We report the first direct observation of gas-phase homogeneous nucleation of GaAs by the thermophoretic collection of an aerosol of nanometer-scale microcrystallite GaAs clusters in an atmospheric organometallic vapor-phase epitaxy (OMVPE) reactor. In addition to shedding light on a fundamental question about the reaction kinetics involved in OMVPE, this is the first report of gas-phase synthesis of nanometer-scale III/V clusters from volatile precursor molecules. We have determined conditions under which nearly defect free, 10-20 nm scale GaAs microcrystallites form. These particles are in the quantum size regime which suggests that the process may be used to form optically active quantum dots. Thus gas-phase homogeneous nucleation in OMVPE, which has been regarded until now as a troublesome parasitic reaction, may ultimately form the basis of an aerosol technology for the fabrication of novel optoelectronic devices such as a quantum dot laser.

The reactor used in these experiments is depicted in Fig. 1. A 34 mm diam, 700 mm long quartz process tube is mounted in a multizone furnace. Trimethyl gallium (TMG) was introduced by bubbling hydrogen through a liquid source held at -10 °C. Heated injectors (Z1, Z2) permit TMG and 10% AsH3 in hydrogen to be introduced into the process tube separately, allowing for the possibility of precracking the precursor molecules. Alternatively, the reactants can be mixed prior to injection by rerouting the arsine flow as shown in Fig. 1. The nucleation zone, Z3, is 18 cm long. Aerosol particles are collected thermophoretically on a holey carbon film which is mounted on a stainless steel fixture aligned coaxially in the collection zone, Z5, of the process tube (Fig. 1). Z5 is held at a temperature of 350 °C in all experiments to prevent the condensation of arsenic on the collection apparatus. Hydrogen, purified with a resin filter, is used as the carrier gas. The composition of the gas exiting the process tube is monitored with a quadrupole mass spectrometer (QMS).

In a typical experiment the TMG and AsH3 are preheated to 400 °C in Z1 and Z2, respectively, before being introduced into the reaction zone. At this temperature negligible pyrolysis of the TMG or AsH3 occurs as previously determined by infrared absorption spectroscopy and mass spectroscopic analysis. This is verified by negligible decreases in the arsine partial pressure and no CH4 partial pressure detectable on the QMS. The total flow is 2 SLM, with the partial pressures of TMG and arsine typically 1.25×10⁻⁴ and 6.25×10⁻⁵ atm, respectively. For these flow conditions, the residence time in the reaction zone (Z3) is 5 s.

When the temperature in the nucleation zone (Z3) is held at 700 °C, solid GaAs deposits on the reactor walls within the first 10 cm of the nucleation zone, indicating that the TMG is entirely consumed within approximately 3 s of entering Z3. Downstream of this point, in Z5, a semitransparent yellow deposit is observed on the walls of the reactor tube and on the collection fixture. A transmission electron microscopy (TEM) micrograph of the collected deposit reveals it to consist of individual GaAs microcrystallites and some chainlike agglomerates (Fig. 2). The electron diffraction pattern of the particles, shown in the inset of Fig. 2, indexes to cubic zincblende structure, as shown in Fig. 3.

Figure 4 shows a high resolution transmission electron micrograph (HRTEM) of an agglomerate of smaller primary particles. In practice, HRTEM is easier to perform on the agglomerated clusters than on individual particles, owing to the fact that the agglomerates are better able to dissipate excess charge and heat induced by the electron beam than are individual particles. Within several of the clusters shown, lattice fringes and faceting indicative of (111) bounding planes are clearly evident. Some of the crystalline faces have as few as 30 atomic rows. The lattice fringes on the particles extend to within 1 nm of the cluster surface, consistent with the formation of an amorphous oxide layer on the particles on exposure to air.

We have investigated the role of growth temperature and reactant concentration on the nucleation and morphology of the clusters. By analyzing low magnification micrographs, such as Fig. 2, we have measured the size distributions of the primary particles for runs at different growth temperatures in the nucleation zone (Z3). We find that the mean particle diameter is strongly influenced by the growth temperature. By fitting the measured particle size distributions to a log-normal distribution, a median particle diameter \( D_{\text{med}} \) and geometric standard deviation \( \sigma_g \) for the distribution are found. Geometric standard deviations...
FIG. 1. Schematic of the experimental apparatus for gas-phase synthesis of nanometer-scale GaAs clusters. The group III precursor (TMG) is introduced into the hot wall reactor by bubbling H₂ carrier gas through a liquid source. The group V precursor, AsH₃, can be mixed with the TMG outside the furnace (at room temperature) or inside the furnace (at elevated temperature). All flows are regulated by mass flow controllers (MFCs). The process runs at atmospheric pressure. A quadrupole mass spectrometer, connected to the exhaust of the process tube by a sampling line, is used to monitor reactant concentrations.

are in the range σₕ = 1.2–1.3. A typical data set and its fit are shown in Fig. 5(a). Figure 5(b) shows the behavior of the median crystallite diameter and σₙ as a function of reciprocal temperature. As the temperature in the nucleation zone (Z3) is increased from 600 °C (1000/T = 1.14 K⁻¹) to 800 °C (1000/T = 0.93 K⁻¹), the median particle diameter rises continuously from 12 to 19 nm. Doubling the input TMG and AsH₃ concentrations relative to the conditions of Figs. 2 and 3, the mean particle diameter is increased by approximately 10% with no noticeable change in particle morphology. Additionally, we have generated clusters by mixing TMG and arsine both at room temperature and at 400 °C and find no difference in particle morphology.

Several important questions remain regarding the growth mechanism of the primary GaAs clusters. Gas-phase infrared absorption measurements showing the presence of TMG—AsH₃ adducts, suggest that GaAs may form homogeneously, nucleating (GaAs)ₙ clusters with n...
The present aerosol technique has several significant advantages over previous methods for producing nanometer-scale GaAs clusters. For example, synthesis in quinoline solution produces GaAs clusters with significant impurities and unequal Ga and As abundances, while GaAs molecular beam epitaxy on silica substrates produces irregular particles in intimate contact with a foreign substrate. Synthesis of nanometer-scale GaAs clusters by homogeneous nucleation from an exploding wire vapor has been recently demonstrated. However, we feel that the process described here is more readily compatible with the incorporation of the GaAs clusters in a passivating medium of AlGaAs both in the aerosol phase and ultimately in a growing epitaxial structure.

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FIG. 5. Analysis of the size distribution of GaAs clusters produced at various temperatures. (a) Representative data for a run with growth temperature 600 °C, plotted as a histogram $\Delta N/\Delta \log D_p$ vs $D_p$, the particle diameter. The solid line is a best fit to a log-normal distribution. (b) Semilog plot of median particle diameter $D_{50}$ vs reciprocal temperature. The geometric standard deviation $\sigma_g$ determined for each point is represented on the plot as a vertical bar of height $2 \sigma_g$.

in the range 10–20. These clusters may then increase to the 10 nm size range by coagulation. Subsequent grain boundary migration through the sintered particle may consolidate the agglomerate into the single crystal, faceted clusters seen in Figs. 2 and 3. Alternatively, further growth to the observed sizes in the 10 nm range may proceed by epitaxial deposition from the supersaturated vapor phase. Since the microscopic crystal planes of the clusters are much smaller than surface diffusion lengths, adsorbed Ga and As atoms can diffuse to fast growing crystal planes, leaving the cluster bounded by slow growth planes as observed in Fig. 4. With the present data, it is impossible to distinguish between these two mechanisms, although the particle size distributions are somewhat narrower than would be expected if coagulation were the dominant growth mechanism ($\sigma_g \approx 1.46$). It is also possible that GaAs cluster growth occurs through a two step process by the reaction of nanometer-scale droplets of Ga or polynorbornyl gallium polymer with the AsH$_3$ ambient. Experiments designed to distinguish among these possible mechanisms are the focus of continuing work.