Fluxes of total reactive atmospheric nitrogen (Sigma N-r) using eddy covariance above arable land
Christian Bruemmer, Oliver Marx, Werner Kutsch, Christof Ammann, Veronika Wolff, Christophe Flechard, Annette Freibauer

To cite this version:
Christian Bruemmer, Oliver Marx, Werner Kutsch, Christof Ammann, Veronika Wolff, et al.. Fluxes of total reactive atmospheric nitrogen (Sigma N-r) using eddy covariance above arable land. Tellus B - Chemical and Physical Meteorology, 2013, 65, 10.3402/tellusb.v65i0.19770. hal-01209139

HAL Id: hal-01209139
https://hal.science/hal-01209139v1
Submitted on 28 May 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License
Fluxes of total reactive atmospheric nitrogen ($\Sigma N_r$) using eddy covariance above arable land

CHRISTIAN Brümmer, OLIVER Marx, WERNER Kutsch, CHRISTOF Ammann, VERONIKA Wolff, CHRISTOPHE R. Flechard & ANNETTE Freibauer

To cite this article: CHRISTIAN Brümmer, OLIVER Marx, WERNER Kutsch, CHRISTOF Ammann, VERONIKA Wolff, CHRISTOPHE R. Flechard & ANNETTE Freibauer (2013) Fluxes of total reactive atmospheric nitrogen ($\Sigma N_r$) using eddy covariance above arable land, Tellus B: Chemical and Physical Meteorology, 65:1, 19770, DOI: 10.3402/tellusb.v65i0.19770

To link to this article: https://doi.org/10.3402/tellusb.v65i0.19770
Fluxes of total reactive atmospheric nitrogen (ΣNr) using eddy covariance above arable land

By CHRISTIAN BRÜMMER1*, OLIVER MARX2, WERNER KUTSCH1, CHRISTOF AMMANN3, VERONIKA WOLFF3, CHRISTOPHE R. FLECHARD4 and ANNETTE FREIBAUER1, 1Thünen Institute of Climate-Smart Agriculture (TI-AK), Bundesallee 50, D-38116, Braunschweig, Germany; 2LI-COR Biosciences GmbH, Siemensstraße 25a, D-61352 Bad Homburg vor der Höhe, Germany; 3Swiss Federal Research Station Agroscope ART, Reckenholzstrasse 191, CH-8046, Zürich, Switzerland; 4INRA, Agrocampus Ouest, UMR 1069 SAS, 65 Rue de St.-Brieuc, 35042 Rennes, France

(Manuscript received 24 September 2012; in final form 22 January 2013)

ABSTRACT
The amount and timing of reactive nitrogen exchange between agricultural land and the atmosphere play a key role in evaluating ecosystem productivity and in addressing atmospheric nitrogen budgets and transport. With the recent development of the Total Reactive Atmospheric Nitrogen Converter (TRANC) apparatus, a methodology has been provided for continuous measurement of the sum of all airborne nitrogen containing species (ΣNr) allowing for diurnal and seasonal investigations. We present ΣNr concentration and net flux data from an 11-month field campaign conducted at an arable field using the TRANC system within an eddy-covariance setup. Clear diurnal patterns of both ΣNr concentrations and fluxes with significant dependencies on atmospheric stability and stomatal regulation were observed in the growing season. TRANC data were compared with monthly-averaged concentrations and dry deposition rates of selected Nr compounds using DELTA denuders and ensemble-averages of four inferential models, respectively. Similar seasonal trends were found for Nr concentrations from DELTA and TRANC measurements with values from the latter being considerably higher than those of DELTA denuders. The variability of the difference between these two systems could be explained by seasonally changing source locations of NOx contributions to the TRANC signal. As soil and vegetation Nr emissions to the atmosphere are generally not treated by inferential (dry deposition) models, TRANC data showed lower monthly deposition rates than those obtained from inferential modelling. Our measurements demonstrate that fertilizer addition followed by substantial Nr emissions plays a crucial role in a site’s annual atmospheric nitrogen budget. As long-term Nr measurements with high temporal resolution are usually cost and labour-intensive, field application of the TRANC helps improve the understanding of ecosystem functioning, atmospheric transport and revising definitions of ecosystem-specific critical loads at a relatively moderate operational cost level.

Keywords: agricultural land, biosphere–atmosphere exchange, converter, ecosystem N saturation, eddy covariance, fertilization, N budget, TRANC, total reactive nitrogen (ΣNr) exchange

1. Introduction
The biosphere–atmosphere interaction of nitrogen (N) containing species has direct impacts on ecosystem nutrient dynamics and on oxidative atmospheric chemistry (Ollinger et al., 2002; Farmer et al., 2006). Fertilizer application, symbiotic fixation of dinitrogen (N2), and atmospheric deposition of gaseous ammonia (NH3), particulate ammonium (NH4+) and oxidized forms of N lead to a net gain of N in ecosystems, whereas soil and plant emissions of nitric oxide (NO) or nitrogen dioxide (NO2) as well as nitrous oxide (N2O) via nitrification and denitrification, cause a release of N into the atmosphere (Vitousek et al., 1997). Elevated deposition rates of reactive nitrogen (Nr) are the result of increased ambient concentrations due to high emissions by intensive farming (mainly reduced Nr) and by
traffic and industry (mainly oxidized N$_r$) (Flechard et al., 2011). The continued boost in anthropogenic emissions of N$_r$ into the atmosphere and N deposition has led to concerns regarding ecosystem N saturation, which may threaten plant health, increase greenhouse gas emissions, and affect water quality (Aber et al., 1998). In a modelling study, Flechard et al. (2011) report on annual N$_r$ dry deposition rates of $<2$ kg N ha$^{-1}$ at remote forest sites in Scandinavia and $>30$–40 kg N ha$^{-1}$ over parts of the Netherlands and Belgium. Generally, turbulent N$_r$ deposition was modelled to be smaller over short vegetation than over forests due to reduced turbulent exchange. The dry deposition of N$_r$, i.e. of various inorganic species such as gaseous NH$_3$, nitric acid (HNO$_3$), nitrous acid (HONO), NO, NO$_2$, and aerosol NH$_4$$_2$ and nitrate (NO$_3$$_2$) as well as in a range of organic molecules in both phases (e.g. gaseous peroxyacetyl nitrate (PAN) and other organic nitrates and amines), typically contributes between one and two thirds of total atmospheric N$_r$ deposition (Erisman et al., 1996; Simpson et al., 2006; Zimmermann et al., 2006; Zhang et al., 2009).

Despite the high relevance of N$_r$ cycling for environmental research, the scientific progress in quantifying ecosystem–atmosphere interactions of N$_r$ has fundamentally depended on advances in chemical measurement techniques (Erisman et al., 2001; Fowler et al., 2001; Sutton et al., 2007). Up to now, measurements of dry, i.e. turbulent, N$_r$ exchange using eddy covariance (EC) (Aubinet et al., 2000) have mainly remained experimental and limited to only a few research sites (Munger et al., 1996) and to short-term campaigns (e.g. Day et al., 2002; Farmer et al., 2006; Sintermann et al., 2011), whereas wet deposition is monitored in regional networks of wet-only or bulk precipitation collectors (Flechard et al., 2011). Reasons for the limitation of dry deposition monitoring of N$_r$ are amongst others the wide range, complexity and labour-intensive of flux measurements of N$_r$ containing compounds as well as their mutual interactions and their different molecular and exchange characteristics.

With the development of a novel converter named Total Reactive Atmospheric Nitrogen Converter (TRANC), a methodology is provided for both the quantification of the sum of all airborne N$_r$ compounds – denoted as $\Sigma N_r$ hereafter – and a device being robust and fast enough to be usable in an EC setup when combined with a fast-response NO detector (chemiluminescence detector, CLD). The basic concept of the TRANC is the full and quasi-immediate conversion of $\Sigma N_r$ to NO within a thermal and a catalytic reaction step (for details see Marx et al., 2012). The specific design of the converter overcomes former detection problems such as surface water effects (Kita et al., 2006; Sintermann et al., 2011) by fast and strong heating of the air sample and quick conversion of N$_r$ compounds to NO, thus allowing for measurements of water soluble and sticky substances like NH$_3$. As NO$_2$ and ozone (O$_3$) are fully converted in the TRANC and therefore total NO is conserved, the use of long tubing between TRANC and CLD – possibly needed for the installation of the TRANC on a tower while the CLD is usually protected and stored in housing on the ground – is feasible. The TRANC was successfully tested for recovery rates of single N$_r$ compounds (NO$_2$, NH$_3$) in the lab and showed excellent agreement during an in-field comparison between the sum of individually measured NO, NO$_2$, NH$_3$, HNO$_3$, HONO, NH$_4$$_2$, and NO$_3$ by commercially available systems and the $\Sigma N_r$ concentration measured by the TRANC-CLD system (Marx et al., 2012). During concentration step changes, the fast response of the system was proved by e-folding times between 0.30 and 0.35 s for different species.

Although a differentiation into single N$_r$ compounds is not possible, the usage of the TRANC-CLD system allows for the determination of the net $\Sigma N_r$ exchange, thereby providing a parameter of major interest for productivity assessment of ecosystems by its N nutrient inputs. The overall aim of this paper is to present the first continuous, multi-month dataset of direct N$_r$ measurements conducted at an agricultural site. Specifically, we (1) investigate environmental controls on diurnal and seasonal progression of N$_r$ concentration and flux patterns, (2) compare TRANC measurements with results from DELTA observations and inferential modelling, and (3) highlight the impact of fertilization on net $\Sigma N_r$ exchange at an agricultural site. In this paper, we define N$_r$ as all nitrogen-containing trace species except for N$_2$ and N$_2$O, which are both inert in the troposphere (Seinfeld and Pandis, 2006) and not measurable by the TRANC-CLD system.

2. Methods

2.1. Site description and management

The research site (51°06.013"N, 10°54.519"E, 162 masl) is located near the village of Gebesee about 20 km northwest of Erfurt in Thuringia, Germany, and was part of the Integrated Projects CarboEurope (2004–2008) and NitroEurope (2006–2011). The measurements presented in this study were conducted from December 2007 to November 2008. Mean annual air temperature ($T_a$) and precipitation in Gebesee during the reference period 1961–1990 was 7.9°C and 500 mm, respectively. The year 2007 was an exceptionally warm (10.45°C) and wet (692 mm) year. In 2008, mean annual air temperature was also significantly above the long-term mean (9.98°C), whereas total precipitation was slightly below average (480 mm).
The soil type is classified as Chernozem (FAO) with granular structure and a texture defined as silty clay loam consisting of approximately 30% clay. Soil pH is 6.6 ± 0.5 in the uppermost 10 cm layer and increases to 7.3 ± 0.2 in 90 cm depth. Soil organic carbon content is 42.6 and 2.9 g kg⁻¹ in the A₀- and B-horizon, respectively. Total carbon stock in <0.6 m depth is 97 t ha⁻¹ (Kutsch et al., 2010) and dry bulk density is 1.29 ± 0.12 g cm⁻³. Carbon to nitrogen ratio in <40 cm depth is ~9 and ~13 in the layers below (Anthoni et al., 2004).

The black soil region of the Thuringian Basin has been farmed for at least 900 years. Since 1970 the research field at Gebesee has been alternately cultivated with winter wheat (Triticum aestivum L.) or winter barley (Hordeum vulgare L.) followed either by potato (Solanum tuberosum L.), sugar beet (Beta vulgaris subsp. vulgaris var. altissima) or rapeseed (Brassica napus L.). Every four years the cereal straw is left on the field after harvest. Farmyard manure is added to the soil about every seven years. Since the 1990, ploughing depth has been reduced and is now usually between 20 and 28 cm.

However, during the observation period, a change in the regular crop rotation was made. Due to the late harvest of sugar beet in 2006, winter wheat seeds were only sown on 9 November 2006, which led to a relatively poor harvest in 2007. As a consequence, the farmers decided to plant winter wheat for a second consecutive year. Seeds in 2007 were sown on October 6. Harvest in the two observational years took place on 7 August 2007 and on 5 August 2008. Four fertilizer applications occurred during this period. The first was farmyard manure three months before the start of measurements at a rate of 260 kg N ha⁻¹. 

2.2. \( \Sigma N_r \) measurements

A tower for conventional EC measurements of carbon dioxide (\( \text{CO}_2 \)), water and energy exchange with auxiliary meteorological measurements was setup in January 2001 (details are given in Anthoni et al. 2004). The same tower was used for the installation of an extra ultrasonic anemometer (model R3, Gill Instruments, Lymington, UK) and the converter for \( N_r \) measurements named TRANC. The sonic and converter were mounted on a boom at a height of 4.8 m facing west. As the converter has a diameter of 12 cm and a length of 71 cm, it was positioned downwind of the sonic as seen from the main wind direction to avoid interferences of the micro-turbulence for most of the time.

Core pieces of the converter are a high-temperature resistant metal tube consisting of an iron–nickel–chrome (FeNiCr) alloy heated up to 870°C followed by a gold (Au) tube heated up to 300°C for thermal and catalytic conversion, respectively, in order to reduce/oxidize all N containing species to NO. A very short distance between the sample air inlet and the two conversion tubes assures a short residence time of the sample air inside the instrument, and minimizes wall sorption problems of water soluble compounds. A current of 80 ampere at a voltage of 5 V was fed to the FeNiCr tube over brass blocks. Provision of electricity was effected by a switch-mode power supply with the temperature of the tube being regulated by a universal modulator and controlled by a thermocouple. The Au tube is indirectly warmed up by the heat loss through the FeNiCr tube. Measurement principle and detailed functionality are given in Marx et al. (2012).

Auxiliary devices required for system calibration and continuous determination of \( N_r \) concentration and fluxes comprise a CLD (780TR, ECO PHYSICS, Dürnten, Switzerland), a multi-gas calibrator (S6100, Environics Inc., Tolland, USA), a pure air generator (PAG 003, ECO PHYSICS, Dürnten, Switzerland), and a dry vacuum scroll pump (BOC Edwards XDS10, Sussex, UK), which were kept in an air-conditioned trailer a few meters away from the bottom of the tower. The TRANC was connected with the CLD over 18 m-long opaque tubing. Gas phase titration in the main chamber of the CLD required a low pressure regime to reduce the probability of energy loss of excited \( \text{NO}_2 \) molecules by collision with others. This was affected by the installation of a critical orifice located at the back end of a stainless steel tube that is leaving the TRANC towards the CLD and also restricted the flow rate to 2.7 L min⁻¹. In our setup, the CLD was running in continuous mode, integrating photons over 0.05 s.

The TRANC-CLD unit was calibrated once every night between 03:30 am and 04:00 am by feeding reference gas with different NO and \( \text{NO}_2 \) concentrations through the system to retain optimal data coverage. The multi-gas calibrator was connected to two reference gas cylinders each containing 10 ppm NO and 10 ppm \( \text{NO}_2 \). The calibrator mixed specified gas concentrations by dilution of the cylinder standard with zero air provided by the pure air generator. Reference gases with concentrations of 0, 6, 10, 20, 50, and 120 ppb NO were used for calibrating the CLD (only one particular concentration per night and the other reference gases in the following nights) and obtaining the respective regression parameters. Additionally, 50 ppb \( \text{NO}_2 \) and 50 ppb NO mixed with \( O_3 \ (<50 \text{ ppb}) \) was put into
the system to determine the conversion efficiency of NO₂. After activating the TRANC calibration valve, the system was calibrated for 30 min with one specific gas concentration. To assure a well-conditioned calibration gas tube and therefore a clearly stable calibration gas concentration at the sample air inlet of the TRANC, only data of the 10-min period from 15 to 25 min after the initial switch were taken to calculate the NO concentrations. If necessary, spikes were removed and the 10-min window was shifted to the most stable interval during the half-hour calibration period. $R^2$ values of the calibration regression for single 8-day periods were ranging between 0.81 and 0.99 with $R^2 = 0.98$ for the averaged 11-month fit (NO₂ was 91%). A possible reason for the mismatch could be positive (Marx et al., 2012). The conversion efficiency for the dominant Nr species, the fast time response of the TRANC-CLD system during step concentration changes between reference gases of NO, NO₂, a gas mixture of NO and O₃, and ambient air (e-folding times of 0.30–0.35 s were found) enables the converter to be used in an eddy-covariance setup. The reaction time of the whole system is mainly limited by the flushing of the CLD reaction chamber as the average residence time of air in the cell ($V = 0.7 l$) at a pressure of 20 mbar and a flow rate of 3 sl min⁻¹ is 0.28 s.

Regular site and instrumental maintenance was carried out every two weeks; filter (Acrodisc® Gelman 0.45 μm PTFE membrane) replacement once a month. The agricultural field where the measurement tower was erected had an approximate semi-rectangular size of 750 m × 850 m. The fetch in the prevailing bimodal wind directions was around 400 m to the west–southwest and around 600 m to the northeast. Estimates of the flux footprint based on an analytic model by Schmid (1993) indicated that for neutral day-time conditions the maximum source area location was at about 80 m distance, 50% contributions were from within 160 m, and 90% within 300 m of the tower. Unstable conditions led to smaller footprints, but very stable conditions at night could have led to larger footprints, bigger than the fetch of the wheat field (Anthoni et al., 2004). However, data from low-turbulence conditions were usually rejected due to low $u*$ (Section 2.3).

2.3. Data analysis and flux calculation

Raw signals of wind vectors, sonic temperature, $\Sigma N_r$ concentration, speed of sound, and system control parameters were recorded on a personal computer at a frequency of 20 Hz using Labview7 software (National Instruments, Texas, USA). Prior to calculating eddy fluxes, high-frequency data were linearly detrended and a 2D rotation of the wind speed vectors $u$, $v$, and $w$ according to Aubinet et al. (2000) was applied. $\Sigma N_r$ fluxes were calculated for half-hourly intervals as

$$F_{N_r} = \Sigma_{0}^{\infty} \rho_{a}[w(t) - \bar{w}] * [c(t + \tau) - \bar{c}]$$  \hspace{1cm} (1)

with $\rho_{a}$ being the molar density of dry air (kg m⁻³), $w$ and $c$ being vertical wind speed and $\Sigma N_r$ concentration (nmol mol⁻¹), respectively, where over bars indicate mean values for each 30-min period. Positive covariances correspond to upward transport. The time lag $\tau$ accounts for the delayed detection of NO by the CLD mainly due to the residence time of the air sample in the converter and in the sampling tube. Time lags were determined by shifting the time series of $c'$ against that of $w'$ until the maximum covariance was reached. In our setup, $\tau$ values were usually between 36 and 46 samples corresponding to 1.8 and 2.3 s, respectively. Possible high frequency damping on trace gas time series (here $\Sigma N_r$) in closed-path systems was analyzed using the ogive method as described in Ammann et al. (2006). Ogives, i.e. cumulative cospectra of $w\Sigma N_r'$ were calculated and scaled to the respective reference ogives of $w'T_s'$ in the lower frequency range. The resulting relative deviations at the high-frequency end are quantitative measures of spectral damping. Over a range of different conditions, e.g. periods with mainly deposition or emission, we found a median of 0.90 indicating a loss of 10% for $\Sigma N_r$ fluxes. As results of this analysis were invariant with changing wind speed and other possible drivers such as $z/L$, we applied a factor of 1/0.9 to all $\Sigma N_r$ flux values. An example of $wT_s'$ and $w\Sigma N_r'$ time series and ogives is given in Fig. 1 (further analyses of high-frequency damping using a similar setup can be found in Ammann et al., 2012).

We also observed a small but significant water vapour influence on the CLD signal, probably due to a quenching of the NO chemiluminescence, which consequently affected $\Sigma N_r$ fluxes. For the NO analyser used, a 0.19% NO sensitivity reduction per 1 nmol mol⁻¹ water vapour increase was found. Thus, we corrected every $\Sigma N_r$ flux value [eq. (1)] by adding the term for the water vapour induced NO interference flux ($F_{NO, int}$) [eq. (2)]. The latter is defined as follows:

$$F_{NO, int} = -0.0019 * c_{\Sigma N_r} * F_{H_2O}$$  \hspace{1cm} (2)

with $c_{\Sigma N_r}$ being the $\Sigma N_r$ concentration (nmol mol⁻¹) and $F_{H_2O}$ being the water vapour flux (mmol m⁻² s⁻¹) as was measured by the conventional EC system on the same tower (Anthoni et al., 2004). For instance, a cross-sensitivity of $-0.038$ nmol N (mmol H₂O⁻¹) for a typical background
SR concentration of 20 ppb and a water vapour flux of 5 mmol m$^{-2}$s$^{-1}$ was determined, which resulted in $F_{\text{NO, int}}$ of $-2.7$ ng N m$^{-2}$s$^{-1}$ (see also Ammann et al., 2012). However, during the presented Gebesee campaign, the overall balance (Section 3.3) was shifted towards less deposition by only 25 g N ha$^{-1}$.

In accordance with Ammann et al. (2012), who used the same system at a grassland site in Oensingen, Switzerland, an upper flux detection limit for the Gebesee setup was estimated by using night-time data under stationary conditions, i.e. relatively high wind speeds between 3 and 7 m s$^{-1}$ and constant wind directions from the homogeneous fetch area southwest of the tower as well as situations with relatively low SR concentrations (<15 ppb). Under these conditions, half-hourly SR fluxes were relatively small. The calculated standard deviation was 3.3 ng m$^{-2}$s$^{-1}$, thus the resulting 2σ-uncertainty range of $\sim 6.6$ ng m$^{-2}$s$^{-1}$ may be regarded as an upper flux detection limit for half-hourly values of the TRANC SR, eddy-covariance system. The lower detection limit, however, solely depends on the sensitivity of the CLD. Using the same analyser model, Rummel et al. (2002) estimated a flux detection limit due to

![Fig. 1.](image)

**Fig. 1.** Upper left and lower left panel: Covariance function of vertical wind speed and temperature and covariance function of vertical wind speed and SR concentration measured by the TRANC-CLD system, respectively. Upper right and lower right panel: Normalized flux ogives, i.e. cumulative cospectra of sensible heat and SR flux, respectively. Data were recorded on 7 June 2008, 11:00–11:30 am.
instrumental noise of 0.07 ng m\(^{-2}\) s\(^{-1}\) for EC measurements of NO.

Night-time data with insufficient turbulence (\(u^*<0.1\)) were discarded. This filter caused a data loss of 24.1%. As exchange characteristics of \(\Sigma N_r\) between ecosystem and atmosphere differ from those of CO\(_2\), where night-time gaps are typically filled by fitting a temperature-based respiration model, we used the mean diurnal variation (MDV) method (Falge et al., 2001; Moffat et al., 2007) to fill all night-time \(\Sigma N_r\) flux gaps. A moving window of five consecutive days before and after the missing value was taken to fill the respective gap. As night-time fluxes were comparatively small (see Section 3.2), we estimated the error being in-respective gap. As night-time fluxes were comparatively small, we estimated the error being in-

Turbulent mixing of the surface layer. Lower daytime \(u^*\) mainly occurred when \(T_a\) was relatively low and surfaces were wetter, i.e. conditions favouring deposition. Those flux cases that passed the \(u^*\)-filter and were recorded at positive atmospheric stability did never exceed \(\zeta=0.05\), thus still indicating near-neutral conditions. These situations mainly occurred during night-time when \(\Sigma N_r\) fluxes were relatively small and did not significantly contribute to the total \(\Sigma N_r\) budget.

Atmospheric stability (\(\zeta\)) was used in the analysis and discussion about controlling factors of \(\Sigma N_r\) exchange and was expressed as described in Foken (2008):

\[
\zeta = \frac{u^*_0}{L}
\]

where \(u^*_0\) is the friction velocity (m s\(^{-1}\)), \(\kappa\) is the von Karman constant (0.4), \(g\) is the gravity acceleration (m s\(^{-2}\)), \(T_a\) is the air temperature (K), \(H\) is the sensible heat flux (W m\(^{-2}\)), and \(c_p\) is the specific heat of air (J g\(^{-1}\) K\(^{-1}\)). Positive values of \(\zeta\) indicate stable conditions, negative values indicate unstable conditions, and values around 0 indicate neutral conditions in the surface layer.

Stomatal conductance \((g_s)\) was used to validate TRANC measurements in the context of canopy resistance modelling. We inferred \(g_s\) from surface conductance to water vapour transfer at canopy scale and leaf area index (LAI) using the Penman-Monteith approach applied to measured eddy covariance data as described in Lu et al. (2003), Kutsch et al. (2008) and Brümmer et al. (2012).

### 3. Results and discussion

To specifically study short-term disturbances as well as longer-term ecosystem functioning, data of emission peak events following mineral fertilizer application on the one hand and background exchange on the other hand were analyzed separately. After the first two fertilizer additions on 18 April and 21 May 2008, \(\Sigma N_r\) fluxes were significantly elevated compared to pre-fertilization levels for only 3–4 days. After the fertilizer application on 6 June 2008, however, a net release of \(\Sigma N_r\) to the atmosphere was observed for several weeks (until August 25) followed by a month of net deposition and another two-week period of net emissions even without previous fertilization. The separation of the entire observation period into net emission and net deposition periods is summarized in Table 1.

#### 3.1. \(\Sigma N_r\) concentrations – temporal patterns and controlling factors

##### 3.1.1. Mean diurnal variation

\(\Sigma N_r\) concentrations on half-hourly basis were usually between 2 and 70 ppb and between 7 and 23 when averaged over a month (Fig. 2). Clear mean diurnal patterns with highest values around 18 ppb at night and lowest values in early afternoon (~3 ppb) were observed during the growing season and late summer (D2 and D4 in Fig. 3A, respectively) indicating distinct turbulent mixing of the surface layer. Lower daytime \(\Sigma N_r\) concentrations during periods like D2, when plants are in a phase of their maximum growth rate, were likely an indicator for atmospheric \(N_r\) – presumably NH\(_3\) – uptake by the wheat plants as was found by Rogers and Aneja (1980) under increased light and temperature conditions. Furthermore, \(\Sigma N_r\) from possible night-time emissions was probably not well mixed into the lower parts of the atmosphere due to a more stable stratification than during

| Period dominated by | Period | DOY |
|--------------------|--------|-----|
| **Deposition**      | D1     | Dec 23–Apr 17 | 357–108 |
| E1                 | Apr 18–Apr 21 | 109–112 |
| D2                 | Apr 22–May 20 | 113–141 |
| E2                 | May 21–May 25 | 142–146 |
| D3                 | May 26–Jun 5 | 147–157 |
| E3                 | Jun 6–Aug 25 | 158–238 |
| D4                 | Aug 26–Sep 20 | 239–264 |
| E4                 | Sep 21–Oct 4 | 265–278 |
| D5                 | Oct 5–Nov 7 | 279–312 |
daytime. Under these conditions, $\Sigma N_r$ could have also been transported through lateral flow paths from sources other than the investigated agricultural field. A smoother and less clear pattern of mean diurnal $\Sigma N_r$ concentrations was found during October (D5). One reason for this observation – particularly for the relatively low nighttime values – might be the fact that already four months passed since the last fertilizer application and a considerable amount of $\Sigma N_r$ had been removed from the atmosphere due to vertical mixing of the planetary boundary layer and relatively low lateral $\Sigma N_r$ input because of homogeneous agricultural land in the whole region without heavy industry and no significant wintertime household heating in residential areas in October/November. Furthermore, manure spreading in the wider vicinity is not done during that time of year, thus keeping the diurnal variability of $\Sigma N_r$ concentrations less dynamic than during the growing season or immediately after fertilization (first part of E3 in Fig. 3B).

3.1.2. Comparison of seasonal patterns by TRANC and DELTA. $\Sigma N_r$ concentrations measured by the TRANC-CLD system were compared with those obtained by DELTA denuders (details are given in Sutton et al., 2001; Tang et al., 2009), which were measured at the same site within the framework of NitroEurope-IP (Flechard et al. 2011). This system integrates gaseous NH$_3$, HNO$_3$, HONO, and aerosol NH$_4^+$ and NO$_3^-$ over a period of one month. As DELTA denuders measure neither NO$_x$ nor organic N compounds (PAN, etc.), we expected TRANC values to be somewhat higher than DELTA values. Fig. 4A shows that this assumption is consistent with the observations, and that the two time series generally revealed similar temporal patterns, which were likely driven by strong seasonal variability of atmospheric turbulent mixing. The difference between the two, however, varied through the year. A discrepancy of up to 10 ppb was observed from December 2007 to May 2008 and from October 2008 to November 2008, whereas the difference during the

![Fig. 2](image-url)
summer months (June 2008 to September 2008) was small (1–2 ppb). We assume that these differences were caused by the NOx contribution to the \( \Sigma N_r \) signal generated from wintertime household heating through domestic coal, which strongly depends on wind direction. A bimodal wind direction pattern at the research site in Gebesee (Fig. 4C) with west–southwest and east–northeast being the main and the second most common wind direction, respectively, was prevalent. The area around the tower (and outside the footprint) is mainly agricultural land with only a few country roads subjected to very low traffic volume. Two kilometres to the north, however, there is the small village of Gebesee, and around 15 km to the south the city of Erfurt with ~200,000 residents is located (Fig. 4B). As shown in Fig. 4D winds are rarely coming from these directions, but if so, \( \Sigma N_r \) concentrations were significantly higher (up to 50 ppb) than by winds from the west or east (usually between 20 and 25 ppb). This was particularly the case in February when the discrepancy between TRANC and DELTA values was relatively high as were \( \Sigma N_r \) concentrations by winds from the south, thus being an indicator for NOx contributions detected at the site through wintertime heating from residential areas of Erfurt.

On an annual basis, the mean difference between TRANC and DELTA \( \Sigma N_r \) concentrations was observed to be ~5 ppb. This magnitude is completely in the range of annual NO2 values obtained from the EMEP model (Simpson et al., 2003; Tuovinen et al., 2009; Flechard et al. 2011) (see Section 3.2.2) and spatially averaged annual NO2 concentrations from a regional air quality monitoring network of the Federal Environment

---

Fig. 3. Mean diurnal variation of \( \Sigma N_r \) concentrations separated in periods dominated by deposition (Panel A) and periods dominated by emission (Panel B). Emission period 3 (E3) was divided into sub-sections due to significant changes four days after fertilization.
Agency (UBA) (Gauger, 2009), who found \( \sim 3-5 \) ppb and \( \sim 5-7 \) ppb from model runs and measurements, respectively.

We found a significant correlation \( (R^2 = 0.40; p < 0.05) \) between \( \zeta \) and \( \Sigma N_r \) concentrations, even when averaged over a month (Fig. 4A). Primarily stable conditions indicated by positive \( \zeta \) values corresponded to relatively high mean \( \Sigma N_r \) concentrations \((\sim 23 \) ppb\), e.g. in February 2008, whereas neutral to unstable conditions indicated by \( \zeta \leq 0 \) corresponded to relatively low \( \Sigma N_r \) concentrations \((\leq 10 \) ppb\). The same relationship was found when comparing \( \Sigma N_r \) concentrations with \( u^* \) values (Fig. 5A). Bin-averaged low turbulence conditions \((u^* < 0.1 \) m s\(^{-1}\)) were observed when mean \( \Sigma N_r \) concentrations were relatively high \((> 20 \) ppb\), whereas relatively low \( \Sigma N_r \) concentrations \((< 18 \) ppb\) were found during unstable atmospheric conditions \((u^* > 0.2 \) m s\(^{-1}\)) revealing a countercyclical course of \( \Sigma N_r \) concentrations (cf. Fig. 3A) and \( u^* \) on a diurnal basis.

### 3.2. \( \Sigma N_r \) fluxes

**3.2.1. Variability and determinants.** Half-hourly \( \Sigma N_r \) fluxes were usually between \(-40 \) to \( 40 \) ng N m\(^{-2}\) s\(^{-1}\) with single uptake values as low as \(-175 \) ng N m\(^{-2}\) s\(^{-1}\) and highest emission values reaching \( 4132 \) ng N m\(^{-2}\) s\(^{-1}\) after the fertilization event on 6 June 2008. The frequency distribution in Fig. 6 shows that the bulk of the fluxes were within \(-20 \) to \( 20 \) ng N m\(^{-2}\) s\(^{-1}\) \((\Sigma = 3.3 \) ng N m\(^{-2}\) s\(^{-1}\), cf. Section 2.3) with mostly zero exchange or a small uptake. Around 60.5\% of the fluxes were negative, corresponding to deposition of \( \Sigma N_r \).

We observed high variability in mean diurnal courses of \( \Sigma N_r \) fluxes (Fig. 7A and 7B) between different periods of both deposition and emission phases. While deposition fluxes during winter (D1) and autumn (D5) were virtually zero during night-time and rarely up to \( 20 \) ng N m\(^{-2}\) s\(^{-1}\) in the morning hours, higher amplitudes of up to \( 60 \) ng N m\(^{-2}\) s\(^{-1}\) (also zero exchange at night, but up to \( 60 \) ng N m\(^{-2}\) s\(^{-1}\) in the morning) were found in the growing season (D2) and after harvest in August to September (D4). In deposition period D2, the diurnal progression of \( \Sigma N_r \) concentration lagged behind that of \( \Sigma N_r \) flux by approximately \( 5-6 \) hours (cf. Fig. 8A and 8B). \( \Sigma N_r \) uptake increased shortly after sunrise when \( \Sigma N_r \) concentrations were relatively high and likely above the canopy compensation point for NH\(_3\) \((\chi)\), thus promoting \( N \) uptake (in the form of NH\(_3\)). Although the N status of crops is usually

---

*Fig. 4.* Monthly mean concentrations of \( \Sigma N_r \) measured by the TRANC-CLD system (red line) and DELTA denuders (blue line) in ppb (left y-axis). Each dot of the DELTA line represents the sum of NH\(_3\), HNO\(_2\), HNO\(_3\), particulate NO\(_3\), and particulate NH\(_4\). Monthly mean atmospheric stability \( \zeta \) is shown in grey (right y-axis) (Panel A). Panel B shows a map of the Thuringian Basin with the field site in Gebesee and the City of Erfurt. Mean frequency distribution of wind direction for selected periods and mean frequency distribution of \( \Sigma N_r \) concentrations measured by the TRANC-CLD system for selected periods are shown in Panel C and Panel D, respectively.
high, we assume that during this period of maximum plant growth the NH$_4^+$ concentration in the apoplast was kept on a relatively low level, because at that time of the year NH$_4^+$ uptake reactions (e.g. N transport and NH$_4^+$ assimilation) outbalance NH$_4^+$-releasing reactions (e.g. deamination, senescence-induced proteolysis, and NO$_3^-$ reduction) thereby favouring NH$_3$ uptake (Schjoerring et al., 1998; Francis et al., 2008). The daily course of $\Sigma$Nr uptake was in phase with that of $g_s$, peaking before noon and recovering to its initial level in the afternoon (cf. Fig. 8B and 8C). Shape and correlation of $\Sigma$Nr exchange and $g_s$ might have been caused by both the fact that water vapour deficit becomes a key limiting factor for stomatal activity around noon and that further uptake of N$_r$ was impeded by already decreased $\Sigma$Nr concentrations at that time of the day. Furthermore, vertical mixing of the planetary boundary layer likely controlled the diurnal pattern of $\Sigma$Nr concentration with highest values during night-time and lowest values in the afternoon (cf. Section 3.1). Beside NO$_2$, which can also be consistently taken up via stomata, the shape of the diurnal $\Sigma$Nr fluxes in the morning hours was – at least on some days – also likely influenced by significant non-stomatal sinks for NH$_3$ such as leaf surface wetness (dew formation, evaporation and rainfall).

Furthermore, we found a significant correlation ($R^2=0.77; p<0.001$) between $\Sigma$Nr fluxes and $\Sigma$Nr concentrations multiplied by $g_s$ for the deposition period D2 with low exchange between $-10$ and $+5$ ng N m$^{-2}$ s$^{-1}$ occurring when values for N$_r$ concentration multiplied by $g_s$ were $<50$ ng N m$^{-2}$ s$^{-1}$ and high uptake ($\sim40$–$50$ ng N m$^{-2}$ s$^{-1}$) corresponding to values of $\Sigma$Nr concentration multiplied by $g_s>150$ ng m$^{-2}$ s$^{-1}$ (Fig. 8D and 8E). This demonstrates that ecosystem–atmosphere matter fluxes – mainly unidirectional deposition fluxes as prevalent in April and May – may be viewed and derived in analogy to Ohm’s law, i.e. the flux of a substance is expressed as the ratio of its concentration at measurement height and the resistance against deposition to the ground. In inferential modelling of trace gas dry deposition, where one-point concentrations of any N$_r$ compound are usually determined

![Fig. 5. Dependency of $\Sigma$Nr concentration (Panel A), $\Sigma$Nr deposition fluxes (Panel B), and $\Sigma$Nr emission fluxes (Panel C) on friction velocity ($u_*$ in m s$^{-1}$). All $u_*$ data are from the net deposition period D2 (DOY113–DOY141, cf. Table 1) and are bin-averaged into 0.05 m s$^{-1}$ classes with the black markers representing the arithmetic mean. Red horizontal lines indicate the median, blue horizontal lines indicate lower and upper quartile values, black whiskers represent the interquartile range and outliers from this range are plotted as red crosses.](image)
at lower temporal resolution and at the same time
information of temperature, wind speed and direction are
available, the resistance term \( R \) is commonly composed of
the sum of \( R_a \), \( R_b \), and \( R_c \), representing aerodynamic,
quasi-laminar layer and canopy resistance, respectively. \( R_c \)
may be further split up into several sub-resistances such as
stomatal, mesophyll, cuticle, surface and soil resistances
(Gru¨nhage and Haenel, 1997; Mohr et al., 2005; Sutton
et al., 2007). This detailed consideration, however, has been
neglected in Fig. 8E, where \( g_s \) only partly represents
\( R_c \), but it is clear that systematic diurnal patterns of \( R_a \)
and \( R_b \) are also expected (mostly controlled by \( u^* \)), with highest
\( R_a \) and \( R_b \) values during night-time.

In total, four net \( \Sigma N_r \) emission periods with significantly
different diurnal patterns were identified (Table 1). While
after the first two fertilizations (18 April and 21 May 2008)
\( \Sigma N_r \) fluxes were relatively low (occasionally up to 50 ng N
m\(^{-2}\) s\(^{-1}\)) and the period of net \( \Sigma N_r \) release lasted only for
a few days, emissions after the fertilization on June 6
reached average peak values of around 1000 ng N m\(^{-2}\) s\(^{-1}\) in
the afternoon during the first four days and decreased
to average values of \( \sim 70 \) ng N m\(^{-2}\) s\(^{-1}\) for the rest of
emission period E3 (DOY265 – 278, Fig. 7B). In emission
period E4, \( \Sigma N_r \) fluxes were ranging on a smaller level
between 1 and 20 ng N m\(^{-2}\) s\(^{-1}\). Half-hourly \( \Sigma N_r \) fluxes
during E3 and E4 are plotted as time series in Fig. 9. The
extensive length of emission period E3 (49 days in Fig. 7B)
revealed an N saturation of the ecosystem during that time.
With the wheat plants reaching maturity, considerable
amounts of NH\(_3\) were likely released to the atmosphere
due to higher concentrations of NH\(_3\) in the apoplast than
ambient values of NH\(_3\) concentrations. In contrast, \( \Sigma N_r \)
fluxes after harvest (on DOY218) were significantly lower
(in E4) than those during the second part of E3 as a
consequence of the missing above-ground biomass pathway
for \( \Sigma N_r \).

The relationship between bin-averaged \( u^*\)-data and
\( \Sigma N_r \) fluxes was analyzed for the net deposition period D2
(Fig. 5B and 5C). As was found for \( \Sigma N_r \) concentrations,
both half-hourly \( \Sigma N_r \), deposition and emission fluxes varied
with changing \( u^*\). Highest mean deposition fluxes around
35 ng N m\(^{-2}\) s\(^{-1}\) were measured at \( u^*\)-values of 0.3 m s\(^{-1}\). Duing low \( (u^* < 0.15 \) m s\(^{-1}\)) and high turbulence condi-
tions \( (u^* \geq 0.35 \) m s\(^{-1}\)), mean deposition rates were
significantly lower \( (< 25 \) ng N m\(^{-2}\) s\(^{-1}\)). A similar pattern
was observed for mean emission \( \Sigma N_r \) fluxes. The \( u^*\)-class
for peak emissions, however, was shifted towards a lower
 turbulence regime (0.2 m s\(^{-1}\)) where mean
\( \Sigma N_r \) release was observed to be \( > 40 \) ng N m\(^{-2}\) s\(^{-1}\). These findings
correspond well with the mean diurnal flux patterns shown
in Fig. 7 where highest emission and deposition rates
occurred during the day when \( u^*\)-values are on average
significantly higher than during night-time.

![Fig. 6. Frequency distribution (Panel A) of 30-min \( \Sigma N_r \) fluxes from the entire observation period (December 2007 to November 2008). Vertical lines indicate the estimated flux detection limit \( (\sigma = 3.314 \) ng m\(^{-2}\) s\(^{-1}\)) at \(-\sigma \) and \(+\sigma\). Panel B shows the cumulative sum of the 30-min \( \Sigma N_r \) flux frequency distribution. As indicated by the vertical and horizontal lines, 30.2% of the measured \( \Sigma N_r \) fluxes are within the 2\(\sigma\)-sector around zero exchange, i.e. the point where deposition changes to emission or vice versa. Although single flux values were ranging from \(-166 \) to \(4145 \) ng m\(^{-2}\) s\(^{-1}\), x-axes are truncated at \(-110 \) and \(+110 \) ng m\(^{-2}\) s\(^{-1}\) due to only marginal contributions outside these ranges.](image-url)
3.2.2. Comparison of TRANC-derived $\Sigma N_r$ fluxes with results from inferential modelling. Monthly sums of $\Sigma N_r$ exchange measured by the TRANC-CLD system were compared with results from four inferential models (Fig. 10). These are four dry deposition routines, which are currently used as modules within chemical transport models at national and continental scales in Europe and North America. They include the UK CBED scheme (Smith et al., 2000; Vieno, 2005), the Dutch IDEM model (Erisman et al., 1994; Bleeker et al., 2004; van Jaarsveld, 2004), the dry deposition module of the Environment Canada model (Zhang et al., 2001, 2003), and the surface exchange scheme of the EMEP model used under CLRTAP (Simpson et al., 2003; Tuovinen et al., 2009). The implemented processes of exchange include the aerodynamic transport from the free atmosphere to the leaf or soil surface, the diffusion through the pseudo-laminar boundary layer, the exchange to/from stomates (diffusion through stomatal apertures, uptake/ emission by the apoplast), the transport through the canopy (in-canopy aerodynamic resistance), the non-stomatal deposition to leaf surfaces (with differential treatment for wet and dry conditions) and to soil surfaces (using soil resistance). Dry N deposition simulated in these atmospheric chemistry and transport models is compensated by values of NH$_3$ and NO taken from national emission inventories when applied at regional scale, thereby not directly modelling soil or plant NH$_3$ or NO release except for the CBED scheme, which does account for stomatal NH$_3$ emissions. In our study, we compare results from model applications at field scale where no NH$_3$ and no NO soil emissions were included. All four models use DELTA-based concentrations of NH$_3$, HNO$_2$, HNO$_3$, aerosol NH$_4$ and NO$_3$ as well as NO$_2$ from modelled EMEP 50 km $\times$ 50 km data as input values. Detailed information about these models and an application to 55 European sites is given in Flechard et al. (2011) and references therein.

Measured monthly $\Sigma N_r$ values from the TRANC-CLD system were only slightly negative (up to 0.2 kg N ha$^{-1}$) in the period December 2007 to May 2008. During this time, simulations of the four models resulted in higher uptake.

![Fig. 7. Mean diurnal variation of $\Sigma N_r$ fluxes separated in periods dominated by deposition (Panel A) and periods dominated by emission (Panel B). Due to the logarithmic scale, some negative night-time flux values in a range of $\sim$50 to 0 ng N m$^{-2}$ s$^{-1}$ are missing in Panel B. As for Fig. 3, emission period 3 (E3) was divided into sub-sections by reason of significant changes in exchange characteristics four days after fertilization.](image-url)
rates with monthly peak values from the IDEM-model between 1.0 and 1.7 kg N ha\(^{-1}\). In June, July, and August 2008, monthly net emissions of up to 0.5 kg N ha\(^{-1}\) were measured by the TRANC-CLD system and even higher release rates of up to 1 kg N ha\(^{-1}\) in July 2008 were simulated by the CBED scheme. During these summer months, modelled deposition rates from the three other models (EMEP, IDEM, Environment Canada model) even increased compared to winter and spring values showing monthly uptake of 1.9 kg N ha\(^{-1}\) in July 2008 (IDEM-model). In the following months from September to November 2008, measured and modelled values showed near-neutral exchange to moderate deposition. Overall, monthly deposition rates from all models were always higher than measured values from the TRANC-CLD system with less discrepancy in winter months. The only exceptions were July and August 2008 where simulations of the CBED scheme resulted in higher net monthly emissions than those obtained by the TRANC measurements.

It turned out that models (EMEP, IDEM, Environment Canada model) without the implementation of NO or NH\(_3\) emission routines significantly overestimate the monthly and consequently the annual sum of N\(_2\) deposition at an agricultural site such as Gebesee in comparison to directly measured N\(_2\) exchange using the novel TRANC methodology. On one hand, this might indicate that the compensation values in these models that are taken from NO and NH\(_3\) emission inventories tend to underestimate the actual N\(_2\) exchange thus leading to higher deposition rates. Furthermore, considerable upward fluxes to the atmosphere are likely to be included in the TRANC data throughout the year – either as soil NH\(_3\) release under high pH conditions or soil NO emissions through nitrification/denitrification or possibly as emissions from the wheat plants when concentrations of NH\(_4^+\) in the leaf tissues are somewhat higher than the ‘compensation point’ concentration for NH\(_3\) \((\chi)\). On the other hand, results from the CBED model with a direct soil and plant NH\(_3\) emission routine, corresponded much better with the measured values – at least when averaged over an entire annual course. In this comparison, higher deposition in winter and spring is compensated by higher emissions during summer when N fertilizer additions (see chapter 3.3) lead to considerable N-enrichment in the ecosystem (cf. chapter 3.2.1). Consequently – although not matching single monthly N\(_2\) exchange values – model types like the CBED scheme appear to be more appropriate and realistic in simulating N\(_2\) exchange as both NH\(_3\) uptake and release are directly accounted for, consistent with the eddy-covariance based TRANC measurements, where every single TRANC \(\Sigma N\) flux is a net signal of emission and deposition of \(\Sigma N\). A partitioning into single N\(_2\) compounds (NH\(_3\), HNO\(_3\), NO\(_2\), pNH\(_4\), pNO\(_3\)) taken from modelled ensemble mean monthly N\(_2\) exchange values (Fig. 10B) revealed that for the most part of the year NH\(_3\) formed the largest fraction

---

**Fig. 8.** Mean diurnal variation of \(\Sigma N\) concentration (Panel A), \(\Sigma N\) flux (Panel B), stomatal conductance \((g_s)\) (Panel C), \(\Sigma N\) concentration multiplied by \(g_s\) (Panel D), and the dependency of \(\Sigma N\) flux on \(\Sigma N\) concentration multiplied by \(g_s\) (Panel E) during deposition phase D2 (DOY113–DOY141).
of total \( \Sigma N_r \), i.e. \( \Sigma N_r \) exchange (between \( \sim 40 \) and 80%). The proportion for \( \text{NH}_3 \) was significantly lower during the summer months from June to August 2008, which was likely due to: (1) lower \( \text{NH}_3 \) concentrations than in spring, (2) plant surfaces were drier and thereby reduced non-stomatal uptake as well as (3) higher \( T_a \) generally favours stomatal and other emissions (the compensation point is doubled every additional 5K). Another reason is the fact that the numbers presented in Fig. 10B are monthly ensemble means of all four above-mentioned models including the results from the CBED model, which are showing \( N_r \) emissions as a consequence of \( N \) fertilizer applications in these months and are therefore shifting the \( \text{NH}_3 \) exchange numbers towards a more neutral exchange regime. Furthermore, higher global radiation favours the deposition of NO\(_2\), whereas higher soil temperatures during summer months leads to enhanced soil respiration, elevated soil organic matter mineralization, nitrification and/or denitrification followed by NO emission (e.g. Rosenkranz et al., 2006). The emitted NO is then oxidized to NO\(_2\), thus offsetting the dry deposition gradient of atmospheric NO\(_2\).

3.3. Effect of fertilization on \( \Sigma N_r \) exchange

During the first five months of measurements, small but permanent uptake of \( \Sigma N_r \), with a slightly higher rate from April to May than from January to March, was observed (Fig. 11). The net gain of the ecosystem was \( \sim 0.6 \text{ kg N ha}^{-1} \) by the end of May 2008. Two shifts in the course of cumulative exchange towards net release of \( \Sigma N_r \) to the atmosphere occurred after fertilizer applications on April 18 and May 21 (emission period E1 and E2). These two emission pulses were relatively low compared to the one observed on June 6 corresponding to the third fertilization. \( \Sigma N_r \) fluxes exceeded 4000 ng N m\(^{-2}\) s\(^{-1}\) immediately after fertilizer addition and were still elevated up to 300 ng N m\(^{-2}\) s\(^{-1}\) on the following days with clear diurnal patterns showing highest release in early afternoon (see Fig. 9 showing time series of \( \Sigma N_r \) fluxes during E3 after fertilization and before harvest as well as parts of E4 corresponding to the post-harvest period). \( \Sigma N_r \) fluxes recorded on June 6 and the following few days led to an almost entirely compensation of the \( \Sigma N_r \) uptake measured until early June. As meteorological conditions on all three fertilization days were similar, the influence of temperature, soil

Fig. 9. Half-hourly \( \Sigma N_r \) fluxes following fertilization on 6 June 2008 (DOY158) (Panel A) as well as half-hourly \( \Sigma N_r \) fluxes in selected net emission periods before (Panel B) and after harvest (Panel C).
moisture, leaf surface wetness, radiation or atmospheric turbulence as drivers for the clear differences in $S_N$ fluxes after fertilizer application can largely be excluded. One possible reason for the considerably different exchange behaviour might be the type of mineral fertilizer applied, i.e. the sole use of calcium ammonium nitrate on June 6, which caused relatively high emissions (Figs. 9 and 11) compared to only little $S_N$ release when a mixture of calcium ammonium nitrate and ammonium sulfate was used and the presence of sulfur in the form of $SO_4^{2-}$ presumably inhibited the release of $N_r$ compounds (primarily $NH_3$). However, contrasting results about the

Fig. 10. Comparison of monthly sums of N exchange as determined by the TRANC-CLD system (black line) and by four inferential models (cf. Flechard et al., 2011), i.e. the UK CBED scheme (CB), the dry deposition module of the Environment Canada model (CD), the surface exchange scheme of the EMEP model used under CLRTAP (EM), and the Dutch IDEM model (ID) (Panel A). Panel B shows the monthly ensemble means of the four models displayed in Panel A with each monthly value being composed of the compounds $NH_3$, $HNO_3$, $NO_2$, particulate $NH_4$, and particulate $NO_3$. Standard deviations of each compound based on these deposition rates are shown in Panel C.

Fig. 11. Cumulative net $S_N$ exchange during the entire observation period from December 2007 to November 2008. The red line indicates the cumulative net $S_N$ loss to the atmosphere during net emission periods, the green line indicates the cumulative net $S_N$ gain of the plant–soil system during net deposition periods, and the blue line represents the cumulative total net $S_N$ exchange. Periods D1–D5 and E1–E4 correspond to naming in Table 1.
emission potential of these fertilizer types were reported in former studies. While Jarvis et al. (1989), Shahandeh et al. (1992) and the FAO (2001) support the reasoning that lower emissions were – at least partly – due to sulfate-containing fertilizer, other authors (e.g. Asman, 1992; Buijsman, 1986; Aneja et al., 2003) report on higher emission factors (EF) for ammonium sulfate (9.7%) compared to ammonium nitrate (2.5%). Thus, it is more likely that the timing and developmental stage of the wheat plants had a considerable influence on the amount of $\Sigma N_r$ released on June 6. While fertilizer N was possibly taken up by the wheat plants with high efficiency during the main growth phase in April and May, $N_r$ could be more readily volatilized in June when the wheat plants were already in a mature state. Similar observations, for example, of $NH_3$ emissions from senescing foliage compared with growing leaves have been shown in studies from Farquhar et al. (1979), Harper et al. (1987) and Husted et al. (1996). Also, canopy capture of soil- and fertilizer-emitted $NH_3$ might have played a role in the variant exchange characteristics after fertilization – particularly between those in April and May where an almost three times higher fertilizer amount applied in May (82.7 kg N ha$^{-1}$) did not lead to significantly elevated emissions compared to the fertilization in April (28.5 kg N ha$^{-1}$). According to studies of Nemitz et al. (2000), Denmead et al. (2008) and Bash et al. (2010), closer plant canopies – as was the case in May – are likely to recapture more $NH_3$ than open plant canopies. However, for the June fertilization with a medium quantity of 44.6 kg N ha$^{-1}$ applied, but followed by the substantial emission rates, maturity effects must have been predominated over canopy recapture effects.

Another emission phase in mid July was observed without prior fertilization (second part of E3, Fig. 11). As net $N_r$ release to the atmosphere, mainly in the form of $NH_3$, has been reported earlier for both agricultural croplands (Sutton et al., 2000) and semi-natural vegetation, e.g. at a dry Dutch heathland site (Erisman and Wyers, 1993), the observation of $N_r$ emission in mid-July at the Geibsee site might be the result of N saturation, raising $\chi_r$ and limiting net deposition, thus being itself an ecosystem response indicating exceedance of the instantaneous crop nitrogen requirement. Also, this short but significant emission phase occurred during a couple of successive dry days, i.e. under conditions favouring emissions in saturated systems as opposed to wet leaf surfaces, which are known for rapid uptake of $NH_3$ (Aneja et al., 1986; Duyzer et al., 1994) due to the large solubility of ammonia in water.

Direction and amplitude of $N_r$ exchange in periods where no measurements were available due to system breakdowns remain speculative. We assume that almost certainly small $N_r$ uptake would have been registered following the commonly observed non-fertilized pattern, particularly before June 2008 when the biosphere–atmosphere exchange was dominated by $N_r$ input into the ecosystem and a supposed saturation state had not been reached.

The cumulative net $\Sigma N_r$ exchange between the wheat field and the atmosphere by the end of the observation period (Fig. 11) resulted in very low deposition of 0.072 kg N ha$^{-1}$. This value is not significantly different from zero and appears to be small given the fact that expected emissions calculated on the basis of literature emission factors of $NH_3$ accounting for the respective amount and mixture of fertilizer types (Jarvis et al., 1989; Shahandeh et al., 1992; FAO, 2001; Aneja et al., 2003) plus reported emissions of NO on fertilized wheat fields (Johansson and Granat, 1984; Ludwig et al., 2001) would lead to a range between 2.7 and 12.1 kg N ha$^{-1}$. However, subtracting the background dry deposition of $\sim$4.1 to 11.75 kg N ha$^{-1}$ as was modelled by Flechard et al. (2011) (cf. Section 3.2.2), the net $\Sigma N_r$ exchange measured by the TRANC-CLD system fits reasonably into the balance between fertilizer-induced $N_r$ emissions and year-round background dry $N_r$ deposition.

Bearing in mind that TRANC-measured fluxes represent the net exchange, i.e. the balance between emission and deposition by turbulent processes, these values cannot be directly compared with pure deposition values determined in other studies, which are usually reported as total, i.e. dry plus wet N deposition. Though highly variable for different land use types and surface characteristics, mean N deposition in Germany was estimated to range between 15 and 18 kg N ha$^{-1}$ yr$^{-1}$ (Gauger et al., 2002; Heinzeller, 2007). N deposition rates at agricultural sites (20 to 48 kg N ha$^{-1}$ yr$^{-1}$) are mainly driven by management intensity and distance to pig and poultry farms, in the vicinity of which values up to 85 kg N ha$^{-1}$ yr$^{-1}$ (Bartels and Wagener, 2005) can be found. Forested areas, particularly those located at edges near adjacent source areas such as cities or agricultural land, also exhibit large N deposition rates due to higher canopy roughness. Friedmann et al. (2001) report on a value of $\sim$32 kg N ha$^{-1}$ yr$^{-1}$ (Bartels and Wagener, 2005) can be found. Forested areas, particularly those located at edges near adjacent source areas such as cities or agricultural land, also exhibit large N deposition rates due to higher canopy roughness. Friedmann et al. (2001) report on a value of $\sim$32 kg N ha$^{-1}$ yr$^{-1}$ for Bavarian forest sites, whereas Mohr et al. (2005) measured 57 kg N ha$^{-1}$ yr$^{-1}$ (13 and 44 kg N ha$^{-1}$ yr$^{-1}$ attributed to bulk and dry deposition, respectively) above a pine forest in Northern Germany.

Regarding the $\Sigma N_r$ exchange pattern and fluxes in Geibsee (Fig. 11), it seems likely that due to the low values found, $N_r$ emissions – either in the form of $NH_3$ after fertilization or as NO release from soil as an intermediate product of nitrification – compensate for a significant part of the $N_r$ deposition. This also shows that the co-occurrence of both stomatal exchange and leaf surface uptake on the one hand, and soil emissions on the other hand, complicate the interpretation of net $N_r$ flux.
measurements. However, as continuous long-term measurements of any N\textsubscript{r} species are extremely rare, ΣN\textsubscript{r} data provided by the TRANC-CLD system are nevertheless of major importance for N\textsubscript{r} budget estimations, possible revisions of N\textsubscript{r} emission factors, and for assessing the productivity of ecosystems by its nutrient N input.

4. Conclusions

With the development of the TRANC, a novel methodology has been provided for both quantifying ΣN\textsubscript{r} concentrations and a device being robust and fast enough to be applicable in an EC setup for the measurement of ΣN\textsubscript{r} fluxes when combined with a fast response NO detector (CLD). In this study, we present the first long-term data set of ΣN\textsubscript{r} net exchange above arable land demonstrating the importance of continuous measurements with high temporal resolution for a better understanding of ecosystem functioning and its controlling parameters. Reasonable agreement and similar seasonal trends were found in a comparison between monthly ΣN\textsubscript{r} concentrations measured by TRANC and DELTA denuders and filters. Differences could be explained by missing NO\textsubscript{x} determination in DELTA measurements. Monthly ΣN\textsubscript{r} net exchange measured by TRANC was fairly in the range of monthly N\textsubscript{r} deposition rates obtained from dry deposition inferential modelling. As TRANC values were closest to those from the UK CBED scheme, the significant differences among the four models used were likely due to inadequate treatment of ecosystem N\textsubscript{r} emissions in the applied inferential model routines. Distinctive diurnal patterns in ΣN\textsubscript{r} concentrations and fluxes were observed to be dependent on turbulence, stomatal regulation and non-stomatal sinks for NH\textsubscript{3}. At the end of the observation period, the net exchange of ΣN\textsubscript{r} at the Gebesee site was quasi neutral (0.072 kg N ha\textsuperscript{-1} deposition) with small net deposition through most parts of the year and substantial emissions after one particular fertilization stressing the likely influence of fertilizer type, environmental conditions and maturity stage of the wheat plants during field application. Offsetting emission factor-based calculations of N\textsubscript{r} release (~2.7–12.1 kg N ha\textsuperscript{-1}) by modelled background dry deposition (~4.1–11.75 kg N ha\textsuperscript{-1}), the cumulative TRANC-derived exchange provides a reasonable estimate of the site’s annual ΣN\textsubscript{r} budget. However, more research and field campaigns like the present one are needed to: (1) conduct this type of measurement in different ecosystems to better understand ΣN\textsubscript{r} dynamics on broader scales and (2) possibly further develop the TRANC methodology to be able to differentiate between oxidized and reduced N species to identify factors and drivers of quantitative N\textsubscript{r} exchange between biosphere and atmosphere, e.g. NO\textsubscript{x} contributions from traffic/industry or NH\textsubscript{3} from agricultural sources.

5. Acknowledgements

This work was supported by the EU project NitroEurope-IP (Contract 017841) under the EC 6\textsuperscript{th} Framework Program for Research and Technological Development and by the Max Planck Institute for Biogeochemistry (MPI-BGC) in Jena, Germany. We gratefully acknowledge Geratal Agrar GmbH (Andisleben, Thuringia), the owners of the field, and the city council for permitting research activities at the Gebesee site. Dirk Lempio and Catharina Don are thanked for technical support and field assistance. We highly appreciate the help of Jürgen Conrad, Kerstin Gilke and the lab team of the Institute of Agricultural Climate Research (vTI-AK) in Braunschweig, Germany, for running and maintaining the DELTA denuder system.

References

Aber, J., McDowell, W., Nadelhoffer, K., Magill, A., Berntson, G. and co-authors. 1998. Nitrogen saturation in temperate forest ecosystems: hypotheses revisited. Bioscience. 48, 921–934.
Ammann, C., Brunner, A., Spirig, C. and Neftel, A. 2006. Technical note: water vapour concentration and flux measurements with PTR-MS. Atmos. Chem. Phys. 6, 4643–4651.
Ammann, C., Wolff, V., Marx, O., Brümmer, C. and Neftel, A. 2012. Measuring the biosphere-atmosphere exchange of total reactive nitrogen by eddy covariance. Biogeosciences. 9, 4247–4261.
Aneja, V. P., Nelson, D. R., Roelle, P. A., Walker, J. T. and Battye, W. 2003. Agricultural ammonia emissions and ammonia concentrations associated with aerosols and precipitation in the southeast United States. J. Geophys. Res. 108(D4), 4152. DOI: 10.1029/2002JD002271.
Aneja, V. P., Rogers, H. H. and Stahel, E. P. 1986. Dry deposition of ammonia at environmental concentrations on selected plant species. J. Air Pollut. Contr. Ass. 36, 1338–1341.
Anthoni, P. M., Freibauer, A., Kolle, O. and Schulze, E.-D. 2004. Winter wheat carbon exchange in Thuringia, Germany. Agric. For. Meteorol. 121, 55–67.
Asman, W. A. H. 1992. Ammonia Emission in Europe: Updated Emission and Emission Variations. National Institute of Public Health and Environmental Protection (RIVM), Bilthoven, The Netherlands. Report No. 228471008.
Aubinet, M., Grelle, A., Ibrom, A., Rannik, Ü., Moncrieff, J. and co-authors. 2000. Estimates of the annual net carbon and water exchange of forests: the EUROFLUX methodology. Adv. Ecol. Res. 30, 114–177.
Bartels, U. and Wagener, S. 2005. Nitrogen Deposition in the Münsterland Region (in German). Reports of the State Environmental Agency, Münster, 21 pp.
Bash, J. O., Walker, J. T., Jones, M., Katul, G., Nemitz, E. and co-authors. 2010. Estimation of in-canopy ammonia sources and
sinks in a fertilized *Zea mays* field. *Environ. Sci. Technol.* **44**, 1683–1689.

Bleeker, A., Reinds, G. J., Vermeulen, A. T., de Vries, W. and Erisman, J. W. 2004. *Critical Loads and Present Deposition Thresholds of Nitrogen and Acidity and Their Exceedances at the Level II and Level I Monitoring Plots in Europe*. Energy research Centre of the Netherlands, Petten, The Netherlands. ECN report ECN-C-04-117.

Brümmer, C., Black, T. A., Jassal, R. S., Grant, N. J., Spittlehouse, D. L. and co-authors. 2012. How climate and vegetation type influence evapotranspiration and water use efficiency in Canadian forest, peatland and grassland ecosystems. *Agric. For. Meteorol.* **153**, 14–30.

Buijsman, E. 1986. *Historical Trends in Ammonia Emission in Europe (1870–1980)*. Institute for Meteorology and Oceanography, State University of Utrecht, The Netherlands. Report R-86-9.

Day, D. A., Wooldridge, P. J., Dillon, M. B., Thornton, J. A. and Cohen, R. C. 2002. A thermal dissociation laser-induced fluorescence instrument for in situ detection of NO2, peroxo nitrates, alkyl nitrates, and HNO3. *J. Geophys. Res.* **107**(D6). DOI: 10.1029/2001JD000779.

Denmead, O. T., Freney, J. R. and Dunin, F. X. 2008. Gas exchange between plant canopies and the atmosphere: case-studies for ammonia. *Atmos. Environ.* **42**, 3394–3406.

Duyzer, J. H., Verhagen, H. L. M., Westraete, J. H., Bosveld, F. C. and Vermetten, A. W. M. 1994. The dry deposition of ammonia onto a Douglas-fir forest in the Netherlands. *Atmos. Environ.* **28**, 1241–1253.

Erisman, J. W., Mennen, M. G., Fowler, D., Flechard, C. R., Spindler, G. and co-authors. 1996. *Towards Development of a Deposition Monitoring Network for Air Pollution in Europe*. RIVM, The Netherlands. Report no. 722108015. Online at: http://rivm.openrepository.com/rivm/bitstream/10029/10432/1/722108015.pdf.

Erisman, J. W., Otjes, R., Hensen, A., Jongejan, P., van den Bulk, P. and co-authors. 2001. Instrument development and application in studies and monitoring of ambient ammonia. *Atmos. Environ.* **35**, 1913–1922.

Erisman, J. W., van Pul, A. and Wyers, P. 1994. Parameterization of surface resistance for the quantification of atmospheric deposition of acidifying pollutants and ozone. *Atmos. Environ.* **28**, 2595–2607.

Erisman, J. W. and Wyers, G. P. 1993. Continuous measurements of surface exchange of SO2 and NH3: implications for their possible interaction in the deposition process. *Atmos. Environ.* **27**, 1937–1949.

Falge, E., Baldocchi, D., Olson, R., Anthoni, P., Aubinet, M. and co-authors. 2001. Gap filling strategies for long term energy flux data sets. *Agric. For. Meteorol.* **107**, 71–77.

FAO (Food and Agriculture Organization of the United Nations) and IFA (International Fertilizer Industry Association). 2001. *Global Estimates of Gaseous Emissions of NH3, NO and NO2 from Agricultural Land*. International Fertilizer Industry Association, Rome, Italy. ISBN 92-5-104689-1.

Farquhar, G. D., Wetselaar, R. and Firth, P. M. 1979. Ammonia volatilization from senescing leaves of maize. *Science*. **203**, 1258–1275.

Flechard, C. R., Nemitz, E., Smith, R. I., Fowler, D., Vermeulen, A. T. and co-authors. 2011. Dry deposition of reactive nitrogen to European ecosystems: a comparison of inferential models across the NitroEurope network. *Atmos. Chem. Phys.* **11**, 2703–2728.

Foken, T. 2008. *Micrometeorology*. Springer, Heidelberg, Germany, 308 pp.

Fowler, D., Coyle, M., Flechard, C., Hargreaves, K. J., Nemitz, E. and co-authors. 2001. Advances in micrometeorological methods for the measurement and interpretation of gas and particle nitrogen fluxes. *Plant Soil*. **228**, 117–129.

Francis, D. D., Vigil, M. F. and Mosier, A. R. 2008. Gaseous losses of nitrogen other than through denitrification. In: *Nitrogen in Agricultural Systems* (eds J. S. Schepers and W. R. Raun). American Society of Agronomy, Madison, WI, pp. 255–280.

Friedmann, L., Rosenwirth, W. and Meier, D. 2001. *Nitrate Report Bavaria, 1996–1999* (in German). Reports of the Bavarian State Office for Water Management, Munich, 35 pp.

Gauger, T. 2009. *Acquisition, Projection and Assessment of Matter Fluxes and Their Impacts on Ecosystems in Germany* (in German). Final report of the MAPESI project ‘Modelling of Air Pollutants and EcoSystem Impact’ by order of the German Federal Environment Agency (UBA), Grant No. 3707 64 200, 32 pp.

Gauger, T., Anshelm, F., Schuster, H., Draaisjers, G. P. J., Bleeker, A. and co-authors. 2002. *Mapping of Ecosystem Specific Long-Term Trends in Deposition Loads and Concentrations of Air Pollutants in Germany and Their Comparison with Critical Loads and Critical Levels. Part 1: Deposition Loads 1990–1999*. Institute of Navigation, Stuttgart University. BMU/UBA, FE No. 299 42 210, 207 pp.

Grünhage, L. and Haenel, H.-D. 1997. PLATIN (PLant-ATmosphere INteraction) I: a model of plant-atmosphere interaction for estimating absorbed doses of gaseous air pollutants. *Environ. Pollut.* **98**, 37–50.

Harper, L. A., Sharpe, R. R., Langdale, G. W. and Giddens, J. E. 1987. Nitrogen cycling in a wheat crop: soil, plant and aerial nitrogen transport. *Agron. J.* **79**, 965–973.

Heinzeller, C. 2007. *Regionalization of Nitrate Export from Forests in Germany* (in German). PhD Thesis. University of Freiburg, Freiburg, Germany.

Husted, S., Mattsson, M. and Schjoerring, J. K. 1996. Ammonia compensation points in two cultivars of *Hordeum vulgare* L. during vegetative and generative growth. *Plant. Cell. Environ.* **19**, 1299–1306.

van Jaarsveld, J. A. 2004. *The Operational Priority Substances Model*. Description and validation of OPS-Pro 4.1. RIVM, Bilthoven, The Netherlands. RIVM report 500045001/2004.
Jarvis, S. C., Hatch, D. J. and Lockyer, D. R. 1989. Ammonia fluxes from grazed grassland – annual losses from cattle production systems and their relation to nitrogen inputs. J. Agric. Sci. 113, 99–108.

Johansson, C. and Granat, L. 1984. Emission of nitric oxide from arable land. Tellus. 36B, 25–37.

Kita, K., Morino, Y., Kondo, Y., Komazaki, Y., Takegawa, N. and co-authors. 2006. A chemical ionization mass spectrometer for ground-based measurements of nitric acid. J. Atmos. Ocean. Tech. 23, 1104–1113.

Kutsch, W. L., Aubinet, M., Buchmann, N., Smith, P., Osborne, B. and co-authors. 2010. The net biome production of full crop rotations in Europe. Agr. Ecosystem. Environ. 139, 336–345.

Kutsch, W. L., Hanan, N., Scholes, B., McHugh, I., Kuhheka, W. and co-authors. 2008. Regulation of carbon fluxes and canopy conductance in a savanna ecosystem. Biogeosciences. 5, 1797–1808.

Lu, P., Yunusa, I. A. M., Walker, R. R. and Müller, W. J. 2003. Regulation of canopy conductance and transpiration and their modelling in irrigated grapevines. Funct. Plant Biol. 30, 689–698.

Ludwig, J., Meixner, F. X., Vogel, B. and Förstner, J. 2001. Soil–air exchange of nitric oxide: an overview of processes, environmental factors, and modelling studies. Biogeochemistry. 52, 225–257.

Marx, O., Brümmer, C., Ammann, C., Wolff, V. and Freibauer, A. 2012. TRANC – a novel fast-response converter to measure total reactive atmospheric nitrogen. Atmos. Meas. Tech. 5, 1045–1057.

Moffat, A. M., Papale, D., Reichstein, M., Hollinger, D. Y., Richardson, A. D. and co-authors. 2007. Comprehensive comparison of gap-filling techniques for eddy covariance net carbon fluxes. Agric. For. Meteorol. 147, 209–232.

Möh, K., Meesenburg, H., Horváth, B., Meiwes, K. J., Schaaf, S. and co-authors. 2005. Determination of Ammonia Input from the Air and Its Effect on Forest Ecosystems (in German). Landbauforschung Völkenrode – FAL Agricultural Research. Braunschweig Völkenrode. Special Issue 279, ISSN 0376-0723, 103 pp.

Munger, J. W., Wofsy, S. C., Bakwin, P. S., Fan, S.-M., Goulden, M. L. and co-authors. 1996. Atmospheric deposition of reactive nitrogen oxides and ozone in a temperate deciduous forest and a subarctic woodland. 1. Measurements and mechanisms. J. Geophys. Res. 101(D7), 639–657.

Nemitz, E., Sutton, M. A., Gut, A., San Jose, R., Husted, S. and co-authors. 2000. Sources and sinks of ammonia within an oilseed rape canopy. Agric. For. Meteorol. 105, 385–404.

Ollinger, S. V., Aber, J. D., Reich, P. B. and Freuder, R. J. 2002. Interactive effects of nitrogen deposition, tropospheric ozone, elevated CO₂ and land use history on the carbon dynamics of northern hardwood forests. Glob. Change Biol. 8, 545–562.

Rogers, H. H. and Aneja, V. P. 1980. Uptake of atmospheric ammonia by selected plant-species. Environ. Exp. Bot. 20, 251–257.

Rosenkranz, P., Brüggemann, N., Papen, H., Xu, Z., Horváth, L. and co-authors. 2006. Soil N and C trace gas fluxes and microbial soil N turnover in a sessile oak (Quercus petraea (Matt.) Lieb.) forest in Hungary. Plant Soil. 286, 301–322.

Rummel, U., Ammann, C., Gut, A., Meixner, F. X. and Andreae, M. O. 2002. Eddy covariance measurements of nitric oxide flux within an Amazonian rain forest. J. Geophys. Res. 107, 8050. DOI: 10.1029/2001JD000520.

Schjoerring, J. K., Husted, S. and Mattsson, M. 1998. Physiological parameters controlling plant-atmosphere ammonia exchange. Atmos. Environ. 32, 491–498.

Schmid, H. P. 1993. Source areas for scalars and scalar fluxes. Boundary-Layer Meteorol. 67, 293–318.

Seinfeld, J. H. and Pandis, S. N. 2006. Atmospheric Chemistry and Physics – From Air Pollution to Climate Change. 2nd ed. John Wiley & Sons, Inc., Hoboken, New Jersey, 1232 pp.

Shahandeh, H., Cabrera, M. L. and Sumner, M. E. 1992. Ammonia volatilization from urea, ammonium sulfate, and nutrasweet sludge in bare and straw-covered soils. Commun. Soil Sci. Plant Anal. 23, 775–786.

Simpson, D., Butterbach-Bahl, K., Fagerli, H., Kesik, M., Skiba, U. and co-authors. 2006. Deposition and emissions of reactive nitrogen over European forests: a modelling study. Atmos. Environ. 40, 5712–5726.

Simpson, D., Fagerli, H., Jonson, J. E., Tsyro, S., Wind, P. and co-authors. 2003. Transboundary Acidification, Eutrophication and Ground Level Ozone in Europe. Part I: Unified EMEP Model Description. Det Meteorologisk Institutt, Oslo. EMEP Status Report, ISSN 0806–4520.

Sintermann, J., Spirig, C., Jordan, A., Kuhn, U., Ammann, C. and co-authors. 2011. Eddy covariance flux measurements of ammonia by high temperature chemical ionisation mass spectrometry. Atmos. Meas. Tech. 4, 599–616.

Smith, R. I., Fowler, D., Sutton, M. A., Flechard, C. and Coyle, M. 2000. Regional estimation of pollutant gas deposition in the UK: model description, sensitivity analyses and outputs. Atmos. Environ. 34, 3757–3777.

Sutton, M. A., Milford, C., Nemitz, E., Theobald, M. R., Hill, P. W. and co-authors. 2001. Biosphere-atmosphere interactions of ammonia with grasslands: experimental strategy and results from a new European initiative. Plant Soil. 228, 131–145.

Sutton, M. A., Nemitz, E., Erisman, J. W., Beier, C., Butterbach-Bahl, K. and co-authors. 2007. Challenges in quantifying biosphere-atmosphere exchange of nitrogen species. Environ. Pollut. 150, 125–139.

Sutton, M. A., Nemitz, E., Milford, C., Fowler, D., Moreno, J. and co-authors. 2000. Micrometeorological measurements of net ammonia fluxes into oilseed rape during two vegetation periods. Agric. For. Meteorol. 105, 351–369.

Tang, Y. S., Simmons, I., van Dijk, N., Di Marco, C., Nemitz, E. and co-authors. 2009. European scale application of atmospheric reactive nitrogen measurements in a low-cost approach to infer dry deposition fluxes. Agr. Ecosystem. Environ. 133, 183–195.

Tuovinen, J.-P., Emberson, L. and Simpson, D. 2009. Modelling ozone fluxes to forests for risk assessment: status and prospects. Ann. Forest Sci. 66(4), 401.
Vieno, M. 2005. *The Use of an Atmospheric Chemistry-Transport Model (FRAME) over the UK and the Development of Its Numerical and Physical Schemes*. PhD Thesis. University of Edinburgh, Edinburgh.

Vitousek, P. M., Aber, J. D., Howarth, R. W., Likens, G. E., Matson, P. A. and co-authors. 1997. Human alteration of the global nitrogen cycle: sources and consequences. *Ecol. Appl.* 7, 737–750.

Zhang, L., Brook, J. R. and Vet, R. 2003. A revised parameterization for gaseous dry deposition in air-quality models. *Atmos. Chem. Phys.* 3, 2067–2082.

Zhang, L., Gong, S., Padro, J. and Barrie, L. 2001. A size-segregated particle dry deposition scheme for an atmospheric aerosol module. *Atmos. Environ.* 35, 549–560.

Zhang, L., Vet, R., O’Brien, J. M., Mihele, C., Liang, Z. and co-authors. 2009. Dry deposition of individual nitrogen species at eight Canadian rural sites. *J. Geophys. Res.* 114, D02301. DOI: 10.1029/2008JD010640.

Zimmermann, F., Plessow, K., Queck, R., Bernhofer, C. and Matschullat, J. 2006. Atmospheric N- and S-fluxes to a spruce forest – comparison of inferential modelling and the throughfall method. *Atmos. Environ.* 40, 4782–4796.