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Iron borate films: Synthesis and characterization


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A B S T R A C T

We report the first successful synthesis of iron borate films. FeBO₃ films on GaBO₃ single crystal substrates have been prepared by a liquid phase epitaxy route. In order to determine optimal crystallization regimes, a series of experiments has been carried out. Electron microscope studies have allowed monitoring different phases of the film formation. The compositions of the film and of the substrate have been determined by energy dispersive spectroscopy. X-ray diffraction analysis has allowed an accurate determination of a mismatch between the lattice parameters of the film and of the substrate. Electron magnetic resonance studies of the FeBO₃ film confirm the existence of magnetic ordering. The values of the effective Dzyaloshinskii field as well as the Néel temperature are in good accordance with those previously determined for FeBO₃ single crystal.

1. Introduction

The research interest in iron borate FeBO₃ is mainly due to its remarkable magnetic, magneto acoustical, optical, magneto optical, resonance, etc., characteristics, e.g., see [1-6]. In particular, iron borate can be categorized as a “transparent magnet”, combining transmission windows in visible spectral range with room temperature magnetic ordering. From the standpoint of magnetic properties, FeBO₃ is an easy plane antiferromagnet with weak ferromagnetism and the Néel temperature $T_N \approx 348$ K. The AFMR studies of FeBO₃ were carried out in a wide range of temperatures and frequencies [6]. From the standpoint of crystalline structure, FeBO₃ is a rhombohedral calcite type crystal of space group $D_3^6$ [7].

In contrast to conventional ferromagnets, in iron borate the surface magnetocrystalline anisotropy, caused by symmetry breaking in the surrounding of near surface iron ions [8], is not suppressed. This is due to the fact that the demagnetizing field, proportional to weak ferromagnetic vector, is small and the anisotropy in the basal plane, (0001) in hexagonal coordinate system, is weak [7]. Consequently, the magnetic characteristics of a thin (0.01-0.1 μm) near surface layer of iron borate drastically differ from those in the volume. Such effects were studied both experimentally using magneto optic Kerr effect [9] and theoretically [9,10]. Another interesting effect is that surface magneto crystalline anisotropy stimulates the formation of bubble magnetic domains in the near surface layer of iron borate [9]. From the standpoint of practical applications, the surface of iron borate single crystals can be considered as a magnetic memory element analogous to thin film magnetic materials containing cylindrical magnetic domains. Undoubtedly, thin FeBO₃ magnetic films deposited on a diamagnetic transparent substrate are appropriate for studying surface magnetism using not only the Kerr effect [9] but also the Faraday effect. In particular, in very thin (less than 0.1 μm) iron borate films one can study “pure” surface magnetic effects, not altered by volume magnetism.

In FeBO₃ single crystals having the shape of basal plates, interesting magneto acoustic effects were found, related to magnetic structure of this crystal [2,11]. The magnetization in this case follows quasi stastically natural longitudinal [2] and transversal [11] acoustic modes. In very thin FeBO₃ film, one can expect the emergence of forced magnetic oscillations at frequencies approaching the natural magnetic oscillation frequency, i.e. the AFMR frequency, resulting in promising new effects.

The aim of the present work is to develop the synthesis technique, to obtain and to characterize FeBO₃ films on a diamagnetic substrate. As the substrate we have used gallium borate, GaBO₃, single crystal. This material is isostuctural with FeBO₃; besides, it is also transparent in the visible range, albeit diamagnetic [12]. Both crystals have similar lattice parameters: in iron borate...
\( a = 4.626, c = 14.493 \), and in gallium borate \( a = 4.568, c = 14.182 \) Å (in the hexagonal coordinate system). As one can see, the relative difference between the corresponding parameters for both crystals is less than 2%, while the structure of deposited film is known to reproduce that of substrate if this difference does not exceed ca. 14% [13]. Thus, GaBO₃ crystal seems to be the best candidate to be used as substrate for depositing FeBO₃ film.

A successful depositing of a high quality film requires using a high quality substrate. On the basis of our previous studies on synthesis of iron based borate crystals, e.g., see [12,14], we have concluded that high quality samples can be obtained by the solution in the melt technique. Thus, this technique has been used for synthesizing the substrate, and for depositing the FeBO₃ film in these conditions, the liquid phase epitaxy (LPE) technique appears to be optimal.

During the sample synthesis, different phases of the film formation and compositions of the film and the substrate have been monitored by electron microscopy and energy dispersive spectroscopy (EDS), respectively. The lattice parameters of the film and of the substrate have been measured by X-ray diffraction (XRD). The magnetic characteristics of the film, such as the Dzyaloshinskii field \( H_D \) and \( T_N \), have been determined by means of electron magnetic resonance (EMR).

2. Synthesis of the film

The synthesis of FeBO₃ film by the LPE technique includes the following steps:

(i) Choosing appropriate charge compositions and temperature modes;
(ii) Preparing a high quality GaBO₃ substrate;
(iii) LPE synthesis of the FeBO₃ film on the substrate.

The crystallizations in the steps (ii) and (iii) were carried out with Ga₂O₃ B₂O₃ PbO PbF₂ and Fe₂O₃ B₂O₃ PbO PbF₂ solution melts, respectively. The most appropriate charge compositions, determined by differential thermal analysis method, are shown in Table 1 [12,14].

The synthesis of the GaBO₃ single crystal the substrate has recently been described by some of the present authors [12].

The synthesized GaBO₃ crystals are shown in Fig. 1. For comparison, we also show previously synthesized FeBO₃ single crystals [14]. Both GaBO₃ and FeBO₃ crystals have the shape of hexagonal plates with the dimensions of 3 7 mm in the basal plane and 0.05 0.1 mm in thickness. Note that gallium borate is colorless while iron borate is green.

The operating mode used to deposit FeBO₃ film by the LPE route was as follows: the GaBO₃ substrate was placed into a metallic supporting cone perforated with small holes and maintained during 30 min in a crucible containing the solution melt for FeBO₃ synthesis, see Fig. 2. The corresponding temperature mode is shown in Fig. 2. It includes the following stages: (i) heating of the furnace, (ii) homogenization of the solution melt, (iii) fast temperature dropping in order to avoid the emergence of spurious phases, e.g., \( \text{Fe}_3\text{BO}_6 \), (iv) nucleation and film growth and (v) cooling the furnace. The moments when the cone with the substrate has been immersed in and extracted from the solution melt are indicated by arrows. Due to the perforation, during the immersion the solution melt was filling the cone and bathing the whole substrate. During the extraction, the solution melt was withdrawn through the holes back into the crucible, and the synthesized sample (substrate with deposited film) remained in the cone, see Fig. 3. As one can see in online version, after the crystallization the sample surface becomes light green, as characteristic of iron borate.

3. Characterization of the synthesized samples

3.1. Electron microscopy

FeBO₃ film formation has been monitored by electron microscopy using REM 106 and field emission SEM JSM 7800F microscopes.

The film formation in this system occurred following the epitaxial island growth mechanism. Consecutive stages of the film growth are shown in Fig. 4. Conventionally, it can be divided into
four stages: (a) formation of film islands, (b) coalescence of the islands with formation of canals, (c) coalescence of fragments of the film and filling the canals in and (d) formation of a continuous film.

Fig. 5 shows: (a) nucleation centers of FeBO₃ on GaBO₃ substrate and (b) a coalescence of these centers, constituting a fragment of well formed film. From Figs. 4 and 5 it is obvious that all nucleation centers have the same orientation on GaBO₃ surface; thus, the structure of FeBO₃ film replicates that of GaBO₃ substrate.

The developed technique allows synthesizing FeBO₃ films of different thickness, including those suitable for studying surface magnetism.

3.2. Energy dispersive spectroscopy studies

The compositions of the film and the substrate have been locally determined by EDS with X Max silicon drift X ray detector of JSM 7800F microscope. Data have been acquired at a low acceleration voltage of 2 kV. A super hybrid lens with gentle beam produced large probe current at low voltage enabling an express analysis of dielectric structures.

Fig. 6 shows EDS spectra of a sample with partly deposited film, corresponding to the stage shown in Fig. 4(b). If the electron beam is focused on the surface of the substrate, see Fig. 6(a), the EDS spectrum shows lines arising from boron, oxygen, gallium and iron. In this case, the iron line is due to iron borate nucleation centers of nanometric size. Otherwise, if the electron beam is focused on the surface of the film, see Fig. 6(b), the line arising from gallium vanishes. Therefore, we confirm that we are indeed dealing with FeBO₃ layer deposited on the substrate.

3.3. XRD studies

The XRD studies of the synthesized samples have been carried out with X ray diffractometer using a monochromatic Kα₁ copper radiation of 1.54051 Å wavelength. The lattice parameters c have been calculated using the following expression [15]:

![Fig. 3. Synthesized sample in the supporting cone. The diameter of the cone basis is ca. 15 mm and those of the holes are ca. 0.5 mm.](image)

![Fig. 4. Electron microscope pictures taken with REM 106, showing different stages of FeBO₃ film formation, see the text for details.](image)
where \( h, k \) and \( l \) are Miller indices of a Bragg plane and \( d \) is the interplanar spacing calculated by means of Bragg’s formula

\[
\frac{1}{d^2} = \frac{4}{3} \frac{h^2 + hk + k^2}{a^2} + \frac{l^2}{c^2}
\]  

(1)

where \( \theta \) is the scattering angle, \( \lambda \) is the radiation wavelength and \( n \) is the reflection order. Thus, for the reflection peaks from the \((00l)\) planes (\( h = 0 \) and \( k = 0 \)) Eq. (1) reduces to

\[
c = n \lambda
\]  

(2)

For the determination of \( c \) in the film and in the substrate reflections from the plane with \( l = 12 \) have been used. Fig. 7 shows XRD patterns for a synthesized sample as well as for FeBO3 and GaBO3 single crystals. From the XRD patterns of the single crystals, containing only one line, we get \( c = 14.476 \pm 0.017 \) Å and \( c = 14.183 \pm 0.016 \) Å for FeBO3 and GaBO3, respectively. From the XRD pattern of the synthesized sample, containing lines arising both from the film and the substrate, we get \( c = 14.479 \pm 0.017 \) Å for the former and for the latter, respectively. The value of \( c \) for the film is in a good accordance with that for FeBO3 single crystal, as expected for a FeBO3 layer on the GaBO3 substrate. One can see that the mismatch between the \( c \) values in the film and in the substrate is \( \Delta c = 0.297 \pm 0.034 \) Å.

3.4. EMR studies: magnetic properties of the film

The EMR studies of synthesized samples with relatively thick (ca. 4 \( \mu \)m) deposited film as well as of an iron borate single crystal have been carried out with laboratory developed spectrometer in the frequency range from 15.0 to 35.7 GHz, the temperature range from 293 to 350 K and the magnetizing field \( H \) up to 10 kOe applied in the basal plane of the sample. To detect the EMR signal,
the field dependence of the microwave power transmitted through the microwave cavity with the sample was measured.

First, we have carried out EMR studies of FeBO₃ single crystal. Fig. 8 shows the EMR spectrum; obviously, only one resonance line is observed. Below $T_N$, this line has been identified as low frequency AFMR mode, e.g., see [6]. No other resonances occur in the temperature range from 4 to 293 K.

In contrast, the EMR spectra of the synthesized sample exhibit two resonance lines (see Fig. 9). The low field line occurs in the same magnetic field range as the AFMR line in FeBO₃ single crystal; moreover, similarly to the latter line it undergoes a temperature dependent shift and disappears at ca. 350 K, in the vicinity of $T_N$ for FeBO₃ single crystals. Thus, this line can be identified as AFMR in FeBO₃ film. The fact that this line is much broader in film than in single crystal can be explained by structural inhomogeneity and internal stresses in the film resulting in statistical distributions of spectroscopic parameters and, consequently, of the resonance fields. The high field line with the effective $g$ factor $g \approx 2$ appears not only above but also below $T_N$, though in the latter case it is much weaker than the AFMR line. The occurrence of this line below $T_N$ can be due to iron borate nanoclusters located in the transition layer between the substrate and the film. Indeed, as local iron concentration in this layer is inhomogeneously distributed, such clusters are expected to be formed and give rise to cluster magnetic resonance.

The above identification can be corroborated by our recent EMR studies of mixed iron gallium borate, FeₓGa₁₋ₓBO₃, single crystals (0.2 ≤ x ≤ 1) in a broad temperature range, in which case we have also observed two resonance lines similar to those described here for FeBO₃ film on GaBO₃ substrate. A detailed account of the results for the mixed borate crystals will be published elsewhere. Here we mention only the interesting fact that in these crystals the intensity of the $g \approx 2$ line does not follow the $1/T$ Curie law, therefore it cannot be a usual paramagnetic resonance line; rather, it can also be due to iron borate nanoclusters.

From field swept EMR spectra recorded at different microwave frequencies, the dependence of the AFMR frequency on the magnetizing field (Dependence of Frequency on Field, DFF) can be obtained. Fig. 10 shows this dependence for FeBO₃ film at different temperatures. In the case where $H$ is applied in the basal plane, the low frequency AFMR mode is expressed as [6]:

$$\nu = \gamma \left[ H(H + H_0) + H_2^2 \right]^{1/2}$$

(4)

where $\gamma$ is the gyromagnetic ratio for the free electron $g$ value, $g \approx 2.00$, $H_0$ is the Dzyaloshinskii field and $H_2^2$ is the isotropic energy gap. (An anisotropic part in the right hand side of Eq. (4) is neglected because for FeBO₃ single crystal at room temperature the magnetocrystalline anisotropy in the basal plane is almost absent [16].)

From fitting the DFF for FeBO₃ film at different temperatures $H_0$ and $H_2^2$ have been obtained and listed in Table 2. The $H_0$ values are in a good accordance with those previously reported for FeBO₃ single crystal at 300 K [6]. In contrast, $H_2^2$ for FeBO₃ film turns out to be several times larger than for FeBO₃ single crystal. As far as elastic and magnetoelastic interactions influence the isotropic energy gap [6,17], this discrepancy can be ascribed to the

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**Fig. 8.** EMR spectrum of FeBO₃ single crystal at 17GHz and 293 K. The magnetizing field is applied in the basal plane.

**Fig. 9.** EMR spectra series for the synthesized sample at $\nu \approx 17$ (a) and 24.5 GHz (b) and different temperatures shown alongside the curves.
The EMR studies have confirmed the existence of anti ferromagnetic ordering in 4 μm FeBO₃ film, the Néel temperature being in a good accordance with that for FeBO₃ single crystal. Besides, these studies suggest that magnetically ordered nanoclusters are formed in the transition layer between the film and the substrate. The effective Dzyaloshinskii field and isotropic energy gap in FeBO₃ film at 300 and 340 K have been determined from fitting DFF data. The effective Dzyaloshinskii field values are close to those previously determined for FeBO₃ single crystal. However, the isotropic energy gap at 300 K in the film is several times larger than in single crystal; we ascribe this difference to a mismatch between the lattice parameters in the film and in the substrate.

In summary, in this work a new magnetic material, FeBO₃ thin film promising for studying surface magnetism as well as for practical applications as a memory element has been synthesized for the first time, to the best of our knowledge, and its magnetic characteristics have been described.

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In summary, the EMR studies confirm the existence of magnetic ordering in iron borate film.

4. Conclusions

We have developed a suitable synthesis technique and prepared FeBO₃ films on GaBO₃ substrates by LPE route. Temperature modes used in synthesizing the film and the substrate have been experimentally determined. The developed technique allows obtaining films of different thickness, appropriate for magnet optical and magneto acoustical studies.

Various stages of the film formation have been monitored by electron microscopy, and the compositions of the film and the substrate have been controlled locally by EDS. The XRD studies have allowed determining the lattice parameter c for both the film and the substrate. These studies have confirmed that the synthesized samples are composed of FeBO₃ layer on GaBO₃ substrate.

![Fig. 10. Dependence of the AFMR frequency on the magnetizing field for FeBO₃ film at 300 (squares, green online) and 340 (diamonds, red online) K. The dashed curves are fittings according to Eq. (4).](image)

### Table 2

<table>
<thead>
<tr>
<th>T, K</th>
<th>H₀, kOe</th>
<th>H₀², kOe²</th>
<th>H₀², kOe²</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>57.7 ± 0.3</td>
<td>3.3 ± 0.3</td>
<td>62.0 ± 0.5</td>
</tr>
<tr>
<td>340</td>
<td>28.3 ± 0.2</td>
<td>1.3 ± 0.4</td>
<td>6.0 ± 0.5</td>
</tr>
</tbody>
</table>

mismatch of the lattice parameters in the film and in the substrate, vide supra.

In summary, the EMR studies confirm the existence of magnetic ordering in iron borate film.

![Graph](image)