Physical controls on CH₄ emissions from a newly flooded subtropical freshwater hydroelectric reservoir: Nam Theun 2

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22 Abstract

- In the present study, we measured independently CH_4 ebullition and diffusion in the footprint of an eddy covariance system (EC) measuring CH_4 emissions in the Nam Theun 2 Reservoir, a recently impounded (2008) subtropical hydroelectric reservoir located in Lao People Democratic Republic (PDR), southeast Asia. The EC fluxes were very consistent with the sum of the two terms measured independently (diffusive fluxes + ebullition = EC fluxes),
- 28 indicating that the EC system picked-up both diffusive fluxes and ebullition from the

1 reservoir. We showed a diurnal bimodal pattern of CH_4 emissions anti-correlated with 2 atmospheric pressure. During daytime, a large atmospheric pressure drop triggers CH_4 3 ebullition (up to 100 mmol m⁻² d⁻¹) whereas at night, a more moderate peak of CH_4 emission 4 was recorded. As a consequence, fluxes during daytime were twice higher than during night-5 time.

6 Additionally, more than 4800 discrete measurements of CH_4 ebullition were performed at a 7 weekly/fortnightly frequency covering water depths ranging from 0.4 to 16 m and various 8 types of flooded ecosystems. Methane ebullition varies significantly seasonally and depends 9 mostly on water level change during the warm dry season whereas no relationship was 10 observed during the cold dry season. On average, ebullition was 8.5 ± 10.5 mmol m⁻² d⁻¹ and 11 ranged from 0 - 201.7 mmol m⁻² d⁻¹.

12 An Artificial Neural Network model could explain up to 46% of seasonal variability of 13 ebullition considering total static pressure (sum of hydrostatic and atmospheric pressure), 14 variations in the total static pressure, and bottom temperature as controlling factors. This 15 model allowed extrapolation of CH₄ ebullition at the reservoir scale and performing gapfilling over four years. Our results clearly showed a very high seasonality: 50% of the yearly 16 CH₄ ebullition occurs within four months of the warm dry season. Overall, ebullition 17 contributed 60-80% of total emissions from the surface of the reservoir (disregarding 18 19 downstream emissions) suggesting that ebullition is a major pathway in young hydroelectric 20 reservoirs in the tropics.

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22 **1** Introduction

Atmospheric methane (CH₄) mixing ratio has recently reached up to 1875 ppb, which is 162% 23 higher than the pre-industrial value (IPCC, 2013), and is the highest mixing ratio ever 24 reported (Dlugokencky et al., 2009). Currently, CH₄ is directly and indirectly responsible for 25 26 43% of the anthropogenic radiative forcing (IPCC, 2013). The emission from aquatic 27 ecosystems (wetlands and inland freshwaters) is the main source of CH₄ on Earth (IPCC, 28 2013) representing 40% of total CH₄ emissions and 75% of natural CH₄ emissions (IPCC, 29 2013). Emissions from inland freshwaters alone would correspond to 50% of the carbon terrestrial sink (Bastviken et al, 2011). The order of magnitude of CH₄ emissions from inland 30 31 waters is probably conservative (Bastviken et al., 2011). However, these estimates are based 32 on dataset characterized by low temporal and spatial resolution (Bastviken et al., 2004; Barros et al., 2011) although a few studies evidenced strong diurnal variations (Bastviken et al., 2010, Sahlée et al., 2014), seasonal variability (e.g., Abril et al., 2005), transient extreme emissions (e.g., Varadharajan and Hemond, 2012; Sahlée et al., 2014) and strong spatial variations (e.g., Del Sontro et al., 2011; Morrissey and Livingstone, 2012). Therefore, it is possible that hot moments and hot spots of emissions were overlooked leading to a potential underestimation of emissions at the global scale.

7 Among the different known CH₄ pathways to the atmosphere, diffusive fluxes and, to a lesser 8 extent, ebullition have been the most studied ones in natural lakes and anthropogenic water 9 bodies (i.e., hydroelectric reservoirs, farm ponds, etc.). Methane ebullition corresponds to 10 vertical transfer of CH₄ from the sediment to the atmosphere with little physical and biological interactions within a shallow (<20m) water column (McGinnis et al., 2006). 11 12 Methane is produced under anoxic conditions in the sediments or the flooded soils during the 13 mineralization of organic matter. CH₄ bubbles can develop if CH₄ concentration in the 14 interstitial water becomes higher than the maximum solubility of this gas in water. Bubbling 15 fluxes mainly occur in shallow part of lakes and hydroelectric reservoirs (Abril et al., 2005; 16 Bastviken et al., 2004; Galy-Lacaux et al., 1997; Keller and Stallard, 1994) where the 17 hydrostatic pressure is low. The release of the bubbles is triggered by atmospheric pressure variations (Casper et al., 2000; Eugster et al., 2011; Fechner-Levy and Hemond, 1996; 18 19 Mattson and Likens, 1990; Tokida et al., 2005; Wik et al., 2013), variations in water current 20 velocity (Martens and Klump, 1980; Chanton et al., 1989), shear stress at the sediment surface 21 (Joyce and Jewell, 2003), variation of the water level above the sediment (e.g., Boles et al., 22 2001; Chanton et al., 1989; Engle and Melack, 2000; Martens and Klump, 1980; Smith et al., 23 2000), increase of temperature that makes the CH_4 solubility decrease (Chanton and Martens, 24 1988) and strong wind events (Keller and Stallard, 1994). Ebullition is episodic, which make 25 it difficult to accurately quantify. Bubbling fluxes are probably always underestimated (Bastviken et al., 2011; Glaser et al., 2004, Wik et al., 2013), thus they must be determined as 26 27 frequently as possible. In most of the ecosystems where it was determined by discrete sampling with funnels or floating chambers, ebullition was shown to dominate diffusive 28 29 fluxes (Bastviken et al., 2011).

Diffusive CH₄ fluxes at the air-water interface depend on the concentration gradient between the surface water and the atmosphere and the gas transfer velocity (Wanninkhof, 1992). They are usually estimated either by calculations or by floating chambers (FC). The calculation by the thin boundary layer (TBL) model (Liss and Slater (1974) is based on the concentration

1 gradient between the water surface and the atmosphere and a gas transfer velocity. In the 2 literature, the gas transfer velocity and thus the diffusive fluxes are related for instance to 3 wind speed (e.g. Borges et al., 2004; Cole et Caraco, 1998; Frost and Upstill-Goddard, 2002, 4 Guérin et al., 2007), rainfall rates (Ho et al., 1997; Guérin et al., 2007), buoyancy fluxes (McIntyre et al., 2011) or water current velocity (e.g., Borges et al., 2004). The limit of this 5 6 approach is that these relationships are site specific (Borges et al., 2004; Kremer et al., 7 2003b), leading to uncertainties when applied without precaution. Fluxes can also be obtained 8 on site by the use of FCs. This technique is frequently criticized because FCs are supposed to 9 either artificially increase turbulence, specially at low wind speed (Matthews et al., 2003; 10 Vachon et al., 2010), or decrease turbulence by isolating the surface water from the wind 11 friction (Liss and Merlivat, 1986). Nevertheless, FCs were shown to give results in fair 12 agreement compared to other methods in some aquatic ecosystems (Kremer et al., 2003a; 13 Guérin et al., 2007; Cole et al. 2010; Gålfalk et al. 2013). FCs capture both diffusive flux and 14 ebullition if present. In low ebullition conditions, these flux components can be separated by variability patterns among replicate chambers (e.g. Bastviken et al 2004). In high ebullition 15 16 environments, bubble shields may be needed to estimate diffusive flux by excluding ebullition 17 from some chambers (Bastviken et al 2010)..

18 Eddy covariance (EC) measurements of CH₄ emissions are becoming feasible with suitable 19 fast-response sensors now available on the market (e.g. Eugster and Plüss, 2010; McDermitt 20 et al., 2010). It is therefore realistic to quantify CH_4 emissions with EC technique at a scale 21 representative of a wide range of ecosystems. Still, very few EC field deployments have been 22 conducted so far to determine CH₄ emissions, whether in freshwaters lakes (Schubert et al., 23 2012) or man-made reservoirs (e.g. Eugster et al., 2011). EC was shown to be able to capture both diffusive flux and ebullition (Eugster et al., 2011), but with no discrimination between 24 25 the two pathways.

The deployment of EC that captures continuously the emissions with a short time resolution (e.g., 30 min), over long periods (day to year), and large areas (typically hectares) in combination with the intensive deployment of classical discrete sampling methodology for the estimation of diffusion and ebullition should allow the determination of the controlling factors on the short term, daily and seasonal variability of CH_4 emissions by its different pathways.

31 In the present study, CH₄ emission was measured with EC, FC and funnels and calculated by

32 TBL at the Nam Theun 2 (NT2) Reservoir in Lao PDR, Southeast Asia. This man-made lake

33 was chosen because of its potential for high CH₄ emissions owing to its recent impoundment

1 (2008) (Abril et al., 2005; Barros et al., 2011) and for it encompasses large and fast water 2 level variations that should enhance ebullition (e.g., Chanton et al., 1989) compared to most 3 of the natural lakes and wetlands. First, the different methods were compared according to the 4 CH₄ pathways they capture. Once all methods were validated, high frequency measurements 5 over diurnal cycles at different seasons obtained with EC were used for the determination of the physical controls on CH₄ emissions and pathways on a daily basis. Based on a weekly 6 7 monitoring of ebullition during one and half year, we examined its controlling factors in order 8 to estimate ebullition at the entire reservoir scale. This was finally achieved with an artificial 9 neural network approach which allowed us to simulate over a four-year period the ebullition 10 for the entire reservoir from the controlling factors.

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12 2 Site description

The Nam Theun 2 (NT2) hydroelectric reservoir (17°59'49"N, 104°57'08"E) was built on the Nam Theun River, in the subtropical region of Lao PDR. Filling of the reservoir began in April 2008, and the full water level of the reservoir (538 m msl) was reached for the first time in October 2009. At maximal water level, the reservoir floods a 489 km² dendritically shaped area which was mainly covered by dense and medium forests (44.7%), light and degraded forests (36.4%), agricultural lands (11%), with the remaining made up by swamps and rivers (Descloux et al., 2011, Fig. 1).

20 The study site is under a subtropical monsoon climate. The classical meteorological years can 21 be separated into three seasons: warm wet season (WW) from mid-June to mid-October, cool drv season (CD) from mid-October to mid-February, and warm dry season (WD) from mid-22 23 February to mid-June (Fig. 2). Since the water inputs to the NT2 Reservoir are directly related 24 to rainfall, filling of the reservoir typically occurs during the WW season when study area 25 receives 80% of annual rainfall (NTPC, 2005). Since the beginning of power plant operation (March 2010), the reservoir water level had varied seasonally, and achieved its maxima 26 27 during the WW season and minima by the end of the WD season. During the period covered 28 by this study, the reservoir water level varied seasonally by up to 9.5 m (Fig. 2), which corresponded to a variation in the reservoir water surface from 168 to 489 km². With an 29 30 annual average depth of 7.8 m, NT2 Reservoir falls among the shallow reservoir category.

1 3 Methods

2 3.1 Sampling strategy

The EC system was deployed in an open water area (17°41.56'N, 105°15.36'E) chosen to 3 4 offer a smooth fetch in all directions. At this location, fetch varied from about 1 km 5 (northeast), to more than 10 km (northwest). Eddy covariance was deployed four times to 6 study the CH₄ emission during a variety of meteorological and environmental conditions. Two 7 deployments (3 days in May 2009 and 5 days in June 2011) were performed during the 8 transition between the WD season and the WW season (Fig. 2). The average water depth was 9 ~10 and ~1.5 m in May 2009 and June 2011, respectively. The two other field campaigns (14 days in March 2010 and 5 days in March 2011) occurred during the transition between the CD 10 and the WD seasons (Fig. 2). Average water depth was ~10.5 m and ~6.5 m in March 2010 11 and March 2011, respectively. During the May 2009 campaign, reservoir water level was 12 increasing with a mean rate of 1.0 cm.d⁻¹, whereas, the other three campaigns were performed 13 during falling reservoir water level with mean rates of -4.5. -4.6 and -6.9 cm d⁻¹ respectively 14 15 for March 2010, March 2011 and June 2011 (Fig. 2). Statistical details of the different 16 meteorological parameters for the four EC deployments are summarized in Table S1.

17 During each EC deployment, independent measurements of the diffusive and ebullitive fluxes 18 were performed in the footprint of the EC set up with FC and funnels, respectively. Each FC 19 measurement was done together with surface water sampling devoted to the determination of 20 the CH_4 concentration. Note that in March 2010, funnel measurements could not have been 21 performed around the EC set up.

Additional CH₄ ebullition measurements were performed with funnels during five field campaigns covering different seasons, from May 2009 to June 2011, and during a weekly monitoring from March 2012 to August 2013. During this monitoring, spatial variation was explored through measurements at 44 locations spread over seven stations (Fig. 1) representative of the different types of flooded ecosystems (dense and medium forests, light and degraded forest and agricultural lands, Descloux et al., 2011), and different depths (from 0.4 to 16 m) at each sampling site.

3.2 Instrumentation of EC system

The basic EC instrumentation included a 3D sonic anemometer (Windmaster Pro, Gill Instruments, Lymington Hampshire, UK, in May 2009 and March 2010 and a CSAT-3, Campbell Scientific, Logan, UT, USA, in March 2011 and June 2011), and a closed-path fast methane analyser (DLT-100 FMA, Los Gatos Research, CA, USA). Data acquisition was carried out at 10 Hz with a Campbell data-logger (CR3000 Micrologger®, Campbell Scientific).

8 Air was carried to the DLT-100 through a 6 m long tube (Synflex-1300 tubing, Eaton 9 Performance Plastics, Cleveland) with an internal diameter of 8 mm. The tube inlet, protected by a plastic funnel to avoid entry of rainwater, was mounted 0.20 m behind the sonic 10 11 anemometer sensors. An internal 2 µm Swagelok filter was used to protect the sampling cell 12 from the dust, aerosols, insects and droplets. High frequency sampling of air was obtained by the use of a dry vacuum scroll pump (XDS35i, BOC Edwards, Crawly, UK) providing a flow 13 rate of 26 L.min⁻¹. Due to the remote location of our study site, a 5 kVA generator running on 14 the gasoline was used for the power supply of the whole EC instrumentation. Possible 15 16 contaminations of the atmospheric CH₄ concentration measurements from the generator were 17 checked using the wind direction and a footprint model (Kljun et al., 2004). The footprints 18 during the four deployments are shown in Figure S1.

During each EC deployment, wind speed, wind direction, atmospheric pressure, atmospheric temperature, relative humidity and rainfall were measured using a meteorological station (Weather Transmitter Model WXT510, Helsinki, Finland). A radiometer (CNR-1, Kipp & Zonen, Delft, The Netherlands) was used to measure upcoming and outgoing short and longwave radiations. The temperature of surface water was measured at 20 cm depth using a thermistor (Pt100 sensor) coupled to the data-logger.

25 3.2.1 Data processing

The 10 Hz raw data were processed using the EdiRe software (Clement, 1999) for the following steps: (1) spike detection using a standard de-spiking algorithm whereby wind vector and scalars values greater than three times the standard deviation are removed, (2) lag correction and tube attenuation relevant to the closed path DLT-100 gas analyser, (3) coordinate rotation using the planar fit method, and (4) high frequency correction factors to take into account the loss at high frequency due to insufficient sampling rate. Differences among the deployments specific variables i.e. sensor separation distance and instrument placement were considered while processing the data. The EC fluxes of CH_4 were calculated as covariance between the scalars and vertical wind speed fluctuations according to commonly accepted procedures (Aubinet et al., 2001). Fluxes were considered positive if they were directed from the water surface toward the atmosphere, and negative otherwise.

6 **3.2.2 EC data quality control**

7 Fluxes were accepted or rejected according to the following criterion. First, a non-stationarity 8 criterion was applied according to Foken and Wichura (1996). Fluxes were considered 9 stationary and therefore accepted only if the difference between the mean covariance of sub 10 records (5 min) and the covariance of the full period (30 min) was less than 30%. Second, a flux was rejected if its intermittency rose above one (Mahrt et al., 1998). Third, for vertical 11 12 wind component, the skewness and kurtosis were used to stay within the range of (-2, 2) and (1, 8), respectively (Vickers and Mahrt, 1997). Fourth, the momentum flux, $\overline{u'w'}$, was 13 14 required to be negative implying a downward directed momentum flux. In addition, fluxes 15 were rejected when the wind was coming from the power generator unit according to the 16 footprint model of Kljun et al. (2004). For footprint analysis, since the roughness length value was unknown, we considered a value of 0.0002 m, as reported for terrain without obstacle 17 18 (WMO, 2008). According to the model, the footprint was different in extension and prevalent 19 wind directions among the different field campaigns. The smallest footprint area was 20 observed during the March 2010 campaign, and the biggest for June 2011, with the greatest 21 values rarely exceeding 500 m. The analysis confirmed that (1) surrounding terrestrial 22 ecosystems were always outside the footprint (Fig. S1), (2) only 2% of the fluxes were 23 rejected because wind was coming from the power generator, and (3) all FC and funnels 24 measurements were conducted within the EC footprint area.

As mentioned by Eugster et al. (2011), the minimum threshold for friction velocity cannot apply as a good criterion for flux rejection since turbulence generated due to heat loss from the water column can contribute significantly to the emissions to the atmosphere (Eugster et al., 2003, 2011; MacIntyre et al., 2002, 2010). In addition, the criteria on atmospheric concentration formulated by Vickers and Mahrt (1997) for CO_2 over terrestrial ecosystems do not apply for CH_4 over an aquatic ecosystem since emissions could be sporadic due to potential CH_4 burst linked to ebullition (Eugster et al., 2011).

- 1 Quality control criteria applied all together resulted in the acceptance of 57% of the flux data,
- 2 with acceptance rates slightly higher during daytime (59%), than night-time (52%) periods.

3 **3.3 Diffusive fluxes**

4 **3.3.1 Measurement by floating chamber (FC)**

5 Diffusive flux measurements around the EC site were performed with two circular floating chambers (FC_{GC}), (surface area = 0.15 m^2 ; volume = 24.6 L) following the same design as in 6 7 Guérin et al. (2007). Moreover, FCs were covered with a reflective surface to limit warming 8 inside the chamber during measurements. Duplicate samples were taken from the FCs at time 9 0 and then every 15 min for 45 min for a total of four samples per chamber deployment. In the 10 chambers, samples were collected in 10-ml glass vials which contained 6M NaCl solution 11 capped with butyl stoppers and aluminium seals as described in Angel and Conrad (2009). All 12 samples were analysed within 48 hours by gas chromatography (GC). Diffusive fluxes (D_{GC}) 13 were calculated from the slope of the linear regression of gas concentration in the chamber 14 versus time. Diffusion chambers will collect diffusive emissions as well as ebullition 15 emissions if they are present. Therefore, if the slope of the linear regression of gas concentration in the chamber versus time was linear with an $r^2 > 0.8$ then the chamber was 16 assumed to be collecting only diffusive emissions (D_{GC}). If $r^2 < 0.8$ then the chamber was 17 assumed to collect total (diffusive + ebullitive) emissions (subsequently noted DE_{GC} , see Sect. 18 19 4.1).

In March 2011, a floating chamber (surface area = 0.16 m^2 ; volume = 17.6 L) connected to a Picarro[®] CH₄ analyser (FC_{GA}) was also deployed to measure diffusive fluxes (D_{GA}). The calculation and rejection procedures are identical to the ones described above for D_{GC}.

23 **3.3.2 Estimate from surface CH₄ concentrations**

Surface water samples for CH_4 concentration were taken with a surface water sampler described by Abril et al. (2007). Water samples were stored in 60 ml glass vials, capped with butyl stoppers, sealed with aluminium crimps and poisoned until analysis (Guérin and Abril, 2007). Before GC analysis for CH_4 concentration, a N₂ headspace was created and the vials were vigorously shaken to ensure an equilibration between liquid and gas phases (i.e. Guérin and Abril, 2007). The specific gas solubility for CH_4 (Yamamoto et al., 1976) as a function of temperature was used for calculation of CH_4 concentrations dissolved in water. 1 The surface CH_4 concentrations were used together with atmospheric concentrations 2 measured on site in order to calculate diffusive fluxes with Eq. (1):

$$3 \qquad D_{\text{TBL}} = \mathbf{k}_{\text{T}} \cdot \mathbf{C}_{\text{w}} - \mathbf{C}_{\text{a}} \tag{1}$$

4 where D_{TBL} is the diffusive flux at water-air interface, k_T the gas transfer velocity for a given 5 temperature (T), C_w the CH₄ concentration in surface water, and C_a the CH₄ concentration in 6 the surface water at equilibrium with the overlying atmosphere. The k_T values were computed 7 with the following Eq.:

8
$$k_{\rm T} = k_{600} \cdot (600/{\rm Sc_T})^{\rm n}$$
 (2)

9 with k₆₀₀, the gas transfer velocity of CO₂ at 20°C; Sc_T, the Schmidt number of CH₄ at a given temperature (T) (Wanninkhof, 1992); n, a number that is either 2/3 for low wind speed (< 3.7 10 m s⁻¹) or 0.5 for higher wind speed (Jahne et al., 1987). The D_{TBL} values were calculated 11 according to the formulation of k₆₀₀ versus wind speed from MacIntyre et al. (2010) and 12 13 Guérin et al. (2007), the average of both formulations being used in the manuscript. These 14 formulations were chosen because MacIntyre et al. (2010) includes the effect of buoyancy 15 fluxes in the gas transfer velocity, and because Guérin et al. (2007) is one of the very few 16 available for tropical hydroelectric reservoirs.

17 3.4 CH₄ ebullition

Clusters of five to ten PET funnels (diameter = 26 cm, height = 30 cm) attached to each other 18 19 at 1 m distance were assembled. Three to six clusters were positioned below the water surface 20 at locations with different water depths around the same site (within 10-30 m). The funnels remained on sites for 24 or 48 hours. Accumulated gas volumes during the deployment period 21 22 were collected manually through a butyl-rubber septum using a 60 ml syringe at the end of the 23 experiment as described in Wik et al. (2013). The gas sample was stored in glass vials which contained 6M NaCl solution. All gas samples were analysed for CH₄ within 48 hours by GC. 24 CH_4 concentration in bubbles was multiplied by the volume of accumulated gas (V_{EB} , mL m⁻² 25 d^{-1}) over the deployment period to determine CH₄ ebullition fluxes (E_{FUN}). 26

The ebullition was also determined from the FC_{GA} measurements in March 2011. The sudden increase of the CH_4 concentrations in the FC_{GA} was attributed to bubbles. Methane ebullition (E_{GA}) was calculated from the increase of CH_4 concentration in the chamber, the deployment

30 time of FC_{GA} measurement (typically 5 to 20 min) and the surface of the chamber.

1 **3.5 Gas chromatography**

Analysis of CH₄ concentration was performed by gas chromatography (SRI[®] 8610C gas chromatograph, Torrance, CA, USA) equipped with a flame ionization detector (FID). A subsample of 0.5 ml from the headspace of water sample vials and 1 ml of air from flux sample vials were injected. Commercial gas standards (2, 10 and 100 ppmv, Air Liquid "crystal" standards and mixture of N₂ with 100% CH₄) were injected after analysis of every 10 samples for calibration. Duplicate injection of samples showed repeatability better than 5%.

8 **3.6** Artificial Neural Network

9 Artificial Neural Networks (ANN) are a branch of artificial intelligence. Multi Layer 10 Perceptron (MLP) is one type of Neural Network. Unlike other statistical techniques, the MLP 11 makes no prior assumptions concerning the data distribution. It can model highly non-linear 12 functions and can be trained to accurately generalise the results when presented with new, 13 unseen data (Gardner & Dorling, 1998).

14 A suitable set of connecting weights and transfer functions will make the MLP approximate 15 any smooth, measurable function between the input and output vectors (Hornik et al., 1989). The learning process of the MLP is called training, which requires a set of training data (series 16 17 of input and associated output data). These training data are repeatedly presented to the MLP 18 and the weights in the network are adjusted until the error between actual and desired output 19 is the lowest. The set of optimal weights determined by the training process will then be 20 applied on the validation set that has not participated to their elaboration (Delon et al., 2007). 21 Once trained with suitably representative training data, the MLP can generalise to new, 22 unseen input data (Gardner & Dorling, 1999). The quality of these processes is assessed through the calculation of training, validation and generalization costs. 23

The ANN used in this study is based on a commercial version of the Neuro One 5.0 ©12 24 25 software, (Netral, Issy les Moulineaux, France). Some details concerning this specific study are given in this paragraph, and in the supplementary material section S1. The whole 26 27 description of the methodology is detailed in Delon et al. (2007). The architecture of the MPL 28 (deduced from the Vapnik-Chervenenkis theory; Vapnik, 1995) is composed of 3 hidden 29 neurons. All inputs and output are normalized and centred in order to avoid artefact in the 30 training process. After normalization, the data have the same order of magnitude. The network 31 is used in a static version where the lines of the database are independent of each other.

1 In this study, an ANN was used to find the best non linear regression between ebullition 2 fluxes and relevant environmental variables. The database of raw data was composed of 4811 3 individual ebullition fluxes. Fluxes from a given station measured the same day and at the 4 same depth were averaged (different fluxes with same depth value and same meteorological 5 data would introduce noise rather than relevant information in the network), leading to a final 6 database for ANN composed of 510 lines, and 4 columns (one output, and 3 inputs, see 7 discussion paragraph for the choice of the inputs). The dataset used by the MLP is separated 8 in two pools, the training one (330 lines) and the validation one (180 lines).

9 Weight values associated to each input are modified a 100 times (optimization process). Ten 10 initializations (10 series of different sets of weights) are tested for each model. This 11 configuration (100 modifications of weights, 10 models) is tested several times, in order to 12 avoid a local minimum solution. The best algorithm within the 10 launched is chosen, by 13 assessing the following criteria: (1) The lowest generalization cost is chosen, (2) Root Mean 14 Square Error (RMSE) of the training set has to be close to the RMSE of the validation set 15 (23.09 and 23.83 in our case), and (3) results giving negative fluxes are discarded. Learning 16 (training) cost is 6.79, validation cost is 6.9, generalization cost is 7.47, and homogeneity is 17 0.93, which are considered as good enough criteria for choosing the equation. The equation, 18 coefficients and weights necessary to calculate the ebullition flux are listed in the 19 Supplementary material Section, and in Table S2 and S3.

20 3.7 Statistical analysis

21 The methodological, spatial and temporal differences in the CH_4 emissions (diffusion, 22 ebullition and total emissions) were explored. Differences among groups of data were 23 examined using either t-test or Analysis of Variance (ANOVA) in GraphPad Prism (GraphPad 24 Software, Inc., v5.04). Choice of the parametric and non-parametric tests (Mann Whitney test 25 or Kruskal-Wallis tests compare median values) was dependent on normal and non-normal 26 behaviour of the data sets. The potential spatial variability of ebullition fluxes was explored 27 on three flooded soils and vegetation clusters: (1) dense forest which includes dense and 28 medium forest, (2) degraded forest which includes light and degraded forest and, (3) 29 agricultural lands. The effect of depth on ebullition was also tested according to the following 30 three depth ranges: shallow zones (0.4-3 m), intermediate (3-6 m) and deep (6-16 m). Finally, 31 CH₄ emissions from the different seasons (WD,WW and CD) were compared in order to 32 evaluate the temporal variability. All statistical tests used a significance level of 5%. The distributions of the volume of gas emitted by ebullition (V_{EB}), CH₄ bubble concentration and flux (E_{FUN}) were characterized using the Anderson-Darling Goodness of fit in EasyFit 5.5 trial version. A multi linear regression (MLR) was used to find the linear relationship between ebullition fluxes and other environmental variables. The MLR used in this study was based on the SPSS 15.0 for Windows.

6 3.8 Reservoir water temperature, meteorological and hydrological variables

7 The temperature at the bottom of the reservoir has been monitored on the fortnightly basis at 8 nine sampling stations in the reservoir, from January 2009 to date. Meteorological (wind 9 speed, atmospheric pressure, atmospheric temperature, relative humidity, wind direction and 10 net radiation) and hydrological data (rainfall and reservoir water level) were obtained from the 11 monitoring conducted by Electricité de France and Nam Theun 2 Power Company Ltd. 12 (NTPC).

13

14 **4** Results and discussion

4.1 Assessment of CH₄ emissions at the reservoir surface by different methods

17 The effectiveness of four methods (EC, FC, funnels and TBL) to measure CH₄ emissions at the water-air interface were explored during four field campaigns at NT2 (Table 1). Using 18 19 these methods, different emission terms were estimated: (1) diffusion (D_{GC} , D_{GA} and D_{TBL}) at 20 the air-water interface from FC_{GC}, FC_{GA} and TBL (Fig. S2a), (2) ebullition (E_{FUN}, E_{GA}) from 21 funnels and FC_{GA}, (Fig. S2b) and (3) the sum of diffusion + ebullition (DE_{EC} and DE_{GC}) 22 emissions from EC and FC_{GC} (Fig. S2c). All methods were only used simultaneously in 23 March 2011 (Fig. S2 and Table 1). No matter the method used nor the pathway measured, the reservoir emitted CH₄ to the atmosphere during the study period (Table 1). 24

25 4.1.1 Diffusive emission

Only 30% of the diffusive fluxes (D_{GC}) measured by the FC_{GC} fulfilled the acceptance criterion ($r^2 < 0.8$) during the four field campaigns. No fluxes were accepted in March 2011, when the water level of the reservoir was decreasing, and only 5% in June 2011, when the water level was at its lowest. In March 2011, 48% of the D_{GA} were accepted. The comparison of the acceptance percentages in March 2011 indicates that short-term deployment of

chambers (5 min for FC_{GA} vs. 45 min for FC_{GC}) limits the risk of a contamination of the 1 measurement by ebullition. Overall, the average D_{GC} is 1.6 \pm 1.1 mmol $m^{\text{-2}}\ d^{\text{-1}},$ which is 2 comparable with the average D_{TBL} of $1.4 \pm 2.0 \text{ mmol m}^{-2} \text{ d}^{-1}$ (Table 1 and 2) for the four field 3 4 campaigns. For all campaigns except June 2011 (only one validated measurement), the D_{TBL} 5 calculations were not significantly different from the diffusive fluxes measured with FC (ttest, p < 0.05; details in Table 2). Combining all diffusive fluxes obtained by different 6 7 approaches (Table 1), our results showed that there is no seasonal variation for the diffusive 8 fluxes measured (t-test, p < 0.05).

9 4.1.2 Methane ebullition

10 The CH₄ ebullition was measured with funnels (E_{FUN}) in May 2009, March and June 2011. In March 2011, ebullition (noted E_{GA}) was also determined using a FC_{GA} (Table 1, and Fig. 11 S2b). One of the major differences between these two methods is the duration of the 12 measurement. E_{FUN} measurements were performed over 24 to 48 hour periods, whereas E_{GA} 13 measurements were conducted during only 5 to 20 min. In June 2011, E_{FUN} (28.0 ± 11.0 mmol 14 $m^{\text{-2}}\ d^{\text{-1}})$ were almost twentyfold and sevenfold higher than E_{FUN} in May 2009 and March 15 2011, respectively (Table 1). In March 2011, E_{FUN} varied by 2 orders of magnitude with an 16 average of 4.2 \pm 3.6 mmol m⁻² d⁻¹, which is not statistically different from E_{GA} during the 17 same field campaign (4.6 \pm 7.1 mmol m⁻² d⁻¹, Table 1 and 2). It has to be noted that ebullition 18 19 was observed in around 50% of FC_{GA} deployment.

20 4.1.3 Total CH₄ emissions

21 We compare here the two techniques that give access to the total emissions, that is the EC 22 technique, and the floating chamber which had captured bubbles (DE_{GC}). The individual 30min DE_{EC} fluxes varied by four orders of magnitude during all EC deployments (from 0.02 to 23 103 mmol m⁻² d⁻¹). On average, DE_{EC} fluxes varied oppositely with the water depth, with the 24 highest mean flux (29 \pm 16 mmol m⁻² d⁻¹) in June 2011 for the shallowest water depth (~1.5 25 m) (Table 1, and Fig. 3). Figure S2c and 3 show total CH₄ fluxes calculated from DE_{GC} data. 26 Altogether, DE_{GC} also varied by four orders of magnitude (from 0.02 to 132 mmol m⁻² d⁻¹) 27 during all deployments, and DE_{GC} fluxes also varied oppositely with the water depth with the 28 highest mean flux (54 \pm 35 mmol m⁻² d⁻¹) in June 2011. For half of the campaigns, DE_{GC} and 29 DE_{EC} were significantly different (Table 2, Fig 3). 30

We then compared continuous DE_{EC} with the sum of the discrete sampling of diffusive (D_{GC} , D_{GA} and D_{TBL}) and ebullitive fluxes (E_{FUN} and E_{GA}) for three field campaigns (May 2009, March and June 2011) among four (no ebullition measurements done within the footprint in March 2010). The sum of independent estimates of diffusive fluxes (D_{TBL} , D_{GC} , D_{GA}) and ebullition (E_{FUN} and E_{GA}) determined on less than a m² were found to be in good agreement with total emissions determined from EC over thousands of m² (Table 2 and Fig. 3). This confirms that the EC system is able to pick-up both diffusive fluxes and ebullition from the reservoir as already shown by Schubert et al. (2012).

7 Even if statistically the comparison of total emissions by the different approaches is good, one 8 should note that in a handful occasions, DE_{GC} exceed DE_{EC} and the sum of diffusive fluxes 9 $(D_{GC}, D_{GA} \text{ and } D_{TBL})$ and ebullition $(E_{FUN} \text{ and } E_{GA})$ by a factor up to 100 (Fig. S2c). These differences can clearly be explained by sudden release of bubbles in these seldom occasions. 10 11 This reveals very strong spatial and temporal heterogeneities of the ebullition process. 12 Because ebullition is highly sporadic and occurs during very short period of time 13 (Varadharajan and Hemond, 2012), its measurement by FC over short period of time and 14 small surface might lead to an over-estimation of this emission pathway if hot spots and hot moments are captured during the deployment of the chamber. Such phenomenon is strongly 15 16 smoothed when using funnels over longer period of time than the typical floating chamber 17 deployment time (typically 12-48 h versus 10-45 min). Globally the EC measurements are 18 ideal for capturing the large spatial and temporal variability of total CH₄ fluxes at the surface 19 of aquatic ecosystems prone to ebullition. However, the discrimination of diffusive fluxes and 20 ebullition requires the deployment of either bubble shielded-FC to obtain the diffusive 21 emissions or the deployment of funnels to obtain the ebullition. The use of recent techniques 22 like the equilibrator technique (Abril et al., 2006) and subsequent TBL calculations for 23 diffusive fluxes, or hydroacoustic measurement which is capable to capture the hot spots of 24 ebullition (DelSontro et al., 2011) combined to the EC might also allow the identification of 25 those hot moments and their controlling factors.

For the four campaigns, the contribution of ebullition to total emissions from EC (DE_{EC}) ranged between 57 and 93% of the total CH₄ emissions from the EC footprint (Table 1 and Fig. 3). As already mentioned in some recent publications (Bastviken et al., 2004, 2010; DelSontro et al., 2010, 2011; Schubert et al., 2012), ebullition is a major CH₄ pathways that is often neglected in aquatic ecosystems, especially in the tropics and subtropics where the high temperature triggers ebullition by both enhancing CH₄ production in the sediments (Duc et al., 2010), and decreasing CH₄ solubility in the water column (Yamamoto et al., 1976).

1 **4.2** Total CH₄ emissions (DE_{EC}) *versus* hydrostatic pressure

Based on the four field campaigns time series from the EC system (Fig. S3), we did not find any correlation between DE_{EC} and the wind speed and the buoyancy fluxes (Fig. S4). As these parameters are known controlling factors of the diffusive fluxes (e.g., Guérin et al., 2007; McIntyre et al., 2010), it indirectly confirms that ebullition dominates the total emissions at the surface of the NT2 Reservoir, as shown in the previous section.

Daily DE_{EC} was plotted against the daily (1) water depth and (2) change in the water depth 7 8 (cm d⁻¹), and (3) specific water level change (water level change normalized by the average 9 water depth) (Fig. 4). DE_{EC} is negatively correlated to the water depth (p <0.0001, Fig. 4a) as 10 it is usually the case for ebullition in lakes (Bastviken et al., 2004; Wik et al., 2013), 11 hydroelectric reservoirs (Galy-Lacaux et al., 1999; Keller and Stallard, 1994), estuaries 12 (Chanton et al., 1989) and the marine environment (Algar and Boudreau, 2010, Martens and 13 Val Klump, 1980). According to our data set, emission can be enhanced by a factor of 5 for a 14 water depth difference of 10 m which corresponds to the observed maximum seasonal water 15 level variations at NT2. Though measured in different seasons, diffusive fluxes measured by 16 FC in the EC footprint are constant for the four deployments (see Table 1). This implies that seasonal variation of the CH₄ emissions at a single site is mostly controlled by water level 17 18 differences and subsequent ebullition. However, this does not exclude that CH₄ emissions are 19 higher during the warm dry season than during cooler seasons as a consequence of enhanced 20 methanogenesis with higher temperature (Duc at al., 2010). It appears that the effect of water 21 level change (6-9 cm) is proportionally stronger in shallow water (2 m) than in deep water 22 (10.5 m) (June 2011, Fig. 4b), meaning that the same water level change could favor higher 23 fluxes in shallow area than in deep waters. This effect is well describes with the specific water 24 level change (Fig. 4c): fluxes were lower when daily variations of the depth were 5 to 7 cm, 25 corresponding to specific water level changes of less than 1% for most of the field campaigns 26 (March 2010 and 2011), than in June 2011 when the same water level variations corresponded to a specific water level change of 4-5% which triggered emissions up to 100 mmol $m^{-2} d^{-1}$. 27 28 Overall, in the context of this subtropical hydroelectric reservoir with a high contribution of 29 ebullition, these results show that the hydrostatic pressure plays an important role in 30 controlling the CH₄ fluxes since (1) the water depth explains about 70% (Fig. 4a) of the 31 variability of the CH₄ emissions, (2) seasonal variations of CH₄ emissions by a factor of 5 are 32 mostly due to the enhancement of ebullition due to the low water level in the WD season and, (3) the effect of change in water level on ebullition is more effective in shallow area than in
 deeper zone of aquatic ecosystems.

3 **4.3** Effect of atmospheric pressure on diurnal cycle of total CH₄ emissions (DE_{EC})

4 In the DE_{EC} time series (Fig. S3), it appeared that two CH₄ peaks of emissions occurred daily. 5 In order to investigate the drivers of these emission peaks, DE_{EC} flux data were binned by 6 time of the day and then averaged for each deployment. A clear diurnal bimodal pattern of 7 DE_{EC} fluxes, with a first peak in the middle of the night (between midnight and 3:00 am) and 8 a second one around noon was observed during all four campaigns (Fig. 5a, b, c, d), and is 9 apparently related to the semidiurnal evolution of the atmospheric pressure (a phenomenon 10 due to global atmospheric tides). Diurnal pattern of CH₄ emissions was also recently 11 evidenced by Sahlée et al. (2014) who measured CH₄ fluxes using a EC system over a natural 12 lake in Sweden. They observed higher fluxes at night-time linked to enhanced diffusion through convective mixing (McIntyre et al., 2010; Sahlée et al., 2014). At NT2, 30 min-13 14 binned DE_{EC} is anti-correlated with atmospheric pressure (Fig. 5a, b, c, d). Furthermore, DE_{EC} was found to be anti-correlated with the change in atmospheric pressure, evidencing a strong 15 16 control of the atmospheric pressure change over the fluxes, most likely through ebullition (Fig. 5e, f, g, h). It is noteworthy to point out that the coefficient of determination is better for 17 18 the campaign with the lower water depth at the EC site (1.5 m, June 2011, Fig. 5e, f, g, h) 19 indicating that the variations of the atmospheric pressure have more effect at low hydrostatic 20 pressure (higher relative change in pressure). We also calculated buoyancy fluxes in order to 21 look for the potential occurrence of high diffusive fluxes due to convective mixing (Fig. 5i, j. 22 k) as in McIntyre et al., (2010) and Sahlée et al (2014). On one hand, night-time peak of CH₄ 23 emission coincides with low but constant buoyancy fluxes (i.e.; most instable water column) 24 and moderate atmospheric pressure drop. The fact that the buoyancy flux does not decrease 25 during the peak of CH₄ indicates a low control on the emissions, if any. On the other hand, 26 daytime peak of CH₄ emissions are linked with maximum buoyancy fluxes which cannot enhance emissions (i.e. most stable water column). These observations tend to prove that CH₄ 27 bursts in the night and around noon (up to 100 mmol m⁻² d⁻¹) could be entirely attributed to 28 the atmospheric pressure drops that triggered ebullition, more than any buoyancy effect. 29

The effect of pressure on ebullition was already shown in natural lakes (Casper et al., 2000; Eugster et al., 2011; Mattson and Lickens, 1990; Wik et al., 2013) and peatlands (Fechner-

32 Levy and Hemond, 1996; Tokida et al., 2005), and the effect of buoyancy fluxes on diffusive

1 fluxes in lakes (McIntyre et al., 2010; Sahlée et al., 2014), but this is the first time that a daily 2 bimodal variation of CH₄ emissions is evidenced. CH₄ emissions around noon were 3 approximately 10 times as high as fluxes near sunset and sunrise (Fig. 51, m, n, o) and 2 times higher than during the night-time for all EC deployments (p = 0.0036, p = 0.0002, p = 0.00154 5 and p <0.0001 respectively for May 2009, March 2010, March 2011 and June 2011, Mann-6 Whitney test). This implies that the quantification of CH₄ emissions by ebullition and 7 diffusion from inland aquatic ecosystems has to be done over 24 h cycles in order to obtain 8 realistic estimates.

9 **4.4** Spatio-temporal variations of CH₄ ebullition (E_{FUN})

By definition, EC systems are not suitable for the exploration of fine spatial variations and of effect of water depth on ebullition within a single season. Because of logistic difficulties, it was not possible to leave the EC system on site for a full year. As a matter of consequence, we also deployed every week funnels at seven stations (4811 measurements) in order to explore the spatial and temporal variability of ebullition, and to identify its controlling factors.

15 The volume of gas emitted V_{EB} averaged 1205 mL m⁻² d⁻¹ and ranged from 0-17587 mL m⁻² d⁻¹ 16 ¹. The positively skewed hyperbolic secant distributions (α = 782.41 and μ = 1205; Fig. 6a) of 17 V_{EB} showed that for most of the records of ebullition (~97%), V_{EB} was below 2000 mL m⁻² d⁻¹ 18 ¹. The V_{EB} in the WW season (median = 732 mL m⁻² d⁻¹) was statistically different (p < 19 0.0001, Kruskal Wallis test) and almost two times lower than in the WD (median = 1330 mL 20 m⁻² d⁻¹) and CD (median = 1254 mL m⁻² d⁻¹) seasons.

21 CH₄ concentration in the bubbles ranged from 0.001 to 69.2% and was most of the time lower than 30% (Fig. 6b). The average concentration was 14.9% that is two to six-fold lower than 22 23 the concentrations reported for subarctic lakes $(34.8 \pm 25.2\%)$, Wik et al., 2013), Siberian 24 thermokarst lakes ($82 \pm 7\%$, Walter et al., 2008), open water and vegetated sites in a beaver pond (47.2 \pm 20.8% and 26.6 \pm 12.4%; Dove et al., 1999) and tropical reservoir (59 - 66%, 25 26 DelSontro et al., 2011). However, the mean CH₄ concentration in bubbles at NT2 is similar to 27 the concentration observed in rive paddies and vegetated wetlands (Rothfuss and Conrad, 28 1992; Frenzel and Karofeld, 2000; Kruger et al., 2002; Chanton et al., 1989; Tyler et al., 29 1997) and well-oxygenated streams (Crawford et al., 2014). The CH₄ concentration in bubbles 30 in these ecosystems are supposed to be low because of a high methanotrophic activity in the rizhosphere of the vegetation permitted by a high ventilation of the soils by active transport of 31 32 air through the stems of the vegetation. In the NT2 reservoir, there is almost no aquatic

1 vegetation rooted in the littoral zone of the reservoir. However, the reservoir floods very 2 compacted soils. As a consequence, bubbles might develop close to the flooded soils/sediment-water interface. The area were bubbles were collected has a maximum depth 3 4 close to the depth of the oxycline most of the year (4-7m) which implies that the first 5 millimetres of the flooded soils are probably oxygenated in the area shallower than 10 m. In addition, during the lake overturn in the CD season and during the sporadic destratification 6 7 events in the WW season, O₂ could reach the flooded soil-water interface. Therefore, CH₄ 8 oxidation could affect the CH₄ concentration in bubbles in the flooded soils before they 9 escape leading to low concentration of CH_4 in bubbles. The statistical test (p < 0.0001, 10 Kruskal Wallis test) suggested that the CH₄ concentrations in bubbles differed significantly 11 seasonally with CH₄ concentrations 3-5 times higher during the WD season ($19.27 \pm 12.43\%$) 12 when the oxygen penetration in the water column is minimum than during the WW (7.30 \pm 13 8.78%) and CD ($4.57 \pm 5.78\%$) seasons. In addition to a potential role of CH₄ oxidation, the 14 effect of temperature on both the CH₄ solubility and the methanogenesis might have 15 influenced the seasonal variations of the CH₄ bubble content. The bubble CH₄ concentration 16 in the WW season ranged from 0.001 to 49% and was similar whatever the depth of the water 17 column (p = 0.08, Kruskal Wallis test), whereas bubble CH₄ concentrations differed among 18 depth zones in WD and CD seasons (p < 0.0001 (WD) and p = 0.0054 (CD)). In WD season, 19 the bubble CH_4 concentration was two times higher in the shallowest (median = 21.52%) than 20 in the deepest zones (12.78%). According to McGinnis et al. (2006) and Ostrovsky et al., 21 (2008), the decrease in the CH₄ concentration in bubbles by the dissolution of CH₄ for a 22 maximum water depth of 10 m can reach up to 20%. Therefore, this process could explain the 23 variation of CH₄ concentration in bubbles according to depth. Overall, we show that the CH₄ 24 concentration in bubbles vary seasonally and spatially by 5 orders of magnitude at the NT2 25 Reservoir, suggesting that precise extrapolation of the ebullition must take into account both 26 the volume of gas released by the sediments at high resolution (e.g., DelSontro et al., 2010), 27 but also the high variability of CH₄ concentration in the bubbles.

Like V_{EB} and bubble CH₄ concentrations, E_{FUN} fluxes varied by five orders of magnitude at the NT2 Reservoir, and showed a large variability (coefficient of variation = 122%). However, E_{FUN} distribution shows that 95% of the ebullition records were below 25 mmol m⁻² d⁻¹ (Fig. 6c). On average, ebullition was 8.5 ± 10.5 mmol m⁻² d⁻¹ and ranged from 0 - 201.7 mmol m⁻² d⁻¹. At NT2, ebullition is in the lower range of ebullition reported for tropical reservoirs (Abril et al., 2005; DelSontro et al., 2011; dos Santos et al., 2006; Galy-Lacaux et al., 1999; Keller and Stallard, 1994). Ebullition was ten times higher in the WD season

(median = 7.9 mmol $m^{-2} d^{-1}$) than in the WW (median = 0.81 mmol $m^{-2} d^{-1}$) and CD season 1 (median = 1.3 mmol $m^{-2} d^{-1}$) (Fig. 7a). This might be related to the potential dependency of 2 CH₄ solubility and production on temperature, and to the dependency of ebullition on water 3 4 depth and change in water depth as explained before. Ebullition from flooded dense forest, 5 degraded forest and agricultural lands was similar during the WW and CD seasons (p = 0.1077 (WW) and p = 0.2324 (CD), Kruskal Wallis test; Fig. 7b) but slightly lower in the 6 dense forest (median = 6.46 mmol $m^{-2} d^{-1}$) than in the degraded forests (median = 8.3 mmol 7 $m^{-2} d^{-1}$) and agricultural lands (8.63 mmol $m^{-2} d^{-1}$) during the WD season. The ebullition 8 9 dependency on water depth varies with season (Fig. 7c). Ebullition decreases significantly 10 with depth in the WD season whereas that decrease was not significant for the low emissions 11 of the CD season. This implies that the annual extrapolation of ebullition must account for the 12 seasonal evolution of the ebullition versus depth relationship.

13 **4.5** Controlling factors on CH₄ ebullition (E_{FUN})

14 According to our results on short term variation of ebullition obtained from EC and previous 15 works based on both EC and funnels, ebullition fluxes were plotted against water level, water 16 level change, specific water level change, atmospheric pressure, change in atmospheric pressure and bottom temperature. The high scatter in different regressions between ebullition 17 18 and the controlling factors is likely due to the fact that ebullition is controlled by a 19 combination of all those factors (Fig.8). Effect of both water depth and the atmospheric 20 pressure were combined by calculating the total static pressure (TSP) and the change in TSP 21 in mH₂O at the bottom of the reservoir, which is the sum of atmospheric and hydrostatic 22 pressure changes. These two parameters were then correlated with ebullition using an exponential decrease regression model (Fig. 8f, g). 23

Ebullition decreased from 203 to 0 mmol $m^{-2} d^{-1}$ for water depth ranging from 0.4 to 16 m 24 (Fig. 8a). The median of all fluxes measured at shallow sites (0.4-3 m: 6.3 mmol $m^{-2} d^{-1}$) was 25 almost twofold higher than the median in the deepest zone (6-16 m: 2.9 mmol $m^{-2} d^{-1}$). The 26 correlation between depth and ebullition is highly significant (p < 0.0001), but still, this 27 parameter alone only explains 4% of the variation of the ebullition ($r^2 = 0.04$; Fig. 8a). The 28 29 dependency of ebullition on the depth could be attributed to two physical processes. First, a 30 deeper water column means higher hydrostatic pressure, which could prevent the formation of bubbles by increasing CH₄ solubility in the sediment pore waters. Second, while the CH₄ 31 32 bubbles escape the sediment, bubbles partly dissolve in the water on their way up to the atmosphere (DelSontro et al., 2010; McGinnis et al., 2006). The percentage of CH₄
dissolution and thereby oxidation increase with the water depth. On the opposite, shallow
zones favour bubble formation because they are generally warmer which both stimulates
methanogenesis (Duc et al., 2010) and makes CH₄ less soluble (Yamamoto et al., 1976).

5 As seen above, water depth has an impact on ebullition, but it appears that water depth change (or hydrostatic pressure change) has a stronger effect ($r^2=0.23$) on this phenomenon. Water 6 depth and hydrostatic pressure decrease trigger ebullition as demonstrated here (Fig. 8b), in 7 8 previous works in marine and estuarine environments, and in freshwater wetlands (Boles et 9 al., 2001; Chanton et al., 1989; Engle and Melack, 2000; Martens and Klump, 1980). During the periods of falling water level, ebullition was fivefold higher (median = 7.5 mmol $m^{-2} d^{-1}$) 10 than the ebullition during increasing water level (median = $1.5 \text{ mmol m}^{-2} \text{ d}^{-1}$). The correlation 11 12 shows that the change in the water level alone explains 23% of the ebullition variability and it 13 evidences why ebullition is significantly higher during the WD season when the water level is 14 falling (negative water level change) than during the WW season when the water level is rising or during the CD season when it is stable. The effect of the specific water level change 15 on ebullition (Fig. 8c) is not as high as expected (p < 0.0001, $r^2 = 0.13$) on this large dataset of 16 funnel measurements encompassing a wider range of environmental conditions and flooded 17 18 ecosystems compared to what we obtained with the EC-derived data only. However, for a 19 given water depth, water depth change and specific water level change, ebullition was in the 20 same range whatever it was obtained from EC or funnels. We hypothesize that EC installed in 21 a zone with a very homogeneous land cover (corresponding to flooded agricultural lands) and 22 covering a large footprint allows to better characterize the controlling factors than discrete sampling with funnels over a few cm^2 in various type of flooded ecosystems. 23

The relationship of ebullition obtained with funnels over 24 hours versus atmospheric pressure and pressure change were highly significant (Fig. 8d, e) but with very low determination coefficients. These much lower r^2 values compared to the one obtained from the EC could be explained by the fact that mean atmospheric pressure change from one day to the other is smaller than the diurnal variations of atmospheric pressure that we observed during the EC deployments.

The magnitude of the atmospheric pressure varied within a small range (9.55 - 9.70hPa), or an equivalent of 0.15 mH₂O). As a matter of consequences, our attempt to combine the effect of hydrostatic and atmospheric pressure (i.e. the so-called Total Static Pressure or TSP) was not highly convincing since we did not improve the correlation coefficient using the TSP (Fig. 8f,
g) compared to what we observed for the hydrostatic pressure alone.

3 Finally, we found a very low correlation between ebullition and reservoir bottom temperature $(r^2 = 0.03, Fig. 8h)$. This shows that, given the hydrodynamics and the temperature range 4 experienced in the NT2 Reservoir, that this physical parameter has a very low predictive 5 6 power. This is due to the co-variation of several factors at the same time hiding a possible 7 effect of temperature on the benthic methanogenesis activity. The absence of correlation 8 between temperature and ebullition is mostly due to the fact that the highest bottom water 9 temperatures were often synchronized with the beginning of the WW season when the 10 ebullition is moderated by the water level increase. This illustrates the complexity of 11 controlling factors interacting at the same time, and one with each other in a non-linear way. 12 As a matter of fact, it is worth trying a non-linear method to represent ebullition through 13 several relevant parameters, identified in this section but not necessarily highly correlated 14 with ebullition.

15 **4.6** Extrapolation of ebullition at the NT2 Reservoir scale by ANN

The extrapolation of ebullition from field measurements to the whole NT2 aquatic ecosystems is challenging. In all studies published so far, the average ebullition is multiplied by the surface area of the shallow zone where ebullition was measured (e.g., Abril et al., 2005; Wik et al., 2013), by type of habitat (e.g., Smith et al., 2000) or by a combination of the two approaches (DelSontro et al., 2011). Our dataset together with the determination of some major controlling factors of ebullition allowed us an attempt for the first time of the extrapolation of this major CH_4 pathway based on physical processes.

23 As a first approach we used multi linear regressions. We obtained good correlations with the 24 change in the total static pressure. However, we were able to explain only 21% of the variance of the ebullition fluxes (data not shown). The relatively low percentage of explained variance 25 26 revealed that the complexity of the interactions between the controlling factors of the 27 ebullition is only partially resolved through simple linear equations. A non-linear approach 28 was used to model ebullition fluxes using an ANN. Taking into account that controlling 29 factors are integrators of several parameters, as shown in the previous section via analyses with TSP, change in TSP, and bottom water temperature, the ANN model resulted in much 30 better agreement between calculate and measured ebullition fluxes ($r^2 = 0.46$, p < 0.0001; Fig. 31 S5). Indeed, a step-by-step study with the ANN revealed that the non linear equation with one 32

input parameter (Total change in TSP) gives an $r^2=0.26$. Two input parameters (Total change 1 in TSP and TSP) gives $r^2=0.39$. The addition of bottom temperature leads to the best result of 2 $r^2=0.46$. The daily time series of the bottom reservoir temperature and atmospheric pressure 3 are shown in Fig. 9a, and the estimated area-weighted modelled ebullition fluxes together 4 5 with the measurements at the NT2 Reservoir from January 2009 till August 2013 are shown on Fig. 9b. Over the span of this study, ebullition remained unexpectedly constant whereas 6 7 total emissions from hydroelectric reservoirs are known to decrease with time (Abril et al., 8 2005; Barros et al., 2011) due to the exhaustion of the source of organic matter fuelling the 9 emissions. The modelled ebullition flux (Fig. 8b) exhibits large seasonal peaks (25.9 ± 9.3) mmol $m^{-2} d^{-1}$) at the transition between the CD and WD seasons. The peaks are anti-correlated 10 11 with the water level variations (Fig. 9b), and occur during the periods when atmospheric 12 pressure is decreasing and water temperature increasing. Due to the high seasonal variations 13 simulated by ANN, 50% of the CH₄ emitted by the NT2 Reservoir each year is released 14 within 4 months even if this period corresponds to the lowest surface of the reservoir. On a 15 yearly basis, ebullition obtained from ANN would represent 60-80% of total emission 16 (diffusion and ebullition) at the surface of the NT2 Reservoir. This further supports the idea 17 that the estimate of ebullition from an aquatic ecosystem with large water level variations 18 requires high frequency measurements over the period of falling water level. This period 19 corresponds to a hot moment of emissions since the water level as well as its variations and 20 the concomitant temperature variations have a strong impact on ebullition and ultimately on 21 total emissions.

22 The ANN model allowed us to simulate the ebullition over a 4-year period by using a few 23 basic meteorological and limnological input data and a one-year intensive monitoring of CH₄ 24 ebullition. This approach constitutes a powerful gap-filling tool allowing the obtaining of past 25 and future ebullition time series for ecosystems in steady state like natural wetlands and lakes 26 receiving constant amount of organic matter from the watershed and under the influence of 27 constant meteorological forcing. However, in the case of an hydroelectric reservoirs, this 28 approach must be taken with caution and can only be applied during short periods of time 29 when the evolution of ebullition is not significant as it is the case for NT2 during our study or 30 once it reaches its steady state (4-15 years after flooding; Abril et al. 2005; Teodoru et al., 31 2012).

1 5 Conclusions

Using a set of classical techniques for the discrete measurements of CH_4 diffusive (FC) and ebullition (funnels), and the recently developed EC techniques for the measurement of total CH_4 emissions over large surfaces, we confirmed that the EC system is able to capture continuously and at a 30 min-frequency the two main pathways of CH_4 release in inland aquatic ecosystems.

The EC system captured a diurnal bimodal pattern of CH₄ emissions following semi-diurnal 7 8 variation of the atmospheric pressure. Daily atmospheric air pressure drops during all seasons 9 and whatever the depth of the water column, triggers CH₄ ebullition, resulting in a first 10 maximum of CH₄ emissions in the middle of the day. At night, a second and moderate peak of 11 CH₄ emission was recorded due to the combination of a smaller pressure drop and a potential 12 enhancement of the diffusive fluxes because of turbulence generated by heat loss. This might 13 be a common feature in wetlands where the methanogenesis is active enough to induce a 14 storage of CH₄ in the sediments or flooded soils. This diurnal pattern implies that precise 15 estimate of CH₄ emissions from aquatic ecosystems require high frequency measurements 16 over 24h in order to capture the daily hot moments of emissions that could contribute up to 17 50% of daily emissions in a few hours.

We have shown that both the concentration of CH_4 in the bubbles reaching the atmosphere and the volume of bubbles are highly variable. The concentration of CH_4 in bubbles exhibited a high seasonality suggesting that estimate of ebullition cannot be done focusing on the volume of bubbles reaching the atmosphere assuming a predetermined concentration of CH_4 in the bubbles for the whole reservoir and all the seasons.

23 The CH₄ ebullition mostly depends on the water level and air pressure variations. The use of 24 these linear regressions did not allow a realistic extrapolation of the flux for the entire 25 reservoir (data not show). This is because of the potential non-linearity of the processes and 26 the complexity of the interactions between the controlling factors. Non-linearity was taken 27 into account using an ANN model with total static pressure, change in total static pressure, 28 and bottom temperature as input parameters. ANN model was able to explain 46% of 29 variation in ebullition CH₄ fluxes, and to perform gap-filling for the ebullition fluxes over a 30 four-year period (2009-2013). Our results clearly showed a very high seasonality with 50% of 31 the yearly CH₄ ebullition occurs within four months of the WD season although the surface 32 water area of the reservoir is at its minimum during this period. Overall, ebullition contributed 33 60-80% of total emissions at the surface of the reservoir (disregarding downstream emissions). Our results on ebullition in this recently flooded reservoir together with the only
other results available in tropical hydroelectric reservoirs (Petit Saut Reservoir, French
Guiana; Abril et al., 2005; Galy-Lacaux et al., 1997) during the first year after impoundment
suggest that ebullition is a major and overlooked pathway in young tropical or subtropical
hydroelectric reservoirs.

6

7 Acknowledgements

Authors are pleased to express their gratitude to Electricité de France (EDF), Nam Theun 2
Power Company Ltd (NTPC) and Hydro-Québec for funding and contribution to the logistic
and all team members of the Aquatic and Environmental Laboratory (AE Lab) for their kind
support during field campaigns. CD benefited from a PhD grant by EDF.

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	Diffusion D		Ebullition E		Diffusion+Ebullition DE			
	Method	Average \pm SD (n)	Method	Average \pm SD (n)	Method	Average \pm SD (n)	Method	Average \pm SD (n)
May-09	D_{GC}^{1}	1.2 ± 0.8 (12)	E_{FUN}^2	1.6±2.9 (9)	$\mathrm{DE_{EC}}^3$	6.5±3.3 (39)	DE _{EC}	6.9±2.6 (2)
	${\rm D_{GA}}^4$	NA	E_{GA}^{5}	NA	${\rm DE_{GC}}^6$	1.9±2.3 (16)	$E_{FUN}+D_{GC}$	2.8±1.6 (2)
	${\rm D_{TBL}}^7$	1.5 ± 2.2 (14)					$E_{FUN} \!\!+\! D_{TBL}$	3.1±1.7 (2)
Mar-10	D _{GC}	0.9±0.5 (9)	E _{FUN}	NA	DE _{EC}	5.8±5.0 (138)	DE _{EC}	5.7±3.7 (14)
	D_{GA}	NA	E _{GA}	NA	DE _{GC}	8.4±17.5 (24)		
	D_{TBL}	1.3 ±0.8 (12)						
Mar-11	D _{GC}	NA	E _{FUN}	4.2±3.6 (95)	DE _{EC}	7.2±2.9 (105)	DE _{EC}	7.2±0.8 (4)
	D_{GA}	1.9±1.2 (28)	EGA	4.6±7.1 (30)	DE _{GC}	8.9±10.5 (58)	$E_{FUN}+D_{GA}$	6.1±1.2 (4)
	D_{TBL}	1.1±2.0 (52)					E_{FUN} + D_{TBL}	5.3±1.2 (4)
Jun-11	D _{GC}	1.5 (1)	E _{FUN}	28.0±11.0 (126)	DE _{EC}	29.1±16.4 (133)	DE _{EC}	26.6±6.7 (5)
	D_{GA}	NA	E _{GA}	NA	DE _{GC}	54.3±35.0 (21)	E_{FUN} + D_{TBL}	29.9±5.5 (5)
	D_{TBL}	1.9±2.5 (19)						
All	D _{GC}	1.1±0.7 (22)	E _{FUN}	17.1±14.7 (230)	DE _{EC}	13.6±14.5 (415)	DE _{EC}	16.0±11.1 (11)
	D_{GA}	1.9±1.2 (28)	E _{GA}	4.6±7.1 (30)	DE _{GC}	15.8±25.2 (121)	$E_{FUN}+D_{FC}$	16.3±13.4 (11)
	D_{TBL}	1.4 ±2.0 (97)					E_{FUN} + D_{TBL}	16.3±13.8 (11)

Table 1. Comparison of different methods to assess CH₄ emissions. All fluxes are in mmol $m^{-2} d^{-1}$ (average ± standard

deviation) and number of measurements (n) given between brackets.

- D_{GC}^{1} : Diffusion from floating chamber (FC) and post-analysis with gas chromatography E_{FUN}^{2} : Ebullition from submerged funnel

- DE_{EC}^{3} : Total emissions measured by eddy covariance D_{GA}^{4} : Diffusion from FC and in situ gas analyser E_{GA}^{5} : Ebullition from FC and in situ gas analyser DE_{GC}^{6} : Total emissions by FC (diffusion + ebullition) affected by bubbling D_{TBL}^{7} : Diffusion calculated by thin boundary layer (TBL) method from surface CH₄ concentrations

	May-09	Mar-10	Mar-11	Jun-11	All
	D _{GC}				
D _{TBL}	0.6027	0.2815	0.0513	-	0.5049
	DE _{EC}				
DE _{GC}	< 0.0001	0.0129	0.1075	< 0.0001	0.0004
$E_{FUN} + D_{GA}$			0.2021		
E_{FUN} + D_{GC}	0.2222	-	-	-	0.5114
E _{FUN} +D _{TBL}	0.2533	-	0.057	0.8413	0.3933

1 Table 2. Statistical test for the comparison of different methods to assess CH_4 emissions. Difference is significant if p < 0.05

Figure captions

Figure 1. Map of the Nam Theun 2 Hydroelectric Reservoir (Lao People': Republic) showing (1) the land cover before flooding (from Descloux et al., 2 dense forest, M: medium forest, L: light forest, DG: degraded forest, R:ripar agricultural soils, S: swamps, O: others and W: water and (2) the location of measurements and the eddy covariance site.

Figure 2. Time series of (a) atmospheris temperature and rainfall rates and (b) th 2 Reservoir water level during the study. The grey bars and shaded area indi experiments and the ebullition monitoring, rspectively. The double arrows seasons (WD: warm dry; WW: warm wet; CD: cold dry).

Figure 3. Inter-comparison of the estimates of CH₄ emissions obtained using 1 methods deployed during the four field campaigns. See text for details about diff the figure.

Figure 4. CH_4 emissions measured by eddy covariance (DE_{EC}) versus (a) wa change in water depth, and (c) specific water level change for the four field can that turbines were not started in May 2009, leading to no water level change du campaign.

Figure 5. (a, b, c, d) 30 min-binned CH₄ emissions measured by eddy covar (circle) and 30 min-binned atmospheric pressure (cross) for the four field campa 30 min-binned buoyancy flux (note that June 2011 data are not available) individual 30-min CH₄ emissions measured by eddy covariance (DE_{EC}) versus atmospheric pressure for the four field campaigns, (l, m, n, o) night and dayti CH₄ emissions measured by eddy covariance (DE_{EC}) for the four field campaig y-axis scale differs for the June 2011 campaign (d, k, o).

Figure 6. Histograms showing the distribution of (a) ebullition rate, (b) concentration, and (c) ebullition measured by funnels.

Figure 7. Ebullition measured by funnels for (a) the three different seasons, (b) the three major different flooded ecosystems (dense/medium forest, light/degraded forest, and agricultural land), and (c) three depth zones.

Figure 8. Ebullition measured by funnels versus (a) water depth, (b) change in water level, (c) specific water level change, (d) atmospheric pressure, (e) change in atmospheric pressure, (f) total static pressure, (g) change in total static pressure, and (h) reservoir bottom temperature.

Figure 9. Time series of the (a) reservoir bottom temperature and atmospheric pressure and, (b) funnels measured and ANN modelled ebullition fluxes along with reservoir water level. In panel b, boxes show the median concentration and the interquartile range, and whiskers denote the full range of all values. Plus sign (+) in the box is showing the mean value.



Figure 1. Map of the Nam Theun 2 Hydroelectric Reservoir (Lao People's Democratic Republic) showing (1) the land cover before flooding (from Descloux et al., 2011) with D: dense forest, M: medium forest, L: light forest, DG: degraded forest, Ririparian forest, A: agricultural soils, S: swamps, O: others and W: water and (2) the location of the ebullition measurements and the eddy covariance site.



Figure 2. Time series of (a) atmospheris temperature and rainfall rates and (b) the Nam Theun
2 Reservoir water level during the study. The grey bars and shaded area indicate the field
experiments and the ebullition monitoring, rspectively. The double arrows indicate the
seasons (WD: warm dry; WW: warm wet; CD: cold dry).



Figure 3. Inter-comparison of the estimates of CH₄ emissions obtained using the variety of methods deployed during the four field campaigns. Note that no ebullition was measured in March 2010. D_{GC}: Diffusion from floating chamber (FC) and post-analysis with gas chromatography, D_{TBL}: Diffusion calculated by thin boundary layer (TBL) method from surface CH₄ concentrations, D_{GA}: Diffusion from FC and in situ gas analyser, E_{FUN}: Ebullition from submerged funnel, E_{GA}: Ebullition from FC and in situ gas analyser, DE_{EC}: Total emissions measured by eddy covariance, DE_{GC}: Total emissions by FC (diffusion + ebullition) affected by ebullition.



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Figure 4. CH_4 emissions measured by eddy covariance (DE_{EC}) versus (a) water depth, (b) change in water depth, and (c) specific water level change for the four field campaigns. Note that turbines were not started in May 2009, leading to no water level change during that field campaign. In all panels, the solid line is the regression line and the dash lines represent the confidence interval



Figure 5. (a, b, c, d) 30 mn-binned CH₄ emissions measured by eddy covariance (DE_{EC}) (circle) and 30 mn-binned atmospheric pressure (cross) for the four field campaigns, (e, f, g); 30 mn-binned buoyancy flux (note that June 2011 data are not available), (h, i, j, k) individual 30-min CH₄ emissions measured by eddy covariance (DE_{EC}) versus change in the atmospheric pressure for the four field campaigns, (l, m, n, o) night and daytime range for CH₄ emissions measured by eddy covariance (DE_{EC}) for the four field campaigns. Note that y-axis scale differs for the June 2011 campaign (d, k, o). In panels e-h, the solid line is the regression line and the dash lines represent the confidence interval



Figure 6. Histograms showing the distribution of (a) ebullition rate, (b) CH₄ bubble
concentration, and (c) ebullition measured by funnels.





3 Figure 7. Ebullition measured by funnels for (a) the three different seasons, (b) the three 4 major different flooded ecosystems (dense/medium forest, light/degraded forest, and 5 agricultural land), and (c) three depth zones.



Figure 8. Ebullition measured by funnels versus (a) water depth, (b) change in water level, (c) specific water level change, (d) atmospheric
pressure, (e) change in atmospheric pressure, (f) total static pressure, (g) change in total static pressure, and (h) reservoir bottom temperature.



Figure 9. Time series of the (a) reservoir bottom temperature and atmospheric pressure and, (b) funnels measured and ANN modeled ebullition fluxes along with reservoir water level. In panel b, boxes show the median concentration and the interquartile range, and whiskers denote the full range of all values. Plus sign (+) in the boxes of panel a and b show the mean value.