Magnetocaloric effect in Gd/W thin film heterostructures

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In an effort to understand the impact of nanostructuring on the magnetocaloric effect, we have grown and studied gadolinium in MgO/W(50 Å)/[Gd(400 Å)/W(50 Å)]₈ heterostructures. The entropy change associated with the second-order magnetic phase transition was determined from the isothermal magnetization for numerous temperatures and the appropriate Maxwell relation. The entropy change peaks at a temperature of 284 K with a value of approximately 3.4 J/kg-K for a 0-30 kOe field change; the full width at half max of the entropy change peak is about 70 K, which is significantly wider than that of bulk Gd under similar conditions. The relative cooling power of this nanoscale system is about 240 J/kg, somewhat lower than that of bulk Gd (410 J/kg). An iterative Kovel-Fisher method was used to determine the critical exponents governing the phase transition to be β = 0.51, and γ = 1.75. Along with a suppressed Curie temperature relative to the bulk, the fact that the convergent value of γ is that predicted by the 2-D Ising model may suggest that finite size effects play an important role in this system. Together, these observations suggest that nanostructuring may be a promising route to tailoring the magnetocaloric response of materials.

INTRODUCTION

Materials exhibiting the giant magnetocaloric effect (MCE) have demonstrated potential for advancing magnetic refrigeration, an energy-efficient and environmentally friendly alternative to conventional refrigeration [1][2]. Historically, magnetic refrigerants have been used only for very specific applications at very low temperatures, and in most cases, the entropy changes were small [3]. In fact, the largest values of the MCE were encountered in the vicinity of a second-order magnetic transition in Gd [4]. The discovery of the giant MCE in Gd (10¯10) and (0001) peaks (higher order peaks of these diffraction for 2θ in the range 20-70° reveals and strong Gd (1010) and (0001) peaks (higher order peaks of these are also present). A simple Scherrer grain size analysis indicates that the Gd layers are structurally coherent throughout their thickness.

Magnetic measurements were performed using a Quantum Design Physical Property Measurement System. The magnetization isotherms were measured in the range of 0–3 T for temperatures of 260–320 K in steps of 10 K. The change in the entropy (S) of a magnetic material in an applied magnetic field (H) is related to the change in magnetization (M) with respect to the temperature (T) through the thermodynamic Maxwell relation:

\[ \left( \frac{\partial S(T, H)}{\partial H} \right)_T = \left( \frac{\partial M(T, H)}{\partial T} \right)_H. \]

For data taken at discrete field and temperature intervals, the change in magnetic entropy, \( \Delta S_M \), due to an applied field from 0 to \( H_0 \) can be approximated as:

\[ \Delta S_M(T, H_0) = \mu_0 \sum_{i} \frac{M_{i+1}(T_{i+1}, H) - M_i(T_i, H)}{T_{i+1} - T_i} \Delta H, \]

where \( \mu_0 \) is the permeability of free space. Figure 1 shows the magnetic entropy change measured for the
W(50 Å)/[Gd(400 Å)/W(50 Å)]₈ multilayer (sample area was 1 cm²). The Δ_S_m peak for low fields is around 284 K, consistent with T_c as determined by the Kovel-Fisher method below [9]. The magnitude of the peak is about 3.4 J/kg-K, about one third of the value for bulk Gd [10]. The shift of the peak with maximum field is approximately linear in the field range studied, with a slope of about 0.16 K/kOe.

Perhaps the most striking feature of the temperature dependence of ΔS_m is that its temperature full width at half max T_{FWHM} is nearly double that of bulk Gd under similar field conditions. In fact, the T_{FWHM} observed for ΔH = 30 kOe exceeds that of bulk Gd for ΔH = 50 kOe. While this is compensated for by a reduction of the ΔS_m peak value, the relative cooling power calculated as ΔS_m × T_{FWHM} [10] is 240 J/kg, which is on the order of that for bulk Gd (410 J/kg). This may indicate that nanostructuring is a potential route for developing magnetic refrigerants with large useful temperature ranges.

An iterative Kouvel-Fisher method was used to determine the critical exponents and Curie temperature of the system [11]. This approach has been quite successful for analyzing amorphous magnetic materials [12, 13]. The data were initially analyzed on a modified Arrott-Noakes plot, M¹/β₀ vs (H/M)¹/γ₀, where β₀ = 2/5 and γ₀ = 4/3 [14]. The high field portions (H > 1 T) of each isothermal data set were extrapolated to determine the M².₅ and (H/M)⁰.₇₅ intercepts with second order polynomials [15], which allows us to determine the spontaneous magnetization (M₀) and the inverse of the initial susceptibility (χ_₀⁻¹), respectively. The slopes of the functions Y(T) = M₀/(dM₀/dT) and X(T) = χ_₀⁻¹/(dχ_₀⁻¹/dT) near T_c are taken as 1/β and 1/γ, respectively. The initial exponents, β₀ = 2/5 and γ₀ = 4/3, led to nonlinear Y(T) and X(T); second order polynomials fit these functions well, and were used to estimate β₁ and γ₁ near T_c. These exponents were then used to create a new modified Arrott-Noakes plot (M¹/β₁ vs (H/M)¹/γ₁), thus beginning another iteration of this process. The data were analyzed in this fashion for four iterations at which point the exponents were unchanging (β₃ = β₄, and γ₃ = γ₄). The convergent exponents were determined to be β = 0.51, and γ = 1.75. Figure 2 shows that the final Y(T) and X(T) obtained by following this procedure are linear on either side of the second order phase transition; the inset shows the values of the two exponents after each iteration. Interestingly, the Curie temperature was relatively insensitive to this iterative process, ranging from 282 K to 284 K for each iteration. This Curie temperature is reduced from that of bulk Gd, which is consistent with finite size effects in Gd thin films [10]. Figure 3 shows the final modified Arrott-Noakes plot using the exponents β = 0.51, and γ = 1.75.

DISCUSSION

The origin of the enhanced T_{FWHM} of the entropy change is currently under investigation. The critical exponent γ determined above to be 1.75 is consistent with the two-dimensional Ising model, potentially suggesting the origin may be related to a change in dimensionality.
FIG. 3: (Color online) Modified Arrott Plot using the critical exponents converged upon after four iterations of the Kouvel-Fisher method: $\beta = 0.51, \gamma = 1.75$.

The suppressed Curie temperature of the Gd does indicate conclusively that finite size effects are playing a significant role in the system, further lending some credence to this possibility. Another potential origin of the enhanced $T_{FWHM}$, as well as the reduced entropy change peak value, is a distribution of Curie temperatures within the Gd. Such a situation could arise, for instance, if the Curie temperature were suppressed near the Gd/W interfaces. Indeed, the interfacial interaction between Fe and Gd in Fe/Gd superlattices causes the Gd moments near the interfaces to exhibit ferromagnetic order above the bulk Curie temperature and with a magnetic moment exceeding that of bulk Gd [18]. It is therefore conceivable that the Curie temperature would be suppressed at the Gd-W interface. An investigation of this sort will likely require depth profiling by polarized neutron reflectometry [19, 20].

**CONCLUSION**

We have investigated Gd/W multilayers in the context of the magnetocaloric effect. It is clear that nanostructuring Gd significantly impacts the behavior of this material. Relative to bulk Gd, 400 Å thick Gd films have a reduced entropy change peak value and enhanced entropy change full width at half maximum. A reduced Curie temperature and susceptibility exponent $\gamma$ of 1.75 suggest that finite size effects are impacting the transition, and may be related to the departure of the magnetic entropy change behavior in thin films from that of bulk material.

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