Colloidal Phase-Change Materials: Synthesis of Monodisperse GeTe Nanoparticles and Quantification of Their Size-Dependent Crystallization

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ABSTRACT: Phase-change memory materials refer to a class of materials that can exist in amorphous and crystalline phases with distinctly different electrical or optical properties, as well as exhibit outstanding crystallization kinetics and optimal phase transition temperatures. This paper focuses on the potential of colloids as phase-change memory materials. We report a novel synthesis for amorphous GeTe nanoparticles based on an amide-promoted approach that enables accurate size control of GeTe nanoparticles between 4 and 9 nm, narrow size distributions down to 9−10%, and synthesis upsizing to reach multigram chemical yields per batch. We then quantify the crystallization phase transition for GeTe nanoparticles, employing high-temperature X-ray diffraction, differential scanning calorimetry, and transmission electron microscopy. We show that GeTe nanoparticles crystallize at higher temperatures than the bulk GeTe material and that crystallization temperature increases with decreasing size. We can explain this size-dependence using the entropy of crystallization model and classical nucleation theory. The size-dependences quantified here highlight possible benefits of nanoparticles for phase-change memory applications.

INTRODUCTION

Phase-change memory is a transistor-free data storage technology that leverages crystallization and melting phase transitions, using the resistivity contrast between the amorphous and crystalline phases of the material as the digital 0 and 1.1,2 A miniaturization of phase-change memory chips will ultimately bring the size of memory cells to sub-10 nm regime, where the phase transition temperatures become a function of size.3 Colloidal nanoparticles are a convenient material system to study size-dependent phase transitions4,5 due to their fast synthesis, providing monodisperse template-free nanoparticles of a specific size.6 Furthermore, colloidal nanoparticles may themselves be used as the building blocks for future phase change memory cells.

This paper focuses on Germanium(II) telluride, GeTe, a prototype phase-change material, which is well studied in bulk and thin films. GeTe offers distinct crystallization and melting temperatures (T_{cryst,bulk} = 170 °C; T_{melt,bulk} = 725 °C),3,5 pronounced resistivity contrast between amorphous and crystalline phases (R_{amorph}/R_{cryst} > 1000),6 and suitable crystallization kinetics (i.e., fast at high temperatures and slow at room temperature, k_{130 °C} ≈ 1 m/s, k_{30 °C} ≪ 1 nm/year).9 GeTe and its ternary alloys are also of interest for other applications, such as ferroelectric and multiferroic, thermoelectric, and infrared semiconductor applications.10,11

Synthesis of colloidal GeTe nanoparticles is not well developed. Depending on the reaction conditions, GeTe nanoparticles can exhibit either an amorphous or crystalline structure upon synthesis. Amorphous GeTe nanoparticles were reported by Caldwell et al. (sizes from 1.8 to 3.4 nm in diameter)12 and by Arachchige et al. (sizes ≥ 8.7 nm in diameter).13 Crystalline GeTe nanoparticles are generally larger in size, and have been reported as 8 and 17 nm dots,14 100 nm octahedrons, or 350 nm cubes.15,16 Since we aim to study the effect of nanoparticle size on the crystallization phase transition, here we wish to obtain amorphous GeTe nanoparticles.

Crystallization temperature of GeTe nanoparticles is known to deviate from the bulk value of 170 °C, however, the size dependent crystallization behavior of colloidal nanoparticles has never been quantitatively explained. Caldwell et al.12 and Arachchige et al.13 observed a general trend—crystallization
temperature gradually increases for smaller GeTe nanoparticles, reaching up to 400 °C for 1.8 nm GeTe clusters. Part of the challenge in explaining the size-dependence of the crystallization temperature comes from the fact that it is opposite to the trend in melting point, which decreases with decreasing nanoparticle size and which is explained by a higher energy per atom due to the increasing number of surface atoms as nanoparticle size decreases.17,18 Furthermore, coalescence of nanoparticles occurs near the crystallization temperatures, and most experimentation methodologies do not separate crystallization and coalescence phenomena.12,13

In this paper, we report a new one-step synthesis of amorphous GeTe nanoparticles with accurate size control between 4 and 9 nm, which closes the size gap between previous reports.12,13 We then use these nanoparticles to perform the first quantitative study of the size-dependent crystallization of template-free GeTe nanoparticles. To separate coalescence and crystallization, we use several complementary methods, such as constant heating ramp X-ray diffraction, differential scanning calorimetry, and ex-situ heating transmission electron microscopy. We then explain the experimentally observed size-dependence of GeTe crystallization with thermodynamic model and classical nucleation theory. Finally, with our results, we calculate the temperature window, reduced crystallization temperature, and the power consumption to evaluate how size effects can be used to tune the properties and performance of phase-change memory devices.

**EXPERIMENTAL SECTION**

**Materials.** Gel2 (99.99%) and Na2Te (99.9%) were purchased from ABCR, tri-octylphosphine (TOP, 97%), Te (broken ingots, 99.999%), and K3S (95%) from STREM, oleic acid (90%), chloroform (99%), ethanol (99.8%), n-butylamine (99.95%), N,N-dimethylethanolamide (DMF, 99.8%), hexane (99.5%), toluene (99.8%), and LiF (99.9%) from Sigma-Aldrich, LiN(SiMe3)2 (95%) from Acros Organics. Oleic acid was dried at 100 °C for 1 h from water residues and all other chemicals were of anhydrous grade and were used as received.

**Synthesis of GeTe Nanoparticles.** In a typical synthesis of 7 nm GeTe nanoparticles, anhydrous Gel2 (107 mg, 0.33 mmol) was dissolved in tri-octylphosphine (TOP; 7.5 mL) in the glovebox and transferred to the prepumped reaction flask, connected to standard vacuum manifold. This mixture was additionally purified under vacuum at 100 °C for 30 min, after which it was filled with N2 and heated up to injection temperature of 280 °C. Meanwhile, an injection mixture was prepared by mixing two TOP-based stock solutions—0.8 mL of 1 M TOP:Te (0.8 mmol of Te, in total) and 0.5 mL of 1.6 M LiN(SiMe3)2 (0.8 mmol of amide, in total)—and swiftly added to the reaction mixture as soon as its temperature reached injection set value. After injection, the temperature of the reaction mixture drops by about 20 °C and was kept as such for another 1–2 min. The solution changes its color to deep brown during this time and reaction was terminated by rapidly cooling the flask with pressurized air and, later, with cold water bath. Once cooled to room temperature, the crude solution of GeTe nanoparticles was transferred air-free to the glovebox, where anhydrous chloroform (10 mL) and dried oleic acid (1 mL) were added to it. The mixture was shaken rigorously and set aside for 5–10 min, allowing for the completion of oleate shell formation. The oleate-covered GeTe nanoparticles were precipitated with anhydrous ethanol (30 mL) and separated by centrifugation at 6000 g for 5 min. The obtained pellet of GeTe nanoparticles was dispersed in anhydrous chloroform, forming a long-term stable colloidal solution.

The size of GeTe nanoparticles can be controlled by the amount of amide salt and by the injection temperature, whereas the selection of growth time between 0.5 and 5 min has a relatively minor effect. The 50X upscaled synthesis of GeTe nanoparticles follows the same procedures as explained above, while using underpressure-governed hot-injection technique for fast addition of injection mixture.59

**Ligand Exchange Process or GeTe Nanoparticles.** In analogy to previously published protocols, several inorganic salts (LiI, Gel2, K3S, or Na2Te) were used to remove initial organic ligands (oleic acid) from the surface of the GeTe nanoparticles.20,21 In a typical ligand-exchange process using Gel2, a starting solution of GeTe nanoparticles (1 mL, approximately 6 mg/mL), was precipitated with ethanol (3 mL), centrifuged at 9000 g for 3 min, and dispersed in toluene (200 μL). This solution was mixed with Gel2/DMF solution (3 mL, 20 mg/mL), and 15 mL of hexane, forming a 2-phase solvent system. The solvent mixture was rigorously shaken for few minutes, during which GeTe nanoparticles transfer to DMF phase. The hexane phase was then decanted and the GeTe/DMF solution was purified 4 more times with hexane to ensure complete removal of oleic acid. In addition, after the second washing step, 3 mL of DMF was added, to ensure that nanoparticles do not cluster. Afterward, Gel2-covered GeTe nanoparticles were precipitated with chloroform (1:3 v:v), centrifuged at 9000 g for 3 min, and redissolved in n-butylamine (0.2 mL).

**High-temperature X-ray diffraction (XRD).** GeTe nanoparticles were precipitated with ethanol and centrifuged. GeTe pellets were mixed with a heat-stable polymer to spatially separate GeTe nanoparticles, and loaded onto an inconel holder. High-temperature XRD was performed on Rigaku SmartLab 9 kW System, equipped with rotating Cu anode and 2D solid state detector (HyPix-3000 SL). High-temperature stage (Anton Paar) consists of a ceramic plate, which is shielded with a carbon dome. The high-temperature XRD measurements were performed under constant heating ramp and with the temperature precision of ±1 °C. All nanoparticle handling and measurements were carried out under N2 atmosphere.

**Differential Scanning Calorimetry (DSC).** Thermal analysis by DSC was carried out using a Mettler Toledo DSC822e instrument calibrated against indium standards. All measurements were performed under N2 blanket. Freshly prepared GeTe nanoparticle dispersions in chloroform were added into aluminum crucibles and immediately heated to 90 °C for 5 min to evaporate the residual solvent, following which the samples were thermally cycled in the −40 to 350 °C range at different heating rates.

**Electron Microscopy.** Energy dispersive X-ray (EDX) spectroscopy was performed on an FEI Quanta 200 SEM microscope (30 kV). Transmission electron microscopy (TEM) images were taken by an FEI Tecnai F30 transmission electron microscope (300 kV). Ex-situ heating TEM analysis was carried out on a series of TEM grids, each heated to a specific temperature for 1 min under N2 atm.

**Fourier-Transform Infrared (FTIR) Spectroscopy.** GeTe nanoparticles were drop-cast on ZnSe transparent windows and measurements were performed on a Bruker Vertex 70 system with a ThGaAs detector.

**Electrical Characterization.** Gel2-covered GeTe nanoparticles in n-butylamine were filtered through 0.2 μm PTFE filter and spin-coated on interdigitated Au electrode structures. Each interdigitated electrode pair consists of 10 electrode fingers, approximately 2.1 mm long, 57 μm wide, and spaced 36 μm apart. Electrode characterization was done using a Keithley 2400 SMU, with voltage double sweeps spanning from −1 to 1 V in 81 steps, with 100 mA current compliance. Electrical characterization of GeTe nanoparticle thin films was done in an inert atmosphere (N2-filled glovebox).

**RESULTS**

**Synthesis.** To prepare monodisperse GeTe nanoparticles, we employ an amide-promoted synthetic approach, which was previously applied to binary and ternary chalcogenide nanocrystals.22–24 An amide salt, injected to the reaction mixture along with the chalcogen precursor, increases the nucleation rate and thus yields smaller sizes of nanoparticles, if compared to analogous reaction conditions without an
addition of amide promoter (Figure 1a). This method is especially useful for telluride nanoparticles because it can effectively separate nucleation and growth of nanoparticles, which is challenging when highly reactive tellurium precursors are employed. Importantly, the amount of amide can be used as a handle for size control—an increase of amide concentration leads to higher nucleation rates and hence smaller sizes of nanoparticles. By tuning the amount of coinjected amide salt, we achieve GeTe nanoparticles sizes between 4 and 8 nm (Figure 1b). In accordance with previous reports, the amide concentration has to be higher than that of the initial iodide anions, enabling full conversion to the germanium-amide intermediate followed by its fast reaction with the chalcogen (tellurium) precursor. Small amounts of LiN(SiMe₃)₂ result in slower nucleation on fewer nucleation centers, leaving sufficient amounts of starting precursors for direct reaction and yielding broadly dispersed large-size GeTe nanoparticles. Figure 1c illustrates these findings, showing a series of TEM images of GeTe products, prepared with different amounts of LiN(SiMe₃)₂ and otherwise identical reaction conditions. In particular, amide-promoted synthesis yields monodisperse GeTe nanoparticles with size distributions as narrow as 9% (Figure S1 of the Supporting Information, SI). Additional size tunability can be attained by regulating the injection temperature. The temperature dependence follows classical kinetics—lower injection (and growth) temperatures result in slower growth and thus smaller GeTe nanoparticles (Figure S2). Changing the growth time has only a minor effect on the size and size distribution of GeTe nanoparticles (Figure S3), which suggests fast completion of the reaction and high chemical yields of the process. The synthesis of GeTe nanoparticles is described in the Experimental Section in more detail.

As-synthesized small GeTe nanoparticles exhibit an amorphous (glass-like) structure, which is indicated by absence of diffraction contrast on TEM images (Figures S2 and S3) and, as shown later, by X-ray diffraction spectra and high-resolution TEM images. From EDX we find that GeTe nanoparticles are close to stoichiometric, with 1:1 Ge:Te atomic ratio, suggesting that the reaction system is well balanced and prevents oxidation of Ge²⁺ ions to its more stable Ge⁴⁺ oxidation state. A typical EDX spectrum of GeTe nanoparticles is shown in Figure S4.

To robustly characterize a material and consider for further applications, it is important to be able to prepare it in large quantities with high chemical yield. To address this challenge, we employed underpressure-governed hot-injection technique that we previously developed. This method allows for fast injection from the addition funnel, run by applying mild vacuum to the reaction flask (TEM analysis and upscaling setup is shown in Figure 2). Using underpressure-governed injection, we successfully upscale the GeTe recipe by a factor of 50 and achieve 2 g of nanoparticles per batch, while maintaining the narrow size distribution of GeTe nanoparticles. For the same precursor ratios, the size of GeTe nanoparticles is slightly smaller than that obtained by small-scale synthesis (Figure S5). This can be explained by larger temperature drop after large-scale injection, associated with faster injection rates and boiling of the reaction mixture under mild vacuum.

Measuring Crystallization Temperature. To demonstrate how we quantify the crystallization temperature, we work with the 6 nm sized GeTe particles produced from the upscaled synthesis. To estimate the crystallization point, we perform high-temperature X-ray diffraction measurements (HT-XRD), heating GeTe nanoparticles under N₂ atmosphere at a constant ramp of 7 °C/min (Figure S6). The results of
HT-XRD are summarized in Figure 3. As-synthesized GeTe nanoparticles retain their amorphous structure up to temperatures >200 °C, as seen by the absence of Bragg reflections on XRD spectra. Upon further heating, X-ray peaks appear and match the rhombohedral modification of bulk GeTe well (Figures 3a and S7). The peak corresponding to the main Bragg reflection (202) is shown in Figure 3b, and its intensity is plotted as a function of sample temperature (Figure 3c). This curve has a sigmoidal growth shape with its two plateau regions corresponding to amorphous and crystalline structures of GeTe nanoparticles (Figure 3d). The shaded region in Figure 3c represents the temperature range where crystallization of GeTe nanoparticles occurs. By analyzing the width of the (202) reflection, we find that crystalline domain size increases up to 25 nm in this region (Figure S8), which indicates that coalescence (i.e., sintering) of neighboring GeTe nanoparticles also occurs in the same temperature range as crystallization.

To distinguish between coalescence and crystallization and to gain a quantitative understanding of crystallization, we perform differential scanning calorimetry (DSC) measurements on 6 nm GeTe nanoparticles, heated under N2 atmosphere at different rates (Figure 4a). We observe an exothermal signal, comprising two closely spaced peaks. Figure 4b plots these peak positions (extracted by taking the second derivatives of the thermograms) as a function of heating rate. On the basis of ex situ heating TEM experiments (Figure S9), we assign the lower temperature process to crystallization and the higher temperature to coalescence. Kissinger analysis can be applied to extract activation energies of the processes (Figure 4c). Crystallization has a higher activation energy (3.29 eV) than coalescence (2.79 eV), which is in agreement with the literature for Ca−Mg alumosilicates and telluride glasses.27,28

We hypothesize that the slightly higher coalescence temperature is due to the presence of an organic shell around the nanoparticles. The shell spaces nanoparticles (by approximately 1 nm, Figures 1c and 2a), which comprises a physical gap for atomic diffusion and coalescence. This assumption is supported by the fact that the relative magnitude (i.e., enthalpy) of crystallization and coalescence peaks in the DSC thermograms is dependent on the heating rate: faster heating increases the contribution of the crystallization exotherm because there is less time for sintering (Figure 4a).

To confirm that the organic shell enables us to observe the temperature-dependent crystallization process, we show that we can instead cause coalescence followed by
crystallization by replacing the steric organic ligands with short inorganic anions (Figure 5a). Fourier-transform infrared spectroscopy confirms nearly complete elimination of organic shell for several different ligand-exchange protocols (Figure 5b).20,21 We perform temperature-dependent resistance measurements of GeI₂-covered GeTe nanoparticle thin films.
and observe a pronounced resistance drop at the bulk crystallization temperature of GeTe of 170 °C. This indicates that coalescence of the film took place prior to or at the crystallization temperature, and that following coalescence the GeTe film behaves as a bulk material (Figure 5c).

**DISCUSSION**

**Size-Dependent Crystallization Temperature.** In the section above, we explain how we determine the crystallization temperature of GeTe nanoparticles using high-temperature XRD, DSC, and ex situ TEM (Figures 3, 4, and S6–S9). In particular, we find that the two processes—crystallization and coalescence of GeTe nanoparticles—happen almost simultaneously. When heating up the nanoparticles with an organic shell, crystallization of GeTe nanoparticles occurs before coalescence (Figures S9 and 6a); however, fast coalescence explains most of the steep growth of XRD reflections (Figure S8). Therefore, crystallization temperature should be taken as the onset temperature of the intensity growth regime. This applies to DCS results (extrapolation of peak temperatures for the lower-temperature exotherms, Figure 4b) and to the analysis of XRD width (Scherrer formula, Figure S8).

We take this approach to quantify the size dependence of the crystallization temperature. Figures 6b–d shows XRD intensity—temperature maps for different sizes of GeTe nanoparticles, from which we extract intensity profiles of the main (202) Bragg reflection (Figure 6e). Taking the onset of each intensity profile, we find that the crystallization temperature increases with decreasing the GeTe nanoparticles size (Figure 6f). While such a trend agrees well with the literature on colloidal GeTe nanoparticles (Figure S10), previous reports did not explain this trend.

Here, we fit the size-dependent crystallization of GeTe nanoparticles with two models. The first model is based on the entropy change during the phase transition. For a melting process, the entropy of fusion, $\Delta S_{\text{fusion}}$, can be linked with size-
dependent melting temperature, $T_{melt}^{NP}$, and physical dimensions of material (i.e., diameter of nanoparticles, $d_{NP}$) using the expression:

$$\frac{T_{melt}^{NP}}{T_{melt}^{bulk}} = \exp \left\{ -\frac{2(\Delta S_{melt} - R)}{3R_k d_{NP} - 1} \right\}$$

where $T_{melt}^{bulk}$ is melting point of bulk material, and $d_0$ is a critical diameter of nanoparticle (defined as the size of nanoparticle, for which all atoms are surface atoms). In analogy, we write the size dependent crystallization temperature, $T_{cryst}^{NP}$:

$$\frac{T_{cryst}^{NP}}{T_{cryst}^{bulk}} = \exp \left\{ -\frac{2(\Delta S_{cryst} - R)}{3R_k d_{NP} - 1} \right\}$$

where we have replaced $\Delta S_{melt}$ with entropy of crystallization, $\Delta S_{cryst}$. Taking the bulk crystallization temperature of GeTe from literature ($T_{cryst}^{bulk} = 443.15$ K), we arrive at a simple model with two fitting parameters: entropy of crystallization, $\Delta S_{cryst}$, and critical diameter, $d_0$. We fit this model to the size-dependent crystallization of GeTe nanoparticles (Figure 6f) and determine important parameters for basic understanding of GeTe crystallization. Namely, we find the entropy of crystallization for GeTe ($\Delta S_{cryst} = -14.5$ J/(mol-K)) that makes sense: (i) it is negative because ordering increases upon crystallization and (ii) its absolute value is smaller than entropy of fusion ($\Delta S_{melt} = 18.0$ J/(mol-K)), which is expected because of gradual decrease of the disorder parameter for material in its supercooled (i.e., amorphous) state (Figure S11). Furthermore, the extracted critical diameter ($d_0 = 0.384$ nm) is consistent with the value expected for GeTe. The volume of the sphere with a critical diameter of 0.384 nm is just slightly smaller than the volume of the smallest GeTe cluster, having at least one nonsurface atom (Figure S12).

We can also explain our data with a second model employing classical nucleation theory, which defines nucleation rate, $J_{cryst}$, as the number of nuclei per unit volume.31 To crystallize a nanoparticle, we need at least one nucleus per nanoparticle. Therefore, $J_{cryst}$ can be expressed as follows:

$$J_{cryst} = \frac{1}{V_{NP}} \frac{6}{\pi d_{NP}^3}$$

where $V_{NP}$ and $d_{NP}$ are volume and diameter of spherical GeTe nanoparticle. Alternatively, $J_{cryst}$ is related to crystallization temperature, $T_{cryst}$, via the Arrhenius-type equation:

$$J_{cryst} = J_0 \exp \left( -\frac{\Delta G_{cryst}^*}{k_B T_{cryst}} \right)$$

Combining 3 and 4, we can relate crystallization temperature and diameter of GeTe nanoparticles via equation, which has two fitting parameters: prefactor, $J_0$, and activation energy of nucleation, $\Delta G_{cryst}^*$. This model displays the observed trend for size-dependent crystallization of GeTe nanoparticles (Figure S11). Smaller sizes of nanoparticles require a higher specific density of nuclei and thus exhibit higher crystallization temperatures. More elaborate nucleation theory models can be built on this basis to account for heterogeneous nucleation, large nucleation rates (i.e., > 1 nucleus per particle), and the effects of surface coverage, etc.

Presented models describe a trend for ultrasmall GeTe nanoparticles (Figure S10),12 while slightly higher crystallization temperature is observed for larger GeTe sizes by...
The latter discrepancy can be explained by the nonstoichiometric effects—when the composition of GeTe deviates from Ge:Te 1:1 ratio, the crystallization temperature increases.\(^7\)

**Nanodimensional Effects for Phase Change Properties.** The size-dependence of the phase transitions in GeTe nanoparticles smaller than 10 nm highlights the opportunities to design memory devices at the nanoscale. Figure 7 summarizes the effect of size on phase-change properties of GeTe nanoparticles. We calculated melting point depression for GeTe nanoparticles using eq 1\(^{29}\) and the critical diameter \(d_k = 0.384 \text{ nm}\) extracted from the size-dependent crystallization fitting. While crystallization temperature increases with smaller nanoparticle size, their melting point decreases to lower temperatures (Figure 7a). This increase of the crystallization temperature should lead to improved data retention at room temperature, while the decrease in melting temperature lowers a power consumption of the memory cell.\(^2\) At the same time, the temperature window (i.e., a difference between \(T_{\text{mel}}\) and \(T_{\text{cryst}}\)) remains wider than 400 K (Figure 7b), ensuring the reliability of memory cell arrays. The size-dependence of these phase transitions may make it possible for materials such as selenides and quaternary telluride compositions that cannot be crystallized or too high bulk melting temperature to be used in thin film phase change memories due to too low bulk crystallization or too high bulk melting temperature to be used in nanoparticle form phase-change memory devices.\(^{33,34}\)

The reduced crystallization temperature (i.e., a ratio between \(T_{\text{cryst}}\) and \(T_{\text{mel}}\)), which indicates the ease of glass formation,\(^35\) increases as the size of GeTe nanoparticles decreases (Figure 7c). A size-dependence of reduced crystallization temperature indicates that (i) smaller GeTe nanoparticles are better glass formers (i.e., they can form glass structure at moderate or slow cooling rates);\(^2\) and (ii) crystallization kinetics of GeTe nanoparticle may attain a size dependence.\(^35\) Importantly, GeTe nanoparticles larger than 5 nm can be classified as marginal glass formers,\(^1\) combining fast crystallization kinetics and stability of supercooled state.

Finally, decreasing the size of phase-change materials leads to improved power efficiency of the device.\(^36,37\) We calculate the power consumption of a hypothetical memory cell, in which a single GeTe nanoparticle represents a switching phase-change volume (Figure S13). As expected, a power efficiency of such a particle-per-bit device improves as the GeTe size decreases. Even after taking into account the 99% power losses due to heat dissipation,\(^36\) the power consumption to crystallize or melt GeTe nanoparticle is on par with energy spent per synaptic event in living organisms (Figure 7d).\(^38\) Such a comparison is provided to emphasize striking opportunities for colloidal phase-change nanoparticles in ultralow power and high-density phase-change memory devices.

**CONCLUSIONS**

This work assessed the potential of colloidal nanoparticles for phase-change memory technology. Using GeTe nanoparticles as a case study, we demonstrated reliable synthesis of colloidal phase change materials and quantified the size-dependence of the crystallization phase transition for the first time. The amide-promoted liquid-phase synthesis enabled accurate size control and large quantity production of GeTe nanoparticles, and it can be extended to other binary and ternary phase change materials. By taking a multipronged approach, we successfully isolated coalescence and crystallization of nanoparticles to quantify the temperature dependence of the crystallization phase transition. We showed that increase in crystallization temperature with decreasing nanoparticle size can be explained using the thermodynamic model or classical nucleation theory. The size-dependent crystallization and melting temperatures in the sub-10 nm size regime offer improved data retention, ultralow power consumption, and fast kinetics of crystallization (i.e., fast write times), highlighting the prospects of nanoparticle phase change materials for memory technology.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.8b02702.

- Size and size distribution of GeTe nanoparticles, Figure S1; size distribution and representative TEM images of GeTe nanoparticles, Figure S2; size distribution and representative TEM images of GeTe nanoparticles, Figure S3; typical energy dispersive X-ray spectrum of GeTe nanoparticles, Figure S4; size distributions of GeTe nanoparticle, Figure S5; intensity of (202) Bragg reflection of GeTe rhombohedral structure while heating GeTe nanoparticles with different ramps, Figure S6; X-ray patterns of GeTe nanoparticles, heated to different temperatures, Figure S7; crystallization temperature estimation for 6 nm GeTe nanoparticles, Figure S8; TEM analysis of 6 nm GeTe nanoparticles, Figure S9; size-dependent crystallization of colloidal GeTe nanoparticles, Figure S10; comparison of two fitting models for the size dependence of GeTe crystallization temperature, Figure S11; octahedral TeGe\(_6\) cluster, Figure S12; and schematics of particle-per-bit device, Figure S13 (PDF)

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The manuscript was written through contributions of all authors.

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**Notes**

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