Mercury cycling between the water column and surface sediments in a contaminated area

Elsa Ramalhosa\textsuperscript{a,*}, Susana Ríó Segade\textsuperscript{b}, Eduarda Pereira\textsuperscript{c}, Carlos Vale\textsuperscript{d}, Armando Duarte\textsuperscript{c}

\textsuperscript{a}CIMO, Escola Superior Agrária de Bragança, Instituto Politécnico de Bragança, 5300 Bragança, Portugal
\textsuperscript{b}Estación de Viticultura e Enoloxía de Galicia (EVEGA), Ponte San Clodio s/n, Leiro, 32427-Ourense, Spain
\textsuperscript{c}Departamento de Química, Universidade de Aveiro, 3810-193 Aveiro, Portugal
\textsuperscript{d}Instituto de Investigación das Pescas e do Mar (IPIMAR), Av. Brasília, 1400 Lisboa, Portugal

\textbf{Abstract}

Mercury cycling in the water column and upper sediments of a contaminated area, the Largo do Laranjo, Aveiro (Portugal), was evaluated after determination of reactive and non-reactive mercury concentrations in the water column and pore waters of sediments, collected in several places of this bay. In the water column, reactive mercury concentrations varied between 10 and 37 pmol dm\textsuperscript{-3}, the highest values being observed near the mercury anthropogenic source. However, reactive mercury was a narrowly constrained fraction of the total mercury, making up only 4–16% of the total, showing evidence of the importance of dissolved organic matter on mercury transport.

In sediments, higher concentrations of mercury were also determined near industrial discharges. Results indicate the existence of an equilibrium between solid and liquid phases, determined by solid sediment/pore water distribution coefficients. Much of the mercury present in the solid fraction is associated with organic matter ($r = 0.837$) and iron oxyhydroxides ($r = 0.919$), but as oxides begin to dissolve in reduced sediments and organic matter decays, the adsorbed mercury is released. In fact, the mercury concentrations in pore waters of those contaminated sediments largely exceeded the values determined in the water column. As molecular diffusion may contribute to the dissolved mercury distribution in the overlying water column, this phenomenon was evaluated. However, the pore waters of Largo do Laranjo do not enrich the water column substantially in terms of reactive and non-reactive mercury. In fact, pore waters can contribute only to 0.2% and 0.5% of the reactive and non-reactive mercury present in the water column, respectively, showing that as long as mercury is being incorporated in sediments, it stays in stable forms.

\begin{thebibliography}{10}

\textsuperscript{a}Corresponding author. Tel.: +351 273 303 262; fax: +351 273 325 405.
E-mail address: elsa@ipb.pt (E. Ramalhosa).

1. Introduction

The fate and reactivity of anthropogenic mercury in the aquatic environment are of major concern owing to their toxicity for most organisms and humans. Trace metals are particle reactive (Paraquetti et al., 2004) and quickly sorbed to the suspended particulate matter (SPM) (Kennish, 1998) or to the particulates in the sediment–water interface. Once they reach the sediment, the major fraction of anthropogenic mercury is ultimately retained (Kennish, 1998). In many situations, the vertical profiles of total mercury in sediments record the historical contamination of the area under study.
However, the coastal sediments are highly dynamic and the conditions existing in the upper layers influence the sediment–water exchanges and have a strong effect on the distribution of trace elements (Sundby, 1994). The remobilisation due to the resuspension of metal enriched on fine bed sediments may increase the dissolved metal content in the overlying water column (Martino et al., 2002), as well as maintain large amounts of particulate mercury (Lacerda and Gonçalves, 2001). The types and the extension of the reactions that take place between the solid sediments and the aqueous phase can be evaluated by the pore water composition (Hong et al., 1994). This may be enriched in mercury in comparison to the water column (Cossa and Gobeil, 2000). This mercury present in the pore waters may escape to the water column and deleteriously affect the biota.

Since decades, a chlor-alkali plant has been discharging mercury into one of the most remote branches of Ria de Aveiro, which is a coastal lagoon with a permanent connection to the sea. Most of the discharged mercury settles in Largo do Laranjo, an inner bay of 1.5 km². Since the 1980s, high mercury concentrations in sediments, SPM and water have been detected and reported in several studies (Lucas et al., 1986, Pereira et al., 1998ab, Ramalhosa et al., 2001a). Generally, the highest mercury concentrations in sediments (maximum 35 μg g⁻¹) were found at 30–40 cm depth, corresponding to the period of maximum industrial production (Pereira et al., 1998a). However, particulate mercury escapes when the surface sediments are resuspended and transported by the tide to the rest of the lagoon (Pereira et al., 1998b). The contribution of the dissolved mercury was not evaluated due to the low concentrations found (Pereira et al., 1998b). In posterior works, the pore water profiles indicated that mercury is being remobilised in the anoxic sediments and these may constitute internal sources of mercury to the water column (Ramalhosa et al., 2001a).

This study presents the distributions of total and reactive mercury in solid sediments, pore waters and water column, as well as iron and organic matter, in Largo do Laranjo. On the basis of these data, associations of mercury in sediments and water column are evaluated and mercury cycling between both compartments is assessed by considering molecular diffusion.

2. Materials and methods

2.1. Sampling

Surface sediments (0-0.5 cm layers) were collected with a grab sampler at 20 sites in Largo do Laranjo contaminated area in February 2000 (Fig. 1). The fluvial flow rate was high in this figure.
month due to rainfall and to the deviation of a significant part of Antuá River flow rate to Esteiro de Estarreja, originating high volumes of circulating water. The water samples were collected at five sites (1, 2, 3, 4 and 5) by hand, using polyethylene gloves and glass bottles of 500 cm$^3$, during low and high tides. All material was previously acid cleaned, following the procedure of Diego et al. (1998). The samples were kept refrigerated and immediately transported to the laboratory.

2.2. Measurements in loco

Several measurements were performed in loco. In the water column, temperature, salinity (WTW, LF 196), pH (Orion, model 720 A) and dissolved oxygen (WTW, Oxi 96A) were measured. On the other hand, pH (WTW, model 320) and redox potential (Mettler Pt 4805/S7, with an Ag/AgCl reference electrode) were determined in the surface sediments.

2.3. Sample treatment

All sediment samples were treated in a nitrogen-filled glove bag to prevent oxidation reactions. The pore water samples were extracted from sediments by centrifugation at 6000 rpm for 20 min (B. Braun, model Sigma 4–10). A portion of the supernatant liquid was filtered through Millipore 0.45 μm porosity filters. The filtrate was collected in acid-cleaned bottles, preserved by acidification to pH<2 with nitric acid (mercury free) and placed at 4°C. Another portion of the supernatant liquid was filtered through Gelman glass fibre filters to evaluate the dissolved organic matter. Previously, the filters were cleaned by heating at 450°C for 24 h. All the water samples were treated in the same way as the pore waters. Identical volumes of ultrapure water were carried through the centrifugation and filtration stages as a blank procedure. Portions of all sediments were freeze dried (Christ, model Alpha 1–4), homogenised and sieved through 1 mm and 63 μm stainless-steel sieves (Fritsch, model Analysette 3).

2.4. Mercury and organic matter determinations in the pore waters and water column dissolved fraction

Both filtered waters were analysed for reactive mercury by cold vapour atomic fluorescence spectrometry (PSA cold vapour generator, model 10.003, associated with a Merlin PSA detector, model 10.023), using SnCl$_2$ as a reducing agent. This mercury fraction includes mostly ionic (Hg$^{2+}$) plus dissolved gaseous mercury, but also one that is weakly bound to inorganic and organic dissolved complexes (Fitzgerald and Mason, 1997). The dissolved mercury detection limit was calculated from 44 calibration curves in the concentration range 10–100 pmol dm$^{-3}$, varying from 1.1 to 8.9 pmol dm$^{-3}$. The detection limit was calculated as the intercept plus three times the standard deviation of the signal. The relative standard deviation ($n=5$) for a 100 pmol dm$^{-3}$ inorganic mercury standard was less than 5%. No peak was detected when the blanks were analysed. The total dissolved mercury concentrations were determined after oxidation with potassium persulphate and ultraviolet radiation, as described by Mucci et al. (1995). Non-reactive mercury was determined by the difference between total dissolved mercury and reactive mercury, and included all dissolved mercury species strongly bound to soluble complexes (Gill and Bruland, 1990). The dissolved organic matter concentration was determined using persulphate/ultraviolet oxidation.

2.5. Mercury, iron and organic matter determinations in the solid fraction of sediments and SPM

The sediments solid fraction was analysed for total mercury, methylmercury, organic matter and hydroxylamine-extractable iron. The total mercury concentrations were determined following the acid digestion procedure reported by Pereira et al. (1998a). The accuracy and precision of the method were evaluated using an estuarine-certified reference material (BCR 277). The determined value 1.73±0.03 μg g$^{-1}$ (mean ± confidence limit at $z=0.05$; $n=10$) was in good agreement (t-test and F-test at $z=0.05$) with the certified value of 1.77±0.06 μg g$^{-1}$ (mean ± confidence limit at $z=0.05$). Methylmercury concentration was determined by the procedure described by Ramalhosa et al. (2001b). Briefly, the methylmercury extraction from sediments is based on a microwave-assisted alkaline digestion with methanolic potassium hydroxide (0.15 g of sediment was suspended into 6 cm$^3$ of 25% methanolic potassium hydroxide and the slurry was exposed to microwave irradiation at 84 W for 2 min). Organomercury species were extracted with dichloromethane in hydrochloric acid medium and back-extracted into ultrapure water. The sediment extracts were injected into an analytical system composed of high-performance liquid chromatography–ultraviolet post-column oxidation–cold vapour atomic fluorescence spectrometry (HPLC–UV-PCO–CV-AFS). The detection limit of this method was 12 ng g$^{-1}$, calculated as the intercept plus three times the standard deviation of the signal. Two sediment reference materials (BCR 580 and IAEA 356), containing different amounts of methylmercury, were analysed to evaluate the accuracy and precision of the method. The analysis of samples with low methylmercury contents required the treatment of several 0.15 g portions and mixing of the dichloromethane extracts obtained for each portion. The determined methylmercury concentrations in both reference materials (BCR 580: 69.5±8.8 ng g$^{-1}$; IAEA 356: 5.53±0.21 ng g$^{-1}$; mean ± confidence limit at $z=0.05$, $n=3$ for each) were in good agreement (t-test and F-test at $z=0.05$) with the certified values (BCR 580: 75.5±3.7 ng g$^{-1}$; IAEA 356: 5.46±0.39 ng g$^{-1}$; mean ± confidence limit at $z=0.05$). The organic matter in sediments was determined by elemental analysis (LECO, model CHNS-392). Iron oxyhydroxides were determined after extraction with hydroxylamine: 0.5 g of sediment was treated with 20 cm$^3$ of 0.1 mol dm$^{-3}$ hydroxylamine solution at pH 2 (obtained by adding acetic acid) for 16 h. The suspension was filtered through Millipore 0.45 μm porosity filters and the filtrate was analysed for iron by flame atomic absorption spectrometry (Perkin Elmer, Aanalyst 100).

SPM was analysed for total mercury and particulate organic carbon (POC). The total mercury concentrations were determined following the procedure described for the sediments solid fraction. POC was determined after potassium dichromate oxidation in presence of sulphuric acid, as suggested by Kramer et al. (1994).
3. Results

3.1. Water column

During the surveys, the Largo do Laranjo water column was oxygenated (51–96% saturation) with pH values smaller at low tide ($7.3 \pm 0.7$) than at high tide ($7.6 \pm 0.1$), and statistically ($\alpha = 0.05$) higher dissolved organic carbon concentrations at low tide ($222 \pm 21\mu\text{mol dm}^{-3}$) than at high tide ($170 \pm 19\mu\text{mol dm}^{-3}$). The reactive mercury concentrations varied between 10 and $37\text{pmol dm}^{-3}$, accounting for 4–16% of total dissolved mercury. At low tide, enhanced reactive mercury contents were found at sites 1, 2 and 3, located close to the industrial effluent. At site 4, similar reactive mercury concentrations were observed at low and high tides, and higher concentrations were observed at site 5 at high tide than at low tide.

In the particulate fraction, SPM showed a similar trend at both tidal conditions, varying in the concentration range $37 + 14\text{mg dm}^{-3}$, as shown in Fig. 2. However, SPM concentration decreased significantly only at low tide, in the farthest place from industrial effluent (site 5). POC showed a different behaviour in comparison with SPM. At low tide, POC concentrations varied slightly ($5.4 \pm 0.3\%$), while they decreased along Largo do Laranjo towards the sea ($5.3 \pm 1.4\%$) at high tide, again with exception of site 5. In terms of total particulate mercury, the highest concentrations were determined at low tide, decreasing from 95 to $19\text{nmol g}^{-1}$ along estuary. At high tide, the total mercury concentrations varied between 8 and $23\text{nmol g}^{-1}$.

3.2. Sediments

The proportion of fine particles (<63 µm), present in the collected sediments, varied between 8% and 61% in the vicinity of the industrial discharge. The pH of the surface sediments ranged between 6.4 and 7.0, decreasing in the sites closest to the effluent. The redox potential irregularly ranged between 8 and 292 mV. The organic carbon content and the hydroxylamine extractable iron in the sediments varied within broad ranges, $0.13–4.6\text{mmol g}^{-1}$ and $5.2–50\text{m mol g}^{-1}$, respectively. The lowest values were found in the samples where sand was visually detected. The organic matter content is linearly correlated ($r^2 = 0.62$) with the proportion of fine particles.

The spatial distribution of total mercury concentrations in sediments is presented in Fig. 3. The highest concentrations (>70 nmol g$^{-1}$) were recorded in the sediments deposited near the industrial discharge and the lowest values (minimum of $0.73\text{nmol g}^{-1}$) were found in the sands of the bay outlet. The total mercury distribution is exponentially related with the fine fraction of the sediments ($r^2 = 0.616$). Methyl-mercury was detected only in the sites closest to the industrial discharge ($61\text{pmol g}^{-1}$, <0.1% of total mercury). The reactive mercury concentrations were higher in the pore waters surface sediments ($17–188\text{pmol dm}^{-3}$) than in the water column ($10–37\text{pmol dm}^{-3}$). Reactive mercury accounts for 3.7–21% of the total dissolved mercury present in the pore waters.

4. Discussion

4.1. Mercury in water column

In Largo do Laranjo, the highest reactive mercury concentrations were found near the industrial discharge and at low tide, due to lower flow rates and the smaller seawater contribution. However, exceptions were observed (site 5), where the highest concentrations corresponded to the high tide, probably due to a higher resuspension of surface sediments originated by the highest flow rates, increasing the SPM from 22 to 45 mg dm$^{-3}$. This resuspension can induce the transfer of mercury associated with the particles to the water column dissolved phase by competition of chloride ions for mercury and/or desorption of mercury from particle’s surface because of magnesium ions present in seawater.

Reactive mercury represents a small fraction (4–16%) of total dissolved mercury, suggesting that the majority of total dissolved mercury is strongly bound to dissolved organic complexes. At low tide, when the dissolved organic carbon content was higher, similar patterns of dissolved organic carbon, total mercury and reactive mercury were observed, as stated in Fig. 4, indicating the importance of this component on mercury transport. However, this role is less pronounced.
at high tide, probably due to the lower dissolved organic carbon concentrations and higher flow rates that alter the existent equilibriums.

In the particulate phase of the water column, total mercury did not show any significant relationship with SPM or POC.

In order to evaluate the contributions of the dissolved and particulate phases on mercury behaviour, distribution coefficients ($K_D$) were calculated for both tidal conditions. At low tide, $K_D$ values were higher than at high tide, indicating that 71–94% of total mercury is associated with particles. At high tide, total mercury was less associated with particles (56–80%), showing that sometimes the dissolved phase is also an important vehicle for mercury. This situation can be more deleterious for biota and must be taken into account when ways of action have to be decided to resolve the mercury contamination problem in Largo do Laranjo.

### 4.2. Mercury in sediments

The surface sediments collected in this study presented mercury concentrations comparable to the values recorded in a previous work (Pereira et al., 1998a). In the present study, the highest mercury contents were found in the area near the industrial discharge and were positively correlated to both the organic carbon ($r = 0.837$) and the iron oxyhydroxides ($r = 0.919$) contents. The organic carbon present in Largo do Laranjo sediments may result from industrial effluents and also from the decomposition of extensive algae coverage and plankton inside the bay, as stated by Monterroso et al. (2003). The triangular representation of total mercury, iron and organic carbon concentrations shows that mercury is more closely associated with iron oxides than with organic matter (Fig. 5). Although it is expected that mercury is strongly linked
to iron oxide forms, only few studies have shown the relationship between mercury and iron oxides in the field (Matty and Long, 1995).

4.2.1. Methylmercury production
In Largo do Laranjo, methylmercury was detected only in one sample and its concentration represented less than 0.1% of total mercury. This local showed a small redox potential (22 mV), indicating that mercury methylation is more favoured in suboxic conditions than in the presence of oxygen, as mentioned by Jackson (1998).

4.2.2. Mercury in pore waters
The organic ligands present in pore waters seem to play an important role, because a positive correlation ($r = 0.690$) was observed between reactive mercury and dissolved organic carbon (Fig. 6). A poorer correlation was obtained for total dissolved mercury ($r = 0.452$). The metal interactions between solid and liquid phases were evaluated by the distribution coefficient ($K_D$, [nmol Hg g$^{-1}$]/[nmol Hg dm$^{-3}$]) (Benoit et al., 1999), calculated as the ratio between molar concentrations of total mercury in solid sediments and of reactive mercury in pore water, and ranged from 14 to $24 \times 10^2$ dm$^3$ g$^{-1}$. Besides, the $K_D$ values were positively correlated to organic carbon content in solids ($r = 0.834$) (Fig. 7), reinforcing the importance of organic matter on mercury partition in sediments.

4.2.3. Estimation of mercury export from the pore waters to the water column
The diffusion first Fick law allows the estimation of the diffusive fluxes ($J$) across water–sediment interface, by constructing a linear gradient of the solute concentration ($C$) in pore water at the 0–0.5 cm layer ($Z$):

$$J = -\phi D_{sed} \frac{\partial C}{\partial Z},$$

where $\phi$ is the porosity calculated as the ratio between water volume and the volume of the corresponding sediment portion, $D_{sed}$ is the whole sediment diffusion coefficient defined as $D_{sed} = D_{sea}\times/\phi^2$, $\theta$ is the tortuosity estimated by the empirical relationship $\theta = 1 - \ln \phi^2$ (Boudreau, 1999) and $(\partial C/\partial Z)$ is the concentration gradient across the sediment–water interface. In Largo do Laranjo, the $D_{sed}$ varied between $1.8 \times 10^{-10}$ and $3.4 \times 10^{-10}$ m$^2$ s$^{-1}$, which is in accordance with the general value of $10^{-10}$ m$^2$ s$^{-1}$ for heavy metal ions and complexes (Li and Gregory, 1974).

The diffusion fluxes were determined for reactive and non-reactive mercury. The mercury concentrations were determined in the water column after considering the existence of a linear gradient of mercury concentrations from low to high tide. After dividing Largo do Laranjo in four zones (Fig. 1), the estimated fluxes for reactive mercury ranged between $0.013$ and $0.67$ nmol Hg m$^{-2}$ (12 h)$^{-1}$, as represented in Fig. 8, the highest values being found in zones A and B due to mercury-enriched pore waters. The reactive mercury fluxes were a small fraction of the total mercury fluxes, however, they can be significant in zones far away from the discharge point, representing almost 30% of the total. In terms of non-reactive mercury, the estimated fluxes ranged between 0.11 and 15 nmol Hg m$^{-2}$ (12 h)$^{-1}$, these fluxes being an order of magnitude higher than the ones related to reactive mercury. However, both mercury species show a similar behaviour.
Mercury fluxes were not determined in locals 14, 15, 16 and 17, because these locals were not strictly important, not being subjected to direct mercury discharge.

In order to evaluate the contribution of pore waters to the enrichment of the overlying water column, mercury was calculated in this compartment by the product of the mercury concentration determined in the water column (nmol dm$^{-3}$) and water volume (dm$^3$) per area unit (m$^2$). The water volume was determined taking into account the variation in the water column height observed in a tidal cycle. This varies between 1.0 and 1.5 m at low and high tides, respectively. These calculations were carried out for reactive and non-reactive mercury, as shown in Table 1. The former varies between 150 and 396 nmol m$^{-2}$ (12h$^{-1}$), the highest values being found in zones A, B and C, all of them with the same order of magnitude. In terms of non-reactive mercury, similar values were determined in all areas, varying between 2.2 and 3.4 μmol m$^{-2}$ (12h$^{-1}$).

Taking into account these values, in Largo do Laranjo, the pore waters contribute only 0.2% to the enrichment of the overlying water with respect to reactive mercury. In terms of non-reactive mercury, pore waters can have a more significant role. However, the contribution does not exceed 0.5%, reinforcing the idea that mercury in Largo do Laranjo surface

![Fig. 8 – Estimation of the reactive and non-reactive mercury fluxes (nmol m$^{-2}$ (12h$^{-1}$)) from the pore waters to the water column, as well as the percentage of the reactive mercury flux, determined for the various sampling sites located on Largo do Laranjo.](image)

<table>
<thead>
<tr>
<th>Zone</th>
<th>Mercury present in water column</th>
<th>Contribution of pore waters to enrichment of water column</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Reactive Hg (nmol m$^{-2}$ (12h$^{-1}$))</td>
<td>Reactive Hg (%)</td>
</tr>
<tr>
<td></td>
<td>Non-reactive Hg (μmol m$^{-2}$ (12h$^{-1}$))</td>
<td>Non-reactive Hg (%)</td>
</tr>
<tr>
<td>Zone A</td>
<td>394</td>
<td>0.016-0.17</td>
</tr>
<tr>
<td>Zone B</td>
<td>396</td>
<td>0.006-0.12</td>
</tr>
<tr>
<td>Zone C</td>
<td>318</td>
<td>0.000-0.012</td>
</tr>
<tr>
<td>Zone D</td>
<td>150</td>
<td>0.027-0.030</td>
</tr>
</tbody>
</table>

Table 1 – Quantities of reactive and non-reactive mercury (nmol m$^{-2}$ (12h$^{-1}$) and μmol m$^{-2}$ (12h$^{-1}$), respectively) present in the Largo do Laranjo water column and estimation of the pore waters contribution to the enrichment of the water column (%).
sediments is in a stable form, limiting its flux to the water column.

5. Conclusions

The results of this study show that mercury concentrations in upper sediments and the water column were significant, in spite of reductions in the emissions in the last years in Largo do Laranjo. The incorporation of anthropogenic mercury in sediments is not a simple function of distance to the source, but depends on sediment characteristics.

In the upper layers of sediment, mercury is retained by iron oxyhydroxides, as well as organic matter. However, it can be released to the pore waters due to the iron oxyhydroxides reduction and the organic matter decay, causing an equilibrium between the solid and liquid phases.

As the mercury concentrations in pore waters largely exceed the values encountered in the water column, mercury remobilisation from sediments to water column can occur. However, the reactive and non-reactive mercury present in pore waters can contribute only 0.2% and 0.5% to the water column enrichment, respectively. Moreover, the mercury diffusion from pore waters is more significant in the areas near the mercury source.

In overall conclusion, the mercury diffusion from pore waters does not contribute significantly to the dissolved mercury distribution in the overlying water column in Largo do Laranjo, demonstrating that the mercury cycling between the water column and the surface sediments is not significant.

References


