The ferroelectric Mott-Hubbard phase of organic \((TMTTF)_2X\) conductors.

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Abstract

We present experimental evidences for a ferro-electric transition in the family of quasi-one-dimensional conductors \((TMTTF)_2X\). We interpret this new transition in the frame of the combined Mott-Hubbard state taking into account the double action of the spontaneous charge disproportionation on the \(TMTTF\) molecular stacks and of the \(X\) anionic potentials.

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Low dimensional electronic systems serve as a workshop on problems of strong correlations. The richest opportunities have been opened by the family of charge transfer salts \(M_2X\) formed of stacks of organic molecules tetramethyltetrathiofulvalene \((M = TMTTF)\) or tetramethyltetraselenafulvalene \((M = TMTSF)\) with anions \(X = PF_6, AsF_6, SbF_6, SCN,\) etc as counterions. The ions sit in loose cavities delimited by the methyl groups of the organic molecules. These materials show almost all known electronic phases: a metal, a paramagnetic insulator (PI), spin/charge density waves (SDW/CDW), the spin-Peierls state and finally the superconducting state \([1]\). In parallel, there is a set of several different structural types due to anion ordering (AO) which are slight arrangements in \(X\) chains \([2]\). Their tran-
osition temperatures $T_a \approx 100 - 200 K$ are much higher than that corresponding to electronic transitions occurring in the range of $T_c \approx 1 - 10 K$.

Very recently, in the already very rich phase diagram of these salts, a new phase transition has been discovered, being revealed by a huge anomaly in the low frequency dielectric constant $\epsilon'$ at $T_0 = 70 K$ in $(TMTTF)_2PF_6$ and at $100 K$ in $(TMTTF)_2AsF_6$ [3]. Very soon NMR studies have proved [4] that the transitions at $T_0$ are related to charge disproportionation, also called charge ordering (CO), dividing molecules within in two non equivalent species. Previous reports on TMTTF salts with larger anions such as $SbF_6$ have suggested the occurrence of a phase transition, sometimes called “structureless” transition, at a temperature of the scale of $T_a$ showing itself in changes of the resistance [5] and the thermopower [6], in weak features in high frequency ($\sim GHz$) dielectric susceptibility [7]. However, X-rays investigations brought no observation of superlattice reflections [5] while some variation of intensity of Bragg reflections could actually have been noticed [10]. NMR allows to identify the very fact of the CO via nonequivalence of molecular sites [4,8] but it cannot distinguish between the $\vec{q} = 0$ state and the known undulating ones (either along the stack like the $4k_F$ (Wigner) condensation in $(DCNQI)_2Ag$ [8] or between stacks like in $(TMTTF)_2SCN)$ [2]). So, the identification of the nature of this “structureless” $\vec{q} = 0$ transition was still unsolved. We show hereafter that $\epsilon'$ in $(TMTTF)_2SbF_6$, as in the case of $(TMTTF)_2PF_6$ and $(TMTTF)_2AsF_6$, exhibits a huge divergence at $154K$. Dielectric experiments are providing thus an unique access to the anomalous collective properties of this mysterious transition identifying the resulting state as a never expected ferroelectric (FE) one.

In the high temperature range $T > T_a, T_0$ the counterions already have a profound effect on the electronic properties: the anions dimerize the intermolecular spacing of the TMTTF molecules along the stacks. As a result, the conduction band is split into a filled lower band separated from a half-filled upper band by a dimerization gap, leading to Mott-Hubbard charge localization. The effects brought by the AOs with $\vec{q} \neq 0$ have been previously discussed [2,9] and refs. therein. The new effects of combined CO-AO with $\vec{q} = 0$ are
the main topic of the present work. Empirically one can see a systematic difference between the usual $\vec{q} \neq 0$ and $\vec{q} = 0$ transitions. The first ones have been always observed for non centrosymmetric (NCS) anions [10]; so, the orientational ordering for which a short contact interaction between the anion and the molecule is established was considered as the main mechanism, positional displacements being only the consequence of it. In a contrary manner to $\vec{q} \neq 0$ AOs, the $\vec{q} = 0$ ones are observed in systems with centrosymmetric (CS) anions.

For such systems, we will present hereafter a possible universal mechanism only related to a positional instability. The remarkable fit of the anomaly in $\epsilon'(T)$ for the three centrosymmetric salts, $X = PF_6$, $AsF_6$ and $SbF_6$, to the Curie law suggests firmly the ferroelectric character of these phase transitions. We account for this FE state from the combined action of charge disproportionation and ionic displacements.

The $(TMTTF)_2SbF_6$ were prepared by a standard electrochemical procedure [11]. We present results on a sample with a regular cross-section of $4 \times 10^{-5} cm^2$ and a length of 3 mm. Its room temperature conductivity was about $10^{-1} cm^{-1}$ close to the value previously published [4]. The complex conductance $G(T)$ was measured in the frequency range $1 kHz - 10 MHz$ using an impedance analyzer HP 4192A. The amplitude of the ac signal was typically $10 mV$, within the linear response. With decreasing temperature, $G$ increases continuously (metallic type of conductivity) to a maximum slightly above $T_0 = 154 K$. Below $T_0$, $G$ sharply decreases with a thermally activated behavior with an activation energy $\Delta = 500 K$. From room temperature to about $200 K$ the magnitude of the $\epsilon'$ is relatively small (below our experimental resolution in this temperature range). With decreasing $T$ below $200K$, we observed a sharp growth of the magnitude of $\epsilon'$ with a tendency to diverge near $T_0 = 154 K$, reaching the huge value of $\sim 10^6$ (at $100 kHz$), and a very deep decrease of the magnitude of $\epsilon'$ below $T_0$. Fig.1 shows the temperature dependence of $1/\epsilon'$ in the $(TMTTF)_2X$ family with centrosymmetric anions $PF_6$, $AsF_6$, $SbF_6$. The generic features are very similar for the three compounds: the two branches above and below the appropriate $T_0$ are very close to be linear, i.e. to follow the Curie law $\epsilon' \sim /|T - T_0|$. The slopes of these branches (i.e. the amplitudes of $A$) at $T < T_0$ are twice that at $T > T_0$, exactly as theoretically
predicted in the description of a second order phase transition.

One can get simply a general idea of the hidden ordering in considering the joint effect of two sources to the dimerization, hence to the charge gap \( \Delta \). The *extrinsic* one is determined by the basic crystal structure and *intrinsic* one is spontaneous being induced by the electronic sub-system itself. (This concept was intensively used in the study of conducting polymers \([12]\)). The gap \( \Delta(W) \) appears as the consequence of both contributions to the Umklapp scattering \( W \). Without CO, there is only the bond contribution \( W_b \) to the amplitude \( W \) from the built-in bond alternation. The CO adds the on-site contribution \( W_s \) from the nonequivalence of sites. The charge gap is a function of the total amplitude \( W = \sqrt{W_s^2 + W_b^2} \) which results from the orthogonality of corresponding matrix elements of electronic scattering \([12]\). (Since the gap is not a linear function of \( W \), then the partial gaps do not add simply in quadrature as the components of \( W \)). The energy change \( F_e \) of the electronic system due to both \( W_b \) and \( W_s \) depends only on the \( \Delta \) and consequently on the total \( W \). But the energy of lattice distortions depends only on the spontaneous site component \( W_s \): \( F_l = 1/2KW_s^2 \). Thus the total energy can be written in terms of the total \( W \) as \( F_{tot} = F_e(W) + 1/2KW^2 - 1/2KW_b^2 \). The ground state is determined by its minimum over \( W \) under the restriction \( W > W_b \). The spontaneous \( W_s \) will appear if the two conditions are satisfied: a) the energy has a minimum at some value \( W = W_0 \) and b) \( W_b \) is not too big to meet the restriction \( W_0 < W_b \). Since the value \( W_0 = W_0(T) \) increases with decreasing temperature, there will be a phase transition at \( W_0(T_0) = W_b \) provided that \( W_0(0) > W_b \).

Now we are going to choose a microscopical description. Recall first a recent theoretical work \([14]\) performed in the frame of a mean field (MF) approximation which has found that the CO can occur as the result of strong enough Coulomb interaction between electrons on the near-neighbor sites. This model has been developed at \( T = 0 \) and it includes ultimately the spin ordering of AFM type which is characteristic for these systems at low temperatures (about 10K). The phase transitions we observe occur at much higher temperatures (100 – 200K) and an appropriate approach is clearly needed. Considering the case of real 3D materials, in our picture the CO is stabilized by interaction with the 3D connected system
of anions. So in the following we will use the bosonization procedure which is suitable for describing low energy and collective processes.

Let consider a 1D electronic system with a mean occupation of 1/2 electron per site (that is per molecule) with sites along the stack labeled as $n$ at $x = na$. The low frequency long wave effects are described by the Hamiltonian for the phase $\varphi$ in the charge channel which is separated from the Hamiltonian for the spin channel. We shall normalize the phase according to conventions for CDWs and SDWs which are $2k_F = \pi/(2a)$ modulations $\sim \cos[\pi/2n + \varphi(x) + \text{cnst}]$. The $4k_F = \pi/a$ fluctuations of the charge density are $\rho_{4k_F} \sim (-1)^n \cos[2\varphi(x) + \text{cnst}]$. For our crystals with two molecules per unit cell the $4k_F$ is projected to $q_x = 0$, then $\rho_{4k_F}$ describes the charge disproportionation within the unit cell.

The Hamiltonian for $\varphi$ (per site) can be written \[ \text{(1)} \] as:

$$H_0 + H_u \sim \frac{\hbar}{4\pi\gamma} \left[ v_\rho (\partial_x \varphi)^2 + v^{-1}_\rho (\partial_t \varphi)^2 \right] - W \cos(2\varphi)$$

Without interactions $v_\rho = v_F$ and $\gamma = 1$. Repulsive interactions reduce $\gamma < 1$ which value carries all necessary information: either about on-site/inter-site interactions of the model \[ \text{(1)} \] or about Coulomb interactions for the WC scenario like in \[ \text{(8)} \]. The Umklapp scattering amplitude $W$ of Luther and Emery or $g_3$ of Dzyaloshinskii and Larkin \[ \text{(15)} \] is a feature of systems with one electron per unit cell. Normally $W$ of the order of other interactions is not small; hence a big gap is opened in the charge $\varphi$– channel, so that only gapless spin degrees of freedom are left for observations. But a specific feature of $(TMTCF)_2X$ crystals is that $W$ is small and appear only as a secondary effect of the anionic sublattice, opening an intriguing crossover to ”charge localization” \[ \text{(1)} \] (to PI in earlier language of \[ \text{(9)} \]).

The $4k_F$ susceptibility $\Pi_{4k_F} \sim T^{4\gamma-2}$ is divergent only for strong enough interactions when $\gamma < 1/2$. For $\gamma > 1/2$ the Umklapp term $\sim W$ is renormalized to zero and the system is a 1D metal, nowadays called the Luttinger Liquid. At the marginal line $\gamma = 1/2$, known in terms of bare interactions as the Luther - Emery line, $\Pi_{4k_F} \sim \ln T^{-1}$ like for the Kohn anomaly leading to the Peierls instability (then the Hamiltonian is reduced to spinless fermions which actually are $\pi$ solitons, with the gap $\Delta$). At arbitrary $\gamma < 1/2$, $W$
is renormalized to a finite value which we shall write as $W^* = \hbar \omega_t^2 / (8\pi \gamma v_p)$, $\omega_t \lesssim \Delta$. The frequency $\omega_t$ is a gap in the spectrum of linear phase excitations $\omega^2 = (v_p k)^2 + \omega_t^2$ and it enters, as a "transverse" frequency of the optical response, the expression for the dielectric susceptibility

$$\epsilon_\Delta = \gamma \frac{v_p \omega_p^2}{v_F \omega_t^2}, \quad \omega_p^2 = \frac{8e^2 v_F}{\hbar s}$$

where $\omega_p$ is the plasma frequency of the parent metal and $s$ is the area per stack. This contribution $\epsilon_\Delta \sim (\omega_p/\Delta)^2$ can be relatively large as $\sim 10^{3-4}$ and it corresponds to the background upon which the anomaly at $T_0$ is developed. But $\epsilon_\Delta$ is regular, showing only a dependence on $\Delta$ which starts to increase below $T_0$ without signs of decrease from above. These features would only add some upward kink in $\epsilon'$ below $T_0$ but cannot account for the observed anomaly.

The degeneracy between $\varphi = 0$ and $\varphi = \pi$ corresponds to the displacement of the electronic system by just one lattice position $x$. Hence the $\pm \pi$ soliton just adds/removes one electron in real space. The elementary excitations registered as charge carriers are these solitons. Their activation energy $\Delta \sim \omega_t$ is determined from our results for $G$ as e.g. $500K$ for $SbF_6$. At finite concentration of solitons as normal carriers $\sim \rho_n$ coming either as thermal excitations or via incommensurability, there is an additional contribution to the low frequency complex $\epsilon$ which we can write in a Drude form $\epsilon_n = -\rho_n \omega_p^2 / (\omega^2 + i\omega/\tau_n)$ to be assigned to the observed conductivity $^3$.

The scaling relations for the gap and the electronic energy read (in units of $E_F$) $F_e \sim -\Delta^2 \sim -W^\zeta$ with $\zeta = 1/(1-\gamma)$ So the instability condition is $\zeta < 2$ that is $\gamma < 1/2$. Indeed, at this condition the value $W$ can appear spontaneously: the gain of the $F_e$ is higher than the loss $\sim KW^2$ of energy due to spontaneous deformations being at the origin of the potential $W$. The minimum is achieved at $W = W_0 \sim K^{\beta}$ with $\beta^{-1} = 2 - \zeta = (1 - 2\gamma)/(1 - \gamma)$.

For our particular system it is, by principle, important to notice the two sources acting for the weak two fold commensurability, that is the two contributions to the Umklapp interaction. The non dimerized system with $1/2$ electron per site has a symmetry $x \rightarrow x + a$
and $x \to -x$ which corresponds to phase transformations $\varphi \to \varphi + \pi/2$ and $\varphi \to -\varphi$. Thus the lowest order invariant contribution to the Hamiltonian is $\sim W_4 \cos 4\varphi$ which is usually negligibly small as studies of CDW have shown. It is small as coming from Umklapp interaction of 8 particles, half of them staying far away from the Fermi energy; but also it is renormalized as $\sim W_4^\kappa$ so that superlow values $\gamma < 1/8$ are required for its stabilization. For the site dimerization the symmetry $x \to x + a$ is broken while the symmetries $x \to x + 2a$ and $x \to -x$ are preserved. The Hamiltonian, invariant under the corresponding transformations of $\varphi$: $\varphi \to \varphi + \pi$ and $\varphi \to -\varphi$ is $H_W^s = -W_s \cos 2\varphi$. The bond dimerization preserves the symmetries $x \to x + 2a$ and $x \to a - x$ (reflection with respect to the bond center). Hence the invariance is required with respect to $\varphi \to \varphi + \pi$ and $\varphi \to \pi/2 - \varphi$ leading to $H_W^b = -W_b \sin 2\varphi$.

Altogether the nonlinear Hamiltonian becomes

$$H_W = -W_s \cos 2\varphi - W_b \sin 2\varphi = -W \cos(2\varphi - 2\alpha);$$

$$W = \sqrt{W_b^2 + W_s^2}, \; \tan 2\alpha = W_b/W_s$$

For a given $W_s$ the ground state is doubly degenerate between $\varphi = \alpha$ and $\varphi = \alpha + \pi$ which still allows for phase $\pi$ solitons. Moreover $W_s$ itself can change its sign between different domains of anionic displacements. Then the electronic system must also adjust its ground state from $\alpha$ to $-\alpha$ or to $\pi - \alpha$, whichever is closer. Thus, at the domain boundary, a phase soliton of $\delta \varphi = -2\alpha$ or $\pi - 2\alpha$ will be placed carrying an non integer charge $q = -2\alpha/\pi$ or $1 - 2\alpha/\pi$ per chain. The CO modulation is $\rho_{co} \sim \cos(\pi x/a + 2\alpha) = (-1)^n W_s/\sqrt{W_s^2 + W_b^2}$.

The MF approach appears appropriate for anionic displacements: critical fluctuations are suppressed by well pronounced 3D correlations, as inferred from the perfect Curie like anomaly in $\epsilon'$ (Fig.1) as well as from X-ray studies on given AOs $[2]$. For electrons quantum fluctuations must be treated exactly as it has been sketched above. The angle $\alpha$ is invariant and only the total gap value is subject to the renormalization from $W$ to $W^\ast(W) \sim \Delta^2$.

Finally the total energy, as a function of the mean phase $\bar{\varphi}$ and $W_s$ at given $W_b$, is

$$F \sim -\Delta^2 \left( \frac{W_s}{W} \cos 2\bar{\varphi} + \frac{W_b}{W} \sin 2\bar{\varphi} \right) + \frac{K}{2} W_s^2 - \frac{e}{\pi} E \bar{\varphi}$$

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which includes contributions from electrons, lattice and electric field \( E \). At \( W_b > W_{cr} \sim K^{1/\beta} \) this energy has only one minimum at \( W_s = 0, \phi = \pi/4 \) (modulo \( \pi \)) but this point becomes unstable at \( W_b = W_{cr} \) from which the anomaly in \( \epsilon \) originates. At \( W_b < W_{cr} \) this minimum is split in two ones which appear, first closely, at \( 2\phi = 2\phi_\pm = \pi/2 \pm 2\delta\phi \), \( W_s = W_b \tan 2\delta\phi_0 \). The resulting ground state will be ferroelectric if the same \( \phi_\pm \) is chosen for all stacks (our case), while the state is antiferroelectric (AFE) if the sign of \( \phi_\pm \) alternates (the case of \((TMTTF)_2SCN\)), see refs. in [2,9]. Above the CO instability the minimization yields

\[
\phi = \pm \frac{\pi}{4} - \frac{eE}{W_b - W_{cr}}, \quad W_s = \frac{E/2\pi}{W_b/W_{cr} - 1}, \quad \epsilon' = \frac{4e^2/s}{W_b - W_{cr}}
\]

In writing \( \epsilon' \) we recall that \( e\phi/s\pi \) is the electronic polarization which equals \( eE \). We come back to the formula (2) but with a modified denominator: \( \omega_t^2 \to \omega_t^2 - \omega_{cr}^2 \). Thus \( \epsilon \to \infty \) at the transition point \( W_b = W_{cr} \) which is the joint electron-ion instability. Importantly, the value of \( \Delta \) is related to the bare value of \( \omega_t \) which stays finite, hence the "normal" carriers (the \( \pi \)-solitons) are gapful.

Doing short, we have supposed above that instead of changing \( T \) one lets the parameters \( K, W_{cr} \) or \( W_b \) to be varied at \( T = 0 \). Actually all of them are functions of \( T \) and the transition temperature \( T_0 \) is determined by \( W_b(T_0) = W_{cr}(T_0) \). Expanding around \( T_0 \) we recover the experimental observation \( \epsilon^{-1} \sim T - T_0 \) and we need only to understand why the Curie law extends over a rather broad region of \( \delta T = T - T_0 \approx 30K \). Clearly the expansion is valid when \( \delta T \) is small in comparing to the lowest energy scales which are the ionic cage energy \( \sim 10^2K \) [2] and \( \Delta \sim 10^2K \). It seems that relative to these scales the expansion within \( \sim 10^4K \) is well assured. Remarkably we arrive even at the right magnitude of the observed effect: \( \epsilon \sim 10^4T_0/(T - T_0) \).

In conclusion, we have shown that \((TMTTF)_2X\) salts with centrosymmetric anions \( X = PF_6, AsF_6 \) and \( SbF_6 \) undergo a phase transition at which the dielectric constant shows a huge divergent peak. The ferroelectric character of this phase transition is demonstrated by the Curie law behavior of \( \epsilon'(T) \) of the three compounds. These results also bring the
nature of the so-called structureless transition of the \((TMTTF)_2SbF_6\) salt with CS anions. The transition has been explained by the combined action of the uniform shift of anions with respect to the oppositely charged organic stacks and of the with charge disproportionation appearing in each \(TMTTF\) molecule yielding thus a macroscopic ferroelectric polarization. Our approach provides a physically transparent phenomenological interpretation in terms of strongly fluctuating \(4K_F\) density wave, i.e. a local Wigner crystal, subjected to a weak two fold commensurability potential. Up to now these FE transitions concern only the salts based on \(TMTTF\) molecules for which \(T_0\) is already in the regime of Mott Hubbard charge localization state. The salts based on TMTSF are less 1\(D\) and are metallic in this temperature range which preclude dielectric measurements. Nevertheless, the same type of the FE transition may exist, just being hidden or existing only in a fluctuating regime. If such a fact would be confirmed by future structural or NMR studies, then the whole analysis of intriguing abnormal properties in the normal state should be revised, as it already needs to be made, see also [4] for the PI phase of \(TMTTF\) salts with centrosymmetric anions as demonstrated by the present work.

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FIG. 1. Temperature dependence of the inverse of the real part of the dielectric permittivity $\varepsilon'$ of $(TMTTF)_2X$ with $X = PF_6, AsF_6, SbF_6$ at the frequency of $100Hz$. 