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Synthesis and characterisation of fluorinated epitaxial films of BaFeO₂F: tailoring magnetic anisotropy *via* a lowering of tetragonal distortion†

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In this article, we report on the synthesis and characterisation of fluorinated epitaxial films of $BaFeO_2F$ via low-temperature fluorination of thin films of $BaFeO_{2.5+d}$ grown by pulsed laser deposition. Diffraction measurements show that fluoride incorporation only results in a contraction of the film perpendicular to the film surface, where clamping by the substrate is prohibitive for strong in-plane changes. The fluorinated films were found to be homogenous regarding the fluorine content over the whole film thickness, and can be considered as single crystal equivalents to the bulk phase $BaFeO_2F$. Surprisingly, fluorination resulted in the change of the tetragonal distortion to a nearly cubic symmetry, which results in a lowering of anisotropic orientation of the magnetic moments of the antiferromagnetically ordered compound, confirmed by Mössbauer spectroscopy and magnetic studies.

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1 Introduction

Perovskite-type and perovskite-related transition metal oxides with stoichiometric ABO₃, A₂BO₄, A₃B₂O₇ and non-stoichiometric ABO_{3-x} compositions, (where A: rare earth or alkali earth ion, and B: 3d, 4d, or 5d transition metal ion) can exhibit fascinating physical and chemical properties, including multiferroicity, ¹⁻³ high temperature superconductivity, ⁴ colossal magnetoresistance, ⁵ and photocatalytic activity. ⁶

One of the most interesting aspects of these oxides is the ability to engineer/tune their physical properties by tailoring the composition of the cation sites (*i.e. the 12-fold coordinated A-and/or the 6-fold coordinated B-site*) and/or partially replacing the oxygen by other anionic species such as hydride,^{7,8} fluoride,⁹⁻¹² or nitride¹³ ions. This most often originates from a tuning of the electronic structure and coordination chemistry of the constituent species. One of the methods that has been used to incorporate fluoride ions to change oxidation states of transition metals within perovskite type (or perovskite related) materials are low-temperature topochemical fluorination routes.^{9-11,14,15}

iron mainly, these compounds show G-type antiferromagnetic

ordering with Néel temperatures above 500 K.

The necessity for using low temperature topotactic fluorination routes originates from the fact that the majority of oxy-

fluoride perovskites are only metastable and will decompose at

elevated temperature to form alkaline earth fluorides or lanthanide oxyfluorides. Therefore, for most of these

compounds it is not possible to grow single crystals at elevated temperatures. Within the last years, adopted approaches have

been reported for fluoride incorporation into epitaxially grown

thin films. 14-19 This does not only provide the opportunity to

Although the structure of BFOF appears simple at first glance, the local structure around the iron seems to deviate significantly from having perfect octahedral symmetry. Heap *et al.*^{27,28} were the first to notice an unusual high atomic displacement parameter of the Fe³⁺ cations, and different configurations of fluoride ions within the FeO₄F₂ octahedra (*cis* = both fluoride ions at the same edge; *trans* = fluoride ions located at opposite corners) were discussed in the context of the compound lacking long-range anion ordering. In combination

investigate such films as a grain boundary free model system to study bulk related properties, but also to use strain engineering²⁰ to further modify materials' properties.

Bulk BaFeO_{2.5} (BFO) crystallizes in a complex vacancy ordered modification of the cubic perovskite type structure.^{21–23} This monoclinic modification is centrosymmetric with space group $P2_1/c$. This compound can be fluorinated using polymer-based routes, and BaFeO_{2.33}F_{0.33} (vacancy ordered, $P2_1/m$)^{24,25} and BaFeO₂F (BFOF) (no significant amounts of vacancies, cubic, $Pm\bar{3}m$) can be formed depending on the amount of polymer used.^{26–28} As is common for perovskites with trivalent

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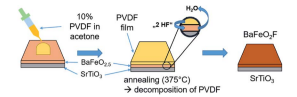


Fig. 1 Representation of drop-fluorination method as carried out on BFO films. The figure is recreated and related to the scheme shown in ref. 14.

with Mössbauer spectroscopic studies, a non-centrosymmetric *cis* environment around a majority of Fe cations from mainly statistic distribution of fluoride ions was concluded.²⁸

The fact that there seems to be chance given for the absence of centrosymmetry around the $\mathrm{Fe^{3^+}}$ cations makes the compound of principle interest for possessing multiferroic properties. To approach such applications, an additional understanding of the structure within fluorinated epitaxially grown thin films seems to be appreciable.

Here, we report two different approaches to topochemical fluorination of epitaxially grown BFO thin films on single crystalline (001) oriented substrates of SrTiO₃.^{29,30} We show that the films can be chemically fluorinated *via* different approaches under formation of BFOF without significant substrate fluorination. *Via* a combination of structural analysis, Mössbauer spectroscopy as well as magnetic studies, an improved understanding of the structural and magnetic behaviour of the films could be gained.

2 Experimental

2.1 Growth of BFO thin films

The BFO target was prepared using a conventional solid state synthesis route. Stoichiometric ratios of solid BaCO₃ (Alfa Aesar, >99.8%) and Fe₂O₃ (Sigma-Aldrich, >99%) were ground, uniaxially pressed and heated twice to 1100 °C for 24 h under flowing Ar with intermediate regrinding.

BFO thin films with a thickness of ~ 85 nm were grown on (001)-oriented SrTiO₃ (STO) substrates *via* pulsed laser deposition (PLD) in the custom made ultra-high vacuum PLD system which is a part of the Darmstadt Integrated System for Battery Research (DaISy-Bat). The BFO target was ablated with a KrF excimer laser ($\lambda = 248$ nm) at a repetition rate of 1 Hz and at a fluence of 1.4 J cm⁻². During deposition, the substrates were heated up to a temperature of 700 °C from their back side, using a high power near infrared diode laser at 938 nm. A background oxygen pressure of 0.018 mbar was used for all depositions. After deposition, the films were cooled down to room temperature at a rate of 50 K min⁻¹. These conditions were chosen in accordance with previous experiments within our group.^{29,30}

2.2 Thin film fluorination

Two fluorination approaches were attempted to fluorinate thin films of BFO to BFOF, which did not show strong differences with respect to the quality, structure and composition of the films reported within this article. In the first approach¹⁴ (see Fig. 1), a solution of 10 wt% of poly-(vinylidene difluoride) (PVDF) in acetone is 'dropped' onto an as-grown oxygen deficient BFO film. This method will be referred to as "drop fluorination" in the following. The film was then heated in air for 24 h at 375 °C.

In a second approach¹⁴ (see Fig. 2), a PVDF powder is placed in a tube furnace heated to 240 °C in front of a BFO film with respect to the gas flow direction. Decomposition products of PVDF, such as HF or small chain components, are then carried over the film and fluorinate it. This approach will be referred to as "vapour transport" fluorination in the following. For this process, a large excess of \sim 2 g of dry PVDF were used (the estimated amount of polymer necessary for full fluorination of the film is in the order of 1–5 µg).

Within this reaction, the fluorine from the decomposition of PVDF under formation of HF can be expected to occupy the interstitial vacancies and replace oxygen from the lattice equally in the film according to

$$O_O + V_i + 2HF \rightarrow F_O + F_i + H_2O$$
 (1)

2.3 Structural characterisation

The X-ray diffraction (XRD) θ – 2θ measurement of the BFO target was performed using a Bruker D8 Advance diffractometer in Bragg–Brentano geometry showing phase purity of the target material as reported previously.²⁹ The thin-film XRD θ – 2θ measurements were performed in parallel beam geometry using a Cu- $K_{\alpha 1}$ Rigaku SmartLab X-ray diffractometer with a Ge(220) 2-bounce monochromator.

2.4 X-ray photoelectron spectroscopy (XPS)

X-ray photoelectron spectroscopy (XPS) measurements were performed within the DAISY-BAT system with a PHI Versaprobe 5000 spectrometer equipped with a monochromated Al Kα X-ray source at 1486.6 eV. All the measurements were carried out at room temperature at a pass energy of 23.5 eV and an energy step size of 0.1 eV. The as-grown BFO films were transferred into the XPS chamber under UHV conditions (<10⁻⁸ Torr) and, therefore, were not exposed to ambient atmosphere prior to the XPS measurements. For fluorinated films, adsorbates such as carbon could not be avoided on the surface of the films as the fluorination reaction was performed outside the UHV system. To probe the inward homogenous incorporation of fluoride into the films, XPS depth profile analysis was performed via argon ion sputtering at an accelerating voltage of 4 kV and an ion current of 3 µA. The adventitious carbon contaminations were detected at the surface of the film only before Ar ion sputtering. Following the first sputtering cycle, no carbon peak could be detected.

2.5 SQUID magnetometry

Magnetic properties of the samples were measured as a function of the applied magnetic field and temperature using a magnetic RSC Advances Paper

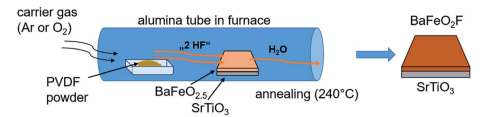


Fig. 2 Vapour transport process carried out on as-grown oxygen deficient BFO placed downstream from PVDF pellets in a heated alumina tube in the presence of a flowing carrier gas. The figure is recreated and related to the scheme shown in ref. 14.

property measurement system (MPMS) by *Quantum Design*, equipped with a superconducting quantum interference device (SQUID). The magnetization data were corrected for diamagnetic contributions from the SrTiO $_3$ substrate. Two different sets of measurements were performed on the same films in non-fluorinated and fluorinated states. Magnetization M for each film was measured as a function of the applied magnetic field H between -2 T and 2 T at 10 K. Magnetization *versus* temperature curves were recorded between 298 and 10 K at H=200 Oe in a field cooled (FC) and zero field cooled (ZFC) regimes.

2.6 Conversion electron Mössbauer spectroscopy (CEMS)

Conversion Electron Mössbauer Spectroscopy (CEMS) was measured with a 57 Co in Rh matrix source (\sim 50 mCi) and the velocity was varied with a constant acceleration drive. The conversion electrons from the resonant absorption of gammarays were detected with a custom-built gas-proportional counter, with a mixture of CH₄ (4%) in He as detector gas. The velocity scale was calibrated with a bcc Fe standard and all isomer shifts are given relative to bcc Fe at room temperature.

3 Results and discussion

3.1 Structural analysis

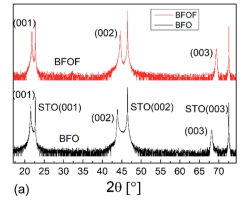
In agreement with our previous findings, 29,30 the XRD measurements showed that the BFO film can be grown in a pseudo-cubic phase (see black curves in Fig. 3a and b). The extracted out-of-plane lattice parameter c=4.13 Å of the BFO

film differs strongly from the lattice parameter of the STO substrate ($a=3.905\,\text{ Å}$). The inset in Fig. 3b shows a rocking curve measurement, showing a full-width at half-maximum (FWHM) of 0.05° for the (002) BFO reflection. The narrow FWHM indicates a high degree of crystallinity and low mosaicity throughout the depth of the film. The value of the lattice parameter $c=4.13\,\text{Å}$ lies between the values of $4.11\,\text{Å}$ observed by Benes $et~al.^{29}$ for 200 nm thick films and of $4.15\,\text{Å}$ reported by Sukkurji $et~al.^{30}$ for 20–30 nm thick films. This indicates that the tetragonal distortion due to the epitaxial strain depends on the film thickness.

After fluorination, the BFOF films do not show any significant change in orientation or strong relaxations within the film, as no additional reflections are observed in the XRD patterns. However, strong shifts of the film (00l) XRD reflections are observed independent of the fluorination method used (see red curves in Fig. 3a and b as well as Fig. S1†), which implies a small BFOF lattice parameter $c \approx 4.066$ Å. Remarkably, this value is similar to what has been reported for the bulk phase of BFOF by Heap $et\ al.^{27}$ ($a = 4.058\ \text{Å}$).

For the BFOF film, the slightly larger FWHM $= 0.08^\circ$ has been extracted from the rocking curve of the (002) reflection (see Fig. S2†) than for the BFO film (FWHM $= 0.05^\circ$), which indicates a small additional broadening after fluorine incorporation. This indicates a small increase of mosaicity after the topochemical incorporation of fluoride ions.

X-ray reciprocal space mapping (RSM) was carried out to determine the in-plane lattice parameter and to understand the topochemical fluorination of the film in better detail. For this



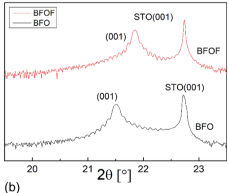


Fig. 3 (a) High resolution X-ray diffractograms of an as-deposited BFO (black) and fluorinated BFOF (red) film on $SrTiO_3$ substrates (b) expanded view of the (001) BFO and BFOF reflections.

purpose, mapping around the STO (103) reflection of the BFO and BFOF films was performed, as shown in Fig. 4a and b.

The shift in the film peak of the BFO film (Fig. 4a) along Q_x in comparison with the shift of the peak of STO substrate indicates that the lattice mismatch is too high for the growth of a fully strained film. This observation is in agreement with our previous study³⁰ which showed the presence of dislocations at the film/substrate interface. The in-plane and out-of-plane lattice parameters of a=4.065 and c=4.128 Å, respectively, were calculated from the RSM measurements (see Table 1).

After fluorination, the BFOF film stays fully strained on the $SrTiO_3$ substrate and the Q_x position of the (103) BFOF film reflection is not changed (see Fig. 4b). This fact gives a strong support for the topotactic nature of the fluorination, which does not break clamping of the film to the substrate and allows only relaxation of the film perpendicular to the substrate plane. Meanwhile, significant changes of the RSM pattern of the BFOF are found along Q_z , implying a strong decrease of the c value to approximately 4.066 Å (Table 1). Thus, it is remarkable that the film fluorination indeed results in basically cubic cell metrics, with an average lattice parameter being close to what is found for powders of BaFeO₂F (a = 4.058 Å (ref. 27 and 28)). Therefore, (taking into account the results of the compositional analysis presented below in Section 3.2), the BFOF films can be considered as being a close approximation to a single crystal of this metastable phase, which cannot be obtained by classic techniques of crystal growth.

3.2 Compositional analysis

Fig. 5 shows a depth profile of the normalized elemental concentration in the BFOF film as extracted from the XPS measurements. In agreement with previous findings, ^{14,16,19,30} fluorine is only observed throughout the complete thickness of the BFOF film. No fluorine is detected in the SrTiO₃ substrate within the resolution limit of about 0.1 at% of the XPS system. This is well plausible as PVDF does not have sufficient reductive power to form low-valent Ti species. ³¹ The detected sudden emergence of the Sr 3d and Ti 2p peaks indicates a sharp interface between the film and the substrate with no sign of interdiffusion of the metal species into the BFOF film. Note that

Table 1 Calculated lattice parameters of the films from RSM

1.120(5) 1
4.128(6) Å 4.066(4) Å 3.903 Å

in the interface region denoted with the vertical black lines, the elemental concentration is imprecise and not given. Within the BFOF film, a rather homogenous distribution of the elements Ba, Fe, O, and F could be confirmed. The Ba: Fe ratio of $\sim 1:1$ in combination with a O: F ratio of $\sim 2:1$ indicates the composition of BaFeO₂F, indicating a fully fluorinated state of the sample, in agreement with the Mössbauer spectra which indicate the presence of Fe³⁺ only (see Section 3.3). Deviations from the ideal composition are within the reliability of the method and should be interpreted neither as anion rich films (*i.e.*, lower oxidation states, as ruled out from Mössbauer

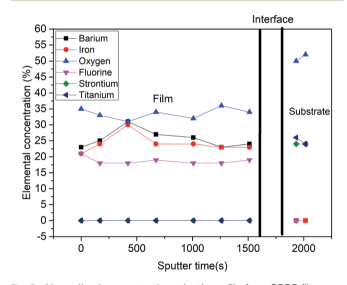
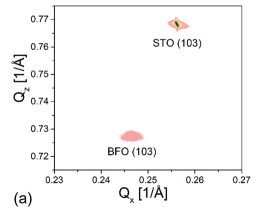


Fig. 5 Normalized concentration—depth profile for a BFOF film corresponding to all the elements observed in XPS spectra.



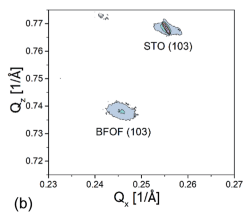


Fig. 4 X-ray reciprocal space mapping of the (a) BFO and (b) drop-fluorinated BFOF films measured around the (103) STO reflection.

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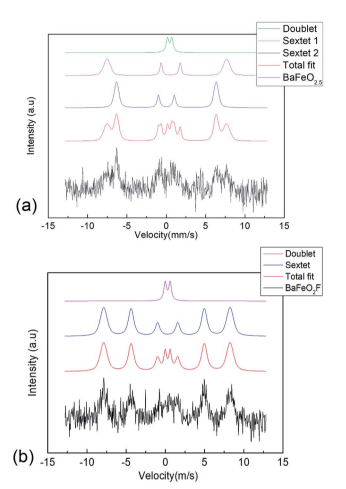


Fig. 6 $\,^{57}$ Fe conversion electron Mössbauer spectrum recorded from (a) BFO and (b) BFOF at room temperature.

spectroscopy, see Section 3.3), nor the presence of, *e.g.* Ruddlesden–Popper related, Ba_2FeO_4 type defects (not indicated within the XRD analysis, see previous section). We would also like to emphasize that a very similar compositional behaviour is indicated the formation of $BaFeO_2F$ films via the second fluorination method (see Fig. S3†). Thus, a $BaFeO_{2.33}F_{0.33}$ (ref. 24 and 25) phase with intermediate fluorine concentration and ordered anion vacancies could not be synthesized with these methods. This phase is either unstable in the investigated strained BFOF films on the $SrTiO_3$ substrates or requires precise control of the fluorine concentration in the film at the level of about 1 μg (only few μg of fluorine are necessary for a full fluorination of the film), which has not been achieved in the presented fluorination methods.

3.3 ⁵⁷Fe mössbauer spectroscopy

 57 Fe conversion electron Mössbauer spectra were recorded for the BFO and drop-fluorinated BFOF films at 298 K (see Fig. 6a and b). It should be noted that the signals from the films were quite weak, which can be explained by the low thickness (\sim 85 nm) of the films. Although the signal-to-noise ratio is low, certain conclusions can still be drawn from the measurements.

- (1) The spectrum observed for BFO is similar to our previous study of 200 nm thick films. 29 It can be fitted with a similar distribution of sextets as reported for bulk BaFeO_{2.5} (ref. 23) (see Table 2 and Fig. 6a), which agrees with tetrahedrally (low isomer shift, sextet 2), square pyramidally, and octahedrally (high isomer shift, sextet 1 & doublet) coordinated iron species within this film. The fit shows that although no long-range ordering of anion vacancies can be obtained within a strained film, the overall distribution of coordination polyhedra must be considered to be similar to the bulk state. 29
- (2) On fluorination of the film to BFOF, the complexity of the spectrum strongly reduces. The spectrum can then be basically fitted with one sextet and one doublet (see Table 3 and Fig. 6b). All isomer shifts are indicative for iron being present in its trivalent oxidation state, in agreement with a composition following $\text{BaFe}^{+\text{III}}\text{O}_{2+x}\text{F}_{1-2x}$ ($x\sim0$). Further, the isomer shift observed for the main sextet is indicative for octahedral environments, where the reduction compared to the octahedrally coordinated FeO₆ species in $\text{BaFeO}_{2.5}$ indicates the presence of the fluoride ions within the coordination environment, and therefore well agrees with the formation of FeO_4F_2 octahedra.³²
- (3) The presence of mainly sextet species clearly evidences magnetic ordering of films. In this respect, we observed that the intensity ratio of the individual sextet lines changed from the non-fluorinated BFO film to the fluorinated BFOF film. For BFO, magnetic anisotropy was already observed in our previous study,29 which is indicated by a change of the relative intensities of the signals of the sextets. Namely a strong decrease of the second and fifth line, which shows a alignment of the magnetic moments parallel to the gamma ray, hence in out-of-plane direction of the thin films along the caxis of the lattice. After fluorination, the BFOF film shows a more isotropic orientation of the magnetic moments indicated by the altered relative intensities of the Mössbauer lines of the sextets towards what would be expected for a randomly oriented powder (intensity ratio of first to second to third line 4:3:1). This agrees with the absence of

Table 2 Fitting parameters for the ⁵⁷Fe Mössbauer spectrum for BFO

	Sextet 1 (pink, 40%) Sextet 2 (blue, 48%)		Doublet (green, 12%)	
	Sextet 1 (pilik, 4070)	Sextet 2 (blue, 4870)	Doublet (green, 1270)	
Isomer shift (IS)	$0.43(4) \text{ mm s}^{-1}$	$0.12(3) \text{ mm s}^{-1}$	$0.54(5) \text{ mm s}^{-1}$	
Quadrupole splitting (QS)	$-0.49(7) \text{ mm s}^{-1}$	$0 \text{ mm s}^{-1} \text{ (fixed)}$	$0.54(8) \text{ mm s}^{-1}$	
Hyperfine field	47.2(4) T	39.3(2) T		
Lorentzian width	$0.30(11) \text{ mm s}^{-1}$	$0.39(13) \text{ mm s}^{-1}$	$0.4 \text{ mm s}^{-1} \text{ (fixed)}$	
Gaussian width	2.3(4)	1.2(4)	_	

Table 3 Fitting parameters for the ⁵⁷Fe Mössbauer spectrum for BFOF

	Sextet (blue, 88%)	Doublet (pink, 12%)
Isomer shift (IS) Quadrupole splitting (QS) Hyperfine field Lorentzian width	0.35(2) mm s ⁻¹ -0.10(4) mm s ⁻¹ 50.0(2) T 0.66(11) mm s ⁻¹	$0.40(3) \text{ mm s}^{-1}$ $0.62(6) \text{ mm s}^{-1}$ 0.4 mm s^{-1} (fixed)
Gaussian line width	1.7(4)	

significant straining after fluorination and a cubic symmetry of the film and is in agreement with literature reports for bulk $BaFeO_2F.^{28}$ The doublet species are commonly found for the various modifications of $BaFeO_2F^{32}$ and most likely are related to magnetically disordered regions within the films. Furthermore, the values of the observed hyperfine field splittings are high and agree with the presence of magnetically ordered high-spin Fe^{3+} at temperatures well below the magnetic ordering temperature. 33

(4) For bulk BaFeO₂F, a random distribution of fluoride ions, forming *cis* (two fluoride ions at the same edge of the octahedron) and *trans* (two fluoride ions at opposite corners of the octahedron) was considered as the most plausible structural scenario.²⁸ It is difficult to determine whether more than one sextet signal is present within the BFOF film. The fact that the sextet is considerably broadened by a Gaussian distribution (width of 1.7 T) might indicate that the different crystallographic scenarios cannot be resolved and are thus compiled in

the broad sextet. However, the small quadrupole splitting of the sextet species might indicate a nearly centrosymmetric environment, which is only the case for the *trans* configuration of anions within FeO_4F_2 octahedra.

3.4 Magnetic analysis

Fig. 7 shows the results of the magnetization measurements of the BFO and BFOF films. For both films, the measured saturation magnetic moments are below 0.015 $\mu_{\rm B}$ per Fe at 10 K in agreement with usually observed antiferromagnetically ordered Fe³⁺-containing perovskites.^{23,27,28,32,34-37} A small difference between the FC and ZFC measurements is similar to the previously reported data for bulk BaFeO2,5 (ref. 23) and BaFeO₂F.²⁸ In agreement with previous discussions with respect to the magnetic hyperfine field splittings observed via Mössbauer spectroscopy,28 these results indicate a canted antiferromagnetic ordering of Fe3+ ions in both BFO and BFOF films with a small canting angle below 0.1°. Overall, no difference could be observed for the magnetic behaviour of the nonfluorinated BFO and the fluorinated BFOF films. Their similar low magnetic moments at low temperatures and curvature of the magnetization temperature dependence indicate that no significant change in the Fe3+ antiferromagnetic ordering occurs after fluorination. Thus, we cannot draw conclusions on potential changes of the Néel temperature, which is known to depend on the fluorine content (e.g. decreasing of the Néel temperatures from 750 K for LaFeO₃ to 645 K for BaFeO₂F, and even further to \sim 480 K for AgFeOF₂).³⁸

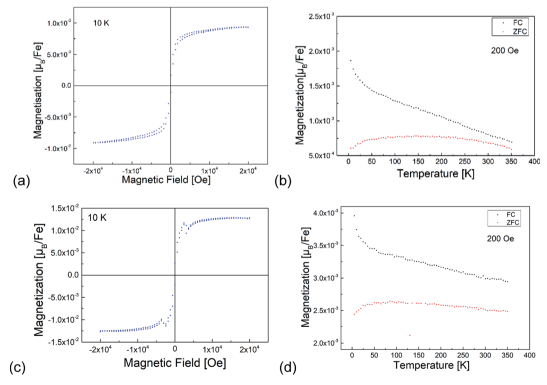


Fig. 7 Dependences of the magnetization on the applied magnetic field at 10 K for (a) non-fluorinated BFO and (c) drop-fluorinated BFOF films. Temperature dependence of the magnetization for (b) BFO and (d) BFOF films, measured at the applied magnetic field of 200 Oe in field-cooled (FC black) and zero-field-cooled (ZFC red) regimes. The data were corrected for the diamagnetic contribution of the SrTiO₃ substrate.

4 Conclusions

In this work we have shown that polymer based routes are efficient means in order to prepare chemically homogenous epitaxial thin films of BaFeO₂F *via* the low temperature top-ochemical fluorination of thin films of BaFeO_{2.5} grown by PLD. No significant differences in film quality were found for the investigated drop-fluorination and vapour transport fluorination methods. The incorporation of fluorine results in a change of the tetragonal distortion to a cubic symmetry, which lowers the magnetic anisotropy of the film of the antiferromagnetically ordered moments. In addition, a small degree of ferromagnetic canting could be observed, similar to what is found for bulk barium ferrates.²⁸

In future works, our group will also focus on the preparation of strained films of $BaFeO_2F$ by using substrates which could allow for more strict cube-on-cube growth, which could then result in additional decrease of symmetry, targeting to form novel multiferroic compounds.

Conflicts of interest

The authors declare no conflicts of interest.

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