Mercury(II) Binding to Metallothionein in *Mytilus edulis* revealed by High Energy-Resolution XANES Spectroscopy

Abstract: Of all divalent metals, mercury (Hg²⁺) has the highest affinity for metallothioneins (MTs) in living cells. Hg²⁺ is considered to be enclosed in the α and β domains as tetrahedral α-type Hg₄Cys₁₁₋₁₂ and β-type Hg₄Cys₇ clusters similar to Cd⁰ and Zn⁰. However, neither the four-fold coordination of Hg nor the existence of Hg–Hg atomic pairs have ever been demonstrated, and the Hg⁻³ partitioning among the two protein domains is unknown. Using high energy-resolution XANES spectroscopy, MP2 geometry optimization, and biochemical analysis, evidence for the coexistence of two-coordinate Hg-thiolate complex and four-coordinate Hg-thiolate cluster with a metacinnabar-type (β-HgS) structure in the α domain of separate metallothionein molecules from blue mussel under in vivo exposure is provided. The findings suggest that the CXXC claw setting of thiolate donors, which only exists in the α domain, acts as a nucleation center for the polynuclear complex and that the five CXC motifs from this domain serve as the cluster-forming motifs. Oligomerization is driven by metallocophilic Hg–Hg interactions. Our results provide clues as to why Hg has higher affinity for the α than the β domain. More generally, this work provides a foundation for understanding how metallothioneins mediate mercury detoxification in the cell under in vivo conditions.

Introduction

How is 5d³ Hg⁺ bonded to the canonical αβ-domains of metallothioneins (MTs) in living cells? This question is important to address of the understanding of the intracellular fate of this potent toxin. If Hg⁺ is tetrahedrally coordinated to four cysteine residues (Hg(Cys)₄), by analogy with 4d¹⁰ Cd⁺ and 3d³ Zn⁺ (Figure 1), how can one explain that extended X-ray absorption fine structure (EXAFS) spectroscopy only detects a linear two-coordination (Hg(Cys)₂) with a prototypical Hg-S distance of 2.33 Å? Also, if the metal tetrahedra are incorporated as tetrnuclear Hg₄(Cys)₁₁₋₁₂ (abbreviated as Hg₄S₁₁₋₁₂) clusters in the α domain and trinuclear Hg₃(Cys)₉ (abbreviated as Hg₃S₉) cluster in the β domain for a total of seven Hg per molecule (Hg₄-MT), similar to Zn⁴ in Znₓ-MT and Cd⁴ in Cdₓ-MT, why does not EXAFS detect a higher Hg shell beyond the thiolate ligands? Have answers to these questions failed because of structural disorder, mixture of the mononuclear and multinuclear bonding environments, or did Hg⁺ alter the tertiary structure of MTs to the point of suppressing the α and β domains to form a supercoiled peptide chain and Hg(Cys)₉ complexes only? Also, do the molecular structures determined in vitro at saturation (i.e., Hg₄-MT) or in excess (e.g., Hg₄S₉-MT) of the seven tetrahedral sites by X-ray absorption and nuclear magnetic resonance (NMR) spectroscopy and by X-ray crystallography represent how Hg is incorporated in vivo?

Here, we show that direct insight into Hg⁺ binding to MTs in living cells is provided by the new application of high energy-resolution X-ray absorption near-edge structure (HR-XANES) spectroscopy at extreme dilution. Compared to EXAFS, XANES spectroscopy provides geometric information, and has superior elemental sensitivity. Therefore, high spectral resolution allows more precise identification of the plurality of the bonding environments in the α and β domains, either within the same MT molecule, if all molecules are structurally equivalent, or among the MT molecules, if various coordination environments coexist in a mixture.

HR-XANES, applied earlier for the determination of the binding site of inorganic and organic mercury in human hair, is employed here to characterize the chemical form of mercury in whole blue mussel Mytilus edulis (ME) and its MT extract. This abundant and widely distributed filter-feeder mollusk is often used as a sentinel organism for marine pollutants. Also, the strong inducibility of its MTs and proximity of the cysteine amino acid sequence to those in mammals make this particular mussel a model organism of choice for studying Hg toxicity. The level of MT in ME was increased previously by exposure to cadmium or mercury, and two isoforms of apparent molecular weight 12 kDa (MT-10) and 20 kDa (MT-20) were isolated by polyacrylamide gel electrophoresis. Here, MTs were induced at Hg(NO₃)₂ concentration (100 μg Hg⁰ L⁻¹ or 0.1 ppm Hg) and exposure time (9 days) broadly similar to those in previous HgCl₂ (0.01 to 0.04 ppm for up to 21 days) and CdCl₂ experiments (0.1 ppm for 11 days) for 3–4 months, 0.2 ppm for 14 days, 0.2 ppm for 3–4 months.
and 0.4 ppm for 22 days\(^{16}\) and extracted by thermal denaturation (see the Experimental Section).

## Results

### Molecular weight and metal content

The metallothionein extracts from the Hg exposed (MT) and unexposed (control) mussels have similar sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) patterns (Figure 2, Experimental Section). Four main bands are observed, one faint at 10 kDa from monomers (MT-10), one smeared at 18 kDa from dimers (MT-20) followed by a faint and narrower band at 21 kDa, another faint at 30 kDa from trimers (MT-30), and an extremely intense last one at 37 kDa from tetramers (MT-40). The MT-20/30/40 oligomers were also detected in the initial mussel homogenate before the MT purification step by thermal denaturation of the cell proteins. The MT oligomer with a molecular weight of 21 kDa is unidentified.

Considering that MTs are metal-inducible proteins, Hg-exposed mussels should contain more MTs than the reference mussels, which we verified with a Bradford protein assay performed on the supernatant after thermal denaturation. The weight concentration of MTs in exposed mussels is 76 nmol g\(^{-1}\) wet weight (w.w.) mussel, compared to 57 nmol g\(^{-1}\) w.w. mussel under basal conditions. The two values are on the same order as those reported previously in the gills of *M. edulis* (35–56 nmol MT g\(^{-1}\) w.w.\(^{19}\)), and whole Pacific oyster *Crassostrea gigas* (38–91 nmol g\(^{-1}\) w.w.\(^{20}\)). The enhancement of MT induction in response to Hg exposure reached (76/57)/57 = 33%. The Hg:Cu:Zn molar stoichiometry, as determined from elemental analysis of the MT extracts using inductively coupled plasma mass spectrometry (ICP-MS), is 0:1:1 in the control and 2:2:5 in the Hg mussels. The constitutive levels of Cu and Zn in naturally occurring MTs are consistent with previ-
and with the role of MTs in the intracellular homeostasis of the two essential metals.\[22\] The total number of metals bound per molecule of MT was calculated by dividing the sum of the metals concentration in the MT extracts by the weight concentration of MTs in the same extracts and taking a MT-10 molecular weight of 7.24 kDa.\[18\] The metal to MT molecular ratios thus obtained are 4% for the control and 7% for the mussels exposed to Hg. Unmetallated apo-MT, or thionein,\[23\] is therefore the major form in solution after thermodenaturation. This form has probably two origins, the thioneins existing in vivo in the mussels and the thioneins generated by the release of metals (Cu, Zn, Hg) during the heating step. Some of the in vivo thioneins were certainly denatured, but their yields have no bearing on XANES results. The proportion of the metallated MTs is even lower if the metals are not randomly distributed among all MT molecules but clustered in the α- and β-domains of a few molecules. HR-XANES results show that the Hg atoms are indeed partly clustered in vivo, despite their vanishingly low amount (3.4 \(10^{-6}\) g Hg mL\(^{-1}\) MT extract, or ppm), supporting the finding that thionein largely prevails.

**Evidence for dithiolate and tetra thiolate Hg complexes**

The HR-XANES spectrum of the MT aqueous extract containing 3.4 \(10^{-6}\) g Hg mL\(^{-1}\) (ppm) is distinctly different from that of the tissues from whole Hg mussels containing 317 \(10^{-6}\) g Hg g\(^{-1}\) tissue or ppm (ME; Figure 3). Two observations stand out. First, the two spectra exhibit a near-edge peak at about 12279.3 eV. This "indicator" region, denoted A in Figure 3, is characteristic of Hg\(^{0}\) linearly coordinated to two thiolate groups [Hg(Cys)\(_2\) complex].\[24\] Its intensity is lower in MT, which indicates that this coordination is less abundant in this sample and coexists in a mixture with either three- or four-coordinate Hg (Figure 4).

![Figure 2](Image 48x569 to 289x775.png)

**Figure 2.** SDS-PAGE electropherograms of the MT extracts after thermal denaturation under argon. Lane 1 = 5 μL and lane 2 = 10 μL from the mussel control, lane 3 = mass markers in kDa, lane 4 = 5 μL and lane 5 = 10 μL from the Hg exposed mussels (MT).

Observation of two-coordinate Hg was expected in ME because it is the most common geometry in mercury chemistry and it occurs in biological systems in complexes with cysteine, peptides, and proteins.\[7a,12,25\] It was somewhat expected in the MT extract because the α domain has a conserved CXXC motif (single-letter amino acid code, where X can be any amino acid) known to bind mercury linearly in metalloproteins (Figure 1). Molecular mechanics (MM2) simulations for a mammalian MT model metallated with seven Hg atoms (Hg- MT consisting of Hg\(_{2}\alpha\), and Hg\(_{2}\beta\) clusters) by isomorphous substitution of Cd and Zn in the X-ray diffraction structure showed that the α cluster is too big to fit inside the protein and that some thiolate groups are exposed to the solvent (Figure S1, Supporting Information). These sulfur atoms could be engaged in trithiolate or tetrathiolate coordination inwardly and dithiolate linear coordination outwardly. The RS-Hg-SR coordination could behave as anchors to intermolecular bridging between MT-10 monomers forming MT-20/30/40.\[14a\] We show below from the quantitative analysis of the HR-XANES spectrum for MT that the amount of two-coordinate Hg is too high to arise significantly from intermolecular RS-Hg-SR crosslinks. In addition, compelling evidence for RS-SR crosslinks (in place of RS-Hg-SR) between monomeric subunits exists.\[14a\]

Second, ME has a broader absorption edge maximum and the trailing spectral edge of the MT extract is shifted to lower energy ("indicator" region D in Figure 3). According to the Natoli rule,\[26\] the energy shift of the post-edge maximum results from an increase of the interatomic distance (i.e., coordination number) between the photoabsorbing ion (Hg) and the neighboring atoms (S). Similar observations regarding the sensitivity of XANES to the nearest ligand distance were reported for the coordination geometry of cadmium in MTs\[9\] and for Hg coordinated to two [d(Hg–S) \(=\) 2.33 Å] and four [d(Hg–S)


Figure 4. Sensitivity of L_{3}-edge HR-XANES to the bonding environment of Hg. (a) Structure of the Hg complexes. (b–d) Spectra of two-, three-, and four-coordinate Hg-thiolate and Hg-sulfide references. The Hg(Cys)\textsubscript{2} complex was prepared at a Cys:Hg molar ratio of 2 and pH 7.5, the Hg(\text{o}-Pen\textsubscript{3})\textsubscript{3} complex at \text{o}-Pen:Hg = 10.0 and pH 11.5, and Hg(Cys\textsubscript{4})\textsubscript{3} at Cys:Hg = 10.0 and pH 11.9. Their structure was optimized geometrically (MP2-RB/def2-TZVP-ecp[67]), and the Hg(SR)\textsubscript{3} and \text{o}-HgS\textsubscript{NP} model are X-ray crystal structures. Peak B is diagnostic of Hg–Hg pairs. Bond lengths are in angstroms. Dark red, Hg; yellow, S; blue, N; red, O; gray, C; light gray, H.

≈2.52 Å sulfur atoms (Figures 4 and S2 in the Supporting Information).[24a] Thus, region D confirms that MT has less dithiolate Hg complex than ME, but still is not conclusive on the coordination of the second Hg species, which can be three- or four-fold. With an average Hg–S distance of 2.44 Å in thiolate complexes,[27] three-coordinate Hg has indeed a railing edge intermediate between those from the two and four coordination modes (Figure 4b). This question can be resolved with a two-component fit of ME and MT. The best fits were obtained with 91% Hg(\text{s}-Cys)\textsubscript{2} + 9% Hg(\text{o}-Pen\textsubscript{3})\textsubscript{3} for ME and 82% Hg(Cys)\textsubscript{2} + 18% Hg(Cys\textsubscript{4})\textsubscript{3} for MT (Figure 5a). The model fits reproduce well the data in regions A, C, and D, but fail to reconstruct peak B. We conclude that Hg is both two- and four-coordinate, and that MT has a higher proportion of the second species.

Evidence for Hg–Hg pairs in the tetrathiolate complex

Peak B is associated with the four-coordinate species because its intensity is higher in MT (Figure 5a). It is however absent in Hg(Cys\textsubscript{4})\textsubscript{3} (Figure 4d). The best shape among the tetrahedral compounds from our extended spectral database (ref.[11] and references therein) that best matched peak B was obtained with nanoparticulate metacinnabar (\text{o}-HgS\textsubscript{NP})\textsubscript{2} (Figure 4d). In \text{o}-HgS, Hg\textsuperscript{II} is coordinated to four sulfide atoms at 2.53 Å and surrounded by higher shells of Hg atoms.[28] The \text{o}-HgS\textsubscript{NP} reference was obtained by aging a Hg(\text{l}-Cys-OEt\textsubscript{2})\textsubscript{3} complex.[24b] The \text{o}-HgS\textsubscript{NP} HR-XANES spectrum is much different from the unaged Hg(\text{l}-Cys-OEt\textsubscript{2})\textsubscript{3} spectrum and cannot be reconstructed with a mixture of Hg(\text{l}-Cys-OEt\textsubscript{2}) and well crystallized \text{o}-HgS or any combination of other references (Figure S3a, Supporting Information). This material has broad \text{o}-HgS X-ray diffraction peaks[24b] and damped HR-XANES structures (Figures S2a and S3b, Supporting Information) characteristic of highly defective material with a wide range of both angles and bond lengths, as commonly observed for metal thiolate clusters in proteins,[7a, 9, 29] \text{o}-HgS\textsubscript{NP} has been identified previously in natural or organic matter and soils,[24b] in plants,[11] and in pyrite (FeS\textsubscript{2}).[30] The \text{o}-HgS\textsubscript{NP} crystallites imaged by HRTEM are about 5 nm in diameter (Figure S3b, Supporting Information).[24b, 30a] \text{o}-HgS\textsubscript{NP} has little bumps in region C (Figure 4d). These features, which result from multiple scattering events of the photoelectron on higher Hg shells (CN = 12), are more intense in well crystallized \text{o}-HgS (Figure S3b, Supporting Information), and should vanish when the \text{o}-HgS\textsubscript{NP} nanocrystals are smaller. We verified this hypothesis with a trithiolate complex [Hg(SR)\textsubscript{3}(XTL)], in which the Hg atoms have two nearest Hg neighbors at 3.89–3.90 Å (Figure 4c). The top edge region C is now loosely structured. Peak B is still present, although it is absent when the trithiolate complex is mononuclear [Hg(\text{d}-Pen\textsubscript{3})\textsubscript{3}] (Figure 4c). We conclude from this that peak B provides a signature for Hg–Hg pairs in three- and four-coordinate Hg.

Best-fit results of the two HR-XANES spectra were obtained with 81% Hg(Cys\textsubscript{2}) + 19% \text{o}-HgS\textsubscript{NP} for ME and 59% Hg(Cys\textsubscript{2}) + 41% \text{o}-HgS\textsubscript{NP} for MT (Figure 5b). The accuracy of estimation of the fit components is 6 mol% of total Hg.[12, 24b] We show...
Intra- and intermolecular Hg may have two origins. One is the elimination during MT purification of thermolabile dithiolate forms from Hg-metalloproteins, such as metallochaperones. SDS-PAGE shows that heat treatment effectively removed undesired high molecular weight proteins. A second is the transformation of some native Hg(Cys)$_2$ forms to Hg$_4$(Cys)$_4$ clusters at the elevated purification temperature of 95°C used here. This hypothesis was tested by thermal denaturation of an air-equilibrated mussel homogenate. In contact with oxygen, nonmetallated cysteine residues from MT are prone to oxidation to cystine and are unlikely to form new Hg$_4$(Cys)$_4$ clusters. In addition, metallochaperones, which in the cell would deliver metals to MTs, are unlikely to do it ex vivo in the homogenate, and even less so in the presence of oxygen. The HR-XANES spectra from the argon and air isolates are essentially the same, differing only in the intensity of the near-edge peak (Figure S4a, Supporting Information). The best two-component fit of the air isolate yielded 63% Hg(Cys)$_2$, +37% β-HgS$_{NP}$ values which coincide closely with those for the argon isolate (Figures 5 and S4b, Supporting Information). The repeatability of the results under two distinct experimental conditions is a strong indication that the enrichment of β-HgS$_{NP}$ in MT (41%) relative to ME (19%) results dominantly, if not only, from the denaturation at 95°C of thermolabile dithiolate forms.

The dithiolate MT complex, represented here by Hg(Cys)$_2$, at pH 7.5, features a digonal coordination to two nearest thiolate ligands at about 2.33 Å completed by one to two secondary amine ligands (Hg(SR)$_2$+N$_2$) coordination, Figures 4 and S2, Supporting Information. Intra- and intermolecular Hg–N bonding complementary to cysteine-derived Hg–S bonds has been observed in keratin and for Zn in metallothioneins, typically with histidine side chains. The dithiolate species could be also an oxothiolate complex, modeled here with the phytochelatine reference Hg[(γ-Glu–Cys)$_3$]$_2$Gly, abbreviated Hg(PC2) (Figure 5b). The Hg(PC2) reference is a bis-six-membered ring chelate, through two cysteinyl S atoms and two carboxyl oxygen atoms (C=O) at 2.6–3.0 Å from the peptide backbone (Figure 6a).

**Discussion**

**Nature and function of the MT-20/30/40 oligomers**

In SDS-PAGE, non-covalent interactions among individual polypeptide chains are disrupted, whereas subunits held together below that the inorganic mercury sulfide reference β-HgS$_{NP}$ is a computationally tractable structural analogue to the mercury thiolate “mineral core” in MT, thus providing the first spectroscopic proof for the existence of Hg–Hg pairs. Metallothioneins are produced in all tissues in bivalves, which explains their detection (19 ± 6 mol%) in whole mussel. Furthermore, the fit of the ME and MT spectra with only two and the same Hg references serves as internal consistency check for the reliability of our analysis and negates the possibility that a new Hg species formed during the purification procedure.
covalently by disulfide bonds between cysteine residues from two different chains remain attached. Mussels MT-10 have 21 Cys residues and the monomeric subunits of MT-20, known to be inducible by Cd-exposure, have 23 Cys residues.\[14a, 18,33\] In mussel Cd-MT-10, twelve out of the 21 Cys residues form tetranuclear Cd\(_4\)S\(_{12}\)-\(\alpha\) clusters, and the other nine form trinuclear Cd\(_3\)S\(_9\)-\(\beta\) clusters (Figure 1).\[28\] In the related Cd-experiment, the two additional cysteinyl sulfurs from MT-20 were considered to bridge the subunits through RS-Cd-SR or disulfide RS-SR links, thus leaving the 21 other Cys residues for the formation of the polynuclear Cd\(_4\)S\(_{12}\)-\(\alpha\) and Cd\(_3\)S\(_9\)-\(\beta\) clusters.\[14a\] Here, stabilization of the quaternary structure of MT-20/30/40 by RS-Hg-SR crosslinks formed during the Hg exposure can be dismissed because the MT oligomers are observed also in the mussel control. Therefore, our results suggest the presence of intermolecular RS-SR links within the mussel cytosol. Care was taken (Ar saturation of all solutions) during the MT purification to avoid the oxidation of thiols.

![Figure 6. Geometry-optimized structure models.](image)

(a) Hg(PC2)
(b) Hg(MT) \(\alpha\) domain
(c) Hg\(_3\)S\(_9\) \(\beta\)-HgS
(d) Hg\(_4\)S\(_{12}\) \(\beta\)-HgS
(e) Hg\(_4\)S\(_{11}\) \(\alpha\) domain
(f) Hg\(_3\)S\(_9\) \(\beta\)-HgS
(g) Hg\(_4\)S\(_{12}\) \(\beta\)-HgS
(h) Hg\(_4\)S\(_{10}\) \(\beta\)-HgS

Figure 6. Geometry-optimized structure models. (a, b) Hg oxothiolate complex in phytochelatine PC2 and the MT-\(\alpha\) domain. The amino acid sequence of the binding site is Cys\(_2\)GluCys(Cys\(_2\)Glu) for PC2 and CysSerGlyCys(CSGC) for MT-\(\alpha\). The two dithiolate sites have approximately the same dimension and therefore bind Hg similarly. Hg is bonded to the two cysteinyl S and O atoms forming a double six-membered bis-oxothiolate ring chelate (Hg(SR + O), coordination). The bis-chelate is further stabilized by an hydrogen bond between a carbonyl oxygen and an amide proton from the cysteine residues. The amine groups for Hg(MT-\(\alpha\)) were not protonated and an amide group has been added to the carboxyl terminus to make the complex neutral. Protons, other than the hydrogen-bonded amide proton, are not represented for clarity. Cartesian coordinates, and atomic charges calculated by natural population analysis (NPA\[40\]) are given in the Supporting Information. (c–e) Connectivity of the HgS\(_2\) tetrahedra from the HgS\(_2\) clusters in the \(\alpha\) and \(\beta\) domains of metallothionein optimized geometrically (MP2/TZVP-ecp). Cys residues from the metallothionein clusters were modeled with methanethiolate (CH\(_3\)S\(^@\)) (Figure S7, Supporting Information). (f–h) Connectivity of the HgS\(_2\) tetrahedra in metacinnabar (\(\beta\)-HgS).\[28\] Dark red, Hg; yellow, S; blue, N; red, O; gray, C.
plain the vast amount of tetrameric MT-40 in the two isolates (Figure 2). In addition, our results agree with the previous observation of Zn- and Cd-rich MT-20, and a Zn-rich MT-40 fraction, in the gel-permeation chromatograms of the cytosol extracts from mussels exposed to Cd.[14a]

The MT-20 and MT-40 oligomers have been observed previously in the Mytilidae family following exposure to Hg[6] and Cd[14ab, 17, 18, 34]. Results on the inducibility of MT expression suggest that MT-10 is specialized in the metabolism of essential Zn[6] and Cu(I/II) ions, whereas MT-20 participates in the detoxification of heavy metal cations and hydroxyl radicals.[35] This suggestion is supported, in particular, by the low concentration of MT-20 in mussels exposed to Cu, and its considerable enhancement in mussels exposed to Cd, and to a lesser extent to Hg.[15b, 35, 36] However, the finding that MT monomers would be involved in physiological functions and MT oligomers in biological effects of metal exposure is controversial,[37] as shown for example by the high Zn content and low Cd content of MT-40 in M. edulis exposed to Cd.[14b] This contradictory view is confirmed here by the detection of MT-20/30/40 in control mussels.

Structure of the dithiolate complex

Divalent mercury is most commonly linearly coordinated to two proximal cysteines (C) separated by two amino acids (X) in living cells.[38] The recurring CXXC motif is found in a wide variety of metalloproteins, including metallochaperones and metal-transporting ATPases. The three-dimensional structures of several Hg-bound proteins have been solved by NMR and X-ray crystallography, including MerP, MerA, and Atx1.[25ab] In these proteins, the dicysteinate metal complex (Hg[4] and Cu[4]) is stabilized by secondary bonding interactions, typically with oxygen atoms from side chains (e.g., from Thr and Ser) and water molecules, and with an extended H-bond network. In M. edulis, the CXXC motif is found at the end of the C domain, just before the short LysValVal linker connecting the two domains (Figure 1). Its sequence from the N-terminus is LysCysSerGlyCysLys (KCSGCK motif).

The geometry of the Hg[4] binding site was modeled computationally at the MP2[39] level of the molecular orbital theory (Figure 6b). The two cysteine residues coordinate the metal ion with their thiolate ligand at 2.31–2.32 Å and their carbonyl oxygen atoms at 2.61 and 2.97 Å. The predicted Hg–O distances are between the sums of the Hg and O covalent radii (1.98 Å) and the Hg and O van der Waals radii (3.07 Å). The closest oxygen atom bends the S–Hg–S angle to 170.8°, a value significantly higher than the 163–165° angular values for the Hg(Cys), and Hg(PC2) references (Figures 4 and 6a). The O ligands are almost perpendicular to the S–Hg–S bond direction, as reported previously for the NH[4] and SR sulphido/thioether secondary ligands.[24b] Bond analysis in terms of natural population analysis (NPA)[40] shows that a carbonyl oxygen is a little more nucleophilic (partial atomic charge −0.6 to −0.7 e) than an amide nitrogen from the -CO–NH- peptide bond (−0.6 e), and these two atoms are much less than an amine nitrogen (−0.9 e) (Figures S2 and S5, see Supporting Information).[12] Therefore, an amine group is more likely to bond Hg than a peptide bond. However, other factors besides the electronic charge of the donor are involved in the stability of a macromolecular complex, such as its conformation and hydrogen bonds. Regarding the chelate structure, the Hg(SR + O) coordination forms a double six-membered bis-oxothiolate ring. The two rings are linked by the two intervening residues Ser and Gly and the cage stabilized by one -NH–O–C- hydrogen bond between the amide group of one cysteine and the carbonyl oxygen of the other cysteine (Figure 6b). In the absence of Hg, the two SH groups protrude from the plane containing the CXXC motif by 1.8 Å like crab-claws (Figure S6, see Supporting Information). This claw setting of thiolate donors has the potential to adapt both geometrically and electronically to the linear coordination requirement of a mercuric ion, which it can capture with extremely high affinity.

The Hg complex in phytochelatine PC2 is also a Hg(6-S/O-ring), chelate (Figure 6a). In PC2, the binding motif is CXC and the intervening amino acid is γ-Glu. Its length is close to that of the SerCys dipeptide, therefore the loops of the PC2 and MT-α complexes have nearly the same size. Phytochelatins mediate metal detoxification in plants.[41–43] Their synthesis is also induced by Hg exposure,[44] but they are enzymatically synthesized from glutathione (GSH) by PC synthase, not transcriptionally induced from DNA like MTs. Still, plants can also express MTs genetically, and plant MTs also possess CXXC motifs like animal MTs.[43] Thus, the same Hg(6-S/O-ring), detoxification chelate appears to exist in the two kingdoms. However, plants have adopted it through two synthesis pathways, one catalyzed by enzymes (PC2) and the other from gene expression (MTs).

A metacinnabar-type (β-HgS) mineral core

The coordination geometry of the tetranuclear CdS[3]-α and trinuclear CdS[3]-β clusters is known for Cd-10 of Mediterranean mussel Mytilus galloprovincialis.[38] They consist of CdS2 tetrahedra connected through their apices by bridging cysteinyl sulfurs for a total of 21 Cys residues (Figure 1 and Figures 6c–e for the equivalent Hg species). In the Cd[4]-β cluster, each tetrahedron comprises two bridging (μ2-SR) and two terminal (μ-SC) thiolates, as commonly observed in animals and plants MTs (Figure 6c).[25a, 45–46] In contrast, in the Cd[4]-α cluster, one of the four tetrahedra has only one bridging thiolate, thus being attached to a single tetrahedron (Figure 6d). This configuration differs from the so-called boat-like conformations[26c] of the Cd[4]-α cluster in vertebrates which, unlike mussels, have 20 MT Cys residues in total (5α-α + 5α-β, Figure 6e). In Cd[4]-α, two tetrahedra are each linked to the other three tetrahedra from the four-metal cluster through three bridging thiolates, and the other two tetrahedra are linked to only two tetrahedra through two bridging thiolates. With its singly-bonded CdS, tetrahedron, the Cd[4]-α mussel cluster is therefore less compact than the Cd[4]-β vertebrate cluster.

The CdS[3]-α and CdS[3]-β clusters have the same tetrahedral association as β-HgS which features a spalactite-type (ZnS) lattice (Figures 6f and g). Thus, poorly crystalline β-HgS is a
good mineral proxy for the Hg$_{2}$-MT mussel structure on the local scale. Similarities between metal thiolate clusters and inorganic structures and complexes are also observed for Cu$^{+}$, Ag$^{+}$, Cd$^{2+}$, and Zn$^{2+}$.[29,40] As an aside, we note that the connectivity in Me$_{6}$S$_{11}$-$\alpha$ has no direct equivalent in ZnS/$\beta$-HgS, in contrast to Me$_{6}$S$_{11}$-$\alpha$ and Me$_{6}$S$_{11}$-$\beta$. In a tetranuclear $\beta$-HgS-type cluster, each tetrahedron shares three bridging sulfurs yielding an adamantane-type cage (Figure 6b). In Me$_{6}$S$_{11}$-$\alpha$, only two tetrahedra have three bridging sulfurs and the two others have two bridging sulfurs. The four tetrahedra and five bridging sulfurs of Me$_{6}$S$_{11}$-$\alpha$ form two fused six-membered rings ($2\times$[S-HgS-Hg-S-Hg]) with a distorted boat conformation (Figure 6e). This topology cannot be represented as a portion of the ZnS/$\beta$-HgS lattice that has only chair conformation. Therefore, the recurrent description in the literature of Me$_{6}$S$_{11}$-$\alpha$ in terms of an adamantane-type structure is incorrect, as pointed out previously.[29,64]

Structure of the mineral core

The geometry of the Hg$_{3}$S$_{12}$-$\alpha$ and Hg$_{3}$S$_{12}$-$\beta$ mussel clusters and the Hg$_{3}$S$_{12}$-$\alpha$ vertebrate cluster were optimized at the MP2 level using the connectivity of the Cd$_{3}$S$_{12}$-$\alpha$ and Cd$_{3}$S$_{12}$-$\beta$ clusters in M. galloprovincialis[38] as structural templates (Figure 1). The model thiolate was methanethiolate CH$_{3}$S$^{-}$, as in our previous computational studies.[11,46] This substitution marginally changes the geometry of a complex or the effective atomic charge on the sulfur and mercury atoms (Figure S7, Supporting Information). The three clusters have a predicted Hg–S bond length of 2.52 ± 0.04–0.08 Å, close to crystallographic values for inorganic Hg(SR)$_{3}$ complexes (2.566 ± 0.047 Å[47]) and well-crystallized $\beta$-HgS (2.53 Å[48]) (Figure 6f). The predicted Hg–Hg distances across the shared corners are 3.78 ± 0.08 Å for Hg$_{3}$S$_{12}$-$\beta$ and 4.09 ± 0.41 Å for Hg$_{3}$S$_{12}$-$\alpha$, compared to 4.14 Å in $\beta$-HgS. The Hg–Hg distances in Hg$_{3}$S$_{12}$-$\alpha$ (4.13 ± 0.19 Å) approach on average the Hg–Hg distance in $\beta$-HgS, but are extremely unequal. Although the Hg$_{3}$S$_{12}$-$\alpha$ and Hg$_{3}$S$_{12}$-$\beta$ clusters do not have the same polyhedral connectivity (Figures 6d and e), and thus the same degree of angular flexibility, this computational study shows that a tetranuclear cluster is more disordered than a trinuclear cluster. The EXAFS photoelectron wave of the Hg–Hg pairs for the Hg$_{3}$S$_{12}$-$\alpha$ cluster was calculated to determine if disorder ($\sigma = 0.19$ Å) could have prevented their detection with this technique by Jiang et al.[5] The large dispersion of the Hg–Hg distances leads to a wave beating at $k = \pi / (2 \Delta R) = \pi / (2 \times 0.16$ Å) = 10 Å$^{-1}$ and, as a result, to the extinction of the Hg-Hg peak on the radial distribution function (Figure S8, Supporting Information). The disordered $\beta$-HgS$_{np}$ reference used in HR-XANES analysis adequately accounts for the distribution of the Hg–Hg distances in MTs. However, being a solid phase $\beta$-HgS$_{np}$ features ideally twelve Hg–Hg pairs at 4.14 and six at 5.85 Å,[48] whereas these numbers are extremely small in nanosized MT clusters, even zero for the second Hg atomic shell at about 5.8 Å in Hg$_{3}$S$_{12}$-$\beta$. Approximating a MT cluster by the $\beta$-HgS$_{np}$ model manifests in the reconstruction of the MT spectra as a significant misfit near the edge maximum at 12300–12320 eV (arrows in Figure 5b), which is the region where $\beta$-HgS$_{np}$ exhibits a modulation of the absorption signal from distant Hg–Hg pairs (Figure 4d). These modulations are absent in the mussel spectra which in contrast have a bell-shape top edge, thus negating the presence of Hg neighbors at a medium distance. Based on XANES calculation,[37] the nuclearity of the MT Hg clusters is less than seven. A source of uncertainty is the effect of the peptide environment on the Hg coordination geometry because the enfolding of the two clusters by the polypeptide chain was not taken into account in our modeling as it would be too costly at the MP2 level of the molecular orbital theory.

Partitioning of the mineral core among the $\alpha$- and $\beta$-domains

Strong compelling evidence from X-ray diffraction, NMR spectroscopy, UV/pH titration, and circular dichroism (CD) spectroscopy, indicate that Cd$^{2+}$ and Zn$^{2+}$ are partitioned among the $\alpha$ and $\beta$ domains of mammals MTS, with Cd$^{2+}$ preferentially bound in the Me$_{6}$S$_{11}$-$\alpha$ cluster and Zn$^{2+}$ in the Me$_{6}$S$_{11}$-$\beta$ cluster:[11,36,70,21] Copper, which forms Cu$_{6}$S$_{11}$-$\alpha$ and Cu$_{6}$S$_{11}$-$\beta$ clusters in mammals,[49] also has a marked preference for the $\beta$ domain when it is mixed with Cd$^{2+}$.[50] With reference to their dynamic behavior, metal positions exchange much faster in the $\beta$ than the $\alpha$ domain.[21,53] The metal-binding selectivity and distinct chemical reactivity of the two domains leads to the hypothesis for a physiological specialization, with the $\beta$ domain having a role in metal homeostasis and the $\alpha$ domain having a role in metal detoxification.[21]

In keeping with the view that metal ions are bound with different affinities in each domain according to their chemical properties and biological functions, we speculate that the polynuclear Hg$_{3}$S$_{12}$ species identified by HR-XANES is Hg$_{3}$(Cys)$_{12}$-$\alpha$. This species would be located in the same domain, but clearly not the same MT molecules, as the Hg$_{3}$(Cys)$_{12}$ species. Mercury atoms have a marked propensity to aggregate through relativistic effects.[52] Mercuration, like auripigment and argentophilic interactions, occurs in solution at room temperature because of the similarity in magnitude of metallophilic interactions with hydrogen bonding.[53] A mercury atom bound to the two cysteines of the CXXC motif is therefore a nucleation center for the formation of Hg$_{3}$S$_{12}$ oligomers through mercuration, similar to the formation of $\beta$-HgS from Hg(SR)$_{3}$ complexes in natural organic matter.[48] Furthermore, the five CXC motifs from the $\alpha$ domain offer the correct spacing for the formation of the Hg$_{3}$S$_{12}$-$\alpha$ cluster, similarly to Cd$_{3}$S$_{12}$-$\alpha$[38] (Figure 1). The suggested nucleation mechanism has a parallel in the case of the Cu$_{6}$S$_{12}$ adamantane-type cluster formation in the human copper chaperone for the superoxide dismutase (CCS)[54] and the yeast copper transporter Ctr1,[55] which both involve CXC motifs. With its 3d$^{10}$ configuration, Cu$^{2+}$ also forms a linear complex with thiolate ligands. One could assert that Cu$^{2+}$ should also be bound preferentially to the $\alpha$ domain rather than the $\beta$ domain. This assertion is undermined by two observations. First, the stability constant of the linear Cu-CXXC complex is two orders of magnitude lower than that of the Hg-CXXC complex.[56] Second, the consensus CXC motif has the
highest metal selectivity for Cu because it is a nucleation center for highly stable Cu clusters.[54,55]

Thermodynamic calculation adds further support to the α domain occupation for the Hg₄S₄ species identified by HR-XANES. The Gibbs free energy (ΔG) of formation of multinuclear n[Hg₄(SR)₄] complexes from the condensation of bis-Hg-thiolate complexes Hg₄(SR)₄ increases with the nuclearity (n) of the cluster.[54,55] Thus, a tetranuclear Hg-α cluster is more stable than a trinuclear Hg-β cluster. The energetic penalty associated with an alteration of neutrality from four to three is probably more elevated for relativistic Hg atoms than non-relativistic Cd and Zn atoms, as corroborated by their co-occurrence in [Cd₄(S₂Cyst)₄]²⁺.[14,54] One could also advocate that the highly defective β-HgS₄⁰ reference, which displays broad X-ray diffraction peaks,[24b] better describes the Hg₄S₄-α cluster because it has more incoherent Hg-Hg distances than the Hg₄S₄-β cluster (Figure 6c). However, this interpretation is not unique because it is not possible by HR-XANES to differentiate a disorder effect from a particle size effect due to both effects producing a damping of the fine structure in the edge spectrum. In summary, we postulate that Hg is incorporated preferentially in the α domain, whereas the constitutive Cu and Zn metal ions are in the β domain in agreement with previous studies on [Cd₄S₄(Cu₂)]−[M₅S₈]−.[48d,50,57] The β domain is unlikely to be heteronuclear because the Cu₄S₄-β and Zn₄S₄-β clusters do not have the same structure, Cu being trihedrally coordinated[49] and Zn tetrahedrally coordinated.[54,48b] According to this scheme, Cu and Zn are bound to different MT molecules. The Cu-MT and Zn-MT molecules may hold a Hg₄S₄ cluster in their α domain, albeit with limited likelihood, given the low proportion of metalled MT (7%) relative to thionine (93%).

Conclusion

Although these findings illuminate significant aspects of Hg²⁺ detoxification in cells, they also give rise to challenging questions and suggest future research directions. Additional studies will be required to investigate the stability of the Me⁶⁺₄S₄-α cluster in bivalves (21 Cys) relative to the far more abundant Me⁴⁺₄S₄-α cluster (20 Cys) present in some invertebrates and all vertebrates, and to know if these differences endow functional differences among the two clusters with regard to metal detoxification.[54] Does the Me⁶⁺₄S₄-α core derive from the Me⁴⁺₄S₄-β core by attachment of a trigonal MeS₄ complex to one terminal cysteiny1 sulfur, as suggested by the conformational similarity of the two cores (Figures 6c and d)? Similarly, does the Me⁴⁺₄S₄-α core derive from the Me⁴⁺₄S₄-β core by attachment of a digonal MeS₄ complex to two terminal cysteiny1 sulfur (Figures 6c and e)? Bridging thiolates are an important factor for the high thermodynamic stability of metal thiolate clusters, and a correlation has been suggested between the ratio of bridging (µ₂-SR) to terminal (µ-SR) thiolates and the stability of a cluster.[55] With a µ₃-µ₄ ratio of 5:6, the Me⁶⁺₄S₄-α cluster would be more stable than the Me⁴⁺₄S₄-α cluster, which has a ratio of 4:8. Under these premises, did evolution select the more closely packed and thermodynamically more stable Me⁶⁺₄S₄-α structure relative to Me⁴⁺₄S₄-α? Structure likely controls stability over the number of cysteine residues. If the Me⁶⁺₄S₄-α cluster structure confers superior resistance to metal toxicity, does it mean that bivalves are more sensitive to Hg²⁺ intoxication than other animals?

How does MT-40 form? Although circumstantial evidence exists for its presence in vivo, an aggregation of the MT-10 units during purification cannot be totally dismissed. What is the amino acid sequence of the MT-10 units (primary structure) and how are they covalently bonded in MT-40 (quaternary structure)? Do the monomeric units from MT-40 contain 21 cysteines, as in canonical MT-10,[56] or 23 cysteines, as in the monomeric MT-10 units of MT-20?[14,55] Lastly, are the biogenic MT clusters more stable than the metallothionein-like β-HgS₄ nanoparticles formed naturally in natural organic materials[24b,48d,60] and plant leaves?[21] All these questions could be addressed similarly to 4d¹⁰ Ag⁰ and 5d¹⁰ Au⁰, which, like 5d¹⁰ Hg⁰, form distorted thiolate clusters in metalloproteins and linear coordination with thiolate ligands when they are embedded in sterically complex and demanding macromolecules. All these important biometals and intriguing questions are now leading to investigations in living matter by HR-XANES down to, and below, the ppm level.[50]

Experimental Section

Materials: Wild blue mussels (Mytilus edulis) were collected from the Charente–Maritime (France) coastline in March 2016. They were cleaned from fouling and acclimatized to laboratory conditions for two weeks in a 150 L aquarium filtered with filtered seawater. Conditions were adjusted to be as close as possible to those in the marine environment: salinity = 35 p.s.u., temperature = 15 ± 0.3 °C, pH 8.1 ± 0.1, light/dark = 12 h/12 h. During the acclimatization and experimental periods, mussels were fed every two or three days with a mix of six marine microalgae (Shellfish Ephyd 1800°). Considering that body size (age) affects metal bioaccumulation in marine organisms, only individuals with homogeneous size were used in the experiment. After the acclimatization period, 60 mussels were equally distributed in two 50 L aquaria, one as control and one for exposure to 100 µg Hg L⁻¹ for 9 days. A stock solution of mercury nitrate at 1 g Hg L⁻¹ was used as the source of mercury (ASTASOL®). Addition of Hg(NO₃)₂ did not change pH value nor salinity. Concentrations of dissolved Hg in the two aquaria were monitored daily by Hg analyses with an Altec AMA 254 spectrophotometer. Thus, the Hg concentration in the experiment tank was readjusted as needed to 100 μg Hg L⁻¹ to compensate losses resulting from bioaccumulation by the mussels and adsorption on the aquaria walls. At the end of the exposure period, mussels were rapidly dissected, frozen with liquid nitrogen and maintained at −20 °C until further analyses. No mortality was observed during our experiment.

Metallothionein purification: Mussels were defrosted and dried on filter paper. MTs being contained in gills, digestive gland, kidney and mantle, the whole mussel tissues were crudely homogenized in a mixer at 4 °C and pH 8.1 in 100 mmol L⁻¹ Tris buffer, then finely mixed in a glass-Teflon® homogenizer. The homogenate was centrifuged at 20000 g for 10 min. The supernatant was decanted with an argon flow, not under gas bubbling to avoid protein denaturation. The Ar saturated solution was gently stirred with a magnetic bar and heated at 95 °C for 15 min under a continuous flow of Ar after previously described procedure.[53,54] Protein denaturation of the enzymes was monitored by UV-Visible spectrophotometry. Enzymes were stabilized at 20 °C in 50 mmol L⁻¹ Tris-HCl, pH 8.1, and their stock solution was used immediately.

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turation at 95 °C for 15 min was considered a suitable trade-off between the extent of MT purification and the amount of remaining MT material needed for spectroscopy (3.4 ppm Hg in 300 µL extract). Thermolabile proteins and lipids precipitated whereas heat stable MTs remained soluble. A control experiment was performed in which the MT was purified in aerated conditions on an air-equilibrated homogenate. After thermal denaturation, the glass vessel was rapidly cooled in ice and the soluble fraction was separated by centrifugation at 30 000 g for 20 min. 5–10 mL of the MT solution was dialyzed against 5 L of pure water in Spectra/Por¨ dialysis tubing of 3.5 kDa molecular weight cut-off. Dialysis was repeated four times every three hours to eliminate salts (mussels were in seawater and Tris buffer was used for the homogenization step), and metals (Cu, Zn, and Hg) which were either free or complexed to small organic molecules, such as cysteine and glutathione, and which may not have precipitated during the heating step. After the dialysis step, all the Hg atoms were bound to metallothioneins. Protein concentration was determined by the Bradford’s method using bovine serum albumin as reference. The Bradford reagent reacts mainly with NH₄⁺ groups, and their content is different in BSA and MT. No correction was applied, however, because the reactivity of the NH₄⁺ groups in the two proteins is unknown and is another source of uncertainty in quantification. The quality control of the protein extract was assessed by dynamic light scattering (DLS). Less than 6% of the protein was in aggregate form. Purity and oligomerization state of the MTs were examined by SDS-PAGE electrophoresis. To this end, the MT solution was concentrated about ten times using Spin-X UF ultracentrifugation concentrators of 5 kDa molecular weight cut-off (Corning). One aliquot of 5 µL and another of 10 µL were mixed with denaturing buffer (BIO-RAD) and heated at 90 °C for 5 minutes. Electrophoresis migration was performed at 180 V for 30 min on Criterion XT (4–12 %) precast gel (BIO-RAD) with 50 mm MES running buffer (pH 6.5). The protein bands were visualized with Coomassie Brilliant Blue.

Metal analysis: Hg, Cu, and Zn concentrations in the control and MT extract were measured by quadrupole ICP-MS (ThermoFisher iCAPQ) at masses 200 and 202, 63, and 64, respectively, after calibration with SPEXCertiprep standards. Analyses were carried out at low concentration to avoid a memory effect and in triplicate with appropriate blank subtraction.

Complex Hg(SR)₂Cr₃X₄: The formulation of the new crystal is [Hg₂(C₆H₅NO₅)₂][SO₄]₂·4H₂O and its structure is represented in Figure S9 (see Supporting Information). Mercury nitrate (Hg(NO₃)₂·H₂O, 238 mg, 0.695 mmol) was dissolved in methanol (6 mL) and added dropwise to 4-mercaptopopyridine (CAS 4556-23-4, 95% purity, 300 mg, 2.56 mmol) in a solution of methanol (15 mL), distilled water (15 mL), and ammonium tetrafluoroborate (CAS 13826-83-0, 99% purity, 210 mg, 1.98 mmol). Then 0.1 M sulfuric acid was added dropwise to reach a pH of 1.0. After 24 h of slow evaporation of the light yellow homogenous solution at room temperature in the dark, a fine yellowish precipitate formed with yellow and colorless crystals. The crystals were isolated by filtration and washed with distilled water. The crystalline quality of the yellow crystals was too poor for structural determination. A colorless single crystal (0.22 x 0.11 x 0.06 mm) was mounted on a Bruker Kappa CCD diffractometer using monochromatic Ag Kα radiation (λ = 0.65867 Å). Crystal data at 293 K: C₅₂H₅₆Hg₂O₄·3.25H₂O, M = 1332.32, space group P-1 (no. 2), a = 14.018(2) Å, b = 14.320(1) Å, c = 15.556(1) Å, α = 77.41(1)°, β = 74.13(1)°, γ = 79.04(1)°, V = 2157.4(7) Å³, Z = 2, Dc = 2.05 g cm⁻³, μ = 41 cm⁻¹, 48326 measured reflections, 9784 unique reflections (Rint = 0.10), R1 = 0.06 (with 6115 > 2σ(I) and wR2 = 0.14, 548 parameters refined, GOF = 1.07, max. min. residual peaks 1.58/−1.77 e Å⁻³. Data were corrected for Lorentz and polarization effects and empirical absorption (SADABS from Bruker/Siemens). The crystal structure was solved by direct methods with the SIR92[11] program and refined by full matrix least-squares, based on F², using the SHELXL[12] software through the WinGX[65] program suite. The refinement was performed with anisotropic thermal parameters for all non-hydrogen atoms. Only (N–H) hydrogen atoms of mercaptopyridinium were localized on difference Fourier map and refined with isotropic thermal parameters. The other hydrogen atoms were generated at idealized positions, riding on the carbon carrier atoms, with isotropic thermal parameters. Hydrogen atoms of water molecules were not localized.

CCDC 1838728 contains the supplementary crystallographic data for this paper. These data are provided free of charge by The Cambridge Crystallographic Data Centre.

HR-XANES spectroscopy: All Hg L₃-edge HR-XANES spectra were measured in high energy-resolution fluorescence yield detection mode with high-luminosity analyzer crystals[40] on beamline ID26 at the European Synchrotron Radiation Facility (ESRF). Three hundred microliters of the concentrated MT extract (MT sample, [Hg] = 3.4 ppm) were placed in a polyether ether ketone (PEEK) holder designed for solutions, and the holder was frozen immediately in liquid nitrogen (LN) and stored in a LN, Dewar until its transfer into the liquid helium cryostat of the ID26. Tissues of whole mussel (ME sample) were freeze-dried to concentrate the sample, placed in a PEEK holder designed for pellets, sealed with poly (4,4’-oxydiphenylene-pyromellitimide) (Kapton®) tape, and stored in a desiccator until transfer into the liquid helium cryostat of the beamline. We checked that the process of freeze-drying a frozen tissue and preparing a pellet did not alter the Hg²⁺ speciation itself by comparing the HR-XANES spectrum of an as-prepared fish sample to the spectrum of a snap-frozen hydrated sample from the same fish. The two data are statistically identical (Figure S10, Supporting Information), in agreement with previous report.[66] Spectra were collected at a temperature of 10–15 K and a scan time of 15 s to reduce exposure, and repeated at different pristine positions on the sample to increase the signal-to-noise ratio. Scans were monitored carefully for any evidence of radiation damage. The incident energy was scanned from 12260 to 12360 eV in 0.2 eV steps and the spectra were normalized to unity at E = 12360 eV. More information on the beamline optics and data acquisition is given in the Supporting Information.

Geometry optimization: Conformations were energy-optimized at the MP2 level with the resolution of identity (RI) approximation using the def2-TZVP basis set and the def2-TZVP/C auxiliary basis set for S, C, and H, and the def2-TZVP-eCP basis set for Hg. ORCA 3.0.3[67] was used for all calculations. Geometries were optimized in the aequous phase with the implicit solvation method COSMO.[66] Thermal energy corrections to the free energy were also calculated at the RI-MP2 level for the Hg₃(SMe)₉ and Hg₉(SMe)₉ clusters (Figure 6) to verify that the conformations corresponded to minima (i.e., no negative/imaginary frequencies). The Hg₉(SMe)₉ frequency calculation was too costly in terms of computation time to be performed. This computational Scheme has been tested previously on the modeling of the structure and stability of mononuclear and multinuclear Hg-thiolate complexes.[24,46]

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Conflict of interest

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