Why productive lakes are larger mercury sedimentary sinks than oligotrophic brown water lakes

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Abstract

Mercury accumulation in lake sediments is a widespread environmental problem due to the biomagnification of Hg in the aquatic food chain. Soil Hg concentrations, catchment vegetation, erosion, and lake productivity are major factors controlling the accumulation of Hg in lakes. However, their influence on the Hg mass balance in lakes with different catchment characteristics and trophic state is poorly understood. In this multilake study, we decipher the effects of catchment vegetation (coniferous vs. deciduous forest), soil Hg content, and trophic state on Hg sedimentation at six lakes in Germany. We investigated Hg concentrations in leaves, soils, and the lake’s water phase. Soils under coniferous stands show slightly higher Hg concentrations than under deciduous forest. Hg concentrations in the water phase were higher in the oligotrophic brown water lakes (8.1 ± 5.6 ng L⁻¹ vs. 3.0 ± 1.9 ng L⁻¹). Lower Hg concentrations in sediment trap material indicate dilution by algae organic matter in the mesotrophic lakes (0.12–0.17 μg g⁻¹ vs. 0.57–0.89 μg g⁻¹). However, Hg accumulation rates in sediment traps were up to 14-fold higher in the mesotrophic lakes (113–443 μg m⁻² yr⁻¹) than in the brown water lakes (32–144 μg m⁻² yr⁻¹), which could not be explained by higher Hg fluxes to the productive lakes. Hg mass balance calculation reveals that water phase Hg scavenging by algae is the major reason for the intense Hg export to the sediments of productive lakes which makes them significantly larger sedimentary sinks than oligotrophic brown water lakes.

Mercury is a toxic pollutant of global concern that is known for its biomagnification in aquatic food chains. Due to the enrichment of methyl-Hg (MeHg) in fish, understanding the biogeochemical cycling of Hg in lakes is crucial. Lakes integrate catchment characteristics and processes and are often a major sink of pollutant (Moser et al. 2019). Thus, most Hg in lakes is received from the atmosphere via direct atmospheric deposition. Subsequent processes as leaching and erosion of catchment soils, or transport via streams and rivers are widely accepted to be the main transport pathways for Hg found in limnic systems. Atmospheric Hg enters forest ecosystems via wet and dry deposition (Hintelmann et al. 2002; St. Louis et al. 2019). Approximately 65–90% of this input is through stomatal uptake of elemental Hg (Hg⁰) (Demers et al. 2007; Jiskra et al. 2015; Enrico et al. 2016). Hence, litterfall accounts for most of the Hg flux to forest soils, although the extent of this process, as well as transport to downstream waters, may differ between coniferous and deciduous forests (Woerdle et al. 2018). Higher Hg concentrations have been observed in the litter of coniferous than those in deciduous forests (Schwesig and Matzner 2000; Obrist et al. 2012), and the highest Hg accumulation is in the organic horizons of coniferous forest soils (Gruba et al. 2019). Moreover, needles of evergreen coniferous trees provide a larger surface area for Hg uptake when regarding the leaf/needle surface area on an annual basis (i.e., no leaf surface area in deciduous forests in winter) (Obrist et al. 2012; Navrátil et al. 2014). This difference may result in higher Hg concentrations in coniferous litter than in litter from deciduous forests, which in turn may result in a higher Hg flux to the forest floor in coniferous forest stands (Grigal et al. 2000; Johnson et al. 2007). However, others have demonstrated that Hg concentrations in coniferous litter are lower than those in deciduous litter (Sheehan et al. 2006; Demers et al. 2007; Navrátil et al. 2016). In a recent study, Wohlgemuth et al. (2020) demonstrated higher foliar Hg uptake per leaf area in broad leaves than in current-season conifer needles. Drenner et al. (2013) found a significant linear relationship between Hg levels in largemouth bass (Micropterus salmoides)
and the percentage of landscape covered by coniferous forest but not with percent land coverage by deciduous forest. Particulate organic matter (POM) and dissolved organic matter (DOM) are major transport vectors of Hg from forests to lakes owing to the enrichment of Hg present in the organic soil horizons (Klaminder et al. 2008; Biester et al. 2012; Jiang et al. 2018). Similarly, a higher percentage of peatlands in the catchment is related to both higher DOM and higher Hg concentrations in lake water (Laudon et al. 2004; Wiener et al. 2006). Previous studies on Holocene palynological and geochemical data derived from lake sediments indicate a significant increase in Hg concentrations when catchment vegetation shifted from deciduous to coniferous vegetation during the Holocene (Rydberg et al. 2015; Schütze et al. 2018).

In a lake’s water phase, Hg is predominately associated with DOM, POM, as well as minerogenic particles and settles to the lake bottom along with the sediments (Gascón Díez et al. 2016). Algae and algal-derived organic matter (OM) have been shown to take up or scavenge Hg from the water column (Mason et al. 1995; Dranguet et al. 2014). The scavenging of Hg by algae has been shown to have a major influence on Hg cycling in lakes (e.g., speciation), being a major vector for Hg accumulation in aquatic sediments (Biester et al. 2018; Zaferani et al. 2018), and a key driver for promoting Hg accumulation in lake sediments since industrialization (Outridge et al. 2007; Faucheur et al. 2014). Some studies have found no relationship between algal productivity and Hg accumulation in sediments (Kirk et al. 2011; Cooke et al. 2012), while others have suggested that terrestrial inputs may obscure the signal from algal scavenging (Sanei et al. 2012; Brazeau et al. 2013). Studies based on Hg mass balance calculations argue that algal blooms may dilute Hg concentrations in algal-derived OM (Pickhardt et al. 2002; Karimi et al. 2007). The idea that Hg scavenging or uptake by algae affects Hg accumulation in lake sediments is widely accepted. However, the extent of Hg scavenging by algae, how this relates to trophic state, and its role in the accumulation of Hg in lake sediments is poorly understood.

In this study, we investigated six lakes in two mountain areas in Germany, which differ in catchment vegetation (coniferous vs. deciduous forest) and in-lake primary productivity. To decipher the effect of these parameters on Hg concentrations and accumulation in the lake sediments, we investigated catchment soils, inflow, and lake water Hg concentrations, along with Hg sedimentation rates in sediment traps over a period of 2 yr. In addition, we used C/N ratios and Fourier-transform infrared spectroscopy (FTIR) to determine the dominant sources of OM in the different lake systems.

Materials and methods

Study areas

The two study areas are located in the eastern Harz Mountains (HZ) and the northern Black Forest (BF) in Germany (Fig. 1A) and are defined as low mountain ranges, a typical geographical feature of Central Europe. In contrast to high-altitude mountains such as the High Alps, the summits at the studied mountain sites do not reach the tree line, which results in seasonal litterfall in the catchments of the forest lakes.

In the Harz Mountains, we investigated three small reservoirs: Großer Siebersteintee (GT), Kleiner Siebersteintee (KT), and Großer Dachsteich (DT). These mesotrophic to eutrophic reservoirs were constructed during the 18th century for fisheries. Nowadays, the lakes are predominantly used for flood control and local recreation, as these waterbodies are located near the provincial town of Ballenstedt (Fig. 1B) where settlements are partly downstream of the lake outflows. The catchment soils are dominated by base-rich brown earth and brown earth podzols from rubble loess over loamy graywacke debris (BGR Geoviewer 2018). The steep flanks of the reservoirs are covered in deciduous stands with oak (Quercus robur), beech (Fagus sylvatica), maple (Acer platanoides), and birch (Betula pendula). Smaller areas with mountain pines (Pinus mugo) are found in the north-west flank of Lake GT.

In the Black Forest, we investigated three small headwater cirque oligotrophic brown water lakes: Herrenwiesersee (HS), Schurmsee (SS), and Glaswaldsee (GS). The anthropogenic pressure on the lakes is relatively low as there are no settlements in the catchments (Fig. 1C). The catchment topography was formed in a cirque excavated by glaciers, so the cupped sections of the three lakes are relatively steep. Acid podzols and Triassic sandstone rubble dominate the catchments at those sites (BGR Geoviewer 2018). The catchments are densely covered by coniferous forests dominated by spruce (Picea abies). An overview of the geographical characteristics of the investigated lakes and their catchments is shown in Table 1. Meteorological data on annual precipitation, frost days, solar radiation, and sunshine duration are shown for both studied areas in the Supporting Information Fig. S1. The graphical representation of all catchment areas is shown in Supporting Information Figs. S2–S7.

Sampling and sample treatment

Litter and soil samples

We collected forest litter and soil samples (n = 69), including a few tree leaves/needles, from five lake catchments (GS,
SS, GT, KT, and DT) from at least three randomly chosen plots per stand in April 2017. For more details, see the Supporting Information (Interactive map file: sampling locations.kmz). Soil samples were taken using a Pürckhauer-corer and five subsamples were combined from each sampling plot (1 × 1 m). Samples encompassed fresh and dry leaves/needles, two litter horizons (Oi: undecomposed surface litter; Oa: fully decomposed humus layer) and one mineral soil layer at a depth of 15–20 cm. These four individual sample compartments are described below as foliage, litter, organic, and mineral soil. Sampling of solid samples was conducted using Nitril gloves and cleaned stainless steel equipment. All samples were packed into plastic bags (Whirl-Pak, U.S.A.), stored at 4°C during the fieldwork and at −20°C in the laboratory until they were freeze-dried, and ground using an agate ball mill (Retsch, Germany).

Lake water phase

In total, 226 water samples were collected from the GS, SS, GT, KT, and DT lakes between June 2016 and November 2017. Samples were taken at 1 m depth intervals from the surface to the maximum water depth using a 1.5-liter water sampler (Uwitec, Austria). In addition, water samples from the direct inflows and outflows of the lakes were taken if permitted by hydrologic conditions. Electrical conductivity (EC), pH, and oxygen concentrations were determined in the field using portable meters (Cond 3110, pH 3110, and Multi 3510 IDS, WTW GmbH, Germany). All samples were filtered in the field using nylon membrane filters (0.45 μm, Carl Roth GmbH, Germany) after preconditioning the filters using a small amount of sample and stored in disposable polypropylene tubes (50 mL, Sarstedt, Germany) at 4°C until analysis. Subsamples for dissolved organic carbon (DOC) and cation determination were not stabilized and measured within 2–3 d after sampling. Subsamples for major and trace element analyses were stabilized by adding 1% (v/v) of distilled nitric acid (HNO₃, Merck AG, Germany). Subsamples for Hg analyses were stabilized following USEPA–Method 1631 using 1% (v/v) ultrapure BrCl–HCl solution (Merck AG).

Fig. 1. (A) Location of the two sampling areas in Germany. (B) Eastern Harz Mountains. (C) Northern Black Forest. The studied lakes are indicated with black stars, mountain summits with black crosses.
Table 1. Geographical characteristics by area and its dominant forest type, lake designation, code, location, altitude, lake depth, surface area, catchment area, and their ratios of all investigated lakes.

| Area | Forest type | Lake            | Code | Location       | Altitude (m a.s.l.) | Lake depth (m) | Surface area (km²) | Catchment area (km²) | Surface/catchment |
|------|-------------|-----------------|------|----------------|---------------------|----------------|--------------------|---------------------|------------------|
| BF   | Coniferous  | Herrenwiesersee | HS   | 48°40’6.84"N  | 8°17’43.90"E       | 830            | 10                 | 0.018               | 0.369            | 1:21             |
|      |             | Schurmsee       | SS   | 48°36’47.61"N | 8°19’7.86"E        | 795            | 13                 | 0.015               | 0.506            | 1:34             |
|      |             | Glaswaldsee     | GS   | 48°25’32.53"N | 8°15’44.40"E       | 839            | 11                 | 0.030               | 0.592            | 1:20             |
| HZ   | Deciduous   | Großer Siebersteineich | GT | 51°42’30.84"N | 11°10’31.98"E      | 319            | 12                 | 0.034               | 2.950            | 1:88             |
|      |             | Kleiner Siebersteineich | KT | 51°43’8.26"N  | 11°11’43.84"E      | 266            | 5                  | 0.017               | 1.200            | 1:70             |
|      |             | Großer Dachsteich | DT   | 51°42’49.28"N | 11°1’25.83”E       | 276            | 5                  | 0.026               | 1.360            | 1:52             |

Sediment traps

Sediment traps (n = 13) were deployed in all six lakes. The traps consisted of two acrylic glass tubes (single tube surface area of 55.4 cm²), two connected sediment containers (2 L, Polypropylene, Nalgene, U.S.A.), a weight anchor, and buoys. At each location, the top edge of the sediment trap was placed approximately 1.5 m above the sediment surface. The sediment traps were installed in the lake water column during spring and autumn (122–208 d) (Supporting Information Fig. S9C). To evaluate the seasonal influence on the sedimentation rate and/or the trap material composition, the sediment traps GS2017 and GS2018 in the Harz Mountains were also installed during the winter period, for 335 d and 487 d, respectively. The sedimentary residues, including the overlying water column, were stored at −20°C. The entire trap content was freeze-dried and ground in an agate mortar. Subsamples of sediment trap material were digested using bi-distilled nitric acid (HNO₃, Merck AG) for 1 h at 180°C (CEM Mars 5 Microwave system, U.S.A.).

Sediment cores

To investigate the temporal variations in element concentrations, four short lake sediment cores (13–29 cm) were taken from the lake centers using a gravity corer (Uwitec, Austria). After sampling, cores were kept upright and stored at 4°C, sliced at a resolution of 1 cm (HS) or 2 cm (SS, GS, GT) using acid-washed plastic tools. Samples (n = 64) were packed into plastic bags (Whirl-Pak, U.S.A.), stored at −20°C, freeze-dried, and ground using an agate ball mill (Retsch, Germany).

Laboratory and data analyses

Solid samples

The Hg content in the solid samples was analyzed by thermal decomposition followed by preconcentration of Hg on a gold trap and cold vapor atomic absorption spectrophotometry using a DMA-80 direct mercury analyzer (Milestone, Italy). The total concentrations of Al as an indicator of mineral matter fluxes from the catchment to the lake sediment cores were analyzed using energy-dispersive X-ray fluorescence (Cheburkin and Shotyk 1996). The major and trace element concentrations in the digests from sediment trap material were determined either by ICP-OES (Varian 715-ES, Agilent, U.S.A.) (Mg, K, Ca, Fe, and Cu) or by ICP-MS (Agilent 7700, U.S.A.) (Al, Zr, V, and Cr). The total carbon and nitrogen concentrations in the solid samples were determined using a Euro EA 3000 elemental analyzer (HEKAtech GmbH, Germany). The samples of trap material from five of the investigated lakes (GS, SS, GT, KT, and GT) were analyzed by FTIR using a Vector 22 FTIR spectrometer (Bruker, Germany) in absorption mode, with subsequent baseline subtraction on KBr pellets (200 mg dried KBr and 2 mg sample). Measurements were recorded from 4,500 to 600 cm⁻¹. Sample imaging of the trap material was conducted using a Phenom XL desktop (Thermo Fischer Scientific, U.S.A.) scanning electron microscope (SEM). The SEM conditions utilized for imaging were a 15 kV accelerating voltage, approximately 8 mm working distance, x5000 magnification, and a backscattering electron detector.

Water samples

Total Hg concentrations in the water samples were measured by means of cold vapor atomic fluorescence spectrometry (mercur, Analytik Jena AG, Germany) after removing excess BrCl by adding hydroxylammonium chloride (0.5% v/v) and stannous chloride Hg²⁺ reduction.

Major and trace element concentrations were determined either by ICP-OES (Varian 715-ES, Agilent, U.S.A.) (Mg, Ca, Fe, and Cu) or by ICP-MS (Agilent 7700, U.S.A.) (Al, Zr, V, and Cr). Filtered water samples were analyzed for total DOC using
thermo-catalytic oxidation at 950°C with subsequent non-dispersive infrared (NDIR) detection of CO₂ using a TOC analyzer (multi N/C 2100, Analytik Jena AG, Germany). Concentrations of nitrate and chloride were analyzed by ion chromatography with chemical suppression and conductivity detection (761 Compact IC, Metrohm, Switzerland).

**Analytical quality assurance**

For each analytical technique, blanks and sample replicates were determined. Standard reference materials (SRMs) were analyzed to assure analytical quality. The following SRMs were used: trace metals in surface water SPS-SW1 and RTC 1-WP, river water NRC SLRS-5, Thames LGC6019, MAURI-09, and ORMS-5, soil NCS DC 73322, sediment CANMET-LKSD-4, IAES-SL-1, and NCS DC 73312, apple leaves NIST 1515, pine needles NIST 1575a, and poplar leaves GBW07604. For more details, see the Supporting Information Table S5.

**Modeled atmospheric deposition**

To compare atmospheric Hg fluxes in the two study areas, we estimated the local Hg atmospheric deposition (HgAT) to the forest soils using an ecosystem-specific modeled data set for 2016 and 2017 (MSC-E 2018). The atmospheric deposition rate of Hg was provided by MSC-E of EMEP (Co-operative Programme for Monitoring and Evaluation of Long-range Transmission of Air Pollutants in Europe). Deposition fluxes of Hg across Germany are based on the chemical transport model “MSCE_HM” (Travnikov and Ilyin 2005). For this study, the data set “evergreen needleleaf forests” was chosen for the locations in the Black Forest. Owing to the presence of mountain pines at the northwest flank of Lake GT, the data set “mixed forest” was used to estimate Hg deposition rates in the deciduous forest-dominated Harz Mountains.

**Results**

**Hg soil concentrations**

The results of soil Hg analyses for all studied sites show that mean Hg concentrations were highest in the organic-rich and lowest in the mineral horizon. The mean Hg concentration in litter was generally lower than in the organic-rich, but higher than that in the mineral horizon. Mean concentrations of Hg in the organic horizon in the Black Forest (0.284 ± 0.128 μg g⁻¹) were slightly higher than those in the Harz Mountains (0.227 ± 0.099 μg g⁻¹). On the other hand, the mineral horizon had lower mean concentrations of Hg in BF soils (0.045 ± 0.017 μg g⁻¹) than in HZ soils (0.090 ± 0.041 μg g⁻¹). The Hg concentration in litter at the BF sites (0.140 ± 0.026 μg g⁻¹) was similar to that at the HZ sites (0.141 ± 0.042 μg g⁻¹). However, the variability of Hg concentrations in the litter from deciduous forest stands was higher (29.8% vs. 18.8% RSD) than in the coniferous stands. In this study, we investigated a few leaves taken directly from the trees. The concentration of Hg in withered beech (0.075 μg g⁻¹) and oak (0.067 μg g⁻¹) leaves sampled in the Harz Mountains in November 2017 was higher than that in 1-yr-old green spruce needles (0.038 μg g⁻¹ at GS and 0.053 μg g⁻¹ at SS). In contrast, the Hg concentration in the shoots of beech leaves sampled in March 2017 was significantly lower (0.003 μg g⁻¹) than that in any other sample, indicating Hg enrichment by up to a factor of ~25 during the vegetation period.

The type of vegetation affects soil formation and cycling of OM and trace elements in terrestrial ecosystems (Smolander et al. 2005; Hansson et al. 2011; Wang et al. 2019). Thus, we determined the relationship between Hg concentrations and the chemical composition (C and N) of the soil samples (Fig. 2). The Hg (μg g⁻¹) to C (g g⁻¹) mass ratio indicates the degree of saturation of the soil OM with Hg. In both study areas, Hg/C ratios increased with depth (Supporting Information Fig. S8). In the Black Forest, average Hg/C ratios in organic (0.92 ± 0.09 μg g⁻¹) and mineral (1.28 ± 0.38 μg g⁻¹) soil horizons are similar to values reported from comparable forest soils in Poland and the Czech Republic (Navrátil et al. 2016; Gruba et al. 2019). In contrast, the average Hg/C ratios in organic (3.10 ± 1.12 μg g⁻¹) and mineral

![Fig. 2](image_url)
Chemical characteristic of the lake water phase

Figure 3 shows water depth profiles for pH, temperature, nitrate (NO$_3^-$), DOC, Hg, and Al of one lake representative for each study area (May and November 2017). The concentrations of dissolved Hg were more than twofold higher and showed higher variance in the BF lakes (8.1 ± 5.6 ng L$^{-1}$) than in the HZ lakes (3.0 ± 1.9 ng L$^{-1}$). Similarly, concentrations of DOC were about twice as high in the BF lakes (13.8 ± 5.4 mg L$^{-1}$) than in the HZ lakes (6.8 ± 1.9 mg L$^{-1}$).

The DOC values in the BF lakes decreased toward the lake's surface, which was not observed in the HZ lakes. Mean pH values of the lake water profiles as well as of the in-and outflows show that pH is lower at the BF sites (4.5 ± 0.3) compared to the HZ sites (7.7 ± 0.5). The high pH values in the upper 5 m of the HZ lakes (up to pH 9) clearly indicate high photosynthesis activity in the HZ lakes, which seems to be low in the BF lakes. EC was lower at the BF sites (23.1 ± 8.5 μS cm$^{-1}$) compared to the HZ sites (373 ± 96 μS cm$^{-1}$). Even greater differences were observed at the two study areas for Al (BF: 557 ± 135 μg L$^{-1}$ and HZ: 14.3 ± 8.8 μg L$^{-1}$). At the coniferous sites, Pearson correlation coefficients (r) for Hg to DOC ($r = 0.82$) and Al ($r = 0.80$) as well as DOC to Al ($r = 0.77$) were significant and positive, while r values for the same elements showed lower positive correlations (i.e., $r < 0.50$) or even no significant relationship to Hg or DOC in the HZ lakes (Supporting Information Table S4). The water depth profiles of Hg and Al corresponded to the DOC concentration at Lake SS (panel A) and demonstrate an enrichment of these elements in the water layer above the sediment, which could not be observed in the HZ lakes. Furthermore, the Hg, DOC, and Al showed higher concentrations in the inflow and outflow in November compared to May, while only minor differences in concentration between the two sampling dates were observed in the HZ lakes. NO$_3^-$ concentrations were generally higher in the BF lakes and did not show a pronounced decline in the photic zone as observed in the HZ lakes, where NO$_3^-$ concentrations were inversely related to pH, both indicating the generally higher phytoplankton productivity in the HZ lakes. Nitrate was the highest in the inflow of the HZ lakes, up to 3 mg L$^{-1}$, while it tended to be less than 1.5 mg L$^{-1}$ in the BF lakes.

Chemical composition of trap material

The trap material obtained from the two sampling areas showed large differences in quantity (Supporting Information Fig. S9C), texture, and color (Supporting Information Fig. S10A), biological residues derived from pollen, spores, and diatoms (Fig. 4A) as well as in their chemical composition. To better visualize these differences, the relative elemental composition is shown in relation to the simplified total amount (100%) of several major (Mg, K, Al, Fe, N, and C) elements. The non-normalized element concentrations of major and trace elements obtained in the sediment traps are given in Supporting Information Table S1. The normalized distribution of major elements indicated that OM is the major component in the traps from both sites. Normalized major elements (Fig. 4B) in the trap material from the BF lakes showed higher proportions of organic carbon (87%) compared to those found in the lakes from the Harz Mountains (62%). The mean relative proportions of nitrogen were similar at both sites (58%), whereas iron (14%), aluminum (12%), magnesium (3%), and potassium (3%) were a factor of 9, 4, 4, and 2, respectively, higher in the traps of the HZ lakes, indicating higher fluxes of soil-derived mineral matter to these lakes.

Trace elements (Supporting Information Fig. S10B) showed site-specific variation in the Black Forest, while element proportions, especially for copper, chromium, vanadium, and calcium, only slightly varied at the Harz Mountains. The highest concentrations (Supporting Information Table S1) of chromium (65 mg kg$^{-1}$) and vanadium (87 mg kg$^{-1}$) were found in the trap material of Lake KT. Both metals may have been emitted from a nearby country road (Supporting Information Fig. S3), which is consistent with the higher concentration of chlorine in the water phase, which may be attributed to the use of road salt during winter (Supporting Information Table S4). However, notable concentrations of copper were also observed in a sediment trap of the Black Forest, which is consistent with the geochemical signals found in the lake sediment of medieval mining activities near the lake catchments (Schütze et al. 2018).

Hg concentrations and accumulation rates in sediment traps

Sediment traps in the BF lakes showed low and relatively constant annual mass accumulation rates compared to the HZ lakes, where mass accumulation rates were up to a factor of 57 higher and varied from 1 yr to the next (Supporting Information Fig. S9C). Similarly, mean Hg accumulation rates in the BF lakes (32–144 μg m$^{-2}$ yr$^{-1}$) were up to a factor of 14 lower than in the HZ lakes (113–443 μg m$^{-2}$ yr$^{-1}$) (Fig. 5C). In contrast, the mean Hg concentrations (Fig. 5B) in trap material derived from the coniferous sites (0.57–0.89 μg g$^{-1}$) were up to a factor of 7.4 higher than at the deciduous sites.
The highest sediment trap Hg concentration was observed at Lake GS in the Black Forest during the summer of 2018, when the mean sunshine duration was high, and the mean precipitation was low (Supporting Information Fig. S9A,B). In comparison, the highest Hg accumulation rate was observed at Lake KT in the Harz Mountains in 2016, a year with an unusually warm springtime in Germany.

To understand the interdependencies between the catchment and in-lake processes, it is essential to differentiate between OM sources. Terrestrial plants have a higher C/N ratio (4.25) than aquatic algae (< 10) (Meyers and Ishiwatari 1993), although decomposed soil OM can have significantly lower C/N ratios than plant material. The mean C/N ratios of the trap material from the BF lakes (13.9 ± 3.0) pointed to a terrestrial dominated OM source, while the low C/N ratios in the trap of the HZ lakes (8.5 ± 0.6) indicate algae as the main source of OM in all three lakes (Fig. 5A). However, it has to be pointed out here, that the C/N-ratios in the catchment soils of the HZ lakes also had values below 10; thus, the C/N-ratios alone do not prove that algae is the main OM source. However, the high pH (> 8) in May (Fig. 3B) and the distribution of nitrate concentrations in the water column relative to the inflow clearly indicated the high phytoplankton productivity in these lakes. Moreover, the FTIR spectra (Fig. 6) also supported a higher influence of algae in the HZ lakes compared to the BF lakes.

Spectral information of trap material

The FTIR spectra analyses were performed using the Peak Analyzer Peak Fit Gaussian function (OriginPro 2018). The spectral information showed that the trap material from the lakes in both study areas was received from both

![Fig. 3. Chemical characteristics (pH, temperature, NO$_3^-$, DOC, Hg, and Al) of lake water, and in- and outflows of two sampling campaigns at two selected representative lakes: Lake SS (A) and Lake GT (B) in May 2017 (black symbols) and November 2017 (open symbols). Notice the different scaling for pH, NO$_3^-$, and Al concentrations at SS and GT.](image-url)
autochthonous and allochthonous sources (i.e., mixture of algal, microbial, and land plant detritus). This assumption was supported by absorptions peaks associated with polysaccharides (1080 cm$^{-1}$), humic and fulvic acids (1410 cm$^{-1}$), secondary amines (1640 cm$^{-1}$), and aliphatic compounds (2850 and 2920 cm$^{-1}$), which could be observed in all six lakes (Fig. 6 and Supporting Information Table S3). Polysaccharides were highly abundant in the HZ lakes, while the BF lake trap material was dominated by secondary amines and humic and fulvic acids. A weak peak associated with lignin (1210 and 1266 cm$^{-1}$) occurred in the BF lakes (HS, GS, and SS). The spectra of the trap material of the HZ lakes (GT, KT, and DT) had peaks associated with (bio-)genic silica (690 and 780 cm$^{-1}$) and a small peak indicating proteins (1543 cm$^{-1}$), typical for microalgae (Dean and Sigee 2006), which was not observed in the BF lakes.

**Sediment geochemistry derived from short cores**

In the BF lakes (Fig. 7A–C), sediment Hg concentrations generally decreased with depth and peaked at the sediment surface layer with 1.55 μg g$^{-1}$ at HS, 0.68 μg g$^{-1}$ at SS, and 0.90 μg g$^{-1}$ at GS. In contrast, the short core of the mesotrophic Lake GT in the Harz Mountains (Fig. 7D) shows the highest Hg concentration (0.25 μg g$^{-1}$) at a depth of 14 cm, with lower concentrations in the surface layer (0.12 μg g$^{-1}$).

The C/N ratios in all sediment cores show relatively constant values indicating that OM sources did not drastically change over time. The three sediment cores from the BF lakes had C/N ratios that varied between 10.9 and 16.1, with a mean value of 13.3, whereas the C/N ratios in the single HZ core were clearly lower (8.3 ± 0.33) than in the BF cores, as found in the sediment traps. Al concentrations were relatively constant in all sediment cores, indicating that strong erosion events were rare. Similar to those observed in the sediment traps, Al concentrations in the BF sediments (1.9 ± 1.2 wt.%) were approximately 50% lower than in the HZ lake sediment (2.9 ± 0.4 wt.%), indicating higher fluxes of lithogenic material at Lake GT.

**Discussion**

**Factors influencing the distributions of Hg in the soil horizons**

Concentrations of Hg in forest soils are highly dependent on tree species. In coniferous forests, the Hg content in OM-
rich horizons is often higher than in deciduous forests (Schwesig and Matzner 2000; Obrist et al. 2012; Navrátil et al. 2016). The results of this study confirm these findings, although the Hg concentrations in the organic-rich soil horizons of the BF soils were only slightly higher than those of the deciduous forests in the HZ. In addition to the uptake by leaves and the extent of litterfall, Hg concentrations in the soils of both sites might be additionally influenced by local climatic factors, such as annual precipitation, rainfall intensity, days with frost, solar radiation, sunshine duration, and wind speed (Hararuk et al. 2012; Rydberg and Martinez-Cortizas 2014; Siudek et al. 2016; Baptista-Salazar and Biester 2019). The number of frost days, solar radiation, and sunshine duration were comparable in both study areas (Supporting Information Fig. S1), whereas the annual precipitation rate is more than threefold higher in the Black Forest (1344 mm) than at the Harz Mountains (406 mm).

Based on the ecosystem-specific modeled data set, the estimated atmospheric Hg deposition rates at the BF study sites were 50% higher ($\sim 27 \mu g \text{ m}^{-2} \text{ yr}^{-1}$) than those at the Harz sites ($\sim 18 \mu g \text{ m}^{-2} \text{ yr}^{-1}$). The ecosystem-specific deposition rates for both study areas were comparable to those reported for other forest sites in the Czech Republic and the northern U.S.A. (9–34 $\mu g \text{ m}^{-2} \text{ yr}^{-1}$) (Blackwell and Driscoll 2015; Navrátil et al. 2016; St. Louis et al. 2019). Blackwell and Driscoll (2015) showed that atmospheric Hg deposition increases with elevation and the dominant deposition pathway shifts from litterfall in low-elevation hardwoods to throughfall in mid-elevation spruce and fir dominated forests. Throughfall in coniferous forests has been shown to increase Hg deposition in soils (Sheehan et al. 2006; Witt et al. 2009; Gerson et al. 2017). Furthermore, the needles of conifers have a higher surface area and roughness (i.e., leaf structure) which reduces air circulation and increases particle adsorption.

Fig. 5. Comparison of the composition of trap material installed during several years in all investigated lakes. Panel (A) shows the C/N ratio, marked with values of organic soil and a range typical for OM derived from algae (Meyers and Ishiwatari 1993). Differences in Hg concentration (B) and Hg accumulation rate (C), marked with a mean Hg atmospheric deposition value ($H_{\text{AT, ecosystem-specific EMEP-model for the year 2016}}$) for both studied areas in panel (C).
This promotes the adsorption and accumulation of Hg on needles and may result in increased Hg concentrations in the needles and the soil of coniferous forests. Based on these findings, the higher throughfall in coniferous BF stands (Demers et al. 2007; Witt et al. 2009) is likely the dominant Hg deposition pathway to

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**Fig. 6.** FTIR spectra of sediment trap material sampled in lakes in the Black Forest (HS, GS, and SS) and in the Harz Mountains (GT, KT, and DT). Extract of wavenumber intensities 600–3000 cm\(^{-1}\) (staggered for better visualization). Bands typical for the trap material are highlighted accordingly.

**Fig. 7.** Down-core profiles of the lake sediments of HS, SS, GS (Black Forest, A–C) and GT (Harz Mountains, D) present carbon/nitrogen ratios (yellow bars) as an indicator for organic matter sources, concentrations of Hg (red line), and concentrations of Al (gray dashed line) as an indicator of mineral matter erosion from the lake catchments.
the soils at the BF sites. Moreover, the mean altitude at the BF sites was ~ 530 m higher (Table 1) than at the HZ sites, which might further support higher atmospheric deposition in the BF. In contrast, deciduous forests are thought to show higher net Hg accumulation in litter than coniferous forests due to differences in litter dynamics (Demers et al. 2007; Navrátil et al. 2016; Wohlgemuth et al. 2020). On the other hand, the generally higher litter mass in deciduous forests leads to a dilution of Hg concentrations (Obrist et al. 2012), which would explain the slightly lower Hg levels in the organic soil of the deciduous forest sites in the HZ.

Coniferous forests produce acidic soils that facilitate higher humic acid-rich DOC production compared to deciduous forests (Navrátil et al. 2016). This is confirmed by the higher DOC concentrations in the BF lakes and their in- and outflows (Fig. 3), which was also reported in a catchment study of Lake HS (Steinhoefel et al. 2017). Due to the high binding affinity of Hg to DOM (Biester et al. 2004; Rydberg et al. 2010; Teisserenc et al. 2011), the higher DOC concentration in the BF lakes explain the high Hg concentration in, and also suggests higher Hg fluxes to the BF lakes as suggested in previous studies (Rydberg et al. 2015).

**Effect of soil erosion on sediment Hg concentrations and accumulation rates**

Although the mean Hg concentrations in the organic soil horizon in coniferous forests of the BF were only slightly higher than those in the deciduous forests of the HZ, Hg concentrations in the trap material and the sediments differed significantly between the two sites. Hg concentrations in the surface sediment layer of Lake GS are more than twofold higher than the highest Hg concentration found in the BF catchment soils. In the HZ, surface sediments of Lake GT show only half the maximum Hg concentration found in the catchment soils, which indicates a dilution process in the Harz lakes. The catchments of the six lakes are considerably larger than the lake surface area (Table 1); thus, sediment Hg accumulation rates depend, to a large extent, on Hg fluxes from the catchment soils (Rydberg et al. 2015). The amount of Hg transported from catchment soils to lakes is regulated by the flux of mineral particles relative to DOM and POM fluxes, where Hg concentrations in DOM and POM are usually higher than in mineral matter (Hermanns and Biester 2013). Previous studies have demonstrated a strong relationship between Hg concentrations in the sediment and DOM mobilization in catchment soils (Kolka et al. 1999; Ouellet et al. 2009). Dilution of organic bound Hg by mineral matter has been shown in several lake sediment studies (Thevenon et al. 2011; Hermanns and Biester 2013; Pompeani et al. 2018) and is also apparent in the sediment cores from our study sites (Fig. 7), where an inverse correlation between Hg and Al (mineral matter) concentrations has been observed. The different chemical compositions of trap material from the two sites clearly indicate that Hg concentrations in the HZ trap sediments were diluted by mineral matter and likely by autochthonous OM resulting from high phytoplankton productivity. In contrast, Hg accumulation rates in the HZ sediment traps were up to a factor of 14 higher than in the BF sediment traps, which cannot be explained by higher Hg fluxes from the catchments. Our results indicate that algae OM appear to dilute Hg concentrations in trap materials but also greatly enhance Hg accumulation rates. High Hg accumulation in sediments as a result of Hg scavenging by algal OM in the water column has been described in previous studies (Outridge et al. 2007, 2019; Zhang et al. 2014; Biester et al. 2018; Zaferani et al. 2018), although the source and mass balance of Hg in these studies was mostly unknown.

Hg concentrations in the HZ sediment traps (Fig. 5B) were within the range of total Hg concentrations found in the sediment traps (0.08–0.26 μg g⁻¹) of Lake Geneva found by Gascón Díez et al. (2016). They concluded that Hg and OM in the settling particles originated primarily from phytoplankton. Gascón Díez et al. (2016) observed also that Hg concentrations strongly decreased in the sediment traps of Lake Geneva during spring and remained low until autumn, when primary productivity decreased, indicating a dilution of Hg concentrations in traps by algal-derived OM similar to what has been observed in the Harz traps (Figs. 5B, 7 and Supporting Information Fig. S10B). In contrast, the chemical composition of the small headwater BF cirque lakes indicated dominant terrestrial Hg sources, predominately DOM-Hg fluxes, and revealed annual differences in Hg concentration in the trap material caused by changes in hydrological conditions. Furthermore, higher OM fluxes from the surrounding catchment resulted in higher Hg to C ratios (Supporting Information Table S1) in the BF sediment traps because terrigenous OM is generally older and more degraded than autochthonous OM (Ouellet et al. 2009; Jiskra et al. 2015).

Outridge et al. (2019) argued in a recent study that inconsistent findings in sediment Hg concentrations vs. the extent of primary productivity relationship in some northern lakes and shallow ponds might be due to differences in lake and catchment properties which limit the amount of Hg that is available for scavenging in the water column by algae. Accordingly, at similar atmospheric Hg deposition rates and comparable soil Hg content, a larger catchment area would result in higher Hg accumulation rates in the lake sediments. Thus, the higher amount of mineral matter found in the traps of the HZ lakes may imply that higher Hg accumulation rates are attributed to higher erosion or larger catchments to lake ratios of these sites.

Figure 8 illustrates the relationship between Hg accumulation in sediment traps and catchment area. Although there were only few lakes, the results indicate a moderate positive catchment area effect for the BF lakes and a negative relationship for the HZ lakes. Lake KT, which had the highest Hg accumulation rate, had a 2.5-fold smaller catchment area than Lake GT, indicating that catchment area does not explain the
high Hg accumulation rates in the HZ lakes. Moreover, precipitation rates in the BF study area were significantly higher than in the HZ, which facilitates high erosional fluxes in the BF lakes.

Role of aquatic productivity on Hg accumulation

Our results showed that differences in atmospheric Hg deposition fluxes, catchment area, or erosional fluxes cannot explain the high Hg accumulation rates in the HZ lakes. Similar to what has been observed in other lakes (Biester et al. 2018; Outridge et al. 2019) or the ocean (Zaferani et al. 2018), we suggest that water column Hg scavenging by sinking algal OM strongly increases Hg export from the water phase to the sediment in the HZ lakes. The chemical composition of the trap material of the HZ lakes (low C/N ratios, peptides, amides, and proteins indicated by the FTIR spectra) as well as in the water phase (high pH, low nitrate) gives a clear indication that OM in the HZ lake sediments is dominated by algal-derived OM in contrast to the BF lakes, where most OM is derived from catchment soils (Fig. 3B, Supporting Information Table S4). Furthermore, SEM images indicated a high abundance of diatom frustules in the trap material of the HZ lakes, while diatoms were found only rarely in BF traps.

To evaluate the effect of water phase Hg scavenging, we calculated the annual number of Hg water column depletions ($n_{\text{Hg}_{\text{dpl}}}$) necessary to balance the Hg flux to the sediment (Eq. 1). We neglected the Hg flux to the lakes by the inflow as well as Hg re-emission from the lake surfaces, which were not determined in this study. Here, we assume that the water column Hg inventory is derived from Hg transported to the lake by leaching/erosion as DOM/POM-Hg from the catchment and via direct atmospheric Hg deposition. Hg contributes to maintaining a steady Hg inventory in the lake. The Hg inventory ($H_{\text{gpwc}}$) was calculated using the mean Hg concentration in the water phase times the volume of the water column ($m^3$) above the trap.

$$n_{\text{Hg}_{\text{dpl}}} = \frac{H_{\text{gtrap}}}{H_{\text{gpwc}}}$$ (1)

Equation 1 is a simplified equation for calculating the number of Hg water column depletion events ($n_{\text{Hg}_{\text{dpl}}}$) necessary to obtain the Hg mass balance in the investigated lakes. $H_{\text{gtrap}}$ and $H_{\text{gpwc}}$ are in $\mu g m^{-2} yr^{-1}$.

Thus, Lake GS had a mean Hg concentration in the water column ($H_{\text{conc}_{\text{w,colum}}} = 11$ ng L$^{-1}$ (2017), which amounts to 105 $\mu g$ Hg in the water column above 1 m$^2$ of the lake bed and a water column ($V_{\text{w,colum}}$) of 9.5 m above the sediment trap. This corresponds to an Hg water inventory of 85 $\mu g m^{-2} yr^{-1}$. Using Eq. 1 results in $n_{\text{Hg}_{\text{dpl}}} = 0.8$, which explains the annual Hg flux of 85 $\mu g m^{-2} yr^{-1}$ to the sediments (Fig. 5C). The calculation of $n_{\text{Hg}_{\text{dpl}}}$ for the other BF lakes even show values below 0.8. Thus, a single Hg scavenging event in the lake water column could explain the entire annual Hg accumulation rate in the BF sediment traps. Due to this ancillary role of water phase Hg scavenging and the fact that the water phase Hg pool in the BF lakes is likely to be continuously refilled by inflow, atmospheric Hg deposition or erosion, and surface run-off, we conclude that the Hg mass balance in the BF lakes is mainly controlled by catchment fluxes, atmospheric deposition, and Hg export through outflows or Hg re-emission to the atmosphere and to a minor extent by export to the sediments.

In contrast, $n_{\text{Hg}_{\text{dpl}}}$ for the HZ lakes ranges between 4 and 55 (Table 2), indicating the large importance of the water phase Hg scavenging needed to build up the high Hg accumulation rates in the HZ lake sediments. Moreover, scavenging is not a process restricted to Hg. Al and trace elements were highly depleted in the water phase (Fig. 3) but accumulated in the sediment traps (Fig. 4 and Supporting Information Fig. S10) of the productive Harz Mountains lakes compared to the brown water lakes in the Black Forest.

The discrepancy in rates of sedimentation and Hg accumulation between Lake GT (113–152 $\mu g m^{-2} yr^{-1}$) with a catchment area of ~3 km$^2$ and Lake KT (384–443 $\mu g m^{-2} yr^{-1}$) with a catchment area of ~1.2 km$^2$ (Fig. 8) emphasizes the role of phytoplankton productivity for Hg accumulation in these lakes. The HZ lakes showed no significant differences in chemical composition or Hg concentrations in the water phase, but Lake GT is twice as deep as Lake KT (Fig. 5B and Supporting Information Fig. S10B, Table S1). Kerimoglu and Rinke (2013) demonstrated that cyanobacteria blooms occurred in a
reservoir, particularly during years with low water levels enriched with nutrients. High production of autochthonous OM in shallow lakes resulted in anaerobic conditions and slow OM mineralization in the water phase, resulting in high sedimentation rates and intense scavenging of trace elements. In contrast to the BF lakes, the environmental fate of Hg in the HZ lakes is characterized by a strong export of Hg from the water phase to the sediment and a relatively low Hg export through the outflow. Thus, the HZ lakes with high primary productivity appear to be large Hg sedimentary sinks, whereas the BF brown water lakes show much lower Hg accumulation in the sediments and higher (DOM-) Hg throughflow. Our results support the findings of a model approach on the changes in Hg cycling in the Baltic Sea attributed to algae bloom events (Soerensen et al. 2016). These authors found that eutrophication enhances Hg export to the sediment and reduces Hg re-emission to the atmosphere (Figure 9).

Conclusions

We compared Hg accumulation in oligotrophic brown water lakes and mesotrophic to eutrophic lakes and catchment soils at two forested areas in Germany which differ in catchment vegetation (coniferous vs. deciduous forest) and weather conditions. Hg concentrations in soils were comparable at both sites but were clearly higher in the sediments of the brown water lakes. However, Hg accumulation rates derived from sediment traps of the productive lakes, where algae blooms are common, were increased by a factor of 14 higher than those derived from brown water lakes. Our data suggest that Hg scavenging and sedimentation by algal OM is

### Table 2. Individual values of a simplified calculation to calculate the number of Hg water column depletion events (n Hgdpl) required to balance the Hg flux to the sediment in the investigated lakes.

| Area | Trap | HgAT (μg m⁻² yr⁻¹) | Hgtrap (μg m⁻² yr⁻¹) | Hg concw.column (ng L⁻¹) | Vw.column (L) | Hgpwc (μg m⁻² yr⁻¹) | n Hgdpl (n) |
|------|------|---------------------|----------------------|--------------------------|--------------|----------------------|-------------|
| BF SS 2016 | 27 | 41 | 4.3 | 11,500 | 49 | 0.8 |
| SS 2017 | 27 | 32 | 4.0 | 11,500 | 46 | 0.7 |
| GS 2016 | 26 | 73 | 12.5 | 9500 | 119 | 0.6 |
| GS 2017 | 26 | 85 | 11.0 | 9500 | 105 | 0.8 |
| HZ GT 2016 | 18 | 152 | 2.8 | 10,500 | 29 | 5.2 |
| GT 2017 | 18 | 113 | 2.7 | 10,500 | 28 | 4.0 |
| KT 2016 | 18 | 443 | 2.3 | 3500 | 8 | 55 |
| KT 2017 | 18 | 384 | 2.2 | 3500 | 8 | 48 |
| DT 2016 | 17 | 316 | 4.2 | 3500 | 15 | 21 |
| DT 2017 | 17 | 186 | 4.1 | 3500 | 14 | 13 |

*Above the sediment trap inlet.

**Fig. 9.** Conceptual transport pathways of Hg from the soil horizons to the lake’s water phase and lake sediments investigated in lake catchments of the Black Forest and the Harz Mountains, respectively.
primarily responsible for the high Hg accumulation rates found in the sediments of the productive lakes. Up to 55 times the water column Hg inventory was required to explain the fluxes in deciduous catchments while less than one depletion was needed for the brown water lakes. From this, we concluded that productive lakes are major sinks for catchment-derived Hg in contrast to brown water lakes, where throughflow and re-emission to the atmosphere are major export pathways for Hg and Hg accumulation in the sediments is less important. This further suggests that increasing eutrophication of aquatic systems as a result of high nutrient fluxes and increasing temperatures will turn more lakes into sedimentary Hg sinks. Due to the high OM content in the sediments and the resulting anaerobic condition, productive lakes are hot spots for methyl-Hg production and transfer to the aquatic food chain, which is subject to future increases.

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Conflict of Interest
None declared.