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1 Forest harvest effects on mercury in streams and biota in Norwegian boreal catchments

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20 Abstract

21 Forest harvesting practices can potentially increase mercury run-off from catchments. A paired 22 catchment experiment was conducted in a boreal forest in southern Norway, to test effects of forest 23 harvest operations on i) concentrations and fluxes of methylmercury (MeHg), total mercury (HgT), 24 nutrients and dissolved organic matter (TOC), and on ii) MeHg bioaccumulation in stream foodwebs. 25 Thirty percent of a catchment was harvested in winter time with snow cover but no soil frost, 26 resulting in wheel tracks and soil compaction. Pre-harvest differences included higher streamwater 27 MeHg, HgT and TOC, and lower pH in the treated catchment compared to the reference. 28 No significant treatment effects on concentrations of MeHg, HgT and TOC were detected, 29 whereas concentrations of nutrients (ammonium, nitrate, phosphorus (P)) increased significantly. 30 Estimated catchment export of nitrate and ammonium increased fourfold, as a combined effect of 31 changed discharge and concentrations. Export of MeHg and HgT increased weakly, primarily because

32 of an increase in discharge.

Levels of MeHg in stream invertebrates mirrored differences in aquatic MeHg between the
two streams, resulting in higher MeHg in biota in the harvest catchment in pre-harvest conditions.
After harvest, MeHg levels in primary consumers (herbivorous stoneflies) were no longer different
between the two streams, despite continued exposure to higher aqueous MeHg in the harvested
catchment. Simultaneously, dietary biomarkers (δ¹⁵N signature, lipid- and algal fatty acid content) in
the stoneflies had changed significantly, consistent with higher nutrient loadings and associated higher
diet availability in the harvested stream.

40 The results of our experiment do not support that forest management has a strong impact on 41 catchment MeHg production. Catchment disturbance through forest harvesting may decrease MeHg in 42 aquatic biota, because of higher stream productivity and associated higher quality of dietary sources, at 43 least in the short-term. Other studies on catchment MeHg production to disturbance have shown a 44 range in responses, from strong to none. So far, no factor has emerged to explain such range in 45 responses. Predictions of forest management effects on MeHg in streamwater and aquatic food webs 46 are hampered by limited understanding of catchment controls on MeHg production.

- 47
- 48 Keywords
- 49 Forest management, catchment manipulation, methylmercury, water chemistry, bioaccumulation,
- **50** primary consumers, clear-cutting

51 1. Introduction

52 Mercury (Hg) is a long-range transported pollutant of great environmental concern in boreal areas 53 across the entire northern hemisphere. Atmospheric deposition of Hg in natural ecosystems leads to 54 long-term accumulation of Hg in soils and wetlands, where transformations of Hg to its highly toxic organic form methylmercury (MeHg) occur with subsequent transport of Hg-species to surface waters 55 56 (Grigal, 2002). MeHg is a neurotoxin with a strong tendency to bioaccumulate in food webs (Morel et 57 al., 1998). Levels of MeHg in the aquatic food web are raised to levels that are potentially harmful for 58 fish and wildlife (Scheulhammer et al., 2007) and, through consumption of fish, to human health (Mergler et al., 2007). 59

60 High Hg concentrations in fish are associated with brown-water streams and lakes in forested regions with a prevalence of wetlands (Nilsson and Håkanson, 1992; Driscoll et al., 2007; Chasar et 61 62 al., 2009). Wetlands are commonly viewed as one of the main suppliers of MeHg to aquatic 63 ecosystems, because of high groundwater levels, creating anoxia and thereby promoting conditions for 64 methylation of Hg (Grigal, 2003). Not just wetlands, but also forests have the potential to be a 65 significant source of MeHg to surface waters. Coniferous forests are highly effective scavengers for 66 atmospheric Hg species, resulting in substantially higher Hg deposition in forests compared to open areas (Graydon et al., 2008; Witt et al., 2009), thereby enriching forest soils with Hg. Forest 67 68 throughfall has been shown to be a significant input of MeHg to boreal catchments (Witt et al., 2009), 69 possibly demonstrating an additional pathway of MeHg from forest canopies to surface waters. 70 Recently, forest management has been suggested to be an important contributor to catchment

export of MeHg, thereby increasing MeHg in the aquatic food chain (Bishop *et al.*, 2009). Forests in
northern Europe (Ostlund *et al.*, 1997) and large parts of North America (Stinson *et al.*, 2011) are
landscapes where forest management practices have left a strong mark. Because of increased interest
in the role of forest for climate mitigation (Jackson *et al.*, 2008), especially as a source of bioenergy
(Schlamadinger and Marland, 1996), forest management might intensify. Thus, further assessment of
environmental impacts of forest harvest practices is needed to protect aquatic ecosystems.

Forest harvesting is known to have a strong impact on catchment hydrology and nutrient 77 78 runoff (Likens et al., 1970; Kreutzweiser et al., 2008). Effects of forest harvesting and soil disturbance 79 on MeHg runoff have been shown in Finland (Porvari et al., 2003) and Sweden (Munthe and Hultberg, 80 2004). However, the mechanisms controlling the increased export of MeHg are not well understood. Soil disturbance with associated increases in mobilization of MeHg from soil pools has been 81 82 hypothesized previously as controlling mechanism (Munthe and Hultberg, 2004), in addition to 83 increased discharge, changed hydrological pathways and higher soil temperatures (Porvari et al., 2003; 84 Eklof et al., 2013). Still, forest operations have not lead to increases in MeHg runoff in all cases. No 85 effect of harvest operation on MeHg runoff was found in catchment manipulations in Sweden, despite 86 small increases in runoff (Sorensen et al., 2009a) and extensive damage to soils from forest 87 machinery (Sorensen et al., 2009a; Eklof et al., 2013) 88 Another type of evidence for relations between forest management and MeHg in aquatic 89 ecosystems comes from synoptic studies. Studies of lake ecosystems in Canada indicated a connection 90 between catchment disturbance and increased levels of MeHg in the aquatic food web (Garcia and 91 Carignan, 1999, 2000; Desrosiers et al., 2006; Garcia et al., 2007). Here, increased levels of MeHg in 92 aquatic biota and periphyton were related to catchment disturbance, while aqueous dissolved MeHg 93 was not investigated. Further, significant relations between MeHg in aquatic biota and dissolved 94 MeHg in waters were found by Hall et al. (2009) in Canadian flooded reservoirs, and by Chasar et al. 95 (2009) in a synoptic study of stream foodwebs in the United States. Thus, relations between catchment 96 disturbance and enhanced levels of MeHg in biota have been implied, but are not well-documented. In 97 addition, the limited number of studies and lack of consistent responses of forest management on **98** MeHg export indicate a strong need for a better understanding of processes underlying catchment 99 MeHg production from experimental settings. 100 In order to test the hypothesis that forest harvest i) increases streamwater MeHg and total Hg 101 concentrations and runoff, and ii) enhances MeHg concentrations in biota, we conducted a paired-

102 catchment study in a Norwegian boreal forest. Streamwater chemistry, hydrology and levels of MeHg

103 in stream invertebrates were investigated, including links between diet and bioaccumulation of MeHg.

104

105 2. Materials and Methods

106 2.1 Site description

The Langtjern study area is located in southeast Norway (60°37' N, 9°73' E) at 500 to 710 m elevation 107 108 approximately 80 km northwest of Oslo (Figure 1). The Langtjern lake catchment is part of the 109 national monitoring programme for effects of acid deposition and consists of two inlet streams and a 110 lake outlet, where streamwater chemistry and discharge have been monitored since 1972. The eastern 111 inlet stream catchment (LAE03) was used as the reference catchment. The treatment catchment 112 (LAE11) is located 1.5 km southeast of LAE03, adjacent to the lake catchment, and is slightly less 113 than one-third of the size of the LAE03 catchment (Table 1). 114 Mean annual discharge from the Langtjern lake outlet between 2008 and 2011 was 702 mm, 115 while mean annual precipitation and temperature were 914 mm and 4.5 °C, respectively (nearby 116 meteorological station Gulsvik II, 132 m elevation, 60°38' N, 9°60' E). Wet sulphur (S) deposition was 5 kg S ha⁻¹ in 1990 (Larssen, 2005) and 3 kg S ha⁻¹ in 2000 (De Wit *et al.*, 2007) respectively and 117 118 is still declining. 119 The vegetation at Langtjern is dominated by low- to unproductive Scots pine forest (Pinus 120 sylvestris L.), interspersed with peatlands (both forested and open Sphagnum mires) and patches of 121 Norway spruce (Picea abies (L.) Karst.) forest. The stands are mature or close to maturity. The 122 geology consists of till of felsic gneisses and granites, where thin mineral soils have developed.

123 Deeper peaty soils are found, being most abundant close to streams. The area proportion of main

124 forest- and vegetation types is similar in the two catchments, the most notable difference being a

higher percentage of forested peatland (forest on peat soils of at least 30 cm depth) in the LAE11

126 catchment. In LAE11, pre-treatment volume proportions of Scots pine, Norway spruce and birch were

127 57%, 34% and 9%, respectively, while the corresponding numbers were 62%, 35% and 3 % in

128 LAE03. LAE03 and LAE11 had a stocking of 78 and 62 m³ ha⁻¹, respectively. These volumes

129 illustrate the low site productivity of the study area.

131 2.2 Experimental design and harvest operation

The paired catchment experiment consisted of two small forested catchments, the reference (LAE03) and the treatment catchment (LAE11). Monitoring started in June 2008. The forest harvest operation in the LAE11 catchment was conducted from January 13 to 16 in 2009. Forest standing volume, water chemistry, discharge and aquatic biota were monitored before and after the harvesting operation. The choice for the timing of the harvest operation and thereby the length of the pre-harvest treatment was partly based on the original period of project funding, i.e. three years.

138 The harvesting operation was done using the 'cut to length' method (harvester and forwarder). 139 The impacted area was confined to the lower and middle part of the LAE11 catchment, affecting about 140 30 % of the catchment area and with a harvest removal corresponding to 38 % of total catchment tree 141 volume. As the forwarder would have to cross several areas with limited bearing capacity on its route 142 between the harvested area and the landing site the harvesting operation was scheduled to take place in 143 winter, when the soil was expected to be frozen. However, due to mild weather prior to harvest, the 144 soil was not frozen. Snow depth was circa 20 cm during harvesting. Thus, harvesting resulted in local 145 soil disturbance in the form of wheel ruts. This was most pronounced along the main extraction tracks 146 and in wetter parts adjacent to the stream in the lower part of the catchment, while the upland parts of 147 the catchment area were less affected. Norwegian guidelines for harvesting close to streams and mires 148 required leaving a buffer zone adjacent to the mire in the central part of the catchment where the 149 buffer zone corresponds with the stream course. With this exception, only scattered trees were retained 150 on the impacted area. Equal volumes of Scots pine and Norway spruce were harvested, whereas 151 birches which only occurred as scattered individuals were mostly retained.

152 2.3 Hydrology

153 V-weirs were installed in the summer of 2008 in the streams of LAE03 and LAE11 for quantification
154 of discharge. Comparison of stream flow estimates at the Langtjern catchment lake outlet indicated
155 that these v-profiles did not supply data of sufficient quality for quantification of stream flow,
156 primarily due to leakage and overflow under high flow conditions. However, the v-profiles did provide
157 information on the variation in discharge in both catchments from 2008 until 2010, and indicated

synchrony in high flow events and low flow periods in both catchments and thus, similar to
hydrographs. Discharge from the LAE03 catchment was estimated instead based on a full water
balance for the entire lake catchment (Figure 1), based on daily discharge in the outlet, temperature
data from a nearby weather station and additional hydrological measurements made during the 1970s
(Wright and Henriksen, 1980).

Water levels in Lake Langtjern may vary with around 60 cm, resulting in variation in lake
water storage. Because of this, the discharge of the inlet and outlet do not follow the same pattern and
the inlet discharge cannot be derived by simple area-scaling of the outlet discharge. A simple water
balance model was used to adjust the impact from lake water storage:

167 where

168 R is total runoff to the lake, calculated from an empirical relation between the change in water 169 storage (ΔS) in the lake and discharge in the outlet (Q), and corrected for precipitation directly on the 170 lake (P) and evapotranspiration from the lake (E). Details on the calculation procedure are given in 171 Wright and Henriksen (1980). In short, ΔS was calculated from an empirical relationship between lake 172 water level (available from weekly measurements between 1976 and 1978) and outlet water level (m) 173 in the stilling pond before the v-profile in the outlet. E was estimated assuming and evaporation of 174 0.15 mm day⁻¹ per °C daily temperature (Lundquist, 1977). Precipitation and daily temperature were 175 derived from nearby meteorological stations. The total runoff (R, in mm) to the lake was assumed to 176 be representative for the subcatchment LAE03.

Discharge from the LAE11 catchment was estimated using area-corrected discharge from
LAE03 for the pre-harvest period. Post-harvest discharge in LAE11 was based on hydrological effects
of harvest in a paired-catchment experiment in in northern Sweden (Sorensen *et al.*, 2009b). The
catchments were covered by coniferous forest and included 3 to 18% wetland, with a slightly colder
and drier climate than our study site. In the Swedish study, two catchments were partially clear-cut,
where 30% and 71% of the catchment areas were logged. Hydrological responses after harvest
between the two catchments were similar. Thus, the Swedish catchments were similar in land cover

and forest harvest, compared to our study site which was also partially clear-cut. Discharge (compared
to a reference, non-harvested catchment) started to increase after July and resulted in a 35% increase
in mean annual runoff. The increase was +45% during base flow conditions (<1 mm day⁻¹); +27%
during intermediate flow (1-5 mm day⁻¹) and +25% for high flow conditions (>5 mm day⁻¹). Daily
discharge in LAE03 (in mm day⁻¹) was adjusted according to these %-wise changes in flow, from
August 2009 onwards.

190 2.4 Stream water sampling

191 Streamwater grab samples for acid-base chemistry were collected biweekly or monthly, according to 192 procedures established in the acid monitoring programme (SFT, 2009). Samples were sent to the 193 Norwegian Institute for Water Research (NIVA) by mail and processed at the NIVA accredited 194 laboratory. From 2008 until December 2011, streamwater grab samples for analyses of total Hg (HgT) 195 and MeHg were collected using 125 mL acid-leached Teflon bottles. The bottles were packed in two 196 plastic bags for ultra-clean handling (USEPA, 1996). Samples were sent to NIVA by mail and 197 forwarded to the Swedish Environmental Research Institute (IVL) in Gothenburg. There were usually 198 4 to 6 days between sampling and preservation with 0.5 ml 37-38% HCl (Baker). From October 2010, 199 streamwater samples were taken by another procedure, using 250 mL fluorinated polyethylene (FLPE) 200 bottles and sent for analysis at NIVA. HCl (concentrated trace level grade, 1 mL) was added to the 201 MeHg bottle to yield a 0.4 % solution during sampling. Water for HgT analysis was sampled in a 202 separate bottle, to which BrCl (bromo monochloride) was added as oxidising agent within 2 days after 203 arrival to the laboratory. All bottles for Hg species determination were packed in two plastic bags for 204 ultra-clean handling. The number of samples for Hg species and acid-base chemistry taken during the 205 pre- and post-harvest period is given in Table 2.

206 2.5 Sampling of biota

207 Streamwater biota was collected from one sampling station in each stream, at October 17 2008, May208 29 2009 and on October 16 2009, thus collecting samples that reflected summer and winter conditions.

209 Each sampling station comprised a stream reach of 30 m length, and was located in the lowermost

210 parts of each catchment. Biofilm was carefully removed from streambed rocks, concentrated by 211 centrifugation and kept in separate glass vials. Macroinvertebrate species composition of the two 212 streams was identical, and comprised two herbivorous stoneflies (nymphs of Nemoura cinerea and 213 Nemurella pictetii) and one predatory caddisfly (larvae of Plectrocnemia conspersa). They were 214 collected by kick-sampling, following the Norwegian Standard Method (EN 37828, 1994), using a 215 hand net (25 x 25 cm opening; 250 µm mesh size). Due to small body size, stoneflies were pooled to 216 samples of 50 to 80 individuals, to obtain enough material for chemical analysis. All samples were 217 shock frozen (-80°C) in the field, transported to the laboratory, freeze-dried and analyzed for MeHg, stable isotopes (δ^{15} N) and fatty acids. Species identification was conducted by the biological 218 219 accredited lab at the NIVA.

220 2.6 Analysis of water chemistry major ions

221 Analyses of pH, conductivity, major anions and cations, total N, total organic C (TOC), Al species,

total P and UV absorbance (at 254 nm) (UVabs₂₅₄) were performed at accredited laboratories at NIVA.

223 Total organic C was analysed by wet chemical oxidation IR-detection (EPA accredited method nr.

415.1). The samples were purged prior to analysis so only non-purgeable organic carbon was

225 measured. Organic carbon in a sample was converted to carbon dioxide by wet chemical oxidations.

226 The carbon dioxide formed was measured directly by an infrared detector. Total organic carbon (TOC)

227 consisted of ca 95% DOC (filtered by 0.45µm) in the LAE03 stream. Specific absorbance at 254 nm

228 (SUVA₂₅₄) was calculated by dividing UVabs₂₅₄ with TOC.

229 2.7 Mercury analysis

Two laboratories were involved in determination of HgT and MeHg; IVL (2008-2011) and NIVA
(2011-2012). Both laboratories follow United States Environmental Protection Agency (USEPA)
Method 1630 (USEPA, 1998) for determination of MeHg in water by distillation, aqueous ethylation,
purge and trap, and cold vapor atomic fluorescence spectrometry (CVAFS). For HgT, USEPA Method
1631 for determining Hg in water by oxidation, purge and trap and CVAFS was followed (USEPA,
2002). The method detection limits (MDL) were 0.02 ng/L (NIVA) and 0.06 ng/L (IVL) for MeHg,

and 0.1 ng/L for HgT (3 standard deviations of blanks). The IVL laboratory determination of Hg
species was done from one bottle (see sampling procedures). Analysis proceeded by the removal of a
sample aliquot for determining MeHg, before BrCl (bromo monochloride) was added as oxidising
agent and the remainder of the sample used for determination of HgT. The NIVA laboratory followed
the same procedure, but samples for MeHg and HgT analysis were taken in two separate bottles.

241 For over a year (October 2010 to November 2011), parallel samples for several locations, 242 including the streams in this study, were run at both laboratories. The IVL laboratory reported 243 significantly higher (p<0.05) HgT concentrations than the NIVA laboratory (Braaten et al., 2014). 244 Braaten et al. (2014) show that difference in HgT is related to the removal of the aliquot for MeHg 245 from the bottle used for both MeHg and HgT analysis, and re-dissolution of HgT species that adhered 246 to the bottle surface. For the LAE03 and LAE11 streams, HgT from IVL was on average 12% higher 247 than from NIVA in the parallel sampling period. Where analytical results were available from both 248 laboratories, we used the value from the NIVA laboratory as default. We tested whether the outcome 249 of the statistical tests (statistical methods described further below) of treatment effects on MeHg and 250 HgT streamwater chemistry was affected by i) using IVL results instead of NIVA results for the 251 parallel sampling of treatment effects, and ii) the change in laboratory, by downwards adjusting IVL 252 results of HgT with 12% from 2008 to September 2010 (see Supplementary Information). The results 253 of the statistical tests were not affected.

Biological samples were treated with hot methanolic potassium hydroxide solution for about
3-4 hours. The samples were then diluted with methanol, separated by ethylation and detected
following the same procedure as described for water samples (see above). The typical detection limit
was 1.5 ng g⁻¹ in MeHg analysis. Reference materials NIST 2977 (Mussel Tissue) and DORM-2

258 (Dogfish muscle) were used, and recovery was 104% and 100%, respectively.

259 2.8 Stable isotope, lipids, and fatty acids analyses

260 Stable nitrogen isotopes (δ^{15} N) of biota were analyzed after transferring freeze-dried samples (1 mg) to

tin capsules and combusted in a Eurovector element analyzer. The N₂ was directly injected online to a

262 Nu Instruments Horizon, Isotope Ratio Mass Spectrometer (Wrexham, UK) for determination of δ^{15} N.

Lipids were extracted from freeze-dried (96 hours) samples using chloroform:methanol (2:1) as 263 described elsewhere (Heissenberger et al., 2010). Fatty acids were esterified from total lipid extracts to 264 265 obtain fatty acid methyl esters (FAME) using toluene (1 mL) and H₂SO₄-methanol (2 mL; 1% v/v). 266 Subsequently, FAME were analysed using a gas chromatograph (TRACE GC THERMO) equipped 267 with flame-ionization detection, a temperature-programmable injector and an autosampler. A Supelco[™] SP-2560 column (100 m, 25 mm i.d., 0.2 µm film thickness) was used for FAME 268 269 separation. Excalibur 1.4TM was used for calculation and, if necessary, manual resetting of the 270 chromatograms. Fatty acid concentrations were calculated using calibration curves based on known 271 standard concentrations. Fatty acids were grouped to characterize bacterial fatty acids (BAFA; i.e., the 272 sum of odd saturated and branched-chain FA: 15:0 and 17:0 and their iso and anteiso series), algal 273 fatty acids (PUFA; i.e., the sum of polyunsaturated fatty acids) as previously presented (Sun et al., 274 2000; Kainz et al., 2002).

275 2.9 Calculation of catchment element and nutrient export

276 Catchment export of elements and nutrients was calculated by linear interpolation of streamwater
277 concentrations of elements to daily concentrations and multiplying with daily discharge, and summed
278 to monthly fluxes.

279 2.10 Statistical analysis

280 Random Intervention Analysis (RIA) was used to analyse treatment effects on water chemistry 281 (Carpenter et al., 1989). For a time series of any given variable, paired differences between the 282 reference and the treated catchment were calculated. The resulting time series of *catchment differences* 283 were used to test the effect of the intervention, by comparing differences in catchment differences before and after the intervention. This was done by random resampling (n=2000) values of *catchment* 284 285 *differences* from the pre-harvest period and the post-harvest period (for one year at a time, and for the 286 entire post-harvest period) and generating new time series. The mean values of 2000 resampled time 287 series were calculated for the pre-harvest and selected post-harvest period, and compared to the 288 statistical distribution of the original time series to determine whether significant treatment effect had

289 occurred. If the value of the mean catchment difference of the original time series before and after the 290 intervention was outside a certain percentile range of the *catchment difference* of the generated time 291 series before and after intervention, we assumed that a non-random effect had occurred as a result of 292 the treatment. The percentiles were 5% and 95% for p=0.10, 2.5% and 97.5% for p=0.05, etcetera. We 293 did not constrain the resampling period to take into account seasonal variation as done in a previous 294 paired-catchment study (Lofgren et al., 2009) by allowing resampling only for a limited number of 295 months for any number of years after the given observation. A test on the effect of sampling interval 296 showed that length of sampling interval did not affect the outcome of the analysis. 297 Pair-wise differences between sample locations (streams LAE03 and LAE11) were tested for 298 concentrations of MeHg, lipid content, stable isotopes and fatty acids in invertebrates and biofilms 299 using Student's t-test.

3 Results

3.1 Streamwater chemistry

303	During the pre-harvest period, both streams had a water chemistry signature typical for small inland
304	acidified catchments with base-poor soils interspersed with peatlands, i.e. a pH below 5, low
305	conductivity (1-3 μ S cm ⁻¹), high TOC (14-22 mg C L ⁻¹), low base cation concentrations (Ca < 1 mg L ⁻¹)
306	¹), low SO ₄ concentrations (<1 mg L^{-1} SO ₄ -S) and low nutrient concentrations (inorganic N-species
307	and total $P < 10 \ \mu g \ L^{-1}$) (Table 2, Figure 2). At almost each sampling occasion in the pre-harvest
308	period, concentrations of MeHg, HgT, TOC, totN and TON and conductivity were higher in the
309	LAE11 (experimental) stream than in the reference stream while pH was lower. By contrast,
310	concentrations of base cations, SO ₄ , species of inorganic N and totP were similar in both streams. The
311	pre-harvest differences in streamwater chemistry were probably related to the higher contribution of
312	peatland in the experimental catchment which lead to more TOC and associated elements. High TOC
313	is commonly associated with a decrease in pH as dissolved organic matter is a weak acid.
314	Concentrations of MeHg varied most in the experimental catchment (LAE11), with winter
315	peak concentrations exceeding 0.5 ng L^{-1} . Interestingly, the highest MeHg concentrations in the
316	LAE11 stream appeared each winter, although by 2011-2012 the winter peak was modest. The peak in
317	MeHg concentration in 2009 started prior to the harvest operation. In the reference stream (LAE03),
318	MeHg varied little from around the detection limit up to $0.2 \text{ ng}/\text{L}$ with a tendency towards higher
319	concentrations in summer. In both catchments, MeHg was lowest during snowmelt. Less than 10% of
320	the variation in MeHg was explained by TOC in the LAE03 stream, while no significant relation
321	between MeHg and TOC was found in LAE11 (LAE03: $r^2 = 0.08$, p<0.01; LAE11: $r^2 = 0.0$, p>0.5).
322	This is also illustrated by the large variation in MeHg to TOC ratio (Figure 2).
323	Seasonal patterns in TOC concentrations were very similar in both catchments, with highest
324	TOC in summer and lowest during snowmelt (Figure 2). Concentrations of HgT were closely
325	correlated with TOC in both catchments (LAE03: $r^2 = 0.51$, p<0.0001; LAE11: $r^2=0.29$, p<0.0001),
326	and HgT was usually highest in September and lowest in early winter. The HgT to TOC ratio was

327 within the same range (roughly 0.15 to 0.35 μ g g⁻¹) in both streams and showed similar temporal 328 variation.

329 Contrary to our hypothesis, no significant effects of the harvest treatment were found for any
330 comparison of pre-harvest period and post-harvest years for streamwater concentrations of MeHg and
331 HgT (p>0.1; Figure 2, Table 2). The ratio of mean MeHg in the LAE11 and LAE03 streams was 2,
332 respectively 2.5 in the pre-harvest and post-harvest periods, respectively, possibly indicating a small
333 but non-significant response to treatment.

334 Nitrate concentrations showed a strong and significant (p<0.01) response to harvest, with peak NO₃-N concentrations between 60 and 120 μ g L⁻¹ in LAE11, while peak concentrations in LAE03 335 were between 20 and 50 μ g L⁻¹ (Figure 2; Table 2). Peak concentrations of NH₄-N in the post-harvest 336 period in LAE11 exceeded 100 μ g L⁻¹, while NH₄-N in LAE03 was below 40 μ g L⁻¹. Total P 337 concentrations in LAE11 were on average almost twice as high as in LAE03 (9 and 5 μ g L⁻¹, 338 339 respectively) in the post-harvest period, while in the pre-harvest period total P concentrations in both streams differed less (7.0 and 5.5 µg L⁻¹, in LAE11 and LAE03 respectively). There was a significant 340 341 (p<0.005) effect of treatment on total P in 2010 only. Organic N (TON), but not total N, showed a weak increase (p<0.1) after harvest in 2009 and 2011, going from 284 to >320 μ g L⁻¹ in LAE11, while 342 TON was $<250 \ \mu g \ L^{-1}$ and remained stable in the reference stream. A consistent and significant 343 344 (p<0.05) treatment effect was found for the CN ratio of DOM in each year of the post-harvest period, 345 where CN ratios in LAE11 decreased from 72 to 57, almost equal to the CN ratio in LAE03 of 54. The 346 decrease in CN ratio suggested enrichment of N in DOM after harvest. No treatment effects were found for UVabs₂₅₄ or for specific UV-absorbance (SUVA₂₅₄). Cation concentrations that increased 347 significantly after harvest were K (p<0.05) and Ca (p<0.1) while pH showed a weak but significant 348 349 (p<0.1) decline in 2009 and 2010.

350 3.2 Hydrology

351 The effect of forest harvest on run-off was estimated using results from a paired-catchment study of
352 forest harvesting in a site with comparable land cover and climate as Langtjern. Only qualitative
353 observations of higher ground water levels and wetter soils after harvest, in the harvested catchment

compared to the reference, were available. Mean annual discharge in the reference catchment LAE03
from 2009 to 2012 was 738 mm. In the post-harvest period from August 2009 until December 2012,
discharge in experimental catchment was calculated as being on average 28% higher than in the
reference. No statistical tests of treatment effect were done on discharge because these results were
obtained by inference and not by in-situ measurements.

359 3.3 Streamwater fluxes

360 Catchment export was calculated for MeHg, HgT, TOC, totP and inorganic N species. In the pre-

361 harvest period, export per unit area of all elements except NO₃ was highest in the experimental

362 catchment (Table 3). Because discharge (in mm) in LAE11 was assumed equal to LAE03 (Table 2),

363 these differences were related to concentrations only. The most noticeable pre-harvest difference was

364 found for MeHg export, which was 75% higher in the experimental catchment than in the reference. In

365 the post-harvest period, export of elements increased more in the harvested catchment than in the

366 reference. The increase in the difference between LAE11 and LAE03 ranged from 24-51% (HgT,

367 TOC, totP), to 74-104% (totN, MeHg), to over 300% (NH₄, NO₃) (Table 3), and related to both

368 increased discharge and increased concentrations. No statistical test was done of treatment effect as

369 the treatment effect on discharge was estimated, not measured.

370 3.4 Stream biota

371 Biofilms covered rocky substrate in both streams and were composed of gelatinous polymers 372 associated with the chlorophytes Tetraspora sp., Microspora sp. and various diatoms (e.g., Eunotia 373 sp.). Detritus and fungi were present in low amounts. The main source of detritus in the streams was 374 Sphagnum, while leaf litter was nearly absent, due to the low presence of deciduous trees. The 375 macroinvertebrate fauna was species-poor and consisted of the same taxa in both streams, also after harvest. Stream water biofilm, stonefly nymphs (two closely related herbivorous Plecopterid species 376 377 *Nemoura cinerea* and *Nemurella pictetii*, which together constituted the principle primary consumers) 378 and caddisfly larvae (the carnivorous Trichopterid Plectrocnemia conspersa, the main predator of the stoneflies) were collected in autumn 2008, spring 2009 and autumn 2009. 379

380 There was no effect on species composition of the harvest operation. However, visual 381 observations of the streambed indicated a strong increase in primary production in the harvested 382 stream. In addition to a higher abundance of algae in the gelatinous biofilms, mats of green thread 383 algae had filled substantial parts of the streambed. This was not observed in the reference stream and 384 was interpreted as an effect of increased nutrient leaching from the catchment after the harvest 385 operation.

Biofilms were low in MeHg (3-7 ng MeHg g⁻¹ dw) (Table 4), and did not reflect differences in 386 aqueous MeHg between the streams (Table 2, Figure 2). In the reference stream, stonefly nymphs 387 contained 35 to 50 ng MeHg g⁻¹ dw, which varied little among sampling events (Table 4, Figure 3). In 388 389 LAE11, stoneflies had significantly (p<0.0001) higher MeHg than in the reference in autumn 2008 and 390 spring 2009, but no differences were observed in the autumn of 2009. The differences in MeHg 391 concentrations of the stoneflies in the first two sampling events were consistent with observed stream 392 differences in aqueous MeHg. However, in the autumn of 2009 stream differences in aqueous MeHg 393 were still present, while stream differences in MeHg in stoneflies had disappeared (Table 4, Figure 3). 394 Similar patterns in stream-wise differences in MeHg levels were observed for the caddisflies, but at a lower significance level (Table 4). The δ^{15} N signatures of stoneflies in the experimental stream 395 396 became significantly higher than in the reference in autumn 2009 (Figure 3, p<0.001), where no such 397 differences were found at earlier sampling events. The other significant changes in chemical content of 398 biota that occurred in the autumn of 2009 were significantly higher algal fatty acids (PUFA) (Figure 3, 399 p<0.05) and total lipids (LAE03, lipid content 0.23±0.03; LAE11 lipid content 0.28±0.01; p<0.05) in 400 stoneflies in LAE11, compared to the reference.

402 4 Discussion

403 4.1 Forest management effects on water chemistry

404 The main hypothesis guiding our paired-catchment experiment was an expected increase in MeHg 405 concentrations as a response to the forest harvest treatment. However, no significant effect of logging **406** was detected for streamwater MeHg concentrations, and our main hypothesis was not supported. **407** There was a substantial increase in MeHg export in our study -50% more MeHg in the 408 harvested catchment than in the reference after harvest - but this was primarily related to the estimated 409 increased runoff after harvest (+28%). The increase in runoff was estimated based on a paired-410 catchment study in Balsjö in Northern Sweden, with similar climate and catchment land cover, and a 411 similar % catchment harvest, i.e. between 30 and 40%, which documented a dominant increase of 412 discharge during low flow (Sorensen et al., 2009b). Increased water yield after harvest and other 413 catchment disturbances is a well-known phenomenon (Hewlett and Helvey, 1970; Guillemette et al., 414 2005; Buttle et al., 2009), which is also described in catchment models (Katsuyama et al., 2009). 415 Katsuyama et al. (2009) simulated a 25% increase in water yield in the first six years after moderate 416 logging operations in a forested catchment with seasonal snow cover. Porvari et al. (2003) reported a 417 doubling of runoff in a Finnish paired-catchment study where a 100% clear-cut was carried out. Our 418 estimated increase in runoff of a moderate logging disturbance appears to be in reasonable agreement 419 with other studies.

420 Interestingly, only two of five published paired-catchment experiments with a focus on 421 catchment disturbance and mercury cycling reported significant treatment effects on MeHg 422 concentrations in surface waters. In southern Finland, the catchment manipulation included a pre-423 harvest period of three years, after which the catchment was clear-cut in one year and soil treatment 424 was conducted the year after. Clear-cutting did not affect MeHg, but after the soil treatment 425 streamwater MeHg concentrations and export rose immediately, a significant effect that lasted for 426 three years (Porvari et al., 2003) and continued for at least seven more years (Porvari, pers.comm.). 427 The increases in streamwater MeHg were especially prominent during the growing season. In 428 Gårdsjön in southwest Sweden, an unintended soil disturbance (wheel tracks of forest machinery)

429 occurred seven years after the start of the monitoring and resulted in three years of increased 430 concentrations and export of MeHg (Munthe and Hultberg, 2004) which continued for at least another 431 five years (J. Munthe, pers. comm.). In both studies, the increase in MeHg export was at least partly 432 related to changes in MeHg concentrations, with peak MeHg concentrations after disturbance exceeding 1 ng L⁻¹. In the paired-catchment manipulation in Balsjö in northern Sweden, no effect of 433 434 logging on MeHg concentrations and MeHg export was found after a year of pre-harvest monitoring 435 (Sorensen et al., 2009a). In Örebro in central Sweden, stump harvesting and site preparation did not 436 affect MeHg concentrations (Eklof et al., 2013). However, only logged catchments were monitored, 437 and therefore this manipulation remains inconclusive with regard to effects of logging on aqueous 438 MeHg.

439 Our study had a relatively short pre-harvest period compared to the catchment manipulations **440** mentioned above, limiting the possibility to detect subtle responses to the treatment as intersite-441 variations may dominate the treatment effect (Buttle et al., 2005). Nevertheless, the experimental **44**2 design of our study allowed detection of a two- to fourfold increase in nitrate, ammonium and totP 443 concentrations, suggesting that if a similarly strong response in MeHg concentration had occurred, we 444 would have detected it. In the two experiments with long pre-disturbance periods, Munthe and 445 Hultberg (2004) and Porvari et al. (2003) found a fourfold and twofold increase in mean 446 concentrations of MeHg, respectively. Such strong responses in MeHg concentration were absent in 447 our study. We conclude that forest management did not strongly impact catchment MeHg production 448 in our study, similar to the results presented by Eklof et al. (2013) and Sorensen et al. (2009a). 449 Summarizing, the conclusion that forest harvest practices may be responsible for 9 to 23% of MeHg 450 loadings to surface waters (Bishop et al., 2009) receives little support from recent catchment 451 manipulations. 452 The lack of consistent responses in MeHg concentrations and export to forest harvest practices

452 The fack of consistent responses in MeHg concentrations and export to forest narvest practices
453 in paired-catchment studies is puzzling, partly because effects of forest harvest on streamwater MeHg
454 have been found in synoptic studies and thus appear to be well-founded (Skyllberg *et al.*, 2009; Eklof
455 *et al.*, 2012). Streamwater and lake MeHg are often found to correlate with the proportion of wetland

456 in catchments (StLouis et al., 1996; Shanley et al., 2005), where MeHg is thought to be produced by 457 sulphate-reducing bacteria using labile organic matter as energy substrate (Morel et al., 1998), 458 possibly also influenced by nutrient status (Tjerngren et al., 2012). However, forest harvest operations 459 are usually not undertaken in wetlands or organic-rich soils, and the increase in MeHg concentrations in southern Finland (Porvari et al., 2003) and Gårdsjön (Munthe and Hultberg, 2004) is related to **460** 461 disturbance of upland, not wetland, soils. Porvari et al. (2003) suggested that the enhanced **462** concentrations of MeHg were possibly related to higher soil temperatures (through increased direct 463 solar radiation) and humidity (from higher ground water levels), favouring methylation. In all referred 464 paired-catchment experiments where logging took place, it is reasonable to assume that increased soil 465 temperatures and humidity in the harvested catchments did occur as this is a common effect of forest clear-cutting (Olchev et al., 2009; Schelker et al., 2013). However, this was clearly not sufficient for **466** 467 increasing streamwater MeHg in three of four cases.

468 Another mechanism relevant to explain forestry effects on MeHg production is through 469 increasing loads of labile organic matter (Kainz et al., 2003; Roy et al., 2009), either in the form of 470 harvest residues or from release of fresh organic matter through soil disturbance, both of which 471 promote microbial activity and thereby Hg-methylation. Additionally, the creation of anoxic spots in 472 the soil related to soil compaction from heavy machinery might also promote methylation. Such 473 compaction is likely to have taken place in all catchment manipulations, but again, this was not 474 sufficient to create increases in streamwater MeHg in all experiments. Possibly, site differences in 475 sulphur (S) deposition could play a role for the susceptibility to logging and soil disturbance as 476 sulphate is a limiting factor for MeHg production (Gilmour et al., 1992; Akerblom et al., 2013). The 477 sites in southern Finland (Porvari et al., 2003) and southwest Sweden (Munthe and Hultberg, 2004) **478** are both located in regions that have historically received considerably higher loads of S deposition 479 (Jenkins et al., 2003; Posch et al., 2012) than our study site in southeast Norway, Örebro in central **480** Sweden (Eklof et al., 2013) and Balsjö in northern Sweden (Sorensen et al., 2009a). 481 The most distinct effect of the harvest operation was the large increase in NO₃ concentration

482 and export, which lasted throughout the entire post-harvest period. Increased runoff of inorganic N

483 species after harvest is common in northern catchments (Likens et al., 1970; Kreutzweiser et al., **48**4 2008), but can be reduced by retaining an intact buffer zone close to the streams (Lofgren et al., 2009). 485 Following common forestry practice in Norway, the stream in our study was too small to include such 486 buffer retention, except for a smaller part of the central catchment area where the stream course **487** followed the border between upland forest and adjacent open mires. Another sign of changes in N 488 cycling was the significant decrease in CN ratio of dissolved organic matter (DOM), suggesting an 489 enrichment of DOM with nitrogen. TOC concentrations and TOC export did not respond to the forest **490** harvest, in contrast to previous findings (Porvari et al., 2003; Laudon et al., 2009). Other responses 491 were increases in total P concentrations which were only significant in the second year after logging, 492 but effective P retention in the streambed was suggested by observations of thread algae and high 493 concentrations of algae in the biofilm. A less distinct response of P compared to N in streamwaters 494 after logging has also been found previously (Kreutzweiser et al., 2008; Lofgren et al., 2009) and 495 could be related to strong biological retention of P in the stream (Valett et al., 2002).

496 4.2 Forest management effects on MeHg in the stream food chain

497 The differences in MeHg concentrations in the streams were reflected in MeHg levels in primary 498 consumers (herbivorous stoneflies) in the autumn of 2008 and the spring of 2009 (de Wit et al., 2012). 499 Trophic enrichment of MeHg in the biota, and the efficiency of MeHg transfer from the stream into 500 the food chain, were similar in both streams and at both sampling occasions. Thus, the mechanisms 501 controlling MeHg levels in aquatic biota in both streams were exactly the same. The different levels 502 in MeHg in the primary consumers in the streams were explained by differences in exposure to 503 aqueous MeHg, where LAE11 had higher MeHg than LAE03. Exposure to MeHg at the base of the 504 food chain is key to the bioaccumulation in the stream food web, as studies by for instance Chasar et 505 al. (2009) also indicate. We also found that fatty acids content of the invertebrates indicated that the 506 ingestion of bacteria was likely to promote MeHg bioaccumulation, while ingestion of algae had the 507 opposite effect. Fatty acids can be used as dietary biomarkers to indicate recent dietary success of 508 biota (Kainz and Fisk, 2009).

509 The surprising observation in the autumn of 2009, compared to the first two sampling events, was 510 that MeHg levels in primary consumers of both streams were similar in the autumn of 2009, despite 511 continued differences in exposure to aqueous MeHg. That indicated that the efficiency of MeHg 512 transfer from the water phase into the base of the food chain had declined in LAE11 compared to 513 LAE03. Upon further inspection, this observation fitted well with the postulated importance of dietary 514 sources for MeHg bioaccumulation in de Wit *et al.* (2012). The significant change in δ^{15} N signature in 515 primary consumers of the harvested stream in the autumn of 2009 was interpreted as a change in 516 baseline N availability, substantiated by the observed increase in streamwater NO₃ and NH₄. 517 Additionally, visual inspection of the biofilm in the harvested stream indicated a much higher 518 abundance of algae than the year before, a strong indication that nutrient access in the stream had 519 increased. Further evidence for higher primary productivity in the harvested stream was found in the 520 significantly higher contents of lipids and algal fatty acids in the primary consumers of LAE11, which 521 indicate a higher dietary access to algae. Possibly, algae are a food source with relatively low 522 contamination of MeHg - consistent with the low concentrations of MeHg in biofilms – or the larger 523 dietary access to algae caused increased somatic growth of consumers leading to lower MeHg per unit 524 biomass, also known as the growth dilution effect (Goedkoop et al., 2007). Algal blooms have been 525 shown to lower MeHg contamination in aquatic food webs in lakes (Pickhardt and Fisher, 2007), and 526 we show here that a similar mechanisms may also exist in stream foodwebs.

527 Our results demonstrate that effects of forest harvest on MeHg in the aquatic food chain should 528 take the following aspects into account: i) changes in MeHg in runoff, ii) changes in in-stream 529 productivity and iii) changes of dietary sources. Previous studies on catchment disturbance effects on 530 MeHg in aquatic biota (Garcia and Carignan, 1999, 2000, 2005; Desrosiers et al., 2006; Garcia et al., 531 2007) have indicated positive correlations between disturbance and MeHg in aquatic biota, but have 532 paid little attention to possible confounding effects of changes in aquatic productivity and diet. 533 Likewise, the documented increase in Hg in fish in Swedish lakes has been suggested to be related to 534 increases in lake DOC with associated higher exposure to MeHg (Akerblom et al., 2012), but our 535 results suggest that changes in dietary sources can drive changes in MeHg levels in aquatic foodwebs.

536 5 Conclusion

537 Contrary to earlier results from paired catchment experiments and synoptic studies, we did not find an 538 effect of forest management on catchment MeHg production. There is little understanding of crucial 539 factors that render MeHg in surface waters sensitive to catchment disturbance, but we speculate that 540 sulphur deposition might be important. We found a strong nutrient release to the streamwaters as a 541 response to harvest. Such additional nutrients promote primary productivity and growth of primary 542 consumers, inducing a decrease of MeHg in biomass in streamwater biota. Our results demonstrate 543 that a short-term effect of forest harvest may be a reduction of MeHg in aquatic biota, because of 544 improved dietary quality for the consumers at the bottom of the stream food webs. An assessment on 545 the effects of forestry on Hg in northern, managed landscapes concluded that one tenth to a quarter of 546 Hg in fish might be attributed to forest harvesting (Bishop et al., 2009). Recent experimental studies, 547 including our own, suggest that this may be an oversimplification. Paired catchment manipulations 548 show that the impact of disturbance on MeHg in streamwaters varies strongly. Predictions of forest 549 management effects on MeHg in streamwater and aquatic food webs are hampered by limited 550 understanding of catchment controls on MeHg production.

551

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561 List of Tables and Figures

562

563 Table 1 Catchment characteristics

564

565 Table 2 Discharge (sum, in mm) and mean water chemistry of streams LAE03 (reference catchment) 566 and LAE11 (harvested catchment) for the pre-harvest period (pre: June 2008 - January 10 2009), post-567 harvest-1 (remainder of 2009) and post-harvest 2 (2010-2012). Numbers show mean / standard 568 deviation / number for observations. Cond, conductivity (μ S cm-1); LAL, labile Al (μ g L-1); other 569 major ions in mg L-1; totP, total P; totN, total N; TOC, total organic C; TON, total organic N; DOM, 570 dissolved organic matter; CN_{DOM}, CN ratio of DOM; UVabs₂₅₄, absorbance of UV at 254 nm; 571 SUVA₂₅₄, specific UV absorbance at 254 nm 572 573 Table 3 Discharge (sum, in mm) and element fluxes (Hg, MeHg, TOC, inorganic N species, totN and 574 totP) of catchment LAE03 (reference catchment) and LAE11 (harvested catchment) as sum for the 575 pre-harvest period (June - December 2008) and as the mean of the post-harvest period (2009-2012). In 576 parentheses for LAE11 are the pre- and post-harvest discharge and fluxes in % of the LAE03 value 577 578 Table 4 Mean concentrations of MeHg (µg g-1 dry weight), standard deviation (std) and nr of 579 observations (n) in biofilm, stoneflies and caddisflies, grouped by sampling event and catchment 580 (LAE03, reference; LAE11, harvested catchment). Significant differences between catchments 581 (Students t-test) are given as a and b when p<0.0001, as c and d when p<0.005, and as e and f when 582 p<0.1. 583 **584** Figure 1 Map of the Langtjern catchments. LAE03 is the reference catchment, LAE11 is the experimental catchment. The dotted line in the LAE11 catchment indicates the border between the 585

- 586 harvested areas in the northwestern, lower catchment area and the non-harvested areas in the upper
- 587 parts of the catchment.

589	Figure 2 Streamwater concentrations of various components, and some of their ratios, in catchment
590	LAE03 (reference) and LAE11 (treated) catchments from June 2008 until December 2012. Dotted
591	vertical line indicates time of harvest, and grey lines indicate 1 st of January. For all components except
592	ratios measured values are shown. For ratio, measured values were averaged to monthly means. The
593	following components are shown in panel A: MeHg, Hg, pH, TOC, NO ₃ , NH ₄ , totP, SO ₄ ; and in panel
594	B: K, Acid neutralizing capacity (ANC), totN, TON, CN ratio of DOM (CN _{DOM}), specific UV
595	absorbance at 254 nm (SUVA $_{254}$), and MeHg to TOC ratio and Hg to TOC ratios. The following
596	symbols denote significance level of treatment effect for any given year in the post-harvest period (i.e.
597	if post-harvest difference in streamwater chemistry is significantly different from pre-harvest
598	difference (RIA analysis); *: p<0.10, **: p<0.05, ***: p<0.01, ****: p<0.005.
599	
600	Figure 3 MeHg concentration in ng g ⁻¹ dry weight (upper panel), and PUFA concentration in $\mu g g^{-1}$
601	dry weight (lower panel) versus $\delta^{15}N$ signature in herbivorous stoneflies in ‰ (primary consumers) in
602	the reference (LAE03) and experimental (LAE11) streams. Three samplings (1= autumn 2008; 2
(0)	

spring 2009; 3 autumn 2009).

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Table 1 Catchment characteristics

		Pine-	Spruce-				Pre-	
		dominated	dominated	Forested	Sphagnum		harvest	Harvest
Catchment	Area	forest	forest	peatland	mire	Other ¹⁾	volume ²⁾	removal
	ha			-% of cate	hment area		$m^3 ha^{-1}$	% ³
LAE03	83	63.9	9.6	4.8	16.3	5.4	77.6	-
LAE11	24	56.8	10.5	14.7	17.9	-	62.3	38

¹⁾ Lakes and powerline; ²⁾ Overall mean for spruce- and pine dominated forest and forested peatland. ³⁾ % of standing volume

Table 2 Discharge (sum, in mm) and mean water chemistry of streams LAE03 (reference catchment) and LAE11 (harvested catchment) for the preharvest period (June 2008 – January 10 2009), post-harvest-1 (remainder of 2009) and postharvest 2 (2010-2012). Numbers show mean / standard deviation / number for observations. Cond, conductivity (μ S cm⁻¹); labile Al (μ g L⁻¹); other major ions in mg L⁻¹; TOC, total organic C; TON, total organic N; DOM, dissolved organic matter; CN_{DOM}, CN ratio of DOM; UVabs₂₅₄, absorbance of UV at 254 nm; SUVA₂₅₄, specific UV absorbance at 254 nm.

		LAE03	LAE11	LAE03	LAE11	LAE03	LAE11
		Preharvest		Post-harvest – 1		Post-harvest – 2	
Discharge (mm)		245	245	684	763	2232	2846
MeHg (ng/L)	MeHg	0.09/ 0.02 /9	0.18/ 0.11 /10	0.08/ 0.04 /28	0.22/ 0.13 /30	0.06/ 0.02 /44	0.15/ 0.10 /44
	Hg	5.0/ 2.6 /9	5.3/ 2.3 /10	3.8/ 1.1 /28	5.0/ 1.3 /30	3.1/ 0.9 /43	4.1/ 1.6 /43
	MeHg/TOC ng/g	6.8/ 3.6 /9	11.7/ 8.3 /8	6.0/ 3.1 /28	11.1/ 7.3 /30	4.7/ 1.9 /43	8.4/ 6.0 /42
Hg,	HgT/TOC ng/g	342/116/9	241/ 70 /8	270/ 62 /28	245/ 53 /30	259/ 52 /42	227/ 71 /41
	рН	4.74 / / 22	4.44 / / 14	4.76 / / 43	4.43 / / 44	4.83 / / 52	4.50 / / 53
	H+	18.3/ 6.5 /22	36.6/ 16.1 /14	17.4/ 10.9 /43	36.8/ 12.0 /44	14.8/ 7.8 /52	31.9/ 11.6 /53
s	Cond	1.53/ 0.27 /22	2.22/ 0.59 /14	1.41/ 0.28 /15	2.18/ 0.50 /15	1.48/ 0.23 /52	2.17/ 0.38 /53
r ior	Са	0.75/ 0.11 /19	0.79/ 0.19 /13	0.78/ 0.18 /15	0.75/ 0.20 /15		
najo	Mg	0.14/ 0.02 /19	0.12/ 0.02 /13	0.14/ 0.02 /15	0.12/ 0.03 /15		
_	К	0.03/ 0.01 /19	0.06/ 0.04 /13	0.06/ 0.03 /15	0.24/ 0.10 /15		
	SO ₄ _S	0.67/ 0.29 /19	0.61/0.22/13	0.78/ 0.31 /15	0.68/ 0.23 /15	0.79/ 0.30 /27	0.67/ 0.27 /28
	Labile Al	19/ 12 /19	8/7/13	23/9/15	9/ 6 /15		
nts μg/L	NO ₃ -N	5.6/ 5.9 /22	5.1/ 5.2 /14	8.3/ 6.9 /43	4.3/ 6.6 /44	10.0/ 12.3 /41	29.9/ 29.5 /42
	NH ₄ -N	3.6/ 1.9 /21	10.1/ 11.2 /9	6.0/ 5.2 /43	10.0/ 11.4 /44	7.2/ 6.3 /41	33.0/ 37.2 /42
ıtrie	Total P	5.5/ 2.1 /22	7.0/ 2.0 /13	4.5/ 1.2 /16	8.3/ 3.9 /16	5.2/ 2.6 /47	9.4/ 5.2 /48
Γ	Total N	258/ 50 /22	320/ 54 /14	256/ 68 /43	342/ 73 /44	246/ 43 /42	382/ 115 /43
DOM	TOC (mg L ⁻¹)	14.2/ 3.7 /22	21.3/ 7.3 /14	13.9/ 5.5 /43	20.3/ 5.4 /44	12.4/ 3.4 /52	18.2/ 5.4 /53
	TON (µg L ⁻¹)	247/ 51 /21	284/ 58 /9	241/ 68 /43	328/ 74 /44	229/ 41 /41	321/ 78 /42
	CN _{DOM} g/g	57/7/21	72/ 15 /9	57/ 8 /43	62/7/44	54/8/41	57/ 12 /42
	UVabs ₂₅₄	0.61/0.14/8	0.93/ 0.18 /8	0.65/ 0.22 /43	0.97/ 0.23 /44	0.61/ 0.19 /52	0.90/ 0.27 /53
	SUVA ₂₅₄	0.045/0.002/8	0.043/0.006/8	0.047/0.003/43	0.048/0.004/44	0.049/0.004/52	0.050/0.004/53

Table 3 Discharge (sum, in mm) and element fluxes (Hg, MeHg, TOC, inorganic N species, totN and totP) of catchment LAE03 (reference catchment) and LAE11 (harvested catchment) as sum for the pre-harvest period (June - December 2008) and as the mean of the post-harvest period (2009-2012). In parentheses for LAE11 are the pre- and postharvest discharge and fluxes in % of the LAE03 value.

			Preharvest	Postharvest	
			June-December 2008	Annual mean 2009-2012	
discharge mm		LAE03	245	729	
		LAE11	245 (100)	903 (124)	
Hg	$\mu g m^{-2}$	LAE03	1.4	2.7	
		LAE11	1.6 (112)	4.1 (156)	
MeHg	د ۲	LAE03	0.019	0.041	
		LAE11	0.034 (175)	0.103 (254)	
TOC	g m ⁻²	LAE03	4.1	9.7	
		LAE11	6.2 (151)	17.1 (177)	
NH ₄ -N	mg m ⁻²	LAE03	0.9	3.7	
		LAE11	1.1 (128)	15.6 (424)	
NO ₃ -N	د ۲	LAE03	0.9	2.5	
		LAE11	0.8 (85)	12.0 (470)	
TotN	د ۲	LAE03	71	176	
		LAE11	85 (119)	311 (213)	
TotP	د ٢	LAE03	1.5	3.4	
		LAE11	1.9 (124)	7.3 (176)	

Table 4 Mean concentrations of MeHg ($\mu g g^{-1}$ dry weight), standard deviation (std) and nr of observations (n) in biofilm, stoneflies and caddisflies, grouped by sampling event and catchment (LAE03, reference; LAE11, harvested catchment). If n=2, single values are given. Significant differences within one group, between catchments, are given as a and b when p<0.0001, as c and d when p<0.005, and as e and f when p<0.1 (Students t-test).

catchment	Autumn '08	Spring '08	Autumn '09
		mean / std / n	
LAE03	5.9, 7.2 / - / 2	6.3 / 1.2 / 3	2.6 / 1.4 / 3
LAE11	5.2 / 0.8 / 4	4.8 / 1.0 / 3	4.0 / 1.0 / 3
LAE03	46 ^a / 7 / 5	43 ^a / 5 / 3	37 / 9 / 3
LAE11	166 ^b / 29 / 4	90 ^b / 5 / 3	36 / 4 / 3
LAE03	90, 91 ^e / - / 2	90° / 20 / 3	120, 130 / - / 2
LAE11	275, 420 ^f / - / 2	233 ^d / 25 / 3	120 / 20 / 3
	catchment LAE03 LAE11 LAE03 LAE11 LAE03 LAE11	catchment Autumn '08 LAE03 5.9, 7.2 / - / 2 LAE11 5.2 / 0.8 / 4 LAE03 46 ^a / 7 / 5 LAE11 166 ^b / 29 / 4 LAE03 90, 91 ^e / - / 2 LAE11 275, 420 ^f / - / 2	catchment Autumn '08 Spring '08 mean / std / n LAE03 5.9, 7.2 / - / 2 6.3 / 1.2 / 3 LAE11 5.2 / 0.8 / 4 4.8 / 1.0 / 3 LAE03 46 ^a / 7 / 5 43 ^a / 5 / 3 LAE11 166 ^b / 29 / 4 90 ^b / 5 / 3 LAE03 90, 91 ^e / - / 2 90 ^c / 20 / 3 LAE11 275, 420 ^f / - / 2 233 ^d / 25 / 3



Figure 2a Click here to download high resolution image



Figure 2b Click here to download high resolution image





Highlights

- Forest harvest may increase MeHg in streamwaters and aquatic food chain
- Streamwater chemistry and MeHg in biota were studied in a paired-catchment design
- Concentrations of nutrients increased but not streamwater MeHg and total Hg
- MeHg in invertebrates was reduced after harvest related to improved nutrient access

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