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Gold bis(dithiolene) radical with fused pyrazine and dithiine rings on dithiolene ligand turns metallic under pressure†

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Conducting molecular materials built on dithiolene complexes, as mixed-valence salts or single component materials, are most often based on planar structures which efficiently stack on top of each other. Herein, we report an original dithiolene ligand, namely, [1,4]dithiino[2,3-b]pyrazine-2,3-bis(thiolate) (hereafter noted as pzdtdt), which combines an electron-withdrawing pyrazine ring and a folded (by 40-50°) dithiine ring. Until now, such strong distortions from planarity have been hindering the isolation of highly conducting materials from dithiine-containing dithiolene complexes. However, in this study, we showed that the gold complex radical [Au(pzdtdt)₂]* obtained via electrocrystallization from the 1e⁻ oxidation of [Ph₄P][Au(pzdtdt)₂] organized into regular, non-dimerized chains in the solid state. Interestingly, [Au (pzdtdt)₂]* exhibited a semi-conducting behaviour at ambient pressure and turned metallic upon application of pressures above 4 GPa. The electronic structure of [Au(pzdtdt)₂]* was investigated in terms of electron localization effects through spin-polarized band-structure calculations.

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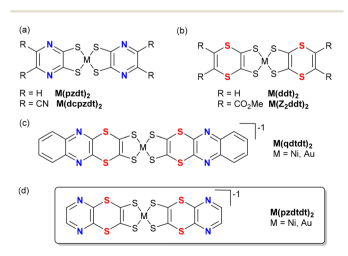
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Introduction

Bis(dithiolene) metal complexes are now more than sixty years old, and among their various applications, their utility as precursors for many highly conducting molecular materials,¹ such as the prototypical [Ni(dmit)₂] mixed valence salts² or their Pd analogs with quantum spin liquid (QSL) behavior, has garnered significant attention.3 Among the most recently studied complexes, the open-shell, neutral gold radical complexes [Au(dt)₂] are being investigated as single component molecular conductors. 4 Such radical complexes often exhibit a tendency to dimerize in the solid state,⁵ leading to a semiconducting behaviour, but recent examples, based on the original

substitution pattern on the dithiolene ligands, enabled the isolation of non-dimerized, regular stack structures with strong intermolecular overlap and associated metallic behaviour, either under high pressure or ambient pressure. The properties of such radical species are strongly dependent on the nature of the dithiolene ligands, often involving fused heterocycles incorporating nitrogen atoms (pyridine and pyrazine) or sulfur atoms (thiophene, dithiole, and dithiine) or both (thiazoline and thiazole). Among these, complexes with fused N-aromatic dithiolene ligands (Scheme 1a), such as pyrazine

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Scheme 1 Molecular structures of dithiolene complexes involving pyrazine, dithiine or both the heterocycles.

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moieties $[Au(pzdt)_2]^{8,9}$ and $[M(qdt)_2]$ (qdt = quinoxaline-2,3dithiolate), 10 have been reported for their ability to coordinate with other metals for the generation of MOF11 and for their proton-dependent solution luminescence properties. 12 However, owing to the strong electron-withdrawing effect of the fused pyrazine, only the reduced monoanionic gold complexes can be isolated, 13 and no single component molecular conductors based on the neutral species have been reported so far. Alternatively, the introduction of sulfur heterocycles at the periphery of the metal bis(dithiolene) complexes is extensively explored to favour intermolecular interactions in the solid state, as illustrated in numerous dmit and dddt complexes. 14 Besides, complexes of the dithiine dithiolate ligand (ddt, Scheme 1b) have also been studied, but only poorly conducting materials were obtained from M(ddt)₂ derivatives so far owing to a lack of sizeable intermolecular interactions, 15,16 which is a consequence of the non-planar structure of the dithiine ring that is strongly folded along the S...S axis. Consequently, no conducting mixed-valence nickel salts or conducting singlecomponent gold complexes have been reported to date from such systems. Only one complex combining the pyrazine and dithiine heterocycles has been reported (Scheme 1c), with two dithiino[2,3-b]quinoxaline-2,3-dithiolates surrounding either the Au³⁺ or Ni²⁺ in M(qdtdt)₂ complexes; however, no conducting materials have been prepared from them so far. 17

Therefore, we decided to investigate the synthesis of another dithiolene ligand with only the pyrazine and dithiine moieties and concentrated our efforts on the [1,4]dithiino[2,3-b]pyrazine-2,3-bis(thiolate) ligand (hereafter pyrazinedithiine-dithiolate is denoted as pzdtdt) (Scheme 1d). Herein, we report the preparation of the ligand precursor, its Ni and Au monoanionic complexes, and the successful isolation of the neutral gold complex radical [Au(pzdtdt)₂]*, behaving as a single-component molecular conductor, which is a remarkable behaviour when considering its non-planar geometry.

Results and discussion

Synthesis of the proligands

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For the synthesis of the dithiolene ligand (Scheme 2), we focused on the synthesis of the corresponding proligands 1 and 2, as 1,3-dithiole-2-thione (1) or 1,3-dithiole-2-one (2) derivatives, via a copper-catalysed coupling reaction between aryl iodides and zinc–thiolate complexes. Thus, reacting 2,3-diiodopyrazine and $[Et_4N]_2[Zn(dmit)_2]$ in the presence of Cu_2O afforded dithiole-2-thione 1 in 63% yield. Trans-chalcogenation with $Hg(OAc)_2$ afforded the desired proligand 2 in 94% yield.

In order to evaluate the possibly different reactivity of $\bf 1$ and $\bf 2$ as precursors of the dithiolate ligand, both were separately engaged with sodium methanolate followed by successive additions of KAuCl₄ and Ph₄PBr, to afford the monoanionic complex [Ph₄P][Au(pzdtdt)₂] in 10 and 40% yields, respectively, demonstrating the usefulness of the transformation of dithiole-2-thione $\bf 1$ into dithiole-2-one $\bf 2$ before treatment with

Scheme 2 Synthetic scheme of 1, 2 and [Au(pzdtdt)₂].

sodium methanolate. Note that the corresponding nickel complex has also been prepared, as Et_4N^+ salt, by the same method (see ESI for experimental details and properties†).

Both 1 and 2 were crystallized from CH_2Cl_2 , and their molecular structures were obtained from X-ray diffraction (Fig. 1). The dithiine ring in both of them strongly deviates from planarity, with a folding along the S···S axis of $49.3(2)^{\circ}$ and $50.0(3)^{\circ}$ in 1 and 2, respectively, a recurrent feature of such dithiine derivatives $(43-53^{\circ})$. 20,21

The redox properties of $[Au(pzdtdt)_2]^-$ were investigated by cyclic voltammetry in CH_3CN and the oxidation potentials are collected in Table 1, together with those of the related complexes. At variance with the pyrazinodithiolate complex $[Au(pzdt)_2]^-$ which oxidizes irreversibly at high potential (+1.12 V νs . SCE), the insertion of the dithiine ring between the pyrazine and the dithiolate moiety suppresses here to a large extent the electron-withdrawing nature of the pyrazine ring on the redox-active dithiolene and provides a -1/0 oxidation potential for $[Au(pzdtdt)_2]^-$ around +0.47 V νs . SCE (Fig. S2 in ESI†). Such an effect was already observed, in a more limited way, from the introduction of a benzo ring (rather than a dithiine one) in a Ni complex with two quinoxaline-2,3-dithiolate ligands²² compared with the complex surrounded by two

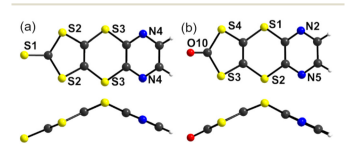


Fig. 1 Molecular structures of (a) 1 and (b) 2, showing the distortion of the central dithiine ring from planarity.

Paper

Table 1	Redox	potentials	of	[Au(pz	dtdt) ₂]	and	its	analogous	com-
plexes, c	btained	in CH ₃ CN	(<i>E</i> i	n V vs.	SCE, s	ee Scł	nem	e 1 for mo	lecular
structure	es). ΔE_{pa}	$=E_3^{0/+1}-$	E_{2}^{-1}	./0					

	$E_1^{-2/-1}$	${E_2}^{-1/0}$	$E_3^{\ 0/+1}$	ΔE_{pa}	Ref.
[Au(pzdtdt) ₂]	-1.01 ^f	+0.54/0.40 ^a	+0.96 ^f	0.42	This work
$[Au(pzdt)_2]$	-1.27^d (-1.66)	$+1.12^{d,f}$ $(+0.73)$	+1.89 ^{d,f} (+1.5)	0.77	8
$[Au(qdtdt)_2]$	-0.89^{b} -1.02^{c}	$+0.49^{b}$ $+0.44^{c}$		_	17
$[Au(Z_2ddt)_2]$	_	+0.86	+1.2	0.34	16 <i>b</i>
[Au(Py-ddt) ₂]	-0.93	+0.74	_	_	16 <i>c</i>
$[Au(Et-thiazdt)_2]^e$	-0.90^{f}	+0.55/+0.49	+0.71/+0.61	0.16	6 <i>a</i>

 $E=(E_{\rm pa}+E_{\rm pc})/2$. $^aE_{\rm pa}/E_{\rm pc}$: anodic and cathodic peak potentials. b As Bu₄N⁺ salt. c As Et₄N⁺ salt. d A correction of 0.39 V has been applied to the given redox potentials (in italics) versus Fc+/Fc. e Performed in CH₂Cl₂. f Irreversible.

Scheme 3 Molecular structures of quinoxaline-2,3-dithiolate and quinoxaline-6,7-dithiolate complexes.

quinoxaline-6,7-dithiolate (Scheme 3) which oxidizes at lower potential (by 300 mV).²³ This effect opens the possibility to isolate the radical [Au(pzdtdt)₂]* species within an accessible potential window. Besides, spectroelectrochemical experiments performed on [Ph₄P][Au(pzdtdt)₂] show, upon oxidation, the appearance of a low-energy NIR absorption band for the neutral complex centred at 1410 nm, characteristic of such radical complexes. 24,25

Accordingly, electrocrystallization experiments of a solution of [Ph₄P][Au(pzdtdt)₂] were conducted with [Bu₄N][PF₆] in THF as the electrolyte and crystals of the neutral radical species [Au(pzdtdt)2] were indeed successfully obtained applying a constant current intensity of 1 µA for 3 days. The molecular structure of the radical [Au(pzdtdt)₂]* can be now compared with that of the monoanionic [Au(pzdtdt)₂] complex, as detailed below, while its solid-state structure will be analysed in a second step, in relation with its conducting properties.

Solid state structures and properties

[Ph₄P][Au(pzdtdt)₂] crystallizes in the triclinic system, space group P1, with two crystallographically independent complexes, both located on inversion centres. The molecular structures of these two independent anions (Mol1 and Mol2) are shown in Fig. 2a, and the bond lengths and angles are shown in Table 2. As observed for the proligands 1 and 2, the 1,4-dithiine rings deviate from planarity with a distortion angle along the S···S hinge of 46.4(1)° and 49.5(4)° in Mol1 and Mol2, respectively. The AuS₂C₂ metallacycles of these

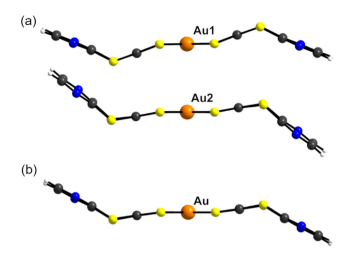


Fig. 2 Side views of (a) two crystallographically independent monoanionic [Au(pzdtdt)₂]⁻ complexes and (b) neutral complex [Au(pzdtdt)₂]^{*}, showing the recurrent distortion of the central dithiine ring from planarity

gold complexes are also distorted from planarity along the S...S axis with a much less pronounced effect (20.2(1)° and $10.4(2)^{\circ}$). The overall shape of this complex is similar to that of the corresponding monoanionic Au complex, with an additional fused benzene ring at each end, [Au(qdtdt)₂]⁻¹ (cf. Scheme 1c), where the dithiine folding angle was found to be 50.6(6)°.17

The neutral [Au(pzdtdt)₂]* complex crystallizes in the monoclinic system, space group $P2_1/n$, with one crystallographically independent complex located on an inversion centre. The neutral gold complex is slightly less distorted than its monoanionic counterpart: for the dithiine ring, the distortion angle decreases from 46-49° to 41.4°, while for the metallacycle, it decreases from 10-20° to 7.5°. Comparison of the intramolecular bond lengths within the monoanionic and neutral states shows that the main modifications concentrate on the AuS₂C₂ metallacycles, with a shortening of the C-S bonds (bond b) and a lengthening of the C=C bonds (bond c) upon oxidation, which is typical of the evolution of such dithiolene complexes upon oxidation. The absence of notable structural effects on the dithiine and pyrazine rings, together with the recurrent strong distortion of the dithiine ring from planarity, clearly indicates that the oxidation is strongly concentrated on the two metallacycles, with limited effects on the outer rings.

As shown in Fig. 3a, the radical complexes organize into layers in the (ac) plane, with very limited interactions between layers. Despite their non-planar structure, these radical complexes form regular stacks along the a axis with a gold-to-gold atomic distance of a = 3.70 Å (Fig. 3b). The shortest S···S contacts within the stacks amount to the same distance of a =3.70 Å, while shorter lateral S···S intermolecular contacts are found between stacks within layers (Fig. 3a), down to 3.49-3.52 Å.

The electrical resistivity and its temperature dependence were measured on single crystals of the neutral complex [Au

Table 2 Bond lengths (in Å) and dihedral angles (in °) in the monoanionic and neutral [Au(pzdtdt)₂]^{-1,0} complexes

Complex		а	b	С	d	e	f	$ heta_1$	θ_2	Ref.
$\overline{\left[\mathrm{Au}(\mathrm{pzdtdt})_2\right]^{-1}}$	Mol1	2.3116(9) 2.3085(8)	1.750(3) 1.751(3)	1.334(4)	1.754(3) 1.766(3)	1.768(3) 1.766(3)	1.394(5)	20.2(1)	46.4(1)	This work
	Mol2	2.3170(7) 2.3114(8)	1.751(3) 1.751(3) 1.754(3)	1.339(4)	1.766(3) 1.768(3)	1.764(3) 1.768(3)	1.404(4)	10.4(2)	49.5(4)	
$\left[Au(qdtdt)_{2}\right] ^{-1}$		2.3114(6) 2.300(4) 2.282(4)	1.734(3) 1.738(12) 1.738(12)	1.397(19)	1.731(12) 1.746(12)	1.733(16) 1.757(14)	1.470(25)	3.6(4)	53.03(6)	17
$[Au(pzdtdt)_2]^{\bullet}$		2.306(2) 2.291(2)	1.725(10) 1.723(9)	1.354(13)	1.764(9) 1.758(9)	1.760(10) 1.768(9)	1.407(13)	7.5(2)	41.4(10)	This work

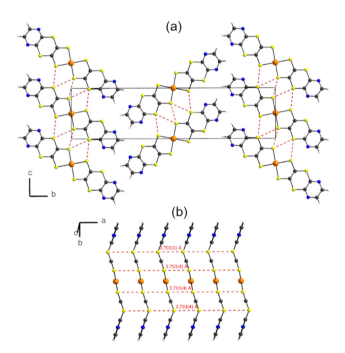


Fig. 3 Structure of $[Au(pzdtdt)_2]^-$ with (a) projection view of the unit cell along the a axis and (b) view of the stack running along a.

(pzdtdt)₂]' by the four-point technique at pressures between 1 bar and 19.6 GPa, using a diamond anvil cell (DAC). At room temperature and ambient pressure, the resistivity amounts to 489.5 Ω cm (σ_{RT} = 2.04 10^{-3} S cm⁻¹), and the compound is a semiconductor with an activation energy of 227 meV (Fig. 4a). Further application of pressure strongly increases the room temperature conductivity up to $\sigma_{RT}(4.2 \text{ GPa})$ = 150 S cm⁻¹ and decreases the activation energy (Fig. 4b). This exponential pressure dependence is observed up to 4.2 GPa to reach a quasi-metallic behaviour. At 4.8 GPa, the complex is metallic in the whole temperature range. Moreover, above 6.2 GPa (Fig. S5 in ESI†), a slight increase in both the resistivity and the activation energy is observed up to 19.6 GPa, with $\sigma_{RT}(19.6)$

GPa) = 2.33 S cm⁻¹ (ρ_{RT} (4.2 GPa) = 0.43 Ω cm) and E_{act} (19.6 GPa) = 22 meV.

Electronic structure

The unit cell of $[Au(pzdtdt)_2]$ contains two symmetry equivalent ac layers built from parallel chains of the $[Au(pzdtdt)_2]$ complex along the a-direction (Fig. 3a). Although there are numerous short S···S contacts within the ac layers, there are no S···S inter-layer contacts along b, so this solid is expected to be at most a 2D conductor. As shown in Fig. 5a, where all intermolecular S···S contacts shorter than 3.9 Å are shown as red dashed lines, there are three different types of intermolecular interactions, one (I) is intra-chain and two (II and III) are inter-chain interactions.

In principle the transport properties of [Au(pzdtdt)₂] should be determined by the SOMO···SOMO (Singly Occupied Molecular Orbital) interactions. However, previous work on such radical gold bis(dithiolene)⁷ complexes has shown that both the SOMO···SOMO and SOMO-1···SOMO-1 interactions should be considered when the complexes interact strongly along one of the two directions of the layer. In that case, there is indeed the possibility of a two-band behaviour, 26 i.e. some electron transfer can occur between the SOMO-1 and SOMO levels. The simplest way to get a hint on the relative strength of the SOMO···SOMO and SOMO-1···SOMO-1 interactions is by evaluating the so-called intermolecular interaction energies (β) . The absolute values of the interaction energies $|\beta_{\text{SOMO}...\text{SOMO}}|/|\beta_{\text{SOMO}-1}...\text{SOMO}-1}|$ calculated for this system are 0.1993/0.1960 (I), 0.0039/0.0375 (II), and 0.0043/0.0030 (III) eV: the inter-chain interactions are either one or two orders of magnitude weaker than the intra-chain interactions. Thus, it is clear that the intra-chain interactions completely control the electronic properties of the solid. These interactions are of similar strength for the SOMO and SOMO-1. The strength of interaction I is easy to understand since, as shown in Fig. 5b, such interaction is associated with four pairs of S...S interactions at 3.703 Å distance. Despite the sliding of adjacent complexes, the interaction has a substantial σ -type character

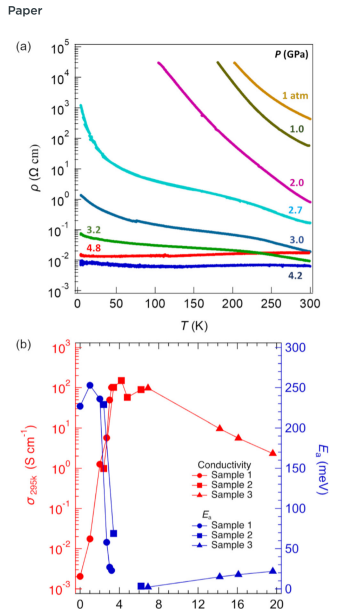


Fig. 4 (a) Temperature dependence of resistivity of [Au(pzdtdt)₂] at different applied pressures; (b) pressure dependence of the conductivity of [Au(pzdtdt)₂], and the activation energy at room temperature.

P (GPa)

and does not almost differentiate between the SOMO and SOMO-1 interactions. More surprising is the very weak character of the inter-chain interactions II and III despite quite short S···S contacts (3.495 and 3.766 (×2) Å for II and 3.528 (×2), 3.640 (×2) and 3.716 Å for III). These interactions are of the usually weak lateral π -type. However, except for the $\beta_{\text{SOMO-1}\cdots\text{SOMO-1}}$ of interaction II, the calculated values are around one order of magnitude too small when compared with the usual values for these interactions. As far as the interactions of the SOMO are concerned, since this orbital is antisymmetric with respect to the plane perpendicular to the inner core of the complex (see Fig. 6) separating the two ligands, several of the S···S interactions almost cancel and the final

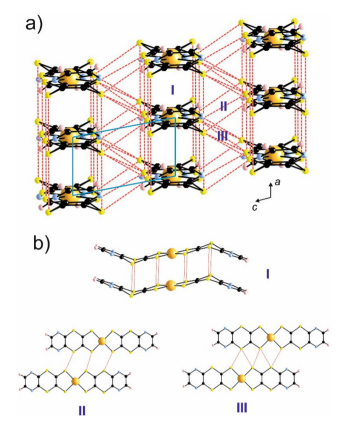


Fig. 5 (a) Layers of the $[Au(pzdtdt)_2]^{-1}$ solid, where the different intermolecular interactions are noted. The S····S bonds shorter than 3.9 Å are shown as red dashed lines. (b) Three different intermolecular interactions.

interaction is very small. However, the SOMO-1 orbital is symmetric with respect to the same plane (see Fig. 6) and the previous effect cannot explain the weakness of the SOMO-1···SOMO-1 interaction III, which can only be due to the poor orientation of the S p_z orbitals. Thus, both poor S orbital interactions and the topology of SOMO orbitals are responsible for very weak inter-chain interactions. In summary, $[Au(pzdtdt)_2]$ is predicted to be a strongly 1D system from the electronic viewpoint with SOMO and SOMO-1 bands of similar width.

The calculated DFT band structure when double occupation of the levels is assumed is shown in Fig. 6, where every band is really the superposition of two bands because the unit cell contains two practically non-interacting layers. The SOMO band is strongly 1D, whereas the SOMO-1 has a small but nonnegligible dispersion along the inter-chain direction (see lines Γ to Z and X to M in Fig. 6). As expected, the band dispersion along the inter-layer direction (Γ to Y) is nil. The two bands have a strong and similar dispersion and exhibit an important overlap. However, the SOMO-1 band does not cross the Fermi level so it is completely full and the SOMO band is half-filled. In other words, the possible two-band behaviour exhibited by other gold bis(dithiolene) solids does not develop at 150 K and ambient pressure for this solid and the SOMO---SOMO inter-

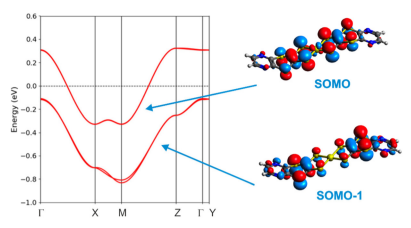


Fig. 6 Band structure calculated for the $[Au(pzdtdt)_2]^*$ solid, assuming the double occupation of levels. The dashed line refers to the Fermi level and $\Gamma = (0, 0, 0), X = (a*/2, 0, 0), Y = (0, b*/2, 0), Z = (0, 0, c*/2)$ and M = (a*/2, 0, c*/2).

actions should govern the transport and magnetic properties. The band structure in Fig. 6 is associated with a Fermi surface with a nesting vector having an $a^*/2$ component (see Fig. S6 in ESI†). Consequently, a dimerization along the chain direction with the consequent opening of a band gap and a semiconducting behaviour is predicted, in contradiction with the observed uniform stack structure. However, strongly 1D gold bis(dithiolene) with such half-filled uniform chains almost invariably exhibits electronic localization. It is thus compulsory to look at possible electronic states with electron localization (i.e. uniform chains of $[Au(pzdtdt)_2]$ radicals) before concluding anything about the nature of the ground state of this solid.

Initial evaluation of the magnetic coupling constants through dimer calculations showed that only the intra-layer ones corresponding to interactions I, II and II were larger than 1 K, with the first being almost two orders of magnitude larger. All inter-layer coupling constants were found to be as weak as 0.05 K so inter-layer interaction can be neglected. We have thus carried out spin-polarized calculations for four different localized states of the $[Au(pzdtdt)_2]$ layers with either ferromagnetic or antiferromagnetic interactions along (i) the intra-chain a direction, and (ii) the inter-chain c direction (see Fig. 7a).

Two of the localized states, AF12 and AF14, having antiferromagnetic interactions along the chains are more stable than the metallic state by 53.3 and 52.7 K, respectively. The two states with ferromagnetic interactions along the chains are found to be even less stable than the metallic state. The AF12 state is found to be the lowest energy state although the energy difference between the AF12 and AF14 states is very small (0.6 K) so we conclude that the ground state of this system should exhibit antiferromagnetic interactions along the chains but no magnetic order along the inter-chain and inter-layer directions. The calculated magnetic coupling constants are $J_{\rm I} = -427.5$ K, $J_{\rm II} = +16.7$ K and $J_{\rm III} = +24.2$ K, respectively (a negative sign means antiferromagnetic interactions). The calculated band structures for the AF12 and AF14 states are shown

in Fig. 7c and d, respectively. For comparison, the band structure of the metallic state calculated with the same cell is shown in Fig. 7b. The set of four bands noted with green asterisks in Fig. 6b is the folded version (because of the larger unit cell) of the SOMO bands in Fig. 6 which are globally halffilled. As shown in Fig. 7c and d, the development of antiferromagnetic spin interactions along the direction results with the opening of a large energy gap, which is the reason for the higher stability of the AF12 and AF14 states. The two upper doubly filled SOMO bands (green asterisks) in Fig. 7b are strongly destabilized around the X point of the Brillouin zone and become two SOMO spin up and two SOMO spin down bands of higher energy. The opposite occurs for the lower doubly filled SOMO bands (green asterisks) in Fig. 7b, which become two SOMO spin up and two SOMO spin down bands of lower energy in the region around Γ for the AF12 and AF14 states. Since these bands are filled, they provide a stabilization to the system. The opening of this band gap in the spin polarized band structure is the equivalent of the development of the well-known Peierls gap for a half-filled band in the metallic state when electronic repulsions are important. The small differences between Fig. 7c and d decode the different interchain magnetic interactions between antiferromagnetic chains. Since J_{II} and J_{III} are very similar, the differences are small and the two states are of similar energy, leading most likely to an absence of magnetic ordering along the inter-chain direction.

Let us now briefly consider the interesting evolution of conductivity under pressure illustrated above in Fig. 4. Pressure application will certainly increase the dispersion of the SOMO bands (both because of the induced shortening of intermolecular contacts and possible decrease of the non-planarity), thus stabilizing the metallic state and increasing the conductivity, as it is observed until around 4–5 GPa. At this point, an almost metallic behaviour is observed. However, a further increase in pressure still leads to activated conductivity and one can foresee three different scenarios behind this behaviour. First, a metallic state is stabilized and a typical Peierls

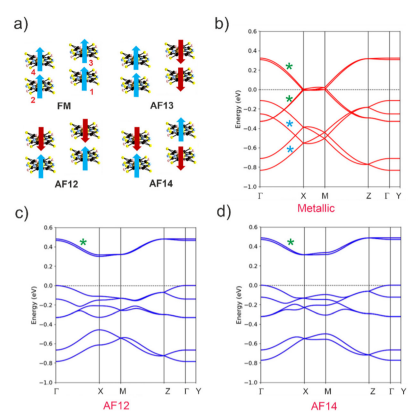


Fig. 7 (a) Four different localized states considered in this study. DFT band structure for the metallic (b) and antiferromagnetic AF12 (c) and AF14 (d) states calculated for a single-layer of $[Au(pzdtdt)_2]$ obtained using a 2 × 1 × 2 supercell and by deleting the four molecules of one of the layers. The dashed line refers to the highest occupied level, and $\Gamma = (0, 0, 0)$, X = (a*/2, 0, 0), Y = (0, b*/2, 0), Z = (0, 0, c*/2) and M = (a*/2, 0, c*/2). The bands in (b) are doubly occupied, and the two sets of four bands noted with green/blue asterisks originate from SOMO/SOMO-1 level. Spin-up and spindown bands in (c) and (d) are shown in blue and red, respectively. The spin-up and spin-down bands are identical. They are located in spatially different but equivalent sites, so only the blue bands are visible.

type instability for a half-filled strongly 1D metal develops, leading to a dimerization along the chains and the appearance of a band gap. Second, pressure leads to an increase of intrachain interactions but also to a change in the magnetic interchain interactions and one of the two becomes clearly dominant, leading to the more stable state at high pressure. As a result, a different magnetic state now with magnetic order also along the inter-chain direction develops. Third, since the Fermi level for the metallic state is not that far from the top of the SOMO-1 band in Fig. 2, the increase in band dispersion due to pressure may lead to the crossing of the top of the SOMO-1 band and the Fermi level, leading to a small electron transfer between the SOMO-1 and SOMO bands (i.e. a twoband behaviour) as it occurs for instance in [Au(Methiazdt)₂].^{4b} In that case, the occurrence of a metallic or semiconducting behaviour subtly depends on the degree of electron transfer. However, if this was the case, a further increase of pressure should stabilize the metallic state, which is apparently not the case. Consequently, one of the first two possibilities seems to be more appropriate. Structural work under pressure is needed in order to clarify this question. Indeed, the pressure dependence of the conductivity of [Au(pzdtdt)₂] at room temperature (Fig. 4b) shows a very abrupt evolution

between 0 and 3 GPa which might indicate a structural transition under pressure and a solid-state organization above 3-4 GPa, different from that determined here by X-ray diffraction at ambient pressure. Work is in progress to tentatively determine the crystal structure above 3-4 GPa.

Conclusions

To date, for dithiolene complexes formed by fusing of dithiine heterocycles, their crystallization into conducting materials has been hindered owing to strong distortions from planarity caused by the dithiine heterocycles. Herein, we reported an original gold bis(dithiolene) radical complex which stacked into regular non-dimerized chains, in the solid-state, with a semi-conducting behaviour at ambient pressure ($\sigma_{RT} = 2.04$ 10^{-3} S cm⁻¹, $E_{act} = 227$ meV). Notably, electron-localization effects of antiferromagnetic nature opened a Mott gap on the SOMO band. Remarkably, this single-component conductor turned metallic upon application of pressures above 4 GPa, which is a consequence of the associated increase in the SOMO band dispersion. A further up-turn to semiconducting behaviour at higher pressures could be the signature of a structural transition, leading to dimerization (Peierls transition) and an increase in inter-stack interactions, which were negligible in the ambient pressure conditions.

Experimental section

General

Chemicals and materials from commercial sources were used without further purification. All the reactions were performed under an argon atmosphere. NMR spectra were obtained in CDCl3 unless indicated otherwise. Chemical shifts are reported in ppm, ¹H NMR spectra were referenced to residual CHCl₃ (7.26 ppm) and ¹³C NMR spectra were referenced to CHCl₃ (77.2 ppm). Melting points were measured on a Kofler hot-stage apparatus and are uncorrected. Mass spectra were recorded by the Centre Régional de Mesures Physiques de l'Ouest, Rennes. Methanol, acetonitrile and dichloromethane were dried using an Inert pure solvent column device. Cyclic voltammetry was performed in a three-electrode cell equipped with a platinum disk working electrode and a glass carbon counter-electrode. CV was carried out in a 10⁻³ M solution of the complex in CH₂Cl₂ with 0.1 M [Bu₄N][PF₆]. Potentials were measured versus Saturated Calomel Electrode (SCE). The spectroelectrochemical study was performed in 0.2 M CH₂Cl₂-[NBu₄][PF₆] using a Pt grid as the working electrode, a Pt wire as the counter electrode and a SCE as the reference electrode. A Shimadzu 3600 Plus spectrophotometer was used to record the UV-vis-NIR spectra. The starting 3,4-diiodopyrazine was prepared according to a published procedure.¹⁹

[1,3]Dithiolo[4',5':5,6][1,4]dithiino[2,3-b]pyrazine-2-thione

(1). 3,4-Diiodopyrazine (0.58 g, 1.74 mmol), $(Et_4N)_2[Zn(dmit)_2]$ (0.62 g, 0.87 mmol), Cu₂O (0.025 mg, 0.174 mmol) and ethyl acetoacetate (41 µL, 0.348 mmol) were dissolved in 6 mL degassed DMF under an inert atmosphere. The suspension was heated at 80 °C for 15 h. After cooling, the solution was diluted in 100 mL of CH2Cl2 and washed three times with water. The organic phase was separated and dried with anhydrous Na2SO4. The crude product was purified by column chromatography on silica gel using CH2Cl2 as the eluent to afford 2 as a bright yellow solid in 63% yield. Mp = 190° C; ¹H NMR (CDCl₃, 500 MHz): 8.53 (2H); ¹³C NMR (75 MHz, CDCl₃) δ 206.89, 151.37, 142.43; HRMS (ASAP) calcd for [M + H]⁺: 274.88943, found: 274.8892. Anal. calcd for C₇H₂N₂S₅: C, 30.64; H, 0.73; N, 10.214. Found: C, 31.08; H, 1.06; N, 9.86.

[1,3]Dithiolo[4',5':5,6][1,4]dithiino[2,3-*b*]pyrazin-2-one Mercuric acetate (0.46 mg; 1.46 mmol) in 10 mL CH₃CO₂H was added to a solution of 1 (0.2 mg; 0.73 mmol) in 30 mL CHCl₃ and the suspension was stirred at room temperature for 2 h. Then the mixture was filtered through Celite® and washed with chloroform. The filtrate was washed with water several times and then with sodium bicarbonate. The organic phase was separated and dried with anhydrous MgSO₄. After removing the solvent, the product was obtained as an orange solid with a 94% yield. Mp = 189 °C; 1 H NMR (300 MHz, CDCl₃) δ 8.42 (s, 2H); 13 C NMR (75 MHz, CDCl₃) δ 190.30, 151.47,

142.60. HRMS (ASAP) Calcd for [M + H]+: 258.91227, found: 258.9126. Anal. calcd for C₇H₂N₂OS₄: C, 32.54; H, 0.78; N, 10.84. Found: C, 32.39; H, 1.00; N, 10.28.

[PPh₄][Au(pzdtdt)₂]. Under an inert atmosphere, a solution of MeONa (Na: 0.21 g, 9.28 mmol) in MeOH (15 mL) was added to the pyrazine-dithiin-dithiol-one (0.3 g, 1.16 mmol). After complete dissolution, the solution was stirred for 30 min at room temperature. Then a solution of KAuCl₄ (0.22 g, 0.58 mmol) in MeOH (5 mL) was added, followed 6 hours later by the addition of Ph₄PBr (0.24 g, 0.58 mmol). After stirring for 15 h, the formed brown precipitate was filtered and recrystallized from acetonitrile to afford the monoanionic complex as brown crystals, 40% yield. Mp > 250 °C. ¹H NMR (300 MHz, $CDCl_3$) δ 8.09 (d, J = 1.9 Hz, 4H), 7.95–7.50 (m, 20H), ¹³C NMR. Anal. calcd for C₃₆H₂₄N₄AuPS₈: C, 43.37; H, 2.43; N, 5.62; S, 25.72. Found: C, 42.72; H, 2.46; N, 5.64; S, 25.01. HRMS (ESI) calcd for C^+ ($C_{24}H_{20}P$): 339.12971, found: 339.1302.

[Au(pzdtdt)₂]. Crystals of [Au(pzdtdt)₂] were prepared electrochemically using a standard U-shaped cell equipped with Pt electrodes (1 cm length × 1 mm diameter). A THF solution (12 mL) of [Ph₄P][Au(pzdtdt)₂] (5 mg) and a supporting electrolyte [Bu₄][NPF₆] (300 mg) were placed in the U-shaped cell. The current intensity between the electrodes was adjusted to a constant value of 1 µA for 3 days. Black plate-like crystals suitable for X-ray diffraction studies were collected at the anode and washed with THF.

Crystallography

Suitable crystals for single crystal X-ray diffraction experiment were selected and mounted on the goniometer head of a D8 Venture (Bruker-AXS) diffractometer equipped with a CMOS-PHOTON70 detector, using Mo-K α radiation (λ = 0.71073 Å, multilayer monochromator). Crystal structures were solved by the dual-space algorithm using the SHELXT program,²⁹ and then refined with full-matrix least-squares methods based on F² (SHELXL program).³⁰ All non-hydrogen atoms were refined with anisotropic atomic displacement parameters. H atoms were finally included in their calculated positions and treated as riding on their parent atom with constrained thermal parameters. Crystallographic data on X-ray data collection and structure refinements are given in Table S1 in the ESI.†

Resistivity measurements

The standard four-probe DC measurements were performed for single crystal [Au(pzdtdt)₂]* by using a diamond anvil cell (DAC) using the same technique as described in ref. 31 A rectangular shape of single crystal sample 1 (0.14 \times 0.03 \times 0.01 mm^3) and sample 2 $(0.11 \times 0.03 \times 0.01 \text{ mm}^3)$ was used for ambient pressure and high-pressure measurements up to 6.2 GPa for sample 2 and 19.6 GPa for sample 3 (0.09 \times 0.03 \times 0.01 mm³). The four contacts were made by four gold wires (10 or 5 µm) with gold paint. The culet size of 0.7 mm (0.56 mm) diamond and tension-annealed stainless still SUS301 (Inconel 625) for sample 1, 2 (sample 3) and pressure-transmitting medium Daphne Oil 7373 was used. The pressure was deter**Paper**

mined by the shift of the ruby fluorescence R1 lines at room temperature. A Cryocooler helium compressor system (Sumitomo Heavy Industries, Ltd) was used for cooling the DAC with a cooling rate of 1.0 K min⁻¹. A KEITHLEY 224 programmable current source and 182 sensitive digital voltmeter were used for all measurements.

Computational details

The first-principles calculations for the solid were carried out using a numeric atomic orbital density functional theory (DFT) approach^{32,33} developed for efficient calculations in large systems and implemented in the SIESTA code. 34-37 We used the generalized gradient approximation (GGA) to DFT and, in particular, the functional of Perdew, Burke, and Ernzerhof.³⁸ To study the relative energies of states with localized electrons, spin polarized band calculations for appropriate supercells have been undertaken. All calculations included a Hubbard correction term $U_{\text{eff}} = U - J = 6.0$ eV for the S 3p states.³⁹ In previous work,⁴⁰ we have found that this U term on the chalcogen atoms is needed for appropriately describing the electronic structure of molecular conductors were accurate experimental information on the bandwidth and charge transfer is available. Only the valence electrons are considered in the calculation, with the core being replaced by norm-conserving scalar relativistic pseudopotentials41 factorized in the Kleinman-Bylander form.42 We have used a split-valence double- ζ basis set including polarization orbitals with an energy shift of 10 meV (ref. 43) for S, C, N and H atoms. For gold atoms we have used a split-valence basis set of double-ζ plus polarization quality, where the 5d electrons of Au were treated also as valence electrons. The basis functions used for Au have been optimized in order to reproduce the geometry and the bulk modulus for the ccp crystal structure of metallic gold.44 The energy cutoff of the real space integration mesh was 300 Ry. The Brillouin zone was sampled using grids⁴⁵ of $(10 \times 3 \times 10)$, $(20 \times 3 \times 20)$ and $(20 \times 20 \times 20)$ k-points. The crystal structure at 150 K was used for the computations.

The intermolecular interaction energies were evaluated by means of extended Hückel type46 calculations with a modified Wolfsberg-Helmholtz formula to calculate the non-diagonal $H_{\mu\nu}$ values.⁴⁷ All valence electrons were considered in the calculations and the basis set consisted of Slater-type orbitals of double- ζ quality for Au 5d and of single- ζ quality for C and N 2s and 2p, S 3s and 3p and H 1s. The ionization potentials, contraction coefficients and exponents were taken from previous works.4c,48

Data availability

The data supporting this article have been included as part of the ESI.† Crystallographic data have been deposited at the CCDC under CCDC 2422633-2422637 for compounds 1, 2, $[Et_4N][Ni(pzdtdt)_2]$, $[Au(pzdtdt)_2]$ and $(Ph_4P)[Au(pzdtdt)_2]$, respectively, and can be obtained directly from https://www. ccdc.cam.ac.uk/products/csd/request/request.php4.

Conflicts of interest

There are no conflicts to declare.

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