Field comparison of nitrous oxide emission measurements using micrometeorological and chamber methods

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Summary – In order to check the reliability of micrometeorological and chamber methods for the measurement of N2O fluxes at the ‘soil–atmosphere’ interface, an experiment using these two methodologies was performed in autumn on a bare soil fertilized with 150 kg N-NH4NO3 ha⁻¹. Micrometeorological measurements were continuously taken using either the gradient or eddy correlation methods, the N2O gas mixing ratio being measured with a tunable diode laser absorption spectrometer. N2O emission fluxes were also monitored in two automatic chambers during two 3-h periods every day, ie, around midday and midnight, whereas 16 manual chambers were monitored only around midday. Areas over which the fluxes were measured ranged from 0.18 and 0.49 m² for the manual and automatic chambers, respectively, to more than 10⁴ m² for the micrometeorological methods. Additional soil variables were obtained, including the soil bulk density profile, water content, temperature and N-NO3⁻ and N-NH4⁺ concentrations. The different methods provided similar estimates of N2O fluxes with an average over the whole of the common measurement period of around 6 ng N-N2O m⁻² s⁻¹. The 16 manual chambers showed coefficients of variation comprised between 30 and 150 % during the 2 weeks of common observations. This variability may explain the unusual shape of the N2O concentration gradient in the atmosphere observed sometimes, which led to the conclusion that the gradient data set contained information allowing the spatial variability in N2O emission to be characterized. Hourly micrometeorological fluxes indicated a temporal variability of about 100 % over daily periods. Microbial processes may be assumed to vary little over such a period and this temporal variability is probably a consequence of physical processes, such as air pressure fluctuations. Slight daily cycle tendencies were observed. The weak levels of N2O emissions were caused by rainfall deficit, which maintained the soil at a low water content, preventing strongly anaerobic conditions and the N fertilizer from leaching from the dry surface to wet soil layers. (© Inra/Elsevier)

nitrous oxide flux / greenhouse effect / denitrification / soil / micrometeorological methods / chamber methods

Résumé – Comparaison au champ des émissions de protoxyde d’azote mesurées par méthodes micrométéorologiques et par méthodes des chambres. Afin de tester la fiabilité des méthodes micrométéorologiques et des méthodes par chambres au sol pour la mesure des flux de N₂O à l’interface « sol-atmosphère », une expérimentation utilisant ces deux méthodologies a été conduite à l’automne sur un sol nu fertilisé préalablement par 150 kg N-NH₄NO₃ ha⁻¹. Les mesures micrométéorologiques ont permis un suivi continu des émissions par la méthode des gradients ou par la méthode des corrélations turbulentes, les rapports de mélange en N₂O étant mesurés par un spectromètre d’absorption

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À diode laser accordable. Ces mêmes émissions ont été aussi observées par deux chambres au sol à fonctionnement automatisé pendant deux périodes journalières de 3 h en milieu de journée et de nuit, et par seize enceintes manuelles en milieu de journée. Les surfaces associées aux méthodes allaient de 0,18 et 0,49 m² respectivement, pour les enceintes manuelles et automatiques, à plus de 104 m² pour les mesures micrométéorologiques. Les profils de densité apparente, de teneurs en eau, températures et concentrations en NO3⁻ et NH₄⁺ ont été suivis simultanément aux mesures de flux de N₂O. Les différentes méthodes ont abouti à des estimations similaires. Le flux de N₂O moyen sur l'ensemble de la période commune de suivi était de l'ordre de 6 ng N-N₂O m⁻² s⁻¹. L'utilisation de 16 chambres manuelles a montré une grande variabilité spatiale des émissions comprise entre 30 et 150 % pendant les deux semaines de mesure. Cette variabilité pourrait expliquer la forme inhabituelle des profils de concentrations atmosphériques observés par les méthodes des gradients. Les flux horaires donnés par les mesures micrométéorologiques présentaient une variabilité de l'ordre de 100 % sur des périodes journalières : on peut penser à de faibles variations des processus microbiens à ces échelles de temps et attribuer cette variabilité à des phénomènes physiques comme les fluctuations de la pression de l'air. Les faibles émissions de N₂O s'expliquent par de faibles précipitations avant et au cours de la période expérimentale, insuffisantes pour créer des conditions anoxiques dans le sol et pour qu'il puisse y avoir un entraînement des fertilisants azotés vers les horizons plus humides. (© Inra/Elsevier)

**INTRODUCTION**

Soils have been identified by the Intergovernmental Panel on Climate Change (IPCC, 1995) as the major source of atmospheric nitrous oxide (Smith et al, 1994). Contributions from cultivated fields are estimated to be 3.5 Tg year⁻¹, with a possible actual value between 1.8–5.3 Tg year⁻¹ (IPPC, 1995). This would represent approximately two thirds of the probable 5.7 Tg year⁻¹ anthropogenic sources of N₂O (3.7–7.7 Tg year⁻¹) and therefore explain most of the observed annual increase in atmospheric N₂O. There is evidence that the flux of N₂O from agricultural land (fertilized fields and grassland) is one of the major and increasing contributors to total global emissions of this gas: emissions increase with cultivation and increasing inputs of N in the form of mineral fertilizers or organic manure (Bouwman, 1990; Granli and Bockman, 1994). However, uncertainty persists as regards the fluxes attributable to cultivated soils (Smith, 1990), and improving assessment of N₂O fluxes from fertilized land is a clear priority for research. Quantitative relationships between soil properties and other environmental factors are not yet sufficiently well understood to permit prediction of emissions over different time scales and extended regions and to assess the global N₂O budget (Granli and Bockman, 1994; Mosier et al, 1996). A variety of techniques have been developed to measure surface–atmosphere gas exchange. Most estimations of N₂O emission rates in the field have been based up to now on the use of various static chamber techniques (Hutchinson and Livingston, 1993; Hénault and Germon, 1995; Arah and Smith, 1990) and many studies have demonstrated a high degree of spatial and temporal variability in denitrification and N₂O fluxes (Ambus and Christensen, 1994; Folorunso and Rolston, 1984; Parkin, 1987). In order to obtain reliable estimates of N₂O fluxes at the field scale, numerous chambers are necessary. Recent technological advances have dramatically improved our ability to perform precise real-time measurement for minor compounds such as N₂O by micrometeorological methods.

Each technique presents its advantages and drawbacks, and no single approach is applicable to all field conditions. Enclosure techniques are relatively low in cost, simple to operate, and especially useful for addressing research objectives based on discrete spatial observations. They are adaptable to a wide variety of field conditions from local to global spatial scales (Hénault et al, 1996; Clayton et al, 1994). Estimating emission at the field scale, however, requires many measurements so that the temporal and spatial variability of emission can be overcome. The chambers also physically disrupt the surface and atmosphere by altering the radiation environment, temperature, humidity and by cutting pressure fluctuations within the chambers (Livingston and Hutchinson, 1995). Such disruptions are likely to affect the rate of N₂O emission.

An alternative non-disruptive approach is offered by micrometeorological techniques (Cellier, 1995), which integrate N₂O emissions at the field scale (1–10 ha). These techniques involve the measurement of meteorological data and gas
concentration at a location site above the soil–vegetation surface. The associated fluxes are representative of the upwind terrain. The conditions required by the method are a large fetch of uniform source surface (100–500 m), which in many applications may be a limitation.

An experiment was carried out to measure N₂O fluxes at different scales, ranging from less than 0.5 m² by traditional manual chamber methods or by automated chamber techniques and to more than 1 ha by micrometeorological methods. In this paper the performances and the sensitivity limits of each technique were evaluated and the various N₂O fluxes obtained with these different approaches were compared. The possible reasons for these differences were also explored.

MATERIALS AND METHODS

Experimental site

The experimental site was a flat cultivated field near Dijon (Burgundy, France), 47°12' latitude north and 5°23' longitude east. The experiment was performed from 23 October to 10 November 1995. The soil (Haplic Luvisol in the FAO classification) was a deep sandy loam brown soil of the Saône Terrace with a sand content of 47% and a clay content of 12.5%. Its organic C content was 1.4% and its water pH was 6. A bulk density profile was carried out with measurements taken every 0.05 m from the soil surface to 0.5 m in depth using a gamma-ray transmission probe from Inra-LCPC (Bertuzzi et al, 1987). The experimental field (∼3 ha) was bordered by a path on one side and a grove on the other. Cultivated fields flanked the other sides. The field was cultivated with wheat in 1995 and ploughed after the harvest. Three days before the experiment began, 150 kg N ha⁻¹ were applied as NH₄NO₃ and the field was then harrowed to a depth of 0.1 m to limit spatial heterogeneity due to N fertilization.

On 25 October 1995 all the equipment for flux measurement was installed. Location of the different equipment and sampling sites in the field are presented in figure 1.

N₂O emission measurements by closed chamber methods

Closed chambers of two designs were used. A set of 16 manual chambers (MA-CH) having a surface of 0.18 m² was used as the first method. The chambers were composed of hollow PVC cylinders, 0.50 and 0.15 m in diameter and height, respectively. They were positioned in the soil for the whole experiment at a depth of approximately 0.08 m at the same location. The 16 chambers were arranged all over the field according to figure 1. During the measurement of N₂O fluxes, these chambers were sealed with a PVC lid fitted with a ring of foam rubber to ensure adhesion to the chambers. The lids were kept in place by a metal cross fixed to the chambers. Tiny holes in the lids maintained the internal pressure of each chamber to atmospheric pressure. N₂O emissions were estimated over a 2 h 15-min period. The atmosphere in the chamber was sampled four times during this period using blood collecting tubes (Terrumo 3 mL). These tubes were previously purged of components interfering with N₂O analysis. N₂O was thereafter analyzed by electron capture chromatography (sensitivity c. 20 ppbv) with Varian Star Workstation integrated software.

The second design consisted of two square automatic chambers (AU-CH) having a surface of 0.49 m² and constructed from galvanized iron. Their characteristics were kindly supplied by K. Smith and A. Scott (University of Edinburgh and Scottish Agricultural Centre, UK, respectively). The chambers were 0.70 and 0.25 m in length and height, respectively, and sunk into the soil at approximately 0.1 m and positioned at around 30 and 100 m from the sonic mast south of the field (fig 1). Opening, closing and gas sampling were performed automatically. Each chamber was connected to a set of 24 loops in copper tubes (40 cm in length, 1/8", ie, 0.9 mL volume approximately) with two Teflon tubes. Commutation between these loops was achieved by two scanning-valves; air was circulated between the chambers and each loop was filled with an air-pump. Kinetics of N₂O emissions were performed for two periods of 3 h every day, ie, between 1:00 and
4:00 and between 13:00 and 16:00 UT. For each period, three gas samplings were taken. N$_2$O concentrations were thereafter analyzed in laboratory by electron gas chromatography with HP 5890 Series II having two Porapak Q columns (10 and 6 ft) coupled to a Hewlett Packard Vectra Workstation integrated software. Its sensitivity was around 20 ppbv N$_2$O.

In both methods, N$_2$O fluxes were estimated by the following equation:

$$F = \frac{\alpha P}{RT} \frac{h_c}{M_{N-N_2O}}$$

where $F$ is the N$_2$O flux (ng N-N$_2$O m$^{-2}$ s$^{-1}$), $\alpha$ is the rate of N$_2$O accumulation of the chambers (m$^3$ m$^{-3}$ s$^{-1}$), $P$ the absolute pressure (Pa), $R$ the gas constant (8.315 J mol$^{-1}$ K$^{-1}$), $T$ the absolute temperature (K), $h_c$ the mean height of the chamber above the soil surface (m) and $M_{N-N_2O}$ the molar weight of nitrogen of N-N$_2$O (ng). The values of $T$ and $P$ were taken to be equal to laboratory temperature and pressure during N$_2$O gas analysis. The accumulation kinetics were considered to be significant when the correlation coefficient was higher than 0.9 ($n = 4$: manual chamber, $n = 3$: automatic chamber, with $n$ the number of samples). If not, the estimated values were still used when the range of the gas mixing fraction during the kinetic was lower than 0.1 ppmv, corresponding approximately to a flux of 1.6 ng N-N$_2$O m$^{-2}$s$^{-1}$ for the automatic chambers. Otherwise, data were not considered as relevant because of the lack of linearity.

The relative flux error may be approximated by a first order limited development of equation [1]. The resulting expression is:

$$\frac{\sigma_F}{F} = \sqrt{\left(\frac{\sigma_\alpha}{\alpha}\right)^2 + \left(\frac{\sigma_P}{P}\right)^2 + \left(\frac{\sigma_T}{T}\right)^2 + \left(\frac{\sigma_{h_c}}{h_c}\right)^2}$$

where $\sigma_\alpha$, $\sigma_P$, $\sigma_T$ and $\sigma_{h_c}$ are the standard deviations of N$_2$O accumulation rate, air pressure, temperature and mean chamber height, respectively. The relative uncertainty of each of these four terms was estimated: the uncertainties on $P$, $T$, and $h_c$ measurements led to a relative emission uncertainty of ±2, 3 and 2 %, respectively. A sensitivity of ±5 % on detection of low accumulation rates ($\alpha$) of N$_2$O was reached according to the two GC characteristics. The mean error on flux measurements with chambers was then less than ±7 %.

N$_2$O emission measurements by micrometeorological methods

Micrometeorological fluxes were determined by two methods: ‘gradient’ and ‘eddy correlation’. The gradient method was used for continuous monitoring of fluxes and the eddy correlation technique was used for 4 d and restricted to short periods (less than 5 h). The N$_2$O mixing ratio observations used for the micrometeorological method were made with a tunable diode laser absorption spectrometer (TDLAS) (Aerodyne Research Inc, Billerica, MA, USA) system (Kolb et al, 1995).

Gas analyser (TDLAS)

The TDLAS uses a multipass absorption Herriott cell with an optical path length of 36 m and a small volume cell (0.27 L), operated at reduced pressure (35 Torr) to minimize interference from nearby spectral lines. Tuning of the laser emission wavelength to the selected absorption line (2207.6 cm$^{-1}$) was accomplished by the appropriate setting of laser temperature (around 88 K) and current through the diode, while small variations in the laser current (ramp) were used to scan absorption across the full infrared transition line shape. The spectral information was analyzed in real time. The diode laser power spectrum (baseline) was represented as a slowly varying polynomial of adjustable order; typically a quadratic or cubic polynomial was used. The area between the absorption line and the polynomial baseline, together with the absorption coefficient for the line and path length, was used to calculate the absolute concentration of N$_2$O.

To avoid the need for density corrections due to water vapour and heat fluctuations (Webb et al, 1980), the air was dried by a counterflow Perma Pure dryer and equilibrated to constant temperature before entering the optical cell. Under field conditions, the instrument precision (RMS) was 0.1 % at 1 Hz and 1 % at 10 Hz. The major problem with this instrument during extended observation times was that a shift in the frequency emission was induced by laser diode temperature regulation drift. In this case the area defined by the baseline and the absorption line for a fixed concentration could be modified causing a drift in the evaluation of the N$_2$O mixing ratio in the long term. For the eddy correlation measurements, this problem was solved by using a low-frequency filter in the data processing.

Gradient measurements

The micrometeorological theory for N$_2$O flux measurement by the gradient method has been described in detail by Thom (1975) and Monteith and Unsworth (1990), and has been widely applied to trace gas fluxes. The following provides a brief overview of the method applied.

In the equilibrated constant-flux layer established over large uniform areas, the flux of any trace gas may be calculated as:

$$F = -u^* c^*$$

where $F$ is the flux of the trace constituent (ng N-N$_2$O m$^{-2}$ s$^{-1}$), $u^*$ and $c^*$ are horizontal velocity (m s$^{-1}$) and N$_2$O concentration (ppbv) scales defined by the wind, $u(z)$ or concentration $c(z)$ vertical profiles according to the equations:

$$u(z) = \frac{u^*}{k} \ln \left(\frac{z}{z_0}\right) - \Phi_n \left(\frac{z}{L}\right)$$

where $k$ is the von Kármán constant, $z_0$ is the surface roughness length, $L$ is the canopy or boundary layer height, $\Phi_n$ is the neutral stability function and $n$ is the gradient Richardson number.
The terms $\Phi_m$ and $\Phi_h$ are empirical non-dimensional stability correction functions for momentum and heat or gas concentration (Brutsaert, 1982); $z_0$ is the roughness length, $k$ is von Karman constant ($k = 0.41$), and $L$ is the Monin-Obukhov length depending on the stability of the atmosphere. In the referentials defined by the bracket terms, the slope coefficients $u^*$, $c^*$ and errors on the slope ($\sigma_{u^*}$ or $\sigma_{c^*}$) are obtained by a linear regression using equations [4] and [5].

By deriving the flux expression (equation [3]), flux error including instrumental and ‘model’ uncertainty will be estimated as follows:

$$\frac{\sigma_F}{F} = \sqrt{\left(\frac{\sigma_{u^*}^2}{c^*}\right)^2 + \left(\frac{\sigma_{c^*}^2}{u^*}\right)^2}$$

[6]

The measurements were made at five heights (0.1, 0.3, 0.7, 1.1 and 1.5 m) above the soil surface. Wind speed was measured with opto-electronic cup anemometers (Cimel, Paris, France) with a stalling speed of about 0.3 m s$^{-1}$ and air temperature with copper-constantan thermocouples (AWG24) placed in a double-shielded aspirated screen. Five inlet tubes allowed the air sample to be directed into the cell of the TDL spectrometer over a distance of 20 m. The switching between heights was performed by five two-way valves. A sequential cycle for N$_2$O measurement was carried out to reduce the temporal variability between scans of each level. The five valves were switched alternately every 36 s, and the data were recorded after 10 s in order to allow residual air from the inlet tube cell to be removed. The average wind speed, temperature and N$_2$O mixing ratio were recorded every 15 min on a CR10 datalogger (Campbell Scientific, Shepshed, UK) together with standard deviation on N$_2$O mixing ratio.

To overcome the variability due to the measurement conditions by the gradient method the hourly mean fluxes, $F_p$, were calculated from 15-min fluxes weighted by the inverse flux error ratio as follows:

$$F_p = \sum_{i=1}^{4} P_i F_i \quad \text{with} \quad P_i = \frac{1}{\Delta F_i} \sum_{j=1}^{4} \frac{1}{\Delta F_j}$$

[7]

$F_i$ and $\Delta F_i$ are the 15-min fluxes and flux errors deriving from equations [3] and [6], respectively. $P_i$ the weight assigned for each 15-min flux.

This treatment allowed the effect of 15-min values with large uncertainties in the hourly flux to be minimized. The resultant hourly mean flux error was:

$$\Delta F_p = \sqrt{\sum_{i=1}^{4} P_i^2 (\Delta F_i)^2}$$

[8]

Eddy correlation measurements

In this method the vertical flux density $F$ of a trace gas is written as the covariance between the vertical wind ($w$) and trace gas density ($c$) fluctuations as follows:

$$F = \bar{w'}c' = \text{cov}(w,c)$$

[9]

where $w'$ and $c'$ are 15-min standard deviations from the mean vertical wind velocity and trace gas density, respectively.

The fluxes were calculated from 10-Hz measurements integrated over 15-min time intervals. A sonic anemometer (Gill Instruments, model 1012R2) mounted at a height of 1.0 m above the soil surface provided the three wind velocity components and air temperature. Concentrations of N$_2$O were measured using the TDLAS. A Teflon tube, whose inlet point was located close to the sonic anemometer, allowed the air to be directed into the TDLAS cell. Since the TDLAS instrument was remote from the sampling mast, there was a time lag between wind and concentration measurements, depending on tube length, its section and the flow rate. The covariance between vertical wind and N$_2$O concentration was calculated for different time shifts. The effective flux was obtained from the shift that gave the maximum covariance between vertical wind and N$_2$O concentration. For synchronization, all the analog outputs of the TDLAS and the sonic anemometer were measured by a datalogger (21X, Campbell Inc, UK) at 10 Hz. The data were continuously recorded by a micro-computer.

The error on the flux measurement by eddy correlation was also established as described for the gradient method, but on the basis of instrumental limitation only. Considering the flux expression it is possible to estimate the flux error according to the resolution of each sensor (see Appendix A for details):

$$\Delta F = \frac{1}{\sqrt{N}} \sqrt{\sigma_{w^*}^2 (\Delta c)^2 + \sigma_{c^*}^2 (\Delta w)^2}$$

[10]

where $N$ is the number of data used for the flux estimation and $\sigma_{w^*}$ and $\sigma_{c^*}$ are the vertical wind and N$_2$O concentration variance, respectively. $\Delta w$ and $\Delta c$ are the resolution of the sonic anemometer and gas analyser, respectively.

In this method flux measurement errors may occur if the gas analysis system induces a cut-off frequency lower than the upper frequency corresponding to turbulence transport. Studies developed by Leuning (1996) and Lenschow (1991) indicated that air flow transport in pipes can cause concentration fluctuation attenuation. The normalized cut-off half-power frequency ($n_p$) in pipes was described as a function of the Reynolds number, $Re = 2a u_\text{mean} / v$, with $a$ the tube radius (m), $u_\text{mean}$ the mean air flow velocity inside the tube (m s$^{-1}$) and $v$ the kinematic viscosity of air (m$^2$ s$^{-1}$). The Reynolds number must be compared to the normalized atmospheric turbulence flux frequency cut-off. Under neutral and unstable conditions, $n_p$ is given by $n_p = f_c / f_{\text{pipe}}$, where $f_c$ is the cut-off frequency of the atmospheric turbulence, $f_{\text{pipe}}$ the cut-off frequency of the pipe flow.

In terms of 'pipe' flow
normalized frequency the critical frequency \( n_c \) may be then written as:

\[
n_c = \left( \frac{ax}{u_s} \right)^{1/2} \frac{4a^{3/2} x^{1/2} u_s}{\nu Re(z-d)} \tag{11}
\]

where \( x \) is the distance along tube (m). Damping of gas concentration will occur only if \( n_0 < n_c \) and will lead to flux underestimation. For the entire gas analysis line, three components could attenuate the \( \text{N}_2\text{O} \) concentration signal: the sampling inlet tube, the Perma Pure dryer and the analysis cell.

Inside the sampling inlet tube (20 m long and 4.2 mm inner diameter) the flow was turbulent with a Reynolds number of 2 800 and a flow velocity of 10 m s\(^{-1}\). In such conditions according to Lenschow (1991) the attenuation of concentration would be characterized by a normalized cut-off frequency of 0.11 (~5.3 Hz); considering a value of \( u_s/(z-d) \sim 1 \) the normalized critical air turbulence flow frequency was equal to 0.04 (~2 Hz).

The Perma Pure dryer consisted of 100, 0.35 long tube with small internal diameter tubes (0.5 mm). The flow inside these small tubes is laminar with a mean speed of 7 m s\(^{-1}\) and Re ~235. This would lead to an attenuation of concentration with a normalized cut-off \( n_0 = 0.06 \) (~45 Hz) according to Lenschow (1991). The critical turbulence flow frequency for this device is then \( n_c > n_0 \) was then widely satisfied.

For the TDLAS cell the time response depended on the air flow '\( V_c \)' through the analysis cell. The working pressure (~40 Torr) inside the cell and its device imposes a response time \( t_0 = \frac{V_c}{V_c} \sim 0.08 \text{ s} \) to sweep one volume of air '\( V_c \)' at a flow rate '\( V_c \)' through the chamber. The half-power frequency is then given by \( f_0 = \frac{3}{2\pi t_0} = 6 \text{ Hz} \) and higher compare to the critical atmospheric turbulence frequency (\( f_c = 2 \text{ Hz} \)).

For this gas analysis system the main limiting factor for the turbulence signal bandwidth is the sampling inlet tube. Nevertheless, the Reynolds number inside was sufficient to maintain a high cut-off frequency and assumed no underestimate of the flux measurements.

### Additional measurements

Measurements were taken in order to describe the variations in soil water content, temperature, nitrate (\( \text{NO}_3^- \)) and ammonium (\( \text{NH}_4^+ \)) concentrations. The average water content between the surface and a depth of 0.3 m was measured every hour with a time domain reflectometry probe (Trace, Soil Moisture, USA). Two parallel probes 0.45 m long were inserted in the soil at an inclination of 45°. Every 3 or 4 d, measurements were made at depths of 0.02, 0.05, 0.1, 0.15 and 0.20 m with three parallel probes 0.2 m long, which were buried horizontally in the soil. The fraction of the pore space (WFPS) saturated with water was deduced from these measurements and the bulk density profiles made with a gamma ray probe (Linn and Doran, 1984).

Temperature was measured every hour at depths of 0.01, 0.02, 0.05, 0.15, 0.2, 0.25, 0.3 and 0.5 m with chromel–alumel thermocouples.

In addition, the soil \( \text{NO}_3^- \) and \( \text{NH}_4^+ \) contents in the 0–0.2-m horizon were periodically measured on six replicates randomly sampled over the whole field surface. The filtrate collected after soil ion extraction in the presence of KCl M (soil solution ratio 1/5) was analysed by colorimetry with a continuous flux Technicon II analyser (Nicolardot, 1988).

### RESULTS AND DISCUSSION

#### Soil conditions

The measurement of soil conditions during the experiment are presented in figure 2. During the experimental period, the soil water content was relatively low near the soil surface. The corresponding WFPS was above 60 % only at depths equal to or higher than 0.15 m suggesting a low denitrification activity (Grundmann and Rolston, 1987) and that both denitrification and nitrification could be involved in \( \text{N}_2\text{O} \) emission (Linn and Doran, 1984). The rainfall amounts were very low (~10 mm, from day of the year (DOY) 298 to 308) and they only increased slightly the water content of the soil near the surface.

\( \text{NO}_3^- \) and \( \text{NH}_4^+ \) contents measured on the 0–20 cm varied from 10 to 70 mg N kg\(^{-1}\) of soil during this experiment. Highest levels were observed for both \( \text{NO}_3^- \) and \( \text{NH}_4^+ \), 6 d after the N fertiliser application. Ten d after application the N-NO\(_3^-\) and N-NH\(_4^+\) levels were between 20 and 40 mg N kg\(^{-1}\) soil.

Variation of N-NO\(_3^-\) and N-NH\(_4^+\) exhibited approximately about the same pattern and the same level. High spatial variabilities were observed for both \( \text{NO}_3^- \) and \( \text{NH}_4^+ \) with coefficient of variation (CV) ranged from 35 to 75 %. In addition, rainfall was not sufficient to permit the N fertilizer to move below 15 cm.

Daily mean temperatures were steady on the seven first days of this experiment (13 °C). Then temperature decreased regularly during 4 d to a values near 5 °C.

#### \( \text{N}_2\text{O} \) emission estimates from the different methods

Fluxes obtained during the experiment with the different methods are presented in figure 3. The two automatic chambers were sampled twice a day
(from 1:00 to 4:00 UT and from 13:00 to 16:00 UT) from DOY 298 to 318. The measurements obtained with both chambers were averaged (fig 3a). Fluxes obtained with automatic chambers ranged from -0.5 to 13 ng N-N\textsubscript{2}O m\textsuperscript{-2} s\textsuperscript{-1}. From DOY 298 to 308, fluxes were low with a mean midday value of 9 ng N-N\textsubscript{2}O m\textsuperscript{-2} s\textsuperscript{-1}. After DOY 308 fluxes decreased with a mean midday value of 3 ng N-N\textsubscript{2}O m\textsuperscript{-2} s\textsuperscript{-1} and we observed a slight increase in N\textsubscript{2}O emission after DOY 315. Night-time fluxes were not always significant. This was notably the case for DOY 303, 304, 305 and 307.

The 16 manual chambers were sampled for 2 h 15 min near midday from DOY 298 to DOY 312. Figure 3b shows the arithmetic means and standard deviations obtained from the 16 measurements. Mean fluxes obtained with the 16 chambers ranged from 0.7 to 12 ng N-N\textsubscript{2}O m\textsuperscript{-2} s\textsuperscript{-1}. For individual chambers the fluxes ranged between 0.6 and 35 ng N-N\textsubscript{2}O m\textsuperscript{-2} s\textsuperscript{-1}. N\textsubscript{2}O emissions between
DOY 298 and 305 had a mean midday value of 10 ng N-N$_2$O m$^{-2}$ s$^{-1}$. After Julian day 305 a large decrease in emissions was observed with a mean value of 2 ng N-N$_2$O m$^{-2}$ s$^{-1}$. The coefficients of variation (CV) ranged from 30 to 150 % showing that spatial variability of N$_2$O emissions was high. These values are in agreement with those published by Clayton (1994) and Ambus and Christensen (1994). The highest CV values were observed in the case of lowest N$_2$O fluxes when non-detectable fluxes were observed on chambers.

Hourly micrometeorological fluxes measured by eddy correlation and gradient methods are presented in figure 3c between DOY 298 and 308. Non-temporal discontinuities on the N$_2$O flux levels have been observed between the data sets of
the two methods. The micrometeorological fluxes were calculated every 15 min and averaged hourly. For the gradient method the hourly mean fluxes were calculated from 15-min fluxes weighted by the uncertainty according to equations [7] and [8].

For eddy correlation, the calculating flux error involves only the metrological uncertainties (which are themselves approximately equivalent). For this method a simple hourly mean was then used.

The hourly fluxes obtained from DOY 298 to 305 ranged from -5 to 50 ng N-\(\text{N}_2\text{O}\) m\(^{-2}\) s\(^{-1}\), with a 6 ng N-\(\text{N}_2\text{O}\) m\(^{-2}\) s\(^{-1}\) mean value. After DOY 307, fluxes reduced with a mean daily value of 4 ng N-\(\text{N}_2\text{O}\) m\(^{-2}\) s\(^{-1}\).

For all three methods, N\(_2\)O fluxes were varied with soil temperature (fig 2b).

**Micrometeorological measurement sensitivity**

**Gradient method**

Figure 4a presents the variation in 15-min averages of N\(_2\)O concentrations during DOY 301 at heights of 0.1 and 1.5 m corresponding to the highest and lowest inlets for the gradient method. For this day the lowest height mean N\(_2\)O concentration was 299 ppbv with a mean standard deviation (obtained with five cycles of 26 s, ie, 130 measurements) of 2.6 ppbv (\(\sigma/m = 0.88\) %). For the upper inlet, the mean daily concentration was 296 ppbv with an average mean standard deviation of 2.3 ppbv (\(\sigma/m = 0.76\) %).

As expected, the relative standard deviation was higher near the soil surface, which indicated the proximity of N\(_2\)O sources and a higher sensitivity to the N\(_2\)O source distribution. In fact, each standard deviation would result from instrumental sensitivity (0.3 ppbv, 0.1 %) and drift, but also and mainly from the temporal and spatial concentration fluctuations over the field.

The mean daily gradient between 0.1 and 1.5 m was 3 ppbv and varied from -0.5 ppbv at midday to 10 ppbv at night. The daily average standard deviation, calculated as the square root of the sum of the squares of the standard deviations of concentration differences, was 3.5 ppbv. Nevertheless, the mean gradient was usually measured more accurately, because mean standard gradient deviation accounted not only for instrumental resolution, but also for the change with time of emissions.

The flux uncertainty for DOY 301 was estimated from equation [6] (fig 4b). The average relative error (\(\sigma_F/F\)) was about 50 % over this day and varied between 5 and 350 %. To discriminate between the contribution of each component, we estimated the relative error linked to the N\(_2\)O profile concentration (\(\sigma_c/c^*\)) and wind speed profile (\(\sigma_u/u^*\)). These values are presented in figure 4c. The average contributions of the two sources of error calculated over 24 h were similar, with a mean value of 33 % for the N\(_2\)O profile and 37 % for the wind speed profile. The error linked to the wind gradient was larger at night because of low wind speed and associated strong stability. Conversely, the error linked to the concentration gradient was larger during the daytime with strong atmospheric unstability. With the gradient methods, the uncertainty with the measured flux depended on the differences of concentration between the highest and the lowest sampling points. The greater the difference in measurement height, the greater this concentration difference was likely to be. In practice, height was limited to 1 % of the fetch, ie, 1.5 m under our conditions. The gradient magnitude also varied with wind speed; the higher the wind speed, the smaller the concentration gradient. At night, when the wind speed was low, the concentration gradient could be large due to low turbulent diffusion. In the daytime, the N\(_2\)O gradient was low due to high wind speed and strong unstability and the uncertainty on the N\(_2\)O profile was very high. The uncertainty might also have been caused by the spatial heterogeneity of the source. According to the variations in wind direction and velocity variation during the 15-min. sampling, the wind might have sampled zones with different source strength which introduced uncertainties into the gradient. Indeed, the gradient method approach requires that the source strength of N\(_2\)O is to be spatially uniform to avoid the development of significant local advection contributions due to horizontal N\(_2\)O flux gradients. The uncertainty in the log linear height versus N\(_2\)O concentration observed during the experiment might be considered to be the consequence of a ‘non-ideal’ emission site.

**Eddy correlation method**

The error on fluxes with the eddy correlation method was studied by using the basis of equation [10]. The Gill sonic anemometer resolution was \(\Delta w \sim 0.03\) ms\(^{-1}\) and the TDLAS resolution was \(\Delta c \sim 1.2\) ppbv. Given the values of N\(_2\)O concentration and vertical wind variance (\(\sigma_c^2, \sigma_u^2\)) during the experiment, the errors for a 15-min time inter-
Fig 4. Example of gradient measurements during DOY 301. a. Atmospheric concentration gradient between 1.5 and 0.1 m above the soil surface. b. 15-min fluxes and error bars obtained with the gradient method. c. Relative contribution of error linked to N₂O concentration and wind vertical profiles.
val ranged from 1 to 9 ng N-N2O m-2 s-1 with a mean value of 4 ng N-N2O m-2 s-1 and a mean relative error of 55 %. Considering the low N2O emission rate, measurements were affected by instrumental TDLAS drift and it was essential to remove low frequency noise from the N2O data.

Intercomparison of flux measurement: effect of temporal and spatial variability

Temporal variability

To show the variations in diurnal N2O emissions, the hourly mean fluxes obtained with micrometeorological measurements over ten successive days are presented in figure 5a. In order to account for the N2O emission variation over the 10 d, each hourly measurement was normalized by the mean daily flux. Also, to obtain mean hourly fluxes symmetrical around the time axis each flux was centered by their daily mean. Each relative flux \( F_{i,j}' \) may be then written as follows:

\[
F_{i,j}' = \frac{F_{i,j} - m_j}{m_j}
\]

where \( F_{i,j} \) is the hourly flux from the \( i \)th hour of the day and \( m_j \) is the mean flux from the \( j \)th day of the 10-d period.

In figure 5a the relative hourly means obtained from the 10 d \( \left( \bar{F}_i = \sum_{j=1}^{10} F_{i,j}' \right) \) are shown. The error bars indicate the standard deviation observed over the 10 days. Except for 3:00 UT when standard deviation was particularly high, a diurnal tendency was observed with maximum N2O emissions between 9:00 and 16:00 UT.

In order to examine the possibility of using flux measurements from chambers as a mean daily flux estimate, chamber measurements were compared with the mean daily flux obtained with the micrometeorological methods (fig 5b). For the automatic chambers, the fluxes were averaged for

![Figure 5a](image-url) - Temporal flux variabilities.

a. Relative diurnal N2O fluxes observed by micrometeorological methods (see text for definition of Y-axis data).

b. Mean daily flux estimates: from 2 h 15-min integration time, corresponding to the 16 manual chambers (grey), from 6-h integration time, corresponding to the two automatic chambers (black) and from 24-h integration time, corresponding to micrometeorological fluxes (white). For these later error bars indicate the standard deviation obtained from 24 hourly measurements.

![Figure 5b](image-url) - Flux measurements.
night and day. In figure 5b, the error bars on the micrometeorological data are the standard deviations for the 24 hourly fluxes of the corresponding day. Their amplitudes have the same order of magnitude as their mean values. Chamber measurements are within the error bars of micrometeorological fluxes. As could be expected, the mean values obtained with the automatic chambers and for the 6-h integration time were more representative of the daily mean emission obtained by the micrometeorological technique compared to the 2 h 15-min integration time measurements of the manual chambers. Fluxes obtained with manual chambers with only 2 h 15-min integration time during midday (except for day 306) are generally greater than the fluxes obtained with other techniques. This result may be expected with respect to the diurnal N$_2$O emission variabilities (fig 5a). Therefore, it is important to monitor the emissions at different times in the day. Nevertheless, such a comparison would be more significant with a larger magnitude of emissions.

**Spatial variability**

The manual chambers at 16 different locations on the field were classified according to their respective flux magnitude. Each value of flux $F_{ij}$ flux from the $i$th chamber was 'centered' and 'normalized' by the mean daily flux $m_j$ obtained from the 16 chambers according to equation [12]. Figure 6a shows the mean and the standard deviation of those relative emissions, obtained from 10 d of measurements. For each chamber a rank was established, error bars are the standard deviations of the 10 d of measurement. This figure suggests that spatial variability is structured with emissions from some chambers always larger than others. This is particularly the case with chamber 14 characterized by large emissions, which suggests a particular set of soil conditions favourable to N$_2$O production; conversely chamber 4 was characterized by persistent low emissions (small error bars).

Figure 6b shows the N$_2$O fluxes measured by the various methods over the same time intervals.
over the ten consecutive days 298–307. Micrometeorological data were averaged over the time intervals when the chambers were covered. The fluxes obtained with the various methods at this temporal scale were closer than for the comparison at daily scale (fig 5b). Fluxes measured by the eddy correlation method are labelled ‘ec’. The spatial variability of the manual chambers was quantified by the standard deviations obtained with the 16 enclosures. This variability is high when compared to the methodological uncertainty (≤ 7 %). Depending on the day, CV ranged between 45 and 150 %. For the micrometeorological gradient method, the mean errors bars obtained theoretically with equation [6] over a 1-h integrating time according to equation [8] were in a similar range: 30–106 %. As explained previously, the error in the gradient method results from the instrumental resolution limits but also for a large part from non-ideal emission conditions with spatially heterogeneous sources. Errors bars obtained by the eddy correlation method (equation [10]) are smaller and vary from 13 to 26 % for 1-h integration times. They include only instrumental accuracy. The spatial variability of the soil NO3- and NH4+ contents obtained with the six samples was of the same order of magnitude as that for N2O emission (fig 2c) suggesting that the heterogeneity in substrates distribution could have induced the existence of hot spots demonstrated by the ranking of chamber sites (fig 6a).

CONCLUSION

The objective of this study was to compare measurements of N2O fluxes under field conditions, which required integrating fluxes representative of different spatial and time scales. Measurements of N2O fluxes from an agricultural soil were obtained by different methods at a common site. Three methods, which measured fluxes over different areas (0.18 m²–105 m²), were used to quantify emission fluxes, and to assess their spatial and temporal representativity.

Fluxes measured using two automated chambers, varied from -0.7 to 14 ng N-N2O m⁻² s⁻¹ during the campaign.

Fluxes measured with 16 manual chambers, varied from 0.6 to 35 ng N-N2O m⁻² s⁻¹, with a coefficient of variation of ~65 % and indicated large-scale (10–100 m) heterogeneity.

Hourly fluxes measured by micrometeorological methods, varied from -5 to 50 ng N-N2O m⁻² s⁻¹ with a relative uncertainty of ~40 % for the gradient method. These measurements indicated large variations with time during the day with a CV of near 100 % correlated with soil temperatures. This study confirmed that the micrometeorological techniques could be employed even for weak N2O emissions. The magnitude of the uncertainty of N2O fluxes was fairly large, because N2O sources were heterogeneous and measurements of small differences were made in a background concentration. Nevertheless, greater relative accuracy may be expected with larger emissions.

Despite weak N2O emissions (around 6 ng N-N2O m⁻² s⁻¹ on average), the correspondence between the different techniques was satisfactory. This study showed the temporal and the spatial variabilities in N2O emissions and showed the necessity of characterizing soil heterogeneity properties in terms of soil water content (which controls the degree of anoxia) and the nitrogen availability to help interpret the N2O flux variations. These results should be confirmed by measurements using a wide range of emission conditions.

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REFERENCES


APPENDIX A. UNCERTAINTY OF THE EDDY CORRELATION METHOD

This assessment includes only instrumental limitation. Given the flux expression it is possible to estimate the flux error according to the resolution of each sensor. The error on the flux is expressed as:

\[ dF^2 = \frac{1}{N^2} \sum_{i=1}^{N} \left[ \left( \frac{\partial F}{\partial c_i} \right)^2 (dc_i)^2 + \left( \frac{\partial F}{\partial w_i} \right)^2 (dw_i)^2 \right] \]

with

\[ F = \frac{1}{N} \sum_{i=1}^{N} F_i = \frac{1}{N} \sum_{i=1}^{N} w_i c_i \]

The following equality was obtained

\[ \frac{1}{N} \sum_{i=1}^{N} \left( \frac{\partial F}{\partial c_i} \right)^2 = \sigma_c^2 \] \[ \text{and} \quad \frac{1}{N} \sum_{i=1}^{N} \left( \frac{\partial F}{\partial w_i} \right)^2 = \sigma_w^2 \]

and

\[ \Delta F = \frac{1}{\sqrt{N}} \sqrt{\sigma_c^2 \cdot (\Delta c)^2 + \sigma_w^2 \cdot (\Delta w)^2} \]