

1 **Temporary stratification promotes large greenhouse gas emissions in a shallow eutrophic lake**

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16 **Abstract**

17 Shallow lakes and ponds undergo frequent temporary thermal stratification. How this affects greenhouse
18 gas (GHG) emissions is moot, with both increased and reduced GHG emissions hypothesised. Here,
19 weekly estimation of GHG emissions, over growing season from May to September, were combined with
20 temperature and oxygen profiles of an 11 hectare temperate shallow lake to investigate how thermal
21 stratification shapes GHG emissions. There were three main stratification periods with profound anoxia
22 occurring in the bottom waters upon isolation from the atmosphere. Average diffusive emissions of
23 methane (CH₄) and nitrous oxide (N₂O) were larger and more variable in the stratified phase, whereas
24 carbon dioxide (CO₂) was on average lower, though these differences were not statistically significant. In
25 contrast, there was a significant, order of magnitude, increase in CH₄ ebullition in the stratified phase.
26 Furthermore, at the end of the period of stratification, there was a large efflux of CH₄ and CO₂ as the lake
27 mixed. Two relatively isolated turnover events were estimated to have released the majority of the CH₄
28 emitted between May and September. These results demonstrate how stratification patterns can shape
29 GHG emissions and highlight the role of turnover emissions and the need for high frequency
30 measurements of GHG emission which are required to accurately characterise emissions, particularly
31 from temporarily stratifying lakes.

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34 Keywords: Climate change; lake stratification; methane; carbon dioxide; nitrous oxide; climate
35 feedbacks

36

37 **1. Introduction**

38 Fresh waters are key sites for the processing of greenhouse gases (GHG), methane (CH₄), carbon dioxide
39 (CO₂) and nitrous oxide (N₂O). Shallow lakes, in particular, have been identified as hot spots of CH₄
40 release, particularly when ebullition is taken into account (Davidson et al., 2018; Aben et al., 2017). The
41 certainty that fresh waters are large emitters of GHGs contrasts with the uncertainties associated with the
42 quantities emitted and this is in large part due to historical paucity of measurements (Cole, 2013). A
43 recent study identified the highly variable emissions from lakes and ponds which contribute to more than
44 half the total emissions (Rosentreter et al., 2021). Whilst different morphometric features and chlorophyll-
45 a explained some of the emission patterns (Deemer and Holgerson, 2021), it is also clear that a dearth of
46 measurement combined with these highly variable emissions makes determining the drivers and controls
47 of those emissions a challenge, which in turn makes predicting future emissions difficult.

48

49 The current and future effects of climate change on lakes in general and on their GHG emissions are
50 relevant questions as there is potential for positive feedbacks and synergies with other human impacts
51 such as eutrophication (Davidson et al., 2018; Beaulieu et al., 2019; Delsontro et al., 2016; Meerhoff et
52 al., 2022). Taking a broad metabolic theory of ecology approach, temperature increases should promote
53 methanogenesis and shift the balance from primary production to respiration increasing CO₂ emission at
54 cellular and ecosystem scale (Yvon-Durocher et al., 2010). However, empirical and experimental data
55 indicate that temperature is not the sole control of primary production and methanogenesis. In particular,
56 eutrophication, and the promotion of large algal crop, has been associated with increased emissions of
57 CH₄ and N₂O (Delsontro et al., 2016) both by diffusion and ebullition (Zhou et al., 2019). Furthermore, in
58 what is globally the most abundant lake type, small shallow lakes, where macrophytes can colonise large
59 areas of the lake bed, trophic state and the dominance of submerged plants or algae may be more
60 important than temperature in shaping GHG dynamics (Davidson et al., 2015; Davidson et al., 2018;
61 Bastviken et al., 2023).

62

63 Climate change effects on lakes are not limited to increases in average temperatures and lengthening of
64 the growing season. Increases in both the frequency and intensity of heat waves are predicted, which will
65 promote the warming of surface waters and in turn make permanent and temporary thermal stratification
66 of lakes more likely (Woolway and Merchant, 2019), even in lakes typically classified as non-stratifying
67 (Kirillin and Shatwell, 2016). A recent study Holgerson et al. (2022) identified stratification and mixing
68 patterns in small water bodies, with permanent summer stratification common and frequent mixing
69 occurring in larger standing waters (>4 ha) lakes. Such periods of stratification and mixing events are
70 likely to have profound effects on GHG dynamics. Emissions of gases, in particular CH₄, that accumulate
71 in the isolated bottom waters of a stratified lake, occurs upon mixing and can make very significant
72 contributions to cumulative emissions (Schubert et al., 2012). High-resolution studies of sites that
73 undergo temporary stratification are, however, rare (Søndergaard et al., 2023). In terms of its effects on
74 GHG dynamics, there are potentially antagonistic processes at work in a stratified lake. On the one hand
75 the ‘shield effect’ results in lower temperatures at the sediment surface slowing down metabolic processes
76 that scale with temperature, i.e. methanogenesis and mineralization of organic carbon (C), reducing
77 emission and promoting C burial. On the other hand, anoxia at the sediment surface may shift processes
78 towards fermentation, increasing the proportion and total amount of CH₄ produced and perhaps reducing
79 C burial (Bartosiewicz et al., 2019). Recent work combining empirical observations and models has
80 suggested that shielding effects are larger than the anoxia effects and that stratification, in general,
81 increases C burial and reduces GHG emissions (Bartosiewicz et al., 2015). The stratification induced
82 isolation of bottom waters was reported to lead to reduced ebullition of CH₄ and a shift to diffusive
83 pathways. It might, however, be predicted that in shallow lakes stratification would lead to much larger
84 CH₄ release as anoxic conditions would limit CH₄ oxidation by CH₄ oxidizing bacteria (MOBs)
85 (Bastviken et al., 2008). There may also be other factors with the potential to increase GHG emission,
86 such as sediment organic content and lake trophic status (Delsontro et al., 2016), which may interact with
87 stratification patterns in shaping GHG emissions.

88

89 In this study, we used data from a shallow lake with high frequency measurements of temperature profiles
90 combined with weekly measurements of dissolved gas concentrations in the surface and bottom waters
91 and continuous measurement of ebullitive emissions of CH₄ to track the effects of lake stratification on
92 GHG emissions. The key question was how ebullitive and diffusive fluxes of the key GHGs: CH₄, CO₂
93 and N₂O respond to temporary thermal stratification.

94

95 **2. Materials and methods.**

96 **2.1 Study site**

97 Ormstrup lake, located in Denmark (lat 56.326°, lon 9.639°) (Fig.1) (depth map with GHG sampling
98 locations), is an 11 ha, shallow lake (average depth 3.4 m), with a maximum depth of 5.5 m, and with a
99 relatively long hydraulic retention time (> 1 year). The lake is eutrophic with high TP and chlorophyll-a
100 (Table 1; Søndergaard et al., 2022) with very sparse occurrence of submerged plants.

101

102 **2.2 Depth profiling and high frequency measurements**

103 In June 2020, a Nexsens (NexSens Technology, Fairborn, OH, USA) CB-450 data buoy system
104 (https://www.nexsens.com/pdf/CB450_datasheet.pdf) was deployed at the deepest point of the lake
105 equipped with a Nexsens TS210 thermistor string https://www.nexsens.com/pdf/TS210_datasheet.pdf
106 with temperature nodes measuring at 4 levels; one sensor “in air”, ca. 5 cm above the water surface, (but
107 shielded from direct light), and three sensors at -1, -2, -3 meters, respectively relative to the water surface.
108 In addition two Aqua TROLL 500 (In-Situ, Fort Collins, CO, USA) multi-sondes were mounted near the
109 surface (-1.0 meters) and at deeper water depth (-3.8 meters). The near surface and deeper water sonde
110 were configured with sensors to measure dissolved oxygen (DO) and water temperature (Tw). The optical
111 sensors were calibrated according to manufacture guidelines and checked on a weekly basis.

112

113 The optical sensors of the Aqua TROLL 500 have a built-in wiper mechanism to clean sensor heads to
114 hamper bio-fouling. The wiper function was enabled to perform cleaning in sync with sensor
115 measurements, hence every 15 minutes. In addition, manual cleaning of sensor heads was done every
116 week, while routine manual field monitoring was carried out at the lake. Prior to the deployment of the
117 buoy, and as a validation exercise for the buoy data, weekly manual profiles of DO and Tw were collected
118 at the deepest point.

119

120 Periods of stratification were defined by a greater than 2 °C difference between the surface and bottom
121 waters and DO below 0.5 mg l⁻¹ at the time of the weekly manual profiling of the system. The high
122 frequency measurements were used to confirm the patterns. During periods of defined as stratified, there
123 were partial mixing events where the depth of the thermocline and oxycline change and there was some
124 mixing of the bottom waters and surface waters. of the water column mixes, whilst the bottom waters
125 remained undisturbed as the DO did not increase the partial mixing above might increase gas exchange.

126

127 **2.3 Water chemistry**

128 Water samples for the analysis of Chlorophyll-a were collected weekly from the 20. April 2020 from
129 surface (-0.5 m) water at station 3 (Fig. (Søndergaard et al., 2005)1). A volume of water ranging from (0.2
130 to 1 litre) was filtered and the GFC papers preserved for chlorophyll-a analysis, which were determined
131 spectrophotometrically after ethanol extraction (Jespersen and Christoffersen, 1987) and alkalinity was
132 measured weekly by gran titration (Søndergaard et al., 2005). Depth profiles of temperature, electrical
133 conductivity (EC) and dissolved oxygen (DO) were measured manually with an Aqua TROLL 500 probe
134 from every -0.5 or -1 m down to -5 m depth).

135

136 **2.4 Greenhouse gas sampling**

137 2.4.1 Dissolved concentration

138 Samples of dissolved concentrations of CH₄, CO₂ and N₂O were collected weekly from the 20. April 2020
139 from surface waters and weekly from surface and bottom water from the 26. May 2020. The samples
140 were taken using head-space equilibration after (Mcauliffe, 1971), where 20 ml of water was collected
141 from just below the water surface and 20 ml of N₂ was introduced as a headspace in a 60-ml syringe and
142 then shaken vigorously for one minute. The 20 ml headspace was then transferred to a 12-ml pre
143 evacuated glass vial.

144

145 Gas concentrations in the headspace were determined on a dual-inlet Agilent 7890 GC system interfaced
146 with a CTC CombiPal autosampler (Agilent, Nærum, Denmark) (Petersen et al., 2012). For the GC,
147 certified CO₂, CH₄ and N₂O standards were used for calibration and validation. Aqueous concentrations in
148 N₂O, CH₄ and CO₂ were calculated from the headspace gas concentrations according to Henry's law and
149 using Henry's constant corrected for temperature and salinity (Weiss, 1974; Weiss and Price, 1980;
150 Wiesenburg and Guinasso, 1979). A recent study (Koschorreck et al., 2021) identified significant bias in
151 the estimate of CO₂ concentrations using headspace equilibration at lower concentrations. We applied their
152 correction using separately measured alkalinity as described in Koschorreck et al. (2021).

153 The fluxes of N₂O, CH₄ and CO₂ between the water and the overlying atmosphere were estimated as

154
$$f_g = k_g(C_{wat,g} - C_{eq,g})$$

155 Where f_g is the flux of a specific gas g , k_g is the piston velocity of the gas and $C_{wat,g} - C_{eq,g}$ is the
156 gradient of concentration between the concentration of gas dissolved in the water ($C_{wat,g}$) and the
157 concentration of gas the water would have at equilibrium with the atmosphere ($C_{eq,g}$).

158 We calculated a gas transfer velocity k_{600} for each sampling occasion using the relationship based on
159 windspeed described in (Cole and Caraco, 1998).

160
$$k_{600} = 2.07 + 0.215U_{10}^{1.7}$$

161 U_{10} is the mean daily windspeed at 10m (m s^{-1}) obtained from the Danish meteorological institute
162 (DMI;20x20 km grid data)

163

164

$$165 \quad k_g = k_{600} \left(\frac{Sc_g}{600} \right)^x$$

166

167 Sc_g is the Schmidt number(Wanninkhof, 1992) of the specific gas g . We chose $x = -2/3$ as this factor is
168 used for smooth liquid surface (Deacon, 1981).

169

170 Daily flux rates were calculated using linear interpolation of the weekly surface measurements from each
171 of the sampling points. The diffusive surface water fluxes were calculated by taking an average of the daily
172 flux rate from the 12. May 2020 to the 13. October 2020 for each location. Then an average of the 3 locations
173 was multiplied by the area of the lake and the number of days covered by the study, here 126 days was
174 chosen to match the period over which ebullition was measured.

175

176 The total content of the gases in the lake's bottom waters were calculated from the concentration of the
177 gases per litre multiplied by an estimate of the volume of the water in the hypolimnion. The volume of
178 water in the hypolimnion was estimated from the lake profiles manually conducted on the day of
179 sampling. The top of the hypolimnion was determined by the depth below which oxygen was less than 0.5
180 mg l^{-1} . A detailed bathymetry of the lake allows the calculation of the area and therefore volume of water
181 that lies below a given depth.

182

183 During the study period two major turnover events occurred, the process of lake turnover and full mixing
184 can take a number of days, and the outgassing even longer. The oxygen data, from the buoy, indicated
185 that it can take up to four days and this provides time for CH_4 oxidation to occur (Søndergaard et al.,

186 2023). In order to estimate the amount of CH₄ oxidised over the course of the multiple days of degassing
187 we directly measured CH₄ oxidation rates in the surface waters of the lake. This was done in June 2023 in
188 five locations using methods outlined in (Thottathil et al., 2019) where five water samples from five
189 different locations and each was incubated over 4 days with and the change in CH₄ concentration used to
190 calculate oxidation rates. We used the minimum (0.267 µg CH₄-C l⁻¹ h⁻¹), mean (0.44 µg CH₄-C l⁻¹ h⁻¹)
191 and maximum (0.58 µg CH₄-C l⁻¹ h⁻¹) oxidation rates to estimate the range of CH₄ oxidation likely to have
192 occurred over the course of the two main turnover events. Assuming that the degassing took four days,
193 these rates would consume between 2 and 8% of the CH₄ contained in the hypolimnion. Using the mean
194 oxidation value the turnover fluxes were reduced by 4.1% on the 30th of June 2020 and by 6% for the 25th
195 August 2022.

196 2.4.2 Ebullition

197 The ebullitive flux of CH₄ was estimated using at total of 40 floating chambers placed on 4 transects of 10
198 chambers each (Fig. 1). The chambers have a volume of 8 litre and a surface area of 0.075 m², similar to
199 those used by (Bastviken et al., 2015). As the existing literature indicated that ebullition is lower as water
200 depth increases (Wik et al., 2013) the transects were placed to maximise the measurement of the low end
201 of the depth gradient on the shallower slopes of the western end of the lake (Fig. 1). The average and
202 maximum depth of each transect was T1: 293 cm and 472 cm; T2: 181 cm and 267, T3: 223 cm and 300
203 cm and T4 166 cm and 220 cm. The chambers were set on the 14. May 2020 and sampled every two
204 weeks from that date, and on one occasion after one week until September 17th, which is a period of 127
205 days. Twenty ml of sample was taken from the floating chamber and injected into a pre-evacuated 12 ml
206 vial (exetainer, Labco). Gas concentrations were determined on the same GC than described above
207 (Petersen et al., 2012)

208 Ebullitive flux of CH₄ was estimated as:

$$209 \frac{p_{gas} \times Vol_{bub}}{t \times A}$$

210 Where p_{gas} is the concentration of CH₄ in the gas that was trapped, Vol_{bub} is the volume of the chamber
211 (i.e. 7L), t is the time during which the samples was collected and A is the area of chamber (i.e. 0.075 m²).
212 A portion of the CH₄ released via ebullition in the chamber will have re-dissolved in the water or might
213 leak through the chamber walls, thus underestimating the ebullitive flux. We have made a number of
214 measurements to constrain this error and to compare estimates based on static chambers with other
215 approaches. The result show that whilst static chambers underestimate ebullition, given the temporal
216 variability of ebullition, static chambers continually deployed provide a better estimate of average ebullition
217 than short term (24-48 hours) deployment using portable gas monitors or flushing chambers.

218
219 Therefore, whilst static chambers method cannot be said to accurately quantify CH₄ emissions, they can be
220 relied upon to compare differences in ebullition between time periods, with the caveat that they are always
221 an underestimate of actual ebullitive flux.

222
223 Total ebullitive flux from the lake was calculated by taking a mean of the emissions from each transect over
224 the 126 day period. Then taking an average of the means of four transects and multiplying this by the time
225 of deployment of the chambers in days, which was 126 days, and by the area of the lake. This gives a total
226 ebullitive flux of CH₄ for the lake over the period of measurement from May to mid September.

227
228 The three different flux types, surface diffusion, ebullition and turnover emission were then converted in
229 comparable units of total lakes emissions (as g or kg of gas) over the studied period and also converted into
230 CO₂-equivalents using a conversion factor related to their 100 year global warming potential (GWP) of 28
231 for CH₄ and 265 for N₂O.

232

233 **2.5 Statistical methods**

234 To test for a significant difference among the emissions from the stratified and mixed phase we used
235 generalised least squares (GLS) with a variance function to account for heterogeneity of variance between
236 the phases. In the case of the ebullitive flux, as the collected phase often covered periods including both
237 mixed and stratified phases there were three categories, mixed, stratified and both mixed and stratified.
238 All analysis was carried out in R version 4.2.1 (R Development Core Team, 2022) and the GLS used the
239 package nlme (Pinheiro et al., 2014).

240

241 **3.0 Results**

242 **3.1 Lake physical and chemical characteristics**

243 Depth profiles measured weekly from April show that stratification was initiated by the 26. May 2020 this
244 may have broken down briefly and established again, visible in the temperature sensors for the buoy on
245 the 5. June 2020 (Fig. 2). There were then 12 days of mixing followed by stable period of stratification
246 with onset the 14. June 2020 and a duration of 16 days until a mixing event around the 30. June 2020. The
247 following two weeks had cooler water and a mixed water column, hereafter a ca. 6 day period of
248 stratification from the 15. to 21. July 2020. A mixed phase of two weeks then followed until stratification
249 reestablished on 4. August 2020 and persisted until the end of August, partial mixing is indicated by the
250 buoy data from the 21. August 2020, but the weekly manual profile to deeper water indicate that full
251 mixing did not occur until after the 25. August 2020. The effects of the stratification and mixing events on
252 the high frequency DO data measured at -3.8 m are clear, with rapid deoxygenation occurring after the
253 onset of stratification and oxic bottom waters returning when the lake mixed (Fig. 2). The pattern in
254 chlorophyll-a also follow, to some degree, those of stratification, with the exception of early spring.
255 Chlorophyll-a values were extremely high in spring peaking at the start of June 2020 and falling gradually
256 (Fig. 2). (Søndergaard et al., 2023) During the periods of stratification chlorophyll biomass was lower, and

257 when a mixing event occurred the values increased, which is particularly evident in the July mixing
258 periods (Fig. 2).

259

260 **3.2 Concentrations of dissolved gases and fluxes from the surface waters.**

261 The concentrations of the dissolved gases showed great variation from near or below atmospheric
262 concentrations in some cases and up to an extremely high concentration (over 5 mg CH₄ C l⁻¹) in the
263 bottom waters on the 30. June 2020. There was some spatial heterogeneity in the surface waters, with the
264 more littoral locations showing the greatest variation and the highest values (Figs. 3,4,5). In particular the
265 most littoral zone, where the water was shallower around 1 m in depth, showed the highest values just
266 prior to, or coincident with, the stratification turnover. Table 2 shows the mean diffusive flux of each gas
267 over the sampling period along with the mean flux in mixed and stratified phases. For CO₂ there was a lot
268 of temporal variation in flux dynamics, though not a large difference between mixed and stratified phases
269 in terms of mean values (Table 2). There were some periods of CO₂ influx in spring and later summer and
270 these tended to coincide with the end of a mixed phase and the start of the stratification phase. Nitrous
271 oxide concentrations were generally low (Figs 4 & 5) with the lake being a source of N₂O in the spring
272 period and a sink or a very small source thereafter. The CH₄ concentration in the surface waters (Fig. 3)
273 and the calculated diffusive emissions are relatively low, but did increase in the stratification periods with
274 higher average values (Table 2 & Fig. 6). There was also some spatial variation with higher CO₂ and CH₄
275 diffusive emissions in the shallower sampling locations, both in stratified and mixed conditions (Fig. 6).

276

277 The most marked patterns in GHG concentration were evident in the bottom waters sampled at -4.5 m,
278 which accumulated to very large concentrations of CO₂ but particularly CH₄ in the periods of
279 stratification (Fig. 3 & 4). The ratio of CO₂ to CH₄ is illustrative in highlighting how stratification has
280 altered the biogeochemical processes in the hypolimnion with CH₄ production becoming more prevalent. .

281 For example on 30. June 2020 after 16 days of stratification the the ratio $\text{CO}_2:\text{CH}_4$ in the bottom waters
282 was 0.8, whereas 7 days later after the mixing event it was 187 at the same depth.

283

284 **3.3 Ebullitive fluxes**

285 The CH_4 bubble flux, presented here as mean values for each of the 4 transects, ranged from 0.303 to 81.1
286 $\text{mg CH}_4 \text{ C m}^2 \text{ d}^{-1}$ for the individual transect over the growing season measurement. There is a very clear,
287 statistically significant impact of stratification on the ebullitive efflux of CH_4 with stratified periods
288 showing significantly markedly higher levels of emission (Fig. 7 and Table 2). In addition, there was a
289 difference in average emissions among the different transects, with those with lower average water depth
290 (T2 & T4) having lower emission than the transects with chambers over deeper water (T1 & T3) (Fig. 7).

291 The samples collected from the chambers reflect two weeks of bubble and diffusion collection and the
292 quantification of the flux is therefore an average of the period of chamber deployment, which was two
293 weeks, or in one case a single week (Fig. 7). This two week period on occasion covered both stratified
294 and mixed phases and on these occasions efflux was intermediate between purely mixed and stratified
295 periods (Table 2 and Fig. 7).

296

297 **3.4 Total lake fluxes**

298 Scaling up the results to total flux of gases from the whole lake over the period of study and including the
299 estimated emissions from two turnover events show a very different effect of stratification on the balance
300 of types of emissions for the three gases. The majority of CH_4 emission (56%) result from the two short-
301 lived turnover events (Fig. 8), whereas their contribution to CO_2 and N_2O emission was 5% and 1%
302 respectively.

303

304 Fluxes of CO_2 and N_2O were mostly diffusive, which represented 95% of emissions of both gases.

305 Methane diffusive flux was 14% of total emission, whereas CH_4 ebullition was more than twice as much at

306 29% of total CH₄ emission. In terms of global warming potential CO₂ and CH₄ emission were
307 comparable, but the contribution of the turnover efflux was the dominant factor for CH₄ emissions.

308

309 **4. Discussion**

310 The emission of the three GHGs showed different degrees of variation between the mixed and stratified
311 phases. The largest and most significant variation was in CH₄ ebullition (Table 2), whilst the difference in
312 diffusive fluxes, though marked for CH₄ was not significant. The mean of the total emissions from
313 Ormstrup in the stratified phase (59.9 mg CH₄-C m⁻² day⁻¹) corresponds relatively closely to the mean of
314 the total emissions (ebullition plus diffusion) reported for lakes in this size range (47 mg CH₄-C m⁻² day⁻¹)
315 from a paper synthesising multiple studies (Rosentreter et al., 2021). The mean emissions for the whole
316 period (26.6 CH₄ -C m⁻² day⁻¹) were lower than Rosentreter et al. (2021) but similar to other studies with
317 mean emissions of 30.9, 20.7 and 22.7 CH₄ -C m⁻² day⁻¹ and were reported by Peacock et al. (2021), Sørensen
318 et al. (2023) and Peacock et al. (2019) respectively. Whereas the average CO₂ (504 mg CO₂-C m⁻² day⁻¹) at
319 Ormstrup was lower than 993.5 mg CO₂-C m⁻² day⁻¹ measured by Peacock et al. (2021) but higher than
320 the 264.6 and 205.1 mg CO₂-C m⁻² day⁻¹ measured by Sørensen et al. (2023) and Peacock et al. (2019)
321 respectively. The different temporal resolution and duration of these studies, eleven single day sampling
322 from April to December (Peacock et al., 2021), five days continuous sampling on one occasion in late
323 September (Sørensen et al., 2023) and a single early summer snapshot (Peacock et al., 2019) make direct
324 comparison difficult. The data here do, however, provide a clear answer to the question of how thermal
325 stratification affects GHG dynamics in shallow eutrophic lakes with an increase in total emissions
326 (diffusion, ebullition and turnover) during the stratified period (Table 2, Fig 9). Previous work, combining
327 observations and modelling suggested the opposite patterns (Bartosiewicz et al., 2019) as the shielding
328 effect of the stratification results in cooler bottom waters which reduces CH₄ production due to the
329 process being temperature dependent (Bartosiewicz et al., 2016). This strong shielding effect may apply
330 in deeper lakes experiencing more stable stratification, or less eutrophic lakes. The result here from a

331 relatively shallow eutrophic lake, indicate that temporary stratification causes increases in GHG
332 emissions.

333

334 Diffusive emissions did not, on average, show a strong stratification effect (Table 2). In particular
335 variation in N₂O emissions did not match patterns of stratification, with emissions more directly related to
336 nitrate concentrations (Audet et al., 2020), as reflected by the fact the lake is a sink of N₂O in late summer
337 when nitrate was below detection limits for several weeks. There were peaks in emission of CH₄ and CO₂
338 at the end of stratification periods, particularly in the shallower water sampling points (Fig. 6). There
339 were periods of influx of CO₂, which coincided somewhat with periods of stratification, but the pattern
340 was not consistent as other factors, for example, chlorophyll-a concentration also play a role.

341

342 Littoral zones can have markedly different GHG dynamics to deeper zones due to shallower water having
343 lower pressure (Wik et al., 2013), less time for CH₄ oxidation (Bastviken et al., 2008) or abundant plants
344 which influence a range of biogeochemical processes (Davidson et al., 2018; Esposito et al., 2023). It is
345 therefore possible that littoral zone dynamics could cause these differences. However, the increase
346 occurred at all three sampling points at the end of June 2020, which indicates a lake-wide driver and the
347 peak may represent the start of mixing after stratification. Strong winds were measured on the 29th and
348 30th June 2020 (Søndergaard et al., 2023) coincident with these increased littoral emissions. These winds
349 would have caused lateral movement of the surface water causing an upwelling of bottom water, rich in
350 CH₄ and CO₂, in the littoral margins at the opposite end of the lake. Thus, whilst we do not have direct
351 evidence it seems more likely that these increased emissions in the littoral zone were driven, at least in
352 part, by the upwelling of GHG rich bottom waters.

353

354 In contrast to the diffusive flux, the ebullitive emission of CH₄ shows a very clear response to
355 stratification with an order of magnitude difference in emissions between periods where the sampling
356 reflected purely mixed or stratified periods (Table 2 & Fig. 7). The two-week resolution of the sampling

357 meant that some samples covered both stratified and mixed phases and these samples had intermediate
358 fluxes, as they cover both low (mixed) and high emission (stratified) periods. The spatial variation in
359 ebullition is also illustrative of the impacts of stratification and the role of anoxia in shaping CH₄ fluxes.
360 The two transects with the largest mean and maximum depths (T1 and T3) had the largest emissions, with
361 the deeper of the two (T1) having the highest emissions and they saw the greatest relative increase during
362 the stratification phases. This pattern is different to that found in other studies where bubble emissions
363 were larger in shallower water, although later in the summer there was an increase in bubble flux (Wik et
364 al., 2013). The deeper water at Ormstrup experienced anoxia earlier and this appears to cause locations
365 with deeper water to have higher ebullition rates than shallower areas. This is at odds with ideas
366 stemming from the metabolic theory of ecology stating that temperature (Yvon-Durocher et al., 2014) in
367 particular at the sediment surface (Bartosiewicz et al., 2019) can be used to predict CH₄ efflux. Whilst it
368 is a fact that CH₄ production is temperature dependent at the cellular level, CH₄ emissions were rather
369 independent of the sediment temperature, for example in the first two weeks of July 2020 emissions were
370 low and the sediment surface temperature was relatively high. Thus, temperature alone is a poor predictor
371 of ecosystem scale CH₄ emissions.

372
373 It should be noted that the methods used to estimate bubble flux here, where floating chambers are
374 sampled every two weeks is a “less than perfect method”, which in nearly all cases will underestimate
375 ebullitive flux. Logistical and financial constraints make continual sampling difficult and here we
376 balanced these constraints against the greater time required to apply more accurate methods, such as
377 bubble traps (Wik et al., 2013), automatic flushing chambers (Bastviken et al., 2015). Such is the
378 variability of bubble flux in space and time that sampling campaigns covering days or weeks would
379 potentially give an even more inaccurate picture of emissions than the method used here (see
380 supplementary methods and supplementary materials). Though eddy covariance approaches offer a means
381 of continuous measurement capable of capturing short term changes and covering a large area (Erkkilä et
382 al., 2018). Thus, the continuous monitoring of ebullition using chambers with known biases was deemed

383 the least worst method available, but we acknowledge the caveat that ebullitive emissions may be
384 underestimated.

385
386 In addition to the diffusive and ebullitive emissions, the turnover flux, which consists of the gases
387 accumulated in the hypolimnion being released on turnover, was also estimated, with a correction of CH₄
388 oxidation applied. There were two major turnover events at the end of June and in late in August 2020,
389 which were preceded by 16 and 22 days of stratification, respectively. It was not possible to directly-
390 measure turnover flux, as they are relatively discrete events where the efflux likely occurs over the course
391 of (Søndergaard et al., 2023) a few hours, or a few days. Thus, the efflux estimation is based on a series of
392 assumptions and thus must be treated with caution. Notwithstanding this uncertainty, we can be confident
393 the turnover flux represents a very large proportion of the total emission of CH₄ emissions from Ormstrup
394 Lake over the growing season. We estimate it contributed more than 50 % of growing season CH₄
395 emissions and 5 % of CO₂ emissions. This highlights a very significant, and difficult to measure,
396 contribution to GHG emissions from lakes undergoing temporary stratification, which are among the
397 most common lake type in Denmark {Søndergaard, 2023 #8812} .

398
399 The results here suggest that GHG dynamics were driven both directly and indirectly by the stratification
400 patterns and the anoxia it induced in the bottom waters. At Ormstrup Lake the thermal stratification of the
401 water column quickly led to anoxia, with only a matter of hours to days for the oxygen to be consumed
402 once the bottom waters were isolated (Fig. 2). The ratios of CO₂:CH₄ evidence how this promotes CH₄
403 over CO₂ production in the stratification phase (see Fig 9). In addition to promoting CH₄ production such
404 conditions would preclude, or severely limit, oxic CH₄ oxidation, which has the potential to consume a
405 large proportion of CH₄ produced in the anoxic sediments (Bastviken et al., 2008), though anoxic CH₄ can
406 still occur (Blees et al., 2014). The raw emission data do not provide any direct information on the
407 balance of production versus oxidation, but the CO₂:CH₄ suggest there was marked shift to conditions
408 where methanogenesis was the dominant process and there was reduced CO₂ production. Studies have

409 shown that CH₄ oxidation can consume large proportions of the CH₄ produced under hypoxia (Saarela et
410 al., 2019) and it is possible that there is intense CH₄ oxidation occurring at the thermocline during the
411 periods of stratification at Ormstrup lake , but this was not directly measured at the lake. In addition to the
412 more direct effects of anoxia there may be some indirect effects of the patterns of stratification and
413 mixing that promote greater GHG emissions. Søndergaard et al. (2023) recently reported how nutrient
414 dynamics at Ormstrup Lake were altered by the lake stratification and full details can be found there, of
415 relevance here is the impact on chlorophyll-a which saw a large spring peak after which the abundance
416 tracked the stratification and mixing regime, with a lag time. There was a general reduction, or at least no
417 increase as the stratification period progressed, perhaps due to nutrient limitation in the epilimnion. Upon
418 mixing there was generally an increase in chlorophyll-a, though the weekly sampling resolution makes
419 this difficult to assess. Chlorophyll-a and the labile dissolved organic carbon (DOC) that result from
420 abundant chlorophyll-a have been shown to be associated with higher diffusive and ebullitive CH₄
421 emissions (Davidson et al., 2015; Beaulieu et al., 2019; West et al., 2012; Zhou et al., 2019). It is not
422 possible to say here whether a stable summer long stratification would have led to decreased chlorophyll-
423 a as nutrients became limiting due to their isolation in the bottom waters and reliable high frequency
424 chlorophyll-a data are required to convincingly demonstrate this phenomenon. Notwithstanding these
425 uncertainties it may be the case that the temporary stratification, interspersed with mixing events,
426 observed here represents a ‘sweet spot’ providing both the resources, i.e. chlorophyll-a and the labile
427 DOC it produces, and optimal conditions (anoxia) for CH₄ production.

428
429 Predicting climate change effects on GHG emissions in a future warmer world is not straightforward, as
430 there are multiple interacting drivers which combine to shape the GHG emissions of lakes. However, this
431 study suggests that temporary stratification, which is increasingly recognised as prevalent in ponds and
432 shallow lakes (Holgerson et al., 2022) and is likely to become more common with continued climate
433 change impacts (Woolway and Merchant, 2019) is likely to increase GHG emissions. This will be

434 particularly the case in more eutrophic systems where abundance algal derived dissolved organic matter
435 can fuel CH₄ production (Zhou et al., 2019).

436
437 The combination of high frequency data on water temperature and dissolved oxygen combined with
438 weekly measurements of GHGs increase the reliability of the findings presented here. Up until relatively
439 recently it has been assumed that for shallow lakes, such as Ormstrup lake, stratification is not an
440 important feature. Sampling has therefore focused on the surface layers of water bodies, using dissolved
441 concentrations of gases or floating chambers to characterise flux, e.g. (Davidson et al., 2015; Audet et al.,
442 2020; Peacock et al., 2021). Thus, most studies have overlooked bottom waters and do not have the
443 temporal resolution required to capture turnover flux emissions from surface measurements. Furthermore,
444 whilst many studies now include estimates of bubble emissions of CH₄ e.g. (Bergen et al., 2019), the
445 necessary temporal resolution to accurately characterise ebullitive emission is not well established. The
446 finding here indicated that in such dynamic systems near continuous measurement is desirable and that
447 short term collection over one or two days could provide massive, over or under estimate of CH₄
448 ebullition.

449
450 Our results show very large temporal variation in emissions of all three gases, but in particular CO₂ and
451 CH₄, and this highlights the need for high frequency measurements to accurately characterize emissions
452 from lakes. Even the weekly frequency of the sampling in this study was not sufficient to directly measure
453 all the emission pathways and turnover flux had to be inferred from bottom water calculations. These data
454 show that to capture the extent of GHG emissions from lakes it is vital we include all forms of flux,
455 including ebullition and turnover flux. Recent work has highlighted the fact that most emissions of CH₄
456 (over 50%) from fresh waters come from highly variable systems (Rosentreter et al., 2021), with the mean
457 and median emission rates of CH₄ differing greatly, indicating a few large emitters are responsible for a
458 large proportion of emissions. The sampling frequency applied here is rare, if a more standard resolution
459 of monthly measurements was applied the emissions estimate of all the gases, but in particular CH₄,

460 would be highly dependent on what phase of the stratification was captured. As an example, a monthly
461 sampling frequency could potentially miss all the stratification peaks - consequently massively
462 underestimating emissions, whereas a different sampling frequency could catch a number of peaks and
463 give a much higher estimate. Thus, the same sampling frequency on the same lake, but timed differently
464 could lead to conclusions of highly variable emissions. Consequently, in these highly dynamic systems
465 where temporary stratification occur in summer, high frequency measurements are required to accurately
466 estimate emissions. This is logistically challenging but the current advances in the use of automatic
467 flushing chambers (Bastviken et al., 2020) may provide the potential for affordable high spatial and
468 temporal resolution measurement of GHG dynamics. This is a requisite for understanding the drivers of
469 GHG dynamic, which is required for being able to predict how they will respond in a range of scenarios
470 related to land use, climate change and management interventions.

471

472 **Code/data availability**

473 The datasets generated during and/or analysed during the current study are not publicly available as they
474 form part of ongoing research projects but are available from the corresponding author on reasonable
475 request and will be made publicly available later in the research project.

476

477 **Author contributions**

478 MS secured the funding for the wider lake restoration research project supplying the data. TAD, MS and
479 JA conceptualized the gas study. TAD and AN established the buoy and sensor system. EL, CE, TAD,
480 TB and JA collected and analysed the data. TAD wrote the paper and all authors commented on earlier
481 versions and read and approved the final draft.

482 **Competing interests**

483 The authors declare that they have no conflicts of interest.

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489 Foundation for providing great support for this work and the Ormstrup project generally. TAD and CE
490 were also supported by GREENLAKES (No. 9040-00195B) and The European Union’s Horizon 2020
491 research and innovation programmes under grant agreement No 869296—The PONDERFUL Project.

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496

497 Table 1. Summary lake information, summer mean values and (standard deviation) of a range of
498 variables

Variable	n	Year 2020
Secchi depth (m)	22	0.86 (0.28)
Chlorophyll a ($\mu\text{g/l}$)	20	53.4 (28.9)
pH	22	8.04 (0.77)
Total phosphorus (mg/l)	22	0.58 (0.11)
Total nitrogen (mg/l)	22	1.50 (0.41)

499
500 Table 2. Mean greenhouse gas flux (units CO_2 : $\text{mg CO}_2\text{-C m}^{-2} \text{ day}^{-1}$, N_2O : $\text{mg N}_2\text{O -N m}^{-2} \text{ day}^{-1}$, CH_4 both
501 diffusive and ebullitive in $\text{mg CH}_4\text{-C m}^{-2} \text{ day}^{-1}$) from the lake from spring to Autumn 2020. The emissions
502 are divided in diffusive, ebullitive emissions. The mean values for all the surface water stations and all
503 four transects of chambers are given. Emissions area separated into mixed versus stratified phases and
504 there SD are also given. Ebullition was collected for a period covering two weeks so on a number of
505 occasion covered both mixed and stratified periods thus ebullition has a third category where both
506 mixed and stratified conditions occurred is given. Ebullition was significantly different across the three
507 phases, diffusive fluxes were not significantly different for p values of 0.05.

508

509

510

Emission type	gas	mean	mixed	Stratified	Strat and mixed
Diffusive	CO ₂	493.7 <i>(529.6)</i>	559.6 <i>(433.1)</i>	449.8 <i>(587.6)</i>	
	CH ₄	9.47 <i>(16.0)</i>	5.9 <i>(4.1)</i>	12.7 <i>(20.2)</i>	
	N ₂ O	0.11 <i>(0.09)</i>	0.09 <i>(0.08)</i>	0.12 <i>(0.11)</i>	
Ebullition	CH ₄	17.28 <i>(19.62)</i>	4.84 <i>(3.44)</i>	47.29 <i>(21.95)</i>	12.74 <i>(10.34)</i>

511

512

513 Figure legends

514 Figure 1. Ormstrup lake bathymetry and sampling stations for surface water greenhouse gas sampling
515 (St1, St2, St3) bottom waters were sampled at S3. Transects of 10 bubble traps were placed on T1- T4.

516 Adapted from the Søndergaard et al. 2023.

517 Figure 2 Temperature profile from June 2020 when the buoy was deployed and surface and bottom water
518 oxygen from June to the end of September 2020. Manual chlorophyll-a ($\mu\text{g L}^{-1}$) values are also given in
519 the top panel.

520 Figure 3. Dissolved CH_4 concentrations from surface and bottom waters – thermal stratification periods
521 highlighted in grey and the white background indicate mixed waters

522 Figure 4 . Dissolved CO_2 concentrations from surface and bottom waters–thermal stratification periods
523 highlighted in grey and the white background indicate mixed waters

524 Figure 5 Dissolved N_2O gas concentrations surface and bottom thermal stratification periods highlighted
525 in grey and the white background indicate mixed waters

526 Figure 6. Ormstrup lake surface fluxes of the CH_4 , CO_2 and N_2O gases based on dissolved concentration ,
527 thermal stratification periods highlighted in grey and the white background indicate mixed waters

528 Figure 7. Plot of CH_4 ebullition averaged for each transect (10 chambers per transect), data collected from
529 40 traps every two weeks. Thermal stratification periods highlighted in grey and the white background
530 indicate mixed waters.

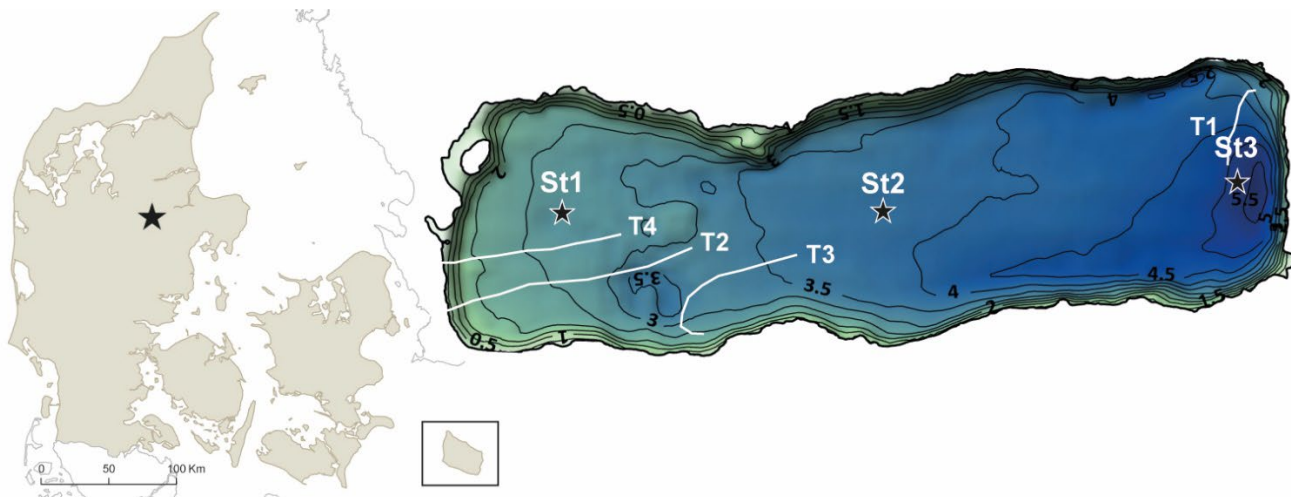
531 Figure 8 – Total lake emissions per gas over the growing season in CO_2 equivalents. The emissions are
532 divided different emission modes: Diffusive, ebullitive and turnover flux. All estimates contain some
533 uncertainty, in particular ebullitive flux is an underestimate and the turnover flux also contains a great deal
534 of uncertainty.

535 Figure 9. Summary of the quantities of the gases present in the water and the volumes emitted from the
536 different pathways. The size of the arrow is proportional to the emissions from each pathway and with the
537 stratified state on the left and the mixed state on the right, with the turnover flux in the centre.

538

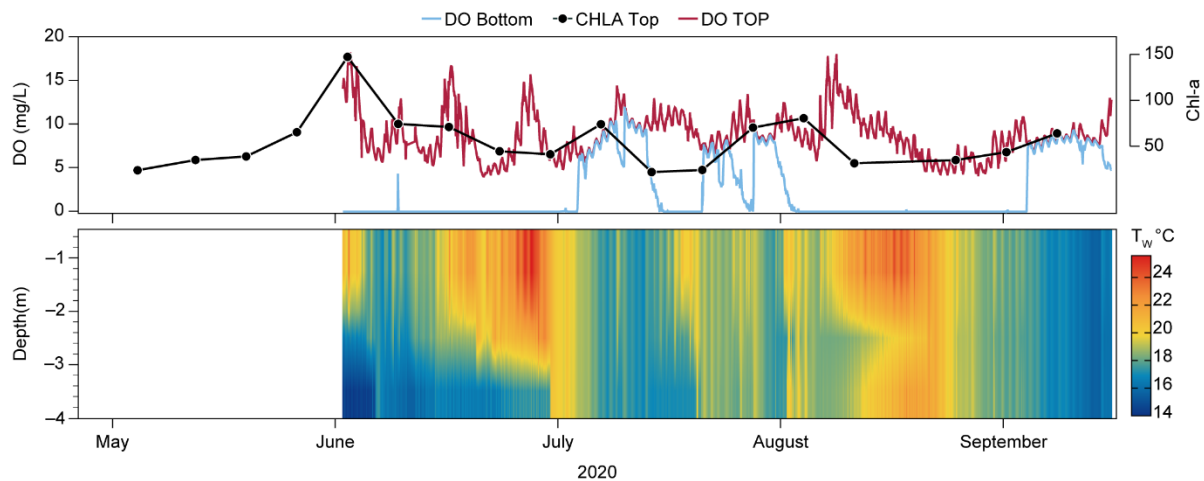
539 Figures and legends

540 Figure 1.



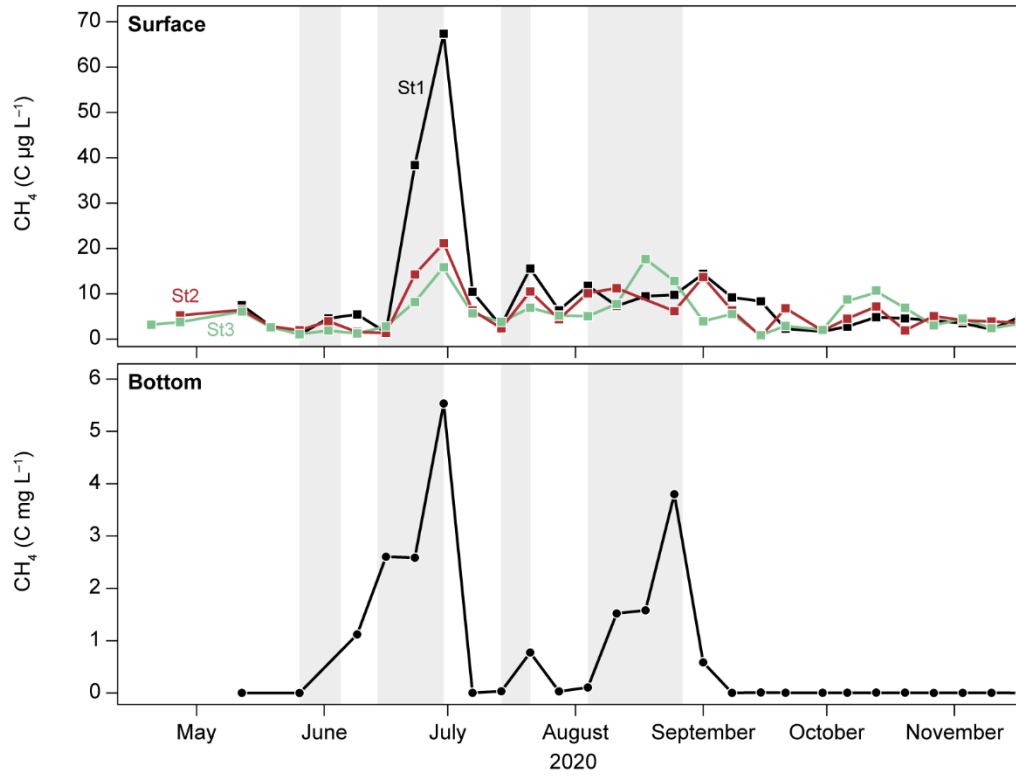
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543 S2, S3) bottom waters were sampled at S3. Transects of 10 bubble traps were placed on T1- T4. Adapted
544 from the Søndergaard et al. 2023.

545



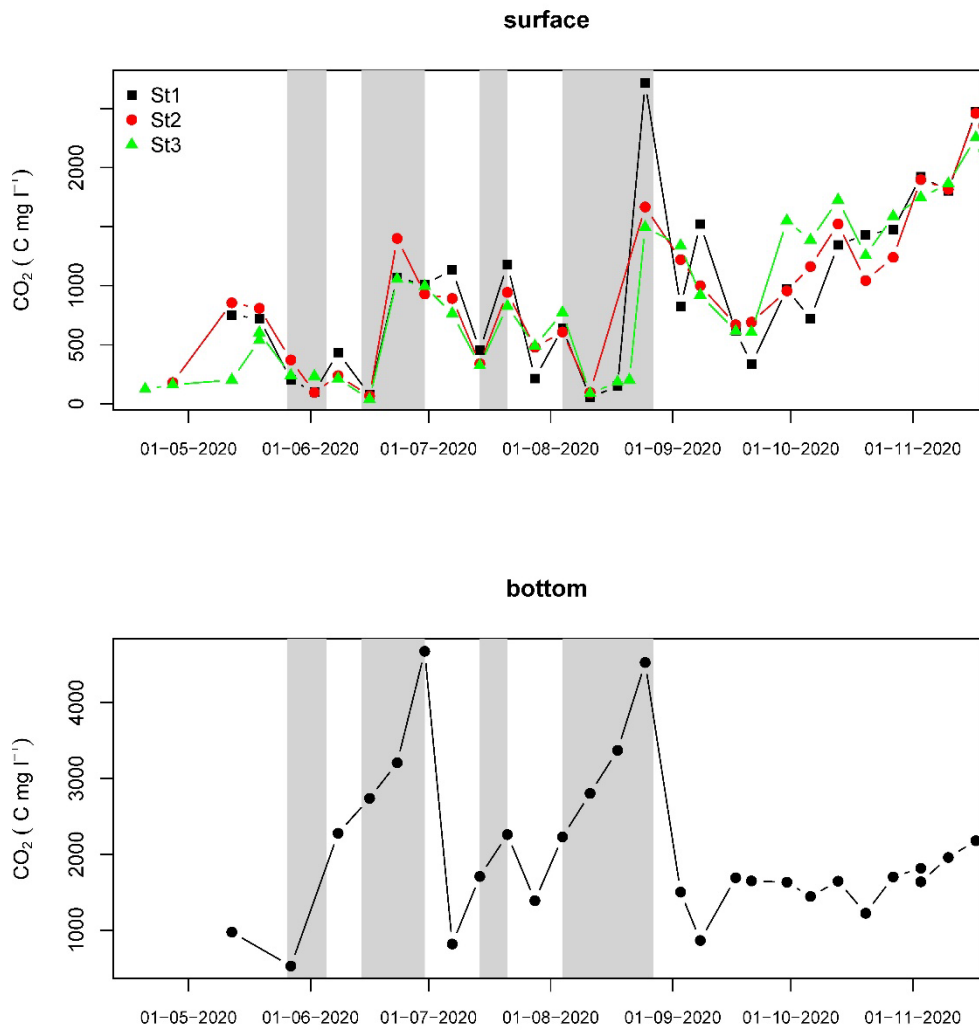
546
547 Figure 2 Temperature profile from June when the buoy was deployed and surface and bottom water
548 oxygen from June to the end of September. Chlorophyll-a ($\mu\text{g L}^{-1}$) values are also given in the top panel
549 and surface (DO TOP) and bottom (DO Bottom) dissolved oxygen (mg L^{-1}) are also given

550



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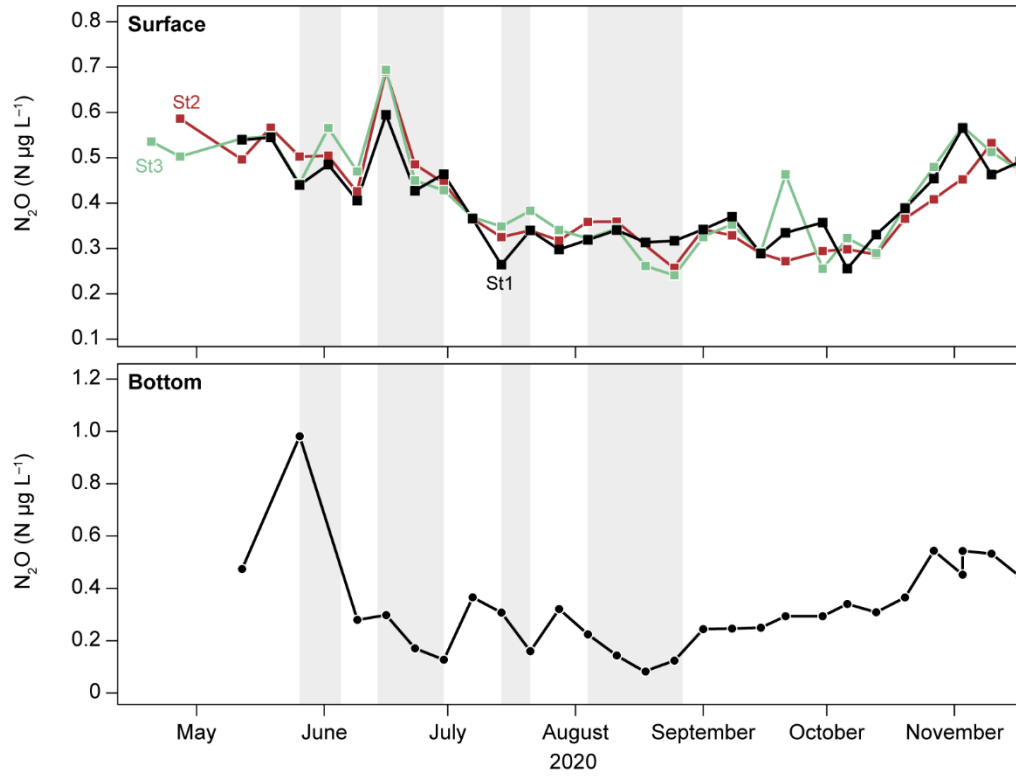
552 Figure 3. Dissolved CH₄ concentrations from surface and bottom waters – thermal stratification periods
 553 highlighted in grey; white background indicate mixed waters. Note different y axis scales



554

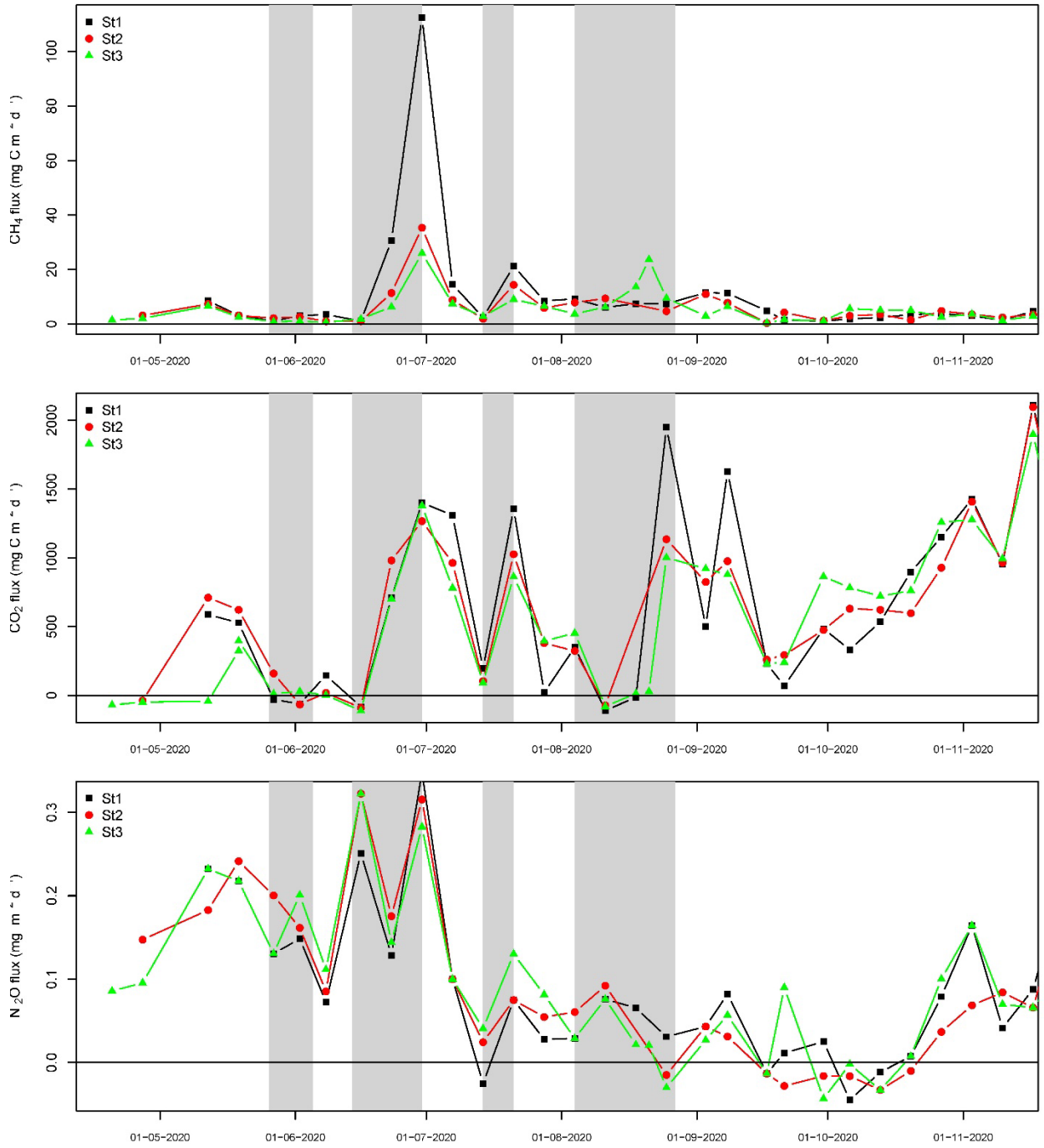
555 Figure 4 . Dissolved CO₂ concentrations from surface and bottom waters–

556 thermal stratification periods highlighted in grey; white background indicate mixed waters



557

558 Figure 5 Dissolved N_2O gas concentrations surface and bottom thermal stratification periods highlighted
 559 in grey; white background indicate mixed waters

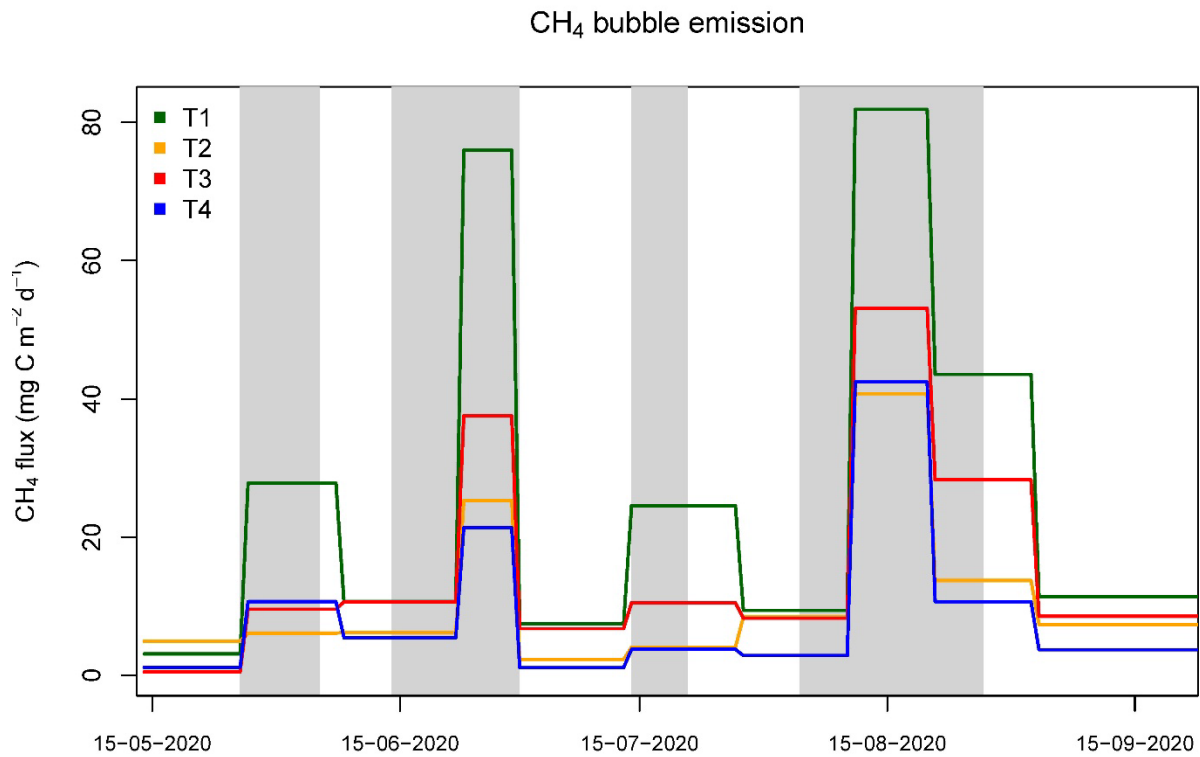


560

561 Figure 6. Omstrup lakesurface fluxes of the CH₄, CO₂ and N₂O gases based on dissolved concentration ,

562 thermal stratification periods highlighted in grey; white background indicate mixed waters

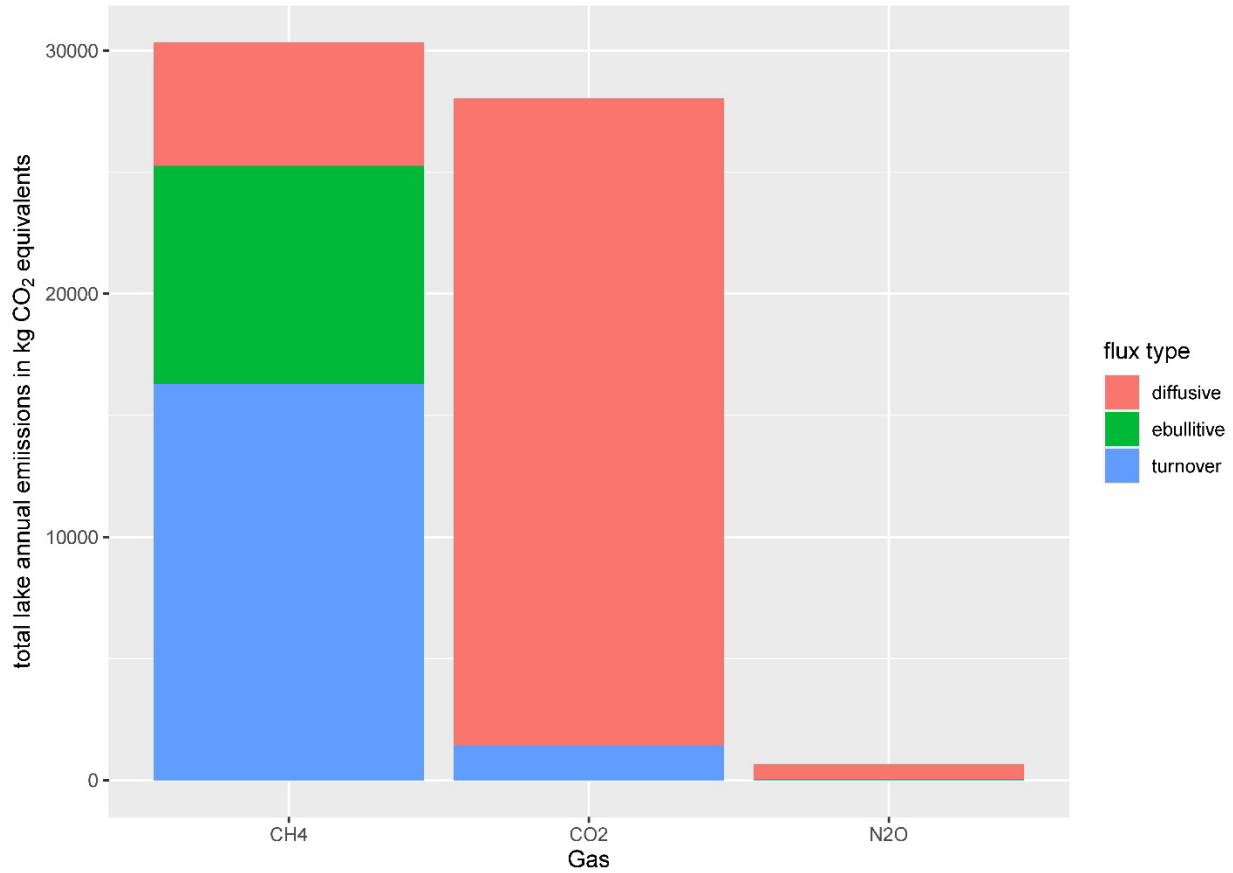
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566 Figure 7. Plot of CH₄ ebullition averaged for each transect (10 chambers per transect), data collected from
567 40 traps every two weeks. thermal stratification periods highlighted in grey; white background indicate
568 mixed waters.

569

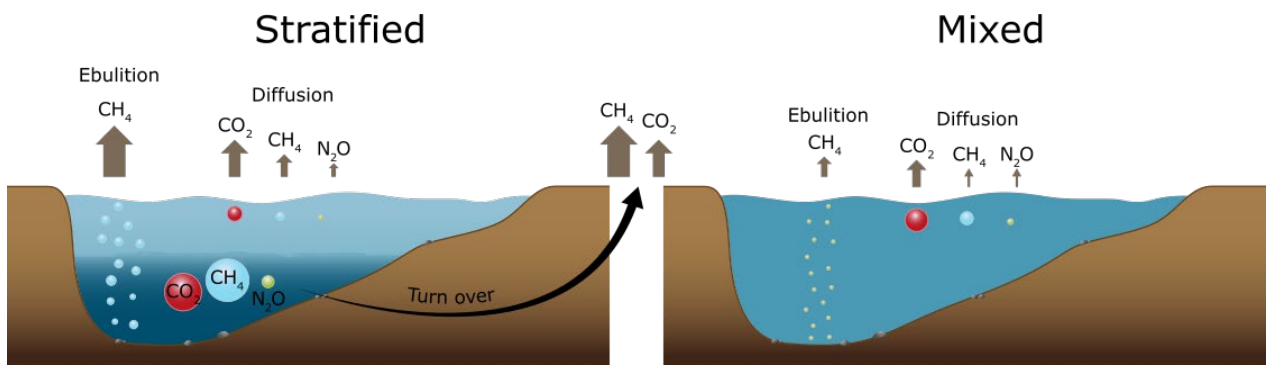


570

571 Figure 8 – Total lake emissions per gas over the growing season in CO₂ equivalents. The emissions are
 572 divided different emission pathways: Diffusive, ebullitive and turnover flux.

573

574



575

576 Figure 9 Summary of different flux types (bubble, diffusive and turnover) for the main greenhouse gases
 577 (CH₄ CO₂ and N₂O) observed between the stratified and mixed phases at Ormstrup lake patterns in the
 578 stratified and mixed phase. The turnover flux of CH₄ and CO₂ is also represented. The size of the arrow

579 represents the relative amount of emission and the size of the circle in the lake represents the
580 concentration of dissolved gases in stratified or mixed water column.

581

582

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