Coupled negative magnetocapacitance and magnetic susceptibility in a Kagomé staircase-like compound \( \text{Co}_3\text{V}_2\text{O}_8 \)

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Abstract
The dielectric constant of the Kagomé staircase-like \( \text{Co}_3\text{V}_2\text{O}_8 \) polycrystalline compound has been measured as a function of temperature and magnetic field up to 14 T. It is found that the application of an external magnetic field suppresses the anomaly for the dielectric constant beyond 6.1 K. Furthermore, its magnetic field dependence reveals a negative magnetocapacitance which is proportional to the magnetic susceptibility, suggesting a common magnetostrictive origin for the magnetic field dependence of the two quantities. This result is very different from that obtained from the isostructural compound \( \text{Ni}_3\text{V}_2\text{O}_8 \) that presents a peak in the dielectric constant at the incommensurate magnetic phase transition coupled to a sign change of the magnetocapacitance.

(Some figures in this article are in colour only in the electronic version)

1. Introduction
Developing new multifunctional materials for electronics of the future is a challenging goal for solid-state chemists and physicists [1]. This kind of research has been illustrated by the multiferroic materials and, in particular, those with a coupling between ferroelectricity and ferromagnetism [2]. Such materials are developed for multiple-state-memory elements based on the writing of ferroelectric data and reading the induced magnetic field change [3]. Another class of multifunctionality is shown by materials with magnetocapacitance effects [4], i.e. for which the dielectric constant (\( \varepsilon \)) depends on the application of an external magnetic field. In the search for the magneto-electric effect in the 1960s, Schmid proposed a classification of systems in terms of symmetry [5]. Since ferromagnetic order does not allow time-inversion symmetry such as spatial polarization inversion symmetry, the number of systems showing both ferromagnetic and spatial polarization is highly restricted [6, 1]. However, the recent
discovery of new multiferroic materials, in which ferroelectricity is induced by magnetic order, has revived research into magneto-electric coupling. These systems are frustrated spin-density-waves magnets in which magnetic order breaks inversion symmetry, leading to ferroelectricity when the spin rotation axis is not parallel to the wavevector [7], as in RMnO₃ (R = Gd, Tb, Dy) [8], RMn₂O₅ (R = Tb, Ho, Er, Dy) [9–12] or Ni₃V₂O₈ [13].

The compound Ni₃V₂O₈ [13] is an extensively studied system belonging to this class of multiferroicity. Magnetism in this compound comes just from the S = 1 spin of Ni²⁺ cations that are distributed in a staircase-like Kagomé lattice [14]. This structure presents two inequivalent sites for Ni²⁺ and magnetic frustration, leading to a complex (H, T) magnetic phase diagram [15]. Four magnetic phase transitions were revealed by specific heat, magnetic measurements [16] and neutron diffraction [17]. The first kind of magnetic order appears below 9 K, consisting of a high-temperature incommensurate magnetic structure (HTI) described by a modulated amplitude in which only one of the two nickel sites is concerned. Then, cooling down further, at T = 6.3 K another low-temperature incommensurate structure is found (LTI), described as a helical order concerning both nickel sites. For the latter, the breaking of inversion symmetry induces the charge displacement responsible for a polarization along the b-axis [18] which can be controlled via an external magnetic field [13]. Below the temperature of appearance for these incommensurate phases, two canted antiferromagnetic commensurate phases take place with a weak ferromagnetic moment along the crystallographic axis c at T = 4 K (Cʹ) and T = 2.3 K (C).

Two other isostructural magnetic compounds have been synthesized Co₃V₂O₈ [19] and β-Cu₃V₂O₈ [20] with spin S = 3/2 and S = 1/2, respectively. In this paper we report on the study of the magnetic and dielectric properties of the spin S = 3/2 compound, Co₃V₂O₈, also characterized by four magnetic phase transitions [21]. These results are compared to the Ni₃V₂O₈ ones.

2. Experimental details

The polycrystalline sample of Co₃V₂O₈ has been prepared by solid-state reaction in a silica tube. The Co₃O₄, V₂O₅ and V₂O₃ in the 1:0.5:0.5 ratios were mixed in an agate mortar. After pressing, the sample was put in a finger-like alumina crucible which was set in an ampoule. The latter, sealed under primary vacuum, was heated at 1100 °C for 12 h and then quenched in air. The structural characterization of the black bars of ceramics that were obtained was performed by x-ray powder diffraction by an X-pert Pro diffractometer. All the diffraction lines could be indexed in the orthorhombic structure of Co₃V₂O₈ (space group Cmca), in good agreement with [16]. The corresponding structure with the Kagomé staircase-like structure is given in figure 1.

To obtain the values for the dielectric constant, complex impedance measurements were performed using a commercial AG4284A LCR meter. In order to measure in a magnetic field, a complete sample holder with four coaxial cables was designed. This allows measurements of the dielectric constant to be made inside a Quantum Design PPMS. The magnetization was measured with the magnetic option (extraction method) of the PPMS. The electrical contacts were deposited by an ultrasonic method, with indium solder covering two parallel faces of the sample in order to approximate the geometry of a parallel-plate capacitor. For accuracy, an ac voltage of 1 V at a frequency of 100 kHz was applied to the sample. Nonetheless, a systematic check was also performed to rule out any frequency and amplitude dependence of the results. In particular, the frequency dependence of the complex impedance allows the modelling of our samples as a capacitor and a resistance in parallel. The dielectric constant has been extracted from the value of the capacitance by using the sample dimensions.
3. Results and discussion

Recent magnetization and heat capacity studies [21] of a single crystal of Co$_3$V$_2$O$_8$ have shown four magnetic phase transitions at 11.2, 8.8, 6.6 and 6.1 K. Following the magnetic phase diagram of the isostructural compound Ni$_3$V$_2$O$_8$, the same progression of magnetic phases was proposed [21]. This interpretation suggests an equivalent ferroelectric LTI phase between 8.8 and 6.6 K, and the presence of transitions with a peak in the dielectric constant. There was no polarization measurement in Co$_3$V$_2$O$_8$, and we did not measure this property either. A neutron scattering study shows that incommensurate phases occur below 6.6 K (LTI) and between 8.8 and 11.2 K (HTI) [22]. In the same paper, preliminary measurements of the dielectric constant as function of temperature show no peak at 8.8 K, but rather an abrupt change at $\sim 6$ K. We have reproduced this anomaly (figure 2, upper panel). The application of a magnetic field softens the anomaly that extends its temperature range of observation up to high temperatures, around 30 K (figure 2, bottom panel), while no effect is observed below 6 K. Furthermore, this anomaly is completely suppressed by the application of a sufficiently large magnetic field ($\geq 4$ T). The behaviour is reminiscent of the anomaly observed in the ferroelectric hexagonal RMnO$_3$ (R = Y, Lu, Sc) [23, 24] when entering in the antiferromagnetic phase, and in the ferroelectric perovskite compound BiMnO$_3$ when crossing the ferromagnetic transition [4].

Isothermal curves of the magnetic field dependence of the dielectric constant have also been collected. Below 6 K, no effect has been observed (not shown). This was interpreted as a ferromagnetic phase where no effect of the magnetic field is observed [22]. In contrast, for $T > 6$ K, very interesting data have been obtained. We observed three types of behaviour presented in figure 3: for temperatures above 30 K (illustrated by $T = 50$ K), a quadratic dependence is observed, similar to what was observed in YMnO$_3$ [25] and BiMnO$_3$ [4]. As the temperature is decreased, the shape evolves continuously towards a peak centred at $H = 0$, whose width decreases (illustrated by 20 K). Below 8.8 K, the peak becomes very sharp (illustrated by 7 K). Remarkably, whatever the temperature is, this evolution follows the
magnetic field dependence of the magnetic susceptibility ($\chi = dM/dH$) which is also shown in the same figure (right, y-axis).

A similar trend observed for the dielectric constant and the magnetic susceptibility suggests a common origin. According to the strong coupling between spins, charges and the lattice often found in transition metal oxides, it is proposed that the application of the magnetic field induces a continuous structural distortion, responsible for the changes in both magnetization and dielectric susceptibility.

Let us start the discussion of the magnetic susceptibility by the measurement of its temperature dependence (figure 4). The data are very well fitted by a Curie law giving $\theta = 4.4$ K and $\mu_{\text{eff}} = 5.14$ $\mu_B$, in agreement with the previous data [21]. For 50 and 20 K, the magnetic field dependence of the susceptibility in the paramagnetic state follows Langevin’s law, where the important control parameter is $g\mu_B H / k_B (T - T_c)$, which exhibits a dome shape as function of magnetic field. At 50 K only the top of the dome is visible, and at 20 K the entire bell shape is observed. In contrast, at lower $T$ (below 8.8 K) there is a sharp peak at $H = 0$ associated with a sharp transition from the zero-field frustrated antiferromagnetic phase [22] to a ferromagnetic component reported by magnetization [21]. In the paramagnetic phase, it is particularly clear that the Zeeman energy and the spin–orbit coupling on a single magnetic site are the only terms which are relevant and are at the origin of the magnetic field variation of both magnetic and dielectric susceptibilities.
Figure 3. Magnetic field dependence of the relative variation of the dielectric constant at different temperatures for Co$_3$V$_2$O$_8$: left axis (black dots), $\Delta \varepsilon /\varepsilon_0$ is defined by $(\varepsilon(H) - \varepsilon(0))/\varepsilon(0)$; right axis (mid-grey/red lines), the magnetic susceptibility is also reported for comparison.

Figure 4. Temperature dependence of the magnetic susceptibility registered at 10 Oe (1000 Hz) of Co$_3$V$_2$O$_8$. A fit by a Curie law (in mid-grey/red) is also presented with the two parameter values $\theta = 4.4$ K and $\mu_{\text{eff}} = 5.14 \mu_B$ for the high-temperature part of the curve (above 15 K).
Figure 5. Temperature dependence of dielectric constant under different magnetic fields for Ni$_3$V$_2$O$_8$. The peak is attributed to the ferroelectric transition at $T_{\text{LTI-HTI}}$.

Figure 6. Magnetocapacitance of Ni$_3$V$_2$O$_8$ for different temperatures, showing positive magnetocapacitance in the LTI and HTI phases and negative in the region of the phase transition.

This is very different from the dielectric susceptibility of Ni$_3$V$_2$O$_8$ which is shown in figure 5 in the incommensurate phases above 4 K. The peak that is observed at $H = 0$ corresponds to the LTI–HTI transition [20]. Under the application of an external magnetic field, the peak broadens and shifts to lower temperatures following the magnetic transition, as already reported [15], up to 8 T. Above 8 T, the new data reveal that the peak broadens very rapidly and disappears. In order to follow the corresponding phase transition, the evolution of the dielectric constant as a function of magnetic field is recorded (figure 6). We have found a positive magnetocapacitance below $T_{\text{LTIT}}$ (6.3 K), while a negative one is found above. Above 7 K (figure 5), the magnetocapacitance is again positive, decreasing slowly up to 20 K in the paramagnetic phase. At 4.5, 5 and 6 K, the LTI–HTI transition appears at about 8.5 T as a cusp.
(figure 6). Small additional features at low field (1–2 T) appear at 4.5 and 5 K, which do not correspond to any reported magnetic transition.

4. Conclusions

The present result shows that the magnetocapacitance of the two isostructural compounds (Co$_3$V$_2$O$_8$ and Ni$_3$V$_2$O$_8$) is very different. In the Ni-containing compound, the majority of the magnetic phases is paraelectric and the only LTI phase is ferroelectric. The dielectric constant exhibits a peak at the LTI–HTI magnetic phase transition. The magnetocapacitance is positive in the LTI and HTI phases, with a sign change observed in the region of the LTI–HTI transition. In contrast, the compound Co$_3$V$_2$O$_8$ always shows a negative magnetocapacitance which is proportional to the magnetic susceptibility. This suggests strongly that a structural distortion is induced by the magnetic field. The existence of a strong spin–orbit coupling in the case of octahedral coordination of the high-spin Co$^{2+}$ could play a crucial role.

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