# Spatial and temporal variability of methane emissions and environmental conditions in a hyper-eutrophic fishpond

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10 Abstract. Estimations of methane  $(CH_4)$  emissions are often based on point measurements using either flux 11 chambers or a transfer coefficient method which may lead to strong underestimation of the total CH<sub>4</sub> fluxes. In 12 order to demonstrate more precise measurements of the CH<sub>4</sub> fluxes from an aquaculture pond, using higher 13 resolution sampling approach we examined the spatiotemporal variability of CH<sub>4</sub> concentration in the water, 14 related fluxes (diffusive and ebullitive) and relevant environmental conditions (temperature, oxygen, chlorophyll-15 a) during three diurnal campaigns in a hyper-eutrophic fishpond. Our data show remarkable variance spanning 16 several orders of magnitude while diffusive fluxes accounted for only a minor fraction of total CH<sub>4</sub> fluxes (4.1– 17 18.5 %). Linear mixed-effects models identified water depth as the only significant predictor of CH<sub>4</sub> fluxes. Our 18 findings necessitate complex sampling strategies involving temporal and spatial variability for reliable estimates 19 of the role of fishponds in a global methane budget. 20

21 Keywords: aquaculture, emissions, fishpond, freshwater, heterogeneity, methane

#### 23 1 Introduction

24 Freshwater aquaculture ponds (fishponds) represent man-made counterparts to natural shallow lakes (Scheffer, 25 2004) which are mainly used for fish production (mostly of common carp, Cyprinus carpio L.) and water retention 26 in the landscape. Fishponds serve also as secondary biotope for various organisms (Kolar et al., 2021), supporting 27 noteworthy animal and plant diversity (Pokorný and Hauser, 2002). However, most fishponds suffer from high 28 fish stock densities, excessive carbon and nutrient loading from supplemental fish feeding, sewage pollution, and 29 fertiliser runoffs from agricultural catchments or nutrient mobilisation from the anoxic sediment layers (Pechar, 30 2000). As a result, the trophic structure of plankton communities has shifted towards a reduction of large 31 zooplankton and massive development of phytoplankton, especially cyanobacterial blooms (Potužák et al., 2007), 32 limiting light penetration in the water column. Rapid changes in the intensity of biological processes such as 33 photosynthesis and respiration often result in pronounced daily or seasonal fluctuations in dissolved oxygen (Baxa 34 et al., 2021), signalling decreasing ecosystem stability. The extent of anoxia, accumulation of organic biomass, 35 and rapid heating of the shallow water during summer result in enhanced production of greenhouse gases (Grasset 36 et al., 2018, Zhang et al., 2021; Bartosiewicz et al., 2021).

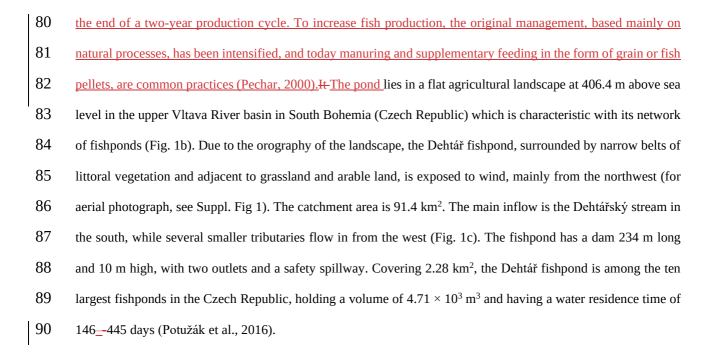
37 Most concerning are CH<sub>4</sub> emissions as freshwater aquaculture systems release more than 6 Tg CH<sub>4</sub> yr<sup>-1</sup> (Yuan et 38 al., 2019). Methane can be emitted via several pathways: simple molecular diffusion, ebullition (in the form of 39 bubbles released from oversaturated sediments), plant-mediated flux (Bastviken et al., 2004), but also through so 40 far neglected pathways including aeration, emissions from dry/drying sediments, or dredged organic material 41 (Kosten et al., 2020). Among all, ebullition is considered the dominant pathway (van Bergen et al., 2019; Kosten 42 et al., 2020), which can contribute 50–96 % (Casper et al., 2000; Xiao et al., 2017; van Bergen et al., 2019; Yang 43 et al., 2020; Zhao et al., 2021) to the total  $CH_4$  flux. Along with the second important pathway – molecular 44 diffusion, both exhibit high spatiotemporal variability due to various physical and biological factors acting on very 45 short time scales, for instance, temperature (van Bergen et al., 2019), nutrient loading (Zhang et al., 2021), CH<sub>4</sub> 46 production rates (Zhou et al., 2019), CH<sub>4</sub> oxidation rates (Sanseverino et al., 2012), dissolved oxygen concentration 47 (Xiao et al., 2017), management regime (Yang et al., 2019), or the quality of organic matter in the sediment 48 (Schmiedeskamp et al., 2021). Recently, the direct potential involvement of phytoplankton in CH<sub>4</sub> production and 49 emissions has been emphasised suggested (Yan et al., 2019; Bižić et al., 2020; Bartosiewicz et al., 2021). The 50 complex interactions between physical and biological factors lead to a dynamic and ever-changing environment, 51 characterised by high spatial and temporal variability of methane fluxes in ponds.

52 Although fishponds are recognised as powerful model systems for studies in ecology and evolutionary or 53 conservation biology (De Meester et al., 2005; Céréghino et al., 2008), the extent of environmental heterogeneity 54 in fishponds and shallow inland small waterbodies remains poorly understood (Ortiz and Wilkinson, 2021), largely 55 because the driving factors are either system-specific or highly variable on short time scales (Laas et al., 2012). 56 Most of current information on lentic ecosystem structure and function comes from single-site sampling, in which 57 measurements are taken over time at the deepest point in the lake, which does not sufficiently account for within-58 lake spatial variation (Stanley et al., 2019). The motivation for our study was the growing concern about the role 59 of fishponds as important sources of  $CH_4$  fluxes to the atmosphere (Wik et al., 2016). Unfortunately, the majority 60 of global CH<sub>4</sub> flux estimates rely on upscaling methods (DelSontro et al., 2018a) based on a limited number of 61 measurements that do not account for diurnal and seasonal variability or ecosystem spatial heterogeneity. Yang et 62 al. (2019) indicates that a larger number of spatial replicates over a number of months is mandatory to improve 63 the accuracy of whole-pond CH<sub>4</sub> flux estimates. The published research from other aquaculture studies have been 64 performed mainly in tropical and subtropical zones in fish or crab aquacultures (e.g., Hu et al., 2016; Ma et al., 65 2018; Yang et al., 2019, 2020; Yuan et al., 2019, 2021). To better understand the spatial dynamics of CH<sub>4</sub> fluxes 66 and environmental heterogeneity in temperate freshwater shallow lake, we conducted a spatial sampling of the 67 hyper-eutrophic Dehtář fishpond (Czech Republic, Europe). Since the seasonal CH<sub>4</sub> production is strongly affected 68 by temperature, we focused on warm summer months where the total CH<sub>4</sub> fluxes were expected to be the highest 69 (Jansen et al., 2019). The objectives of our study were (i) to determine the spatial heterogeneity of CH<sub>4</sub> diffusive 70 and total fluxes and fundamental limnological variables (oxygen, temperature, chlorophyll-a) and their change 71 daily and monthly in the hyper-eutrophic pond, and (ii) to identify the factors that influence CH<sub>4</sub> fluxes to improve 72 our understanding of the importance of spatiotemporal variability for global estimates of CH4 efflux to the 73 atmosphere.

# 74 2 Material and Methods

# 75 2.1 Study site description

The Dehtář fishpond (<u>49°00'24.4"N 14°17'39.3"E49° N, 14° E</u>) is a shallow man-made lake (average and maximum depth: 2.4 and 6 m) constructed in 1479 and used for polycultural, semi-intensive production of common carp-(Potužák et al., 2016). It is used for polycultural, semi-intensive production of common carp, which account for 90–95 % of the fish biomass (Rutegwa et al., 2019). The pond is stocked with two-year old carp harvested at



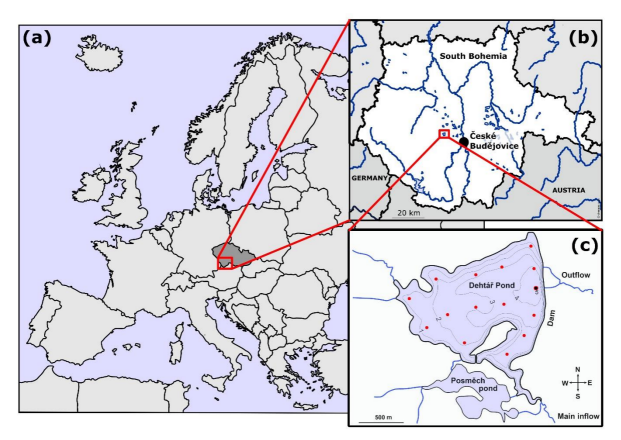


Figure 1. Location (a, b; copyright www.d-maps.com; https://d-maps.com/carte.php?num\_car=2232&lang=en and https://d-maps.com/carte.php?num\_car=265046&lang=en; modified) and bathymetric map (c; credit Jiří Jarošík) of the sampled Dehtář fishpond: <u>blue</u> lines indicate hydrological connections; red dots representing the sampling points. Highlighted sampling point at the dam depicts the deepest site where vertical profiles were measured. Numbers indicate isobath depth.

#### 97 2.2 Sampling design and measurement

98 To measure spatial heterogeneity and temporal changes in limnological parameters and methane fluxes, we 99 conducted three 36-hour surveys in summer 2019 (July 2-3, August 13-14, September 19-20). In the morning 100 (between 5-6 a.m.), we first measured surface values and vertical profiles of temperature, oxygen, and 101 chlorophyll-a concentration at the deepest point (see below for details). We subsequently installed 15 floating 102 polyethylene chambers (as shown in Fig. 1c), serving as fixed sampling sites and at the same time for accumulation 103 of CH<sub>4</sub> fluxes (see further), starting in the western part of the fishpond. During installation (and further during 104 each sampling), temperature, pH, and oxygen concentration were measured at 0.3 m depth using the WTW 330i 105 pH meter and Oximeter (WTW, Weilheim, Germany). Vertical chlorophyll-a profiles were measured at each 106 sampling site using a submersible fluorescence probe (FluoroProbe, bbe Moldaenke, Kiel, Germany). From each 107 site, the average chlorophyll-a concentration in the surface layer (0-1 m depth) was used to assess the 108 phytoplankton spatial heterogeneity.

To minimise the chance that the differences observed among sites were due to time of day, we conducted repeated measurements at the deepest point at the end of each sampling. This was relevant mainly to the initial measurement, when the installation of all floating chambers took a total of 3 hours and 50 minutes. All other measurements, i.e., the interval between the first and last sampling point, required approximately two hours each. If there was a change, all values were corrected for the sampling time by linear interpolation:

114 
$$P_{corr} = P_t + (P_{end} - P_0) \times \frac{(t - t_0)}{(t_{end} - t_0)}$$
(1)

115 where  $P_{corr}$  is the corrected value of a parameter,  $P_t$  is its value measured at the time t,  $P_0$  and  $P_{end}$  are parameter 116 values measured at the deepest point at the start (time t<sub>0</sub>) and at the end (t<sub>end</sub>) of the sampling. In the evening and 117 morning of the second day (roughly at 12 h intervals), we performed additional measurements of spatial 118 heterogeneity, allowing us to assess diurnal and nocturnal changes. In addition, samples for measuring CH<sub>4</sub> 119 concentration in the surface water were collected at each site and analysed as described below. To assess diurnal 120 variations in thermal structure and oxygen concentration in the water column, we made vertical profile 121 measurements at the deepest point (Fig. 1c) at 3-6 h intervals using the YSI EXO 2 multiparametric probe (YSI 122 Inc., Yellow Springs, USA).

# 123 **2.3 Methane measurements**

Water samples for determining CH<sub>4</sub> concentration in the surface water were collected at all 15 sampling sites in triplicates into 20 ml glass bottles. The bottles were capped bubble-free under water with black butyl rubber stoppers (Ochs, Germany) and sealed with aluminium crimps. Immediately after sampling, the water samples were preserved by injecting 100  $\mu$  of concentrated sulfuric acid to stop the microbial activity (Bussmann et al., 2015). The samples were processed within one week in the laboratory using a headspace technique according to McAuliffe (1971). Methane concentration in the headspace was measured using an HP 5890 Series II gas chromatograph (Agilent Technologies, USA) and calculated with the solubility coefficient given by Yamamoto et al. (1976).

Methane diffusive fluxes (F) were then calculated for each sampling site indirectly using the 2-layer model withthe equation:

$$134 F = k(C_{sur} - C_{eq}) (2)$$

where  $C_{sur}$  is the CH<sub>4</sub> concentration in surface water in µmol L<sup>-1</sup>,  $C_{eq}$  is the CH<sub>4</sub> concentration in surface water in equilibrium with the atmosphere in µmol L<sup>-1</sup>, and k is the CH<sub>4</sub> exchange constant (cm h<sup>-1</sup>). The atmospheric partial pressure of CH<sub>4</sub> was set to 1.8 ppm. To compute k <u>values</u>, we first derived k<sub>600</sub> estimates using a wind speed-based relationship according to Crusius and Wanninkhof (2003):

139 
$$k_{600} = 1.68 + (0.228 \times U_{10}^{2.3})$$
 (3)

where  $U_{10}$  represents the wind speed at 10 m height (in  $m_{\overline{r}}s^{-1}$ ; obtained from the nearby gauging station) approximated by  $U_{10} = 1.22U$ , where U is the wind speed at 1.5 m height. We then converted  $k_{600}$  to k using the eq. 4 according to Crusius and Wanninkhof (2003):

143 
$$k = k_{600} \left(\frac{sc}{600}\right)^n$$
 (4)

where  $k_{600}$  is the gas transfer velocity for a Schmidt number (Sc) of 600; n is a wind speed-dependent conversion factor, for which we used -2/3 for  $U_{10} < 3.7 \text{ m s}^{-1}$  (Jähne et al., 1987). The Schmidt number for CH<sub>4</sub> was calculated according to Wanninkhof (2014):

$$147 \quad Sc = 1909.4 - 120.78t + 4.1555t^2 - 0.080578t^3 + 0.000658t^4 \tag{5}$$

148 where t (°C) is the water temperature at the time of  $CH_4$  extraction. The parameter  $C_{eq}$  in Eq. (1) was determined

149 from the equation:

$$150 \qquad C_{eq} = \beta \times pCH_4 \tag{6}$$

- where  $\beta$  is the solubility coefficient of CH<sub>4</sub> as a function of temperature according to Wiesenburg and Guinasso (1979), and pCH<sub>4</sub> is the partial pressure of CH<sub>4</sub> in the atmosphere.
- To estimate total  $CH_4$  fluxes from the water column to the atmosphere (i.e., diffusive and ebullitive fluxes), we measured  $CH_4$  accumulation in open-bottom floating polyethylene chambers (volume 3.1 L; area 0.024 m<sup>2</sup>). Each gas chamber was anchored at individual 15 fixed sampling sites, but allowed to float freely on the water surface. Gas was accumulating for approximately 12 h (each incubation had a start and end point) during particular

157 sampling period, i.e., during the day and night periods. Afterwards, 30 ml of gas was carefully taken from each 158 chamber, after mixing the headspace in the chamber, and stored in evacuated Exetainers<sup>®</sup> (Labco Limited, UK). 159 Chambers were ventilated after each sampling period to reset the incubation conditions. Methane fluxes were 160 calculated as the difference between initial background and final concentration in the chamber headspace and 161 expressed on the 1 m<sup>2</sup> area of the surface level per day according to Bastviken et al. (2004).

## 162 2.4 Background limnological parameters

163 During each campaign, water samples for analysis of nutrient concentration and phytoplankton composition were 164 collected from the surface at the deepest point using a Friedinger sampler. Water transparency was measured using 165 a Secchi disk. Total phosphorus (TP) and soluble reactive phosphorus (SRP) were analysed spectrophotometrically 166 according to Kopáček and Hejzlar (1993) and Murphy and Riley (1962), respectively. Concentrations of NH<sub>4</sub><sup>+</sup> and 167 NO<sub>3</sub><sup>-</sup> were determined according to the procedure of Kopáček and Procházková (1993) and Procházková (1959), 168 respectively. Phytoplankton samples were preserved with Lugol's solution and examined for species composition 169 with an inverted microscope (Olympus IMT-2). Weather data were obtained from the gauging station at the 170 fishpond dam.

#### 171 2.5 Statistical analyses

172 Two-tailed paired Student's t-tests and Two-way ANOVA with post-hoc Tukey's multiple comparison test (Prism 173 9.3, GraphPad Software Inc., La Jolla, USA) tested for differences between diffusive and total CH<sub>4</sub> fluxes between 174 day and night and among three sampling campaigns, respectively. The percentage of data variability explained by 175 different factors (daytime, month and site) was calculated with the Two-way RM ANOVA. Contour graphs 176 illustrating changes in spatial heterogeneity of measured parameters were constructed in Surfer 10 (Golden 177 Software, Inc., Colorado, USA) using the kriging contouring method. Spatial heterogeneity was quantified for 178 each sampling by calculating the spatial variance (i.e., coefficient of variation of values measured at 15 sampling 179 sites; see, e.g. Fig 2):

$$180 \qquad CV\% = 100 \times \frac{SD}{mean} \tag{7}$$

Higher spatial variance indicates increasing ecosystem patchiness. Linear mixed-effects models were used to analyse the effects of  $O_2$ , pH, temperature, and water depth on the  $CH_4$  diffusive fluxes with the random effect of time of day nested within the effect of sampling date. The most parsimonious model was obtained by a manual backward selection, where we sequentially removed all insignificant predictors (p > 0.05) using likelihood ratio tests implemented in the drop1 function (Zuur et al., 2009). We also compared the slopes of the month-specific regression lines produced by the model using analysis of covariance (Zar, 1984). Linear mixed-effects models were implemented in the lme4 package version 1.1-21 (Bates et al., 2015), and Kenward-Roger F-tests were computed using the ANOVA Type II function from the pbkrtest package version 0.4-7 (Halekoh and Hojsgaard, 2014). The prediction of the resulting final model was visualised in the package ggeffects version 0.14.1 (Lüdecke,

- 190 2018). Package performance version 0.4.4 (Lüdecke et al., 2020) was used to calculate Nakagawa's R<sup>2</sup> of the linear
- 191 model. The statistical analyses were performed using R software (v. 3.5.2, R Core Team, 2018).

#### 192 3 Results

# 193 **3.1** Weather and background fishpond characteristics

Weather parameters varied among sampling campaigns. In July, clear skies prevailed with the daily air temperature above 30 °C (Table 1). During the August and September measurements, it was very cloudy, and daily air temperatures decreased to 22 and 18 °C, respectively. The water level was stable during the whole studied period with a monthly fluctuation of ~ 10 cm. Water transparency was low (15–40 cm), with an increasing trend towards the end of summer (Table 1). Concentrations of total phosphorus and soluble reactive phosphorus were high (Table 1), consistent with a hyper-eutrophic state of the fishpond. In contrast, nitrogen concentrations were rather low, with ammonium nitrogen being the predominant form of inorganic N in the water (Table 1).

201 **Table 1**: Basic characteristics of the Dehtář fishpond during the studied period, measured at the surface at the deepest point.

	July	August	September	
Weather	Clear sky, windy	Partly cloudy, no wind	Partly cloudy, no wind	
Air temperature (°C)	2532	2022	11_18	
Water temperature (°C)	2429	2223	1617	
Maximum wind speed (m s <sup>-1</sup> )	3.2	0.8	0.9	
PHAR (mol m <sup>-2</sup> day <sup>-1</sup> )	9.5	3.4	5.0	
Secchi depth (cm)	15	30	40	
TP (μg <u>L</u> I <sup>-1</sup> )	568	527	406	
SRP (µg LI <sup>-1</sup> )	100	200	107	
N-NH4 <sup>+</sup> (μg <u>L</u> <sup>-1</sup> )	23	783	560	
N-NO <sub>3</sub> <sup>-</sup> (μg <u>L</u> <sup>1-1</sup> )	14	23	46	

Chl-a ( $\mu$ g L <sup>t-1</sup> )	456	156	185	
Phytoplankton composition	Cyanobacteria	Cyanobacteria, green algae, cryptophytes	Cryptophytes, green algae	

Chlorophyll-a concentrations were highest in July due to the dense cyanobacterial bloom accumulated at the
surface (Table 1). The phytoplankton consisted of only three cyanobacterial taxa: *Dolichospermum flos-aquae*, *Planktothrix agardhii*, and *Raphidiopsis mediteranea*. In August, phytoplankton was more diverse but also
dominated by cyanobacteria: *P. agardhii*, *Aphanizomenon issatschenkoi*, and *D. flos-aquae*. In September,
cyanobacteria were absent and instead, cryptophytes (*Cryptomonas reflexa*), green algae (*Pediastrum, Coelastrum*and *Desmodesmus*) and dinoflagellates (*Ceratium hirundinella*) prevailed.

# 209 **3.2 Methane concentration and fluxes**

210 The CH<sub>4</sub> concentration in surface water was highly supersaturated over the whole studied period. The obtained

211 values varied from 0.003 up to 3.75  $\mu$ mol L<sup>-1</sup> (Fig. 2), which corresponded to saturation levels of 108-12 834%.

It is obvious, that the obtained data show remarkable variance: the mean ( $\pm$  SD) values were 0.22  $\pm$  0.18 for July,

213  $0.34 \pm 0.45$  for August, and  $1.61 \pm 0.61 \mu mol L^{-1}$  for September (Suppl. Fig. 11).

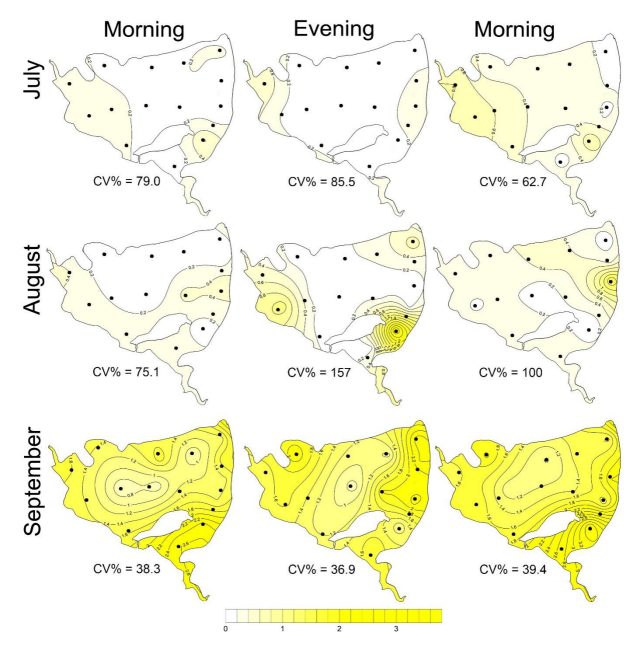
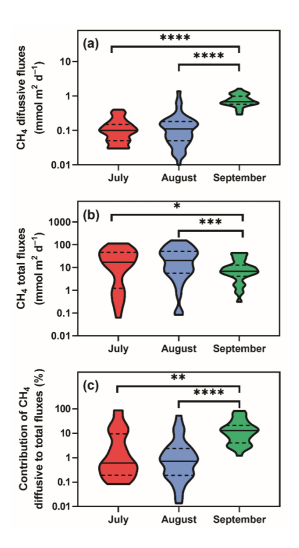


Figure 2: Surface methane concentrations (μmol L<sup>-1</sup>). Contour graphs illustrating both seasonal and daily changes in spatial
 heterogeneity (indicated by the coefficient of variation, CV%) in the fishpond. Black dots representing the sampling sites.

Diffusive fluxes (i.e., calculated from CH<sub>4</sub> concentration, see Eq. 2) showed the lowest values in July and August (average 0.12 and 0.16 mmol m<sup>-2</sup> d<sup>-1</sup>, respectively) and pronouncedly peaked in September (average 0.78 mmol m<sup>-2</sup> d<sup>-1</sup>, Fig. 3a). By contrast, in July and August, the average total CH<sub>4</sub> fluxes (obtained with floating chambers) showed the highest values (average 31.8 mmol m<sup>-2</sup> d<sup>-1</sup>; ranging from 0.08 to 152 mmol m<sup>-2</sup> d<sup>-1</sup>) while in September, total CH<sub>4</sub> fluxes were three times lower than before (average 11.8 mmol m<sup>-2</sup> d<sup>-1</sup>, range 0.3 to 43.5 mmol m<sup>-2</sup> d<sup>-1</sup>, Fig 3b). As a result, diffusive fluxes accounted for only a minor fraction of total CH<sub>4</sub> fluxes to the atmosphere (on average, 9.2 % in July, 4.1 % in August, 18.5 % in September, Fig. 3c).



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Figure 3: Violin plots of CH<sub>4</sub> diffusive (a) and total fluxes (b) during the studied period. Panel (c) depicts differences in the percentage contribution of diffusive to total fluxes. Solid lines are medians, while dashed lines denote quartiles. Asterisks indicate significant differences (\* p<0.05, \*\* p<0.01, \*\*\* p<0.001, \*\*\*\* p<0.001) between sampling dates determined by two-way ANOVA with Tukey's multiple comparison test. Note that a log scale is used here for clarity.</p>

The total CH<sub>4</sub> fluxes show spatial variability within the fishpond that range four orders of magnitude (Fig. 3, 4; Suppl. Fig. 11; Suppl. Table 1). The observed spatial pattern showed high temporal variability on both daily and monthly scales (Fig. 2, 4, Suppl. Table 1). Most of the variability in CH<sub>4</sub> diffusive fluxes was explained by sampling date (62.4 %), while for the total CH<sub>4</sub> fluxes, spatial heterogeneity accounted for 87.2 % of data variability (Table 2). Using linear mixed-effects models, we identified water depth as the only significant predictor of total CH<sub>4</sub> fluxes (Df = 1, p < 0.0001, marginal Nakagawa's R<sup>2</sup> = 0.348; Fig. 5).

- 239
- 240

241 **Table 2**: The percentage of data variability explained by different factors (daytime, month = sampling date, and site)

	% of variability				Significance		
	Daytime	Month	Site	Unexplained	Daytime	Month	Site
CH4 diffusive flux	2.3	62.4	13.2	22.1	0.0123	< 0.0001	n.s.
CH4 total flux	0.19	2.4	87.2	10.2	n.s.	n.s.	< 0.0001
рН	4.4	64.9	11.1	19.6	0.0001	< 0.0001	n.s.
Water temperature	3.3	92.3	2.5	1.9	< 0.0001	< 0.0001	< 0.0001
<b>O</b> <sub>2</sub>	21.7	48.1	13.8	16.4	< 0.0001	< 0.0001	0.0135
Chl-a	0.019	74.9	16.7	8.4	n.s.	< 0.0001	< 0.0001

242 calculated with the Two-way RM ANOVA. <u>Statistically</u> significant values (p < 0.01) are bold.

243 Interestingly, slopes of the linear regressions differed significantly among individual sampling campaigns (Fig. 5),

244 indicating an additional season-related factor that affects CH<sub>4</sub> fluxes in the fishpond. Calculated CH<sub>4</sub> diffusive

245 fluxes were not correlated with total fluxes. Linear mixed-effects models did not identify any significant predictor

of the fluxes, indicating that factors and processes out of the study's scope are involved. We found no significant

247 difference in either diffusive or total CH<sub>4</sub> fluxes between day and night.

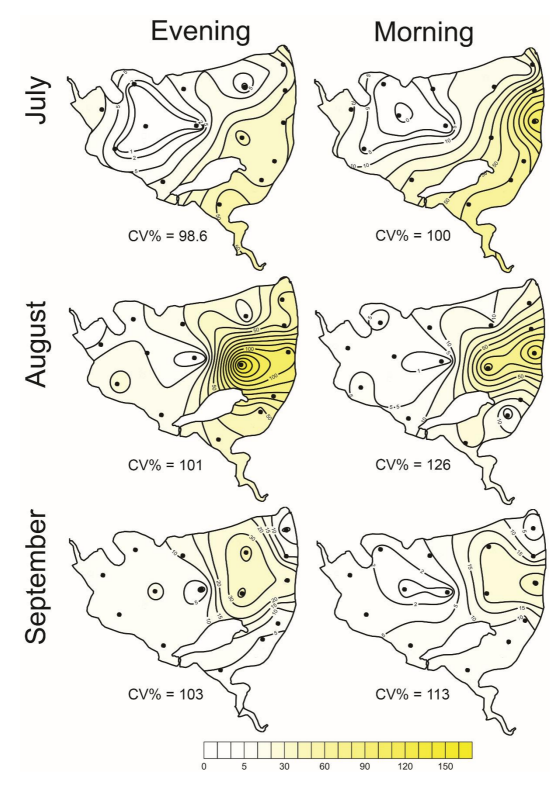


Figure 4: Contour graphs of methane total fluxes in the Dehtář fishpond. Isopleths connect sites with the same value of methane fluxes (mmol m<sup>-2</sup> day<sup>-1</sup>). CV% is a measure of spatial heterogeneity. Black dots representing the sampling sites.

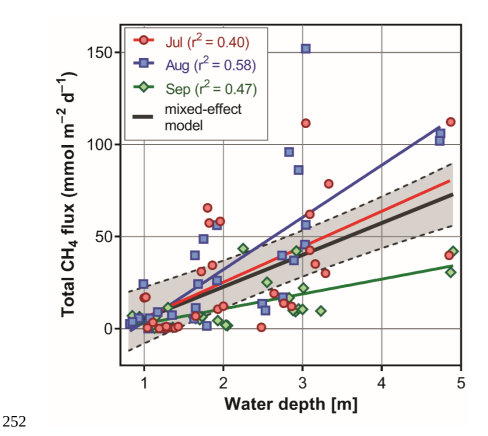
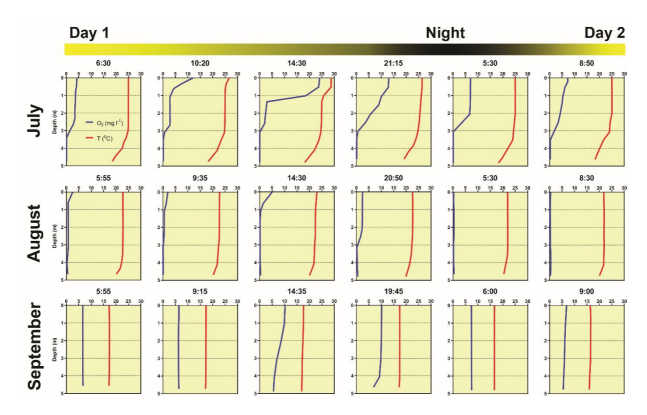


Figure 5: The most parsimonious linear mixed-effect model of methane total fluxes showing the water depth as the only significant predictor. Symbols are the measured values, the solid black line is the prediction, and dashed lines are 95th confidence intervals. Colours indicate month specific relation between total methane fluxes and water depth. Differences in slopes were tested using the F-test. In September, the slope of the regression line was significantly different from that in July and August.

# 258 **3.3 Diurnal changes in vertical profiles of oxygen and temperature**

Several contrasting patterns of vertical temperature and oxygen profiles occurred during summer 2019. Diurnal changes were most pronounced in July (Fig. 6). Surface temperatures varied from 25 °C in the morning to nearly 30 °C in the afternoon. Thermal stratification of the water column was weak in the morning but became strongest at 14:30 with a thermocline at 0.5 m depth (Fig. 6). Later in the afternoon, the water column began to be mixed by wind. The morning vertical oxygen profile was characterised by a surface value of 4.3 mg L<sup>-1</sup>, corresponding to 51 % saturation and anoxia below 3 m.



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Figure 6: Diurnal changes in vertical profiles of temperature and oxygen concentration measured at the deepest point of thefishpond. Numbers above each graph indicate the time of measurement.

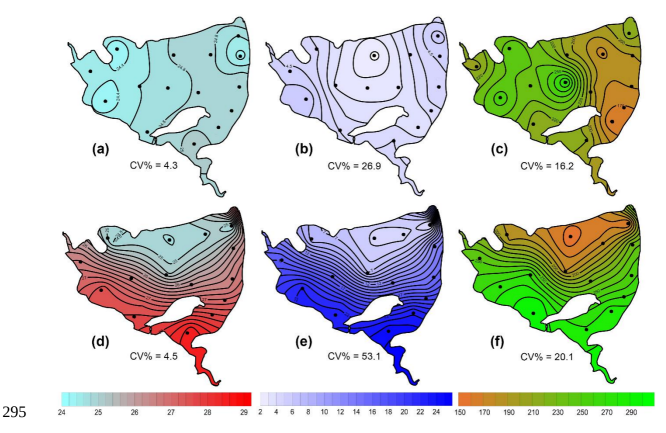
268 Due to the high photosynthetic activity of cyanobacteria, the surface oxygen concentration increased to 24 mg L<sup>-</sup> 269 <sup>1</sup> (320 % saturation, Fig. 6), and a steep oxycline was established at a depth of 0.5\_1.5 m with no effect on the 270 anoxic conditions at the deeper layers. Wind action eroded both the oxy- and thermoclines in the evening, and by 271 the next morning, the vertical profiles were similar to those at the beginning.

272 In August, the water column was almost entirely mixed and low in oxygen in the morning, with only 2.6 mg  $L^{-1}$ 273 (30 % saturation) of oxygen at the surface. Due to cloudy weather, the daily photosynthetic activity of 274 phytoplankton resulted in only a slight increase in oxygen concentration at 0-1.5 m depth (4 mg L<sup>-1</sup>, 47 % 275 saturation). By the morning of the next day, the entire water column turned very close to anoxic (0.4 mg L<sup>-1</sup>, 4 % 276 saturation; Fig. 6), which in turn affected the spatial distribution of zooplankton, as evidenced by the formation of 277 dense zooplankton clouds accumulated in the thin layer just at the surface (see Suppl. Fig. 3). In September, the 278 water column was completely mixed, and we observed only weak daily changes in thermal and oxygen vertical 279 structures (Fig. 6).

# 280 3.4 Effect of wind on spatial heterogeneity of temperature, oxygen and chlorophyll-a

During the summer, all measured parameters showed remarkable within-lake spatial heterogeneity (Suppl. Fig. 5<sub>-</sub>
-8). In July, meteorological conditions allowed for demonstrating the effect of wind on fishpond spatial

283 heterogeneity. In the morning, there were no substantial differences in the surface temperature and oxygen 284 concentrations (Fig. 7ab). Phytoplankton biomass was accumulated mostly in the shallow western part, with the 285 maximum in the centre (Fig. 7c). At 14:00, a light breeze started to blow from the northwest, achieving a maximum 286 of  $3.2 \text{ m s}^{-1}$  (Suppl. Fig. 9). This episode lasted till the evening measurement, and the wind ceased by 21:00. The 287 wind was strong enough to change spatial distribution substantially (Fig. 7d\_-f, Suppl. Fig. 4). In the evening, the 288 surface water temperature on the windward (south) side of the fishpond was ~ 4 °C higher than in the north (Fig. 289 7d). The wind also induced order of magnitude differences in oxygen concentration along the north-south axis of 290 the fishpond (3 mg  $L^{-1}$  of O<sub>2</sub> at the north, 24 mg  $L^{-1}$  of O<sub>2</sub> at the south; Fig. 7e) and affected phytoplankton 291 distribution in the fishpond, resulting in remarkable bloom accumulation in the south (Fig. 7f, Suppl. Fig. 8). 292 During the calm night after the disturbance, the north-south gradient substantially weakened. In August and 293 September, the thermal heterogeneity of the pond was rather low, but the spatial distribution of oxygen and 294 chlorophyll-a remained highly variable (Suppl. Fig. 5-8, Suppl. Table 1).



**Figure 7:** Contour graphs of surface temperature (a, d; °C), oxygen concentration (b, e; mg L<sup>-1</sup>) and chlorophyll-a concentration (c, f;  $\mu$ g L<sup>-1</sup>) measured on July 2 at different times of day: a, b and c are the morning measurements; d, e and f are evening measurements following a wind disturbance. Coefficient of variation (CV %) is a measure of spatial heterogeneity of measured parameters. Black dots representing the sampling sites.

# BOO <u>4</u>Discussion

#### **301 4.1 Methane fluxes**

302 Assessing spatial heterogeneity of the CH<sub>4</sub> fluxes within a fishpond is critical for a reliable estimate of its 303 contribution to the global CH<sub>4</sub> budget. In our study, the variability in total CH<sub>4</sub> fluxes spanned several orders of 304 magnitude (ranging from 0.06 up to 1 121.3 mmol m<sup>-2</sup> d<sup>-1</sup>), which is in agreement with similar studies (Casper et 305 al., 2000; DelSontro et al., 2016; Natchimutu et al., 2016). However, most system-specific CH<sub>4</sub> flux estimates rely 306 on upscaling from a limited number of sites (Bastviken et al., 2004; Rasilo et al., 2015; Wik et al., 2016) because 307 obtaining spatial variability in CH<sub>4</sub> emission is methodologically challenging. In general, spatial heterogeneity 308 may reflect differences in water sources, physical mixing, local transformations and biogeochemical processes and 309 rates among lake habitats (Loken et al., 2019). In deep lakes, littoral areas can contribute disproportionately to 310 total lake CH<sub>4</sub> fluxes (Hofmann et al., 2010; Hofmann 2013, Natchimuthu et al., 2016; Schilder et al., 2013) and 311 are often missed by traditional sampling approaches (Wik et al., 2016). According to Wik et al. (2016), low 312 temporal and spatial resolutions are unlikely to cause overestimates. On the other hand, DelSontro et al. (2018b) 313 suggested that horizontal transport of CH<sub>4</sub> produced in littoral zones and the interaction between physical and 314 biological processes (e.g. air-water gas exchange, water column mixing, the interplay between CH<sub>4</sub> production 315 and microbial oxidation) may result in an underestimation of whole-lake CH<sub>4</sub> fluxes based on centre samples. 316 Similarly, Natchimuthu et al. (2016) found that up to 78 % underestimation would occur if samples obtained only 317 from the lake centre are used to extrapolate the total CH<sub>4</sub> flux. However, extrapolating our data from the deepest 318 point of the Dehtář fishpond would lead to an overestimation of the CH<sub>4</sub> fluxes by a factor of 2.9 (Suppl. Fig. 12). 319 The bias introduced by the deepest point measurement appears to be highly variable among systems with different 320 morphology, geographical location, mixing regimes or trophic states. For instance, analysis of 22 European lakes 321 during late summer has shown that spatially resolved CH<sub>4</sub> diffusive fluxes were highly variable for individual 322 lakes, yielding 55-300 % differences in the whole-lake estimates (Schilder et al., 2013). Schmiedeskamp et al. 323 (2021) observed an increase in CH<sub>4</sub> fluxes from the shore towards the centre in response to increasing sediment 324 C-content in two shallow German lakes. In line with these findings, our results provide further evidence that 325 spatially resolved data are needed to validate the uncertainties that come from using single-point samples to 326 represent whole-lake processes in hyper-eutrophic systems. As stated by Loken et al. (2019), rather than assuming 327 spatial homogeneity, scaling-up exercises of global carbon budgets should acknowledge the uncertainty that comes 328 from extrapolating from spatially limited data sets.

329 In the Dehtář fishpond, the total CH<sub>4</sub> fluxes increased with water depth, and this relationship was month specific. 330 The highest CH<sub>4</sub> fluxes at the deepest points may seem contradictory to previous studies, in which the highest 331 fluxes were typically observed in littoral areas (e.g., DelSontro et al., 2018b; Hofman et al., 2010; Natchimuthu et 332 al., 2016; Schilder et al., 2013). However, these findings are based on studying mostly large, shallow to medium-333 deep oligotrophic lakes whose morphology, trophic state or oxygen regime sharply contrast with the hyper-334 eutrophic Dehtář fishpond, where the upper two meters of the water column were oxygen-saturated while the 335 deepest strata were mostly anoxic, i.e., the extent and duration of bottom anoxia could be the most influential 336 factor contributing to the highest methane fluxes at the deepest point of the pond. In such-hyper-eutrophic systems, 337 high nutrient loading increases autochthonous primary production (Potužák et al., 2007; Rutegwa et al., 2019) and 338 promotes oxygen consumption and anaerobic decomposition in the sediments (Baxa et al., 2020), leading to 339 enhanced CH<sub>4</sub> production (Bastviken et al., 2004; Grasset et al., 2018). In aquaculture ponds in Southeast China, 340 CH<sub>4</sub> fluxes exhibited considerable spatial variations and peaked in the relatively deep feeding zone, where the 341 large loads of sediment organic matter fueledfuelled CH<sub>4</sub> production (Yang et al., 2020). Furthermore, sediment 342 temperature was the strongest predictor of CH<sub>4</sub> fluxes in <u>shallow</u> ponds with anoxic hypolimnion (DelSontro et 343 al., 2016; Yang et al., 2020). It is, therefore, reasonable to assume that both temperature and oxygen concentration 344 in the sediment likely contributed to changes in observed CH<sub>4</sub> fluxes during the studied period in our study. 345 Although both parameters were not directly measured in the sediment, it can be deduced from their vertical profiles 346 that the probability of sediment anoxia was highest in August and lowest in September, and the sediment 347 temperature was lowest in September (see Fig. 5).

348 Our results agree with the generally accepted view that processes other than diffusive fluxes,— most likely 349 ebullition<sub>4</sub>— represent the major CH<sub>4</sub> pathway to the atmosphere in hyper-eutrophic ponds used for intensive fish 350 production (Kosten et al., 2020). Although freshwaters with high primary production are more likely to have high 351 CH<sub>4</sub> ebullition rates (DelSontro et al., 2016), the dominant role of ebullition was also found across lentic systems 352 differing in size, trophic status or geographical location (Aben et al., 2017). Ebullition accounted on average for 353 56 % of total CH<sub>4</sub> fluxes in northern ponds in Canada (DelSontro et al., 2016), 49 and 71 % in two different zones 354 of Lake Taihu (Xiao et al., 2017) and 48-83 % in three Swedish lakes (Natchimuthu et al., 2016; Jansen et al., 355 2019). The highest contribution was found in the small hyper-eutrophic Priest Pot (UK), where ebullition 356 represented 96 % of the total CH<sub>4</sub> flux from the pond (Casper et al., 2000). Apparently, the contribution of 357 ebullition can vary among systems and will remain uncertain until measurement designs cover enough 358 spatiotemporal variability to yield representative values for the whole ecosystem.

359 In shallow water bodies, a semi-stable flux of microbubbles was suggested to account for a significant portion of 360 the total CH<sub>4</sub> flux (Prairie and del Giorgio, 2013). When CH<sub>4</sub> concentration in the water column is above a certain 361 threshold of microbubble density, these microbubbles likely aggregate, fuse, and escape to the atmosphere from 362 buoyancy (Prairie and del Giorgio, 2013). Even a small fluctuation in hydrostatic pressure (e.g., due to changes in 363 atmospheric pressure) or lake water level was shown to trigger enhanced CH<sub>4</sub> ebullition (Bastviken et al., 2004; 364 Casper et al., 2000; Varadharajan and Hemond, 2012). Since ebullition rates increase exponentially with 365 temperature, CH<sub>4</sub> fluxes tend to peak in warm summer months (van Bergen et al., 2019). In our study, 1 % lower 366 air pressure in July and August than in September, along with bottom anoxia and higher water temperature, could 367 account for the enhanced release of CH<sub>4</sub> bubbles from the sediment (31.7 mmol  $m^{-2}d^{-1}$ , >90 % of total CH<sub>4</sub> fluxes; 368 Suppl. Fig. 2). In September, when we observed the lowest water temperatures from the studied period and the 369 oxygen profile was rather uniform, ebullition accounted for 81 % (11 mmol m<sup>-2</sup>-d<sup>-1</sup>) of the total CH<sub>4</sub> fluxes. The 370 spatially pooled data of the total CH<sub>4</sub> fluxes measured in the Dehtář fishpond varied from 11.8 to 34.5 mmol m<sup>-2</sup> 371  $d^{-1}$ , which is comparable with similar systems elsewhere (e.g., Bastviken et al., 2010; van Bergen et al., 2019; 372 Baron et al., 2022). To sum up, both diffusive fluxes and ebullition must be addressed to understand the magnitude 373 of total aquatic CH<sub>4</sub> fluxes and how their relative contributions vary across and within aquatic systems (Kosten et 374 al., 2020). Moreover, with an improved determination of  $CH_4$  hot-spots and its causes, the management of ponds 375 could be changed accordingly and so the overall emissions reduced for example by decreasing P-availability and 376 dredging (Nijman et al., 2022).

# 377 **4.2 Effect of wind event on ecosystem spatial structure**

378 Sudden changes in ecosystem spatial structure in response to meteorological forcing have rarely been documented 379 (Loken et al., 2019) since they are hard to predict. Research into them using traditional methods requires intensive 380 effort or expensive instrumentation (Ortiz and Wilkinson, 2021), and it remains a matter of luck to obtain a relevant 381 dataset. In the July sampling campaign, we observed a strong impact of the wind on environmental heterogeneity 382 in the fishpond, which was apparent at a sub-daily time scale. Due to the methodological constraints, i.e., lack of 383 initial measurement, we can only speculate about the effect of wind on the total CH<sub>4</sub> fluxes. The northwest wind 384 during the day advected warmed surface water with cyanobacterial bloom from the north basin to the south. In the 385 evening, it resulted in bloom accumulation on the upward side and a north-south gradient of more than 4 °C and 386 4-2420 mg L<sup>-1</sup> oxygen. After the winds fell off, the observed gradients declined during cooling at night. We assume 387 that the wind blowing across the pond surface drove buoyant cyanobacteria and surface water downwind and 388 caused an upwelling of deeper, colder, and hypoxic water on the upwind side. This wind-related circulation pattern

has been described as a "conveyer belt" in classical textbooks (Reynolds et al., 2006), held responsible for a
disruption of the thermal structure of the water column and the non-uniform spatial distribution of pH, oxygen,
CO<sub>2</sub> or CH<sub>4</sub> and also plankton assemblages (e.g., Loken et al., 2019; Natchimuthu et al., 2016; Rinke et al., 2009;
Ortiz and Wilkinson, 2021).

393 Similar to our study, mild winds (~4 m s<sup>-1</sup>) were strong enough to redistribute heat and induce lake-wide 394 circulations driving upwelling and downwelling in 24 m deep Lake Pleasant (Czikowsky et al., 2018). As the wind 395 blows harder and lasts longer, the effects on ecosystem functioning may target higher trophic levels and become 396 more complex (Rinke et al., 2009). In Lake Constance, a three-day storm event with wind velocities of ~10 m s<sup>-</sup> 397 <sup>1</sup> resulted in a lake-wide displacement of water masses and the formation of the 6–15 °C horizontal surface water 398 gradient, which in turn changed the spatial distribution of phytoplankton, zooplankton and juvenile fish (Rinke et 399 al., 2009). After several stormy days (wind velocities of 12-15 m s<sup>-1</sup>), Čech et al. (2011) observed negative effects 400 of wind-driven changes in water temperature and wave action on perch (Perca fluviatilis) spawning in the Lake 401 Milada. Although wind events affect shallow and deep lakes differently, there is growing evidence that they can 402 have far-reaching consequences on the functioning of aquatic ecosystems by disrupting energy flows, nutrient 403 fluxes, productivity and reproduction, and consequently altering community composition and trophic interactions 404 in the short and long term (Stockwell et al., 2020). As the frequency, intensity, spatial extent and duration of these 405 extreme meteorological events are projected to increase due to ongoing climate change (Comou and Rahmstorf, 406 2012), there is an urgent need to better understand the mechanisms underlying their impacts on the maintenance 407 of the ecosystem services.

## 408 **4.3 Summer changes in the oxygen regime**

409 Our data demonstrate that shallow, hyper-eutrophic ponds have disrupted oxygen regimes (Baxa et al., 2021) with 410 anoxic hypolimnion and may experience severe whole-water column hypoxia critical for aquatic biota (Miranda 411 et al., 2001). The hypoxic periods may result, for example, from sudden weather change (Jeppesen et al., 1990) 412 and last several days, during which physical processes and phytoplankton photosynthesis cannot compensate for 413 intense community respiration (Baxa et al., 2021). This became obvious in August when severe oxygen depletion 414 was measured at the surface across the whole pond, mostly far below a critical level of 4.5 mg  $L^{-1}$ , when adverse 415 effects came into play (Baneriee et al., 2019). However, oxygen surface concentrations in shallow parts of the 416 pond were substantially higher regardless of the time of day, which contrasts with findings of Miranda et al. (2001), 417 who emphasised shallow waters as the most sensitive parts of lakes, where hypoxic events can occur due to the 418 respiration of sediment biota. The observed spatial gradients of oxygen may create temporal refugia which allow 419 fish to survive harsh conditions that occur in the deepest part of the pond. To minimise economic losses and 420 negative impacts on the ecosystem, future research should identify the interplay between meteorological forcing, 421 trophic status and anthropogenic pressures (e.g., management practices) that affect oxygen fluctuations at various 422 time scales.

# 423 **4.4 Study limitations**

424 Like in other research, there are some limitations in the current study. Since our measurement had only a limited 425 temporal resolution (three samplings during the summer season), it is not appropriate to extrapolate CH<sub>4</sub> emissions 426 for annual values. Noticeably, future research must increase the frequency of the sampling and include also 427 innovative techniques to measure CH<sub>4</sub> fluxes at multiple fishponds, with different management regime. In our 428 study, the 12-h deployment time of the floating chambers could have led to extensive gas accumulation, which in 429 turn might have resulted in an underestimation of the total CH<sub>4</sub> fluxes due to the dissolution of the CH<sub>4</sub> from the 430 chamber into the water once the equilibrium concentration in the chamber is overcome (Bastviken et al., 2010). 431 However, CH<sub>4</sub> concentrations in water corresponded to a supersaturation of several orders of magnitude, so the 432 introduced bias appears to be of minor importance. In any case, our daily spatially pooled total CH4 fluxes (11.8-433 34.5 mmol m<sup>-2</sup> d<sup>-1</sup>) represent a rather conservative estimate for the global methane budget. In our study, we also 434 did not address the important processes that could shed light on the lake CH<sub>4</sub> budget, such as CH<sub>4</sub> oxidation rates 435 (Bastviken et al., 2008) or biological interaction (e.g., protistan grazing on CH<sub>4</sub> oxidising bacteria) in aquatic food 436 webs (Sanseverino et al., 2012) that can affect the overall CH<sub>4</sub> fluxes. We also lack information about spatial 437 differences in sediment microbiota and organic carbon content and compositions, which were found to affect CH<sub>4</sub> 438 production rates (Berberich et al., 2020; Emerson et al., 2021). Despite the limitation mentioned above, our results 439 show that complementary spatial surveys help contextualise the fixed station dynamics and provide additional, 440 management-relevant information about the fishpond.

441 For improved monitoring strategies, however, a continuous measurement approach like eddy covariance would be 442 generally more efficient than traditional sampling at regular intervals. Eddy covariance accounts for temporal 443 variability and provides high temporal resolution data by continuously measuring wind speed, gas concentration, 444 and vertical turbulent fluxes to estimate methane emissions (Erkkillä et al., 2018). More importantly, it also offers 445 spatially integrated measurements, averaging emissions over a larger area and therefore accounts for pond spatial 446 heterogeneity. However, it's worth noting that the choice of monitoring approach depends on various factors, 447 including the specific objectives, available resources, and the characteristics of the emission sources. To accurately 448 capture both short-term variability and lake spatial heterogeneity of methane ebullition and diffusion fluxes, the most efficient approach was found to be a combination of continuous measurements with traditional methods
including floating chambers, anchored funnels and boundary model calculations (Schubert et al., 2012; Podgrajsek
et al., 2014; Erkkiłlä et al., 2018). This integrated approach would provide a comprehensive understanding of
methane emissions, enabling better estimation and more effective mitigation efforts.

### 453 **5** Conclusions

454 Many fishponds are hundreds of years old (Potužák et al., 2007), and as such, they are an integral part of our 455 cultural heritage. Nowadays, ponds face a variety of conflicting interests often leading to a focus on maximising 456 fish production that comes at the expense of other ecological services. Intensification of fish production has 457 brought a transition from the traditional management based on natural processes to practices involving 458 supplementary feeding, fertilisation, and overstocking (Pechar, 2000). These changes coupled with the impacts of 459 climate change has resulted to frequent anoxic events and cyanobacterial blooms that reduce biodiversity and limit 460 recreational activities increasingly valued by the public. Our study not only illustrates common water quality 461 problems in fishponds but also provides compelling evidence that methane emissions in these degraded ecosystems 462 further exacerbates negative climate feedbacks and should be considered in discussions to advance the 463 development of sustainable management.

464 Deciphering the mechanisms that drive spatial and temporal heterogeneity in aquatic ecosystem structure and 465 function not only expands our understanding of pond ecology but also provides insights to improve the 466 management of these ecosystems and the services they provide. Our results suggest that spatial heterogeneity needs 467 to be considered when designing experiments and monitoring programs. Without the spatially resolved sampling, 468 we introduce bias into our datasets, hampering our limnological understanding of the ecosystem's functioning and 469 impeding our ability to accurately estimate rates such as methane emissions on a global scale (DelSontro et al., 470 2018a). In agreement with Kosten et al. (2020), we demonstrated that neglecting ebullition leads to a considerable 471 underestimating of the total CH<sub>4</sub> fluxes. Since there are thousands of these intensively managed fishponds, we 472 argue for changing the management practices toward sustainable use of natural resources to mitigate the overall 473 emissions of greenhouse gases from these ecosystems. Future studies are needed to characterise CH<sub>4</sub> fluxes over 474 a greater number and diversity of aquaculture ponds and examine the mechanisms controlling CH<sub>4</sub> emissions in 475 aquatic ecosystems.

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#### 481 Data availability

482 Dataset associated with the manuscript can be found in the GitHub Repositories under 483 https://zenodo.org/badge/latestdoi/587640213.

#### 484 Author contributions

- 485 All authors contributed to the study conception and design. PZ planned the campaign; PZ, AM and JN performed
- 486 the sampling and analyzed the data; AM performed the gas-measurements; VK performed statistical analyses and
- 487 modelling; PZ and AM wrote the manuscript. All authors read and approved the final manuscript.

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