Catalyst Composition and Impurity-Dependent Kinetics of Nanowire Heteroepitaxy

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ABSTRACT The mechanisms and kinetics of axial Ge–Si nanowire heteroepitaxial growth based on the tailoring of the Au catalyst composition via Ga alloying are studied by environmental transmission electron microscopy combined with systematic ex situ CVD calibrations. The morphology of the Ge–Si heterojunction, in particular, the extent of a local, asymmetric increase in nanowire diameter, is found to depend on the Ga composition of the catalyst, on the TMGa precursor exposure temperature, and on the presence of dopants. To rationalize the findings, a general nucleation-based model for nanowire heteroepitaxy is established which is anticipated to be relevant to a wide range of material systems and device-enabling heterostructures.

KEYWORDS: nanowire · Ge–Si heteroepitaxy · nucleation barrier · environmental transmission electron microscopy · AuGa catalyst alloy · dopants

Heterojunctions play a central role in modern semiconductor devices, ranging from quantum well optoelectronics, † two-dimensional electron gas devices,‡ and thermoelectric devices to tunnel field effect and high mobility strained transistors.³ Small diameter semiconductor nanowires (NWs) offer an unprecedented range of new material combinations⁴–⁶ to form device-relevant heterostructures,⁷–¹² due to their ability to accommodate lattice-misfit strain¹³ and the ease of their integration via bottom-up growth. A key prerequisite is thereby the controlled formation of NW junctions during growth, in particular, compositionally abrupt and structurally perfect axial NW heterostructures.⁹,¹⁴ Whereas sharp, defect-free heterojunctions can readily be achieved within group III–V NWs by vapor–liquid–solid (VLS) growth where the group V constituent typically has a low solubility in the conventionally used Au catalyst,¹⁵ this turns out to be much more challenging for group IV NWs with significantly higher solute solubilities¹⁶. In particular for Si/Ge, the amount of growth solute in the Au catalyst nanoparticle,¹⁶ which acts as a reservoir during VLS growth,¹⁷ generally limits the junction abruptness, creating a current limitation to the further exploitation of the inherent advantages of these NWs. This reservoir effect can be minimized by catalyst alloying.⁵,¹⁸,¹⁹ A particular versatile process is thereby to alloy the catalyst via the vapor phase during chemical vapor deposition (CVD), for example, the use of trimethylgallium (TMGa) to create AuGa alloy catalyst nanoparticles.¹⁹ Ga lowers the solubility of Ge and Si in the catalyst,²⁰ enabling sharper Ge–Si heterojunction NWs. However, currently there is limited understanding of the underlying mechanisms of such advanced axial heterostructure formation, the signature of which are unexpected side effects such as the NW diameter becoming nonuniform at the Ge–Si heterointerface, creating a “bulge” (Figure 1).

In this study, we use environmental transmission electron microscopy (ETEM) to study in situ the morphological details of Ge–Si NW heterojunction formation based on the tailoring of the Au catalyst composition via TMGa addition. We establish a nucleation-based model which allows us to rationalize the observed dependencies of bulging and...
related morphological changes at the Ge–Si interface with respect to the catalyst composition and the presence of impurities, such as dopants. Rather than just looking at the catalyst reservoir effect\(^\text{17}\) and the surface energy balance at the heterointerface\(^\text{6}\) as in previous literature, we propose that the nucleation barrier for the heteroepitaxy (i.e., the Si precipitation on Ge) depends on the solubility of the growth species in the catalyst. The observed bulging is thereby a physical manifestation of the Si supersaturation in the alloy catalyst and reflects a transient change of surface energies. The presence of dopants can depress the nucleation barrier and consequently eliminate the formation of a bulge near the Ge–Si heterointerface. Hence the kinetics of heterointerface nucleation can be controlled through the catalyst composition and the addition of impurities during growth. Due to the generality of our arguments, we anticipate that our model is relevant to a wide range of material systems and device-enabling heterostructures, for some of which similar morphologies have been reported already.\(^\text{21,22}\)

RESULTS

Figure 1A–F shows post-growth scanning electron microscopy (SEM) and transmission electron microscopy (TEM) images of \textit{ex situ} grown Au-catalyzed Ge–Si heterojunctions, highlighting in particular the morphological differences (i.e., bulge size) based on the level of TMGa addition to the catalyst. In brief, the nanowire growth process consists of initial epitaxial GeNW growth using a GeH\(_4\) precursor, followed by the addition of TMGa to alloy the Au catalyst particle and
then growth of a SiNW segment using SiH₄ (see Methods). The addition of Ga to the liquid Au catalyst serves to decrease the Au–Ge eutectic composition (28 atom % Ge for pure Au; ~8 atom % for ~2:1 Au/Ga alloy), thus significantly lowering the concentration of Ge in the liquid catalyst prior to SiNW growth. This reduction of the reservoir of Ge atoms in the catalyst prior to Si growth leads to a much sharper Ge–Si compositional transition, and details of this process have been demonstrated previously. Here we focus on the underlying mechanisms and kinetics of this heteroepitaxy, which for instance manifest themselves in the formation of a bulge at the Ge–Si interface, that is, a locally asymmetric increase in NW diameter at the heterointerface. Figure 1 shows that the relative size of this bulge increases with increasing Ga catalyst composition. Ge–Si NWs grown from elemental Au, without TMGa addition, show no bulge (Figure 1B), whereas a clear bulge is observed for 23% Ga (Figure 1C), which gets even more pronounced for 34% Ga (Figure 1D). These values refer to the Ga mole fraction in the catalyst alloy based on SEM- and TEM-based EDX analysis of the catalyst particle. This bulge morphology is not an isolated curiosity, but rather is observed for >90% of the NWs for a given sample. This reflects a conservative estimate of abundance, noting that for the remaining 10% of the NWs, a bulge may be present but is not visible due to the projected nature of the imaging. We note that smaller regularly spaced sawtooth sidewalls can be a common feature of [111]-type wires, in particular, for UHV-based growth conditions, but the appearance of such a pronounced bulge has not been reported and more importantly its formation cannot be rationalized with previously reported growth models.

The size of the bulge at the heterointerface also depends on the temperature at which the TMGa exposure and catalyst alloying is carried out. For a lower TMGa exposure temperature of ~280 °C, the bulge becomes more pronounced and protrudes in a more radially symmetric way (Figure 1E). In addition, we find the size of the bulge to be also influenced by the presence of dopant gases during NW growth. Figure 1F shows a Ge–Si heterojunction which was grown with an additional doping modulation from p-type Ge to n-type Si, implemented using B₂H₆ and PH₃ precursors during the growth of the respective NW segments (see Methods). No bulge is visible at the heterointerface for the many NWs observed, although the Ga fraction in the catalyst alloy was ~20%. A more in-depth discussion is given further below as to the influence of the dopants on the growth kinetics governing the formation of the observed bulge.

In order to gain unique insight into the formation of the observed bulge at the Ge–Si interface, we use an ETEM to observe in situ the growth of the heterostructure from a Au–Ga alloy catalyst. Figure 2A–F are representative of a video sequence (video S1 in Supporting Information) recorded at 9 frames per second. After exposure to TMGa, Figure 2A,B shows a GaNW with a liquid AuGa catalyst alloy at the tip during disilane exposure, moments before the nucleation of a Ge₁₋ₓSiₓ heterojunction. We note that the liquid AuGa catalyst/Ge interface (dashed line) is not planar but truncated toward the triple phase boundary (TPB). We previously highlighted the importance of this TPB morphology in the context of the step flow cycle that constitutes axial NW growth. In particular, the
truncated edges act as a preferential attachment site for dissolved semiconductor material leading up to step nucleation and thus provide a direct visualization of the level of supersaturation in the catalyst.27–29 At 19 s, a distortion is observed in the upper right of the projected TPB (Figure 2C), leading to the filling of the truncated edge at the TPB (Figure 2D), presumably with precipitated Ge$_{1-x}$Si$_x$. The limited time resolution of the ETEM movie here does not allow us to resolve any oscillatory behavior of the truncated TPB morphology. However, we can clearly observe how, after the initial precipitation event, NW growth accelerates and the Si segment temporarily increases in diameter (Figure 2D,E) to then return to its original size as growth continues, leaving behind a small bulge (Figure 2F).

Note that we cannot directly measure the Ge composition after Ga exposure or the Si composition prior to the resumption of growth. However, as the NW did not appear to grow during TMGa exposure, we believe a significant fraction of the Ge remained in the catalyst prior to regrowth. In addition, since the NW diameters were all above 10 nm, we do not expect a Gibbs–Thomson effect to significantly alter the alloy catalyst composition.

Despite different CVD conditions, the observation of a bulge from the in situ data is consistent with ex situ grown observations (Figures 1 and 2G). A fast Fourier transform (FFT) analysis of representative phase contrast TEM images (see inset Figure 2G) shows that the upper and lower bulge facets are positioned approximately normal to the [111] and [002] directions. Hence we suggest that the bulge can be seen as an enlarged sawtooth facet consisting of [111]/[100]-type planes on the [111]-orientated NWs. Supporting Information Figure S1 gives further high-resolution TEM analysis of the Ge–Si heterostructure (Figure 2G), highlighting the as-grown crystalline quality.

Further to our ETEM data showing a truncated growth interface, post-growth imaging of ex situ grown NWs reveals a nonplanar heterojunction interface for both intentionally doped and undoped NWs. Figure 3 compares ex situ transmission electron microscopy data of an undoped Ge/Si heterojunction NW grown with a 34% Ga alloyed catalyst (Figure 3A) and a boron-doped Ge/phosphorus-doped Si heterojunction grown with a 20% Ga alloyed catalyst (Figure 3B). Details of the CVD growth process for these samples are described in Methods below. In both cases, the Ge–Si interface shows a sharp compositional transition but is nonplanar with a clearly truncated edge. Whereas for the undoped NW a clear bulge can be seen in the Si region near the truncated edge (Figure 3A), there is no such bulge for the doped NW (Figure 3B). We note that for both the ex situ doped and undoped NWs, only a single truncated edge is observed in projection. As shown in Figure 2, the truncated edge forms at the TPB which serves as a preferential site for solute incorporation likely leading to an enhanced incorporation of dopants at the truncated edges as shown in a similar case by Connell et al.30 A formal discussion as to the formation of the truncated edge and the possible enhanced incorporation of dopants in this case is beyond the scope of this paper and will be addressed separately.

**DISCUSSION**

We next turn to a discussion of the kinetics governing the formation of the bulge for which we rely on three key observations from both the in situ ETEM and ex situ growth data. First, we observe that the size of the bulge is dependent on the Ga composition of the catalyst, the TMGa exposure temperature, and the presence of dopants (Figure 1G). Second, we find initially preferred Ge$_{1-x}$Si$_x$ precipitation at the truncated TPB edge.31 Third, the AuGa catalyst requires a long incubation time despite high precursor pressures. Our data, in particular, the rapid NW growth that occurs after the nucleation of Ge$_{1-x}$Si$_x$ (i.e., after Figure 2B), do not support a possible reduced activity of the catalyst as
the principle explanation for this long incubation time.\(^{19}\)

We propose here the long incubation time indicates that a relatively high supersaturation in the AuGa catalyst is needed to nucleate Ge\(_1\text{−}x\)Si\(_x\) on Ge, suggesting the presence of a significant nucleation barrier for Ge\(_1\text{−}x\)Si\(_x\) precipitation on the GeNW. To reconcile all key observations, we introduce a model that couples the kinetics of Ge−Si heterostructure nucleation to catalyst composition and the presence of dopants during growth.

The premise of our model is centered on the nucleation barrier for the NW heteroepitaxy. As a consequence of the relatively low solubility of Ge in the AuGa alloy\(^{20}\) (the very aspect that produces a sharper Ge−Si transition relative to growth from elemental Au\(^{19}\)), a more Si-rich Ge\(_1\text{−}x\)Si\(_x\) layer must first nucleate at the liquid/solid interface of the GeNW. We assert that the more Si-rich the layer to be nucleated, the higher is the expected nucleation barrier for heteroepitaxy. A widely used, simple model for heteroepitaxy of growing material B on A is to consider the balance of the relevant surface \((\alpha_B/A\), \(\alpha_A/A)\) energies: \(\Delta \sigma = \sigma_B - \sigma_A\). A layer-by-layer growth mode is expected to occur for \(\Delta \sigma < 0\), whereas island growth of B on A is expected for \(\Delta \sigma > 0\).\(^{12}\) This simple idea has been previously expanded to NW heteroepitaxy,\(^6\) noting that, for any given material combination, layer-by-layer growth will lead to a straight NW morphology, while island growth will lead to a kinked morphology. We note that Si epitaxy on Ge is energetically unfavorable\(^{33}\) (i.e., \(\Delta \sigma > 0\)). Nonetheless, our data here show straight SiNW segments to grow on GeNWs (Figures 1−3). The existence of an energy barrier to deposit Ge\(_1\text{−}x\)Si\(_x\) on Ge, however, is consistent with our observation of a long catalyst incubation time, indicating that a high Si supersaturation is required to start growth. Hence an excess of Si must build up in the AuGa catalyst to nucleate Ge\(_1\text{−}x\)Si\(_x\) layer-by-layer growth on the GeNW.

We hypothesize that the solubility of Ge in the alloy catalyst is the key variable in the formation of the bulge, and this is reinforced by two key experimental observations. First, the bulge size is proportional to the Ga composition in the catalyst where an increase in bulge size is observed with increasing Ga composition from 17 to 34% (Figure 1G). Second, the TMGa exposure temperature (i.e., temperature at which the AuGa alloy catalyst is formed) influences the bulge size (Figure 1E), despite the catalyst still having 34% Ga. At 280 °C before TMGa exposure, we estimate the liquid AuGe catalyst to have 22.7% Ge (mole fraction)\(^{34,35}\) compared to 28.4% Ge at 380 °C. Therefore, there is less Ge initially dissolved in the liquid AuGe catalyst at lower temperatures, and adding Ga to the catalyst (via TMGa) results in an even lower Ge mole fraction in the catalyst nanoparticle. The low Ge composition of the catalyst following the TMGa exposure at 280 °C suggests an even larger barrier to nucleate Ge\(_1\text{−}x\)Si\(_x\), requiring a larger Si supersaturation to restart NW growth, resulting in a larger bulge at the Ge−Si heterointerface (Figure 1E) compared to 380 °C. We thus infer that the bulge is a physical manifestation of the Si supersaturation in the liquid AuGa catalyst.

As highlighted in Figure 1F and 3B, with the addition of dopant gases, it is possible to grow sharp Ge−Si heterojunctions without heterointerface deformities (i.e., without a bulge) from liquid AuGa catalyst particles. This suggests that the boron impurities do not raise the Ge solubility of the AuGa catalyst, and instead that the introduction of dopant impurities (phosphorus during Ge growth and boron during Si growth) lowers the energy penalty for Ge\(_1\text{−}x\)Si\(_x\) precipitation on Ge, that is, lowers the nucleation barrier for the heteroepitaxy. This hypothesis is supported by evidence which directly shows dopants to preferentially incorporate at the truncated edge of the TPB and suggests that impurities play a role in the nucleation process in VLS grown nanowires.\(^{36}\) Impurities are known to lower the nucleation barrier in phase transformations, depressing the barrier for homogeneous nucleation and reducing the supersaturation needed to achieve solidification.\(^{57}\) Such impurities are referred to as inoculants and are commonly used to speed up nucleation in undercooled melts.\(^{38}\) If impurities are lowering the nucleation barrier, then a lower supersaturation is required to nucleate Ge\(_1\text{−}x\)Si\(_x\), resulting in a smaller bulge. Although the exact atomistic mechanisms by which dopants may lower the nucleation barrier remain unclear, we speculate that the dopants may relax some of the strain in the Ge\(_1\text{−}x\)Si\(_x\) cluster nuclei during precipitation on Ge. In order to reduce the formation of the bulge without the introduction of impurities, while at the same time be able to create a sharper junction, a suitable catalyst system is needed which lowers the solute solubility and simultaneously modestly changes the nucleation barrier. To our knowledge, no such catalyst system (pure or alloy) has been identified for VLS growth which will lead to abrupt junctions without the formation of a bulge. However, by switching to a VSS growth approach, this can be achieved as demonstrated by Wen et al.\(^7\)

We note that transient changes in the surface/interface free energy terms could provide an alternative explanation for the observed bulge formation. In terms of Young’s wetting equation,\(^{37}\) a decrease in the liquid−vapor surface energy and/or increase in the liquid−solid surface energy during the Ge−Si transition would result in a decreased wetting angle, allowing the alloy droplet to cover a larger liquid−solid interface and to effectively increase the NW diameter (assuming the droplet’s volume is constant). Our data show, however, that the maximum of the bulge occurs at some point after the Ge−Si heterointerface when Si can be assumed to be the dominant dissolved semiconductor material in the catalyst. A simple surface energy argument thus fails to explain the observed heterointerface contour. Further, such argumentation is also inconsistent with the observed dependence on the TMGa exposure temperature (Figure 1E). To explain our observations, we propose that the initial high supersaturation needed to initiate
heterostructure formation temporarily alters the surface energies and, consequently, the equilibrium force balance at the TPB. Upon initiation of growth of the $\text{Ge}_{x} \cdot \text{Si}_{1-x}$ segment on Ge, the high supersaturation in the liquid alloy droplet appears to either lower the liquid–vapor energy or raise the liquid–surface energy, decreasing the droplet wetting angle and causing the NW diameter to increase accordingly (see Figure 2E). However, because the supersaturation in the droplet then decreases after nucleating $\text{Ge}_{x} \cdot \text{Si}_{1-x}$ on Ge, the surface energies return to their steady-state growth values, resulting in the NW diameter to contract according to the changed force balance at the TPB. The result of the initial heterostructure formation process is the formation of a “bulge” which has its maximum slightly after the actual heterointerface (see Figure 2F).

In order to go beyond purely qualitative arguments, we employ a continuum nucleation model of depositing an $\text{A}_{1-x}\text{B}_{x}$ alloy on pure A from a liquid catalyst C (analogous to depositing $\text{Ge}_{x} \cdot \text{Si}_{1-x}$ on Ge from a liquid AuGa catalyst). Note that we ignore the added complexities of step flow, 2D nucleation, and nanoscale effects. The surface energy difference $\Delta \gamma(x)$ of depositing $\text{A}_{1-x}\text{B}_{x}$ on A is treated as a linear function of the material composition being deposited: $\Delta \gamma(x) = x\Delta \sigma$, where $x = N_B/(N_A + N_B)$ and $\Delta \sigma = \sigma_B - \sigma_A$. Note $\Delta \sigma > 0$, as for Si growth on Ge.\(^3\) $N_A$, $N_B$, and $N$ are the number of A, B, and C atoms in the liquid catalyst, respectively. Taking $\Delta \gamma(x) = x\Delta \sigma$ as the surface energy, we can construct an expression for the Gibbs free energy change associated with nucleating $\text{A}_{1-x}\text{B}_{x}$ on pure A (eq 1):

$$\Delta G = \beta x \Delta \sigma (\Omega N)^{2/3} - \Delta \mu N$$

where $\Omega$ is the atomic volume of A or B, $\beta$ is a geometric prefactor which governs the shape of the critical nucleus, and $N$ is the number of atoms in the $\text{A}_{1-x}\text{B}_{x}$ cluster that will be precipitated on the A crystal surface. Treating the liquid as an ideal solution, the supersaturation is given by $\Delta \mu = k_B T [(1 - x)\ln(c^A) + x \ln(c^B)] - \mu_s$, where $c^A = N^A/ (N^A + N^B + N^C)$ and $c^B = N^B/ (N^A + N^B + N^C)$ are the mole fraction of A and B in the liquid $\text{A} - \text{B} - \text{C}$ catalyst, respectively; $\mu_s$ is the chemical potential of solid $\text{A}_{1-x}\text{B}_{x}$.

We rewrite eq 1 to get the following dimensionless expression for the free energy (eq 2):

$$\frac{\Delta G}{k_B T} = bxN^{2/3} - N[(1 - x)\ln(c^A) + x \ln(c^B) - a]$$

where $a$ and $b$ are dimensionless coefficients defined as $a = \mu_s/k_B T$ and $b = (\beta x(2/3)\Delta \sigma)/k_B T$.

We want to understand how raising/lowering the composition of A in the catalyst, $c^A$, and the dimensionless surface energy term $b$ determine if a nucleation barrier $\Delta G/k_B T$ is present in eq 2. The presence of a nucleation barrier is important, indicating a significant supersaturation must be reached to deposit $\text{A}_{1-x}\text{B}_{x}$ on A. Figure 4A shows regions where a nucleation barrier is present for various values of $b$ and the cut off composition $c^A$ where $\Delta G/k_B T = 0$. In the dark blue region of Figure 4A, there exists a barrier to nucleate $\text{A}_{1-x}\text{B}_{x}$ on A (i.e., $\Delta G/k_B T > 0$). In the white region of Figure 4A, no such barrier exists (i.e., $\Delta G/k_B T = 0$) and growth of $\text{A}_{1-x}\text{B}_{x}$ on A proceeds readily, even under small supersaturations. The underlying free energy functions are plotted in Figure 4B,C for pathways $X \rightarrow Y$ and $Y \rightarrow Z$, respectively. Point $X$ in Figure 4A corresponds to an alloy catalyst rich in A and consequently no nucleation barrier.

Traveling from $X$ to $Y$ illustrates how progressively higher supersaturations are needed to achieve nucleation and growth of $\text{A}_{1-x}\text{B}_{x}$ on A as $c^A$ falls. As $c^A$ falls along pathway $X \rightarrow Y$, we must nucleate more B-rich $\text{A}_{1-x}\text{B}_{x}$ on A, caus-
In Figure 5, we illustrate how the observed morphology of our Ge–Si heterostructured nanowires can be explained in the framework of our kinetic model (eq 2 and Figure 4). In Figure 5, we again have points X, Y, and Z which correspond to Ge–Si heterostructure formation under different growth conditions. Point X corresponds to Ge–Si heterostructure growth from a Au catalyst (Figure 1B): as the Ge solubility in the catalyst is relatively high, Ge-rich Ge1−xSi0x readily precipitates on the GeNW with no significant nucleation barrier for heteroepitaxy, and growth is possible at low supersaturations. Point Y corresponds to lowering the Ge solubility in the catalyst through the creation of a AuGa alloy (Figure 1C,D). As the Ge catalyst composition falls, we expect the nucleation barrier for Ge1−xSi0x precipitation to rise (as illustrated in our A–B model along pathway X–Y in Figure 4A), requiring a higher supersaturation to achieve heteroepitaxial growth and producing a bulge in the NW. Point Z corresponds to growing doped Ge–Si heterostructure NWs with AuGa catalyst particles (Figure 1F). In this case, the addition of the dopant impurities has reduced the overall energy penalty needed to achieve epitaxial growth, allowing sharp Ge–Si heterostructures without bulges at the heterointerface.

CONCLUSIONS

In conclusion, the mechanisms and kinetics of advanced axial Ge–Si nanowire heteroepitaxy based on the tailoring of the Au catalyst composition via TMGa exposure and the addition of dopants have been studied by environmental transmission electron microscopy combined with systematic ex situ CVD experiments. The morphology of the Ge–Si heterojunction, in particular, the extent of a local, pronounced bulge in the NW, is found to critically depend on CVD parameters, including the Ga composition of the catalyst, the TMGa exposure temperature, and the presence of dopants. Such pronounced NW bulging at a heterointerface has not been reported before, and more importantly, it cannot be rationalized by prior simple models for heteroepitaxy, such as considering only the balance of the relevant surface/interface energies. For our Ge–Si model system, we infer that the NW bulging is a physical manifestation of the Si supersaturation in the liquid AuGa catalyst. We hypothesize that the lowered solubility of Ge in the AuGa alloy, that is, the very aspect that produces an improved sharp Ge–Si transition, invokes a higher nucleation barrier for the heteroepitaxy based on the necessity of nucleating a more Si-rich Ge1−xSi0x on the GeNW. The presence of dopants (specifically boron) can in turn depress this nucleation barrier and consequently enable the growth of sharp Ge–Si heterojunctions without heterointerface deformities. Our results highlight how the kinetics of NW heteroepitaxy can be controlled through the catalyst composition and the addition of impurities. While this work explores the control of the influence of nucleation kinetics over diameter and bulge formation with respect to the formation of heterojunctions, it is not possible to directly address the issue of heterojunction abruptness without additional studies. Due to the generality of our arguments, we anticipate that our nucleation-based model for heteroepitaxy will be relevant to a wide range of material systems and device-enabling heterostructures.

METHODS

In Situ TEM. A modified Tecnai F20 ETEM operated at 200 kV with a differential pumping system and a digital video camera was used. Specimen heating was accomplished using a TEM holder mini furnace with a directly attached thermocouple to measure the growth temperature. The ETEM experiments began with Au-catalyzed Ge nanowires grown ex situ in a cold-walled chemical vapor deposition reactor at the Center for Integrated Nanotechnologies user facility at Los Alamos National Laboratory. The GeNWs were subsequently transferred via sonication in isopropyl alcohol from the Si(111) substrate and dispersed onto a 200 mesh Cu TEM grid coated with both a holey carbon film and a 30 nm sputtered SiO2 layer. The samples were then exposed to a heat lamp for 30 min in air before loading into the ETEM for in situ time-resolved imaging of Ge–Si heterojunction formation. To form the AuGa catalyst within the ETEM, the Au-catalyzed GeNWs were heated to 380 °C and exposed to an atmosphere of ≈10 mTorr of pure TMGa for 45 s. Ga droplets were visible on the sputtered SiO2 (not shown), confirming the addition of Ga to the system. After forming the AuGa alloy, the GeNWs with the AuGa catalyst were heated to ≈450 °C and exposed to ≈1 Torr of pure Si2H6 to reinitiate growth.
Nanowire Growth (Ex Situ)\(^{15}\). Both intrinsic Ge/Si axial heterostructured nanowires and boron-doped Ge/phosphorus-doped Si heterostructured nanowires were synthesized using GeH\(_4\), SiH\(_4\), B\(_2\)H\(_6\), and PH\(_3\) as the gas sources, respectively, and H\(_2\) as the carrier gas. Nanowire growth occurred on either Si(111) or Ge(111) substrates that were initially solvent-cleaned and native-oxide-etched prior to deposition of 30–50 nm Au colloid or e-beam-deposited Au. Boron-doped Ge(111) nanowire growth \((P_{\text{B2H6}}/P_{\text{GeH4}} = 1.6 \times 10^{-2})\) was initiated from Au by nucleation at 380 °C for 3 min followed by growth at 280 °C for 90 min. In situ catalyst alloying was accomplished during growth by ramping from 280 to 380 °C with GeH\(_4\) flow followed by simultaneously flowing vapor-phase trimethylgallium (TMGa) from a bubbler line and GeH\(_4\) B\(_2\)H\(_6\) was not flowing. The germanium nanowire growth at 380 °C results in unintentional vapor–solid (VS) sidewall growth leading to a marked increase in diameter and a tapered structure near the junction. The phosphorus-doped Si segment of the heterostructure was grown after Au, \(_{1-x}\)Ga\(_x\) alloy formation by removing both TMGa and GeH\(_4\) and introducing SiH\(_4\) and PH\(_3\) \((P_{\text{SiH4}}/P_{\text{PH3}} = 6.6 \times 10^{-3})\), while the temperature was ramped from 380 to 495 °C. A total pressure of 2 Torr was maintained throughout the Ge growth and alloy formation, while the total pressure of 0.5 Torr was maintained for the Si growth. Undoped Ge/Si heterostructures were also grown using the same procedure as for the doped wires with the exception that the GeNWs were grown at 380 °C. We note the possibility of Ga incorporation into the NW during growth analogous to what has recently been reported for pure Al catalysts\(^{17}\), which could create a p-type region in the Si segment. Such possible minor Ga incorporation and related doping effects are not addressed here but left as subjects for further studies. Ex situ TEM imaging was performed on nanowires transferred to either copper grids with lacey carbon or to Si pillars using a dual-beam focused ion beam/scanning electron microscope.

Conflict of Interest: The authors declare no competing financial interest.

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Supporting Information Available: Bright-field, 9 fps ETEM video showing Ge–Si NW heteroepitaxy (corresponding to Figure 2A–F) and additional post-growth TEM analysis of Ge–Si heterostructure shown in Figure 2G. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES AND NOTES

24. The highest Ga composition we could achieve in the alloy catalyst was 34% Ga. Further exposure to TMGa does not lead to higher Ga compositions in the catalyst.


31. We do not speculate here about the detailed nucleation site(s) of step flow across the catalyst–NW interface, which remains experimentally undetermined.


36. Further discussion of the APT data in doped Ge–Si heterostructure NWs will be left for a future publication.
