The Influence of Sulphate Deposition on the Seasonal Variation of Peat Pore Water Methyl Hg in a Boreal Mire

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Abstract

In this paper we investigate the hypothesis that long-term sulphate (SO4\(^{2-}\)) deposition has made peatlands a larger source of methyl mercury (MeHg) to remote boreal lakes. This was done on experimental plots at a boreal, low sedge mire where the effect of long-term addition of SO4\(^{2-}\) on peat pore water MeHg concentrations was observed weekly throughout the snow-free portion of 1999. The additions of SO4\(^{2-}\) started in 1995. The seasonal mean of the pore water MeHg concentrations on the plots with 17 kg ha\(^{-1}\) yr\(^{-1}\) of sulphur (S) addition (1.3 ± 0.08 ng L\(^{-1}\), SE; n = 44) was significantly (p < 0.0001) higher than the mean MeHg concentration on the plots with 3 kg ha\(^{-1}\) yr\(^{-1}\) of ambient S deposition (0.6 ± 0.02 ng L\(^{-1}\), SE; n = 44). The temporal variation in pore water MeHg concentrations during the snow free season was larger in the S-addition plots, with an amplitude of > 2 ng L\(^{-1}\) compared to +/− 0.5 ng L\(^{-1}\) in the ambient S deposition plots. The concentrations of pore water MeHg in the S-addition plots were positively correlated (r\(^2\) = 0.21; p = 0.001) to the groundwater level, with the lowest concentrations of MeHg during the period with the lowest groundwater levels. The pore water MeHg concentrations were not correlated to total Hg, DOC concentration or pH. The results from this study indicate that the persistently higher pore water concentrations of MeHg in the S-addition plots are caused by the long-term additions of SO4\(^{2-}\) to the mire surface. Since these waters are an important source of runoff, the results support the hypothesis that SO4\(^{2-}\) deposition has increased the contribution of peatlands to MeHg in downstream aquatic systems. This would mean that the increased deposition of SO4\(^{2-}\) in acid rain has contributed to the modern increase in the MeHg burdens of remote lakes hydrologically connected to peatlands.

Introduction

Historically, human activities have caused large emissions of mercury (Hg) to the atmosphere. This has contributed to widespread Hg pollution and increased concentrations in the biota e.g. fish. Deposition of Hg in the boreal region modelled by EMEP [1] show relatively low rates in the Nordic countries, deposition rates that also are verified by moss monitoring [2]. The concentrations of Hg in fish are higher, though, than in other regions with similar or even higher deposition rates [3]. One key to the difference between Hg deposition and the high degree of bioaccumulation is one particular species of Hg, methyl mercury (MeHg) that is most prone to bioaccumulation [4–6]. The cycling of MeHg in boreal catchments is complex and involves numerous biogeochemical and physical controls. Therefore, it is important to elucidate the impact of these factors on MeHg production.

Important sources of MeHg loadings to lakes are direct precipitation [7,8], run-off from wetlands [9–11], and in-lake methylation [12–14]. Mass-balance calculations, however have revealed that often neither precipitation, nor in-lake methylation can account for the total amount of MeHg in the catchment runoff [15] or in lakes [16–18]. The runoff of MeHg from catchments and the levels found in fish can vary considerably between different catchments with similar levels of atmospheric deposition [5,19]. This suggests that the MeHg species is produced within the catchments.

Boreal wetlands have been identified as an important source of MeHg to lakes [16,20–23] though there is considerable variability in the strength of these hotspots across the landscape [24–26]. One factor affecting the significance of wetland sources can be the proportion of mires [27] or riparian zones [28,29] in the catchment. While wetlands generally are considered sources of MeHg one alder swamp have actually been found to be a sink of MeHg [25]. Another possible factor in the catchment production of MeHg is sulphate (SO4\(^{2-}\)) deposition which stimulates the production of MeHg [9,12,30].

The importance of sulphate-reducing bacteria (SRB) for methylation of Hg is well-documented [12,31,32]. The possibility that SO4\(^{2-}\) in “acid-rain” has enhanced in-lake methylation was suggested by Gilmour and colleagues [12,33]. A number of studies now suggest that the availability of SO4\(^{2-}\) for SRB is a major factor regulating the concentration of MeHg in mire pore water. Branderem et al. [30] suggested SO4\(^{2-}\) reduction in the anoxic zone of mires and riparian wetlands as the functional explanation.
for the increased concentrations of MeHg in runoff from boreal wetlands. In that study addition of $SO_4^{2-}$, corresponding to 28 kg $SO_4^{2-}$ ha$^{-1}$, to a boreal peatland increased the pore-water concentrations of MeHg from 2 to 4 ng L$^{-1}$ within 24 hours. The MeHg concentrations, however, returned to ambient levels after 5 days. In constructed wetlands, Harmon et al. [34] found that $SO_4^{2-}$ amendments gave significantly higher MeHg pore water concentrations over the course of one year. In a mesocosm experiment, Mitchell et al. [35] found that $SO_4^{2-}$ additions significantly increased MeHg pore water concentrations, while additions of different carbon substrates alone did not have an effect. Combined $SO_4^{2-}$ and carbon (C) additions, however, gave the largest increases, providing an explanation of why hotspots of MeHg appear in mires where there are inputs of both C and $SO_4^{2-}$ [23].

An experiment to simulate the effect of atmospheric $SO_4^{2-}$ at the ecosystem scale by sprinkling $SO_4^{2-}$ solutions systematically across a mire for one snow-free season also found a significant influence on MeHg, both in pore water and in the runoff from the mire [9]. $SO_4^{2-}$ additions in that experiment were made on five occasions, and the effects on pore water concentrations varied considerably. This may have to do with the relatively short term additions of $SO_4^{2-}$ to the system, a feature common to most other studies on how $SO_4^{2-}$ influences MeHg in mires.

Pore [36] and run off [9] water from wetlands responded to experimentally elevated sulphur (S) deposition with an increased potential to produce MeHg. Branfireun et al. [36] found a tripling of peat pore water MeHg in plots exposed to two years of elevated S deposition (20 kg S ha$^{-1}$ yr$^{-1}$) compared to plots with the ambient deposition in N. Sweden (3 kg S ha$^{-1}$ yr$^{-1}$), five weeks after the most recent experimental $SO_4^{2-}$ amendment had been made. An almost equally large increase (2.4 times) in stream water MeHg concentrations were also found after experimental additions of $SO_4^{2-}$ of an experimental wetland in northeastern Minnesota [9]. Such long-term effects would greatly increase the potential significance of S deposition for MeHg-loading to the aquatic ecosystem.

Observation of elevated MeHg at a single point in time is not, however, sufficient to prove a persistent effect since other factors are also important for the regulation of $SO_4^{2-}$ reduction (and hence any sustained effect of $SO_4^{2-}$). These factors include the occurrence of competing electron acceptors e.g. oxygen which is mainly determined by the prevalence of saturated conditions [37,30], availability of easily degradable C for the SRB [35,39,40] and temperature [41]. To investigate the importance of these factors for the persistence and magnitude of the “acid-rain” effect, we studied the differences in temporal variation of peat pore-water MeHg between plots exposed to ambient (ca 3 kg S ha$^{-1}$ yr$^{-1}$) and five years of enhanced $SO_4^{2-}$ deposition (20 kg S ha$^{-1}$ yr$^{-1}$) in a low sedge mire in northern Sweden during one snow-free period of five months duration. On three occasions we sampled these plots at four depths to get a deeper profile of pore water MeHg. Our main hypothesis is that elevated pore-water MeHg concentrations are maintained by an increased pool of S (due to enhanced deposition) which can be recycled at the mire surface through reduction of $SO_4^{2-}$ (to e.g. HS$^{-}$) and subsequent re-oxidation of reduced S species to $SO_4^{2-}$. Since we are interested in ascertaining the effect of this process on MeHg loading to lakes and their biota, we have sampled the uppermost ten centimetres below the water table every week during the study period, regardless of the water table depth relative to the mire surface. This is the water most likely to compose the runoff from wetlands to downstream lakes [42].

Materials and Methods

Field Experimental Setup and Sampling

The pore water samples were collected from a low sedge mire, Degero Stormry (Lat. 64°11’N, Long. 19°33’E, altitude 270 m a s l) situated 70 km from the coast of the Gulf of Bothnia, Sweden. The experimental site is part of the Vindeln Research Forests, Swedish University of Agricultural Sciences. The experimental setup was established in 1995 and is based on the manipulation of 20, 4 m$^2$ plots, each of which are isolated from the surroundings by a 0.5 m deep plastic frame inserted to a depth of 0.4 m in the peat. Together with one extra control plot without plastic frames, there were a total of 21 experimental plots. In this study we utilised four of these plots, two receiving ambient levels of S (3 kg S ha$^{-1}$ yr$^{-1}$) from the precipitation, and two plots receiving experimental additions, applied at five occasions during the snow free period, which together with the ambient input equals 20 kg S ha$^{-1}$ yr$^{-1}$. A thorough description of the experiment is given in Granberg et al. [43] and Wiedermann et al. [44].

Pore water from these plots was sampled every week between May 26th and October 20th in 1999, 0–10 cm beneath the groundwater level (GWL) at the time of sampling. In addition, pore water was sampled from 10–20 cm, 20–30 cm, and 30–40 cm beneath the GWL on June 23rd, July 20th, and August 25th, 1999. The actual GWL was measured at 5 evenly distributed points within each plot. Runoff water in the stream draining the mire was also sampled each week, starting June 23rd.

Subsamples for dissolved organic carbon (DOC) were kept cool and in the dark before analysis within one month after sampling. The subsamples for anions were preserved by freezing, and the subsamples for cation analysis were preserved with suprapur sulphuric acid. The anions and cations were all analyzed within six months of sampling.

The DOC was determined by analytical combustion (Shimadzu, DOC-500). Pore water concentration of $SO_4^{2-}$ was determined by suppressed ion chromatography (Shimadzu, UV detector SPD-6A) using 0.5 mM sulfosalicylic acid as the mobile phase. After filtration, pH was determined in all pore water samples. Total S, $H_2SO_4$ and MeHg in the solid peat were determined on freeze-dried and ground peat samples taken at 0–5 cm, 5–10 cm, 10–20 cm, and 20–40 cm beneath the surface of the sphagnum moss. The solid samples were collected on June 24th, 1999. For total S, the samples were digested with HNO$_3$+ HClO$_4$ and thereafter analysed using Inductive Coupled Plasma – Atomic Emission Spectrometer (ICP – AES, Perkin Elmer).

Pore water sampling procedure. Pore water samples were collected with a 70 cm long, custom-made, Teflon sampler (13/6 mm, outer/inner diameter) equipped with 4 holes (3 mm diameter) for every 1 cm of the lower 10 cm i.e. a total of 40 holes. At the end of the sampler there was a removable conical tip to facilitate penetration of the peat surface. The sampler was connected to a 500 mL transfer bottle (Pyrex®) by Teflon tubing (8/6 mm, outer/inner diameter), and the transfer bottle was connected to a custom-made, portable vacuum pump. Even though the air flows in the direction from the transfer bottle to the vacuum pump, diffusion of Hg originating from the pump constitute a mass transfer in the opposite direction that potentially might contaminate the pore water samples. To avoid potential contamination from the vacuum pump, gold traps were installed between the vacuum pump and the transfer bottle or the filter holder.

Samples were collected by inserting the sampler 10 cm below the actual GWL and then pumping the pore water into the transfer bottle. Approximately 100 mL of water was collected from
5 evenly distributed spots within each experimental plot to create a pooled sample of 500 mL. Approximately 150 mL of the pooled sample was then filtered through an acid washed membrane filter (45 mm diameter, 0.45 µ, cellulose nitrate 100%, sterile, Millipore®) using a 500 mL filter holder (Nalgene®). Filter holders were connected to a manifold which in turn was connected to a portable vacuum pump (Prenart equipment ApS, Frederikssberg, Denmark). This set-up allowed four samples to be filtered simultaneously. Immediately after filtration, each sample was transferred to a Teflon bottle (Nalgene®), put in a plastic bag, and then kept cool on dry ice in the dark. For stream water, a grab sample was taken directly in a similar Teflon bottle without filtration due to the low content of particulate matter. The samples were stored in the laboratory at +4°C until analysis. The MeHg and total Hg in the pore water were always determined within a maximum of five days.

Separate tests before the start of the investigation showed negligible losses of MeHg during five days of storage in the dark and at 4°C. In an earlier study [45] the stability of spiked (1.0 ng/ L) MeHg in river water was determined and only minor losses of MeHg during the storage of samples was found. After 15 days storage in Telfon containers without additions of preservation reagents the mean concentration of MeHg was decreased by 6%. (The concentration of MeHg originally present was 0.35 ng L⁻¹ and 4.1 ng Hg²⁺ L⁻¹; TOC was 15 mg L⁻¹ and pH was 5). Prior to each sampling occasion, the Telfon bottles, transfer bottles, filtering units, filters, Telfon tubing and the Telfon sampler were acid washed with HCl (p.a. quality, pH = 2) overnight at 70°C, rinsed three times with Millipore water and thereafter kept in plastic bags prior to the sampling. Plastic gloves were always used during washing and sampling, or when otherwise handling the equipment.

Determination of Hgtot and MeHg in Peat, Pore Water and Stream Samples

Chemicals were of analytical-reagent grade unless indicated otherwise. All standards and solutions were prepared using Millipore deionized water (Millipore, Bedford, MD, USA).

Digestion and Determination of Hgtot

**Peat samples.** Some 200 mg of peat (wet weight) were weighed and placed into Telfon tubes. Then 4 ml of HNO₃ and 4 drops of HCl were added to samples that were left overnight on a clean bench. After this 10 drops of H₂O₂ were added and samples were subsequently heated to 80°C for 8 hours. 4 ml of H₂O were added and after centrifugation the clear solution was used for quantification of Hg. The Hg species in the samples were reduced using SnCl₂ (30%) to Hg⁰ that was purged onto gold traps using argon gas. The Hg adsorbed to gold traps was thermally desorbed followed by analysis using cold vapour atomic absorption spectrometry (CVAAS) at 254 nm (FIMS, Bodenseewerk Perkin Elmer, Überlingen, Germany) [46]. All samples were analysed at least two times. Tests with addition of 50 and 100 µg HgCl₂ L⁻¹ to one of the samples before acid treatment gave recoveries of 100.9% and 98.7%. Certified reference material (Light sandy soil, BCR 142 with a certified content of Hg of 67±11 ng g⁻¹) was analysed in the series and gave a value of 74.0±11.0 ng g⁻¹ (n = 3). The SO₄²⁻ salt used in the field application treatments was analyzed for Hg³⁺, which was found to be below the detection limit of 40 pg Hg g⁻¹ [36 criterion [47]].

**Pore and stream water.** Pore water (30 mL), HNO₃ (3 mL, ultra pure), and HCl (0.5 mL ultra pure) were added to quartz flasks, which thereafter were subjected to UV radiation for seven hours. After digestion, 5 mL of a solution containing 30% SnCl₂ and 5% H₂SO₄ was added to the sample. The Hg⁰ formed was thereafter removed during 10 minutes under a constant flow of Argon. Hg⁰ was amalgamated on a gold trap and thermally desorbed and detected by CVAAS as described above [46].

**Extraction and Derivatization Procedure for the Determination of MeHg**

**Pore water and streamwater.** A modified in situ ethylation method using tetrathylborate [48,49] was employed for analysis of MeHg concentration in pore water samples [50]. Derivatized Hg species, trapped to Tenax (Supelco, Bellefonte, PA, USA) columns was thermo-desorbed at 200°C and injected into the GC for determination by Microwave Induced Plasma – Atomic Emission Spectrometer (MIP-AES) [51].

**Peat Samples**

A solvent extraction (0.03 M CuSO₄ (3 mL), 0.38 M KBr (3 mL), and CH₂Cl₂ (5 mL)) method [52] followed by ethylation using tetrathylborate [49] and GC separation and MIP-AES detection were used for analysis of the total MeHg content in peat samples [51]. The absolute detection limit for MeHg determined with the in situ ethylation method was 1.7 pg L⁻¹ as Hg based on three times the relative standard deviation of repeated blank measurements [47]. For quality assurance of the in situ ethylation procedure standard additions of MeHg were performed to determine recoveries. These were found to be 0.90–1.05 for an increase of MeHg concentrations in the range of 1 ng L⁻¹. The ethylation method provided accurate results (within 2% of the certified value) for MeHg when analysing certified reference materials (Tort 2; Dolt 2) and it provided good agreement with results for MeHg in brackish water 0.12±0.009 ng/L compared to 0.1±0.002 ng L⁻¹ for flow injection-liquid chromatography-CVAAS. For the comparison three determinations were performed with each method. The precision of the in situ ethylation method was typically better than 10% for the pore water samples. Normally one analysis was performed for each pore water sample.

**Statistics**

To compare the pore water MeHg concentrations between the plots with 20 kg S ha⁻¹ y⁻¹ (HighS) and 3 kg S ha⁻¹ y⁻¹ (LowS) plots, General Linear Model (GLM) statistical analysis (SYSTAT inc.) was run with treatment (HighS, LowS) as the category variable, groundwater level (GWL) as the independent variable and MeHg concentration as a dependent variable. The GLM was also used to compare the pore water MeHg concentrations at different depths, with treatment, depth, and date as category variables, while the concentration of MeHg was the dependent variable.

Statistical comparison of the concentration of MeHg, Hgtot, and S₄ (dependent variables) in the peat organic matter between the HighS and LowS plots was also made using GLM. Here, the category variables used were: treatment (HighS or LowS) or depth (0–5 cm, 5–10 cm, 10–20 cm, 20–40 cm). GLM was also used for comparing pore water pH (dependent variable) between the HighS and LowS plots (category variable). All pairwise comparisons within each variable were made using Tukey’s test.

**Results**

The concentrations of pore water MeHg were always measured in the top ten centimetres below the water table. The depth of the mire water table relative to the mire surface varied 15–20 cm during the measurement period. The variation in pore water MeHg concentrations may therefore result from both seasonal
variation in production/consumption of MeHg at each depth as well as differences in production/consumption between depths. The seasonal mean of the pore water MeHg concentrations (1.3 ng L⁻¹±0.08, SE; n = 42) from the plots receiving additional SO₄²⁻ (HighS) was significantly (p<0.0001) higher than the MeHg concentrations (0.6 ng L⁻¹±0.02 SE; n = 42) in the plots receiving ambient levels of SO₄²⁻ (Fig. 1A). A pronounced, systematic seasonal pattern in MeHg concentrations was found in the HighS plots, but not in the plots receiving ambient S deposition (3 kg ha⁻¹, LowS). Generally for both HighS plots the pore water MeHg level was highest from the beginning of the summer until late July. Thereafter the MeHg concentrations decreased to the lowest levels from the middle of August to the beginning of September. The pore water concentrations of MeHg then increased again (Fig. 1A). In the LowS plots, the amplitude of MeHg variation was lower (a standard deviation of 0.11 ng L⁻¹, with a range of 0.48 ng L⁻¹ between the highest and the lowest values) than in the High S plots which had a standard deviation of 0.53 ng L⁻¹ and a range of 2.55 ng L⁻¹ (Fig. 1A).

The concentrations of pore water MeHg in the HighS plots were positively correlated (r² = 0.21; p = 0.001) to the GWL at the time of sampling, with the lowest concentrations of MeHg during the period with lowest GWL (Fig. 1A, B). For the LowS plots there was no correlation between MeHg concentrations and the local groundwater table (p = 0.87) (Fig. 1A,B).

The mean pore water concentrations of MeHg decreased with depth from the current GWL down to 40 cm below, for both the LowS and HighS plots (i.e. samples were taken at a maximum depth of 60 cm below the vegetation surface; Fig. 2). The decrease in MeHg concentrations with depth were more pronounced at the plots with enhanced SO₄²⁻ deposition than those with ambient SO₄²⁻ deposition levels (Fig. 2). For the LowS plots the MeHg concentrations at the deepest sampling depth (30–40 cm) were significantly lower than the three more superficial sampling depths for all three dates of sampling (Fig. 2, upper panel). In the HighS plots the pore water concentrations of MeHg were significantly (p<0.0001) higher in the two upper sampling depths compared to the two lower sampling depths (Fig. 2, lower panel).

The MeHg concentrations in the stream runoff (Fig. 1A, mean 0.6 ng L⁻¹±0.06, SE; n = 13) were very similar to the concentrations measured just below the fluctuating groundwater surface on the ambient deposition plots (0.6 ng L⁻¹±0.02, SE; n = 42) both with regards to absolute concentrations and variability over the course of the year.

The pore water concentrations of HgSO₄ varied between 0.3 ng L⁻¹ (detection limit) and 30 ng L⁻¹ during the sampling period. No correlations (p>0.1) between the concentrations of MeHg and total Hg were found in any of the four plots, and there were no differences in these concentrations between the HighS plots and the LowS plots (p>0.1).

The pore water concentrations of SO₄²⁻ were analysed for the period between June 16th and September 22nd. Until the beginning of August, during the period with the highest pore water MeHg, the concentrations of SO₄²⁻ in the HighS plots were very low, often below the detection limit (0.5 mg SO₄²⁻ L⁻¹) (Fig. 3A). However, during the period when the pore water MeHg concentrations in the HighS plots were lowest (and thus closest to the more stable MeHg concentrations in the LowS plots), the highest SO₄²⁻ concentrations in the HighS plots were measured. The SO₄²⁻ concentration in the HighS plots reached their highest levels when the GWL’s were deepest, between 15 and 21 cm below the mire surface (Fig. 3A, B). In the LowS plots the pore water SO₄²⁻ concentrations were always below the detection limit (data not shown).

A seasonal trend in pH was observed in the HighS plots (Fig. 3B) with a slow decrease in pH from 4.4 ±0.1 SE, n = 2) to 3.8 ±0.02 SE, n = 2) between June 16th and October 13th (Fig. 3B). The corresponding pH values in the pore water samples from the LowS plots were 4.0 ±0.02 SE, n = 2), and 3.9 ±0.02 SE, n = 2), respectively (data not shown). The seasonal mean pH in the pore water from the HighS plots (4.0±0.2 SE, n = 36) was significantly different (p = 0.002; N = 72) from the seasonal mean pH in the LowS plots (3.9±0.1 SE, n = 36).

There was a small but steady increase in the concentrations of DOC during the sampling period except in the beginning of June when the DOC concentrations in the LowS plots increased to approximately twice the value of the HighS plots (Fig. 4). The highly elevated DOC values on August 25th coincided with the deepest sampling depth for each of the plots (Fig. 1B).

The amount of S in the peat organic matter ranged between 0.05% and 0.16% (Table 1). There was no significant correlation between total S content in the peat and pore water MeHg. There was no significant difference in the concentration of MeHg in the peat between the LowS and the HighS plots (Table 1). However, in one of the HighS plots the concentration of MeHg in the organic matter was substantially larger compared with the LowS plots (Table 1).

The concentrations of total Hg were several orders of magnitude higher than the concentration of MeHg in the peat organic matter (Table 1). In the LowS plots there were significantly (p<0.001) lower concentrations of Hg in the upper peat (0–5 cm) compared to the lower depths (Table 1). The concentrations of HgSO₄ in the HighS plots were similar down to 20 cm peat depth, and thereafter the HgSO₄ concentrations increased significantly (p = 0.026) (Table 1). The peat organic matter HgSO₄ concentrations were significantly (p<0.001) different between the HighS and the LowS plots. The amount of Hg-tot in the solid phase of the upper 5 cm was significantly (p = 0.001) higher in the HighS plots compared to the LowS plots. The salt used (Na₂SO₄) to experimentally increase the load of S to the plots would cause an increase of 40 pg assuming that all Hg in the salt added over five years is retained. This amount is three orders of magnitude less than the difference between the Hg in the upper 5 cm of the HighS plots relative to the LowS plots. So the content of HgSO₄ in the HighS plots do not emanate from the Na₂SO₄ additions.

Discussion

Boreal mires have earlier been identified as an important source of MeHg loading to surface waters [20,21,27,53]. There is increasing awareness that SO₄²⁻ is an important control on the strength of mires as a MeHg source. A number of short term studies have seen this after one to half a dozen SO₄²⁻ amendments, during the course of up to one year [9,30,35]. The potential for chronic (multiple years) of SO₄²⁻ addition to increase the mire pore water MeHg concentration was demonstrated by Branfireun et al. [36] based on data from one point in time during 1997 from the same experimental site used in this study. Our results from the entire snow-free period of 1999 after five years of SO₄²⁻ treatment provide evidence that the elevation of superficial peat pore water MeHg by long-term SO₄²⁻ deposition can be persistent and endure throughout the snow-free period. The plausibility of peat pore water just below the water table being the source of the MeHg in runoff from the study wetland was confirmed by the close correspondence between the MeHg concentrations in the superficial pore water of the LowS plots receiving ambient atmospheric deposition and the stream draining.
the study mire (Fig. 1A). Furthermore, the water table level is shown to be a major factor controlling the magnitude of the MeHg concentration enhancement in peat pore water on the sites with increased S deposition.

We believe that the persistent elevation of pore water MeHg concentrations and the connection to water table fluctuation can be explained according to the following conceptual model. Pore water MeHg is produced by SRB's through the reduction of $\text{SO}_4^{2-}$ [12,31,54]. Sulphur deposited from the atmosphere to the mire surface is cycled between $\text{SO}_4^{2-}$ and hydrogen sulphide ($\text{H}_2\text{S}$), resulting from the changing redox conditions caused by the vertical movement of the GWL and the presence of SRB microbial communities. The GWL fluctuations depend on variations in evapotranspiration, precipitation and surface water runoff. One aspect of this conceptual model is a transient reduction of the recently deposited $\text{SO}_4^{2-}$ by the SRB's with a concomitant transient Hg methylation [30,35,55]. The current study indicates that a chronic deposition of anthropogenic $\text{SO}_4^{2-}$ also results in

Figure 1. Weakly variations in A: concentrations of MeHg in pore water collected 10 cm below the groundwater surface at the time of sampling from HighS plots (filled symbols) or LowS plots (open symbols), as well as in the main stream draining the mire where the study was conducted (shaded squares). B: mean groundwater level from the HighS plots (filled symbols) or LowS plots (open symbols). The bars indicate SE. The arrows in A refer to the date on which the plots were fertilised with $\text{SO}_4^{2-}$.

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a more long-term reduction/oxidation cycle of the previously deposited S which sustains elevated MeHg levels for weeks after the last precipitation input of SO$_4^{2-}$ to the mire. That long-term redox cycle is driven by the microbial decomposition of organic matter and changing redox-conditions. The possibility of strong internal cycling of S due to variations in redox conditions has been shown earlier for peatlands [38,56,57].

This conceptual “SRB” model is supported by the temporal variation in pore water MeHg found in the HighS plots that correlates significantly with the GWL over the season (Fig. 1).

Figure 2. Depth profiles of MeHg pore water concentrations sampled 0–10 cm, 10–20 cm, 20–30 cm and 30–40 cm below the groundwater level at the LowS (upper panel) and HighS (lower panel) plots. The vertical axis refers to the actual peat depths at which the water was sampled. Each bar represents the mean value (±SE) of MeHg from two plots for the particular treatment. For each treatment (LowS or HighS), significant differences (p < 0.05) in the average pore water MeHg concentration between the sampling dates are designated by different upper case letters. Different lower case letters indicate significant differences (p < 0.05) in MeHg concentrations between depths for each treatment and sampling date.

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During one distinct period the pore water concentrations of MeHg in the HighS plots were close to the concentrations found in the plots receiving ambient levels of SO$_4^{2-}$. This period coincided with the lowest GWL and the highest pore water concentrations of SO$_4^{2-}$ measured at the HighS plots (Fig. 3). We interpret this temporal variation in MeHg as a response to the vertical distribution of the anaerobic heterotrophic bacteria delivering the carbon precursors used by sulphate reducing bacteria. The vertical distribution for other heterotrophic microorganisms in mires, e.g. methanogenic archaea and methanotrophic bacteria is related to the average growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61]. The methanogenic archaea, utilizing the same carbon and energy source as the SRB, has its maxima just below the average depth for the growing season mire water table level and does not change directly in response to the actual GWL [58–61].

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A decrease in MeHg concentrations with depth (Fig. 2) also suggests that the methylation activity of the SRB’s decreases with depth, and that the effect of enhanced SO$_4^{2-}$ deposition on Hg methylation is highest within the area of regular groundwater fluctuations. This zone of regularly fluctuating water tables is also the zone that is likely to contribute most to runoff from this type of wetland [63,64]. Altogether this indicates that SO$_4^{2-}$ reduction is restricted at depth. This would be expected and results mainly from the vertical distribution of high quality carbon substrate, acting as an electron donor to the SRBs, which is highest close to the mire surface. Thus, the abundance of SRB’s would be relatively lower at these greater depths. (Note that the absolute depth of pore water sampling relative to the mire surface varies with the water table).

While most of the data were consistent with the “SRB” hypothesis, there was one piece of evidence in this study that does not fit with this theory. That is the lack of correlation between solid phase S in the peat, and pore water MeHg (Table 1). The
experimental additions of SO$_4^{2-}$ every 4 or 5 weeks during the snow-free period are expected to be reduced within hours or days after addition [30], and indeed SO$_4^{2-}$ was below detection limit for most of the study period, with the one exception during the period of low GWL from mid-August to mid-September. Therefore, according to the hypothesis, it is increased concentrations of S in the organic matter that should feed the persistent effect from the SO$_4^{2-}$ addition found in the HighS plots. Such a correlation between total S in the peat and elevated pore water MeHg levels (or even S additions) was not found in our study. However, in the study from 1997 [36] done on the same site, a positive correlation between pore water MeHg and S in the solid phase was found, supporting the hypothesis described above. We believe that more needs to be known about the cycling of S in the mire to devise measurements better suited to testing this aspect of the SRB hypothesis.

The factors which pore water MeHg was not related to are also important arguments for the SRB hypothesis. To begin with, there was no correlation between Hg$_{tot}$ and MeHg. Such a lack of correlation between pore water MeHg and Hg$_{tot}$ was also found in a variety of aqueous samples [65,66].

High DOC and low pH [67–70] have been shown to correlate to increased MeHg concentrations in lake water and fish. Neither of these correlations are a direct outcome of the SRB hypothesis, and no such correlations between DOC and MeHg concentrations were found in this study of peat pore water that is a precursor to runoff. In late August, water was sampled from the lowest peat depth during the season, yielding DOC concentrations twice as high as on the other summer and autumn sampling occasions. This increase in DOC concentration was accompanied by only a very limited increase in MeHg in the HighS plots. The pH in the HighS plots was approximately 0.2–0.4 pH units higher than in the LowS plots at the beginning of the snow free season when the HighS plots had the greatest elevation of pore water MeHg relative to the LowS plots. The microbial reduction of SO$_4^{2-}$ consumes H$^+$ [71] and theoretically, the complete reduction of the added SO$_4^{2-}$ would be sufficient to generate the higher pH in the

### Table 1. Content of total S (S$_{tot}$ %), Hg$_{tot}$ (ng g$^{-1}$), and MeHg (ng g$^{-1}$) in the organic matter at different depths from the two plots receiving 3 kg S ha$^{-1}$ yr$^{-1}$ (LowS) or 20 kg S ha$^{-1}$ yr$^{-1}$ (HighS).

| Analysis | LowS | LowS | HighS | HighS |
|----------|------|------|-------|-------|
| S$_{tot}$ (%) | | | | |
| 0–5 cm | 0.05 | 0.06 | 0.06 | 0.07 |
| 5–10 cm | 0.05 | 0.06 | 0.05 | 0.07 |
| 10–20 cm | 0.07 | 0.09 | 0.10 | 0.12 |
| 20–40 cm | 0.10 | 0.16 | 0.10 | 0.12 |
| Hg$_{tot}$ (ng g$^{-1}$, S.E., n = 2 or 3) | | | | |
| 0–5 cm | 22(6.5) | 27(14) | 50(8.7) | 64(4.0) |
| 5–10 cm | 68(15) | 45(1.3) | 59(3.2) | 63(2.3) |
| 10–20 cm | 65(17) | 68(2.2) | 66(3.2) | 57(0.9) |
| 20–40 cm | 79(5.0) | 67(0.6) | 89(8.6) | 80(0.7) |
| MeHg (ng g$^{-1}$) | | | | |
| 0–5 cm | 0.17 | 0.30 | 0.20 | 0.76 |
| 5–10 cm | 0.21 | 0.22 | 0.18 | 3.70 |
| 10–20 cm | 0.26 | 0.34 | 0.42 | 1.90 |
| 20–40 cm | 0.20 | 0.93 | 0.18 | 0.26 |

The solid samples were collected on June 24th 1999.

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HighS plots. Thus, neither acidification, DOC nor $H_{SO_4}$ could account for the enhanced concentrations and temporal variation of MeHg after $SO_4^{2-}$ additions.

Another alternative explanation for the enhanced levels of pore water MeHg found in the HighS plots might be that the microbially produced MeHg sulphides which increase the partitioning of MeHg to the water phase [72,73]. However, calculations based on the amount of added $SO_4^{2-}$ revealed that a negligible concentration of MeHg bound to peat organic matter would be exchanged due to the low pH in the peat.

Conclusion

This study demonstrates that chronic deposition of $SO_4^{2-}$ to a boreal mire can result in large and persistent increases in the MeHg of peat pore water located near the surface of the water table. The increase in pore water MeHg caused by the addition of 17 kg S ha $yr^{-1}$ beyond the ambient deposition of 3 kg ha $yr^{-1}$ varied strongly over the snow free season in conjunction with seasonal fluctuations of the water table. The conceptual model we present to explain the HighS effect on MeHg postulates that enhanced pore water MeHg concentrations are maintained by the cycling of S as a result of fluctuating redox conditions that stimulate, and presumably increase the SRB community in the immediate vicinity of the redox fluctuation associated with the fluctuating water table.

Since the pore water pore just below the water table can be an important source of MeHg in runoff from a wetland, we believe these results support the hypothesis that $SO_4^{2-}$ deposition has increased the contribution of peatlands to MeHg in downstream aquatic systems. This would mean that the increased deposition of $SO_4^{2-}$ in acid rain has contributed to the modern increase in the MeHg burdens of remote lakes hydrologically connected to peatlands, as has also been suggested by other studies [9,36]. This would help explain why the catchment yields of MeHg from small boreal catchments are not well correlated to the atmospheric input of Hg [19,74]. According to this “sulphur rain” hypothesis, the large catchment MeHg yields are enhanced by the stimulation of Hg methylation by $SO_4^{2-}$ in acid rain, rather than directly from the Hg deposition itself or as a consequence of surface water acidification.

The cycling of MeHg in catchments is complex, and involves numerous biogeochemical and physical controls, many of which remain to be satisfactorily quantified. The findings presented here are not likely to be applicable to all catchments, but if they are applicable to many high-latitude catchments containing peatlands or other areas of anoxic organic sediments, then they represent an important link in our understanding of catchment-scale Hg cycling with implications for Hg uptake by fish that extend across the boreal and sub-boreal zones of North America, Europe and Asia.

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Author Contributions

Conceived and designed the experiments: IB MN. Performed the experiments: IB MN. Analyzed the data: IB KB WF MN. Contributed reagents/materials/analysis tools: QT WF. Wrote the paper: IB KB SA... MN.

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