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Nitrogen oxides and ozone fluxes from an oilseed-rape management cycle: the influence of cattle slurry application

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Abstract. This study reports NO, NO₂ and O₃ mixing ratios and flux measurements using the eddy covariance method during a 7-month period over an oilseed-rape field, spanning an organic and a mineral fertilisation event. Cumulated NO emissions during the whole period were in agreement with previous studies and showed quite low emissions of $0.26 \text{ kg N ha}^{-1}$ with an emission factor of 0.27 %, estimated as the ratio between total N emitted in the form of NO and total N input. The NO emissions were higher following organic fertilisation in August due to conditions favouring nitrification (soil water content around 20% and high temperatures), while mineral fertilisation in February did not result in high emissions. The ozone deposition velocity increased significantly after organic fertilisation. The analysis of the chemical and turbulent transport times showed that reactions between NO, NO2 and O3 below the measurement height occurred constantly throughout the 7-month period. Following organic fertilisation, the NO ground fluxes were 30 % larger than the NO fluxes at the measurement height (3.2 m), while the NO₂ fluxes switched from deposition to emission during certain periods, being negative at the surface and positive at the measurement height. This phenomenon of "apparent NO₂ emissions" appears to be significant during strong NO emissions and high O₃ ambient mixing ratios, even on a bare soil during August.

1 Introduction

Agricultural soils represent an important source of atmospheric nitric oxide (NO), especially in highly fertilised regions (Oikawa et al., 2015). Global estimates of total NO_x $(NO + NO_2)$ emissions from soils range between 4 and 21 Tg N yr^{-1} , which represents between 10 and 15 % of the global NO_x budget (Davidson and Kingerlee, 1997; IPCC, 2001; Yienger and Levy, 1995). NO_x inventories are subject to error in magnitude and especially in spatial distributions (Martin et al., 2003), which can be constrained by satellite observations and range around 30% at the global scale (Toenges-Schuller et al., 2006). NO_x emissions are of considerable interest also for atmospheric photochemistry, and acting as ozone (O₃) precursors, they indirectly have an impact on climate. O₃ is indeed an important greenhouse gas, contributing to 25 % of the anthropogenic net radiative forcing (IPCC, 2007).

 NO_x , and especially NO_2 , are toxic gases for humans, exposure to which increases risks of various respiratory diseases. The World Health Organization gives guidelines for NO_2 exposure limits, both annual means ($40 \,\mu g \,m^{-3}$) and 1 h mean ($200 \,\mu g \,m^{-3}$) (Prüss-Üstün et al., 2016). For O_3 , only a short-term threshold is given ($100 \,\mu g \,m^{-3}$ for the 8 h mean) because there are fewer studies on long-term exposure. These thresholds are established both in epidemiological and toxicological studies on humans and animals. Similarly, nitrogen deposition leads to serious adverse effects on the ecosystem (eutrophication, biodiversity erosion and acidification being

the most serious ones), while O_3 has a direct adverse effect on plant health through oxidation of photosynthesis pathways and direct tissue destruction above large thresholds. For nitrogen, the concept of critical load has been developed which gives the amount of nitrogen deposition above which an ecosystem is impacted. These critical loads range from $5 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ for sensitive habitats to $20 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ for less sensitive ones (APIS, 2016). For these reasons, national and international authorities regulate atmospheric levels of these pollutants.

 NO_x emissions from soils are primarily the by-products of nitrification and denitrification processes and the chemical decomposition of HNO₂ (Laville et al., 2005; Meixner, 1997; Remde et al., 1989). Many authors emphasise that for most agricultural soils, nitrification is the dominant process of NO emissions (Bollmann et al., 1999; Dunfield and Knowles, 1999; Godde and Conrad, 2000). Organic and mineral fertilisers, rich in ammonium, increase NO emissions both by stimulating NO production by nitrification and by decreasing NO consumption.

There is a significant knowledge gap in understanding NO_x exchanges between agricultural ecosystems and the atmosphere, partly due to a lack of direct measurements over long periods. NO emissions by soils can either be measured by dynamic chambers (Breuninger et al., 2012; Laville et al., 2009, 2011; Pape et al., 2009), aerodynamic gradient (Kramm et al., 1991), or eddy covariance methods (Rummel et al., 2002; Stella et al., 2013a). Each method has its drawbacks and challenges. The dynamic chamber method may change the surface exchange parameters (Pape et al., 2009), and modify the fluxes due to fast reactions between the triad O₃-NO-NO₂, but thoroughly designed Teflon chambers can overcome this problem (Skiba et al., 2009). The aerodynamic gradient method (AGM) is a well-established method applicable to water-soluble compounds such as NH₃ (Milford et al., 2009), but has several biases of which flux divergence due to chemical reactions is the most limiting for NO-NO₂-O₃ (Duyzer et al., 1995; Kramm et al., 1991; Loubet et al., 2013). Non-stationarity and integration time are also limiting (Lenschow et al., 1994; Stella et al., 2012). The eddy covariance method is adapted for measuring NO fluxes. It is however also vulnerable to flux divergence issues due to NO–NO₂–O₃ chemical reactions. It is therefore essential to measure the fluxes and mixing ratios of the three compounds together.

The eddy covariance (EC) method is the state-of-the-art flux measurement method for energy and CO_2 fluxes (Baldocchi, 2003), and due to the development of new analysers such as fast chemiluminescence, quantum cascade lasers, absorption spectroscopy, or proton time-of-flight mass spectrometers (Ammann et al., 2012; Brodeur et al., 2009; Ferrara et al., 2012; Li et al., 2013; Müller et al., 2010; Park et al., 2014; Peltola et al., 2014; Sintermann et al., 2011; Stella et al., 2013a; Wolfe et al., 2009) it can currently be applied to several other trace gases. The main advantage of

the EC method is that it is a "direct" measurement of the flux at a given height, which depends on fewer assumptions than the AGM, namely the Reynolds averaging and ergodicity hypothesis requiring that "the averaging time must be much larger than the timescales of variation of the air velocity" (Corrsin, 1975, see also Kaimal and Finnigan, 1994). This method has been successfully applied for measuring NO fluxes in a limited number of studies (Eugster and Hesterberg, 1996; Lee et al., 2015; Marr et al., 2013; Min et al., 2014; Rummel et al., 2002; Stella et al., 2013a). The main difficulties of EC measurements are the losses that appear at high frequencies due to adsorption of the gas to the tubing system, which depends also on the size of the absorption cell (Eugster and Senn, 1995) and differential advection caused by the radial variation of the mean velocity and simultaneous radial diffusion of the sample gas (Lenschow and Raupach, 1991). Moreover, since NO₂-to-NO photolytic converters typically applied in combination with chemiluminescence analysers have a conversion efficiency below 100 %, measuring both NO and NO₂ with such a converter remains a challenge that requires the use of two analysers simultaneously (Lee et al., 2015).

Due to these limitations, simultaneous measurements of NO, NO₂ and O₃ fluxes by EC have hence seldom been made. To our knowledge, only a few studies report such measurements and none over an arable crop. There is therefore a gap in knowledge as to whether the reactions between NO, NO₂ and O_3 significantly influence the fluxes above crops and how nitrogen application modifies these fluxes and their interactions. Eugster and Senn (1995) report NO2 fluxes by EC and analyse the errors of the method. Most studies conducted over forests show moderate to large in-canopy reactions: Andreae et al. (2002) report comprehensive flux measurements in the Amazonian forest showing evidence of within-forest cycling of the nitrogen oxides emitted from the soil. Horii et al. (2004) report NO_x and O₃ fluxes over a temperate deciduous forest showing consistent NO_x deposition. Geddes and Murphy (2014) report such measurements over two mixed hardwood forests in North America, under a very low NO concentrations regime, which show mainly NO_x deposition with evidence of chemical reactions in the canopy. Min et al. (2014) report such flux measurements over ponderosa pines which shows evidence of within-canopy chemical removal of NO_x . Ammann et al. (2012) report total reactive nitrogen fluxes by EC above grassland which compared well with dynamic chamber NO and NO2 fluxes during periods with no NH₃ emissions. Lee et al. (2015) and Marr et al. (2013) report fluxes of NO and NO₂ over urban areas which differ in their comparison with national emissions inventories: while Lee et al. (2015) found fluxes 80 % higher than national inventories, the second study found similar fluxes but with large disparities at the local scale.

In this study we are addressing the following questions: (1) is the EC method suitable for quantifying the seasonal dynamics and diurnal cycles of the NO, NO_2 and O_3 fluxes

above a crop rotation? (2) How are organic and mineral fertilisations affecting these fluxes and their dynamics? (3) To what extent are the chemical reactions between NO, NO_2 and O_3 modifying the fluxes above the ground? And finally, (4) why is O_3 deposition increasing following organic fertilisation? Is that a consequence of interactions with NO emissions?

To answer these questions we report measurements of NO, NO_2 and O_3 fluxes by EC using a system similar to Lee et al. (2015) for 1 month following slurry spreading over a bare soil at the FR-GRI FLUXNET and ICOS site (Loubet et al., 2011). The NO and O_3 fluxes were measured over an additional 6-month period to study the seasonality of these fluxes and to measure the fluxes following application of mineral fertiliser.

2 Materials and methods

2.1 Site description and management

The experiment took place in a 19 ha field located at Grignon (48°51' N, 1°58' E), 40 km west of Paris (France) and lasted more than 7 months from 7 August 2012 to 13 March 2013. The field was surrounded by heavy traffic roads on the east, south and southwest. The field belongs to a large farm (buildings at around 450 m to the southwest) with around 210 dairy cows, 500 sheep, and a production of approximately 900 lambs. The terrain has a gentle slope of $\sim 1 \%$ and the mean annual temperature and precipitation were 10.9 °C and 575 mm between 2005 and 2013. The main wind directions are northwest during clear days and southwest during cloudy and rainy days. The soil type is a Luvisol or loamy clay (25 % clay, 70 % silt, 5 % sand in the top 15 cm). The soil organic carbon content was $\sim 20 \text{ g C kg}^{-1}$, pH (in water) = 7.6, and bulk soil density was 1.3 g m^{-3} , in agreement with previous measurements on the same site (Laville et al., 2009, 2011; Loubet et al., 2011). High pH values are common in soils over calcareous layers and with high fine fraction content (clay and silt) as is the one of the Grignon site. Indeed, alkalinity fosters the nitrification process and this range of pH is optimum for it to occur (e.g. Nieder and Benbi, 2008). A detailed description of the site can be found in Laville et al. (2009, 2011) and Loubet et al. (2011).

The field was cultivated with winter wheat (a mix of Atlass and Premio species), which was harvested on 3 August 2012 (16.7 Mg ha⁻¹ of dry matter). Cattle slurry was applied on the field with a trailing hose from 18 to 19 August 2012, at a rate of 42 kg N ha^{-1} of which 78 % was ammonium (NH₄⁺). The slurry had a very low dry matter content of 3.2 % and a C / N ratio of 15.7. The total C applied was 666 kg C ha⁻¹. A gentle tillage was performed on 29 August 2012 to incorporate the crop and slurry residues and prepare the soil for oilseed rape sowing (variety Adriana) at a density of 35 plants per square metre. The crop developed slowly during the winter with a dry matter above ground (leaf area index) of 37 g m^{-2} (0.65 m² m⁻²) on 25 October 2012 and 104 g m⁻² (0.7 m² m⁻²) on the 18 February 2013. The canopy height stayed below 10 cm during the whole winter. Ammonium nitrate pellets were applied on the oilseed-rape field on 20 February 2013 at a rate of 54 kg N ha⁻¹. Two selective herbicides were applied on the 2 (Springbok: 200 g L⁻¹ of metazachlor; 200 g L⁻¹ of DMTA-P at 3 L ha⁻¹) and 31 October 2012 (Devin/cycloxydim: 100 g L⁻¹ at 1 L ha⁻¹) which only destroyed the weeds. In December 2012 slug-repellent pellets were applied.

2.2 Micrometeorological and ancillary measurements

Meteorological measurements included wind speed, wind direction, air and soil temperatures and humidity as well as rainfall, global, net and photosynthetic active radiation. The meteorological measurements were performed on a mast (3.17 m high) near the centre of the field and close to the flux measurement site (Fig. 1). Soil was sampled approximately once a month for water content, total nitrogen and mineral nitrogen analysis. Measurements are described in Loubet et al. (2011) and will not be detailed here.

A simplified sketch of the EC measurement system is shown in Fig. 1. Three-dimensional wind and temperature fluctuations were measured near the centre of the field at 3.17 m above ground by a sonic anemometer (Gill R3 3axis anemometer, Gill Instruments Limited, UK). A fast response open-path CO₂ / H₂O infrared gas analyser (IRGA LI-7500A, LI-COR, USA) installed at a lateral distance of around 0.2 m to the sonic path measured CO₂ and H₂O fluctuations. O₃ mixing ratios were measured by a highfrequency, dry chemiluminescence O₃ detector (NOAA, USA) and its Teflon PFA inlet tube (length = 2.8 m, internal diameter = 0.32 mm) was positioned in-between the sonic path and the IRGA at a lower height. The high-frequency signals were recorded at 20 Hz by a Labview program developed in the laboratory. In accordance with Lee et al. (2015), highfrequency (10 Hz) time series of NO and NO₂ were determined by two fast-response and closed-path chemiluminescence NO analysers (CLD 780TR, EcoPhysics, Switzerland), one being coupled to a photolytic converter (blue light converter, BLC, Droplet Measurement Technologies Inc, USA) for the detection of NO_2 (see Fig. 1). The horizontal separation of the trace gas inlets from the sonic path was 20 cm. Air was sampled through two heated and shaded PFA tubes with a length of 20 m and an inner diameter of 9.55 mm. The first CLD was used for measuring NO and the second one connected to the BLC was used for detecting NO₂. Conversion efficiencies for NO₂ to NO of around 30% were achieved. The high-frequency signals of NO, NO₂ and O₃ were calibrated with mixing ratios measured at 30 min time resolution by slow-response analysers (Thermo Scientific, Waltham, USA) (Fig. 1). These instruments were calibrated every 3 to 6 weeks using the gas-phase titration (GPT) method and



Figure 1. (a) Simplified map of the FR-GRI field site showing the mast and surrounding roads. The colours correspond to elevation. (b) Simplified sketch of the instrumental setup to measure EC fluxes. Gill R3 is the ultrasonic anemometer, Li7500 is the open-path infrared CO₂ and H₂O gas analyser, the rain cup is the air sampler for NO and NO₂ detection. CLD780-TR NO and NO_x are the fast-response NO analysers (Ecophysics) operating in parallel, one connected to a BLC measuring NO + α NO₂. The NO, NO₂ and O₃ slow analysers (Thermo Scientific, Waltham, USA) are placed behind a Teflon pump ensuring atmospheric pressure at the inlet. A large pallet pump ensured a flow rate of 80 NL min⁻¹ in the heated inlet line. Teflon filters (1 µm) were installed at the front of the instrument inlets (purple cylinders).

a 17 ppm NO standard (Air Liquide, FR). The fetch of the field site extended at least to 150 m in all directions and a footprint analysis showed that 90% of the time the entire field was in the footprint during neutral and moderately stable or unstable conditions (Loubet et al., 2011). NO and O₃ fast sensors were functioning during the whole campaign (7 August 2012 to 13 March 2013) together with NO, NO₂ and O₃ slow-response analysers and the meteorological station. High-frequency NO₂ measurement was performed from 14 August to 30 September 2012. In this study we focus on two periods: (1) from 14 to 29 August 2012 during which all fluxes were measured and NO fluxes were the highest, in order to investigate the interactions between the fluxes and mixing ratios of the NO–NO₂–O₃ triad, and (2) over the whole period for NO fluxes analysis.

2.3 Eddy covariance flux computations

The turbulent fluxes were computed as the covariance between the fluctuations of the scalar of interest and the vertical component of the wind. As the EC method and its theoretical background are described in the literature – e.g. Foken (2008) – details will not be provided here.

For closed-path sensors (NO, NO₂ and O₃), the lag time between w' and the dry mole fraction χ had to be determined. This was done by searching for the maximum of the covariance function

 $(\chi \overline{w'(t)'(t-\text{lag})}.$

The lag for NO was 3.1 s [2.4-3.65 s] (Q_{50} [$Q_{25}-Q_{75}$]), for NO₂ it was 4.0 s [3.65-4.55 s] and for O₃ it was 2.9 s [2.5-3.25 s]. The lag was filtered for outliers (points outside of median lag ± standard deviation were considered as outliers)

and the covariance was computed as the value of the covariance function at the filtered lag.

As fast-response sensors for NO, NO₂ and O₃ were not absolute, the fluxes were computed following the ratio method for O₃ described by Muller et al. (2010), and in accordance with Lee et al. (2015) for NO and NO₂:

$$\chi F_{\rm O_3} = \frac{\overline{\chi_{\rm O_3}}}{V_{\rm dry}} \frac{\overline{w'O_3'}}{\overline{O_3}},\tag{1}$$

$$F_{\rm NO} = \frac{\overline{w' \rm NO'}}{S_{\rm NO} V_{\rm drv}},\tag{2}$$

$$F_{\rm NO_2} = \frac{1}{\alpha V_{\rm dry}} \left(\frac{\overline{w' \rm NO_x'}}{S_{\rm NO_x}} - \frac{\overline{w' \rm NO'}}{S_{\rm NO}} \right), \tag{3}$$

where O₃ (in mV), NO and NO_x (in counts s⁻¹) are the uncalibrated fast signals, $\overline{\chi_{O_3}}$ is the 30 min average of the slow-sensor reference O₃ mixing ratio (in ppb), while S_{NO} and S_{NO_2} are the sensitivity of the analysers (in counts s⁻¹ ppb⁻¹). The α is the blue light converter conversion efficiency, and V_{dry} is the molar volume of dry air (in m³ mol⁻¹). All fluxes (momentum, heat, CO₂, H₂O, NO, NO₂, O₃) were computed by the EddyPro softwareversion 5 (http://www.licor.com/eddypro) and final flux data were averaged for 30 min intervals. Evaluation methodologies from the CarboEurope project were applied – see Aubinet et al. (2000) and Loubet et al. (2011).

2.4 Spectral corrections and flux uncertainties

Spectral attenuation of the flux is due to differential transport time of the compound in the tube and interaction with tube walls and filter surfaces (Massman and Ibrom, 2008).



Figure 2. Hourly averaged high-frequency loss correction factors for O_3 , NO and NO_x over the 15 August–7 September 2012 period determined with the in situ ogive method. The corrected flux equals the measured flux multiplied by the correction factor. Black bars are medians, boxes show the interquartile, error bars show the minimum and maximum of the whisker and empty dots shows the outliers.

We tend to minimise this effect by ensuring a large flow rate in the tubes with a Reynolds number well above the critical threshold for turbulence – see Lenschow and Raupach (1991) – as well as heating the tubes to around 5 °C above ambient temperature. The residence time of the air samples inside the tubing was around 1 s, ensuring low chemical conversions, and the Reynolds number was 11 700, hence largely in the turbulent range (Re > 4000). However, water vapour interaction is still expected, and sensor separation also generates high-frequency losses.

The NO, NO₂ and O₃ random instrument noises were estimated as the 1σ random uncertainty of the signals as in Lenschow et al. (2000), Langford et al. (2015) and Mauder et al. (2008). This is assumed to be "white noise" and hence uncorrelated with itself apart from at lag = 0 s. It is therefore estimated as the difference between the autocorrelation at lag = 0 s and at lag = ± 0.05 s. The flux random uncertainty was itself evaluated as the covariance detection limit. It was determined as the root mean square error of the covariance function over 60 s periods at lag times well away from the position of the time lag. In practice, these were taken at lags larger than 120 s as absolute values, as proposed by Langford et al. (2015).

2.5 Chemical reactions, timescales and flux divergence

Chemical reactions between NO, NO₂ and O₃ are important to consider when interpreting the measured fluxes as they can affect the fluxes above the ground. A common way to determine whether these reactions may indeed affect the flux is through comparison of chemical and transport timescales. Details of the reactions rates, timescales and flux divergence calculations are given in Sects. S1–S3 in the Supplement.

3 Results and discussion

3.1 Quality check and uncertainties in NO, NO₂ and O₃ flux measurements

 NO_x and O_3 half-hourly fluxes were filtered by the quality check test included in EddyPro (http://www.licor.com/ eddypro), according to the 0–1–2 labelling proposed by Mauder and Foken (2006), which includes tests for stationarity and for well-developed turbulence. As recommended in the framework of the CarboEurope project, we discarded fluxes with a quality check index value of 2. This led to keeping 74, 84 and 76 % of the records for NO, O₃ and NO₂, respectively. The total records of NO and O₃ half-hourly fluxes were 11 329 (from 7 August 2012 to 13 March 2013), while for NO₂ they were 2257 (during the period 14 August to 30 September 2012).

The largest systematic uncertainties were the high-frequency losses, which were estimated with the in-situ ogive method (Ammann et al., 2006), and amounted to 10 % for O₃, 20 % for NO and 30 % for NO₂ on average over the August–September period (when all fluxes were measured, see Fig. 2). As a bias, they can be corrected for, as in the following sections of this paper.

The second largest uncertainties were the random uncertainties which were lower than 20 % in most cases for O_3 , NO (and similar to H_2O) and around 30 to 40 % for NO₂ (Fig. 3). For NO and NO₂ the random uncertainties peaked during the morning traffic hour around 06:00–08:00 UTC, which is explained by the non-stationarity generated by the local traffic on the mixing ratios. Hence overall the EC method proved to be usable for measuring NO fluxes over part of the season with an overall uncertainty similar to H_2O . A higher random uncertainty was found for NO₂ fluxes which were smaller than NO fluxes and with a relatively low conversion ratio from NO₂ to NO (30 %).



Figure 3. Daily variations of the ratio of the random uncertainty to the flux for H_2O , O_3 , NO and NO_x during August 2012 (15 August to 9 September). Black bars are medians, boxes show the interquartile, error bars show the minimum and maximum of the whisker and empty dots shows the outliers.

3.2 Meteorological conditions

Daily averages of the air temperature decreased during the measurement period, starting from about 20 °C in summer and reaching minima around -5 °C from December to March. Daily averages of global radiation decreased from $250 \text{ W} \text{ m}^{-2}$ in August to around $0 \text{ W} \text{ m}^{-2}$ in December, back to around $150 \,\mathrm{W}\,\mathrm{m}^{-2}$ by the end of March. The daily average of the relative humidity was around 65 % in August and September, and it increased to about 85 % for the rest of the period (Fig. 4). The wettest period was between October and November, and cumulative rain was 319 mm over the 7-month period, which is quite high. The prevailing wind direction was southwest while the most intense winds were observed from north and south (Fig. S2 in the Supplement). Figure S2 also shows that wind regimes were quite different in summer and winter: prevailing wind directions during August and February were from the southwest and northeast, respectively. Soil water content (SWC) ranged between 20 and 40 % (volume) (Fig. 4), with a long period between October and January with values around 28%, and increased further in January to 35 %, with sharp decrease during some periods.

3.3 Seasonal dynamics and diurnal cycles of the NO, NO₂ and O₃ fluxes above the crop rotation

3.3.1 Seasonal dynamics of NO–NO₂–O₃ mixing ratios

Average daily NO, NO₂ and O₃ mixing ratios were 3.6, 6.9 and 24.8 ppb, respectively. The NO and NO₂ mixing ratios were higher when winds blew from the east (from the direction of Paris), while O₃ showed the opposite behaviour, which can be explained by depletion of O₃ by NO sources from the surrounding traffic (as shown in Fig. S2) and by reactions (Sects. S1–S2). Daily NO₂ / NO_x ratios were on average 66 %, which is typical for traffic and urban pollution (Carslaw, 2005; Minoura and Ito, 2010), and ranged from 4 to 93 % during the entire period. The NO₂ mixing ratios were significantly higher (Student *t* test *p* value lower than 8×10^{-11}) than the NO mixing ratios in August and early September, end of January and mid-February, and end of March. During sporadic episodes, NO peaks were of the same order or even higher than NO₂ peaks (Fig. 4).

3.3.2 Seasonal dynamics of NO, NO₂ and O₃ fluxes

The daily averaged NO fluxes were very small, except during a period of strong emission following organic fertilisation over two days in August (18-19 August 2012), with maximum daily average fluxes of around 1.5 nmol $m^{-2} s^{-1}$ (Fig. 4). Other emission episodes, including mineral fertilisation in February (20 February 2013), were characterised by mean daily fluxes below 0.5 nmol m⁻² s⁻¹. The NO fluxes were slightly negative for some events (Q_{25} , Q_{50} and Q_{75} equal to -0.013, 0.031 and 0.11 nmol m⁻² s⁻¹, Fig. S3). The O₃ fluxes ranged between -13.8 and $0 \text{ nmol m}^{-2} \text{ s}^{-1}$, and averaged to $-3.12 \text{ nmol m}^{-2} \text{ s}^{-1}$. The largest O₃ deposition fluxes were observed following organic fertilisation in August, and were correlated with the highest NO emissions. This period also corresponded to large daily O_3 mixing ratios (Fig. 4). The NO₂ fluxes were only measured during the first one and a half months (14 August to 30 September 2012) and were mostly negative (indicating deposition), except during the first week following organic fertilisation (Q_{25} , Q_{50} and Q_{75} equal -0.11, -0.07 and $0.08 \text{ nmol m}^{-2} \text{ s}^{-1}$) (Fig. S3). O₃ fluxes were in the same range of magnitude, typically between -20 and $0 \text{ nmol m}^{-2} \text{ s}^{-1}$, as those reported by previous studies at the same site (Stella et al., 2013b, 2011b; Tuzet et al., 2011) and in the literature over various canopies such as grassland (Stella et al., 2013a), barley (Gerosa et al., 2004), potato field (Coyle et al., 2009) or forests (Fares et al., 2010; Gerosa et al., 2005), although O₃ flux magnitude is sharply dependent on local O3 mixing ratio. We found similar magnitudes of ozone fluxes in August and September as



Figure 4. Meteorological and soil conditions (daily averages, sums for rainfall), NO, NO₂ and O₃ mixing ratios and fluxes during the entire measurement period from 7 August 2012 to 13 March 2013 at the Grignon field site. Averages for night-time and daytime are also given as dotted lines. R_g is the global radiation, T_a and T_g the air and ground temperature, SWC the soil water content, we the wind speed, RH the air relative humidity.

those reported by Stella et al. (2013a) over a meadow during the summer. We also found similar nocturnal O₃ deposition velocity as found by Stella et al. (2011a) over bare soil during summer, but with a higher daily maximum (0.8 cm s^{-1}) instead of $0.5-0.6 \,\mathrm{cm}\,\mathrm{s}^{-1}$). Seasonal and daily dynamics of O₃ deposition velocity are shown in Fig. 5. We further find a similar midday magnitude as Stella et al. (2011a) found in April with wetter soils. Night-time ozone deposition velocity did not go lower than around 0.2 cm s^{-1} in our study, as also found by Zhu et al. (2015) over a growing wheat in China, Stella et al. (2011a) over bare soil in summer, and Lamaud et al. (2009) over maize. These authors as well as Huang et al. (2016) clearly show that this is due to non-stomatal deposition being primarily driven by u_* which does not reach zero at night during these periods. We can hence conclude that we found consistent ozone deposition in August and September compared to other studies at that site or in other geographical areas. When compared to previous years at the same site the deposition velocity measured during the winter in this study was clearly smaller. We interpret this as being primarily due to u_* being smaller that winter compared to other winters, as well as due to a bad development of the winter crop due to soil drought in September (SWC = 20% in the 15 cm horizon).

25

-8

40

0

[qdd] ⁶ 20

0 -10

្ច 15

3.3.3 Comparison of ozone fluxes to the Stella et al. (2011a) parameterisation over soil

²s-1

In order to compare to previous studies of ozone deposition onto bare soil on the same site, we have calculated the soil surface resistance as in Stella et al. (2011a) and deduced the ozone deposition velocity as $V_{\rm dO_3} = (R_{\rm soilO_3} + R_{\rm bO_3} + R_{\rm a}(z_{\rm ref}))^{-1}$. In this way, we can compare the two studies while excluding any confounding factors (roughness and turbulent exchange intensity). We can see in Fig. 6a that the measured ozone deposition velocity during August follows the parameterisation of Stella et al. (2011a) most of the time except for some days including 18 and 19 August which correspond to slurry application and 24, 25, and 26 August, which follow a small rainfall. We also see an overestimation of the Stella parameterisation before the 18 August, which we interpret as being due to the straw and wheat residues being present on the ground before slurry incorporation. This comparison hence demonstrates that the ozone deposition was indeed increased slightly following slurry application and subsequently following rainfall. This may be either due to a physical reason (increased surface exchange on the soil due to tillage or humidity change due to slurry) or a chemical reason (surface reactivity changes due to added organic matter or volatile organic compound (VOC) emissions from the slurry). Figure 6b further shows that the main differences are observed for wet soils and relatively low



Figure 5. Seasonal changes of ozone deposition velocity V_{dO_3} and NO fluxes. Blue lines show median and grey area inter-quantiles.

temperatures (this is after rainfall) and to a lesser extent for drier and hotter situations (following slurry spreading).

3.3.4 Diurnal cycles of mixing ratios and fluxes over periods of interest

 O_3 mixing ratios exhibited a typical diurnal cycle that was governed by photochemistry and convective mixing within the boundary layer and from the free troposphere during daytime. It started to increase with sunlight around 07:00, and declined in the evening starting from 18:00 due to lack of photochemical formation in the absence of sunlight, as well as deposition and destruction with NO in this high NO_x emission area. In general, NO mixing ratios featured a marked peak in the early morning and remained high until around 13:00 UTC (Fig. 7b). During the early afternoon, the O₃ increase was correlated with the NO decrease. NO₂ mixing ratios showed a bi-modal diurnal cycle with its maxima in correspondence with morning and evening traffic peaks, i.e. around 06:00 and 19:00.

The NO fluxes also showed a diurnal cycle similar to the one of soil temperature with an emission peak around 00:00 (Fig. 7a and b). This suggests that NO emissions are related to nitrification, for which the emission rate is an exponential function of soil temperature (Henault et al., 2005). This was already shown for the Grignon soil by Laville et al. (2011). The fact that NO fluxes decrease earlier than soil temperature is most likely due to titration of NO by O_3 in the late morning and early afternoon, causing the NO emissions at the reference height to be reduced with respect to ground emissions.

After correction for chemical reactions the NO emissions diurnal cycle is indeed at a peak later in the day, more in phase with ground temperature (see Fig. 11). This is also indicated by the positive NO₂ flux observed during the same time of the day. The O₃ flux was mainly negative (deposition) and follows the diurnal dynamics of measured mixing ratios. In terms of deposition velocity, the ozone deposition velocity followed a clear diurnal cycle with a maximum during the day and a minimum at night. The measured NO₂ deposition velocity showed slightly negative values, but slightly positive ones when corrected for reactions with NO and O₃.

3.4 Influence of organic and mineral fertilisations on NO emissions

The NO flux averaged over the whole period was $0.09 \text{ nmol m}^{-2} \text{ s}^{-1}$ (mean), which is in the range of previous findings for the same site. Laville et al. (2011) and Loubet et al. (2011) reported yearly averaged NO fluxes varying between 0.07 and 0.15 nmol m⁻² s⁻¹ for 2007–2009. The NO flux distribution was shifted towards positive values after the organic fertilisation in August (Fig. S3), with the mean NO flux during the 2 weeks following the fertilisation (0.49 nmol m⁻² s⁻¹) being six times larger than the one for the whole period. For the same period, the ozone flux distribution was shifted towards more negative values. Figure S3 also shows that flux distributions after mineral fertilisation do not differ much from the ones relative to the whole period. During the 2 weeks following the February mineral fertilisation the NO flux increased less and was only 1.7 larger than



Figure 6. (a) Comparison of ozone deposition velocity from this study (black dots), and from the parameterisation of Stella et al. (2011a) (red line) based on surface temperature. (b) Response of ozone deposition velocity to surface humidity $RH(z_0)$ and surface temperature $T(z_0)$. Shown are data from this study and from the parameterisation of Stella et al. (2011a). The period covered is from 14 August to 6 September which is before and after slurry spreading and corresponds to Fig. S5.

over the whole period $(0.14 \text{ nmol m}^{-2} \text{ s}^{-1})$. These numbers are also in line with those reported following fertilisation on the same soil in the 2007–2009 period by Laville et al. (2011) and Loubet et al. (2011), which also showed some periods with slightly negative NO fluxes. Stella et al. (2012) measured a larger peak of NO emissions following slurry spreading, but only lasting 2 to 3 days, which was probably due to a drier soil in our study compared to Stella et al. (2012).

Following the slurry application, the NO emissions amounted to 0.1 kg N ha^{-1} , which represents 0.24 % of the applied nitrogen (42 kg N). Following the mineral fertilisation, the NO emissions amounted to $0.02 \text{ kg N ha}^{-1}$, which represents 0.037 % of the applied nitrogen (54 kg N). Over the whole period from August 2012 to March 2013, we evaluate a loss of $0.26 \text{ kg N ha}^{-1}$. With a total N input of 96 kg N ha⁻¹, this gives an estimate of the NO emission factor of 0.27 %, which is similar to values reported earlier for the same site (Laville et al., 2011) but one order of magnitude larger than the EMEP/IPCC default value of 0.04. Nevertheless, this is an average value calculated with the Tier 1 approach, which does not take into account correction factors depending on soil pH or fertiliser type. This more detailed approach, the Tier 2, has not been developed for NO (EEA, 2016).

The reasons for lower emissions following winter mineral fertilisation than following summer manure application are manifold. Even if the amount of applied nitrogen was similar for the two cases (42 and 54 kg N ha^{-1}), meteorological and soil conditions were much more favourable for nitrification in summer than in winter (Davidson, 1992; Williams and Fehsenfeld, 1991). Indeed, NO emissions from agricultural soils are primarily the by-products of nitrification, and this hypothesis was tested for the Grignon site by Laville et al. (2011). Nitrification is inhibited by low soil temperature and high water content that causes anoxia. Soil temperature was much lower in February than in August (2.5 compared to 20 °C on average). February was particularly humid, with a total precipitation of 10 mm, while in August no significant rain event occurred after the first week. In this period indeed, the soil was only humidified by the organic manure supply (on a 4.8 mm thick layer) that was applied on a dry soil. The soil water content at 5 cm depth in September 2012 was around 21 % in volume, while in February it was 33 % in volume. These two factors led to more favourable conditions for nitrification in August than in February.

3.5 Influence of surrounding roads on the measured fluxes and concentrations of the NO–NO₂–O₃ triad

Using the FIDES flux and concentration footprint model (Loubet et al., 2010) we evaluated the footprint of nearby roads. Overall the flux footprint from the nearby roads was smaller than 1% (which means that only 1% of the road emissions contributes to the flux at the mast) most of the time, but the concentration footprint reaches up to 10 % during some episodes, with separate roads contributing differently depending on the period (Fig. S1). Assuming a conservative emission of 250 mg km⁻¹ vehicle⁻¹ and an average vehicle count of 10 000 vehicles day⁻¹ (2010 counts, "Statistiques du département des Yvelines pour 2010" ranges between 5000 and 15 000), we evaluate that the roads contribute from 4 to 40 % to the measured fluxes. However, since vehicles emissions of NO_x have a sporadic nature, 10000 vehicles day⁻¹ means a maximum of ~ 1 vehicle every 2 s (if we consider, conservatively, that most of the traffic is condensed during 9h only). These vehicles are also moving at about 90 km h^{-1} (25 m s⁻¹), hence leading to a moving point source of NO_x . We therefore expect that the signal of this moving and sporadic source is not captured by the EC method, and would be filtered out by despiking and flux calculation procedures (Foken, 2008; Mahrt, 2010).

3.6 Chemical interactions: the NO–NO₂–O₃ triad and effect on the fluxes

In order to investigate the interactions between the fluxes and mixing ratios of the NO–NO₂–O₃ triad, we focus on the pe-



Figure 7. (a) Diurnal cycles of global irradiance and net radiation, air and soil temperatures, relative humidity and wetness index averaged over the three periods of interest at the Grignon field site. The shaded areas represent the interquartile range. (b) Diurnal cycles of NO, NO₂ and O₃ mixing ratios and fluxes as well as the deposition velocities of NO₂ and O₃, averaged over the three periods of interest at the Grignon field site. The deposition velocity of NO₂ and O₃ based on the fluxes accounting for chemical reactions above ground are also shown (V_{dO_3} and V_{dNO_2} corrected).



Figure 8. Meteorological variables and NO_x - O_3 mixing ratios and fluxes measured during the period 14 to 29 August 2012 at the Grignon field site. Ticks on the *x* axis correspond to midnight.

riod from 14 to 29 August 2012, during which all fluxes were measured and NO fluxes were the highest.

The 2 weeks following the organic manure application (from 18 to 19 August) are characterised by hot sunny days, with maximal global radiation above 800 W m^{-2} , except for 24 August when the only rain event occurred (Fig. 8). The period of 18 to 23 August was the warmest, with soil surface temperatures above 40 °C at noon during most days, while the air temperature decreased from around 35 to around 20 °C during the same period. The soil temperature at 5 cm depth followed the same trend, but with a lower daily maximum and a higher night-time minimum. Due to sensor breakdown the soil water content was not measured during this period. The small latent heat flux (LE) after the 19 August, $(17 \text{ W m}^{-2} \text{ on average between 19 and 31 August)}$ the large sensible heat flux (60 W m^{-2} on average) and radiation $(212 \,\mathrm{W}\,\mathrm{m}^{-2})$ on average) indicate that the soil humidity of the top soil layer was low. Hence, we assume that the SWC was probably similar to what was measured in September (around 20% in volume), which is ideal for nitrification to occur (Laville et al., 2011; Oswald et al., 2013).

The 18 August was the first day when NO emissions from the soil occurred. The emissions lasted around 2 weeks following the organic fertilisation (Fig. 4), during which the NO flux during daytime exceeded 0.5 nmol m⁻² s⁻¹, peaking around 00:00. The nocturnal NO flux usually decreased to zero, except for the night of 25 August, characterised by strong winds (Fig. 8). The maximum of the NO emissions was 2.7 nmol m⁻² s⁻¹ observed 6 days after fertilisation on 21 August.

The NO₂ flux daily pattern was different during the two weeks following organic manure application compared to the period before (Fig. 8). It was in general positive during the day and around zero at night during the period from 18 to 29 August, except for the night of 25 August when it was large and negative. Positive NO₂ fluxes might be explained by chemical reactions between NO and O₃ in the surface layer (De Arellano et al., 1993), which will be discussed in the next section.

The O₃ flux was also significantly higher (Student *t* test *p* value lower than 2×10^{-16}) following organic fertilisation (mean $-10.7 \text{ nmol m}^{-2} \text{ s}^{-1}$) than during the rest of the experimental campaign (mean $-3.1 \text{ nmol m}^{-2} \text{ s}^{-1}$) (Fig. S3). Since the mixing ratio of O₃ was quite variable during the campaign (Fig. 4), it is more interesting to look at the deposition velocity which underpins the surface exchange processes (Figs. 7b and 8). The median V_{dO_3} during the organic fertilisation event exceeded the median over the rest of the experimental campaign by a factor of 2. However, this increase in O₃ deposition velocity cannot be explained by reaction with soil-emitted NO alone as the O₃ flux is an order of magnitude larger than the NO flux.

Different pathways for the near-surface O_3 removal are likely: (i) photolysis of O_3 by ultraviolet light in the presence of water vapour forming OH radicals, (ii) gas phase re-



Figure 9. Diurnal cycles of the O₃ penetration depth in the soil (Δ_{dryO_3}), the aerodynamic ($R_a(z_{ref})$), boundary layer (R_{bO_3}) and soil resistances (R_{soilO_3}) for O₃, the chemical reaction time τ_{chem} and transport time τ_{trans} , the chemical reaction rates for NO₂ photolysis J_{NO_2} and NO depletion by O₃ ($k_r \times [O_3]$), and the Damköhler number (Da), averaged over the periods of interest at the Grignon field site. The shaded areas represent the interquartile range.

actions with reactive VOCs and (iii) heterogeneous reactions with the soil or with molecules adsorbed on soil.

The NO mixing ratio was well correlated with the NO flux, with a normal correlation coefficient of 40% for the two weeks following the organic fertilisation (excluding 24-25 August), while it was only 2% for the 7-month period. This suggests that, following fertilisation, the ambient NO levels were mainly due to local emissions. The NO₂ mixing ratio was less correlated with the NO₂ flux, suggesting that NO₂ levels were more related to advection from surrounding road traffic than from local emissions. Indeed, both NO and NO₂ are emitted from road traffic and urban pollution, but the NO₂ component quickly becomes prevalent as the plume is advected, especially in the presence of high O₃ levels, as in our case (Carslaw, 2005; Minoura and Ito, 2010). The minimum night-time mixing ratio is mainly controlled by night-time wind velocity: the higher the night-time velocity, the higher the mixing ratio, due to a better mixing in the atmospheric surface layer. During conditions with lower wind speed, deposition and reaction with local NO_x sources lead to a high depletion of O₃ during the night.

3.7 To what extent are the chemical reactions between NO, NO₂ and O₃ modifying the fluxes above the ground?

Measured mixing ratios and fluxes of NO, NO₂ and O₃ are affected by chemical reactions (Reactions SR1 to SR4 in the Supplement) in addition to emissions and deposition processes. Especially, the diurnal fluxes of NO₂ observed from 18 to 23 August, were positive (emissions) and of the same order of magnitude as the NO fluxes, while they were negative afterwards. The simultaneous observation of positive NO and NO₂ fluxes are typical for the NO-to-NO₂ transformation below the flux observation level in the presence of high O₃ mixing ratios. This phenomenon is called "apparent NO₂ emissions" and was observed in other studies mainly above dense or tall canopies (Ammann et al., 2012; Min et al., 2014; Plake et al., 2015). For the Reactions (SR1)-(SR2) to occur below the measurement height, the turbulent transport time (τ_{trans}) needs to exceed the chemical reaction time (τ_{chem}) (Arellano and Duynkerke, 1992; De Arellano et al., 1993; Lenschow and Delany, 1987; Plake et al., 2015; Stella et al., 2011a, 2012, 2013a). The Damköhler number $Da = \tau_{\text{trans}} / \tau_{\text{chem}}$ is often used to determine the conditions favourable for chemical reactions: in cases when Da is higher



Figure 10. Half-hourly values of photo-stationary state ratio (PSS) and $Q = k_r[NO][O_3] - J_{NO_2}[NO_2]$ (s); chemistry and transport timescales (τ_{chem} and τ_{trans}) and Damköhler number (*Da*); measured NO, NO₂ and O₃ fluxes and surface fluxes as computed by assuming a logarithmic flux divergence profile (F_{NO} , F_{NO_2} and F_{O_3}) at the Grignon field site.

than unity chemical reactions are faster than the transport (flux divergence), whereas Da values smaller than 0.1 indicate that the influence of chemical reactions was negligible. The aerodynamic resistance $R_a(z_{ref})$ (Eq. S8) was overall quite small and ranging from 45 to 128 sm^{-1} (first and third quantiles), hence leading to a quite short transport timescale (but larger than 100 s most of the time). The boundary layer resistance was around 22 and 43 s m⁻¹ (first and third quantiles) for O₃ (Fig. 9). The surface resistance for O₃ was estimated as $R_{\text{soilO}_3} = V_{\text{dO}_3}^{-1} - R_{\text{a}}(z_{\text{ref}}) - R_{\text{bO}_3}$, and dominated the other resistances (100 to $480 \,\mathrm{s}\,\mathrm{m}^{-1}$). The O₃ penetration depth in the soil was estimated as the depth necessary to explain the measured R_{soilO_3} if molecular diffusion in the soil pores is the main limitation factor. In practice this corresponded to the dry soil layer used in Personne et al. (2009). This depth ranged from 2 to 10 mm on average and was smaller at noon than during the night (Fig. 9). Overall, the chemical time τ_{chem} and the transport time τ_{trans} were of the same order of magnitude at any time of the day between applications and during mineral fertilisation, and τ_{chem} was smaller than τ_{trans} during the organic fertilisation. As a consequence, the Damköhler number was around unity most of the time and larger than unity during the organic fertilisation period, showing that the reaction between O₃, NO and NO₂ happened during transport from the ground to the EC measurement height at all times at this site. During the fertilisation event, the Damköhler number was especially high at night, when the transport time increased more substantially than the chemical timescale. These results are similar to findings by Stella et al. (2012) for the same site over bare soil. During the periods with vegetation, the increase of the transport timescale above the canopy was less than that of the chemical timescale during night-time, as the presence of vegetation increases the mixing, and, hence diminishes $R_a(z_{ref})$.

The Damköhler number shows that NO reacts with O3 and that photolysis also plays a role. How does this affect the NO flux measured at the reference height compared to the one at the ground? We quantified this variation by numerically solving Eq. (S13), based on the model of Duyzer et al. (1995). Due to the reaction with O₃, the calculated NO flux at the ground surface was on average 32 % larger than that at the measurement height during the period 17-29 August (0.93 instead of 0.63 nmol $m^{-2} s^{-1}$). This would represent an increase of 37 g of N emission following slurry spreading. For NO₂, the calculated flux at the ground surface was mostly negative while it was mainly positive at the reference height during the period 18-22 August. On average the NO₂ flux at the ground was $-0.33 \text{ nmol m}^{-2} \text{ s}^{-1}$ over the period 17– 29 August while it was $-0.03 \text{ nmol m}^{-2} \text{ s}^{-1}$ at the reference height. For NO fluxes, the major discrepancy between fluxes at the surface and the measurement height occurs during periods with relatively large and stable values of the Damköhler number (Fig. 10), as this is the case when chemical reactions consume NO before it reaches the measurement height.

The derivation of surface fluxes with the Duyzer model also leads to a diurnal cycle of the NO flux that is closer to the one observed for ground temperature, corroborating the hypothesis that ground emissions are mostly due to nitrification for our site (Fig. 11).

Since the O_3 deposition flux was much larger than the NO flux, the reaction with NO changed the absolute value by only 3% when comparing the flux at the measurement height



Figure 11. Diurnal cycles of ground temperature, NO flux at measurement height and at surface determined by the logarithmic profile in August 2012.

to the ground surface. Indeed, as only Reactions (SR1) and (SR2) are considered in Eqs. (S12) and (S13), which we used to numerically evaluate surface fluxes, we obtain: $\Delta [F_{NO}] = \Delta [F_{O_3}] = -\Delta [F_{NO_2}] = 0.3 \text{ nmol m}^{-2} \text{ s}^{-1}$ where Δ stands for the difference between surface and measurement height.

3.8 Why is O₃ deposition increasing following organic fertilisation?

We observed that following organic fertilisation (performed by injection and hence soil tillage), O_3 deposition increased by a factor of 2 (as shown by the deposition velocity, Figs. 9 and 10). Several hypotheses may explain this increase: (1) the increased surface exchange due to soil tillage, (2) the reaction with NO emitted by the ground, and (3) the reaction with VOCs emitted by the ground.

The first hypothesis is that the increase in deposition velocity following the organic fertilisation could be due to a change in physical characteristics of the soil surface. Indeed, the application of cattle slurry with a trailing hose modifies the soil structure at the surface which consequently increases the available surface for O₃ deposition, and therefore the deposition velocity. This hypothesis is consistent with the comparison of measured deposition velocities and modelled deposition velocities using the Stella et al. (2011a) R_{soilO_3} parameterisation (see Sect. 3.3.3 and Fig. 6a). Indeed, while there is a good agreement between measured and modelled $V_{\rm dO_3}$ after the 26 August (i.e., after the rainfall event), modelled V_{dO_3} systematically underestimates measured V_d between slurry application and the rainfall event. Since the parameterisation of R_{soilO_3} was obtained for the Grignon site over different periods, that means R_{soilO_3} accounts for the mean soil structure of the Grignon site. Therefore, it can be hypothesised that (i) R_{soil} is underestimated from slurry application to the rainfall event due to the change of soil surface structure, and (ii) after the rainfall event, the soil surface recovers its mean structure corresponding to the R_{soilO_3} parameterisation.

The second hypothesis is that O₃ would react with NO emitted by the soil. Although the reactions with NO during transport are shown to be small compared to the NO flux (Fig. 10), reactions in the soil surface layer may be more significant due to large NO concentrations in the soil, despite the fact that this layer is very small. A graph of the difference between the measured and the modelled ozone flux following fertilisation (Fig. S4) seems to show that the additional O_3 deposition is correlated with the NO flux. This would mean that the NO₂ produced in the soil by reaction with NO would be adsorbed on the soil surface either in the mineral phase or dissolved in the water phase as NO₂. To evaluate this assumption further, we evaluated the Damköhler number in the soil surface layer by assuming that the layer depth is equal to the O₃ penetration depth δ_{O_3soil} (Fig. 9). In this layer the transport time is equal to soil resistance for O₃ times the penetration depth $R_{soilO_3} \times \delta_{O_3soil}$. We can evaluate the NO mixing ratio that would explain the additional O₃ destruction at the surface, by searching for the value of [NO]soil that satisfies $\tau_{\text{trans}}(\text{Soil}, O_3) = \tau_{\text{chem}}(\text{Soil}, O_3)$. By doing so, we found that [NO]soil would need to reach 5 to 40 ppm to explain the increase in O₃ deposition following organic fertilisation. Gut et al. (1998, 1999) measured NO mixing ratios at a 2 cm depth in the soil under wheat with the membrane tube technique and report mixing ratios around 100 ppb and always below 400 ppb following fertilisation, which is 1–2 orders of magnitude below the mixing ratio which would be needed to explain the observed O₃ flux. Moreover, the rate of NO production in the soil surface layer would have to be equal to the O₃ flux to the ground (around 20 nmol $m^{-2} s^{-1}$) which is an order of magnitude larger than what Gut et al. (1998) or Laville et al. (2009) report as maximum NO flux. However, we should stress that both Gut et al. (1998) and Laville et al. (2009) report NO fluxes that were measured in the presence of O₃ and hence would have been depleted by reaction with it in a similar way as here.

A third hypothesis is that O₃ would react with VOCs emitted by the ground. Reactive VOCs such as sesquiterpenes and monoterpenes have been previously found to be emitted from soils (Horvath et al., 2012; Penuelas et al., 2014), and some of these sesquiterpene species react with O₃ in the order of a few seconds. The reactions of O_3 with larger terpenes are important sources of OH, as well as the ozonolysis of simpler unsaturated compounds (Donahue et al., 2005). Currently, there are few or no data available on the emission of VOCs from slurry application. However, a recent study mainly focusing on quantification of odour emissions from soil application of manure slurry showed the formation of a certain number of VOCs, included organic sulfur compounds, carboxylic acids, alcohols, carbonyl compounds (ketones and aldehydes), aromatic compounds (phenols and indoles) and nitrogen compounds (Feilberg et al., 2015). Based on their analyses, the compound most responsible for the overall odour impact from the VOC emissions was 4-methylphenol. These authors also showed the emission of trimethylamine, a compound that can react quickly with O₃, leading to formation of secondary organic aerosols (Murphy et al., 2007). Furthermore, these authors suggest that a large part of these VOCs are formed through ozonation reactions (i.e. byproducts of ozonation: methanol, acetone and acetaldehyde). Indeed, the slurry would be transported downwards through the soil, where efficient heterogeneous reactions can take place at particle interfaces. It has been shown that the heterogeneous reaction probabilities may be much greater than anticipated. For example, measurements on oxide surfaces with a chemical structure commonly found in VOCs (i.e. alkenes, terpenes, carbonyls) showed that the O₃ reaction probability of a surface-attached alkene can be up to 5 orders of magnitude greater than for the same reaction in the gas phase (Stokes et al., 2008). In the same way, Fick et al. (2005) observed that ozonolysis reaction rates of some terpenes were much higher than predicted, possibly as a result of reactions on the surfaces used in their experiments. These results suggest that terpenes can remain on the surfaces, enhancing the O₃ reactivity. Similarly, some other authors observed that surface reaction probabilities with O3 were 10 to 120 times greater than their corresponding gas-phase values (Dubowski et al., 2004; Springs et al., 2011). It is also known that soils can act as a sink of VOCs, by their adsorption to soil mineral particle surfaces and humic substances (Penuelas et al., 2014). Hence, it is likely that surface chemistry including photo-enhanced O₃ uptake on organic matter (Jammoul et al., 2008; Reeser et al., 2009) may explain the increase in O₃ deposition, a process not yet described in the literature. It may also be likely that O₃ is destroyed by very reactive VOCs in the gas phase as hypothesised by Wolfe et al. (2011). These gas-phase reactions, however, require that the chemical reaction time be shorter than the turbulence transport time (Plake et al., 2015; Stella et al., 2012).

However, our study does not allow us to conclude definitively which of the three hypotheses is the most likely.

4 Conclusions

Eddy covariance flux measurements of the NO–NO₂–O₃ triad during a 7-month period allowed evaluating several mechanisms controlling the exchange of these reactive trace gases with an agricultural soil. The eddy covariance technique proved to be suitable at capturing seasonal and diurnal dynamics of the fluxes, and allowed us to interpret flux behaviour according to meteorological variables, fertilisation practices and chemical reactions. Nevertheless, random uncertainty was particularly important (> 20 %) during morning traffic peaks due to the non-stationarity of NO_x and O₃ mixing ratios. As concerns NO₂, uncertainty was even higher (up to 40 %) due to the indirect measurement method. We thus recommend caution in the use of the method in non-

stationary conditions, and combined measurements of horizontal gradients of mixing ratios to quantify the effect of advection. Also, additional measurements of surface mixing ratios would be useful to check the reconstruction of surface fluxes that we performed by using the logarithmicprofile model of Duyzer. Finally, high NO₂ to NO conversion efficiency should be assured to reduce uncertainty of NO₂ fluxes.

In particular, the magnitude and temporal variability of NO emission fluxes following two fertilisation episodes were analysed, one in summer and the other one in winter. Mean NO emissions during the whole period were in agreement with previous studies on the same site. Emissions were significantly higher (Student t test p value lower than 2×10^{-16} , and a factor of 7 difference on the mean) during 2 weeks following organic fertilisation in August than during the rest of the experimental period. These large emissions are mainly due to favourable conditions for nitrification: soil water content around 20% and high temperatures. In February, following mineral fertilisation, the increase of NO emissions was less pronounced, although the same amount of N was applied. This difference is likely due to less favourable conditions for nitrification in February (low temperature and higher soil water content), rather than to the different form of fertiliser. On average over the whole period, we derived a loss of $0.26 \text{ kg N ha}^{-1}$ as NO from the field. With a total N input of $96 \text{ kg} \text{ N} \text{ ha}^{-1}$, this results in an NO emission factor of 0.27 %, which is in the lower range of earlier reported values on this site (Laville et al., 2011).

Our findings show that NO emissions from agricultural soils are limited (0.27% of the N-NO applied over the 7-month period, which with a conservative estimation can be extended to a yearly amount). When hypothetically extended to France with an averaged nitrogen fertiliser use of $80 \text{ kg N} \text{ ha}^{-1}$ over a fertilised area of around 26 Mha, this would lead to a total emission of NO_x of around 5.6 kt N– NO. This is negligible compared to national emissions which are around 240 kt N-NO (CITEPA, 2015). The seasonality and spatial distribution of these emissions may, however, lead to air quality issues during spring and late summer-autumn which are the main fertiliser application periods in rural environments. Indeed, most of the emissions we measured occurred within a few weeks following fertilisation. In terms of ozone, our findings are in accordance with previous ones, showing that ozone is efficiently deposited throughout the year. This means that crops are participating through this process in the reduction of the atmospheric oxidising capacity.

The O₃ deposition velocity was significantly higher following organic fertilisation than during the rest of the experiment (Student *t* test *p* value lower than 2×10^{-16} and a factor 3 difference on the mean), despite the fact that vegetation was absent. This increase in O₃ deposition could not be explained by the reaction of O₃ with NO in the atmosphere as the NO flux was an order of magnitude smaller than that of O₃. The process behind this ozone deposition increase remains to be discovered. We hypothesised three underlying processes: (1) increase in soil surface due to soil tillage, (2) reaction with NO in the soil layer and (3) reactions of O_3 with VOCs emitted by the slurry. None of these hypotheses can be dismissed and further investigation is required. Experiments in controlled conditions are desirable to better understand these processes.

The evaluation of the chemical and turbulent transport times showed that reactions between NO, NO₂ and O₃ below the measurement height occurred during the whole measurement period, leading to a depletion of NO and a build-up of NO₂ from the ground to the measurement height. Following organic manure application, NO fluxes were reduced by 30 % from the surface to measurement height, while the NO₂ fluxes switched from deposition to uptake, being negative at the surface and positive at the measurement height. This phenomenon of "apparent NO₂ emissions" was reported in other studies, especially above forests. Here it also appears to be important above a bare soil and at moderate measurement heights, during conditions of strong NO emissions and high ambient O₃ mixing ratios.

Data availability. Data can be found in the Supplement.

The Supplement related to this article is available online at doi:10.5194/bg-14-2225-2017-supplement.

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

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