About the Role of Fe-Ions in the Formation of Magneto caloric Effect in Ho(Co$_{1-x}$Fe$_x$)$_2$ Compounds

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A study of crystalline structure, magnetic and magneto caloric properties of Ho(Fe$_{0.9}$Co$_{0.1}$)$_2$ ($x = 0.09, 0.12$) intermetallic compounds has been undertaken. Phase composition was controlled by X-ray diffraction analysis. Magnetic properties were measured within the temperature range 4.2-350 K in magnetic fields up to 7 T. Magnetic ordering temperatures corresponding to paramagnetic-ferrimagnetic phase transitions were found to be 199 K and 258 K respectively. Temperature dependences of heat capacity for these compounds have been inferred for the temperature interval 77-340 K. Comparison of magneto caloric effect (MCE) values determined by direct measurement and by calculation was carried out as well. It was found that significant MCE peak broadening occurs for higher iron concentration in the compound.

DOI: 10.12693/APhysPolA.127.635
PACS: 75.30.Sg, 75.50.Gg

1. Introduction

RT$_2$ type intermetallic compounds (where R — rare earth, T = Fe, Co, Ni) or Laves phases with cubic crystalline structure have been studied intensively for more than 3 decades, primarily because of the giant magnetostrictive values observed for the iron-based alloys [1].

Some RT$_2$ alloys based on Co, along with the magnetostrictive effect, demonstrate the significant magneto caloric effect (MCE) near the Curie temperature ($T_C$), which is applicable for magnetic refrigeration technology [2]. Early MCE studies on RCo compounds by direct measurements were carried out on the samples of binary HoCo$_2$ alloy [3].

Later, many authors [4-8] have been carried out indirect MCE estimations on RCo$_2$ compounds using isotherms of magnetization curves and specific heat data.

The RT$_2$ type intermetallic alloys are ferrimagnets with antiparallel orientation of R and T magnetic moments in case of heavy R. The magnetic behavior and magneto caloric properties of these compounds are determined by competitive exchange interactions within subsystem (rare earth or 3d-metal) as well as between R and T subsystems.

Recently, the study of Ho(Ni$_{1-x}$Fe$_x$)$_2$ quasibinary system revealed that the nickel substitution by iron leads to abnormal MCE [9]. This anomaly manifested as MCE presence in much wider temperature range, contrary to the many known magneto caloric materials. Later, a similar MCE behavior was observed in Tb(Ni$_{1-x}$Fe$_x$)$_2$ [10] and Er(Co$_{1-x}$Fe$_x$)$_2$ [11] compounds. Authors [12], investigating structure and magnetothermal properties of the Tb(Co$_{1-x}$Fe$_x$)$_2$ system also found a significant magnetic entropy temperature range broadening in the TbCo$_{0.9}$Fe$_{0.1}$ compound compared with that in TbCo$_2$.

The neutron diffraction studies carried out in wide temperature range indicated existence of the structural phase transition (from rhombohedral to cubic with decreasing temperature) associated with the magnetic phase transition and broadening of the transition temperature interval when Fe is added.

These results prompted us to undertake the study of structural, magnetic and magneto-thermal properties of alloys similar to mentioned above - Ho(Fe$_{0.9}$Co$_{0.1}$)$_2$ system with low Fe content ($x = 0.09, 0.12$), which have not been investigated considered in literature from such point of view.

2. Experimental details

Ho(Fe$_{0.9}$Co$_{0.1}$)$_2$ alloys ($x = 0.09, 0.12$) were melted in induction furnace using a quartz crucible with an argon protective atmosphere. A homogenizing annealing of alloys was made in a vacuum furnace at 1220 K for six hours. Powders of the alloy were obtained by mechanical milling in a mortar with the 200-500 μm fraction sieved. Phase composition control and crystalline structure studies were carried out by means of X-ray diffraction technique (diffractometer Bruker D8 Advance) with CuK$_{α1}$-radiation source. For X-ray diffraction patterns analysis FullProf v.2.05 software was used.

Magnetic phase analysis and magnetization curves measurements were made using SQUID-magnetometer MPMS-XL-7 EC (Quantum Design) in the temperature range from 4.2 K to 350 K in magnetic field up to 7 T. Heat capacity was measured at zero magnetic field in the temperature range from 77 K to 340 K.

Direct measurement of the Δ$T$ effect was carried out with the help of a proprietary experimental apparatus in the magnetic field with 0.46 T magnetic field applied. To exclude the displacement of powder particles under study in the process of measurement, these were yielded into tablets, by mixing powders with epoxy adhesive and
pressing. To account the influence of epoxy adhesive on the MCE, in a similar way the reference sample of Gd metal with the known value of the MCE for the chosen magnetic field was made [13]. A copper-constantan thermocouple was inserted into this sample. The signal from it was sent to a DC millivoltmeter, and from there to a computer.

3. Results and discussion

Analysis of the room temperature X-ray diffraction data showed that all samples within the study were almost single-phase. The 1:2 composition phase had a cubic structure of MgCu2 type with Fd3m space group. Its crystal lattice parameter (a) has been determined (Table).

Samples were confirmed to contain only 1:2-phase by thermomagnetic analysis performed in the external magnetic field (H) of 0.5 T (Fig. 1). Specific magnetization values corresponding to the magnetic field μ0H = 7 T (M5) at 4.2 K and the magnetic moment per formula unit (M_FU) were calculated from M5 values (in Bohr magneton — μB), which are shown in Table. Temperatures of magnetic transitions for different samples (Table) were determined in the position of the first derivative’s (dM/dT) peaks on the temperature axis, taken from the specific magnetization temperature dependencies (M(T)) in the magnetic field, with induction of 0.01 T.

Fig. 1. Temperature dependencies of Ho(Co1-xFe2x)2 compounds magnetization measured in the magnetic field ∆μ0H = 0.5 T. Arrot curves for these compounds at corresponding Curie temperatures are shown in inset.

Type of the magnetic phase transition was determined using magnetic criterion proposed by Banerjee [14]. According to that, a positive value of the tangent angle to the Arrot isotherms at any point in the ferromagnetic state indicates the second order magnetic phase transition. Thus, based on the analysis of the experimental data (inset in Fig. 1) it can be argued that the phase transition in these compounds is a second order one.

For the calculation of isothermal entropy change(ΔS), which can be made using formulae (1) [12, 13], the samples magnetization field dependencies in the external magnetic fields (0–7 T) in the wide temperature range with a step of 5–10 K were performed

\[
ΔS(H, T) = \sum_i M_{i+1}(T_{i+1}, H) - M_i(T_i, H) \frac{T_{i+1} - T_i}{T_{i+1} - T_i} ΔH. \tag{1}
\]

Figure 2 shows the calculated values of ΔS data for the change of magnetic field ∆μ0H, = 7 T. The figure shows that with iron concentration increasing, value Δ\(T_{\text{FWHM}}\), characterizing the difference between higher and lower temperatures at half maximum of the −ΔS(T) peak, increases. Similar trend of \(\Delta T_{\text{FWHM}}\) increasing with magnetic field is observed for all Ho(Co1-xFe2x)2 samples. ΔS/ΔS_Tc(T), where ΔS_Tc is ΔS at Tc for the sample with x = 0.12 at various magnetic fields is shown in the inset I of Fig. 2. The inset II shows the values of Δ\(T_{\text{FWHM}}\) for R(M1-xFe2x)2 compounds, where M = Co or Ni. Data have been taken from the papers [9-12, 15] and from the present study. The figure shows that for all R(M1-xFe2x)2 compounds with low Ni or Co substitutions by Fe, a significant increase of ΔS(T) peak width is observed. For sample with x = 0.12Δ\(T_{\text{FWHM}}\) = 255 K. Availability of ferromagnets with MCE in the wide temperature range

<table>
<thead>
<tr>
<th>x</th>
<th>a [Å]</th>
<th>(M_s) [Am²/kg]</th>
<th>(M_{FU}) [µB per formula unit]</th>
<th>(T_c) [K]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.09</td>
<td>7.1877</td>
<td>140.4</td>
<td>7.10</td>
<td>199</td>
</tr>
<tr>
<td>0.12</td>
<td>7.1908</td>
<td>137.5</td>
<td>6.95</td>
<td>258</td>
</tr>
</tbody>
</table>

Table: Crystal lattice parameters (a), magnetization values in the magnetic field of 7 T (\(M_s\)), magnetic moment of formulæ unit in 7 T (\(M_{FU}\)), and magnetic transition temperatures (\(T_c\)), for Ho(Co1-xFe2x)2 compounds on Fe concentration (x).
is essential for potential use in magnetic refrigerants.

Figure 3 shows the temperature dependence of the magnetocaloric effect \( \Delta T \) in \( \text{Ho}(\text{Co}_{1-x}\text{Fe}_x)_2 \) compounds at various magnetic fields. For calculation of MCE values (\( \Delta T_{\text{calc}} \)), the following formulae were used [13]:

\[
\Delta T_{\text{calc}}(T) = -\frac{T}{C_{P, H}(T)} \Delta S_H(T),
\]

where \( C_{P, H} \) is the heat capacity of the sample material at the DC magnetic field, \( \Delta S_H \) is the change of magnetic entropy part. Because of the absence of data on the temperature dependences of heat capacity for the DC magnetic field of 0.5 T, the calculation of \( \Delta T_{\text{calc}} \) was performed using the value of \( C_{P, H} \) under \( \mu_0 H = 0 \) (inset Fig. 3). For comparison, in this figure values of \( \Delta T(T) \) measured by direct method (\( \Delta T_{\text{direct}} \)) are shown. It can be concluded that the calculated values are close to those obtained by direct measurements. Differences between calculated and measured \( \Delta T \)-effect values may appear due to not taking into account the magnetic contribution to the specific heat.

To give more detailed magnetocaloric characteristics of the studied compounds, a comparison of their relative cooling capacity (RCP) [12] with the RCP of the similar compound was made:

\[
\text{RCP} = \frac{T_2}{T_1} \int_0^{T_1} \Delta S(T) dT,
\]

where \( T_1 \) and \( T_2 \) are the lowest and the highest temperatures on the \( T \)-axis, corresponding to half-height maximum of \( \Delta S(T) \) dependence. RCP value of the \( \text{Ho}(\text{Co}_{0.86}\text{Fe}_{0.14})_2 \) compound in magnetic field of 5 T is about 612 J/kg, which is significantly higher than for the \( \text{Tb}(\text{Co}_{0.97}\text{Fe}_{0.03})_2 \) compound, where RCP = 299 J/kg (\( \Delta \mu_0 H = 5 \) T) [12].

A significant increase in \( \Delta S(T) \) peak width observed for \( \text{Ho}(\text{Ni}_{1-x}\text{Fe}_x)_2 \), \( \text{Tb}(\text{Ni}_{1-x}\text{Fe}_x)_2 \), and \( \text{Tb}(\text{Co}_{1-x}\text{Fe}_x)_2 \) compounds according to the authors [9, 10, 12] may be due to both randomization of rare-earth magnetic moments orientations in the R sublattice and crystal structure transition in the mentioned temperature range.

Presented study shows similar behavior of the change of magnetic part of the entropy in quasi-binary \( \text{Ho}(\text{Co}_{1-x}\text{Fe}_x)_2 \) compounds. Such behavior of magneto-thermal properties of magnetic materials is very important from both practical and scientific points of view. In order to clarify unambiguously the role of substitute of base Co or Ni atoms by Fe in \( \text{R}(\text{M}_{1-x}\text{Fe}_x)_2 \) intermetallic compounds with heavy Rare Earth elements, additional studies are required.

**Acknowledgments**

This work has been supported by the UrFU State contracts Nos. 2582 and 1362.

**References**


