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ABSTRACT

To evaluate the hypothesis that plant-mediated oxygen supplies decrease methane (CH$_4$) production and total global warming potential (GWP) in a tropical peatland, the authors compared the fluxes and dissolved concentrations of greenhouse gases (GHGs; CH$_4$, carbon dioxide (CO$_2$), and nitrous oxide (N$_2$O)) and dissolved oxygen (DO) at multiple peatland ecosystems in Central Kalimantan, Indonesia. Study ecosystems included tropical peat swamp forest and degraded peatland areas that were burned and/or drained during the rainy season. CH$_4$ fluxes were significantly influenced by land use and drainage, which were highest in the flooded burnt sites (5.75±6.66 mgC m$^{-2}$ h$^{-1}$) followed by the flooded forest sites (1.37±2.03 mgC m$^{-2}$ h$^{-1}$), the drained burnt site (0.220±0.143 mgC m$^{-2}$ h$^{-1}$), and the drained forest site (0.0084±0.0321 mgC m$^{-2}$ h$^{-1}$). Dissolved CH$_4$ concentrations were also significantly affected by land use and drainage, which were highest in the flooded burnt sites (124±84 µmol L$^{-1}$) followed by the drained burnt site (45.2±29.8 µmol L$^{-1}$), the flooded forest sites (1.15±1.38 µmol L$^{-1}$), and the drained forest site (0.860±0.819 µmol L$^{-1}$). DO concentrations were influenced by land use only, which were significantly higher in the forest sites (6.9±5.6 µmol L$^{-1}$) compared to the burnt sites (4.0±2.9 µmol L$^{-1}$). These results suggested that CH$_4$ produced in the peat might be oxidized by plant-mediated oxygen supply in the forest sites. CO$_2$ fluxes were significantly higher in the drained forest site (340±250 mgC m$^{-2}$ h$^{-1}$ with the water table level of −20 to −60 cm) than in the drained burnt site (108±115 mgC m$^{-2}$ h$^{-1}$ with the water table level of −15 to +10 cm). Dissolved CO$_2$ concentrations were 0.6 to 3.5 mmol L$^{-1}$, also highest in the drained forest site. These results suggested the enhanced CO$_2$ emission by aerobic peat decomposition and plant respiration in the drained forest site. N$_2$O fluxes ranged from −2.4 to −8.7 µgN m$^{-2}$ h$^{-1}$ in the flooded sites and from 3.4 to 8.1 µgN m$^{-2}$ h$^{-1}$ in the drained sites. The negative N$_2$O fluxes might be caused by N$_2$O consumption by denitrification under flooded conditions. Dissolved N$_2$O concentrations were 0.005 to 0.22 µmol L$^{-1}$ but occurred at <0.01 µmol L$^{-1}$ in most cases. GWP was mainly determined by CO$_2$ flux, with highest levels in the drained forest site. In spite of almost the same CO$_2$ flux, GWP in the flooded burnt sites was 20% higher than that in the flooded forest sites due to the large CH$_4$ emission (not significant). N$_2$O fluxes made little contribution to GWP.

Key words: Dissolved oxygen (DO), greenhouse gas (GHG), Indonesia, tropical peatland, and water table level.
1. INTRODUCTION

Tropical peatland ecosystems in Southeast Asia cover 24.8 million hectares, comprising 56% of the tropical and 6% of the global peatland area. This ecosystem’s high carbon density allows for a large regional peat carbon store of 68.5 Pg, equivalent to 77% of the tropical and 11–14% of the global peat carbon store (Page et al., 2011), which is comparable to the total fossil fuel emissions released throughout the world over 9 years (van der Werf et al., 2008). Given the size of this carbon reservoir, tropical peatlands play a critical role as a global sink for gaseous carbon. Inversely, if environmental conditions (e.g., land use type, groundwater level) change, these peatlands may become a major source of gaseous carbon and greenhouse gases (GHGs), such as CO$_2$, CH$_4$ and N$_2$O.

In recent decades, considerable portions of peatland in Southeast Asia have been reclaimed to support agroforestry and rice paddy development. The deforestation and development of peatlands is usually accompanied by the draining of water from these areas, releasing much carbon through aerobic peat decomposition. In addition, the water table level can drastically decrease during exceptionally long dry periods, which frequently occur in El Niño and Southern Oscillation (ENSO) years (Hirano et al., 2007; 2012; van der Werf et al., 2008).

Several trials have been conducted to restore lowland tropical peatlands (Page et al., 2009). For example, restoration efforts have included raising and stabilizing the water table level by blocking drainage canals, which reduces aerobic peat decomposition in the flooded peatlands. However, flooding can also increase anaerobic CH$_4$ production. Soil denitrification, which is a major contributor to N$_2$O production, is also generally promoted under reduced conditions. The radiative forcing effects of CH$_4$ and N$_2$O are 25 and 298 times higher, respectively, than that of CO$_2$ (over a 100-yr time horizon; IPCC, 2007). Consequently, the increases in the production of CH$_4$ and N$_2$O under flooded conditions could possibly offset decreases in aerobic peat decomposition and associated CO$_2$ emissions. A large portion of previously drained peatland has lost its original vegetation and surface layer through deforestation and/or fire. Under such conditions, the recovery of the water table level will not be able to restore ecosystem function and carbon balance to those associated with original vegetation levels. The change in GHG dynamics in tropical peatland soils after drainage, deforestation, peat fire, and re-wetting has not been fully studied to date.
After reviewing the existing data, Couwenberg et al. (2010) have demonstrated that the range of \( \text{CH}_4 \) fluxes in tropical peatlands was one-fifth that observed in temperate and boreal peatlands. They attributed low \( \text{CH}_4 \) emissions to the poor quality of woody tropical peat, which contains higher levels of recalcitrant materials (e.g., lignin). Williams and Yavitt (2010) reported that the biochemical compositions of lignin affect soil methanogenesis. Another explanation involved the oxygen supply from plant roots. Vascular plants in wetlands usually possess special morphological adaptations, such as hypertrophied lenticels, adventitious roots and enlarged aerenchyma (Joabsson et al. 1999), which promote gas exchange between the atmosphere and the rhizosphere and allow for the entry of oxygen to the root zone (Kozlowski, 1997; Megonigal and Day, 1992). According to a recent review by Laanbroek (2010), the percentage of \( \text{CH}_4 \) oxidized before entering the atmosphere ranges from 0 to up to >90% of the potential \( \text{CH}_4 \) efflux. Several studies conducted in temperate and boreal wetlands reported that an increase in measurement depth was associated with a decrease in the amount of root tissue and dissolved oxygen (DO) concentrations and an increase in dissolved \( \text{CH}_4 \) concentrations (van der Nat and Middelburg, 1998; Fritz et al., 2011; Liebner et al., 2012). Thus, if plant-mediated oxygen transport and \( \text{CH}_4 \) oxidation also occur in tropical peatlands, significant differences in soil \( \text{CH}_4 \) fluxes, dissolved \( \text{CH}_4 \) concentrations, and DO concentrations in peat pore water would be expected between vegetated and non-vegetated lands.

Given these relationships, we hypothesized that 1) DO supplied from plant roots should oxidize \( \text{CH}_4 \) produced in saturated tropical peat soil and 2) \( \text{CH}_4 \) oxidation should reduce total GWP in a tropical peat swamp forest. To evaluate these hypotheses, we compared DO and GHG fluxes and dissolved concentrations in an ecosystem containing submerged forest vegetation with those in degraded peatland areas that have been burned and/or drained during the rainy season.

2. MATERIALS AND METHODS

2.1. Study site
This study was conducted in a tropical peat swamp area located near Palangka Raya City (02° 19’ 25.20” S, 113° 54’ 16.86” E), Central Kalimantan Province, Indonesia. Most of the peat
swamp forest in this region was previously deforested and drained during a national project (i.e., the Mega Rice Project) in the late 1990s. Although the project was terminated in 1999, it left vast tracts of devastated peatland (Notohadiprawiro, 1998; Muhanmad and Rieley, 2002). A large portion of the devastated peatland was subsequently abandoned because it was unsuitable for the intended agricultural production. In these areas, several large fires have occurred during the dry season, severely damaging both the vegetation and soil. These fires were mainly ignited as a result of agricultural activities, including land clearance and ash production by local farmers. Some of these fires spread uncontrollably to become wildfires (Usup et al., 2004).

Six locations were established as study sites. Three sites were placed in a peat swamp forest that was not affected by deforestation or fire (FW1, FW2, and FD; forest sites). The other three sites were located in a degraded peatland affected by fire (BW1, BW2, and BD; burnt sites). The burnt sites were deforested in 1995 and burned in 1997, 2002 and 2009 during El Niño years. The flooded forest sites (FW1 and FW2) and the flooded burnt sites (BW1 and BW2) were close to the flood plains of the Sebangau and Kahayan Rivers, respectively. Sites FD and BD were located in the middle of a peat dome formed between these two rivers. These two sites are influenced by a large drainage canal, which runs through the peat dome southward to the Java Sea (>100 km long).

In the forest sites, the ground surface was covered by a dense canopy of peat swamp forest, although the composition of tree species was previously affected by selective logging until the late 1990s (Hirano et al., 2012). Dominant species present included *Tristania whittiana*, *Combretocarpus rotundatus*, *Palaquium leiocarpum*, and *Stemonurus scorpioides*. In the burnt sites, the ground surface was covered by ferns and grasses, such as Kalakai (*Stenochlaena palustris*) and Pakis/Paku-pakuan (*Nephrolepis* sp.), with patchy depressions. Trees, such as Tumih (*Combretocarpus rotundatus*), were found sparsely. Detailed information about the vegetation around the study sites is provided elsewhere (Jauhiainen et al., 2005; Page et al., 1999; Sundari et al., 2012; Takakai et al., 2006; Tuah et al., 2001; 2003). Based on the USDA Soil Taxonomy, the study sites were all classified as Typic Tropofibrists (Takakai et al., 2006). The peat depths were 2–3 m at FW1 and FW2 and about 4 m at the other sites. The volumetric carbon density of the peat was 71.5±17.3 kg m$^{-3}$ (Shimada et al., 2001). Annual precipitation and annual mean air temperature over the nine study years (2002–2010) were 2540±596 mm and 26.2±0.3°C, respectively (Sundari et al., 2012). The dry season
normally begins in June and ends in October. Field observations were conducted during the wet season from December 2011 to May 2012.

2.2. GHG fluxes

Greenhouse gas fluxes at the ground surface were measured monthly using a closed-chamber method (Morishita et al., 2003; Takakai et al., 2006; Toma and Hatano, 2007). A set of cylindrical stainless steel chambers (18.5–21 cm in diameter and 25 cm in height), each with an acrylic lid fixed to a sample collector and a pressure-regulating bag, was used for the flux measurement. Stainless steel collars were permanently planted at three locations at each site. First, 400 mL of the air inside the chamber was taken into a Tedlar® bag as the sample at 0 min before closing the chamber lid. The chamber was then quickly capped with the lid. After 6 min, 250 mL of the air inside the chamber was transferred into another Tedlar® bag. CO₂ concentrations in these bags were determined on the day of sampling with a non-dispersive infrared (NDIR) analyzer (ZFP9; Fuji Electric, Japan). The lid was then removed temporarily, and the air inside the chamber was ventilated. Next, 20 mL of the air inside the chamber was periodically collected into pre-evacuated vial bottles with butyl rubber stoppers at 0, 20 and 40 min after closing the lid for CH₄ and N₂O analyses. CH₄ concentrations were determined using a gas chromatographer (GC-8A, Shimadzu, Kyoto, Japan) equipped with 1) a flame ionization detector (FID) maintained at 130°C and 2) a 2-m-long activated carbon column (80/100 mesh; Stainless column, Serial No. D-0015, Shimadzu, Kyoto, Japan) maintained at 70°C with pure nitrogen gas as a carrier. N₂O concentrations were determined with a gas chromatographer (GC-14B, Shimadzu, Kyoto, Japan) equipped with 1) an electron capture detector (ECD) maintained at 340°C and 2) a 1-m-long Porapak N column maintained at 60°C with proportional gas (5% CH₄ in Ar). We did not measure fluxes when water table levels exceeded the height of the chamber. As a result, GHG fluxes at several sites were not measured in December 2011 (FW1, BW1 and BW2), February 2012 (BW1), and March 2012 (FW1, FW2 and BW1). Due to a separate technical issue, we could not measure GHG fluxes in January 2012 for any site, with the exception of FW2.

Gas flux was calculated as the change in gas concentration inside the chamber (ΔCᵢ) against closure time (Δt) according to the following equation:

\[
F = \rho_g \times (V / A) \times (\Delta C_i / \Delta t) \times [273 / (273 + T)]
\]  

(1)
where $F$ is the gas flux (gC or gN m$^{-2}$ h$^{-1}$), $\rho_g$ is the gas density (0.536 \times 10^3$ g m$^{-3}$ for CO$\_2$-C and CH$\_4$-C; 1.259 \times 10^3$ g m$^{-3}$ for N$_2$O-N), $V/A$ is equivalent to the height of the chamber from the water or ground surface (m), and $T$ is the air temperature during the sampling period ($^\circ$C). Air temperature was measured with a digital thermometer (TESTO 625, Lenzkirch, Germany) before closing and after opening the chamber, and the mean of both temperatures was used in Eq. 1. A positive flux is indicative of gas emissions from the soil or water surface into the atmosphere, while a negative flux indicates gas uptake from the atmosphere. The total flux in global warming potential (GWP) was calculated as the CO$_2$-equivalent carbon flux (CO$_2$-eq gC m$^{-2}$ h$^{-1}$) according to the following equation:

$$\text{GWP} = F_{\text{CO}_2} + F_{\text{CH}_4} \times (16/12) \times 25 \times (12/44) + F_{\text{N}_2\text{O}} \times (44/28) \times 298 \times (12/44)$$  \hspace{1cm} (2)

where 25 and 298 are the factors used for converting CH$_4$ and N$_2$O fluxes, respectively, to their GWPs over a 100-yr time horizon (IPCC 2007); and (16/12), (44/28), and (12/44) are the conversion factors for C to CH$_4$, N to N$_2$O, and CO$_2$ to C, respectively.

### 2.3. Dissolved GHG concentrations

To determine the concentrations of GHGs dissolved in peat water, two types of in-situ headspace methods were applied. We used an in-situ equilibration chamber for CO$_2$ and a rapid equilibration method for CH$_4$ and N$_2$O. An in-situ equilibration chamber (i.e., a small chamber designed to equilibrate air with the surrounding dissolved gases) was developed according to previous studies (Faulkner et al., 1989; Yu and DeLaune, 2006). An inverted 60-mL plastic syringe was used as the chamber and was connected aboveground to a thin tube, which was equipped with a three-way stopcock at the upper end. A chamber was installed at depths of 20, 40, 60, and 80 cm. First, 60 mL of O$_2$-free air was injected through the tube, and the stopcock was closed to keep the injected air from escaping. Two to four weeks following the injection, the air was extracted from the chamber and collected into an aluminum sample bag. CO$_2$ concentrations were determined on the day of sampling according to the same method used to measure CO$_2$ flux described above. The dissolved gas concentration in pore water ($C_l$, mol mol$^{-1}$) was given by Henry’s law:

$$C_l = \frac{C_g \times 10^5 \times P}{K_H}$$  \hspace{1cm} (3)
where $C_g$ is the concentration of the sample air (ppmv), $P$ is the total pressure inside the chamber (atm), and $K_H$ is Henry’s constant (mol mol$^{-1}$ atm$^{-1}$). The effect of water pressure above the chamber was reflected in $P$. The value of $K_H$, a function of soil temperature, was cited from a chemical engineering handbook (Society of Chemical Engineers, Japan, 1999). Soil temperature was preliminarily observed at depths of 5, 10, 30 and 50 cm (see below). The average temperatures between 10 and 30 cm and between 30 and 50 cm were used as the soil temperatures at 20 and 40 cm, respectively. Soil temperatures at 60 and 80 cm were assumed to be equivalent to that at 50 cm.

The $C_i$ in the molar ratio was converted to $C_i$ (mol L$^{-1}$) according to the following equation:

$$C_i [\text{mol L}^{-1}] = \frac{C_i [\text{mol mol}^{-1}] \times \rho_w}{m_w}$$  \hspace{1cm} (4)

where $\rho_w$ is the water density (1000 g L$^{-1}$), and $m_w$ is the molecular weight of water (H$_2$O, 18.01 g mol$^{-1}$). The time required to reach CO$_2$ equilibrium was investigated beforehand at the same study sites as follows. The CO$_2$ concentrations in the air samples collected following different equilibration durations (5 to 100 days) were compared, and no statistically significant differences were observed. Therefore, CO$_2$ in the collected samples was assumed to be in equilibrium with CO$_2$ dissolved in the surrounding water.

Gas samples for the analyses of dissolved CH$_4$ and N$_2$O were obtained by rapidly equilibrating a fixed volume of air with pore water in a syringe. Water samples were collected from the same depths as those used for dissolved CO$_2$ measurements (i.e., 20, 40, 60, and 80 cm). At the beginning of field observations, a water sampler consisting of a stainless steel pipe (0.7 cm in diameter), a plastic tube and a three-way stopcock was installed at each depth. During the observation intervals, the sampler was filled with water, and the stopcock was kept closed. Before sampling, the water remaining in the sampler was removed. First, 30 mL was collected from the sampler directly into a 60-mL plastic syringe. Next, the same volume of O$_2$-free air was injected into the same syringe. The syringe was then shaken by hand for two minutes (100 times). After that, 20 mL of the air that was equilibrated with the water sample was stored in a pre-evacuated vial. The concentrations of CH$_4$ and N$_2$O in the bottles were determined in the same manner used for flux measurements for these gases. According to this
rapid equilibration method, the dissolved gas concentrations in pore water could be calculated using the following equation (Alberto et al. 2000):

$$C_i = \frac{C_g(\alpha V_g - C_{bg} V_g)}{V_i}$$  \hspace{1cm} (5)

where $C_{bg}$ is the initial concentration of the gas in the O$_2$-free air, $\alpha$ is the water:air partition coefficient, and $V_g$ and $V_i$ are the volume of the gaseous and liquid phases, respectively. Eq. 5 can be simplified provided that $C_g \gg C_{bg}$ and $V_g = V_i$ in the following equation:

$$C_i = C_g (1 + \alpha)$$  \hspace{1cm} (6)

Here, $\alpha$ is defined as $C_i/C_g$, where $C_i$ is the dissolved gas concentration in the liquid phase inside the syringe. The variable $C_i$ was calculated in the same manner as $C_i$ in Eqs. 3 and 4, assuming that the temperature inside the syringe was equal to $T$. The variable $C_g$ was also converted to the same unit of $C_i$ as follows:

$$C_g [\text{molL}^{-1}] = \frac{C_g [\text{ppmv}] \times 10^{-6}}{V}$$  \hspace{1cm} (7)

where $V (\text{L mol}^{-1})$ is the volume of 1-mol air at total pressure $P (= 1.0 \text{ atm})$, and temperature $T (\text{°C})$ can be calculated as:

$$V = 22.4 \times \frac{1}{P} \times \frac{T + 273}{273}$$  \hspace{1cm} (8)

**2.4. DO concentration**

Dissolved oxygen concentrations were measured in situ by extracting pore water directly into a closed cell equipped with a DO sensor to avoid contact between the water sample and the atmosphere. A luminescent DO probe (LDO10101; Hach, Loveland, USA) was used because, in principle, it neither consumed oxygen nor required water flow toward the sensor during measurement. The closed cell consisted of the tip of the probe, a 60-mL plastic syringe, and a connecting PVC adapter (Fig. 1). Pore water was extracted from the same type of water sampler used for the measurement of dissolved GHGs, separately installed at depths of 10, 20,
40, and 80 cm. The water was first stored in the preparatory syringe and then pushed into the closed cell. As the water entered, the cell plunger was synchronously pushed out. After measurement, the water was drained from another cell port. DO was measured in December 2011 and March, April, and May 2012.

Dead space in the closed cell was initially filled with ambient air or pore water at different depths. To approximate the true value of the DO concentration in the cell, 50 mL of pore water was repeatedly extracted until the measured value was stabilized. The stroke of the extraction was limited to 10 times to reduce the influence of the inflow of pore water from other depths. Finally, DO was determined based on the sequential data of measured DO by solving a numerical convergence model with Excel® Solver.

2.5. Environmental parameters

Water table levels were measured and recorded with a pressure logger (Hobo U20, Onset, Bourne, USA) settled near the bottom of a perforated PVC pipe (ca. 1.5 m from the ground surface). The influence of the change in atmospheric pressure was canceled based on the pressure data observed at the ground surface level inside the well of FD, where the water table level was lower than −20 cm during the observation period (see Results). The water table level in the well was converted to that at the position of the flux chambers by canceling the small differences in the altitude of the ground surface. Precipitation was measured in the vicinity of the study sites with a tipping bucket rain gauge linked to a HOBO Pendant® event data logger.

Values for pH and electric conductivity (EC) were measured at depths of 10, 20, 40, and 80 cm. Water samples were extracted from the same samplers used for DO measurements. The values of pH and EC were determined in situ with portable sensors (pH: B-212, Horiba, Japan; EC: B-173, Horiba, Japan).

For the measurement of dissolved nitrogenous ions (NO$_3^-$ and NH$_4^+$), 50 mL of water were collected into a plastic bottle at depths of 10, 20, 40, and 80 cm using the same samplers used for DO measurements. These water samples were stored in a refrigerator until analysis. Concentrations of NO$_3^-$ were measured by ion chromatography (Dionex QIC Analyzer, Thermo Fisher Scientific, Waltham, USA). Concentrations of NH$_4^+$ were determined by
colorimetry based on the indophenol-blue method and an ultraviolet-visible spectrophotometer (UV mini 1240, Shimadzu, Kyoto, Japan).

Prior to the flux observations made in this study, hourly monitoring of soil temperatures was conducted in February, June, and December 2011 at the flooded sites and in July 2011 for all study sites. Soil temperatures were measured at depths of 5, 10, 30, and 50 cm with a T-type thermocouple and recorded to a 4-ch data logger (47SD Thermo Recorder, Sato Shoji, Japan). One monitoring effort was usually continued over 4 to 10 consecutive days.

2.6. Statistics
A one-way analysis of variance (ANOVA) was conducted to test for differences in the environmental parameters among the six study sites. In addition, the effects of land use (forest or burnt area) and drainage (flooded or drained) on GHG fluxes were tested by two-way ANOVA; and the effects of land use, drainage and depth on dissolved gas concentrations were tested using three-way ANOVA. Multiple comparison tests among the sites and depths were conducted using the Bonferroni method. A probability level less than 0.05 (P<0.05) was considered statistically significant. All analyses were performed in IBM SPSS Statistics Ver. 21.

3. RESULTS

3.1. Environmental parameters
Fig. 2 shows the temporal changes in precipitation and water table level that occurred during the observation period. The water table levels occasionally increase at all sites following large amounts of rainfall, but they were generally stable from December 2011 to April 2012. In late May, the water table levels were at their lowest due to lower monthly precipitation amounts (94 mm compared to 205–371 mm in other months). Sites FW1, BW1, and BW2 generally remained flooded during the study period. At BW1, a temporary rise in water table level was recorded in late March despite the fact that heavy rainfall was not recorded and that similar increases at other sites were not observed. The rise in water table at BW1 may have been caused by an increase in the Kahayan River’s water level. Sites FW2 and BD remained flooded for 45% and 60% of the observation period, respectively. With a water table level at below −20 cm during the study period, site FD was never flooded.
The air temperatures observed following flux chamber measurements and other environmental parameters are summarized in Table 1. Air temperatures at the burnt sites ranged from 32 to 34°C and were higher than those at the forest sites (27–28°C); however, a significant difference in air temperature was not detected between the same land use types. Forest sites likely had lower temperatures because they were more shaded by dense vegetation. Diurnal soil temperatures varied by as much as 6°C at a depth of 5 cm, but variations were less than 1°C at 50 cm. Although we observed large differences in diurnal variation in soil temperatures, differences in daily mean soil temperatures were small at all depths and study intervals. The largest difference in daily mean soil temperature was 3.3°C and occurred at BW1 at a depth of 5 cm (February to December 2011); temperatures differed by < 2°C for all other depths and sites. The average soil temperatures were 24.5–25.5°C at the forest sites and 27.6–28.1°C at the burnt sites, reflecting differences in air temperature between the different land use types. The vertical distributions in average soil temperatures are shown in Fig. 5b.

The pH of peat water ranged from 3.5 to 4.0 among the study sites, indicating acidic conditions. The electric conductivity (EC) of peat water ranged from 35 to 125 μS cm$^{-1}$ among the study sites and was higher in the forest sites. Variations in pH and EC between measurement depths and months were small. NO$_3$-N concentrations ranged from 0.0519 to 0.260 mg L$^{-1}$ and were relatively higher in the burnt sites. NH$_4^+$-N concentrations ranged from 0.061 to 0.205 mg L$^{-1}$, with no significant differences among the study sites.

3.2. GHG fluxes

Temporal changes in GHG (i.e., CH$_4$, CO$_2$, and N$_2$O) fluxes are shown in Fig. 3. Fluxes in GHGs and CO$_2$-equivalent GWP were averaged for each group (classified by land use and drainage) and are summarized in Table 2, with the results of two-way ANOVA. The relationships between water table levels and these fluxes are shown in Fig. 4.

The site-specific averages of CH$_4$ fluxes ranged from 0.0084 to 5.75 mgC m$^{-2}$ h$^{-1}$ (Table 2). We periodically observed high CH$_4$ emissions at BW1 and BW2 (Fig. 3a) and negative CH$_4$ fluxes (= CH$_4$ absorption) at the forest sites. At FD, 7 of 15 measured fluxes were negative. However, on average, all sites were a net CH$_4$ source during the observation period. In the burnt sites, higher average CH$_4$ fluxes were associated with higher average water table levels.
Relative to the burnt sites, the CH₄ fluxes at the forest sites were consistently lower regardless of water table levels. The results of two-way ANOVA indicated that both main effects (land use and drainage) and their interaction were significant, and CH₄ flux at the flooded burnt sites was significantly higher than those of the other sites (Table 2).

The site-specific averages of CO₂ fluxes ranged from 108 to 340 mgC m⁻² h⁻¹ (Table 2). CO₂ fluxes tended to be relatively low (<150 mgC m⁻² h⁻¹) and stable until February 2012, with higher values (>350 mgC m⁻² h⁻¹) occurring after March 2012 at FW2, FD and BD (Fig. 3b). Among the study sites, CO₂ flux was highest at FD, except for in April 2012. In the forest sites, slightly lower average CO₂ fluxes were associated with higher average water table levels (Fig. 4b). In contrast, the largest CO₂ flux was associated with the highest water table level at BW1. However, this trend may have been an artifact of sampling effort; CO₂ flux was not measured at BW1 from December 2011 to February 2012, when this flux was generally low at all other sites. According to the results of two-way ANOVA, the effects of land use and the interaction of land use and drainage were significant, and CO₂ flux at the drained forest site was significantly higher than that at the drained burnt sites (Table 2).

The site-specific averages of N₂O flux ranged from −8.7 to 8.1 µgN m⁻² h⁻¹ (Table 2). Although we observed variation in N₂O fluxes in early April at BD and BW2, little variation was observed in this flux at other sites during the observation period (Fig. 3c). On average, the N₂O fluxes at the drained sites tended to be positive, whereas they tended to be negative at flooded sites. This difference suggested that drainage influenced whether a site acted as a sink or a source of N₂O in tropical peatlands. However, the relationship between N₂O flux and water table level was unclear (Fig. 4c). The results of two-way ANOVA suggested that there were no significant differences in N₂O fluxes between sites based on either land use or drainage (Table 2).

GWP was mostly affected by CO₂ flux, followed by CH₄ flux. At BW1 and BW2, 18–25% of the GWP were attributed to CH₄, while most of the GWP (>92%) was due to CO₂ at other sites. As a result, average GWP at the flooded burnt sites was 20% larger compared to the flooded forest sites, although the CO₂ fluxes at both sites were similar (Table 2). The results of two-way ANOVA were similar to those for the analysis of CO₂ flux, except that the simple main effect of drainage in the burnt area was significant (Table 2). This exception was probably due to the fact that CH₄ emissions in the flooded burnt sites were larger than those in
the drained burnt site. According to the post hoc test, GWP in the drained forest site was significantly higher than the GWPs of the drained burnt sites, as was observed for CO$_2$ flux.

### 3.3. Dissolved concentrations of GHGs and DO

The vertical distributions of the dissolved concentrations of GHGs (i.e., CH$_4$, CO$_2$, and N$_2$O) and oxygen are shown in Fig. 5. Soil temperatures preliminarily observed in 2011 are also plotted in Fig. 5b. The results of three-way ANOVA on these dissolved gases based on land use, drainage and depth are summarized in Table 3.

Dissolved CH$_4$ concentrations in the burnt sites ranged from 50 to 250 µmol L$^{-1}$ (except at 20 cm in BD), 1–4 orders of magnitude higher than concentrations measured in the forest sites (where dissolved CH$_4$ was <2 µmol L$^{-1}$; Fig. 5a). From 20 to 60 cm depth, dissolved CH$_4$ concentrations increased with increasing depth at BD.

Dissolved CO$_2$ ranged from 0.5 to 3.5 mmol L$^{-1}$ (Fig. 5b). Dissolved CO$_2$ concentrations at FD were highest at depths of 40, 60, and 80 cm out of all the study sites, showing an increasing trend with depth. In contrast, BW1 showed a decreasing trend with depth, with dissolved CO$_2$ being highest at 20 cm out of all the study sites. Other sites did not show clear vertical trends. Similar to CO$_2$ flux, dissolved CO$_2$ showed an increasing trend during the observation period at all sites, especially after March 2012 (time series data are not shown).

We applied the daily mean soil temperature averaged for each depth and site (Fig. 5b) to the calculation for dissolved CO$_2$ concentration (Eq. 3). The largest diurnal variation in soil temperature at a depth of 20 cm was 3.2°C, recorded at FD in July 2011. This means that the difference between the daily average and the actual hourly soil temperature was as much as ±1.6°C. According to the sensitivity analysis for the dissolved CO$_2$ calculation, errors of ±1.6°C in soil temperature produced differences of ±4.2% in dissolved CO$_2$. Because diurnal variations in soil temperatures were less than 3.2°C for all other sites and at the lower depths, errors associated with the use of daily mean soil temperatures in calculations were less than ±4.2% for all other sites.

The dissolved N$_2$O concentrations were typically <0.01 µmol L$^{-1}$ (142 data/167 in total; Fig. 5c). Only five data points for dissolved N$_2$O were >0.1 µmol L$^{-1}$; four of these points were obtained at depths of 20 and 40 cm in FD, and the fifth point was at 20 cm in BW2. As a
result, dissolved N₂O concentrations in the drained forest site were higher compared to other sites at 20 and 40 cm.

Dissolved oxygen concentrations were generally <10 µmol L⁻¹, much lower than DO in a solution which is in equilibrium with the atmospheric oxygen level (220–280 µmol L⁻¹; Fig. 5d). In the burnt sites, DO concentrations decreased between depths of 10 and 20 cm and became stable below 20 cm, suggesting the presence of an oxygen supply from the ground surface. According to the results of three-way ANOVA, only the main effect of land use was significant (forest; 6.9±5.6 µmol L⁻¹ > burnt; 4.0±2.9 µmol L⁻¹).

4. DISCUSSION

4.1. CH₄ fluxes and dissolved concentrations

In this study, CH₄ fluxes were mainly influenced by drainage conditions (Table 2; Fig. 4a). Previous studies have reported positive relationships between mean water table level and seasonal CH₄ emissions in non-flooded northern peatlands (e.g., Bubier and Moore, 1993; Pelletier et al., 2007). In tropical peatlands, Jauhiainen et al. (2005) found that CH₄ fluxes were between −75 and 260 µgC m⁻² h⁻¹ and generally increased with increasing water table level (−70 to +20 cm). Melling et al. (2005a) showed that CH₄ fluxes ranged from −4.53 to 8.40 µgC m⁻² h⁻¹ in a mixed peat swamp forest (water table level: −60 to −20 cm), from −7.44 to 102 µgC m⁻² h⁻¹ in a sago plantation (−40 to 0 cm), and from −32.8 to 4.17 µgC m⁻² h⁻¹ in an oil palm plantation (−80 to −30 cm) and also increased with increasing water table levels. We observed similar CH₄ fluxes and trends in the drained sites as those reported in these previous studies (Table 2).

Based on a comprehensive review on CH₄ flux observation in tropical peatlands, Couwenberg et al. (2010) has recently reported that CH₄ fluxes are generally low and often distinctly negative for water levels below −20 cm, while tend to be higher and more variable at higher water levels. Jauhiainen et al. (2005) found that CH₄ fluxes were positive at water table levels >−50 cm and were negative at levels ≤−60 cm at locations that were in the vicinity of our study area. Jauhiainen et al. (2008) also reported that CH₄ fluxes began to increase as the water table levels rose higher than −40 to −20 cm in a drained forest or higher than −30 to 0 cm in a deforested, burned site. Considering the relationship between CH₄ flux and water
table level, there is an empirical depth of water table level at which CH$_4$ production and oxidation are balanced. In this study, the drained forest site was both a weak sink and source of CH$_4$ (Table 2; Fig. 3a) with a water table level that ranged from −39.7 to −7.0 cm at the time of flux measurement, similar to the water levels observed in the previous studies.

In contrast to the CH$_4$ fluxes, dissolved CH$_4$ concentrations were mainly affected by land use (Fig. 5a). Ueda et al. (2000) found that dissolved CH$_4$ concentrations in groundwater in a coastal peat swamp in Thailand varied from 0.01 to 417 µmol L$^{-1}$, with average values of 48 and 226 µmol L$^{-1}$ in wet seasons spanning 4 years at two sampling sites. Koschorreck (2000) reported that CH$_4$ in pore water at the top 8 cm of a silty loam sediment on an island in the Amazon River ranged from 0 to 900 µmol L$^{-1}$. Terazawa et al. (2007) found that dissolved CH$_4$ concentrations in groundwater ranged from 5.6 to 28.4 µmol L$^{-1}$ in a floodplain forest located in northern Japan. Pangala et al. (2013) recently reported that dissolved CH$_4$ concentrations observed in the vicinity of the flooded forest sites in this study ranged from 113–1539 µmol L$^{-1}$, much higher than our results. Several differences between the two studies, including sampling depths (50–150 cm in Pangala et al., 2013 compared to 20–80 cm in this study) and procedures, may have resulted in this discrepancy in dissolved CH$_4$ concentrations. However, the primary reason for this difference is unclear.

The CH$_4$ flux in the drained forest site was the lowest out of all sites (Table 2), likely because this site had the lowest water table level (20–60 cm below the ground surface; Fig. 2). Under such aerobic conditions, methanotrophic activity would have been promoted over methanogenesis. Hanson and Hanson (1996) indicated that anoxic soils produce CH$_4$, while well-drained soils act as a sink for atmospheric CH$_4$ due to CH$_4$ oxidation. In the drained burnt site, CH$_4$ fluxes were lower than in the flooded forest sites, although dissolved CH$_4$ concentrations were much higher in the burnt sites compared to the forest sites (Fig. 5a). Previous studies suggested that CH$_4$ diffusing toward the atmosphere is oxidized to CO$_2$ by methanotrophic bacteria when oxic conditions are present in the upper peat profile (Couwenberg et al., 2010; Inubushi et al., 2003; Jauhiainen et al., 2005; 2008). In the drained burnt site, the occasionally non-flooded condition (Fig. 2) would create an aerobic layer near the ground surface. This would allow CH$_4$ produced in deeper peat layers to be oxidized and CH$_4$ emissions and dissolved CH$_4$ concentrations at a depth of 20 cm to decrease. However, in the flooded burnt sites, the lack of an aerobic surface layer prohibited CH$_4$ oxidation, resulting
in a higher CH$_4$ flux compared to other sites and a higher dissolved CH$_4$ concentration at 20 cm compared to deeper layers.

4.2. CO$_2$ fluxes and dissolved concentrations

In tropical peatlands, Melling et al. (2005b) reported that soil CO$_2$ fluxes ranged from 100–533 mgC m$^{-2}$ h$^{-1}$ in a mixed peat swamp forest, from 63–245 mgC m$^{-2}$ h$^{-1}$ in a sago plantation, and from 46–335 mgC m$^{-2}$ h$^{-1}$ in an oil palm plantation in Sarawak, Malaysia. Jauhiainen et al. (2005) also reported that CO$_2$ fluxes were 132–166 mgC m$^{-2}$ h$^{-1}$ in hummocks and 37.9–188 mgC m$^{-2}$ h$^{-1}$ in hollows in a tropical peat swamp forest in Central Kalimantan. The CO$_2$ fluxes obtained in this study were comparable to those found in previous studies. Ueda et al. (2000) reported that the dissolved CO$_2$ concentrations in groundwater in a coastal peat swamp in Thailand varied from 0.240 to 3.29 mmol L$^{-1}$ during the wet season, similar to the range observed in this study.

The high CO$_2$ fluxes (Table 2 and Fig. 3b) and dissolved CO$_2$ concentrations at depths of 40–80 cm (Fig. 5b) in the drained forest site suggested that CO$_2$ production was enhanced due to root respiration at this site. In addition, the water table level at this site was consistently at 20–60 cm below the ground surface, much lower than the other sites (Fig. 2). The aerobic conditions that resulted from the low water table level in the drained forest site should promote peat decomposition and contribute to high CO$_2$ emissions at this site. Dissolved CO$_2$ concentrations at 20 cm were lower than those at deeper layers, which could be attributed to diffusive CO$_2$ loss into the atmosphere through the unsaturated top layer in the drained forest site.

The CO$_2$ fluxes and dissolved concentrations in the drained burnt site, which were significantly lower than those in the drained forest site (Table 2; Figs. 3b and 5b), suggested a low root respiration rate from poor vegetation cover. Insufficient drainage relative to the drained forest site (Fig. 2) may have also inhibited CO$_2$ production through peat decomposition in the drained burnt site. In the flooded forest sites, the inhibition of aerobic peat decomposition would suppress CO$_2$ emissions and result in lower dissolved CO$_2$ concentrations. In the flooded burnt sites, a continuously flooded condition (Fig. 2) would inhibit CO$_2$ diffusion into the atmosphere and cause the high dissolved CO$_2$ concentrations at 20 cm.
The increasing trends in CO$_2$ flux and dissolved concentration during the observation period (Fig. 3b), which differed from the trends in CH$_4$, N$_2$O, and other environmental factors, could be attributed to the increase in plant root respiration as opposed to microbial peat decomposition. Although we could not determine the cause of this trend from data collected in this study, the acclimation of plant roots to saturated conditions may be one possible explanation (Drew et al., 1994; Mommer et al., 2004).

4.3. Flux and dissolved concentration of N$_2$O

Like the fluxes in CH$_4$, the N$_2$O fluxes observed in this study were generally related to drainage condition. Melling et al. (2007) reported that the N$_2$O flux ranged from −3.4 to 19.7 µgN m$^{-2}$ h$^{-1}$ in a mixed swamp forest, from 1.0 to 176.3 µgN m$^{-2}$ h$^{-1}$ in a sago plantation, and from 0.9 to 58.4 µgN m$^{-2}$ h$^{-1}$ in an oil palm plantation. The N$_2$O fluxes in the mixed peat swamp forest in that study were comparable to those observed in the drained sites of this study. Takakai et al. (2006) also measured N$_2$O flux at locations that were identical several of those observed in this study. According to their results, the average N$_2$O fluxes in the wet season (2002 to 2004) were 49±63 µgN m$^{-2}$ h$^{-1}$ in the drained forest site and 55±100 µgN m$^{-2}$ h$^{-1}$ in the drained burnt site, comparable to the observations made in the wet season (2011–2012) in this study. Ueda et al. (2000) found that dissolved N$_2$O concentrations in the groundwater of a coastal peat swamp in Thailand varied from 0 to 0.012 µmol L$^{-1}$, with average values of 0.006 and 0.008 µmol L$^{-1}$ (in the wet seasons of 4 consecutive years) at two sampling sites. These results were similar to those observed in this study.

Given that methanogenesis is an anaerobic process that tends to occur after all other electron donors have been consumed (Burgin and Groffman, 2012), the flooded sites in this study were likely anaerobic. Under anaerobic conditions, even atmospheric N$_2$O can be absorbed into peat water and consumed by denitrification, which oxidizes organic carbon by reducing NO$_3^-$ (Burgin and Groffman, 2012). A recent review indicated that net negative N$_2$O fluxes have been reported in numerous previous studies, showing that low mineral N and high moisture content are favorable for N$_2$O consumption (Chapuis-Lardy et al., 2007). In this study, the flooded burnt sites consumed more N$_2$O than the flooded forest sites (Table 2). This could be attributed to differences in DO concentrations, with significantly higher concentrations in the forest sites compared to the burnt sites (Table 3; Fig. 5d).
NO$_3^-$ concentration is one of the important controlling factors for soil N processes, including nitrification and denitrification. In this study, the effect of NO$_3^-$ on N$_2$O flux was unclear, although there were significant differences in dissolved NO$_3^-$ concentrations in peat water among the study sites (Table 1). Takakai et al. (2006) found that the N$_2$O flux in a cropland in the vicinity of our study area increased with increasing NO$_3^-$-N content in the top 10 cm of soil. In that study, however, the NO$_3^-$-N content was significantly higher in cropland (200–300 mg kg$^{-1}$ dry soil) than in the drained forest or in the drained burnt sites (0.43–91 and 0.79–5.9 mg kg$^{-1}$ dry soil, respectively). In this study, the total NO$_3^-$-N content was likely too small to influence N$_2$O emissions.

4.4. Effect of plant roots on dissolved GHGs and DO

Ueda et al. (2000) measured DO concentrations in the groundwater of a coastal peat swamp in Thailand, finding that most concentrations were below the detection limit and had a maximum value of 14 µmol L$^{-1}$. Liebner et al. (2012) observed a rapid decrease (from >80% to 0% of air saturation) in DO in the top 20 cm of soil in an alpine wetland and a constant DO profile below the top layer. Given these observations, it is likely that DO in saturated peat soils remains very low even near the ground surface, similar to DO concentrations observed in this study (Fig. 5d). Most of DO in peat water would be consumed in the top 10 cm, thus there was no significant differences among depths at each study site.

The high dissolved CH$_4$ (Fig. 5a) and the low DO (Table 3) in the burnt sites may be due to a lack of large trees, which have large and deep root systems that can supply oxygen into the rhizosphere. The increase in the concentration of dissolved CH$_4$ with depth in the flooded forest sites suggested that CH$_4$ oxidation rate by plant-derived O$_2$ may depend on the amount of plant root biomass, which usually decreases with increasing depth. Previous studies have also reported increased dissolved CH$_4$ as well as decreased DO (Liebner et al., 2012), redox potential (Fritz et al., 2011; Koschorreck, 2000), and root density (Fritz et al. 2011) in saturated peat profiles. Thus, O$_2$ supply via plant roots would reduce net CH$_4$ production in forest sites even under flooded conditions. The decline in DO by CH$_4$ oxidation could be compensated by this plant-mediated oxygen supply.

In addition, dissolved N$_2$O concentrations at depths of 20 and 40 cm in the drained forest site (Fig. 5c) could also be explained by the effect of drainage and soil O$_2$ on denitrification. Namely, a water table level that was consistently 20–60 cm below the ground surface (Fig. 2)
and high DO concentrations (Fig. 5d) created slightly oxic conditions in the peat profile, which is favorable for N\textsubscript{2}O production. Burgin and Groffman (2012) found that N\textsubscript{2}O production in intact soil cores collected from a riparian wetland in the northeastern US increased with increasing O\textsubscript{2} concentration. However, nitrification is also an important microbial N process that produces N\textsubscript{2}O. Bollmann and Conrad (1998) suggested that the main source of N\textsubscript{2}O was through nitrification when soil moisture is low and through denitrification when soil moisture is high. Therefore, the high concentrations of dissolved N\textsubscript{2}O in the drained forest site may have been due to nitrification in the drained surface layer.

5. CONCLUSIONS

Our first hypothesis was supported given that 1) CH\textsubscript{4} emissions in the flooded burnt sites were significantly larger than those in the flooded forest sites, 2) dissolved CH\textsubscript{4} concentrations in the burnt sites were much higher than those in the forest sites, and 3) DO concentrations in the forest sites were significantly higher than those in the burnt sites. In this study, however, CH\textsubscript{4} fluxes were affected by drainage conditions rather than land use. The CH\textsubscript{4} flux and the dissolved CH\textsubscript{4} concentration at a depth of 20 cm in the drained burnt site were similar to values observed in the forest sites, suggesting that CH\textsubscript{4} oxidation in the surface soil layer occurred in the drained burnt site. Given the high dissolved CH\textsubscript{4} concentrations observed in the deep layers, CH\textsubscript{4} emissions in the drained burnt site will likely rapidly increase when this area is flooded again.

Our second hypothesis was weakly supported by the observed GWPs in the flooded burnt sites, which were 20\% higher than those in the flooded forest sites (no significant difference; \(P=0.493\)). CO\textsubscript{2} fluxes in both sites were almost equivalent, and high CH\textsubscript{4} emissions in the flooded burnt sites actually increased GWPs at these sites. In this study, however, GWP was mainly determined by CO\textsubscript{2} flux. Consequently, GWP and CO\textsubscript{2} flux in the drained forest site were the highest for all study sites, and N\textsubscript{2}O flux made little contribution to GWP.
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Captions for Figures

**Fig. 1.** Diagram of a closed cell for DO measurement.

**Fig. 2.** Seasonal changes in precipitation (vertical bars) and water table level at each site (lines with symbols).

**Fig. 3.** Seasonal variations in the fluxes of a) CH$_4$, b) CO$_2$, and c) N$_2$O at the ground surface. Error bars show standard deviations.

**Fig. 4.** Relationship between water table level and the fluxes of a) CH$_4$, b) CO$_2$, and c) N$_2$O averaged for the observation period. Error bars show standard deviations.

**Fig. 5.** Vertical distribution in dissolved concentrations of a) CH$_4$, b) CO$_2$, c) N$_2$O, and d) oxygen (DO) averaged for each depth. Soil temperature preliminarily observed in 2011 was plotted with dissolved CO$_2$ (b). Error bars show standard deviations. For better identifiability, the depths in the profiles were slightly shifted from their actual depths (20, 40, 60, and 80 cm for CH$_4$, CO$_2$, and N$_2$O; 10, 20, 40, and 80 cm for DO; 5, 10, 30, and 50 cm for soil temperature).
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Fig. 5. Vertical distribution in dissolved concentrations of a) CH₄, b) CO₂, c) N₂O, and d) oxygen (DO) averaged for each depth. Soil temperature preliminarily observed in 2011 was plotted with dissolved CO₂ (b). Error bars show standard deviations. For better identifiability, the depths in the profiles were slightly shifted from their actual depths (20, 40, 60, and 80 cm for CH₄, CO₂, and N₂O; 10, 20, 40, and 80 cm for DO; 5, 10, 30, and 50 cm for soil temperature).
**Table 1.** Environmental factors at all study sites averaged for the observation period

<table>
<thead>
<tr>
<th>Site</th>
<th>Description</th>
<th>Air temperature (°C)</th>
<th>pH</th>
<th>EC (µS cm⁻¹)</th>
<th>NO₃⁻-N (mg L⁻¹)</th>
<th>NH₄⁺-N (mg L⁻¹)</th>
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<td>Flooded forest site #1</td>
<td>28.2 ± 1.0 ab (3)</td>
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<td>3.6 ± 0.2 ab (31)</td>
<td>86 ± 14 c (31)</td>
<td>0.0519 ± 0.0319 a (13)</td>
<td>0.143 ± 0.214 (13)</td>
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<tr>
<td>FD</td>
<td>Drained forest site</td>
<td>28.9 ± 1.5 ab (5)</td>
<td>3.5 ± 0.1 a (17)</td>
<td>125 ± 16 d (17)</td>
<td>0.0781 ± 0.0325 ab (9)</td>
<td>0.086 ± 0.130 (9)</td>
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<tr>
<td>BW1</td>
<td>Flooded burnt site #1</td>
<td>33.9 ± 4.0 c (2)</td>
<td>3.7 ± 0.2 b (31)</td>
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<td>35 ± 5 a (28)</td>
<td>0.0915 ± 0.0401 ab (13)</td>
<td>0.061 ± 0.110 (13)</td>
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1. Values are means and standard deviations of environmental factors.
2. Numbers in parentheses represent sample size at each site.
3. Values within the same column with different lowercase letters differ significantly ($P<0.05$; corrected by Bonferroni method).
Table 2. Comparison of GHG fluxes among different land use and drainage conditions

<table>
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<tr>
<th>Sites</th>
<th>CH₄ flux (mgC m⁻² h⁻¹)</th>
<th>CO₂ flux (mgC m⁻² h⁻¹)</th>
<th>N₂O flux (µgN m⁻² h⁻¹)</th>
<th>GWP (CO₂-eq mgC m⁻² h⁻¹)</th>
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<tr>
<td>Flooded forest sites (FW1 and FW2)</td>
<td>1.37 ± 2.03 a (24)</td>
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ANOVA

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Simple main effects

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<td>Drainage in forest vegetation</td>
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<td>5.32 *</td>
<td>0.18</td>
<td>4.34 *</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Drainage in burnt area</td>
<td>20.05 **</td>
<td>1.86</td>
<td>1.23</td>
<td>4.26 *</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. Fluxes are shown in means ± standard deviations. Positive fluxes mean net emission from peat soil.
2. Numbers in parentheses represent sample sizes in each level.
3. Values within the same column with different lowercase letters differ significantly (P<0.05; corrected by Bonferroni method).
4. Symbols * and ** mean P<0.05 and P<0.01, respectively.
Table 3. Results of three-way ANOVA on dissolved gas concentrations among different land use, drainage and depth

<table>
<thead>
<tr>
<th></th>
<th>Dissolved CH\textsubscript{4}</th>
<th>Dissolved CO\textsubscript{2}</th>
<th>Dissolved N\textsubscript{2}O</th>
<th>DO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(df)</td>
<td>(F)</td>
<td>(df)</td>
<td>(F)</td>
</tr>
<tr>
<td><strong>Main factors</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Land use</td>
<td>1</td>
<td>125.95 **</td>
<td>1</td>
<td>8.54 **</td>
</tr>
<tr>
<td>Drainage</td>
<td>1</td>
<td>28.25 **</td>
<td>1</td>
<td>14.48 **</td>
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<tr>
<td>Depth</td>
<td>3</td>
<td>0.41</td>
<td>3</td>
<td>4.09 **</td>
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<tr>
<td>Error</td>
<td>151</td>
<td>137</td>
<td>151</td>
<td>73</td>
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<tr>
<td><strong>Interactions</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Land use*Drainage</td>
<td>1</td>
<td>27.84 **</td>
<td>1</td>
<td>47.98 **</td>
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<tr>
<td>Land use*Depth</td>
<td>3</td>
<td>0.50</td>
<td>3</td>
<td>6.70 **</td>
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<tr>
<td>Drainage*Depth</td>
<td>3</td>
<td>5.26 **</td>
<td>3</td>
<td>9.36 **</td>
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<tr>
<td>Land use<em>Drainage</em>Depth</td>
<td>3</td>
<td>5.30 **</td>
<td>3</td>
<td>1.67</td>
</tr>
</tbody>
</table>

Symbols * and ** mean \(P<0.05\) and \(P<0.01\), respectively.