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Introduction

Metallophilic interactions are a unique variant of weak attractive forces in chemical bonding, acting between atoms with interatomic distance less than the sum of their van der Waals radii. These interactions lead to novel physicochemical properties in the systems. Typically, metal ions with closed-shell (d¹⁰ and s^2) or pseudo-closed shell (d^8) electronic configurations are known to exhibit metallophilic interactions.1-7 Owing to their unusual nature, which is distinct from the conventional concepts of chemical bonding, the past few decades have seen a growing interest in generating and understanding such metallophilic interactions, especially involving univalent coinage metal (Au(1), Ag(1) and Cu(1)) compounds.⁸⁻¹⁵ Initially, significant research focused on exploring aurophilic (Au(I)···Au(I)) interactions, followed by argentophilic $(Ag(I) \cdots Ag(I))$ interactions and cuprophilic $(Cu(1) \cdots Cu(1))$ interactions. Argentophilicity is known to impart various interesting properties in systems and has applications such as dynamic chirality inversion in metal organic supramolecular polymers, circularly polarized luminescence in alkynyl-stabilized clusters, capturing and

Ultralow thermal conductivity and thermallydeactivated electrical transport in a 1D silver array with alternating δ -bonds[†]

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We report the synthesis of a (TMA)AgBr₂ (TMA = tetramethylammonium) crystal, which comprises inorganic anionic chains of $-(AgBr_2)_{\alpha}$ - stabilized by columnar stacks of organic TMA cations with a periodic arrangement of shorter and longer Ag(i)···Ag(i) bonds, even though all the Ag(i) ions are chemically equivalent. The presence of two chemically non-equivalent bridging Br ions is attributed to the primary cause of such an unusual arrangement, as clearly visualized in the charge density plot of (TMA)AgBr₂ extracted from the theoretical calculations based on density functional theory. Remarkably, we identified from the orbital-projected density of states the existence of alternate δ -like bonding involving d_{xy} orbitals of 4d¹⁰ Ag(i), which was attributed to the cause for ultralow thermal conductivity and thermallydeactivated electrical transport in (TMA)AgBr₂. Barring the energetics, our observations on the existence of a δ -bond will shed new light in understanding the nature of metal–metal chemical bonding and its unprecedented implications.

> photocatalysis of mustard-gas simulant in a porphyrinic cluster, chemical fixation of carbon dioxide in a two-dimensional metal-organic framework, combating antibiotic resistance against Staphylococcus aureus, as bioprotective materials for combating superbacteria, and artificial light-harvesting.16-22 However, argentophilic interactions have so far been observed in discrete complexes, clusters, and supramolecular assemblies, except a report on an uninterrupted one-dimensional silver array stabilized by specific interactions with the base pairs of DNA.23 The implications arising from the long-range ordering of such weak forces of attraction can be intriguing; for example, the silver-DNA hybrid nanowire is expected to facilitate electrical conduction, similar to earlier observations of the metallophilic interactions in linear-chains imparting electrical conductivity. Notably, all these materials can be considered as semiconductors viz. they exhibit the thermally-activated electrical transport property. The question is: what would happen if the long-range metallophilic interactions are periodically interrupted? Herein, we synthesized an organic-inorganic hybrid crystal, (TMA) $AgBr_2$, (TMA = tetramethylammonium cation), exhibiting a one-dimensional array of Ag(1) ions bridged with Br ions, and we identified an alternating arrangement of the argentophilic interactions, which is unprecedented.

Results

Single crystals of (TMA)AgBr₂ were synthesized following a controlled cooling crystallization process. The crystal structure was obtained by a single-crystal X-ray diffraction (SCXRD)

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technique using an orthorhombic crystal system with the space group *Immm*, having the unit cell parameters as follows: a =6.7744(13) Å, b = 9.1274(19) Å, c = 15.003(3) Å, V = 927.675 Å³ (Tables S1–S3[†]). The crystal is made up of polymeric $-(AgBr_2)_n$ anionic chains stabilized by stacked TMA cations (Fig. 1a and S1a-c[†]). The $-(AgBr_2)_n$ - chains are isolated from each other, and a single chain is surrounded by six stacks of TMA cations. The Ag(1) ions in the chains are tetrahedrally coordinated to four Br⁻ ions, with each Br⁻ ion being connected to two neighboring Ag(1) cations (Fig. 1b and S2a-c[†]). Interestingly, every Ag(1)cation exhibits an additional $Ag(I) \cdots Ag(I)$ bond with one of its nearest neighbors, with a bond distance of 3.151 Å (well below the sum of their van der Waals radii) - indicating the presence of argentophilic interactions. In general, metallophilic interactions are considered to exist in a compound if the metal-metal separation is shorter than the sum of their van der Waals radii¹⁰ and it is not related to Shannon ionic or crystal radii,²⁴ which are usually considered for metal-halide bonds. The $Ag(I) \cdots Ag(I)$ distance involving the opposite nearest neighbor is measured to be 3.623 Å and consequently, no bonding was observed (viz. argentophilic interactions are absent). Therefore, a one-



Fig. 1 Crystal structure of (TMA)AgBr₂. (a) Extended arrangements of TMA cations and $(AgBr_2)_n^{n-}$ anionic chains viewed along the crystallographic *a*-axis (perspective view, H atoms are omitted for clarity). All the $(AgBr_2)_n^{n-}$ polymeric chains are segregated from one another, and each chain is surrounded by six stacks of TMA cations running along the *a*-axis. (b) Zoomed-in view of a single inorganic $(AgBr_2)_n^{n-}$ chain along the crystallographic *a*-axis, showing the alternate Ag···Ag (argentophilic) interactions.

dimensional silver array with alternating argentophilic interactions is clearly identified in (TMA)AgBr $_2$.

Such an alternating arrangement of Ag ions is completed with the presence of two Br⁻ ions, here assigned as Br1 and Br2. The Br1 ion is bridged to the $Ag(I) \cdots Ag(I)$ bonding pair with a distance of 2.692 Å and angle of 71.66°, while the Br2 ion is bridged to the $Ag(I) \cdots Ag(I)$ non-bonding pair with a distance of 2.738 Å and an angle of 82.86° (Fig. S2[†]). Another important consideration of the overall energetics that could influence the contribution of ionic interactions in the system is the closest distance between Br⁻ ions and the TMA cation: 5.039 Å for Br1 and 4.203 Å for Br2. The powder X-ray diffraction (PXRD) pattern matched very well with the simulated XRD pattern from SCXRD analysis, thereby confirming the phase-purity of our bulk sample (Fig. S3[†]). X-ray photoelectron spectroscopy (XPS) was employed to probe the oxidation state of Ag. The Ag 3d photoemission signal, with peaks for Ag 3d_{3/2} and Ag 3d_{5/2} (due to spin-orbit coupling) at respective binding energy values of 373.4 eV and 367.4 eV ensured the exclusive presence of Ag(1) in (TMA)AgBr₂ (Fig. S4[†]).^{25,26} Thermogravimetric analysis (TGA) revealed no thermal degradation up to 600 K, attributed to the excellent tolerance of (TMA)AgBr₂ to thermal energy (Fig. S5⁺).

Electronic structure calculations on the (TMA)AgBr₂ system were performed using density functional theory (DFT). A direct band gap (E_g) of 3.53 eV with the valence band maxima (VBM) as well as the conduction band minima (CBM) located at the gamma (G) point was identified (Fig. S6[†]). This theoretically estimated band gap (E_{α}) is in reasonable agreement with the experimentally obtained value of 4.1 eV (Fig. S7†). The orbital projected density of states (DOS) plots suggest that the primary contributions towards VBM and CBM originate from the Ag4d-Br4p and Ag5s-Br4p orbitals, respectively (Fig. S8⁺). Notably, the organic part, TMA, has negligible contribution towards VBM and CBM, further implying the importance of the inorganic $-(AgBr_2)_n$ - chain in predominantly shaping the electronic band structure. The partial charge density plots clearly depict periodic accumulation of electronic charge between the alternate Ag atoms, viz. a direct visualization of the alternate argentophilic interactions in the one-dimensional Ag chain (Fig. 2a). Critically looking at the orbital contributions in generating the 4d¹⁰... 4d¹⁰ Ag(1)...Ag(1) bond, as reflected in the orbital projected density of states, the overlap between the d_{xy} orbitals was predominantly identified to be the origin of the argentophilic interactions in (TMA)AgBr $_2$ - resembling a classical δ -like bonding scenario (Fig. 2b and c).27-29 Earlier, in view of the energetics,3 argentophilic interactions were primarily assigned to be weak non-covalent interactions, even weaker than ionic and dispersion interactions; however, our DFT calculations clearly reveal the involvement of δ -bonding in these argentophilic interactions. Being very much sensitive to distance, the presence or absence of δ -bonding can have a significant impact on the electronic structure,28 rendering such systems to behave as topological materials.

The unique electronic and crystal structure of (TMA)AgBr₂ inspired us to measure its thermal transport properties. The variable temperature thermal diffusivity (α) as well as the specific heat capacity (C_p) for the (TMA)AgBr₂ system were

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Fig. 2 Visualization of argentophilic interactions in (TMA)AgBr₂. (a) Charge density plot illustrating the distribution of electronic charge in the alternate Ag···Ag region depicting interrupted argentophilic interactions (electron accumulation is represented by yellow isosurfaces). (b) Possible orbital overlapping scenarios involved in the M···M σ -bond (top; $d_z^2 \cdots d_z^2$), π -bond (middle; $d_{yz/zx} \cdots d_{yz/zx}$) and δ -bond (bottom; $d_{xy} \cdots d_{xy}$). (c) Orbital-projected density of states (DOS) plots showing the orbital contributions in the Ag···Ag region.

collected in the temperature range \sim 300 K to \sim 380 K. The α was $\sim 1.83 \times 10^{-3} \text{ cm}^2 \text{ s}^{-1}$ at 307 K, which gradually decreased with increasing the temperature and reached a value of $\sim 1.57 \times 10^{-3}$ $\rm cm^2~s^{-1}$ at 375 K (Fig. 3a). $C_{\rm p}$ exhibited the opposite trend, with a lower value of \sim 1.68 J g⁻¹ K⁻¹ at 300 K and a higher value of ~2.05 J g⁻¹ K⁻¹ at 380 K (Fig. 3b). The thermal conductivity (κ) of the material was then calculated using the following equation: $\kappa = \alpha \rho C_p$, where ρ is the density.³⁰ Remarkably, an ultralow value of κ (~0.6 W m⁻¹ K⁻¹) was reproducibly achieved at near room-temperature for the (TMA)AgBr₂ system (Fig. 3c), resembling the benchmark values of all-inorganic halide perovskites CsPbI₃ (~0.45 W m⁻¹ K⁻¹), CsPbBr₃ (~0.42 W m⁻¹ K⁻¹) and $CsSnI_3$ (~0.38 W m⁻¹ K⁻¹).³¹ An intrinsically low thermal conductivity in CuBiI₄ (\sim 0.34–0.28 W m⁻¹ K⁻¹) was attributed to strong antibonding I(p)-Cu(d) states32 and likewise the primary involvement of Br(p)-Ag(d) states in the formation of the valence band and conduction band of (TMA)AgBr₂. Further, bonding heterogeneity in AgPbBiSe₃ (ref. 33) leading to an ultralow thermal conductivity resembles an alternating arrangement of δ -bond facilitating ultralow thermal conductivity in (TMA)AgBr₂. Additionally, the organic cations could induce structural modulations, further lowering the thermal conductivity.34

The electrical transport properties of the (TMA)AgBr₂ system were also evaluated upon recording the temperature-dependent current-voltage (I-V) profiles on the pelletized sample using our two-probe configuration¹³ (Fig. S9[†]). Upon gradually increasing the temperature from 298 K to 333 K, the current value gradually increased, which therefore contradicts the traditional conduction of electrons in semiconductors and suggests a thermallydeactivated electrical transport mechanism in the (TMA)AgBr₂ system, which is typically observed in metals, viz. metallic conduction (Fig. 4a).35 The electrical conductance values were extracted from the I-V profiles, and the conductance value decreased from $\sim 8 \times 10^{-10}$ S at 298 K to $\sim 5 \times 10^{-13}$ S at 333 K (Fig. S10a[†]). Such a remarkable change in the electrical conductance value (by ca. 10³ times) could be associated with a structural phase transition in the system, and in order to probe it, variable temperature SCXRD data and PXRD patterns were collected for the (TMA)AgBr₂ sample. Interestingly, the SCXRD data and the PXRD patterns did not show any appreciable change in the lattice parameters or crystal symmetry (Tables S4–S6[†] and Fig. 4b). This established the fact that the electrical transport behavior of (TMA)AgBr₂ is inherent in the solitary phase and that internal electronic and phononic changes might be responsible for the thermally-deactivated



Fig. 3 Possibility of electron-phonon coupling. (a) Thermal diffusivity of (TMA)AgBr₂ as a function of temperature. (b) Heat capacity (C_p) of (TMA) AgBr₂ as a function of temperature. (c) Variable temperature thermal conductivity of (TMA)AgBr₂ yielding an ultra-low value.



Fig. 4 Metallic conduction without structural change. (a) Variable-temperature current–voltage (I-V) diagram of (TMA)AgBr₂ in the temperature range from 298 K to 333 K, illustrating thermally deactivated charge transport. (b) Variable temperature PXRD patterns showing the retention of the structural phase over the temperature range.

charge transport phenomenon. *I–V* measurements were also carried out to estimate the room-temperature electrical conductivity of (TMA)AgBr₂, using a collinear four probe set-up, and the value was calculated to be \sim 5.90 × 10⁻⁶ S m⁻¹ (Fig. S10b†).

Unlike normal semiconductors, upon light irradiation (450 nm) (possibly due to trapped and/or defect state mediated absorption within the band gap, *cf.* Fig. S7†), the electrical conductance value of (TMA)AgBr₂ dropped by more than an order of magnitude (Fig. 5) – a phenomenon known as negative photoconductivity (NPC). Various explanations have been reported for NPC; to mention a few: the generation of secondary hot electrons, formation of scattering centers, excitation into deep defect levels, surface plasmon resonance, charge carrier trapping due to moisture adsorption, trion formation, the presence of DX (D: donor atom, X: unspecified lattice defect) like centers, light-activated trap states, and thermal broadening of electron distribution.³⁶ Particularly, in halide perovskites and



Fig. 5 Current–voltage (I-V) plots of (TMA)AgBr₂ in the dark and under the influence of 450 nm laser irradiation.

related systems, NPC was attributed to the creation of lightactivated metastable trap states³⁷ and the self-trapping of charge carriers.³⁸ Similar mechanisms could also be responsible for the negative photoconductivity in our (TMA)AgBr₂ system. Notably, in our earlier-reported (TMA)Ag₂Br₃ system,¹³ under identical conditions of light irradiation, the electrical conductivity value was observed to be unaltered, which clearly highlights the implications of a distinctive structure and chemical bonding scenarios, *viz.* the structure–property relationship.

To further emphasize the importance of δ -bonding, *i.e.*, argentophilic interactions behind the metal-like electrical conduction in the (TMA)AgBr2 system, we have collected variable temperature I-V profiles for commercially available AgBr using a conventional two-probe setup on a pelletized sample. The phase-purity of AgBr was confirmed by collecting the PXRD pattern, which matched well with the simulated pattern (ICSD 1627211), without the presence of any impurity peaks (Fig. S11[†]). Notably, in the cubic crystal structure of AgBr, the closest Ag…Ag distance is identified as 4.059 Å (Fig. S12a[†]), which is beyond the upper limit of argentophilic interactions. Upon increasing the temperature, the electrical conductance value of the AgBr sample gradually increased - a clear signature of the thermally-activated electrical transport typically valid for semiconductors (Fig. S12b[†]). The thermal conductivity value of AgBr was estimated to be \sim 1.34 W m⁻¹ K⁻¹ at near roomtemperature (Fig. S13a–c[†]). Such a significantly higher κ value of AgBr highlights the influence of Ag…Ag interaction in imparting a comparatively lower thermal conductivity value for the (TMA)AgBr₂ system. Thermal conductivity values of relevant systems, starting from the conventional all inorganic and inorganic-organic halide perovskites to more recently reported diverse Cu(1) and Ag(1) based compounds, are summarized in Table S7.†

Raman spectra of the (TMA)AgBr₂ sample were recorded at 298 K and 353 K to investigate the possible origin of the ultralow thermal conductivity as well as the thermally-deactivated charge transport; the spectral features were almost unaltered, expect in

two regions: 1350–1450 cm⁻¹ and 100–150 cm⁻¹ (Fig. S14†). The Raman peaks at 1370 cm⁻¹ and 1400 cm⁻¹ (which can be denoted as the $\nu_2^{CH}(a_1)$ -CH₃ symmetric deformation in the organic part of (TMA)AgBr₂)³⁹ were respectively blue-shifted to 1381 cm⁻¹ and 1411 cm⁻¹ upon increasing the temperature, demonstrating an alteration in the specific phonon mode as a function of temperature (Fig. S15b[†]). For the Raman peak at 130 cm^{-1} , upon elevating the temperature, there is a significant enhancement in the intensity (almost twice compared to that between the normalized spectra with respect to the highest intensity peak at 3020 cm^{-1}); this peak can be a characteristic of the $Ag(I) \cdots Ag(I)$ bond (Fig. S15a⁺). Typically, Raman intensity is directly related to the polarizability (and symmetry); so as to say, a higher spectral intensity reflects an increase in the bonding covalence of the system. Therefore, on the one hand, there is a possibility of an exotic electron-phonon coupling; on the other hand, upon increasing the temperature, an extended electron delocalization along the Ag chain is anticipated, which altogether led to the unusual metallic conduction in the (TMA) AgBr₂ system.

Conclusions

Looking at the trends of both electrical conductance (and so the electrical conductivity) and thermal conductivity values of (TMA)AgBr₂, the ratio of the thermal conductivity (κ) to the electrical conductivity (σ) would increase with increasing temperature. Such an important observation apparently validates the applicability of the Wiedemann–Franz law ($\kappa/\sigma \propto T$) to our (TMA)AgBr₂ system, viz. metal-like conduction; key to this observation was the presence of alternating argentophilic interactions in a one-dimensional array of the Ag(1) ions, which is unprecedented. Our study highlighting the concept of δ bonding within metallophilic interactions will stimulate experimental as well as theoretical investigations on the electron-phonon coupling in such unique Ag-based systems; further, it may open the door for development of organicinorganic hybrid materials with ultra-low thermal conductivity as well as elusive 'plasmonic salts'.

Data availability

All data supporting the findings of this study are available within this article and its ESI.[†] The X-ray crystallographic details for structures reported in this article have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition nos. CCDC2352448, CCDC2352449, and CCDC2352450 for (TMA)AgBr₂ at 150 K, 340 K and 298 K, respectively.

Author contributions

N. H. prepared the samples, carried out the experiments, and performed electrical transport measurements along with S. S. under the guidance of N. B.; S. N. performed DFT calculations under the supervision of K. T.; N. H. and S. S. wrote the first draft, which was finalized by N. B. with inputs from the coauthors; N. B. supervised the research project.

Conflicts of interest

The authors declare no competing financial interest.

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