Interplay between bandwidth-controlled and filling-controlled pressure-induced Mott insulator to metal transition in the molecular compound \([\text{Au(Et-thiazdt)}_2]\)

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Optical properties of the quasi 2D single-component molecular Mott insulator [Au(Et-thiazdt)$_2$] (Et-thiazdt = N-ethyl-1,3-thiazoline-2-thione-4,5-dithiolate) have been investigated under pressure at room temperature. At 1.5 GPa, [Au(Et-thiazdt)$_2$] undergoes an insulator to metal transition (IMT). Optical conductivity spectra exhibit a clear Drude peak at high pressure. In addition, we observed a clear anisotropy of pressure induced modifications of the electronic structure. With increasing pressure, along the molecules stacks, a strong increase of the spectral weight below 1 eV is observed, while in the transverse direction, it remains barely constant with a redistribution from mid-infrared to low energy. Besides the increase of SOMO bandwidth (Singly Occupied Molecular Orbital), calculations show that the SOMO-1 bands cross the Fermi level at the transition. Moreover, we have calculated the optical conductivity as function of pressure to provide a picture of the compound physics under 1 eV. Our results indicate that the pressure induced IMT is simultaneously due to a bandwidth and a bandfilling phenomenon that imply both Mott physics and uncorrelated charge carriers.

One fascinating phenomenon in strongly correlated fermion systems is the transition from an insulator to a metallic state driven by electronic correlations. An electron in a crystal goes from localized to itinerant (or vice versa) when a control parameter such as doping, pressure or temperature is modified. For a half-filled system, this problem is known as the Mott transition [1]. Understanding the properties of Mott insulators has become of primary importance since the discovery of molecular and high-Tc superconductors [2-5] which have in common the fact that from half filled band insulator they become metallic and superconductor with doping (so called 'doped Mott insulator'). On this basis, extensive efforts have been devoted to develop new types of molecular metals and superconductors by enhancing the interstack interactions [6]. One solution to raise conductivity and to overcome the charge repulsion issue is to increase the electronic bandwidth $W$ with respect to the on-site Coulomb repulsion $U$ which can easily be done by applying pressure [7, 8], especially in organic material due to their soft nature. It has been extensively studied in Mott organic salts like the one-dimensional (1D) tetrathiafulvalene (TTF) [9,11] or the two-dimensional (2D) $\kappa$-(Bis(ethylenedithio)tetrathiafulvalene)$_2$X (BEDT-TTF) [12, 13]. Moreover, crystals of molecular conductors present numerous advantages for organic electronics compared to traditional inorganic materials: low fabrication cost, high mechanical flexibility, light weight and ease of fabrication [14]. For this purpose, the use of neutral organic radicals as building blocks is a promising track to design material with unpaired electrons which serve as charge carriers without any doping process [15].

Among single-component organic conductors based on bis(1,2-dithiolene) ligands [3, 16, 21], [Au(Et-thiazdt)$_2$] (Et-thiazdt = N-ethyl-1,3-thiazoline-2-thione-4,5-dithiolate) is the first well characterized molecular metal without TTF dithiolate ligands. It also is the first molecular member of a new class of Resistive Random Access Memory (RRAM) called 'Mott Memories'. RRAM applications are made possible by the resistive switching phenomenon (resistive transition induced by electric pulses). Whereas the resistive switching is usually obtained by electro-chemical effects or charge trapping, it is based on a pure electronic intrinsic effect in inorganic Mott GaTa$_2$Se$_3$ [22] and in [Au(Et-thiazdt)$_2$] [23]. This phenomenon could constitute a generic feature of Mott insulators. Indeed, with a 2D structure made of molecules stacked along $b$-axis and interstacked along $a$-axis (see supplemental material [24], part I.), [Au(Et-thiazdt)$_2$] is a half-filled electron system with significant electronic correlations that place it close to an IMT controlled by U/W. Pressure reduces U/W sufficiently to induce the IMT at 1.3 GPa [19].

In this paper we report a pressure-dependent optical study of [Au(Et-thiazdt)$_2$] single crystals made possible by use of a home-made high-vacuum micro-spectrometer and a synchrotron radiation source (see supp. mat. [24] and reference [25]). In order to understand the mechanisms of the IMT, we performed first-principles Density Functional Theory calculations (DFT). The study highlights the 'Mottness' of the compound and describes a new type of Mott transition which involves two types of charge carriers.

In order to elucidate the ambient pressure electronic structure at room temperature, we measured polarized optical conductivity (see Fig[1] [20, 27]. The spectra exhibit an insulating behavior in both $E||a$ and $E||b$...
Figure 1. Optical conductivity spectra between 43 meV and 1.2 eV for $E_{||a}$ (a) and $E_{||b}$ (b) at ambient pressure and room temperature. Dotted lines provide the gap. Red circles curves represent the sum of all the contributions.

directions. Although both spectra can be decomposed in the mid infrared (MIR) with at least two absorption bands roughly located at 0.43 eV and 0.74 eV, they differ in shape and level. Along $a$-axis, the amplitude of the 0.74 eV band is higher than the 0.43 eV one. Along $b$-axis, it is the contrary and the spectrum also exhibits an overall lower absolute level. Despite these differences, the optical gap is roughly the same in both directions (250 meV), in good agreement with transport experiments and previous optical measurements [23] (see supp. mat. [24] part I for the gap extraction procedure).

In order to assign the absorption features to electronic transitions, we performed ab-initio calculations [25, 28]. Calculation details are provided in the supplemental material [24]. Although DFT does not fully include electronic correlations, it provides valuable information as evidenced in various studies [15, 22]. The band structure with 'fatband' representation shown in Figure 2 allows to identify the Singly Occupied Molecular Orbital bands (SOMO) (green), the SOMO-1 (cyan), the SOMO-2 (blue) and the Lowest Unoccupied Molecular Orbital bands (LUMO) (orange). Remarkably, two SOMO bands cross the Fermi level $\epsilon_F$ driving the state metallic contrary to experiments. This is a typical signature of strong electronic correlations insofar as DFT underestimates on-site interactions. Moreover the SOMO-1 bands remain below $\epsilon_F$ and the bandwidth of SOMO bands $W = 0.37$ eV. Note that the band dispersion along the $\Gamma$-$Y$ direction is directly related to electron hopping along $b$-axis. The $\Gamma$-$X$ direction is close to the interstack direction $a$ since $\hat{a}\cdot \hat{c} = 91.83(5)^\circ$ [19]. Nevertheless, due to the monoclinic symmetry, the optical conductivity tensor is non-diagonal. The non-zero elements of the tensor are provided in Figure 3. The experimental conductivity along the $a$-axis (Fig.2.a), involves both $\sigma_{xx}$ and $\sigma_{xz}$ components (see supp. mat. [24] part II and reference [34]). $\sigma_{yy}$ is directly compared to the experimental conductivity along $b$ (Fig.1.b) [1].

Remarkably, $\sigma_{xx}$ and $\sigma_{xz}$ are dominated by an interband contribution SOMO-1 $\rightarrow$ SOMO-1 at 0.4 eV (cyan curves Fig.3). Note that a small intraband contribution (green curve) appears in $\sigma_{xx}$ due to the presence of SOMO bands at $\epsilon_F$. Moreover, two interband transitions SOMO-2 $\rightarrow$ SOMO and SOMO $\rightarrow$ LUMO are predicted around 1 eV. SOMO-1 $\rightarrow$ LUMO is predicted around 1.5 eV (above the experimental range). The calculated conductivity for the $E_{||b}$ case ($\sigma_{yy}$) is dominated by the SOMO intraband contribution. In addition, $\sigma_{xz}$ indicates an insulating character in agreement with the 2D character of the system.

It is crucial to notice that the experiment exhibits a first contribution around 0.43 eV (green contribution Fig.1) and a second one around 0.74 eV (cyan contribution) whereas the DFT predicts an intraband (green curve Fig.3) and an interband contribution SOMO-1 $\rightarrow$ SOMO around 0.4 eV (cyan curve). This discrepancy can be understood by taking into account the on-site correlations $U$. For a large enough $U$, SOMO bands are split into a filled Lower Hubbard Band (LHB) and an empty Upper Hubbard Band (UHB), resulting in an Mott insulator (Fig.4.a). By considering the correlations, the DFT intraband SOMO contribution should shift of $U$, towards higher energies [24]. Consequently the contribution at 0.43 eV, which has to correspond to the DFT intraband contribution, is assigned to electronic transitions LHB $\rightarrow$ UHB. Moreover, the DFT interband contribution SOMO-1 $\rightarrow$ SOMO should shift of $U/2$ towards higher energies in presence of correlations. Therefore, the experimental contribution at 0.74 eV could be assigned to SOMO-1 $\rightarrow$ UHB (empty states of SOMO) excitations. Hence, with an overall good agreement to the experimental data, we deduce $U = 0.43$ eV. Such a value of $U$ is commonly found in this class of Mott organic compounds [25, 26]. Note that, $U - W = 0.43 - 0.37 \approx 0.1$ eV which is the same order of the electronic gap. The higher energy excitations SOMO $\rightarrow$ LUMO and SOMO-2 $\rightarrow$ SOMO above the experimental spectral range will not be discussed.

The experimental polarized conductivity spectra presented in Figure 4 from 0 to 5 GPa (at room temperature), exhibit an obvious rising Drude-like response at low frequency above 1.5 GPa, a decrease and a closing of the gap between 1 and 1.5 GPa (Fig.4.d). Hence, by applying pressure, [Au(Et-thiazdt)$_2$] undergoes an IMT.

Remarkably, the low frequency data extrapolation represented in Figure 4.c are qualitatively consistent...
with room temperature transport measurements realized up to 2.1 GPa by Tenn et al. [19]. Indeed, as shown in Figure 5, they both exhibit the same pressure dependence and order of magnitude. Note that this extrapolation was obtained by fitting the low frequency reflectivity \( R_{sd} \) (measured through the diamond) with a Drude model. Also, error bars were extracted by evaluating the order of magnitude and the pressure dependence.

Figure 5. Comparison between room temperature \( \sigma_{DC} \) values (up to 2.1 GPa) obtained by transport measurements by Tenn et al. [19] and by extrapolation of the optical conductivity data at \( \omega = 0 \). They are qualitatively consistent according to both the order of magnitude and the pressure dependence.

In order to rule out possible structural change (from which could originate the IMT), we performed vibrational properties using infrared and Raman spectroscopies. The phonon frequencies as a function of pressure obtained by fitting the infrared and Raman spectra with Drude-Lorentz oscillators and Gaussian-Lorentzian profiles respectively are reported in the supplemental material [24]. Within experimental resolution, apart the usual blue shift of phonon modes (due to the volume lattice shrinking) no phonon anomaly (disappearance, appearance, splitting, kink in the phonon frequency pressure dependence) has been noticed at 1.5 GPa and beyond, in both infrared and Raman spectra. This indicates that no structural phase transition has been detected under pressure in agreement with the Mott transition picture.

Note that the anisotropy still persists at high pressure as it is clearly evidenced by the higher absolute level of the low frequency conductivity along \( b \) than \( a \) and the spectral weight (SW) values (see Fig 4e). In the \( b \) direction, SW increases between 0 and 3 GPa and is constant above 3 GPa, whereas along \( a \), it slightly decreases between 0 and 1.5 GPa and is constant above 1.5 GPa.

In order to understand the pressure effects on the electronic structure and to disentangle the IMT mechanisms, we performed a full optimization of cell parameters and internal coordinates at different pressures between 0 and 4 GPa within DFT [39], as successfully done for other organic molecular compounds [15, 10].

Pressure effects on band structure are depicted in Figure 6. Note that the lack of dispersion along \( \Gamma-Z \) indicates that the system mainly remains 2D at high pressure. In addition, the bands shape is almost unchanged between 0 and 4 GPa but the bandwidth increases from \( W = 0.37 \) to 1.05 eV for the SOMO and from 0.70 to 1.05 eV for the SOMO-1 bands. As a conse-
sequence, \( U/W \) goes from 1.16 at 0 GPa to 0.41 at 4 GPa, well below the critical value of \( U/W \approx 1 \) for which IMT is theoretically expected to occur \[11, \ 42\]. Consequently, the role of correlations is expected to be weaker when pressure increases and charge carriers are supposed to be less correlated. In addition, as shown in Figure 7, the evolution of the bandwidth \( W \) and the pressure dependence of the damping \( \gamma \) of the optical conductivity contribution assigned to LHB \( \rightarrow \) UHB transitions are compatible with an usual bandwidth enlargement observed in correlated 2D organic conductors through the Mott transition \[12, \ 43\]. The damping was obtained by fitting the Hubbard mid-infrared absorption band from 0 to 5 GPa with a Drude-Lorentz model. Remarkably, the increase of dispersion in both \( \Gamma-X \) and \( \Gamma-Y \) directions raises the energy level of the SOMO-1 bands going at \( \Gamma \) from 0 to 0.10 eV between 0 and 2 GPa. As a result, the SOMO-1 bands cross \( \epsilon_F \) and become hole filled. In contrast, the SOMO bands (previously assigned to UHB and LHB when correlations are included) go slightly deeper under \( \epsilon_F \) with pressure, getting electron filled. Hence, part of the IMT is achieved by tuning the electronic filling of the UHB suggesting a pressure induced doping of the Mott insulator. Thus, two mechanisms are driving the IMT: the bandwidth enlargement and the bandfilling. Pressure induces a bandwidth enlargement and an electronic doping of the SOMO bands (UHB) that implies physics related to correlated electrons ('Mottness'). On the contrary, since the SOMO-1 bands are far from half-filling at 0 and 4 GPa, the charge carriers arising from hole doping of SOMO-1 bands are expected to be less- or un-correlated.

Comparing the conductivity tensors calculated at 4 GPa (Fig. 8) and at 0 GPa (Fig. 9) we notice an enhancement of the low frequency conductivity for both \( \sigma_{xx} \) and \( \sigma_{yy} \) driving the IMT. Moreover, the SOMO-1 \( \rightarrow \) SOMO contribution is shifted of \( \approx 0.2 \) eV towards low frequency and its amplitude is also clearly lower than at ambient pressure. In contrast, for \( \sigma_{zz} \), apart from the decrease of the magnitude of the SOMO-1 \( \rightarrow \) SOMO contribution between 0 and 4 GPa, there is no significant changes at low frequency. Hence the system remains 2D as previously observed. Remarkably, the Drude contribution present in \( \sigma_{xx} \) and \( \sigma_{yy} \) is composed of two different intraband contributions: one due to SOMO bands and an other one produced by the hole filling of the SOMO-1 bands (cf. green curves of Fig. 8). Note that the SW of the total Drude is lower in \( \sigma_{xx} \) than in \( \sigma_{yy} \), underlining the remaining anisotropy at high pressure.

In order to get a global picture of the IMT, we both plot in Figure 9 the calculated (deduced from the non-diagonal optical conductivity tensor) and experimental...
optical conductivity at 0 and 4 GPa along a and b-axis. Calculations at 4 GPa in both directions are in good agreement with the experiment in many respects. Along the a-axis, the DFT SW (see inset Figure 9b) decreases slightly from 0 to 4 GPa and along b, it increases between 0 and 4 GPa. Moreover, the low frequency DFT conductivity level is consistent with the anisotropy evidenced experimentally since $\sigma_{ab}^{DFT} > \sigma_{ab}^{DFT}$ at high pressure. Noticeably, the agreement between DFT and experiments is clearly better at high pressure since the role of electronic correlations is expected to be weaker than at ambient pressure.

Naively, in correlated systems, we could consider that the electronically active bands at $\epsilon_F$ are ruled/dominated by electronic correlations. However, for [Au(Et-thiazdt)$_2$], other uncorrelated interactions seem to occur. The high pressure experimental conductivity can be fitted under 1 eV with a minimal model consisting of three Lorentz oscillators and one Drude contribution at 0.4 eV (green contribution Fig.10) and unpreceded by DFT, should correspond to a reminiscence of LHB→UHB transitions as predicted by DMFT.

In conclusion, a novel Mott IMT induced by pressure has been evidenced in the organic compound [Au(Et-thiazdt)$_2$]. The originality of the transition lies in the existence of two cooperative mechanisms (bandwidth and bandfilling) that drive the state metallic. On the one hand, SOMO-1 bands are filled with holes underlining the existence of uncorrelated charge carriers. On the other hand, the bandwidth enlargement of Hubbard bands and the filling (doping) of the UHB imply the presence of correlated carriers. Therefore, two types of charge carriers are expected to co-exist in the metallic phase as it appears in other intriguing systems [15-17]. Moreover considering the suggested ‘doped Mott insulator’ character of the system at high pressure, we could expected rich physics at low temperature as encountered for example in cuprates.

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