Multiferroicity in an organic charge-transfer salt: Electric-dipole-driven magnetism

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Multiferroics, showing simultaneous ordering of electrical and magnetic degrees of freedom, are remarkable materials as seen from both the academic and technological points of view.¹² A prominent mechanism of multiferroicity is the spin-driven ferroelectricity, often found in frustrated antiferromagnets with helical spin order.¹³,¹⁴,¹⁵ There, similar to conventional ferroelectrics, the electrical dipoles arise from an off-centre displacement of ions. However, recently a different mechanism, namely purely electronic ferroelectricity, where charge order breaks inversion symmetry, has attracted considerable interest. Here we provide evidence for this exotic type of ferroelectricity, accompanied by antiferromagnetic spin order, in a two-dimensional organic charge-transfer salt, thus representing a new class of multiferroics. Quite unexpectedly for electronic ferroelectrics, dipolar and spin order arise nearly simultaneously. This can be ascribed to the loss of spin frustration induced by the ferroelectric ordering. Hence, here the spin order is driven by the ferroelectricity, in marked contrast to the spin-driven ferroelectricity in helical magnets.

In the present work, we have investigated single crystalline κ-(BEDT-TTF)₂Cu[N(CN)₂]Cl (κ-CI), where BEDT-TTF stands for bis(ethylenedithio)-tetrathiafulvalene (often abbreviated as ET). Two crystals with different geometries and contact materials were investigated (see methods section). In these compounds, dimers of ET molecules form an anisotropic triangular lattice with a half-filled dimer band, where the strong on-dimer Coulomb interaction drives the system to a Mott insulating state.³ The importance of intra-dimer degrees of freedom and inter-site interactions has been pointed out.²⁴,²⁵-k-CI consists of alternating conducting ET layers and insulating anion sheets (see Supplementary Information, SI, Fig. S1). Within the ET layers, adjacent molecules form dimers on which a single electron hole is located. Below T_N ≈ 27 K, intralayer antiferromagnetic and interlayer ferromagnetic ordering of hole spins occurs, followed by weak ferromagnetic canting below 23 K (refs. 12, 13). κ-CI becomes superconducting below 12.8 K, when applying pressures of 300 bar.¹⁴

Figure 1 shows the temperature dependence of the conductivity σ of κ-CI, measured at 2.1 Hz, providing a good estimate of the dc conductivity σac, see SI. Aside of the well-known overall semiconducting characteristics of σac(T)¹⁴,¹⁵, we find a jump-like decrease by two decades at around 27 K, about the same temperature where long-range antiferromagnetic ordering is reported.¹⁶ A corresponding jump was also found in sample 2 (see SI). A distinct drop in σac has not been observed in most ac and dc studies reported so far (except in ref. 16), which may be ascribed to the higher precision of dielectric measurements at high resistances, compared to standard dc setups. This abrupt reduction of the conductivity within the Mott insulating state is a first hint to a charge-order (CO) transition, i.e. a further localization of charge carriers.

Figure 2 shows the dielectric constant ε(T) of sample 1 for various frequencies. Pronounced peaks reaching absolute values up to several hundreds are revealed. While the peak positions are nearly frequency independent, their amplitudes become strongly suppressed with increasing frequency. The overall behaviour is typical for order-disorder-type ferroelectrics, where the electric dipoles that are disordered at high temperatures order with net polarization below a phase-transition temperature T_ε (refs. 17, 18). Identifying the dipoles with the holes fluctuating within the dimers at T > T_ε and cooperatively locking in to one of the two ET molecules at T < T_ε, provides a plausible scenario for the present material (see insets of Fig. 1). Similar behaviour as

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observed in Fig. 2 was also found for other ferroelectric charge-transfer salts. The peaks are superposed to a rather high background contribution \( \omega_0 \) of about 120, partly arising from stray-capacitance contributions due to the small sample dimensions. The dashed line in Fig. 2 demonstrates that the high-temperature plateaus of \( \varepsilon(T) \) at low frequencies (i.e., the static dielectric constant \( \varepsilon_s(T) \)) are consistent with a Curie-Weiss behaviour with an additional background term, \( \omega = C(T-T_{CW}) + \omega_0 \) with a Curie-Weiss temperature \( T_{CW} = 25 \, \text{K} \approx T_{CW} \). Sample 2 showed similar dielectric response to that seen in Fig. 2 (see SI), despite having significantly different geometry and when different contact material was used – clear evidence for the intrinsic nature of the observed behaviour. In a previous dielectric investigation of \( \kappa\text{-Cl}^{17,18} \), relaxational behaviour was deduced from frequency-dependent dielectric measurements below 30 K, consistent with the notion of order-disorder ferroelectricity. Due to the restricted temperature range in that work, no \( \varepsilon(T) \)-peaks were detected. Likewise, measurements of the in-plane dielectric properties showed no signatures of ferroelectric order.25

Further evidence for ferroelectric ordering in \( \kappa\text{-Cl} \) is provided by so-called “positive-up-negative-down” (PUND) measurements,31 where a sequence of trapezoid pulses (Fig. 3a) is applied to the sample. The resulting current response (Fig. 3b) shows strong peak-like features occurring when the electric field \( |E| \) exceeds about 10 kV/cm for the first and third pulse. Obviously, here the macroscopic polarization of the sample is switched, i.e., the dipoles within the ferroelectric domains reorient along the field, implying a motion of charges and, thus, a peak in \( I(t) \). As expected for ferroelectrics, at the second and forth pulse no corresponding features show up since the polarization was already switched by the preceding pulse. (The horizontal sections in \( I(t) \) represent the trivial charging and discharging currents of the sample capacitor.) The typical PUND response was clearest for low frequencies, \( \nu < 100 \, \text{Hz} \).

Stimulated by these results, we have also performed field-dependent polarization measurements, \( P(E) \). Typical results are shown in Fig. 3c. Above \( T_{RT} \), elliptical curves are observed (crosses), typical for linear polarization response (paraelectricity) with additional loss contributions from charge transport. However, at lower temperatures, a clear onset of non-linearity above about 8.5 kV/cm occurs and saturation at the highest fields is observed (circles), as expected for ferroelectrics. The saturation polarisation of several tenths of \( \mu \text{C/cm}^2 \) compares well to that of the ferroelectric charge-transfer salt tetrathioualene-p-bromanil (0.2 \( \mu \text{C/cm}^2 \)),32 and the electronic ferroelectric and multiferroic magnetite (0.5 \( \mu \text{C/cm}^2 \)).33 In the electronic ferroelectrics \( \text{LuFe}_2\text{O}_4 \) (25 \( \mu \text{C/cm}^2 \)) and \( \text{Pr}_{1-x}\text{Ca}_x\text{MnO}_3 \) (44 \( \mu \text{C/cm}^2 \)), somewhat higher values are found. Interestingly, at higher fields, \( |E| > 12 \, \text{kV/cm} \), breakdown effects were observed. This may indicate that the sample resistance has become too low and that interplane charge transport sets in. Thus there is a rather restricted range of roughly 9–12 kV/cm for nonlinear polarization response in \( \kappa\text{-Cl} \).
driven ferroelectrics. Could this be the mechanism generating multiferroicity in \(\kappa\)-Cl? Indeed, it has been pointed out that a finite DM interaction is present in \(\kappa\)-Cl\(^{11,12}\). However, to our knowledge no experimental evidence for a helical magnetic structure in this material has been found until now.

To check for a spin-driven mechanism, we have performed magnetic field-dependent dielectric measurements. Usually, spin-driven multiferroics show a strong dependence of the dielectric properties on externally applied magnetic fields (so-called magnetocapacitive effects)\(^{12,27}\). Moreover, in \(\kappa\)-Cl a spin-flow transition at 0.25 Tesla into a direction perpendicular to the field has been found below 23 K (refs. 12,13). In the spin-driven scenario, the spin-flow transition would affect the dielectric properties through \(P \propto Q \times (S_i \times S_j)\). Remarkably enough, no variation of the dielectric properties in fields up to 9 T was found (see SI), ruling out a spin-driven mechanism for multiferroicity in \(\kappa\)-Cl\(^{3}\).

This suggests that for \(\kappa\)-Cl a purely electronic, CO-driven ferroelectric state, as also considered for other ferroelectric charge-transfer salts\(^{8,19,28}\), is realized. In fact, in the seminal article by van den Brink and Khomskii\(^8\), charge-transfer salts were considered as good candidates for multiferroics with a ferroelectric state driven by CO. The mechanism discussed in that work is based on a combination of bond- and site-centred CO leading to the breaking of inversion symmetry. This provides a plausible scenario for \(\kappa\)-Cl: Just as in the structurally related \(\kappa\)-(ET)\(_2\)Cu\(_2\)(CN)\(_3\), at high temperatures each hole can be assumed to be delocalized on an ET dimer\(^{10,11}\). This corresponds to pure bond centring (cf. Fig. 1c of ref. 6). Below \(T_{FE}\), the holes lock in to one of the two molecules of the dimers, establishing additional site centring (Fig. 1d in ref. 6). This will primarily lead to a polarization parallel to the planes, and, due to the inclined spatial orientation of the ET molecules\(^8\), a polarization along \(b\) is also induced.

In contrast to the spin-driven systems, for CO multiferroics, the ferroelectric and magnetic transition temperatures can differ significantly\(^{8,25,29}\). How can the nearly identical transition temperatures in \(\kappa\)-Cl be explained within the above framework? The magnetism in \(\kappa\)-Cl arises from the holes’ spins, which, at \(T > T_{FE}\), are already well defined on each dimer. As the holes fluctuate within the dimers, their spins – on average – sit at the centres of the dimers, forming an anisotropic triangular lattice within the \(ac\)-planes (indicated by lines in the schematic drawing in the inset of Fig. 1), giving rise to geometrical frustration. Indeed, in \(\kappa\)-(ET)\(_2\)Cu\(_2\)(CN)\(_3\), where frustration is even stronger, a spin-liquid state is formed\(^{29}\). There, dielectric measurements\(^9\) have revealed relaxor ferroelectricity, i.e. short-range cluster-like ferroelectric order. However, in \(\kappa\)-Cl we find long-range ferroelectricity, caused by CO, which implies the collective off-centre positioning of holes within the dimers below \(T_{FE}\). Consequently, the holes (and spins) no longer reside on a highly-frustrated triangular lattice (inset of Fig. 1). The frustration should therefore be reduced and magnetic order sets in (see SI).

In summary, the observed conductivity, dielectric and polarization properties of \(\kappa\)-Cl provide evidence for CO-driven electronic ferroelectricity, making this material the first example of a multiferroic charge-transfer salt. Moreover, our measurements indicate that ferroelectric order drives the magnetic order, in contrast to the well-known spin-driven mechanism in the helical magnets! Thus \(\kappa\)-Cl not only represents a new class of multiferroics but also provides the first example for a new coupling mechanism of magnetic and ferroelectric ordering.

Methods
Crystall growth. Crystals of \(\kappa\)-(BEDT-TTF)\(_2\)Cu[N(CN)\(_2\)]Cl were grown according to literature methods\(^{12,27}\). Sample 2 was grown on the surface of a platinum electrode, as is typical for growth by electrocrysallization. It was roughly plate-like (\(b\)-axis vertical to the surface) with an area of about 0.3 \(\times\) 0.5 mm\(^2\) and a thickness of 0.2 mm. Occasionally during the electrocrysallization process, crystals grow on the bottom of the electrochemical cell, removed from the electric field at the electrode surface, and have a rod-like morphology. X-ray diffraction measurements show both morphologies are the same crystalline phase. Sample 1 was grown by this method. The long crystal axis (parallel to the \(b\) crystallographic axis) was about 0.4 mm and a cross section of about 0.07 mm\(^2\).

Dielectric measurements. For the dielectric measurements, contacts of graphite paste (sample 1) or evaporated gold (sample 2) were applied at opposite sides of the crystals, ensuring an electrical field direction parallel to \(b\), i.e. in all measurements, the electrical field was directed perpendicular to the ET-planes. The dielectric constant and conductivity were determined using a frequency-response analyzer (Novocontrol \(\alpha\)-Analyzer) and an autobalance bridge (Hewlett-Packard HP4284). For sample cooling at zero magnetic field, a \(^{3}\)He-bath cryostat (Cryovac) was used. Magnetic field-dependent dielectric measurements with \(H / b\) up to 9 T were performed in a Quantum Design physical properties measurement system (PPMS). Additional measurements up to 1.5 T, with the magnetic field directed parallel to the \(ac\) plane, were performed using a conventional electromagnet. In this case, sample cooling was achieved by a closed-cycle refrigerator. For the non-linear measurements, a ferroelectric analyzer (aixACCT TF2000) was used where cooling was also being provided by a closed-cycle refrigerator.

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Supplementary Information

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Crystal structure and magnetic frustration

Figure S1 shows the crystal structure of \( \kappa-(ET)_2Cu[N(CN)_2]Cl \) consisting of alternating layers of ET molecules (C10S8H8) and insulating anion sheets. For materials with sufficiently large overlap of the molecular orbitals, the ET layer is conducting, providing a quasi-two-dimensional electronic band structure1. In the \( \kappa \)-type packing motif, the ET molecules form dimers, where adjacent dimers are arranged orthogonal to each other. On the basis of the crystal structure, the hopping energies (transfer integrals \( t \) and \( t' \)) of charges between adjacent molecules have been derived by ab initio calculations2,3. These studies yield a ratio of hopping energies for the holes between dimers, forming an anisotropic triangular lattice (see insets of Fig. 1 of main paper), of \( t'/t \sim 0.44 \) for \( \kappa-(ET)_2Cu[N(CN)_2]Cl \) and \( t'/t \sim 0.83 \) for \( \kappa-(ET)_2Cu_2(CN)_3 \) (ref. 2) Thus the degree of frustration \( t'/t \) is distinctly smaller for \( \kappa-(ET)_2Cu[N(CN)_2]Cl \) as compared to \( \kappa-(ET)_2Cu_2(CN)_3 \).

For \( T > T_{FE} \) the electrons reside on average between the dimers’ molecules, forming an only weakly distorted triangular lattice. Upon entering the charge-ordered state below \( T_{FE} \), however, the spins are located on a highly distorted triangular lattice, the symmetry of which is significantly reduced as compared to \( T > T_{FE} \). In fact, when the calculation for \( \kappa-(ET)_2Cu_2(CN)_3 \) in ref. 10 of the main paper is applied to \( \kappa-(ET)_2Cu[N(CN)_2]Cl \), the symmetry of the anion layer is significantly reduced as compared to \( T > T_{FE} \). In fact, when the calculation for \( \kappa-(ET)_2Cu_2(CN)_3 \) in ref. 10 of the main paper is applied to \( \kappa-(ET)_2Cu[N(CN)_2]Cl \), the symmetry of the anion layer is significantly reduced as compared to \( T > T_{FE} \).

Figure S1 | Crystal structure of \( \kappa-(ET)_2Cu[N(CN)_2]Cl \). Left: side view displaying the layered structure of ET molecules separated by thin insulating anion sheets. The unit cell is indicated by blue lines. Here, the hydrogen atoms are omitted for clarity. Right: top view of the anion layer (upper panel) and the ET layer (lower panel), selected dimers are highlighted by grey ellipses showing the \( \kappa \)-type packing motif of the ET molecules. The direction perpendicular to the planes with the ET molecules is the crystallographic b axis. The anion layers are parallel to the ac plane. The polymeric anion chains are running along the a direction.
(ET)₂Cu[N(CN)₂]Cl, the value of J'/J = (t'/t)¹/₂ in the charge-ordered state decreases to about 1/3 of that found in the dimer-Mott insulating state.

Conductivity and Néel temperature

Figure S2 shows the temperature dependence of the conductivity σ of crystal 1, measured at various frequencies for T < 50 K. We found no significant frequency dependence of σ at T > Tₙ ≈ 27 K. The frequency dependence observed at lower temperatures, revealed in Fig. S2, follows a power law, σ = σ₀ + σ₁Tⁿ, with n < 1, typical for hopping conductivity. However, at T < 100 Hz there is no detectable frequency dependence of σ down to the lowest temperature. Thus, the curve at 2.1 Hz shown in Fig. 1 of the main article provides a good estimate of the dc conductivity σ₀. Below the CO transition close to Tₙ, which reduces the dc conductivity by two decades, obviously hopping becomes the dominant charge-transport process.

Dielectric response of sample 2

Figure S4 shows the dielectric constant ϵ(T) of sample 2 for various frequencies. In contrast to sample 1, for sample 2 measurements below 1 kHz were impeded by its higher conductivity. Otherwise, very similar behaviour to sample 1 is observed (cf. Fig. 2 of main article). The characteristic temperatures of sample 2 are somewhat higher (peak at ~30 K, T_{CW} = 31 K), than in sample 1. This may indicate different levels of defects, acting as dopands, also mirrored by its higher conductivity. The peaks are superposed to a somewhat lower background contribution than in sample 1 of about 60, mainly arising from stray-capacitance contributions. The dashed line in Fig. S4 demonstrates that the static dielectric constant ϵ₀(T) is consistent with a Curie-Weiss behaviour with an additional background term, ϵ₀(T) = C(T-T_{CW}) + ϵ₀, leading to a Curie-Weiss temperature T_{CW} = 31 K ~ T_{N}.

Note that slightly different Néel temperatures have been reported, depending on the technique applied and the sample studied: While dc magnetization and ¹H-NMR relaxation rate in the work by Miyagawa et al. yield a Tₙ ~ 26 – 27 K (ref. 5), the ¹³C-NMR studies by Smith et al. suggest a Tₙ slightly in excess of 30 K (ref. 6). The latter value is consistent with anomalies observed in the resistivity by Ito et al. (ref. 7). Lefèbvre et al. report a Néel temperature of Tₙ ~ 25 K at ambient pressure deduced from a peak in the ¹H-NMR relaxation rate.

Figure S3 shows σ(T) at 2.1 Hz in two representations for temperatures above the phase transition (T > 30 K). Obviously σ₀(T) neither follows a simple thermally activated behaviour nor a Mott’s T¹/₄-law predicted by the variable-range-hopping model. One may speculate on the presence of two different activated regions as indicated by the broken lines in Fig. S3a, but their significance should not be overemphasized.
The absolute values of ε(T) of samples 1 and 2 are of similar order \(10^7\) at 100 kHz. These numbers are distinctly smaller than those observed for the (TMTTF)_2X salts, where the dielectric anomalies reach values of order \(10^9\) at 100 kHz (see, e.g., ref. 19 of the main paper). For the latter materials, the charge disproportionation could be directly observed through distinct line splittings in NMR and infrared (IR) spectroscopy. The comparatively small dielectric anomalies detected for the present material suggest that the corresponding features in NMR and IR spectroscopy are very small.

Dielectric measurements in magnetic field

Figure S5 compares ε(T) at 56.2 kHz of sample 2 measured at a magnetic field of 9 T with \(H || b\) and at zero field. Obviously, the applied magnetic field has no effect on the dielectric constant. Similar results were obtained for fields up to 1.5 T, directed parallel to the planes. The inset shows field-dependent measurements performed at 30 K and several frequencies. Again no indications of any magnetocapacitive effects are found negating a spin-driven multiferroicity mechanism in κ-Cl.

There are some variations in the absolute values of the peak amplitude and position shown in Figs. S4 and S5. This may be ascribed to inner strains arising during cooling or strains produced from a small amount of "Apiezon N" grease applied to one side of the sample to ensure good thermal coupling to the sample mount of the PPMS. κ-Cl already becomes superconducting at 300 bar and a high sensitivity of the dielectric properties to pressure may be suspected. Pressure-dependent dielectric measurements are being prepared in order to address this point in detail.

Lattice effects

The observed ferroelectric transition implies that the inversion symmetry of the high-temperature structure is lost. The scenario suggested here, i.e., a combination of site- and bond-centred CO, naturally breaks the inversion symmetry. As pointed out in the main text, such an effect may be very small, beyond the resolution of structural investigations. However, as demonstrated for the (TMTTF)_2X salts\(^1\), lattice effects accompanying the charge-order transition can be well resolved by means of thermal expansion measurements.

In fact, high-resolution thermal expansion measurements, performed both along the out-of-plane and in-plane direction of κ-Cl\(^7\), give clear evidence for lattice effects in the out-of-plane direction accompanying the CO transition, see Fig. S6. The results are qualitatively similar to those in the (TMTTF)_2X salts\(^1\), where the charge modulation along the TMTTF stacks gives rise to displacements of the counterions.

![Figure S5](image)

**Figure S5** | Magnetic field dependence of the dielectric constant. The main frame presents ε(T) of sample 2, measured for \(H || b\) without and with a magnetic field of 9 T. The inset shows the dielectric constant at 30 K and various frequencies as a function of the magnetic field. For all measurements shown, the magnetic field was oriented perpendicular to the planes, i.e., \(H || b\).

![Figure S6](image)

**Figure S6** | Thermal expansion coefficient perpendicular to the planes. The main frame presents the linear coefficient of thermal expansion measured along the out-of-plane direction, \(\alpha_T\), of κ-Cl around the ferroelectric CO transition. The inset shows the same data in a plot \(\alpha_T/T\) vs. \(T\) highlighting a distinct peak slightly above the reported Néel temperature of 27 K.

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