1 Methane and nitrous oxide exchange over a managed hay

2 meadow

3

4 L. Hörtnagl^{1,*} and G. Wohlfahrt¹

- 5 [1]{Institute of Ecology, University of Innsbruck, Austria}
- 6 [*]{Department of Environmental Systems Science, Institute of Agricultural Sciences IAS,
- 7 ETH Zurich, Switzerland}
- 8 Correspondence to: L. Hörtnagl (lukas.hoertnagl@usys.ethz.ch)
- 9

10 Abstract

11 The methane (CH_4) and nitrous oxide (N_2O) exchange of a temperate mountain grassland near 12 Neustift, Austria, was measured during 2010 - 2012 over a time period of 22 months using 13 the eddy covariance method. Exchange rates of both compounds at the site were low, with 97 % of all half-hourly CH₄ and N₂O fluxes ranging between ± 200 and ± 50 ng m⁻² s⁻¹, 14 15 respectively. The meadow acted as a sink for both compounds during certain time periods, but was a clear source of CH₄ and N₂O on an annual time scale. Therefore, both gases contributed 16 17 to an increase of the global warming potential (GWP), effectively reducing the sink strength 18 in terms of CO₂-equivalents of the investigated grassland site. In 2011, our best guess estimate showed a net GHG sink of -32 g CO₂-equ. m⁻² yr⁻¹ for the meadow, whereby 55 % of 19 the CO₂ sink strength of -71 g CO₂ m⁻² yr⁻¹ was offset by CH₄ / N₂O emissions of 7 / 32 g 20 CO_2 -equ. m⁻² yr⁻¹. When all data were pooled, the ancillary parameters explained 27 / 42 % of 21 22 observed CH₄ / N₂O flux variability, and up to 62 / 76 % on shorter time scales in-between 23 management dates. In case of N₂O fluxes, we found highest emissions at intermediate soil water contents and at soil temperatures close to zero or above 14 °C. 24

In comparison to CO₂, H₂O and energy fluxes, the interpretation of CH₄ and N₂O exchange was challenging due to footprint heterogeneity regarding their sources and sinks, uncertainties regarding post-processing and quality control. Our results emphasize that CH₄ and N₂O fluxes over supposedly well-aerated and moderately fertilized soils cannot be neglected when evaluating the GHG impact of temperate managed grasslands.

1 **1 Introduction**

Methane (CH₄) and nitrous oxide (N₂O) are the most important anthropogenic greenhouse gases (GHG) after carbon dioxide (CO₂). Due to their long atmospheric lifetimes of approx. 9 and 131 years (Prather et al., 2012), respectively, both compounds are well-mixed in the atmosphere and can influence atmospheric chemistry directly and indirectly. The emission or deposition strength of terrestrial ecosystems is possibly influenced by climate change, which may trigger important feedbacks to the global climate system (Xu-Ri et al., 2012).

8 Methane has a major influence on climate and chemistry of the atmosphere (Crutzen and 9 Lelieveld, 2001; Khalil et al., 2007). CH₄ can react with hydroxyl radicals, resulting in a 10 reduction of the oxidizing capacity of the atmosphere and the production of ozone (O_3) in the troposphere. Methane can influence the lifetime or production of other atmospheric 11 12 constituents like stratospheric water vapor and CO₂ (Boucher et al., 2009; Collins et al., 2010; 13 Shindell et al., 2009). Its global warming potential over a 100 year lifespan (GWP) and on a 14 per molecule basis is 25 times that of CO₂ (Forster et al., 2007) or higher when the production of CO₂ from CH₄ oxidation is taken into account (Boucher et al., 2009). 15

16 The main portion of global CH₄ originates from single-celled archaea (methanogens) found in 17 anaerobic microsites in the soil, in water-saturated zones rich in carbon and in the digestive systems of ruminants (Baldocchi et al., 2012; Whalen, 2005). CH₄ is also emitted from 18 19 organic waste deposits, e.g. manure, and from thermogenic and pyrogenic sources (Kirschke 20 et al., 2013). Highest emissions were previously reported from regions with intensive 21 agriculture and animal husbandry (Schulze et al., 2009). Atmospheric CH₄ increased 22 significantly since the industrial revolution until the end of the 1990s, remained constant for 23 nearly a decade and again began to increase after 2007 (Bousquet et al., 2011; Dlugokencky et 24 al., 2009; Nisbet et al., 2014).

25 The main sink of methane is through its reaction with the hydroxyl radical OH in the troposphere (Ehhalt and Heidt, 1973). Other, minor sinks are methanotrophic bacteria in 26 27 aerated soils and reactions with atmospheric constituents in the stratosphere and the marine 28 boundary layer (Allan et al., 2007; Cicerone and Oremland, 1988). Previous studies reported 29 reduced CH₄ deposition in a forest and in a temperate grassland due to elevated CO₂ (Dubbs 30 and Whalen, 2010; Ineson et al., 1998; Phillips et al., 2001) and increased CH₄ uptake due to 31 warming in a temperate forest and several subarctic ecosystems (Peterjohn et al., 1994; Sjogersten and Wookey, 2002). 32

Nitrous oxide can deplete O_3 in the upper and increase O_3 in the lower regions of the 1 2 stratosphere (Revell et al., 2012). It can therefore influence tropospheric chemistry by 3 increasing the stratosphere-troposphere exchange of O_3 and odd nitrogen species, and by increasing OH formation (Prather and Hsu, 2010). Similar to CH₄, N₂O has a high warming 4 5 potential, 298 times that of CO₂ over a 100 year lifespan (Forster et al., 2007). The dominant source of N₂O is microbial production through nitrification and denitrification processes in 6 7 soils, which is fueled by accelerated use of nitrogen fertilizers in agriculture (Davidson, 2009; 8 Fowler et al., 2009). As a consequence of fertilization agricultural soils are unlikely to act as a 9 sink for N₂O (Syakila and Kroeze, 2011).

The production of N_2O by bacteria in soils is controlled by a number of factors, for example soil water content, temperature and labile carbon availability (Barnard et al., 2005; Holtan-Hartwig et al., 2002; Xu-Ri and Prentice, 2008). Food production was described as the largest single source of N_2O (Syakila and Kroeze, 2011), while photolysis and oxidation reactions in the stratosphere are the main processes involved in N_2O depletion (Prather et al., 2012).

15 Denitrification is an anaerobic process (Zumft and Kroneck, 2007) that is likely exclusively 16 responsible for N₂O uptake in the soil (Vieten et al., 2008). On a global scale, the uptake of 17 N_2O by soils may be limited (Chapuis-Lardy et al., 2007). Schlesinger (2013) estimated that 18 the global N₂O sink in soils is not more than 2 % of current estimated sources in the 19 atmosphere. Deposition fluxes to the soil were reported before, e.g. for grasslands, forests, 20 low-nitrogen soils, wetlands and peatlands (Dijkstra et al., 2013; Flechard et al., 2005; 21 Goldberg and Gebauer, 2009a, 2009b; Schlesinger, 2013; Syakila et al., 2010; Wu et al., 22 2013).

Over managed grasslands, CH₄ and N₂O fluxes are characterized by high spatial and temporal variability (Baldocchi et al., 2012; Imer et al., 2013), with emissions of both compounds greatly influenced by land use, management events and animal husbandry. As a consequence, long-term year-round GHG measurements are indispensable when it comes to assessing the effectiveness and feasibility of GHG mitigation strategies.

In this work we present long-term eddy covariance CH_4 and N_2O fluxes above a temperate mountain grassland near Neustift, Austria. To this end we investigated 22 months of diurnal, seasonal and interannual exchange rates of both compounds at ecosystem scale and in relation to biotic and abiotic drivers under *in situ* conditions.

The objective of this study is to (1) quantify eddy covariance CH_4 and N_2O fluxes, (2) couple 1 2 exchange patterns to independent driving variables, (3) determine the annual total GHG 3 balance and (4) compare our findings to previous results from chamber and eddy covariance 4 measurements at ecosystem scale and from laboratory measurements. In line with these 5 objectives and based on earlier studies we hypothesized for both compounds that (1) the investigated grassland, due to generally well-aerated soils and modest fertilizer input, is 6 7 characterized by low fluxes and (2) exchange patterns are predominantly driven by soil 8 parameters. In addition we assumed that (3) despite their low fluxes, CH₄ and N₂O exchange 9 significantly contribute to the GHG balance of the meadow.

The study site Neustift, a managed temperate mountain grassland in Austria that is cut three times per year for hay production, was selected because it has been the focus of numerous studies over the last ten years and is therefore well described in terms of management effects, net ecosystem CO₂, H₂O, energy (Brilli et al., 2011; Hammerle et al., 2008; Wohlfahrt et al., 2008b) and VOC exchange (Bamberger et al., 2010, 2011; Brilli et al., 2012; Hörtnagl et al., 2011, 2014; Müller et al., 2010; Ruuskanen et al., 2011).

16

17 2 Methods

18 **2.1 Site description**

19 The study site is an intensively managed meadow in the middle of the flat valley bottom of the Stubai Valley in the Austrian Alps, in proximity of the village of Neustift (47°70' N, 20 11°19' E) at an elevation of 970 m a.s.l. The climate is humid continental with alpine 21 22 influences, with an average annual temperature of 6.5 °C, the average annual precipitation 23 amounts to 852 mm. The fetch is homogeneous up to 300m to the north-northeast (the 24 dominant daytime wind direction) and 900m to the south-south-west (nighttime) of the 25 instrument tower, parallel to the Valley's orientation. Typically, higher wind speeds and 26 unstable conditions result in a smaller footprint during daytime than during nighttime, where 27 the footprint of the site is larger due to the stable stratification of the atmosphere (Bamberger 28 et al., 2010). The vegetation of the meadow is dominated by a few graminoids (Dactylis glomerata, Festuca pratensis, Phleum pratensis, Trisetum flavescens) and forbs (Ranunculus 29 30 acris, Taraxacum officinale, Trifolium repens, Trifolium pratense, Carum carvi), while the 31 slopes of the surrounding mountains are covered mainly by coniferous forest. The soil was classified as a Fluvisol (FAO classification) and is approximately 1 m deep, with a thin
organic layer (0.001 m), followed by an A horizon that extends down to 0.02 m and a B
horizon, best described as a sandy loam. The organic volume fraction of the A horizon is
approximately 14 %.

5 Measurements of CH₄ and N₂O for this work were conducted from 13 April 2010 - 29 February 2012 (684 days). In each year, the meadow was cut three times, with the 1st cut on 5 6 / 6 June in 2010 / 2011, respectively, the 2nd cut on 31 July / 1 August and the 3rd cut on 20 / 7 26 September. In addition, the meadow was fertilized by manure spreading between 18 - 228 9 October in 2010 and on 18 and 19 October in 2011. The meadow was snow-covered from 1 January – 28 February 2010, from 26 November 2010 – 10 March 2011 and from 7 December 10 2011 – 24 March 2012, resulting in a total of 246 snow days for this analysis. During the 11 12 measurement campaign, no cows were present on the meadow.

13

14 **2.2 Eddy covariance measurements**

15 The net ecosystem exchange for CH₄ and N₂O was calculated by combining the 20 Hz threedimensional wind speeds quantified by a sonic anemometer (R3IA, Gill Instruments, 16 Lymington, UK) at a height of 2.5 m above ground with the simultaneously detected volume 17 mixing ratios (VMRs) of CH₄ and N₂O, which were both measured by a commercially 18 available continuous-wave quantum cascade laser (QCL; CWQC-TILDAS-76-D, Aerodyne, 19 20 USA). Fluxes were then calculated using the virtual disjunct eddy covariance method (vDEC) 21 method proposed by Karl et al. (2002), which is based on the eddy covariance (EC) method 22 (Baldocchi et al., 1988; McMillen, 1988). The intake tube for the QCL was mounted at 0.2 m 23 below the sonic anemometer and displaced laterally perpendicular to the predominating wind direction in order to minimize flux loss due to vertical and longitudinal sensor separation 24 25 (Massman, 2000). Sample air was drawn from the inlet through a filter (1-2 µm, PTFE) and heated (35 °C) PFA Teflon tubing (1/4" inner diameter) of 12 m length to the QCL at a flow 26 27 rate of around 8 SLPM (standard liter per minute; air volume normalized to standard 28 temperature and pressure conditions: 273K, 1013 hPa). Sonic anemometer data were stored to 29 the hard drive of a personal computer (PC) using the Eddymeas software (O. Kolle, Max Planck Institute for Biogeochemistry, Jena, Germany). More details regarding the CO₂, H₂O 30

and energy flux measurements are given in Wohlfahrt et al. (2008) and Hammerle et al.
 (2008).

3

4 2.3 QCL setup

5 Ambient air was analyzed for CH₄, N₂O and H₂O at time resolutions of 10 Hz (13 March - 16 6 August 2010), 5 Hz (16 – 24 August 2010) and 2 Hz (26 August 2010 – 29 February 2012). 7 The QCL and associated hardware (vacuum pump and thermo cube) were housed in a 8 climate-controlled instrument hut next to the field site. During the last five minutes of every 9 half-hour, CH₄- and N₂O -free air and air with known, close-to-ambient, VMRs were 10 switched into the sampling line to determine zero and span of the OCL, respectively. The 11 QCL was operated at a pressure of 4 kPa using a built-in pressure controller and temperature of the optical bench and housing controlled to 35°C. Fitting of absorption spectra, storing of 12 calculated VMRs, switching of zero/calibration valves, control of pressure lock and other 13 14 system controls were realized by the TDLWintel software (Aerodyne, USA) run on a PC synchronized with the main PC collecting anemometer data using the NTP software 15 (Meinberg, Germany). 16

17

18 **2.4 Despiking**

Similar to observations by Baldocchi et al. (2012) for methane, we experienced elevated 19 VMRs of both compounds, but especially CH₄, at night. We attributed these increased VMRs 20 21 to atmospheric phenomena in the calm and stable nocturnal boundary layer rather than to 22 elevated biogenic emissions. Therefore, VMRs of both compounds were subjected to a 23 rigorous outlier removal routine before entering flux calculations (Fig. 1a). The despiking method in this study is based on a median filter that runs through each half-hourly VMR time 24 25 series data point by data point. In comparison to the arithmetic mean, the median value of a time series is relatively insensitive to outlier values. For each 30 min period, (1) a smoothed 26 27 time series of the original VMR time series was created. This was done by replacing each 28 original data point with the median value of a moving time window of \pm 500 values around 29 the respective VMR value. In order to enable the calculation of median values also for data points at the start and end of the measured time series, the first and last 500 values were 30

copied and repeated at the start and end of the smoothed time series, respectively. (2) Each 1 2 data point in the smoothed time series was then subtracted from the respective measured data 3 point, generating a time series of differences between the two data matrices. (3) When the difference exceeded the empirically determined outlier threshold of 100 ppb, the data point in 4 5 the measured time series was marked as an outlier. This outlier threshold was tailored to the CH₄ variability, but worked also well for removing extreme values in the N₂O time series. (4) 6 7 The arithmetic mean without these outliers was then calculated and used to (5) replace 8 outliers in the respective half-hourly time series. As turbulent fluctuations for final flux 9 computations are calculated using block averaging, the contribution of these substituted data 10 points to resulting half-hourly fluxes is minor. To better account for natural variability in the 11 time series, three different runs with varying window sizes (\pm 500, 250, 150 values) and 12 outlier thresholds (100, 80, 60 ppb) were performed for each 30 min period.

During daytime / nighttime, at least one outlier was removed in 30 % / 66 % of half-hourly
CH₄, but only in 1 % / 1 % of all recorded N₂O VMRs.

15

16 **2.5 Flux calculations**

17 Half-hourly fluxes of CH₄ (F_{CH4}) and N₂O (F_{N2O}) were then calculated using the virtual 18 disjunct eddy covariance (vDEC) method (Karl et al., 2002) as the covariance between 19 turbulent fluctuations of the vertical wind speed and the VMRs derived from Reynolds averaging of 30 min blocks of data. The time lag between the high-resolution wind data and 20 21 the disjunct OCL time series was removed using a homemade program, resulting in a 22 subsample of the wind data corresponding to the sampling rate of the QCL. In the same step, 23 CH₄ and N₂O fluxes were corrected for the effect of air density fluctuations and laser bandbroadening following *Neftel et al.* (2010), using the OCL H₂O VMR. It was shown previously 24 25 that flux estimates using the vDEC method are characterized by a larger random uncertainty 26 compared to the true EC, but are unbiased (Hörtnagl et al., 2010). The tubing induced delay 27 time between the wind and the QCL concentration time series was determined in a procedure comprising multiple steps. First, the correlation coefficient between the H₂O time series 28 measured concurrently by the QCL and a closed-path infrared gas analyzer (Li-7000, LiCor, 29 USA), the data of which were acquired together with the sonic anemometer wind data, was 30 optimized to remove potential time differences between the two PCs caused by deviating 31

internal clocks, effectively adjusting the starting points of the two time series. Due to 1 2 generally low values of F_{CH4} and F_{N20} at our study site, the determination of lag times 3 between the CH₄ / N₂O time series and the wind data was difficult, but worked well between the QCL H₂O signal and the wind data. Therefore, secondly, the time delay between the wind 4 5 components and the QCL H₂O was determined by identifying the maximum/minimum of the cross-correlation function in a time window of +/- 7 s. The frequency distribution of this 6 7 search revealed a peak around 2 s. Thirdly, a second time window of +/- 2 s (daytime) and +/-8 5 s (nighttime) was then applied around this peak and used for the final lag search between 9 the CH₄ / N₂O signal and the vertical wind velocity.

10 Final fluxes were then calculated using the post-processing software *EdiRe* (University of 11 Edinburgh). Frequency response corrections were applied to raw fluxes of both compounds, 12 accounting for high-pass (block averaging, finite impulse response filter) and low-pass (lateral 13 sensor separation, dynamic frequency response, scalar and vector path averaging, frequency response mismatch and the attenuation of concentration fluctuations down the sampling tube) 14 filtering according to Massman (2000), using a site-specific cospectral reference model 15 (Wohlfahrt et al., 2005a). The high pass, non-recursive, finite impulse response (FIR) filter 16 17 was applied digitally to account for an overestimation of the flux contributions of lowfrequency eddies. Best results were achieved by applying the FIR filter using a Hamming 18 19 window, whereby time constants of respectively 50 and 100s for CH₄ and N₂O sufficiently 20 filtered out unwanted flux contributions at frequencies < 0.05 Hz (Fig. 1b). Missing low-21 frequencies were then back-corrected based on the site-specific reference model co-spectrum 22 (Wohlfahrt et al., 2005b). Exchange rates of CH₄ and N₂O calculated with these settings 23 represent our final best guess fluxes that were used for all analyses in this manuscript.

24 Two days in April 2011 are used to exemplify the effect of different FIR filters, applied to the CH₄ and N₂O time series, on the resulting flux estimates (Fig. 1b). The largest difference 25 26 between unfiltered and filtered data as well as between the different filter time constants was 27 found during nighttime. In contrast, during turbulent conditions e.g. around noon, fluxes calculated with different time constants exhibited exchange patterns of comparable magnitude 28 (Fig. 1b, left panels). FIR filtering had a larger effect on CH₄ than on N₂O fluxes. As an 29 example, over the course of one day unfiltered CH₄ exchange rates fluctuated between -217 30 and 780 ng m⁻² s⁻¹ (average: 4 ± 260 ng m⁻² s⁻¹), while best guess fluxes ranged between -96 31 and 87 ng m⁻² s⁻¹ after FIR filtering (-7 \pm 51). Similarly, unfiltered N₂O fluxes were between -32

1 38 - 146 ng m⁻² s⁻¹ (11 ±46), with best guess fluxes of -33 - 18 ng m⁻² s⁻¹ (-5 ±15). Cospectral 2 analyses revealed that lower frequencies of the CH₄ and N₂O fluxes were overrepresented 3 compared to the sensible heat flux (Fig. 1b, right panels).

4 In total, 28891 raw flux values were calculated for CH₄ / N₂O, which corresponds to a data 5 coverage of 88 % over the whole measurement period between 13 March 2010 and 29 6 February 2012. Flux results of each FIR run required separate quality control. When applying 7 a FIR filter with a time constant of 50 s / 100 s / 150 s to the data, 57 / 55 / 55 % of all raw 8 CH₄ fluxes and 66 / 64 / 63 % of all raw N₂O fluxes passed all quality tests, respectively. However, only 28 % and 39 % of all raw CH₄ and N₂O fluxes, respectively, passed all tests 9 10 when no FIR filter was used in the flux calculations. Only data that passed all quality tests in 11 a respective scenario were used in the present study. All fluxes in this manuscript are 12 expressed as molecular mass per unit time and ground surface area.

13 In order to calculate the annual balance of CH₄ and N₂O in 2011, the respective quality-14 controlled half-hourly flux dataset was gap-filled. Gaps less than or equal to two hours were 15 filled by linear interpolation. For the filling of larger gaps a lookup table was generated, using flux data in a time window of 14 days around the missing flux value and T_{soil} bin widths of 16 17 1°C. If no lookup table could be generated, e.g. no flux data were available within the time 18 window, the mean diurnal variation (±14 days) was used to fill the gap. For the calculation of 19 the annual GWP of the meadow in Neustift, CH₄ and N₂O fluxes were converted to CO₂-20 equivalents using the respective compound warming potential as given by Forster et al. 21 (2007).

Instrumentation, data treatment and quality control of CO₂, sensible and latent heat fluxes
have been described at length by Wohlfahrt et al. (2008) and Hammerle et al. (2008).

24

25 **2.6 Quality control**

Half-hourly methane and nitrous oxide fluxes were excluded from the analysis if (i) the deviation of the integral similarity characteristics was larger than 60 % (Foken and Wichura, 1996), (ii) the maximum of the footprint function (Hsieh et al., 2000) was outside the boundaries of the meadow, (iii) fluxes were outside a specific range (F_{CH4} : +/- 800 ng m⁻² s⁻¹, F_{N20} : +/-220), (iv) half-hourly VMRs were outside a specific range (CH₄: 1800 – 3500 ppb, N_2O : 280 – 450 ppb), (v) the stationarity test for the respective flux exceeded 60 % (Foken and Wichura, 1996), (vi) the third rotation angle exceeded 10° (McMillen, 1988), (vii) the
number of half-hourly VMR values was below 3000 or (viii) more than 20% of data were
classified as spikes in any half-hourly period.

4

5 2.7 Ancillary data

6 Major environmental parameters were measured continuously at the field site, including air 7 temperature (T_{air}) , soil temperature (T_{soil}) at 0.05 m depth (TCAV thermocouple, Campbell 8 Scientific, Logan, UT, USA), volumetric soil water content (SWC) (ML2x, Delta-T Devices, 9 Cambridge, UK), soil heat flux (SHF) quantified by means of heat flux plates (3 replicates at 0.05 m depth, corrected for the change in heat storage above that depth; HFP01, Hukseflux, 10 11 Delft, Netherlands), total photosynthetically active radiation (PAR) (BF3H, Delta-T, 12 Cambridge, UK) and precipitation (52202, R. M. Young, Traverse City, MI, USA). All data were collected continuously by a data logger (CR10X, Campbell Scientific, Logan, UT, 13 14 USA). The green plant area index (GAI) was assessed (i) in a destructive fashion by 15 harvesting the plant matter of square plots (0.09 m², 3-5 replicates) and subsequent plant area determination (Li-3100, LiCor, Lincoln, NE, USA) and (ii) from measurements of canopy 16 17 height which was related to destructively measured GAI (Wohlfahrt et al., 2008b). Continuous time series of the GAI were derived by fitting appropriate empirical functions to 18 19 measured data separately for each growing phase before and after cutting events. A more 20 detailed list of all auxiliary parameters measured at this site is given by Wohlfahrt et al. 21 (2008b) and Hammerle et al. (2008).

22

23 **2.8 Statistical Analyses**

Statistical analyses were done using *Statistica 9* (StatSoft, Inc.), *SigmaPlot 12.5* (Systat Software, Inc.) and *Excel 2010* (Microsoft, Inc.). The natural logarithm (ln) of the observed CH₄ and N₂O fluxes was calculated and used in the simple (SLR) and multiple linear regression (MLR) analyses as the dependent variable. The *partial correlation* in the MLR analysis gives the correlation between two variables after controlling for the effect of all other variables in the equation. To determine significant differences between group means in a repeated measures analysis of variance (ANOVA) setting, the Unequal N HSD *post hoc* test, a

modification of the Tukey's HSD test, was used. For statistical analyses, only days or half-1 2 hours where all parameters were available were included. In case of ancillary data, the daily 3 average of the respective parameter was calculated when at least 40 half-hours of data were present for the respective day. In comparison, fewer values were available for CH₄ / N₂O 4 5 fluxes and VMRs due to the strict quality criteria. For CH₄ / N₂O data, the daily average was regarded as representative for the day when at least 14 half-hours were available after quality 6 7 control. In total 91 and 95 % of the presented CH₄ and N₂O daily average values, 8 respectively, were calculated from at least 20 half-hourly values.

9

10 **3 Results**

11 Daily average values of F_{CH4} / F_{N20} were calculated for 567 / 574 out of 684 days, respectively (Fig. 2). While fluxes of both compounds fluctuated around zero towards the end 12 13 of the vegetation period and during snow cover, net emission and deposition on a daily basis occurred for both compounds during certain time periods. Daily net uptake (negative sign) 14 15 was recorded on 162 / 203 days, whereby time periods characterized by clear deposition were found especially for N₂O, for example some weeks after snowmelt in spring 2011 (Fig. 2). 16 Highest daily average emissions for both compounds were found around the 2nd cutting of the 17 meadow at the end of July 2010 (123.5 / 33.4 ng m⁻² s⁻¹). CH₄ VMRs were highest during 18 19 snow cover and lowest during periods of strong growth (Fig. 2). We attribute the sudden drop of N₂O concentration values around the 1st cut in 2010 to a problem with the zero-calibration 20 21 of the QCL. Over all two years, the median VMR was 2.02 / 0.32 ppm for CH₄ / N₂O, respectively, the median flux amounted to 9.6 / 0.9 ng m⁻² s⁻¹ (Fig. 2). 22

Daily average PAR was found between approx. 40 µmol m⁻² s⁻¹ in winter and 674 µmol m⁻² s⁻¹ 23 ¹ in summer, with a median value of 215 μ mol m⁻² s⁻¹. In 2010, the yearly average T_{air} at the 24 25 field site of 6.1 °C was colder than the long-term average (2001 – 2007) of 6.7 °C, while 2011 26 was warmer than average (7.1 °C). During this study, the maximum daily average T_{air} was 27 22.7 °C in July 2010, the minimum of -17.3 °C was recorded in February 2012 (Fig. 2). T_{soil} 28 was similar in both years, about 8.5 °C on average and values just above 0 °C when snow 29 covered the ground. SWC was highest immediately after snow melt, with a maximum daily average value of 0.44 m³ m⁻³ at the end of February 2010, and lowest in May 2011 after a 30 period of only little precipitation (0.08 m³ m⁻³). In 2011, SWC was generally low (0.25 m³ m⁻³) 31 averaged over the growing season) and significantly lower (p < 0.001) than in 2010 (0.32 m³) 32

m⁻³). Over the duration of the flux measurements, precipitation was detected on 262 days, 1 amounting to 525 and 537 mm in 2010 and 2011, respectively, and 46 mm over the first two 2 3 months in 2012 (Fig. 2). Relative air humidity (RHA) was around 80 % on average over the whole measurement campaign, with minima below 50 % in June 2010 (Fig. 2). In 2010 and 4 5 2011, highest VPD values of more than 1 kPa were recorded during the warmer months between the end of May and August. GAI was below 1 m² m⁻² right after snow melt, reached 6 maximum values of up to 8 m² m⁻² right before the 1^{st} cut and was then reduced to below 1.5 7 m² m⁻² as a consequence of the cutting. GAI maxima before the 2nd and 3rd cut were lower 8 compared to the 1st cut. Towards the end of the year after the 3rd cut, GAI first increased and 9 later decreased due to vegetation regrowth and senescence, respectively (Fig. 2). 10

The meadow was a source for CO_2 during snow cover and became a net sink for CO_2 some weeks after snowmelt and until the 1st cut (Fig. 3). The cutting event turned the meadow into a CO_2 source for about two weeks before it again became a net sink. This behavior recurred after the 2nd and 3rd cut, however the CO_2 uptake after the last cutting was less pronounced than after the previous cuttings. More information about CO_2 fluxes at the site was given by Wohlfahrt et al. (2008).

17 Fluxes of both CH₄ and N₂O showed high variability on a half-hourly time scale, especially 18 during the first two months of the measurements and during the night (Fig. 3). However, 97 % 19 of all half-hourly CH₄ and N₂O fluxes during the vegetation period were found between ±200 and ± 50 ng m⁻² s⁻¹, respectively. During snow-free conditions and including only days not 20 influenced by management events, the average CH₄ / N₂O flux was found at 14.0 \pm 80.7 / 2.6 21 22 ± 21.6 ng m⁻² s⁻¹, respectively (Fig. 3). Compared to these undisturbed conditions, average fluxes were higher on days where the meadow was influenced by cutting events (17.5 \pm 83.7 / 23 4.8 ± 20.7 ng m⁻² s⁻¹) and lower on days characterized by snow cover (2.1 $\pm 82.8 / 0.9 \pm 20.7$). 24 The day of manure spreading and the two days thereafter were covered by our measurements 25 26 only in October 2011. On the day of fertilization and two days later, average N₂O fluxes were elevated (3.5 \pm 17.2 ng m⁻² s⁻¹) when compared to the rest of the same month (1.8 \pm 13.6), 27 while CH₄ fluxes remained virtually unaffected (24.7 ±91.0 vs. 27.0 ±88.9). In total, emission 28 fluxes were observed in 56 / 57 % of all recorded CH_4 / N_2O half hour periods (Fig. 3). 29

30 Average diurnal cycles of F_{CH4} and F_{N20} were often characterized by high variability with 31 large fluctuations around zero, but followed a clear diurnal cycle during certain time periods 32 (Fig. 4). Methane fluxes showed weak diurnal cycles after snowmelt and before the 2nd cut in

2011, with peak average uptake rates of -31.0 ± 41.4 ng m⁻² s⁻¹ around noon. The uptake of 1 2 CH₄ before the 1st cut coincided with strong N₂O deposition during daytime, with average peak rates of up to -12.3 ± 23.8 ng m⁻² s⁻¹ in the early afternoon. While CH₄ fluxes continued 3 to exhibit a very similar deposition pattern up until the 2nd cut, N₂O fluxes switched in sign 4 and showed a clear diurnal cycle of constant emission during daytime, up to 15.4 ± 18.9 ng m⁻² 5 s⁻¹ on average just before noon. The N₂O flux pattern after the 1st and before the 2nd cut was 6 7 very similar in both years, whereby peak emission rates in 2010 occurred earlier in the day (Fig. 4). In contrast to CH₄ fluxes, which showed no clear diurnal pattern after the 2nd cut in 8 both years, the meadow constantly emitted N₂O during daytime and before the 3rd cut in 2011, 9 on average up to 26.8 \pm 23.3 ng m⁻² s⁻¹ around noon, while during daytime after the 3rd cut in 10 2010 N₂O was transported to the meadow, peak deposition amounted to -7.5 ± 8.8 ng m⁻² s⁻¹ 11 on average. During snow cover, fluxes of both compounds fluctuated around zero (Fig. 4). 12

13 When all data were pooled, a MLR analysis explained 27 / 42 % of the variability in daily 14 average $\ln(F_{CH4}) / \ln(F_{N2O})$ during snow-free conditions (Table 1). Over all years, the partial 15 correlation (PC) of the net ecosystem exchange of CO₂ (NEE) and T_{air} with ln(F_{CH4}) was high and positive in sign, while SHF was negatively correlated with ln(CH₄); all three PCs were 16 17 highly significant (p < 0.001). During shorter time periods in-between, before and after 18 cutting events in single years the chosen set of parameters explained between 23 and 62 % of the observed flux variability, with r^2 being highly significant only once, namely in a period of 19 high CH₄ uptake before the 1st cut 2011, with NEE and H as the dominant regressors (Table 20 21 1). Explaining the $ln(F_{CH4})$ variance during the same time periods but using data of both years worked best during the vegetation period until the 2nd cut, and again after the 3rd cut until 22 23 snow cover, explaining up to 40 % of observed ln transformed CH₄ fluxes. The PC of SHF 24 and NEE were significant during the early vegetation period and towards the end of the year, 25 respectively. LE was a significant regressor towards the end of the vegetation period and 26 during snow cover (Table 1). We expanded on these findings by performing a forward step-27 wise MLR analysis using the same data, effectively reducing the number of variables in the regression equation but yielding similar results. In this analysis NEE, SHF, Tair and VPD were 28 29 identified as the most significant regressors (all p < 0.05), explaining 25 % of the observed 30 $ln(F_{CH4})$ variability over all years excluding snow periods (data not shown). The SLR analysis 31 found highly significant positive correlations for NEE and RHA, and highly significant 32 negative correlations for LE, H, and PAR (Table 1).

Generally, the MLR analysis resulted in r^2 being considerably higher for $ln(F_{N2O})$ than for 1 2 ln(F_{CH4}) (Table 1). The partial correlations were highly significant for multiple regressors. A 3 positive PC was found for the ecosystem fluxes NEE and LE, and in addition for RHA, Tair and N₂O VMR, while significant negative PCs were found for SWC, H and T_{soil}. All 4 5 regressors combined were able to explain between 55 and 76 % of the $ln(F_{N2O})$ variance during shorter time periods in single years, with the exception of the time period before the 1st 6 cut 2010 when r² was found to be statistically not significant (Table 1). The chosen set of 7 8 parameters performed well with pooled data during the same time periods and especially after 9 the 1st cut, explaining between 66 and 73 % of observed daily average $ln(F_{N2O})$ values. SWC 10 was the most dominant regressor towards the end of the year, featuring a highly significant, 11 negative PC (Table 1). Similarly, T_{soil} was an important parameter in the MLR analysis after the 1st cut, being first positively, later negatively correlated with ln transformed N₂O 12 exchange. Seven parameters were highly significant (p < 0.001) in a forward step-wise MLR 13 14 analysis and explained 41 % of the ln(F_{N2O}) variance during snow-free conditions, with T_{air}, N₂O VMR, RH, NEE and LE being positively correlated, SWC and H negatively (data not 15 16 shown). In a simple linear regression eight out of 11 parameters were significantly correlated 17 with the $ln(F_{N2O})$, with T_{air} and T_{soil} as the highest positively and SWC as the highest negatively correlated regressors, respectively (Table 1). 18

19 A closer look at the two most prominent soil related regressors, T_{soil} and SWC, and $ln(F_{N2O})$ under snow-free, undisturbed conditions revealed a clear pattern. Daily average N₂O 20 21 exchange showed a bell-shaped relationship with SWC with highest emissions during periods 22 of intermediate soil water content (Fig. 5, top panel). Even clearer was the correlation between T_{soil} and N₂O flux: days with a daily average T_{soil} above 14 °C showed an almost 23 consistent net emission of N₂O. This was also observed for days where T_{soil} was close to zero, 24 25 whereas N₂O exchange fluctuated around zero with no clear pattern between 0 and 14 °C (Fig. 5, middle panel). Taking both SWC and T_{soil} into account, days characterized by low to 26 27 intermediate SWC with T_{soil} close to 0 °C or above 14 °C generally resulted in a net emission 28 of N₂O, while deposition was mainly observed during cool conditions with high SWC (Fig. 5, 29 lower panel). In contrast to N₂O, comparably clear exchange patterns were not found for CH₄ 30 fluxes.

31 On a daily average time scale, a repeated-measures ANOVA revealed statistically significant 32 differences among environmental conditions on days with net uptake (group f-), net emission

(f+) or close-to-zero exchange (f0) of CH₄ and N₂O (Table 2). In case of CH₄, T_{air} was 1 significantly colder on low-flux days than on emission and deposition days. Generally, 2 3 environmental conditions were most different between high deposition days and days resulting in emission or close-to-zero exchange of CH4 (Table 2). In group f-, the ecosystem 4 5 fluxes LE and H, SHF, PAR, VPD and RHA were all significantly higher compared to f+ and f0, while also the net uptake of CO₂ was larger. Although results were less clear for N₂O 6 7 fluxes, the meadow tended to act neither as a source or sink on days when air and soil 8 temperatures as well as LE were low (Table 2). In addition, SWC was significantly lower in 9 f+, while H was significantly higher on deposition days.

Cumulative fluxes for 2011 resulted in a net CO₂-uptake of -70.5 g CO₂ m⁻² (Fig. 6). CH₄ and 10 N₂O fluxes were converted to CO₂-equivalents, with cumulative fluxes being calculated for 11 each of the different FIR filter time constants. In 2011, the meadow acted as a source for both 12 compounds. When no FIR filter was applied, i.e. the overestimation of the low frequency 13 14 eddy flux contribution was not corrected for, cumulative methane fluxes amounted to an emission of 54.5 g CO₂-equ. m⁻². With FIR filters of varying time constants, cumulative 15 16 fluxes were considerably lower, in the range of 6.8 - 19.3 g CO₂-equ. m⁻², whereby the lower number was obtained using a FIR filter time constant of 50 s and constitutes our best guess 17 18 estimate. Results were very similar for N₂O, the cumulative fluxes of which resulted in a net emission of 97.9 g CO₂-equ. m⁻² without FIR filter, and 25.2 – 39.8 g CO₂-equ. m⁻² using 19 filters with different time constants. In case of N₂O, a time constant of 100 s was considered 20 to give the most representative flux results, yielding 32.0 g CO_2 -equ. m⁻² over the whole year 21 22 (Fig. 6).

The total GHG budget can be calculated by summing up the different cumulative contributions of CO₂, CH₄ and N₂O. Based on the best guess estimates, the meadow acted as a GHG sink (-31.7 g CO₂-equ. m⁻²) in 2011. However, when no FIR filter was applied to neither CH₄ nor N₂O data, the sum of the two compound fluxes more than compensated for the sink effect of CO₂, turning the meadow into a GHG source (81.9 g CO₂-equ. m⁻²; Fig. 6).

1 4 Discussion

2 **4.1 Methane**

3 It was shown recently that plants do not contain a known biochemical pathway to synthesize 4 methane (Nisbet et al., 2009), a finding that contradicts observations of methane emissions from terrestrial plants under aerobic conditions in an earlier study (Keppler et al., 2006). 5 6 Methane emissions from plant tissue may be due to the transpiration of water that contains 7 dissolved CH₄ or due to the abiotic breakdown of plant material as a consequence of high UV 8 stress conditions (Nisbet et al., 2009), but the contribution of terrestrial plants to the global methane emission is considered to be small (Dueck et al., 2007). Based on these earlier 9 10 findings it is feasible to regard observed eddy covariance emission fluxes in this study as a 11 direct (methanogen microorganisms) or indirect (transpiration of soil CH₄) consequence of 12 processes in the soil, an important player in the global methane cycle (Kirschke et al., 2013; 13 Smith et al., 2000).

14 Therefore, one might expect clear relationships between soil environmental parameters such 15 as temperature or moisture and CH₄ exchange, which were also reported by other studies (Dijkstra et al., 2013; Hartmann et al., 2010; Imer et al., 2013; Jackowicz-Korczyński et al., 16 17 2010; Kroon et al., 2010; Liebig et al., 2009; Rinne et al., 2007; Schrier-Uijl et al., 2010). However, when all data were pooled no clear correlation between soil parameters and eddy 18 covariance CH₄ exchange at the grassland site in Neustift was observed. Although the 19 20 explanatory power of T_{soil} in the MLR was relatively high and significant between the 1st and 2^{nd} cutting of the meadow in 2011 – a period when small quantities of CH₄ were taken up by 21 22 the meadow around noon – no consistent relationship between soil parameters and the CH₄ 23 flux was observed (Table 1). SHF was significantly higher on days with net deposition 24 compared to zero-flux and net emission days (Table 2), which might be an indication of soil 25 processes as possible drivers for observed exchange patterns. The partial correlations of SWC 26 with CH₄ exchange, however, were statistically not significant throughout the measurement 27 campaign and close to zero when all data were pooled (Table 1). This is in contrast to 28 chamber studies that identified soil moisture as a key driver for methane exchange (e.g. 29 Dijkstra et al., 2013b).

30 One explanation for this lack of correlation between soil parameters and methane fluxes 31 might be that half-hourly eddy covariance fluxes represent an integral signal, averaged over

30 minutes over a possibly heterogeneous area of methane sources and covering both "hot 1 2 spots" of high methane emission and areas of relatively high uptake within the same flux 3 footprint (Baldocchi et al., 2012). Therefore, SWC may be high in certain patches of the meadow and create environmental conditions conducive for methanogenic microorganisms, 4 5 but low in other microsites across the grassland. Half-hourly fluxes reflect this heterogeneity 6 across the footprint to a varying degree, mainly depending on wind direction, wind speed and 7 atmospheric stability. In addition, the direct effect of certain drivers on CH₄ exchange may 8 smear out at ecosystem scale, especially if associated fluxes are generally low. Recently 9 Yvon-Durocher et al. (2014) found an average temperature dependence of CH₄ emissions 10 from aquatic, wetland and rice-paddy ecosystems similar to that of CH₄ production derived 11 from pure cultures of methanogens and anaerobic microbial communities in the laboratory. 12 No such relationship was found in the present study, which may be a direct consequence of a 13 heterogeneous footprint with regards to CH₄ sources and generally low CH₄ fluxes at the 14 measurement site in Neustift.

15 The observation of weak CH₄ uptake around noon between March and July 2011 (Figure 2) is most likely a consequence of methanotrophic microorganisms in the soil, a process enhanced 16 17 by increased soil temperature. However, it is difficult to observe this temperature dependence at ecosystem scale, as the whole footprint regardless of emission / deposition hot spots is 18 19 sampled. In addition, it was shown that both methanotrophic and methanogenic activity in the 20 soil are temperature dependent (von Fischer and Hedin, 2007; Yavitt et al., 1995), whereby 21 the latter tends to be more responsive to temperature (Topp and Pattey, 1997). Imer et al. 22 (2013) reported nearly consistent methane uptake throughout the year except for winter at 23 three different grassland sites along an altitudinal and management gradient using static chambers, with flux rates of generally below 10 ng m^{-2} s⁻¹. Three pastures investigated by 24 25 Liebig et al. (2009) were identified as minor CH₄ sinks.

Daily average CH₄ emissions in this study generally ranged ranged up to 100 ng m⁻² s⁻¹ and were relatively similar to eddy covariance results over a drained and grazed peatland pasture during dry periods, when fluxes were often below 160 ng m⁻² s⁻¹ (Fig. 2; Baldocchi et al., 2012). However, the maximum CH₄ flux and concentration of more than 5700 ng m⁻² s⁻¹ and 3500 ppb, respectively, at the peatland site were much higher than the 128 ng m⁻² s⁻¹ and 2300 ppb recorded at Neustift. Higher maximum methane fluxes were also observed by Schrier-Uijl et al. (2010) over a grass ecosystem on peat (1604 ng m⁻² s⁻¹).

In comparison to CO₂ and energy fluxes, there are only few long-term EC methane exchange 1 2 studies. However, year-round measurements are indispensable for accurately estimating the 3 CH₄ budget of an ecosystem. Baldocchi et al. (2012) give a three-year mean annual methane efflux at a peatland pasture of 11.6 \pm 9.0 g m⁻² vr⁻¹ without any discrimination for cattle or 4 elongated footprints during the night, and 3.6 ± 1.9 g m⁻² yr⁻¹ when only daytime data 5 representing the well-drained portion of the pasture, additionally filtered for favorable wind 6 7 directions and the presence of cows, were used. This latter number is relatively similar to the methane efflux of 2.1 g m⁻² yr⁻¹ in Neustift in 2011. In comparison, Hendriks et al. (2007) 8 reported $14.2 + -26.1 \text{ g m}^{-2} \text{ yr}^{-1}$ from the relatively dry portions of an abandoned peat meadow 9 using chamber measurements, and 42.5 \pm 27.7 g m⁻² vr⁻¹ when the whole meadow, including 10 11 water-saturated land and ditches, was considered. Mander et al. (2010) conducted a literature survey and reported median fluxes of 0.16 g m⁻² yr⁻¹ for fertilized grasslands on hydromorphic 12 soils in Estonia, similar to Neustift (0.27 g m^{-2} vr⁻¹). Methane emissions reported by Merbold 13 et al. (2014) from a grassland after restoration where one order of magnitude higher (3.6 g m^{-2} 14 yr⁻¹). Using eddy covariance measurements, methane emissions between 24 - 29 g m⁻² yr⁻¹ 15 were reported from a subarctic peatland (Jackowicz-Korczyński et al., 2010), 12.6 g m⁻² yr⁻¹ 16 from a boreal fen (Rinne et al., 2007) and 16.5 g m⁻² yr⁻¹ from a managed fen meadow (Kroon 17 18 et al., 2010).

19 Baldocchi et al. (2012) reported mean diurnal patterns characterized by lowest methane efflux densities during midday and elevated methane emission throughout the night, a pattern very 20 similar to Neustift during certain time periods, e.g. between the $1^{st} - 2^{nd}$ cut 2010 (Fig. 4). We 21 22 mainly attributed this observation to meteorological factors, i.e. intermittent exchange during calm and stable nighttime conditions, which was also the reasoning behind the outlier 23 24 handling in our despiking procedure (Fig. 1a). Another reason might be the preferential 25 sampling of an elevated methane source in combination with a larger nighttime footprint as 26 described by Baldocchi et al. (2012). It is possible that methane emissions from a small 27 stream and adjacent wet patches of the meadow, that are normally not part of the footprint, 28 have contributed disproportionally to observed methane emissions. Unfortunately we lack 29 detailed high-resolution spatial data (e.g. vegetation, soil) about small areas and patches 30 within the sampled flux footprint in Neustift, which would be required for a meaningful footprint analysis. Therefore, we are currently not able to further discuss potential emission 31 32 hotspots, their impact on calculated CH₄ balances and the problem of possibly preferential sampling within this manuscript. Hot spot footprint analysis merits its own research and
 would provide important insights in how to interpret eddy covariance flux data.

Several studies reported that 81 – 90 % of the total annual methane emission occurred during
the snow free period or between spring – autumn (Jackowicz-Korczyński et al., 2010; Rinne
et al., 2007), which is very similar to Neustift in 2011, where 84 % of the yearly net CH₄
emission occurred during snow free conditions.

7

8 4.2 Nitrous oxide

9 Despite occasional uptake, the meadow was a source of N₂O, in accordance with previous studies over managed grasslands. Half-hourly emission rates of N₂O, mostly below 50 ng 10 N₂O m⁻² s⁻¹, were similar to exchange rates reported by Neftel et al. (2010) for an 11 12 experimental farm site and Imer et al. (2013) from a mountain rangeland. N₂O fluxes in 2011 amounted to an emission of 107 mg m⁻² yr⁻¹. For comparison, Mander et al. (2010) reported 13 14 approx. 94 and 723 mg m⁻² yr⁻¹ for unfertilized and fertilized grasslands, respectively. 15 Considerably higher emissions were found by Kroon et al. (2010) for a managed fen meadow (2.4 g N₂O m⁻² yr⁻¹), and by Merbold et al. (2014) for a grassland after restoration (4.6 g m⁻² 16 17 vr⁻¹).

18 Many of the observations made for CH_4 were also valid for N_2O , with generally low fluxes, a 19 possibly heterogeneous flux footprint with respect to emission / deposition hot spots and soil 20 processes as the driving force behind N₂O exchange patterns. In contrast to CH₄ exchange, 21 N₂O fluxes on a daily scale could be well explained by environmental parameters during 22 specific time periods. The important role of temperature in soil processes was shown 23 previously, as N mineralization, nitrification, denitrification and N₂O emissions all increase with temperature (Barnard et al., 2005), while reduced soil moisture as a result of high air 24 25 temperatures and increased plant transpiration can decrease N₂O emissions (Li et al., 1992). 26 These findings are comparable to observations in the present study, where N₂O exchange 27 tended to emission during warm and relatively dry soil conditions (Figure 5, lower panel).

28 N_2O consumption in the soil occurs when N_2O reduction exceeds N_2O production (Chapuis-

29 Lardy et al., 2007). Soil water is probably the key driver regulating N_2O consumption in soils,

30 as it can act as a temporary storage body that entraps N₂O, effectively hindering its diffusion

from the soil matrix to the surface. As a consequence, the time for potential reduction of N₂O 1 2 to N₂ through anaerobic denitrification is increased (Clough et al., 2005). This can result in a 3 low N_2O / N_2 ratio during wet conditions, which favors N_2O consumption (Ruser et al., 2006; Wu et al., 2013). These observations agree with our findings at ecosystem scale. When all 4 5 data were pooled, N₂O uptake was highest during relatively wet conditions (Figure 5, top panel) and SWC was significantly lower on days with clear net emission of N₂O (Table 2). 6 7 The latter finding is further highlighted by a clear positive correlation between daily average 8 ln(F_{N2O}) and T_{soil} in the soil temperature range 12-16 °C as long as SWC was low (data not 9 shown).

10 In October 2011, manure application resulted in a pulse of N₂O emission one day later, after 11 which fluxes rapidly decreased and reached pre-fertilization rates two days after manure 12 spreading. Similar behavior of N₂O fluxes returning to background levels within 2-6 days 13 after fertilization has been observed by Jones et al. (2011) for a Scottish grassland and Neftel et al. (2010) for an experimental farm site. Pulses of N₂O emissions after fertilizer application 14 15 were also described in other studies (e.g. Granli and Bockman, 1994; Jones et al., 2011) and 16 might be the result of animal manure – the most concentrated form of anthropogenic N input 17 (Davidson, 2009) – directly fueling nitrifying and denitrifying bacteria in the soil, which are most active when N is abundant (Firestone and Davidson, 1989). Over the weeks following 18 19 fertilization, N₂O emissions increased with air temperature, which is in-line with the 20 temperature dependence of the involved processes. We observed a sharp increase of N₂O 21 emissions once the daily average air temperature fell below the freezing point, approx. four 22 weeks after manure spreading in November 2011. During this time period the meadow 23 remained snow-free, with soil temperatures close to 0°C. The combination of reduced plant 24 metabolism (low nitrate demand by plants) and prior manure spreading could result in an 25 abundance of soil NO₃₋ at the end of the vegetation period. Wertz et al. (2013) showed that denitrification can still occur at very low temperatures and even below the freezing point 26 27 when NO₃₋ and C are present. The observation of high N₂O emissions from frozen or nearly frozen soil was also made by earlier studies (Röver et al., 1998; Teepe et al., 2001). 28

Production and subsequent emission of N_2O remained high after the beginning of the snow cover in December 2011. Zhu et al. (2005) described a similar situation where microbial activity in the soil of a lowland tundra did not cease during snow cover and N_2O continuously diffused to the atmosphere through the snowpack. In Neustift, high N₂O emissions were not
observed one year earlier during similar conditions.

3

4 **4.3 Global warming potential**

5 The availability of year-round data allows for the calculation of a yearly GWP balance over a 6 specific ecosystem. In this study, year-round CH₄, N₂O and CO₂ flux data were available for 7 2011. When expressing the net exchange of the three compounds in terms of CO₂-equivalents 8 and adding up these different contributions, the resulting GWP of the meadow in Neustift was 9 -32 g CO₂-equ. m⁻² yr⁻¹ in 2011, whereby a yearly NEE of -71 g CO₂ m⁻² yr⁻¹ was offset by 10 CH₄ and N₂O emissions of 7 and 32 g CO₂-equ. m⁻² yr⁻¹, an offset of approx. 55%.

Liebig et al. (2009) investigated three years of CH₄ / N₂O static chamber fluxes, soil organic 11 12 carbon change, CO₂ emissions associated with N fertilizer production and CH₄ emission from enteric fermentation for three grazing management systems. The resulting net GWP between -13 78 - 40 g CO₂-equ. m⁻² yr⁻¹ is similar to results in this study. Hendriks et al. (2007) reported -14 86 g CO₂-equ, m^{-2} vr⁻¹ from an abandoned peat meadow. Merbold et al. (2014) give the full 15 GHG flux budget of an intensively managed grassland after restoration, including ploughing. 16 GHG emissions reported in their study were much higher than in Neustift, amounting to 2.9 17 kg CO₂-equ. m⁻², and relatively similar to the balance of 1.6 kg CO₂-equ. m⁻² found by Kroon 18 et al. (2010) for a managed fen meadow. Zona et al. (2013) reported a GHG balance of -260 g 19 CO₂-equ. m⁻² yr⁻¹ for a poplar plantation in 2011, taking into account CO₂ fluxes of -351 g 20 CO₂-equ. m⁻² vr⁻¹, and CH₄ and N₂O fluxes of 49 and 42 g CO₂-equ. m⁻² vr⁻¹, respectively, 21 22 with CH₄ and N₂O offsetting the NEE sink by 26 %. Soussana et al. (2007) investigated the 23 GHG budget of nine European grassland sites over two years, covering a major climatic 24 gradient and a wide range of management regimes. On average, the investigated grassland plots were a net sink of -879 g CO_2 m⁻² yr⁻¹, and a net source of 117 and 51 g CO_2 -equ. m⁻² yr⁻¹ 25 ¹ for CH₄ and N₂O, respectively, with emissions of the latter two compounds resulting in a 19 26 27 % offset of the NEE sink activity. Tian et al. (2014) reported offset ratios of 73 % for the whole North American continent, with the grassland GWP being nearly neutral. 28

Rinne et al. (2007) reported a GWP balance of $+108 \text{ g CO}_2$ -equ. m⁻² when taking into account CO₂ and CH₄ fluxes from a boreal fen, with respective fluxes amounting to -156 and +264 g CO₂-equ. m⁻². Although the GWP calculated from CO₂ and CH₄ fluxes was much lower in Neustift (-64 g CO₂-equ. m⁻²), the situation was similar in that the carbon uptake of the meadow through CO₂ was partially offset by carbon loss through CH₄ emission. The number for Neustift may change drastically on a year-to-year basis, as the meadow can act both as a source and sink of CO₂ (Wohlfahrt et al., 2008a), while it is supposedly a constant source of CH₄. Dijkstra et al. (2013) used static chambers to calculate the GWP for five years of CO₂ and CH₄ data in a semiarid grassland, ranging between -3 and -6 g CO₂-equ. m⁻².

7

8 5 Conclusion

9 The grassland site in Neustift is characterized by low fluxes of CH_4 and N_2O . Although the 10 meadow can act as a source and sink for both compounds during certain time periods, it is a 11 clear source of CH_4 and N_2O on an annual time scale. As a consequence, both gases 12 contribute to an increase of the GWP, effectively reducing the sink strength in terms of CO_2 -13 equivalents.

14 Our analyses showed that daily average N₂O exchange during most of the vegetation period can be well explained with simultaneously recorded ancillary data, especially in the time 15 16 period after the 1st cut in June up until snow cover towards the end of the year. In contrast, 17 modeling daily average exchange with the same ancillary data worked considerably worse for 18 CH₄, a finding that suggests the possibility of a more heterogeneous footprint in regard to 19 methane sources and sinks. For both compounds it was not possible to single out one driving 20 variable as the most important one, which is to be expected due to the nature of the eddy 21 covariance flux signal in combination with generally low CH₄ and N₂O fluxes at the 22 investigated grassland site.

In comparison to CO₂, H₂O and energy fluxes, the interpretation of CH₄ and N₂O exchange is 23 24 challenging due to uncertainties regarding post-processing, quality control and footprint 25 heterogeneity. Knowledge about emission and deposition hotspots within the footprint area 26 would allow for a more comprehensive interpretation of the bulk EC flux. Additional 27 information about GHG producing and consuming patches within the flux footprint could be achieved for example via chamber measurements, another possibility would be to perform a 28 29 detailed statistical analysis of EC fluxes and underlying footprint information in combination with detailed spatial data of the sampled area. 30

We conclude that CH₄ and N₂O fluxes over supposedly well-aerated and moderately fertilized 1 2 soils cannot be neglected when evaluating the GHG impact of temperate managed grasslands. 3 Both compounds can significantly influence the GWP balance of a meadow and be determining if a grassland is acting as a source or sink of CO₂-equivalents. In order to reliably 4 5 assess GHG budgets on a local and global scale, long-term measurements of CH4 and N2O 6 fluxes in combination with CO₂ exchange are necessary, especially over ecosystems that are 7 normally characterized by low GHG fluxes. In addition, we recommend to carefully check 8 flux results and underlying cospectra for an overestimation in the low spectral range and 9 correct for this effect if necessary.

10

11 6 Acknowledgements

This study was financially supported by the Austrian National Science Fund (FWF) under contract P23267-B16, the Tyrolean Science Fund under contract Uni-404/1083 and the EU framework 7 project GHG Europe (EU contract no. 244122). Family Hofer (Neustift, Austria) is acknowledged for granting us access to the study site.

16

17 7 References

18 Allan, W., Struthers, H. and Lowe, D. C.: Methane carbon isotope effects caused by atomic

19 chlorine in the marine boundary layer: Global model results compared with Southern

20 Hemisphere measurements, J. Geophys. Res., 112(D4), D04306, doi:10.1029/2006JD007369,

21 2007.

22 Baldocchi, D. D., Hincks, B. B. and Meyers, T. P.: Measuring Biosphere-Atmosphere

- Exchanges of Biologically Related Gases with Micrometeorological Methods, Ecology, 69(5),
 1331, doi:10.2307/1941631, 1988.
- 25 Baldocchi, D., Detto, M., Sonnentag, O., Verfaillie, J., Teh, Y. A., Silver, W. and Kelly, N.
- 26 M.: The challenges of measuring methane fluxes and concentrations over a peatland pasture,
- 27 Agric. For. Meteorol., 153, 177–187, doi:10.1016/j.agrformet.2011.04.013, 2012.
- 28 Bamberger, I., Hörtnagl, L., Ruuskanen, T. M., Schnitzhofer, R., Müller, M., Graus, M., Karl,
- 29 T., Wohlfahrt, G. and Hansel, A.: Deposition Fluxes of Terpenes over Grassland., J. Geophys.
- 30 Res. Atmos. JGR, 116(D14), D14305, doi:10.1029/2010JD015457, 2011.
- 31 Bamberger, I., Hörtnagl, L., Schnitzhofer, R., Graus, M., Ruuskanen, T. M., Müller, M.,
- 32 Dunkl, J., Wohlfahrt, G. and Hansel, A.: BVOC fluxes above mountain grassland,
- 33 Biogeosciences, 7(5), 1413–1424, doi:10.5194/bg-7-1413-2010, 2010.

- 1 Barnard, R., Leadley, P. W. and Hungate, B. A.: Global change, nitrification, and
- 2 denitrification: A review, Global Biogeochem. Cycles, 19(1), GB1007,
- 3 doi:10.1029/2004GB002282, 2005.
- 4 Bijoor, N. S., Czimczik, C. I., Pataki, D. E. and Billings, S. A.: Effects of temperature and
- 5 fertilization on nitrogen cycling and community composition of an urban lawn, Glob. Chang.
- 6 Biol., 14(9), 2119–2131, doi:10.1111/j.1365-2486.2008.01617.x, 2008.
- 7 Boucher, O., Friedlingstein, P., Collins, B. and Shine, K. P.: The indirect global warming
- 8 potential and global temperature change potential due to methane oxidation, Environ. Res.
- 9 Lett., 4(4), 044007, doi:10.1088/1748-9326/4/4/044007, 2009.
- 10 Bousquet, P., Ringeval, B., Pison, I., Dlugokencky, E. J., Brunke, E.-G., Carouge, C.,
- 11 Chevallier, F., Fortems-Cheiney, A., Frankenberg, C., Hauglustaine, D. A., Krummel, P. B.,
- 12 Langenfelds, R. L., Ramonet, M., Schmidt, M., Steele, L. P., Szopa, S., Yver, C., Viovy, N.
- and Ciais, P.: Source attribution of the changes in atmospheric methane for 2006–2008,
- 14 Atmos. Chem. Phys., 11(8), 3689–3700, doi:10.5194/acp-11-3689-2011, 2011.
- 15 Brilli, F., Hörtnagl, L., Bamberger, I., Schnitzhofer, R., Ruuskanen, T. M., Hansel, A., Loreto,
- 16 F. and Wohlfahrt, G.: Qualitative and quantitative characterization of volatile organic
- 17 compound emissions from cut grass., Environ. Sci. Technol., 46(7), 3859–65,
- 18 doi:10.1021/es204025y, 2012.
- 19 Brilli, F., Hörtnagl, L., Hammerle, A., Haslwanter, A., Hansel, A., Loreto, F. and Wohlfahrt,
- 20 G.: Leaf and ecosystem response to soil water availability in mountain grasslands, Agric. For.
- 21 Meteorol., 151(12), 1731–1740, doi:10.1016/j.agrformet.2011.07.007, 2011.
- 22 Chapuis-Lardy, L., Wrage, N., Metay, A., Chotte, J.-L. and Bernoux, M.: Soils, a sink for
- 23 N2O? A review, Glob. Chang. Biol., 13(1), 1–17, doi:10.1111/j.1365-2486.2006.01280.x,
- 24 2007.
- Cicerone, R. J. and Oremland, R. S.: Biogeochemical aspects of atmospheric methane, Global
 Biogeochem. Cycles, 2(4), 299–327, doi:10.1029/GB002i004p00299, 1988.
- Clough, T. J., Sherlock, R. R. and Rolston, D. E.: A Review of the Movement and Fate of
 N2O in the Subsoil, Nutr. Cycl. Agroecosystems, 72(1), 3–11, doi:10.1007/s10705-004-7349z, 2005.
- Collins, W. J., Sitch, S. and Boucher, O.: How vegetation impacts affect climate metrics for ozone precursors, J. Geophys. Res., 115(D23), D23308, doi:10.1029/2010JD014187, 2010.
- Crutzen, P. and Lelieveld, J.: Human Impacts on Atmospheric Chemistry, Annu. Rev. Earth
 Planet. Sci., 29(1), 17–45, doi:10.1146/annurev.earth.29.1.17, 2001.
- 34 Davidson, E.: The contribution of manure and fertilizer nitrogen to atmospheric nitrous oxide 25 = 1860 Net Council 2(0) (50 (62 dai 10.1028/mars 608.2000)
- 35 since 1860, Nat. Geosci., 2(9), 659–662, doi:10.1038/ngeo608, 2009.

- 1 Dijkstra, F. A., Morgan, J. A., Follett, R. F. and Lecain, D. R.: Climate change reduces the net
- 2 sink of CH4 and N2O in a semiarid grassland., Glob. Chang. Biol., 19(6), 1816–26,
- 3 doi:10.1111/gcb.12182, 2013.
- 4 Dlugokencky, E. J., Bruhwiler, L., White, J. W. C., Emmons, L. K., Novelli, P. C., Montzka,
- 5 S. A., Masarie, K. A., Lang, P. M., Crotwell, A. M., Miller, J. B. and Gatti, L.V.:
- 6 Observational constraints on recent increases in the atmospheric CH4 burden, Geophys. Res.
- 7 Lett., 36(18), L18803, doi:10.1029/2009GL039780, 2009.
- 8 Dubbs, L. L. and Whalen, S. C.: Reduced net atmospheric CH4 consumption is a sustained
- 9 response to elevated CO2 in a temperate forest, Biol. Fertil. Soils, 46(6), 597–606,
- 10 doi:10.1007/s00374-010-0467-7, 2010.
- 11 Dueck, T. A., de Visser, R., Poorter, H., Persijn, S., Gorissen, A., de Visser, W.,
- 12 Schapendonk, A., Verhagen, J., Snel, J., Harren, F. J. M., Ngai, A. K. Y., Verstappen, F.,
- 13 Bouwmeester, H., Voesenek, L. A. C. J. and van der Werf, A.: No evidence for substantial
- 14 aerobic methane emission by terrestrial plants: a ¹³C-labelling approach., New Phytol.,
- 15 175(1), 29–35, doi:10.1111/j.1469-8137.2007.02103.x, 2007.
- 16 Ehhalt, D. H. and Heidt, L. E.: Vertical profiles of CH4 in the troposphere and stratosphere, J.
- 17 Geophys. Res., 78(24), 5265–5271, doi:10.1029/JC078i024p05265, 1973.
- 18 Firestone, M. and Davidson, E.: Microbiological basis of NO and N2O production and
- 19 consumption in soil, in: Exchange of trace gases between terrestrial ecosystems and the 20 atmosphere, edited by M. Andreae and D. Schimel, pp. 7–21, Wiley., 1989.
- 21 Von Fischer, J. C. and Hedin, L. O.: Controls on soil methane fluxes: Tests of biophysical
- mechanisms using stable isotope tracers, Global Biogeochem. Cycles, 21(2), GB2007,
 doi:10.1029/2006GB002687, 2007.
- 24 Flechard, C. R., Neftel, A., Jocher, M., Ammann, C. and Fuhrer, J.: Bi-directional
- soil/atmosphere N2O exchange over two mown grassland systems with contrasting
- 26 management practices, Glob. Chang. Biol., 11(12), 2114–2127, doi:10.1111/j.1365-
- 27 2486.2005.01056.x, 2005.
- Foken, T. and Wichura, B.: Tools for quality assessment of surface-based flux measurements,
 Agric. For. Meteorol., 78(1-2), 83–105, doi:10.1016/0168-1923(95)02248-1, 1996.
- 30 Forster, P., Ramaswamy, V., Artaxo, P., Berntsen, T., Betts, R., Fahey, D. W., Haywood, J.,
- Lean, J., Lowe, D.C., Myhre, G., Nganga, J., Prinn, R., Raga, G., Schulz, M. and Van
- 32 Dorland, R.: Changes in Atmospheric Constituents and in Radiative Forcing, in: Climate
- 33 Change 2007: The Physical Science Basis. Contribution of Working Group I to the Fourth
- 34 Assessment Report of the Intergovernmental Panel on Climate Change [Solomon, S., D. Qin,
- 35 M. Manning, Z. Chen, M. Marquis, K.B. Averyt, M.Tignor and H.L. Miller (eds.)].
- 36 Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2007.
- 37 Fowler, D., Pilegaard, K., Sutton, M. A., Ambus, P., Raivonen, M., Duyzer, J., Simpson, D.,
- 38 Fagerli, H., Fuzzi, S., Schjoerring, J. K., Granier, C., Neftel, A., Isaksen, I. S. A., Laj, P.,
- 39 Maione, M., Monks, P. S., Burkhardt, J., Daemmgen, U., Neirynck, J., Personne, E., Wichink-

- 1 Kruit, R., Butterbach-Bahl, K., Flechard, C., Tuovinen, J. P., Coyle, M., Gerosa, G., Loubet,
- 2 B., Altimir, N., Gruenhage, L., Ammann, C., Cieslik, S., Paoletti, E., Mikkelsen, T. N., Ro-
- 3 Poulsen, H., Cellier, P., Cape, J. N., Horváth, L., Loreto, F., Niinemets, Ü., Palmer, P. I.,
- 4 Rinne, J., Misztal, P., Nemitz, E., Nilsson, D., Pryor, S., Gallagher, M. W., Vesala, T., Skiba,
- 5 U., Brüggemann, N., Zechmeister-Boltenstern, S., Williams, J., O'Dowd, C., Facchini, M. C.,
- 6 de Leeuw, G., Flossman, A., Chaumerliac, N. and Erisman, J. W.: Atmospheric composition
- 7 change: Ecosystems–Atmosphere interactions, Atmos. Environ., 43(33), 5193–5267,
- 8 doi:10.1016/j.atmosenv.2009.07.068, 2009.
- 9 Goldberg, S. D. and Gebauer, G.: Drought turns a Central European Norway spruce forest soil
- 10 from an N2O source to a transient N2O sink, Glob. Chang. Biol., 15(4), 850–860,
- 11 doi:10.1111/j.1365-2486.2008.01752.x, 2009a.
- 12 Goldberg, S. D. and Gebauer, G.: N2O and NO fluxes between a Norway spruce forest soil
- and atmosphere as affected by prolonged summer drought, Soil Biol. Biochem., 41(9), 1986–
 14 1995, doi:10.1016/j.soilbio.2009.07.001, 2009b.
- Granli, T. and Bockman, O. C.: Norwegian Journal of Agricultural Science Supplement, 12thed., 1994.
- 17 Hammerle, A., Haslwanter, A., Tappeiner, U., Cernusca, A. and Wohlfahrt, G.: Leaf area
- 18 controls on energy partitioning of a temperate mountain grassland, Biogeosciences, 5(2), 421-
- 19 431, doi:10.5194/bg-5-421-2008, 2008.
- 20 Hartmann, A. A., Buchmann, N. and Niklaus, P. A.: A study of soil methane sink regulation
- in two grasslands exposed to drought and N fertilization, Plant Soil, 342(1-2), 265–275,
 doi:10.1007/s11104-010-0690-x, 2010.
- 23 Hendriks, D. M. D., van Huissteden, J., Dolman, A. J. and van der Molen, M. K.: The full
- 24 greenhouse gas balance of an abandoned peat meadow, Biogeosciences, 4(3), 411–424,
- 25 doi:10.5194/bg-4-411-2007, 2007.
- 26 Hörtnagl, L., Bamberger, I., Graus, M., Ruuskanen, T. M., Schnitzhofer, R., Müller, M.,
- 27 Hansel, A. and Wohlfahrt, G.: Biotic, abiotic, and management controls on methanol
- 28 exchange above a temperate mountain grassland, J. Geophys. Res., 116(G3), 1–15,
- 29 doi:10.1029/2011JG001641, 2011.
- 30 Hörtnagl, L., Bamberger, I., Graus, M., Ruuskanen, T. M., Schnitzhofer, R., Walser, M.,
- 31 Unterberger, A., Hansel, A. and Wohlfahrt, G.: Acetaldehyde exchange above a managed
- 32 temperate mountain grassland., Atmos. Chem. Phys., 14, 5369-5391, doi:10.5194/acp-14-
- 33 5369-2014, 2014.
- Hörtnagl, L., Clement, R., Graus, M., Hammerle, A., Hansel, A. and Wohlfahrt, G.: Dealing
- 35 with disjunct concentration measurements in eddy covariance applications: A comparison of
- 36 available approaches, Atmos. Environ., 44(16), 2024–2032,
- 37 doi:10.1016/j.atmosenv.2010.02.042, 2010.

- 1 Holtan-Hartwig, L., Dörsch, P., Bakken, L.R.: Low temperature control of soil denitrifying
- 2 communities: kinetics of N₂O production and reduction, Soil Biol. & Biochem., 34, 1797-
- 3 1806, 2002.
- 4 Hsieh, C.-I., Katul, G. and Chi, T.: An approximate analytical model for footprint estimation
- 5 of scalar fluxes in thermally stratified atmospheric flows, Adv. Water Resour., 23(7), 765–
- 6 772, doi:10.1016/S0309-1708(99)00042-1, 2000.
- 7 Hu, Y., Chang, X., Lin, X., Wang, Y., Wang, S., Duan, J., Zhang, Z., Yang, X., Luo, C., Xu,
- 8 G. and Zhao, X.: Effects of warming and grazing on N2O fluxes in an alpine meadow
- 9 ecosystem on the Tibetan plateau, Soil Biol. Biochem., 42(6), 944–952,
- 10 doi:10.1016/j.soilbio.2010.02.011, 2010.
- 11 Imer, D., Merbold, L., Eugster, W. and Buchmann, N.: Temporal and spatial variations of soil
- 12 CO2, CH4 and N2O fluxes at three differently managed grasslands, Biogeosciences, 10(9), 13 5931–5945, doi:10.5194/bg-10-5931-2013, 2013.
- 14 Ineson, P., Coward, P. A. and Hartwig, U. A.: Soil gas fluxes of N2O, CH4 and CO2 beneath
- 15 Lolium perenne under elevated CO2 : The Swiss free air carbon dioxide enrichment
- 16 experiment, , 89–95, 1998.
- 17 Jackowicz-Korczyński, M., Christensen, T. R., Bäckstrand, K., Crill, P., Friborg, T.,
- 18 Mastepanov, M. and Ström, L.: Annual cycle of methane emission from a subarctic peatland,
- 19 J. Geophys. Res., 115(G2), G02009, doi:10.1029/2008JG000913, 2010.
- 20 Jäger, N., Duffner, A., Ludwig, B. and Flessa, H.: Effect of fertilization history on short-term
- 21 emission of CO2 and N2O after the application of different N fertilizers a laboratory study,
- 22 Arch. Agron. Soil Sci., 59(2), 161–171, doi:10.1080/03650340.2011.621420, 2013.
- 23 Jones, S. K., Famulari, D., Di Marco, C. F., Nemitz, E., Skiba, U. M., Rees, R. M. and Sutton,
- 24 M. A.: Nitrous oxide emissions from managed grassland: a comparison of eddy covariance
- and static chamber measurements, Atmos. Meas. Tech., 4(10), 2179–2194, doi:10.5194/amt-
- 264-2179-2011, 2011.
- 27 Karl, T. G., Spirig, C., Rinne, J., Stroud, C., Prevost, P., Greenberg, J., Fall, R. and Guenther,
- A.: Virtual disjunct eddy covariance measurements of organic compound fluxes from a
- subalpine forest using proton transfer reaction mass spectrometry, Atmos. Chem. Phys., 2(4),
- 30 279–291, 2002.
- 31 Keppler, F., Hamilton, J. T. G., Brass, M. and Röckmann, T.: Methane emissions from
- 32 terrestrial plants under aerobic conditions., Nature, 439(7073), 187–91,
- 33 doi:10.1038/nature04420, 2006.
- Khalil, M. A. K., Butenhoff, C. L. and Rasmussen, R. A.: Atmospheric methane: trends and
 cycles of sources and sinks., Environ. Sci. Technol., 41(7), 2131–7, 2007.
- 36 Khalil, M. A. K. and Rasmussen, R. A.: Climate-induced feedbacks for the global cycles of
- 37 methane and nitrous oxide, Tellus B, 41B(5), 554–559, doi:10.1111/j.1600-
- 38 0889.1989.tb00141.x, 1989.

- 1 Kirschke, S., Bousquet, P., Ciais, P., Saunois, M., Canadell, J. G., Dlugokencky, E. J.,
- 2 Bergamaschi, P., Bergmann, D., Blake, D. R., Bruhwiler, L., Cameron-Smith, P., Castaldi, S.,
- 3 Chevallier, F., Feng, L., Fraser, A., Heimann, M., Hodson, E. L., Houweling, S., Josse, B.,
- 4 Fraser, P. J., Krummel, P. B., Lamarque, J.-F., Langenfelds, R. L., Le Quéré, C., Naik, V.,
- 5 O'Doherty, S., Palmer, P. I., Pison, I., Plummer, D., Poulter, B., Prinn, R. G., Rigby, M.,
- 6 Ringeval, B., Santini, M., Schmidt, M., Shindell, D. T., Simpson, I. J., Spahni, R., Steele, L.
- 7 P., Strode, S. A., Sudo, K., Szopa, S., van der Werf, G. R., Voulgarakis, A., van Weele, M.,
- 8 Weiss, R. F., Williams, J. E. and Zeng, G.: Three decades of global methane sources and
- 9 sinks, Nat. Geosci., 6(10), 813–823, doi:10.1038/ngeo1955, 2013.
- 10 Kroon, P.S., Schrier-Uijl, A.P., Hensen, A., Veenendaal, E.M. and Jonker, H.J.J.: Annual
- balances of CH₄ and N₂O from a managed fen meadow using eddy covariance flux
- 12 measurements, Eur. J. Soil Sci., 61, 773-784, doi: 10.1111/j.1365-2389.2010.01273.x, 2010.
- 13 Lam, S. K., Lin, E., Norton, R. and Chen, D.: The effect of increased atmospheric carbon
- 14 dioxide concentration on emissions of nitrous oxide, carbon dioxide and methane from a
- 15 wheat field in a semi-arid environment in northern China, Soil Biol. Biochem., 43(2), 458–
- 16 461, doi:10.1016/j.soilbio.2010.10.012, 2011.
- 17 Li, C., Frolking, S. and Frolking, T. A.: A model of nitrous oxide evolution from soil driven
- 18 by rainfall events: 2. Model applications, J. Geophys. Res. Atmos., 97(D9), 9777–9783,
- 19 doi:10.1029/92JD00510, 1992.
- 20 Liebig, M. A., Gross, J. R., Kronberg, S. L., Phillips, R. L. and Hanson, J. D.: Grazing
- 21 management contributions to net global warming potential: a long-term evaluation in the
- 22 Northern Great Plains., J. Environ. Qual., 39(3), 799–809, doi:10.2134/jeq2009.0272, 2009.
- 23 Mander, Ü., Uuemaa, E., Kull, A., Kanal, A., Maddison, M., Soosaar, K., Salm, J.-O., Lesta,
- 24 M., Hansen, R., Kuller, R., Harding, A. and Augustin, J.: Assessment of methane and nitrous
- 25 oxide fluxes in rural landscapes, Landsc. Urban Plan., 98(3-4), 172–181,
- 26 doi:10.1016/j.landurbplan.2010.08.021, 2010.
- 27 Massman, W. J.: A simple method for estimating frequency response corrections for eddy
- covariance systems, Agric. For. Meteorol., 104(3), 247–251, doi:10.1016/S01681923(00)00164-7, 2000.
- McMillen, R. T.: An eddy correlation technique with extended applicability to non-simple terrain, Boundary-Layer Meteorol., 43(3), 231–245, doi:10.1007/BF00128405, 1988.
- 32 Merbold, L., Eugster, W., Stieger, J., Zahniser, M., Nelson, D. and Buchmann, N.:
- 33 Greenhouse gas budget (CO2, CH4 and N2O) of intensively managed grassland following 34 restoration., Glob. Chang. Biol., doi:10.1111/gcb.12518, 2014.
- 35 Müller, M., Graus, M., Ruuskanen, T. M., Schnitzhofer, R., Bamberger, I., Kaser, L.,
- 36 Titzmann, T., Hörtnagl, L., Wohlfahrt, G., Karl, T. and Hansel, A.: First eddy covariance flux
- 37 measurements by PTR-TOF., Atmos. Meas. Tech., 3(2), 387–395, doi:10.5194/amt-3-387-
- 38 2010, 2010.

- 1 Neftel, A., Ammann, C., Fischer, C., Spirig, C., Conen, F., Emmenegger, L., Tuzson, B. and
- 2 Wahlen, S.: N2O exchange over managed grassland: Application of a quantum cascade laser
- 3 spectrometer for micrometeorological flux measurements, Agric. For. Meteorol., 150(6), 775-
- 4 785, doi:10.1016/j.agrformet.2009.07.013, 2010.
- 5 Niboyet, A., Brown, J. R., Dijkstra, P., Blankinship, J. C., Leadley, P. W., Le Roux, X.,
- 6 Barthes, L., Barnard, R. L., Field, C. B. and Hungate, B. A.: Global change could amplify fire
- 7 effects on soil greenhouse gas emissions., PLoS One, 6(6), e20105,
- 8 doi:10.1371/journal.pone.0020105, 2011.
- 9 Nisbet, E. G., Dlugokencky, E. J., and Bousquet, P.: Methane on the Rise Again, Science,
 343, 493-495, 2014.
- 11 Nisbet, R. E. R., Fisher, R., Nimmo, R. H., Bendall, D. S., Crill, P. M., Gallego-Sala, A. V.,
- 12 Hornibrook, E. R. C., López-Juez, E., Lowry, D., Nisbet, P. B. R., Shuckburgh, E. F.,
- 13 Sriskantharajah, S., Howe, C. J. and Nisbet, E. G.: Emission of methane from plants., Proc.
- 14 Biol. Sci., 276(1660), 1347–54, doi:10.1098/rspb.2008.1731, 2009.
- Peterjohn, W., Melillo, J. and Steudler, P.: Responses of trace gas fluxes and N availability to experimentally elevated soil temperatures, Ecol. Applications, 4(3), 617–625, 1994.
- 17 Phillips, R. L., Whalen, S. C. and Schlesinger, W. H.: Influence of atmospheric CO2
- enrichment on methane consumption in a temperate forest soil, Glob. Chang. Biol., 7(5), 557–
 563, doi:10.1046/j.1354-1013.2001.00432.x, 2001.
- Prather, M. J., Holmes, C. D. and Hsu, J.: Reactive greenhouse gas scenarios: Systematic
 exploration of uncertainties and the role of atmospheric chemistry, Geophys. Res. Lett., 39(9),
- 22 L09803, doi:10.1029/2012GL051440, 2012.
- Prather, M. J. and Hsu, J.: Coupling of nitrous oxide and methane by global atmospheric
 chemistry., Science, 330(6006), 952–4, doi:10.1126/science.1196285, 2010.
- 25 Revell, L. E., Bodeker, G. E., Smale, D., Lehmann, R., Huck, P. E., Williamson, B. E.,
- Rozanov, E. and Struthers, H.: The effectiveness of N2O in depleting stratospheric ozone,
 Geophys. Res. Lett., 39(15), L15806, doi:10.1029/2012GL052143, 2012.
- 28 Rinne, J., Riutta, T., Pihlatie, M. and Aurela, M.: Annual cycle of methane emission from a
- boreal fen measured by the eddy covariance technique, Tellus B, 59(3), 449–457,
- 30 doi:10.1111/j.1600-0889.2007.00261.x, 2007.
- 31 Röver, M., Heinemeyer, O. and Kaiser, E.-A.: Microbial induced nitrous oxide emissions
- from an arable soil during winter, Soil Biol. Biochem., 30(14), 1859–1865,
- 33 doi:10.1016/S0038-0717(98)00080-7, 1998.
- 34 Ruser, R., Flessa, H., Russow, R., Schmidt, G., Buegger, F. and Munch, J. C.: Emission of
- 35 N2O, N2 and CO2 from soil fertilized with nitrate: effect of compaction, soil moisture and
- 36 rewetting, Soil Biol. Biochem., 38(2), 263–274, doi:10.1016/j.soilbio.2005.05.005, 2006.

- 1 Ruuskanen, T. M., Müller, M., Schnitzhofer, R., Karl, T., Graus, M., Bamberger, I., Hörtnagl,
- 2 L., Brilli, F., Wohlfahrt, G. and Hansel, A.: Eddy covariance VOC emission and deposition
- 3 fluxes above grassland using PTR-TOF., Atmos. Chem. Phys., 11(2), 611–625,
- 4 doi:10.5194/acp-11-611-2011, 2011.
- 5 Schlesinger, W. H.: An estimate of the global sink for nitrous oxide in soils., Glob. Chang.
- 6 Biol., doi:10.1111/gcb.12239, 2013.
- 7 Schrier-Uijl, A. P., Kroon, P. S., Hensen, A., Leffelaar, P. A., Berendse, F. and Veenendaal,
- 8 E. M.: Comparison of chamber and eddy covariance-based CO2 and CH4 emission estimates
- 9 in a heterogeneous grass ecosystem on peat, Agric. For. Meteorol., 150(6), 825–831,
- 10 doi:10.1016/j.agrformet.2009.11.007, 2010.
- 11 Schulze, E. D., Luyssaert, S., Ciais, P., Freibauer, A., Janssens et al., I. A., Soussana, J. F.,
- 12 Smith, P., Grace, J., Levin, I., Thiruchittampalam, B., Heimann, M., Dolman, A. J., Valentini,
- 13 R., Bousquet, P., Peylin, P., Peters, W., Rödenbeck, C., Etiope, G., Vuichard, N., Wattenbach,
- 14 M., Nabuurs, G. J., Poussi, Z., Nieschulze, J. and Gash, J. H.: Importance of methane and
- 15 nitrous oxide for Europe's terrestrial greenhouse-gas balance, Nat. Geosci., 2(12), 842–850,
- 16 doi:10.1038/ngeo686, 2009.
- 17 Shindell, D. T., Faluvegi, G., Koch, D. M., Schmidt, G. A., Unger, N. and Bauer, S. E.:
- 18 Improved attribution of climate forcing to emissions., Science, 326(5953), 716–8,
- 19 doi:10.1126/science.1174760, 2009.
- 20 Sjogersten, S. and Wookey, P. A.: Spatio-temporal variability and environmental controls of
- 21 methane fluxes at the forest-tundra ecotone in the Fennoscandian mountains, Glob. Chang.
- 22 Biol., 8(9), 885–894, doi:10.1046/j.1365-2486.2002.00522.x, 2002.
- 23 Smith, K. A., Dobbie, K. E., Ball, B. C., Bakken, L. R., Sitaula, B. K., Hansen, S., Brumme,
- 24 R., Borken, W., Christensen, S., Priemé, A., Fowler, D., Macdonald, J. A., Skiba, U.,
- 25 Klemedtsson, L., Kasimir-Klemedtsson, A., Degórska, A. and Orlanski, P.: Oxidation of

26 atmospheric methane in Northern European soils, comparison with other ecosystems, and

- 27 uncertainties in the global terrestrial sink, Glob. Chang. Biol., 6(7), 791–803,
- 28 doi:10.1046/j.1365-2486.2000.00356.x, 2000.
- 29 Soussana, J. F., Allard, V., Pilegaard, K., Ambus, P., Amman, C., Campbell, C., Ceschia, E.,
- 30 Clifton-Brown, J., Czobel, S., Domingues, R., Flechard, C., Fuhrer, J., Hensen, A., Horvath,
- 31 L., Jones, M., Kasper, G., Martin, C., Nagy, Z., Neftel, A., Raschi, A., Baronti, S., Rees, R.
- 32 M., Skiba, U., Stefani, P., Manca, G., Sutton, M., Tuba, Z. and Valentini, R.: Full accounting
- 33 of the greenhouse gas (CO2, N2O, CH4) budget of nine European grassland sites, Agric.
- 34 Ecosyst. Environ., 121(1-2), 121–134, doi:10.1016/j.agee.2006.12.022, 2007.
- 35 Syakila, A. and Kroeze, C.: The global nitrous oxide budget revisited, Greenh. Gas Meas.
- 36 Manag., 1(1), 17–26, doi:10.3763/ghgmm.2010.0007, 2011.
- 37 Syakila, A., Kroeze, C. and Slomp, C. P.: Neglecting sinks for N2O at the earth's surface:
- 38 does it matter?, J. Integr. Environ. Sci., 7(sup1), 79–87, doi:10.1080/1943815X.2010.497492,
- 39 2010.

- 1 Teepe, R., Brumme, R. and Beese, F.: Nitrous oxide emissions from soil during freezing and
- thawing periods, Soil Biol. Biochem., 33(9), 1269–1275, doi:10.1016/S0038-0717(01)000840, 2001.
- 4 Tian, H., Chen, G., Lu, C., Xu, X., Hayes, D. J., Ren, W., Pan, S., Huntzinger, D. N. and
- 5 Wofsy, S. C.: North American terrestrial CO2 uptake largely offset by CH4 and N2O
- 6 emissions: toward a full accounting of the greenhouse gas budget, Clim. Change,
- 7 doi:10.1007/s10584-014-1072-9, 2014.

- 10 Vieten, B., Conen, F., Seth, B. and Alewell, C.: The fate of N2O consumed in soils,
- 11 Biogeosciences, 5, 129–132, 2008.
- 12 Wertz, S., Goyer, C., Zebarth, B. J., Burton, D. L., Tatti, E., Chantigny, M. H. and Filion, M.:
- 13 Effects of temperatures near the freezing point on N2O emissions, denitrification and on the
- 14 abundance and structure of nitrifying and denitrifying soil communities., FEMS Microbiol.
- 15 Ecol., 83(1), 242–54, doi:10.1111/j.1574-6941.2012.01468.x, 2013.
- Whalen, S. C.: Natural Wetlands and the Atmosphere, Env. Engineering Sc., 22(1), 73-94,
 2005.
- 18 Wohlfahrt, G., Anderson-Dunn, M., Bahn, M., Balzarolo, M., Berninger, F., Campbell, C.,
- 19 Carrara, A., Cescatti, A., Christensen, T., Dore, S., Eugster, W., Friborg, T., Furger, M.,
- 20 Gianelle, D., Gimeno, C., Hargreaves, K., Hari, P., Haslwanter, A., Johansson, T., Marcolla,
- 21 B., Milford, C., Nagy, Z., Nemitz, E., Rogiers, N., Sanz, M. J., Siegwolf, R. T. W., Susiluoto,
- 22 S., Sutton, M., Tuba, Z., Ugolini, F., Valentini, R., Zorer, R. and Cernusca, A.: Biotic,
- 23 Abiotic, and Management Controls on the Net Ecosystem CO2 Exchange of European
- 24 Mountain Grassland Ecosystems, Ecosystems, 11(8), 1338–1351, doi:10.1007/s10021-008-
- 25 9196-2, 2008a.
- 26 Wohlfahrt, G., Anfang, C., Bahn, M., Haslwanter, A., Newesely, C., Schmitt, M., Drosler, M.,
- 27 Pfadenhauer, J. and Cernusca, A.: Quantifying nighttime ecosystem respiration of a meadow
- using eddy covariance, chambers and modelling, Agric. For. Meteorol., 128(3-4), 141–162,
- 29 doi:10.1016/j.agrformet.2004.11.003, 2005a.
- 30 Wohlfahrt, G., Bahn, M., Haslwanter, A., Newesely, C. and Cernusca, A.: Estimation of
- 31 daytime ecosystem respiration to determine gross primary production of a mountain meadow, 32 $A_{11} = E_{12} + E_{$
- 32 Agric. For. Meteorol., 130(1-2), 13–25, doi:10.1016/j.agrformet.2005.02.001, 2005b.
- 33 Wohlfahrt, G., Hammerle, A., Haslwanter, A., Bahn, M., Tappeiner, U. and Cernusca, A.:
- 34 Seasonal and inter-annual variability of the net ecosystem CO2 exchange of a temperate
- 35 mountain grassland: Effects of weather and management, J. Geophys. Res., 113(D08110),
- 36 doi:10.1029/2007JD009286, 2008b.
- 37 Wu, D., Dong, W., Oenema, O., Wang, Y., Trebs, I. and Hu, C.: N2O consumption by low-
- nitrogen soil and its regulation by water and oxygen, Soil Biol. Biochem., 60, 165–172,
 doi:10.1016/j.soilbio.2013.01.028.2013
- 39 doi:10.1016/j.soilbio.2013.01.028, 2013.

⁸ Topp, E. and Pattey, E.: Soils as sources and sinks for atmospheric methane, Can. J. Soil Sci.,
9 77(2), 167–177, doi:10.4141/S96-107, 1997.

- 1 Xu-Ri and Prentice, I.: Terrestrial nitrogen cycle simulation with a dynamic global vegetation
- 2 model, Glob. Chang. Biol., 14(8), 1745–1764, doi:10.1111/j.1365-2486.2008.01625.x, 2008.
- 3 Xu-Ri, Prentice, I. C., Spahni, R. and Niu, H. S.: Modelling terrestrial nitrous oxide emissions
- 4 and implications for climate feedback., New Phytol., 196(2), 472–88, doi:10.1111/j.1469-5 8137.2012.04269.x, 2012.
- 6 Yavitt, J. B., Fahey, T. J. and Simmons, J. A.: Methane and Carbon Dioxide Dynamics in a
- 7 Northern Hardwood Ecosystem, Soil Sci. Soc. Am. J., 59(3), 796,
- 8 doi:10.2136/sssaj1995.03615995005900030023x, 1995.
- 9 Yvon-Durocher, G., Allen, A. P., Bastviken, D., Conrad, R., Gudasz, C., St-Pierre, A., Thanh-
- 10 Duc, N. and del Giorgio, P. A.: Methane fluxes show consistent temperature dependence
- 11 across microbial to ecosystem scales, Nature, doi:10.1038/nature13164, 2014.
- 12 Zhu, R., Sun, L. and Ding, W.: Nitrous oxide emissions from tundra soil and snowpack in the
- 13 maritime Antarctic., Chemosphere, 59(11), 1667–75,
- 14 doi:10.1016/j.chemosphere.2004.10.033, 2005.
- 15 Zona, D., Janssens, I. A., Aubinet, M., Gioli, B., Vicca, S., Fichot, R. and Ceulemans, R.:
- 16 Fluxes of the greenhouse gases (CO2, CH4 and N2O) above a short-rotation poplar plantation
- 17 after conversion from agricultural land, Agric. For. Meteorol., 169, 100–110,
- 18 doi:10.1016/j.agrformet.2012.10.008, 2013.
- 19 Zumft, W. G. and Kroneck, P. M. H.: Respiratory transformation of nitrous oxide (N2O) to
- 20 dinitrogen by Bacteria and Archaea, Advances in Microbial Physiology, 52, 107-227, doi:
- 21 10.1016/S0065-2911(06)52003-X, 2007.

Table 1. Partial correlations of a multiple linear regression analysis and correlation coefficients (r) of a simple linear regression analysis using daily average values of ln transformed CH₄ (F_{CH4}) and N_2O (F_{N2O}) flux rates as dependent variables and air temperature (T_{air}), soil temperature (T_{soil}) and soil water content (SWC) in 5 cm depth, soil heat flux (SHF), net ecosystem CO₂ exchange (NEE), latent (LE) and sensible (H) heat flux, photosynthetically active radiation (PAR), vapor pressure deficit (VPD), relative air humidity (RHA) and CH₄ / N_2O volume mixing ratios (VMR) as independent variables. Management events were excluded from the analysis. Bold numbers highlight p < 0.05, except bold underlined numbers resulted in p < 0.001. Results shown for the "vegetation period" do not include time periods with snow cover on the meadow.

		MULTIPLE LINEAR REGRESSION partial correlations								SIMPLE LINEAR REGRESSION r						
<u>_</u>		vegetation period	snow melt – 1st cut			1st cut – 2nd cut		2nd cut – 3rd cut		3rd cut – snow cover		snow cover	vegetation period			
		2010-11	2010	2011	2010-2011	2010	2011	2010-2011	2010	2011	2010-2011	2010	2011	2010-2011	2010-2012	2010-2011
ln(F _{CH4})																
	T _{air}	<u>0.19</u>	0.07	0.07	0.25	-0.02	-0.35	0.11	-0.05	0.32	0.02	0.20	0.01	-0.02	0.17	0.13
	T _{soil}	-0.04	0.10	-0.07	-0.16	-0.13	0.57	0.10	-0.08	-0.12	-0.09	0.01	0.15	0.11	-0.11	0.16
	SWC	0.07	0.06	-0.24	0.04	-0.20	-0.13	0.13	0.03	0.33	0.05	0.33	-0.05	-0.09	-0.13	0.10
	SHF	<u>-0.22</u>	-0.14	-0.14	-0.26	0.12	0.22	-0.16	0.02	-0.28	0.01	-0.29	-0.08	-0.10	0.04	-0.09
	NEE	<u>0.20</u>	0.12	0.38	0.18	0.24	0.10	0.19	-0.04	0.05	0.01	0.33	0.18	0.32	0.20	<u>0.30</u>
	LE	-0.05	-0.16	-0.17	-0.12	0.09	-0.23	-0.05	-0.21	0.10	-0.17	0.20	0.48	<u>0.44</u>	0.28	<u>-0.19</u>
	н	-0.06	-0.08	-0.38	-0.13	-0.25	-0.03	-0.09	-0.08	0.13	0.10	0.01	-0.33	-0.18	-0.09	-0.19
	PAR	0.10	0.23	0.16	0.25	-0.08	-0.16	0.00	0.25	-0.20	0.06	-0.17	-0.07	-0.13	0.00	-0.19
	VPD	-0.07	0.08	0.02	-0.09	-0.01	0.10	-0.01	0.20	-0.26	0.19	-0.11	-0.16	-0.08	-0.12	-0.09
	RHA	0.03	0.12	0.07	0.06	0.12	0.05	0.05	0.30	-0.28	0.21	-0.31	0.03	0.02	-0.08	<u>0.23</u> 0.02
	CH ₄ VMR	0.01	0.08	0.00	0.02	0.15	0.39	0.06	-0.35	0.11	-0.15	0.35	-0.12	-0.11	0.01	0.02
	multiple r ²	<u>0.27</u> 356	0.31	<u>0.54</u>	0.20	0.43	0.62	<u>0.36</u>	0.41	0.23	0.18	0.55	0.53	<u>0.40</u>	0.22	
	Ν	356	47	67	114	50	36	86	44	40	84	35	37	72	82	365-397
ln(F _{N20})																
	\mathbf{T}_{air}	0.14	-0.04	0.27	0.03	0.28	-0.06	0.03	0.10	0.14	0.21	0.03	0.05	0.05	0.17	0.29
	T _{soil}	-0.11	0.09	-0.16	0.06	-0.16	0.22	0.30	-0.07	-0.06	-0.27	-0.22	-0.18	-0.33	-0.12	0.24
	SWC	<u>-0.24</u>	-0.13	-0.15	-0.24	-0.18	-0.27	-0.21	-0.38	-0.31	<u>-0.51</u>	0.01	-0.45	<u>-0.47</u>	-0.05	-0.33
	SHF	-0.02	0.03	-0.23	0.02	-0.26	0.16	-0.11	-0.12	-0.11	-0.14	0.42	0.19	0.24	-0.10	0.15
	NEE	<u>0.23</u>	-0.16	0.31	0.13	0.10	-0.10	0.10	0.35	0.24	0.32	0.32	-0.12	0.09	0.02	0.00
	LE	0.19	-0.16	-0.11	-0.08	-0.10	0.07	-0.03	0.04	0.19	0.11	0.41	0.05	0.17	0.18	0.22
	Н	-0.14	-0.24	-0.16	-0.23	-0.08	-0.20	-0.14	0.45	0.00	0.18	0.22	-0.10	0.13	-0.25	-0.20
	PAR	-0.02	0.21	0.21	0.13	0.37	-0.03	0.22	-0.22	0.18	0.02	-0.37	-0.04	-0.32	0.06	0.05
	VPD	0.01	-0.24	-0.11	-0.11	0.09	-0.04	0.13	0.26	0.05	0.23	-0.47	-0.03	-0.17	-0.10	0.16
	RHA	$\frac{0.24}{0.25}$	-0.21	0.18	-0.01	0.45	0.03	0.33	0.37	0.23	<u>0.37</u> -0.13	-0.60	0.07	-0.13	0.00	0.08
	N ₂ O VMR	0.25	-0.05	-0.11	0.02	0.39	0.28	0.26	-0.15	0.11	0.20	-0.26	-0.06	-0.04	<u>0.39</u>	<u>0.17</u>
	multiple r ² N	<u>0.42</u> 360	0.19 49	<u>0.55</u> 67	<u>0.26</u> 116	<u>0.76</u> 50	<u>0.73</u> 36	<u>0.66</u> 86	<u>0.72</u> 44	0.56 41	<u>0.68</u> 85	<u>0.73</u> 36	<u>0.68</u> 37	<u>0.73</u> 73	<u>0.44</u> 83	369-401

Table 2. Daily average means in three different groups of daily net CH₄ / N₂O exchange. Significant differences between group means were determined in a repeated measures ANOVA setting, using the Unequal N HSD post hoc test. Group labels to the right of a given group mean show to which flux group the respective value was significantly different. Bold numbers mark group means that were significantly different from one other group, except bold underlined numbers denote group means that were significantly different from both other groups. f+... daily average CH₄ / N₂O emission fluxes > 3 / 0.4 ng m⁻² s⁻¹, f0... fluxes between 3 / 0.4 and -3 / -0.4 ng m⁻² s⁻¹, f-... deposition fluxes < -3 / -0.4 ng m⁻² s⁻¹. 5

*	Unit	Mean values, standard deviations and significant differences											
compound			CH ₄		N ₂ O								
flux class		f+	f-	f0	f+	f-	f0						
T_{air}	°C	9.2 ±7.0 f0	9.6 ±6.0 f0	<u>5.9 ±8.5</u> f+,f-	10.1 ±7.5 f0	8.4 ±5.1 f0	<u>4.2 ±7.3</u> f+,f-						
T _{soil}	°C	10.8 ± 6.5	10.9 ±5.3	8.2 ± 6.7	11.5 ±6.8 f0	10.1 ±4.8 f0	<u>6.5 ±5.5</u> f+,f-						
SWC	m ³ m ⁻³	0.29 ± 0.06	0.28 ± 0.06	0.29 ± 0.09	<u>0.27 ±0.07</u> f-,f0	0.31 ±0.05 f+	0.32 ±0.06 f+						
SHF	W m ⁻²	1.0 ±6.9 f-	<u>3.9 ±6.6</u> f+,f0	0.5 ±6.4 f-	2.2 ±7.3 f0	1.4 ± 6.0	-1.5 ±6.1 f+						
NEE	μg CO ₂ m ⁻² s ⁻¹	-70 ±224 f-	-220 ±229 f+	-119 ±220	-106 ± 246	-141 ±220	-40 ± 172						
LE	W m ⁻²	55 ±53 f-	<u>86 ±58</u> f+,f0	50 ±57 f-	67 ±64 f0	64 ±44 f0	<u>30 ±37</u> f+,f-						
Н	W m ⁻²	6.9 ±22.0 f-	<u>20.2 ±22.7</u> f+,f0	8.8 ±19.8 f-	7.5 ±22.8 f-	<u>16.7 ±21.9</u> f+,f0	4.4 ±16.8 f-						
PAR	µmol m ⁻² s ⁻¹	271 ±158 f-	<u>372 ±169</u> f+,f0	250 ±168 f-	$293 \pm \! 180$	314 ±149 f0	217 ±139 f-						
VPD	kPa	0.33 ±0.28 f-	<u>0.42 ±0.26</u> f+,f0	0.28 ±0.29 f-	0.36 ± 0.30	0.35 ± 0.25	0.23 ±0.22						
RHA	%	81 ±10 f-	<u>75 ±10</u> f+,f0	82 ±11 f-	81 ±10 f-	77 ±11 f+	82 ±12						
VMR	ppb	$2014 \pm \! 59$	2004 ±53	2021 ±60	319 ±6 f-	317 ±4 f+	319 ±4						
Ν	days	294	96	48	261	138	44						

1 2

3

4

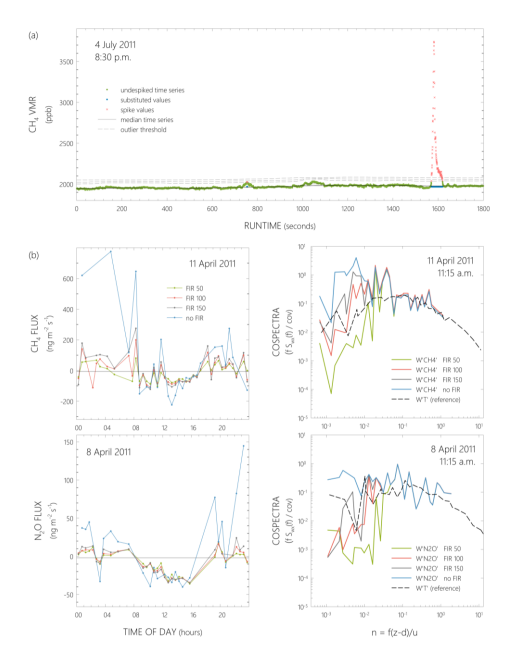


Figure 1. (a) Despiking example of 2 Hz methane VMRs using median filters. (b) Diurnal courses (left panels) and normalized co-spectra (right panels) illustrating the effect of highpass filtering CH_4 (upper panels) and N_2O (lower panels) time series with a non-recursive finite impulse response (FIR) filter with different time constants (50, 100 und 150 s). Sensible heat cospectra are shown in the right panels for reference.

- 7
- 8

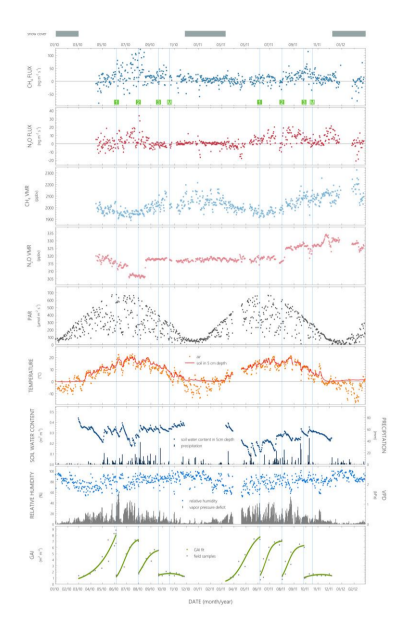


Figure 2. Daily average CH₄ and N₂O fluxes and volume mixing ratios (VMR), photosynthetically active radiation (PAR), air temperature, soil temperature at 5 cm depth, soil water content at 5 cm depth, relative air humidity, vapour pressure deficit, green plant area index (GAI) and daily sums of precipitation over 22 months of measurements between April 2010 and February 2012. Vertical lines show management dates, numbers 1, 2 and 3 in green squares indicate the 1st, 2nd and 3rd cutting of the meadow, respectively, while M denotes manure spreading.

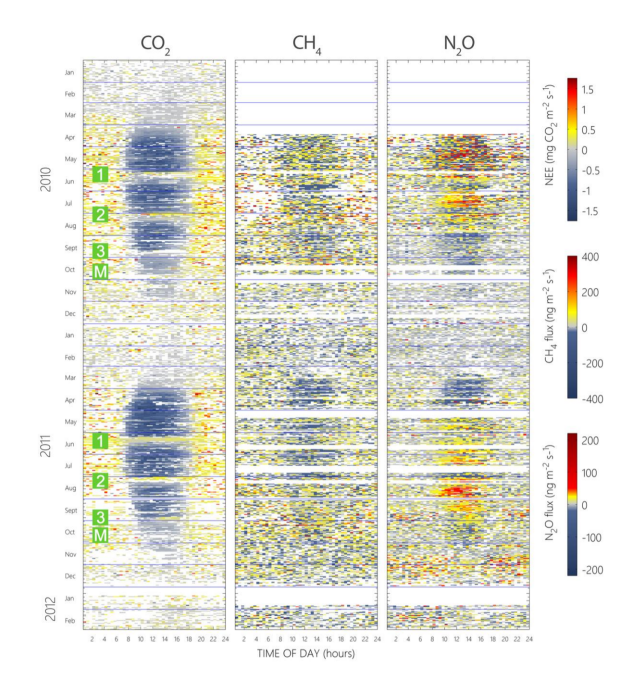


Figure 3. Half-hourly CO₂, CH₄ and N₂O fluxes over two years of GHG flux measurements.
Numbers 1, 2 and 3 in green squares indicate the 1st, 2nd and 3rd cutting of the meadow,
respectively, while M denotes manure spreading. Horizontal blue lines show the start and end
of months. White color marks missing data.

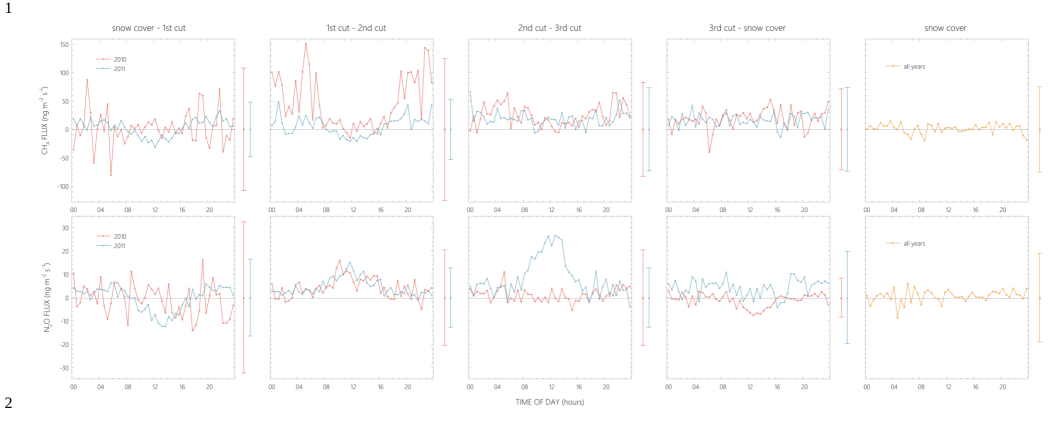
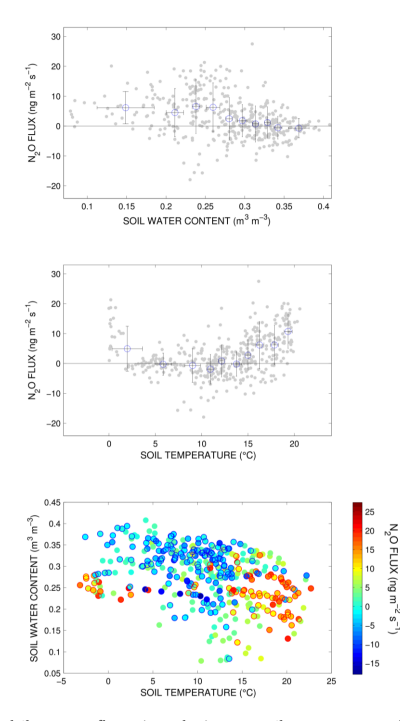
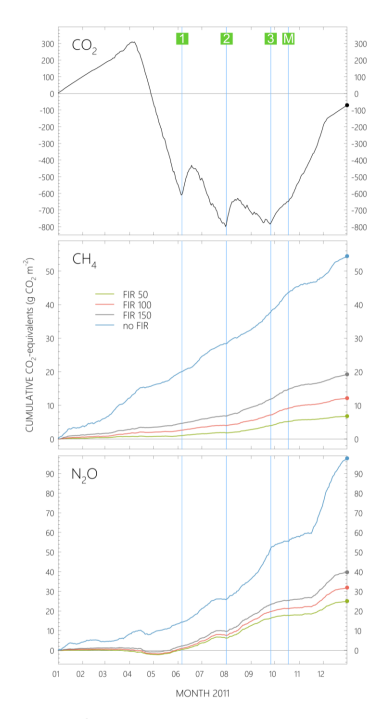


Figure 4. Diurnal cycles of CH_4 and N_2O fluxes during different time periods in 2010 and 2011. Whiskers to the right of each plot show the average standard deviation during the respective time period. Management data were excluded from the analysis.



1

Figure 5. N₂O daily average fluxes (grey dots) *versus* soil water content and soil temperature. Blue circles in the upper two panels show bin averages (40 days per bin), with error bars representing the standard deviation within each bin. In the lower panel, fluxes < 0 ng m⁻² s⁻¹ are circled in blue, fluxes > 9 ng m⁻² s⁻¹ are circled in red. Management events were excluded from the analysis.



1

Figure 6. Cumulative GHG fluxes in 2011 expressed as CO_2 -equivalents. The effect of the finite impulse response (FIR) filter with different time constants is shown for CH_4 and N_2O budgets. Vertical lines show management dates, numbers 1, 2 and 3 in green squares indicate the 1st, 2nd and 3rd cutting of the meadow, respectively, while M denotes manure spreading.