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#### Management Effects on Greenhouse Gas Dynamics in Fen Ditches

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9

#### 10 Abstract

11 Globally, large areas of peatland have been drained through the digging of ditches, generally 12 to increase agricultural production. By lowering the water table it is often assumed that drainage reduces landscape-scale emissions of methane  $(CH_4)$  into the atmosphere to 13 negligible levels. However, drainage ditches themselves are known to be sources of CH<sub>4</sub> and 14 other greenhouse gases (GHGs), but emissions data are scarce, particularly for carbon 15 16 dioxide (CO<sub>2</sub>) and nitrous oxide (N<sub>2</sub>O), and show high spatial and temporal variability. Here, we report dissolved GHGs and diffusive fluxes of CH<sub>4</sub> and CO<sub>2</sub> from ditches at three UK 17 18 lowland fens under different management; semi-natural fen, cropland, and cropland 19 restored to low-intensity grassland. Ditches at all three fens emitted GHGs to the atmosphere, but both fluxes and dissolved GHGs showed extensive variation both 20 seasonally and within-site. CH<sub>4</sub> fluxes were particularly large, with medians peaking at all 21 three sites in August at 120-230 mg m<sup>-2</sup> d<sup>-1</sup>. Significant between site differences were 22 detected between the cropland and the other two sites for CO<sub>2</sub> flux and all three dissolved 23 24 GHGs, suggested that intensive agriculture has major effects on ditch biogeochemistry. Multiple regression models using environmental and water chemistry data were able to 25 26 explain 29-59% of observed variation in dissolved GHGs. Annual CH<sub>4</sub> fluxes from the ditches were 37.8, 18.3 and 27.2 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> for the semi-natural, grassland and cropland, and 27 annual CO<sub>2</sub> fluxes were similar (1100 to 1440 g CO<sub>2</sub>  $m^{-2}$  yr<sup>-1</sup>) among sites. We suggest that 28 fen ditches are important contributors to landscape-scale GHG emissions, particularly for 29 30 CH<sub>4</sub>. Ditch emissions should be included in GHG budgets of human modified fens, particularly where drainage has removed the original terrestrial CH<sub>4</sub> source, e.g. agricultural 31 peatlands. 32

34 Keywords: peatland, carbon dioxide, methane, nitrous oxide, ditch flux, restoration

35

#### 36 **1. Introduction**

Northern peatlands store approximately 547 Pg of carbon (Yu et al., 2010) and 37 38 contribute to the global atmospheric balance of GHGs through the release and uptake of 39 carbon dioxide (CO<sub>2</sub>), methane (CH<sub>4</sub>) and nitrous oxide (N<sub>2</sub>O). Intact peatlands are typically net sinks for CO<sub>2</sub>, and sources of CH<sub>4</sub> and N<sub>2</sub>O (Freeman et al., 1993, Nykänen et al., 1995, 40 41 Smith *et al.*, 2004, Kirschke *et al.*, 2013). On a 100-year timescale CH<sub>4</sub> and N<sub>2</sub>O have global 42 warming potentials (GWP) of 28 and 298, respectively, relative to CO<sub>2</sub> (IPCC, 2013). Insights 43 into biogeochemical cycling in peatlands are therefore important in developing 44 understanding of global GHG dynamics and future climate change.

45 Globally, peatlands have been extensively drained for conversion to agriculture, 46 forestry and peat extraction. Drained lowland fens, such as those of Eastern England, the 47 Netherlands and the Southern Baltic coast are extremely fertile, and are therefore principally converted to intensive agricultural use (Morris et al., 2000). Conversion to 48 agricultural use often includes strict hydrological management, such as the use of 49 50 subsurface irrigation and, in part due to the long-term subsidence which is an inevitable consequence of peat drainage, the active pumping of water around fields (e.g. Morrison et 51 al., 2013). There is now growing interest in the restoration of agricultural fens to wetlands 52 53 (e.g. Höll et al., 2009, Peh et al., 2014), although there are strong commercial factors, as well as food security considerations, that favour their continued agricultural use (Glenk et 54 al., 2014). 55

Drainage and conversion of fens to agricultural use has the capacity to alter the cycling of GHGs. It is generally considered that peatland drainage leads to a decrease in CH<sub>4</sub> emissions (to near-zero values), but increases in CO<sub>2</sub> and N<sub>2</sub>O emissions (Glenn *et al.*, 1993, Martikainen *et al.*, 1995, Alm *et al.*, 1999, Haddaway *et al.*, 2014). Upon draining, peatlands therefore become a diminishing carbon reservoir, releasing carbon into the atmosphere that was fixed over thousands of years.

62 CH<sub>4</sub> fluxes from drained peatlands were previously assumed to be insignificant (IPCC,
63 2006). However, a number of studies have shown that the ditches created during drainage
64 can themselves be significant CH<sub>4</sub> sources (Best & Jacobs, 1997, Sundh *et al.*, 2000,

- 65 Minkkinen & Laine, 2006, Hendriks et al., 2007, Hyvönen et al., 2013), contributing 60-70%
- of total CH<sub>4</sub> emissions in one study (Schrier-Uijl *et al.*, 2010), over 84% in another (Teh *et al.*,
- 67 2011) and with measured fluxes as high as 366 mg CH<sub>4</sub> m<sup>-2</sup> hr<sup>-1</sup> (Schrier-Uijl *et al.*, 2010).
- 68 Where the space between ditches is small, drainage could in theory actually result in a net
- 69 increase in landscape-scale CH<sub>4</sub> fluxes compared to undrained sites (Roulet & Moore, 1995).





71 Figure 1. Schematic of methane transport pathways within ditch systems and surrounding peat.

73 Large ditch CH<sub>4</sub> fluxes are usually associated with productive, high-nutrient, sites 74 with low water flow and high labile carbon inputs (e.g. agricultural grasslands; Best & 75 Jacobs, 1997). Conversely, faster-flowing ditches in nutrient-poor upland bogs typically 76 have small fluxes; Cooper *et al.* (2014) recorded an annual mean CH<sub>4</sub> flux of 59.7 kg CH<sub>4</sub> ha<sup>-2</sup> v<sup>-1</sup> from an open ditch in a blanket bog, and Sirin *et al.* (2012) measured a growing season 77 flux of 9.9 mg CH<sub>4</sub> m<sup>2</sup> d<sup>-1</sup> from ditches in a forested bog. A recent review found mean fluxes 78 for different peat/land-use types varied from approximately 30 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> in forest/semi-79 80 natural peatlands, to 200 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> in tropical deforested peatlands (Evans et al., 2016a). It is important to recognise that methane emissions can occur via different 81 pathways, and the rates of flux via these pathways will have different controls (fig.1). 82 Diffusive/steady emissions result from the CH<sub>4</sub> concentration differential between the ditch 83 and the atmosphere. Wetland plant aerenchyma may provide a chimney through which 84 oxygen is transported into sediment and CH<sub>4</sub> escapes to the atmosphere. Finally, steady 85 emissions may be punctuated by temporally and spatially heterogeneous ebullition, which 86 87 can contribute significantly to net CH<sub>4</sub> fluxes (Vermaat et al., 2011). The importance of 88 ditches in GHG cycling has therefore been recognised by the IPCC and incorporated into their guidelines (IPCC, 2014). 89

90 As well as CH<sub>4</sub>, drainage ditches emit N<sub>2</sub>O (Reay et al., 2003, Teh et al., 2011, Hyvönen et al., 2013). Some ditches have been found to emit CO<sub>2</sub> (Best & Jacobs, 1997, 91 Sundh et al., 2000, Teh et al., 2011, Hyvönen et al., 2013), which others with emergent 92 93 vegetation have sometimes been observed to fix CO<sub>2</sub> (e.g. Vermaat et al., 2011). However, 94 whilst ditches appear to be consistent hotspots for CH<sub>4</sub> emissions, CO<sub>2</sub> and N<sub>2</sub>O fluxes are of 95 a considerably smaller magnitude in terms of their overall contribution to GHG emissions, and are typically more similar to fluxes from drained peat adjacent to ditches (Evans et al., 96 2016a). For example, Hyvönen et al. (2013) found ditches in a boreal cutaway peatland 97 98 being used to cultivate *Phalaris arundinacea* contributed just 1% and 5% of total ecosystem 99 emission of  $N_2O$  and  $CO_2$ .

100 Internationally, there is a lack of information on GHG emissions from drainage 101 ditches; in a recent review of published studies, a total of just 19 studies were identified in 102 which peatland CH<sub>4</sub> emissions had been reported, for a total of 69 individual peatland sites 103 where CH<sub>4</sub> was measured (Evans et al., 2016a). The same analysis suggested that studies of 104 CO<sub>2</sub> and N<sub>2</sub>O are still too few to allow the data to be collated in a meaningful way. Just two studies to date have reported CH<sub>4</sub> fluxes from ditches in the UK. In contrast to this dearth of 105 106 information on ditches, numerous studies have looked at GHG emissions associated with 107 other freshwaters. For instance, Cole et al. (2007) noted that carbon emissions from lakes and rivers could be approximately 0.8 Pg C y<sup>-1</sup>; enough to exert effects on regional budgets, 108 despite these features occupying small areas. Similarly, Bastviken et al. (2011) suggested 109 110 that CH<sub>4</sub> emissions from inland waters have the capacity to offset 25% of the terrestrial carbon sink, whilst Deemer et al. (2016) calculate that reservoirs emit 1.5% of global 111 anthropogenic CO<sub>2</sub>-equivalent emissions from CO<sub>2</sub>, CH<sub>4</sub> and N<sub>2</sub>O. Considering N<sub>2</sub>O, rivers 112 and estuaries could account for 20% of global anthropogenic emissions (Seitzinger & Kroeze, 113 114 1998).

To help address this knowledge gap, we carried out seasonal fieldwork for one year in ditches at three lowland fens in East Anglia, England. Each site was under a different management regime: 1) a semi-natural fen under conservation management; 2) former cropland that has been restored to extensive grassland, and; 3) intensive deep-drained cropland. We measured dissolved GHGs within ditches, diffusive fluxes of CO<sub>2</sub> and CH<sub>4</sub> from ditches, and a variety of physical ditch attributes and water chemistry determinands. Our aim was to quantify the differences in GHGs between and within sites, and across seasons,

and to attempt to elucidate the drivers behind GHG dynamics.

123

#### 124 **2. Materials and methods**

#### 125 *2.1. Field sites*

126 All three field sites were located in East Anglia, in Eastern England. This region was once the largest area of lowland fen peatland in the UK, covering several thousand square 127 kilometres. Since the 17<sup>th</sup> century, drainage of the land resulted in the loss of most of the 128 129 natural fenland, with only a handful of intact fragments remaining. The principal land use of 130 the drained areas is intensive arable and horticultural agriculture. The drainage and 131 conversion of the fens has resulted in extensive peat wastage, with much of the original deep peat area now reduced to a dense, thin intermixed organic and mineral layer 132 133 (Hutchinson, 1980, Burton and Hodgson, 1987). The altitude of the land is close to (and in 134 many areas below) sea level. Mean annual rainfall in the area is 574 mm, and mean annual temperature is 10.1 °C (data from UK Met Office station in Mepal, within 30 km of all study 135 136 sites). The sites were:

137

138 1. Sedge Fen (semi-natural fen). 52.31 N, 0.28 E. Area = 61 ha. Sedge Fen is part of the Wicken Fen National Nature Reserve. Peat depth is 3.8 m, bulk density is 0.37 g cm<sup>-3</sup>, C/N is 139 15.8 (Evans et al., 2016b). Vegetation comprises reedbeds dominated by Cladium mariscus 140 141 and Phragmites australis, with some Phalaris arundinacea and Calamagrostis canescens (Eades, 2016), as well as areas of fen carr dominated by Rhamnus cathartica and Frangula 142 143 alnus (Rowell, 1986). The fen cannot be considered to be 'pristine' as it contains numerous internal ditches, and the reedbeds are cut on a three year rotation. However, the site 144 145 contains vegetation and peat that is characteristic of an intact site, and has never been converted to other land-uses. 146

147

2. Baker's Fen (extensive grassland). 52.30 N, 0.29 E. Area = 56 ha. Baker's Fen is part of the
wider Wicken Fen area. Historically, the fen was drained and used for arable agriculture,
resulting in extensive peat wastage and loss of organic soil. Soil depth is now less than 50
cm, bulk density is 1.06 g cm<sup>-3</sup>, C/N is 19.7 (Evans *et al.*, 2016b), and organic content is low
(measured as 13-18 % loss on ignition by Stroh *et al.*, 2013). The site was removed from

arable use and re-seeded with an unknown "grass mixture" in 1995 and 1996, and is
undergoing "open-ended" restoration (Hughes *et al.*, 2011); river water is pumped onto the
site in autumn and winter to inundate it, and highland cattle and wild horses graze it. Much
of the fen consists of species-poor, flood-plain pasture. Plant species vary across the site
according to variations in hydrology and nutrient status, but include *Carex otrubae*, *Arrhenatherum elatius*, *Agrostis stolonifera*, *Cirsium arvense*, *Poa trivialis* and several *Juncus*species (Eades, 2016). *C. mariscus* and *P. australis* occur in some of the ditches.

160

161 3. Rosedene (cropland). 52.52 N, 0.49 E. Field area = 8.7 ha. The cropland site consists of 162 ditches that surround a field near Methwold Hythe. The field is part of a much larger area (~ 163 90 km<sup>2</sup>) of drained fen that is now under intensive arable cultivation, and is bounded by rivers and canals. Peat depth is 1 m, bulk density is 0.32 g cm<sup>-3</sup>, and C/N is 15 (Evans *et al.*, 164 165 2016b). The hydrology of the site is highly managed; the fields contain subsurface pipes at 1 166 m depth to aid irrigation and drainage, and water is actively pumped round field perimeter 167 ditches in order to maintain water levels within the field, removing water during wet periods and providing irrigation water during dry periods. During 2015, the study site was 168 used to cultivate celery (Apium graveolens). This site is 28 km from the other two sites. 169 170

All three sites formed part of a larger study of GHG emissions from a total of fifteen lowland
peatland sites at located across six regions of England and Wales, which included a broad
suite of eddy covariance and static chamber gas flux, hydrological and water quality
measurements. The results of this large-scale study are reported elsewhere (Evans et al.,
2016b).

176

#### 177 *2.2. Sampling*

The sites were visited on four occasions in 2015 in March, May, August and October. Because of proximity, the intact and restored site could be visited on the same day, whilst the agricultural site was visited within three days. The sampling dates were as follows: 11<sup>th</sup> March – semi-natural/grassland, 12<sup>th</sup> March – cropland; 5<sup>th</sup> May – semi-natural/grassland, 6<sup>th</sup> May – cropland; 17<sup>th</sup> August – cropland, 20<sup>th</sup> August – semi-natural/grassland; 12<sup>th</sup> October – cropland, 15<sup>th</sup> October – semi-natural/grassland. At each site, ditch sampling locations were selected with the aim of covering a large area, and were selected on a non185 random basis according to where measurements from the ditch could easily be taken. For the semi-natural site, we sampled along a 910 m length of ditch network (i.e. all sampling 186 points were hydrologically connected) and then onto a ditch that bounded the edge of the 187 188 fen (fig. 2). Similarly, all ditch locations at the grassland site were hydrologically connected, 189 with a ditch distance of 1200 m between farthest sampling points. At the cropland site the 190 ditch ran continuously round a field, with junctions connecting to other ditches at field 191 corners. The sampling locations here ran for 1200 m. The number of sampling locations for each site was: semi-natural = 13, grassland = 11, cropland = 10. The same sampling 192 193 locations were used for each of the four seasonal visits.

194 A range of measurements were taken at each sampling location. Environmental and 195 physical measurements were: air temperature, water temperature, atmospheric pressure, and water depth. A 50 ml water sample for water chemistry analysis was collected in a 196 197 polypropylene vial. A sample for dissolved GHG analysis was collected using the headspace 198 method (Hope et al., 2004); 30 ml of ditch water was collected in a 60 ml plastic syringe and 199 equilibrated with 30 ml of ambient air by shaking for approximately 60 seconds, and 12 ml of headspace was then collected in a 12 ml borosilicate glass vial. Fluxes of CH<sub>4</sub> and CO<sub>2</sub> 200 201 were measured in real time in the field using a floating chamber (0.6 x 0.6 x 0.3 m) that was 202 shrouded to exclude light. Buoyancy for the chamber was provided by two 2 l plastic bottles filled with air, and the chamber was placed carefully on the water to minimise disturbance. 203 Emergent vegetation was excluded (e.g. P. australis), but some sampling points contained 204 205 floating algae that will have contributed to fluxes. The chamber was connected to a Los Gatos Ultraportable Greenhouse Gas Analyzer. The chamber was deployed until a linear 206 flux was observed, and this was typically 1-5 minutes. Whilst there has been some criticism 207 of the use of floating chambers, flow rates in the ditches we studied where either extremely 208 209 low or absent (i.e. chambers did not drift away) and therefore our measurements are likely to be robust (see Lorke et al., 2015). 210



N Micken Code 1 2 3 4 5 6 1 0 m



215 Figure 2. Maps of the semi-natural site (top panel), extensive grassland site (middle panel) and cropland site 216 (bottom panel). Red dots mark numbered sampling locations, blue lines mark ditches/watercourses. For the 217 semi-natural and extensive grassland sites, blue arrows mark where water is pumped onto site from Wicken 218 Lode.

219

#### 2.3 Analysis 220

Electrical conductivity (EC) and pH were measured on the 50 ml water sample. The 221 sample was then passed through a nylon filter at 0.45 µm for further analysis. Dissolved 222 223 organic carbon (DOC) and inorganic carbon (DIC) were analysed using a Shimadzu TOC Analyzer. DOC was measured as non-purgeable organic carbon (NPOC). Absorbance was 224 measured at 280 nm using a Thermo Spectronic Helios Gamma Spectrophotometer. This 225 226 was normalised against DOC concentration to give the specific ultraviolet absorbance (SUVA). SUVA is commonly measured at 254 nm, although high nitrate (NO<sub>3</sub><sup>-</sup>) 227 concentrations can interfere at wavelengths < 250 nm (Wang & Hsieh, 2001). Considering 228

the potential for high NO<sub>3</sub><sup>-</sup> concentrations in surface waters in areas of intensive agriculture a SUVA wavelength of 280 nm was selected. NO<sub>3</sub><sup>-</sup> was measured using a NICO 2000 ionselective electrode and appropriate standards. Dissolved CH<sub>4</sub> and CO<sub>2</sub> were analysed using a Los Gatos Ultraportable Greenhouse Gas Analyzer equipped with a sampling loop (Baird *et al.*, 2010). Dissolved N<sub>2</sub>O was analysed on an Ai Cambridge GC94 equipped with an Electron Capture Detector (ECD).

Floating chamber fluxes of CH<sub>4</sub> and CO<sub>2</sub> fluxes were calculated according to
 Denmead (2008), using the modified formula:

$$F_{g} = \frac{1}{A} \frac{dg_m}{dt}$$

where  $F_g$  is the flux of CH<sub>4</sub> or CO<sub>2</sub> (M L<sup>-2</sup> T<sup>-1</sup> – mg m<sup>-2</sup> day<sup>-1</sup>), A is the area inside the 238 chamber  $(L^2 - m^2)$ , gm is the mass of gas in the chamber (M - mg), and t is time (T - days). 239 240 Fluxes were calculated using a linear regression between time and chamber gas mass, and 241 accepted if this regression was significant ( $p \le 0.05$ ). Fluxes that were not significant were assumed to be zero. Although it is usual to specify a cut-off value for the R<sup>2</sup> of the flux 242 regression (below which value fluxes are rejected) we did not take this approach, because 243 the high-frequency measurements provided by the analyser allowed detection of small but 244 clearly non-zero (significant) fluxes despite high short-term scatter (low R<sup>2</sup>). However, of 245 the 253 fluxes that were significant, only 12 had an R<sup>2</sup> under 0.7. Fluxes were corrected for 246 atmospheric pressure and temperature measured during each individual chamber 247 248 deployment. Because of the short deployment time we assumed that pressure and temperature remained steady during flux measurement. Piston velocity was calculated 249 using the standard formula (e.g. Gålfalk et al., 2013): 250

251

$$F = k x \left( C_{aq} - C_{eq} \right)$$

where F is the CH<sub>4</sub> flux, k is the piston velocity, C<sub>aq</sub> is the dissolved concentration of CH<sub>4</sub>, and C<sub>eq</sub> is the theoretical dissolved concentration if the water is in equilibrium with the air (calculated via Henry's Law). The formula was rearranged to give k.

255

#### 256 2.4 Statistics

257 Statistical analysis was carried out in SPSS to determine if differences in GHGs and 258 piston velocity (CH<sub>4</sub> and CO<sub>2</sub> flux, and dissolved concentrations of CH<sub>4</sub>, CO<sub>2</sub> and N<sub>2</sub>O) were 259 present between sites. All six variables failed Levene's test for homogeneity of variance. 260 Kolmogorov-Smirnov tests were used to check for normal distributions. All six variables were not normally distributed, so transformations were sought to resolve this. CO<sub>2</sub> flux was 261 transformed by square root transformation, and dissolved CO<sub>2</sub> was normalised by cube root 262 263 transformation. Remaining variables could not be transformed to fit normal distributions. 264 As such, a linear mixed model was used to test for differences between sites, using time as a 265 repeated measure, and with Bonferroni correction for pairwise comparisons. Stepwise regression analysis was used as an exploratory test to look for relationships between 266 267 dissolved CH<sub>4</sub> and CO<sub>2</sub> and the following variables: ditch water temperature, ditch depth, 268 EC, absorbance at 280 nm, NO<sub>3</sub><sup>-</sup>, DOC, SUVA, DIC, peat depth of the terrestrial fen, C:N, and 269 water table in the terrestrial fen at the time of sampling (data for this was taken from Evans 270 et al., 2016b). For the dissolved CO<sub>2</sub> model, pH was not used as an explanatory variable due to the fact that dissolved CO<sub>2</sub> and pH are interlinked (e.g. Abril et al., 2015). Differences 271 272 were considered significant when  $p \le 0.05$ .

273

#### 274 **3. Results**

#### 275 3.1. Water chemistry, ditch depths and environmental data

276 Table 1 displays a range of environmental and biogeochemical/physical data for the 277 three sites through the year. Ditch water depths at the semi-natural site were consistently deep through the year (60 cm and above). Depths at the grassland site were generally 278 shallow (~ 20cm), as were those at the cropland site, except during August when the mean 279 280 was 60 cm. For all sites water and air temperature was highest during August. Mean ditch pH at all three sites was between 7.2 and 8.0, but EC was more variable both seasonally and 281 between sites (intact < agricultural < restored).  $NO_3^-$  concentrations peaked in May at the 282 cropland site (18 mg l<sup>-1</sup>), presumably due to the use of fertilisers. At the grassland site NO<sub>3</sub><sup>-</sup> 283 284 was low ( $\leq 5 \text{ mg l}^{-1}$ ) except in March when the mean was 19 mg l<sup>-1</sup>. The fen is rewetted during autumn and winter using high- NO<sub>3<sup>-</sup></sub> river water, and the high concentration in March 285 is a legacy of this rewetting. DOC concentrations were moderately high at the semi-natural 286 and grassland sites (mean ~ 30 mg  $l^{-1}$ ), but were lower by a third at the cropland. 287

288

#### 289 3.2. Differences in ditch fluxes between and within sites

There was no significant difference in  $CH_4$  flux between sites, but a significant (p < 0.001) difference was found for  $CO_2$  flux between the cropland and other two sites (fig.3).

Median CH<sub>4</sub> fluxes were relatively low in March ( $\leq 10.5 \text{ mg m}^{-2} \text{ d}^{-1}$ ). Fluxes stayed low in 292 May at the grassland and cropland fens, but were higher (80 mg m<sup>-2</sup> d<sup>-1</sup>) at the semi-natural 293 fen. Median CH<sub>4</sub> fluxes peaked in August at all three sites, at 120-230 mg m<sup>-2</sup> d<sup>-1</sup>. Highest 294 individual fluxes at each site were: 3650, 25400 and 7430 mg m<sup>-2</sup> d<sup>-1</sup> for the semi-natural 295 (May), grassland (August) and cropland (August) site respectively. CO<sub>2</sub> flux was relatively 296 stable at the semi-natural site at 2050-3250 mg m<sup>-2</sup> d<sup>-1</sup>, but fluctuated at the other two sites, 297 peaking at 6600 mg m<sup>-2</sup> d<sup>-1</sup> in August at the grassland site, and at 4760 mg m<sup>-2</sup> d<sup>-1</sup> in October 298 at the cropland site. Highest individual fluxes at each site were: 9580, 16800 and 13800 mg 299 m<sup>-2</sup> d<sup>-1</sup> for the semi-natural, grassland and cropland sites respectively, and were all recorded 300 301 in August. Differences were also apparent within sites, and median fluxes for each 302 individual sampling location are shown in fig.4.

There was considerable variation apparent in piston velocities between sites and months, but none of these differences was significant (table 2).

305

#### 306 *3.3. Annual ditch fluxes*

To calculate annual mean fluxes for 2014, a simple time-weighted median approach was used, using the medians from fig.3. For CH<sub>4</sub>, these produced estimates of 37.8, 18.3 and 27.2 g CH<sub>4</sub> m<sup>-2</sup>yr<sup>-1</sup> for the semi-natural, grassland and cropland sites respectively, with respective standard errors of 74.6, 244, and 97.3 g CH<sub>4</sub> m<sup>-2</sup>yr<sup>-1</sup>. For CO<sub>2</sub> the annual fluxes were 1100, 1170 and 1440 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup> for the semi-natural, grassland and cropland sites respectively, with respective SEs of 225, 340 and 312 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup>.



315 Figure 3. Median ditch fluxes of CH<sub>4</sub> and CO<sub>2</sub> measured using floating chambers at the three sites. Error bars

represent first and third quartiles. . Note that the error bar for CH<sub>4</sub> in August at the cropland site exceeds the

317 scale. There was a significant difference ( $p \le 0.001$ ) for CO<sub>2</sub> fluxes between the cropland and other two sites.



318

Figure 4. Median CH<sub>4</sub> (left) and CO<sub>2</sub> (right) fluxes for each individual numbered sampling point, grouped by
 site. Fig.2 displays numbered sampling points on site maps.

#### 322 3.4. Differences in dissolved gases between sites

Significant differences were observed for dissolved CO<sub>2</sub> between the cropland and 323 other two sites (p < 0.001). For dissolved CH<sub>4</sub>, significant differences were found between 324 the cropland and semi-natural fen (p < 0.01) and the cropland and grassland (p < 0.05). For 325  $N_2O$ , a significant difference was found between the cropland and other two sites (p < 326 0.001) (fig.5). Median CH<sub>4</sub> concentrations were below 0.1 mg  $l^{-1}$ , except for a spike of 0.43 327 mg l<sup>-1</sup> at the cropland site in August. Median dissolved CO<sub>2</sub> at the semi-natural site showed 328 no obvious seasonal variation (range 4.8-9.0 mg l<sup>-1</sup>), whilst there was an increase through 329 330 the year at the grassland site (1.7-7.5 mg  $l^{-1}$ ). Dissolved CO<sub>2</sub> at the cropland site also peaked later in the year (15 mg l<sup>-1</sup> in August and October). Median N<sub>2</sub>O concentrations were under 331

1.5 μg l<sup>-1</sup> at the semi-natural and grassland sites. At the cropland site N<sub>2</sub>O concentrations
were generally higher. Differences were apparent within sites, and median concentrations
for each individual sampling location are shown in fig.6.

- 335
- 336





338 Figure 5. Median ditch dissolved concentrations of CH<sub>4</sub>, CO<sub>2</sub> and N<sub>2</sub>O at the three sites. Error bars represent

first and third quartiles. Note that the error bar for CH<sub>4</sub> in August at the cropland site exceeds the scale.

340 There were significant differences between the cropland and other two sites for CO<sub>2</sub> (p < 0.001) CH<sub>4</sub> (cropland

341 vs semi-natural p < 0.01, cropland vs grassland p < 0.05) and  $N_2O$  (p < 0.001).





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#### 346 3.5. Drivers of dissolved GHGs

347 Significant regression models were produced for both dissolved  $CH_4$  and  $CO_2$ , with 348 respective  $R^2$  values of 0.29 and 0.50. Table 3 displays the p values and slope coefficients 349 used in each model.

There was a significant positive linear relationship between dissolved N<sub>2</sub>O concentrations and NO<sub>3</sub><sup>-</sup> (p < 0.001, R<sup>2</sup> = 0.33) but an improved fit was found between dissolved N<sub>2</sub>O and the DOC:NO<sub>3</sub><sup>-</sup> ratio, with N<sub>2</sub>O concentrations increasing as the ratio decreased (fig.7). Apart from three clear outliers (which were not removed from the analysis), dissolved N<sub>2</sub>O concentration did not rise above 1.5 µg l<sup>-1</sup> until DOC:NO<sub>3</sub><sup>-</sup> fell below 5. This relationship was consistent across all three sites.



Figure 7. Relationship between dissolved N<sub>2</sub>O and DOC:NO<sub>3</sub><sup>-</sup> ratio for all individual samples. Red diamonds =
 semi-natural fen, black circles = grassland, blue triangles = cropland. Trend line is 3<sup>rd</sup> order polynomial.
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#### 360 4. Discussion

#### 361 *4.1. Site characteristics*

There were physical and biogeochemical differences in the ditches of the three fen sites. The ditches at the semi-natural site were deepest whilst those at the grassland site were shallowest, reflecting the difficulties in keeping this grassland site wet, as noted by Peh *et al.* (2014). Ditch water levels at the cropland site were also shallow, but were raised for irrigation during the peak of the growing season; this demonstrates the high degree of water management to maximise arable production (Morrison *et al.*, 2013).

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#### 369 *4.2. CO*<sup>2</sup> *and CH*<sup>4</sup> *fluxes*

We no significant differences between sites for CH<sub>4</sub> fluxes, but fluxes were CO<sub>2</sub> fluxes were significantly higher at the cropland compared to the grassland and semi-natural sites.. There were seasonal patterns in fluxes of CH<sub>4</sub>; emissions peaked at all three fens in August at which time they were not significantly different. It is likely that these high fluxes are due to the effect of summer temperatures on methanogenesis (Dunfield *et al.*, 1993). There was 375 extensive within-site variation in gas fluxes, particularly for CH<sub>4</sub>. For instance, at the seminatural fen, sample point 1 had CH<sub>4</sub> fluxes an order of magnitude higher than the adjacent 376 sample point 2. Sample point 1 was close to the wind pump that pumps river water onto 377 378 the fen, and it could be that the mixing between low DOC/high NO<sub>3</sub><sup>-</sup> river water and high 379 DOC/low NO<sub>3</sub><sup>-</sup> fen water produces a 'hotspot' of organic carbon processing resulting in CH<sub>4</sub> 380 production (sensu Palmer et al., 2016). The lowest CH<sub>4</sub> fluxes were recorded at a ditch that 381 bounded the edge of the fen, which displayed lower DOC concentrations and higher NO<sub>3</sub><sup>-</sup> 382 concentrations, suggesting that this ditch was connected to the river, and contained less 383 organic substrates for methanogenesis. At the grassland site, CH<sub>4</sub> fluxes were highest at 384 adjacent sample points 7 and 8 though this was not obviously related to any measured 385 variables, e.g. EC, DIC and DOC were not elevated at these locations. At the cropland site, sample point 10 was extremely high compared to the other points. This ditch was shaded 386 387 by a dense cover of trees and was near to a dead-end in the ditch system. This point had 388 elevated levels of EC (25% higher compared to the mean of the other sample points) and 389 DOC (114% higher), and it is likely that standing water here leads to an accumulation of 390 organic matter and stagnation, and hence higher rates of methanogenesis (fig.1). This 391 hypothesis is supported by the fact that dissolved CH<sub>4</sub> concentrations at this location were 392 higher than any other sampling point at any site (fig. 6).

393 Although we only sampled four times within a year, our design featured large numbers of sampling points per site, and different sites were sampled at the same times of 394 395 year. Calculated mean fluxes may not therefore be an accurate representation of the annual values, but should provide a reasonable representation of between-site differences. 396 Mean CH<sub>4</sub> fluxes followed the order semi-natural>cropland>grassland. The mean flux for 397 the semi-natural fen, 38 g  $CH_4$  m<sup>-2</sup> yr<sup>-1</sup>, falls within the range of other reported fluxes from 398 ditches in semi-natural peatlands; e.g. 12 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> and 164 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> from drained 399 boreal fens (Glagolev et al., 2008, Minkkinen & Laine, 2006). The only reported annual ditch 400 CH<sub>4</sub> flux from a temperate semi-natural site are 5.5 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> from a UK upland blanket 401 bog (Cooper et al., 2014). The considerably higher flux reported from our semi-natural site 402 therefore shows the effect of nutrient status on ditch emissions. It has been suggested that 403 ditch CH<sub>4</sub> emissions increase as land-use intensity increases (Evans et al., 2016a) but our 404 405 data do not show this. The flux from our grassland site, 18 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> is low compared to values such as 43, 66, 77 and 70 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> from other low-intensity grasslands (Schrier-406

407 Uijl et al., 2010, Vermaat et al., 2011, McNamara, 2013, Hendricks et al., 2007), although van den Pol-van Dasselaar et al. (1999) recorded an annual flux of just 11 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup>. To 408 409 our knowledge, our annual ditch flux calculation of 27 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> for the cropland is, along 410 with the fluxes in our broader project report (Evans et al., 2016a), the first annual flux 411 estimate for a temperate peatland under agriculture. There are several possible reasons for 412 the highest annual flux being observed at the semi-natural site. Firstly, subsidence at our grassland site has resulted in the loss of the majority of peat soil, and it may be that the low 413 organic content of the soil has led to a reduction in CH<sub>4</sub> production. This could be especially 414 415 relevant if CH<sub>4</sub> is produced in the saturated peat, then transported laterally and degassed 416 from ditches (e.g. fig.1); the grassland site dries out completely, presumably resulting in zero 417 methanogenesis, whilst the water table remains in the peat at a deep level at the cropland, making this a plausible hypothesis. Secondly, the semi-natural site is likely to have a well-418 419 established methanogenic community compared to the other two sites where severe 420 drainage and loss of peat (Stroh et al., 2013) may have disrupted the microbial communities 421 (Jerman et al., 2009). Thirdly, the ditches at the semi-natural site were relatively deep, and depth fluctuations were minimal compared to the other two sites. This could lead to the 422 423 formation of anoxic conditions, thus stimulating CH<sub>4</sub> emissions and reducing oxidation in the 424 water column (O<sub>2</sub> measurements on future sampling campaigns would help to resolve this). Finally, Vermaat et al. (2011) recorded more ebullition in ditches sheltered by reed beds. It 425 is therefore possible that steady ebullition contributed to the high fluxes at the semi-natural 426 fen, as well as being responsible for the individual high fluxes (e.g. 25400 and 7430 mg m<sup>-2</sup> d<sup>-</sup> 427 <sup>1</sup>) that were observed at the grassland and cropland. Ebullition from ditches in a Finnish 428 mire measured using bubble traps was 3-37 mg m<sup>-2</sup> d<sup>-1</sup>, and was negligible (0.2-2.3% 429 430 compared to diffusive emissions) in flowing ditches but substantial in ditches with standing 431 water (10-22% of diffusive flux) (Minkkinen et al., 1997, 2006). Vermaat et al. (2011) calculated ditch ebullition by interpreting steep, short-term increases in CH<sub>4</sub> concentration 432 433 in a floating chamber as evidence of bubbling, and stated that approximately 50% of total flux was due to ebullition. Other research using bubble traps has shown that ebullition in 434 wetland and agricultural streams can equal the diffusive flux (Wilcock & Sorrell, 2008, 435 Crawford et al., 2014). More measurements of ebullition in ditches are clearly needed. 436 437 Unlike CH<sub>4</sub>, annual CO<sub>2</sub> fluxes did increase with land-use intensity, in the order semi-438 natural<grassland<cropland. Estimates of annual ditch CO<sub>2</sub> fluxes are lacking from the

literature, but scaling up the measurements of Vermaat et al. (2011) would produce annual 439 fluxes of 1050 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup> for ditches in reed beds, and 1310 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup> for ditches in 440 rough pasture. Our semi-natural site is therefore similar, with a flux of 1100 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup>, 441 although our grassland annual flux was 1170 g CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup>. Our median CO<sub>2</sub> fluxes ranged 442 from 488 mg m<sup>-2</sup> d<sup>-1</sup> to 8000 mg m<sup>-2</sup> d<sup>-1</sup>, and are therefore similar to those reported by 443 444 Schrier-Uijl et al. (2011), Teh et al. (2011) and Hyvönen et al. (2013). CO<sub>2</sub> fluxes at the seminatural site displayed less seasonality which may be a function of the deeper ditches 445 minimising temperature increases in the basal peat, and therefore suppressing productivity 446 447 (McEnroe *et al.*, 2009).

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#### 449 *4.3. Dissolved GHGs*

For dissolved CH<sub>4</sub>, CO<sub>2</sub> and N<sub>2</sub>O, we found significant differences between the cropland compared to the grassland and semi-natural sites. This suggests that intensive agriculture has affected the biogeochemistry of the cropland ditches. Some sampling locations showed similar concentrations of dissolved gases, and this could be due spatial autocorrelation in dissolved GHGs (e.g. Chapra & Di Toro, 1991). This was most obvious for CH<sub>4</sub> at the semi-natural site, and N<sub>2</sub>O at the semi-natural site and grassland.

456 Once pH had been removed as a predictive variable, we were able to account for 29% of temporal and spatial (within and between site) variability in CH<sub>4</sub>, and 50% of 457 variability in CO<sub>2</sub>. For dissolved CO<sub>2</sub> there were positive relationships with depth to water 458 459 table, DIC concentration, SUVA, and ditch depth. A deeper water table within the peat should result in increased decomposition, with CO<sub>2</sub> then exported laterally into ditches. A 460 negative relationship between CO<sub>2</sub> and water depth has been found for pools in natural 461 peatlands (McEnroe et al., 2009), and so our contrary finding could be due to the high 462 463 degree of management at these fens; e.g. irrigation at the cropland reversed the natural seasonality in ditch depth and doubled the water level of the ditches in August, which 464 465 coincided with the growing season increase in dissolved CO<sub>2</sub>.

Dissolved CH<sub>4</sub> concentrations fell within the same range as those in agricultural
streams (0.001-0.4 mg L<sup>-1</sup>, Wilcock & Sorrell, 2008), and ditches in agricultural peatlands
(maximum of 0.04 mg L<sup>-1</sup>, Schrier-Uijl *et al.*, 2011). They were of the same magnitude as
0.022 mg L<sup>-1</sup> which was the calculated mean fluvial CH<sub>4</sub> concentration from 111 published
studies (Stanley *et al.*, 2016). Dissolved CH<sub>4</sub> correlated positively with air temperature, NO<sub>3</sub><sup>-</sup>,

DIC, and depth to water table (in the fen/field), and negatively with EC. Higher 471 temperatures could stimulate methanogenesis, leading to increased concentrations of CH<sub>4</sub>. 472 The positive correlation between dissolved CH<sub>4</sub> and depth to water table may, in part, be 473 474 due to the confounding effect of seasonality; i.e. water tables were lower in the growing 475 season when ditches become depleted in oxygen, leading to higher rate of methanogenesis. 476 The positive correlation between  $NO_3^-$  and  $CH_4$  is unexpected, as  $NO_3^-$  inhibits 477 methanogenesis (Watson & Nedwell, 1998) and, as an electron acceptor, allows denitrifying bacteria to favourably out-compete methanogens (Le Mer & Roger, 2001). One possible 478 479 explanation is that increased NO<sub>3</sub><sup>-</sup> levels are associated with increased ammonium 480 concentrations at the semi-natural fen (Conrad & Rothfuss, 1991), and the inhibitory effect 481 of ammonium on methanotrophy is larger than the inhibitory effect of NO<sub>3</sub><sup>-</sup> on methanogenesis. It may be that high nutrient levels associated with NO<sub>3<sup>-</sup></sub> could coincide 482 483 with inputs of labile organic matter, particularly at agricultural sites, thus stimulating 484 methanogenesis when other electron acceptors have been depleted in the sediment. 485 Alternatively, as discussed in section 4.2, it could be that CH<sub>4</sub> is produced in the saturated peat and then transported laterally into the ditch; i.e. methanogenesis occurs in zones 486 487 distant from potential NO<sub>3</sub><sup>-</sup> inhibition. Schade *et al.* (2016) did find a weak negative 488 correlation between NO<sub>3</sub><sup>-</sup> and CH<sub>4</sub> in a low NO<sub>3</sub><sup>-</sup>/high DOC stream but found no correlation in a high NO<sub>3</sub><sup>-</sup>/low DOC stream or in a high NO<sub>3</sub><sup>-</sup>/high DOC stream. Similarly, Crawford *et al.* 489 (2016) found no evidence that NO<sub>3<sup>-</sup></sub> inhibited CH<sub>4</sub> production or emission in streams, and, in 490 491 line with our hypothesis above, suggested that methanogenesis could be spatially removed from high NO<sub>3</sub><sup>-</sup> concentrations. The absence of ditch depth from the CH<sub>4</sub> model is 492 interesting as negative relationships between CH<sub>4</sub> flux and depth have been noted 493 494 previously, although these are sometimes low; e.g. McEnroe et al. (2009) reported an R<sup>2</sup> value of 0.23 for pools, and Vermaat et al. (2011) found an R<sup>2</sup> of 0.15 for ditches. Pelletier 495 et al. (2007) found both negative and positive relationships between CH<sub>4</sub> flux and depth in 496 497 pools at different peatlands, and postulated that ebullition could be a confounding variable. The active water management at some sites could also be a confounding factor; as 498 previously mentioned this management removes the natural seasonality in ditch depth. 499 Finally, it is worth considering that wind speed may play a role in GHG dynamics. However, 500 501 the ditches at our sites are predominantly sheltered by reedbeds or banks and, as previously noted, the floating chamber did not drift, suggesting that wind speed was low on samplingdays.

504 Dissolved  $N_2O$  was present in the ditches at all three fens, but was low at the semi-505 natural site. Concentrations were only high at the grassland site in March, but were high for 506 most of the year at the cropland site, presumably due to the application of fertilisers to 507 adjacent fields. Positive relationships between dissolved N<sub>2</sub>O and N<sub>2</sub>O flux have been demonstrated in rivers (Yang et al., 2011). Diffusive fluxes of N<sub>2</sub>O have been shown to occur 508 in oxygenated waters and it therefore seems highly probable that ditches at all three fens 509 510 were sources of  $N_2O$  to the atmosphere. Wilcock & Sorrell (2008) measured N2O 511 concentrations in agricultural streams between 0.26-28.5  $\mu$ g  $l^{-1}$ , considerably higher than our maximum individual measurements of 6.15 µg l<sup>-1</sup>, whilst concentrations in a eutrophic 512 river have been reported as 0.66-1.14  $\mu$ g l<sup>-1</sup> (Silvennoinen *et al.*, 2008). Sturm *et al.* (2014) 513 recorded average concentrations of N<sub>2</sub>O in lake surface water as 0.61  $\mu$ g l<sup>-1</sup> and 0.74  $\mu$ g l<sup>-1</sup>, 514 515 similar to median concentrations at our grassland and semi-natural site, although 516 concentrations at our cropland were higher. The authors also measured N<sub>2</sub>O fluxes, with averages of 3.7 and 5.3  $\mu$ g m<sup>-2</sup> hr<sup>-1</sup>. Reay *et al.* (2003) reported a relationship between N<sub>2</sub>O 517 fluxes and dissolved N<sub>2</sub>O in UK agricultural ditches; applying that relationship to our data 518 519 allows estimates of median flux for each fen to be calculated as 300, 210, and 1150  $\mu$ g m<sup>-2</sup> hr<sup>-1</sup> for the semi-natural, grassland and cropland sites respectively. These fluxes at the 520 semi-natural and grassland sites are similar to those reported by Teh et al. (2011) for ditches 521 522 in a peatland pasture in the USA.

We found a statistically significant relationship between dissolved N<sub>2</sub>O and NO<sub>3</sub><sup>-</sup>, in 523 agreement with others (e.g. Reay et al., 2003, Hinshaw & Dahlgren, 2013, Schade et al., 524 2016). However, a better fit was found between dissolved  $N_2O$  and the DOC:NO<sub>3</sub><sup>-</sup> ratio. 525 526 Aquatic systems generally show an inverse relationship between DOC and NO<sub>3</sub><sup>-</sup> concentrations, which reflects a gradient from nitrogen limitation of microbial processes in 527 carbon-rich systems to labile organic matter limitation in carbon-poor systems (Goodale et 528 al., 2005; Taylor and Townsend, 2010). Our observation that dissolved  $N_2O$  only increases 529 above ambient atmospheric concentrations when DOC/NO<sub>3</sub><sup>-</sup> ratios are low suggests both 530 that NO<sub>3</sub><sup>-</sup> concentrations need to be high enough to allow denitrification to occur, and that 531 labile organic matter concentrations need to be low enough to favour this process over 532 533 other microbial processes such as NO<sub>3</sub><sup>-</sup> reduction or assimilation. There were three samples 534 that appeared to deviate from the observed relationship, and it may be that higher concentrations of ammonium cause elevated N<sub>2</sub>O concentrations, particularly if dissolved 535 oxygen is not limiting (Liikanen & Martikainen, 2003). 536

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#### *4.4. Implications for GHG accounting and conclusions*

539 Our data support previous studies in showing that ditches in both semi-natural and agricultural peatlands act as sources of CH<sub>4</sub>, CO<sub>2</sub> and N<sub>2</sub>O emissions. It is widely recognised 540 that intact fens are important emitters of CH<sub>4</sub> (Turetsky et al., 2014). Although not intact, 541 542 our semi-natural fen is under conservation management, and therefore the vegetation is 543 similar to intact fens. As such, the annual terrestrial flux from our semi-natural site is 11.7 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> (Evans *et al.*, 2016b) , compared to 37.8 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> from the ditches. When 544 weighted by area for the entire fen, ditches would therefore be responsible for 0.53 g CH<sub>4</sub> 545 m<sup>-2</sup> yr<sup>-1</sup>, approximately 5% of total emissions. Although ditches occupy only a fraction of the 546 547 landscape, the magnitude of the fluxes observed here suggest that ditches in modified fen 548 landscapes must be considered when calculating carbon balances, particularly for studies relying on static chamber, rather than eddy-covariance, methods, since this component of 549 CO<sub>2</sub> and CH<sub>4</sub> emissions will otherwise be missed. In drained peatland systems, the 550 551 contribution of ditches to the overall CH<sub>4</sub> budget is even more marked, because CH<sub>4</sub> fluxes from drained peat surfaces tend to be near zero (Willison, 1998, IPCC, 2014). At our 552 cropland site, the field surface acted consistently as a small net sink for CH<sub>4</sub> (Evans et al., 553 554 2016b) and ditches were thus responsible for the entirety of CH<sub>4</sub> emissions from the system as a whole, which would give an areally-weighted flux of 0.44 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup> compared to the 555 field sink of -0.17 g CH<sub>4</sub> m<sup>-2</sup> yr<sup>-1</sup>. This is probably true for agriculturally drained peatlands in 556 general (IPCC, 2014, Evans et al., 2016a). Terrestrial fluxes at our grassland restoration site 557 558 show that both uptake and emission of CH<sub>4</sub> occur, but the annual flux is approximately zero (Evans et al., 2016a). Thus, the ditches here are responsible for the majority of CH<sub>4</sub> 559 emissions to the atmosphere, calculated on an areal basis as  $0.31 \text{ g CH}_4 \text{ m}^{-2} \text{ yr}^{-1}$ . 560

Our  $CO_2$  fluxes were similar to others reported in the literature from ditches, which 561 have often been of the same magnitude as fluxes from terrestrial fen (e.g. Schrier-Uijl et al., 562 2011, Hyvönen et al., 2013). Our median estimated N<sub>2</sub>O fluxes for the semi-natural and 563 grassland sites are slightly higher than terrestrial fluxes from a Finnish drained 564 565 minerotrophic fen (Martikainen et al., 1995) whilst our calculated N<sub>2</sub>O fluxes were in the

- same range as mean terrestrial fluxes from a German agricultural fen (Flessa *et al.*, 1998).
  These observations support previous suggestions that ditches do not act as hotspots for CO<sub>2</sub>
  and N<sub>2</sub>O in the same way that they do for CH<sub>4</sub> (Evans *et al.*, 2016a, Teh *et al.*, 2011).
- 569 Future work should continue to examine the role that ditches play in releasing GHGs 570 to the atmosphere, but a particular focus should be on CH<sub>4</sub>. It is likely that high-frequency 571 measurements combined with sampling replication on both small (i.e. field) and large (i.e. 572 regional) scales would elucidate in greater detail the drivers between both dissolved GHGs 573 concentrations and their efflux to the atmosphere. It is apparent that neglecting to consider 574 ditches in drained peatlands will lead to significant errors when calculating landscape-scale 575 GHG budgets.
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Table 1. Mean environmental, physical and water chemistry measurements for ditches at the three sites on the four sampling occasions. Numbers in brackets are standard error of the

mean. Depth is ditch water depth.

		March	May	Aug	Oct		March	May	Aug	Oct
Air temp (°C)	Semi-natural	8	12.5	12.7	12.2	Water temp (°C)	7.3 (0.2)	15.5 (0.2)	18.6 (0.2)	10.2 (0.1)
	Grassland	8	12.5	12.7	12.2		10.7 (0.3)	15.2 (0.3)	17.6 (0.4)	10.1 (0.2)
	Cropland	6.9	13.7	15.9	9.1		9.9 (0.6)	12.6 (0.2)	15.5 (0.4)	9.8 (0.2)
Depth (cm)	Semi-natural	85.6 (4.8)	86.9 (3.5)	64.0 (6.7)	70.1 (5.5)	рН	7.7 (0.02)	7.9 (0.05)	7.6 (0.04)	7.6 (0.02)
	Grassland	23.3 (7.3)	36.4 (3.9)	16.2 (1.4)	21.1 (2.8)		8.0 (0.06)	7.8 (0.09)	7.3 (0.06)	7.6 (0.03)
	Cropland	18.4 (3.1)	32.7 (4.6)	60.6 (3.5)	20.5 (3.9)		7.5 (0.04)	7.8 (0.05)	7.2 (0.07)	7.2 (0.04)
EC (µS cm <sup>-1</sup> )	Semi-natural	921 (62)	907 (18)	810 (73)	965 (51)	NO <sub>3</sub> - (mg l <sup>-1</sup> )	11.6 (4.2)	6.7 (1.8)	5.2 (0.8)	6.5 (1.2)
	Grassland	994 (32)	1117 (102)	1306 (80)	1584 (99)		19.1 (5.8)	4.1 (0.1)	3.9 (0.1)	5.0 (0.2)
	Cropland	1263 (87)	968 (69)	888 (28)	1134 (65)		9.9 (1.4)	18.2 (4.6)	4.8 (0.7)	5.3 (0.4)
DOC (mg l <sup>-1</sup> )	Semi-natural	28.7 (0.9)	28.9 (2.1)	27.4 (0.8)	37.4 (2.2)	SUVA	2.8 (0.1)	2.8 (0.1)	3.3 (0.1)	2.8 (0.1)
	Grassland	19.7 (2.7)	28.8 (1.1)	30.3 (1.6)	37.6 (2.4)		2.3 (0.1)	2.2 (0.0)	2.4 (0.1)	2.2 (0.2)
	Cropland	27.4 (3.3)	15.2 (4.8)	18.0 (2.8)	25.5 (3.7)		2.1 (0.1)	2.1 (0.1)	2.8 (0.1)	2.3 (0.1)
DIC (mg l⁻¹)	Semi-natural	92 (2)	80 (3)	70 (2)	97 (2)					
	Grassland	72 (2)	59 (6)	79 (4)	102 (3)					
	Cropland	69 (2)	52 (6)	68 (4)	77 (5)					

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able 2. Median p	oiston velocities (	m s <sup>-+</sup> ) for each sit	e and each mont	h, with first and l	third quartiles
		March	May	Aug	Oct
Semi-natural	Median	5.87E-06	1.16E-04	2.65E-04	8.58E-06
	1st	2.97E-06	1.33E-05	8.72E-05	3.96E-06
	3rd	1.33E-05	2.42E-04	4.17E-04	1.23E-04
Grassland	Median	4.89E-06	3.80E-05	9.18E-06	8.89E-06
	1st	2.23E-06	3.75E-06	7.99E-06	2.21E-06
	3rd	9.76E-06	7.87E-04	6.75E-05	2.33E-05
Cropland	Median	2.25E-06	2.26E-06	4.31E-06	8.01E-06
	1st	1.62E-06	1.67E-06	2.88E-06	4.39E-06
	3rd	3.40E-06	3.18E-06	7.20E-05	3.61E-05

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- 918 Table 3. Results of the multiple linear regressions to determine the relationships between dissolved CO<sub>2</sub>/CH<sub>4</sub> and other measured variables for the intact, restoration, and agricultural site.
- 919 Note that depth to WT refers to the water table depth in the terrestrial part of the fen.

Diss CO <sub>2</sub>	Slope coefficient	р
Intercept	-15.1	<0.001
Depth to WT	0.101	<0.001
DIC	0.133	<0.001
SUVA	2.594	0.001
Ditch depth	0.032	0.02
Diss CH <sub>4</sub>	Slope coefficient	р
Intercept	-0.63	<0.001
Air temp	0.0148	<0.001
NO <sub>3</sub> -	0.0066	<0.001
DIC	0.0066	<0.001
Depth to WT	0.0019	<0.001
EC	-0.0002	0.02