provides some insights into how these differences arise. First, hydrogen-shift reactions in the arsine pathway are more exothermic and have lower activation energies than the corresponding reaction in the phosphine pathway. This results in a much faster rate of dissociation for AsH₃. Second, a characteristic of the phosphine pathway is that hydrogen-shift dissociation and PH₂-fragment diffusion are in close competition; this becomes manifest in the great diversity of PH₃ dissociation species formed, and in the observed diffusion of the P atom away from the initial adsorption site. This competition no longer exists for arsine due to the much reduced hydrogen-shift activation barrier. In effect, an adsorbing AsH₃ molecule rapidly falls apart where it lands, undergoing minimal rearrangement of the fragments. The result is that the arsenic atom does not have opportunity to diffuse away from the initial landing site and a single structure, namely, the inter-row end-bridge As + 3H structure, is formed as the majority species.

**Substitutional Incorporation of Arsenic Atoms into the Silicon Lattice.** Following molecular dissociation of AsH₃ on the Si(001) surface, the next step critical to the STM lithography fabrication process, is incorporation of the arsenic atoms into the silicon lattice. Figure 2a and b show a clean Si(001) surface exposed to a low dose (0.015 L) of arsine at room temperature (Figure 2a), and then subsequently annealed to promote the incorporation of the arsenic atoms into the surface (Figure 2b). Before annealing, the surface exhibits many As + 3H features (type-1), two of which have been highlighted by arrows. High-resolution filled- and empty-state images of the same surface (inset to Figure 2a) provide a close view of a type-1 feature (arrow) and a nearby C-defect. Figure 2b shows the same surface (but a different area) after a 1 min thermal anneal at 500 °C. Arrows in Figure 2b highlight examples that, by their appearance and by analogy to the behavior of PH₃ on Si(001),48 we assign to Si−As heterodimers formed during the anneal. High-resolution filled and empty state images of two such As−Si heterodimers (inset of Figure 2b) reveal an appearance that is very similar to that of P−Si heterodimers, confirming that arsenic incorporation into the surface has taken place.

At the temperature of 500 °C used in the annealing experiment above, all hydrogen atoms are expected to desorb from the silicon surface, and thus Figure 2b shows a surface completely depleted of hydrogen. At this temperature, the substituted silicon atoms, which are ejected from the surface as the As−Si dimers are formed, rapidly diffuse away from the ejection site and are eventually captured at the existing step edges, or newly formed ad-dimer rows.49 This process of controllably substituting arsenic atoms into the silicon lattice is a requirement for compatibility of arsenic and arsenic with STM hydrogen resist lithography, as substitution of isolated dopant atoms is necessary if the dopants are to be activated as electron donors in STM fabricated electronic devices.

**Saturation Arsine Coverage.** Having resolved the elementary AsH₃ single-molecule processes and arsenic surface incorporation in the low-coverage regime, we now move on to the saturation-coverage regime. High-density regions of arsenic dopants are important for STM lithography device fabrication, as these conductive 2D areas serve as device interconnects and contacts. Figure 2d shows a large-scale STM image of the Si(001) surface dosed to saturation by 1.5 L of arsine. The image shows a disordered pattern of protrusions and depressions as expected for a surface densely covered with the products of AsH₃ dissociation.

Hydrogen atoms bonded to Si(001) are immobile at room temperature, and in the stable configuration of the inter-row end-bridge As + 3H structure, the arsenic atom is locked into place by the three hydrogen atoms surrounding it. Activation barrier considerations reveal that arsenic incorporation can only take place when the thermal conditions are such that hydrogen atoms can diffuse away from the arsenic atom. The result of annealing the arsenic saturated surface to 350 °C for S
min is shown in the STM images in Figure 2e. That incorporation has occurred is evidenced in Figure 2e by the bright linear protrusions running perpendicular to the underlying dimer rows (also see inset to Figure 2b). These bright linear protrusions correspond to short ad-dimer chains, formed from the silicon atoms ejected during arsenic incorporation. These ad-dimer chains are a familiar sight from phosphorus incorporation experiments in the context of PH₃ lithographic fabrication. Because there is one ejected silicon atom for every incorporated arsenic atom, one can interpret the density of ejected silicon atoms observed in STM images as a direct measure of the density of incorporated arsenic. Using this approach we estimate the incorporated arsenic coverage in samples prepared by a saturation arsine dose of 1.5 L and thermal anneal of 350 °C to be 0.24 ± 0.04 ML.

We find that this incorporated arsenic coverage is independent of the arsenic saturation dose rate, which is interesting since this is not the case for phosphine (discussed further below). Saturation dosing at different rates can be achieved by changing the arsine partial pressure, with simultaneous adjustment of the total exposure time. When the arsenic dose rate is varied by decreasing the arsine partial pressure from 5 × 10⁻⁷ mbar to 5 × 10⁻¹⁰ mbar (along with a dose equalizing change in total exposure time from 5 to 50 min), the resulting total arsenic coverage is found to be constant within measurement error. This dose-rate independent saturation arsenic coverage can be explained as follows. (i) At low coverages (∼2 × 10⁻¹⁴ molecule/cm²), AsH₃ moieties dissociate rapidly (∼10 s) and completely at random locations. (ii) As the density of surface adsorbates increases, the occupied sites begin to sterically hinder the complete dissociation of further incoming AsH₃, resulting in the subsequent adsorption of a distribution of partially dissociated AsH₂ moieties—a distribution determined by site availability, not dissociation rate. (iii) Eventually all adsorption sites are blocked and coverage becomes saturated, leaving a final surface comprised of all possible AsH₂ moieties.

The constant incorporated arsenic coverage value, measured across a range of arsine dose rates, is also found to be consistent with predictions from simple kinetic Monte Carlo (KMC) simulations that utilize activation energies from our DFT-derived dissociation pathway (Figure 1g). The results of these simulations are plotted in Figure 2c. Here, we use KMC calculations to determine the total arsenic coverage at saturation, for a range of arsine partial pressures, spanning five orders of pressure magnitude, in which the distribution of AsH₃ fragments is allowed to vary. We assume that isolated molecules will dissociate in the type-1 structure, with a 2.5 dimer footprint discussed in the Dissociation of Isolated AsH₃ Molecules section, but that as the surface fills in, and silicon site availability becomes reduced, AsH₃ dissociation will be halted at different stages leaving behind AsH and AsH₂ fragments. Although the ratio of As, AsH, and AsH₂ varies as a function of partial pressure, the total number of arsenic atoms present on the surface remains in the 0.24–0.26 ML range, and the arsenic surface concentration only increases by 2–3% as the dose rate is increased across the full range.

This dose rate independence of arsenic coverage in the saturated AsH₃/Si(001) system is in direct contrast to the PH₃/Si(001) system, where phosphorus coverage at saturation is found to be dose-rate dependent at room temperature. When Si(001) is saturated with PH₃ there is competition over bare Si surface dimers between the adsorbing PH₃ molecules and the dissociating PH⁺ + H species. For AsH₃, this competition no longer exists, due to a dissociation rate that is 3 orders of magnitude more rapid than that of PH₃. In effect, any adsorbing AsH₃ molecule breaks apart on the Si(001) surface to a degree limited only by the availability of bare Si dimer sites surrounding the molecule.

The much more rapid dissociation of AsH₃ also manifests in a lack of ordering in this system. The saturated surface and magnified inset shown in Figure 2d show very little evidence of long- or short-range ordering. This is once again in direct
In the AsH3/Si(001) system, the H atoms are stripped from initial adsorption site to energetically preferred, ordered sites. PH3, dissociation and removal of successive H atoms from the PH moieties aligned along Si dimer rows.50 In the case of dangling bonds, introduced in the hydrogen mask by the STM fabrication process, AsH3 molecules must selectively adsorb only on depassivated Si(001) regions and not on the passivated Si(001):H surface. Furthermore, to allow for maximum in-plane confinement and fabrication flexibility, arsenic should also incorporate into the silicon lattice at anneal temperatures below the onset of H2 desorption (380 °C). Fulfillment of both these requirements is demonstrated in Figure 3. Each panel in this figure corresponds to one step in the STM lithography patterned doping process, with the corresponding steps indicated by the inset schematics on each STM image.

In the first step, a hydrogen passivated Si(001) surface is prepared, as shown in Figure 3a. Each silicon atom has one or two hydrogen atoms bonded to it, except in a few cases where the hydrogen atom is missing, leaving a dangling bond (DB). The DBs have the appearance of bright dots in the image and the DB density is <1%. Selective removal of a region of the hydrogen “resist” layer is achieved by scanning over a 50 × 50 nm region of the surface using an increased bias/current of +7.0 V and 1.0 nA, at a tip raster speed of 100 nm/s (Figure 3b). The patterned region, in the center of the image, shows the density of bright dots now approaches 100% coverage of the surface, indicating that most of the hydrogen atoms have been desorbed from the surface, leaving behind the bare silicon.

Dosing the patterned surface of Figure 3b with 1.5 L of arsine results in AsHx moieties (where x = 0–3) selectively adsorbing within the patterned area, producing a saturation arsenic density of 0.24 ML. An image of this surface is shown in Figure 3c. Following arsine dosing, the exposed region of the surface has a different appearance, consisting of a high density of features, whose identity is hard to distinguish due to their tight and disordered packing. An enlargement of the upper right corner of the patterned and dosed area is shown in Figure 3e. It is clear from these images that arsenic containing molecules from the gas phase have attached to the silicon surface where the hydrogen resist has been removed, while the resist protects the surface from AsHx desorption elsewhere.

In the final step, shown in Figure 3d, annealing the surface to 350 °C for 2 min results in substitutional incorporation of arsenic and the ejection of silicon onto the surface, exclusively within the patterned area. Additional bright spots now present in the image outside of the patterned region are silicon dangling bonds, introduced in the hydrogen mask by the contrast to the PH2/Si(001) system, where PH2 moieties are found to assemble in small patches of local \( p(2 \times 2) \) ordering, with PH moieties aligned along Si dimer rows.50 In the case of PH3, dissociation and removal of successive H atoms from the PH2 moieties occurs on the time scale of minutes, and the moieties survive long enough to undergo diffusion from the initial adsorption site to energetically preferred, ordered sites. In the AsH3/Si(001) system, the H atoms are stripped from the molecule much more rapidly, with isolated molecules completely dissociating on the time scale of tens of seconds. The AsHx moieties do not have opportunity to diffuse before fully dissociating, and at saturation coverage they remain immobilized at the initial, random landing site, producing the completely disordered surface shown in Figure 2a.

**Adsorption and Incorporation of Arsenic through a Hydrogen Resist.** We now demonstrate that the arsine adsorption, dissociation, saturation, and incorporation on the clean Si(001) surface, as described above, is compatible with hydrogen desorption lithography. For compatibility with the STM fabrication process, AsH3 molecules must selectively adsorb only on depassivated Si(001) regions and not on the passivated Si(001):H surface. Furthermore, to allow for maximum in-plane confinement and fabrication flexibility, arsenic should also incorporate into the silicon lattice at anneal temperatures below the onset of H2 desorption (380 °C). Fulfillment of both these requirements is demonstrated in Figure 3. Each panel in this figure corresponds to one step in the STM lithography patterned doping process, with the corresponding steps indicated by the inset schematics on each STM image.

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incorporation anneal. Once again, an enlargement of the upper right-hand corner of the patterned and dosed area (this time postanneal) is shown in Figure 3d. Two examples of short dimer chains consisting of ejected silicon atoms are highlighted with arrows. Confirmation of the successful execution of this final step demonstrates that an arsenic dopant layer can be substitutionally incorporated into an arbitrarily defined region of a silicon surface, where the shape of the region is defined by the hydrogen desorption pattern written by an STM tip.

Finally, the full patterning process is repeated on the atomic scale in Figure 3g. Here we demonstrate hydrogen desorption, AsH$_3$ adsorption/dissociation, and finally As incorporation along a patterned line of exposed dangling bonds only 2 silicon dimer rows wide. Arrows in these three images highlight a silicon dangling bond (DB) dimer in the first image, an adsorbed AsH$_3$ moiety in the second, and an ejected silicon adatom (ad-Si) adsorbed on the surface. The resulting patterned line of incorporated arsenic atoms is at most a few atoms wide.

**Encapsulation of Arsenic Layers: Low Temperature Silicon Epitaxy.** The final, *in situ*, ultrahigh vacuum step in the STM device fabrication is encapsulation with crystalline silicon. Once the arsenic atoms are incorporated into the silicon surface through a patterned hydrogen resist, the dopant structure must then be buried beneath a few nanometers of epitaxial silicon. This protective capping layer is grown via molecular beam epitaxy, using a solid intrinsic silicon sublimation source (SUSI-40, MBE Komponenten GmbH).

The thermal budget of the sample during the epitaxy stage is tightly restricted by two competing requirements: (1) dopant segregation and diffusion must be minimized so that the substitutional arsenic atoms remain in position—demanding a minimization of sample temperature; and (2) crystallographic defect formation must be minimized to ensure that arsenic donor atoms remain electrically activated—usually achieved by elevated temperature overgrowth. In the case of phosphorus, these two requirements can be reasonably satisfied by holding the sample temperature constant at 250 °C, throughout. In the case of arsenic we find (via secondary ion mass spectrometry; SIMS) that this approach is inadequate. We attribute this to the fact that arsenic is considerably more prone to surface segregation. As described below, we have determined a variable sample temperature program during overgrowth that suppresses arsenic surface segregation during silicon epitaxy, allowing for the growth of both well confined and electrically activated arsenic δ-layers.

To achieve thermal control of arsenic segregation and diffusion during silicon epitaxy, a number of saturation arsenic delta layers were overgrown with 15 nm of silicon at a rate of 1 ML/min, using different sample temperature programs.
Arsenic concentration depth profiles were subsequently measured using SIMS, and Figure 4 shows SIMS data for four different temperature programs as described by the temperature vs time insets in each SIMS profile. Figure 4a shows the result of silicon overgrowth at a constant sample temperature of 240 °C. This arsenic distribution is dominated by a high and narrow peak located at a depth of ~1 nm. Beneath this narrow surface peak the arsenic concentration reduces to a much lower, but not an insignificant constant level (>1 × 10^{18} atoms/cm^3) before terminating in a slight peak at the δ-layer origin depth, 15 nm below the surface. Beyond this depth, the arsenic concentration rapidly drops below the instrument’s detection limit (1 × 10^{15} atoms/cm^3). From this we conclude that the arsenic layer is not effectively buried beneath the silicon, but rather, a large percentage of the 0.24 ML of arsenic atoms undergo continual cycles of surface segregation, i.e., they exchange positions with newly arrived silicon adatoms, and assume energetically preferred positions within the top layer.55,56

Figure 4b shows the arsenic profile for overgrowth at a constant but lower temperature of 190 °C. Here we observe a shift in weight from the surface to the subsurface arsenic peak, and find that the arsenic concentration is no longer uniform in the region between peaks. Below the surface peak, it slowly rises to an asymmetric peak at the arsenic origin depth of ~15 nm, before rapidly falling off below the detection limit. Segregation is an activated process, and by lowering the constant growth temperature the segregation is found to be partially suppressed.

This arsenic segregation suppression can be further enhanced through a greater reduction in substrate temperature during encapsulation. Figure 4c shows the arsenic profile resulting from overgrowth at the lowest achievable sample temperature given our silicon source—sample geometry. In this case, there is no Joule heating of the sample, only the unavoidable radiative heating from the silicon MBE source. The substrate begins overgrowth at 40 °C increasing to a stable 100 °C after the first 12 min of growth (12 ML). From the SIMS profile, we see that arsenic surface segregation is highly suppressed in this sample, resulting in a very well confined (<2 nm fwhm) and symmetric peak (~ 5 × 10^{20} atoms/cm^3) located at the origin depth of 15 nm. This peak, which falls rapidly to below the instrument’s detection limit within 3 nm in either direction, represents the best achievable confinement of the arsenic δ-layer grown in our MBE system. However, due to the reduced substrate temperature during overgrowth, and the high concentration of arsenic surface atoms, the critical thickness at which the epitaxial layer becomes amorphous will also be reduced considerably.57 At a temperature of 100 °C, and deposition rate of 1 ML/min, the critical thickness is expected to reduce below 2 nm,53 meaning the upper 13 nm or more of silicon will be amorphous, and unsuitable for device encapsulation.

In order to facilitate suppression of arsenic segregation during reduced temperature silicon overgrowth, yet still achieve epitaxy and avoid amorphization, we have developed the strategy (adapted from a similar concept used first for phosphorus δ-layers in germanium,58–60 and later for phosphorus δ-layers in silicon61–63) of growing a very thin, reduced temperature “locking layer” that is kept below the critical amorphization thickness. With a reduced temperature locking layer in place, the substrate is then heated to a temperature that can support epitaxial growth for the remainder of the overlayer deposition. Figure 4d shows an example of locking layer assisted epitaxial growth applied to an arsenic δ-layer. Here the sample is not actively heated for the first 10 min (10 ML for a 1 ML/min growth rate) after which it is then rapidly heated to 250 °C, and held constant for the remaining 230 min of overgrowth. Comparing the SIMS profiles of Figure 4c and d, we see that increasing the temperature to 250 °C following growth of a 10 ML locking layer, has no detrimental effect on the confinement. The locking layer strongly suppresses arsenic surface segregation, producing epitaxially capped arsenic δ-layers with very tight confinements—confinements which are limited primarily by the solid-state diffusivity of arsenic in silicon. It is interesting to note here that arsenic has a lower diffusion coefficient than phosphorus in bulk silicon.53 Arsenic structures buried in silicon should therefore be less prone to pattern smearing at increased processing temperatures compared to phosphorus structures.

**Electrical Activation and Contacting of Arsenic δ-Layers.** To establish full compatibility of arsine and arsenic with the STM device fabrication process, the buried arsenic dopant structures must be electrically activated and Ohmic contacts achieved with the arsenic layer beneath the silicon cap. Satisfaction of this requirement has been verified by measurement of linear and symmetric IV curves during standard magnetotransport characterization of the arsenic δ-layers. As explained in detail in the Supporting Information, Hall bars were fabricated from locking layer encapsulated arsenic δ-layers, and Ohmic contacts achieved through deposition of aluminum directly onto the δ-layer edge. We note that phosphorus doped, n-type substrates (donor concentration N_D = 5 × 10^{14} to 5 × 10^{15} cm^{-3}) are used in this work. This doping level allows for direct current sample heating and room temperature STM measurements, and does not interfere with the low temperature transport measurements of the delta layers.

In addition to fabricating Ohmic contacts, we have also optimized the transport properties of the arsenic δ-layers for improved device performance. The optimizations were achieved by improving the epitaxial silicon quality and minimizing crystallographic defects, resulting in a maximization of carrier density and mobility, and a minimization of sheet resistance. Despite the fact that the locking layer thickness used is below the critical amorphization limit, including this reduced temperature step in the overgrowth is detrimental to crystal quality, and thus to the electrical performance of the δ-layers. To overcome this issue, we include a rapid thermal anneal (RTA) stage in the overgrowth sample temperature program, as has been done for phosphorus δ-layers.51–63

Figure 4e shows the SIMS profile of an arsenic δ-layer grown using both locking layer and RTA stages in the substrate temperature program. The temperature vs time profile used here is shown in the inset of Figure 4e. It is similar to the profile shown in the inset of Figure 4d, with the exception of the additional 500 °C × 10 s RTA, occurring between the locking layer growth and constant temperature epitaxy stages. Comparing the SIMS profiles of Figure 4d and e, we find that introduction of this RTA stage is not detrimental to the enhanced confinement provided by the locking layer in any significant way. The effect of the RTA on transport properties is however significant.
Figure 4f and g show the effect of the RTA temperature maximum on δ-layer carrier concentration, sheet resistance, mobility, and confinement. The data for these figures of merit were obtained from Hall measurements on five arsenic δ-layers, with heating profiles identical to that of Figure 4e, except with each using a different RTA peak value, covering the range from 250 to 560 °C. For a RTA greater than 500 °C, the electronic layer thickness increases beyond 2 nm, while the improvement in sheet resistance and mobility has leveled out. For this reason, the RTA temperature maximum is considered optimum at ~500 °C. Despite the fact that both the mobility and sheet resistance values plateau at 28 ± 1 cm²/V-s, and 1.6 ± 0.1 kΩ/square, respectively, the carrier density continues increasing beyond this optimum RTA temperature. At the highest RTA temperature maximum examined, 560 °C, we measure a carrier density of 1.45 × 10¹⁴ cm⁻², which corresponds to an arsenic donor atom density of only 0.21 ML. Given our estimate of arsenic saturation coverage at 0.24 ML, this maximum value represents an activation of 88% of the donor atoms. At the optimized RTA temperature of 500 °C, this activation percentage is even lower at 81%. Nevertheless, this carrier concentration is well above the metal insulator transition, and greater than 1/2 of the typical carrier density of 2.4 × 10¹⁴ cm⁻² found in phosphorus δ-layers. It is not entirely surprising that less than 100% of the arsenic δ-layer donor atoms are electrically activated, even at the highest RTA temperature used, as arsenic is known to suffer from clustering at high concentrations,⁶⁴,⁶⁵ resulting in incomplete activation at concentrations >10¹⁵ cm⁻² (or 2 × 10¹³ cm⁻²) in traditional implantation or diffusion doping of silicon.⁵³

Given the large parameter space involved in optimizing arsenic δ-layer growth for both confinement and electrical transport, there is potentially room for improvement in performance values beyond those presented here. However, comparing the figures of merit achieved for arsenic δ-layers with those typically achieved for phosphorus δ-layers,⁶¹,⁶³ we find that they are at most a factor of 2 worse, and that arsenic and arsenic are 100% compatible with atomic-scale STM device fabrication, and therefore represent a viable alternative to phosphine and phosphorus.

CONCLUSIONS

We have demonstrated the full compatibility of arsenic as a precursor gas for the atomic-scale positioning of arsenic donors in silicon using STM-based hydrogen-desorption lithography, and, when combined with low temperature silicon epitaxy, the capability to fabricate buried, atomic-scale, dopant structures in silicon. Room temperature dissociation of arsenic on Si(001) differs from the dissociation of phosphine on Si(001) in that full dissociation of arsenic occurs much more rapidly at room temperature. We find also that saturation arsenic coverage is self-limiting at 24% ML, invariant with dose rate, and that monolayer hydrogen termination acts as an effective resist layer for the adsorption of arsenic on Si(001). We show that arsenic can be incorporated in the silicon surface lattice by annealing at 350 °C. We demonstrate successful low temperature silicon epitaxial overgrowth by growing arsenic δ-layers, where segregation was suppressed through the use of an unheated, ultrathin, locking layer. A rapid thermal anneal stage of 500 C × 10 s immediately following locking layer growth improves electronic transport properties resulting in 2 nm thick, conductive layers with sheet resistance, and carrier concentration of 1.5 kΩ/square, and 1.3 × 10¹⁵ cm⁻², respectively. It has been a decade and a half since phosphorus dopant placement using STM H-lithography was first demonstrated, and the work presented here shows that using arsenic as the n-type dopant in place of phosphorus is possible. Finally, the parameters used for arsenic and for phosphorus device fabrication are completely compatible with one another, demonstrating the possibility of multispecies dopant devices.

EXPERIMENTAL METHODS

Sample Preparation and Scanning Tunneling Microscopy. Si(001) samples were diced to 2 × 9 mm from a 0.3 mm thick, Czochralski grown, bulk phosphorus doped wafer with a resistivity of 1−10 Ω-cm. Samples were cleaned ultrasonically ex situ in acetone followed by isopropyl alcohol, then thermally outgassed in situ (base pressure <1 × 10⁻¹⁰ mbar) for >8 h at 600 °C, and finally flash annealed multiple times at 1200 °C, using direct current resistive sample heating. Sample temperature was monitored using an infrared pyrometer (IMPAC IGA50-LO plus), with a total estimated measurement uncertainty of ±30 °C. Silicon epitaxy was performed at a base pressure of 1 × 10⁻¹⁰ mbar, using an all silicon, solid sublimation source (SUSI-40, MBE Komponenten GmbH) operated at a deposition rate of 1.5 Å/min. During silicon overgrowth, sample temperature was indirectly monitored by measuring sample resistance, while the sample was heated using direct current resistive sample heating. All scanning tunneling microscopy measurements were performed in an Omicron variable temperature series STM at room temperature with a base pressure of <5 × 10⁻¹¹ mbar.

Density Functional Theory Calculations. All DFT calculations reported here were performed using the Gaussian 09 software and methods of energy computation and geometry optimization implemented therein.⁶⁶ We calculate adsorption energies for various arsine configurations on the silicon (001) surface using an approach that we refer to as the cluster composite model (CCM).¹⁵ This approach uses the B3LYP hybrid exact-exchange density functional and a large 6-311G(2df,2pd) atom-centered basis set, and the Si(001) surface is represented using hydrogen-truncated cluster models, such as Si₆H₁₆ or Si₁₆H₄₄ with correction terms applied to minimize the effects of finite cluster size. The CCM approach has an established track record of describing the dissociation pathways of small molecules on the Si(001) surface and providing an explanation of the adsorbate species observed in STM experiments (see, e.g., refs 13, 46, 68, 69). The specific approach used here for the AsH₃/Si(001) system is exactly the same as in our previous work describing the complex dissociation pathways of the analogous PH₃/Si(001) system,¹³ and the reader is referred to this article for the full technical details. While this approach may be computationally expensive, it has been proven for the PH₃ system, and its use here allows direct and transparent comparison between the two systems. Reaction step (i) in Figure 2g was calculated using what is referred to as the 3D2R cluster model in ref 13, while reaction steps (ii), (iii), and (iv) used the 2D1R cluster model with the hemihydride dimer represented by a separate cluster (see eq 5 in ref 13).

Kinetic Monte Carlo Simulations. KMC methods are explained in detail in the Supporting Information.

Secondary Ion Mass Spectrometry. Time of flight (ToF) SIMS measurements were made using an IONTOF ToF-SIMS⁷ system with a 25 keV Bi⁺ primary ion beam in high current bunch mode (HCBM), and a 500 eV, 35 nA Cs⁺ sputter beam. Depth profiles were made with a 300 μm² sputter crater, and the analytical region was gated to the central 50 μm² of the sputter region.

Hall Bar Fabrication and Magnetotransport Measurements. Arsenic delta layer samples were fabricated into 6 terminal Hall bars using standard UV photolithography, with electrical contact made via aluminum thermally evaporated over the edge of the mesa structures. Magnetotransport measurements were made at a temperature of 5 K in a Cryogenics Ltd. cryogen free measurement system using magnetic fields up to 5 T. Details of these measurements, and quantification of the delta layer electrical transport characteristics, are explained in detail in the Supporting Information.
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