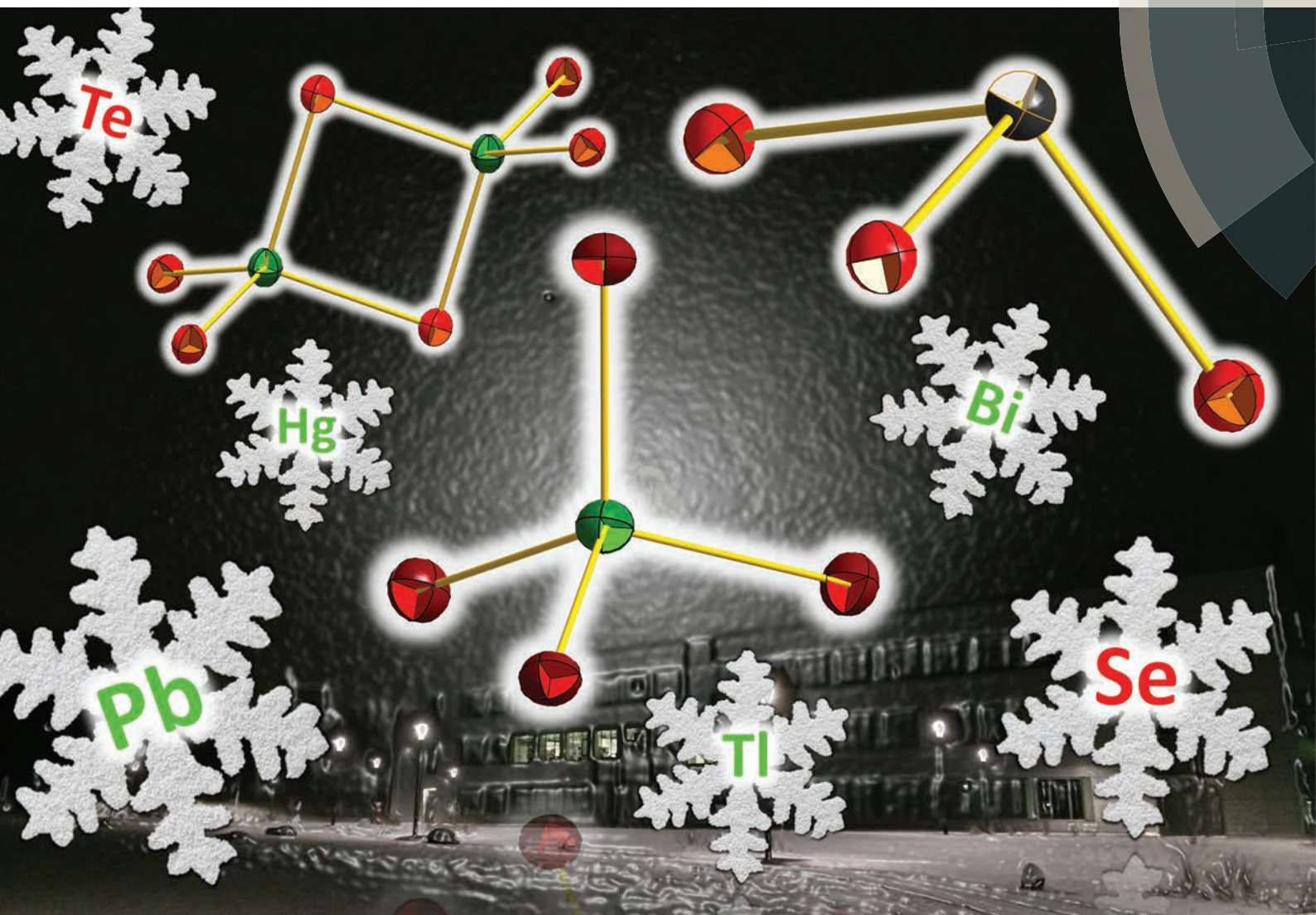


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Smallest molecular chalcogenidometalate anions of the heaviest metals: syntheses, structures, and their interconversion†

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The syntheses of the first molecular *meta*-selenidomercurate(II), *ortho*-telluridothallate(III) and a hydrate of an *ortho*-selenidoplumbate(IV) are presented alongside an improved and facile synthesis of the selenidobismuthate(III) with almost quantitative yields. By means of quantum chemical calculations, the energetics of the interconversions of small metalate anions is discussed and the existence of the heaviest homologues of $[\text{NO}_2]^-$, $[\text{NO}_3]^-$, $[\text{PO}_4]^{2-}$ and $[\text{CO}_3]^{2-}$ are predicted.

1. Introduction

Chalcogenides of the heavy metals possess intriguing optoelectronic¹ and thermo-electric² properties and have therefore been used in numerous applications.³ Those properties can be fine-tuned and adjusted *via* dimensional reduction,⁴ the choice of elemental combination,⁵ or further incorporation of additional metal species towards multinary compounds.⁶ Conventional solid state syntheses, solvothermal reaction pathways and solution-based methods have been shown to facilitate the generation of a large variety of such compounds, including chalcogenidometalate salts.⁷

Whereas the class of oxidometalates has extensively been studied – regarding both their fundamental properties and their potential applications – only a few reports have been dealing with the elemental combination of atoms of the heaviest metals with heavier chalcogen ligands. Here, we will concentrate on salts of the smallest of the heaviest chalcogenidometalate anions, hence with the general formulae $[\text{M}_x\text{Ch}_y]^{z-}$ ($\text{M} = \text{Hg}, \text{Tl}, \text{Pb}, \text{Bi}$; $x = 1$ or 2 ; $\text{Ch} = \text{Se}$ or Te , $y = 2-4$), of which only the following are known to date. Besides the mononuclear *ortho*-chalcogenidomercurate anion $[\text{HgCh}_4]^{6-}$, received from solid state reactions,⁸ only the linear $[\text{HgCh}_2]^{2-}$ anion, synthesized in solution,⁹ has been reported thus far. Molecular chalcogenidothallates and chalcogenidoplumbates have only been obtained from solution. Molecular chalcogenidothallates have so far been restricted to Schrobil-

gen's dinuclear $[\text{Tl}_2\text{Te}_2]^{2-}$.¹⁰ The largest number of small anions is currently found among the chalcogenidoplumbates. Here, one knows the *ortho*-selenidoplumbate anion $[\text{PbSe}_4]^{4-}$,¹¹ the dinuclear, trigonal bipyramidal $[\text{Pb}_2\text{Ch}_3]^{2-}$ anion,¹² and the mononuclear trigonal pyramidal species $[\text{PbTe}_3]^{4-}$.¹³ For bismuthates, only the trigonal pyramidal $[\text{BiCh}_3]^{3-}$ is known from solid state¹⁴ and solution-based approaches.¹⁵

Based on a broad experience in chalcogenidogermanate and chalcogenidostannate chemistry,¹⁶ we are currently about to extend the investigations on the heaviest tetrel congener as well as related anions of the neighbouring heavy metal elements. For this, we are addressing the synthesis of new small and soluble metalate anions, besides the exploration of their physical properties as well as their reactivity in solution or under solvothermal conditions. During the course of our investigations, we were able to synthesize a number of according compounds comprising new such anions with unprecedented elemental combinations, and were thus interested in general tendencies of the energetics of such species and their hypothetical interconversion. Here we report about our findings for the first time.

2. Experimental and computational details

2.1 General remarks and synthesis details

All manipulations and reactions were performed in an argon atmosphere using standard Schlenk or glovebox techniques under strict exclusion of air and external moisture. Multinary solid mixtures of the nominal parent composition “ $\text{A}_m\text{A}'_n\text{M}_o\text{M}'_p\text{Ch}_q$ ” (A, K ; $\text{A}' = \text{Na}$; $\text{M} = \text{Hg}, \text{Tl}, \text{Bi}$; $\text{M}' = \text{Pb}$; $\text{Ch} = \text{Se}, \text{Te}$) were prepared by fusion of the elements and K_2Ch ,

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unless stated otherwise, with an oxygen/methane burner for at least 15 min until homogeneity was achieved. The $m:n:o:p:q$ ratios, which are specified in the Discussion section, are those that actually produced the named products, usually after testing a variety of different ratios. If necessary, metallic by-products were manually removed. The multinary product was pestled and further worked-up as described below. Solvents ethane-1,2-diamine (en), *N,N*-dimethylformamide (DMF) and pyridine were freshly distilled from CaH_2 , H_2O was degassed at least three times to a final pressure of $p < 10^{-5}$ mbar and successively saturated with argon atmosphere. K_2Ch was synthesized from the elements in liquid NH_3 . Metal salts were evacuated to $p < 10^{-3}$ mbar for at least 8 h unless otherwise noted.

In the case of solvothermal reaction conditions, the reactants and 2 mL of a chosen solvent were placed in a glass vial within a Teflon inlay which was then placed in a stainless steel autoclave. The autoclave has been tightly closed and heated to 150 °C ($\text{Ch} = \text{Se}$) or to 130 °C ($\text{Ch} = \text{Te}$) for 7 days and was allowed to slowly cool down to room temperature over 1 day. The products were hand-selected under a light microscope unless stated otherwise. The product mixture for the synthesis of thallate and plumbate compounds always contained amorphous (side-)products that were not further investigated, and K_2Ch_2 and K_2Ch_3 that were identified *via* single crystal diffraction. Consequently, the provision of yields is afflicted with high uncertainty. Energy dispersive X-ray spectroscopy was performed on all samples to verify the given heavy atom composition. All yields are given with respect to the heaviest element involved in the respective case.

Synthesis of $\text{K}_4[\text{HgSe}_3]\cdot\text{H}_2\text{O}$ (1). 0.8 g (5.1 mmol, 3 eq.) of K_2Se , 0.5 g (1.7 mmol, 1 eq.) of HgSO_4 and 2 mL of en were treated according to the solvothermal reaction protocol. **1** was obtained as yellow blocks in approx. 80% yield beside K_2SO_4 .

Synthesis of $\text{K}_4[\text{HgTe}_3]\cdot\text{H}_2\text{O}$ (2). 2.10 g (10.2 mmol, 3 eq.) of K_2Te , 1 g (3.4 mmol, 1 eq.) of HgSO_4 and 2 mL of en were treated according to the solvothermal reaction protocol. **2** was obtained as metallic black blocks in approx. 20% yield beside K_2SO_4 and polytellurides.

Synthesis of $\text{K}_{10}\text{Na}_2[\text{HgTe}_3]_2(\text{HCO}_2)_3$ (3). 1 g of a phase of the nominal composition $\text{K}_3\text{NaHgTe}_3$ and 2 mL of DMF were treated according to the solvothermal reaction protocol. A few crystals of **3** were obtained as metallic black blocks.

Synthesis of $\text{K}_2[\text{HgSe}_2]\cdot\text{H}_2\text{O}$ (4). 0.5 g of a phase of the nominal composition $\text{K}_2\text{HgPbSe}_2$ and 2 mL of pyridine with 0.5 vol% of H_2O were treated according to the solvothermal reaction protocol. **4** was obtained as orange blocks in approx. 30% yield.

Synthesis of $\text{K}_6[\text{Tl}_2\text{Se}_6]\cdot 2\text{H}_2\text{O}$ (5). 0.5 g of a phase of the nominal composition K_5TlSe_4 and 2 mL of en with 0.5 vol% of H_2O were treated according to the solvothermal reaction protocol. **5** was obtained as red plates in approx. 15% yield.

Synthesis of $\text{K}_6[\text{Tl}_2\text{Se}_6]\cdot 2\text{K}_2\text{CO}_3$ (6). 0.5 g of a phase of the nominal composition K_5TlSe_4 , 0.05 g of K_2CO_3 and 2 mL of en were treated according to the solvothermal reaction protocol. A few crystals of **6** were obtained as orange blocks.

Synthesis of $\text{K}_{0.94}\text{Tl}_{0.06}[\text{TlSe}_2]$ (7). 0.5 g of a phase of the nominal composition K_2TlSe_2 and 2 mL of en were treated according to the solvothermal reaction protocol. **7** crystallizes as black blocks beside $\text{Tl}[\text{TlSe}_2]$, K_2Se_2 and K_2Se_3 .

Synthesis of $\{\text{K}_5[\text{TlTe}_4]\cdot 2\text{KOH}\}_3\cdot 10\text{H}_2\text{O}$ (8). 1 g of a phase of the nominal composition KTlTe and 2 mL of en with 3 vol% of H_2O were treated according to the solvothermal reaction protocol. **8** was obtained as black blocks in approx. 15% yield beside several known polytelluride salts.

Synthesis of $\text{K}_4[\text{PbSe}_4]\cdot\text{en}\cdot\text{H}_2\text{O}$ (9). 0.5 g of a phase of the nominal composition $\text{K}_2\text{HgPbSe}_2$ and 2 mL of en with 0.5 vol% of H_2O were treated according to the solvothermal reaction protocol. **9** was obtained as red blocks in approx. 20% yield.

Synthesis of K_3BiSe_3 (10). A phase of the nominal composition K_3BiSe_3 was washed with en until no coloured contaminants were observable any more. **10** was obtained in up to 90% yield as orange microcrystalline powder. Crystals suitable for single crystal diffraction were obtained *via* recrystallization according to the solvothermal reaction protocol in 2 mL of en.

2.2 General computational details

All molecular electronic structure calculations were carried out with the TURBOMOLE program package V6.6.¹⁷ COSMO was used for the compensation of negative charges, applying the default parameters with $\epsilon = \infty$.¹⁸ Structure optimizations were undertaken by employment of the RIDFT program, using the BP86 functional¹⁹ and def2-TZVP basis sets with respective ECPs.²⁰ The optimized structures were confirmed to be local minima on the energy hypersurface by means of calculations of the 2nd derivative of the energy. Further details are given in the ESI.†

3. Results and discussion

Our primary objective was the synthesis and characterization of small and heavy chalcogenidometalate homologues of their lighter congeners, *e.g.* stannates or indates.²¹ We therefore systematically investigated solvothermal reactions of phases of the nominal composition “ $\text{A}_x\text{M}_y\text{Ch}_z$ ” (A = alkali(ne earth) metal, $\text{M} = \text{Hg}, \text{Tl}, \text{Pb}, \text{Bi}$; $\text{Ch} = \text{Se}, \text{Te}$), in most cases with $x:y:z$ such that the ratio corresponds with a charge balance for *ortho*-metalates ($x:1:4$) or *meta*-metalates ($x:1:3$). We started out with selenium as the chalcogen component, since according phases are generally thermally more stable and less sensitive. Upon identification of suitable reaction conditions for the selenium containing compounds, analogous reactions were attempted with tellurium phases. Due to the lower thermal stability of the latter, the reaction temperatures were then decrease by at least 20 °C, and the amount of starting materials was increased by a factor of two or more due to higher solubility.

The phases were then treated under solvothermal reaction conditions with diverse solvents, such as amines, alcohols, thiols and ethers. Until now, only the application of amines yielded products that were suitable for single crystal diffraction



experiments, apart from the formation of a vast variety of single-crystalline polyselenides from other solvents.^{22,23}

We also tested whether water, amines like NH₃, 2,6-dimethylmorpholine (dmmp) or 4-(dimethylamino)pyridine (dmap), ammonium or phosphonium salts (NMe₄Cl, NBu₄Cl, PPh₄Cl), or Brønstedt bases (AOH, A₂CO₃) might serve as auxiliary reactants for enhanced hydrogen bonding, and/or to change the basicity. However, only addition of water, or (in reactions of thallium compounds) addition of Brønstedt bases led to the formation of crystalline products (see below).

Upon solvothermal treatment of phases that contained mercury as K_xHg_ySe_z (with all combinations of $x = 2-6$, $y = 1-10$, $z = 1-12$), we exclusively observed the formation of known compounds K₂Hg₃Se₄,²⁴ K₂Hg₂Se₃,²⁵ and polyselenides.²² We thus changed the mixture of starting materials to potassium monochalcogenides, K₂Ch, and mercury salts. This way, we crystallized K₄[HgSe₃]·H₂O (1) or K₄[HgTe₃]·H₂O (2) upon utilization of K₂Ch and HgSO₄·*n*H₂O (0.1 – *n* – 2.7 according to Karl Fischer titrations) in en as a solvent. The reaction towards the molecular *meta*-mercurate 1 is straight forward, according to a reaction scheme given in eqn (1):



The precipitation of K₂SO₄ seems to be the driving force, leading to an almost quantitative yield. Furthermore, the reaction can be scaled-up to 25 g per batch, only restricted by the volume of the glass vial within the autoclave. Solvent water in 1 stem from the crystal water in HgSO₄·*n*H₂O. The use of dried HgSO₄ (*n* = 0) results in the precipitation of a yellow amorphous powder, which can, however, be recrystallized in the presence of water to yield 1. The amount of crystal water in HgSO₄·*n*H₂O varies with the time of storage from newly purchased (*n* ≈ 0.1) to years of storage under ambient conditions and use (*n* ≈ 2.7).

The synthesis of the homologous *meta*-telluridomercurate K₄[HgTe₃]·H₂O 2 can be carried out with the quoted, typical variations for a replacement of Se with Te: treatment of K₂Te and HgSO₄·*n*H₂O in en under solvothermal reaction conditions at slightly reduced temperature (130 °C). However, the yields are lower, since K₂Te₂,²⁶ K₂Te₃,²⁷ and amorphous side products are obtained along with 2. Intending to increase both yield and purity, we changed the reaction conditions to the use of a solid mixture containing sodium besides potassium. This way, we thought to allow for more flexibility in crystallization conditions according to various ionic radii. Further, we used DMF as solvent, as it is known that [Na(dmf)₆]⁺ cations are easily formed and well soluble. However, in the present case, not 2 or its sodium salt was formed, but the *meta*-telluridomercurate K₁₀Na₂[HgTe₃]₂(HCO₂)₃ (3). 3 comprises a formate moiety and Na⁺ as well as K⁺ counter ions. The formation of sodium formate is often observed as a by-product due to decomposition of DMF under the given conditions; for this, the presence of the additional anion was not unexpected.

With a solvothermal reactions starting from a phase of the nominal composition K₂PbHgSe₂ and pyridine as solvent, we

investigated whether we might access a mixture of metalate anions, either in a double salt or a salt of a ternary anion, or not. The latter turned out to be the case, as small amounts of K₂[HgSe₂]·H₂O (4) emerged from the experiment upon addition of 0.5 vol% of water to the reaction mixture. Complete absence of water, or other volume ratios than 995 : 5 led to no or lower yields.

For us, such hydrate compounds are particularly interesting for their apparent stability in water and thus the potential for subsequent reactions in aqueous media, which are, however, beyond the scope of this report.

The crystallization of the hydrate 4 inspired us to revisit the formation of heavy main-group *ortho*-chalcogenidometalates, which have either not been identified at all (Tl, Bi) or at extremely low yields (Pb)¹¹ to date. Upon solvothermal treatment of a phase of the nominal composition K₅TlSe₄ and en/H₂O as solvent, K₆[Tl₂Se₆]·2H₂O (5) has been obtained. The formation of such dinuclear species was reported previously for some indates to be dependent on various reaction parameters, including the pH value.^{6,21,28} We thus explored the utilization of different amines, addition of various amounts of NH₃ or H₂O, and Brønstedt bases. However, for the heavier congener of indium, none of these variations yielded the mononuclear *ortho*-selenidothallate. Additional change of the nominal composition of the used phases afforded further compounds instead, K₆[Tl₂Se₆]·2K₂CO₃ (6) and K_{0.94}Tl_{0.06}[TlSe₂] (7). Compound 6 again comprises dinuclear aggregates of the *ortho*-selenidothallate anion, while 7 incorporates infinite 1D strands {[TlSe₂][−]}_{*n*} of corner sharing [TlSe₄] tetrahedra. So far, such chalcogenidothallate strands were only obtained as K_{*x*}Tl_{1−*x*}[TlSe₂] with $x < 0.44$,^{29,30} but they are also known for the lighter homologues, for instance from KInTe₂.³¹

Only when changing the chalcogen from selenium to tellurium, and increasing the water content within the en/H₂O mixture to 3 vol%, the first known *ortho*-chalcogenidothallate(III) was obtained as {K₅[TlTe₄]·2KOH}₃·10H₂O (8). The high water content is apparently necessary for the formation and/or crystallization of 8; replacement with NH₃, for instance, did not lead to crystalline products. However, at the same time the presence of water triggers the formation of polytellurides and large amounts of amorphous side products, hence, the product had to be hand-separated from the reaction mixture.

Application of similar reaction conditions, including a 95 : 5 en/H₂O solvent mixture, to the generation of respective lead compounds resulted in the formation of a hydrate salt of the rare *ortho*-selenidoplumbate anion, K₄[PbSe₄]·en·H₂O (9). The structure is isostructural to the ammoniate K₄[PbSe₄]·en·NH₃, which was published only recently.¹¹ The crystallographic discrimination from the ammoniate was unambiguous upon collection of a data set of very high quality (see below). Further analytical methods fail here, as both ammonia N–H and water O–H vibrations are overlaid by the en N–H vibrations, and as elemental analyses tend to produce erroneous results for these heavy metal salts. The existence of a hydrate is not less amazing than the discovery of the [PbSe₄]^{4−} anion itself – the latter is anticipated to be highly



redox active, due to the relatively poor stabilization of formal Pb(IV) by selenide and furthermore supposed to be highly oxophilic. Its relative stability in the presence of H₂O, instead of spontaneous formation of [PbO₄]^{4−} or even spontaneous reduction, was thus unexpected. Counterintuitively again, the yield was dramatically increased from a few crystals of the ammoniate, to approx. 20% of the hydrate **9**, although separation of the crystals by hand from the reaction mixture remained necessary.

Finally, we attempted solvothermal treatment of phases of the nominal composition K_xBi_ySe_z. Due to its thermodynamic preference, K₃[BiSe₃] (**10**) was obtained from all batches. The structure was reported in the literature, however upon a rather cumbersome experimental procedure: Bi₂O₃ was reduced with K₂[CO₃] in a selenium-charged hydrogen stream at 850 °C.¹⁵ **10** can alternatively be obtained from simple fusion of the elements and successive work up *via* excessive washing with en until the filtrate turns colorless. **10** is then obtained as micro-crystalline powder in almost quantitative yields. Single crystals suitable for X-ray analysis were obtained *via* recrystallization by solvothermal treatment in en.

3.1 Crystal structure description

Crystal structure of K₄[HgSe₃]·H₂O (1). **1** crystallizes in the monoclinic space group *P*2₁ with four formula units in the unit cell. The crystal features orthorhombic geometry, but monoclinic symmetry upon pseudo-merohedral twinning pretending orthorhombic symmetry. In the [HgSe₃]^{4−} anions in **1**, the Hg²⁺ ions are coordinated by three Se atoms in a distorted trigonal planar manner (deviation from ideal Se–Hg–Se 120° angles by up to 10°; sum of angles around central Hg atom 359.97°). Within the crystal structure, the anions are stabilized by K⁺ and [K(μ-H₂O)K]²⁺ counter ions and thus well separated from each other (see Fig. 1).

Each K⁺ ion is situated in a distorted octahedral environment of five Se atoms and one O atom, with the exception of K4 with coordination number (c.n.) 7 (see ESI†).

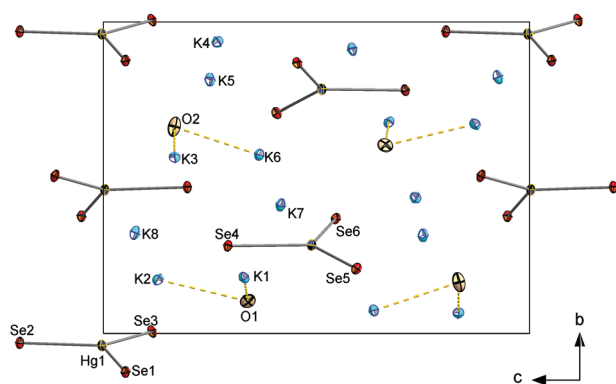


Fig. 1 Fragment of the crystal structure of **1**. Ellipsoids are drawn at 50% probability. Selected structural parameters [Å, °]: Hg–Se: 2.552(2)–2.621(2), K...Se 3.233(5)–3.620(5), K...O 2.78(2)–3.19(2), Se–Hg–Se 114.02(6)–129.76(7).

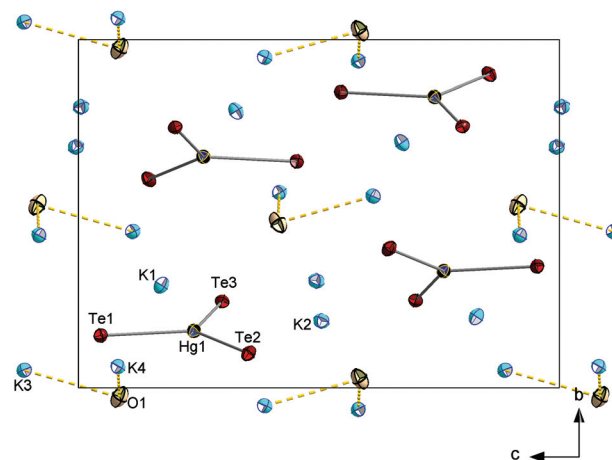


Fig. 2 Fragment of the crystal structure of **2**. Ellipsoids are drawn at 50% probability. Selected structural parameters [Å, °]: Hg–Te: 2.7256(6)–2.7775(6), K...Te 3.468(2)–3.774(2), K...O 2.800(7)–3.258(7), Te–Hg–Te 114.427(18)–126.824(18).

Crystal structure of K₄[HgTe₃]·H₂O (2). **2** crystallizes in the orthorhombic space group *P*2₁2₁2₁, a minimal non-isomorphic supergroup to that in **1**, with four formula units in the unit cell. The [HgTe₃]^{4−} anions also adopt a distorted trigonal planar shape (deviation from ideal angles by approx. 6°, sum of angles around central Hg atom 359.94°, see Fig. 2).

K⁺ and [K(μ-H₂O)K]²⁺ ions separate the anions from each other, again with distorted octahedral coordination environments of K⁺ ions (see ESI†).

Crystal structure of K₁₀Na₂[HgTe₃]₂(HCO₂)₃ (3). **3** crystallizes in the monoclinic space group *C*2/*m* with 2 formula units in the unit cell. Maximum deviation from the ideal Te–Hg–Te angle within the [HgTe₃]^{4−} anions amounts to 2°, the sum of angles around the central Hg atom is 359.98°. Formate anions are linked to adjacent formate anions in two different modes: first *via* one K⁺ ion to form a nearly linear O...K...O connection (K...O 2.756(3) Å), and second *via* two slightly bent O...Na...O bridges (Na...O 2.898(4)–3.058(5) Å; O...Na...O 45.66(14)°, 77.19(11)°) and one additional O...K...O linkage (2.701(3)–2.854(4) Å), see Fig. 3. This way, a complicated 3D cationic network is formed to embed well-separated [HgTe₃]^{2−} anions (see ESI†).

Crystal structure of K₂[HgSe₂]·H₂O (4). **4** crystallizes in the monoclinic space group *P*2₁/*c* with four formula units in the unit cell. Hg²⁺ ions are coordinated by three Se atoms, of which two are μ-bridging to adjacent Hg²⁺ ions, while one is a terminal ligand. This way, {[HgSe₂]^{2−}}_n 1D anion strands are formed that extend along [001] (see Fig. 4). One of the two crystallographic independent K⁺ ions (K4) is situated in close vicinity to its symmetry equivalent, forming a trapezoidal [K₂(μ-H₂O)]²⁺ cation *via* bridging by two μ-H₂O. Further saturation of the coordination sphere is achieved by three Se atoms. The second K⁺ ion is coordinated by six Se atoms (K2, see ESI†).



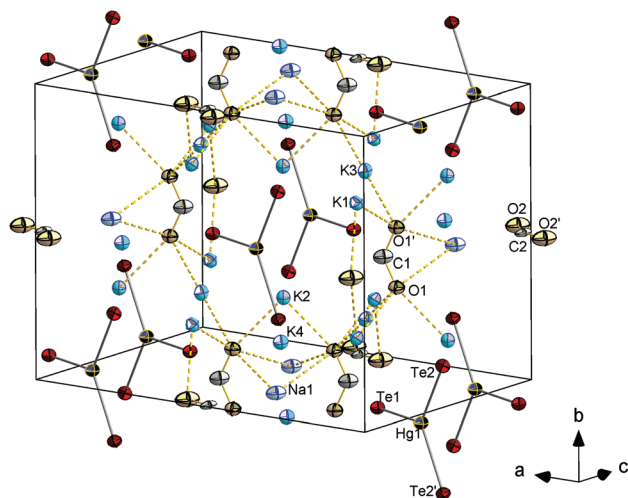


Fig. 3 Fragment of the crystal structure of 3. Ellipsoids are drawn at 50% probability. Selected structural parameters [$\text{\AA}$, $^\circ$]: Hg–Te: 2.7397(4)–2.7795(5), K–Te 3.5028(10)–3.7766(10), K...O 2.701(4)–2.854(4), Na...Te 3.5342(13), Na...O 2.898(4), Te–Hg–Te 119.246(8)–121.482(16).

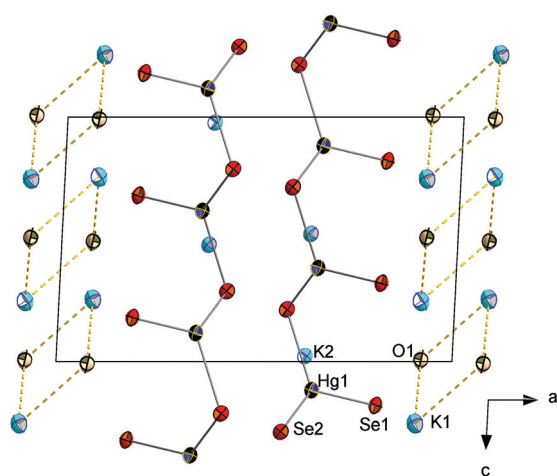


Fig. 4 Fragment of the crystal structure of 4. Ellipsoids are drawn at 50% probability. Selected structural parameters [$\text{\AA}$, $^\circ$]: Hg–Se1 2.5097(10), Hg–Se2 2.5908(10)–2.6090(11), K...Se 3.193(2)–3.497(2), K...O 2.789(9)–3.426(9), Se–Hg–Se 107.19(3)–132.38(3).

Crystal structure of $\text{K}_6[\text{Ti}_2\text{Se}_6]\cdot 2\text{H}_2\text{O}$ (5). 5 crystallizes in the triclinic space group $P\bar{1}$ with one formula unit in the unit cell. Each (formal) Ti^{3+} ion is coordinated by two terminal Se atoms and two μ -bridging Se atoms, thus forming a molecular, dinuclear $[\text{Ti}_2\text{Se}_6]^{6-}$ anion. The anions are separated by K^+ ions and $[\text{K}_2(\mu\text{-H}_2\text{O})]^{2+}$ units (see Fig. 5). K^+ ions are each coordinated by five Se atoms and one O atom in a distorted octahedral fashion (see ESI†).

Crystal structure of $\text{K}_6[\text{Ti}_2\text{Se}_6]\cdot 2\text{K}_2\text{CO}_3$ (6). 6 crystallizes in the triclinic space group $P\bar{1}$ with one formula unit in the unit cell. The constitution of $[\text{Ti}_2\text{Se}_6]^{6-}$ anions is the same as found in 5. Carbonate ions are coordinating to eight K^+ ions,

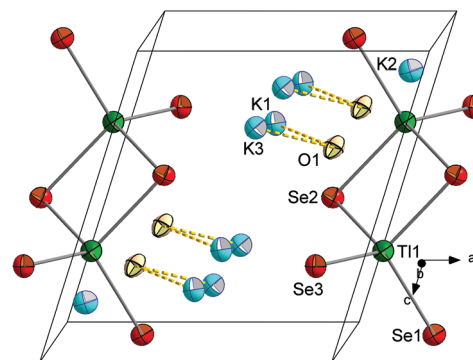


Fig. 5 Fragment of the crystal structure of 5. Ellipsoids are drawn at 50% probability. Selected structural parameters [$\text{\AA}$, $^\circ$]: Ti–Se(13) 2.5905(15)–2.5918(15), Ti–Se(2) 2.6876(15)–2.7132(15), K...Se 3.223(4)–3.908(4), K...O 2.766(11)–2.901(12), Se(1)–Ti–Se(3) 114.53(5), Se(1,3)–Ti–Se(2) 106.85(5)–116.66(5), Se(2)–Ti–Se(2') 96.17(4), K...O...K 96.5(3).

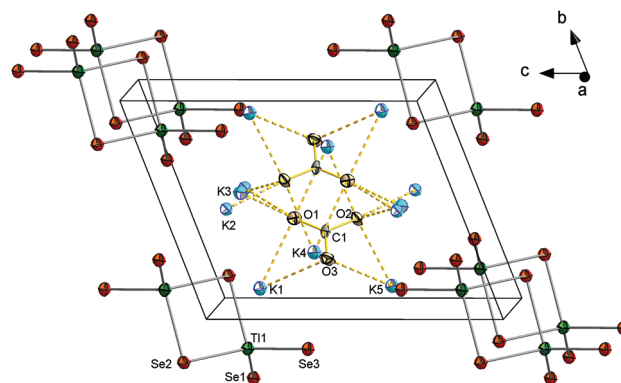


Fig. 6 Fragment of the crystal structure of 6. Ellipsoids are drawn at 50% probability. Selected structural parameters [$\text{\AA}$, $^\circ$]: Ti–Se(1,3) 2.5824(11)–2.5828(12), Ti–Se(2) 2.6629(10)–2.7273(11), K...Se 3.246(2)–3.919(3), K...O 2.681(9)–3.366(9), C–O 1.270(13)–1.327(12), Se(1)–Ti–Se(3) 115.80(3), Se(1,3)–Ti–Se(2) 103.64(4)–116.71(4), Se(2)–Ti–Se(2') 93.17(3).

three of which are chelated by the same one carbonate ion. Five further terminal K^+ ions, in turn, are situated between two adjacent carbonate ions to form a complicated 2D $[\text{K}_{10}(\text{CO}_3)_2]^{6+}$ sheet extending between layers of the anions parallel to the ab plane by interconnecting strands of $[\text{K}_5(\text{CO}_3)]^{3+}$ along the crystallographic a direction (see Fig. 6 and ESI†).

Crystal structure of $\text{K}_{0.94}\text{Ti}_{0.06}[\text{TiSe}_2]$ (7). 7 crystallizes in the tetragonal space group $I4/mcm$ with four formula units in the unit cell, being isostructural with $\text{Ti}[\text{TiSe}_2]$.³⁰ Hence, $[\text{TiSe}_4]$ tetrahedra form 1D strands $\{[\text{TiSe}_2]^{-}\}_n$ along c by edge sharing (see Fig. 7). The voids between the strands accommodate the counterions, which in 7 represent a statistical mixture of K^+ and Ti^+ ions. Refinement with free second variables results in a 94 : 6 ratio for K : Ti on the positions.

Crystal structure of $\{\text{K}_5[\text{TiTe}_4]\cdot 2\text{KOH}\}_3\cdot 10\text{H}_2\text{O}$ (8). 8 crystallizes in the trigonal space group $R\bar{3}c$ with six formula units in



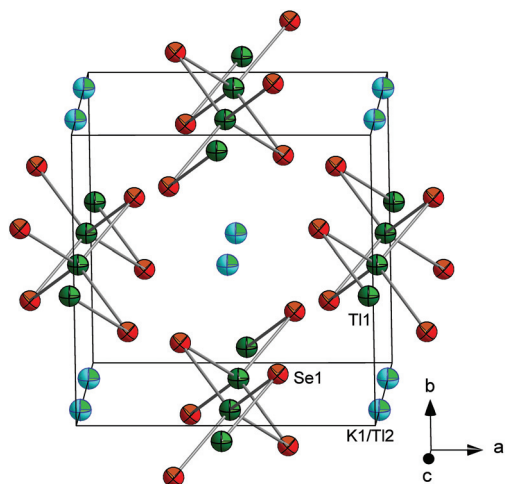


Fig. 7 Fragment of the crystal structure of 7. Ellipsoids are drawn at 50% probability. Two-colored K1/Tl2 ellipsoids indicate statistical disorder on the respective positions. Selected structural parameters [$\text{\AA}$, $^\circ$]: Tl–Se 2.6653(7), K...Se 3.4299(4), Tl–Tl 3.5052(5), Se–Tl–Se 97.77(3), 115.619(14).

the unit cell. The (formal) Tl^{3+} ions are surrounded by Te atoms in a tetrahedral mode, thus forming molecular $[\text{TlTe}_4]^{3-}$ anions that are embedded in and separated by a complex 3D $\{\text{K}\cdots(\text{OH})\cdots\text{K}\cdots(\text{H}_2\text{O})\}_n$ substructure (see Fig. 8). Most of the K^+ ions are hexacoordinate in an octahedral manner ($\text{K5}\cdots\text{Te}_6$), a distorted octahedral manner ($\text{K3}\cdots\text{Te}_4\text{O}_2$), an distorted tetrahedral ($\text{K2}\cdots\text{Te}_3\text{O}$) or a trigonal prismatic manner ($\text{K1}\cdots\text{Te}_3\text{O}_3$). One of the cations is nonacoordinate ($\text{K4}\cdots\text{Te}_2\text{O}_7$). O atoms in turn are either surrounded by 6 K^+ ions in a distorted octahedral fashion (O1, O4) or possess a tetragonal pyramidal environment by 5 K^+ ions (O2, O3) (see ESI†). As illustrated in Fig. 8, the crystal structure of this first *ortho*-chalcogenidothallate salt is extreme complex, with a c axis of 39.226(6) $\text{\AA}$ and a unit cell volume of 9718(2) $\text{\AA}^3$, which is very large for an *ortho*-chalcogenidometalate salt. Maybe the steric requirements or even the electrostatic properties of the $[\text{TlCh}_4]^{5-}$ anion impede a suitable and efficient packing of alkali(ne earth) metal cations along with the anions; hence, co-crystallization with two formula units of KOH might be essential, however leading to the shown structural complexity.

Crystal structure of $\text{K}_4[\text{PbSe}_4]\cdot\text{en}\cdot\text{H}_2\text{O}$ (9). 9 crystallizes in triclinic space group $P\bar{1}$ with two formula units in the unit cell. Compound 9 is isostructural with $\text{K}_4[\text{PbSe}_4]\cdot\text{en}\cdot\text{NH}_3$.¹¹ The presence of water instead of ammonia was unambiguously confirmed by structure solution and refinement using a very precise data set: the refinement of nitrogen atoms on the respective positions led to not positive-defined displacement parameters; furthermore, the two hydrogen atoms per water molecule were localized on the difference Fourier map and refined freely. Thus, 9 indeed represents the second known *ortho*-chalcogenidoplumbate(IV) salt (see Fig. 9). Bond lengths and angles are in good agreement with the reported structure parameters of the ammoniate.

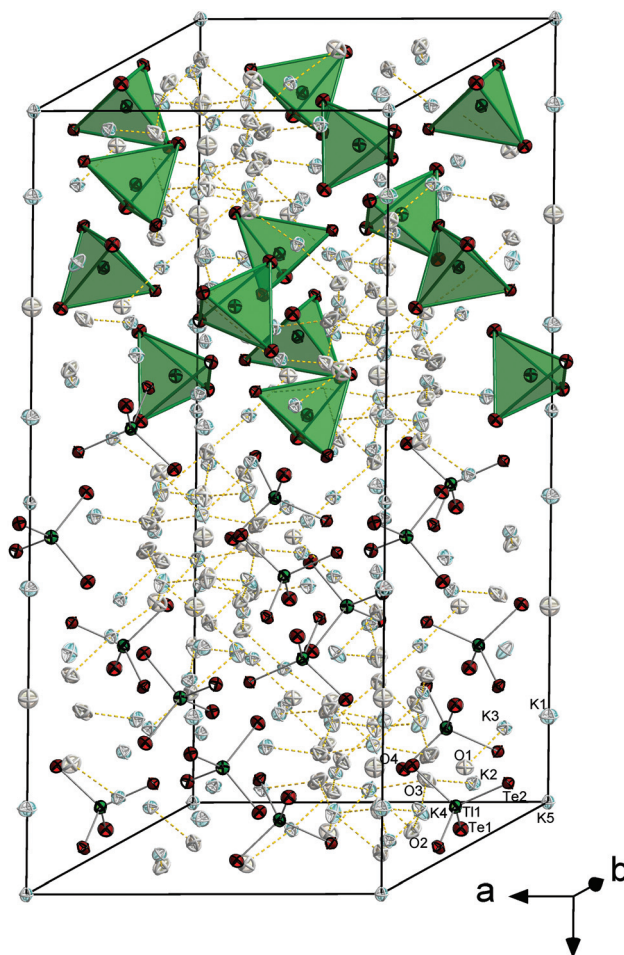


Fig. 8 Fragment of the crystal structure of 8. Ellipsoids are drawn at 50% probability. K and O atoms are drawn in opaque mode, and some of the tetrahedral anions are shown as polyhedra for clarity. Selected structural parameters [$\text{\AA}$, $^\circ$]: Tl–Te 2.8386(12)–2.8458(13), K...Te 3.537(4)–3.916(5), K...O 2.753(15)–3.053(6), Te–Tl–Te 107.26(3)–122.43(3).

3.2 Quantum chemical calculations

The compounds presented above served to fill some of the existing gaps in the spectrum of known mononuclear or dinuclear chalcogenidometalate anions of the heaviest metals (see Table 1 below). However, many combinations have still not been observed experimentally. We were thus interested, whether according, yet missing species are (a) local minima on the respective energy hypersurfaces, and whether they are (b) accessible *via* chemical reactions in principle.

This might then allow for recommendations for future synthetic approaches: in the case of endoenergetic reactions, classical solid state reaction would more likely be successful, while in the case of an exoenergetic reaction, a reaction in solution should in principle be also possible or even preferred.

In a first step, we simultaneously optimized electronic and geometric structures of all mononuclear anions (*meta* and *ortho* type) and the pseudo-dimeric ones by means of DFT calculations, followed by the determination of second derivatives



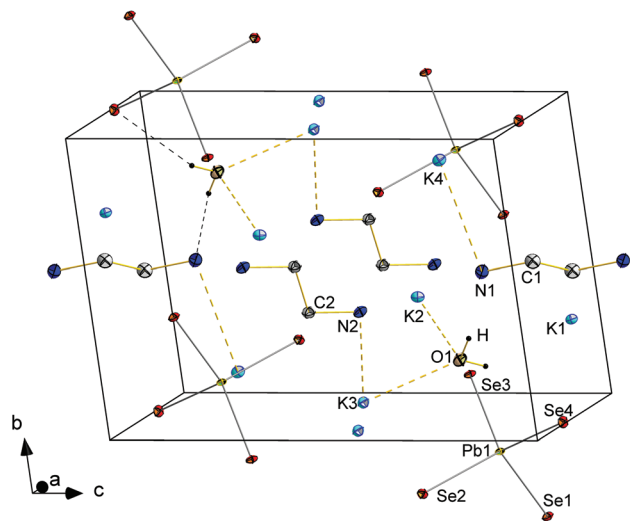


Fig. 9 Fragment of the crystal structure of **9**. Ellipsoids are drawn at 50% probability. H atoms are omitted for clarity. Selected structural parameters [Å,°]: Pb–Se 2.5985(4)–2.6157(4), K...Se 3.2861(9)–3.4920(9), K...O 2.776(3)–3.038(3), K...N 2.801(3)–3.384(4), Se–Pb–Se 107.02(2)–110.94(2).

of the energy to verify the local minimum character of the shown anions on the energy hypersurface (non-minimum structures to be found in the ESI†). In a second step, we compared the energies of the respective anions and potential elimination products to calculate hypothetical reaction energies.

The accuracy of the calculated structures are within the expected range, *i.e.*, bond lengths by a maximum of 0.067 Å larger than experimentally found ones. Hence, the chosen method is reliable enough for investigation of reaction energies.

It should be noted that the energy values that result upon application of COSMO are afflicted with systematic errors. This is due to a mismatch of gradient and absolute energy conver-

gence in all cases, and even more pronounced if the numbers and/or charges of educts and products differ. Still, the errors are comparably small compared to the errors that are made during calculations of anions without the use of COSMO. In the present case, the errors are identical or at least very similar for all calculations as all species are highly related, such that a discussion of relative energies is reasonable. Furthermore, for the calculations, we chose appropriate reactants to account for the named problems with the quantum chemical treatment of anions: by using H_2Se and $(\text{HSe})^-$ for balancing the number of selenium ions (instead of Se^{2-}), and by the choice of synthesis steps that do not require the consideration of naked metal cations Hg^{2+} , $\text{Tl}^{1/3+}$, $\text{Pb}^{2/4+}$, or $\text{Bi}^{3/5+}$ we cared for a minimum of charges and charge differences to be dealt with. The results are presented in the following, on the example of the respective selenides.

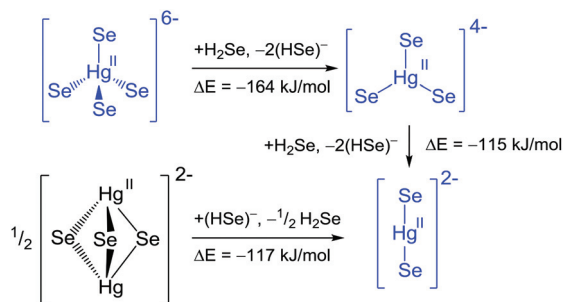
Mercurates. Including our own contributions of the synthesis of **1–3**, three mononuclear chalcogenidomercurate anions have been experimentally observed so far: $[\text{HgCh}_4]^{6-}$, $[\text{HgCh}_3]^{4-}$, and $[\text{HgSe}_2]^{2-}$. However, attempts to generate the so far unknown $[\text{Hg}_2\text{Ch}_3]^{2-}$ anion, in analogy to $[\text{Pb}_2\text{Ch}_3]^{2-}$ (ref. 13) by *in situ* reduction of chalcogen-rich metalchalcogenides with alkali metals in solution¹⁵ have not succeeded, but led to the formation of the well-known, linear $[\text{HgCh}_2]^{2-}$ anions⁹ in all cases. This could be rationalized by the calculations illustrated in Scheme 1.

Starting out from the *ortho*-mercurate anion in a hypothetical transformation pathway through the mononuclear *meta*-mercurate towards the linear $[\text{HgSe}_2]^{2-}$ anion under release of an Se^{2-} ligand in each step (considered as $(\text{HSe})^-$ for the reasons given above), both reaction steps are exoenergetic. This is in line with the synthesis of $[\text{HgCh}_2]^{2-}$ from solution and $[\text{HgCh}_4]^{6-}$ from solid state reactions, and does not contradict an access of the molecular *meta*-mercurate under solvothermal conditions. Starting from a hypothetical $[\text{Hg}_2\text{Se}_3]^{2-}$ anion in a reverse reaction scheme produces two equivalents of the linear $[\text{HgSe}_2]^{2-}$, again in an exoenergetic reaction.

Table 1 Overview of previously known (x) or newly synthesized (✓) mononuclear selenidometalates or telluridometalates of Hg, Tl, Pb, or Bi as central atoms. Both signs together indicate significant additions to a known species (see text). Empty boxes indicate so far unknown species. Circles or dashes, respectively, indicate whether the calculated structures of the selenides represent local minima on the energy hypersurfaces or not (or did not converge at all). According tellurides have not been calculated in this study

	Higher formal oxidation state of M		Lower formal oxidation state of M		Higher formal oxidation state of M		Lower formal oxidation state of M		Higher formal oxidation state of M	
	Se	Te	Se	Te	Se	Te	Se	Te	Se	Te
Hg	⊗	x	—	—	✓	✓	—	—	⊗	x
Tl	○	—	○	—	○	—	○	—	○	—
Pb	—	—	—	—	○	—	○	x	✓	—
Bi	○	—	○	—	○	—	⊗	x	○	—





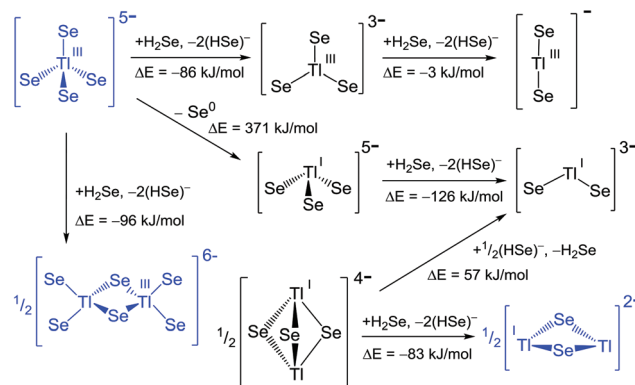
Scheme 1 Reaction pathways and energies for the transformation of chalcogenidomercurates. Blue colour indicates experimentally observed anions.

Thus, although $[\text{Hg}_2\text{Se}_3]^{2-}$ is a minimum on the energy hypersurface, it becomes obvious, why no such species has been reported so far: in solution the transformation towards $[\text{HgSe}_2]^{2-}$ is expected, while a solid state reaction employing the respective 2:2:3 stoichiometric ratio rather produces an extended mercurate framework structure, such as $\text{K}_2\text{Hg}_2\text{Se}_3$.²⁵ Obviously, this structure type is not available for the 6:1:4 ratio, hence solid state reactions allow for the isolation of the tetrahedral $[\text{HgSe}_4]^{6-}$ subunit. A comparison of the Hg–Se bond lengths for the planar $[\text{HgSe}_3]^{4-}$ (2.64 Å) versus $[\text{HgSe}_3]^{5-}$ (3.01 Å) is in agreement with the formal +II oxidation state at the Hg atoms in 1–3.

To the best of our knowledge, no chalcogenidomercurate(I) anion has been characterized so far. Nevertheless, we calculated the corresponding anions $[\text{Hg}_2\text{Se}_3]^{4-}$ (trigonal bipyramidal) and $[\text{HgSe}_3]^{5-}$ (trigonal planar or pyramidal). The trigonal bipyramidal anion does not converge into a reasonable structure. The *meta*-mercurate(I) is a minimum on the energy hypersurface in the planar and the pyramidal conformation, with the latter being favoured, but both compounds are open shell systems, which is expected to hamper their isolation.

Thallates. The situation for chalcogenidothallates is somewhat more complex in terms of the availability of different oxidation state. Both, +I and +III, are known within experimentally observed metalate anions, as *ortho*-chalcogenidothallate(III) anion $[\text{TlCh}_4]^{5-}$ (5–8) and the butterfly-like chalcogenidothallate(I) anion $[\text{Tl}_2\text{Ch}_2]^{2-}$. However, as shown in Scheme 2, even more of the small anions turned out to be minima on the energy hypersurface.

Hypothetic reaction cascades starting out from the *ortho*-thallate anion through a molecular *meta*-thallate anion $[\text{TlSe}_3]^{3-}$ and further on to a linear $[\text{TlSe}_2]^{-}$ anion are exoenergetic, but especially the second step is only slightly exoenergetic. Principally, the existence of the respective species under the simulated conditions should hence be possible. However, this pathway is obviously excelled by the shown and observed dimerization, which is even more exoenergetic. A possible redox reaction yielding a pyramidal $[\text{TlSe}_3]^{5-}$ anion is energetically highly disfavoured, such that the exoenergetic step to the bent $[\text{TlSe}_2]^{3-}$ species is not achievable this way. Thus, it

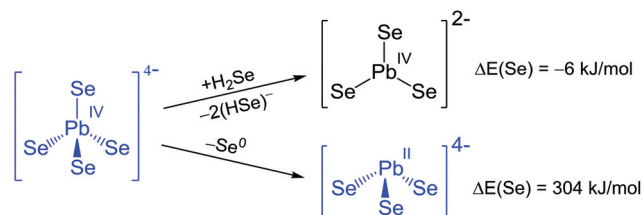


Scheme 2 Reaction pathways and energies for the transformation of selenido-thallates. Blue colour indicates experimentally observed anions.

becomes perspicuous why an interconversion between thallate(III) and thallate(I) anions have not yet been observed. Regarding the possible thallate(I) species, both $[\text{TlSe}_2]^{3-}$ and the butterfly-like $[\text{Tl}_2\text{Se}_2]^{2-}$ anion should be accessible from an energetic point of view starting out from a (hypothetical) $[\text{Tl}_2\text{Se}_3]^{4-}$ bipyramid, but only the second was observed so far. Neither Schrobilgen's comprehensive NMR studies on such systems,¹⁰ nor our own attempts¹⁶ have so far yielded any indication of further thallate(I) anions. However, a related compound, $[\text{TlTe}_3\text{Pb}]^{3-}$,^{12b} was published, hence further extension and variations of the synthetic approach should eventually allow for the synthesis of the other species, maybe even by the use of the quoted ternary anion.

Plumbates. As for the lighter homologues, the only possible mononuclear chalcogenidoplumbates are those given in Scheme 3, besides $[\text{Pb}_2\text{Ch}_3]^{2-}$. Reaction energies for transformations starting from $[\text{Pb}_2\text{Ch}_3]^{2-}$, however, require the elimination of Pb^{2+} or the simulation of PbSe formation, both of which are not feasible with the methods used here (see also above). Bent $[\text{PbCh}_2]^{2-}$, as found in the anionic Tl/Ch system, possess two imaginary vibrational modes, thus does not represent a minimum on the energy hypersurface.

The degradation of *ortho*-selenidoplumbate(IV) anion towards a carbonate homologue under release of a selenide ligand is not associated with a significant energy change,



Scheme 3 Reaction pathways and energies for the transformation of selenidoplumbates. Blue colour indicates experimentally observed anions, however, in the case of the $[\text{PbCh}_3]^{2-}$ anion, only the tellurium analogue has been isolated so far.



hence differences either considering a solvent shell or a counterion environment may easily overcompensate it to an overall preference of the tetrahedral anion. An oxidative elimination of selenium to form the pyramidal $[\text{PbSe}_3]^{4-}$ anion is considerably endoenergetic, apparently contradicting the experimental observation of the *meta*-telluridoplumbate(II), but in agreement with a synthesis starting out from a Pb(II) precursor and thus avoiding the disfavoured redox reaction step. Moreover, the distinct energetic preference of the *ortho*-selenidoplumbate(IV) over its dissociation or dechalcogenation is in line with its formation in and crystallization from solution, even as a hydrate like **9**.

Bismuthates. For chalcogenidobismuthates, the list of experimentally observed anions is the smallest along the elemental combinations investigated in the current study. However, the list of hypothetical anions comprises four minima on the energy hypersurface (see Scheme 4). Further minima on the energy hypersurface with intriguing electronic features are the neutral trigonal bipyramidal Bi_2Ch_3 , which however rather exist as the extended solid state compounds paraganajuatite³² or tellurobismuthite,³³ and two chalcogen-bismuth cations $[\text{Bi}_2\text{Ch}_2]^{2+}$ with butterfly-like geometry and linear $[\text{BiCh}_2]^+$. For the reasons given above, these shall not be discussed in the context of chalcogenidometalate anion chemistry.

All reactions starting from a hypothetical *ortho*-bismuthate(V) anion including stepwise elimination and/or dissociation of $\text{Se}^0/\text{Se}^{2-}$ are endoenergetic. Thus the existence of $[\text{BiSe}_4]^{3-}$ does not seem to be generally unrealistic. However, very obviously, so far only chalcogenidobismuthates of Bi(III) have been produced that seem to be kinetically stable enough not to undergo the hypothetical, exoenergetic oxidation towards the tetrahedral species. $[\text{BiSe}_3]^-$ would be the second heaviest possible homologue of the nitrate $[\text{NO}_3]^-$ anion, yet its formation is not favourable according to energetic considerations. Also a bent $[\text{BiSe}_2]^-$ species represents a minimum on the energy hypersurface, yet its formation from the observed $[\text{BiSe}_3]^{3-}$ anion would require another 21 kJ mol⁻¹. For this, although not isolated yet, the list of potentially observable Bi/Se anions definitely include the homologues of $[\text{NO}_3]^-$, $[\text{PO}_4]^{3-}$ and $[\text{NO}_2]^-$ in addition to the observed $[\text{BiCh}_3]^{3-}$ anions. The

Table 2 Overview of previously known (x) or newly synthesized (✓) dinuclear selenidometalates or telluridometalates of Hg, Tl, Pb, or Bi as central atoms. Empty boxes indicate so far unknown species. Circles indicate that the calculated structures of the selenides represent local minima on the energy hypersurfaces. The entry n/a indicates that an according species was not considered as it would be either neutral or cationic. According tellurides have not been calculated in this study

						
	Lower formal oxidation state of M		Lower formal oxidation state of M		Higher formal oxidation state of M	
	Se	Te	Se	Te	Se	Te
Hg	n/a		○		○	
Tl	○	x	○		○	✓
Pb	n/a		⊗	x	○	
Bi	n/a		n/a		○	

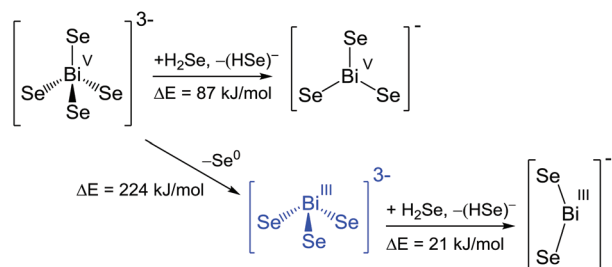
remaining challenge is finding an appropriate synthetic approach, such as described recently and once more herein for salts of $[\text{PbSe}_4]^{4-}$.

4 Conclusions

We presented the synthesis and structure of novel *ortho*- and *meta*-chalcogenidometalates of mercury, thallium and lead, including the first molecular *meta*-chalcogenidomercurate(II), the first *ortho*-chalcogenidothallate(III) and the second *ortho*-selenidoplumbate(IV). The latter emerged for the first time and unexpectedly as its hydrate salt. In addition, we reported a new and facile synthetic pathway for the generation of the *meta*-selenidobismuthate in almost quantitative yield. The existence of all those anions inspired quantum chemical investigations of structures and energetics of the diverse species with different elemental combinations. We discussed hypothetical reaction energies, which helped to rationalize the observations and to predict further species to be isolated in future work. In the case of the chalcogenidoplumbates, the heaviest $[\text{CO}_3]^{2-}$ homologues are generally stable, however most likely not accessible *via* solution based techniques. The same is valid for the chalcogenidobismuthate homologues of $[\text{NO}_3]^-$. Tables 1 and 2 summarize the experimental and theoretical findings for all smallest molecular chalcogenidometalate anions of the heaviest metals, including the new ones.

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Scheme 4 Reaction pathways and energies for the transformation of selenidobismuthates. Blue colour indicates experimentally observed anions.



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