Use of biogenic and abiotic elemental selenium nanoparticles to sequester elemental mercury released from mercury contaminated museum specimens


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Mercuric chloride solutions have historically been used as pesticides to prevent bacterial, fungal and insect degradation of herbarium specimens. The University of Manchester museum herbarium contains over a million specimens from numerous collections, many preserved using HgCl₂ and its transformation to Hg⁰ represents a health risk to herbarium staff. Elevated mercury concentrations in work areas (~1.7 μg m⁻³) are below advised safe levels (<25 μg m⁻³) but up to 90 μg m⁻³ mercury vapour was measured in specimen boxes, representing a risk when accessing the samples. Mercury vapour release correlated strongly with temperature. Mercury salts were observed on botanical specimens at concentrations up to 2.85 wt% (bulk); XPS, SEM–EDS and XANES suggest the presence of residual HgCl₂ as well as cubic Hg₅ and HgO. Bacterially derived, amorphous nanospheres of elemental selenium effectively sequestered the mercury vapour in the specimen boxes (up to 19 wt%), and analysis demonstrated that the Hg⁰ was oxidised by the selenium to form stable HgSe on the surface of the nanospheres. Biogenic Se⁰ can be used to reduce Hg⁰ in long term, slow release environments.

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1. Introduction

The preservation of botanical specimens in museums across the world has traditionally proven difficult. A wide range of organic and toxic metal biocides have been used to deter the onset of decay caused by bacteria, fungi, insects and rodents; the efficiency of these treatments can be seen by the excellent state of preservation in many specimens today. The use of mercury salt solutions in the preservation of botanical specimens goes back to 1687, and continued in the UK up until the 1980s [1,2], ultimately giving way to organic pesticides including naphthalene, p-dichlorobenzene, thymol, lindane and DDT [2,3]. The preparation of mercury bearing pesticide solutions is detailed by Briggs et al. [4] and a concentration of 30 g l⁻¹ HgCl₂ and 30 g l⁻¹ phenol dissolved into methylated spirits was used in the Cambridge University Herbarium, UK. The low rate of sublimation of HgCl₂ and careful storage ensure that botanical specimens dating back hundreds of years still hold significant concentrations of Hg; as Hg⁰, residual HgCl₂, HgS and 2HgO·HgS [2,4–10]. Amalgamation of botanical collections into large herbaria, each having undocumented preservation techniques, has lead to inhomogeneous distribution of mercury contaminated specimens within collections. Staining of sample mounting paper due to the formation of crystalline Hg phases has been used to identify the presence of Hg-bearing compounds by fluorescence using UV irradiation [2].

Health risks associated with Hg-bearing botanical samples are not confined to dermal contact with contaminated specimens; Briggs et al. [4] report the evolution of Hg⁰ vapour leading to elevated local Hg concentrations of 25 μg m⁻³ at the Cambridge University Herbarium. Investigations in other herbaria indicate that this problem is widespread [8,9]. The concentration of Hg⁰ is dependent on ambient temperature; Oyarzun et al. [8] show that the Hg vapour concentration in the MAF Herbarium, Spain rises from 404–727 ng m⁻³ in late winter (23 °C) to 748–7797 ng m⁻³ in early summer (31 °C). Oyarzun et al. [8] also show that in a well insulated herbarium such as the relatively modern MA Herbarium, Spain, Hg⁰ concentrations exceeded 40 μg m⁻³.

Regulatory standards for workplace exposure limits have been summarised by Baughman [11], with guideline exposure limits varying between 25 μg m⁻³ and 100 μg m⁻³ Hg. It is also noted that children are much more susceptible to Hg toxicity than adults, which is of concern for public access museums.

The transformation mechanism of the relatively stable HgCl₂ into Hg⁰ is not well understood. Oyarzun et al. [8] surmised that microbial enzymatic reduction of Hg⁴⁺ to Hg⁰ as part of a Hg detoxification mechanism may be responsible, and work by Roane and
Snelling [12] has identified bacteria isolated from museum specimens that display a high Hg²⁺ tolerance (in excess of 10 mg l⁻¹), which may also be used as a possible remediation strategy. Complicating this process, however, is the presence of a range of other pesticides often found alongside Hg.

Briggs et al. [4] and Oyarzun et al. [8] both report that an increase in ventilation significantly drops the Hg₀ concentration of the air. There are circumstances where increasing ventilation cannot be considered due to climate, expense or positioning of the herbarium within buildings, and so an alternative method for decreasing the Hg content of herbarium and museum specimens must be identified.

Johnson et al. [13] studied the capture of Hg₀ by nanoscale sorbents, and note that elemental α-Se nanoparticles sequester Hg more efficiently than many commercially available sorbents. Se nanoparticles can be produced biogenically by bacterial reduction of soluble Se oxyanions. Se pollution, in the form of the oxyanions selenate [SeO₄²⁻] and selenite [SeO₃²⁻], is associated with waste materials from a broad spectrum of anthropogenic operations, including mining, agricultural, petrochemical, and industrial manufacturing operations [14]. Water-soluble forms of selenium can be microbiologically reduced to elemental selenium nanoparticles, which are less bioavailable and generally less toxic than other selenium species. The elemental selenium nanoparticles can potentially be separated from the aqueous waste stream [15]. The biologically recovered selenium nanoparticles can then be used in subsequent applications, such as the sequestration of elemental mercury released from mercury contaminated museum specimens, to offset the cost of the biological treatment [16]. This highlights how biomineralisation approaches can be applied to convert metal-containing wastes into new nanomaterials for environmental protection [17].

This research reveals the nature and extent of Hg contamination associated with museum specimens and measurement of Hg content of airspaces at the herbarium of the Manchester museum (The University of Manchester, UK), which has a botanical collection approaching a million specimens from all over the world, representing collections spanning hundreds of years. The collection is subdivided into British and European collections, as well as several smaller collections. The level of Hg contamination is inhomogeneous and poorly documented. This work also investigates the use of Se bionanominerals, which display an increased stability in comparison with their chemically synthesised counterparts [14,18,19] for the capture of Hg₀ released from specimens within herbaria.

2. Materials and methods

All chemicals used were of analytical grade and obtained from Sigma–Aldrich (UK).

2.1. Determination of herbarium air Hg₀ concentration

The Hg content of the herbarium air was determined using a portable mercury vapour indicator (MVI, Shawcity, Faringdon, UK). The Hg₀ concentration of the air was determined around the central workspaces of the herbarium and in sample boxes from a number of collections. The boxes analysed were chosen at random and were analysed immediately for Hg₀ concentration.

2.2. Sampling

The museum’s herbarium specimens are mounted onto A3 sized cardboard sheets and stored in fabric lined cardboard boxes on shelves within the herbarium, as shown in Fig. 1. Samples from the British and European collections were analysed for Hg concentration and oxidation state. Due to the irreplaceable nature of the specimens, milligram sized samples were taken and non-destructive analysis techniques (X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), scanning electron microscopy (SEM) coupled to energy dispersive spectroscopy (EDS) and X-ray absorption spectroscopy (XAS)) were used prior to chemical dissolution to determine bulk Hg values. Samples include leaf segments, mounting paper and plant sample debris (Table 1).

2.3. Mercury sequestration experiments

The Hg absorption capacity of dry powders of both biogenic and abiotic Se was tested. Biogenic red elemental α-Se was formed by the reduction of Na₂SeO₃ by Geobacter sulfurreducens coupled to the oxidation of H₂. G. sulfurreducens (ATCC 51573) was obtained from the Geomicrobiology Laboratory (University of Manchester) collection and grown to late log phase in modified freshwater medium amended with sodium acetate and fumarate as the electron donor and acceptor, respectively [20].

Fig. 1. Stacks of herbarium samples in containing boxes (left) and a typical herbarium specimen mounted on A3 paper with dark staining visible towards the top (right).
were harvested, washed and re-suspended in 20 mmol l\(^{-1}\) 3-((N-morpholino)propanesulfonic acid (MOPS) buffer at pH 7. Aliquots of the cell suspension were used to inoculate sterile, anaerobic solutions (final OD\(_{600}\) \sim 0.2) of 1 mmol l\(^{-1}\) Na\(_2\)SeO\(_3\) in 20 mmol l\(^{-1}\) MOPS amended with 2 \mu mol l\(^{-1}\) of the electron shuttle compound antheraquinone-2,6-disulfonate (AQDS). The headspaces of the bottles were replaced with H\(_2\) as an electron donor. Cultures were incubated at 30 °C and red Se\(^0\) was precipitated. To produce abiotic Se\(^0\), a solution containing 1 mmol l\(^{-1}\) Na\(_2\)SeO\(_3\) and 4 mmol l\(^{-1}\) glutathione (reduced) was titrated against a 1N NaOH solution until the formation of red elemental Se was observed [13]. For the sequestration experiments, biogenic and abiotic red Se\(^0\) suspensions (5 ml) were filtered through 0.22 \mum polycarbonate microfibre filters, washed with deionised H\(_2\)O (5 ml) and allowed to air dry.

### 2.3.1. Open system experiments

The Se\(^0\)-containing membrane filters were tested to determine the rate at which Hg\(^0\) could be sequestered from an Hg\(^0\)-containing N\(_2\) gas flow. The source of Hg\(^0\) in these experiments were solutions containing 10 ppm HgCl\(_2\) and 100 ppm SnCl\(_2\) [21]. N\(_2\) was bubbled through the HgCl\(_2\)/SnCl\(_2\) solutions at a varied flow rate (10–50 ml min\(^{-1}\)) and into 1 l vessels containing the Se\(^0\) membrane filters. Effluent was passed through an acidified KMnO\(_4\) solution prior to release under a fume hood [22]. The filters were exposed to the Hg\(^0\)-laden N\(_2\) flow for 3 h and stored at \sim\sim 80 °C prior to acid digestion and Hg determination using ICP-MS.

Real-time determination of the effect of Se-bearing membrane filters on a metered Hg-laden gas flow was analysed via a modified Cetac M6000A Cold Vapour Atomic Absorption Spectrometer (CVAAS) mercury analyser. Modifications to this equipment involved the removal of the auto sampling stage and inline attachment of Se\(^0\)-bearing membrane filters, altering analyses from aqueous, single point Hg determination to a real-time analysis of Hg concentration in the gas phase.

### 2.3.2. Sealed environment experiments

The ability of the Se\(^0\)-containing membrane filters to sequester Hg\(^0\) was assessed under idealised laboratory conditions. Se\(^0\) laden membrane filters were inserted into air tight 11 jars for 1 week at constant temperature (19 °C). HgCl\(_2\)/SnCl\(_2\) solutions served as the Hg\(^0\) source for these experiments. HgCl\(_2\) concentrations ranged from 10 ppb to 10 ppm and SnCl\(_2\) concentrations were kept in the ratio 1:100 Hg:Sn. After 1 week, Se\(^0\) membrane filters were removed from the sealed jars and stored at \sim\sim 80 °C prior to acid digestion and Hg determination using ICP-MS.

### 2.3.3. In situ experiments

Biogenic Se\(^0\)-containing membrane filters were inserted into herbarium specimen boxes known to contain significant Hg\(^0\) concentrations (\sim\sim 25 \mu g/m\(^3\)). The samples were left for four weeks, during which the time ambient temperature rose from 19 °C to 24 °C due to weather conditions. Following collection, samples were stored at \sim\sim 80 °C until acid digestion for Hg determination.

### 2.4. Hg-species characterisation

Samples described in Table 1 and Se\(^0\)-containing membrane filters from the mercury sequestration experiments were analysed by SEM using a Philips XL30 FEG-ESEM with elemental analysis by EDAX Gemini EDS. All samples were mounted onto adhesive C-coated Al stubs to remove interference in Se determination caused by overlapping Al emission peaks. XRD of mounting paper samples (Table 1) was carried out on a Bruker D8 Advance with a Cu-K\(_\alpha\) source. Spectra were compared to samples published by the International Centre for Diffraction Data (ICDD). XPS was carried out on sample 7 (Table 1) with a Kratos AXIS Ultra using monochromated Al K\(_\alpha\) radiation and a wide scan pass energy of 80 eV with narrow scans recorded at 20 eV pass energy for greater chemical state resolution. The intense C 1s peak was used to normalise the energy of the spectra.

XAS was performed on samples described in Table 1 and Se\(^0\)-containing membrane filters at the Diamond Light Source, Oxfordshire, UK, on the core XAS beamline B18 where samples were analysed at the Hg L\(_3\) absorption edge at 12.287 keV. Hg standards were used for absorption edge energy normalisation. Sample identification was via determination of inflection point difference (IPD) as described by Huggins et al. [23].

Acid digestion using \textit{aqua regia} (1 ml) with a molar ratio 1:4 HCl:HNO\(_3\) was used to determine the concentration of Hg in all sample materials. Excess HNO\(_3\) was used to encourage formation of Hg\(^2+\) cations to decrease losses due to volatility. Samples were run on an Agilent 7500cx ICP-MS, with quoted Hg detection down to 0.1 ppm. A 10 ppm Au solution was run concurrently to inhibit Hg retention. To ensure complete recovery of Hg, a 1 ml HF–HCl–HNO\(_3\) digestion (5:1:4) was used on any remaining material. The HF was neutralised using a 4 wt% solution of H\(_2\)BO\(_3\). Owing to the necessary high dissolved solid content, these samples were analysed using a Perkin-Elmer Optima 5300 dual view ICP-AES.

### 3. Results and discussion

The Hg\(^0\) concentration of air around the work area was 1.7 \mu g/m\(^3\) at 21 °C, which did not significantly alter throughout the herbarium. These results are similar to those observed in previous studies by Oyarzun et al. [8] and Kataeva et al. [9], which
recorded readings of up to 0.7 μg·m⁻³ and 1.1 μg·m⁻³ under similar conditions (≈21–23 °C).

The Hg⁰ concentration within the specimen boxes was determined for 70 boxes from 5 collections, of which 55 came from either the British or European collections. The Hg⁰ concentrations were highly variable with the highest concentrations in the European collection with Hg⁰ ranging from below detection limits up to 52 μg·m⁻³, with 5 boxes tested recording >25 μg·m⁻³ (Fig. 2). The British collection was largely free of significant Hg-contamination; however the Hg⁰ readings for the Rubus sp. type samples box were in excess of 40 μg·m⁻³ (data not shown).

High Hg⁰ concentrations were not observed outside of the specimen boxes prior to opening, indicating that the boxes inhibit the release of Hg into the working environment.

3.2. Characterisation of museum Hg-contaminated specimens

3.2.1. Electron microscopy and X-ray diffraction

Fig. 3A shows representative SEM images of samples collected from specimen boxes containing grains of sample debris (sample 4). EDS revealed that most of the samples examined did not have significant quantities of Hg. However, analysis of one particular grain (Fig. 3B and C), showed ~1 μm particles composed of ~87 wt% Hg with the counter ions S²⁻ and Cl⁻. The S component could not be quantified due to the overlap of the Hg M and S K emission lines, but the Hg wt% observed is similar to that expected from pure HgS (86 wt% as opposed to 74 wt% and 93 wt% Hg in HgCl₂ and HgO, respectively).

Samples of the A3 specimen mounting card (samples 2 and 8) were analysed by XRD (Fig. 4), indicating the presence of crystalline Hg phases. Sample 2 displayed peaks corresponding to unaltered HgCl₂ (ICDD PDF No. 00-026-0315) whilst sample 8 displayed a reflection at 26.4° 2θ, suggesting crystalline metacinnabar (ICDD PDF No. 00-006-0261). Hawks et al. [6] and Purewal et al. [2] have previously observed crystalline HgO, HgS and HgCl₂ phases on the mounting card.

3.2.2. XPS and XAS analysis

XPS analysis was used to identify the chemical forms of Hg present on the sample surface. Fig. 5 shows XPS spectra obtained from a Eucalyptus sp. leaf (sample 7) historically treated with Hg-based pesticides. XPS analysis showed that Hg was inhomogeneously distributed, with some areas falling below detection limits and others up to 1100 ppm. For the high Hg-containing areas, the...
Fig. 4. XRD data from analysis of samples 2 and 8 (mounting card). (A) Indicating remnant HgCl₂ and (B) diffraction peaks suggesting metacinnabar.

Fig. 5. XPS wide scan and inset, narrow scan spectra of *Eucalyptus* sp. leaf. Peaks correlating to Hg, C, N and O are all clearly discernable. Hg concentration determined to be ~1100 ppm.

binding energy for the Hg 4f emission was 101 eV. Comparison with published standards indicates that the Hg could be present as HgS, HgCl₂, HgO or as an organo-mercury compound, but precludes the presence of Hg⁰ [24].

Fig. 6. XANES Hg L₃ absorption edge spectra obtained from (A) *Eucalyptus* sp. leaf; (B) box dust (sample 4) and (C) Se-laden polycarbonate membrane filter exposed to Hg⁰. Inset: 1st derivatives of the XANES spectra.
The XANES spectrum derived from the *Eucalyptus* sp. leaf is presented in Fig. 6A, with the 1st derivative of the XANES spectrum inset. An IPD value of 7.88 eV was recorded for the 1st derivation, consistent with published HgS species [25]. Linear combination fitting of published HgS standards [23,25,26] suggests the XANES profile is consistent with metacinnabar, concordant with XRD findings. An IPD value of 7.89 eV was derived from the XANES analysis (Fig. 6B, inset) of the Hg-rich dust (sample 4), and is concordant with the EDS indicating the presence of a dominant HgS phase. The presence of multiple peaks in the 1st derivation of all XANES spectra obtained shows no Hg⁰ is present in these samples.

### 3.2.3. Acid digestion of specimens in Table 1

Total extraction of Hg from the museum specimens was used to determine absolute Hg concentrations. The results of the sequential acid extractions using *aqua regia* and HF-*aqua regia* showed that Hg concentration in the specimens varied greatly (Table 1). Interestingly, the digests show that the *Eucalyptus* sp. leaf and plant stem (samples 7 and 9) do not contain significant concentrations of Hg (0.08 and 0.07 wt%). Dust and debris collected from two of the specimen wrapping sheets (samples 1 and 3) show Hg concentrations of ∼0.2 wt%, but debris collected from the box containing the specimens (sample 4) had a Hg concentration of 2.85 wt%, over an order of magnitude larger. These results highlight the heterogeneity of Hg contamination within the specimens and may relate to varying pesticide coating practices. Hg contamination was found in all samples.

### 3.3. Mercury sequestration experiments

#### 3.3.1. Selenium nanoparticle and filter characterisation

Selenium impregnated membrane filters were characterised prior to use in laboratory experiments by ICP-AES/-MS, SEM with EDS and particle sizing. Total acid digests showed that membrane filters were loaded with 500–600 μg of Se per filter for both abiotic and biogenic Se nanoparticles. SEM analyses of the loaded membranes prior to experimentation show smooth, spherical particles approximately 100–200 nm in diameter (Fig. 7A). EDS analyses of the Se phases (Fig. 7Di) show a dominant Se L edge peak at 1.4 eV along with smaller C and O emissions from the filter material, indicating a pure Se phase. Size distribution information was obtained (Fig. 8E) and was concordant with SEM analyses show particles ranging from 40 to 700 nm, similar to that as reported by Johnson et al. [13]. Size distribution information was used to calculate surface area assuming spherical particles; abiotic Se⁰ had a surface area of 11.64 m² g⁻¹ and biogenic Se⁰ 9.64 m² g⁻¹ equating to 5.8–7.0 × 10⁻³ m² and 4.8–5.8 × 10⁻³ m² per filter, respectively.

#### 3.3.2. Open system experiments

Biogenic Se⁰-containing membrane filters were assessed using the modified CVAAS to follow the evolution of Hg⁰ from a 10 ppb Hg²⁺/Sn²⁺ solution and to determine the effect of Se on the Hg⁰ release (Fig. 9). Evolution of Hg⁰ at a flow rate of 50 ml min⁻¹ shows an initial spike corresponding to release of Hg⁰, followed by an exponential decay over time (r² = 0.98). The addition of biogenic Se⁰ to the inline membrane filter drastically decreased initial Hg⁰.

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**Fig. 7.** (A) Initial biogenic Se⁰ starting materials; (B) formation of HgSe crust following Hg⁰ exposure; (C) increased magnification view of HgSe phase; (D) EDS of biogenic Se⁰ before (i) and following (ii) exposure to Hg⁰.
Fig. 8. Characterisation of the Se particles. (A) Crystalline Se⁰ formed following crystallisation of abiotic Se⁰; (B) higher magnification view of hexagonal abiotic Se⁰ crystals; (C) biologically precipitated Se⁰ following aging under the same conditions as (A); (D) EDS of biological (i) and abiotic (ii) Se⁰ precipitates showing high purity following washing; (E) size distribution of biogenic (solid) and abiotic (dashed) Se⁰ precipitates (abiotic data from Johnson et al. [13]); (F) XRD data for initial Se⁰ materials (i), amorphous biogenic Se⁰ after 58 days, hexagonal crystalline abiotic Se⁰ after 53 days (iii).
Fig. 9. CVAAS spectra for Hg concentration of gas phase with (Se) and without (No Se) Se-laden membrane filters, showing the absorption of Hg\(_0\) by α-Se\(^0\) is immediate and decreases in efficiency over time.

emission (a decrease of 78% in peak intensity), and shows an overall decrease of 47% in Hg emission in the first 40 min. The extent of Hg capture decreases over time until Hg levels approach those of the system without biogenic Se\(^0\).

Experiments comparing biogenic and abiotic Se\(^0\)-containing membrane filters at varying gas flow rates and reaction times using a 10 ppm Hg source show a strong correlation between the Hg sequestered and flow rate (Fig. 10), with a maximum Hg\(^0\) sequestration of 3.4 mg m\(^{-2}\) for biogenic Se\(^0\) at 10 ml min\(^{-1}\), nearly quadruple that of Hg\(^0\) sequestered at a flow rate of 50 ml min\(^{-1}\). Abiotic Se\(^0\) tested under the same conditions was capable of sequestering up to 7.7 mg m\(^{-2}\) of Hg\(^0\) at 25 ml min\(^{-1}\), more than four times that of biogenic Se\(^0\).

3.3.3. Sealed environment experiments

Biogenic α-Se\(^0\) was exposed to a range of HgCl\(_2\) concentrations between 10 ppb and 10 ppm with SnCl\(_2\) at a 1:100 Hg:Sn ratio, and an increasing trend of Hg\(^0\) absorption was noted (Fig. 10C), suggesting that the saturation limit of the biogenic Se\(^0\)-containing membrane filters was not reached. The maximum Hg\(^0\) absorption of 2.2 mg m\(^{-2}\) (equivalent to 2.1% Hg by mass of Se), occurred at 10 ppm HgCl\(_2\) and represented only 10% of the maximum Hg/Se mass ratio found by Johnson et al. [13]. Abiotic Se\(^0\)-containing membrane filters were able to capture >4 mg Hg\(^0\) m\(^{-2}\), double the capacity of biogenic Se\(^0\).

Stock solutions of abiotic α-Se\(^0\) were seen to change colour from red to black. XRD analysis showed the crystallisation of amorphous α-Se\(^0\) to hexagonal Se\(^0\) (ICDD PDF Card No. 00-006-0362), whereas biological Se\(^0\) remained stable in suspension for over a year following synthesis (Fig. 8F). Similarly, a colour change was noted in the abiotic Se\(^0\) impregnated membrane filters exposed to 10 ppm Hg, and subsequent SEM analyses (Fig. 8A and B) show that crystallisation of the dried Se\(^0\) powders had occurred. EDS analysis confirmed that the crystalline Se was pure Se with no Hg, indicating recrystallisation of Se rather than neocrystallisation of an HgSe phase (Fig. 8D). Biologically precipitated α-Se\(^0\) on the impregnated filters did not change (Fig. 8C).

Johnson et al. [13] highlighted the effect of the protein bovine serum albumin (BSA) on Se\(^0\) powders; Hg\(^0\) capture was hindered relative to Se\(^0\) synthesised without BSA, despite a large increase in the available surface area. This was attributed to surface passivation by BSA, reducing the density of available reactive sites on the Se\(^0\) particle surface. Results published by Pearce et al. [14] demonstrate the presence of surface-associated proteins on Se\(^0\) produced by G. sulfurreducens, and work published by Prakash et al. [27] show that efforts to remove the biological coatings on bacterially precipitated Se\(^0\) resulted in the formation of similar hexagonal form seen for abiotic Se\(^0\) (Fig. 8A). Similar organic coatings have also recently been characterised by C\(_{60}\) TOF-SIMS on other complimentary biomonomaterials synthesised by G. sulfurreducens [28]. It is likely this surface-bound bacterially derived organic layer is increasing the

Fig. 10. The Hg content of Se-laden membrane filters following acid extraction for the biologically formed Se filter membranes (A) and abiotic Se membranes (B) under varying carrier gas flow rates. (C) The Hg concentration of biological Se membrane filters under sealed conditions.

![Graph](image-url)
bionanominerals. Potentially link bioremediation strategies with production of functional materials are accessed. Hg-contamination of specimens is variable between collections, within collections and even within specimens stored in the same box. Hg is found on specimens predominantly as metacinnabar. However, indications of a surface coating of HgO and the presence of HgCl2 suggest that the form of Hg in this system is affected by a number of factors including biological activity, oxidation and original Hg concentration. Significant quantities of Hg (2.5 wt%) were found in some specimens. Hg0 was not found to oxidise into the air. The observed absorption capacity of up to 20% Hg by mass is affected by a number of factors including biological activity, oxidation and original Hg concentration. Significant quantities of Hg (2.5 wt%) were found in some specimens. Hg0 was not found to oxidise into the air. The observed absorption capacity of up to 20% Hg by mass is affect

3.3.4. In situ experiments

Biogenic Se0-containing membrane filters were placed in five boxes of the European collection with Hg0 concentrations greater than 25 µg m⁻³ (Fig. 2) and left for 4 weeks. The effect of the biogenic Se0-containing membrane filters on the concentration of Hg0 in the air within herbarium specimen boxes could not be determined due to the changes in ambient air temperature during the course of the in situ experiment; a temperature increase from 19 °C to 24 °C occurred during the 4 weeks, leading to an increase in Hg0 release as previously observed by Oyarzun et al. [8]. An increase from 42 µg m⁻³ Hg0 to 90 µg m⁻³ in one specimen box was seen. SEM images of the biogenic Se0 before and after exposure to Hg0 in the specimen boxes are shown in Fig. 7. The initial form of the α-Se0 is as clearly defined spheres with smooth surfaces with particle sizes ranging from 35 to 680 nm (Figs. 7A and 8E). Following 4 weeks in Hg contaminated herbarium boxes, there was a morphological change of some spheres to a rough, textured surface (Fig. 7B). EDS spectra of remaining smooth and rough spheres showed that smooth particles contained little Hg, whereas the rough textured particles show a significant Hg concentration (Fig. 7D). EDS analysis showed that these particles were composed of Hg and Se (31–33% Hg by mass). As pure HgSe is ~72% Hg by mass, these results suggest that there is a significant surface coating of an HgSe phase.

Following exposure of biogenic Se0 to Hg0 in the specimen boxes, a sample was analysed using XAS at the Hg L3-edge (Fig. 6C). Linear combination fitting of the recorded XANES profiles against published standards suggests the presence of HgSe, concurrent with EDS findings. An IPD value of 7.47 eV was determined from the 1st derivation of the XANES spectra, which differs from published HgSe IPD values [23], likely as a result of the low Hg concentrations in the sample, resulting in a high signal:noise ratio (Fig. 6C, inset).

Acid digestion of the biogenic Se0-containing membrane filters in aqua regia was carried out to determine total Hg concentrations (Fig. 11). All membrane filters tested showed Hg concentrations in the range 0.02–0.19 µg Hg per µg Se, corresponding to the sequestration of up to 19% Hg by mass, which is comparable to the sequestration results obtained for abiotic Se0 by Johnson et al. [13].

The normalised concentration of Hg0 sequestered by the biogenic Se0-containing membrane filters shows a linear correlation with the Hg0 concentration of the air within the herbarium specimen boxes.

4. Conclusion

Measurement of Hg concentrations in the Manchester museum herbarium has identified the presence of Hg0 both within well-sealed specimen boxes and in the air of the herbarium workspaces. The concentration of Hg0 in workspaces did not exceed 1.7 µg m⁻³, which is well below the lowest advised workplace mercury exposure limit of 25 µg m⁻³. The Hg0 content observed within specimen boxes was variable and sometimes very high, increasing from 43 µg m⁻³ at 19 °C to 90 µg m⁻³ at 24 °C in one box. The construction of the specimen boxes is an advantage as it prohibits the build up of significant concentrations of Hg0 within boxes, which is released when materials are accessed. Hg-contamination of specimens is variable between collections, within collections and even within specimens stored in the same box. Hg is found on specimens predominantly as metacinnabar. However, indications of a surface coating of HgO and the presence of HgCl2 suggest that the form of Hg in this system is affected by a number of factors including biological activity, oxidation and original Hg concentration. Significant quantities of Hg (2.5 wt%) were found in some specimens. Hg0 was not found to be directly associated with the specimens, but a strong correlation between temperature and release of Hg0 was observed.

Biogenic α-Se0 efficiently sequesters Hg0, with a 47% initial reduction in Hg emission. Contact time between the Hg0 and the biogenic Se0 nanoparticles has a significant effect on the reaction rate, with the biogenic Se0 capturing Hg0 more effectively at lower gas flow rates, similar to the conditions of low level, continuous Hg0 release expected in the sealed specimen boxes. The sequestration of the Hg0 as a stable layer of HgSe on the surface of the nanoparticles represents a safe option to limit the release of Hg0 into the air. The observed absorption capacity of up to 20% Hg by mass compares favourably with that observed for abiotically synthesised α-Se0. A major advantage of biogenic α-Se0 over abiotic α-Se0 is the improvement in long term stability, which offsets the initial low reaction rates with improved longer term performance. Amorphous elemental red Se nanospheres produced both biogenically and abiotically represent a promising new way to capture Hg0 released from Hg- contaminated herbarium specimens. The potential to use biogenic Se nanospheres, formed as a by product of biological treatment of selenium contaminated wastewater, also provides an opportunity to link bioremediation strategies with production of new nanomaterials for environmental protection.

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References


