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Synthesis, structure, and magnetic and dielectric properties of magnetoelectric BaDyFeO4 ferrite

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Abstract

BaDyFeO₄ was prepared by a conventional solid-state method in air at 1573 K. It crystallizes in space group *Pnma* (No. 62) with a = 13.16861(1) Å, b = 5.70950(1) Å, and c = 10.26783(1), and it is isostructural with BaYFeO₄. Three magnetic transitions were found in BaDyFeO₄ at $T_{N3} = 9$ K, $T_{N2} = 23$ K, and $T_{N1} = 47$ K in zero magnetic field in comparison with two magnetic transitions observed in BaYFeO₄. Magnetic-field-induced transitions were also detected in BaDyFeO₄ at 18 and 28 kOe (at T = 1.8 K). Frequency-dependent broad dielectric peaks were observed in BaDyFeO₄ spanning between T_{N2} and T_{N1} and centred at 35 K – this temperature does not coincide with any T_N . No dielectric anomalies were found at T_{N1} and T_{N3} , while very weak frequency-independent dielectric anomalies were detected at T_{N2} . Positive and negative magnetodielectric effects were measured in BaDyFeO₄ (within a range of -0.8 and +0.4 % up to 90 kOe) reflecting magnetic-field dependence of dielectric constant. Pyroelectric current measurements did not detect any ferroelectricity in BaDyFeO₄ under measurement conditions used. No dielectric anomalies and no magnetodielectric effects were found in BaYFeO₄.

Keywords: ferrites; magnetodielectric effect; multiferroics; crystal structure

1. Introduction

A phenomenon of spin-induced ferroelectricity has recently received a lot of attention [1-3]. It usually takes place when the symmetry of a long-range magnetic structure is polar. In more general words, it requires non-trivial, complex magnetic structures, for example, sinusoidal-type and cycloid-type orderings [3]. Non-trivial magnetic structures could be realized (1) in systems with very complex networks of physical connections among magnetic ions and (2) in systems with simple networks of physical connections among magnetic ions (for example, in perovskites with three-dimensional connections of corner-shared octahedra) but with the presence of spin frustration or complex networks of spin exchange interactions [1-3]. Spin-induced ferroelectricity has been discovered in many previously known magnetic materials.

The exploration of new materials with complex connection networks of magnetic ions is a way to expand the family of spin-induced multiferroics. A new barium yttrium ferrite, BaYFeO₄, was recently reported with a complex magnetic sublattice [4, 5]. In BaYFeO₄, there are [Fe₄O₂₈] clusters or rings formed by corner-shared FeO₅ square pyramids and FeO₆ octahedra; the rings, in turn, share corners to form one-dimensional chains along the *b* axis (Figure 1b). BaYFeO₄ is isostructural with Ba₂CuPtY₂O₈ and related compounds [6]. Neutron powder diffraction and magnetic susceptibility measurements showed the existence of two magnetic transitions in BaYFeO₄ with $T_{N2} = 33$ K and $T_{N1} = 48$ K [4, 5]. Below T_{N1} , a spin-density wave structure was suggested, and below T_{N2} - an incommensurate cycloid structure [5, 7]. Very weak spin-induced ferroelectricity was found below T_{N2} [8]. Spin-glass-like behavior and relaxation of magnetization below 17 K were reported in Ref. 9. There are also some puzzling behaviours of BaYFeO₄, for example, (1) the absence of any specific heat anomalies at T_{N1} and T_{N2} despite a large spin of Fe³⁺ (*S* = 5/2) indicating a very small entropy release at the magnetic transition temperatures [4] and (2) the absence of Curie-Weiss behavior, reported in all papers [4, 5, 8, 9], up to 650 K [5].

Ferrites show rich crystal chemistry and magnetism [10]. There are many ferrites with different compositions in Ba-R-Fe-O systems, where R is a rare-earth cation and Y [11], for

example, YBaFe₄O₇, Ba₃YFe₂O_{7.5}, YBa₂Fe₃O₈, YBa₂FeO_{5.5}, and YBaFe₂O₅. In Ba-Dy-Fe-O system, DyBaFe₄O₇ [10], DyBaFe₂O₅ [12], and DyBa₂Fe₃O₈ [13] have been reported. Ferrites with such a simple composition, BaRFeO₄, are reported only for R = La and Y, and BaLaFeO₄ crystallizes in a different crystal structure [14]. This fact suggests that at least two different structure types are realized in BaRFeO₄ depending on the size of rare-earth elements, and BaRFeO₄ ferrites can exist for other rare-earth elements.

Therefore, in the present work, we extended the family of BaRFeO₄ simple ferrites and report on the synthesis, crystal structure, and properties of BaDyFeO₄. We selected Dy^{3+} as it has the close ionic radius with that of Y^{3+} . In addition, Dy^{3+} has the largest magnetic moment among rare-earth cations, and the appearance of an additional magnetic sublattice could significantly modify properties of BaRFeO₄ ferrites. We also report on some properties of BaYFeO₄. First, in order to understand effects of the Dy³⁺ sublattice in BaDyFeO₄ it is necessary to make comparison between BaDyFeO4 and BaYFeO4, but some properties of $BaYFeO_4$ have not been reported yet, such as, magnetic-field dependence of specific heat and magnetodielectric effects. Second, more experimental information is needed to understand intrinsic properties of BaYFeO₄, especially the influence of synthesis conditions and routes and the sample quality on properties. We found that BaDyFeO4 shows a more complex behavior than that of BaYFeO4 from the viewpoint of magnetism and lowtemperature dielectric properties. We observed positive and negative magnetodielectric effects in BaDyFeO₄ due to magnetic-field dependence of dielectric constant, but no magnetodielectric effects in BaYFeO4. No signs of spin-induced ferroelectricity were detected in BaDyFeO₄ within measurement conditions used and the sensitivity of our methods.

2. Experimental

BaDyFeO₄ and BaYFeO₄ were prepared from stoichiometric mixtures of BaCO₃, Y₂O₃, Dy₂O₃, and Fe₂O₃, where we used about 25 % of Fe₂O₃ enriched by ⁵⁷Fe (95.5 %) in case of BaDyFeO₄ (for future Mössbauer studies). Y₂O₃ and Dy₂O₃ were dried at 1270 K before use, Fe₂O₃ – at 1170 K, and BaCO₃ – at 870 K. The mixtures were pressed into pellets and

annealed in air on Pt foil at 1430 K for 48 h, 1520 K for 18 h, and 1570 K for 38 h in case of BaDyFeO₄ and at 1430 K for 48 h and 1520 K for 48 h in case of BaYFeO₄ (with grinding after each step). After such procedures, BaDyFeO₄ contained about 2 weight % of DyFeO₃ impurity; however, BaYFeO₄ contained about 7.7 wt. % of BaFeO₃ and 2.2 wt. % of Y_2O_3 impurities. Therefore, BaYFeO₄ was additionally annealed at 6 GPa and about 1420 K for 90 min in an Au capsule using a belt-type high-pressure machine (heating time to the synthesis temperature was 10 min). After the high-pressure heat treatment, the sample was quenched to room temperature (RT), and the pressure was slowly released. After the high-pressure treatment, BaYFeO₄ contained about 2.0 wt. % of YFeO₃ impurity. We note that additional annealing of BaYFeO₄ at 1570 K for 48 h in air (after annealing at 1430 K for 48 h and at 1520 K for 48 h in air, the sample was split into two batches: the first one was for the HP treatment, the second one – for the additional annealing in air) did not significantly improve the sample quality in our case as it contained about 3.9 wt. % of BaFeO₃, 0.6 wt. % of Y₂O₃, and 1.4 wt. % of YFeO₃ impurities.

X-ray powder diffraction (XRPD) data were collected at RT on a RIGAKU MiniFlex600 diffractometer using CuK α radiation (2 θ range of 8–110°, a step width of 0.02°, and a scan speed of 1 deg/min). XRPD data were analysed by the Rietveld method using *RIETAN-2000* [15]. Weight fractions of impurities were estimated from refined scale factors by *RIETAN-2000* after the Rietveld analysis [15].

Synchrotron XRPD data of BaDyFeO₄ were measured at RT on a large Debye-Scherrer camera at the undulator beamline BL15XU of SPring-8 [16, 17]. The intensity data were collected between 3° and 71.40° at 0.003° intervals in 20; the incident beam was monochromatized at $\lambda = 0.65298$ Å. The sample was packed into a Lindemann glass capillary (inner diameter: 0.1 mm), which was rotated during the measurement. The absorption coefficient was also measured. The Rietveld analysis was performed using the *RIETAN-2000* program [15].

Magnetic measurements were performed on SQUID magnetometers (Quantum Design, MPMS-1T and MPMS-XL-7T) between 2 and 350 K (or 400 K) in different applied fields under both zero-field-cooled (ZFC) and field-cooled on cooling (FCC) conditions. Isothermal magnetization measurements were performed between -70 and 70 kOe at

different temperatures. Specific heat, C_p, at different magnetic fields (0-90 kOe) was recorded between 2 and 300 K on cooling and heating by a pulse relaxation method using a commercial calorimeter (Quantum Design PPMS). Dielectric properties were measured using a NOVOCONTROL Alpha-A High Performance Frequency Analyzer between 3 and 300 K on cooling and heating in the frequency range of 100 Hz and 2 MHz and at different magnetic fields; both magnetic and electric fields were perpendicular to the flat surfaces of pellets. Pyroelectric current measurements were done with a Keithley 6517B electrometer. Poling in different positive and negative electric fields was performed on cooling from 100 K to 2 K at different magnetic fields; at 2 K, the electric field was removed and electrodes were shorted. Measurements (with a heating rate of 8.1 K/min for BaDyFeO₄ and 4.3 K/min for BaYFeO₄) were started after the background current was below 1 pA for more than 5 min. In case of BaDyFeO₄, bias electric field measurements were also used, when the sample was cooled to 2 K under a zero electric field; at 2 K, an electric field of 390 kV/m was applied; and current was measured on heating with a rate of 8.1 K/min. Temperature and magnetic fields were controlled by a PPMS. Pieces of pellets were used in all magnetic, specific heat, dielectric, and pyroelectric current measurements.

Time-dependent magnetization relaxation was measured on MPMS 1T under the following protocol: a sample was cooled from 100 K to a measurement temperature in a nominal zero magnetic field (after a reset magnet procedure). At the measurement temperature, a magnetic field of 1 kOe was applied (in the no-overshoot mode), and magnetization was measured as a function of time. In Ref [9], a different protocol was used, when a sample was cooled under 1 kOe, and magnetization was measured under nominal zero magnetic field. However, we did not use this protocol because there are always trapped fields inside magnetometers. Values and signs of trapped fields depend on a magnetometer, magnetic-field history, and how the zero field was set (there are oscillate and no-overshoot modes on Quantum Design MPMS). These features deteriorate the reproducibility. Moreover, magnetization should be zero under real zero magnetic field in pure antiferromagnets, and any measurable magnetization will originate from non-zero real magnetic field or extrinsic contributions.

3. Results and Discussion

BaDyFeO₄ was found to be isostructural with BaYFeO₄. Therefore, we used structural data of BaYFeO₄ [4] as the initial model for the crystal structure refinement of BaDyFeO₄ by the Rietveld method. The crystal structure parameters of BaDyFeO₄ are summarized in Table 1. Primary bond lengths and bond-valence sums (BVS) [18] and given in Table 2. Figure 2 shows the fitting results with experimental, calculated, and difference synchrotron XRPD patterns. The BVS values of +2.1 (for Ba²⁺), +3.1 (for Dy³⁺), and +2.8 (for Fe³⁺) are close to the expected values. The crystal structure of BaDyFeO₄ is illustrated on Figure 1. The iron sublattice was described in the introduction. There is a Dy magnetic sublattice in BaDyFeO₄, illustrated on Figures 1c and 1d, in comparison with BaYFeO₄. The Dy1 and Dy2 sites have seven-fold oxygen coordination environments. The Dy1O₇ and Dy2O₇ polyhedra are joined by edges forming one-dimensional chains along the *b* axis. While chains of the Fe1O₅ and Fe2O₆ polyhedra are isolated from each other (Figure 1a), there are corner-shared connections between chains of the Dy1O₇ and Dy2O₇ polyhedra (Figure 1c).

Magnetic susceptibilities of BaYFeO₄ and BaDyFeO₄ are shown on Figure 3. Two magnetic transitions were observed in BaYFeO₄ at $T_{N1} = 47$ K and $T_{N2} = 35$ K at magnetic fields from 100 Oe to 70 kOe. The transitions could be more clearly seen on the $d(\chi T)/dT$ versus *T* curves (the inset of Figure 3a). No Curie-Weiss behavior was observed in BaYFeO₄, also in agreement with previous reports, and magnetic susceptibilities were field-dependent up to 400 K. This behavior is a strong indication of the presence of a ferromagnetic-like impurity. We could clearly detect YFeO₃ impurity in our sample by XRPD. YFeO₃ has strong ferromagnetic-like properties (due to canting of antiferromagnetically aligned spins), and it has $T_N \approx 655$ K [19]. Traces of YFeO₃ impurity (or other magnetic impurities) in the previous reports could be a reason for the absence of the Curie-Weiss behavior up to 650 K [5], and the absence of the Curie-Weiss behavior could in our work ($T_{N1} = 47$ K and $T_{N2} = 35$ K) are in very good agreement with previous reports.

This fact gives strong evidence that the reported χ versus *T* curves of BaYFeO₄ are sampledependent (or more precisely, strongly impurity-dependent), and the intrinsic behavior has yet to be determined.

Five anomalies could be seen on the $d(\chi T)/dT$ versus T curves of BaDyFeO₄ (the inset of Figure 3b). No difference was observed between ZFC and FCC curves above 1 kOe; therefore, only one of the two curves is shown (Figure 3). The anomalies near 59 K (and a small difference between the ZFC and FCC curves at 100 Oe) could originate from a Fe³⁺ spin reorientation transition in DyFeO₃ impurity, and the anomalies at 4 K – from a Dy³⁺ ordering transition in DyFeO₃ impurity [20]. The anomalies near 59 K disappeared above 1 kOe in agreement with strong suppression of the spin reorientation transition in DyFeO₃ by magnetic fields [21]. Other anomalies at 9, 23, and 47 K should be related to magnetic transitions in the main BaDyFeO₄ phase. The transitions at $T_{N1} = 47$ K and $T_{N2} = 23$ K could originate from orderings of the Fe³⁺ sublattice. The transition at $T_{N3} = 9$ K is probably due to ordering of the Dy³⁺ sublattice. A noticeable field-dependence was present in the paramagnetic temperature range of BaDyFeO₄ up to 400 K (Figure S1) similar to BaYFeO₄; this behavior could be caused by the presence of DyFeO₃ impurity. Nevertheless, the Curie-Weiss behavior was observed in the paramagnetic temperature range with the estimated effective magnetic moment of $11.28 \mu_{\rm B}/f.u.$ and the Curie-Weiss temperature of -34 K (we used an FCC χ^{-1} versus T curve measured at 70 kOe from 395 to 250 K for fitting), where the calculated effective magnetic moment of BaDyFeO₄ is $12.14 \mu_B/f.u.$ (Figure S1a). The observation of the Curie-Weiss behavior is probably caused by a large moment of Dy³⁺ cations, which dominates over DyFeO₃ impurity at high magnetic fields. γ versus T curves of BaDyFeO₄ strongly depended on the measurement magnetic field (Figure 4a). Anomalies on the $d(\chi T)/dT$ versus T curves remained the same at T_{N3} up to about 18 kOe; they were gradually smeared at T_{N1} with the increase of the measurement magnetic field; on the other hand, the anomalies showed strong field-dependence at T_{N2} with sharp peaks at 23 K at 1 kOe, 20 K at 5 kOe, 13 K at 10 kOe, and 10.5 K at 14 kOe (Figure 4b). Above 18 kOe, sharp peaks at T_{N2} disappeared; and above about 26 kOe, the χ versus T curves became featureless. We note the presence of a kink on the $d(\gamma T)/dT$ versus T curves at 19 K at 22 kOe, which

was still visible at 20.5 K at 9 kOe (Figure S1b). Therefore, BaDyFeO₄ should show a complex temperature–magnetic-field phase diagram in comparison with BaYFeO₄. However, the construction of such a phase diagram is out of the scope of the present work, and single crystals are desirable for the construction of a precise phase diagram.

Isothermal magnetization curves (M versus H) of BaDyFeO₄ at 5, 15, and 30 K are shown on Figure 5. The M versus H curves at 1.8 K and 5 K were very similar to each other (Figure S2). However, the dM/dH versus H curve at 1.8 K (inset of Figure 5) shows more clearly the presence of three field-induced transitions at 6, 18, and 28 kOe. The weak transition at 6 kOe matches well with a field-induced transition in DyFeO₃ and could be caused by this impurity [20]. Moreover, the field-induced transition at 6 kOe disappeared at T = 5 K above $T_{N,Dy} = 4$ K of DyFeO₃ (Figure S2). Much stronger transitions at 18 and 28 kOe should originate from the main BaDyFeO₄ phase, and these values coincide with the field values where the χ versus *T* curves qualitatively changed (Figure 4b). The M versus H curve of BaYFeO₄ at 5 K demonstrated a small hysteresis near the origin, which could originate from YFeO₃ impurity (Figure S2), without any field-induced transitions. Therefore, the presence of the magnetic Dy³⁺ sublattice noticeably enriches magnetic behaviors.

Spin-glass-like magnetic properties below 17 K and magnetization relaxation reaching about 35 % at T = 9 K and H = 0 Oe were observed in BaYFeO₄ [9]. However, we emphasize that magnetization relaxation was only measured between 2 and 9 K, and no results were reported above 9 K [9]. We also observed magnetization relaxation in our BaYFeO₄ sample, but the magnetization relaxation reached maximum 3 % under H = 1 kOe (Figure S1c). In addition, the relaxation was observed at all temperatures between 5 and 50 K, that is, above T_{N1} . Therefore, we conclude that the magnetization relaxation in our BaYFeO₄ sample is an extrinsic property. No magnetization relaxation was observed in BaDyFeO₄ at 5, 10, and 20 K (Figure S1c).

Specific heat data of BaYFeO₄ and BaDyFeO₄ are shown on Figure 6. No anomalies were observed in BaYFeO₄ in agreement with one of the previous reports [4]. Surprisingly, a magnetic field of 90 kOe had no effects on specific heat data of BaYFeO₄ suggesting that AFM states of BaYFeO₄ are very robust. On the other hand, a very small specific heat anomaly was observed at T_{N2} in BaDyFeO₄, no anomalies at T_{N1} , and a very strong and

broadened anomaly at T_{N3} (an anomaly was also observed at $T_{N,Dy} = 4$ K from the Dy³⁺ ordering transition in DyFeO₃ impurity). Therefore, the entropy release at the Fe³⁺ ordering transitions was also very small in BaDyFeO₄ similar to BaYFeO₄. A magnetic field of 90 kOe showed its effect on specific heat of BaDyFeO₄ below a very high temperature of about 130 K, far above T_{N1} . This fact could indicate that a magnetic field strongly polarizes the Dy³⁺ sublattice far above T_{N1} . A magnetic field of 90 kOe completely suppressed the anomaly at T_{N3} . The broadened anomaly at T_{N3} and its suppression by a magnetic field indicate that it is caused by Dy³⁺ ordering.

Frequency-dependent dielectric data of BaDyFeO4 are presented on Figure 7a and Figures S3-S6. Dielectric constant starts increasing below about 50 K, which is close to $T_{\rm N1}$ = 47 K, but without any clear anomalies at $T_{\rm N1}$, and it shows maxima near 35 K. Then, dielectric constant decreases below 35 K, and a broad peak suddenly disappears at $T_{N2} = 23$ K. There are very small dielectric peaks at T_{N2} (which look frequency-independent), especially on heating curves (Figures 7a and 8a). The peak positions near 35 K are slightly frequency-dependent. Dielectric loss also demonstrates peaks spanning from 23 to 38 K (Figure 7b). Therefore, broad dielectric anomalies are present in a magnetic phase between T_{N2} and T_{N1} with clear hysteresis on cooling and heating at H = 0 Oe, suggesting a first-order phase transition at T_{N2} . Magnetic fields have noticeable effects on temperature dependence of dielectric constant (Figure 8a) resulting in the observation of positive and negative magnetodielectric effects (Figure 8b). For example, broad dielectric anomalies move to lower temperatures at H = 10 kOe, especially from the low temperature side in agreement with the magnetic susceptibility data (Figure 4b) and the movement of T_{N2} by magnetic fields. In other words, broad dielectric anomalies span again between T_{N2} and T_{N1} at H = 10 kOe with hysteresis on cooling and heating. On the other hand, no dielectric anomalies were observed in BaYFeO₄ (Figure 9a), and the magnetodielectric effect was negligible (Figure 9b). Therefore, the presence of the magnetic Dy^{3+} sublattice noticeably enriches dielectric behaviors of BaDyFeO₄ in comparison with those of BaYFeO₄.

Temperature dependence of dielectric constant of $BaYFeO_4$ was qualitatively similar with that of Ref. 8, including the observation of a small kink near 25 K (Figure 9a). However, the absolute values were different probably because of uncertainties in pellet and

electrode dimensions. Therefore, dielectric properties of BaYFeO₄ are not so sample sensitive in comparison with the χ versus *T* curves. In Ref. 8, difference in dielectric constant of the order of 2×10^{-3} at $T_{N2} = 35$ K was found between the curves measured at H = 0 and 70 kOe, while no dielectric constant anomalies were detected at H = 70 kOe. Our measurements were less sensitive; this could be a reason why we did not observe any dielectric anomalies in our BaYFeO₄ sample at $T_{N2} = 35$ K and H = 0 Oe and other magnetic fields.

We did not observe any sharp pyroelectric current anomalies in BaDyFeO₄ and BaYFeO₄ (Figures 10 and 11 and Figure S8), which could be assigned to a ferroelectric transition. There were some very broad anomalies, and they reproduced very well during the change of the sign of the poling field. Therefore, they were not caused by the appearance of ferroelectric polarization. The bias electric field measurements, which should detect peaks at every ferroelectric transitions (even if a ferroelectric phase appears in intermediate temperature ranges), also did not show any peaks (Figure S9). Therefore, we conclude that dielectric anomalies in BaDyFeO₄ are not caused by spin-induced ferroelectricity or spininduced ferroelectric polarization is below the detection limit of our measurements. In case of BaYFeO₄, polarization of about 2 μ C/m² was measured under a poling field of 800 kV/m [8]. We could apply only 235 kV/m to our BaYFeO₄ sample. If the electric polarization is a linear function of an applied electric field, we could miss polarization of about 0.6 μ C/m², which is near the detection limit. Therefore, the origin of broad dielectric peaks observed between T_{N2} and T_{N1} and centred near 35 K (at H = 0 Oe) in BaDyFeO₄ remains unknown at the moment. Additional experiments will be required to understand their origin, for example, measurements using single crystals and magnetic structure determinations. We emphasize that dielectric peak positions sometimes do not match precisely with $T_{\rm N}$, for example, in o-LuMnO₃ [22]. Dielectric peaks near T_N and magnetodielectric effects were observed in Sc₂NiMnO₆, where no pyroelectric current anomalies were also detected [23]. Therefore, it was suggested that an antiferroelectric transition takes place in Sc₂NiMnO₆. The similar picture could be realized in BaDyFeO₄.

4. Conclusion

We prepared a new BaDyFeO₄ ferrite by a conventional solid-state method in air and investigated its structural, magnetic, and dielectric properties. Three temperature-driven magnetic transitions and two magnetic-field-driven transitions were found in BaDyFeO₄. This compound shows peculiar temperature-dependent and field-dependent dielectric constant and magnetodielectric effects in comparison with BaYFeO₄. However, no pyroelectric current anomalies were found suggesting the absence of spin-induced ferroelectric polarization within the sensitivity of our methods and the used pyroelectric current measurement conditions. The observation of dielectric anomalies in magnetic phases without the appearance of ferroelectric polarization is intriguing and will require further clarification.

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Appendix A. Supplementary data

Supplementary data to this paper contain details of magnetic and dielectric properties of BaDyFeO₄ and BaYFeO₄ and can be found online. CCDC 1899326 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB21EZ, UK; fax: +44 1223 336033.

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Table 1

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Structure parameters of BaDyFeO4 at 295 K from synchrotron X-ray powder diffraction data.

Site	WP	X	У	Z.	$B(\text{\AA}^2)$
Ba1	4 <i>c</i>	0.21160(4)	0.25	0.67406(5)	0.715(9)
Ba2	4 <i>c</i>	0.41492(4)	0.25	0.39532(4)	0.760(8)
Dy1	4 <i>c</i>	0.41447(3)	0.25	0.01463(3)	0.536(6)
Dy2	4 <i>c</i>	0.14330(3)	0.25	0.30953(4)	0.540(6)
Fe1	4 <i>c</i>	0.46928(8)	0.25	0.71533(11)	0.554(18)
Fe2	4 <i>c</i>	0.18974(9)	0.25	0.02269(10)	0.488(18)
01	4 <i>c</i>	0.5870(4)	0.25	0.6153(4)	0.38(8)
O2	4 <i>c</i>	0.2927(4)	0.25	0.1791(5)	1.08(10)
03	8 <i>d</i>	0.0050(3)	0.5090(8)	0.3591(3)	1.33(8)
O4	8 <i>d</i>	0.2194(3)	0.5090(7)	0.4410(4)	1.36(8)
05	8 <i>d</i>	0.1113(3)	0.9977(7)	0.1314(3)	0.81(7)

WP: Wyckoff position. The occupation factor of each site is unity. Space group *Pnma* (No 62); Z = 8.

a = 13.16861(1) Å, b = 5.70950(1) Å, c = 10.26783(1) Å, and V = 771.9985(11) Å³; $R_{wp} = 3.20$ %, $R_p = 2.44$ %, $R_B = 3.70$ %, and $R_F = 2.22$ %; $\rho_{cal} = 7.222$ g/cm³. DyFeO₃ impurity: 1.9 wt. %.

Table 2

Selected bond lengths (l (Å) < 3.25 Å) and bond valence sums (BVS)^a in BaDyFeO₄ at 295 K

Ba1 – O1	2.715(5)	Ba2 - O2	2.741(5)
Ba1 – O5 (×2)	2.763(4)	Ba2 – O5 (×2)	2.827(4)
Ba1 – O4 (×2)	2.815(4)	Ba2 – O1 (×2)	2.857(4)
Ba1 – O2 (×2)	2.856(4)	Ba2 – O5 (×2)	2.973(4)
Ba1 – O3 (×2)	3.185(4)	Ba2 – O4 (×2)	3.006(4)
Ba1 – O4 (×2)	3.199(4)	Ba2 - O1	3.200(5)
		Ba2 – O3 (×2)	3.228(4)
BVS(Ba1 ²⁺) ^a	2.14		
		BVS(Ba2 ²⁺) ^a	2.05
Dy1-O3 (×2)	2.300(4)	Dy2 – O4 (×2)	2.239(4)
Dy1 - O2	2.329(5)	Dy2-O5 (×2)	2.367(4)
Dy1-O3 (×2)	2.360(4)	Dy2 O2	2.380(5)
$Dy1 - O4 (\times 2)$	2.361(4)	Dy2 – O3 (×2)	2.401(4)
BVS(Dy1 ³⁺) ^a	3.10	BVS(Dy2 ³⁺) ^a	3.11
Fe1 – O1	1.860(5)	Fe2 - O1	1.958(5)
Fe1 – O5 (×2)	1.967(4)	Fe2 – O4 (×2)	2.007(4)
Fe1 – O3 (×2)	2.046(4)	Fe2 – O5 (×2)	2.095(4)
		Fe2 - O2	2.103(5)
BVS(Fe1 ³⁺) ^a	2.82		
		BVS(Fe2 ³⁺) ^a	2.81

^{*a*} BVS = $\sum_{i=1}^{N} v_i$, $v_i = \exp[(R_0 - l_i)/B]$, *N* is the coordination number, B = 0.37, $R_0(Ba^{2+}) = 2.29$, $R_0(Dy^{3+}) = 2.036$, and $R_0(Fe^{3+}) = 1.759$ [18].



Figure 1. Crystal structure of BaDyFeO₄. (a) Connection of the Fe1O₅ and Fe2O₆ polyhedra viewed along the *b* axis. (b) One chain of the Fe1O₅ and Fe2O₆ polyhedra along the *b* axis. (c) Connection of the Dy1O₇ and Dy2O₇ polyhedra viewed along the *b* axis. (d) One chain of the Dy1O₇ and Dy2O₇ polyhedra along the *b* axis.



Figure 2. A fragment (9-35°) of experimental (black crosses), calculated (red line), and difference (blue line) synchrotron XRPD patterns of BaDyFeO₄ at T = 295 K. The tick marks show possible Bragg reflection positions for BaDyFeO₄ (the first row) and DyFeO₃ impurity (the second row). The inset shows an enlarged fragment (45-50°).



Figure 3. ZFC (filled symbols) and FCC (empty symbols) dc magnetic susceptibility ($\chi = M/H$) curves of (a) BaYFeO₄ and (b) BaDyFeO₄ at different magnetic fields. The insets show the $d(\chi T)/dT$ versus *T* curves to emphasize magnetic anomalies. *T*_N: Néel tempereature, *T*_{SR}: spin-reorientation transition temperature.



Figure 4. (a) FCC dc magnetic susceptibility ($\chi = M/H$) curves of BaDyFeO₄ at different magnetic fields from 1 kOe to 30 kOe. (b) $d(\chi T)/dT$ versus *T* curves at the same magnetic fields to emphasize magnetic anomalies.



Figure 5. M versus H curves at T = 5, 15, and 30 K for BaDyFeO₄. (f.u.: formula unit). The insets show the dM/dH versus H curves to emphasize field-induced phase transitions at T = 1.8 K (left) and T = 5, 15, and 30 K (right).



Figure 6. Specific heat, plotted as C_p/T versus *T*, of BaDyFeO₄ and BaYFeO₄ at H = 0 (black) and 90 kOe (red). Measurements were performed on cooling. The inset shows C_p/T versus *T* curves below 60 K at different magnetic fields (measured on cooling) for BaDyFeO₄.



Figure 7. (a) Dielectric constant and (b) dielectric loss of BaDyFeO₄ as functions of temperature. Measurements were performed on heating at zero magnetic field at different frequencies.



Figure 8. (a) Temperature dependence of dielectric constant in BaDyFeO₄. Measurements were performed on cooling (filled symbols) and heating (empty symbols) at H = 0, 10, 30, 50, 70, and 90 kOe. Curves at one frequency of 665 kHz are shown. (b) Magnetic-field dependence of relative changes of dielectric constant (magnetodielectric effects) in BaDyFeO₄ at T = 5, 10, 15, 20, 30, 35, 40, and 50 K. Curves at one frequency of 665 kHz are shown.



Figure 9. (a) Temperature dependence of dielectric constant in BaYFeO₄. Measurements were performed on cooling (filled symbols) and heating (empty symbols) at H = 0, 30, 50, 70, and 90 kOe. Curves at one frequency of 665 kHz are shown. An error bar of 2×10^{-3} is shown – a dielectric constant anomaly of this order was detected in Ref. [8]. (b) Magnetic-field dependence of relative changes of dielectric constant (magnetodielectric effects) in BaYFeO₄ at T = 10, 20, 30, 40, 50, and 60 K. Curves at one frequency of 665 kHz are shown.



Figure 10. Results of pyroelectric current measurements in (a) BaDyFeO₄ and (b) BaYFeO₄. The poling was performed from 100 K to 2 K under the poling fields $E_P = +390$ and -390 kV/m for BaDyFeO₄ and $E_P = +235$ and -235 kV/m for BaYFeO₄ at H = 0 Oe.

Highlights:

- BaDyFeO₄ ferrite was prepared by a solid-state method.
- BaDyFeO₄ shows three magnetic transitions.
- BaDyFeO₄ shows two magnetic-field-induced transitions.
- BaDyFeO₄ demonstrates complex behaviors of dielectric constant.

Graphical Abstract

