Supercooling of nanoscale Ga drops with controlled impurity levels

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(Received 7 October 2011; published 23 November 2011)

We use in situ observations by variable temperature transmission electron microscopy on Ga drops at the tips of GaAs nanowires to investigate the phase behavior of nanoscale Ga. Experiments on pure Ga drops are compared with drops containing well-defined levels of impurities. Our controlled experiments show that the crystallization temperature, and hence the ultimate achievable supercooling, strongly depends on the concentration of impurities. All drops show predominant β- and γ-Ga correlations in the liquid phase and ultimately crystallize to solid β- and γ-Ga, which provides support for a scenario in which impurities limit the achievable supercooling without significantly templating the crystalline phase.

DOI: 10.1103/PhysRevB.84.193303 PACS number(s): 64.70.dg

Melting and crystallization are fundamental processes by which most substances change between a disordered liquid state and an ordered solid state. When a liquid droplet is cooled, it reaches a point at which it transforms into a solid. This point is not necessarily the equilibrium melting point. The possibility of significant supercooling in small droplets has been investigated extensively following the pioneering work of Turnbull.1 It has been shown that small particles can be achieved in small particles.14 However, the Ga drops undergo significantly higher supercooling,4 and crystallize along different pathways5 and in structures different from the bulk.5–11 The extent of achievable supercooling can presumably vary with droplet size, purity,12 interactions with supports, etc. For most substances, the supercooling is limited to \( \Delta T / T_M \sim 0.1–0.2 \), where \( \Delta T = T_M - T_C \) is the difference between the melting and the crystallization temperatures.4 Ga is a particular metal for which this value can be exceeded significantly; thus, Ga droplets have become a model system for studying deep supercooled liquids. Ga droplets can be supercooled to half of the melting temperature, \( \Delta T / T_M \sim 0.5 \), as demonstrated in different experiments on micro- and nanometer-sized Ga in emulsions13 and films of supported Ga droplets.8,14 Most investigations considered small encapsulated or confined Ga particles and thus were affected by several factors simultaneously: the effects of confined geometry and interactions with the confining matrix,2,9,15 uncontrolled levels of impurities, nonuniform droplet sizes, etc. An investigation of supercooled liquids and their intrinsic crystallization pathway requires eliminating heterogeneous crystallization, e.g., due to contact with confining matrices and foreign materials. Limited ensemble measurements on films of supported Ga droplets indicate that very high-supercooling may be achieved in small particles.14 However, the Ga drops in these films were polydisperse and covered with SiO₂, and an ensemble measurement would not capture possible effects of particle size, shape, and support/matrix interactions on the highest achievable supercooling and crystallization pathway.

Here we investigate the crystallization and melting of Ga drops at the tips of GaAs nanowires (NWs). This particular geometry avoids any matrix effects and limits the interactions between liquid the Ga nanodroplets and the support surface to a well-defined solid–liquid interface. The use of in situ, variable temperature transmission electron microscopy (TEM) allows us to follow the phase behavior of individual Ga droplets among a large ensemble of similar NWs/drops, confirming the generality of the observations on single drops. This unprecedented level of control allows us to perform identical experiments on pure Ga nanodroplets and on liquids containing different levels of impurities and thus identify the effect of well-defined impurity concentrations on the achievable supercooling and crystallization of nanoscale Ga.

Our variable temperature in situ experiments were carried out in a JEOL 2100F field emission TEM equipped with a Gatan liquid He cooling sample holder in the temperature range between 380 and 10 K, at pressures below 2 × 10⁻⁷ torr, and at low electron irradiation intensities (<0.1 A/cm²). Specimen temperatures were measured by a thermocouple, calibrated via the melting points of In and eutectic Au-Ge,10,16,17 and confirmed here via the melting temperature of Ga. Heating and cooling rates were slow (~5 K/min) to ensure efficient heat transfers and equilibration. The Ga/GaAs NW structures were synthesized by molecular beam epitaxy as described in detail elsewhere.18 Briefly, Ga-catalyzed growth was carried out at 630 °C on GaAs substrates covered with SiO₂. P-type doping of the NWs and different levels of Si in the Ga drops were achieved with the addition of a controlled silicon flux during the NW growth. In situ TEM experiments were performed with Si-doped Ga droplets obtained using Si fluxes of 2.2 × 10¹⁰ and 1.6 × 10¹¹ Si/cm² s, resulting in measured silicon concentrations in the NWs of 5.5 × 10¹⁸ and 4 × 10¹⁹ cm⁻³, respectively.18,19 The Si concentration in the Ga droplets \( c_{Si(Ga)} \) is calculated from that of the NW \( c_{Si(NW)} \) via the distribution coefficient \( k = c_{Si(NW)}/c_{Si(Ga)} \), which is well established for the Si-Ga(liquid)/GaAs(solid) system.18,20 Using values obtained in liquid phase epitaxy for our two doping levels gives Si concentrations in the liquid Ga droplets \( c_{Si(Ga)} \) of 0.06 and 0.47 at.%, which we later refer to as low- and high-doped Ga droplets, respectively. The NWs were transferred through air and dispersed on amorphous C films supported by Au grids. A thin amorphous oxide layer, initially visible on the Ga surface, desorbed during annealing of the samples to 450 K under the electron beam in the TEM.21

Figure 1(a) shows a characteristic TEM image of an undoped GaAs NW with a Ga drop at the tip. The striped
FIG. 1. (Color online) Crystallization and melting of pure Ga drops at the tips of GaAs NWs. Crystallization of a Ga drop: (a) TEM image of the GaAs NW with a Ga drop at its tip during cooling from room temperature \((T = 94 \text{ K})\). (b) DP from the Ga drop at 94 K, exhibiting diffuse rings. (c) Simulated DP for a liquid with a \(\beta\)-Ga short-range structure. (d) TEM image of the single-crystalline \(\beta\)-Ga particle at 12 K and (e) the corresponding DP along the \([-101]\) zone axis. Melting of the Ga particle: (f) TEM image and (g) DP of the Ga drop after melting \((T = 260 \text{ K})\) during heating from 12 K.

contrast of the GaAs NW stems from its morphology, consisting of a sequence of segments with wurtzite and zincblende structure. The Ga drops show darker contrast than the GaAs wires and appear as homogeneous spheres without any internal contrast at the NW tips. The NWs are initially heated to 350 K and then cooled slowly to 85 K while recording TEM images and electron diffraction patterns (DPs). The TEM image in Fig. 1(a), obtained at 94 K, is representative of the Ga drops over the entire range from 350 to 85 K. DPs over this temperature range [Fig. 1(b)] consist of diffuse rings that indicate either a liquid or a glassy amorphous phase of the Ga drop. Further cooling of the drops to 12 K, using liquid He, results in crystallization of the Ga drop [Figs. 1(d) and 1(e)] at a temperature between 85 and 12 K. In the TEM image [Fig. 1(d)], a change of contrast in the drop and facets on its surface are seen. The DP of the solidified Ga drop [Fig. 1(e)] indicates that it is single crystalline and can be indexed to the metastable \(\beta\)-Ga phase with a monoclinic structure. For our pure Ga drops at the tips of the GaAs NWs, a crystallization temperature below 84 K implies a supercooling of at least \(\Delta T/T_M \sim 0.7\) \((T_M^{\beta\text{-Ga}} = 256 \text{ K}, \text{as explained later})\). Investigation of a large number of crystallized drops shows that the resulting solid Ga nanoparticles are mostly monocristalline and adopt either the metastable \(\beta\)- or the metastable \(\gamma\)-Ga phase. Some particles are polycrystalline. Ga drops on planar surfaces and Ga encapsulated in carbon nanotubes were found to crystallize in the same two metastable Ga phases that can even coexist in the same particle.

Following crystallization at \(T < 84 \text{ K}\), the temperature is increased and the Ga particles are heated slowly to room temperature. Melting generally occurs \(\sim 240 \text{ K}\). Upon melting, the Ga drops recover their spherical shape with homogeneous contrast [Fig. 1(f)]. The DPs lose the discrete Bragg spots and again consist of diffuse rings, characteristic of the liquid state [Fig. 1(g)]. The DPs after melting are identical to the DPs of the initial drops before crystallization and closely match the simulated DP for a liquid Ga drop with \(\beta\)-Ga–like short-range correlations [Fig. 1(b)]. Real-space imaging never shows the characteristic contrast associated with an amorphous solid phase during heating or cooling cycles; hence, we conclude that the Ga particles change directly between liquid and crystalline solid phases.

To investigate the effect of controlled concentrations of impurities on the supercooling, crystallization, and melting, Ga drops with two different Si concentrations, 0.06 and 0.47 at.%, are subjected to identical experiments. Even such small impurity concentrations influence significantly the crystallization temperatures of the Ga drops. Figure 2 shows variable temperature TEM images of highly doped Ga:Si drops with impurity levels of 0.47 at.%. The drops are originally liquid [Figs. 2(a) and 2(b)], with DPs in the liquid state identical to those of our pure Ga drops. Upon decreasing the temperature, crystallization occurs much sooner, i.e., at significantly higher temperatures than in the pure Ga drops. Ga drops with 0.47 at.% Si content crystallize at 155 \(\pm 5 \text{ K}\), whereas drops containing 0.06 at.% Si crystallize at 144 \(\pm 5 \text{ K}\) Fig. 3. In repeated experiments, the crystallization temperature \(T_C\) was determined by lowering \(T\) in 5-K steps, allowing the sample to equilibrate, and surveying an ensemble of Ga drops. Generally, all drops are still liquid at 5 K above the stated \(T_C\), whereas all are crystalline at 5 K below the \(T_C\), thus giving an error bar of \(\pm 5 \text{ K}\). The melting temperatures \(T_M\) were determined analogously. A survey of a large number of drops...
shows the crystalline Ga particles again single crystalline or polycrystalline, in either the metastable $\beta$- or the metastable $\gamma$-Ga phase. For instance, the Ga drop shown in Fig. 2 crystallizes as $\gamma$-Ga [Fig. 2(e) and 2(f)]. Upon increasing the temperature, all Ga particles (independent of Si content) melt in the interval between 230 and 240 K. Figure 2(g), a TEM image at 230 K, shows the melted Ga drops, with a DP of diffuse rings in Fig. 2(h).

Our experiments on Ga drops with controlled impurity levels show that the concentration of Si impurities affects primarily the crystallization temperature of the Ga drops but does not lead to systematic changes in the melting temperature of the solid Ga particles Fig. 3. Hence, the degree of supercooling $\Delta T / T_M$ that can be achieved depends sensitively on the impurity level of the melt, varying from $\Delta T / T_M \sim 0.7$ for pure Ga drops to $\Delta T / T_M \sim 0.35$ for drops with the highest Si content. The observation of melting temperatures only marginally below the melting temperatures of the bulklike metastable $\beta$- and $\gamma$-Ga phases is consistent with the notion that melting, initiated as surface premelting, should not be affected strongly by size, support, and purity.\textsuperscript{16,24}

The observed differences in supercooling between pure and doped Ga drops imply that the ultimate supercooling in the doped drops is limited by the presence of impurities. What, then, determines the achievable supercooling of the pure Ga melt? The NW–drop interface could provide a template for heterogeneous nucleation of the crystalline phase in the drop. However, several factors argue against this scenario. First, the facet at the NW tip, which can have either a wurtzite or a zincblende structure, does not induce a particular orientation of the crystallized Ga particles. Analyzing the DPs of many particles after crystallization, we detected no particular alignment of the Ga lattice relative to the NW. In some cases, the particles are even polycrystalline after freezing, suggesting several crystallization centers. Second, to further investigate possible support effects, some Ga drops were severed from the tips of the GaAs NWs and distributed on the amorphous C films of the TEM grid Fig. 4. Such amorphous C-supported drops show crystallization and melting identical to the drops held at the NW tips. In particular, the freezing and melting temperatures, and hence the extent of supercooling, are the same as for NW-supported drops, suggesting that the observed

![FIG. 2.](image)

**FIG. 2.** (Color online) Crystallization and melting of highly Si-doped Ga drops at the tips of GaAs NWs. Crystallization of the Ga drop: (a) TEM image of the GaAs NW and the Ga drop at its tip at room temperature. (b) DP from the Ga drop at room temperature exhibiting diffuse rings, characteristic of a liquid. (c) TEM image of and (d) DP from the Ga drop during brief heating to 350 K, prior to cooling. (e) TEM image of the single crystalline $\gamma$-Ga particle at 150 K and (f) the corresponding DP along the [110] zone axis. Melting of the Ga particle: (g) TEM image and (h) DP of the Ga drop after melting ($T = 230$ K) during heating from 90 K.

![FIG. 3.](image)

**FIG. 3.** (Color online) Dependence of the melting $T_M$ and crystallization $T_C$ temperature of Ga drops at the tips of GaAs NWs on the Si impurity concentration.
differences in supercooling are robust and independent of the support. Thus, the supercooling of our pure Ga drops is determined by heterogeneous nucleation involving a very low level of residual impurities, by homogeneous nucleation, or by surface-induced crystallization.

Both pure and Si-doped Ga drops—at the NW tips, as well as on amorphous C—variably crystallize in either the β- or the γ-Ga metastable phases but not in the stable bulk phase α-Ga. As the crystal structure and orientation are not determined by the support interface, it is possible that short-range correlations in the Ga liquid provide a template for the crystalline phase, especially for the pure Ga drops that show extreme supercooling. A diffraction analysis on individual liquid drops is used to investigate this possibility. Figure 5(a) shows the radial distributions of the diffracted intensity of pure and doped Ga drops at two temperatures. The principal feature is a peak with a shoulder toward larger wave vectors, consistent with x-ray DPs of liquid Ga. The diffraction features do not change significantly with temperature or purity, except for a slight thermal (Debye-Waller) broadening at higher temperatures. Figure 5(b) shows calculated structure factors of the α-, β-, and γ-Ga phases, computed with an ab initio powder pattern simulation as implemented in the software package JEMS. Both β- and γ-Ga have structure factors nearly identical to the experimentally observed diffraction intensities, in particular the characteristic primary peak ~4 nm^-1 and a shoulder toward higher spatial frequencies. α-Ga is clearly different and shows a minimum at 4 nm^-1 with peaks on either side. If α-Ga correlations exist in our Ga drops, Fig. 5 shows that they clearly constitute a minority structure, with the majority being β- and γ-Ga–like. The short-range structural motifs observed in the liquid Ga drops in our experiments could act as precursors of the crystalline β- and γ-Ga phases. All of our drops—irrespective of impurities—show β- and γ-Ga correlations in the liquid phase and ultimately crystallize to solid β- and γ-Ga, which provides strong support for a scenario in which the crystalline phase is templated by a short-range order in the liquid phase and not by external factors such as a particular support structure (both GaAs and amorphous C supports give the same results) or a nucleation center involving impurities in the bulk or on the surface. We can thus conclude that impurities in the melt (e.g., Si in liquid Ga) can constitute crystallization centers that limit the achievable supercooling, while the structure of the incipient solid is determined by the particular structural motifs of the liquid metal itself. Our results show strong similarities with previous work on Ti-Zr-Ni alloys, which concluded that the icosahedral short-range order in the supercooled liquid is responsible for the nucleation of a metastable crystalline icosahedral (quasicrystal) phase instead of the stable solid phase. While this previous work supported Frank’s hypothesis that the icosahedral short-range order stabilizes the liquid phase and promotes deep supercooling, our results on deep supercooled elemental Ga suggest that other structural motifs in the liquid—similar to the structure of the metastable β- and γ-Ga—may play an analogous role. This raises the possibility that short-range motifs other than the accepted icosahedral one (and perhaps especially those close to a metastable structure of the solid) can stabilize deep supercooled liquids.
This work was performed at the Center for Functional Nanomaterials under the auspices of the US Department of Energy, under Contract No. DE-AC02-98CH10886. A.F.iM. thanks G. Abstreiter and M. Bichler for experimental support, as well as funding through the Marie Curie Excellence Grant project Semiconductor Nanowires and Their Field Effect Devices and the Swiss National Science Foundation via Grant No. 2000021-121758.

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