- Technical Note: Preventing CO₂ overestimation from mercuric or 1
- copper (II) chloride preservation of dissolved greenhouse gases in 2
- freshwater samples

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Abstract

The determination of dissolved gases (O2, CO2, CH4, N2O, N2) in surface waters allows to estimate biological processes and greenhouse gas fluxes in aquatic ecosystems. Mercuric chloride (HgCl₂) has been widely used to preserve water samples prior to gas analysis. However, alternates are needed because of the environmental impacts and regulation prohibition of mercury. HgCl2 is a weak acid and interferes with dissolved organic carbon (DOC). Hence, we tested the effect of HgCl2 and two substitutes (copper (II) chloride - CuCl₂ and silver nitrate - AgNO₃), as well as storage time (24h to 3 months) on the determination of dissolved gases in low ionic strength and high DOC lake-water from a typical boreal lake. Furthermore, we investigated and predicted the effect of HgCl₂ on CO₂ concentrations in periodic samples from another lake experiencing pH variations (5.4-7.3) related to in situ photosynthesis. Samples fixed with inhibitors generally showed negligible O_2 consumption. However, effective preservation of dissolved CO₂, CH₄ and N₂O for up to three months prior to dissolved gas analysis, was only achieved with AgNO3. In contrast, HgCl2 and CuCl2 caused an initial increase in CO_2 and N_2O from 24h to 3 weeks followed by a decrease from 3 weeks to 3 months. The CO₂ overestimation, caused by HgCl₂-acidification and shift in the carbonate equilibrium, can be calculated from predictions of chemical speciation. Errors due to CO2 overestimation in HgCl2preserved water, sampled from low ionic strength and high DOC freshwater that are common in the northern hemisphere, could lead to an overestimation of the CO₂ diffusion efflux by a factor of >20 over a month, or a factor of 2 over the ice-free season. The use of HgCl2 and CuCl2 for freshwater

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preservation should therefore be discontinued. Further testing of AgNO3 preservation should be

performed under a large range of freshwater chemical characteristics.

- 36 **Key-words**: lake, greenhouse gases, water sample preservation, mercuric chloride, metal toxicity,
- 37 carbon dioxide
- **Running tile**: CO₂ overestimation from HgCl₂ fixation

1 Introduction

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73 74 The determination of dissolved gases by gas chromatography from water samples collected in the field allows the estimation of biological processes in aquatic ecosystems such as photosynthesis and oxic respiration (O2, CO2), denitrification (N2, N2O) and methanogenesis (CH4). This technique is also useful to test the calibration of in-situ sensors in long term deployment. However, the accuracy of this approach largely depends on the effectiveness of sample fixation. In fact, 7the partial pressure of these dissolved gases will continue to evolve in the water sample from the time of collection to the time of analysis unless biological activity is prevented. This is an issue when field sites are far from laboratory facilities, and it may be more efficient to process thewhen samples need to be stored in large batches at until the end of the field season for more efficient processing in large batches. Hence, before using a given biocide to preserve water samples, it must be ensured that it is efficient in inhibiting biological activity without changing the sample's chemistry. Mercuric (II) chloride (HgCl₂) has been widely used as an inhibitor of the above-mentioned biological processes to preserve water samples for the determination of dissolved CO2 in seawaters (e.g. Dickson, Sabine & Christian, 2007) and several dissolved gases in natural and artificial freshwater bodies (e.g. O₂, CO₂, CH₄, N₂ and/or N₂O; (Guérin et al., 2006; Hessen et al., 2017; Hilgert et al., 2019; Okuku et al., 2019; Schubert et al., 2012; Xiao et al., 2014; Yan et al., 2018; Yang et al., 2015) because it is extremely toxic at very low concentrations compared to other reagents (e.g. Horvatić & Peršić, 2007; Hassen et al., 1998). Worldwide efforts have sought to reduce the use of mercury because it is considered toxic to the environment and exposure can severely affect human health (Chen et al., 2018). Therefore, alternative preservation techniques to HgCl₂ amendment have been tested for dissolved inorganic carbon (DIC) and δ^{13} C-DIC such as acidification with phosphoric acid (Taipale & Sonninen, 2009) or a combination of filtration and exposure to benzalkonium chloride or sodium chloride (Takahashi et al., 2019). At least two studies, one also including dissolved organic carbon (DOC) and $\delta^{13}\text{C-DOC}$, showed that simple filtration (and cooling), fixation (precipitation) or acidification were better effective in preserving water samples than the use of toxic inhibitors, including HgCl₂ (Wilson, Munizzi & Erhardt, 2020). Another solution is to sample the headspace out in the field, and bring back gas samples (e.g., Cole et al., 1994; Karlsson et al., 2013; Kling et al., 1991). However, these techniques were not tested for the simultaneous determination of several dissolved gases, including CH₄ which is subject to rapid degassing during handling or storage if samples are not preserved because of its low solubility in water (Duan & Mao, 2006). In addition, some of the existing alternatives, such as filtration or field headspace equilibration, are difficult to operate in remote areas in the field under harsh weather conditions and prone to potential ambient air contamination. Solutions for water sample preservation should therefore involve a minimum of

manipulation steps in the field to avoid gas exchange with ambient air. Biocide amendments into

sealed water bottles appears as one of the most efficient methods. Copper(II) chloride (CuCl ₂) and	
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silver nitrate (AgNO ₃), the most toxic form of Silver, are relevant alternatives to HgCl ₂ given their	Formatted: Subscript
known toxicity (e.g., Ratte 2009; Amorim and Scott-Fordsmand 2012) and wide application in water	Formatted: Subscript
treatments and water purification (Larrañaga et al., 2016; Nowack et al., 2011; NPIRS, 2023; Ullmann	
et al., 1985). Nevertheless, the efficiency of these alternative biocides has never been tested for	
dissolved gas samples preservation.	
The addition of $HgCl_2$ to water is known to produce hydrochloric acid through hydrolysis (Ciavatta &	
Grimaldi, 1968) and to form complexes with many environmental ligands, both inorganic (Powell et	
al., 2004) and organic (Tipping, 2007; Foti et al., 2009; Liang et al., 2019; Chen et al., 2017). The	
complexation of Hg^+ with the carboxyl or thiol groups of $\frac{\text{dissolved organic carbon}}{\mathrm{DOC}}$ in oxic	
environments could further increase the concentration of H ⁺ (Khwaja et al., 2006; Skyllberg, 2008).	
This acidification could-can be an issue in poorly buffered water (low ionic strength) with high	
concentration of DOC where a shift in the pH and carbonate equilibrium equilib	
case, the estimated CO ₂ concentration would be higher after HgCl ₂ fixation than the in situ	Formatted: Subscript
concentration, and if the shift in pH is not accounted for, can thus resulting in an overestimation of	
dissolved CO ₂ and bicarbonate concentrations. A similar acidification effect is also expected with	
CuCl ₂ amendments (Rippner et al., 2021), but not for AgNO ₃ amendments. Such effects would not be	Formatted: Subscript
expected in marine water due to the high ionic strength of the water (Chou et al., 2016) or freshwater	Formatted: Subscript
with low pH (<5.5) under which conditions nearly all dissolved inorganic carbon is CO_2 (Stumm &	
Morgan, 1981). Thus, there are clear limits of the application of HgCl ₂ -, and possibly CuCl ₂ , for	Formatted: Subscript
freshwater sample preservation given its risk of leading to overestimation of CO2 and bicarbonate	Formatted: Subscript
concentrations, in addition to exposing field workers to the risks of its high toxicity.	
The use of HgCl₂ to preserve water samples prior to dissolved gas analyses is part of the current	
guidelines for greenhouse gas measurements in freshwater reservoirs (Machado Damazio et al., 2012;	
UNESCO/IHA, 2008, 2010). Hence, there is a risk of overestimating CO ₂ concentrations and	
emissions, in absence of discrete measurement of emissions, from hydropower reservoirs with	
consequence on the present and expected greenhouse gas footprint from hydroelectricity. To ensure	
precise estimation of greenhouse gas emission from hydropower, the largest present and future	
renewable source of electricity (IEA, 2020), the use of HgCl ₂ should therefore be discontinued.	
Here we combine data from laboratory experiments (i) and field work (ii) to illustrate risks of mis-	
estimation of dissolved gas concentrations in freshwaters with some preservatives and provide	
recommendation for best practices in the field. First, we (i) performed some short-term and long-term	
incubations of water from a typical heterotrophic unproductive boreal lake with circumneutral pH,	
incubations of water from a typical heterotrophic unproductive boreal lake with circumneutral pH, low ionic strength (poor buffering capacity) and high DOC concentration to tested the effect of	
	treatments and water purification (Larrañaga et al., 2016; Nowack et al., 2011; NPIRS, 2023; Ullmann et al., 1985). Nevertheless, the efficiency of these alternative biocides has never been tested for dissolved gas samples preservation. The addition of HgCl ₂ to water is known to produce hydrochloric acid through hydrolysis (Ciavatta & Grimaldi, 1968) and to form complexes with many environmental ligands, both inorganic (Powell et al., 2004) and organic (Tipping, 2007; Foti et al., 2009; Liang et al., 2019; Chen et al., 2017). The complexation of Hg' with the carboxyl or thiol groups of dissolved organic earbonDOC in oxic environments could further increase the concentration of H' (Khwaja et al., 2006; Skyllberg, 2008). This acidification eould-gan be an issue in poorly buffered water (low ionic strength) with high concentration of DOC where a shift in the pH and carbonate equilibrium eould-gan be induced. In that case, the estimated CO ₂ concentration would be higher after HgCl ₂ fixation than the in situ concentration, and if the shift in pH is not accounted for, can thus resulting in an overestimation of dissolved CO ₂ and bicarbonate concentrations. A similar acidification effect is also expected with CuCl ₂ amendments (Rippner et al., 2021), but not for AgNO ₃ amendments. Such effects would not be expected in marine water due to the high ionic strength of the water (Chou et al., 2016) or freshwater with low pH (<5.5) under which conditions nearly all dissolved inorganic carbon is CO ₂ (Stumm & Morgan, 1981). Thus, there are clear limits of the application of HgCl ₂ -and possibly CuCl ₂ , for freshwater sample preservation given its risk of leading to overestimation of CO ₂ and bicarbonate concentrations, in addition to exposing field workers to the risks of its high toxicity. The use of HgCl ₂ to preserve water samples prior to dissolved gas analyses is part of the current guidelines for greenhouse gas measurement of emissions, from hydropewer reservoirs with consequence on the present and expec

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111 dissolved gases (O2, CO2, CH4, N2 and N2O) by headspace equilibration and gas chromatography. in 112 The preservatives were the presence of mercuric chloride (HgCl₂) and, two alternative inhibitors, 113 chosen for their wide and effective application in water treatments and water purification (copper (II) 114 chloride - CuCl₂ and silver nitrate - AgNO₃; Xu & Imlay, 2012; Rai, Gaur & Kumar, 1981). 115 Unamended water samples, where only ultrapure water was added, were also included for 116 comparison, and a control (samples chilled in the dark at +4°C). In addition, we (ii) analysed 117 dissolved CO₂ concentration data obtained from a typical productive boreal lake using two 118 independent methods, one by gas chromatography following HgCl2 fixation, and one through 119 dissolved inorganic carbon determination without fixation. We further investigated the effect of 120 preservation with HgCl2 on the determination of dissolved CO2 from water samples collected weekly 121 during a full ice free season from a lake experiencing varying pH conditions related to photosynthesis. 122 Finally, wWe show that the overestimation of dissolved CO₂ concentrations caused by HgCl₂ fixation 123 can be predicted based on chemical equilibria. 124 125 2. Methods 126 2.1. Effects of storage time and inhibitors on the quantification determination of dissolved gases 127 Study areasite and sampling <u>Surface w</u>Water samples wereas collected from Lake Svartkulp (59.9761313 N, 10.7363544 E; 128 129 Southeast Norway) north of Oslo, Norway, on the 4th of September 2019. Lake water was earefully 130 collected into A 5 L plastic bottles was gently pushed into the water and progressively tilted to let the 131 water flow into the bottles without bubbling. The bottle aperture was covered with a 90 μm plankton 132 net to avoid sampling large particles. This procedure was repeated five times to yield a total water 133 volume of 25 L. The 5 L water bottles_andwere immediately brought back to the lab. Upon arrival at 134 the laboratory, after temperature equilibration, water from the 5 L bottles were was slowly poured, to 135 avoid limit gas exchange with the ambient airloss, into a 25 L tank to provide a single bulk sample to 136 start the incubation experiment. Filtration, e.g., with 0.45 or 0.2 µm filters, was avoided to minimize changes in dissolved gas concentrations (e.g., Magen et al., 2014). The mixed water sample (25 L) 137 138 was sub-sampled (0.5 L) for the determination of alkalinity (127 μmol L⁻¹), pH (6.73), ammonium (3 139 μg N L⁻¹), nitrate (5 μg N L⁻¹), total N (230 μg N L⁻¹), phosphate (1 μg P L⁻¹), total P (9 μg P L⁻¹) and, 140 TOC (8.9 mg C L⁻¹) and colour (59 mg L⁺Pt), all analysed by standard methods at the accredited 141 NIVA lab (see Table Table S1). In situ Ttemperature of the lake water was measured with a handheld 142 thermometer and was 18.5 °C. Note that particulate organic carbon is a negligible fraction of TOC in 143 Norwegian lake waters, representing on average less than 3% (de Wit et al., 2023).

low ionic strength (poor buffering capacity) and high DOC concentration on the determination of five

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144 Lake Svartkulp was selected for this experiment because it is representative of low ionic strength 145 Northern Hemisphere lakes, typically found in granitic bedrock regions in North-East America and 146 Scandinavia. It is a typical low-productivity, heterotrophic, slightly acidic to neutral, moderately 147 humic lake. Similar lakes are found in Southern Norway (de Wit et al., 2023), large parts of Sweden 148 (Valina et al. 2014), Finland, Atlantic Canada (Houle et al., 2022), Ontario, Québec, and North-East 149 USA (Skjelkvåle and de Wit 2011; Weyhenmeyer et al., 2019). 150 Note that particulate organic carbon is a negligible fraction of TOC in Norwegian lake waters, 151 representing less than 3% (de Wit et al., 2023). 152 Laboratory <u>incubation</u> experiment <u>with different preservatives and storage times</u> 153 154 The experimental design involved to incubate 72 borosilicate glass bottles (120 mL) filled with lake 155 water from our 25 L bulk sample and submitted to four different treatments: addition of 240µL of a 156 preservative solution of -((i) HgCl₂, (ii) CuCl₂, or a (iii) AgNO₃-, or addition of 240 μL of (iv) MilliQ 157 water. The bottles amended with MilliQ water are hereafter referred to as "unfixed".and control, i.e., 158 no added preservative), The 72 bottles were divide into three groups three time points of dissolved gas 159 analysis (t=0, i.e., within 24hwhich were incubated cold (+4°C) and dark for 24h, three weeks or three 160 months respectively, before being processed for dissolved gas analysis by gas chromatography. These 161 incubation times were selected to represent situations where samples are processed directly upon 162 return to the laboratory (24h), or after medium (3 weeks) to long (3 months) -term storage, 163 respectively., t=3 weeks and t=3 months) and six replicates, yielding in total 72 experimental 164 units At each time point and for each treatment, a group of 6 bottles were further processed for 165 dissolved gas analysis. Concentrations of O2, N2, N2O, CO2 and CH4 were determined by gas 166 chromatography (see below) using the headspace technique following Yang et al. (2015). 167 Unfortunately, pH was not measured at the end of the storage period. The same day as the water 168 samples were collected 169 In details, within 3h of lake water sampling, the 120mL bottles were gently filled with water from the 170 mixed sample (25 L). Each 120mL bottle was slowly lowered into the water and progressively tilted 171 to let the water flow into the bottle without bubbling. The bottle was then capped under water with a 172 gas tight butyl rubber stopper after ensuring that there were no air bubbles in the bottle. was added 173 gently to 72 borosilicate glass bottles (120 mL) and Tthe bottles were randomized prior to 174 preservative or MilliQ amendment. The preservative solution or MilliQ amendment was pushed in 175 each bottle with a syringe and needle through the rubber septum. To avoid overpressure, another 176 needle was placed through septum at the same time, at least 2 cm above the other needle, to allow an 177 equivalent volume of clean water to be released.

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Stock solutions of HgCl₂, CuCl₂ and AgNO₃ were prepared according to Table Tab. 1 using high accuracy chemical equipment (e.g., high accuracy scale, volumetric flasks). The Ag (Silver nitrate EMSURE® ACS; Merck KGaA, Germany), Cu (Copper(II) chloride dihydrate; Merck Life Science ApS, Norway) and Hg (Mercury(II) chloride; undetermined) salts were dissolved in MilliQ ultrapure water (>18 M Ω cm). For measurement of CO₂ in seawater samples, the standard method involves poisoning the samples by adding a saturated HgCl₂ solution in a volume equal to 0.05-0.02% of the total volume (Dickson 2007). We used this as a starting point and added 0.02 % saturated HgCl₂ solution to 1/4th18 bottles of the samples (240 μL of HgCl₂ 10× diluted saturated solution), resulting in a sample concentration of 14 µg HgCl₂ mL⁻¹ (51.6 µM; Table-Tab. 1). Based on estimated toxicity relative to Hg (Deheyn et al., 2004; Halmi et al., 2019), the silver and copper salts were added in molar concentrations equal to two and three times the molar concentration of HgCl2, respectively (Table Tab. 1), although it varies between species of microorganisms and environmental matrices (Hassen et al., 1998; Rai, Gaur & Kumar, 1981). CuCl2 and AgNO2, the most toxic form of Silver, were chosen because of their wide application in water treatments and water purification (Larrañaga et al., 2016; Nowack et al., 2011; NPIRS, 2023; Ullmann et al., 1985). Equal volume of MilliQ water was added to 1/4th of the samples as a control. The bottles were capped with gas tight butyl rubber stoppers after ensuring that there were no air bubbles in the samples. The samples for measurement of starting concentrations (t=0) were stored dark and cold (+4°C) overnight and analyzed within 24h of preparation, while the rest were stored dark and cold (+4°C) until measurement after 3 weeks (t= 3w) and 3 months (t= 3m) of storage. Concentrations of O2, N2, N2O, CO2 and CH4 were determined by phromatography (see below) using the headspace technique following Yang et al. (2015).

Table 1. Stock and sample concentrations of HgCl₂, CuCl₂ and AgNO₃.

Salt	Stock solution	Sample concentration	Rationale
HgCl ₂	70 g/L (saturated)	14.0 μg/mL (51.6 μM)	Dickson, Sabine & Christian, 2007
$CuCl_2$	131.9 g/L	$26.4~\mu g/mL~(154.7~\mu M)$	$3 \times Hg$
$AgNO_3$	87.6 g/L	$17.5~\mu g/mL~(103.1~\mu M)$	$2 \times Hg$

Additional 24h incubation experiment with different preservatives for pH measurements

Since pH was not measured at the end of the first incubation experiment, we performed an additional experiment to document any potential rapid (within 24h) impacts of preservative on pH. A total of 48 borosilicate glass bottles (120 mL) filled with lake water were submitted to the same four different treatments as the first experiment described above: HgCl₂, CuCl₂, AgNO₃ or MilliQ water amendments. To this end, a 20L water tank was filled with surface water from Lake Svartkulp on the 14th of December 2023. The water tank was immediately returned to the laboratory and left for 24h to

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CO_2 overestimation from $HgCl_2$ fixation – Clayer et al.

209	equilibrate to the room temperature. On December 15 th , 120mL bottles were gently filled with water	Formatted: Superscript
210	from the bulk 20L sample, as described above. The bottles were randomized prior to preservative or	
211	MilliQ amendment performed as described above. The bottles were then incubated at room	
212	temperature for 2h or 24h. pH was measured in the initial unamended lake water, in 24 bottles opened	
213	after 2h incubation, and in 24 bottles opened after 24h incubation. pH measurements were performed	
214	with a WTW Multi 3620 pH meter calibrated using a two-point calibration at pH = 4 and pH = 7. All	
215	pH measures were corrected for temperature. Water temperature of the water samples during pH	
216	measurements ranged between 19.1 and 21.2°C,	Formatted: Font: Not Italic
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218	2.2. Effects of HgCl ₂ on dissolved CO ₂ analyses over a range of pH values	
219	Lake Lundebyvannet time seriesStudy site and sampling	
 220	Water samples were collected from Lake Lundebyvannet located southeast of Oslo (59.54911 N,	
221	11.47843 E, Southeast Norway). The lake has a surface area of 0.4 km ² and a maximum depth of	
222	5.5 m. Two sets of Ssamples were taken from 1, 1.5, 2 and 2.5 m depth using a water sampler once or	
223	twice a week between April 2020 and January 2021 for the determination of (i) dissolved CO2 by GC	Formatted: Subscript
224	analysis following fixation with HgCl2 and (ii) DIC analysis with a TOC analyser. Samples for GC	Formatted: Subscript
225	analysis were filled into 120 mL glass bottles (as described above for the 72 incubation bottles),	·
226	which were sealed with rubber septa under water immediately without air bubbles. Samples for GC	
227	analysis were preservation preserved prior to GC analysis (within 24h) was ensured by adding in the	
228	<u>field by adding</u> a half-saturated (at 20°C) solution of $HgCl_2$ (150 μL) through the rubber seal of each	
229	bottle using a syringe, as described above the 72 incubation bottles directly after sampling, resulting	
230	in a concentration of 161 μM similar to previous studies (Clayer et al., 2021; Hessen et al., 2017;	
231	Yang et al., 2015). Samples for DIC analysis were filled without bubbles in 100 ml Winkler glass	
232	bottles that were sealed airtight directly after sampling. These samples were not fixed in any way and	
233	were analysed by a TOC analyzer and a pH meter within two hours. Lake water tTemperature and pH	
234	were measured in-situ using a HOBO pH data loggers placed at 1, 1.5, 2 and 2.5 m (Elit, Gjerdrum,	
235	Norway).	
236	The lake Lundebyvannet has a surface area of 0.4 km² and a maximum depth of 5.5 m. It often	
237	experiences large blooms of G. semen over the summer between May and September (Hagman et al.,	
238	2015; Rohrlack, 2020). The lake water was is characterised by high and fluctuating concentrations of	
239	humic substances (with DOC concentrations ranging from 8 to 28 mg C L^{-1}), ammonium (5 to 100 μg	
240	$N L^{-1}$), nitrate (20 to 700 μ g N L^{-1}), total N (average of 612 μ g N L^{-1}), phosphate (2 to 4 μ g P L^{-1}),	Formatted: Superscript
241	total P (average of 28 µg P L ⁻¹ ; Rohrlack et al., 2020; Hagman et al., 2015), a fluctuating pH (from 5.5	
242	to 7.3), weak ionic strength with alkalinity ranging between 30 and 150 μ mol L ⁻¹ , and electric	
243	conductivity varying from 40 to 70 μS cm ⁻¹ . Temperature and pH were measured in situ using a	
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244 placed at 1, 1.5, 2 and 2.5 m (Elit, Gjordrum, Norway). For more details, see 245 Rohrlack et al. (2020). 246 Lake Lundebyvannet was selected for this experiment because it is representative of productive, low-247 ionic strength Northern Hemisphere lakes typically found in the southern part of granitic bedrock 248 regions in North-East America and Scandinavia. 249 2.3. Analytical chemistry 250 Gas chromatography 251 Headspace was prepared by gently backfilling sample bottles with 20-30 mL helium (He; 99,9999%) 252 into the closed bottle while removing a corresponding volume of water. Care was taken to control the 253 headspace pressure within 5% of ambient and a slight He overpressure was released before equilibration. The bottles were shaken horizontally at 150 rpm for 1 h to equilibrate gases between 254 255 sample and headspace. The temperature during shaking was recorded by a data logger. Immediately after shaking, the bottles were placed in an autosampler (GC-Pal, CTC, Switzerland) coupled to a gas 256 257 chromatograph (GC) with He back-flushing (Model 7890A, Agilent, Santa Clara, CA, US). 258 Headspace gas was sampled (approx. 2 mL) by a hypodermic needle connected to a peristaltic pump (Gilson Minipuls 3), which connected the autosampler with the 250 µL heated sampling loop of the 259 260 261 The GC was equipped with a 20-m wide-bore (0.53 mm) Poraplot Q column for separation of CH₄, CO2 and N2O and a 60 m wide-bore Molsieve 5Å PLOT column for separation of O2 and N2, both 262 263 operated at 38°C and with He as carrier gas. N2O and CH4 were measured with an electron capture 264 detector run at 375°C with Ar/CH₄ (80/20) as makeup gas, and a flame ionization detector, 265 respectively. CO₂, O₂, and N₂ were measured with a thermal conductivity detector (TCD). Certified 266 standards of CO2, N2O, and CH4 in He were used for calibration (AGA, Germany), whereas air was 267 used for calibrating O2 and N2. The analytical error for all gases was lower than 2%. For the Lake Lundebyvannet time series, CO₂ was separated from other gases using the 20 m wide-bore (0.53 mm) 268 269 Poraplot Q column while the other gases were not measured. 270 The results from gas chromatography give the relative concentration of dissolved gases (in ppm) in 271 the headspace in equilibrium with the water. For the lab experiment with Svartkulp samples (section 272 2.1), the concentration of dissolved gases in the water at equilibrium with the headspace were 273 calculated from the solubility of gases intemperature corrected Henry constant in water using Carroll, 274 Slupsky and Mather (1991) for CO₂, Weiss and Price (1980) for N₂O, Yamamoto, Alcauskas and 275 Crozier (1976) for CH₄, Millero, Huang and Laferiere (2002) for O₂, Hamme and Emerson (2004) for 276 N₂. For the Lake Lundebyvannet time series (section 2.2), the concentration of CO₂ in the water

samples were determined using temperature-dependent Henry's law constants given by Wilhelm,

278 Battino and Wilcock (1977). The quantities of gases in the headspace and water were summed to find

- the concentrations and partial pressures of dissolved gases from the water collected in the field as
- 280 <u>follows:</u>

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$$[gas] = \frac{p_{gas}HV_{water} + \frac{p_{gas}V_{headspace}}{RT}}{V_{water}}$$
 (Eq. 1)

- where [gas] is the gas aqueous concentration, p_{gas} is the gas partial pressure, H is the Henry constant,
- V_{water} is the volume of water sample during headspace equilibration, $V_{headspace}$ is the headspace gas
- volume during equilibration, R is the gas constant and T the temperature during headspace
- 285 <u>equilibration (recorded during shaking)</u>. The calculations were similar to Yang *et al.* (2015).
- 286
- 287 DIC analyses
- 288 DIC analysis was performed for the Lake Lundebyvannet time series using a Shimadzu TOC-V CPN
- 289 (Oslo, Norway) instrument equipped with a non-dispersive infrared (NDIR) detector with O2 as a
- 290 carrier gas at a flow rate of 100 mL min⁻¹. Two to three replicate measurements were run per sample.
- 291 The system was calibrated using a freshly prepared solution containing different concentrations of
- NaHCO3 and Na₂CO3. CO₂ concentrations in water samples ($[CO_2]$) were calculated on the bases of
- 293 temperature, pH and DIC concentrations as follows (Rohrlack et al., 2020):

[
$$CO_2$$
] = $\frac{[H^+]^2 C_T}{Z}$ (Eq. $\frac{12}{2}$

- where $[H^+]$ is the proton concentration (10^{-pH}) , C_T is the dissolved inorganic carbon concentration
- and Z is given by:

$$Z = [H^{+}]^{2} + K_{1}[H^{+}] + K_{1}K_{2}$$
 (Eq. 23)

- where K_1 and K_2 are the first and second carbonic acid dissociation constant adjusted for temperature
- 299 (pK₁ = 6.41 and pK₂ = 10.33 at 25°C; Stumm & Morgan, 1996).
- 300
- 301 2.4. Data analysis
- 302 pCO_2 and saturation deficit
- 303 Lake Lundebyvannet CO2 concentrations provided by GC and DIC analyses were converted to pCO2
- 304 (in µatm) as follows:

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$$pCO_2 = \frac{[CO_2]}{0.987 \times K_H P_{atm}}$$
 (Eq. 34)

where K_H is Henry constant for CO₂ adjusted for in-situ water temperature (Stumm & Morgan, 1996) and P_{atm} is the atmospheric pressure in bar given by:

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$$P_{atm} = (1013 - 0.1 \times altitude) \times 0.001$$
 (Eq. 45)

where altitude is the altitude <u>above sea level</u> of Lake Lundebyvannet (158 m). Finally, the CO₂
 saturation deficit (Sat_{CO₂} in μatm) was <u>approximatedgiven</u> by

Sat_{CO₂} =
$$pCO_2 - [CO_2]_{air}$$
 (Eq. 56)

where $[CO_2]_{air}$ is the pCO₂ in the air (416 µatm for 2020 in Southern Norway retrieved from EBAS database; NILU, 2022; Tørseth et al., 2012). Sat_{CO_2} gives the direction of CO₂ flux at the air-water-atmosphere interface, and its product with gas transfer velocity determine the CO₂ flux at the water-atmosphere interface, i.e., whether lake ecosystems are sink ($Sat_{CO_2} < 0$) or source ($Sat_{CO_2} > 0$) of atmospheric CO₂.

318 Statistical analyses

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- The effect of storage time and treatment on five dissolved gases $(O_2, N_2, CO_2, CH_4, N_2O)$ from the
- 320 Lake Svartkulp samples was tested with a two-way ANOVA at an alpha level adapted using the
- Bonferroni correction for multiple testing, i.e., $\alpha = 0.05/5 = 0.01$. To evaluate the impact of Hg fixation
- 322 on Lake Lundebyvannet samples, $[CO_2]$ values determined by headspace equilibration and GC
- analysis of HgCl₂-fixed samples were compared with those calculated from DIC measurements of
- 324 unfixed samples with a paired t-test.
- 325 A regression analysis was performed to describe the overestimation of CO₂ concentrations caused by
- 326 HgCl₂ fixation in Lake Lundebyvannet samples as a function of pH. The total CO₂ concentration in
- 327 the HgCl₂-fixed samples ($[CO_2]_{HgCl_2}$) can be expressed as:

[
$$CO_2$$
] _{$HqCl_2$} = [CO_2] _{i} + [CO_2] _{ex} (Eq. 67)

- where $[CO_2]_i$ is the initial CO₂ concentration prior to HgCl₂ fixation, i.e., CO₂ concentration in the
- unfixed samples, and $[CO_2]_{ex}$ is the excess CO_2 concentration caused by a decrease in pH following
- HgCl₂ fixation. The relative CO₂ overestimation ($E \underline{\text{in } \%}$) is given by:

332
$$E = \frac{[CO_2]_{HgCl_2} - [CO_2]_i}{[CO_2]_i} = \frac{[CO_2]_{ex}}{[CO_2]_i}$$
 (Eq. 78)

- The impact of pH (or $[H^+]$) on E was mathematically described by running a regression analysis
- using MATLAB®. The *fminsearch* MATLAB function from the Optimization toolbox was used to
- find the minimum sum of squared residuals (SSR) for functions of the form of: $E = A/[H^+]$ or $E = A/[H^+]$

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empirically from observed $[CO_2]_i$ and $[CO_2]_{ex}$. 338 339 340 Chemical speciation, saturation-index calculations, and prediction of CO₂ overestimation 341 The speciation of solutes and saturation index values (SI) of selected minerals were calculated with the program PHREEQC developed by the USGS (Parkhurst & Appelo, 2013), neglecting the effect of 342 dissolved organic matter. This was used to assess the impact of the addition of preservative on shifting 343 344 the carbonate equilibrium as well as dissolved inorganic carbon losses due to carbonate mineral 345 precipitation. For each PHREEQC simulation, two files, respectively the database (with input 346 reactions) and input files, were used to define the thermodynamic model and the type of calculations 347 to perform. The database of MINTEQA2 (e.g., minteq.dat, Allison et al., 1991) was used to describe 348 the chemical system because it includes, inter alia, reactions and constants for Ag, Cu and Hg 349 complexation with Cl, NO₃ and carbonates. In total, three simulations were run representing the 350 addition of each preservative solution to sample water from Lake Syartkulp. The input files described 351 the composition of two aqueous solutions: (i) the preservative solution assumed to contain only the 352 preservative and (ii) sample water from Lake Svartkulp with observed major element concentrations 353 (pH, Al, Ca, Cl, Cu, Fe, Mg, Mn, N as nitrate, K, Na, S as sulfate, Zn; Table Tab. S1) and Hg and Ag concentration assumed to be 10⁻⁵ mg/L. The output file provided the activities of the various solutes in 354 355 the preserved samples, i.e., simulating the mixing of 120 mL of lake water with 240 μ L of the AgNO₃, 356 CuCl₂ and HgCl₂ preservative solutions, as described in section 2.1. This procedure allows to estimate 357 the pH of the preserved samples as well as SI for various mineral phases. The SI is calculated by 358 PHREEQC comparing the chemical activities of the dissolved ions of a mineral (ion activity product, 359 IAP) with their solubility product (Ks). When SI > 1, precipitation is thermodynamically favourable. 360 However, PHREEQC does not give information about precipitation kinetics. PHREEQC was also used to estimate the decrease in pH caused by adding 150 μL of a half-saturated 361 362 HgCl₂ solution to Lake Lundebyvannet samples prior to GC analyses, as described in section 2.2. In absence of data on the chemical composition of Lake Lundebyvannet, we assumed that it had the 363 same composition as Lake Svartkulp water samples. This assumption is supported by the fact that 364 365 waters from both lakes have circumneutral pH, low ionic strength (poor buffering capacity) and high 366 DOC concentration and would therefore behave similarly in presence of acids. Briefly, for each 0.1 pH value between pH of 5.4 and 7.3, the carbonate alkalinity was first adjusted by increasing HCO₃ 367 concentrations in the input files for PHREEQC to confirm that the water was at equilibrium at the 368 given pH value. Then, the effect of adding 150µL of a half-saturated HgCl₂ solution was simulated as 369 described above for Lake Svartkulp. Knowing the new equilibrated pH, after addition of HgCl2, the 370

 $A \times 10^{-B \times pH}$. For each optimal solution, the root-mean-square error (RMSE) and coefficient of

determination (\mathbb{R}^2) were calculated against observed values of E, i.e., values of E determined

371 overestimation of CO₂ concentration in Hg-fixed samples relative to unfixed samples (E, described in

- Eq. $\frac{78}{8}$ above) can be predicted as described below.
- Adapting Eq. $(\frac{12}{2})$, we obtain:

$$[CO_2]_{HgCl_2} = \frac{[H^+]_{HgCl_2}^2 c_T}{Z_{HgCl_2}}$$
 (Eq. 89)

375 and

[
$$CO_2$$
]_i = $\frac{[H^+]_i^2 C_T}{Z_i}$ (Eq. 910)

- where $[H^+]_i$ is the proton concentration measured in the initial water samples prior to HgCl₂ fixation,
- and $[H^+]_{HgCl_2}$ is the proton concentration estimated by PHREEQC following HgCl₂ fixation, and
- similarly for Z_i and Z_{HgCl_2} from Eq. (23). Combining Eqs. (67), (89) and (910) we obtain:

380
$$[CO_2]_{ex} = C_T \left(\frac{[H^+]_{HgCl_2}^2}{Z_{HgCl_2}} - \frac{[H^+]_i^2}{Z_i} \right)$$
 (Eq. $\frac{10}{1}$

381 Hence:

383

382
$$E = \frac{[CO_2]_{ex}}{[CO_2]_i} = \frac{\left(\frac{[H^+]_{HgCl_2}^2 - [H^+]_i^2}{Z_{HgCl_2}^2}\right)}{\frac{[H^+]_i^2}{Z_i}}$$
(Eq. $\frac{1112}{C_i}$)

Alternatively, *E* can also simply be predicted based on the carbonic acid dissociation:

385
$$CO_2 + H_2O \stackrel{K_1}{\Leftrightarrow} HCO_3^- + H^+$$
 (Reaction 1)

386 At equilibrium, we have:

387
$$K_1 = \frac{[HCO_3^-][H^+]}{[CO_2]}$$
 (Eq. $\frac{1213}{}$)

- When pH is decreased upon addition of $HgCl_2$, a fraction (α) of the initial bicarbonate concentration
- [HCO_3^-]_i is turned into CO_2 . This fraction, expressed as $[CO_2]_{ex}$ in Eq. (67) above, can be estimated
- 390 with Eq. <u>12-13</u> as follows:

391
$$[CO_2]_{ex} = \alpha [HCO_3^-]_i = \frac{\alpha K_1 [CO_2]_i}{[H^+]_i}$$
 (Eq. 1314)

Introducing the expression of $[CO_2]_{ex}$ from Eq. 13-14 into Eq. 7-8 yields:

393
$$\frac{[CO_2]_{ex}}{[CO_2]_i} = E = \frac{\alpha K_1}{[H^+]_i}$$
 (Eq. 1415)

394 When the decrease in pH, or acidification, is greater than the buffering capacity of the water: $\alpha=1$. 395 The value of α cannot exceed 1 because the amount of CO_2 produced by a decrease in pH cannot 396 exceed the amount of HCO_3^- initially present. In all the other cases, we have: $\alpha<1$. For both 397 predictions of E, i.e., with Eqs. $\frac{11-12}{2}$ and $\frac{1415}{2}$, the root-mean-square error (RMSE) and coefficient 398 of determination (R^2) were calculated.

Finally, additional sources of CO₂ overestimation were investigated by analysing the residuals of the model described by Eq. $\frac{1+12}{2}$, i.e., the difference between E predicted with Eq. $\frac{1+12}{2}$ and E determined empirically with Eq. $\frac{78}{2}$. Briefly, residuals were plotted against pH and in situ temperature. Residuals were separated in two groups based on the empirical value of $[HCO_3^-]_i - [CO_2]_{ex}$, i.e., the first group had values of $[HCO_3^-]_i - [CO_2]_{ex} \ge a$ while the second group had values of $[HCO_3^-]_i - [CO_2]_{ex} \le -a$ where different values for a were used: 20, 10 or 5 μ M. The justification for separating residuals in two groups is that: (i) the first group represents samples for which bicarbonate alkalinity in the original sample is, as expected, higher than CO₂ overestimation after HgCl₂-fixation, while (ii) the second group represents samples for which bicarbonate alkalinity is not sufficient to explain CO₂ overestimation after HgCl₂-fixation.

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- 410 CO2 diffusion fluxes from Lake Lundebyvannet
- The diffusive flux of CO₂ (F_{CO_2} in mol m⁻² d⁻¹) from Lake Lundebyvannet surface water was estimated according to:

413
$$F_{CO_2} = \frac{k_{CO_2}([CO_2] - [CO_2]_{eq})}{1000}$$
 (Eq. $\frac{1216}{100}$)

where k_{CO_2} is the CO₂ transfer velocity in m d⁻¹, [CO_2] is the surface water CO₂ concentration (μ M), and 1000 is a factor to ensure consistency in the units and [CO_2]_{eq} is the theoretical water CO₂ concentration (μ M) in equilibrium with atmospheric CO₂ concentration calculated with Eq. (3) and pCO₂ of 416 μ atm (see above).

The CO₂ transfer velocity (k_{CO_2}) was estimated as follows (Vachon & Prairie, 2013):

419
$$k_{CO_2} = k_{600} \left(\frac{600}{5c_{CO_2}}\right)^{-n}$$
 (Eq. 4417)

where k_{600} is the gas transfer velocity (m d⁻¹) estimated from empirical wind-based models and Sc_{CO_2} is the CO₂ Schmidt number for in situ water a given temperature (unitless; Wanninkhof, 2014). We used n values of 0.5 or 2/3 when wind speed was below or above 3.7 m s⁻¹, respectively (Guérin et al., 2007). Empirical k_{600} models included those from Cole & Caraco (1998; $k_{600} = 2.07 + 0.215U_{10}^{1.7}$), Vachon & Prairie (2013; $k_{600} = 2.51 + 1.48U_{10} + 0.39U_{10}\log_{10}LA$) and Crusius & Wanninkhof

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(2003; power model: $k_{600} = 0.228U_{10}^{2.2} + 0.168$ in cm h⁻¹). U_{10} and LA refer to mean wind speed at 10 m in m s⁻¹ and lake area in km², respectively. Sub-hourly U_{10} data for 2020 was retrieved from a 426 427 weather station of the Norwegian Meteorological Institute located 1.5 km west of Lake Lundebyvannet (station name: E18 Melleby; ID: SN 3480; 59.546 N, 11.4535E) using the Frost 428 429 application programming interface (Frost API, 2022). Daily, monthly, and yearly (only covering the 430 ice-free season: April-November) F_{CO_2} was estimated using Eq. (12). Daily [CO₂] was interpolated from weekly data using a modified Akima spline (makima spline in Matlab® based on Akima, 1974). 431 432 This interpolation method is known to avoid excessive local undulations. 433 3. Results and discussion 434 3.1. Effects of inhibitors preservatives and storage time on dissolved gases 435 In the control unfixed samples from Lake Svartkulp, the concentration of O₂ declined while CO₂ 436 increased over time in a close to 1:1 molar ratio, likely reflecting the effect of microbial respiration 437 activity and mineralisation of organic matter (Fig. 1, Table-Tab. S2). Concentration of O2 in the 438 control unfixed decreased from near 300 to below 200 µM (Fig. 1). In the presence of inhibitors, O2 439 concentrations tended to be slightly higher at t=0.24h and remained constant or declined only slightly 440 over time to generally remain at or above saturation (280 to 300 µM). Thus, the inhibitors were 441 effective in reducing oxic respiration. 442 The concentration of CO_2 in the presence of $AgNO_3$ at $t = \frac{0.24h}{0.200}$ was not significantly different to the 443 control unfixed at t = 0 (Fig 1; paired t-test, P > 0.1). At $t = \frac{0.24h}{0.00}$, CO₂ concentrations were however 444 much higher in the presence of HgCl₂ (135 μM) or CuCl₂ (131 μM) than in the control unfixed (89 445 μM; Fig 1, Table Tab. S2). This is likely due to an acidification of the poorly buffered (alkalinity 127 446 µM) and near neutral water (pH=6.73), shifting the carbonate equilibrium from HCO₃ to CO₂-as also 447 shown by Borges et al. (2019). The increase of CO_2 further increases from 130 μ M to ~160 μ M after 448 3 weeks in both sample sets preserved with HgCl2 and CuCl2 while a decrease in O2 is less 449 pronounced not mirrored by a similar decrease in O₂ for samples fixed with CuCl₂ and completely 450 absent for samples fixed with HgCl2. This suggests that oxic respiration is not the main source for this 451 additional 30 µM of CO2 but rather points towards additional acidification of the samples caused by 452 kinetically controlled complexation of Hg2+ with dissolved organic matter (Miller et al., 2009). In fact, 453 the relatively slow complexation of Hg²⁺ with organic thiol groups can release two protons 454 (Skyllberg, 2008) and up to three, with some participation of a third weak acid group (Khwaja et al., 455 2006). The following decrease in CO₂ after 3 months (down to ~145 µM) points to other processes. 456 Overall, the addition of HgCl₂ or CuCl₂ following sampling increased CO₂ concentrations by 47% 457 within the firstafter 24h compared to the eontrol unfixed and caused further changes over the three-458 month storage time, while preservation with AgNO3 yielded CO2 concentrations consistent with the

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control unfixed and caused negligible changes over time (Fig. 1; paired t-test, P>0.1).

460	The concentration of CH4 $\underline{across\ all\ samples}$ ranged between 0.017 and 0.377 μM (Fig. 1), as
461	expected two orders of magnitude smaller than CO ₂ . At $t = \frac{024h}{100}$, the concentration of CH ₄ was over
462	$0.2~\mu M$ in the presence of inhibitors while it was below saturation in the <u>control unfixed</u> (0.03 μM ;
463	Fig. 1). CH ₄ oversaturation in the preserved samples persisted after three weeks and three months of
464	storage and CH ₄ concentration remained constant unchanged (Fig. 1, Table Tab. S2). Both
465	observations are consistent with the fact that the three preservatives were effective in preserving CH ₄
466	from oxidation. Even at t = 0, i.e., for samples analysed within 24h after sampling, preservatives are
467	required to preserve CH4 in oxic samples.

 CO_2 overestimation from $HgCl_2$ fixation – Clayer et al.

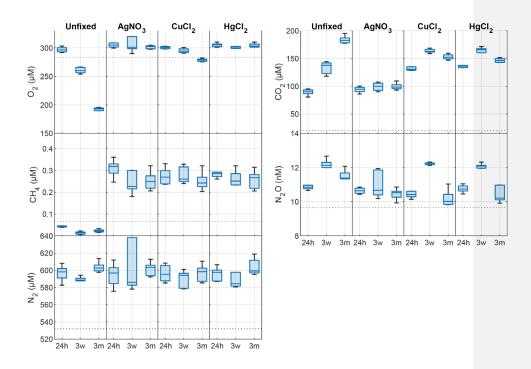


Fig 1. Changes in dissolved O₂, CO₂, CH₄, N₂O and N₂ concentrations (nM or μM) in the absence (controlunfixed) and presence of different preservatives (AgNO₃, CuCl₂, HgCl₂) at three times (θ24h, day after sample collection24h after incubation start; 3w, three weeks after collection; 3m, three months after collection). The horizontal dotted line is the saturated gas concentration corresponding to 100% gas saturation at *in situ* lake temperature. Box plots show the median, 25th and 75th percentiles and the whiskers display all data coverage.

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CO_2 overestimation from $HgCl_2$ fixation – Clayer et al.

475	In fact, oxic methanotrophy typically show rates in the order of μM day ¹ (Thottathil et al., 2019; van	
476	Grinsven et al., 2021). Hence, a CH ₄ consumption of 0.3 μM within 24h in the unpreserved, control	
477	water samples is realistic (Fig. 1).	
 478	The concentration of N ₂ O ranged between 9.8 and 12.7 nM with only samples preserved with AgNO ₃	
479	showing negligible changes over time (Fig. 1; paired t-test, P>0.1). All the other samples showed	
480	consistent patterns with storage time. N ₂ O concentrations initially increased within the first 3 weeks,	
481	followed by a decrease after 3 months. Similar patterns of net N ₂ O production followed by net	
482	consumption were also reported in short term incubations of seawater from the high latitude Atlantic	
483	Ocean, although over much shorter timescales, i.e., 48 and 96h (Rees et al., 2021). The lack of	
484	inhibition of N ₂ O production and consumption in the samples preserved with HgCl ₂ and CuCl ₂ can be	
485	attributed to the fact that N ₂ O production tends to increase under increasing acidic conditions	
486	(Knowles, 1982; Mørkved et al., 2007; Seitzinger, 1988). In fact, the mole fraction of N ₂ O produced	
487	during denitrification increases compared to N ₂ as pH decreases (Knowles, 1982). In summary,	
488	AgNO ₂ appears to be the only preservative inhibiting N ₂ O cycling, although further tests with a larger	
489	range of N ₂ O concentrations are required to confirm its efficiency.	
490	The changes in N ₂ were likely within handling and analytical errors and not different in the presence	
491	or absence of inhibitors (Fig. 1; Table Tab. S2; paired t-test, P>0.1).	
	<u> </u>	
492		
493	3.2. Effects of preservatives on pH	
494	In the samples amended with ultrapure water or AgNO ₃ , the pH did not show any significant changes	Formatted: Subscript
495	after 2h or 24h. In contrast, both groups with HgCl ₂ and CuCl ₂ amendments show significant	Formatted: Subscript
496	decreases of pH after 2h, -0.12 and -0.19, respectively, and 24h, -0.16 and -0.21, respectively. In	Formatted: Subscript
497	addition, they showed a significant decrease in pH from 2h to 24h. Samples amended with CuCl ₂	Formatted: Subscript
498	show that strongest decrease in pH.	

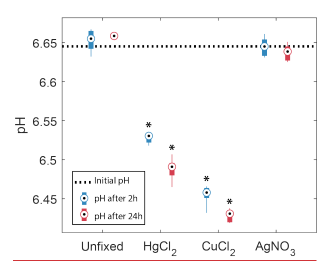


Fig 2. Changes in pH in the absence (unfixed) and presence of different preservatives (AgNO₃, CuCl₂, HgCl₂) at two times, 2h and 24h after the start of the incubation. The horizontal dotted line represents the initial pH of the bulk water sample. Box plots show the median, 25th and 75th percentiles and the whiskers display all data coverage of the 6 replicates. Stars indicate groups that are significantly different from each other and from the initial pH (two-way ANOVA).

3.23. Contrasting impacts of $HgCl_2$, $CuCl_2$ and $AgNO_3$ on dissolved CO_2 estimation revealed by chemical speciation modelling

The PHREEQC simulation of unpreserved samples, based on concentrations of all major elements (Table Tab. S1), predicted a pH of 6.72 (Table Tab. 2) which is very close to the measured pH of 6.73 (Table Tab. S1). This suggests that chemical information provided to PHREEQC is likely sufficient to describe the system, without having to invoke more complex reactions with dissolved organic matter. The addition of HgCl₂ and CuCl₂ both caused a significant decrease in pH to 6.40 and 6.45, respectively (Table Tab. 2). The proximity of these pH values to the first carbonic acid dissociation constant (pK₁ = 6.41 at 25°C; Stumm & Morgan, 1996) implies a significant shift in the carbonate equilibrium from HCO₂-to CO₂. In fact, introducing pH and CO₂-concentration values of 6.41–6.45 and 130 μ M, respectively, for the samples preserved with HgCl₂-and CuCl₂-into Eqs. 1 and 2 yields DIC concentrations (C₁) of about 270 μ M at t=0. These DIC concentrations are almost equal to those calculated for the control and samples preserved with AgNO₂-at t = 0, i.e., with a pH of 6.73 and CO₂ concentration of 88 μ M. Interestingly, the concentration of CO₂ in the samples preserved with HgCl₂ and CuCl₂-continue to increase up to ~160 μ M after 3 weeks. Given that oxic respiration is inhibited (Fig. 1), this additional CO₂ is believed to originate from progressive release of protons following relatively slow complexation of Hg²⁺ with dissolved organic matter (Khwaja et al., 2006; Miller et al.,

2009; Skyllberg, 2008). Note that PHREEQC could not predict complexation of Hg²⁺ with dissolved organic matter given that we neglected the effect of dissolved organic matter.

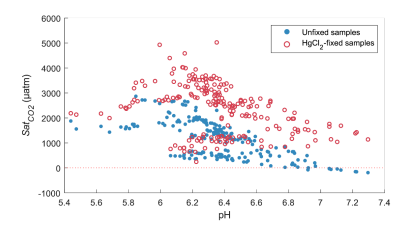
In absence of preservatives, none of the common carbonate minerals, including calcite, were associated with a saturation index higher than 1, i.e., dissolution was thermodynamically favourable for all these minerals and no DIC loss was expected (Table Tab. 2). However, upon addition of HgCl₂ or CuCl₂, some carbonate minerals, e.g., HgCO₃ or malachite and azurite, respectively, were expected to spontaneously precipitate given their relatively high saturation index values. This observation can explain the consistent decrease in CO₂ concentrations observed between three weeks and three months for sampled treated with HgCl₂-and CuCl₂- Calcite precipitation is typically observed in supersaturated solutions within 48h (Kim et al., 2020). Hence, it is realistic to consider that Hg and Cu carbonate precipitation influenced the CO₂ concentration within the preserved samples over the three months of storage time. Impacts of Hg or Cu carbonate precipitation is not evident after three weeks likely because of slow but persistent CO₂ production in presence of HgCl₂ and CuCl₂ related to acidification as described above (Fig. 1). However, after three weeks, this production likely weakens and is counterbalanced by increasing carbonate precipitation.

Table 2. pH and saturation indicesexes of selected carbonate minerals estimated by PHREEQC for the unpreserved and preserved samples

			Saturation	ind <u>ices</u> exes	
Preservatives	pН	HgCO ₃	Cu ₂ (OH) ₂ CO ₃	Cu ₂ (OH) ₂ CO ₃	Ag ₂ CO ₃
			 Malachite 	- Azurite	
<u>Unpreserved</u> <u>Unfixed</u>	6.72	-2.31	-4.96	-8.71	-16.42
$HgCl_2$	6.40	3.64	-5.89	-10.10	-17.20
$CuCl_2$	6.45	-2.55	2.26	2.11	-17.44
$AgNO_3$	6.71	-2.31	-4.97	-8.73	-4.33

3.3. Effects of HgCl₂ on dissolved CO₂ concentration under a range of pH

CO₂ concentrations in unfixed water samples from Lake Lundebyvannet were significantly lower than in the HgCl₂-fixed samples (mean difference: 52 μM; paired t-test; P<0.0001; Table-Tab. 3). Fixation with HgCl₂ caused a general overestimation of CO₂ concentration and the saturation deficit (Fig. 23), thus missing out events of CO₂ influx (carbon sink) under high photosynthesis activity (high pH; Fig. 23). As for the samples from Lake Svartkulp described above, the overestimation in CO₂ concentration likely stems from the acidification by HgCl₂ (161 μM added) shifting the carbonate equilibrium towards CO₂. In parallel fact, PHREEQC predicted a decrease of 0.6 to 1.8 units of pH related to HgCl₂ addition (Fig. S1).



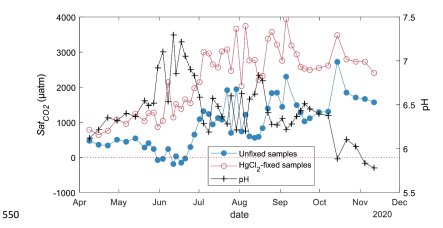


Figure 23. CO₂ saturation deficit in Lake Lundebyvannet as a function of <u>in situ</u> pH for all unfixed <u>(obtained from DIC analysis)</u> and HgCl₂-fixed <u>(obtained from GC analysis)</u> samples (top panel). Timeseries of pH and CO₂ saturation deficit of <u>Lake Lundebyvannet</u> surface water (1-m deep) for unfixed and HgCl₂-fixed samples (bottom panel).

Table 3. CO₂ concentrations ([CO₂], μM) and diffusion fluxes (F_{CO2}, mol m⁻² d⁻¹) from Lake Lake Lundebyvannet estimated from HgCl₂-fixed and unfixed samples following Cole and Caraco (1998). Ice-free season spans April to November. Data are also shown in Fig. 5

Prese	ervatives	Apr.	May	Jun.	Jul.	Aug.	Sep.	Oct.	Nov.	Ice-free season
[CO ₂]	HgCl ₂ No ne	45.3	39 .0	18.7 <u>19</u>	6 <u>8</u> 7.9	59 .2	85.1	123.4	419120 . 8	66.867
	HgCl ₂ No	67.9 <u>68</u>	75 .1	74.0	13 <u>32.9</u>	129.913 0	149.1	179.3	178 .1	121.5
	Diff (%)	<u>+50 %</u>	<u>+93 %</u>	<u>+</u> 296 %	<u>+96 %</u>	<u>+</u> 119 %	<u>+</u> 75 %	<u>+</u> 45 %	<u>+</u> 49 %	<u>+82</u> %
F_{CO2}	HgCl₂No ne,	0.10	0.07	0.01	0.15	0.11	0.23	0.37	0.48	0.17
	HgCl ₂ No	0.20	0.21	0.16	0.34	0.29	0.47	0.57	0.77	0.35
	Diff (%)	+97 - %	+188 - %	<u>+2163</u>	<u>+,130 %</u>	<u>+</u> 162 %	<u>+99 %</u>	<u>+55 %</u>	<u>+62%</u>	<u>+108</u>

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> The pH value of water samples from Lake Lundebyvannet varied between 5.4 and 7.3 (Fig 23 and 34), mainly due to marked variations in phytoplankton photosynthetic activity (Rohrlack et al., 2020). The relative overestimation of CO₂ (E) follows an exponential increase with pH and is well reproduced by a simple exponential function $(2.56 \times 10^{-5} \times 10^{1.015 \times pH}, RMSE=44\%, R^2=0.81,$ p<0.0001; Fig. 34). This exponential increase likely reflects the corresponding decrease in absolute CO2-concentration with pH (Stumm & Morgan, 1981) concomitant here to phytoplankton photosynthesis (Fig. 2). In fact, the exponential increase in CO2-overestimation is easily predicted by Eq. (8) with an equivalent level of accuracy than the optimized exponential function (Fig. 3). Consistently, the relative overestimation of $CO_2(E)$ shows an inverse decrease with $[H^+]$ that is well reproduced by a simple inverse function $(3.25 \times 10^{-5}/[H^+]; RMSE=44\%, R^2=0.81, p<0.0001; Fig.$ 3) and predicted by Eq. (14), with an α value of 1. Combining Eqs. 7 and 14 and solving it with pH values estimated from PHREEQC (Fig. S1) for α yields values ranging between 0.72 and 0.89 with an average of 0.85. Unexpectedly, this average a value is almost equal to the ratio of the inverse function = 0.87. Hence, the relative overestimation of CO₂ (E) caused by HgCl2 fixation is easily predicted by the change in bicarbonate equilibrium knowing the proton release from HgCl2 addition.

575 In waters with higher ionic strength, the release of proton from HgCl₂ addition will likely be smaller.

576 PHREEQC can be used to predict it, if sufficient knowledge is gathered on the ionic water

577 composition. Proton release during HgCl₂ fixation can be represented by the following reaction:

$$HgCl_2 + 2H_2O = Hg(OH)_2 + 2H^+ + 2Cl^-$$
 (Reaction 2)

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579	From reaction 2, it becomes evident that the initial concentration of chloride in the water samples will		
580	likely limit HgCl ₂ -dissociation and proton release. This is a likely mechanism occurring in seawater		
581	where HgCl ₂ -has been shown to cause a decrease in pH, although at a negligible level (Chou et al.,		
582	2016).		
583	4. Discussion		
303	H D ISCUSSION		
584	Prior to using dissolved gas concentrations in freshwater to estimate the magnitude of biological		
585	aquatic processes such as photosynthesis and oxic respiration, denitrification and methanogenesis, we		
586	must ensure that biological activity between sampling and laboratory analyses was efficiently		
587	inhibited without significant impacts on the sample's chemistry. Here we report a unique dataset on		
588	the impact of three preservatives on water samples from a typical low-ionic strength, unproductive		
589	boreal lake to inform on potential risks of mis-estimation of dissolved gas concentrations. We further		
590	show, using CO2 concentration data from a typical productive boreal lake, that using HgCl2 can lead		Formatted: Subscript
591	to negligence of the role of photosynthesis in lake C cycling.		Formatted: Subscript
592	4.1 Best preservative for the determination of dissolved gas concentrations		
593	Given that none of the four treatments (unfixed, HgCl ₂ , CuCl ₂ or AgNO ₃) applied to Lake Svartkulp		Formatted: Subscript
594	water samples during the 3-month incubation offer an independent control, a first challenge is to		Formatted: Subscript
595	determine which of the treatment represent the most realistic dissolved gas concentrations close to		Formatted: Subscript
596	real condition. For CO ₂ and O ₂ , a few studies have used unfixed samples (only preserved dark at		Formatted: Subscript
597	±4°C) up to 48h after sampling to determine CO ₂ or DIC concentrations (e.g., Sobek et al. 2003,		Formatted: Subscript
598	Kokic et al., 2015). So, the CO ₂ and O ₂ concentrations in the unfixed samples collected after 24h		Formatted: Subscript
599	incubation are the most representative of the initial real concentrations. Biological activity might have		
600	had an impact, but this is likely negligible over the first 24h. In addition, the fact that the CO ₂ and O ₂		
601	concentrations in the samples fixed with AgNO ₃ after 24h, three weeks and three months are equal to		Formatted: Subscript
602	those from unfixed samples after 24h (Fig. 1) confirms that the unfixed samples after 24h can be used		
603	as a control. In fact, only samples fixed with AgNO3 are trustful given the expected toxicity of Ag, the		Formatted: Subscript
604	absence of impact on pH (Fig. 2), and unchanged concentrations over the three-month experiment for		
605	all gases. Similarly, N ₂ O and N ₂ concentrations in the unfixed samples after 24h can be used as		Formatted: Subscript
606	control. However, for CH _d , Fig. 1 shows that already after 24h, the CH _d concentration in the unfixed		Formatted: Subscript
607	samples is below atmospheric saturation while it is consistently much higher in all three sets of fixed	1	Formatted: Subscript
608	samples. Boreal lakes are typically over saturated with respect to CH_4 (Valiente et al., 2022) and it is		Formatted: Subscript
609	very unlikely that CH ₄ could have been produced in lake water incubated under high concentration of	_	Formatted: Subscript
610	oxygen and toxic preservatives. Hence, unfixed samples do not represent real CH ₄ concentrations.		Formatted: Subscript
611	These observations are all consistent with the fact that the three preservatives were effective in		Formatted: Subscript
612	preserving CH ₄ from oxidation. Even at t = 24h, preservatives are required to preserve CH ₄ in oxic		

 $\underline{samples.\ In\ fact,\ oxic\ methanotrophy\ typically\ show\ rates\ in\ the\ order\ of\ \mu M\ day^{-1}\ (Thottathil\ et\ al.,$

614 615	2019; van Grinsven et al., 2021). Hence, a CH ₄ consumption of 0.3 μM within 24h in the unfixed water samples is realistic (Fig. 1).
616	In summary, preservation with AgNO ₃ is the only method that offered robust determination of all five
617	dissolved gases with negligible changes in concentration over time.
618	4.2 Risks of mis-estimating dissolved gas concentration with HgCl2 and CuCl2 preservation
619	Both sets of samples preserved with either HgCl ₂ and CuCl ₂ showed CO ₂ concentrations that were
620	much higher that the unfixed (after 24h) or the AgNO ₃ -fixed samples. This is due to an acidification
621	of the poorly buffered (alkalinity 127 μM) and near neutral water (pH=6.73), shifting the carbonate
622	equilibrium from HCO ₃ to CO ₂ as also shown by Borges et al. (2019). In fact, a rapid decrease in pH
623	was observed upon HgCl ₂ and CuCl ₂ amendments (Fig. 2). The increase of CO ₂ from about 130 μM
624	to ~160 μM after 3 weeks in both sample sets preserved with HgCl ₂ and CuCl ₂ is not mirrored by a
625	similar decrease in O ₂ (Fig. 1) This suggests that oxic respiration is not the main source for this
626	additional 30 μM of CO ₂ but rather points towards additional acidification of the samples caused, e.g.,
627	by kinetically controlled complexation of Hg ²⁺ with dissolved organic matter (Miller et al., 2009). In
628	fact, the relatively slow complexation of Hg ²⁺ with organic thiol groups can release two protons
629	(Skyllberg, 2008) and up to three, with some participation of a third weak-acid group (Khwaja et al.,
630	2006). The transient nature of acidification caused by HgCl ₂ and CuCl ₂ is also evident in the pH
631	impacts showing higher acidification after 24h than after 2h incubation (Fig. 2). The following
632	decrease in CO ₂ after 3 months (down to ~145 μM) points to other processes. The precipitation of Hg
633	and Cu carbonates, given their high saturation index values (Tab. 2), would be consistent with the
634	decrease in CO ₂ concentrations observed between three weeks and three months. Calcite precipitation
635	is typically observed in supersaturated solutions within 48h (Kim et al., 2020). Hence, it is realistic to
636	consider that Hg and Cu carbonate precipitation influenced the CO ₂ concentration within the
637	preserved samples over the three months of storage time. Impacts of Hg or Cu carbonate precipitation
638	is not evident after three weeks likely because of slow but persistent CO2 production in presence of
639	HgCl ₂ and CuCl ₂ related to acidification as described above (Fig. 1). However, after three weeks, this
640	production likely weakens and is counterbalanced by increasing carbonate precipitation.
641	Overall, the addition of HgCl ₂ or CuCl ₂ following sampling increased CO ₂ concentrations by 47%
642	within the first 24h compared to the unfixed consistent with the -0.16 to -0.21 pH-unit acidification
643	observed over the same time in the pH incubation experiment (Fig. 2) and the pH estimated with
644	PHREEQC without the interaction with dissolved organic matter (Tab. 2). In fact, introducing pH and
645	CO ₂ concentration values of 6.40–6.45 and 130 μM, respectively, for the samples preserved with
646	HgCl ₂ and CuCl ₂ into Eqs. 1 and 2 yields DIC concentrations (C _T) of about 270 μM at t=24h. These
647	DIC concentrations are almost equal to those calculated for the unfixed samples and those preserved
648	with AgNO ₃ at $t = 24h$, i.e., with a pH of 6.73 and CO ₂ concentration of 88 μ M. Interestingly, the

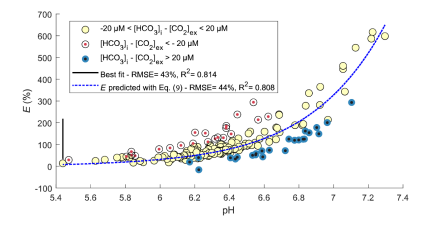
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649	$\underline{\text{concentration of CO}_2 \text{ in the samples preserved with } \underline{\text{HgCl}_2 \text{ and } \text{CuCl}_2 \text{ continue to increase up to } \sim 160}$
650	μM after 3 weeks. Given that oxic respiration is inhibited (Fig. 1), this additional CO ₂ is believed to
651	originate from progressive release of protons following relatively slow complexation of Hg ²⁺ with
652	dissolved organic matter (Khwaja et al., 2006; Miller et al., 2009; Skyllberg, 2008). Note that
653	PHREEQC could not predict complexation of Hg ²⁺ with dissolved organic matter given that we
654	neglected the effect of dissolved organic matter.
655	Unlike the AgNO ₃ -fixed samples, all the other samples showed an initial increase in N ₂ O
656	concentration from 24h to 3 weeks, followed by a decrease from three weeks to 3 months. Similar
657	patterns of net N ₂ O production followed by net consumption were also reported in short-term
658	incubations of seawater from the high latitude Atlantic Ocean, although over much shorter timescales,
659	i.e., 48 and 96h (Rees et al., 2021). The large difference in kinetics between the latter experiment
660	(Rees et al., 2021) and our incubation might be attributable to differences in incubation temperature
661	where the seawater from the high latitude Atlantic Ocean was incubated at ambient temperatures
662	while our samples were kept at +4°C. Other difference in the experimental setup might have also
663	played a role. The lack of inhibition of N ₂ O production and consumption in the samples preserved
664	with HgCl ₂ and CuCl ₂ can be attributed to the fact that N ₂ O production tends to increase under
665	increasing acidic conditions (Knowles, 1982; Mørkved et al., 2007; Seitzinger, 1988). In fact, the
666	mole fraction of N ₂ O produced during denitrification increases compared to N ₂ as pH decreases
667	(Knowles, 1982).

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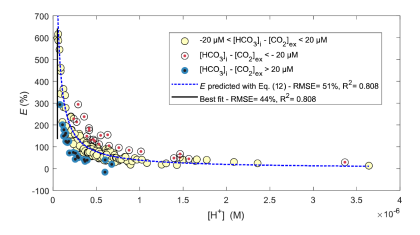


Figure 34. Comparison of observed (circles) and predicted (blue line) relative overestimation (E) of CO₂ concentrations caused by HgCl₂ fixation in Lake Lundebyvannet samples as a function of pH (top panel) or proton concentration (bottom panel). The black line shows the best fit of the regression analysis. White symbols represent samples for which the bicarbonate concentration in the unfixed samples ([HCO_3^-]_i) is nearly equal to CO₂ overestimation ([CO_2]_{ex}), i.e., \pm 20 μ M (equivalent to a pH error of 0.05), while red and blue symbols represent samples for which initial bicarbonate concentration was lower and higher than the CO₂ overestimation, respectively.

	CO2 OVERSHINGTON FIGURE 13 INCOME CHAPTER CHAPTER 1	
677	4.3 Using PHREEQC to estimate acidification caused by HgCl2 in samples from Lake Lundebyvannet	Formatted: Subscript
678	As for the samples from Lake Svartkulp as described above, the overestimation of CO ₂ concentration	Formatted: Subscript
679	in the samples from Lake Lundebyvannet fixed with HgCl ₂ (161 μM added; Fig. 3) likely stems from	Formatted: Subscript
680	the acidification shifting the carbonate equilibrium from bicarbonate to CO ₂ . In fact, PHREEQC	Formatted: Subscript
681	predicted a decrease of 0.6 to 1.8 units of pH related to HgCl ₂ addition in these samples (Fig. S1).	Formatted: Subscript
682	The relative overestimation of CO ₂ (E in Fig. 4) followed a typical exponential increase reflecting the	Formatted: Subscript
683	decrease in absolute CO ₂ concentration with increasing pH (Stumm & Morgan, 1981) caused here by	Formatted: Font: Italic
684	phytoplankton photosynthesis. In fact, the exponential increase in CO ₂ overestimation is easily	Formatted: Subscript
685	predicted by Eq. (9) with an equivalent level of accuracy as the optimized exponential function (Fig.	
686	4). Consistently, the relative overestimation of CO ₂ (E) shows an inverse decrease with [H ⁺] that is	
687	well reproduced by a simple inverse function (3.25 \times 10 ⁻⁵ /[H^+]; RMSE=44%, R ² =0.81, p<0.0001;	
688	Fig. 4) and predicted by Eq. (15), with an α value of 1. Combining Eqs. 8 and 15 and solving it with	
689	pH values estimated from PHREEQC (Fig. S1) for α yields values ranging between 0.72 and 0.89	
690	with an average of 0.85. Unexpectedly, this average α value is almost equal to the ratio of the inverse	
691	function coefficient and K_1 , i.e., $\frac{3.25 \times 10^{-5}}{K_1} = 0.87$. Hence, the relative overestimation of CO_2 (E)	
692	caused by HgCl ₂ fixation is easily predicted by the change in bicarbonate equilibrium knowing the	
693	proton release from HgCl ₂ addition, here estimated with PHREEQC.	
694	Hence, PHREEQC can be used to predict decrease in pH caused by HgCl2 fixation, if sufficient	Formatted: Subscript
695	knowledge is gathered on the ionic water composition. Proton release during HgCl ₂ fixation can be	
696	represented by the following reaction:	
697	$HgCl_2 + 2H_2O = Hg(OH)_2 + 2H^+ + 2Cl^-$ (Reaction 2)	
698	From reaction 2, it becomes evident that the initial concentration of chloride in the water samples will	
699	likely limit HgCl ₂ dissociation and proton release. This is a likely mechanism occurring in seawater	
700	where HgCl ₂ has been shown to cause a decrease in pH, although at a negligible level with a	
701	maximum decrease in pH of -0.01 (Chou et al., 2016).	
702	Figure $\underline{32}$ shows that a range of water samples were associated with a relative CO_2 overestimation (E)	
703	that substantially deviated from the overestimation predicted with Eq. 142 (red and blue symbols in	
704	Fig. $\frac{34}{2}$). In fact, some samples had a higher initial bicarbonate content ($[HCO_3^-]_i$) than the excess	
705	CO_2 concentration ($[CO_2]_{ex}$), while other showed the opposite. The former case (blue symbols in Fig.	
706	34) can easily be explained by a higher buffering capacity of the sampled water, i.e., a higher pH after	
707	HgCl ₂ -fixation than that predicted by PHREEQC related to a different water composition. Indeed, the	

concentration of major elements in the water from Lake Lundebyvannet may vary significantly over

time, and in absence of data, we considered that the water composition, except for DIC, pH and

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710 HgCl₂, was constant over time. By contrast, samples associated with $[CO_2]_{ex}$ being larger than 711 $[HCO_3^-]_i$ are more enigmatic. In order to shed light on possible explanations, we visually inspected 712 trends between empirical deviations from predictions, i.e., residuals, and in situ temperature or pH. 713 Absolute values of residuals showed a progressive increase with pH and in situ temperature which is in agreement with decreasing precision of the headspace method with increasing temperature and pH 714 715 (Koschorreck et al., 2021). In fact, CO₂ is less soluble at higher temperature, hence more gas can 716 evade during sampling, and thus the error increases with in situ temperature. In addition, at higher pH, 717 CO₂ concentration decreases and consequently the absolute error on CO₂ quantification becomes 718 larger relative to measured CO2 concentration. Interestingly, many of the high residual values were 719 not evenly distributed across the year, nor across the summer and were rather associated with only a 720 few specific sampling events during summer (Fig. S2). This suggests that degassinggas loss could 721 have occurred due to high ambient temperature in the field. Water associated with $[CO_2]_{ex}$ being 722 larger than $[HCO_3^-]_i$ (red symbols in Fig. 43 and S4) could have been subject to a larger degassing gas 723 loss-in the samples collected for DIC analysis than the samples for GC analysis. On the other hand, 724 degassing loss of gas was likely larger for samples for GC analysis than for DIC analysis for water 725 associated with $[HCO_3^-]_i$ being larger than $[CO_2]_{ex}$ (blue symbols in Fig. 43 and Fig. S2). In addition 726 to degassing gas losses and temperature effects, errors in pH measurements can also cause a large 727 misestimation of CO₂ concentration from DIC analysis, and this error increases exponentially with pH 728 following the shift in carbonate equilibrium. In summary, our analysis is consistent with that of 729 Koschorreck et al. (2021) showing that errors in the determination of CO2 concentrations are smaller 730 at lower pH and lower temperature (Fig. S2). 731 34.44. Implications for the estimation of lake and reservoir C cycling and recommendations 732 Using HgCl₂ (or CuCl₂) to preserve dissolved gas samples in poorly buffered water samples would 733 hasve large impacts on CO2 concentrations with considerable risk of leading to incorrect 734 interpretations. The risk of mis-estimating CO₂ concentration due to HgCl₂ and CuCl₂ preservation is 735 the highest when natural water pH is close to the first carbonic acid dissociation constant (pK_d = 6.41 Formatted: Subscript 736 at 25°C; Stumm & Morgan, 1996). It implies that any small shift in pH will have a significant impact 737 in the carbonate equilibrium between bicarbonate to CO2. The risk is also the highest in the lowest 738 ionic strength waters. In that respect, low-ionic strength, slightly acidic to neutral, moderately humic 739 lakes commonly found in Norway (de Wit et al., 2023), large parts of Sweden (Valina et al. 2014), 740 and Finland, Atlantic Canada (Houle et al., 2022), Ontario, Québec, and North-East USA (Skjelkvåle 741 and de Wit 2011; Weyhenmeyer et al., 2019) are the most prone to errors in CO2 concentrations Formatted: Subscript 742 related to HgCl2 or CuCl2 preservation. A significant part of these low-ionic strength lakes become Formatted: Subscript 743 increasingly sensitive to changes in nutrients with strong impacts on their role in carbon cycling Formatted: Subscript 744 (Myrstener et al., 2022). In this context, it is crucial to avoid mis-estimation of CO₂ concentrations Formatted: Subscript 745 and thus avoid use of HgCl2 or CuCl2 to ensure a robust understanding of the role of autotrophic Formatted: Subscript

746 processes in lake C cycling. Below we describe the implications for the lake C budget of 747 Lundebyvannet as an example of a mis-estimation of the role of photosynthesis in a typical productive 748 boreal lake. 749 In fact In Lake Lundebyvannet, over the ice-free season, average CO2 concentrations in Lake 750 Lundebyvannet determined following HgCl₂-fixation and GC analysis were 82% higher than those 751 obtained from DIC analyses (Table Tab. 3; Fig. 45 and S3). CO₂ concentrations obtained from HgCl₂-752 fixed samples created the illusion that Lake Lundebyvannet was a steady net source of CO2 to the 753 atmosphere over the ice-free season with large CO2 saturation deficit (Fig. 3) while, in reality, the lake 754 switched from being a net source in May, to a net sink over a few weeks in June, and returning to a 755 net source in July (Fig. 45 and S3). Indeed, monthly CO2 overestimation related to HgCl2-fixation 756 reached about 300% in June (Table Tab. 3). Propagating this overestimation into the estimates of CO2 757 diffusion fluxes with typical wind-based models yields overestimation of CO2 fluxes of 108-112% 758 over the ice-free season and up to 2100% in June (Tables-Tab. 3 and S3). Hence, interpreting CO2 759 data without correcting for CO2 overestimation caused by HgCl2-fixation leads to negligence of the 760 role of photosynthesis in lake C cycling with major implications for current and future predictions of 761 lake CO2 emissions. 762 The use of HgCl₂ to preserve water samples prior to dissolved gas analyses is part of the current 763 guidelines for greenhouse gas measurements in freshwater reservoirs (Machado Damazio et al., 2012; 764 UNESCO/IHA, 2008, 2010). Hence, there is a risk of overestimating CO₂ concentrations and 765 emissions, in absence of discrete measurement of emissions, from hydropower reservoirs with 766 consequence on the present and expected greenhouse gas footprint from hydroelectricity. To ensure 767 precise estimation of greenhouse concentration and, possibly, emission from hydropower, the use of 768 HgCl₂ should therefore be discontinued. 769 5. Conclusion 770 Mercury is a potent neurotoxin for humans and toxic for the environment and its use should be 771 discouraged, notably following the Minamata convention on mercury, a global treaty ratified by 126 772 countries (16 December 2020) to protect human health and the environment from the adverse effects 773 of mercury. This study further questions the use of HgCl2 for preservation of poorly buffered (low 774 ionic strength) water samples with high DOC concentration for analysis of dissolved gases in the 775 laboratory. Although CuCl2 is less toxic, it behaved similarly to HgCl2 and cannot be recommend. In 776 fact, both chlorinated inhibitors caused a significant decrease in pH shifting the carbonate equilibrium 777 towards CO2 and are also suspected to promote carbonate precipitation over long-term storage. The 778 only promising inhibitor tested in this study was AgNO3 notably for dissolved CO2, CH4 and N2O. 779 Silver nitrate may beis a suitable substitute for HgCl2 in low-ionic strength waters, but further tests 780 should be carried out with a range of inhibitor concentration and more diverse water samples. The use

 CO_2 overestimation from $HgCl_2$ fixation – Clayer et al.

of chemical inhibitors may not be the best approach. Alternatives exist, such as directly measuring gas concentrations *in situ* with sensors, or sampling the headspace out in the field, and bringing back gas samples (e.g., Cole et al., 1994; Karlsson et al., 2013; Kling et al., 1991; Valiente et al., 2022), rather than water samples, to the lab for gas chromatography analyses. However, care must be taken to know the exact equilibration temperature (Koschorreck et al., 2021) and to avoid gas exchange with the atmosphere as well as to use a clean background gas during headspace equilibration which can be challenging in remote environments under harsh meteorological conditions.

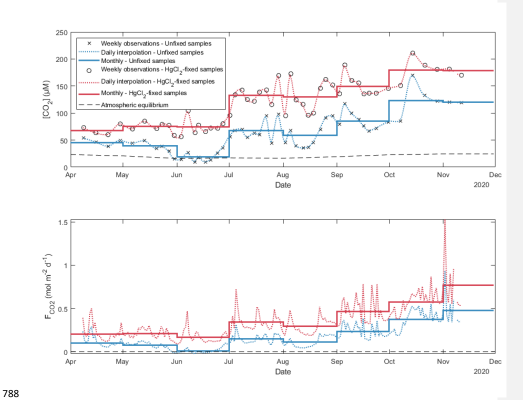


Figure 54. Daily and monthly surface CO₂ concentrations ([CO₂]; top panel) and diffusion fluxes (F_{CO2}; bottom panel) at the water-atmosphere interface from Lake Lundebyvannet (also in Tab. 3). Unfixed samples were obtained by DIC analysis. Daily [CO₂] was interpolated from weekly data using a modified spline (see text for details). Diffusion fluxes were calculated following Cole & Caraco (1998).

794	We further advise against interpretation of CO ₂ concentration data from low ionic strength, circum-
795	neutral water samples preserved with HgCl ₂ or CuCl ₂ . The overestimation of CO ₂ concentration
796	caused by HgCl ₂ can mask the effect of photosynthesis on lake carbon balance, creating the illusion
797	that lakes are net CO ₂ sources when they are net CO ₂ sinks. Our analysis from Lake Lundebyvannet
798	shows that HgCl ₂ fixation led to an overestimation of the CO ₂ concentration by a factor of 1.8, on
799	average, but approaching a factor of 4 during the peak photosynthetic period. An even larger impact is
800	expected on CO ₂ diffusive fluxes which were overestimated by a factor of 2 on average and up to a
	factor of >20 during peak photosynthesis. Interpreting such data would have underestimated the
801	
802	current and future role of aquatic photosynthesis.
803	Data availability
804	All data supporting this study will be made available on a permanent repository upon acceptance, e.g.,
805	Hydroshare.
806	Author contribution
000	Author Contribution
807	JET, AK, and TR supervised and PD, KN and FC contributed to the study design. JET, KN and TR
808	carried out the experiments. PD and TR performed the chemical analyses. JET and FC wrote the first
809	draft. FC performed the modelling, data, and statistical analyses, and drafted the figures. All co-
810	authors edited the manuscript.
811	Competing interests
812	The contact author has declared that none of the authors has any competing interests.
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