The effect of mercuric chloride treatment as biocide for herbaria on the indoor air quality

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Abstract
One of the most previous conservation treatments for plant specimen in herbarium collections was mercuric chloride (HgCl₂). However, due time HgCl₂ may decompose and it may cause (metallic) mercury (Hg) emission. Hg vapour in indoor air should be avoided as mercury poisoning can already occur at levels of 6 μg/m³. Both field and laboratory studies were carried out while the Hg concentration was analysed in the air. Field studied included analysis of the indoor air in the repository and analysis of the concentration inside a herbarium box. Subsequently lab experiments were carried out on measuring the Hg emission using a small scale emission chamber. The lab studies proved that the emission of Hg from collections is according to the theories of Hoetjer–Berger–Fuji and therefore it can be seen as a continuous emission. Field studies concluded that workers and visitors may be subjected to a high mercury concentration in the repositories. Opening a herbarium box may even cause a peak concentration of over 80 μg/m³. The highest concentration observed in the repositories was 13 μg/m³. Although this value is lower than the official Dutch threshold levels of 20 μg/m³ (time weight average over 8 h for work places), this concentration has to be considered as high. Based on the lab studies, it can be concluded that by adjusting the ventilation rate in work rooms and repositories, the mercury concentration indoor can be lowered drastically and improve the indoor air quality. However care should to be taken when opening herbarium boxes, due to possible high mercury emission.

Keywords: Herbarium, Mercury emission, Indoor air quality, Occupational health

Background
Mercuric chloride and conservation
One of the most used biocides for disinfection of natural history collections and in special herbarium collections is mercuric chloride (or mercury(II)chloride, HgCl₂, CAS number 7487-94-7). HgCl₂ forms solid white crystals, and once it was applied as a medicine against e.g. syphilis. However, HgCl₂ is extremely poisonous and has the ability to sublimate. HgCl₂ decomposes into metallic mercury (Hg⁰, or just Hg) by reduction. Hg is the only metal that is a liquid at room temperature and due its sublimation a high concentration of Hg vapour in the air may easily occur [1].

Both HgCl₂ and Hg (vapour) are extremely dangerous for our health, as for example Hg taken up by the lungs will accumulate in the kidney and brains [2]. The World Health Organisation, WHO, is classifying Hg as ‘extremely hazardous class 1a’ [1, 3]. In the European Union the advisory value for health of Hg in the indoor air is set to 50 ng/m³ (average annual concentration) and therefore it is lower than the official Dutch threshold level of 20 μg/m³ (time weight average over 8 h) [4].

Up to 1980, herbarium collections were disinfected by means of the use of mercuric chloride [5] and, for example, in the Netherlands it was last used in 1984 [6]. Based on surveys it has become clear that different methods were applied [7], for example by soaking or brushing materials using a solution of HgCl₂ with ethanol. Herbarium samples can continue to emit metallic mercury many decades after their last treatment with mercuric chloride [5, 8–11].
Release of mercury
The emission of mercury (Hg) from herbarium collection forms a serious occupational health hazard, not only for the workers at the collection, but also for the visitors and researchers who need to work with the collection materials. In the Netherlands (National Herbarium collection repositories, now dismounted) steady state concentrations in the indoor were found to be over 10 µg Hg/m³ while for example other collections were causing concentrations of mercury indoor of about 25 µg Hg/m³ (Cambridge herbarium) and 40 µg Hg/m³ (MAF, Madrid). However, opening collection boxes or cabinets where specimens are stored can cause a peak concentration of 50 µg Hg/m³ and higher [5, 8–11].

Different mechanisms can be hypothesised as responsible for the emission of mercury from herbarium collections. First the ionogenic reactions, uptake of electrons by the mercuric ion, that may occur even at the normal moisture content in cellulose materials [12]:

\[ \text{Hg}^{2+} + 2e^- \rightarrow \text{Hg} \quad (E^0 = 0.851 \text{ V}) \quad (1) \]

\[ 2\text{Hg}^{2+} + 2e^- \rightarrow \text{Hg}_2^{2+} \quad (E^0 = 0.920 \text{ V}) \quad (2) \]

\[ \text{Hg}_2^{2+} + 2e^- \rightarrow 2 \text{Hg} \quad (E^0 = 0.797 \text{ V}) \quad (3) \]

Practically there is moisture present during the treatment, and therefore this may cause the initiation of oxidation and reduction reaction in cellulose-containing specimens. These reactions may enhance in case, e.g., more hemi-cellulose is present as these polyoses contain more reactive carbonyl groups (C≡O) than cellulose [13, 14]. A carbonyl group may be oxidized resulting in a carboxyl group (–COOH) (Fig. 1).

For example, the oxidation of cellulose produces electrons according to

\[ \text{Cellulose} - \text{C}_\text{H} = \text{O} \rightarrow \text{Cellulose} - \text{COOH} + 2e^- \quad (4) \]

\[ \text{O}_2 + 2e^- \rightarrow 2\text{O}^- \quad (5) \]

Summarized:
\[ \text{Cellulose} - \text{C}_\text{H} = \text{O} + \frac{1}{2} \text{O}_2 \rightarrow \text{Cellulose} - \text{COOH} \quad (6) \]

It was also suggested that the reduction of \( \text{Hg}^{2+} \) can occur as a two-step reaction [15]. This is based on the \( \text{Hg}^{+} \) determination in the specimen and the fact that Hg is found in the indoor air. Also Purewal demonstrated that, due to the occurring reactions, the mount paper used to display the specimens will shine fluorescent orange under a UV lamp if mercury is present, however it has not be fully proven if the fluorescent is due to the mercury-cellulose complex or due to paper degradation.

The analysis and prediction of the emission of volatile from materials was firstly reported in European Committee for Standardization (CEN) in 1984, [16]. Other important work discussing the emission of volatiles was published by the European Cooperation in Science and Technology Network platform (COST) by COST action 613 “indoor air Quality and its impact on man” and described in several reports on how to measure the steady state concentration in situ, how to predict the emission and also how to make interpretation of the observed emission results [17–20]. Only in the last decade has the role of the emission for objects of cultural value come under discussion. This not only because of the quality of the material where the cultural artefact has been made of, but more related to the conservation treatment and its impact on occupational health.

It has been proven frequently that there is a relation between the emission rate and the ventilation rate [21, 22]. Once this relation is found, the concentration of the indoor air can be modelled. The principle of the measurement is as above, however the steady state concentration is measured using at least 3 datapoints (different ventilation rates). The relation between the concentration of a compound and the ventilation rate can be described according to the Hoetjer equation [21, 23] and later more known as the Hoetjer–Berge–Fujii equation [24]:

\[ \frac{1}{C} = \frac{1}{C_e} + \left( \frac{1}{ksC_e} \right) x \left( \frac{n}{L} \right) \quad (7) \]

Here \( C \) = steady state concentration (µg m⁻³), \( C_e \) = equilibrium concentration at a ventilation rate of 0 (h⁻¹), \( k \) = mass transfer coefficient, \( n \) = air exchange rate (h⁻¹), \( L \) = loading factor (m² m⁻³)

Experimental
Concentration of mercury inside a herbarium box
As the herbarium materials were subjected to a mercuric chloride it is assumed that the herbarium boxes, having several treated sheets inside, are the main source for the mercury concentration indoor. In order to determine

Fig. 1 A suggested pathway on the formation of volatile mercury.
The carbonyl group of, e.g., cellulose reacts with mercuric ions to form an organic acid while metallic mercury is formed and can be emitted.
the initial concentration of mercury inside a herbarium box the concentration was measured using a Mercury Vapour Monitor MV-3000 from Mercury Instruments (Anonymous 2013). We applied the most simple way, i.e., to put directly the inlet of the mercury monitor inside the box as shown in Fig. 2. As the monitor has an air flow of approx. 0.6 L/min reading was done after 30 s to minimize the amount of outside air entering the box.

Concentration mercury in the repositories
Repository workers are subjected to the indoor air quality present in the repositories and thus to the present concentration of mercury in the air. For the concentration analyses and measurements of mercury a Mercury Vapour Monitor MV-3000 from Mercury Instruments was used [25]. For the mercury concentration measurements in the repositories, a random walk was carried out and on different locations readings were taken. The walk was from the corridor, where a copy machine is located, to a connecting corridor with two repositories (Fig. 3).

Mercury emission
For the evaluation of the mercury concentration of the content of a herbarium box at different ventilation rates a small emission chamber was used. The chamber was developed within the EU-PaperTreat project [26]. Within this chamber not only the ventilation rate could be adjusted, but also the circulation rate, comparable to archival storage rooms having a condition of a ventilation rate of 0.2 l/h and a recirculation rate of 2.0 l/h.

The chamber is made of stainless steel and has a volume of 0.1 m³. This size has an advantage that it is simple to reach a high loading (i.e., emitting surface of the material compared to the volume of the chamber). The chamber is connected to an air inlet, provided with a filter to assure non-polluted air and the outlet is connected to two pumps and the mercury monitor. One of the pumps is responsible for bringing the air back to create the circulation rate. The ventilation rate is created by the sum of the flows of one pump plus the analysers.

Figure 4 shows the experimental set up including how the materials are placed inside the chamber. For our research we used a circulation rate of 2.0 l/h and three ventilation rates: 1, 1.5 and 2 l/h.

Results and discussion
Figure 5 shows a frequency distribution of the concentrations measured using 29 randomly selected herbarium boxes. Based on this graph it is suggested that the concentration of mercury inside a herbarium box will be mainly between 10 and 40 µg/m³, while concentrations of higher than 80 µg/m³ are possible. Therefore opening a herbarium box should be done with care and preferable precautions should be taken to avoid inhalation of high concentration of mercury.

The fact that there is a concentration of mercury inside a herbarium box indicates that mercury should also emit from the boxes into the storage rooms. And this is indeed what we observed in the building that was housing the collection. This building housing the collection had two floors and both floors were housing parts of the collection. Both floors are separated by means of doors that are closed in a normal situation. The repositories have also doors that should be closed in normal situations, however during the days we were measuring all doors were open. The ground floor is in direct open connection with the outdoors, via the main entrance and side entrances. The first floor can be entered by a corridor and stair with closed doors on the ground and first floor. Entering the first floor of the building, near the copy machine, we observed between 6.1 and 6.7 µg mercury per m³. Walking through the corridor we observed 6.5–8.5 µg mercury per m³ and finally inside the repository we observed 10–13 µg mercury per m³, indeed indicating that mercury emits from the collection boxed to the indoor air of the building. The rooms where administrative and research
Fig. 3 Entrance of one of the repositories. Here work places are present for external visitors.

Fig. 4 The experimental set up for the mercury emission measurements. Left the stainless steel chamber, right an example on how specimen can be placed in the chamber.
work is being carried out were all connected by a door
directly with the corridor. In these rooms we observed
concentrations of mercury varying from 2 µg/m³, where
the door of the room was closed, up to 8.5 µg/m³ where
doors were open.

Although it looks obvious, that the mercury concentra-
tion in the working areas comes from the repositories, we
checked the flows of air by means of smoke-tubes. The
generated smoke could be easily followed and showed the
stream of air was from the repositories towards the cor-
ridor and subsequently the working rooms.

Although the found concentrations are below the offi-
cial Dutch threshold levels of 20 µg/m³ (time weight
average over 8 h for work places) they have to be consid-
ered as high as it is known that serious mercury poison-
ing can already occur at levels of 6 µg/m³, especially for
young workers and women who are pregnant [27]. Finally
looking at the advisory maximum concentration given by
Boerstra et al. of mercury in living houses, i.e., 0.05 µg/
m³, it has to be concluded that the found values can be
seen as high.

Based on this work, serious actions were undertaken
by the ‘herbarium’, for example, closing doors where the
doors should be closed and improving the quality of
the ventilation. By improving the ventilation, thus the
amount of fresh air led into the building increased, may
result in diluting the mercury concentration. This was
also demonstrated by the experiments on the effect of the
ventilation rate on the mercury emission.

Figure 6 shows an example of the results on the emis-
sion of mercury from a content of a herbarium box in
relation to the ventilation rate. Here the ratio of n/L in
m/h (ventilation rate divided by the loading) is expressed
on the x-axis while the reciprocal concentration in m³/µg
has been placed on the y-axis. The bundle tested consists
of 41 folders filled with total 87 sheets of herbarium spec-
imens. All calculated concentrations were corrected for
a blank, i.e., concentration of mercury in the sampled air
when no specimens were present in the emission cham-
ber. The measured point at n/L = 0 m/h has been taken
by connecting the outlet of outgoing air with the inlet of
the incoming air of the emission chamber.

As can be seen from Fig. 6, the emission of mercury
follows very well the theories of Hoetjer. Based on the
results of this experiment it is suggested that the emis-
sion from mercury is according to a continuous emis-

Fig. 5 Distribution of the mercury concentration inside a herbarium box, measured directly inside the box using a Mercury Vapour Monitor MV-3000

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follows very well the theories of Hoetjer. Based on the
results of this experiment it is suggested that the emis-
sion from mercury is according to a continuous emis-
ion process, i.e., the emission of mercury from the
specimen will be due to the presence of relatively high
concentration of bound mercury on the specimen.

Purewal [15] already suggested that it is due to the
uptake of mercury by the mount paper, however based on our work we suggest that it is mainly due to the presence of paper deterioration products, because mercuric chloride was found as particles on the specimen (Fig. 8). Some fallen leaves from the tested materials were put under a scanning electron microscope coupled with energy dispersive X-ray spectroscopy (SEM/EDX). In this example white spots on the leaf could be clearly observed by means of a backscattered electrons detector. The EDX analysis indicated the presence of both mercury and chloride, which suggests that the mercury is combined with chloride (Fig. 7).

The fact that Purewal [15] found a fluorescence effect may indicate that this is not only due to presence of mercury but also due to occurring reactions, for example the Fenton reaction of cellulose caused by the presence of transition metals as mercury(I)- and (II)-ions. The degradation of the cellulose may result in discoloration as it is found on the flyers present between the specimens (see Fig. 8).
Further research suggestion

Based on our findings and literature reading, we hypothesised that mercuric chloride treatment may also cause deterioration of the collection. This deterioration can be already observed by the naked eye, i.e., discoloration and fluorescent effect of the paper substrate. Therefore we recommend further research on the occurring mechanism of cellulose degradation by mercuric chloride.

Authors’ contributions

JH supervised the experimental field and lab studies, performed the calculations and drafted the manuscript. RD described the mercury problem, co-supervised the field studies, delivered the original materials for lab-studies and helped to draft the manuscript. RS had a serious part in the discussion on the feasible possibilities for mercury reduction and helped to draft the manuscript. All authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

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