Photoinduced Perturbations of the Magnetic Superexchange in Core-Shell Prussian Blue Analogues

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Cubic heterostructured (BA) particles of Prussian blue analogues, composed of a shell of ferromagnetic K$_{0.3}$Ni[Cr(CN)$_6$]$_{0.8}$·1.3 H$_2$O (A), $T_c \sim 70$ K, surrounding a bulk core of photoactive ferrimagnetic Rb$_{0.4}$Co[Fe(CN)$_6$]$_{0.8}$·1.2 H$_2$O (B), $T_c \sim 20$ K, have been studied. Below $T_c \sim 70$ K, these samples exhibit a persistent photoinduced decrease in low-field magnetization, and these results resemble data from other core-shell particles and analogous ABA heterostructured films. This net decrease suggests that the photoinduced lattice expansion in the B layer generates a strain-induced decrease in the magnetization of the A layer, similar to a pressure-induced decrease observed by others in a pure A-like material and by us in our BA cubes. Upon further examination, the data also reveal a significant portion of the A material whose superexchange, J, is perturbed by the photoinduced strain from the B constituent.

PACS numbers: 75.50.Xx, 75.30.Et, 75.70.Cn
Keywords: Prussian blue analogues, photoinduced magnetism, heterostructures

INTRODUCTION

Molecule-based magnets present an intriguing field of materials due to their combination of magnetism with various other physical phenomena. Specifically, those materials which exhibit photomagnetic properties provide a vast potential for technological applications. In recent years, Prussian blue analogues (PBAs) have piqued the interest of the molecular magnetism community due to their range of magnetic properties. Although examples exist that are known to order magnetically at or above room temperature [1–2], the feature that distinguishes a particularly appealing subset of PBAs is the phenomenon of a charge-transfer-induced spin transition (CTIST) [3], which is linked with that of persistent photoinduced magnetism (PPIM) [4]. When these properties are combined with long-range magnetic order, the potential for information storage applications begins to emerge.

A quintessential example of such a material is the CoFe-PBA, A$_x$Co[Fe(CN)$_6$]$_y$·nH$_2$O, where A is a monovalent alkali cation. CoFe-PBAs are well-known to exhibit a photoinduced CTIST up to a temperature of 150 K [5–7], as well as a similar photoinduced transition within the hysteresis of the thermal CTIST near, and even at, room temperature for certain stoichiometries [8–9]. However, these photoinduced effects are weaker above 20 K, which is the critical temperature, $T_c$, for the long-range magnetic ordering. The fact that the photoeffect in the CoFe-PBA is also structural in nature has been utilized to effect an increase in the temperature at which photocontrol is possible, by incorporating this material into heterostructures [9–11] with PBAs having higher ordering temperatures and known to be pressure-sensitive [12]. To guide work pursuing room temperature photocontrolled magnetism, we seek a deeper fundamental understanding of the mechanism leading to these novel effects in the heterostructures, focusing here on quantifying the amount of material, in the non-photoactive layers, affected by this photoinduced strain from the CoFe-PBA.

In the present study, core (B) and core-shell (BA) particles of PBAs (Fig. 1), composed of shells of K$_{0.3}$Ni[Cr(CN)$_6$]$_{0.8}$·1.3 H$_2$O (A), which orders ferromagnetically with $T_c \sim 70$ K, surrounding bulk cores of Rb$_{0.4}$Co[Fe(CN)$_6$]$_{0.8}$·1.2 H$_2$O (B), which is photoactive and orders ferrimagnetically with $T_c \sim 20$ K, have been investigated. Below $T_c \sim 70$ K of the shell, these samples exhibit a persistent photoinduced decrease in low-field susceptibility, similar to those results from other core-shell particles [10] and analogous ABA heterostructured films [9–11]. This net decrease is attributed to the photoinduced lattice expansion in the B layer, which gen-

![FIG. 1: (Color online) TEM images of (a) core (B) and (b) core-shell (BA). Each scale bar is 200 nm. The arrows denote a selected particle in each image that has been colorized to highlight the core-shell contrast.](image-url)
generates a strain-induced decrease in the magnetization of the A layer, similar to a pressure-induced decrease observed by others in a pure A-like material, attributed to spin-canting \[12\]. Upon further examination, the core-shell susceptibility data also reveal a significant portion of the A material whose superexchange, \(J\), is perturbed with the application of this photoinduced strain from the B material.

**EXPERIMENTAL DETAILS**

**Synthesis**

K\(_3\)Cr(CN)\(_6\) was synthesized by treating aqueous solutions of potassium cyanide with CrCl\(_3\) · 6 H\(_2\)O and was used after recrystallization from methanol \[13\]. Deionized water used in synthetic procedures was obtained from a Barnstead NANOpure. All of the other reagents were purchased from Sigma-Aldrich or Fisher-ACros and were used without further purification. The filters used during the synthesis were Fast PES Bottle Top Filters with 0.45 \(\mu\)m pore size (Nalgene).

The synthesis of the core particles was performed at room temperature. An aqueous solution (100 mL) of CoCl\(_2\) · 6 H\(_2\)O (0.40 mmol, 4.0 mM), and an equal volume of an aqueous solution containing K\(_3\)Fe(CN)\(_6\) (0.45 mmol, 4.5 mM) were simultaneously added dropwise to 200 mL of nanopure water. To substitute the Rb cation, RbCl (0.8 mmol, 8 mM) was added to the CoCl\(_2\) solution. The final solution was kept under vigorous stirring for one hour after all additions were complete. The particles were subsequently filtered under vacuum using a 0.45 \(\mu\)m filter before being washed with ultrapure water and redispersed in 100 mL of water for use in the next step. To isolate the particles, instead of water, the particles were concentrated by centrifugation and dried under a flow of air.

To prepare the core-shell particles, the previously synthesized core particles solution was diluted with water to 400 mL. An aqueous solution (50 mL) of NiCl\(_2\) · 6 H\(_2\)O (0.095 mmol, 1.9 mM) and an aqueous solution (50 mL) of K\(_3\)Cr(CN)\(_6\) (0.105 mmol, 2.1 mM) were added dropwise (10 mL/hour, peristaltic pump) under stirring at room temperature. The final solution was kept under vigorous stirring for one hour after all additions were complete. Once the addition was complete, the particles were filtered using a 0.45 \(\mu\)m filter and rinsed with ultrapure water. To isolate the particles, instead of water, the particles were concentrated by centrifugation and dried under a flow of air.

**Instrumentation**

Transmission electron microscopy (TEM) was performed on a JEOL-2010F high-resolution transmission electron microscope at 200 kV. TEM grids used were carbon film on a holey carbon support film, 400 mesh, copper from Ted-Pella, Inc. Grids were prepared by dropping 20 \(\mu\)L of a solution containing 5 mg of sample dispersed in 2 mL of EtOH by sonication for 30 min. Energy dispersive x-ray spectroscopy (EDS) was performed with an Oxford Instruments EDS X-ray Microanalysis System coupled to the TEM.

Room temperature powder x-ray diffraction (XRD) was performed on a Bruker DUO diffractometer. Powder samples were packed into 0.3 mm diameter boron-rich thin walled capillary tubes purchased from the Charles Supper Company. Diffraction patterns were collected in the range 5° ≤ \(2\theta\) ≤ 89° with a 600 sec/image collection time. The raw data from the CCD detector were converted to powder patterns using PowDLL converter.

Fourier-transform infrared (FTIR) spectroscopy was performed on a Nicolet 6700 Thermo Scientific spectrophotometer. Powder samples were mixed with KBr and pressed into pellets under 20 MPa. Scans (typically 64) were performed between 2200 and 1900 cm\(^{-1}\), in the region of the CN stretching bands, with a precision of 0.482 cm\(^{-1}\). Corresponding scans of a pure KBr pellet were taken as the background measurement.

Magnetization measurements were performed on a Quantum Design MPMS XL-7 superconducting interference device (SQUID) magnetometer using a homemade fiber optic sample rod with a bundle of 10 optical fibers to bring light into the sample space \[14\]. Powder samples were spread between two pieces of transparent tape.
and held in transparent drinking straws directly below the optical fibers. Visible white light was supplied by a 150 W tungsten-halogen lamp located at room temperature. The temperature dependence of the low-field susceptibility was measured in an applied field of 100 G while warming the sample from base temperature, 2 K. Isothermal magnetization was measured at 2 K while sweeping the field between 70 kG and −70 kG. All light state measurements were performed subsequent to 8 hours of isothermal irradiation at 40 K, after irradiation had been ceased. The sample was then cooled again to base temperature, where the same protocol that was used for the dark state measurements was repeated to acquire the light state data set.

RESULTS AND DISCUSSION

The TEM images reveal contrast between core and shell in the BA particles not present for the pure B particles (Fig. 1). Additionally, EDS line scans of individual particles show Co and Fe content concentrated in the cores, with Ni and Cr content predominately occupying the edges [13]. The FTIR spectra display a new CN stretching frequency, near 2175 cm⁻¹, arising after addition of the A shell, consistent with observations for the pure A material. Powder XRD patterns also present two distinct lattice constants consistent with the single-phase B and A materials, respectively, while only the B phase is exhibited in the core particles (Fig. 2).

The low-field susceptibility of the BA sample exhibits a persistent photoinduced decrease (Fig. 3). Although the B material is a well-established photomagnet [5–7], these heterostructures invert the sign of the effect in addition to increasing the temperature of photocontrol [9–11]. This photoeffect possesses a significant magnitude up to $T_c \sim 70$ K of the A material, attributed to a strain-induced canting of the A spins, similar to pressure-induced effects in a similar material [12].

Additionally, a subtle shoulder near the $T_c$ of the A material emerges in the photoinduced decrease, $\Delta \chi$, indicating a shift in the superexchange of some portion of the shell. In order to quantify the amount of affected material, the derivative of the susceptibility was taken and normalized to the same area, in the range 40 K to 80 K, for each data set (Fig. 4). The validity of normalizing the light and dark data sets to equal areas is substantiated by the corresponding saturation magnetizations, which indicate no loss of spins upon irradiation. Thus, taking the area under these curves to represent the total number of spins in the shell, this area must remain constant for a given sample between the dark and light states. Upon inspection of these derivatives, an aspect of the effect not immediately obvious in the raw data (Fig. 3 inset) becomes apparent. Namely, the magnetic transition of the A material sharpens significantly when the B material is transitioned to the light state. Again, this observation indicates that a significant portion of the A material experiences a change in its superexchange.

Guided by these observations, we apply a simple model of the shell comprising two distinct regions: an inner, strain-sensitive interface region and an outer, rigid bulk-like region, which remains unaffected by switching be-
between the dark and light states. Taking these regions to represent distinct superexchange values, the areas under the curves in Fig. 4, representing the total number of spins in the shell constituent, must account for these two regions. Fitting these curves with two peaks, corresponding to the two assumed superexchange values, provides a measure of the percentage of the shell in each region. The bulk-like value is taken to be the higher $T_c$, which does not shift, and this value remains constant between the fits, while the total area is constrained to unity. These fits reveal 50% of the material with each $T_c$, where the bulk-like material has $T_{c_{\text{bulk}}} \approx 68$ K and the strained material has $T_{c_{\text{dark}}} \approx 62$ K shifting to $T_{c_{\text{light}}} \approx 65$ K, which is closer to that attributed to the bulk.

The direction of this shift may be explained phenomenologically in the following manner. Upon cooling, the B material undergoes a lattice contraction from a 10.3 Å lattice constant to 10.0 Å, thus stretching the A lattice near the interface. With irradiation, the B lattice again expands to its room temperature size of 10.3 Å, allowing the interfacial A lattice to relax closer to its bulk state.

CONCLUSION

By analyzing the first derivative of the susceptibilities in the dark and light states of the BA particles, we have quantified the amount of the A material affected by the strain arising from the photinduced expansion of the B lattice. Although the shift in superexchange is found to be small ($\Delta T_c \approx 3$ K), nearly half of the volume of the A shell experiences this superexchange perturbation upon photoswitching the B core. A forthcoming report will detail the investigation of the same cores with increasing shell thicknesses.

This work was supported by NSF DMR-1005581 (DRT), DMR-1202033 (MWM), the National High Magnetic Field Laboratory via cooperative agreement NSF DMR-0654118, and the state of Florida. The authors would like to thank Dr. Khalil A. Abboud, for the use of the Bruker DUO instrument, and Kerry Sieben, at the University of Florida Major Analytical Instrumentation Center, for TEM and EDS work.

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