



Rhizosphere to the atmosphere: contrasting methane pathways, fluxes, and geochemical drivers across the terrestrial–aquatic wetland boundary

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Abstract. Although wetlands represent the largest natural source of atmospheric CH₄, large uncertainties remain regarding the global wetland CH₄ flux. Wetland hydrological oscillations contribute to this uncertainty, dramatically altering wetland area, water table height, soil redox potentials, and CH₄ emissions. This study compares both terrestrial and aquatic CH₄ fluxes in permanent and seasonal remediated freshwater wetlands in subtropical Australia over two field campaigns, representing differing hydrological and climatic conditions. We account for aquatic CH₄ diffusion and ebullition rates and plant-mediated CH₄ fluxes from three distinct vegetation communities, thereby examining diel and intra-habitat variability. CH₄ emission rates were related to underlying sediment geochemistry. For example, distinct negative relationships between CH₄ fluxes and both Fe(III) and SO₄²⁻ were observed. Where sediment Fe(III) and SO₄²⁻ were depleted, distinct positive trends occurred between CH₄ emissions and Fe(II)/acid volatile sulfur (AVS). Significantly higher CH₄ emissions ($p < 0.01$) in the seasonal wetland were measured during flooded conditions and always during daylight hours, which is consistent with soil redox potential and temperature being important co-drivers of CH₄ flux. The highest CH₄ fluxes were consistently emitted from the permanent wetland (1.5 to 10.5 mmol m⁻² d⁻¹), followed by the *Phragmites australis* community within the seasonal wetland (0.8 to 2.3 mmol m⁻² d⁻¹), whilst the lowest CH₄

fluxes came from a region of forested *Juncus* spp. (−0.01 to 0.1 mmol m⁻² d⁻¹), which also corresponded to the highest sedimentary Fe(III) and SO₄²⁻. We suggest that wetland remediation strategies should consider geochemical profiles to help to mitigate excessive and unwanted methane emissions, especially during early system remediation periods.

1 Introduction

Wetlands are considered one of the most valuable ecosystems on Earth (Costanza et al., 2014) and may be classified as both permanently inundated (i.e. lakes and shallow waters) and seasonally inundated (i.e. vegetated) biomes. They are biodiversity hotspots that provide ecosystem services such as water filtration, sediment trapping, floodwater retention, and carbon (C) storage (Bianchi, 2007). Wetlands account for ~ 5.5 % of terrestrial surfaces (Melton et al., 2013) and have been estimated to store ~ 4% (Bridgham et al., 2014) to ~ 30 % (Mitsch et al., 2013) of Earth's estimated 2500 Pg soil C pool (Lal, 2008). Pristine wetlands have long been considered net C sinks due to their high rates of productivity and low rates of decomposition (Petrescu et al., 2015); however, due to their waterlogged nature and anaerobic soils, wetlands are ideal environments for the production of methane (CH₄), a potent greenhouse gas. As such, wet-

lands are recognised as Earth's largest natural source of CH₄ to the atmosphere ($185 \pm 21 \text{ Tg C yr}^{-1}$) (Saunois et al., 2016).

Resolving the drivers, pathways, and effects of seasonal weather oscillations on wetland CH₄ sink or source behaviours is important to enable more accurate climate model projections and to reduce uncertainties in the global wetland CH₄ budget (Kirschke et al., 2013; Saunois et al., 2016). Weather oscillations affect the total wetland areal extent and inundation periods, with wet conditions facilitating anaerobic conditions favouring methanogenesis, while the opposite is seen during dry periods, which potentially mitigates CH₄ emissions (Wang et al., 1996; Whiting and Chanton, 2001). Mitsch et al. (2013) estimated that the average ratio of freshwater wetland CO₂ sequestration to CH₄ emissions was 25.5 : 1, though this was later refuted by Bridgman et al. (2014). As CH₄ is 34 times more potent than carbon dioxide (CO₂) over a 100-year timescale (Stocker et al., 2013), this suggests that many freshwater wetlands may have a net positive radiative forcing effect on climate (Petrescu et al., 2015; Hemes et al., 2018). However, variability in geomorphology, wetland maturity, salinity, and underlying geochemical composition contributes to variable CH₄ dynamics (Whiting and Chanton, 2001; Bastviken et al., 2011; Poffenbarger et al., 2011). The lack of latitudinally resolved wetland CH₄ emission data, the limited number of studies constraining the multiple wetland CH₄ flux pathways (i.e. ebullition, diffusion, and plant-mediated), and the ongoing anthropogenic conversion of wetland systems (Bartlett and Harriss, 1993; Neubauer and Megonigal, 2015; Saunois et al., 2016) further contribute to uncertainties around CH₄ regional- to global-scale budgets.

Extensive clearing and drainage of many coastal wetlands has occurred over the previous 2 centuries in order to accommodate agriculture, aquaculture, and urban development (Armentano and Menges, 1986; White et al., 1997; Villa and Bernal, 2018). Drained wetlands can lead to rapid soil organic matter oxidation and transform systems to net CO₂ sources (Deverel et al., 2016; Pereyra and Mitsch, 2018). Drainage systems can also reduce wetland inundation periods and alter sediment redox-dependant geochemistry and microbially mediated reactions (Johnston et al., 2014), particularly those involving bioavailable iron (Fe(III)), sulfate (SO₄²⁻), and nitrate (NO₃⁻). Importantly, anaerobic carbon metabolism employing these terminal electron acceptors (Fe(III), SO₄²⁻, NO₃⁻) competes thermodynamically with methanogenic bacteria and Archaea and can thereby inhibit CH₄ production (Lal, 2008; Burdige, 2012; á Norði and Thamdrup, 2014; Karimian et al., 2018). With increasing value placed on the ecosystem services provided by wetlands, many degraded systems are now undergoing remediation and re-flooding (Johnston et al., 2014). However, the ecosystem benefits, such as enhanced biodiversity and water quality, may come at a price in the form of higher initial CH₄ flux rates and predicted net radiative forcing for several

centuries post-remediation – thus posing a “biogeochemical compromise” (Hemes et al., 2018).

Within Australia, it has been estimated that more than 50% of natural wetlands have been lost to land use change, drainage, and degradation since European settlement (ANCA, 1995; Finlayson and Rea, 1999). By comparing and reviewing pristine Australian wetland carbon stocks to drained sites and greenhouse gas dynamics, Page and Dalal (2011) estimated that through biomass loss, enhanced soil respiration, N₂O production, and a reduction in CH₄ emissions, Australian wetland loss equated to ~1.2 Pg of CO₂ equivalent emitted to the atmosphere. Much of eastern Australia's freshwater coastal wetlands are underlain by Holocene-derived sulfidic sediments (i.e. pyrite – Fe₂S, known as coastal acid sulfate soils; CASSs) formed during periods of higher sea levels (Walker, 1972; White et al., 1997). When CASSs are drained, pyrite is oxidised, producing sulfuric acid (H₂SO₄). This results in highly acidic soils with pH levels as low as 3 (Sammut et al., 1996; Johnston et al., 2014). After rainfall events, groundwater transports H₂SO₄ from the CASS landscapes into nearby creeks and estuaries (Sammut et al., 1996). The low pH groundwater discharge also mobilises iron and aluminium, fuels aquatic deoxygenation, and can lead to large fish kills and degradation of infrastructure (White et al., 1997; Johnston et al., 2003; Jeffrey et al., 2016; Wong et al., 2010). Drained CASS wetlands typically contain abundant reactive Fe(III) and exhibit complex sulfur and Fe cycling (Burton et al., 2006, 2011; Boman et al., 2008). Wetland iron and sulfur cycling can profoundly influence CH₄ production and consumption via a series of complex redox reactions coupled with organic matter mineralisation (Holmkvist et al., 2011; Sivan et al., 2014). As such, terminal electron acceptor availability is critical when considering wetland remediation and the biogeochemical compromise paradigm.

Here we assess CH₄ emissions from a remediated freshwater CASS wetland in subtropical eastern Australia and compare fluxes from the permanent wetland and the adjacent seasonal wetland ecotypes. We hypothesise that wetland CH₄ emissions will differ significantly between the campaigns and between the four wetland communities due to differences in soil chemistry, hydrology, and plant physiology. We account for three atmospheric flux pathways for methane; ebullition, diffusion, and plant-mediated fluxes over diel cycles and within different hydrological conditions. CH₄ fluxes were also assessed in relation to the underlying soil properties, including sulfate, reactive iron III and iron II, acid volatile sulphur, chloride, and organic carbon.

2 Methods

2.1 Study site

Cattai Wetlands are located on the mid-coast of New South Wales, Australia. The reserve covers 500 ha, featuring a shallow permanent wetland covering an area of approximately 16 ha that is adjacent to a seasonal wetland and floodplain located to the south (Fig. 1). Both sites discharge into the nearby Cooperbrook Creek, a tributary of the larger Manning River estuary. The site was extensively cleared and low-lying areas drained during the early 1900s in order to aid agriculture and development in the region. As a result of this anthropogenic drainage, the oxidation of CASS produced sulfuric acid and episodic acidic discharge to adjacent creeks for many years (Tulau, 1999). To ameliorate acidic discharge, the natural hydrology of the site was restored in 2003 through the decommissioning of agricultural drains and removal of floodgates. Re-flooding of the CASS landscape has reduced the production of sulfuric acid, acid discharge, and aluminium and iron mobilisation, hence improving the downstream water quality (GTCC, 2014).

The region receives a mean annual rainfall of 1180 mm with the majority falling during early autumn with an average maximal monthly rainfall occurring in March (152 mm). The lowest rainfall generally occurs during the winter months with average minimal rainfall during September (60 mm). Average minimum and maximum summer temperatures range from 17.6 to 29 °C (January) and in winter range from 5.9 to 18.5 °C (July) (BOM, 2018). The dominant vegetation type within the permanent wetland is an introduced water lily species (*Nymphaea capensis*), while the fringes of the wetland consist of wetland tree species: *Casuarina* spp. and *Melaleuca quinquenervia*. The seasonal wetland to the south is dominated by the sedge *Juncus kraussii* (“Juncus” from here on) and features scattered stands of *Phragmites australis* (“Phragmites” from here on) with areas of slightly higher elevation dominated by *Juncus kraussii* below *Casuarina* spp. (“Juncus–forest” from here on) (Fig. 1).

2.2 The aquatic CH₄ flux of the permanent wetland

To quantify CH₄ ebullition rates, up to 12 ebullition domes were deployed under two different hydrological conditions (detailed below) at ~20 m intervals along a longitudinal transect, from the edge of the permanent wetland towards the centre. Each dome was carefully suspended below the water level by flotation rings, ensuring minimal disturbance of sediment and the water column. Gas samples were extracted from the headspace of each dome using a 300 mL gas-tight syringe at periods of ~48 h. The volume was recorded and each sample then diluted using ambient air (1 : 729 ratio) and analysed in situ using a manufacturer-calibrated cavity ring-down spectrometer (Picarro G2201-*i*) to determine CH₄ concentrations (ppm). Diffusive CH₄ fluxes from the

permanent wetland were measured using a floating chamber with a portable greenhouse gas analyser (UGGA, Los Gatos Research). To account for spatial and temporal variability, measurements were conducted during both daytime and night-time, and sampling was within vegetated areas featuring lilies (*Nymphaea capensis*) that were only present during the second campaign, forested areas (*Melaleuca* spp.), and in areas where no aquatic vegetation was present (i.e. open water). A total of 39 CH₄ floating chamber incubations averaging ~8 min in duration were recorded over the two campaigns, with 19 during C1 (nine at night) and 30 during C2 (12 at night). The average r^2 value of linear regressions of CH₄ concentrations versus time during chamber incubations was 0.97 ± 0.05 . One chamber measurement was disregarded as an outlier (as it was more than 3 times the standard deviation of the mean) and any chambers capturing ebullition bubbles (determined by a non-linear increase in concentration) were also disregarded. Examples of these, in addition to the ebullition and diffusive CH₄ flux methods and measurements from the permanent wetland, have previously been reported elsewhere (Jeffrey et al., 2019).

2.3 Plant-mediated CH₄ fluxes

Simultaneous time series chamber experiments were conducted over a minimum of 24 h to measure diel CH₄ fluxes during each campaign from the three different wetland vegetation ecotypes. These ecotypes were *Juncus kraussii*, *Phragmites australis*, and *Juncus kraussii* amongst *Casuarina* spp. forest (Fig. 1). In each ecotype, three acrylic bases (65 × 65 × 30 cm) were installed 4 months before the first time series experiment to minimise disturbance to the sediment profile and vegetative rhizosphere. Vegetative flux chambers were constructed of an aluminium frame with clear Perspex walls and a roof that matched the areal footprint of the pre-inserted acrylic bases. The chambers were 100, 150, and 50 cm high at Juncus, Phragmites, and Juncus–forest sites, respectively. The custom sizes were tailored for the different vegetation heights, whilst minimising chamber volume as much as possible. Each chamber was leak-tested under laboratory conditions prior to fieldwork.

Before each field incubation, chambers were flushed with atmospheric air and then carefully lowered over the vegetation and onto the acrylic base, ensuring an airtight seal. A small fan circulated internal air within each chamber. Air within the chamber was pumped through a closed loop from the top of the chamber using gas tubing (BEV-A-Line), passing through a drying agent (Drierite desiccant), and then analysed in situ using a calibrated cavity ring-down spectrometers (Picarro G2201-*i* or Los Gatos), recording the flux rate of CH₄ (ppm s⁻¹). The gas flow was returned near the base inside each vegetation chamber, closing the loop. Vegetation incubation times ranged from 6 to 15 min depending on the flux rate and were taken from triplicate chambers to account for heterogeneity within each ecotype. During the

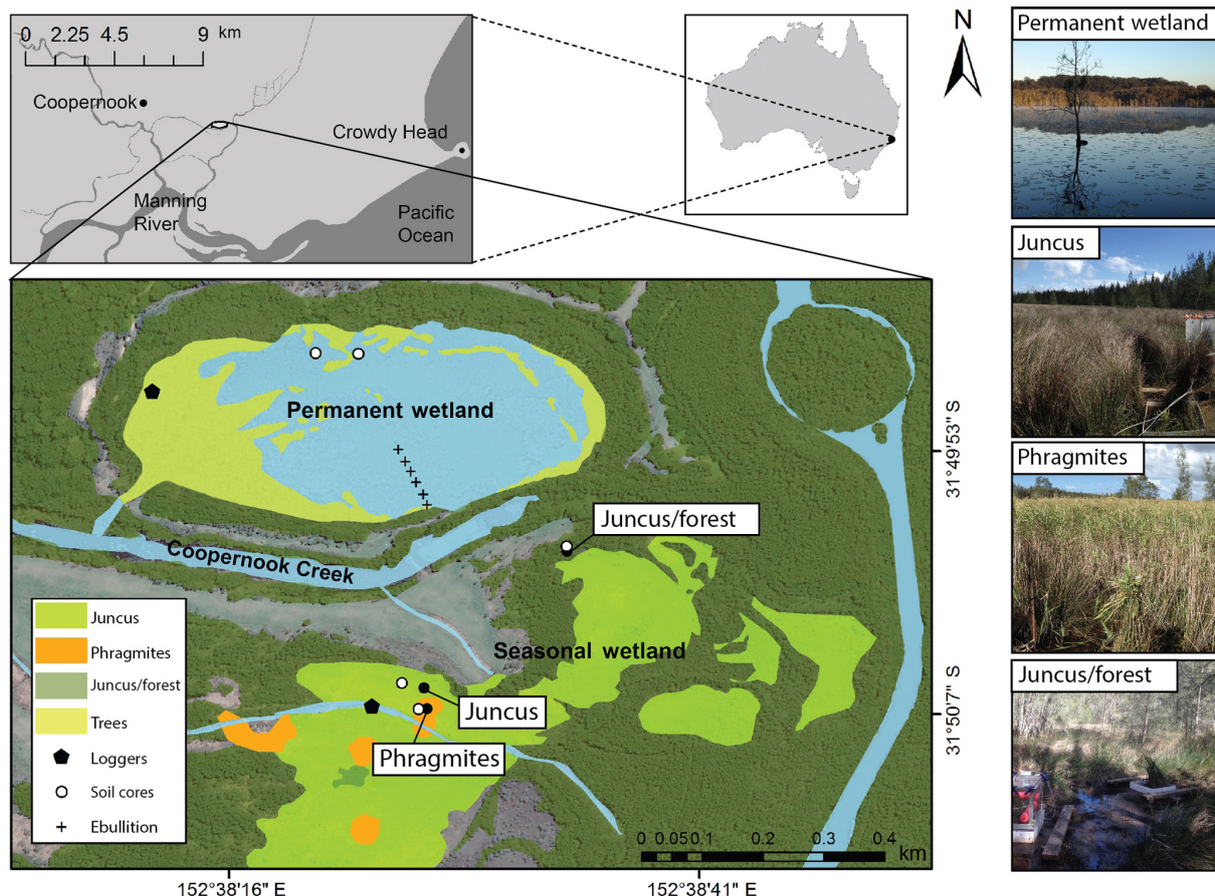


Figure 1. The seasonal wetland study sites consisting of *Juncus* (*Juncus kraussii*), *Phragmites* (*Phragmites australis*), and *Juncus*–forest (*Juncus kraussii* below *Casuarina* spp.); the permanent wetland and sediment coring sites, ebullition replicate transect, 24 h vegetation time series sites, and imagery of vegetation ecotypes.

first time series (C1), an average of 16.7 ± 2.9 daytime flux measurements (i.e. after sunrise) and 7.3 ± 1.6 night-time (i.e. after sunset) were recorded within each habitat. During the second campaign (C2) an average of 27.7 ± 2.9 (daytime) and 10.3 ± 1.5 (night-time) flux measurements were recorded within each habitat. In addition, CH_4 fluxes from the adjacent exposed soils or shallow overlying water at each site were also measured at ~ 4 h intervals to determine the influence and role of plant-mediated CH_4 fluxes compared to non-vegetated CH_4 fluxes. Light and temperature loggers (Onset Hobo) measured the changes in diel air temperature ($^{\circ}\text{C}$) and photosynthetically active radiation (PAR) at each site.

2.4 Soil geochemistry and redox conditions

A water logger (Minidiver, Van Essen Instruments) was deployed in the permanent wetland before the first campaign to monitor changes in water depth (cm) and temperature ($^{\circ}\text{C}$). Field pH (pH_F) and the redox potential (Eh_F ; reported against standard hydrogen electrode) were determined in situ by directly inserting the electrode into the soils (5 cm of

depth, eight replicates on average) at each site. A composite sampling approach (three cores) was used to collect sediment samples from each site to determine organic C content, $\text{Fe(III)}_{\text{HCl}}$, $\text{Fe(II)}_{\text{HCl}}$, Cl , SO_4^{2-} , and acid volatile sulfur (AVS). The cores were sampled in close proximity to the time series habitats (5 to 15 m) in December 2016, but within the permanent wetland the cores were taken from elsewhere to avoid disturbance of the shallow water column and sediments. The cores were extracted by inserting a 4.0 cm diameter acrylic tube into the sediment to a depth of up to 50 cm. Cores were immediately sectioned into 2 cm increments to a depth of 20 cm, and 5 cm increments thereafter, ensuring higher vertical resolution in the organic-rich near-surface sediments. Samples were immediately placed into airtight bags, then frozen within 12 h of collection at -16°C in a portable freezer and transferred to a -80°C freezer in the laboratory.

For analysis, the frozen samples were thawed in an oxygen-free anaerobic chamber (1%–5% H_2 in N_2) using an oxygen-consuming palladium (Pd) catalyst. The defrosted samples were homogenised using a plastic spatula. AVS con-

ment was determined by adding 1–2 g of wet sediment with 6 M HCl : 1 M L-ascorbic acid. The liberated H₂S was captured in 5 mL of 3 % Zn acetate in 2 M NaOH and then quantified using iodometric titration. The reactive Fe fractions were determined using a sequential extraction procedure optimised for acid sulfate soils based on Claff et al. (2010). Poorly crystalline solid-phase Fe(II) and Fe(III) were determined by extracting 2 g wet subsamples with cold N₂-purged 1 M HCl for 4 h. Aliquots of 0.45 µm filtered extract were analysed for Fe(II) [Fe(II)_{HCl}] and total Fe [Fe_{HCl}] using the 1,10-phenanthroline method with the addition of hydroxylammonium chloride for total Fe (APHA, 2005). The Fe(III) [Fe(III)_{HCl}] was determined by the difference of [Fe_{HCl}] – [Fe(II)_{HCl}]. Total organic carbon (TOC) and total S (S_{Tot}) were determined via a LECO CNS-2000 carbon and sulfur analyser. Chloride and sulfate concentrations were measured using filtered (0.45 µm) aliquot from a 1 : 5 water extract of freshly defrosted wet soil, as per Rayment and Higginson (1992), via ion chromatography using a Metrosep A Supp4-250 column, an RP2 guard column, and eluent containing 2 mM NaHCO₃, 2.4 mM Na₂CO₃, and 5 % acetone, in conjunction with a Metrohm MSM module for background suppression.

2.5 Calculations

Both the air–water and vegetative CH₄ fluxes were calculated for the chamber deployments in the permanent wetland and seasonal wetland using the equation

$$F = (s (V/RT_{\text{air}}A))t, \quad (1)$$

where s is the regression slope for each chamber incubation deployment (ppm s⁻¹), V is the chamber volume (m³), R is the universal gas constant (8.205 × 10⁻⁵ m³ atm K⁻¹ mol⁻¹), T_{air} is the air temperature inside the chamber (K), A is the surface area of the chamber (m²), and t is the conversion factor from seconds to days and to millimoles. We assume that atmospheric pressure is 1 atm. Ebullition rates (E_b) (mmol m⁻² d⁻¹) were calculated using the equation

$$E_b = ([\text{CH}_4]\text{CH}_{4\text{Vol}}) / A V_m T_d, \quad (2)$$

where [CH₄] is the CH₄ concentration in the collected gas (%), CH_{4Vol} is the gas volume sampled (L), A is the funnel area (m²), V_m is the molar volume of CH₄ at in situ temperature (L), and T_d is deployment time (days).

2.6 Statistical analysis

As the CH₄ flux data was non-parametric we used a Kruskal–Wallis one-way analysis of variance (ANOVA) on ranks to test for significant differences between each campaign, between flux pathways, and between diel variability, where $p < 0.001$. Dunn's multiple pairwise comparisons were then used to analyse specific sample pairs ($p < 0.05$).

3 Results

Prior to the first campaign in April 2017 (C1), an extreme hot–drying summer period occurred (Fig. 2). This resulted in an average wetland water column temperature of 23.3 ± 0.7 °C and a water depth in the permanent wetland as low as ~ 7.3 cm, with exposed sediments along the wetland perimeter during the preceding month. There was a high rainfall event prior to C1 with 342 mm of rainfall recorded over the preceding 2 weeks and an additional 35 mm of rain occurring during C1 fieldwork (Fig. 2), thus raising the water column depth in the permanent wetland to 77.2 cm in less than 4 weeks. This C1 deployment was therefore categorised as the “post-dry–flooded” period, during which air temperatures ranged from 13.3 to 22.8 °C and the average water column temperature in the permanent wetland was 20.4 ± 0.5 °C. The second fieldwork campaign was conducted in September 2017 (C2) under cool–drying conditions, in which air temperatures ranged from as low as 3.4 to 34.9 °C (Fig. 2), with cooler average water temperatures of 12.6 ± 0.4 °C in the permanent wetland (Fig. 2). The depth of the permanent wetland at this time had dropped slightly to ~ 33 cm (Fig. 2).

3.1 Sediment core profiles and soil redox potentials

Average concentrations from soil cores (Table 1, Fig. 3) were based upon the top 20 cm of the profile, in which the highest organic carbon concentrations were found. The Fe(III)_{HCl} concentrations were greater than Fe(II)_{HCl} at all three seasonal wetland sites; however, the permanent wetland showed an opposite trend with low concentrations of both Fe(III) (5.6 ± 10.7 mmol kg⁻¹) and SO₄²⁻ (1.5 ± 1.0 mmol kg⁻¹) (Fig. 3, Table 1). The highest average concentrations of Fe(III)_{HCl} were found at the Juncus–forest site (204.0 ± 51.6 mmol kg⁻¹) and the highest and similar concentrations of SO₄²⁻ were in Phragmites and Juncus–forest sediments (45.4 ± 41.0 and 43.3 ± 16.7 mmol kg⁻¹) (Fig. 3, Table 1). Net positive redox potential was found at all four sites during C1 (under post-dry–flooded conditions), indicating a lag time between recent flooding and the onset of reducing conditions. In contrast, a negative redox potential was found within the permanent wetland and Phragmites during C2, indicating reduced conditions under cool–drying conditions (Table 1). The TOC concentrations (%) were highest in the upper profiles and similar across all sites (Fig. 3, Table 1), averaging 13.4 ± 7.6 %.

3.2 Permanent and seasonal wetland CH₄ fluxes

The vegetation time series revealed that diel variability of plant-mediated CH₄ emissions occurred at most ecotypes, with the highest CH₄ fluxes occurring during daytime around midday and the lowest CH₄ fluxes during the night-time (Fig. 4, Table 1). The lowest CH₄ fluxes were found in the Juncus–forest habitat with a net negative CH₄ flux observed

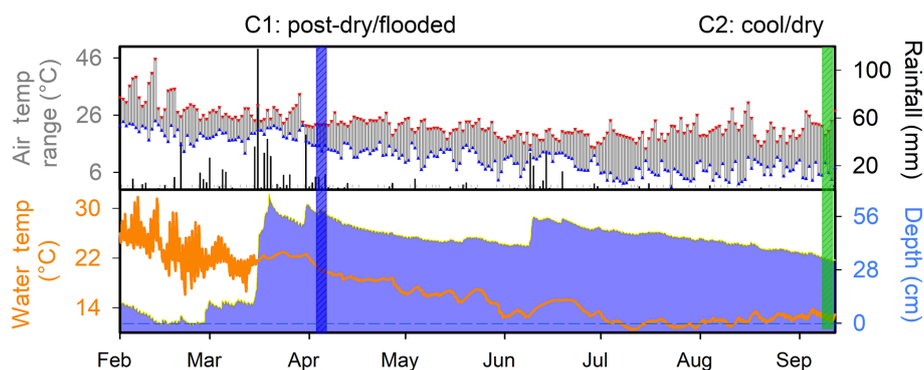


Figure 2. Hydrograph for 7 months in 2017 indicating daily rainfall, maximum–minimum air temperature, water temperature, and antecedent hydrology. Vertical coloured bands represent the two fieldwork campaigns.

Table 1. Summary of plant-mediated CH_4 fluxes from the seasonal wetland time series and diel CH_4 diffusive fluxes and ebullition from the permanent wetland during C1 (post-dry–flooded) and C2 (cool–drying). The corresponding sediment core data are average concentrations from 0 to 20 cm below ground level.

CH_4 flux ($\text{mmol m}^{-2} \text{d}^{-1}$)	Ebullition	Diffusion	Juncus	Phragmites	Juncus–forest
Sediment flux – C1			0.06	0.04	0.10
Daytime flux – C1		0.57	1.79	2.64	0.13
Night-time flux – C1		2.07	1.50	1.59	0.10
Daily average flux – C1	2.02	1.49	1.70	2.27	0.12
Sediment flux – C2			0.00	0.20	0.00
Daytime flux – C2		11.72	0.06	0.94	0.13
Night-time flux – C2		8.39	0.04	0.48	0.10
Daily average flux – C2	2.10	10.46	0.05	0.77	–0.01
FeHCl(II) (mmol kg^{-1})		202.3	11.6	15.4	1.5
FeHCl(III) (mmol kg^{-1})		5.6	83.3	56.1	204.0
SO_4^{2-} (mmol kg^{-1})		1.5	17.6	45.4	43.3
Cl : SO_4^{2-}		14.8	8.4	13.9	7.4
AVS ($\mu\text{mol g}^{-1}$)		18.5	0.7	0.9	0.3
TOC (% C)		11.6	14.3	14.8	14.6
C1 – redox Eh (mV)		71.7	46.5	9.6	54.4
C2 – redox Eh (mV)		–216.3	11.9	–89.3	424.5

during C2 time series. The CH_4 sediment fluxes measured amongst each vegetation time series were consistently much lower than the plant-mediated CH_4 fluxes, indicating that the vegetation was indeed the main conduit for CH_4 to the atmosphere (Fig. 4, Table 1). The CH_4 fluxes were highly variable between the replicates at each site. Temperature and PAR followed similar diel trends to each other and had positive correlations with CH_4 emissions (Fig. 4).

CH_4 fluxes from the three vegetation types were significantly higher during C1 than during C2 ($p < 0.001$). During C1, the CH_4 fluxes from Juncus and Phragmites were not significantly different from each other but were both significantly higher ($p < 0.001$) than Juncus–forest; however, during C2 the CH_4 fluxes of each seasonal wetland habitat were significantly different between all habitats ($p < 0.05$)

(Fig. 5). The highest average CH_4 fluxes in each of the vegetation types always occurred during the daytime but were not significantly different to night-time fluxes (Fig. 5, Table 1). Phragmites consistently emitted the highest CH_4 fluxes ($2.27 \pm 1.42 \text{ mmol m}^{-2} \text{d}^{-1}$ during C1 and $0.77 \pm 0.46 \text{ mmol m}^{-2} \text{d}^{-1}$ during C2). The Juncus–forest ecotype within the seasonal wetland consistently produced the lowest CH_4 fluxes of all sites, with a negligible flux that was not significantly different from zero occurring during C2 ($-0.01 \pm 0.08 \text{ mmol m}^{-2} \text{d}^{-1}$).

The permanent wetland showed an inverse trend with 7-fold and significantly higher ($p < 0.001$) diffusive fluxes during the cool–drying C2 when lilies were present ($10.46 \pm 15.81 \text{ mmol m}^{-2} \text{d}^{-1}$) compared to the post-dry–flooded C1 when no lilies were present ($1.49 \pm 2.75 \text{ mmol m}^{-2} \text{d}^{-1}$),

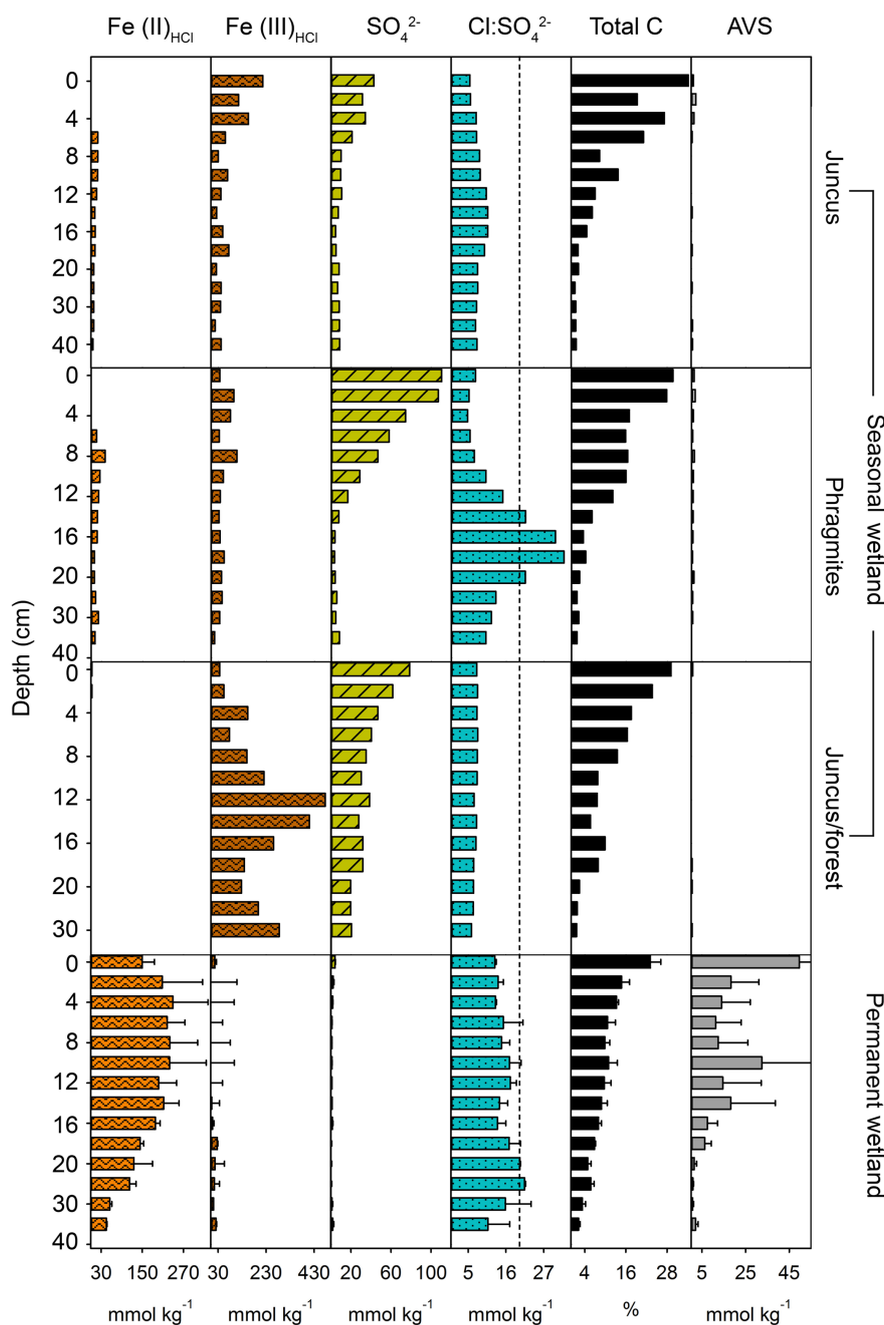


Figure 3. Soil profiles of the permanent and seasonal wetland sites indicating $\text{Fe(II)}_{\text{HCl}}$, $\text{Fe(III)}_{\text{HCl}}$, SO_4^{2-} , $\text{Cl} : \text{SO}_4^{2-}$ (a proxy for depletion of marine-derived sulfate, where > 20 is broadly indicative of SO_4^{2-} reduction and < 8 CASS pyrite oxidation; Mulvey, 1993), total C, and acid volatile sulfur (AVS). Note: the permanent wetland profiles are averages from two adjacent sites with error bars representing the standard deviation.

while the ebullition rates were similar during both campaigns (Fig. 5, Table 1). Overall, the diffusive fluxes of the permanent wetland were within the range of CH_4 fluxes from the three seasonal wetland habitats but were significantly higher than Juncus–forest during both campaigns and Juncus during C2 (Fig. 5). Diel diffusive flux variability was not significant between daytime and night-time (Table 1, Fig. 5).

3.3 Temperature and PAR

Correlation plots for both temperature ($^{\circ}\text{C}$) and sunlight (PAR) versus CH_4 emissions from the three vegetation ecotypes showed no distinct relationships with the exception of Phragmites during C2 for PAR ($r^2 = 0.18$, $p < 0.01$) and temperature ($^{\circ}\text{C}$) ($r^2 = 0.35$, $p < 0.001$). No clearer trends

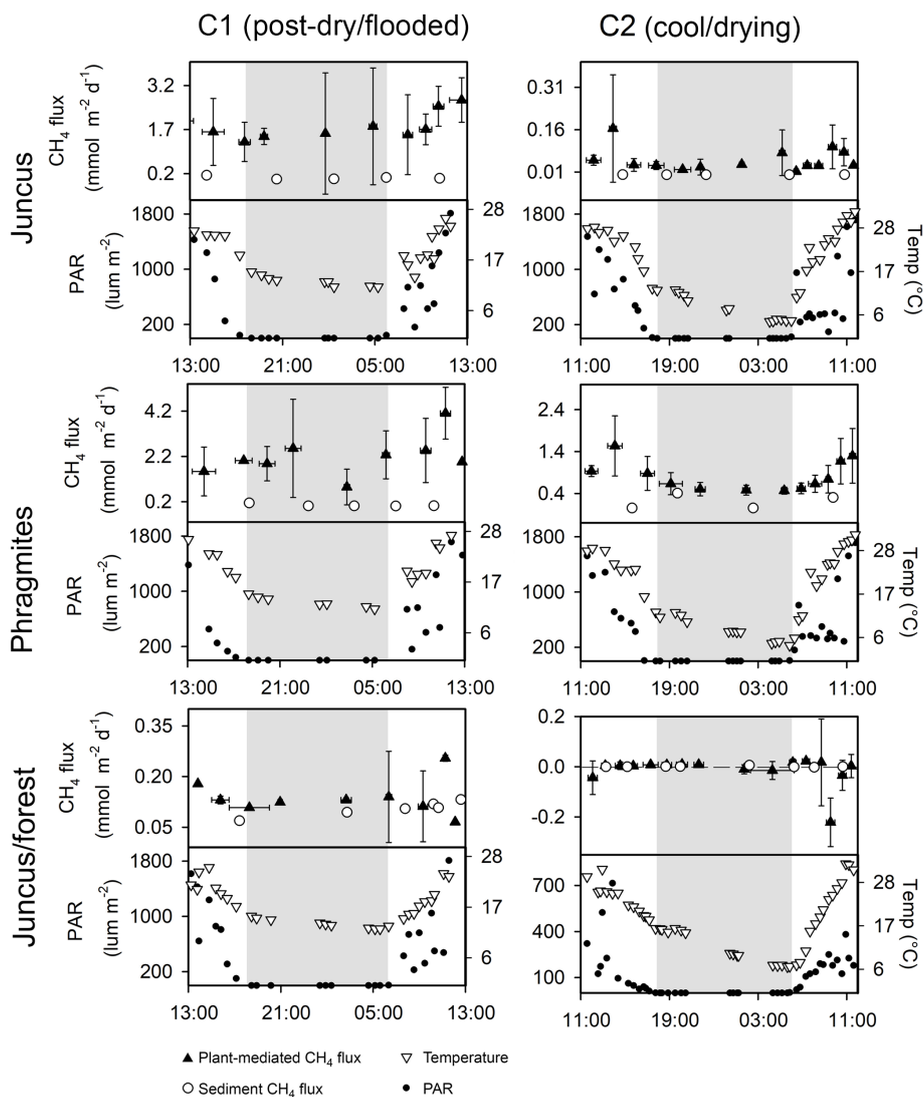


Figure 4. Simultaneous 24 h time series of vegetative CH_4 fluxes from the seasonal wetland ecotypes at Cattai Wetlands during C1 (post-dry–flooded, April 2017) and C2 (cool–drying conditions, September 2017). The vertical error bars of the plant-mediated CH_4 flux ($\text{mmol m}^{-2} \text{d}^{-1}$) represent the standard deviation of the triplicate time series measurements taken from each site, and the horizontal bars represent the total aggregated time period represented by replicate chambers. The grey shading indicates night-time. Note: different y axis scales for CH_4 to highlight diel trends.

were observed by combining all site measurements or separating daytime fluxes and drivers from night-time fluxes and drivers.

4 Discussion

4.1 Geochemistry of the CASS landscape

Sediment profiles provide insights into the historical geochemical changes that have occurred across the CASS landscapes of the four Cattai Wetlands sites (Fig. 3). We base our results and discussion on the upper rhizosphere depth zone (20 cm) as this featured the highest organic carbon concen-

trations and is therefore assumed to be an active area of carbon metabolism and CH_4 production and consumption (Nedwell and Watson, 1995). If we assume that relatively uniform deposition of late Holocene materials occurred, the differences between present-day profiles are related to historical changes in hydrology and land use, topographic elevation, geochemical trajectories, and vegetative carbon inputs. For example, the permanent wetland shows distinct differences to the adjacent seasonal wetland sites, with divergent geochemical signatures of both iron and sulfate that reflect sustained inundation (Table 1, Fig 5). The permanent wetland had significantly lower Fe(III) ($p < 0.001$) and 11- to 30-fold lower SO_4^{2-} concentrations within the upper soil pro-

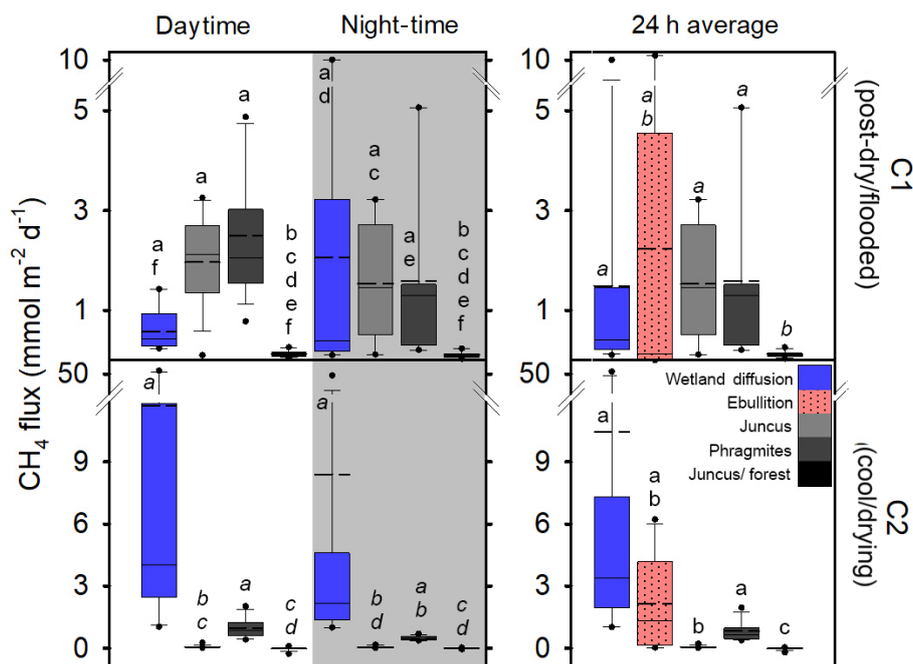


Figure 5. Fluxes of CH₄ from diel sampling and ebullition over two campaigns from the permanent wetland and adjacent 24 h time series of the seasonal wetland vegetation types. Note: diffusive fluxes during C2 include chambers featuring lilies, dashed line represents the average, solid line represents the median, and dots represent the 5th and 95th percentiles. Letters show groups that did not differ significantly ($p > 0.05$) using ANOVA on ranks and Dunn's pairwise comparisons within each campaign.

file compared to the seasonal wetland. The ratio of Fe(III)_{HCl} to Fe(II)_{HCl} from the flooded soils of the permanent wetland was 0.03, indicating the sediments were almost completely depleted of Fe(III). Under reducing conditions in which there is low SO₄²⁻ and little to no Fe(III) to competitively exclude methanogenesis, CH₄ production becomes more favourable. Indeed, CH₄ production was on average the highest from the permanent wetland, especially when considering the dual CH₄ pathways of ebullition and air–water diffusion (Table 1).

In addition to sulfate reduction, some depletion of the sulfur pool from the permanent wetland may have occurred due to drainage exports of sulfuric acid (H₂SO₄) discharging from the CASS landscape throughout the last century. Alternatively, reducing conditions induced by re-flooding freshwater wetlands is known to encourage the re-formation of AVS and pyrite (FeS₂) and produce alkalinity, thereby attenuating acid production and discharge (Burton et al., 2007; Johnston et al., 2012, 2014) and reducing the total SO₄²⁻ pool of CASS landscapes. While the AVS concentrations found within the permanent wetland (up to 18.5 μmol g⁻¹) were a result of sulfate reduction induced by CASS wetland restoration, they nonetheless represent a relatively volatile form of sulfur, which is at risk of rapid oxidation during drought periods (Johnston et al., 2014; Karimian et al., 2017). The AVS concentrations of the permanent wetland sites were more than 20-fold higher than the three adjacent seasonal wetland sites and represent a potentially volatile by-product and con-

sequence of re-flooding CASS soil landscapes, in addition to leading to increases in CH₄ emissions (Table 1).

The soil profile from the seasonal wetland Juncus–forest habitat featured abundant Fe(III)_{HCl} (with an Fe(III)_{HCl} to Fe(II)_{HCl} ratio of 136) and also SO₄²⁻. This was associated with the lowest fluxes of CH₄ for both sampling periods (Fig. 3, Table 1). Relatively low CH₄ fluxes from Juncus–forest are likely due to the more oxidising conditions present at this site and the surfeit of thermodynamically favourable terminal electron acceptors (i.e. Fe(III) and SO₄²⁻), which would competitively exclude organic matter degradation by methanogenic Archaea (Postma and Jakobsen, 1996). At the other seasonal wetland sites, the average Fe(III) and SO₄²⁻ concentrations were intermediate (i.e. lower than Juncus–forest, but higher than the permanent wetland), although in the upper profile the Phragmites had more SO₄²⁻, while Juncus had more Fe(III) (Fig. 3, Table 1). CH₄ flux values from these sites were also intermediate (Table 1). Sediment profiles from both Juncus and Phragmites indicated a degree of Fe reduction based on the ratios of Fe(III) : Fe(II), which were 7.2 and 3.6, respectively. The redox potentials from Phragmites during both the C1 and C2 campaigns (9.6 and –89.0 mV, respectively) were consistently lower than Juncus during the C1 and C2 campaigns (46.5 and 12.0 mV, respectively), which is consistent with the more reducing conditions encouraging CH₄ production in the Phragmites habitat. Further, as iron reduction yields more free energy

than SO_4^{2-} reduction (Burdige, 2012), then Fe reduction at the *Juncus* site may outcompete CH_4 production ahead of SO_4^{2-} reduction in *Phragmites*, which may help explain some of the differences in CH_4 production between the two sites. The positive significant trends between Fe(II), AVS, and the $\text{Cl} : \text{SO}_4^{2-}$ ratios with CH_4 flux rates ($r_s = 0.88$, $p < 0.01$) further support our hypothesis that reducing conditions and a smaller pool of sediment Fe(III) and SO_4^{2-} facilitate higher CH_4 production rates (Fig. 7). Alternatively, the negative trends observed between soil redox potentials, SO_4^{2-} , Fe(III), and CH_4 fluxes affirm that the abundance of thermodynamically favourable terminal electron acceptors plays a role in attenuating CH_4 production at each site.

4.2 Plant-mediated CH_4 fluxes from the seasonal wetland

Plant-mediated CH_4 fluxes were significantly higher ($p < 0.001$) during C1 under post-dry–flooded conditions with 20–30 cm of standing waters in the seasonal wetland (Table 1). While waterlogged conditions are an obvious driver of higher CH_4 production rates from saturated sediments in addition to the geochemical differences (previously discussed), other drivers which may explain these trends include differences in diel variability in temperature, PAR, and plant physiology, which may influence CH_4 gas transport pathways.

In vegetated seasonal wetlands, plant-mediated gas transport is recognised as a dominant pathway for CH_4 emission to the atmosphere and accounts for up to 90 % of total wetland fluxes (Whiting and Chanton, 1992; Sorrell and Boon, 1994). For plant survival in near-permanent inundation environments, oxygen transport occurs via the aerenchyma downwards to the rhizome. This increases plant performance by mitigating (i.e. oxidising) the accumulation of phytotoxins such as sulfides and reducing metal ions around the roots (Penhale and Wetzel, 1983; Armstrong and Armstrong, 1990; Armstrong et al., 2006). As oxygen transfer to the rhizosphere occurs, an exchange of sedimentary CH_4 can be efficiently transported from the rhizosphere to atmosphere, bypassing sedimentary oxidative processes along the way. This process in plants can be either convective (i.e. pressurised) or via passive diffusive gas flow, both of which are adaptive traits of many wetland species (Armstrong and Armstrong, 1991; Konnerup et al., 2011).

During both campaigns the highest CH_4 fluxes from seasonal wetland vegetation were emitted from *Phragmites* and always occurred during daylight (Table 1, Fig. 8). In *Phragmites australis*, the presence of pressurised lacunar leaf culms drive a mass flow of oxygen to the rhizome and back to the atmosphere via older (non-pressurised) efflux culms (Sorrell and Boon, 1994; Henneberg et al., 2012). This process has been widely studied in wetlands featuring *Phragmites australis*, as it is one of the most productive and widespread flowering wetland species (Tucker, 1990; Clevering and Liss-

ner, 1999; Brix et al., 2001; Chanton et al., 2002). Milberg et al. (2017) found no apparent diel patterns of CH_4 fluxes from *Phragmites australis* during seven campaigns within the Swedish growing season. In a mid-latitude prairie wetland, Kim et al. (1998) showed that CH_4 emissions peaked around midday and that daytime emissions were about 3-fold higher than night-time emissions, positively correlating with temperature and PAR. These were similar to our findings with the highest CH_4 fluxes of each time series occurring near midday local time ($4.88 \text{ mmol m}^{-2} \text{ d}^{-1}$ at 10:50 LT during C1 and $2.06 \text{ mmol m}^{-2} \text{ d}^{-1}$ at 12:15 LT during C2) (Fig. 4). We also found a positive significant relationship between CH_4 flux and both temperature and PAR during C2 (Fig. 6). The often high diel variability in CH_4 fluxes from *Phragmites australis* occurs as convective gas transport increases rhizospheric oxygen and CH_4 exchange via living culms during the daytime, whereas molecular diffusion during the night-time facilitates a more passive and lower CH_4 flux pathway through dead culms (Armstrong and Armstrong, 1991; Chanton et al., 2002).

One possible reason CH_4 fluxes were lower from *Juncus* than *Phragmites* despite their close geographical location may be due to the passive gas diffusion mechanism utilised by *Juncus* spp. (Henneberg et al., 2012). Unlike the pressurised conductive gas flow mechanisms of *Phragmites*, many wetland rush species (such as *Juncus* spp.) employ passive diffusive gas flow to survive within waterlogging environments (Brix et al., 1992; Konnerup et al., 2011). Despite diffusion being a less efficient gas transport mechanism (Konnerup et al., 2011), plant-mediated CH_4 diffusion is recognised as the dominant pathway for CH_4 emissions from many seasonal wetland species. During C1 and C2, daytime fluxes (diffusive) from *Juncus* were only 19 % and 33 % higher than night-time fluxes (diffusive). In comparison, from *Phragmites* these day-to-night ratios were almost triple this (67 % and 94 % higher) during the same periods. This may potentially be due to the more efficient daytime conductive gas transfer pathway of CH_4 through *Phragmites australis* compared to the more passive diffusive CH_4 gas transfer pathway of *Juncus kraussii* and/or the effectiveness of these different species in altering sedimentary redox conditions. This suggests that non-pressurised pathways may result in lower net rhizosphere–atmosphere gas exchange of CH_4 from seasonal wetland vegetation. Alternatively, root depth and root density differ between these two species (De La Cruz and Hackney, 1977; Moore et al., 2012), which may further influence redox dynamics in the rhizosphere and the potential extent of net gas exchange.

The *Juncus*–forest habitat emitted significantly lower fluxes of CH_4 during both time series campaigns and was a net sink for CH_4 during C2 (Table 1, Fig. 8). Although wetland trees have recently been shown to contribute significantly to CH_4 fluxes from flooded environments (Pangala et al., 2017), we could not quantify or constrain the role of trees as a conduit of methane to the atmosphere at this

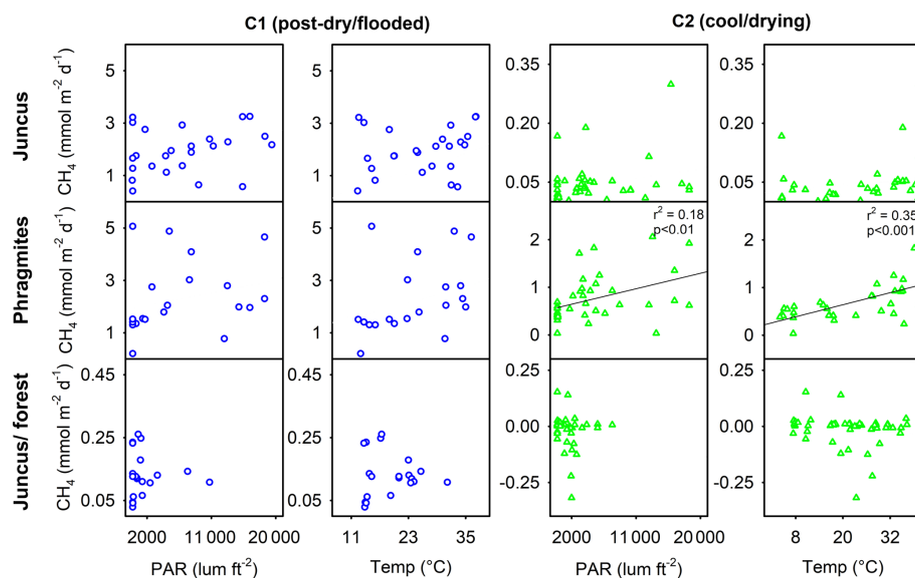


Figure 6. Correlations of CH₄ with temperature (°C) and photosynthetically active radiation (PAR) (lum ft⁻²) for the three wetland vegetation sites of Cattai Wetlands during two field campaigns.

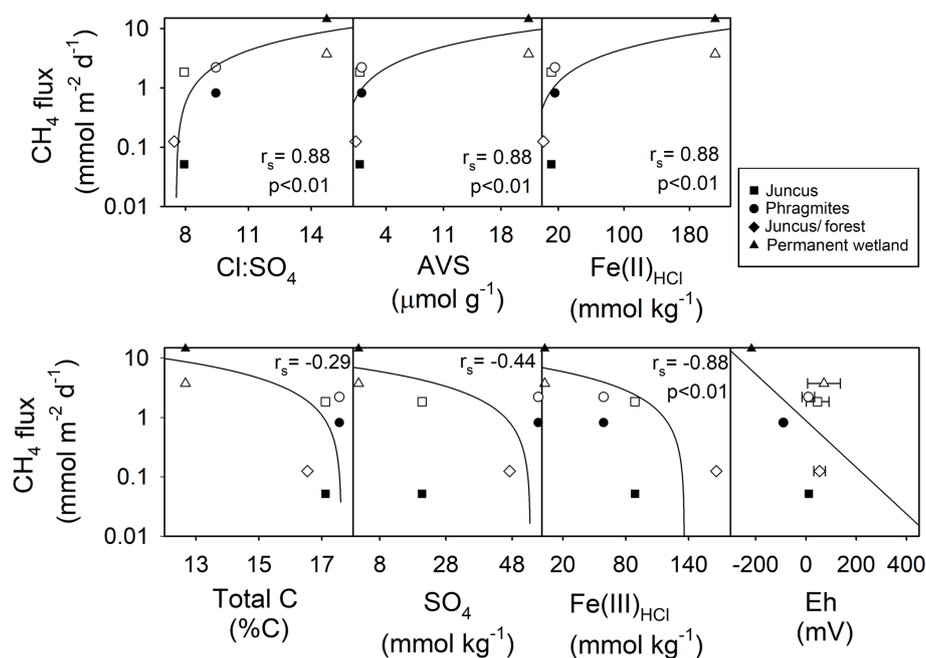


Figure 7. Regression analysis of average daily CH₄ fluxes (mmol m⁻² d⁻¹) vs. subsoil parameters of 0–20 cm core depth (i.e. CH₄ “active” zone). Note: log scale y axis of CH₄ fluxes from the four wetland ecotypes over two campaigns. Note: the r_s values calculated using Spearman rho are for C1 (black shapes) and C2 (white shapes).

site. Regardless, there were clearly lower CH₄ fluxes through the *Juncus kraussii* at the Juncus–forest habitat compared to the Juncus-only habitat. As the species at ground level were identical, these differences are not related to vegetative gas transport mechanisms or organic carbon content (Table 1). Shading by the overhanging trees may inhibit daytime diffusive CH₄ gas transport through the Juncus–forest habitat,

presumably due to lower rates of photosynthesis; however, PAR was only lower during C2 (Fig. 7) and therefore does not appear to explain the CH₄ flux differences observed during C1. The differences are therefore likely explained by the higher positive redox potentials (Table 1) that may be partially attributable to rhizome aeration by the nearby trees and more abundant thermodynamically favourable terminal elec-

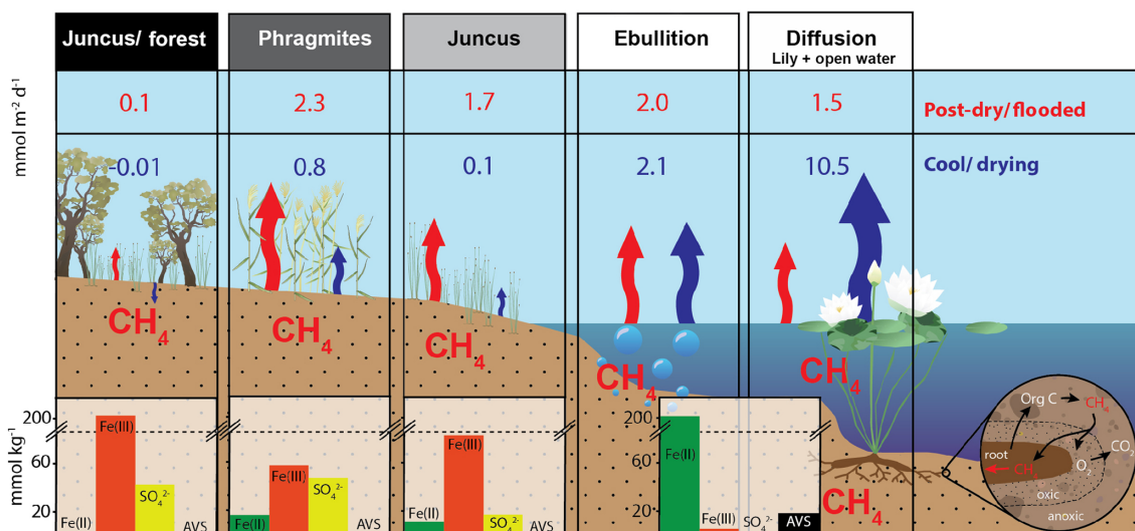


Figure 8. Conceptual model summarising the terrestrial and aquatic CH₄ fluxes (mmol m⁻² d⁻¹) and sediment core profile parameters (mmol kg⁻¹) of the permanent and seasonal wetlands during C1 (post-dry–flooded conditions) and C2 (cool–drying conditions) of Cattai Wetlands. Conceptual diagram rhizome process insert adapted from Conrad (1993). Note: dashed line highlights y axis break.

tron acceptors (i.e. Fe(III) and SO₄²⁻) (Fig. 3), all of which can inhibit methane production within the sediments (Burdige, 2012).

4.3 Permanent wetland CH₄ fluxes

Diffusive CH₄ fluxes from the permanent wetland varied considerably between campaigns; however, ebullition fluxes were similar (Table 1, Fig. 8). The highest CH₄ fluxes for both ebullition and diffusion (2.1 and 10.5 mmol m⁻² d⁻¹, respectively) occurred during C2 despite cooler conditions (Figs. 2 and 8). This, however, was the opposite trend to the seasonal wetland CH₄ fluxes (Table 1, Fig. 8). One reason may be the antecedent hydrological conditions before C1 (Fig. 2). Jeffrey et al. (2019) reported that a water level drawdown of the permanent wetland after a hot and drying summer period exposed some of the permanent wetland sediments to oxidative conditions. This may have oxidised a portion of the labile sedimentary carbon pool prior to C1 sampling of the permanent wetland, therefore reducing the total CH₄ pool observed during C1 sampling. A lag time (ranging from weeks to months) for recovery of the CH₄ pool post-drought has been observed in other systems (Boon et al., 1997) and also during lab-based experiments (Freeman et al., 1992; Knorr et al., 2008). Further, during C2 the return of macrophyte species *Nymphaea capensis* most likely enhanced CH₄ gas transport from the rhizosphere to the floating chambers, as discussed in detail in Jeffrey et al. (2019). Therefore, this combination of drivers most likely explains the higher CH₄ fluxes during C2 when the system (and lilies) had sufficient time to recover, despite lower water column temperatures that would normally reduce microbial metabolism rates. This hypothesis is also supported by the

shift of net positive redox potential of the permanent wetland during C1 (71.7 ± 65 mV) to a strong negative redox potential during C2 (−216 ± 42 mV), indicating that there was a time lag for reducing conditions to recover within the permanent wetland for C2. Further, although aquatic vegetation can facilitate root zone aeration, therefore increasing sedimentary redox potentials, as no aquatic vegetation was present in the permanent wetland during C1, this suggests that water level drawdown was the main driver of the observed redox conditions. This highlights the critical role of antecedent hydrological conditions and how dynamic weather oscillations of drought and floods (a common occurrence of many Australian wetland systems) strongly influence redox potentials, soil geochemistry, and ultimately CH₄ fluxes.

4.4 Implications and conclusions

Within the global wetland CH₄ budget, both subtropical systems and Southern Hemisphere systems are poorly represented (Bartlett and Harriss, 1993; Bastviken et al., 2011) (Fig. 9). Further, the fluxes from seasonal wetlands are poorly constrained (Pfeifer-Meister et al., 2018) due to their intermittent nature and variability of intra-seasonal areal extent, which may compound the fact that natural wetlands have the largest uncertainty of the global methane budget (Kirschke et al., 2013; Saunio et al., 2016). Although the temporal resolution of our study cannot be upscaled to realistic annual estimates, our high-resolution sampling strategy provided insights into daily CH₄ flux rates, revealing distinct differences between different vegetation types across the terrestrial–aquatic wetland boundary. Our CH₄ emissions rates were at the low end of the scale of measurements made in Southern Hemisphere subtropical systems but within the

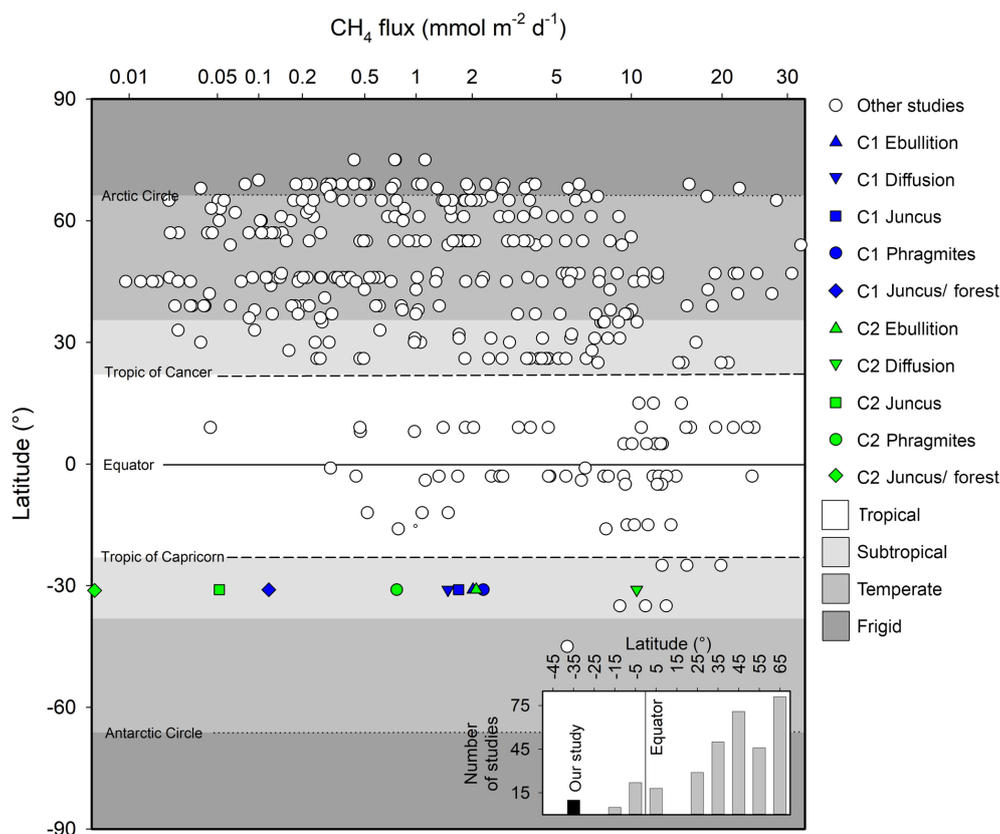


Figure 9. Summary of major CH_4 wetland reviews by Bartlett and Harriss (1993) and Bastviken et al. (2011), as well as modelled fluxes by Cao et al. (1998) adapted from Jeffrey et al. (2019), highlighting latitudinal trends and bias from a variety of wetland systems. Inset figure highlights the number of studies in these reviews by latitudinal increments of 10° poleward of the Equator. Note: x axis scaled to highlight subtle differences between studies.

range of Northern Hemisphere subtropical systems of similar latitudes (Fig. 9).

Although remediating degraded wetlands through re-flooding is a common technique to improve biodiversity, increase C sequestration, and improve downstream water quality issues (Johnston et al., 2004, 2014), our results propose a nuanced dilemma for land use managers, as wetland remediation can potentially have net positive radiative forcing effects on the Earth's climate due to high rates of CH_4 production (Petrescu et al., 2015). This has also been shown to be particularly high during early remediation periods (Hemes et al., 2018). Our results suggest that seasonal wetlands emit less CH_4 on an areal basis than permanent wetlands, yet carbon accumulation in these soils may be lower (Brown et al., 2019). Longer-term studies over annual cycles encompassing seasonal drivers and CH_4 fluxes would further test this hypothesis of the different drivers between seasonal and permanent wetland systems.

Our results also suggest that selective hydrological restoration of wetlands featuring sediments with abundant thermodynamically favourable terminal electron acceptors (i.e. Fe(III) or SO_4^{2-}) may be a (partial) biogeochemical solution

(also suggested by Hemes et al., 2018) to both remediate degraded sites whilst simultaneously mitigating some CH_4 emissions. When Fe(III) and SO_4^{2-} are abundant in anaerobic environments they provide preferential terminal electron acceptors for microbial metabolism and thus limit methanogenesis via competitive exclusion (Acht nich et al., 1995). However, high rates of sulfate reduction coupled with Fe reduction can also lead to the accumulation of metal sulfide minerals, e.g. pyrite and AVS (Johnston et al., 2014). Under permanently saturated and low oxygen conditions, metal sulfides will steadily accumulate and remain relatively benign. However, if the saturated state of remediated sites cannot be maintained, AVS may react with oxygen, resulting in the undesirable production of acidity and low pH conditions. Therefore, the remediation of wetlands for carbon storage should involve careful site selection to both limit CH_4 production and to avoid redox-related geochemical by-products with detrimental environmental effects.

This study has highlighted how sediment geochemistry is intimately related to CH_4 production and consumption. While high sulfate and Fe(III) favour lower CH_4 production, sites featuring more reducing conditions and depleted sul-

fate and Fe(III) favour the highest CH₄ fluxes. Results reveal distinct differences between the areal CH₄ fluxes of four different ecotypes located within a remediated subtropical Australian wetland and indicate high variability between campaigns. By combining novel and well-established techniques we delineated several CH₄ pathways of both seasonal and permanent wetland sources (ebullition, diffusion, and plant-mediated pathways) and linked these to hydrological drivers. This provided evidence that soil geochemistry is an important factor to consider for wetland remediation in the context of CH₄ production and mitigation strategies. The CH₄ emissions results were comparable to other wetlands of similar latitudes and contribute important data for both the understudied Southern Hemisphere wetlands and seasonal subtropical wetland ecotypes.

Data availability. Data used for the composition of this paper can be obtained from <https://data.mendeley.com/datasets/st4vyjc73f/1> (last access: 24 April 2019) (Jeffrey, 2019).

Author contributions. LCJ, DRT, SGJ, and DTM conceived the study. LCJ wrote the first draft. LCJ, DRT, SGJ, and KM conducted the fieldwork. LCJ, DRT, SGJ, DTM, ADLS, and KM reviewed and edited the paper.

Competing interests. The authors declare that they have no conflict of interest.

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