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CONDENSED MATTER PHYSICS | RESEARCH ARTICLE

Near-zero thermal expansion in magnetically ordered state in dysprosium at high pressures and low temperatures

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Abstract: The atomic volume of rare earth metal dysprosium (Dy) has been measured up to high pressures of 35 GPa and low temperatures between 200 and 7 K in a diamond anvil cell using angle dispersive X-ray diffraction at a synchrotron source. The hexagonal close-packed (hcp), alpha-Samarium (α -Sm), and double hexagonal close-packed (dhcp) phases are observed to be stable in Dy under high-pressure and low-temperature conditions achieved in our experiments. Dy is known to undergo magnetic ordering below 176 K at ambient pressure with magnetic ordering Néel temperature (T_N) that changes rapidly with increasing pressure. Our experimental measurement shows that Dy has near-zero thermal expansion in the magnetically ordered state and normal thermal expansion in the paramagnetic state for all the three known high pressure phases (hcp, α -Sm, and dhcp) to 35 GPa. This near-zero thermal expansion behavior in Dy is observed below the magnetic ordering temperature T_N at all pressures up to 35 GPa.

ABOUT THE AUTHORS

The high-pressure structural and magnetic phase transition study on rare earth metal dysprosium at high pressures and low temperatures is carried out by Yogesh Vohra's research group at the University of Alabama at Birmingham (UAB) and is supported by the US Department of Energy -National Nuclear Security Administration. This is part of a larger study on structural, electronic, and magnetic transitions in transition metals, rare earth metals and their alloys under extreme conditions of pressure and temperatures. Yogesh Vohra is the founding director of UAB Center for Nanoscale Materials and Biointegration (CNMB) that houses a Diamond Microfabrication Laboratory. Yogesh Vohra's research group has also pioneered several enabling technologies used in studies of materials under extreme conditions including diamond anvils with embedded sensors (designer diamonds) and nanocrystalline diamond micro-anvils grown by chemical vapor deposition techniques to study materials to pressures exceeding 500 GPa.

PUBLIC INTEREST STATEMENT

Rare earth elements are extensively utilized in a variety of applications including strong permanent magnets, lasers, lighting industry, automobiles, nuclear industry, and medicine. With one of the highest intrinsic magnetic moments (10.6 Bohr Magnetron) among the heavy rare earth elements, dysprosium (Dy) exhibits a rich magnetic phase diagram under high-pressure and low-temperature conditions. On cooling below 176 K, Dy shows spatial ordering of its magnetic moments on ions at the crystalline sites that are accompanied by unusual thermal expansion properties. Our paper reports on the relationship between magnetic ordering and near-zero thermal expansion observed in Dy at low temperatures. Furthermore, our X-ray diffraction measurements combined with magnetic ordering studies at high pressures and low temperatures reveal that the correlation between magnetic ordering and near-zero thermal expansion is observed in all known high pressure phases of Dy to 35 GPa. The connection between lack of thermal expansion and magnetic ordering is likely a universal phenomenon for rare earth metals under extreme conditions.









Subjects: Material Science; Materials Science; Physics

Keywords: magnetic ordering; rare earth metals; high pressures; low temperatures; thermal expansion

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1. Introduction

With one of the highest intrinsic magnetic moments (10.6 Bohr Magnetron) among the heavy rare earth elements, dysprosium (Dy) exhibits a rich magnetic phase diagram under high-pressure and low-temperature conditions. The structural phase transitions in 4f-lanthanide metals have been recently reviewed to ultra-high pressures up to 200 GPa and the crystal structure trends with increasing pressure have been well documented (Samudrala & Vohra, 2013). The crystallographic sequence $hcp \rightarrow \alpha$ -Sm $\rightarrow dhcp \rightarrow fcc$ – distorted fcc is observed with increasing pressure or decreasing atomic number in all lanthanide metals and is explained on the basis of number of d-electrons facilitated by electron transfer from the s-conduction band to the d-conduction band under high pressures (Duthie & Pettifor, 1977; Johansson & Rosengren, 1975). At the same time, lanthanide metal ions contain an un-filled f-shell and localized magnetic moments that show magnetic ordering at low temperatures due to interplay between strong correlation effects and indirect exchange effects involving their felectrons. The magnetic ordering transitions in heavy lanthanides have been studied by magnetic susceptibility measurements in a diamond anvil cell up to modest pressure of 10-20 GPa and a systematic decrease in magnetic ordering temperature with increasing pressure has been documented (Jackson, Malba, Weir, Baker, & Vohra, 2005). The recent innovations in measurements of magnetic ordering temperature in rare earth metals by electrical transport techniques have revealed complex magnetic behavior in Dy and have extended magnetic measurements to megabar pressures (Lim, Fabbris, Haskel, & Schilling, 2015; Samudrala, Tsoi, Weir, & Vohra, 2014). It has been shown previously (Samudrala et al., 2014) that the electrical transport measurement is a very sensitive technique for measuring magnetic ordering temperatures in rare earth metals and these measurements have been validated by direct observation of magnetic structures by neutron diffraction studies at high pressures and low temperatures (Thomas, Tsoi, Wenger, & Vohra, 2011; Thomas et al., 2013). The magnetic ordering temperature information coupled with atomic volume measurements by X-ray diffraction at high pressures and low temperatures allows for direct correlation between the thermal expansion and the magnetic ordering in 4-f metals under compression.

At ambient pressure, the paramagnetic (PM) to anti-ferromagnetic (AFM) transition at $T_{\rm N}=176~{\rm K}$ and AFM to ferromagnetic (FM) transition at $T_{\rm C}=87~{\rm K}$ are well documented in Dy (Legvold, Spedding, Barson, & Elliott, 1953). A helical magnetic structure is observed between 87 and 176 K at ambient pressure whereby the magnetization is always in the basal plane perpendicular to the c-axis but changes in direction from one hexagonal close-packed plane to the next with a turn angle that is temperature and pressure dependent. This turn angle has been measured at ambient pressure by neutron diffraction studies and increases almost linearly with temperature from 27 at 87 K to 43 per layer at 176 K (Wilkinson, Koehler, Wollan, & Cable, 1961). The magnetic ordering transition temperature in Dy shows a decrease with increasing pressure in the hcp-phase followed by a discontinuous drop in transition to the α -Sm phase (Samudrala et al., 2014). The magnetic ordering transition then undergoes through a minimum and then increases with further increase in pressure in the dhcp-phase to 35 GPa (Samudrala et al., 2014). Dy also shows unconventional magnetic state under ultra-high pressures as the magnetic ordering temperature increases rapidly above 73 GPa (above pressure of volume collapse) and attains a value greater than ambient temperature of 300 K at pressures above 107 GPa (Lim et al., 2015).



The anomalous thermal expansion and magnetostriction in single-crystal Dy at ambient pressure are well established (Clark, DeSavage, & Bozorth, 1965). In the paramagnetic phase, the thermal expansion is normal and as the temperature is lowered and the spins align and terms in thermal expansion attributed to thermal vibrations are dwarfed by those arising from the magneto elastic energy (Clark et al., 1965). As a result, c-axis expands on cooling below the anti-ferromagnetic phase, while a-axis shows normal contraction with decreasing temperature as reveled by direct X-ray diffraction studies at low temperatures and ambient pressure (Darnell & Moore, 1963). This gives rise to anomalous thermal expansion in Dy below the Néel temperature $T_{\rm N}$ and discontinuous changes in c-axis expansion and a-axis contraction on further cooling to ferromagnetic phase below 87 K (Darnell & Moore, 1963). Our experimental study is motivated by studying the effect of high pressure on this anomalous thermal expansion behavior in Dy. In particular, we are investigating the role of phase transitions to various close-packed structures under compression in Dy and their effect in anomalous thermal expansion behavior. Our experiments provide for a direct correlation between the structural data and the magnetic transition temperature measurements in Dy under simultaneous high-pressure low-temperature conditions.

2. Experimental details

X-ray diffraction experiments on polycrystalline Dy were carried out at the beam-line 16-BM-D, HPCAT, Advanced Photon Source at Argonne National Laboratory to low temperatures of 7 K using a diamond anvil cell. The Dy sample of 99.99% purity was placed in a sample chamber of 80 micron in diameter and 40 micron in thickness in a spring steel gasket along with a ruby pressure sensor in a diamond anvil cell. An angle-dispersive technique with an image-plate area detector was employed using an X-ray wavelength $\lambda = 0.4246$ Å. For simultaneous temperature- and pressure-dependent X-ray diffraction experiments, the diamond anvil cell (DAC) was cooled in a continuous helium flowtype cryostat, and the pressure in the cell was measured in situ with the ruby fluorescence technique (Tsoi et al., 2009; Yen & Nicol, 1992). The image plate XRD patterns were recorded with a focused X-ray beam of 6 µm by 13 µm (FWHM) on sample mixed with ruby, which served as the pressure marker. The Dy sample-to-image plate detector distance was calibrated using CeO, diffraction pattern and was measured to be 341.12 mm and was held constant throughout the entirety of the experiment. The software package FIT2D (Hammersley, 1995) was used to integrate the collected MAR345 image plate diffraction patterns which were analyzed by GSAS (Larson & Von Dreele, 2004) software package with EXPGUI interface (Toby, 2001) employing full-pattern Rietveld refinements to extract lattice parameters.

3. Results and discussion

The methodology we have utilized for determining Néel temperature (T_N) from electrical transport measurements and the effectiveness of this technique have been described in detail elsewhere (Lim et al., 2015; Samudrala et al., 2014; Thomas et al., 2011, 2013). Figure 1 shows four-probe electrical resistance data for Dy as a function of temperature at pressures of 1.1 GPa. The magnetic transition temperature has been determined by finding out the lowest point in the temperature derivative of resistance data (dR/dT) as shown in Figure 1. It can be readily seen that the derivative of the resistance has a clear minimum and this has been taken as the Néel temperature (T_N). The experimentally determined value of 175 K agrees very well with the ambient pressure T_N value of dysprosium, which is 179 K.

Figure 2 shows the magnetic ordering temperature as a function of temperature in various phases of Dy to 35 GPa. Dy shows an initial decrease in Néel temperature (T_N) with pressure at a rate of -5.7 Kelvin/GPa in agreement with previously published data (Jackson et al., 2005). The structural phase transition $hcp \rightarrow Sm$ occurs at 7 GPa, and $Sm \rightarrow dhcp$ transition occurs at 17 GPa as illustrated in the magnetic phase diagram in Figure 1. There is an abrupt drop in T_N on transition to Sm-phase at 7 GPa followed by a gradual decrease till 17 GPa before transition to dhcp-phase. T_N increases with



Figure 1. The four-probe electrical resistance measurement on the *hcp* phase of Dy as a function of temperature at a pressure of 1.1 GPa (left vertical axis).

Note: The anti-ferromagnetic transition temperature $(T_{\rm N})$ is marked by a resistance anomaly that is best represented by a minimum in the derivative of resistance with temperature (dR/dT) curve also shown (right vertical scale).

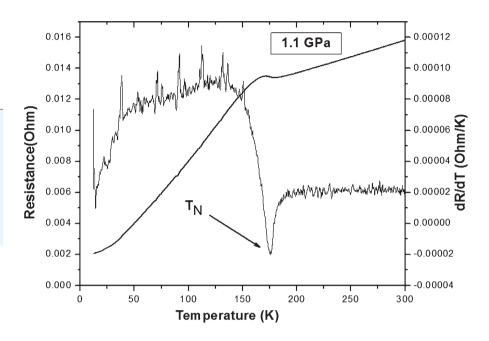
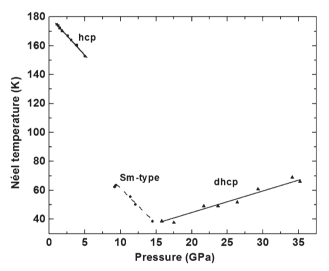


Figure 2. Magnetic ordering Néel temperature for Dy as a function of pressure up to 35 GPa with data adapted from Samudrala et al. (2014).

Note: The solid lines are the linear fit to the data as described in the text. The three known crystal structures *hcp*, *Sm-type*, and *dhcp* at high pressures and low temperatures for Dy are indicated in the graph.



increasing pressure in the *dhcp*-phase. The magnetic ordering Néel temperature (T_N) for Dy has been fitted as a linear function of pressure in Figure 2 for various phases as shown below for pressures up to 35 GPa.

$$T_{\rm N} = -5.7221P + 181.28 (hcp - phase)$$

$$T_{\rm N} = -4.6002P + 105.96(\alpha - Sm - {\rm phase})$$

$$T_{\rm N} = 1.6094P + 11.834 (dhcp - phase)$$

Figure 3 shows the image plate angle-dispersive X-ray diffraction patterns at high pressures up to 35 GPa and temperatures down to 7 K for hcp, α -Sm, and dhcp-phases for Dy. Figure 4 shows the integrated X-ray diffraction patterns for the hcp, α -Sm, and dhcp-phase of Dy that were analyzed using Rietveld refinement to obtain atomic volume at high pressures and low temperatures. The measured lattice parameters and refinement parameters at high pressures and low temperatures



Figure 3. Image plate angle-dispersive X-ray diffraction data for Dy at high pressures and low temperatures utilized in the measurements of thermal expansion. The X-ray wavelength λ = 0.4246 Å and Dy sample-to-image plate distance is 341.12 mm (a) Temperature 7 K and pressure 1.7GPa in the *hcp* phase, (b) temperature 7 K and pressure 8.6 GPa in the *Sm*-type phase, and (c) temperature 6.8 K and pressure 35.2 GPa in the *dhcp* phase.

Note: The fitted crystal structure data are shown in Table 1.

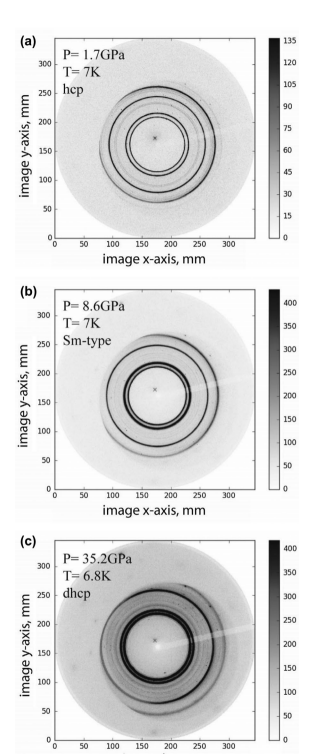


image x-axis, mm



Figure 4. Integrated X-ray diffraction patterns at high pressures and low temperatures for (a) hcp, (b) Sm-type, and (c) dhcp phases of Dy. The diffraction peaks are labeled with the corresponding Miller indices (hkl) and the measured lattice parameters are described in Table 1.

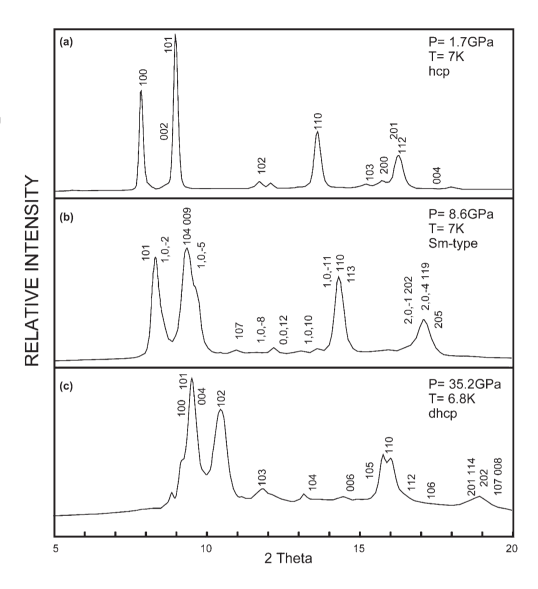


Table 1. The fitted lattice parameters for dysprosium at high pressures and low temperatures			
Crystal Structure	Experimental conditions (P, T)	Lattice parameters	Refinement parameter (reduced χ^2)
hcp	1.7 GPa, 7 K	a = 3.587, c = 5.601	0.658
		c/a = 1.562	
Sm-type	8.6 GPa, 7 K	a = 3.415, c = 23.574	2.626
		c/a = 6.903	
dhcp	35.2 GPa, 6.8 K	a = 3.100, c = 10.103	0.825
		c/a = 3.259	

for the three phases of Dy are shown in Table 1. The X-ray diffraction data collected at various pressures and low temperatures between 7 K and 200 K were collated to produce atomic volume maps for Dy.

Figure 5. The normalized change in volume ($\Delta V/V_s$) where V_o is the measured volume at the lowest temperature plotted as a function of temperature (T) at various pressures. The dashed lines represent the Néel temperature (T_N) at various pressures obtained from data in Figure 1. (a) Dy at 1.5 GPa in the hcp phase with $V_0 = 31.20 \text{ Å}^3/$ atom at 7 K, (b) Dy at 8.2 GPa in the α -Sm phase with $V_0 = 26.44 \text{ Å}^3/\text{atom at 7 K, (c)}$ dysprosium at 17.2GPa in the dhcp phase with $V_0 = 21.89 \text{ Å}^3/$ atom at 9 K, and (d) dysprosium at the highest pressure of 33.9GPa in the dhcp phase with $V_0 = 21.03 \text{ Å}^3/\text{atom at 9 K}.$

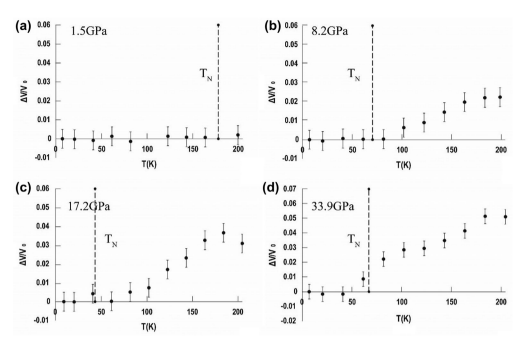


Figure 5 shows the normalized change in atomic volume ($\Delta V/V_o$) as a function of temperature (T) for various pressures as indicated. The ΔV is the change in volume relative to measured volume V_o at the lowest temperature (generally at 7 K). The data in Figure 5 cover four different pressures of 1.5 GPa (T_N = 173 K), 8.2 GPa (T_N = 68 K), 17.2 GPa (T_N = 40 K), and 33.9 GPa (T_N = 66 K). The experimentally measured magnetic ordering temperature (T_N) obtained from fits described earlier (Figure 2) at these pressure is shown as dashed lines in Figure 5. The experimental data at 1.5 GPa for the hcp phase show near-zero thermal expansion almost in the entire temperature range with some increase above 173 K. It is clear from Figure 5 that the change in atomic volume with temperature is near-zero for temperature below T_N and rises gradually above the magnetic ordering temperature in the paramagnetic phase. This phenomenon appears to be universal for all high-pressure phases including hcp, α -Sm, and dhcp-phase of Dy to 35 GPa. It should be added that the details of the magnetic ordering in Sm-phase and dhcp-phase of Dy have not yet been established by neutron diffraction at high pressures and low temperatures. However, the phenomenon of near-zero thermal expansion in Dy appears to be broadly connected with magnetic ordering and is independent of the details of the magnetic structures at high pressures and low temperatures.

In summary, we have shown that the rare earth metal dysprosium exhibits near-zero thermal expansion with temperature in the magnetically ordered phase below Néel temperature (T_N). This correlation is obtained by combining electrical resistance data and X-ray diffraction data obtained at high pressures and low temperatures. This anomalous thermal expansion phenomenon is universal and is observed for all high-pressure phases including hcp, α -Sm, and dhcp to 35 GPa. The anomalous thermal expansion behavior tracks the magnetic transition temperature in dysprosium at high pressures. It is now possible to correlate thermal expansion data at high pressure and low temperature for rare earth metals to the magnetic ordering temperatures determined by electrical transport measurements.

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