We thank both referees for their valuable comments and appreciate the associated improvements to the manuscript.

Referee#1:

"1. Despite continuous monitoring of N2O fluxes (Fig. 3) the authors give no values of cumulative losses per year and don't address this aspect in the discussion (see also details below). However it is quite important to interpret the impact of their findings, i.e. to which extent observed isotope fluxes are representative to agricultural ecosystems. So these numbers should be reported and discussed" *Answer: We addressed this comment by including a corresponding paragraph in section 4.7 (see below in the response to the specific comment "Section 4.7")*.

"2. The authors use the relationship between isotope values (average d15N, d18O and SP) to estimate and discuss possible N2O reduction. While this is justified, it should be better illustrated. Instead of showing isotope maps of d18O vs d15Nbulk only, they should also show SP vs d18O and also illustrate their postulated reduction events. Here I suggest to give isotope maps (SP/d18O and d15Nbulk/d18O) showing the change in values during estimated reduction events. Reduction vectors could be added in these figures to show agreement or deviation of observed data from previously reported reduction dynamics. These figures might be given in the appendix."

Answer: We changed the manuscript accordingly and included a SP/d18O map (see below in response to the specific comment "P1590, Section 4.4").

P1576 L 13.14: this reasoning is not exactly correct here: independence of SP from precursors is due to the fact that N20 is the first molecule with two N atoms and thus SP is not existent in precursor compounds *Answer: in order to be unequivocal we reworded to "…, as SP remained constant <u>in the de novo produced</u> <u>N₂O even though</u> …"*

L 26 suggest to address N2O isotopocules (or isotopologues) instead of SP since d15Nbulk and not SP is used in the cited examples

Answer: Agreed and changed to "In addition, N₂O isotopocules can be used for …"

P1577 L 5see previous comment

Answer: Agreed and changed to "N₂O isotopomers can be ...".

L7 flask sampling with chambers is better in spatial resolution than atmospheric measurement (N = 1) which gives not info on spatial resolution at all

Answer: We agree that measurements in the surface layer cannot be replicated and that for some experimental designs (e.g. different treatments on an experimental field), a chamber setup is indispensable. However, we want to raise the point that chamber measurements are representative for the usually small chamber area and are therefore limited in spatial representation. As a consequence, we changed the section to "... is limited in temporal resolution and spatial representation of a given site".

L 21 how about soil properties?

Answer: In this study, measurements were made above one intensively managed grassland site for which soil properties are assumed to be constant on the timescale of this study. Therefore, an analysis with respect to variations in soil properties is not possible.

L 22 goal (iii) can clearly not be achieved with this approach as there is no way to check the process information from isotopomers independently. Please modify accordingly or explain how you can test this with your approaches

Answer: We agree and removed (iii) from the objectives.

Study site: please report numbers on soil texture, Corg, C/N, pH and bulk density since these are very important to compare findings to other sites.

Answer: We included information on bulk density, texture, pH as well as C and N content in section 2.1

P 1579 L 16 suggest "(increase of 0.31: : : per mil.." Answer: Changed to "(increase by 0.31 per mille ...")

P 1583 L 21 please better explain "surface layer", lowest ten of m is quite vague, maybe add a reference here?

Answer: The term surface layer is now explained at the beginning of section 3.2: "Air samples were taken at 2.2 m height which is in the lowest 10% of the atmospheric boundary layer (ABL) where mechanical generation of turbulence exceeds buoyant generation or consumption. This part of the ABL is called surface layer, hence corresponding air samples are referred to as surface layer air samples.".

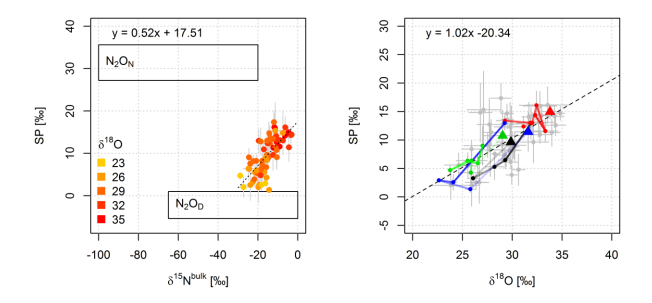
P 1587 L 4 data are representative for this site, but not for agricultural land in general, please clarify Answer: This sentence was meant to explain that simple averages do not represent the isotopic composition of any given treatment, site or ecosystem, but we agree that our wording might be misleading. Therefore we changed to a more general term: "Representative isotopic composition of N_2O emitted from a given site or treatment can be"

P1590 L 8 Please add that the preferred cleavage of N-O bonds between lighter isotopes leads to increasing d18O and SP in residual N2O

Answer: The explanation is included now: "However, in the terminal step of denitrification, namely the reduction of N₂O to N₂, N-O bonds between lighter isotopes are cleaved preferentially, leading to an increase in SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$ in the remaining N₂O".

P 1590, Section 4.4 suggest to add an SP/d18O plots (see discussion on problems of SP/d15N plots in Well et al 2012, Geochimica et Cosmochimica Acta 90, 265–282 Lewicka-Szczebak et al., 2014 Geochimica et Cosmochimica Acta, 134, 55-73, Lewicka-Szczebak et al 2015 Rapid Comm Mass Spectrometry 29:269-282). The extremely large range in d15N of the endmember areas make the use of d15N really difficult. Moreover, these ranges might even be larger than reported in the literature due to the unknown d15N of NO3 in active microsites. Conversely, d18O of N2O produced by denitrification is mostly governed by O exchange with soil water and thus less variable.

Answer: We agree with the referee and figure 7's right panel was substituted by a panel showing SP as a function of δ^{18} O (see uploaded figure). We also indicated the development of isotopic composition during management events and the rewetting event as suggested by the referee in the general comments to the manuscript. The text in section 4.4 was adapted accordingly.



L 15 to 20: please compare your endmember areas with those given and discussed in Zou et al. 2014 Soil Biology and Biochemistry, 77, 276–291.

Answer: Zou et al. give endmember areas in a SP/d15N map for the processes "Nitrification", "Nitrifierdentirification", "Fungal denitrification" and "Denitrifier-denitrification". While the SP-values representing the areas' corner nodes are literature values, the corresponding d15N-values were calculated from the analyzed isotopic composition of N2O precursors (NO3- / NH4+) and the range of fractionation factors derived from literature for the four process groups (see table below). As our isotopocule measurements integrate over a large and (depending on wind direction) variable area, it was not possible within the scope of the presented study to sample and analyze the isotopic composition of N2O substrates. We appreciate the referee comment and the approach chosen by Zou et al., but due to the lack of knowledge on the substrate's isotopic composition, we had to decide for a more general approach. To cover the whole range of potential d15Nbulk values, minimum and maximum values of fractionation factors reported in literature for the process groups N2O_N (nitrification and fungal denitrification) and N2O_D (denitrification and nitrifier denitrification) were selected and combined with the endpoints of literature values reported for the N2O precursors. Overall, as can be derived from the table below, these approaches are very much alike, but the lack of the substrate's isotopic composition does not allow for an application of the approach chosen by Zou et al..

Study	SP _{nit}	SP _{fungalDen}	SP _{N2ON}	SP _{den}	SP nit-den	SP _{N2OD}
Zou	31.4 (31.4-35.6)	37 (34.1-39.6)		-2 (-6.9-1.4)	-3.8 (-13.6-5)	
This			32.8 ± 4			-1.6 ± 3.8
study						
	Enit	$\mathcal{E}_{fungalDen}$	E _{N2ON}	Eden	Епнз/пог, Епог/пго	E _{N2OD}
Zou	Е nit -68 to -46,9	<i>E</i> fungalDen -36.2 to -24.1	Enzon	<i>Е</i> _{den} -37 to -11	Елнз/лог, Елог/лго -39.8 to -10.6	E _{N2OD}
Zou	-	, ,	En20N			E _{N2OD}
Zou This	-	, ,	£ _{N2ON} -90 to -40		-39.8 to -10.6	£ _{N2OD} -40 to -15

L 28 here and elsewhere: BACTERIAL denitrification, since isotopologues of fungal denitrification are close to NH2OH-N2O.

Answer: "bacterial" was added to revoke the unclear wording.

P 1591 L 6-7 you might refer to the discussion of this aspect in Well et al 2012, Geochimica et Cosmochimica Acta 90, 265–282.

Answer: Reference was added

P1592 L10 note that N2O_N includes Nitrification AND fungal denitrification. You can't exclude the fungi (see Sutka et al., 2008 Rapid Commun Mass Spectrom. 2008 3989-96,. Rohe et al., 2014 Rapid communications in mass spectrometry, 28, 1893-1903, please discuss).

Answer: The section was corrected to "...due to increased contribution of nitrification or fungal denitrification, ...".

P1593 L18 please show linearity of Keeling plots

Answer: Keeling plots for each single day are provided as supplementary file S1, and a corresponding reference to the supplementary material was placed in the text.

P 1994 L 6-9 not clear to me why the footprints are different, please explain

Answer: An explanation was added in section 4.6: "Alternatively, the variation in isotope values associated with small overnight concentration increase may result from other land use or land cover. The EC fluxes are calculated from the turbulent fluctuation of concentration and vertical wind speed (i.e. the covariance of the concentration and wind speed deviations from the half-hourly mean) and therefore account for the modulation of concentration around a short term (30 min) mean caused by locally emitted N₂O. Isotopic composition based on Keeling plots, however, is determined from total N₂O accumulated in the nocturnal boundary layer and, thus, this approach also contains molecules that had been emitted outside the flux footprint, which almost exclusively comprised our grassland site (Zeeman et al. 2010), within the larger concentration footprint (Griffis et al., 2007)."

Section 4.7 pleas discuss cumulated N2o fluxes of this site in comparison to other grassland sites in order to interpret to which extent flux weighted isotope values can be seen representative. In view of high mean fluxes it seems to me that this site is a real hot spot. . (rough look at Fig 3 suggests that flux level is at least higher tha 5 kg N/ha and year). This needs to be taken into account in the discussion here or at least addressed somewhere.

Answer: We address this issue at the end of section 4.6: "Secondly, the CHA grassland can be characterized as a site with vigorous N₂O emission and therefore may dominate the determined N₂O isotopic composition as the influence of a source area increment scales with the source strength. The grassland was restored in 2012 which lead to extraordinary high N₂O-N emission of 29.1 kg ha⁻¹ year⁻¹ (Merbold et al 2014). In the following year, 2.5 kg N ha⁻¹ were released. This value is still in the range of maximum emissions reported for another intensively managed Swiss grassland, emitting 1.5-2.6 kg N ha⁻¹ year⁻¹ and at least a factor of five compared to an extensively managed grassland with less than 0.5 kg N ha⁻¹ year⁻¹ (Ammann et al 2009). With regard to distant land use and land cover, the 2.5 kg N ha⁻¹ are also more than double the median of all reported values for cultivated temperate sites and higher than the maximum value reported for forests presented in a study containing 1008 N₂O emission measurements from agricultural fields (Stehfest and Bouwman, 2006)".

P 1595 L 5 and elsewhere: add fungal nitrification since it is not distinguishable from NH2OH oxidation. Answer: fungal <u>denitrification was added</u>.

Referee#2:

The manuscript reports a technical feat: the on-line isotopic characterisation of N2O emitted from an agricultural area. All descriptions, data and discussion relating to the isotopic characterisation are excellent science. A weak point is the relation of isotopic compositions to N2O flux (and soil parameters) measured on the experimental grassland plot. There are probably something like four orders of magnitude in size difference between the concentration footprint (in the order of 10 x 10 km; from which isotopic compositions were derived) and the N2O flux footprint (in the order of 0.1×0.1 km).

By relating changes in isotopic composition to N2O flux (and soil conditions) on the experimental grassland plot, the implicit assumption is made that N2O emitted from the grassland plot is representative, in terms of isotopic composition, for a much larger area. Looking at aerial photographs of Chamau (e.g. Google Earth), it seems there is a large proportion of arable crops and also forest within the concentration footprint (I am not familiar with this site, but think to have located it at 47 degrees 12' and 24" N and 8 degrees, 24' and 32" E). This mix of different landuse constitutes the concentration footprint and is the source of observed changes in the isotopic composition observed during nocturnal inversions. In contrast, the N2O flux measured by eddy covariance relates to the grassland site, where also the soil parameters (soil temperature and moisture, inorganic N) were measured. I would propose to drop the N2O flux part of the manuscript and relate observed changes in isotopic composition during nocturnal inversions solely to meteorological parameters ("wet phase" and "dry phase", as in section 3.5), which are much more likely to have been homogenous within the concentration footprint, than N2O flux or soil parameters (in particular NH4+, NO3-, DOC,) or management events.

Answer: We agree with referee#2 that there is a remarkable difference in size between flux and concentration footprint and therefore the concentration footprint associated with our measurements of N_2O isotopic composition certainly comprises adjacent areas differing in land use or land cover. In summary, however, we are confident, that the grassland represents a major contribution to the analyzed isotopic signature of the N_2O accumulated above the grassland in the nocturnal boundary layer. To better illustrate our argumentation, but also to present potential limitations, we added the following changes to the manuscript:

- 1) The manuscript title was changed from "First on-line isotopic characterization of N₂O emitted from intensively managed grassland" to "First on-line isotopic characterization of N₂O above intensively managed grassland".
- 2) As given above in response to referee#1 (see comment to P 1994 L 6-9) an explanation was added why flux and concentration footprint are different and how this could influence N₂O isotope analysis. In addition we added the following wording to the end of section 4.6: "However, it cannot be excluded that N₂O isotopic signatures analyzed above the grassland were influenced by adjacent ecosystems". As well as section 4.7: "One has to keep in mind, however, that part of the observed variability may be attributed to the fact that the footprint area of the N₂O isotopic composition includes areas with other land use or land cover".
- 3) As detailed above in response to referee#1 (see comment to section 4.7) the weight of influence scales with source strength and distance so that areas of high source strength and low distance to the sample inlet have a higher influence on the determined isotopic composition. The grassland on which our measurements were carried out was restored in 2012 which caused extraordinary high N₂O emission in 2012 (29.1 kg ha⁻¹). Emissions during the measurement campaign were also distinctly elevated with up to 500 µg m⁻² h⁻¹ and the grassland site CHA can be considered as a site with vigorous N₂O emission. In the study by Griffis et al. (2007) the shift of source signature was calculated for different concentration footprint sizes, assuming an increasing area of contributing C3 canopy. However, Griffis et al. postulated a constant source strength for C3 and C4 canopy, which is certainly not the case for CHA.
- 4) In addition, the linearity of the Keeling plots, given in the supplementary file S1, indicates a constant source process. During the overnight concentration increase, which was used for the Keeling plots, wind speed and direction were not constant over the several, approx. 16 minute

intervals during which surface layer air was pre-concentrated. Consequently, influences from other grasslands, land use or land cover with different isotopic composition would result in Keeling plots that show a deviation from a linear relation of isotopic composition with 1/concentration. This is especially the case for d15Nbulk as this value is most variable due to its dependence on precursor composition which can be expected to vary significantly in space. To highlight this, the following phrase was added to the text: "However, two facts indicate a major influence of the studied grassland on the determined N_2O isotopic composition: First, the N_2O isotopic composition is very stable for a noon-to-noon period as indicated by a linear relationship between individual measurements (supplementary file S1). This relationship persists even though wind speed and direction are changing and, therefore, individual N_2O isotope measurements integrating over 16 minutes sampling interval originate from different source areas."

Owing to the considerable source strength in combination with the strictly linear Keeling plots (given as supplementary material), we assume that the grassland studied has a major influence on the determined isotopic composition. Nonetheless, we clearly give credit to the possibility that adjacent regions may have influenced the determined N₂O isotopic composition.

Page 1575, lines 13-15 state: "Hence, the development of adequate mitigation strategies is pertinent and requires a better understanding of the processes driving N2O fluxes."

Please return in your discussion to this statement and try to show how the study has contributed to this goal (maybe as a follow-up to sections 4.4 and 4.5).

Answer: As stated at the end of section 4.4, a better understanding of processes driving N₂O emission cannot be reached to date by determination of N2O isotopic composition alone, but will need to be combined with other methods determining isotopic composition in the substrates and quantification of N₂ emission.

Minor:

Page 1576, line 22: insert space between "in" and "Toyoda".

Answer: done

Page 1579, line18, and page 1594, line 22: maybe "comparability" instead of "compatability"? Answer: The term "compatibility" is used to refer to the agreement between results from different laboratories, in accordance with the vocabulary for metrology (see GAW report No. 213, 2013).

Page 1584, line 22: Results of DOC measurements are presented here, without the DOC measurements having been explained in the Methods section.

Answer: Section 2.7 was supplemented by a corresponding paragraph

1	First on-line isotopic characterization of N_2O emitted from above intensively managed
2	grassland
3	
4	Running head: Real-time grassland N2O isotopic signature
5	
6	^{1,2} Benjamin Wolf, ³ Lutz Merbold, ³ Charlotte Decock, ¹ Béla Tuzson, ¹ Eliza Harris, ³ Johan Six,
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17	Keywords: N ₂ O, isotopomers, QCLAS, source partitioning, grassland, nitrification,
18	denitrification, N ₂ O reduction
19	Paper type: primary research

21 Abstract

The analysis of the four main isotopic N₂O species (${}^{14}N^{14}N^{16}O$, ${}^{14}N^{15}N^{16}O$, ${}^{15}N^{14}N^{16}O$, ${}^{14}N^{14}N^{18}O$) and especially the intramolecular distribution of ${}^{15}N$ (site preference, SP) has been suggested as a tool to distinguish source processes and to help constrain the global N₂O budget. However, current studies suffer from limited spatial and temporal resolution capabilities due to the combination of discrete flask sampling with subsequent laboratory-based mass spectrometric analysis. Quantum cascade laser absorption spectroscopy (QCLAS) allows selective high-precision analysis of N₂O isotopic species at trace levels and is suitable for in-situ measurements.

Here, we present results from the first field campaign, conducted on an intensively managed 29 30 grassland in central Switzerland. N₂O mole fractions and isotopic composition were determined in 31 the atmospheric surface layer (2.2 m height) at high temporal resolution with a modified state-ofthe-art laser spectrometer connected to an automated N_2O preconcentration unit. The analytical 32 33 performance was determined from repeated measurements of a compressed air tank and resulted in measurement repeatability of 0.20, 0.12 and 0.11% for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$ and $\delta^{18}O$, respectively. 34 35 Simultaneous eddy-covariance N₂O flux measurements were used to determine the flux-averaged isotopic signature of soil-emitted N₂O. 36

Our measurements indicate that in general, nitrifier-denitrification and denitrification were the prevalent sources of N₂O during the campaign, and that variations in isotopic composition were rather due to alterations in the extent to which N₂O was reduced to N₂, than other pathways such as hydroxylamine oxidation. Management and rewetting events were characterized by low values of the intra-molecular ¹⁵N site preference (SP), $\delta^{15}N^{bulk}$ and $\delta^{18}O$, suggesting nitrifier denitrification and incomplete heterotrophic bacterial denitrification responded most strongly to the induced disturbances. Flux-averaged isotopic composition of N₂O from intensively managed grassland was 6.9 ± 4.3, - 17.4 ± 6.2 and 27.4 ± 3.6 ‰ for SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$, respectively. The approach presented here is capable of providing long-term datasets also for other N₂O emitting ecosystems, which can be used to further constrain global N₂O inventories.

48 1 Introduction

Atmospheric nitrous oxide (N₂O) mole fraction is increasing since pre-industrial times 49 50 predominately due to increased agricultural activity (Davidson, 2009; Mosier et al., 1998). Owing to the approximately 300 times higher global warming potential (GWP) compared to CO₂, this 51 52 greenhouse gas (GHG) currently accounts for 6% of total anthropogenic radiative forcing (Myhre 53 et al., 2013). Recent estimates showed that N_2O is in addition the single most important ozone-54 depleting substance (Ravishankara et al., 2009). Because at least 60% of total anthropogenic N_2O 55 emissions is attributed to food production (Syakila and Kroeze, 2011), growing human 56 population and meat consumption per capita as well as biofuel production will accelerate the rate of increase in atmospheric N₂O concentration. Hence, the development of adequate mitigation 57 strategies is pertinent and requires a better understanding of the processes driving N_2O fluxes. To 58 59 date, nitrification, nitrifier denitrification and denitrification are considered to constitute the dominant N₂O producing processes, especially in agricultural soils (Wrage et al., 2001). Other 60 61 N_2O source-processes such as abiotic N_2O production, co-denitrification and heterotrophic 62 nitrification have also been observed; a concise overview of observed processes is given 63 elsewhere (Butterbach-Bahl et al., 2013). This complexity inherent in the N cycle and associated transformation processes is a major challenge in developing mitigation strategies, as attribution of 64 65 N_2O production to the respective processes is required to tailor target-oriented actions (Baggs, 66 2008). Approaches for apportioning of N₂O emissions to nitrification, denitrification, and N₂O reduction to N_2 (source partitioning) have mostly relied on acetylene (C_2H_2) inhibition and 67 isotope labeling (Groffman et al., 2006), but denitrification rates are underestimated by the C₂H₂ 68 method (Butterbach-Bahl et al., 2013; Groffman et al., 2006; Watts and Seitzinger, 2000). Isotope 69 labeling approaches are vulnerable to incomplete diffusion of the tracer and to stimulation of 70 71 process rates by the addition of the labeled substrates themselves (Groffman et al., 2006).

72	Changes in natural abundance of 15 N and 18 O in N ₂ O have been explored to investigate N ₂ O
73	production processes, but the determined $\delta^{15}N$ and $\delta^{18}O$ depend on both fractionation factors and
74	isotopic composition of precursors, which in turn exhibit strong variations (Baggs, 2008; Bedard-
75	Haughn et al., 2003; Heil et al., 2014; Toyoda et al., 2011).
76	N ₂ O is a linear molecule and four main isotopic species can be discerned: ¹⁴ N ¹⁴ N ¹⁶ O, ¹⁴ N ¹⁵ N ¹⁶ O,
77	${}^{15}N^{14}N^{16}O$ and ${}^{14}N^{14}N^{18}O$. The isotopic species ${}^{14}N^{14}N^{16}O$, ${}^{14}N^{14}N^{18}O$ and ${}^{14}N^{15}N^{16}O$ (or
78	${}^{15}N^{14}N^{16}O$) are isotopologues, while ${}^{14}N^{15}N^{16}O$ and ${}^{15}N^{14}N^{16}O$ are isotopomers and will be
79	termed ${}^{15}N^{\alpha}$ -N ₂ O and ${}^{15}N^{\beta}$ -N ₂ O (Toyoda and Yoshida, 1999). The umbrella term isotopocule is
80	used for both isotopomers and isotopologues. The intra-molecular distribution of ^{15}N in N_2O
81	('site preference'; SP = $\delta^{15}N^{\alpha} - \delta^{15}N^{\beta}$) has been reported to be independent of the substrate's
82	isotopic composition, as SP in the de novo produced N ₂ O remained constant even though $\delta^{15}N$
83	and $\delta^{18}O$ values of both N ₂ O and substrates changed markedly during experiments with pure
84	cultures (Heil et al., 2014; Sutka et al., 2003, 2006, 2008; Toyoda et al., 2005). Therefore, SP can
85	be considered as a tracer conserving the source process information (Ostrom and Ostrom, 2011).
86	The SP of different processes has been characterized in a number of pure-culture, mixed culture
87	(Ostrom et al., 2007; Sutka et al., 2003, 2006; Toyoda et al., 2005; Wunderlin et al., 2012, 2013),
88	and soil-incubation studies (Köster et al., 2011, 2013a; Lewicka-Szczebak et al., 2014; Well et
89	al., 2006, 2008) with a compilation of data in Toyoda et al. (2011). A recent review on source
90	partitioning and SP (Decock and Six, 2013b) concluded that SP is capable of distinguishing
91	between the process groups N_2O_N (NH ₂ OH-oxidation, fungal denitrification and abiotic N_2O
92	production; SP = 32.8 ± 4.0 ‰) and N ₂ O _D (nitrifier-denitrification and denitrification; SP = -
93	1.6 ± 3.8 ‰). In addition, the intramolecular distribution of ${}^{15}NN_2O$ isotopocules can be used as
94	an independent validation of the global, measurement-based bottom-up N_2O budget and has

already confirmed that the isotopically light sources such as agriculture and industry contribute to 95 96 the increase in atmospheric N₂O (Toyoda et al., 2013; Yoshida and Toyoda, 2000). Owing to the temporal and spatial variability of isotopomer ratios, it is indispensable to derive flux-weighted 97 average values from different sources (such as ecosystems) for later use in budget analysis using 98 99 box models (Kim and Craig, 1993; Perez et al., 2001; Yoshida and Toyoda, 2000). Intramolecular distrubution of ¹⁵N in N₂ON₂O isotopomers can be measured by mass 100 101 spectrometry, but it requires discrete flask sampling with subsequent laboratory analysis. Hence, 102 this approach is limited in temporal resolution and spatial resolution representation of a given site. 103 Additionally it is indirect, as information on the site-specific isotopic composition is derived from 104 the analysis of the NO⁺ fragment and N_2O^+ molecular ion. Recently, a quantum cascade laser 105 absorption spectrometer (QCLAS) capable of selective analysis of the three most abundant N2O 106 isotopocules has been presented (Waechter et al., 2008) and its potential for in-situ measurements in conjunction with an automated pre-concentration unit has been shown (Mohn et al., 2010, 107 2012). Here we present the results obtained from a, to our knowledge worldwide first, campaign 108 in which the isotopic composition of N₂O (SP, δ^{15} N, δ^{18} O) in the atmospheric surface layer was 109 determined on-line by using an optimized state-of-the-art laser spectrometer. With the 110

111 combination of N_2O isotopic analysis by QCLAS, accompanying eddy-covariance based N_2O

112 flux measurements as well as monitoring of environmental conditions and inorganic nitrogen

113 concentrations, our specific objectives for this study were: i) to demonstrate the capability of

114 QCLAS systems for high precision isotopic analysis of (soil emitted) N₂O in ambient air; ii) to

investigate management and weather effects on isotopic composition and source processes; iii) to

116 test the capability of the N_2O isotopic composition for source partitioning; and ivand iii) to

- 117 characterize the flux-averaged isotopic composition of N₂O emitted from an intensively managed
- 118 grassland.

119 2 Material and Methods

120 2.1 Study site

The agricultural research station Chamau (CHA) is located in Central Switzerland at an elevation 121 122 of 400 m a.s.l.. The experiment was conducted on an intensively managed grassland belonging to CHA which is primarily used for fodder production and occasional winter grazing by sheep 123 (Zeeman et al., 2010). The soil type is a cambisol with a <u>bulk density of 0.97g cm⁻³, 30.6 % sand</u>, 124 125 47.7 % silt and 21.8 % clay in the top 10 cm and pH of 5.7-6.2. Soil carbon and nitrogen content in the top 10 cm was 37.9 g kg⁻¹ and 4.1 g kg⁻¹ (Roth, 2006). Mean annual temperature and 126 annual precipitation are 9.1°C and 1151 mm, respectively (Zeeman et al., 2010). Management 127 128 practices aim at fodder production and consist of mowing followed by slurry application, with up to six mowing/slurry applications per year and occasional grazing of sheep and cattle in October 129 and November. During the campaign in summer 2013, three management cycles were carried 130 out. Harvest dates were June 6th, July 11th and August 21st and slurry was applied within 10 days 131 132 after each mowing event. Nitrogen input was calculated from the applied amount of slurry 133 brought to the field and the N concentration determined (Labor für Boden- und Umweltanalytik, Eric Schweizer AG, Thun, Switzerland) in a sample drawn from the supply to the trailing hose 134 applicator. The applied N amounted to 30, 40 and 43.3 kg N ha⁻¹ for the first, second and third 135 136 application, respectively. The grassland is re-established via ploughing and resowing approximately every 10 years. The last re-establishment event took place in 2012 (Merbold et al., 137 2014). 138

139 2.2 Instrumental setup for analysis of N_2O isotopocule ratios

140 The four most abundant N₂O isotopic species were quantified using a modified QCLAS

141 (Aerodyne Research Inc., Billerica MA, USA) equipped with a continuous wave quantum

142 cascade laser (cw-QCL) with spectral emission at 2203 cm⁻¹, an astigmatic Herriott multi-pass

143	absorption cell (204 m path length, AMAC-200), and reference path with a short (5 cm) N_2O -
144	filled cell to lock the laser emission frequency (Tuzson et al., 2013). During the campaign, the
145	QCLAS was operated in an air-conditioned trailer located 60 m west of the eddy-covariance (EC)
146	tower. This trailer position contributes < 20 % to the main flux and is at the far side of prevailing
147	wind direction (Zeeman et al., 2010). The sample air inlet was installed next to the inlet of the EC
148	tower (2.2 m height). Sample air was drawn through a PTFE tube (4 mm ID) by a membrane
149	pump (PM 25032-022, KNF Neuberger, Switzerland). Upstream of the pump, the sample air was
150	pre-dried with a permeation drier (MD-050-72S-1, PermaPure Inc., USA). Following the pump,
151	the pressure was maintained at 4 bar overpressure using a pressure relieve valve. Humidity, as
152	well as CO ₂ , were quantitatively removed from the gas flow by applying a chemical trap filled
153	with Ascarite (7 g, 10 – 35 mesh, Fluka, Switzerland) bracketed by Mg(ClO ₄) ₂ (2 x 1.5 g, Fluka,
154	Switzerland). Finally, the sample gas was passed through a sintered metal filter (SS-6F-MM-2,
155	Swagelok, USA) and directed to a preconcentration unit described in detail previously (Mohn et
156	al., 2010, 2012). For an increase of N ₂ O mixing ratios from ambient level to around 50 ppm N ₂ O,
157	approx. 8 litres of ambient air were preconcentrated. Afterwards, the preconcentrated N_2O was
158	introduced into the evacuated multi-pass cell of the QCLAS. Isotopic fractionation during
159	preconcentration (increase by 0.31 ± 0.10 , 0.34 ± 0.16 and 0.29 ± 0.07 ‰ for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$ and
 160	δ^{18} O, respectively) was quantified by preconcentration of N ₂ O with a known isotopic
161	composition and subsequently corrected. Compatibility of N2O isotopomer analysis by QCLAS
162	with isotope ratio mass spectrometry (IRMS) laboratories was recently demonstrated in an inter-
163	laboratory comparison campaign (Mohn et al., 2014).
164	2.2 Macaurant and calibration starts are

164 2.3 Measurement and calibration strategy

165 To ensure high accuracy and repeatability of the analytical system, a measurement and

166 calibration strategy similar to the one presented by Mohn et al. (2012) was applied. It is based on

167	two standard gases differing in N2O isotopic composition, which were produced by dynamic
168	dilution of pure medical N ₂ O (Pangas, Switzerland) with defined amounts of isotopically pure
169	(>98 %) 14 N 15 N 16 O (Cambridge Isotope Laboratories, USA) and (>99.95 %) 14 N 14 NO (ICON
170	Services Inc., USA). Subsequent gravimetric dilution with high purity synthetic air (99.999 %,
171	Messer Schweiz AG) resulted in pressurized gas mixtures with 90 ppm N ₂ O (parts per million,
172	10 ⁻⁶ moles of trace gas per mole of dry air). Both standards were calibrated against primary
173	standards which were previously measured by the Tokyo Institute of Technology (TIT, Toyoda
174	and Yoshida) to anchor δ -values to the international isotopic standard scales. The first standard
175	(S1, Table 1) was used as an anchor point to international δ -scale and used as input data for data
176	analysis algorithms (see data processing). Therefore, the N_2O isotopic composition of S1 was
177	targeted to closely resemble background air. As the N2O isotopic composition of surface layer air
178	is mainly a mixture of soil-derived and background composition, the second standard (S2, Table
179	1) used for span correction was depleted in $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$ and $\delta^{18}O$ compared to background air in
180	accordance with the expected terrestrial source signatures.
181	The measurement protocol started with the injection of S1, dynamically diluted to 50 ppm, the
182	mole fraction of ambient N_2O after preconcentration. After flushing the absorption cell with
183	synthetic air, S2 was injected, also diluted to 50 ppm. For determination of the slight
184	concentration dependence already reported (Mohn et al., 2012), S1 was injected again but at a
185	higher mole fraction of 67 ppm (later referred to as $S1_h$). This mole fraction represents the mole
186	fraction expected after preconcentration of high concentration surface layer air. Subsequently, S1
187	was injected again, diluted to 50 ppm, before the cell was filled with preconcentrated ambient
188	N_2O (A). This subroutine (S1+A) of injection of S1 and preconcentrated ambient N_2O took 35
189	minutes and was repeated three times. For an independent determination of repeatability, the

190 fourth sample was preconcentrated compressed air (target gas). During the campaign, two 191 compressed air cylinders (C1 and C2, referred to as target gas) were used. Isotopic composition and N₂O mixing ratio of both cylinders were determined in the laboratory prior to campaign start 192 (Table 1). N_2O mole fractions and isotopic composition analysed in the laboratory and at the field 193 194 site agreed within their analytical uncertainty. Following target gas analysis, S1 and S1_h were analyzed again. Another set of three subroutines S1+A completed one run. One complete cycle of 195 6 ambient air samples and one compressed air sample took 340 minutes, leading to approx. 25 196 ambient air samples being analysed during 24 hours. N₂O mole fractions were determined 197 198 according to Mohn et al. (2012).

199 2.4 Data processing

200 Data processing is based on individual mixing ratios of the four main N₂O isotopic species and spectrometer characteristics as recorded by the instruments's software (TDLWintel, Aerodyne 201 202 Research Inc., Billerica, MA, USA). In the first step, variations in the isotope ratios induced by drifts in the instrument working parameters during the field operation were corrected. A linear 203 additive model explaining the deviation of isotope ratios R^{α} , R^{β} and R^{180} for repeated 204 205 measurements of standard S1 from their mean value by absorption cell temperature (T1), laser 206 temperature (T2), line position (LP) and pressure (p) was calibrated based on S1 injections. For isotope ratios of $S1, S1_{h}, S2$, sample air and compressed air, these systematic deviations were 207 208 corrected based on the respective values of T1, T2, LP and p. In a second step, concentration dependence of isotope ratios, determined using the measurements of S1 and S1_h, was addressed 209 with corrections (0.013, 0.028 and 0.004 % ppb⁻¹ for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$ and $\delta^{18}O$) being in the same 210 211 range as described earlier (Mohn et al., 2012). Subsequently, remaining drifts were corrected based on analysis of S1. Finally, isotope ratios were converted to δ -values using a 2-point 212 calibration derived from corrected values of S1 and S2. 213

214 2.5 Determination of soil-emitted N₂O isotopic composition

Isotopic composition of the source process "soil N₂O emission" was derived using the Keeling 215 plot approach (Keeling, 1958), where δ -values measured (here in 2.2 m height) are plotted versus 216 the inverse of N₂O mole fractions. The intercept of the linear regression line can be interpreted as 217 the isotopic composition of soil emitted N_2O (Pataki et al., 2003). Therefore, determination of 218 soil N₂O isotopic composition requires an increase in N₂O mole fraction. During the day, 219 220 turbulence mixes surface layer air to the atmospheric background. At night, the surface layer becomes more stable and the N₂O mole fraction increases, shifting isotopic composition towards 221 its source composition. As a consequence, Keeling plots were based on noon-to-noon periods. 222 This approach is discussed in section 4.6. 223

224 2.6 N_2O Flux measurement

At CHA, greenhouse gas mole fractions, including N₂O, are measured continuously since 2012 225 by means of the eddy covariance (EC) method (Baldocchi and Meyers, 1998). The system 226 227 consists of a three-dimensional sonic anemometer to measure wind speed and direction (2.41 m height, Solent R3, Gill Instruments, Lymington, UK) and a QCLAS (mini-QCLAS, Aerodyne 228 229 Research Inc., Billerica, MA, USA) to determine N₂O mole fractions at a temporal resolution of 230 10 Hz. Both data streams are merged near-real time within a data acquisition system (MOXA embedded Linux computer; Moxa, Brea, CA, USA) via an RS-232 serial data link (Eugster and 231 232 Plüss, 2010). The setup has been described in detail previously (Merbold et al., 2014). Postprocessing of N₂O fluxes included screening for obvious out-of-range values (+/- 100 nmol m⁻ 233 2 s⁻¹). N₂O fluxes were further aggregated to noon-to-noon daily averages to smoothen the large 234 variability in the 30 min flux averages. Daily averages were calculated for days where more than 235 236 30 half-hour values were available, with this filter excluding three days from analysis.

237	2.7	Soil inorganic	N,	dissolved	organic	<u>C</u>	and	environmental	conditions
		•							

Ammonium (NH₄⁺) and nitrate (NO₃⁻) concentrations were determined from soil (0-20 cm depth) 238 239 sampled at 10 positions along a transect within the footprint of the EC measurements following the predominant wind direction. Samples were taken weekly throughout the campaign or daily 240 241 during mowing and slurry application events. Per sample, ~ 15 g of fresh soil were added to 242 specimen vessels containing 50 ml 1M KCl. After 1 hour on a shaker, the supernatant was 243 filtered (Whatman no.42 ashless filter paper, 150 mm diameter) and analysed colorimetrically for NH_4^+ and NO_3^- . For a subset of extracts, we determined dissolved organic carbon (DOC) 244 concentrations by combustion of KCl extracts using a total organic C analyzer (Shimdazu TOC-245 246 V, Columbia, MD, USA). 247 Soil temperatures and volumetric soil moisture contents at 10 cm depth were measured at the 248 same 10 locations along the transect (5TM-sensors, Decagon Devices Ltd., Pullman, USA). Data were stored as 10 minute averages on a data logger (EM50, Decagon Devices Ltd., Pullman, 249 USA). The volumetric water content was converted to water filled pore space (wfps) using a bulk 250 251 density of 1.09 g cm⁻³. Precipitation was measured with a tipping bucket rain gauge (Type 10116, 252 Toss GmbH, Potsdam, Germany) and stored as 10 min averages on a data logger (CR10X-2M, Campbell Scientific Inc., Logan, USA). 253

256	3 Results
257	3.1 Long term precision for target gas analysis
258	System performance for N_2O mole fractions and isotopic composition was determined based on
259	repeated analysis of compressed air from target gas tanks (C1, C2). There was no significant drift
260	in the δ -values and N ₂ O mole fractions, indicating stability of the applied measurement
261	technique. Repeatability, calculated as the standard deviation (σ) of 331 target gas measurements,
262	amounted to 0.20, 0.12, 0.10, 0.12 and 0.22 ‰ for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$, $\delta^{18}O$, $\delta^{15}N^{bulk}$ and SP,
263	respectively (Figure 1). Standard deviation for the N_2O mole fraction of the target gas was
264	0.25 ppb.
265	3.2 N_2O mole fractions and isotopic composition at 2.2 m height
266	N2O isotopic composition of the surface layer (lowest tens of meters above ground)Air samples
267	were taken at 2.2 m height which is within the lowest 10% of the atmospheric boundary layer
268	(ABL) where mechanical generation of turbulence exceeds buoyant generation or consumption.
269	This part of the ABL is called surface layer, hence corresponding air samples are referred to as
270	surface layer air samples. N ₂ O isotopic composition of the surface layer air samples ($n = 2130$)
271	ranged from 2.5 to 16.1 ‰, -11.9 to -2.4 ‰, 37.6 to 44.6 ‰, -4.6 to 6.6 ‰, and 14.3 to 19.3 ‰
272	for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$, $\delta^{18}O$, $\delta^{15}N^{bulk}$ and SP, respectively (Figure 2). Surface layer N ₂ O mole fractions
273	varied between 325 and 469 ppb and followed a diurnal cycle with highest values during the
274	night when the boundary layer became more stable. Increasing N_2O mole fractions were
275	associated with decreasing δ -values, indicating that soil emitted N ₂ O that mixed into the surface
276	layer was depleted in ¹⁵ N as compared to N ₂ O in the atmospheric background.

277 3.3 Auxiliary measurements

Half hourly N₂O fluxes were averaged from noon-to-noon (f_{N2O}), and ranged from -1 to 5 nmol 278 $m^{\text{-}2} \, \text{s}^{\text{-}1}.$ Maximum N_2O fluxes coincided with an overnight build up in N_2O mole fractions 279 $(\Delta N_2 O)$ as analysed by QCLAS and could not be attributed to slurry application events alone 280 (Figure 3). Among the correlations of f_{N2O} and auxiliary variables, only the one with nitrate 281 concentration ($r^2 = 0.18$) was significant (p<0.01). Soil water content (wfps) was modulated by 282 283 precipitation and two clear states could be identified. During the "wet" part of the campaign lasting until July 7th, average wfps was with 62 ± 4 % significantly (t-test, p < 0.001) higher than 284 the average of 37 ± 4 % calculated for the remainder of the campaign (referred to as the "dry") 285 part). Soil temperature did not show such a clear two-phase pattern, however temperatures during 286 the first, "wet" part were with 16.7±4 °C significantly (p<0.001) lower than during the "dry" 287 phase with 21.2 ± 2 °C. 288 Background NH₄⁺ and NO₃⁻ concentrations were smaller than 3 μ g g_{soil}⁻¹ and clearly responded to 289 mowing and slurry application in the second and third management events. The NO_3^{-1} 290 concentration was higher than the NH_4^+ concentration and peaked at 16 and 50 µg g_{soil}^{-1} , while 291 NH_4^+ concentration peaked at 9 and 15 µg g_{soil}^{-1} for these two management events. In contrast, 292 dissolved organic carbon concentrations (DOC) did not respond to management events, but were 293 higher during the "dry" phase of the campaign (p < 0.001). 294

295 3.4 Isotopic composition of soil-emitted N₂O

296 The uncertainty of the determined source isotopic composition was estimated based on the

- standard error of the Keeling plot intercept and depends on the degree to which soil air
- 298 accumulated in the surface layer (ΔN_2O , Figure 4). For instance, the intercept (source) standard
- error ranged from 0.3 to 82 ‰ for SP. To apply the Keeling plot approach only to situations in

300 which soil air accumulated in the surface layer, only source isotopic compositions for overnight

301 increases in N₂O mole fractions of more than 12 ppb were considered in this study. This filter

lead to a maximum and average (μ) standard error of 6.8 (μ =2.2) ‰, 4.5 (μ =1.4) ‰ and 2.2

303 (μ =1) ‰ for SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$ isotopic source signatures, respectively.

304 During the field campaign, Keeling plot derived isotopic composition of soil-emitted N₂O ranged

from 1.4 to 17.3 ‰, -29 to -3 ‰ and 22.6 to 34.8 ‰ for SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$, respectively. All

explanatory variables except NH_4^+ and NO_3^- were found to significantly correlate with SP (Table

2). For δ^{15} N^{bulk}, correlations with Δ N₂O, wfps, soil temperature, DOC and NO₃⁻and for δ^{18} O

308 correlations of f_{N2O} , ΔN_2O , precipitation, soil temperature and NO_3^- were significant. However,

the adjusted r^2 for all regressions was below 0.4; in addition, multiple explanatory variables such

- as NH_4^+ and NO_3^- or wfps and temperature (Figure 5) did not increase the explained variance
- 311 above this value.

312 3.5 Event-based data aggregation

As already described in the section "Auxiliary measurements", there was a "wet" phase (n=27 313 Keeling-plot derived N₂O isotopic compositions) in the beginning of the campaign, which lasted 314 315 about one month and a "dry" phase lasting about two months (n=38). Therefore, the dataset was split in two corresponding parts with averages of 7.4 \pm 3.6 ‰ versus 11.1 \pm 4.2 ‰ for SP, -316 19 ± 3.8 % versus -12.5 \pm 5.9 % for $\delta^{15}N^{bulk}$ and 28.7 \pm 2.2 % versus 29.7 \pm 3.4 % for $\delta^{18}O$ in 317 the wet versus the dry phase, respectively. Averages of SP and $\delta^{15}N^{\text{bulk}}$ were significantly 318 different (p < 0.001) but δ^{18} O averages were not. Based on this simple classification, the dry 319 phase contains rewetting events. A rewetting event was defined as a two day period starting at the 320 day for which wfps increased. Exclusion of these rewetting events during the dry phase increased 321

average δ -values (n=30) as well as decreased standard deviations for SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$ to

 12.5 ± 3.4 , -10.8 ± 4.5 and 30.7 ± 2.8 ‰. Moreover the difference in δ^{18} O was significant (p < 0.001).

325	In addition to the dry/wet classification, we also defined three subsets representing the N_2O
326	emission associated with management events of mowing followed by fertilization ("Mana I" -
327	"Mana III"), one subset representing a rewetting event between Mana II and III ("Rewetting")
328	and one subset representing background ("BG", all remaining measurements). There were two
329	distinct rewetting events between management events II and III, but N2O isotopic composition is
330	only available for the first one (07/29/2014 - 07/31/2014). Isotopic compositions of soil-emitted
331	N ₂ O were assigned to subsets of management or rewetting if the associated flux or nutrient
332	concentration was elevated. This classification scheme led to 3 to 7 measurements for
333	management and rewetting events (Figure 3, underlaid in transparent blue) while 47 measurements
334	were assigned to class BG. Boxplots for SP, $\delta^{15}N^{bulk}$, $\delta^{18}O$, and wfps (Figure 6) showed
335	characteristic δ -values and wfps for management and rewetting, but not for subset BG.
336	Measurements assigned to BG covered practically the whole range of values observed across all
337	the other classes. Therefore, standard deviations for class BG were one order of magnitude larger
338	than for the four other classes.
339	Statistical analysis is confounded by low and unequal sample size so that we compared
340	exclusively the subsets management and rewetting using multiple non-parametric Wilcoxon tests
341	after having checked homogeneity of variances using Bartlett test. For all investigated δ -values,
342	only differences between groups Mana II and Mana III were significant.
343	3.6 Averages of N ₂ O isotopic signature for intensively managed grassland
344	Simple averages of daily isotopic composition of soil-emitted N ₂ O were 9.6 \pm 4.4, -15.2 \pm 6.0
345	and 29.3 \pm 3 ‰ for SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$, respectively (n=62). Representative isotopic signature

346	for agricultural landcomposition of N ₂ O emitted from a given site or treatment can be estimated
347	based on flux-weighted averages of daily signatures isotopic composition. For some noon-to-noon
348	periods included in the above average, thus with an overnight increase in N_2O mole fractions of
349	at least 12 ppb, negative N ₂ O fluxes were detected by the EC system (-0.17 \pm 2.1 nmol m ⁻² s ⁻¹ ;
350	n=14). This might be due to the uncertainty of N_2O flux measurements, temporal averaging over
351	positive and negative fluxes in a noon-to-noon period or different footprint regions for N_2O flux
352	and isotopic analysis (flux vs. concentration footprint). To avoid bias to the flux-weighted
353	average of emitted N_2O by either one of the above mentioned possible reasons, the weighted
354	averages were calculated for positive flux events only. Flux weighted averages were 6.9 ± 4.3 ,
355	- 17.4 \pm 6.2 and 27.4 \pm 3.6 ‰ for SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$ respectively (n=48).

357 4 Discussion

358 4.1 Analytical performance

To our knowledge, only two pilot studies exist demonstrating the potential of QCLAS based 359 360 analytical techniques for on-line and high-precision analysis of N₂O mole fractions and isotopic composition in surface layer air. While Mohn et al. (2012) analyzed the three most abundant ¹⁵N-361 isotopocules (¹⁴N¹⁴N¹⁶O, ¹⁵N¹⁴N¹⁶, ¹⁴N¹⁵N¹⁶O), Harris et al. (2014a) included the ¹⁸O 362 isotopologue (¹⁴N¹⁴N¹⁸O). In both studies, however, the instrument was located in the laboratory. 363 Based on three weeks of measurements, Mohn et al. (2012) reported a precision of 0.24 and 364 0.17 ‰ for $\delta^{15}N^{\alpha}$ and $\delta^{15}N^{\beta}$, respectively and Harris et al. (2014a) reported 0.17, 0.19 and 365 0.32 % for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$ and $\delta^{18}O$, respectively, for a twelve days period. In both studies, 366 analytical performance was determined, in accordance with the presented study, based on 367 368 repeated analysis of compressed air samples. Thereby, the analytical precision reached in the presented study, was distinctly higher for $\delta^{15}N^{\beta}$ and $\delta^{18}O$ and similar for $\delta^{15}N^{\alpha}$ compared to these 369 370 two previous studies, even though the measurements were done under field-conditions and over a 371 much longer, three months, period. This confirms the high level of precision associated with 372 QCLAS based determination of N₂O isotopic composition. Standard errors for Keeling plot 373 intercepts (Figure 4) confirm that this precision is sufficient to resolve the variability of atmospheric N₂O sampled close to the ground. As our instrument was located directly at the field 374 site and measurements were conducted over a period of more than three months, our study 375 376 indicates that this level of repeatability can be achieved both at long time scales and in the field. 4.2 N₂O isotopic composition in the atmospheric surface layer (2.2 m height) 377 In our study, δ -values of single preconcentrated air samples were between atmospheric 378 background and 14.3 ‰ (SP) and -4.7 ‰ ($\delta^{15}N^{bulk}$). Mohn et al. (2012) reported similar values 379

between atmospheric background and 12 ‰ (SP) and -4 ‰ ($\delta^{15}N^{bulk}$). Therefore the variation

381	observed in both studies is much higher compared to the measurements by Harris et al. (2014a)
382	where the N_2O isotopic composition deviated only slightly from atmospheric background. A
383	consistent decrease in $\delta^{15}N^{bulk}$ in parallel with increasing N ₂ O mole fractions (accumulation of
384	soil-derived N ₂ O) confirms that the soil N ₂ O source is depleted in 15 N-N ₂ O relative to ambient
385	N_2O (Toyoda et al., 2013). A similar pattern was found for $\delta^{18}O$; an increase in N_2O mole
386	fraction was associated with a decrease in ¹⁸ O-N ₂ O, again indicating that soil emissions were
387	depleted in ¹⁸ O-N ₂ O with respect to the atmospheric background. In contrast, Harris et al. (2014a)
388	reported a decoupling of δ^{18} O and δ^{15} N ^{bulk} . This may have been due to only marginal influence of
389	soil-emitted N ₂ O since the measurements were carried out in urban area and approx. 95 m above
390	the ground. Studies on N_2O derived from combustion processes indicate that some of these
391	sources might be less depleted or even enriched in 15 N-N ₂ O compared to ambient N ₂ O (Harris et
392	al., 2014b; Ogawa and Yoshida, 2005).

393 4.3 Isotopic composition of soil-emitted N₂O

394 SP of soil-emitted N₂O observed in our study (1 to 17 ‰) is within the ranges expected for a 395 mixture of the two process groups N₂O_N and N₂O_D, and does not necessarily indicate significant 396 contribution of N₂O reduction, an effect which is discussed further below. Isotopic composition

397 of soil-emitted N₂O has been predominately determined in laboratory incubation studies (Köster

et al., 2013a, 2013b; Perez et al., 2006; Well and Flessa, 2009b; Well et al., 2006, 2008).

Additionally, results from field experiments using static chambers (Opdyke et al., 2009; Ostrom

400 et al., 2010; Toyoda et al., 2011; Yamulki et al., 2001) and N₂O accumulation below a snowpack

- 401 have been published (Mohn et al., 2013). Based on pure culture studies SP values from 19.7 to
- 402 40 % and -8.7 to 8.5 %, were observed for N₂O_N and N₂O_D, respectively (Decock and Six,
- 403 2013b). In field experiments SP was found to range between -1 and 32 ‰ (Opdyke et al., 2009), -
- 404 3 and 18 ‰ (Yamulki et al., 2001), -14 and 90 ‰ (Toyoda et al., 2011) and 0 and 13 ‰ (Ostrom

et al., 2010). The very high SP values detected by Toyoda et al. (2011) may have resulted from 405 extensive N₂O reduction to N₂, a process increasing SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$ (Ostrom et al., 2007). 406 For δ^{15} N^{bulk} and δ^{18} O, a much wider variation as compared to SP is expected, because these 407 variables depend both on fractionation factors, which vary among different microbial 408 409 communities and depend on reaction conditions, as well as on the isotopic composition of the substrate (Baggs, 2008). Under field conditions, $\delta^{15}N^{\text{bulk}}$ was reported to range between -17 and 410 9 ‰ (Opdyke et al., 2009), -27 and 1 ‰ (Yamulki et al., 2001), -44 and 34 ‰ (Toyoda et al., 411 2011) and -18 and -15 ‰ (Ostrom et al., 2010), covering the range of -29 to -3 ‰ observed in 412 this study. With respect to δ^{18} O, the values of 22.6 to 34.8 % detected for grassland in this study 413 are at the lower end of measurements under field conditions (4-82 ‰). 414 415 4.4 Changes in N₂O source signatures induced by N₂O reduction to N₂. 416 Quantitative source partitioning between process groups N₂O_N and N₂O_D based on SP is possible only when no other processes except those contained in the process groups have an influence on 417 418 the site-specific N_2O isotopic composition. However, in the terminal step of denitrification, 419 namely the reduction of N_2O to N_2 , where N_2O is the substrate, the N-O bonds between lighter isotopic species is consumed isotopes are cleaved preferentially, leading to an increase in SP, 420 δ^{15} N^{bulk} and δ^{18} O in the remaining N₂O. Consequently, part of the N₂O originating from a 421 422 combination of the two process groups, i.e. N_2O_N and N_2O_D , may have been consumed by N_2O to N₂ reduction prior to emission. 423 For identification of processes determining N_2O isotopic composition, isotopocule maps were 424

425 suggested in which site preference is plotted versus the difference in substrate and product

426 isotopic composition (Koba et al., 2009). Determination of isotopic composition in the substrates

427 is time consuming and additionally confounded in our study by the large and varying footprint

428	area. Therefore, we present a modified isotope map of SP versus $\delta^{15}N^{bulk}$ (Figure 7, left panel)
429	instead of $\Delta \delta^{15}$ N, the δ^{15} N differences between substrate and product (i.e. N ₂ O gas). Rectangles
430	for process groups N_2O_N and N_2O_D are defined by SP values given by Decock and Six (2013b)
431	and by $\delta^{15}N^{\text{bulk}}$ values calculated based on process fractionation factors and substrate isotopic
432	composition. For nitrification and denitrification minimum and maximum fractionation factors of
433	-90 to -40 ‰ and -40 to -15 ‰ were assumed (Baggs, 2008), for the isotopic compositions of the
434	N_2O precursors (i.e., NH_4^+ and NO_3^-) a range of -20 to +10 ‰ and -25 to 15 ‰ were assumed.
435	Koba et al. (2009) attributed a concurrent decrease in $\delta^{15}N^{bulk}$ with increasing SP values as
436	indicative for an increasing contribution of N_2O_N . In contrast, an increase in $\delta^{15}N^{bulk}$ in parallel to
437	increasing SP values (enrichment of 15 N in the α -position relative to the β -position), as observed
438	in the present study, was allocated to a substantial increase in N_2O reduction to N_2 . For
439	ε ¹⁵ N ^{bulk} /εSP of N ₂ O reduction, Koba et al. (2009) assumed a factor of 1.2 based on previous
440	publications. Our results (Our results (Figure 7) indicate that N2O is predominately formed by
441	denitrification, and that deviations in the isotope values from denitrification may have been
442	caused by variations in the extent to which N ₂ O was reduced to N ₂ , left panel) indicate that N ₂ O
443	is predominately formed by bacterial denitrification, and that deviations in the isotope values
444	from denitrification may have been caused by variations in the extent to which N ₂ O was reduced
445	to N ₂ Additionally, δ^{18} O was found to be positively correlated with δ^{15} N ^{bulk} , which enforces the
446	interpretation that varying shares of N_2O reduction occurred because it acts on both N and O
447	isotopic composition (Koehler et al., 2012). It is noteworthy that based on such modified isotope
448	maps, systematic changes in $\delta^{15}N^{bulk}$ induced by systematic changes in N isotopic composition of
449	one of the precursors NH_4^+ or NO_3^- could be misinterpreted as reduction events- <u>(Well et al.</u> ,
450	<u>2012).</u>
I	

451	The ratios of fractionation factors for $\delta^{18}O$ and $\delta^{15}N^{bulk}$ (r_{o-n}) and SP and $\delta^{18}O$ (r_{sp-o}) during N_2O
452	reduction were suggested for estimation of the share of N_2O reduction to N_2 since these ratios
453	were found to be 2.5 and 0.2 to 0.5, respectively in laboratory incubation experiments In addition
454	to the SP/ δ^{15} N ^{bulk} maps, SP/ δ^{18} O maps have been suggested to trace N ₂ O reduction to N ₂
455	(Lewicka-Szczebak et al., 2014, 2015; Well et al., 2012). While $\delta^{15}N^{\text{bulk}}$ depends on the isotopic
456	composition of the precursor (e.g. NO ₃ ⁻) and, thus, may vary considerably, δ^{18} O-N ₂ O is expected
457	to be more stable as during both nitrification and denitrification, oxygen (O) later found in N_2O
458	may almost completely originate from water (Kool et al., 2009). Due to this almost complete O-
459	exchange with water, relatively stable δ^{18} O in soil water, and the observed constant ratio of
460	fractionation factors for SP and δ^{18} O-N ₂ O (r_{sp-o}), variation in the share of N ₂ O reduced to N ₂
461	should be reflected by a linear relationship between SP and δ^{18} O-N ₂ O with a slope of 0.2-0.5
462	(Jinuntuya-Nortman et al., 2008; Ostrom et al., 2007; Well and Flessa, 2009a). In this study, a
463	linear relationship with a slope of 1.02 was found (Figure 7, right panel). Tracking the
464	management events (ManaI to ManaIII) and the rewetting event in SP/ δ^{18} O space revealed that
465	the onset of such an event is associated with a decrease of both SP and δ^{18} O, gradually increasing
466	back to approximately initial values, except for ManaII. During ManaII, no significant change in
467	<u>SP/δ^{18}O occurred (Figure 7, right panel, red trace). The gradual increase in isotopic composition</u>
468	supports the conclusion from the SP/ $\delta^{15}N^{bulk}$ map that N ₂ O was mainly produced by bacterial
469	denitrification and that variations in isotopic composition may have been caused predominately
470	by N_2O reduction to N_2 . This interpretation is in agreement with observations of isotopoic
471	composition of N ₂ O, NO ₃ ⁻ and NH ₄ ⁺ during a rewetting event in an agricultural field (Decock and
472	Six, 2013a). Additionally, δ^{18} O was found to be positively correlated with δ^{15} N ^{bulk} , which

473 enforces the interpretation that varying shares of N₂O reduction occurred because it acts on both 474 N and O isotopic composition (Koehler et al., 2012). We calculated these ratios As introduced above, the ratios of fractionation factors for δ^{18} O and δ^{15} N^{bulk} (r_{o-n}) and SP and 475 δ^{18} O (r_{sp-0}) during N₂O reduction were 2.5 and 0.2 to 0.5 in laboratory incubation experiments 476 (Jinuntuya-Nortman et al., 2008; Ostrom et al., 2007; Well and Flessa, 2009a). In our study, ro-n 477 and r_{sp-o} were 0.5 and 1, respectively for the whole dataset. We calculated these ratios also for a 478 subset of data for which all δ -values (SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$) increased for two consecutive days, 479 480 indicating that N₂O reduction may have occurred. Such events were observed on 8 occasions. If 481 source processes (N_2O_D, N_2O_N) contributed constantly over two consecutive measuring days, changes in the isotopic composition of emitted N₂O were solely attributed to changes in the 482 483 fraction of N₂O reduction. Under such conditions one would expect that the ratio of the changes in δ^{18} O and δ^{15} N^{bulk} (r_{o-n}) is around 2.5 and that the ratio of the changes in SP and δ^{18} O (r_{sp-o}) is 484 between 0.2 and 0.5. The mean (median) ratios for r_{o-n} and r_{sp-o} for these selected events were 485 486 0.69 (0.44) and 2.1 (1.16), respectively. While the high values of r_{sp-0} indicate that for instance changing physical conditions such as soil moisture may play a role in field measurements, the 487 deviation of r_{o-n} from the value of 2.5 could either indicate that the fractionation factor for ¹⁸O 488 might be smaller than the one for ¹⁵N or that there is no correlation of fractionation factors in 489 natural environments. This is in line with recent findings showing that apparent isotope effects 490 491 associated with N₂O reduction are sensitive to experimental conditions which influenced diffusive isotope effects (Lewicka-Szczebak et al., 2014, 2015). The same study also showed that 492 fractionation factors during N₂O reduction for ¹⁵N and ¹⁸O were variable (from -11 to +12 ‰ and 493 494 from -18 to +4 ‰, respectively), and not predictable for field conditions yet. Therefore, to date,

495 the amount of N_2O reduction prior to emission cannot be inferred with sufficient robustness from 496 field measurements alone, without the knowledge of isotopic composition of the substrates.

497 4.5 Controls on isotopic composition and event based data aggregation

498 The high temporal resolution of N_2O isotopic and auxiliary measurements allowed us to investigate controls on N₂O isotopic composition over the 3 months campaign period. 499 500 Correlations with isotopic composition were highest and positive for DOC and soil temperature 501 (Table 2). The significant correlation with temperature for the whole campaign was due to a significant correlation during the "dry" part of the campaign. If the increase in SP was due to 502 increased contribution of nitrification, $\delta^{15}N^{\text{bulk}}$ should decrease due to the higher isotopic 503 fractionation during this process. The simultaneous increase in SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$ revealed in 504 Figure 7, however, indicates an increased share of N₂O reduction to N₂ which might have been 505 triggered by increased substrate availability (DOC) for heterotrophic denitrification. The reported 506 effect of temperature on the N₂O:N₂ ratio is not without any doubt, but a decrease has been 507 observed with increasing temperature, supporting the hypothesis that N_2O reduction increased as 508 temperature rose throughout the measurement period (Saggar et al., 2013). 509 510 Though substrate availability has been identified as a major control on N_2O source processes (see references in Saggar et al., 2013), correlations between N₂O isotopic composition and NO_{3⁻} and 511 NH_4^+ concentrations were low, except for the correlation with $\delta^{15}N^{\text{bulk}}$. The reason might be both 512

the number of measurment points for substrate concentrations being lower compared to other

explanatory variables and substrate concentrations not necessarily reflecting process or turnover

515 rates (Wu et al., 2012).

The low explanatory power of all linear regressions underlines that drivers for N_2O emissions are

517 highly variable and may even change from event to event. In absence of management or

rewetting events (group BG), isotopic composition covered the whole range of measured values, while management or rewetting events were characterized by lower variability in isotopic composition. Values for SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$ were low for Mana I, rewetting and Mana III, whereas event Mana II showed increased SP, $\delta^{15}N^{bulk}$ and $\delta^{18}O$. This indicates that processes must have been different for Mana II, although management was almost identical.

523 4.6 Short term variation of isotopic composition

The Keeling plot approach is based on conservation of mass and assumes that the atmospheric 524 concentration of a gas in the surface layer is a mixture of background atmospheric concentration 525 526 and a variable amount of gas added by a source, raising the atmospheric concentration above background. The source's isotope value can be determined given that its isotope value remains 527 528 constant during the observation period. In this study, we used noon-to-noon data in the Keeling plots to determine isotope values of soil-derived N₂O for the respective noon-to-noon period. 529 Hence, the source processes underlying these N_2O emissions have to be constant on this time 530 531 scale. Currently, little is known about the rate of change of N_2O source processes over time-steps of minutes to hours. However, changing relative contributions of source processes, which change 532 the isotopic composition in soil-emitted N_2O , would be reflected by deviations from a linear 533 534 relation between inverse concentration and isotopic composition. As the Keeling plots showed no 535 obvious deviations from a linear relation within our measurement precision, we conclude (1) that 536 the use of the Keeling plot approach was valid in our study, and (2) that changes in N_2O source 537 processes in our study site occurred at a time step of one day or more. While our data suggests 538 that there are little or no changes in source processes underlying N₂O emissions within a noon-to-539 noon period, clear and distinct day to day variation in isotope values of soil derived N_2Q , 540 especially in SP, were observed. Such changes were often strong and abrupt following management events (ManaI & III, Rewetting), indicating a significant response of microbial 541

processes to the imposed disturbance. Larger than expected variability in isotope values was 542 543 observed in between management events (class BG), when no obvious variation in environmental drivers occurred. Since noon-to-noon concentration increases were very small during these 544 periods, part of this variability may be attributed to increased uncertainty around the intercept of 545 546 the Keeling plot. This is also reflected in the relatively large error bars around isotope values on days when N₂O fluxes were low (Figure 3). Alternatively some of the variation in isotope values 547 associated with these small fluxes may result from air masses not representative of the grassland 548 site as the concentration footprint influencing the N₂O source signature is larger than the flux 549 footprint (Griffis et al., 2007). As the Keeling plots showed no obvious deviations from a linear 550 relation within our measurement precision (See supplementary file S1), we conclude (1) that the 551 552 use of the Keeling plot approach was valid in our study, and (2) that changes in N_2O source processes in our study site occurred at a time step of one day or more. While our data suggests 553 554 that there are little or no changes in source processes underlying N₂O emissions within a noon-tonoon period, clear and distinct day-to-day variation in isotope values of soil derived N_2O_1 . 555 especially in SP, were observed. Such changes were often strong and abrupt following 556 557 management events (ManaI & III, Rewetting), indicating a significant response of microbial processes to the imposed disturbance. Larger than expected variability in isotope values was 558 observed in-between management events (class BG), when no obvious variation in environmental 559 drivers occurred. Since noon-to-noon concentration increases were very small during these 560 periods, part of this variability may be attributed to increased uncertainty around the intercept of 561 the Keeling plot. This is also reflected in the relatively large error bars around isotope values on 562 days when overnight N_2O concentration increase was low (Figure 3). Alternatively, the variation 563 in isotope values associated with small overnight concentration increase may result from other 564 565 land use or land cover. The EC fluxes are calculated from the turbulent fluctuation of

566	concentration and vertical wind speed (i.e. the covariance of the concentration and wind speed
567	deviations from the half-hourly mean) and therefore account for the modulation of concentration
568	around a short term (30 min) mean caused by locally emitted N ₂ O. Isotopic composition based on
569	Keeling plots however is determined from total N2O accumulated in the nocturnal boundary layer
570	and, thus, this approach also contains molecules that had been emitted outside the flux footprint,
571	which almost exclusively comprised our grassland site (Zeeman et al., 2010), within the larger
572	concentration footprint (Griffis et al., 2007). However, two facts indicate a major influence of the
573	studied grassland on the determined N_2O isotopic composition: First, the N_2O isotopic
574	composition is very stable for a noon-to-noon period as indicated by a linear relationship between
575	individual measurements (supplementary file S1). This relationship persists even though wind
576	speed and direction are changing and, therefore, individual N ₂ O isotope mesurements integrating
577	over 16 minutes sampling interval originate from different source areas. Secondly, the CHA
578	grassland can be characterized as a site with vigorous N_2O emission and therefore may dominate
579	the determined N ₂ O isotopic composition as the influence of a source area increment scales with
580	the source strength. The grassland was restored in 2012 which lead to extraordinary high N_2O-N
581	emission of 29.1 kg ha ⁻¹ year ⁻¹ (Merbold et al., 2014). In the following year 2.5 kg N_2 O-N ha ⁻¹
582	were released. This value is still in the range of maximum emissions reported for another
583	intensively managed Swiss grassland, emitting 1.5-2.6 kg N ha ⁻¹ year ⁻¹ and at least a factor of five
584	compared to an extensively managed grassland with less than 0.5 kg N ha ⁻¹ year ⁻¹ (Ammann et
585	al., 2009). With regard to distant land use and land cover, the 2.5 kg N ₂ O-N are also more than
586	double the median (between the 70 and 75 percentile) of all reported values for cultivated
587	temperate sites and higher than the highest value reported for forests presented in a study
588	containing 1008 N ₂ O emission measurements from agricultural fields (Stehfest and Bouwman,

589 2006). However, it cannot be excluded that N₂O isotopic signatures analyzed above the grassland 590 were influenced by adjacent ecosystems.

591 4.7 Flux weighted averages of source isotopic compositions

592 N₂O isotopic composition can be used to calculate and further constrain the global N₂O budget (Kim and Craig, 1993; Yoshida and Toyoda, 2000). The analysis of emissions from different 593 594 sources such as agricultural soils or managed grasslands based on box models and isotopic 595 composition is complicated by distinct temporal and spatial variability of isotopic composition 596 (Kim and Craig, 1993; Toyoda et al., 2011; Yoshida and Toyoda, 2000); hence, flux weighted 597 averages are required to obtain representative values for agricultural N₂O (Perez et al., 2001). Our flux weighted averages of 6.9 ± 4.3 , -17.4 ± 6.2 and 27.4 ± 3.6 % for SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$ are 598 well within the range of values 2.9 to 36.6, -41.5 to -1.9 and 23.2 to 51.7 ‰ for agricultural soils 599 600 (Park et al., 2011; Toyoda et al., 2011). The comparison with other grassland soils (Opdyke et al., 601 2009; Park et al., 2011) indicates that the variability of isotopic composition within a group, such as grassland, may be considerable (for SP: 2.2 to 11.1 %). PartOne has to keep in mind, however, 602 that part of the observed variability may be attributed to the fact that the footprint area of the N₂O 603 604 isotopic composition includes areas with other land use or land cover. Another part of the 605 variability might be also explained by a limited compatibility of laboratory results, as recently 606 demonstrated in an inter-laboratory comparison campaign (Mohn et al., 2014). The uncertainty in 607 budgets derived by isotopic composition depends on the uncertainty of the representative isotopic composition for a single source, which can be reduced by a quasi-continuous measurement 608 approach, as shown in this study. 609

610 5 Conclusion

Our field observations indicate that nitrifier-denitrification and denitrification (process group 611 N₂O_D) dominated throughout the measurement period and that variation in isotopic composition 612 was more likely due to variation in the extent of N2O reduction rather than contributions of 613 614 NH₂OH oxidation, or fungal denitrification. High temporal resolution of isotopic composition in soil-emitted N₂O showed that at the beginning of the growing season, medium wfps and low 615 temperature induced low isotope values (representative for process group N_2O_D), whereas in the 616 617 second part of the measurement period, higher temperature and DOC stimulated N₂O reduction to N₂, although wfps was lower. Management or rewetting events were mostly characterized by low 618 SP, $\delta^{15}N^{\text{bulk}}$ and $\delta^{18}O$, but the event Mana II indicated that processes underlying N₂O emissions 619 can vary even under similar management conditions. With this study, a new method is available 620 621 that can provide real-time datasets for various single N₂O emitting (eco)systems, such as as 622 grasslands or agriculturally used regions, which will help in further constraining the global N₂O budget based on box model calculations. However, future campaigns should be accompanied by 623 624 footprint modeling for optimization of the inlet height and associated concentration footprint size. 625

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Tables

876 877

 Table 1: Reference gas and compressed air tanks used during the campaign. S1 and S2 represent the anchor and calibration standard. C1 and C2 are the target gases used for determination of system performance. The reported

precision is the 1σ standard deviation.

Tank	$\delta^{15}N^{\alpha}$	$\delta^{15}N^{\beta}$	$\delta^{15}N^\beta \qquad \delta^{18}O$		SP	mixing ratio	
	[%0]	[%0]	[%0]	[%0]	[‰]	[ppm] / [ppb]*	
S1	15.66 ± 0.03	-3.22 ± 0.13	34.89 ± 0.05	6.22 ± 0.07	18.88 ± 0.13	90.09 ± 0.01	
S2	10.38 ± 0.03	-10.55 ± 0.1	25.44 ± 0.06	-0.09 ± 0.05	20.93 ± 0.10	87.28 ± 0.003	
C1	15.40 ± 0.08	-3.04 ± 0.06	43.65 ± 0.08	6.18 ± 0.05	18.44 ± 0.10	327.01 ± 0.05	
C2	15.65 ± 0.17	-4.27 ± 0.08	44.20 ± 0.07	5.69 ± 0.09	19.92 ± 0.19	327.45 ± 0.05	

* ppm for S1 and S2, ppb for C1, C2

882 Table 2: Adjusted r² and p-values for regression analysis of Keeling-plot derived isotopic compositions in soil-emitted N₂O

883 versus auxiliary variables N₂O flux (f_{N2O}), difference of maximum and minimum concentration over a noon-to-noon

884 period (ΔN₂O), precipitation (prcp), soil moisture (wfps) and nutrient concentrations (NO₃⁻, NH₄⁺ and DOC).

885

explanatory	SP	SP	$\delta^{15} N^{bulk}$	$\delta^{15} N^{bulk}$	δ ¹⁸ O	δ ¹⁸ Ο	Ν
	r ²	р	\mathbf{r}^2	р	\mathbf{r}^2	р	
f _{N2O}	0.14	**	0.04	0.06	0.16	**	62
$\Delta N_2 O$	0.09	*	0.1	*	0.11	*	65
prcp	0.24	**	0.03	0.08	0.24	**	62
wfps	0.14	*	0.29	**	-0.009	0.52	65
Т	0.22	**	0.30	**	0.12	*	65
DOC	0.23	*	0.30	*	0.03	0,23	18
NO ₃ -	0.04	0.14	0.27	*	0.16	*	31
$\mathbf{NH_{4}^{+}}$	-0.03	0.75	-0.03	0.89	-0.03	0.93	31

 $Significance \ codes: \ *: \ p < 0.05; \ **: \ p < 0.001. \ sample \ size \ (n) \ differs \ due \ to \ data \ availabilities.$

886

888 Figure legends

Figure 1: Long-term stability (standard deviation σ) derived by target gas injections (n=331) over a 3-month period. As two target gas tanks were used, histograms show deviation of respective tank means, \bar{x} , for $\delta^{15}N^{\alpha}$, $\delta^{15}N^{\beta}$, $\delta^{18}O$, $\delta^{15}N^{\text{bulk}}$ and SP, respectively

892

Figure 2: Target gas (red) and surface layer (black) N₂O mole fractions (top) and δ-values (three bottom panels)

measured in the atmospheric surface layer in 2.2 m height during the field campaign. Each couple of vertical dashed

blue lines indicates the management events mowing (first line) and fertilization (second line).

896

897 Figure 3: Noon-to-noon averaged N₂O flux (f_{N2O}), overnight increase in N₂O mole fractions (difference in minimum 898 and maximum N₂O concentration in a noon-to-noon period; $\delta N_2 O$), Keeling-plot derived isotopic composition of soil-emitted N₂O (SP, $\delta^{15}N^{\text{bulk}}$, $\delta^{18}O$), nutrient concentrations (ammonium, nitrate and dissolved organic carbon; 899 900 DOC), water filled pore space (wfps), precipitation (prcp) and soil temperature (T) over the measurement period. 901 Each couple of vertical dashed blue lines indicates the management events mowing (first line) and fertilization 902 (second line). Transparent blue boxes represent periods of N₂O emission influenced by management or rewetting 903 (third box). 904 905 **Figure 4:** Standard error for SP (ε_{SP}) of soil-derived N₂O estimated by the Keeling plot approach as function of

906 overnight N₂O accumulation in the surface layer. The red dashed lines show 12 ppb increase in N₂O mole fractions.

907 Red full circles represent the selected subset.

908

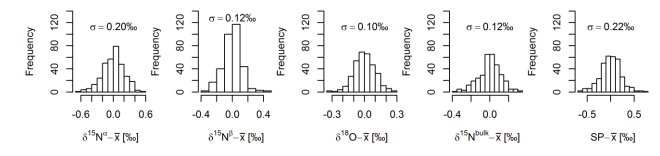
Figure 5: SP - NH_4^+ / NO_3^- and SP - wfps / soil temperature maps. The size of the points is inversely scaled to Keeling plot intercept standard error so that biggest points are those with lowest uncertainty.

912 **Figure 6:** Boxplots for Keeling-plot derived SP, $\delta^{15}N^{\text{bulk}}$, $\delta^{18}O$ of soil-emitted N₂O and wfps of management events 913 (Mana I – III), rainfall after a dry period (Rewetting), and the remaining measurement period (BG).

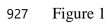
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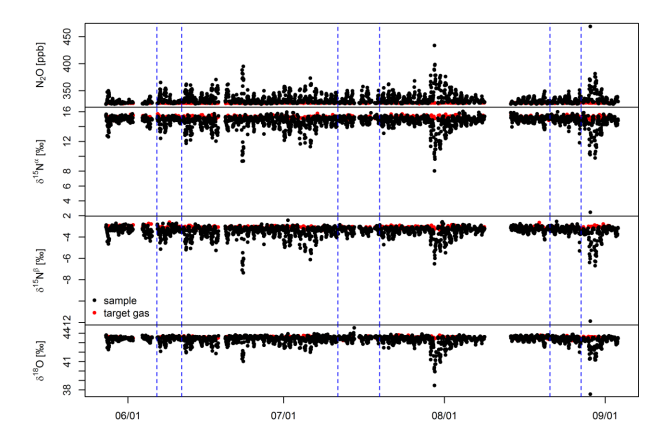
- 915 **Figure 7:** MapLeft panel: map of N₂O isotopic composition SP/ δ^{15} N^{bulk} with rectangles representing process groups
- 916 N₂O_N and N₂O_D based on SP values in Decock and Six ($\frac{20132013b}{2013b}$) and δ^{15} N^{bulk} estimated from minimum and
- 917 maximum fractionation factors reported in Baggs (2008) and substrate isotopic compositions reported by Bedard-
- Haughn et al (2003), Pörtl et al. (2007) and Toyoda et al. (2011). <u>Right panel: map of SP/ δ^{18} O with traces of</u>
- 919 <u>management events (ManaI in black, ManaII in red, ManaIII in green) and the rewetting event (blue). Isotopic</u>
- 920 compositions are plotted for the transparent blue boxes in Fig. 3 including one preceding and one following
- 921 composition. The preceding composition is represented by the enlarged filled triangle and transparency of the line
- 922 <u>connecting the compositions decreases with event duration.</u>

923



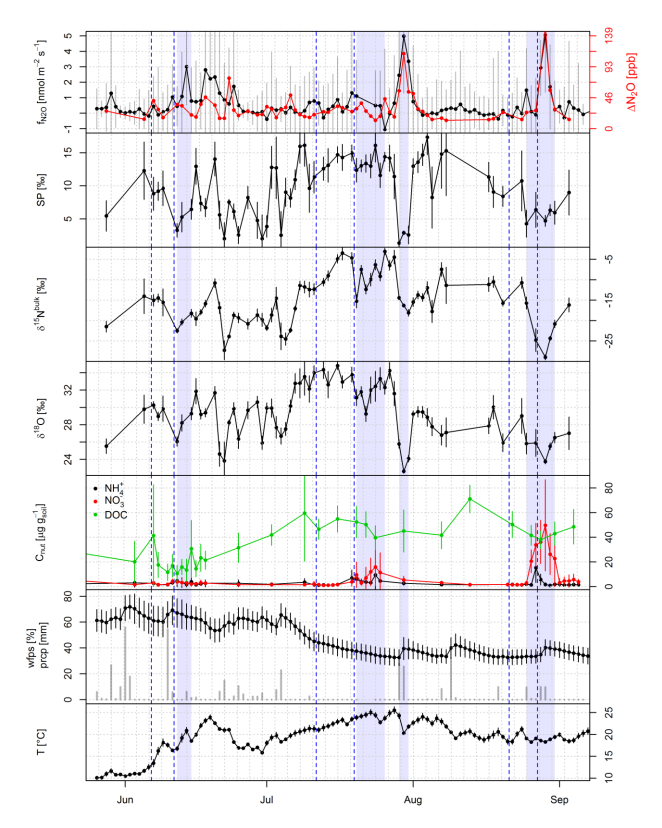




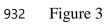


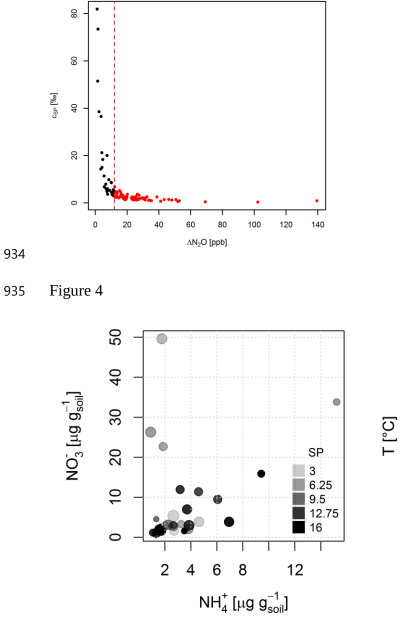


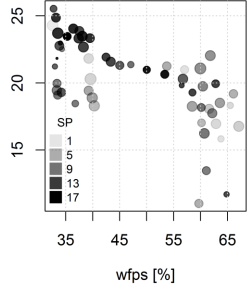
929 Figure 2











937 Figure 5

