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[Hg₄Te₈(Te₂)₄]⁸⁻: A Heavy Metal Porphyrinoid Embedded in a Lamellar StructureCarsten Donsbach,^[a] Kevin Reiter,^[b] Dage Sundholm,^[c] Florian Weigend,^[b] and Stefanie Dehnen*^[a]Dedicated to Professor Bernt Krebs on the occasion of his 80th birthday.

Abstract: By use of ionic liquids (C_nC₁Im)[BF₄] with long alkyl chains (n = 10, 12) at the ionothermal treatment of Na₂[HgTe₂], lamellar crystal structures were obtained, with molecular macrocyclic anions [Hg₈Te₁₆]⁸⁻ (**1**) representing the heaviest known topological relative of porphyrin. [Hg₈Te₁₆]⁸⁻ naturally differs from porphyrin by the absence of an electronic π-system, which prevents a “global” aromaticity. Quantum chemical studies instead reveal small ring currents in the pyrrole-type five-membered rings that indicate weak local (σ) aromaticity. Due to their layered nature, the compounds may be promising candidates for the generation of 2D chalcogenidometalate sheets.

Dimensional reduction of semiconductor materials is currently actively pursued to control and tune their electronic properties.^[1] In this context, post-synthetic methods beyond nano-particle formation, like stamping or exfoliation techniques, have picked up pace.^[2] Preconditions for the latter are twofold, (a) the presence of suitable semiconductor substructures and (b) relatively weak interactions in two dimensions within the parent compound. Both is fulfilled in layered elements like graphite or phosphorus,^[3,4] binary chalcogenides,^[5] or various inorganic-organic hybrid salts.^[6]

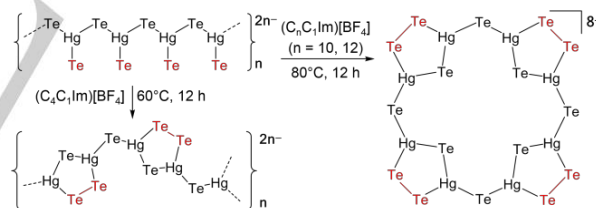
Semiconductor nano-structures are also found in heavy-metal chalcogenidometalates that exhibit a large variety of elemental combinations and architectures. High-temperature reactions usually lead to neat structures, whereas solution-based routes afford more open frameworks with solvated or organic cations. The anionic substructures may comprise lower dimensionalities, down to molecular chalcogenidometalate units.^[7]

An elegant way to transform semiconductor-based solids into hybrid compounds with weakly interacting organic cations is provided by ionothermal treatment of inorganic chalcogenidometalate salts.^[8] Resulting substructures can be as uncommon as the [Cu₅Ga₃₀S₅₂(SH)₂(C₄C₁im)₂]¹¹⁻ cluster (C₄C₁im = 1-butyl-3-methylimidazolyl),^[9] or the 0D-[Ge₂₄Sn₃₂Se₁₃₂]²⁴⁻

supersphere,^[10] the formation of which largely depends on the chosen temperatures, further auxiliaries, and the nature of the ionic liquid.^[11]

One parameter to vary is the chain lengths of the alkyl substituents at the (C_nC₁Im)⁺ cations in imidazolium-based ionic liquids. For n = 1-4, the cations behave similar to spherical ones, hence their structure-directing effect does not generally differ from the variation of ionic radii of atomic cations. By contrast, longer alkyl chains cause the ionic liquid cations to assemble, for instance into herring-bone or lamellar structures.^[12]

In the course of our studies regarding dimensional reduction of heavy-metal chalcogenidometalates, we are currently focusing on Hg/Te substructures formally derived from the important II-VI semiconductor material HgTe.^[13] By ionothermal treatment of Na₂[HgTe₂], comprising a one-dimensional [-Te-Hg(Te)-Te-] substructure, in the short-chain ionic liquid (C₄C₁Im)[BF₄] (C₄C₁Im = 1-butyl-3-methylimidazolium) at 60°C, a salt comprising another one-dimensional strand was obtained upon partial Te oxidation (Scheme 1, left).^[14] However, the use of 1-(do)decyl-3-methylimidazolium ionic liquids (C_nC₁Im)[BF₄] (n = 10, 12) affords salts of the unprecedented molecular anion [Hg₈Te₁₆]⁸⁻ (**1**; Scheme 1, right) within a lamellar structure.



Scheme 1. Diagram of the strand-like anion in Na₂[HgTe₂] (top left), and its treatment in ionic liquids (C_nC₁Im)[BF₄], yielding another strand-like anion in the presence of short-chain imidazolium cations (n = 4, bottom left),^[14] or the molecular anion [Hg₈Te₁₆]⁸⁻ (**1**) in ionic liquids with long-chain imidazolium cations (n = 10, 12; this work).

(C₁₀C₁Im)₈ crystallizes as thin, dark red plates in the monoclinic space group *P*2₁/*c*, with four formula units per unit cell. The inorganic macrocycle in **1** is built by four [Hg₂Te₃] five-membered rings comprising Te²⁻ and (Te₂)²⁻ ligands, which are bridged by four Te²⁻ ligands to form a cyclic tetramer. Hence, the anion **1** exhibits a close structural relation to organic porphyrins, which are composed of four methylene-bridged pyrrole rings, although the total valence electron count differs (120 in **1** versus 114 in porphyrin). **1** is thus related to the isoelectronic [B₈E₁₆] structures of BE₂ (E = S, Se),^[15] yet with the difference of being charged and with notable structural differences. Figure 1 illustrates the crystal structure of (C₁₀C₁Im)₈**1**.

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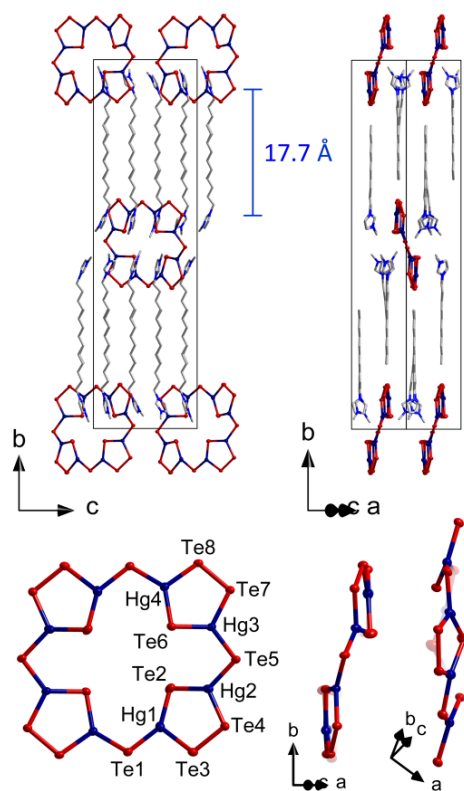


Figure 1. Top: Cut-out of the crystal structure of $(C_{10}C_1Im)_3 1$ in different orientations. Bottom: molecular structure of the anion $[Hg_8Te_{16}]^{3-}$ (**1**) in different views (Hg, Te: thermal ellipsoids at the 50% probability level. C, N: wireframes. H atoms omitted for clarity). External diameters of the anion range from 12.3 Å (Te1...Te1') to 16.2 Å (Te7...Te7').

Whereas Te-bridged $[Hg_2Te(Te_2)]$ motifs have been aligned in one-dimensional chains, as found in the $(C_4C_1Im)^+$ salt mentioned above¹⁴ and salts with $[N(C_2H_5)_4]^+$, $\{[Mn(en)_3]_2Cl_2\}^{2+}$, or $[M(trien)(N_2H_4)]^{2+}$ ($M = Mn, Zn$) cations,^[16] a cyclic oligomer as found in **1** has not been known yet. The bond lengths in **1** [$Hg-(\mu-Te)$ 2.692(2)–2.710(1), $Hg-(\mu:\eta^1:\eta^1-Te_2)$ 2.762(1)–2.803(1), $Te-Te$ 2.728(2)–2.750(1)] are in good agreement with those of the related 1D- $\{[Hg_2Te_4]^{2-}\}$ anions [$Hg-(\mu-Te)$ 2.654(2)–2.729(5) Å; $Hg-(\mu:\eta^1:\eta^1-Te_2)$ 2.750(2)–2.807(2) Å; $Te-Te$ 2.736(2)–2.788(3) Å]. All metal atoms possess a roughly trigonal planar coordination by three Te atoms (angle sums of $\geq 359.35(2)^\circ$), but the individual $Te-Hg-Te$ angles cover a wide range ($102.73(4)^\circ$ – $142.39(5)^\circ$), with somewhat more obtuse $(\mu-Te)-Hg-(\mu-Te)$ angles [$116.84(7)^\circ$ – $137.12(6)^\circ$ in the 1D strand] and slightly more acute $(\mu-Te)-Hg-(\mu:\eta^1:\eta^1-Te_2)$ angles [$105.45(6)^\circ$ – $127.73(7)^\circ$ in the 1D strand]. This indicates a significant coordinative flexibility around the Hg atoms that reacts on the respective arrangement of the counterions that vary in size, structure, charge, polarizability and hydrogen-bonding properties.

As a consequence of the coordinative flexibility of the Hg atoms, the macrocyclic anion **1** as a whole is not planar (Figure 1, bottom right). In order to optimize the interaction and structural match with the imidazolium cations, opposite pairs of $[Hg_2Te_3]$ rings are inclined against each other by about $24.8(1)^\circ$. This way,

the endocyclic Te atoms are relatively far apart from each other ($Te\dots Te$ 7.0808(7)–7.4130(6) Å across the center of **1**). We note that the five-membered rings in the mentioned 1D- $\{[Hg_2Te_4]^{2-}\}$ anionic chains are also not co-planar. Angles between mean planes ($1.2(1)$ – $35.0(6)^\circ$) span an even larger range, while in **1** this is naturally limited by its macrocyclic structure.

At the formation of **1**, the anionic substructure of the starting material is re-organized, accompanied by (partial) oxidative coupling of the telluride ligands (dichalcogenide anions are relatively readily formed in ionic liquids, which are less “innocent” in this regard than originally anticipated)^[8,14]. In the telluridomercurate anions of the starting material, $[HgTe_3]$ triangles are linked by edge-sharing into one-dimensional $[-Te-Hg(Te)-Te-]$ chains with all of the terminal Te atoms pointing towards the same side of the chain (Scheme 1). Thus, generation of $Te-Te$ bonds to form the $[Hg_2Te(Te_2)]$ units may, in theory, occur without deconstruction of the chain. However, for the assembly of the oligomeric structure of **1**, at least every tenth $Hg-Te$ bond needs to be cut, and the five-membered rings need to rearrange by rotation about $Hg-Te$ bonds (Scheme S1). We ascribe the preference of $[Hg_8Te_{16}]^{3-}$ macrocycles over chain-like 1D- $\{[Hg_2Te_4]^{2-}\}$ anions to the perfect fit of the molecular anions within the cationic template.

A closer look at the crystal structure of $(C_{10}C_1Im)_3 1$ reveals segregation into ionic and nonpolar parts. The latter result from *van-der-Waals* interactions of the decyl chains that form a membrane-like interlocked double-layer with a width of 17.7 Å (Figure 1, top). The positively charged imidazolium rings of the cations perfectly accommodate the anions as follows: two imidazolium rings are located in close proximity to one of the $[Hg_2Te(Te_2)]$ rings each, four from the cationic layer above and four from the one below, and thus compensate for the anions' relatively high charge within the ionic layers. In this arrangement, the imidazolium rings are arranged orthogonal to the $[Hg_2Te(Te_2)]$ units, thereby maximizing the anion...anion distance. The second compound obtained with longer alkyl chains, $(C_{12}C_1Im)_3 1$, is isostructural to $(C_{10}C_1Im)_3 1$. However, due to even thicker non-polar layers within the solid state (20.2 Å, Figure S4), it forms very thin and fragile crystals of relatively poor crystallographic quality (see the Supporting Information). The habitus of the plates points towards the potential of exfoliation, which is currently under exploration.

Beside segregation of long alkyl chains of such cations into polar/non-polar sections in the solid state,^[17] similar effects of nanoscale segregation into micro phases of homogeneous polarity was also observed in the liquid state of ionic liquids and confirmed by means of molecular dynamic simulations.^[18,19]

Aside from the idea of dimensional reduction of semiconductors, the formation of heavy homologs of well-known molecules is a fascinating area of chemical research, informing about the correlation of intrinsic atomic properties, and the structural, chemical, and physical properties of molecules. Prominent examples include the series of ethane, ethene, or ethyne homologs formed by heavier main group atoms, with the typical impact on their molecular structures.^[20] Another recent example is PbSe acting as a CO-like bridging ligand in $\{[(Ph_3P)_2Rh][[(Ph_3P)(CN)Rh]_2Se_2(\mu-PbSe)]\}^{3-}$.^[21] Particularly worth noting are topological mimics of organic compounds by purely inorganic homologs – both with isoelectronic situations as well as

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under toleration of other electron counts. Molecules like the benzene analog borazine, $B_3N_3H_6$,^[22] or the P_5^- anion as an isoelectronic equivalent of cyclopentadienide, Cp^- ,^[23] have been well-known since decades, while more complex structures have remained rare. Besides the above-mentioned porphyrin-like structures of BE_2 ($E = S, Se$), an inorganic double-helix mimicking DNA-like features was recently reported with the ternary, semi-conducting phase $SnIP$.^[24]

Quantum chemical studies were applied to explore electronic similarities and dissimilarities of this porphyrin-related molecule to porphyrin itself (TURBOMOLE,^[25] TPSSh functional,^[26] def2-TZVP bases,^[27] charge compensation with COSMO^[28] employing default parameters). Ring currents were obtained from GIMIC,^[29] based on the magnetic response calculated with a local version^[30,31] of TURBOMOLE. Currents for the two systems are shown in Figure 2.

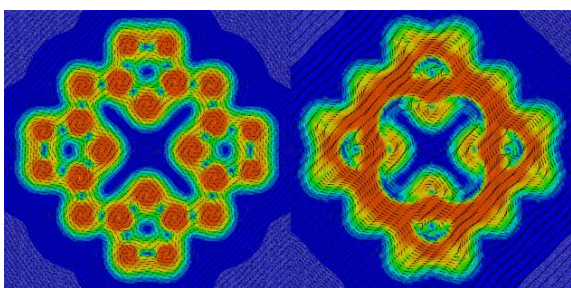


Figure 2. Ring currents, 1 bohr above the molecular plane of the inorganic anion **1** (left) and of organic porphyrin (right), drawn between 0 a.u. (blue) and 0.07 a.u. (red).

1 exhibits (weak) ring currents in the pyrrole-type five-membered rings. These currents consist of diatropic (clockwise, outside the ring) and paratropic (counterclockwise, inside) contributions of similar size but with a small surplus of the former leading to net currents of +5.8 nA/T. This is about half as much as for benzene,³² but – in contrast to the latter – they arise from sigma contributions only. The global net ring current (also consisting of diatropic and paratropic contributions) is essentially zero (0.24 nA/T). Organic porphyrines, in contrast, exhibit a global ring current of ca. 27 nA/T, which in the five-rings is split into two sub-currents, each amounting to ca. 13 nA/T.^[32]

These differences correspond to very different electronic structures for the two systems (despite identical topology). Aromaticity and according currents in porphyrin is based on the delocalized π -electrons. Also in **1**, there are delocalized canonical MOs, simply for symmetry reasons. The corresponding cluster orbitals (s-, p-, d-, f-, g-type) are the highest-energy MOs representing σ -bonding (Figure S7). However, in contrast to porphyrin, these MOs can be combined into two-center-two-electron bonds and lone pairs by application of a localization procedure.^[33] This yields two lone pairs per Te atom, and single σ -type bonds between each pair of neighbored atoms (in total, 32 lone pairs and 28 bonds).

Optical absorption measurements were recorded on a single-crystal of $(C_{10}C_{11}Im)_3\mathbf{1}$ (Figure 3). The onset of absorption is found at 2.2–2.8 eV, with a small shoulder of lower intensity at 2.1 eV.

This in excellent agreement with molecular TD-DFT calculations, with the lowest energy electronic transitions at 2.18 eV (HOMO to LUMO), followed by two more intense transitions at 2.48 eV (HOMO-4 to LUMO+3 and LUMO+4). During excitation, the electron density is essentially shifted from the μ -Te lone pairs to antibonding Te–Te MOs.

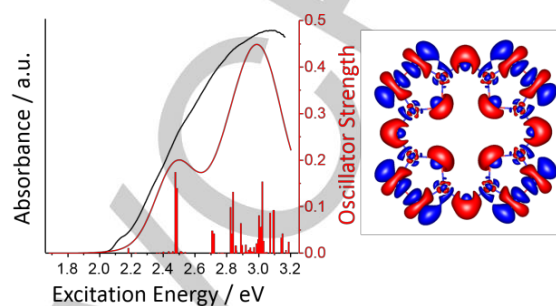


Figure 3. Comparison of the optical absorption spectrum measured on single crystals of $(C_{10}C_{11}Im)_3\mathbf{1}$ (black line) with calculated singlet excitation energies and oscillator strengths (lowest 250 excitations), plotted as vertical red lines with superimposed Gaussians of $fwhm = 0.3$ eV (red curve) to simulate the spectrum. The character of the entire band (up to 3.0 eV) is visualized using the nonrelaxed difference densities as described previously.^[34] The contributions of the occupied orbitals are plotted in red, those of the unoccupied orbitals in blue.

Due to the dimensional reduction, the optical gap is naturally blue-shifted from solid HgTe (-0.26 eV) and 1D- $[Hg_2Te_4]^{2-}$ (1.63 eV) to 0D- $[Hg_8Te_{16}]^{8-}$ (2.16 eV). It is still relatively narrow in the light of the given molecular size (24 atoms), as a consequence of the mixed-valence $Te^{2-}/(Te_2)^{2-}$ situation. As the absorption energy is likely to change upon complexation of transition metal atoms, we analyzed the possibility of using **1** as ligand. The lack of π -electrons in $[Hg_8Te_{16}]^{8-}$ comes along with a high structural flexibility (see above). As modelled by quantum chemistry, the macrocycle may accommodate metal ions of different sizes and coordination demands (e.g., Ti^{4+} , Cu^+ , Ce^{4+}) by adopting other shapes of the ring, which indeed show a different absorption behavior (Figure S8). It may be possible this way to form ternary complexes and clusters with tunable band gaps within lamellar arrangements. Yet, as the bonding energies are naturally lower than for corresponding porphyrin complexes the experimental isolation of according complexes remains a challenge.

In summary, we pursued a new approach for dimensional reduction of Hg/Te motifs by ionothermal treatment of $Na_2[HgTe_2]$ in long-chain alkyl imidazolium-based ionic liquids. The obtained substructure represents purely inorganic macrocycles, $[Hg_8Te_{16}]^{8-}$ with porphyrin-related topology, that assemble in lamellar crystal structures. The anion notably derives from planarity in order to optimize its situation in between the imidazolium units of the lamellar counterion structure. As confirmed by quantum chemical studies, the lack of π -electrons prevents significant aromaticity but leads to high structural flexibility, which may allow the coordination of metal cations despite the macrocycle's size by adopting its structure to the requirement of the cation. This way, lamellar arrangements of all-heavy-element complexes with tunable electronic properties may be accessible.

Experimental Section

Details on experimental procedures (syntheses, single-crystal X-ray diffraction including CIF, spectroscopy, and quantum chemical studies) are available as Supporting Information free of charge on the Wiley-VCH Publications website.

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Keywords: Ionothermal synthesis • Chalcogenidometalates • Inorganic Macrocycles • Dimensional Reduction • Ring Currents

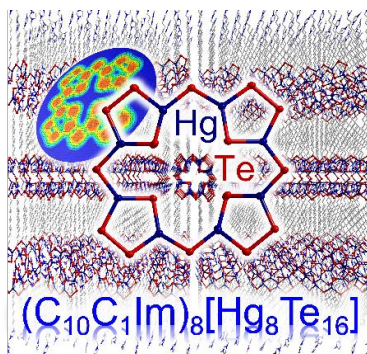
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lonothermal treatment of $\text{Na}_2[\text{HgTe}_2]$ with $(\text{C}_n\text{C}_1\text{Im})[\text{BF}_4]$ ($n = 10, 12$) affords lamellar crystal structures embedding macrocyclic anions $[\text{Hg}_8\text{Te}_{16}]^{8-}$ (**1**), the heaviest known topological relative of porphyrin. In contrast to the latter, **1** does not exhibit a π -electron system, in accordance with its high structural flexibility, but possesses small ring currents in the pyrrole-type five-membered rings that indicate weak local (σ) aromaticity.



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$[\text{Hg}_4\text{Te}_8(\text{Te}_2)_4]^{8-}$: A Heavy Metal Porphyrinoid Embedded in a Lamellar Structure

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