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Determination of field scale ammonia emissions for common slurry spreading practice with two independent methods

J. Sintermann¹, C. Ammann¹, U. Kuhn^{1,*}, C. Spirig^{1,**}, R. Hirschberger², A. Gärtner³, and A. Neftel¹

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Abstract. At a cropland and a grassland site field scale ammonia (NH₃) emissions from slurry application were determined simultaneously by two approaches based on (i) eddy covariance (EC) flux measurements using high temperature Chemical Ionisation Mass Spectrometry (HT-CIMS) and on (ii) backward Lagrangian Stochastic (bLS) dispersion modelling using concentration measurements by three optical open path Fourier Transform Infrared (FTIR) systems. Slurry was spread on the fields in sequential tracks over a period of one to two hours. In order to calculate field emissions, measured EC/HT-CIMS fluxes were combined with flux footprint analysis of individual slurry spreading tracks to parameterise the NH₃ volatilisation with a bi-exponential time dependence. Accordingly, track-resolved concentration footprints for the FTIR measurements were calculated using bLS. A consistency test with concentrations measured by impingers showed very low systematic deviations for the EC/HT-CIMS results (<8 %) but larger deviations for the bLS/FTIR results. For both slurry application events, the period during fertilisation and the subsequent two hours contributed by more than 80 % to the total field emissions. Averaged over the two measurement methods, the cumulated emissions of the first day amounted to $17 \pm 3\%$ loss of applied total ammoniacal nitrogen over the cropland and $16 \pm 3\%$ over the grassland field.



Correspondence to: J. Sintermann (joerg.sintermann@art.admin.ch)

1 Introduction

The growing demand for food and energy products has lead to highly intensified agriculture with increasing emissions of nitrogen-containing compounds that pose environmental risks. One of the particularly important trace gas species in emissions associated with agriculture is ammonia (NH₃) (Aneja et al., 2008). This anthropogenic NH₃ release contributes to a large extent to the harmful effects of high reactive nitrogen loads (Galloway et al., 2003; Erisman et al., 2007). In central Europe, agricultural NH3 volatilisation accounts for more than 90% of the release (Erisman et al., 2008; Reis et al., 2009) and NH₃ emissions following organic livestock waste application on fields have been identified to amount for roughly a third to half of the agricultural NH₃ losses (Reidy et al., 2008a,b; EEA, 2009). A detailed quantification of NH₃ emissions with high accuracy is essential for a better knowledge about the factors controlling NH₃ volatilisation after application of organic fertiliser (Erisman et al., 2008; Zhang et al., 2008). Such measurements are vital for the characterisation of the agricultural nitrogen budget (Ammann et al., 2009) as well as to link emissions and monitoring, and hence to assess abatement strategies (Bleeker et al., 2009; Erisman et al., 2009).

In the literature, a very large range of NH₃ loss factors related to the application of slurry to agricultural surfaces (as percentage of the applied total ammoniacal nitrogen: TAN) is found. It comprises values between 4 % to almost 100 % (e.g. Pain et al., 1989; Braschkat et al., 1997; Vandre et al., 1997; Génermont et al., 1998; Menzi et al., 1998; Huijsmans et al., 2001; Søgaard et al., 2002; Huijsmans et al., 2003; Misselbrook et al., 2002, 2005a; Sanz et al., 2010; Spirig et al., 2010; Uusi-Kämppä and Mattila, 2010). The strong

¹Swiss Federal Research Station Agroscope Reckenholz-Tänikon ART – Air Pollution and Climate, Zürich, Switzerland

²Engineering consultants Niemann and Partner, University of Bochum, Bochum, Germany

³Northrhine-Westphalian Agency for Nature, Environment and Consumer Protection, Recklinghausen, Germany

^{*}now at: Institut für Energie- und Klimaforschung Troposphäre (IEK-8) Forschungszentrum Jülich, Jülich, Germany

^{**}now at: Swiss Federal Office of Meteorology and Climatology MeteoSwiss, Zürich, Switzerland

stickiness of the polar NH3 molecule complicates measurements of ambient NH3 concentrations (Parrish and Fehsenfeld, 2000; von Bobrutzki et al., 2010) and consequently fluxes. Varying instrumental performance (Milford et al., 2009; Sutton et al., 2009) and limited signal range of sensors (Spirig et al., 2010) introduce large uncertainties into NH₃ emission quantifications. The possibility for accurate field scale assessments under common agricultural practice is important for the validation of emission levels (Sommer et al., 2003; Spirig et al., 2010). Agricultural practice means that the slurry spreading is not performed instantaneously, but as a sequence of dispersals over a period of typically one to several hours. Thus, the fertilised field does not represent a homogeneous area source for NH3. In addition, the emission rate of freshly applied slurry can show a fast decrease (Sintermann et al., 2011). These spatial and temporal inhomogeneity effects have to be considered when evaluating emission losses on the field scale.

In the present paper, we report on two experiments in August 2009, devoted to quantify the field scale NH₃ emissions associated with spreading of slurry in high temporal resolution. We present two methods to determine these emissions. The first is based on direct flux measurements by the eddy covariance (EC) method using fast high temperature Chemical Ionisation Mass Spectrometry (HT-CIMS) with a high temperature inlet line (Sintermann et al., 2011) in combination with a detailed flux footprint attribution. The HT-CIMS instrument derives from the common Proton Transfer Reaction-Mass Spectrometer (PTR-MS) and uses electron transfer reactions for NH3 ionisation. The second approach uses open path line concentration measurements by optical Fourier Transform Infrared (FTIR) systems in combination with a backward Lagrangian Stochastic (bLS) dispersion model (Flesch et al., 2004) for concentration footprint attribution. For NH₃ flux measurements, the EC/HT-CIMS approach has the advantage that wall interaction are minimised by strong heating of all surfaces and their impact on the EC flux can be quantified. The bLS/FTIR method is based on inlet-free measurements and links the downwind NH₃ concentration to its source. The two methods have been applied simultaneously on two slurry spreading events, one on a cropland and the other on a grassland field in Oensingen, Switzerland. The results of both approaches are intercompared and discussed in view of previous micrometeorological measurements at the site with a wet chemical gradient system (Spirig et al., 2010).

2 Methods

2.1 Analytical techniques for NH₃ detection

2.1.1 HT-CIMS

The HT-CIMS, based on PTR-MS (Hansel et al., 1995; Lindinger et al., 1998), is a chemical ionisation technique making use of electron transfer reactions to on-line ionise continuously sampled gas with subsequent detection of selected ion products (Norman et al., 2007, 2009). It operates with positively charged oxygen instead of protonated water as a source for charge transfer. A detailed description of the instrumental principles is given by Norman et al. (2007). In order to obtain a fast time response (\sim 1 s) for NH₃ we employed the instrument in a modified way including an adopted inlet scheme and a prototype drift tube of reduced volume and altered materials, heated to 180 °C. Measuring ambient NH₃ concentrations and EC fluxes, the HT-CIMS sub-sampled gas at the downstream end of a 23 m 1/2" PFA (OD) tube, heated to 150 °C and flushed with 100 STP 1min^{-1} . The air intake was located 1 m above ground level (m a.g.l.) adjacent to an ultrasonic anemometer (HS Research Anemometer, Gill Instruments Ltd, Lymington, UK) mounted at a height of 1.25 m a.g.l. The operation of the instrument and the entire flux setup, along with analyses confirming a sufficiently fast time response of the system for EC flux measurements are described by Sintermann et al. (2011). The HT-CIMS was calibrated before every fertilisation against an NH3 permeation device (LN Industries, Geneva, Switzerland) equipped with a permeation tube (VICI, Metronics Inc., Poulsbo, WA, USA). The permeation rate was determined with impingers.

2.1.2 FTIR

With the purpose to monitor the NH₃ concentration profile downwind of the slurry emissions, three open path FTIR systems (K300, Kayser-Threde GmbH, München, Germany) in bi-static configuration were installed at three heights: 0.8, 1.8 and 3.0 m a.g.l. on 4 August 2009 and 0.8, 1.9 and 3.0 m a.g.l. on 6 August 2009, respectively. Path integrated NH₃ concentrations with a time resolution of about 2 min over lengths of 109 m on 4 August and 58 m on 6 August were obtained. The FTIR measurements (e.g. Gärtner et al., 2008) are based on interferometric analysis of infrared NH3 absorption spectra using Fourier transformation (Hirschberger, 2000). The specific configuration of the applied systems is described in detail by Heise et al. (2001). The detecting units were of MCT (Mercury Cadmium Telluride) and light source was a GLOBAR (glowing bar, silicium carbide) operated at 1500 °C, resulting in broad band infrared irradiation. The systems were calibrated 2 weeks prior to the experiment. This was done with a multi-reflection cell (white cell, Bastian Feinmechanik, Wuppertal, Germany) using calibration

position sample height sample time amount of sampled air NH_3 $[\mu g \, m^{-3}]$ [CET] [moles] [m a.g.l.] 4 August 2009 0.45/1.45 12:47-14:50 5.41/5.73 688/365 (a) (a) 0.45/1.45 14:58-18:55 10.42/10.34 153/79 12:55-15:05 0.45/1.45 5.61/5.35 649/303 (b) 0.45/1.45 15:06-19:12 10.61/10.13 (b) 131/68 6 August 2009 (a) 0.45/1.45 10:15-12:30 6.02/5.72 660/320 (a) 0.45/1.45 12:35-16:15 9.82/9.23 141/52 (b) 0.45/1.45 10:17-12:37 6.18/6.00 728/261 (b) 0.45 14:23-16:35 5.83 82 29 (b) 1.45 12:38-16:35 10.15

Table 1. Impinger measurements; positions as displayed in Fig. 1.

gas (Messer, Griessheim, Germany). Concentration calculations were performed by the software CLSEVAL (Müller, 2000). The detection limit for NH₃ was about 15 μ g m⁻³ and the relative accuracy was estimated to better than 10 %.

2.1.3 Cavity ring-down spectroscopy

As with FTIR spectroscopy, cavity ring-down spectroscopy utilises the light absorption of NH₃ in the infrared, measuring the ring-down time of a multiple reflected laser pulse (Berden et al., 2000). The instrument (G1103; Picarro Inc., California, USA) was housed in an air-conditioned container sampling ambient air at 5 m a.g.l. A 9.5 m long 1/4" OD PFA inlet tube was provided with a PTFE membrane filter (5 µm pore size; Whatman Ltd, Maidstone, UK) and the instrument sampled at a flow rate of 0.5 STP 1 min⁻¹. The analyser was run continuously during the experimental period to monitor NH₃ background levels recording data in 3 s intervals. In field measurements, a setup including a comparable instrument had an effective time resolution of roughly 5 to 10 min (von Bobrutzki et al., 2010). In the present study, the recorded NH₃ concentration were averaged over 10 min. The instrument was calibrated before each fertilisation against the same permeation source as used for the HT-CIMS gas-phase calibrations (Sintermann et al., 2011). The relative accuracy was 10%. In a side-by-side field intercomparison with the HT-CIMS under background conditions, absolute deviations were generally below $2 \mu g m^{-3}$.

2.1.4 Sampling by impingers and laboratory analysis

For comparison, NH₃ concentrations over the fields were recorded with wet chemical impingers and subsequent laboratory analysis. Ambient air was directed at a controlled flow rate of $1\,\mathrm{STP}\,1\,\mathrm{min}^{-1}$ through acidic solution (0.01 M H₂SO₄) that ideally strips all gaseous NH₃ and particulate NH₄⁺ into dissolved NH₄⁺. The solution was spiked with CH₂O to suppress microbial activity during sampling and storage. The air was aspirated by flow-controlled pumps

(Gilair-5, Sensidyne, Florida, USA). In addition, sample air flow was calibrated prior to and after each experiment. The impingers collected over periods of one to several hours. They had been tested for breakthrough of NH₃ in the laboratory and during the field campaign by applying two impingers in series. The efficiency was > 99 %. A short (15 cm) 1/4" PFA inlet tube was added in front. Collected samples were immediately cooled to 4 °C and analysed off-line within 3 days by means of ion-chromatography. The device was calibrated using liquid NH₄+ standards. Sampling periods, heights and sampled air volumes are shown in Table 1. Based on in-field side-by-side measurements and the precision of the laboratory calibration of the ion-chromatograph, the uncertainty of the impinger derived NH₃ concentrations was estimated to be 3 %.

2.2 Field experiment

The experiments were conducted at an agricultural site (longitude 7°44′ E, latitude 47°17′ N, elevation 450 m a.s.l.) close to the town of Oensingen, located in the central lowland of Switzerland. Two adjacent fields cultivated as cropland (wheat) and intensively managed grassland (grass-clover mixture) were fertilised with liquid cattle slurry. The grassland is one of the level 3 sites of the NitroEurope project (www.nitroeurope.eu) and NH3 concentrations as well as exchange patterns had been investigated here previously (Norman et al., 2009; Spirig et al., 2010; Wolff et al., 2010; Flechard et al., 2010, 2011). The climate is temperate continental with mean annual temperature and rainfall of 9.5 °C and 1200 mm, respectively. The soil is classified as Eutri-Stagnic Cambisol (FAO et al., 1998) developed on clayey alluvial deposits. Slurry was spread on the arable field (4 August) and on the grassland (6 August) using a tank trailer with splash plate. This is currently the most common slurry spreading technique in Swiss agriculture. Table 2 lists key parameters of the applied slurry. It consisted of a mixture of cattle (80%) and aged pig (20%) slurry on 4 August and

	A _{fert} [ha]	$V_{\rm slurry}$	pН	DM	N_{tot}	TAN		
	[ha]	$[m^3]$		[%]	$[gl^{-1}]$	$[g l^{-1}]$		
4 August 2009	1.23	41.0	7.82 ± 0.10	0.99 ± 0.09	1.07 ± 0.04	0.87 ± 0.01		
6 August 2009	0.77	22.5	7.49 ± 0.19	2.03 ± 0.35	1.57 ± 0.13	1.18 ± 0.05		

Table 2. Characteristics of the applied slurry: A_{fert} = fertilised area, V_{slurry} = volume of applied slurry, DM = dry matter content, N_{tot} = total nitrogen content, TAN = slurry total ammoniacal nitrogen = [NH₄⁺] + [NH₃]; \pm standard deviation of the analytical sample replicates.

of cattle slurry on 6 August. Corresponding to the typical practice on Swiss farms the slurry had been diluted with rain water during storage, resulting in a low dry matter (DM) content. The arable field had been harvested a few days earlier with stubbles of $\sim 20\,\mathrm{cm}$ height remaining in very low density, and the grassland field had been previously cut with a resulting average vegetation height of 5 cm.

In order to distribute the slurry over the entire field, the farmer needed to spread several tank volumes and each refilling required 20 to 25 min. Thereby, a sequence of fertilisation tracks composed of spatial and temporal displaced emission areas was produced (Fig. 1, chronologically numbered). The complete period of fertilisation took almost two hours on 4 August and about one hour on 6 August while individual slurry tracks were dispensed within 3 min.

Figure 2 provides an overview over the predominant meteorological conditions for the days of fertilisation. Warmest air temperatures were 24 °C and 27 °C, respectively, during daytime and no rainfall occurred. High solar radiation intensity was accompanied by relative humidity down to below 50 % during the day with highest values in the early morning. Winds prevailed from north-easterly directions speeding up to 4 m s⁻¹ and up to 2.5 m s⁻¹, respectively.

The EC flux system (ultrasonic anemometer and sample gas intake connecting to the HT-CIMS) was placed on the field immediately after the distribution of the first slurry track had been completed. On 6 August it became necessary to subsequently move the system to the adjacent second track to provide a better fetch for the flux measurement. Hence, as soon as the second track was cast the EC system was positioned at its final location (Fig. 1). The FTIR systems measured NH₃ concentrations spatially integrated over the length of the infrared paths. The data were averaged to obtain 10 min intervals. The optical paths were situated parallel to the south-western border of the respective field, stacked at the three heights. The paths were thus arranged orthogonal to the site's main wind axis, downwind of the expected wind direction. In addition, the cavity ring-down NH3 analyser was placed on the upwind edge of the fertilised field inside a temperature controlled container to monitor the NH3 background concentration, i.e. unaffected by local emissions. NH₃ was sampled by the wet chemical impingers with subsequent laboratory analysis in order to provide a robust concentration measurement over the fertilised fields (Sect. 3.4). Immediately following both fertilisations – after the distribution of the slurry on the entire field – two towers equipped with impingers at two heights were positioned at each field. They captured NH₃ concentrations integrated over one to several hours (Table 1).

2.3 Determination of NH₃ fluxes and quantification of field emissions

2.3.1 Eddy covariance method with fast HT-CIMS detection (EC/HT-CIMS)

The procedure to calculate the EC fluxes (Dabberdt et al., 1993) of NH₃ measured by HT-CIMS is described by Sintermann et al. (2011). The EC fluxes were corrected for the amount of high-frequency attenuation in the used closed path system by an empirical ogive approach (Ammann et al., 2006). Fluxes were calculated in 10 min intervals and were rejected if stationarity was violated according to Foken and Wichura (1996) (using 2.5 min sub-intervals). Data derived from all applied measurement systems are explicitly shown for the period when integral turbulence characteristics (Foken and Wichura, 1996) indicated conditions with turbulent exchange.

The application of slurry creates a non-stationary setting where areas of high slurry emissions are surrounded by plots with very small background NH3 fluxes close to zero. This evokes vertical flux divergence influencing the flux measured at a specific height (Fowler and Duyzer, 1989; Loubet et al., 2009). In order to deduce surface emissions from the EC measurements the flux divergence has to be considered. The flux footprint describes the upwind area determining an EC flux. It reflects the spatial density distribution of the flux at the measurement location (Schmid, 2002). In our experiment, several of the individually emitting slurry tracks intersected with the footprint area as well as fractions of unfertilised regions beyond the field boundaries, and footprint analysis (Neftel et al., 2008) was used to correct for the divergence. For the situation of high slurry NH₃ emissions it was reasonable to assume negligible surface fluxes outside the fertilised field. The applied footprint model is based on the analytical algorithm by Kormann and Meixner (2001). It has been tested in a tracer experiment (Tuzson et al., 2010)

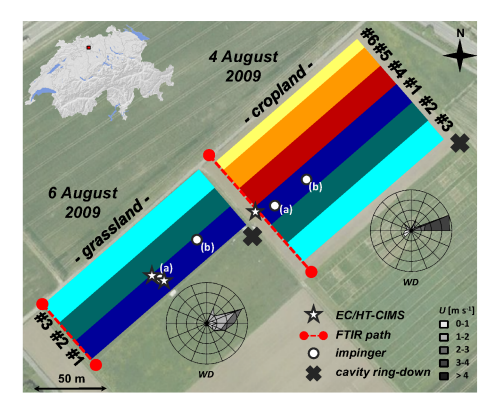


Fig. 1. Application of liquid cattle slurry by sequential spreading of six tanks (tracks #1...#6 on the cropland on 4 August 2009) and three tanks (tracks #1...#3 on the grassland on 6 August 2009), respectively, including wind direction (WD) and wind speed (*U*) from the start of slurry application until the end of the respective day; locations of measurement systems are displayed, (**a**) and (**b**) denote different impinger positions, EC/HT-CIMS position on 6 August on #1 only temporary until #2 was dispensed; satellite image: ©Google Inc., 2010.

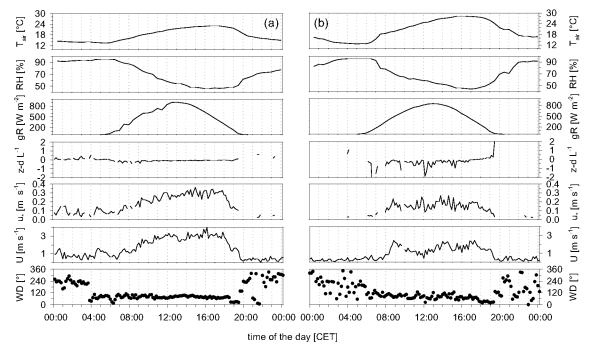


Fig. 2. Air temperature (T_{air}) , relative humidity (RH), global radiation (gR), stability $(z - dL^{-1}, z - d = 1 \text{ m})$, where z = measurement height, d = displacement height, L = Obukhov length), friction velocity (u_*) , wind speed (U) and wind direction (WD) (T_{air}, RH, gR, WD) and U measured at 0 = 1 m a.g.l.; U = 1 m a.g.l. on (a) 4 August 2009 and (b) 6 August 2009.

and has performed well in a state-of-the-art footprint intercomparison (Kljun et al., 2003). The footprint correction for flux divergence is valid, given (i) negligibly small flux interference of chemical conversion of NH_3 to particulate NH_4^+ between the surface and measurement level with the high emissions following fertilisation (Nemitz et al., 2009), and (ii) the small impact of storage of NH_3 in the corresponding air column on the measured flux (<1 %, determined following Spirig et al., 2010).

Specifically, with the footprint analysis the relative contributions of each slurry track and outer regions to the measured EC fluxes were quantified. By linear combination of the various track footprint fractions with a representative individual track emission over time we calculated the whole field's emissions. As proposed by Denmead et al. (1977) a characteristic time course of NH₃ volatilisation, represented by a function $F_{\text{volat}}(t)$, was assumed to be equal for each track. The equality assumption was reasonable because the site conditions were about uniform for the field as a whole (regarding soil, vegetation cover, surface roughness and also meteorological properties) and the applied slurry was taken from the same source and was homogenised before bringing it to the field. The NH₃ flux at the position of the EC system $(F_{\rm EC})$ could then be written as the sum (i = 1 to the number of tracks $n_{\rm T}$) of the emission from each track adjusted for the individual application time (t_{0i}) and weighted with the corresponding footprint fractions (Φ_i):

$$F_{\text{EC}}(t) = \sum_{i=1}^{n_{\text{T}}} \Phi_i(t) \cdot F_{\text{volat}}(t - t_{0i}). \tag{1}$$

Denmead et al. (1977) chose an exponential decrease as shape for $F_{\text{volat}}(t)$ due to emissions from tracks fertilised with injected anhydrous NH₃. At the Oensingen grassland site, Spirig et al. (2010) estimated the NH₃ emissions associated with slurry application to decrease about exponentially to bi-exponentially in the period during and following the spreading, albeit with some uncertainty in the first up to three hours. In the present study, supported by visual inspection of the measured fluxes, we used a bi-exponential decay function (for a discussion of possible related processes see Sect. 4.3) with a total of four fittable parameters as course of $F_{\text{volat}}(t)$:

$$F_{\text{volat}}(t - t_0) = F_1 \cdot \exp\left(-\frac{t - t_0}{\tau_1}\right) + F_2 \cdot \exp\left(-\frac{t - t_0}{\tau_2}\right), \quad (2)$$

where the NH₃ emission F_{volat} at a time t after the moment of slurry application t_0 is a combination of two exponential functions with coefficients F_1 and F_2 , and decay times τ_1 and τ_2 . $F_1 + F_2$ yields the theoretical initial flux F_{ini} directly at the very beginning of the emissions.

Equation (2) was best-fitted to the respective measured EC fluxes by adjusting the four constants F_1 , τ_1 , F_2 , τ_2 in combination with the relative footprint fractions. The function's coefficients were determined by iterative minimisation (Newton type algorithm) of the sum of square deviations of the fit-

ted to measured flux values. The choice of the specific function for the course of F_{volat} will be justified by good correspondence of fit and measurements. For this procedure, values were taken into account over the period of the day when meteorological drivers for slurry NH₃ volatilisation, like U and u_* remained about constant. In the following, we use the term $F_{\text{volat,EC}}$ to refer to the parameterised individual track emissions derived from the EC/HT-CIMS measurements and the fitting procedure. The fluxes calculated for the position of the EC system, derived by the combination of Eq. (1) and $F_{\text{volat,EC}}$ are denomiated $F_{\text{EC,fit}}$. To estimate the field's average emission strength over time $F_{\text{field}}(t)$ the track emissions, weighted with the corresponding track areas (track area $A_{\text{T,i}}$, hence field area $A_{\text{field}} = \sum_{i=1}^{n_{\text{T}}} A_{\text{T,i}}$, A_{in} m²), were combined considering the individual timing of slurry spreading:

$$F_{\text{field}}(t) = \frac{1}{A_{\text{field}}} \sum_{i=1}^{n_{\text{T}}} F_{\text{volat}}(t - t_{0i}) \cdot A_{\text{T},i}. \tag{3}$$

In the situation when only the first track had been applied and thus a single source area could be identified by the footprint evaluation, the measured EC flux combined with the respective footprint was used to compute the average field emissions at that time.

2.3.2 Dispersion method using FTIR concentration measurements (bLS/FTIR)

The FTIRs at the downwind field border measured mean path concentrations representing horizontally exported NH3 originating from the emissions. Since the infrared paths were on average not perpendicular to the prevailing wind direction, only a fraction of the field's NH₃ export was comprised by the FTIR measurements. Correcting for the missing fraction has to account for the spatial inhomogeneity of the emissions due to the sequential slurry spreading. In order to relate the FTIR concentration measurements to surface emission fluxes a tool to determine the spatial dispersion of a tracer is necessary that can calculate concentration footprints (in analogy to the flux footprint). The bLS method (Flesch et al., 1995, 2004) is a powerful mean to determine emission rates employing a single (or multiple) downwind concentration measurement and vice versa. It is based on Lagrangian Stochastic and uses Monin-Obukhov similarity theory (MOST). The model calculates an ensemble of particle trajectories where the particles are being released at a given location and traced backward to determine the resulting particle-ground intersections at one or several areas. The applied bLS model is implemented in a freely available software called Wind-Trax (version 2.0.8.3, Thunder Beach Scientific, Halifax, Canada; www.thunderbeachscientific.com) that exhibits a graphical user interface (see review by Denmead, 2008). In the present study, a fixed number of 50 000 particles was released to calculate particle-ground intersections. The software demands information about the atmospheric state and accepts input variables of gradual complexity levels, with preference to the most detailed inputs. The quantities are either specified directly using measured data and/or are estimated by WindTrax with the help of MOST. The software does not allow simultaneous input of U and u_* , although it can be important to have U and u_* corresponding to the measured values in order to describe turbulence most accurately (Neftel et al., 2008). WindTrax rather computes U from knowledge of u_* , L and the roughness length of the underlying surface (z_0 : height where the vertical profile of U approaches zero) according to MOST. Therefore, we directly provided WD, u_* , L, the standard deviations of the three wind vectors u, v, w, and z_0 . Before, z_0 had been determined numerically with MOST using measured u_* , L and U(z-d) that define the vertical profile of U. Thus, it was assured that within WindTrax, u_* and U satisfied the actual measurements. The magnitude of z_0 could then be consulted to identify situations when atmosphere might not have sufficiently obeyed MOST (Laubach, 2010), which could lead to modelling errors (Flesch et al., 2004). All input variables were derived from the ultrasonic anemometer measurements of the EC system, and 10 min averaging intervals were used for the computations.

In *WindTrax*, the normalised concentration field B(x, y, z) is determined (Laubach, 2010) from the downwind (c) and background (c_{bgd}) concentrations and the emission rate (F_{volat}):

$$B(x, y, z, t) = \frac{c(x, y, z, t) - c_{\text{bgd}}}{F_{\text{volat}}}.$$
 (4)

Calculating field emissions with bLS/FTIR consisted of two steps. First, *WindTrax* was applied to determine the relative contributions of individual slurry track emissions to the measured FTIR line concentrations. For this purpose, Eq. (4) was evaluated for the setup of the FTIR systems and the individual slurry track sources: the tracks were thus given unity emissions ($F_{\text{volat}} = 1 \, \mu \text{g m}^{-2} \, \text{s}^{-1}$) and in separate runs for each track the according $B_{\text{FTIR},i}(t)$ was computed. Next, the actual emissions were determined in a way similar to the procedure used above in case of the EC fluxes: the measured line concentrations (c_{FTIR}) were described as the sum of c_{bgd} and presumed individual track emissions ($F_{\text{volat}}(t)$) adjusted by their footprint contributions ($B_{\text{FTIR},i}(t)$),

$$c_{\text{FTIR}}(t) = c_{\text{bgd}}(t) + \sum_{i=1}^{n_{\text{T}}} \left(B_{\text{FTIR},i} \cdot F_{\text{volat}}(t - t_{0i}) \right). \tag{5}$$

As with the EC fluxes, the time course of $F_{\rm volat}$ (see Eq. 2) was assumed to be equal for each track, allowing the iterative determination of the parameters of $F_{\rm volat}$ from the measured line concentrations and concentration footprints. In such, each fit for $F_{\rm volat}$ obtained from a distinct height of FTIR measurement yielded one emission estimate, which should theoretically correspond to the emissions calculated from the other heights, provided equal performance of the single FTIR systems and realistic representation of dispersion in the bLS model. This parameterisation of individual

track emissions derived from bLS/FTIR is henceforth termed $F_{\text{volat,FTIR}}$, while $c_{\text{FTIR,fit}}$ is used to refer to the concentrations calculated for the FTIR paths based on the bLS concentration footprints (see Eq. 2) and $F_{\text{volat,FTIR}}$.

2.4 Estimating initial volatilisation from liquid slurry characteristics

By knowledge of the chemical slurry constituents and the physical parameters driving the NH₃ volatilisation from solution one can calculate the theoretical flux arising from the initial NH₃ volatilisation at the moment when the slurry is freshly exposed on the surface. Contrasting this slurry derived initial flux ($F_{\rm ini}$) to the corresponding initial flux determined from the respective $F_{\rm volat}(t_0)$ (see Sect. 3.1) one may judge whether this initial flux was of a reasonable physical-chemical magnitude. Assuming immediate liquid-gas phase equilibrium and ideal solution, the initial NH₃ concentration above the hypothetical slurry surface $c_{\rm ini}(z_0')$ (for the concept of z_0' see e.g. Sutton et al., 1993) was inferred with the help of Henry's law (requiring slurry pH, [NH₄+] and surface temperature $T(z_0')$) (Spirig et al., 2010):

$$c(z_0') = \frac{\left[NH_4^+\right] \cdot 10^{4.1218 - 4507/T(z_0')}}{\left[H^+\right] \cdot 10^{-9}},\tag{6}$$

 $c\left(z_0'\right)$ in ppb and $T\left(z_0'\right)$ in K. $c_{\text{ini}}\left(z_0'\right)$ can be translated into the initial surface flux F_{ini} . A flux F_c relates to $c\left(z_0'\right)$ via the corresponding air concentration at a second height $c\left(z-d\right)$ and the aerodynamic and the viscous sublayer resistance R_a and R_b as defined in Flechard et al. (2010):

$$F_c = \frac{c(z_0') - c(z - d)}{R_2(z - d) + R_b}. (7)$$

The NH₃ concentration measured with the cavity ring-down system upwind of the fertilised fields was approximated as background concentration c(z-d) at 20 m a.g.l. $F_{\rm ini}$ was calculated from the initial slurry properties and atmospheric transport capacity using the gradient in NH₃ concentrations. The surface temperature $T(z_0')$ was derived equivalent to Eq. (7), using the air temperature T(z-d) and sensible heat flux, both measured by the ultrasonic anemometer at 1.25 m a.g.l.

3 Results

3.1 Concentrations, fluxes, and emission rates by the EC/HT-CIMS method

Over both fields, NH₃ concentrations observed by the HT-CIMS showed a typical pattern of a fast increase during the spreading of the slurry (Fig. 3). The highest recorded 10 min averaged concentrations were 817 μ g m⁻³ on 4 August and 1543 μ g m⁻³ on 6 August. Unfortunately, failures of the

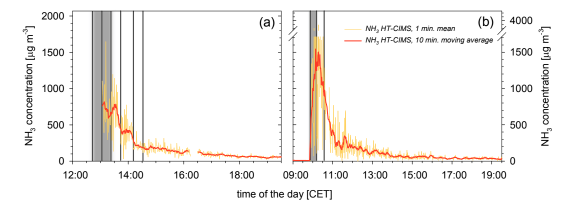


Fig. 3. NH₃ concentrations measured by the HT-CIMS on (a) 4 August 2009 and (b) 6 August 2009; vertical bars represent the spreading of the individual slurry tracks (#1... #6 on 4 August 2009 and #1... #3 on 6 August 2009) while grey shaded areas indicate the period of slurry spreading that mainly affected the fetch of the measurement (Sintermann et al., 2011).

combined sonic and HT-CIMS data acquisition system resulted in short data gaps on 4 August, therefore during the very beginning of this fertilisation no measurement data were available. The concentration maximum was followed by a fast decrease down to about $60 \, \mu \mathrm{g \, m^{-3}}$ and $30 \, \mu \mathrm{g \, m^{-3}}$, respectively, in the evening.

On the days of both fertilisation events, the course of observed EC fluxes featured a similar pattern as the concentrations: an initial fast decrease followed by a slower decline dominating from roughly one hour after the slurry distribution for the rest of the day (Fig. 4). The measured NH₃ fluxes were a composite of the emissions of the sequentially spread tracks. As shown in Fig. 4, the footprint analysis revealed that only tracks #1 to #3 on 4 August and tracks #1 and #2 on 6 August (as well as smaller fractions from outside the field) contributed with an approximately steady proportion to the EC flux. In the evenings, shifts in wind directions promoted increasing influence of the remaining tracks. It becomes evident that track #1 contributed most to the respective measured EC flux on both days. The course of observed EC fluxes indicated an exponential to bi-exponential emission decrease as assumed for F_{volat} (Eq. 2). Consequently, the fit function according to Eq. (1) could well reproduce the measured fluxes during almost the whole day (Fig. 5). The temporal fluctuations of the fitted fluxes represent changes in track footprint contributions influencing the flux at the EC location (see Fig. 4). Associated with a less constant WD, the fluctuations were stronger on 6 August. Median deviations between fitted and measured fluxes were about 11 % on 4 August and 13 % on 6 August considering values until 18:00 and 17:30, respectively. Afterwards, when the differences became larger, the field was assumed to emit homogeneously and average field emissions were calculated from the measured fluxes and a footprint analysis considering the field boundaries as a whole. Table 3 summarises the bi-exponential function parameters of $F_{\text{volat,EC}}$ as derived from the fitting procedure. The initial fast decays showed decay times τ_1 of about 30 and 20 min while the slower decreases had time constants τ_2 of roughly 2 and 3 h. With 332 µg m⁻² s⁻¹ the fitted initial flux $F_{\rm ini}$ immediately at the start of the volatilisation was almost twice as high on 6 August than with 170 µg m⁻² s⁻¹ on 4 August.

3.2 Concentrations and emission rates by the bLS/FTIR method

The vertical concentration profiles measured by the FTIR systems (Fig. 6) showed maximum NH $_3$ concentrations (at 0.8 m a.g.l.) of 773 µg m $^{-3}$ on 4 August and 1446 µg m $^{-3}$ on 6 August, respectively. The course of concentrations approximately reflected the one observed with the HT-CIMS measurements. On 4 August, the values at the highest measurement level (3 m a.g.l.) mostly fell below the limit of detection at 18:00 in the evening. This happened to the NH $_3$ measurements at all heights on the evening of 6 August.

The FTIR measurements were combined with respective bLS concentration footprints to parameterise NH₃ volatilisation $F_{\text{volat,FTIR}}$ from the various tracks using Eq. (5). The time course of measured NH₃ concentrations was well reproduced by $c_{\text{FTIR,fit}}$ before 18:00. $c_{\text{FTIR,fit}}$ began to deviate from measured values only in the evening, when the meteorological regime started to change. Then, the whole field was regarded to emit homogeneously and *WindTrax* was applied to calculate the respective field emissions. Apart from that evening period, median deviations between fitted and measured concentrations for the three heights (low to high) accounted for 6 %, 6 % and 22 % on 4 August and for 19 %, 21 % and 28 % on 6 August.

Table 3 contains the characteristics of the derived biexponential functions describing the time course of a single track emission. The time constants τ_1 of the first exponential function ranged from about 50 to 70 min on 4 August and were around 30 min on 6 August for the results inferred from the three measurement heights. The second

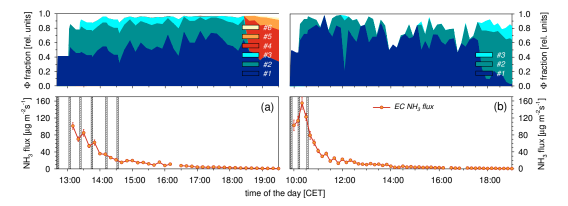


Fig. 4. Influence of the slurry tracks on the EC flux on (a) 4 August 2009 and (b) 6 August 2009 (6 and 3 tracks, respectively); upper panels: relative footprint (Φ) contributions of the individual slurry tracks to the EC flux measurement, white area displays the fraction of the flux footprint covering areas outside the fertilised tracks; lower panels: measured EC NH_3 flux, striped bars represent the actual time of the distribution of each track.

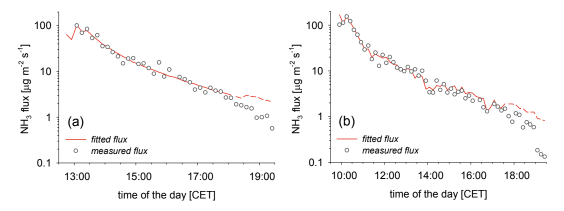


Fig. 5. EC NH₃ fluxes measured and corresponding $F_{EC,fit}$ on (a) 4 August 2009 and (b) 6 August 2009; dashed lines indicate the periods when the parameterisation was regarded not to reflect representative surface emissions anymore, but the field was considered to emit homogeneously.

exponential function was of minor influence for the time course of the emissions on 4 August and had time constants τ_2 larger than 4 h on 6 August. The initial flux $F_{\rm ini}$ was between 144 and 202 µg m⁻² s⁻¹ on 4 August and between 206 to 252 µg m⁻² s⁻¹ on 6 August.

3.3 Spatially averaged and cumulative field emissions

The average field emissions (Eq. 3) increased step-wise during the actual period of fertilisation, which was of course associated with the distribution of the individual slurry tracks (Fig. 7). That was followed by an overall decline of emissions for the rest of the day. In the evening, when turbulence broke down, the NH₃ field emissions became small (i.e. the concentration measurements by FTIR systems reached their detection limit and EC fluxes became very small and instationary). On both events, around 18:00 the field was considered to emit homogeneously over the whole field extent. In this transition regime the emissions decreased faster

than would have been described by the bi-exponential time course. Looking at Figs. 2 and 7 it becomes clear that the change was caused by a shift in meteorological drivers. At around 18:00 essentially U and u_* sharply began to decrease, followed by a delayed increase in atmospheric stability. Almost at the same time, $T_{\rm air}$ started to fall and RH began to rise.

At the cropland fertilisation (4 August), the track emissions $F_{\rm volat,FTIR}$ were higher than $F_{\rm volat,EC}$, and $F_{\rm volat,FTIR}$ had a course closer to a single exponential function (Table 3). The resulting bLS/FTIR field emissions remained higher for the whole day. The parameters of the individually fitted functions $F_{\rm volat,EC}$ and $F_{\rm volat,FTIR}$ showed some considerable differences. The deviations were especially associated with the second exponential sub-function (Table 3), for which there is no reasonable explanation. However, these differences had only a smaller effect on the overall temporal course of emissions (see Fig. 7). Peak field emissions ranged from 88 to 72 μ g m⁻² s⁻¹ for the bLS/FTIR fluxes and were

Table 3. Parameters of the best-fitted bi-exponential functions, calculated surface temperature $(T(z'_0))$, and the according initial NH₃ fluxes (F_{ini}) as derived from Eq. (2) and as expected from initial slurry equilibrium conditions.

		fit					
		F_1 [µg m ⁻² s ⁻¹]	τ ₁ [min]	F_2 [µg m ⁻² s ⁻¹]	τ ₂ [min]	$T(z_0')$ [K]	$F_{\rm ini}$ [µg m ⁻² s ⁻¹]
4 August 2009	EC/HT-CIMS	134	32	36	129		170
	bLS/FTIR 0.8 m	192	49	10	2724674		202
	bLS/FTIR 1.8 m	150	60	6	74 951		156
	bLS/FTIR 3.0 m	142	73	2	74 951		145
	slurry equil.					302.9	564-894
6 August 2009	EC/HT-CIMS	295	23	37	161		332
S	bLS/FTIR 0.8 m	207	29	23	258		230
	bLS/FTIR 1.9 m	237	28	15	367		252
	bLS/FTIR 3.0 m	201	30	5	788		206
	slurry equil.					299.6	186–446

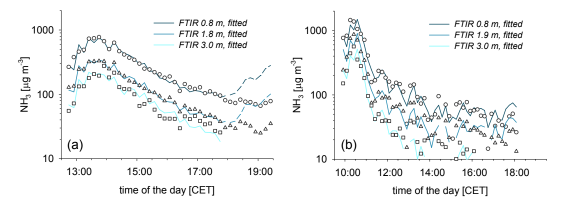


Fig. 6. NH₃ concentrations measured by FTIR systems and corresponding $c_{\text{FTIR,fit}}$ on (a) 4 August 2009 and (b) 6 August 2009; open symbols show the measured values at the various measurement heights: circles = 0.8 m, triangles = 1.8 and 1.9 m, squares = 3.0 m a.g.l.; dashed lines indicate periods when the parameterisation was regarded not to reflect representative surface emissions anymore, but the field was considered to emit homogeneously.

 $66 \, \mu g \, m^{-2} \, s^{-1}$ for the EC/HT-CIMS emissions. Over grassland (6 August), the bLS/FTIR track emissions and hence the corresponding average field emissions were characterised by similar bi-exponential functions as those derived from the EC/HT-CIMS fluxes. The temporal behaviour of $F_{\text{volat,EC}}$ obeyed to the bi-exponentiality on both events in a similar way (except that F_{ini} was much higher over the grassland). In contrast to 4 August, the EC/HT-CIMS field emissions were larger than the bLS/FTIR field emissions. Peak emissions occurred immediately after fertilisation. They were in maximum $174 \,\mu\text{g m}^{-2} \,\text{s}^{-1}$ and for the bLS/FTIR derived emissions, highest values amounted to 114 to $139 \,\mu \mathrm{g m^{-2} s^{-1}}$. Although theoretically equivalent, the fluxes calculated from the FTIR concentrations at the three heights differed from each other. On 4 August, the emissions inferred from the concentrations of the lowest height exceeded those calculated

from the measurements at both remaining heights, whereas on 6 August the fluxes derived from the uppermost height were smaller than those derived from the other two.

In terms of cumulated emissions (Fig. 8) the fertilised fields lost 5.35 kg N on 4 August and 4.88 kg N on 6 August with regards to the EC/HT-CIMS derived results. On 4 August, the bLS/FTIR based losses were higher (6.69 to 7.90 kg N) and lower (3.21 to 4.33 kg N) on 6 August. Over the arable site, the bLS/FTIR cumulated emissions were about 20 % (middle and highest measurement height) to 32 % (lowest height) larger than the cumulated EC/HT-CIMS emissions. Over the grassland, they deviated from EC/HT-CIMS by -13 % (middle and lowest height) and -48 % (highest height). According to both measurement systems over 80 % of the total emissions occurred in the period during the slurry spreading and the two subsequent hours.

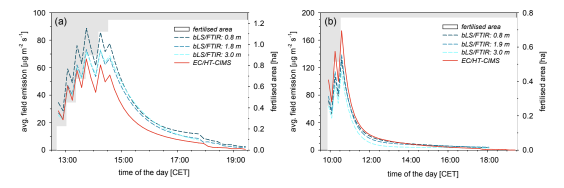


Fig. 7. Average field emissions on (a) 4 August 2009 and (b) 6 August 2009 derived from EC/HT-CIMS and bLS/FTIR.

Table 4 summarises the various estimates of overall NH₃ field emission losses. Cumulated emissions over several days were calculated based on ongoing EC/HT-CIMS measurements over the second days after slurry spreading (whole field regarded to emit homogeneously). The field emissions were expected to decrease exponentially during 10 days at maximum (Flechard et al., 2010). The period subsequent to the days of fertilisation then contributed only to a very small degree to the overall NH₃ losses, namely to about 7 % and 4% of the first day's losses, respectively. Taking these fractions into account, the EC/HT-CIMS derived emissions represented 16% loss of the applied TAN for the cropland and 19% for the grassland site. The FTIR measurements were close to or below the detection limit on the days following the fertilisations and the systems' application period was restrained by their timely relocation between the measurement sites. The corresponding emission estimates therefore account exclusively for the first day of emissions. They comprise 18 % to 22 % loss of applied TAN on 4 August and 12 % to 16% on 6 August, and when averaged over both measurement methods they amounted to $17 \pm 3\%$ and $16 \pm 3\%$, respectively.

3.4 Verification of concentrations and derived emissions

3.4.1 Comparison with impinger concentration measurements

The bLS model was used to calculate the expected NH $_3$ concentration at the sampling locations of the impingers, using the fitted volatilisation functions $F_{\text{volat,EC}}$ and $F_{\text{volat,FTIR}}$ as source terms. Figure 9 illustrates the comparison of the observed impinger concentrations to the calculated concentrations, averaged over the different impinger sampling periods. Based on theoretical consideration and on visual inspection, we analysed the (dis-)agreement between the plotted concentrations by linear regression through zero (see slope and RMSE results in Table 5). With respect to the emissions based on EC/HT-CIMS, calculated and directly measured impinger concentrations showed a very good agree-

ment without systematic deviations (Fig. 9, panel a). The regression slopes were close to 1 with an average uncertainty range of $\pm 8\%$. The root mean squared error (RMSE) of individual deviations was about $30 \,\mu g \, m^{-3}$. In contrast, the NH₃ concentrations calculated from the bLS/FTIR emissions showed generally larger deviations from the measured impinger concentrations (Fig. 9, panel b). There were systematic differences into opposite directions in the two events. In addition, the inconsistency found between the three FTIR measurement heights is clearly visible again. On 4 August, the concentrations obtained from the lowest height deviated most from the reference with the reverse picture on 6 August, when the uppermost height produced emissions least corresponding to the impinger measurements. Using bLS/FTIR emissions, the impinger concentrations were systematically overestimated by 23 % to 43 % on 4 August and underestimated by 10 % to 23 % on 6 August (Table 5).

Concerning EC measurements, the high-frequency attenuation of fast NH₃ fluctuations inside the sampling and analytical system causes systematic under-determination of the fluxes. We corrected the EC fluxes for this amount as described by Sintermann et al. (2011). As this is an empirical approach comparing the ogives of the NH₃ fluxes to those of sensible heat fluxes, an incomplete correction cannot principally be ruled out. To provide an additional check, the EC/HT-CIMS derived emissions were also used to calculate the concentrations at the HT-CIMS location via bLS. These were compared to the NH₃ concentrations measured by HT-CIMS averaged over 10 min and one hour intervals, sufficiently long to exclude damping influences. Figure 10 shows that there was good agreement on both days.

3.4.2 Plausibility of initial volatilisation

A physical-chemical upper limit of the initial NH_3 volatilisation, expressed as $F_{\rm ini}$, was derived from the slurry analysis and meteorological properties as described in Sect. 2.4. There was a considerable range of $F_{\rm ini}$ calculated from the slurry analysis (Table 3), mainly because of limited analytical precision in pH determination (Table 2). While the initial

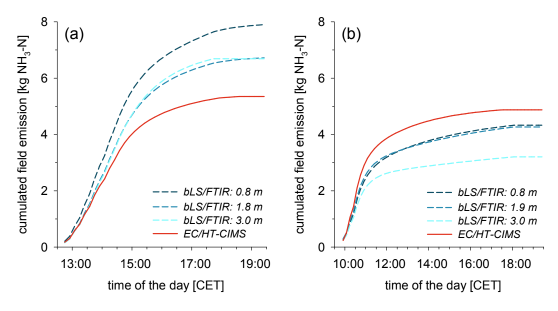


Fig. 8. Cumulative field emissions derived from EC/HT-CIMS and bLS/FTIR on (a) 4 August 2009 and (b) 6 August 2009.

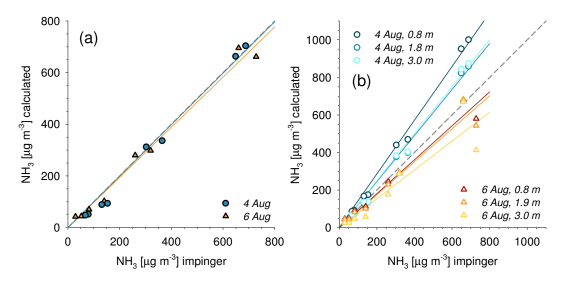


Fig. 9. Comparison of NH₃ concentrations measured by impingers and calculated with bLS, using (a) $F_{\text{volat,EC}}$ and (b) $F_{\text{volat,FTIR}}$; dashed lines show 1:1 relationship and solid lines depict linear regressions through zero (regression parameters see Table 5).

EC/HT-CIMS emissions bLS/FTIR emissions 6 August 2009 4 August 2009 day 4 August 2009 6 August 2009 $[kg NH_3-N][\% \text{ of TAN}]$ [kg NH₃-N][% of TAN] [kg NH₃-N] [% of TAN] [kg NH₃-N] [% of TAN] (II)(II) (II) (II)1 5.35 14.6 4.88 18.0 7.90 6.72 6.69 21.6 18.4 18.3 4.33 4.27 3.21 16.0 15.8 11.8 2 0.83 1.04 0.18 0.68 3 - 100.03 0.08 0.01 0.03 4.33 4.27 3.21 5.76 15.7 5.07 18.7 7.90 6.72 6.69 21.6 18.4 18.3 16.0 15.8 11.8

Table 4. Cumulated field emissions calculated by EC/HT-CIMS and by bLS/FTIR on (I) lower, (II) middle and (III) upper measurement height.

Table 5. Parameters of linear regressions of NH₃ concentrations measured by impingers and corresponding concentrations calculated with bLS, using the emissions F_{volat} derived from the respective methods (Fig. 9); RMSE = root mean squared error (in μ g m⁻³).

	EC/HT-CIMS		bl	bLS/FTIR _{low}		bLS/FTIR _{middle}			bl	bLS/FTIR _{high}		
	slope	R^2	RMSE	slope	R^2	RMSE	slope	R^2	RMSE	slope	R^2	RMSE
4 August 2009	1.00	0.985	31.2	1.43	0.993	28.4	1.23	0.990	31.0	1.25	0.989	32.8
6 August 2009	0.97	0.985	30.4	0.90	0.967	41.4	0.88	0.950	50.8	0.77	0.859	81.3

fluxes derived from the flux determination of all field-applied systems were below the maximum possible values on 4 August, the values on 6 August were in range suggesting that the inferred initial fluxes had been close to the theoretical maximum.

4 Discussion

4.1 Uncertainty of the EC/HT-CIMS approach

The largest difficulty and strongest potential limitation of the EC approach for NH₃ is the correct quantification of the attenuation of fast and turbulent high-frequent concentration fluctuations between the sample location and actual measurement (Brodeur et al., 2009). To our knowledge, the only other study of NH₃ EC flux measurements that simultaneously compared the results to the fluxes obtained from an established (gradient) method discovered substantial high-frequency attenuation losses that could not be quantified by inherent methods (Whitehead et al., 2008). Sintermann et al. (2011) quantified the attenuation empirically and corrected the raw EC fluxes for high-frequency losses. Figure 10 demonstrates that the EC derived emissions $F_{\text{volat,EC}}$ were consistent with NH3 concentrations recorded by the HT-CIMS, averaged over timescales when high-frequency attenuation is considered not to play a role. As well, the comparison to the measured impinger concentrations was good. This underlines that the applied correction was appropriate and systematic underestimation of the fluxes due to highfrequency attenuation could be avoided.

Analytical difficulties in NH₃ measurements may originate from gas-phase calibration uncertainties (von Bobrutzki et al., 2010), drifting instrumental stability (Milford et al., 2009), selective sampling and analysis of gas and aerosol phase and NH₃ sorption in filters, tubes and devices (Parrish and Fehsenfeld, 2000). Flux measurements with the aerodynamic gradient method using AMANDA instruments are associated with uncertainties of 20 % to 76 % (Sutton et al., 2000; Milford et al., 2009). After fertilisation, relaxed eddy accumulation approaches have underestimated NH3 fluxes compared to the AMANDA gradient systems by 20% to 70 % (Hensen et al., 2009) while EC measurements based on laser absorption spectrometry have exhibited biases in the order of -50% when related to the AMANDA fluxes (Whitehead et al., 2008). Emissions from field application of organic fertiliser determined with simultaneously replicated mass balance measurements using passive flux samplers (Leuning et al., 1985) showed unexplained variations between 23 % to 52 % (Misselbrook et al., 2005b). Considering these differences as representative for the range of expected uncertainties in NH₃ flux measurements, the consistency between EC/HT-CIMS derived emissions and the impinger concentrations, found here, is excellent.

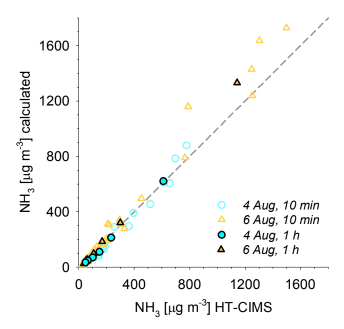


Fig. 10. Measured HT-CIMS NH₃ concentrations (averaged over 10 min and 1 h) vs. corresponding concentrations calculated by bLS using $F_{\text{volat,EC}}$; dashed line shows 1:1 relationship.

4.2 Uncertainty of the bLS/FTIR approach

Inspection of the bLS/FTIR emissions revealed two distinct features: deviations between the results derived from the three measurement heights within one fertilisation event (Figs. 7, 8, and 9) and, when compared to the impinger concentrations and EC/HT-CIMS results, an apparent overestimation on 4 August and an underestimation on 6 August (Fig. 9; Table 5). The relative deviation with respect to the impinger concentrations were around +24% (best two measurement heights) and +43% (least suitable measurement height) for the cropland fertilisation and around -11% (best two measurement heights) to -23% for the grassland fertilisation. These values are within the typical uncertainty range of NH₃ flux determination as reviewed above.

In the past years the applied bLS method has been proven to determine emissions from concentration measurements with accuracies around 10% under most circumstances (Flesch et al., 2004, 2005; McBain and Desjardins, 2005; Gao et al., 2009a, 2010). It is considered to be currently among the most accurate micrometeorological techniques to calculate dispersion and determine emission rates (Denmead, 2008; Laubach, 2010; Loubet et al., 2010). It has been applied to assess methane and/or NH₃ emissions from agricultural fields fertilised with slurry (Sanz et al., 2010) and urea (Sommer et al., 2005), grazed fields (Denmead et al., 2004; Laubach and Kelliher, 2005; Laubach et al., 2008; Laubach, 2010), cattle feedlots (Flesch et al., 2007; McGinn et al., 2007; van Haarlem et al., 2008; Loh et al., 2008), and even complete farms (Flesch et al., 2009). The bLS calcu-

lates emissions accurately provided homogeneously emitting source areas (or well represented point sources), a precise monitoring of c_{bgd} and a largely undisturbed wind field, i.e. an obstacle-free downwind fetch longer than 5-10 (Flesch et al., 2005; Gao et al., 2010) to 25 (McBain and Desjardins, 2005) times z - d, depending of the complexity of the disturbance. The accuracy can be negatively affected by extreme atmospheric instability and is sensitive to low U and nonstationarity indicated by low u* (Flesch et al., 2004; McBain and Desjardins, 2005; Gao et al., 2009b). In the present study, all mentioned quality criteria were fulfilled. The differences in bLS/FTIR emission estimates depending on measurement height (Figs. 7 and 9) may be explained by another specific limitation of the model. Laubach (2010) found that the accuracy of the bLS, implemented in WindTrax, can depend on the ratio of z - d to the mean fetch length with resulting differences of up to 20 % under unfavourable conditions. This is probably due to an overestimation of the speed of vertical dispersion as it relates on the uncertain parameterisation of energy dissipation. Laubach (2010) identified a specific relative crossover height (z - d/mean fetch length)at which the effect vanishes. Below, potential overestimation occurs, reversing into underestimation above. It was not possible to identify a representative crossover height in the present study as this would have required winds blowing perpendicular to the FTIR paths in order to determine the mean fetch length. It is, however, very likely that on 4 August the lowest FTIR height was significantly below the crossover height due to the orientation of the field setup relative to the predominant wind direction. This could qualitatively explain the overestimation of the emissions derived from the lowest height relative to the other heights. The reverse picture occurred on 6 August when probably the highest measurement height exceeded the respective crossover height. This was the case because the emission fetch for the measurement was quite small that day, as the relative influence of source areas outside the fertilised field (due to easterly wind components) were larger for the short FTIR path on that day, producing a smaller effective fetch for the line concentration measuruement (see Fig. 1). Taking these aspects into consideration, it is likely that the respective single outliers of the bLS/FTIR emissions were an effect of the shifting bLS performance due to the experimental setting. In addition, on 6 August the wind direction fluctuated significantly and thus the angle of the FTIR paths often happened to be close to the direction of the wind which can increase the uncertainty in the emission calculation (Flesch et al., 2004). The preceding reflections suggest that within one event, differentiating between the three measurement heights, the two bLS/FTIR emissions showing agreement were more plausible than the deviating value. The different biases observed in the two fertilisation events might be attributed to shifting instrumental performance. For example, in a recent intercomparison, von Bobrutzki et al. (2010) characterised eleven state-of-the-art instruments for NH3 concentration measurements based on eight analytical methods. Agreements better than 25 % can hardly be achieved and deviations can be much larger.

4.3 Dynamics of NH₃ volatilisation

The fluxes presented in our study show that highest emissions from individual tracks occurred immediately after slurry application. It is difficult to assess the emission course over the first minutes after the spreading. The determination of the function parameters relied on 10 min averages and Eq. (2) allowed to extrapolate to the beginning of fertilisation. The spreading of the slurry itself took about 3 min, thus the initial timing was not sharply defined. On 4 August, there was a data gap of about 20 min in the HT-CIMS measurements at the beginning of the fertilisation introducing additional uncertainty into the back-extrapolation.

According to Sommer et al. (2003) there are two distinct stages in the NH₃ volatilisation which could explain a biexponential decrease. In the first period, immediately after fertilisation, the slurry is exposed at the soil/vegetationatmosphere interface as it has not undergone complete soil infiltration and drying. The emissions depend only on the characteristics of the slurry on the surface and of the atmospheric transport. At the second stage, the emissions are governed by slurry-soil interactions. They include evaporation, sorption and microbial activity provoking a time course that exhibits lowered emissions with a longer time constant. Considering the first stage, an upper limit of the initial volatilisation rate can be calculated from the slurry and atmospheric properties. Volatilisation of fatty acids as well as fast mineralisation with carbon dioxide volatilisation can increase the slurry pH (Vandre and Clemens, 1997) promoting high emissions. On the other hand, volatilisation of NH₃ decreases pH (Sommer et al., 2003). In slurry, a temporal increase in pH has been observed with a high total inorganic carbon (TIC) content of the slurry (Sommer and Sherlock, 1996). We have no information about TIC in the applied slurry, but since it was rather thin (low DM content, most of the N in form of TAN: 81 % and 75 %, respectively) and as infiltration happened fast we must assume that a change in slurry pH was not driving NH₃ volatilisation in our experiment. On 6 August the slurry was applied to the cut grassland where the canopy intercepted a larger fraction of the slurry exposing more liquid to direct volatilisation. The emission enhancing effect of slurry interception on a short canopy was shown by Rochette et al. (2008) and Thorman et al. (2008). Therefore, emissions in the initial period can be expected to peak stronger on grassland than on harvested arable land. It has been demonstrated that increasing DM content dampens initial NH₃ emissions but prolongs them in the following (Braschkat et al., 1997). The slurry spread in our experiments generally had a low DM content and thus promoted quick volatilisation as reflected in the fast initial decrease of the emissions. Increased wind speed and air temperatures as well enhance initial emissions (Søgaard et al., 2002), and in the experiments wind speed and especially the high air temperatures (though in a slightly different combination between the two events) favoured a high emission rate during the initial period following fertilisation (Sommer et al., 1991). Considering the high infiltration and sorption potential of the applied slurry with the bare soil on 4 August (see below), it is likely that equilibrium in NH₃ volatilisation described by Henry's law did not occur over an initial period of several minutes during that event. Consequently, the initial flux derived from the bi-exponential fit was lower than the potential flux derived from slurry properties, surface temperature and transfer velocity. On 6 August, the initial NH₃ volatilisation as determined from the slurry properties matched the initial values identified by the field measurements (Sect. 3.4.2). The stronger slurry-canopy interception and thus slower soil infiltration during this experiment favoured slurry-atmosphere equilibrium conditions. We conclude therefore that the initial emissions inferred from the field measurements were not prone to severe underestimation as they were close to a physical-chemical plausible value. Employing the Henry equilibrium like in Eq. (6) to calculate $c(z'_0)$ ignores the fact that slurry is not really an ideal solution. The activity of other dissolved ions can influence the ionic strength of the slurry solution which may decrease the expected volatilisation. When considering the applied rather thin slurry to have an ionic strength at the higher end of the average range reported in the literature, the volatilisation would be reduced by about one quarter (Sommer et al., 2003) and the main findings summarised in Table 3 would not very much change. It becomes only more obvious that F_{ini} derived from the various measurement systems were very close to the maximum on 6 August.

The soil at the Oensingen site has a cation exchange capacity of more than 20 cmol kg⁻¹. Compared to the grassland, the arable field's sorption potential of the soil acts more effectively in buffering emissions from the low DM containing slurry since bare, dry soil without canopy interception promotes fast penetration below the surface (Sommer and Jacobsen, 1999; Sommer and Hutchings, 2001; Misselbrook et al., 2005c; Sommer et al., 2006). In addition, slurry TAN content drives to a large extent the NH₃ emissions (Menzi et al., 1998). The applied slurry on 6 August had a higher TAN and DM content (albeit a lower pH) than on 4 August. Therefore, the emissions could be expected to be higher on 6 August relative to the event on 4 August (Sommer and Olesen, 1991; Braschkat et al., 1997; Søgaard et al., 2002; Sommer et al., 2003) when they declined with a less pronounced bi-exponential course.

It is known for the investigated site that the emissions decrease exponentially over only a few days after slurry spreading. The vast majority of NH₃ loss (at least 80 % of the total emission) has always been observed at the day of fertilisation (Spirig et al., 2010). Such a course with the main part of emissions occurring in the first ten to twenty hours is quite common after slurry spreading and has often been documented (e.g. Pain et al., 1989; Mannheim et al., 1995;

Braschkat et al., 1997; Vandre et al., 1997; Génermont et al., 1998; Menzi et al., 1998; Misselbrook et al., 2002; Huijsmans et al., 2003; Rochette et al., 2008; Sanz et al., 2010). The conditions at the Oensingen grassland revert back to a small, potentially bi-directional flux regime after about 5 days and even the canopy compensation point returns to prefertilisation levels after about 10 days (Flechard et al., 2010).

5 Conclusions

Field scale NH₃ emissions from slurry application were determined over a cropland and a grassland field by two different analytical approaches. For both methods, the respective detailed footprint analysis in combination with the high time resolution of 10 min averages enabled to attribute the measured fluxes and concentrations, respectively, to individual slurry track emissions. In this way, the very important initial period of emissions could be described in detail.

The cumulated EC/HT-CIMS and (the two most plausible) bLS/FTIR emissions agreed within 20 %, a difference typical for NH₃ flux quantification. Both methods were further tested for consistency with independent impinger concentration measurements. These analyses showed good agreement without systmatic deviations for the EC/HT-CIMS results (uncertainty ± 8 %) but some systematic over- or underestimation for the bLS/FTIR results. Therefore we attribute the observed differences between the two flux methods mainly to uncertainties of bLS/FTIR. The overall NH₃ loss (EC/HT-CIMS) during the day of slurry spreading was quantified to amount for 14.6 % of the applied TAN at the cropland and 18.0 % at the grassland with contributions \leq 1 % loss of TAN on the subsequent day.

Over two years, Spirig et al. (2010) determined NH₃ emissions following slurry application at the Oensingen grassland field using wet chemical AiRRmonia instruments (Erisman et al., 2001) in a gradient approach. They had to estimate the initial phase's (first one to four hours) field emissions due to limited resolution of the instruments and the unsteady, inhomogeneous emission conditions associated with slurry spreading, concluding their overall flux uncertainty was dominated by the vagueness of this period. The approaches and resulting emissions in the present study now provide a clearer, less uncertain picture over the whole period of field emissions yielding similar losses as described by Spirig et al. (2010). Over three years, seasonally distributed between April and October, the grassland field's slurry NH₃ emissions have been quantified in total by means of four different approaches. They kept persistently small in a range between about 4 % and 19 % of the applied TAN.

Generally, flux measurements on the field scale under realistic slurry spreading practice are only feasible with techniques that provide a fast flux determination and can measure a large dynamic concentration range with sufficient accuracy. These requirements were particularly met with the two methods applied in the present study, having the advantage that analytics were either based on open path measurements or were applied with excess heating of all inner walls minimising damping effects and including procedures for quantifying the influence of the remaining wall interactions on the flux. For both methods applied here, the analytical instruments are relatively expensive and need a high level of maintenance in the field. Thus, an important future challenge will be to establish either of the two methods with easier-to-use (and cheaper) analytical instruments. For the bLS method, also point concentration measurements instead of open-path line measurements could be used. However, a high temporal resolution (about 15 min or better) is necessary for properly evaluating the temporal dynamics. Another important future challenge will be the modelling of the observed course of fluxes with a mechanistic approach and to validate such a model at differing site conditions.

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