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“Spin flop transition and magnetocaloric effect through disconnected magnetic blocks in Co^{III}/Co^{IV} oxybromides”

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**Abstract:** The magnetic properties of the layered $\text{Ba}_{n+1}\text{Co}_n\text{O}_{3n-1}\text{Br}$ ($n=5$ and 6) oxybromides $\text{Ba}_7\text{Co}_6\text{BrO}_{17-\delta}$ have been determined by means of magnetization measurements and neutron diffraction under variable magnetic field. For $n=6$, the magnetic phase diagram has been built on the basis of several features, including the notions of short range ordering, spin reorientation and spin-flop transitions. In those compounds, the competition between magnetic exchanges arise from to the existence of $[\text{Ba}_2\text{O}_2\text{Br}]^-$ double-layers that separate perovskite 2D-blocks. The latter are dominated by ferromagnetic (FM) intra-exchanges while the inter-block exchanges are antiferromagnetic (AFM), but of the same order of magnitude than the perturbation created by the external field. It leads to a complex set of spin reorientations versus H and T. From the point of view of magneto-crystalline anisotropy, the AFM system with moments parallel to the c-axis turns into a FM system with moments aligned in the perpendicular (a,b) planes. The magnetic entropy is distributed within at least three competing phenomena, spread out in a wide range of temperature. In addition, the influence of the magneto-crystalline anisotropy on the magnetocaloric effect is unambiguously shown.

**PACS:**

75.30.Sg Magnetocaloric effect, magnetic cooling

75.25.+z Spin arrangements in magnetically ordered materials (including neutron and spin-polarized electron studies, synchrotron-source x-ray scattering, etc.)

61.05.F- Neutron diffraction and scattering

75.30.Kz Magnetic phase boundaries including magnetic transitions, metamagnetism, etc.

75.60.-d Domain effects, magnetization curves, and hysteresis
I. Introduction

Nowadays, the need for low cost energy stimulates the development of better-adapted prototypes and/or to find new energy sources. Among them, extensive studies about the magnetocaloric effect (MCE) are being carried out since the discovery of a giant magnetocaloric effect in Gd$_5$(Si$_x$Ge$_{1-x}$)$_4$. The MCE is characterised by a temperature change of a magnetic material under an applied magnetic field$^2$ and is believed to be a good alternative for future magnetic cooling units$^3$. In that frame, a large class of materials have been studied including intermetallics, La-Fe-Si and Gd-Si-Ge families, Mn/Fe-based arsenic materials and manganites compounds$^4$-$^6$. To our knowledge, the largest MCE has been found in the compound Mn$_{1.1}$Fe$_{0.9}$P$_{0.8}$Ge$_{0.2}$ with 74 J/Kg.K around 255K$^7$.

Quantitatively, from the thermodynamics and using the Maxwell formalism, the isothermal field-induced magnetic entropy change from 0 to $H$ can be written as$^8$

$$\Delta S_M(T,H) = \int_0^H \frac{\delta M}{\partial T} \partial H$$

eq. (1)

Determining the field-induced isothermal specific heats change, the field-induced adiabatic temperature change ($\Delta T_{ad}$) at the magnetic transition from 0 to $H$ can be evaluated by

$$\Delta T_{ad} = - \int_0^H \frac{T}{C_{P,H}} \frac{\partial M}{\partial T} \partial H$$

eq. (2)

Thus, there is an immediate correlation between the sign of the magnetic entropy change and the capability of the concerned compound to be used as a cooling or heating medium. Such opposite behaviours are referred as inverse or normal MCE respectively and are usually associated with antiparallel or parallel spin arrangements. More recently, a quantitative parameter suitable for application has been introduced: the Relative Cooling (or Heating) Power, hereafter RCP$^8$. In the case of a refrigerator describing a thermodynamic cycle, it would indicate how much heat can be transferred from the cold to the hot end. It can be estimated by the product $\Delta S_M(T)^{\text{max}} \times \delta T_{\text{FWHM}}$.
where $\delta T_{\text{FWHM}}$ is the full width at half maximum of the field induced magnetic entropy change $\Delta S_M(T)$.

Up to now, further routes are being investigated to find new materials showing MCE and/or RCP the highest as possible. One possible way to enlarge the RCP is to elaborate materials showing multiple magnetic transitions. In the present paper, we suggest an another way taking advantage of the possible tuning of the magnetocrystalline anisotropy as already proposed by Isnard et al. We report herein the fine magnetic investigation and the MCE effect of the $18R-$Ba$_6$Co$_5$BrO$_{14-\delta}$ and $14H-$Ba$_7$Co$_6$BrO$_{17-\delta}$ compounds which belong to a series of hexagonal perovskite compounds recently isolated in the Ba-Co-O-X chemical systems (X=F, Cl, Br). Those compounds show disconnected 2D-blocks at both sides of central [Ba$_2$O$_2$Br] double-layers. They can also be described as n=5 and n=6 members of the series Ba$_{n+1}$Co$_n$O$_{3n-1}$Br, underlying the layer's aspect of the compounds. This topology is such that the nature of the exchanges between the blocks and the magneto-crystalline anisotropy is strongly depending on the external field. Strikingly, an external applied magnetic field promotes spin-flop and spin reorientation transitions which leads (i) to enlarge the full width at half maximum of $\Delta S(T)$ and then the refrigerant capacity and (ii) to tune the sign of the magnetic entropy. It follows that those compounds could be considered as cooling or heating media at constant temperature depending on the amplitude of the applied magnetic field.

II. Experimental procedures

Neutron diffraction: Pure powder of Ba$_{n+1}$Co$_n$O$_{3n-1}$Br (n=5 and 6) compounds have been prepared as described in a previous work dedicated to the crystal structure of those recent compounds. Their magnetic structures versus temperature, at zero field, have been refined from neutron diffraction (ND) data collected on the G41 diffractometer, $\lambda=2.425$ Å (LLB, Saclay,
France). The influence of an applied field has been investigated through different ND data sets collected at the high flux D1B diffractometer, \( \lambda = 2.52 \) Å (ILL, Grenoble France). In a first experiment, an electromagnet with H<1.2 T was used on free powder. In a subsequent stage, we benefited of a cryo-magnet with H going up to 2.5 T, which was used on dense pellets. In both cases, the applied magnetic field was oriented vertically, that is to say perpendicular to the scattering vector. The temperature and applied field have been varied from 1.4K to 90K and from 0 to 2.5T, respectively.

**Magnetic measurements:** Magnetization experiments have been performed using a Superconducting Quantum Interference Device (SQUID, Quantum Design) under a dc field. For both temperature and field dependence studies, the sample was introduced at room temperature and cooled down in zero-field at 5K/min before the magnetic field was applied. M(T) measurements were performed on warming the sample. M(H) magnetic hysteresis loops were from -5T to 5T

### III. Structural aspects of the related series of compounds

Starting from the 1D 2H-BaCoO₃ with isolated columns of face-sharing CoO₆ containing low spin (LS) Co⁴⁺, \( S=\frac{1}{2} \) the introduction of oxygen deficient \([\text{BaO}_2]^{2-}\) layers yields the two 5H- and 12H-BaCoO₃-δ polymorphs. They are formed of face sharing octahedral oligomers, namely trimeric-[Co₃O₁₂]₅H or tetrameric-[Co₄O₁₅]₁₂H, capped by terminal CoO₄ tetrahedra \( S=\frac{1}{2} \). In the BaCoO₃-δ polytypes, the so-formed 2D-blocks are not connected by their terminal tetrahedra. The replacement of the central deficient \([\text{BaO}_2]^{2-}\) layers by either \([\text{BaOX}]^{-}\) layers (X=F,Cl)\( ^{12,17-20} \) or \([\text{Ba}_2\text{O}_2\text{Br}]^{-}\) double-layers\(^{11}\) enables various connections between the common 2D-blocks. See ref.\(^{[12]}\) for a review on the preliminary electric and magnetic investigations. In the title compounds \( \text{Ba}_6\text{Co}_5\text{BrO}_{14-\delta} \) (a=5.6578(3) Å, c=43.166(4), space group R-3m) and \( \text{Ba}_7\text{Co}_6\text{BrO}_{17-\delta} \) (a=5.666(1) Å, c=33.367(2) Å, space group P6\(_3/mmc\)), respectively the trimeric and tetrameric 2D-blocks are
pushed aside the c-axis and shifted in the (a,b) plane due to the central \([\text{Ba}_2\text{O}_2\text{Br}]\) double layer, fig.1a-b.

**IV. Magnetic ordering at zero-field**

The preliminary magnetic characterizations of the two compounds are shown on the figure 2. This figure emphasizes similar behaviours for both compounds. From the M(T)/H plots measured at H =1 T, the characteristics of the paramagnetic domain have been extracted from the typical Curie-Weiss behaviour at high temperature, i.e.\(\text{Ba}_7\text{Co}_6\text{BrO}_{17-\delta}: \mu_{\text{eff}} = 3.12 \mu_B/\text{Co} \) and \(\theta_{\text{CW}} = 48.7 \text{ K} \); \(\text{Ba}_6\text{Co}_5\text{BrO}_{14-\delta}: \mu_{\text{eff}} = 3.56 \mu_B/\text{Co} \) and \(\theta_{\text{CW}} = 45.1 \text{ K} \). The values of \(\theta_{\text{CW}}\), close to those found for other hexagonal cobaltites \(^{22,23}\), emphasize the similitude existing in this class of materials. As for the oxides, the positive values of \(\theta_{\text{CW}}\) show dominating ferromagnetic (FM) exchanges. However for both compounds, an antiferromagnetic (AFM) ordering exists below \(T_N \sim 66 \text{ K} \) for the former and \(T_N \sim 48 \text{ K} \) for the latter. On further cooling an additional transition occurs characterized by an abrupt increasing of M/H. This transition takes place at \(T_t \sim 20 \text{ K} \) for both compounds and will be assigned latter to a cone-spin structure. For the two compounds, the M(H) plots show strong field and temperature dependences( Fig2.c-d) at the origin of our interest for these two compounds.

The magnetic structures refined at 1.4 K have already been shortly reported in a previous work \(^{12}\) which corresponds to the moments pictured in the figure 1. We have reinvestigated the neutron diffraction data, not only at 1.4K, but also as a function of the temperature. In \(\text{Ba}_6\text{Co}_5\text{BrO}_{14-\delta}\) magnetic satellites appear below 50 K in good agreement with the susceptibility data. They can be indexed using the propagation vector \(k= (0,0,1/2)\) which corresponds to the setting of an AFM ordering between FM 2D-blocks. The local moments are lying parallel to the c-axis. At 1.4K our reviewed refinement leads to \(M(\text{Co})_{\text{tetra}}=3.25(5)\mu_B\), \(M(\text{Co})_{\text{central octa}}=0.87(9)\mu_B\) and \(M(\text{Co})_{\text{edge octa}}=0.61(6)\mu_B\) with final \(R_{\text{magn}} = 5.8\%\) for \(\text{Ba}_6\text{Co}_5\text{BrO}_{14-\delta}\). Since the unit cell contains two
blocks in Ba₇Co₆BrO₁₇₋δ, the magnetic contributions are superimposed to nuclear ones, and thus no additional peaks (implying a k=(0,0,0) propagation wave vector). In the tetrameric compound, the magnetic ordering is similar to the ordering of the trimeric ones. The refinement at 1.4K leads to M(Co)tetra=3.22(5)μB, M(Co)central octa=0.57(9)μB and M(Co)edge octa=0.51(6)μB with final R_magn=4.9%. The experimental and calculated patterns for the two compounds are shown on the figure 3.a-b. In fact, the assignment of the cobalt valence states in the series of related compounds has already been intensively discussed suggesting octahedral Co³⁺ and tetrahedral Co⁴⁺ in an intermediate spin state. However this last point is contested by some recent DFT calculations on oxo-fluorides containing the same blocks which unambiguously lead to high spin (HS) Co³⁺, S=2 in the tetrahedra and a mixture of low-spin (LS) Co³⁺ (S=0) and Co⁴⁺ (S=1/2) in the face sharing octahedral. The similarity between the individual blocks and their internal ferromagnetic ordering suggest an identical distribution for the oxo-bromides. This is also consistent with the refined values of the local moments, and the experimental μ_eff values matches rather well this proposed charge distribution, i.e.

\[
\text{Ba}_6\text{Co}_5^{+3.4}\text{BrO}_{14} : 2 \text{HS-Co}^{\text{III}} + 1 \text{LS-Co}^{\text{III}} + 2 \text{LS-Co}^{\text{IV}} : \mu_{\text{eff exp.}} = 3.56 \mu_B/\text{Co}, \mu_{\text{eff calc.}} = 3.28 \mu_B/\text{Co}.
\]

\[
\text{Ba}_7\text{Co}_6^{+3.5}\text{BrO}_{17} : 2 \text{HS-Co}^{\text{III}} + 1 \text{LS-Co}^{\text{III}} + 3 \text{LS-Co}^{\text{IV}} : \mu_{\text{eff exp.}} = 3.12 \mu_B/\text{Co}, \mu_{\text{eff calc.}} = 3.08 \mu_B/\text{Co}.
\]

Note that, the theoretical values are calculated within a spin-only picture (g=2). However, we cannot exclude the contribution of oxygen vacancies which would lead to significant change in the octahedral Co³⁺/Co⁴⁺ partition. Furthermore, if one considers the strong magnetocrystalline anisotropy discussed in the next section, one should also take into account a significant amount of orbital contribution for cobalt cations probably responsible for the slightly underestimated calculated μ_eff.

The evolution of the moments versus the temperature is shown on the figure 4.a-b for Ba₇Co₆BrO₁₇₋δ and Ba₆Co₅BrO₁₄₋δ. For the two compounds, the moments are rapidly increasing below T_N. The magnetic moment of Co1 and Co2 refer to those of the Co atoms on the face
sharing octahedra in good agreement with the moments proposed from the Curie-Weiss law. A.
delocalized spin density within the trimeric-[Co₃O₁₂]₅H or tetrameric-[Co₄O₁₅]₁₂H species and then
direct Co-Co hybridization. Most of the magnetic moment is carried by Co3 on tetrahedral
environment as illustrated figure 1. In fact, a fine examination of the intensities of some magnetic
satellites displays an unexpected decrease below ~10 K, fig. 5. It yields slight anomalies in the
evolution of the moments at low temperature as seen on the figure 4. This phenomenon may be
correlated with the opening of a M(H) hysteresis loop. However these phenomena are relatively
weak in absence of an applied field and ND intensities do not show the appearing of extra magnetic
satellites. It is such that in the Rietveld refinements, our attempts to deviate the moments out of the
c-axis under zero-field failed. As we will see in the next section, the concerned spin reorientation
will be best characterized under an applied magnetic field.

V. Effect of an applied field

The magnetization plots observed in the figure 2 for the two compounds is consistent with
the appearing of some kind of metamagnetism, likely present in those systems due to the strong
disconnection between the blocks. The topology of the two compounds is such that the AFM inter-
block exchanges are weak compared to related “connected” systems. It was attempted to verify the
existence of a field induced spin reorientation using ND under an applied field. Precisely, our main
goal is to distinguish between two possible magnetic configurations that could occur under an
applied field: (1) the progressive reversal of magnetic moments, conserving their orientation
parallel to c. This would involve the setting of variable ferrimagnetic ordering until the FM
saturation. (2) an increasing tilt of the spins leading to canted structures leading to a magnetic
contribution parallel to the (a,b) planes. The evolution of the ND-magnetic lines or the appearance
of new magnetic peak should give clear indications on the magnitude and direction of the moments
versus H. The first collected data sets using free-powder samples below Tₜ are shown on figure 6,
For \(\text{Ba}_7\text{Co}_6\text{BrO}_{17.5}\). For the two compounds, a strong reorientation of the crystallites was observed for applied fields greater than \(> 0.5\) T. It is evidenced by a strong modification of the full diffraction pattern, i.e. strong decreasing of \(I_{00l}\), strong increasing of the \(I_{hkl}\) and variable behaviour of \(I_{h0l}\) depending on the \(h/l\) ratio. It pictures a drastic re-orientation of the crystallites due to the alignment of the induced ferromagnetic component aligning parallel to the vertical applied field.

Most of the crystals are reoriented with 00l faces in diffraction positions and suggests the occurring of the canted structure of the above-model 2. Indeed, in an AFM structure the magnetization is larger when applying the field perpendicularly to the direction of the moments since a slight tilt of the spins can lead a FM component.

To lock the crystallite re-orientation, we decided to use, instead of free powder, some sintered pellets. However, we have not been not able to properly densify pellets of the tetrameric \(\text{Ba}_7\text{Co}_6\text{BrO}_{17.5}\) since it transforms into the trimeric \(\text{Ba}_6\text{Co}_5\text{BrO}_{14.5}\) at 990°C \(^{11}\). Then only dense cylinders of this latter (compaticy greater than 80\%) have been used for ND experiments under applied field. As expected, in that case the crystallites reorientation is blocked. At 1.4 K, the evolution of the ND patterns varying \(H\) from 0 T to 2.5 T is shown on the figure 7. The intensities of the AFM satellites, e.g. \(h00 + k\), with \(k=(0,0, \frac{1}{2})\) progressively decrease above \(H=1.2\)T while extra magnetic components are growing superposed to some Bragg diffraction lines, e.g. the 104 and 105 peaks. They highlight a deviation of the moments away from the c axis. Several magnetic models have been tested by Rietveld refinement in order to characterize the nature of the spin-deviation. Actually, a canting structure from block to block lead to the best agreement and has been selected. In that case, to describe the magnetic structure, it was not possible anymore to use a propagation vector since the reversal of the moment by the \((0,0,\frac{1}{2})\) translation is not respected in the canted structure. Thus, the unit cell was doubled and each magnetic atom was introduced in association with its moment \(M_{\text{Co1 to Co3}}\) and a tilting angle \(\theta\) or \(-\theta\) from the c-axis. These models have converged leading to a best matching between the experimental and calculated data and to the
successful generation of the FM peaks while reducing AFM satellites. Depending on the value of H, the final magnetic $R_F$ factors are between $\sim5$ and $\sim10\%$. The M-Co(H) and $\theta$(H) are shown on the figure 8. Strikingly, the tuning of $\theta$ reaches 42° at 2.5 T. The progressive spin-deviation is accompanied by a slight exaltation of the magnetic moments of the octahedral Co1 and Co2 and a small reduction of those on the tetrahedral Co3. Note that the slight differences in the refined moments at H=0 T and T=1.4 K between our experiments on D1B, ILL (ie. with cryo-magnet) and on G4.1 LLB (ie. without cryo-magnet), $2.79(3)\mu_B$ -D1b- compared to $3.26(5)\mu_B$-G4.1-, can be attributed to the most narrow resolution range in the former due to the use of this cryo-magnet which unfortunately brings parasitic lines for $d_{hkl}>2.58$ Å. However the main picture of the magnetic structure remains identical.

VI. Spin flop transition

In this section we will examine in detail the magnetization of the tetrameric compound, Ba$_7$Co$_6$BrO$_{17-\delta}$, on the basis of the previously refined magnetic structures. Once more, it is clear from the figure 2 that phenomena occurring in this compound should be also qualitatively suitable for the parent trimeric Ba$_5$Co$_6$BrO$_{14-\delta}$. Figure 9 represents the Zero Field-cooled M(T)/H for Ba$_7$Co$_6$BrO$_{17-\delta}$ under variable field. It enhances its strong field dependence. On cooling under 0.9 T, we recognize the two maxima briefly discussed for both compounds in the section IV.

i) The first one centred at T=56K corresponds to the Néel temperature and comforts the neutron diffraction experiment. However it displays a large broadening when the field increases until it almost vanishes (but is still observed on the derivative dM/dT(T) curve at 60K). Thus, even at $T_N$, an applied external field induces the setting of magnetic reorientations. Above $T_N$, M(T)/H corrected from the Curie-Weiss susceptibility changes its sign within a critical field range between 0.9T and 1.4 Tesla (figure 10). It involves a drastic change of the magnetic correlations from anti-
parallel to parallel alignment when H increases. ii) The second transition is centred at 22K for H=0.9T. As shown by ND, it corresponds to the appearance of resulting ferromagnetic component in the (a,b) plane by canting. On increasing H, this transition can also be followed by the position of the local minimum of the dM/dT(T) derivative curve. The transition temperature increases continuously with the field, until being equal to 30K at 5T. At high fields, FM exchanges predominate.

At this step of the discussion, from the Zero Field-cooled M(T)/H measurements, two kinds of ordering are observed:

(i) the 3D anti-parallel ordering of the 2D units along the c axes, as shown by the first Néel-like transition temperature at 56K for H=0.9T for Ba$_7$Co$_6$BrO$_{17-\delta}$ and from our neutron diffraction data.

(ii) the appearance under high external magnetic field and/or on cooling of a FM -like component as supported by the positive $\theta_{CW}$. For Ba$_7$Co$_6$BrO$_{17-\delta}$, the transition, which is the field-driven reorientation, suggested by our ND experiments has been analyzed from the isothermal magnetization experiments. A selection of them is shown in figure 11.

The Magnetization loop at 66K is typical of a paramagnetic behaviour. However no Brillouin law adjusts the experimental data using the average spin state deduced from the effective moment fitted by the Curie-Weiss law (see section IV). Short range magnetic ordering (SRO) likely occurs at such temperature. Indeed, similar superparamagnetism as observed in the “parent compound” 2H-BaCoO$_3$ could be suggested. Here, magnetic clusters would consist on bunches of linear tetramers already coupled above $T_N$. It is reminiscent of the 3D-ordering of the related oxo-halogenides (X=F, Cl) for which similar tetramers/trimers also play the role of magnetic connectors at higher temperature, $T_N$~110-135 K and, we already pointed out that a drastic change of the magnetic correlations occurs (from anti-parallel to parallel alignment) when H increases. On the isotherm measured at 53K, a first linear part is shown until 0.5T as expected from the anti-parallel
alignment along the c-axis from neutron data refinement. This linear evolution is followed by a significant upturn above 0.8 T showing a magnetic transition. No hysteresis is observed pointing out a reversible phenomenon. In term of magnetocrystalline anisotropy, an important spin/orbit coupling is expected from Jahn-Teller cobalt cations and might be at the origin of the magnetic transition in such crystallographic topologies. The ‘critical field’ $H_c$ responsible for it is defined as the field for which the fastest change in the Magnetization occurs. It coincides with a local maximum in the $dM/dH$ curves. Its temperature dependence will be discussed later. Typically, $H_c$ increases significantly as the temperature is lowered, following the continuous lost of the linear part on isothermal M(H) curves at low magnetic field. The opening of the magnetization loop at 22K coincides with the loss of the linear M(H) dependency at low field. Consequently, between ~20K and 55K, a field-driven spin flop transition from canted antiferromagnetic to weak ferromagnetic ordering definitively occurs, as characterized by the value of $H_c$.

Similarly, the second sharp transition on M/H(T) around 10K (see figure 9) is also related to the continuous lost of the linear part on isothermal M(H) curves at low magnetic field, and is also assumed to be due to a spin-canting phenomenon. It is highlighted by the small remnant moment when temperature further decreases below 12K. We already stated that the coinciding spin-tilting at H=0 and T=1.5K could not been refined from our powder ND experiments, even if it is correlated with the low-temperature anomalies shown on the figure 2.

At this stage, it is worth noting that the value of the first M(H) plateau at 12K roughly corresponds to one third of the magnetization at saturation. This behaviour is reminiscent of comparable phenomena occurring for related materials such as Ca$_3$Co$_2$O$_6$ made of 1D- cobalt ising chains$^{27}$. In the series of related compounds, such behaviour corresponds to the competition between strong in-chain FM's and weaker interchain AFM exchanges. It gives clues for similar effects in Ba$_7$Co$_6$BrO$_{17-\delta}$ in which the competition exists between FM blocks containing similar 1D-segments and weaker AFM inter-block couplings. In addition, similarly to the Ca$_3$Co$_2$O$_6$ case, it
is probable that the exchanges between adjacent tetramers of a same block are negative but dominated by the cooperative FM alignment imposed by the terminal tetrahedral connections.

**VII. Ba$_7$Co$_6$BrO$_{17.8}$ magnetic phase diagram**

Figure 12 summarizes the final (H,T) phase diagram deduced from the systematic M(T) and M(H) screening of the system combined with analysis of the ND data. The temperature for which the inverse of the magnetic susceptibility and the Curie Weiss law behaviour diverge is correlated to the appearance of short range ordered (SRO) domains and it is defined as the disorder/order temperature (full square). Also shown are the long range antiferromagnetic and weak ferromagnetic ordering temperature (empty square) related to the magnetic transition observed by neutron diffraction, $H_c$ (arrow) and $T_{sp}$ (circle) at which spin reorientation or spin-flop occur. They are defined as the local maximum of $dM/dH$ and as the fastest change on the curvature of the M(T) respectively. As the temperature is lowered, $H_c$ grows up almost linearly until reaching a plateau at $1.2T$ in the range [25K-40K]. It is worth recalling that $H=1.2T$ corresponds to the value for which a change of magnetic correlation occurs below $T_N$ (see fig. 10). It follows that the divergence of $\chi^{-1}$ from the Curie-Weiss law signs up the evolution of the field-induced spin flop transition.

*For magnetic field lower than 1.2T*, irregular (SRO) occurs above the Néel ordering. Below $T_N$, the moments align along the c easy axis. Further cooling leads to a continuous transition from AFM to a canted AFM state resulting in an imperfect compensation of the magnetic moment along the c axis. At $T_{sp}$ a spin-flop transition occurs responsible for the second sharp transition on the temperature dependent magnetization. The magnetic anisotropy is kept along the c-axis as observed by ND experiment.

*For magnetic field greater than 1.2T*, a transition from SRO to canted antiferromagnetic ordering occurs on cooling, responsible for the smoothing of M(T). On further cooling the
progressive transition into a weak ferromagnet is signed by the opening of an hysteresis loop which depends on the (H,T) coordinates. For example, the isothermal magnetization at 53K is perfectly reversible just above Hc while a coercive field of 250 Oe is observed on the isothermal magnetization at 42 At lower temperature and for a applied magnetic field larger than Hc, Ba7Co6BrO17-δ likely turns into a weak FM with a net moment in the (a,b) plane.

It is clear that the disconnection of the individual blocks by the central [Ba2O2.δBr]− double-layers enables a competition between various inter and intra-blocks exchanges, by comparison to parent BaCoO3-δ and other oxy-halides polymorphs. A large uniaxial magnetic anisotropy related to the quasi-one-dimensional arrangement of the interacting Co atoms is observed on the 1D- 2H BaCoO3 and gradually lifted from 5H or 12H BaCoO3-δ to Ba7Co6BrO17-δ. In the 5H- BaCoO3-δ polymorphs FM exchanges predominate below 50K and a canted angle of 63° with respect to the c-axis was announced. A lowering of the magneto-crystalline anisotropy is supported for strongly disconnected blocks of the two examined Ba_{n+1}Co_{n}O_{3n-1}Br (n=5 and 6) compounds. The intra-blocks exchanges have already been intensively discussed in the parent Ba6Co6(Cl, F)O16. Here, a competition between FM metal−metal bonding and inter-block AFM correlations may also occur. Then the magneto-crystalline anisotropy turns to become of the same order of magnitude than the applied magnetic field which explains such fascinating (H,T) phase diagram.

**VIII. Magnetic entropy changes**

Figure 13a shows the temperature dependence of the magnetic entropy $\Delta S_M(T,H)$ calculated using the temperature dependence of $(\partial M/\partial T)_H$ determined at H=0.9, 2 and 5 T, multiplied by the field (see equation 1). Distinct features depending on the temperature and field values are observed. They are analyzed considering the four magnetic phases described in the (H,T) phase diagram (H,T).
Let remember that for FM materials a large peak centered at the Curie temperature is expected whereas a symmetrical-sign-reversed-phenomenon is expected at $T_N$ for AFM materials. When the temperature decreases, $\Delta S_M(T, H=0.9T)$ shows a large distribution of sign. Two different structures are observed. At temperature higher than 40K, $\Delta S_M$ is first positive and then negative at higher temperature. The peak areas are almost symmetric which support the magnetic transition in the antiferromagnetic state at $T_N$. The second specific feature is the large magnetic entropy change centered at 20K in association with the smallest one at 5K. Here, the system changes its arrangement from AFM to a weak ferromagnetic ordering. However, the large dissymmetry between positive and negative $\Delta S_M(T, H=0.9T)$ domains, suggest that the spin flop transition is not entirely completed and that some canted AFM are still giving in average a ferrimagnetic-like ordering. Under increasing field, two broad peaks are promoted. The one centered at 25K is significant of the FM arrangement. The second peak around 64K at 2T and 55K at 5T is characteristic of the magnetic entropy change due to the spin-flop transition. In summary, the magnetic entropy change is mediated by the competition between the internal magnetic mean field which supports AFM arrangement along the c axis and the external applied magnetic field promoting the FM-like spin-flop transition. At 50K, it results in a field-tuned sign of the magnetic entropy which indicates the possibility of the material to be used as cooling or heating medium depending of the magnetic field amplitude.

If one now considers the intrinsic values of the magnetic entropy for $\text{Ba}_7\text{Co}_6\text{BrO}_{17-\delta}$, an average spin value per mole could be obtained from the Curie–Weiss law. A theoretical magnetic entropy of about 10 J/kg.K is thus expected using the expression $S_M=R \ln(2S+1)$. If one sums the maximum magnetic entropy change observed at 60K (0.8 J/K.Kg from 13.a) and 30K (1.1 J/K.Kg from 13.a) when 5T is applied, almost 80% of magnetic entropy is missing in contrast with the study of $\text{Ba}_5\text{Co}_5\text{O}_{14}$ for which more than 50% of the magnetic entropy is missing. In contrast with $\text{Ba}_5\text{Co}_5\text{O}_{14}$ which shows canted (ab)-plan ferromagnetic behaviour below 50K, the magnetic
entropy for Ba$_7$Co$_6$BrO$_{17-\delta}$ is spread out in a wide range of temperature and distributed within at least three phenomena (i) the short range ordering (ii) the spin reorientation (iii) the spin-flop transition.

If the total relative cooling power (RCP) as defined in section I is now considered, it is enhanced by the two distinct transitions. In a first approximation $\Delta S_M(K,5T)$ is fitted using two Gaussian functions in order to separate the different contribution of the 3 phenomena listed before (see figure 13b). The RCP for applied magnetic field change of 5T is estimated to 38 J/Kg and 15 J/kg for the spin flop transition along the applied magnetic field and for the spin reorientation transition respectively. The total RCP could be here estimated to $(1.1+0.8)/2 \times 80 \approx 76$J/kg, where the first term represents the mean $\Delta S_{\text{max}}$ at 5T in J/K.Kg, and 80 represents an estimated full width at half maximum in Kelvin, see fig. 13b. Interestingly is the use of the total area under the three distinct phenomena responsible for the magnetic entropy which expresses the total refrigerant capacity of the system. The short range ordering is responsible for the offset of 0.1J/K.kg considered before fitting the curve. Then the refrigerant capacity of the short range ordering ($Q_{\text{SR}}$) is estimated to about 15 J/kg on the temperature range [0K- 150K]. It is about the same order than the refrigerant capacity due to the spin reorientation transition ($Q_{\text{SR}}\approx 19$ J/kg) but three times smaller than $Q_{\text{SR}} \approx 48$ J/kg the refrigerant capacity due to the spin flop transition. Most of the magnetic entropy in Ba$_7$Co$_6$BrO$_{17-\delta}$ is due to the spin flop. Once, the influence of both the magneto-crystalline anisotropy $^{30}$ and of field induced order/order transition$^{31}$ on the magnetocaloric effect is unambiguously shown.

**IX CONCLUSIONS**

The magnetic phase diagram of Ba$_7$Co$_6$BrO$_{17-\delta}$ has been experimentally investigated by neutron diffraction and magnetic measurements. Even if the fundamental magnetic state is an AFM
arrangement along the c axis, this study shows that an applied magnetic field induces a spin reorientation transition toward the (a,b) plane, with variable magnitude depending on the temperature. By comparison with the parents layered 5H or 12H-BaCoO$_{3.\delta}$ hexagonal polytypes, the spin reorientation is explained by a reduction of the magnetocrystalline anisotropy when a extra degree of disconnection between the 2D-blocks is introduced by the [Ba$_2$O$_2$Br]$^-$ double-layers. It enables various magnetic paths between the blocks. A clear competition between the inter-tetrameric blocks antiferromagnetic exchange interaction along the c axis $J_c$ and intra-tetrameric blocks ferromagnetic exchange interaction $J_{ab}$ through the (ab) plane is suggested. As soon as any magnetic field is applied, the two dimensional spin-correlations in (ab) plane are promoted.

The second new insight of this study concerns the preponderance of the magnetocrystalline anisotropy on the magnetocaloric properties. It highlights the potentialities of materials with intrinsic multiple magnetic transitions. Of course, efforts are now provided to modify the title compound toward higher temperature transition and local magnetic moments. Here the chemical substitution of iron for cobalt appears promising.

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8 Tishin A M and Spichkin Y I 2003 The Magnetocaloric Effect and its Applications (Bristol: Institute of Physics Publishing)
27 Y. B. Kusadov, EuroPhysics letters 78 78 (2007) 57005
Figures Caption:

Figure 1: Crystal and magnetic structure under zero magnetic fields for a) Ba$_6$Co$_5$BrO$_{14-\delta}$; b) Ba$_7$Co$_6$BrO$_{17-\delta}$.

Figure 2: a) Temperature dependence of the normalized Magnetization measured under 1T for Ba$_6$Co$_5$BrO$_{14-\delta}$. b) Temperature dependence of the normalized Magnetization measured under 1T for Ba$_7$Co$_6$BrO$_{17-\delta}$. c) Isothermal Magnetization versus magnetic field at the indicated temperature for Ba$_6$Co$_5$BrO$_{14-\delta}$. d) Isothermal Magnetization versus magnetic field at the indicated temperature for Ba$_7$Co$_6$BrO$_{17-\delta}$.

Figure 3: experimental and calculated neutron diffraction patterns at 1.4 K (G4.1, LLB, $\lambda=2.425\text{Å}$) for a) Ba$_6$Co$_5$BrO$_{14-\delta}$; b) Ba$_7$Co$_6$BrO$_{17-\delta}$ The vertical bars indicate calculated Bragg peak positions for both nuclear (first row) and magnetic structures (second row).

Figure 4: Thermal evolution of the magnetic moments per cobalt atom for a) Ba$_6$Co$_5$BrO$_{14-\delta}$; b) Ba$_7$Co$_6$BrO$_{17-\delta}$. The values were obtained from neutron diffraction data Rietveld refinement recorded on G4.1 line (LLB, Saclay).

Figure 5: Temperature evolution of selected magnetic diffraction peaks for Ba$_7$Co$_6$BrO$_{17-\delta}$. Note that intensity at 10K is higher than at 1.4K.

Figure 6: Neutron diffraction pattern of a non-densified powder sample of Ba$_7$Co$_6$BrO$_{17-\delta}$ (a) without Magnetic field; (b) with H= 1.2T. The magnetic field clearly induces preferential orientations.
Figure 7: Evolution of the neutron diffraction pattern with the magnetic field for Ba$_7$Co$_6$BrO$_{17-\delta}$. The data were measured on D2B line (ILL, Grenoble) equipped with a cryomagnet allowing field variation from 0 to 2.5T.

Figure 8: Evolution of the magnetic moment held per cobalt atom (bottom) and of the flopping angle for Ba$_7$Co$_6$BrO$_{17-\delta}$. The values were obtained from neutron diffraction data Rietveld refinement recorded on D2B line (ILL, Grenoble) equipped with a cryomagnet allowing field variation from 0 to 2.5T.

Figure 9: Temperature dependence of the normalized Magnetization measured under different field for Ba$_7$Co$_6$BrO$_{17-\delta}$.

Figure 10: M(T)/H curves under different fields corrected from the Curie-Weiss Law susceptibility extracted on the lower temperature range. The inset show a magnification of the curve in the region near the ordering temperature. The arrow point out the change in sign observed when the applied field increases.

Figure 11: Isothermal Magnetization versus magnetic field at the indicated temperature.

Figure 12: Diagram H-T built from the experimental data (Temperature dependence of the Magnetization and Isothermal Magnetization versus magnetic field). SRO is a short range ordering phase; AF is the antiferromagnetic phase; CAF is a canted antiferromagnetic phase; Weak FM is a weak ferromagnetic phase resulting from the spin-flop transition. Solid lines are guides for eyes.
Figure 13: a) Magnetic entropy change ($-\Delta S_M$) as function of temperature under magnetic field changes deduced from the data presented figure 9 and the use of equation 1. 13, b) The fit of $\Delta S_M(T,5T)$ assuming two Gaussian functions centred at 25.5 K and 57.8 K.
Figure 2
Figure 3

\[ \text{Ba}_2\text{Co}_2\text{Br}_6 \]  

\[ R_f = 0.045 - R_{\text{Bragg}} = 0.056 \]
\[ R_{\text{Mag}} = 0.058 \]

\[ \text{Ba}_2\text{Co}_2\text{Br}_7 \]  

\[ R_f = 0.029 - R_{\text{Bragg}} = 0.039 \]
\[ R_{\text{Mag}} = 0.049 \]
Figure 4

(a) and (b) show the temperature dependence of magnetization for different samples labeled Co1, Co2, and Co3.
Figure 6
Figure 9
Figure 10
Figure 11

[Graph showing magnetisation (μB/f.u.) versus magnetic excitation (μ₀H - Tesla) at different temperatures: 66K, 53K, 22K, and 12K.]
Figure 12

Graph showing the relationship between temperature (K) and magnetic susceptibility ($\mu_0 H$) for weak ferromagnetic (FM) and antiferromagnetic (AF) phases transitions.
Figure 13