

Fig. 4. Dispersion data for the sample containing both donors and acceptors with structure $Alq_3:PMMA|60$ nm Ag|R6G:PMMA deposited onto a corrugated substrate. Data were obtained by recording the transverse-magnetic polarized (A) transmittance and (B) PL emission as a function of the in-plane wave vector of light, $k_x/2\pi$. Both are plotted with a logarithmic gray scale, where for (A), transmittance ranges from T=0.001 (black) to T=0.1 (white), whereas for (B), white represents minimum and black maximum PL emission. In each case, the data are dominated by the scattered features corresponding to the excitation and emission of both symmetric and antisymmetric coupled SPPs.

donor molecules on the far side of the silver film, whereas above $\lambda = 550$ nm there are two components to consider. In addition to the strong coupled SPP reradiation, which now arises from SPPs generated by the relaxation of acceptor molecules on the near side of the silver film (primarily excited by energy transferred from the donors), there is a broad emission band from $\lambda = 550$ to 675 nm, peaking at ~565 nm, which does not disperse with angle. This band is an order of magnitude more intense than for the acceptor-only sample (27) and corresponds to the enhanced acceptor emission previously seen in Fig. 2. This emission is direct radiation from acceptor molecules rather than grating-scattered reradiation from acceptor-excited SPP modes. Figure 4B clearly shows that both donors and acceptors strongly excite the coupled SPP

modes of the structure, that strong acceptor emission arises from excitation of the remote donor layer, and that the total emission from the structure can be strongly enhanced by recovering coupled SPP emission from both donor and acceptor through scattering from the grating microstructure.

A direct application of the strategy outlined above would be to top-emitting organic light-emitting diodes. Such devices are attractive for display applications where emission occurs through a metallic cathode. We have shown that depositing a dielectric layer on the surface of the device may lead to greater efficiency (7), and by dye-doping this layer SPP-mediated energy transfer could increase the spectral coverage of the output. Another potential area of application is synthetic light harvesting structures; SPP modes

could act to channel energy from absorbing species to reaction centers and more generally in photochemistry near surfaces.

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Enhancement of Ferroelectricity in Strained BaTiO₃ Thin Films

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Biaxial compressive strain has been used to markedly enhance the ferroelectric properties of ${\rm BaTiO_3}$ thin films. This strain, imposed by coherent epitaxy, can result in a ferroelectric transition temperature nearly $500^{\circ}{\rm C}$ higher and a remanent polarization at least 250% higher than bulk ${\rm BaTiO_3}$ single crystals. This work demonstrates a route to a lead-free ferroelectric for nonvolatile memories and electro-optic devices.

Enormous strains can exist in thin films when one material is deposited on another (1), resulting from differences in crystal lattice parameters and thermal expansion behavior between the film and the underlying

substrate or arising from defects formed during film deposition (2, 3). As a result, the properties of thin films can be markedly different than the intrinsic properties of the corresponding unstrained bulk materials (4-9).

Although such strain often leads to degraded film properties, if judicious use is made of substrates and growth parameters, strain offers the opportunity to enhance particular properties of a chosen material in thin film form, namely strain engineering.

Strain engineering could facilitate the introduction of more environmentally benign ferroelectric random-access memories (FeRAM). Large shifts in the paraelectric-to-

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ferroelectric transition temperature (T) and remanent polarization (P_r) are expected (10-14)and have been observed (15-17) in ferroelectrics, signaling the viability of a strainengineered advance for FeRAM. The major disadvantages of the two materials most widely being pursued for FeRAM (18), Pb(Zr,Ti)O₃ and SrBi₂Ta₂O₉, are (i) the volatility of the lead and bismuth constituents of these materials, which complicates their introduction into semiconductor fabrication facilities, and (ii) environmental issues associated with the toxicity of lead. We demonstrate that the ferroelectric properties of BaTiO₃ can be enhanced with the use of strain to make them viable for ferroelectric memory applications. The widespread use of (Ba,Sr)TiO₂ in semiconductor fabrication facilities for dynamic random-access memories (DRAM) greatly simplifies the introduction of this related material into silicon-based devices.

To predict the $T_{\rm c}$ enhancement and the temperature dependence of the lattice parameters of BaTiO₃ thin films under large biaxial strains with the use of Landau thermodynamic theories (10), we determined a new set of phenomenological coefficients, because existing ones are only applicable to small compressive strains ($<\sim 0.4\%$) (11). Figure 1 shows the T_c enhancement predicted from thermodynamic analysis for a BaTiO₃ thin film under biaxial strain ε_s = $(a_{\parallel} - a_0)/a_0$, where a_0 is the lattice parameter of free-standing cubic BaTiO₃ and a_{\parallel} is the inplane lattice parameter of a biaxially strained (001) BaTiO₃ film. The green region shows the range in predicted $T_{\rm c}$ resulting from the range of reported property coefficients for BaTiO₂ that enter into the thermodynamic analysis (19-21). Figure 1 implies that a biaxial compressive strain of only ~1% should be sufficient to produce strained (001) BaTiO₃ films with a T_c comparable to or higher than unstrained Pb(Zr,Ti)O3 films.

Although Fig. 1 might seem to imply that T_c can be enhanced without bound, there are limits to strain engineering. The driving force for film relaxation increases with strain and film thickness. When films are grown to thicknesses greatly exceeding their critical values, relaxation toward a zerostrain state by the introduction of dislocations begins. Thus, for strain engineering to be effective, it is important to grow films below, or at least close to, their critical thickness for relaxation. Because the critical thickness at which dislocations begin to form varies approximately inversely with lattice mismatch (1), lower mismatch is desired to allow the growth of strained BaTiO₃ films that are thick enough to allow their ferroelectric properties to be conveniently probed or used in devices. Notably, Fig. 1 only applies to thick strained ferroelectrics; as ferroelectrics get thin ($<\sim$ 100 Å), their ferroelectric properties can be substantially diminished by finite-size effects (17, 22–24). Optimizing the trade-off between strain and film thickness depends on the particular application. For FeRAM, films several hundred angstroms in thickness are needed (25). Based on the equilibrium critical thickness (1, 26) for BaTiO₃, this would constrain ε_s to be less than about 0.5%; however, we experimentally found that it is possible to grow coherent BaTiO₃ films at ε_s = -1.7% that are 500 Å thick.

We used the single-crystal substrates $GdScO_3$ and $DyScO_3$, because they are structurally (27), chemically (27), and thermally (28) compatible with $BaTiO_3$, and they have appropriate lattice constants to impart ε_s of about -1.0 and -1.7%, respectively, on coherent (001) $BaTiO_3$ films (21). Epitaxial $BaTiO_3$ thin films were grown on (110) $GdScO_3$ and (110) $DyScO_3$ substrates by reactive molecular beam epitaxy (MBE) and by pulsed-laser deposition (PLD) with in situ high-pressure reflection high-energy electron diffraction (21).

The lattice parameters of the strained (001) $BaTiO_3$ thin films are summarized in Table 1. These films are epitaxial, purely c-axis oriented (the c axis of all $BaTiO_3$ domains is perpendicular to the wafer surface), and, with the exception of the $BaTiO_3$ film on $DyScO_3$ that is 2000 Å thick, are

Table 1. Results from high-resolution x-ray diffraction measurements on the films. The in-plane (a) and out-of-plane (c) lattice constants and full width at half maximum (FWHM) of rocking curves of various peaks (002 of BaTiO₃, 200_{pseudo-cubic} of SrRuO₃, and 200_{pseudo-cubic} of GSrGO₃ and DyScO₃) at room temperature are given. The a- and c-lattice parameters of single-crystalline BaTiO₃ are 3.992 and 4.036 Å, respectively (20).

	(±0.002) (Å)	(±0.0005) (Å)	FWHM (°)
Molecular beam epitaxy			
BaTiO ₃ (1000 Å) on GdScO ₃			
BaTiO ₃	3.964	4.0693	0.080
GdScO ₃	3.965	3.9638	0.009
BaTiO ₃ (500 Å) on DyScO ₃			
BaTiO ₃	3.940	4.0953	0.120
DyScO ₃	3.943	3.9396	0.009
Pulsed-laser deposition			
BaTiO ₃ (2000 Å) on SrRuO ₃ (1000 Å) on GdScO ₃			
BaTiO ₃	3.965	4.0692	0.042
SrRuO ₃	3.968	3.9052	0.036
GdScO ₃	3.964	3.9646	0.008
$BaTiO_3$ (500 Å) on $SrRuO_3$ (2000 Å) on $DyScO_3$			
BaTiO₃	3.939	4.0989	0.045
SrRuO ₃	3.943	3.9110	0.022
DyScO ₃	3.944	3.9396	0.009
$BaTiO_3$ (2000 Å) on $SrRuO_3$ (1000 Å) on $DyScO_3$			
BaTiO ₃	3.958	4.0819	0.135
SrRuO ₃	3.947	3.9187	0.047
DyScO ₃	3.944	3.9398	0.009

fully coherent with the substrates, without any resolvable lattice relaxation.

To identify the ferroelectric phase transition, the temperature dependence of the in-plane and out-of-plane lattice parameters of the films and substrates was measured with a variable-temperature four-circle x-ray diffractometer equipped with a twodimensional (2D) area detector with an angular resolution of ~0.02°. Unstrained BaTiO₃ undergoes a ferroelectric transition at about 130°C from the high-temperature cubic phase $(Pm\overline{3}m)$ to the low-temperature tetragonal phase (P4mm) (20). Figure 2A shows 2D images of the 002 and 202 diffraction peaks at selected temperatures from a BaTiO₃ single crystal as well as from coherent BaTiO₂ thin films grown on (110) GdScO₂ and (110) DyScO₂ substrates. There is no substantial change in the diffraction peaks of the BaTiO3 thin films that are at or above $T_c \sim 130^{\circ}$ C. As expected, the single diffraction spot of the BaTiO₃ single crystal splits into two below 130°C, corresponding to a and c domains of the tetragonal (P4mm) ferroelectric phase.

The in-plane and out-of-plane lattice parameters of the strained BaTiO₃ films grown by MBE were determined from the 202 and 002 diffraction peaks and are plotted as a function of temperature in Fig. 2B. The inplane lattice parameters of the BaTiO₃ are coherent to the underlying substrates over the entire temperature range (25° to 700°C). There are marked differences in the evolution of the lattice parameters with temperature between the unstrained BaTiO₃ single crystal and the strained BaTiO₃ thin films. Notably, the BaTiO₃ thin films never become cubic; they remain tetragonal as a re-

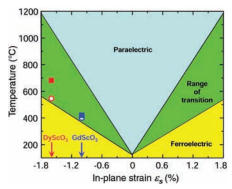


Fig. 1. Expected T_c of (001) BaTiO₃ under biaxial in-plane strain ($ε_s$), based on thermodynamic analysis (10, 21). The green region represents the range (error bars) in the predicted T_c resulting from the spread in reported property coefficients (20, 21) for BaTiO₃ that enter into the thermodynamic analysis. The data points show the observed $ε_s$ and T_c values of coherent BaTiO₃ films grown by MBE on GdScO₃ (blue circle) and DyScO₃ (red circle) substrates and by PLD on GdScO₃ (blue square) and DyScO₃ (red square) substrates.

sult of the biaxial substrate constraint. The predicted dependence of the c-lattice parameter of biaxially strained BaTiO3, with and without a ferroelectric phase transition, was calculated from thermodynamic analysis (10) and is shown by the green solid and dashed curves in Fig. 2B, respectively. Because the BaTiO3 film is clamped inplane, all structural changes resulting from the phase transition and thermal expansion are accommodated through changes in the out-of-plane lattice parameter only. The agreement between the solid green prediction and the measured c-axis lattice parameters in Fig. 2B is strong evidence that the change in slope in the c-axis lattice parameter at high temperature corresponds to a ferroelectric phase transition. Analogous lattice constant behavior has been observed in other constrained ferroelectric films (15, 17), is consistent with theory (13, 17), and has been used to determine T_c . The T_c of the coherent BaTiO₃ thin films shown in Fig. 2B is ~400°C on GdScO₃ and ~540°C on DyScO₃.

To confirm the huge shifts in $T_{\rm c}$, we attempted to measure polarization hysteresis loops on a 2000-Å-thick coherent BaTiO₃ film grown by PLD on a coherent SrRuO₃ bottom electrode on (110) GdScO₃. At temperatures up to about 200°C, hysteresis loops were clearly seen, but at higher temperatures the dielectric losses in the films became too high for reliable measurements. We made

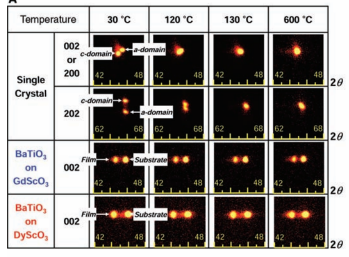
second harmonic generation (SHG) measurements as a function of temperature on this PLD-grown BaTiO₃/SrRuO₃/GdScO₃ sample as well as the MBE-grown BaTiO₃/ GdScO₃ sample, whose lattice constants versus temperature behavior is shown in Fig. 2B (21). An SHG signal is only exhibited by materials that lack inversion symmetry. All ferroelectrics must lack inversion symmetry, but there are many materials that lack inversion symmetry and are not ferroelectric. This makes SHG a necessary but insufficient probe for ferroelectricity. Nonetheless, SHG (Fig. 2C) shows that the phase we know from hysteresis loops to be ferroelectric at room temperature remains noncentrosymmetric to the same high temperature (29, 30) at which x-ray diffraction indicates a phase transition. The interpretation consistent with all our analyses—x-ray diffraction, SHG, and hysteresis measurements—is that biaxial compressive strain increases the $T_{\rm c}$ of BaTiO₃.

Hysteresis measurements were made on 200-μm-diameter capacitors of strained BaTiO₃ thin films sandwiched between epitaxial top and bottom electrodes of the conducting perovskite oxide SrRuO₃ (31). High-resolution x-ray diffraction measurements (summarized in Table 1) revealed the BaTiO₃ ferroelectric layers as well as the 1000-Å-thick SrRuO₃ bottom electrodes to be fully coherent with the underlying substrates. No relaxation was observed even for

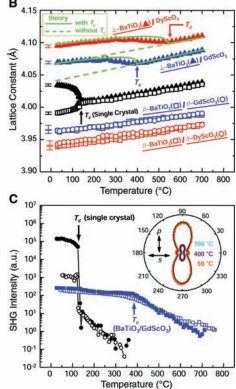
BaTiO₂ films as thick as 500 Å on DyScO₂ and 2000 Å on GdScO₃. The critical thicknesses of BaTiO3 thin films grown on coherent SrRuO₃ bottom electrodes on GdScO₃ and DyScO₃ are higher than those of BaTiO₃ films grown directly on GdScO₃ and DyScO₃. This observation is consistent with critical thickness theory, in which the difference arises from strain partitioning between the layers as well as the altered geometry of misfit dislocations in a single layer versus a bilayer (26). Because the leakage in the coherent stack containing a 500-Å-thick BaTiO3 layer on DyScO3 was too high for good ferroelectric hysteresis measurements, a SrRuO₃/BaTiO₃/SrRuO₃/DyScO₃ stack was grown with a 2000-A-thick BaTiO₃ layer. This latter stack had low leakage; however, it was partially relaxed.

Figure 3A shows the ferroelectric hysteresis loops measured on the ferroelectric stacks grown on GdScO₃ and DyScO₃ substrates with 2000-Å-thick BaTiO₃ layers, together with results from a BaTiO₃ single crystal (32). The hysteresis loops are shifted in the positive voltage direction. This imprint effect is probably due to the asymmetric interfacial properties of the top and bottom electrodes to the BaTiO₃ films. Even though we used SrRuO₃ for both electrodes, the growth temperature (350°C) of the top electrode was much lower than that of the bottom electrode (680°C), which might lead to poor crystallinity of the top electrode and

Fig. 2. (A) 2D images at selected temperatures of the x-ray diffraction peaks from a BaTiO₃ single crystal and strained BaTiO₃ thin films. (B) Temperature dependence of the lattice parameters of single-crystal BaTiO₃, and strained BaTiO₃ thin films grown by MBE on DyScO₃ substrates and GdScO₃ substrates. The in-plane (//) and out-of-plane (\bot) lattice constants of the BaTiO₃ thin films and underlying substrates are shown. The change in slope at high temperature



signals a phase transition. The error bars $(\pm SD)$ of the measured lattice constants are shown to the left of each curve. The measured values of the out-of-plane lattice constant of biaxially strained BaTiO₃ are compared with theoretical predictions (10, 21) with and without a ferroelectric transition. (C) Optical SHG signals from a BaTiO₃ single crystal [heating (black open circles) and cooling (black filled circles)] and the same strained BaTiO₃ on GdScO₃ film as (B) [heating (blue open squares) and cooling (blue filled squares)]. The fundamental wave (wavelength 900 nm, 65-fs laser pulses) is incident on the sample at an angle of $\phi = 23^{\circ}$ on the substrate side of the thin film and at normal incidence ($\phi = 0^{\circ}$) for the BaTiO₃ single crystal. SHG signals of 1 arbitrary unit (a.u.) or lower are at the noise level of our lockin-based measurement system. The inset shows polar plots of SHG intensity (radius) versus fundamental polarization (azimuth). Circles are experiment and solid lines are theory (21, 35).



asymmetric interfaces. The P_r and coercive field (E_r) were determined to be ~50 μ C/cm² and 80 kV/cm for the fully coherent BaTiO₃/ GdScO₃ sample and $\sim 70 \,\mu\text{C/cm}^2$ and 25 kV/ cm for the partially relaxed BaTiO₃/DyScO₃ sample, respectively. This $P_{\rm r}$ value is almost 270% of the 26 μC/cm² (32) of single crystal BaTiO₃, 3.5 times higher than the maximum switching charge density (20 μC/cm²) assumed in the scaling analysis of FeRAM (18), and comparable to the P_r of unstrained Pb(Zr,Ti)O3 films (33). As this $P_{\rm r}$ of ~70 μ C/cm² was observed in a partially relaxed sample with ε_s of -1.3%, a coherently strained BaTiO3/DyScO3 sample with ε_s of -1.7% could have an even higher P_r .

Another important issue for the application of ferroelectric capacitors to memory devices is the loss of switched polarization after repeated switching, i.e., fatigue. We performed fatigue measurements by applying 8.6-µs-wide pulses with a repetition frequency of 10 kHz to the top and bottom SrRuO₃ electrodes of the SrRuO₃/BaTiO₃/SrRuO₃/ $GdScO_3$ structure at $V_{max} = 4$ V, where V_{max} is the amplitude of the voltage pulse. $V_{\rm max}$ = 4 V corresponds to 200 kV/cm of the electric field. The switched polarization decreased by 10% of its original value after 106 fatigue cycles, but recovered its original value after 1010 cycles. This is consistent with previous observations of fatigue-free behavior when conducting oxide electrodes are used (34, 35).

As a check that the enhancement of T_c observed in coherently strained BaTiO₃ thin films grown by MBE (Fig. 2B) is inherent and applicable to a device structure with a

conductive bottom electrode, we performed high-temperature x-ray diffraction measurements on the coherent BaTiO3 thin films with SrRuO₃ bottom electrodes grown by PLD. Figure 3B shows the evolution of the inplane (a) and out-of-plane (c) lattice parameters of the BaTiO₃ film and the GdScO₃ and DyScO₃ substrates as a function of temperature. The in-plane lattice parameters reveal that both the BaTiO₃ and SrRuO₃ layers are coherently strained to the underlying substrates over the entire temperature range. This is consistent with the absence of misfit dislocations along the interface between GdScO3 and SrRuO₃ and along the interface between SrRuO₃ and BaTiO₃, as shown by the crosssectional transmission electron microscope images in figs. S1 and S2 (21). As seen in the figure, the transition behavior of the PLD samples is quite similar to those grown by MBE. T_c was determined to be $\sim 420^{\circ}$ and ~680°C for samples grown on GdScO₃ and DyScO₃, respectively. The green solid and dashed lines in Fig. 3B are theoretical predictions of c-lattice parameters with and without the ferroelectric phase transition, which are fairly consistent with the experimentally measured values. The agreement in the results for films grown by MBE and PLD indicates that the observed shifts in ferroelectric properties with strain represent the intrinsic behavior of strained BaTiO3. This experimental dependence of T_c on ε_s is also consistent with the expectations shown in

In summary, we have demonstrated that the ferroelectric properties of BaTiO₃ can be markedly enhanced through strain engineering. These strain-engineered heteroepi-

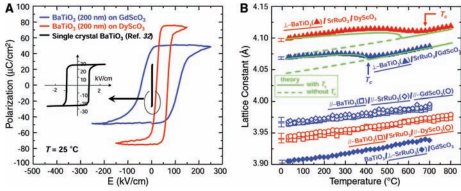


Fig. 3. (A) Polarization-electric field hysteresis loops of BaTiO₂ thin film capacitors (2000 Å) grown by PLD on GdScO₃ and DyScO₃ with SrRuO₃ top and bottom electrodes. The inset shows the hysteresis loop of an unstrained bulk BaTiO₃ single crystal for comparison (32). (B) Temperature dependence of the lattice parameters of strained SrRuO₃/BaTiO₃/SrRuO₃ capacitor structures grown by PLD on DyScO₃ substrates and GdScO₃ substrates. The in-plane (//) and out-of-plane (⊥) lattice constants of the BaTiO₃ thin films, SrRuO₃ bottom electrode, and underlying substrates are shown. The lattice parameters of the SrRuO₃ film on DyScO₃ could not be resolved because SrRuO₃ and DyScO₃ are isostructural with very similar lattice parameters. The change in slope at high temperature indicates a phase transition. The error bars (±SD) of the measured lattice constants are shown to the left of each curve. The measured values of the out-of-plane lattice constant of biaxially strained BaTiO₂ are compared with theoretical predictions (10, 21) with and without a ferroelectric transition.

taxial thin films provide a broad range of operating temperatures as well as higher remanent polarization for improved noise immunity and the ability to scale FeRAM to smaller cell sizes. Another application of strain-engineered BaTiO3 films is highspeed electro-optic modulators, in which the sizeable electro-optic coefficients of BaTiO₃ can be enhanced by appropriate strain engineering. The ability to withstand huge strains gives thin films a degree of freedom absent from bulk. This can be exploited to enhance the ferroelectric properties of any ferroic system, including multiferroics (8, 22, 36), whose ferroic order parameter has a strong coupling to strain.

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- 19. For biaxial tensile ($\varepsilon_s \ge 0$) or biaxial compressive ($\varepsilon_s \le 0$) strains, T_c is expressed as follows: $T_c = \theta + 2\epsilon_0 C \times \frac{Q_{11} + Q_{12}}{S_{11} + S_{12}} \epsilon_s$ ($\epsilon_s \ge 0$) or $T_c = \theta + 2\epsilon_0 C \frac{2Q_{12}}{S_{11} + S_{12}} \epsilon_s$ ($\epsilon_s \le 0$). In the equations, θ is the Curie-Weiss temperature of unstrained BaTiO₃, ε_0 is the permittivity of free space, C is the Curie constant, Q_{11} and Q_{12} are electrostrictive coefficients, and s_{11} and s_{12} are elastic compliances. The breadth of the green region in Fig. 1 for T_c is due to the variation in what are considered the most accurate reported values of these constants for BaTiO₃ single crystals (20, 21). Because $(Q_{11}+Q_{12})>0$ and $Q_{12}<0$, T_c is predicted to increase for both positive and negative strains.
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well as the polycrystalline films used in today's FeRAM. This leads to an upper bound in film thickness of about 1000 Å. The thinness of the ferroelectric film is also constrained. First, there is an intrinsic finite-size effect in which the T_c begins to decrease at a thickness of about 100 Å (17, 22-24) and eventually vanishes for thicknesses in the 10 to 30 Å range (23, 24). The second reason is that extrinsic effects (pinholes and nonuniform thickness over the capacitor area) lead to unacceptably high leakage currents for FeRAM device operation.

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Supporting Online Material

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Materials and Methods

Figs. S1 and S2

References

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Cation Exchange Reactions in Ionic Nanocrystals

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Cation exchange has been investigated in a wide range of nanocrystals of varying composition, size, and shape. Complete and fully reversible exchange occurs, and the rates of the reactions are much faster than in bulk cation exchange processes. A critical size has been identified below which the shapes of complex nanocrystals evolve toward the equilibrium shape with lowest energy during the exchange reaction. Above the critical size, the anion sublattice remains intact and the basic shapes of the initial nanocrystals are retained throughout the cation exchange. The size-dependent shape change can also be used to infer features of the microscopic mechanism.

Chemical transformations from one solid to another via insertion and exchange of atoms can be used to modify the properties of crystalline materials (1). Recent developments have enabled the production of many technologically important crystalline materials in nanometer sizes, with a wide range of size- and shape-tunable properties (2-8). Of particular interest is the creation of nanocrystals with nonequilibrium shapes and with higher structural and compositional complexity (9-13). In extended solids, reactions involving chemical transformation are in general very slow because of high activation energies for the diffusion of atoms and ions in the solid. For this reason, typical solid-phase reactions require very high temperatures or pressures (14-16) and therefore would seem to be incompatible with kinetically controlled nonequilibrium nanostructures.

However, in crystals only a few nanometers in size, both the thermodynamics and kinetics of reactions can change with size. For example, a large surface-to-volume ratio can be accompanied by a lowering of phase transition temperatures (17, 18). With the decrease in the volume, statistical averaging of the kinetics and mechanisms over a distribution of heterogeneous reaction sites intrinsic to the bulk solid is also reduced, leading to more homogeneous molecule-like reaction kinetics and even different reaction mechanisms in nanocrystals (19). The optimal use of various chemical transformation methods to broaden the range of nanocrystalline materials depends on an understanding of how chemical transformations in a crystalline solid will be affected by a reduction in size. We show that cation exchange reactions can occur completely and reversibly in ionic nanocrystals at room temperature with unusually fast reaction rates. We also show that the crystal structure and morphology of the reaction products are strongly dependent on the size and shape of the nanocrystals.

The prototypical semiconductor nanocrystal system of CdSe reacts with Ag^+ ions to yield Ag_2 Se nanocrystals by the forward cation exchange reaction, and vice versa for the reverse cation exchange reaction. We chose to work with CdSe nanocrystals because of the high degree of control over size and shape that has been achieved (2, 3). The conversion to Ag_2 Se is strongly favored by a thermodynamic driving force of about

–1000 kJ/mol in the bulk (20, 21). Ag₂Se also exhibits an interesting temperature-dependent polymorphism: The superionic conducting phase transition occurs at a relatively low temperature of 133°C in the bulk phase (22). Thus, it may be possible to prepare Ag₂Se with unusually high cation mobility. These two factors favor complete cation exchange in nanocrystals and may prove sufficient to overcome the fact that the exchange reaction is completely kinetically hindered at ambient temperature and pressure in the bulk.

We investigated the reaction by mixing a solution of CdSe nanocrystals (diameter 4.2 nm) in toluene with a small amount of methanolic solution of AgNO3 under ambient conditions. The volume fraction of methanol in the solution mixture is about 1%; the solution contains Ag+ ion in a slightly larger amount than necessary to replace all the Cd²⁺ ions in the nanocrystals. Methanol more strongly binds to any free binary cations in solution and thus favors the forward reaction. A rapid («1 s) change of solution color and complete disappearance of fluorescence is observed upon mixing the solutions. Measurements of the x-ray diffraction (XRD) patterns and optical absorption spectra confirm that the reaction product is Ag₂Se (Fig. 1). The reverse reaction is done under ambient conditions by mixing Ag₂Se nanocrystals with an excess amount (typically 50 to 100 times the initial Cd²⁺ content) of Cd(NO₃)₂ in a mixture of toluene and methanol in the presence of tributylphosphine (volume fraction <3%). A slower color change back to that of CdSe nanocrystals and the reappearance of fluorescence are observed over a period of 1 min. XRD patterns, optical absorption, and fluorescence spectra all indicate that CdSe is recovered from the reverse cation exchange. The XRD linewidths of the initial and recovered case are nearly identical. Moreover, the absorption and fluorescence peak positions, which show strong size dependence due to the quantum confinement effect (2), are also nearly identical for the initial and recovered CdSe nanocrystals. Finally, transmission electron micrograph (TEM) images of the initial and recovered

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