

Surface-atmosphere exchange of ammonia over peatland using QCL-based eddy covariance measurements and inferential modeling

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Abstract. Recent advances in laser spectrometry offer new opportunities to investigate ecosystem-atmosphere exchange of environmentally relevant trace gases. In this study, we demonstrate the applicability of a quantum cascade laser (QCL) absorption spectrometer to continuously measure ammonia concentrations at high time resolution and thus to quantify the net exchange between a semi-natural peatland ecosystem and the atmosphere based on the eddy-covariance approach. Changing diurnal patterns of both ammonia concentration and fluxes were found during different periods of the campaign. We observed a clear tipping point in early spring with decreasing ammonia deposition velocities and increasingly bi-directional fluxes that occurred after the switch from dormant vegetation to CO₂ uptake, but was triggered by a significant weather change. While several biophysical parameters such as temperature, radiation, and surface wetness were identified to partially regulate ammonia exchange at the site, the seasonal concentration pattern was clearly dominated by agricultural practices in the surrounding area. Comparing the results of a compensation point model with our measurement-based flux estimates showed considerable differences in some periods of the campaign due to overestimation of non-stomatal resistances caused by low acid ratios. The total cumulative campaign exchange of ammonia after nine weeks, however, differed only in a 6 % deviation with 911 and 857 g NH₃-N ha⁻¹ deposition being found by measurements and modeling, respectively. Extrapolating our findings to an entire year, ammonia deposition was lower than reported by Hurkuck et al. (2014) for the same site in previous years using denuder systems. This was likely due to a better representation of the emission component in the net signal of eddy-covariance fluxes as well as better adapted site-specific parameters in the model. Our study not only stresses the importance of high-quality measurements for studying and assessing land surface-atmosphere interactions, but also demonstrates the potential of QCL spectrometers for continuous observation of reactive nitrogen species as important additional instruments within long-term monitoring research infrastructures such as ICOS or NEON at sites with strong nearby ammonia sources leading to relatively high mean background concentrations and fluxes.

1 Introduction

Increased agricultural production and energy consumption over the last century led to a dramatic increase in anthropogenic reactive nitrogen (N) production (Erisman et al., 2008). Atmospheric N deposition can be a major driver of change in most natural and semi-natural ecosystems and may considerably alter species composition, biodiversity, and ecosystem functioning with regard to causing nutrient imbalances. As ammonia (NH₃) mainly originates from agricultural activities, it has received more and more attention in the past 20 years (e.g., Sutton et al., 2011). It is estimated that in 2008 65 Mt NH₃-N were emitted globally to the atmosphere (Sutton et al., 2013), a large proportion (60 %) of it from anthropogenic sources. Around 1910 the creation and therefore release of NH₃ experienced a steep rise due to the Haber-Bosch-process (Galloway et al., 2003). While ammonia emissions doubled since 1950 (Asman et al., 1998), Sutton et al. (2013) predict the emission of 132 Mt NH₃-N yr⁻¹ by 2100. Ammonia is an essential part of the nitrogen cascade with reactive nitrogen (N_r) tending to accumulate in both, ecosystems and atmosphere, mainly because denitrification in soils cannot balance out industrial N creation (Galloway et al., 2003) causing a severe environmental problem for future generations. Additionally, reactive N is widely spread through hydrologic and atmospheric transport processes (Galloway et al., 2003). Ammonia can cause foliar injury, increase sensitivity to drought, reduce frost hardiness, lead to structure change in plant communities and loss in biodiversity, especially regarding sensitive nutrient-poor ecosystems like peatbogs (Krupa, 2003). Furthermore, deposited ammonia causes ecosystem acidification, fertilization, and eutrophication (Galloway et al., 2003), while air and water quality deterioration also impact human health (Erisman et al., 2013).

Until now only little is known about the temporal and spatial variability of NH₃ concentrations and exchange fluxes between different ecosystems and the atmosphere. Estimates to what extent NH₃ is being emitted from plant canopies under common environmental conditions remain highly uncertain (e.g., Flechard et al., 2013). This is due to the fact that continuous half-hourly, micrometeorological measurements of NH₃ exchange, e.g. based on the aerodynamic gradient or eddy-covariance (EC) technique, have largely remained experimental and were limited to selected research sites and to measurement campaigns of typically a few weeks to a few months due to technical complexity and to the large equipment and operational costs involved (e.g., Sutton et al., 2007; Flechard et al., 2011; Marx et al., 2012; Brümmer et al., 2013). Other well-established techniques like denuder, impinger or filter sampling in combination with ion chromatography analysis usually provide concentration values and flux rates at poor time resolution and require labor- and cost-intensive chemical analyses (e.g., Dämmgen and Zimmerling, 2002; Tang et al., 2009; Hurkuck et al., 2014). Over the last few years, substantial progress has been made in the use of tunable diode laser absorption spectrometers (TDLAS) and quantum cascade lasers (QCL). The precision and fast response of these approaches have allowed first EC measurements of field scale N₂O and CH₄ fluxes (Denmead et al., 2010; Kroon et al., 2010; Neftel et al., 2010; Merbold et al., 2014) and are expected to become a standard method within integrated observation networks such as ICOS in Europe or NEON in North America. Eddy-covariance measurements of NH₃ fluxes, however, have been extremely limited and are still subject to considerable uncertainty (Famulari et al., 2004; Ellis et al., 2010; Sintermann et al., 2011; Ferrara et al., 2012) mainly due to issues regarding the tube

inlet design, sampling high frequency flux losses and tube wall and air chemistry (e.g., Marx et al., 2012 and references therein).

In this study, we demonstrate the applicability of a quantum cascade laser (QCL) absorption spectrometer to continuously and fast measure turbulent fluctuations at background-level NH_3 concentrations and thus the net exchange between a semi-natural peatland ecosystem and the atmosphere, with the eddy-covariance technique. Specifically, we (1) test the QCL performance to measure NH_3 concentration fluctuations and calculate NH_3 fluxes and net deposition during the observation period, (2) investigate the biophysical controls on NH_3 concentrations and fluxes, and (3) compare the measured fluxes with results from a local application of a two-layer NH_3 canopy compensation point model using the parameterization after Massad et al. (2010). The general objective is to better understand the mechanisms of peatland-atmosphere NH_3 exchange under the influence of highly intensive agricultural land management in the surrounding area.

2 Materials and methods

2.1 Site description and local climate

Tower-based fast response measurements of ammonia concentrations were conducted from 18 February to 8 May 2014 at an ombrotrophic, moderately drained peatland located in Northwest Germany near the city of Meppen ($52^{\circ}39'19.98''\text{N}$, $7^{\circ}10'56.65''\text{E}$, 14 m.a.s.l.). The site is surrounded by intensive agricultural land and livestock holdings. Local vegetation is dominated by bog heather (*Erica tetralix*), purple moor-grass (*M. caerulea*), cotton grass (*Eriophorum vaginatum*, *Eriophorum angustifolium*) and is further characterized by some smaller, mostly solitary trees like birches (*B. pubescens*), and Scots pines (*Pinus sylvestris*). The peat layer depth is approx. 4 m. The area around the tower has a fetch of 650 m including some small paths, tree lines and hedgerows. Facing towards the main wind direction (SSW) 230 m can be regarded as totally homogenous. Details on fetch and footprint analysis are given in Hurkuck et al. (2014).

From 1981 to 2010 the annual average air temperature (T_a) and the mean annual precipitation in the area were 10.0°C and 800 mm, respectively (German Weather Service, station Lingen, 2015). For the respective months of observation (February to May), the long-term average air temperatures were 2.7°C , 5.7°C , 9.3°C , and 13.6°C , respectively. Except for May, monthly mean air temperatures in 2014 were higher (4.1°C , 6.3°C , 10.9°C , and 12.3°C) than the 30-yr averages. Monthly precipitation (36 mm, 31 mm, 39 mm, and 124 mm) was lower in February, March and April than the long-term observations (53 mm, 67 mm, 45 mm and 58 mm). As most of the precipitation was recorded after the campaign on 8 May, the measurement period was considerably drier than the long-term mean.

2.2 Measurements of ammonia and microclimate

Ammonia concentrations were measured at high temporal resolution with a QCL absorption spectrometer (model mini QC-TILDAS-76) from Aerodyne Research, Inc. (ARI, Billerica, MA, USA). They were recorded as mol NH_3 per mol dry air, also termed as mixing ratio, due to an online internal correction for water vapor dilution provided by the QCL. Laser

detectors were thermoelectrically cooled at 25 °C. A 76-m path length and 0.5 L multiple pass absorption cell for sampling at an operation pressure of approx. 40 Torr was used. The QCL is able to offer up to 10 Hz resolution with the detection limit in the sub-ppb range (McManus et al., 2008). The main frequencies of the flux contributing eddies were in a range of 0.01 to 1 Hz. Precision, i.e. measurement sensitivity, of the instrument is 0.042, 0.021, 0.016, and 0.010 ppb in 1, 10, 20, and 60 s, respectively. Along with its compact design, the QCL forms an appropriate basis for eddy-covariance measurements in the field (Ellis et al., 2010, Ferrara et al., 2012). A dry vacuum scroll pump (TriScroll 600, Agilent Technologies, Santa Clara, United States) was used to transport the sample air with a flow rate of approx. 17 l min⁻¹ through 3-m long tubing (1 cm inner diameter) to the laser cell and back outside through an exhaust device. For further conceptual details see McManus et al. (2008) and Zahniser et al. (2005) as well as Ellis et al. (2010) for a performance test.

To prevent damage to the laser cell from particles, it is necessary to filter the ambient air. Because of the stickiness of NH₃ and its high reactivity, it is not possible to use any conventional membrane filter. Aerodyne Research, Inc. developed a specific inertial inlet (Fig. 1) that removes particles with an aerodynamic diameter larger than 300 nm and that reduces pressure inside tubing and laser cell. After a short PTFE inlet, a critical orifice made of glass ensures that the pressure in the system is decreased to a range of approx. 5.3 to 8 kPa, which significantly reduces wall adsorption effects (Warland et al., 2001) and is required for the operation of the laser cell. After passing the critical orifice, the sample air is forced to make a sharp turn, thereby losing 10 % of its volume, going straight to the pump, and approx. 50 % of the particles (> 300 nm) (Ellis et al., 2010; Ferrara et al., 2012; von Bobruzki et al., 2010). Heating the inertial inlet box (at approx. 40 °C) and the anti-adhesive PFA tubing that leads the sample air to the QCL's analyzer cell minimize water condensation or absorption (Massman and Ibrom, 2008, Ibrom et al. 2007), thus avoiding interactions of ammonia with surfaces (Walker et al., 2006; Norman et al., 2009; Ellis et al., 2010). Ellis et al. (2010) compared different tubing variants and reported significant improvement when using heated tubing at 40 °C, which reduced unintended interactions to around 10 % at 30 ppb. A side effect is, that at low atmospheric pressures NH₄NO₃ aerosols can be volatilized, if not removed by the inertial inlet, and thereby produce NH₃. DELTA denuder measurements (details below) showed a NH₄⁺ to NH₃ ratio of 0.17 up to 0.24 during the campaign. Assuming that 100 % of the NH₄⁺ aerosols passed the inlet and were volatilized, there was on average an overestimation of 21% of NH₃ concentration, which is the same rate Walker et al. (2006) found in their investigation and is an acceptable compromise between NH₃ loss due to wall sorption effects and ammonia alterations through NH₄NO₃ volatilization. This affects not only the concentration but also the flux, because NH₄NO₃ particles are also deposited. In this study, we performed calibration with zero air every 8 hours, i.e. at 00:00, 08:00 and 16:00 local time for 60 s per interval (data not shown) as well as internal system calibration through the laser itself by aligning the NH₃ absorption peak of the sampled air to the standard of the HITRAN database (Rothman et al., 2009).

An ultrasonic anemometer (model R3, Gill Instruments, Lymington, UK) was installed at 2.5 m above ground. The inertial inlet box was mounted next to it on the north side, a less common wind direction at the site (5.6 % of the campaign), placing the sample air inlet westward 40 cm below the center of the sonic anemometer array. To protect the inlet from rain, a tee-inlet was attached, which allowed to feed in calibration gas (see above).

Additional measurements of NH_3 concentrations with monthly time resolution were conducted by means of passive samplers (e.g., Dämmgen et al., 2010) and DELTA denuder (**DE**nuder for **L**ong-**T**erm **A**tmospheric sampling, e.g., Tang et al., 2009). A detailed description of the measurement setup of meteorological parameters such as air and soil temperature, radiation components, precipitation, water table depth as well as the operation of carbon dioxide and water vapor eddy-covariance measurements is presented in Hurkuck et al. (2014, 2016).

2.3 Data acquisition, analysis and flux calculation

Sonic anemometer data were recorded by the EddyMeas software, which is an embedded application of the software package EddySoft (Kolle and Rebmann, 2009). The QCL was controlled by Aerodyne's TDL Wintel software with ammonia concentration data being recorded at a frequency of 10 Hz on the QCL computer. Anemometer and concentration time series needed alignment to a reference timestamp, before the software EddyPro (LI-COR Inc.) could be used to compute half-hourly exchange fluxes. Block averaging and 2-D coordinate rotation were applied.

The effective time lag between the data series of the vertical wind speed (w) and the NH_3 concentration was determined by searching the maximum (absolute) covariance as a function of the time shift. A strong and irregular drift in the form of a sawtooth wave was found for the lag time of maximum covariance (Fig. 2b), which is attributed to diverging drifts of the system clocks of the sonic anemometer's data acquisition computer and the QCL clock. Thus, the drifting time lag was divided into subsections (one for each tooth) and a linear trend was fitted to the apparent time lag in each section. Then each half-hourly NH_3 time series was shifted with this estimated time lag using a specifically designed R-script (R Core Team, 2012) resulting in near-zero time lags in the corrected datasets. After a further covariance maximization procedure, a smaller less varying time lag remained. In EddyPro a range of -2 to 4 s was chosen.

The ogive method from Ammann et al. (2006) was applied to empirically investigate the high-frequency damping of NH_3 concentration measurements which usually occurs due to imperfect turbulent flow regime and possible wall sorption effects in the sampling line. Cumulative cospectra of $\overline{w'\text{NH}_3'}$ were scaled to the corresponding $\overline{w'T'}$ cospectra in the medium-low frequency range (see example ogives in Fig. 2a) for quality filtered cases. The resulting relative deviations at the high-frequency end are quantitative measures of the flux damping factor. A median damping factor of 0.67 was found which is similar to the damping factors Ferrara et al. (2012; 2016) reported for their NH_3 QCL system. As no clear dependency of the damping factor on horizontal wind speed (Fig. 2c) nor atmospheric stability could be observed, a constant correction factor of $1/0.67$ was applied to all NH_3 flux values as a simplified approach. The random uncertainty of the correction factor was estimated to 15%, but a potential systematic deviation (part of the damping not appropriately detected by the ogive method) cannot be fully excluded.

The random flux error was computed according to the method after Finkelstein & Sims (2001) and the corresponding limit of detection was determined after Langford et al. (2015) as 1.96 times the flux error (95% confidence limit) resulting in a median value of $7.75 \text{ ng N m}^{-2} \text{ s}^{-1}$. Alternatively, an upper flux detection limit for half-hourly values can be calculated by using only night-time data under stationary conditions ($3 - 7 \text{ m s}^{-1}$ wind speed) and wind from the west-southwest sector,

where local ammonia sources were negligible and concentrations were low (< 15 ppb). The standard deviation was $16.5 \text{ ng N m}^{-2} \text{ s}^{-1}$, so the 2σ -uncertainty range is $33.0 \text{ ng N m}^{-2} \text{ s}^{-1}$.

To guarantee a high level of data quality, fluxes were flagged according to criteria presented in Mauder and Foken (2006). Data from April 11 to 28 were excluded from analysis due to several technical difficulties such as power outages, pump failure or insufficient temperature control inside the analyzer housing. 33.9 %, 44.3 %, and 10.9 % of data were flagged with grade 0, 1, and 2, respectively. Data with quality flags 0 and 1 were used for further analysis, while flag 2 data were discarded. Because the stationarity tests included in the flagging protocol by Mauder and Foken (2006) might not be applicable for NH_3 due to possibly higher variability of the flux over short time scales, flag 1 was generally not discarded. The remaining 10.9 % of data were not used due to insufficient turbulence ($u_* < 0.1 \text{ m s}^{-1}$). The data gaps (except for the lost data in April) were filled with the mean diurnal variation (MDV, window= ± 5 days) method (Falge et al., 2001; Moffat et al., 2007). Campaign data were grouped into four periods (Tab. 1) of 9 to 25 days depending on different concentration patterns and meteorology.

2.4 Modeling ammonia exchange

A state-of-the-art dry deposition inferential model driven by measured NH_3 concentrations and local micrometeorological conditions was applied to assess plausibility of the flux measurements. We used the parameterization of a two-layer canopy compensation point model from Massad et al. (2010), which simplifies to a one-layer big leaf model for unmanaged ecosystems (i.e., no soil layer is considered explicitly below the canopy; cf. Nemitz et al., 2000). Aerodynamic-, quasi-laminar-, and cuticular resistance ($R_a(z-d)$, R_b , and R_w , respectively) were parameterized as described in Massad et al. (2010) for semi-natural/moorland vegetation as well as the stomatal compensation point (N-input dependent). Stomatal resistance (R_s) was modeled using the simple global radiation and temperature dependent formulation of Wesely (1989, see supplement for further details), with a minimum stomatal resistance for H_2O of 200 s m^{-1} , since detailed measurements of vegetation characteristics – which are necessary for more complex approaches (e.g., Emberson et al., 2000) – were not available for the study period. R_s and R_w were calculated with temperature and relative humidity at the mean notional height of trace gas exchange, estimated from measurements at the reference height and measured turbulent sensible (H) and latent (LE) heat fluxes as described in Nemitz et al. (2009).

3 Results and discussion

3.1 Diurnal and seasonal pattern of ammonia concentrations

On a half-hourly basis, ammonia concentrations ranged from 2 to 85 ppb with short term (10 Hz) maxima reaching up to 110 ppb. The highest values were found in the beginning of March (Period II) and in the beginning of April (end of Period III), which coincided well with the peak of fertilization activities on nearby agricultural fields including the spreading of organic manures from livestock farming (Fig. 3; for details on farming practices see Hurkuck et al., 2014). The base concentration

level outside the fertilization periods ranged mostly between 7 and 15 ppb and is well represented by mean values of Periods I and IV.

The highest mean diurnal variability of ammonia concentrations was found in Period II with peak concentrations being observed in the late afternoon (>30 ppb from 5 to 15 March; Fig. 4). Concentrations were lowest at night during the whole observation period. While in Period III the average mean diurnal course exhibited less variability than in Period II, almost stable concentrations on a low level (7 to 12 ppb) were found in Periods I and IV.

The frequency distributions of wind directions and ammonia concentrations for the whole observation period are shown in Fig. 5. The typical main wind direction from the 200 to 280° sector is clearly visible. However, only the lowest concentrations (<15 ppb, average for 10° wind sectors) were observed under southwesterly winds, whereas peak concentrations were found under winds from the east and northeast. This finding is consistent with observations from previous years using denuder systems (*cf.* Hurkuck et al., 2014). The reason for the concentration dependency on wind direction is the land use in close vicinity to the measurement site. While southwest of the tower and outside the protected zone, the area is characterized by active peat cutting, thus no elevated ammonia concentrations can be expected, east and northeast of the tower a number of farm houses with livestock buildings, manure storage areas and adjacent fertilized land are located in a distance of approx. 2 km. The pattern of Fig. 5 is also revealed in the progression of wind direction and concentrations over time (data not shown). At the beginning of the campaign, i.e. Period I and early Period II, the predominant wind direction was south and southwest while concentrations were relatively low (Fig. 3). Later on, i.e. from late Period II onwards, wind direction was much more variable with sporadic episodes from the northeast sector, when peak concentrations between 60 and 110 ppb occurred. Despite the fact that frequent winds from the northeast were recorded in Period IV, it was also the time of the lowest concentration levels ($\bar{c}_{NH_3} = 8.0$ ppb). This is an indication that the main fertilization activities on the adjacent fields had been terminated during that time.

Similarly diurnal variability of ammonia concentrations with peak values during afternoon over a variety of ecosystems has been observed by other authors (e.g., Sutton et al., 2000; Wolff et al., 2010). There are most likely several reasons for the observed pattern in this study. First and foremost, concentration levels are highly influenced by agricultural activities in the surrounding area. Farmers usually fertilize their land during the day, thereby causing ammonia volatilization, which is then transported and detected at the study site. With a stable nighttime planetary boundary layer, ammonia in the lower atmosphere is likely being deposited causing decreasing concentrations. With no further penetration from higher layers containing higher loads, concentrations remain low from midnight to sunrise (Fig. 4). When temperatures rise, turbulent mixing of the planetary boundary layer starts and vertical exchange with higher layers increases, which, consequently, leads to rising ammonia concentrations over the day. On the other hand, continuous sources like mechanically ventilated stables could cause an opposite pattern with the planetary boundary layer acting as a lid and leading to a concentration build up at night. As no information was available of ventilation types of stables, we assume that land applied manure during the day dominates the concentration signal in Fig. 4. The conspicuous afternoon peak cannot be explained by turbulent mixing, but

it is strongly linked to the northeast wind sector, where agricultural sources are closest (approx. 1.5 km distance, see also Fig. S1 in supplement). Therefore this is assumed to be an artifact caused by these agricultural point sources.

Beside the strong influence of agricultural management on seasonal concentration variability, temperature is usually a substantial driver. Higher temperature indirectly leads to higher NH_3 concentrations (Fig. 7), because it is often related to low relative humidity and thus favors ammonia release from the condensed phase towards the gas phase. But increasing air temperature can also be linked to lower NH_3 concentrations, especially observed in this study in late April and early May (Period IV). With higher temperature a larger amount of acidic gas or particle species are present in the atmosphere, which usually leads to reactions of ammonia to ammonium salts such as ammonium nitrate (Kim et al., 2011). We observed higher NH_3 concentrations when it was dry and/or cold, whereas rainy conditions led to lower ammonia concentration levels, which confirms findings reported by Mosquera et al. (2001). In our study, high concentrations were at the same time triggered by local sources northeast of the tower as described above. This phenomenon was particularly observed in the second half of Period III under decreasing air temperatures.

Another driver for the observed concentration pattern might be leaf surface wetness. Peatlands in general, particularly during colder parts of the year, are moist environments where ammonia can easily be taken up by wet surfaces. On the other hand, it is released back to the atmosphere when surface water, e.g. dew, evaporates during morning and midday hours (Walker et al., 2006; Wentworth et al., 2014; Flechard et al. 1999; Wu et al. 2009; Burkhardt et al. 2009), which then causes again rising ammonia concentrations as observed in this study (Fig. 4). Other authors observed the concentration peak earlier in the morning, e.g., Walker et al. (2006) and Wolff et al. (2010) at arable land and grassland, respectively. The observed peak in our study might have been shifted because of the much higher humidity at our peatland site, indicated by small water pools, causing a longer duration of the evaporation process.

The monthly integrated ammonia concentration of 16.8 ppb in March from QCL measurements was in good agreement with those values measured by DELTA denuder and passive samplers. The latter approaches resulted in 14.5 and 15.2 ppb for DELTA and passive samplers, respectively, indicating their robustness and validity as low-cost methodologies for longterm air quality monitoring. As the time of exposure of DELTA denuders and passive samplers was not consistent with our QCL measurements due to instrument failures and campaign duration in February, April, and May, as well as due to highly variable concentrations during that time, we could not directly compare QCL numbers with those from the monthly integrating methods.

3.2 Ammonia exchange and its biophysical controls

Half-hourly measured ammonia fluxes ranged mainly within -80 and $20 \text{ ng N m}^{-2} \text{ s}^{-1}$ with only very few values as low as -300 and as high as $300 \text{ ng N m}^{-2} \text{ s}^{-1}$ (upward fluxes positive, Fig. 3, lower panel). On average the peatland was a sink with a mean flux of $-17.4 \text{ ng N m}^{-2} \text{ s}^{-1}$. At the beginning of the campaign (Period I and II), ammonia deposition was consistently

recorded ($<-20 \text{ ng N m}^{-2} \text{ s}^{-1}$; Fig. 3), while in Periods III and IV the average deposition decreased to values $>-10 \text{ ng N m}^{-2} \text{ s}^{-1}$ and the exchange became clearly bi-directional.

We observed considerable diurnal variability in ammonia fluxes throughout the campaign. The average diurnal flux showed moderate uptake around $-25 \text{ ng N m}^{-2} \text{ s}^{-1}$ from midnight to 10 a.m., near-neutral exchange around noon, and highest uptake of $-40 \text{ ng N m}^{-2} \text{ s}^{-1}$ in the late afternoon and early evening hours (Fig. 6 and 7). Separated by episodes, Period II showed the largest amplitude, whereas Period IV revealed only little variability over the daily course. Separated by surface wetness, considerably higher uptake was observed in the late afternoon and early evening when there was no precipitation recorded at the site than during times when it was raining. Furthermore, ammonia exchange shifted from around zero to net emissions at noon during rain events. Typically rain events were mostly associated with winds from the south-west, where the influence of the agriculture is lowest. We also used the elapsed time after the last recorded rain as a proxy for leaf surface wetness (Fig. 6). We found that higher ammonia uptake coincided with a larger number of days passed since last rain.

Springtime ammonia uptake at sites that were highly influenced by fertilization and other local sources of ammonia in the surrounding area has been reported earlier, e.g. by Mosquera et al. (2001), who found considerable average deposition fluxes at their semi-natural grassland site. Beside management, they showed that higher surface wetness in spring amplified local ammonia deposition, while net emission was typically found only in summer (see also Wichink Kruit et al., 2007). An undisturbed peatland site is likely to be a higher ammonia sink than managed grasslands due to a lower nitrogen status and therefore lower ammonia compensation point. However, with the chronically high atmospheric nitrogen loads caused by agriculture over several decades our peatland is presumably not such an efficient sink anymore. With this assumption and the decreasing uptake over the progression of the measurement campaign (box plots in Fig. 3), saturation effects might have played a role in biosphere-atmosphere exchange characteristics. Flechard and Fowler (1998) already showed that peatland vegetation under its common wet conditions may not necessarily be an 'almost perfect sink' as was reported by Duyzer (1994) due to nitrogen saturation effects in heathland plants caused by persistently high ammonia deposition (approx. $13 \text{ kg N ha}^{-1} \text{ yr}^{-1}$, 2.5 times more than in this study) induced by local sources.

Regarding the diurnal flux patterns observed in this study, significantly different exchange characteristics have been reported elsewhere. For example Horvath et al. (2005) and Wichink Kruit et al. (2007) found highest deposition rates in early morning hours due to dew formation at their grassland sites with decreasing deposition and even emission afterwards through drying leaves and stomatal release. Their finding, i.e. the ecosystem emits only under dry conditions is contrary to our observations (Fig. 6 and 7).

Regarding the entire campaign, a significant tipping point in ammonia exchange was found on 15 March, where higher deposition rates changed to much lower deposition and bi-directional exchange (*cf.* Fig. 8 and 11; Section 3.3). This tipping point followed 10 days after the onset of spring, occurring on 5 March, as indicated by CO_2 uptake shown as significant gross primary productivity (GPP) in Hurkuck et al. (2016) and was mainly triggered by a significant weather change. A huge pressure drop and daily minimum air temperatures exceeding 5°C for the first time in that particular year were recorded.

While conditions were dry until 15 March, 25 mm precipitation was recorded during the following nine days. Thus, a likely reason for decreasing uptake after 15 March might have been a shift of the stomatal compensation point of the local active peatland vegetation due to higher temperatures as was shown by Milford (2004). Similar to findings in Kim et al. (2011), precipitation after 15 March significantly reduced ammonia concentrations in ambient air while probably the ratio of wet to dry deposition considerably increased.

The tipping point from higher to much lower ecosystem ammonia uptake was also observed in the course of deposition velocities (v_d) (Fig. 8 upper panel, tipping point in this depiction not visible, see also supplement Fig. S3). Mean v_d is defined as the ammonia flux divided by concentration over a given period and is still a valuable indicator to verify dry deposition inferential models such as AUSTAL2000 (Janicke, 2002), even in the case of bi-directional exchange (in the case of emission flux, v_d is negative). In our study mean v_d decreased from approx. 0.5 cm s^{-1} at the beginning of the campaign to approx. 0.2 cm s^{-1} on 15 March and remained relatively constant until the end of the campaign. Compared to $v_{d,\max}$ ($v_{d,\max} = (R_a + R_b)^{-1}$) the deposition velocity is, with a few exceptions, always smaller, confirming the plausibility of the data (see Fig. 8 upper panel). The deposition velocity values are significantly lower than those reported by Schrader and Brümmer (2014), who found a median and weighted average of 0.7 and 0.9 cm s^{-1} , respectively, for the category ‘semi-natural’ ($n=19$) in their literature review. In that study, single site values of v_d were ranging between 0.1 and 1.8 cm s^{-1} .

A continuous time-series of the canopy compensation point was derived (Fig. 8, lower panel), using the relation after Nemitz et al. (2000) for a single-layer canopy compensation point resistance model $\chi_c = F_t \cdot (R_a\{z - d\} + R_b) + \chi_a\{z - d\}$, where χ_c ($\mu\text{g m}^{-3}$) is the canopy compensation point at the notional height of trace gas exchange z_0' (m), $\chi_a\{z - d\}$ ($\mu\text{g m}^{-3}$) the air NH_3 concentration measured at the aerodynamic reference height $z-d$ (m), F_t ($\mu\text{g m}^{-2} \text{ s}^{-1}$) is the total NH_3 flux measured by the eddy covariance system, $R_a\{z-d\}$ (s m^{-1}) is the aerodynamic resistance at the reference height, and R_b (s m^{-1}) is the quasi-laminar resistance to NH_3 exchange. From this, also the canopy emission potential could be calculated. The results indicate that there appears to be a very large canopy compensation point (compared for example with Loubet et al. (2012)) that closely follows the air NH_3 concentration, which triggers emission events and effectively reduces deposition in the way that it prevents NH_3 from depositing at the maximum allowed deposition velocity allowed by turbulence (see Fig. 8 upper panel). The observed tipping point on March 15 is also likely another indicator of increased canopy resistance (R_c) over time, which effectively reflects the increase of the canopy compensation point, which may be related to both stomatal and non-stomatal influences. As can be seen in Fig. 8 (middle panel) the canopy resistance increases from the beginning and is already on a high level in period II. Note that here R_c is derived from a deposition- only model approach (cf. Sutton et al., 1993). Over a coniferous forest, Wyers and Erisman (1998) described an increase in nocturnal R_c , which was interpreted as a consequence of pre-deposited ammonia leading to an alkaline saturation of leaf surfaces. For drying canopy, the same authors also found an increase in R_c in a heathland study (Erisman and Wyers (1993), which is consistent with Walker et al. (2006). Our observations showed that the tipping point occurred in a relatively dry period and also that ammonia uptake remained low during rain events ten days later. This might indicate that at a certain point ammonia exchange was partly no

longer controlled by surface wetness but by the pH on non-stomatal surfaces. However, stomatal plant physiological effects in the form of the seasonal onset of CO₂ uptake as mentioned above was likely be caused by the change in weather and therefore be probably the main reason for the tipping point.

The statistical significance of the ammonia flux dependency on meteorological variables when classified into different ranges of values was checked by means of a Kruskal-Wallis-Test (Tab. 2, see also Tab. S1 for deposition velocity, canopy resistance, canopy compensation point and emission potential). The null hypothesis of identical population was rejected in all cases when the p-value was below the commonly used significance level of $\alpha = 0.05$. A Post-hoc-test confirmed that the distributions in all groups, except for '1–2 days' and '2–5 days' in the category 'days after last rain', were significantly different. Thus, all criteria, i.e. the biophysical factors air temperature, precipitation, surface wetness, and radiation, had a statistically significant influence on ammonia exchange. However, fluxes were not well correlated with air temperature, radiation or concentration ($R^2 < 0.1$) when using simple regression analysis. Milford et al. (2001) reported that these variables usually regulate the ammonia exchange but could not find a good correlation between flux and temperature or radiation either because of many non-linear interrelations (*cf.* Milford, 2004; Yamulki et al., 1996). This can be confirmed by visually inspecting the diurnal cycles shown in Fig. 7. The course of ammonia fluxes is more closely coupled to net radiation and u_* , whereas peaks in concentration, although rather forming a bimodal pattern, follow the shape of air temperature with a 2–3 hour lag. Furthermore, solubility of ammonia is related to temperature, which in turn drives the opening of stomata (Fowler et al., 1998), although the response of radiation is much stronger and the temperature effect on stomatal conductance is often confounded by the stronger vapor pressure deficit effects. With increasing temperature ammonia is less dissociated in the available water reservoirs and the plants are able to easier release it to the atmosphere.

Radiation and temperature are also driving local turbulence, i.e. u_* , which appears also to be a controlling factor for both ammonia concentration and fluxes (Fig. 9). While concentrations were lowest under low turbulence, even when analyzed for single periods, maxima were observed in the 0.1 to 0.2 m s⁻¹ class with slightly decreasing values under increasing turbulence, which is a natural phenomenon due to better mixing and transport in the boundary layer, whereas low concentrations under low turbulence at night are likely an artefact of the local fertilizer management that is usually applied during the day. Highest uptake fluxes, however, were found under low u_* , whereas lowest uptake was found under high u_* . This is confirmed by separating day and nighttime data with highest uptake of ammonia clearly occurring during the night (left and middle panel of Fig. 9). The same patterns were found when plotting u_* vs. normalized fluxes, e.g. v_d (not shown). Decreasing uptake with increasing u_* is rather an unusual finding and could be indicating that the control of turbulence is significantly masked by the strong influence of plant physiological regulation (VPD response of stomata) and agricultural practices at our study site with co-occurring gradual ammonia saturation of the leaf surface. On the other hand it could be a consequence of a stomatal or ground compensation point which follows the daily surface temperature course. The temperature itself increases with increasing u_* , also during nights with clear sky, therewith the compensation point increases, which leads to less uptake and less deposition respectively. Further, at low turbulence local transport from nearby fields and

farms into the peatland occurs, at high turbulence a high proportion is going directly from the sources into long-distance transport. At daytime this is mirrored by the flux, thus it seems like the influence of u_* on the ammonia flux is small compared to concentration. At nighttime there is no tendency except a generally higher flux compared to daytime, which may be an effect of wet surfaces rather than any turbulence effect. It has been reported – even for agricultural sites – that
5 medium to high turbulence favors the magnitude of exchange fluxes, regardless of direction, i.e. emission or deposition (*cf.* Brümmer et al., 2013).

The interdependency of the ammonia flux and concentrations is shown in Fig. 10. We separated emission from deposition periods and bin-averaged the concentration data (for an extract without bin-averaging see supplement Fig. S2). Our observations are consistent with Milford (2004) who also found both increasing emission and deposition under elevated
10 concentration, also when separating by wind direction (data not shown). It remains a matter of speculation whether the flux controls the concentration or vice versa as the relationship is highly controlled by plant nitrogen status and at least to some extent by biometeorological variables as mentioned above. However, Milford's (2004) statement that the concentration may still determine the flux during deposition periods, whereas during emission periods it may be the ammonia flux itself, which is controlling the concentration, is also likely to be applicable for our site during deposition periods. In emission periods, . a
15 coincidence of flux drivers and high concentration levels which were high due to advection from the local sources, is the more realistic reason for the relationship in Fig.10 (right panel). The nearest agricultural ammonia point source was 1.5 km away from the tower. With a measurement height of 2.5 m none of the sources were located within the flux footprint, thus we can largely exclude effects from flux heterogeneity such as a direct contribution of the sources to the measured vertical fluxes. However, there might be still large scale transport processes as outlined in a study by Mohr et al. (2015) that
20 influence ammonia concentrations at the site. What is needed at this point is not just an observation on one field and one ecosystem, but a landscape-scale or regional-scale model of emission, dispersion, chemistry, exchange and deposition, which makes it possible to work on this question interactively.

3.3 Measured vs. modeled fluxes and cumulative exchange

The comparison of measured and modeled daily mean and cumulative half-hourly ammonia fluxes is given in Fig. 10 and 11
25 (for one week with half-hourly values see supplement Fig. S4). We found a considerable mismatch between modeled and measured fluxes with the latter showing higher uptake in Period I and lower uptake in Period III than model outputs. In contrast, during Periods II and IV measured and simulated fluxes run fairly parallel, with the exception of a short period of overestimated deposition during last week of March, indicating that, on average, during these times the model is able to reproduce the measured fluxes well. The interdependency of modeled ammonia fluxes and measured concentrations is very
30 similar to the measured ones except that the model does not exhibit any emission and the fluxes are generally lower with increasing concentrations (Fig. 10). We do not see the larger measured deposition fluxes during Periods I and II as being indicative of faulty measurements. Instead, we suspect that under the local pollution climate during the measurement period, the model predicts a too large non-stomatal resistance (R_w). The Massad et al. (2010) parameterization uses a so-called acid

to ammonia ratio (AAR), i.e. the molar ratio of the sum of HNO_3 , SO_2 , and HCl concentrations to the NH_3 concentration, to scale the minimum allowed R_w in the model. The very low AAR (0.07 to 0.11) measured at the peatland site increases the baseline (acid ratio = 1) minimum R_w of 31.5 s m^{-1} roughly 9- to 14-fold. A comparison with non-stomatal resistances inferred from night-time measured fluxes and modeled $R_a(z-d)$ and R_b (cf. Wichink Kruit et al., 2010) reveals that this is too large for periods I and II (not shown here); however, in Periods III and IV, when temperatures rise and thermodynamic equilibria are shifted towards ammonia in the gas phase – increasing apparent R_w and decreasing deposition on the external leaf surfaces – the model is able to reproduce the measured R_w well. Furthermore, HNO_3 , SO_2 and HCl concentrations were only measured on a monthly basis using DELTA denuders, thus introducing some uncertainty to the estimated acid ratio and therefore modeled R_w . A detailed investigation on R_w parameterizations can be found in Schrader et al. (2016, under review).

Regarding the canopy compensation point (Fig. 8, lower panel) it must be considered that in a unidirectional framework, this high canopy compensation point increases the effective canopy resistance R_c (Fig. 8, middle panel), and it appears to have a much larger influence on the observed fluxes than the atmospheric resistances. The constant (stomatal) emission potential from the Massad et al. (2010) model is much lower than the observed canopy emission potentials, and the stomatal compensation point is only a function of temperature, not of the ambient NH_3 concentration, which may be an indicator that, at this site, there is another, ambient concentration-dependent bidirectional pathway that is not being modelled (e.g. wet surfaces; as described, for example, by Burkhardt et al. (2009) for the case of leaf wetness).

The cumulative exchange after approx. 9 weeks of observation resulted in total deposition of 911 and 857 $\text{g NH}_3\text{-N ha}^{-1}$ for measurements and model output, respectively, thus matching relatively well over a longer period despite considerable deviation in Periods I and II. Hurkuck et al. (2014) estimated annual ammonia deposition of 8.5 kg N ha^{-1} at the same study site using denuder filter systems in combination with inferential modeling. They found considerable seasonal variability and also accounted agricultural practices in the surrounding area as the main driver for ammonia exchange variability. Extrapolating our measurement-based campaign total time proportionally to an entire year results in a net deposition estimate of approx. $5.3 \text{ kg NH}_3\text{-N ha}^{-1}$. Reasons for the mismatch might be the fact that our measurements are based on a much higher time resolution leading to a more accurate representation of emission components in the net signal of the eddy-covariance fluxes. Additionally, R_w is based on a much more complex parameterization in Massad et al. (2010), including AAR and a correction for temperature and leaf area index, than those methods used in Hurkuck et al. (2014). However, even with an approx. $3.2 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ lower ammonia deposition, our study site remains a substantial total nitrogen sink. Adding the numbers presented in Hurkuck et al. (2014), i.e. another $2.4 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ dry deposition of HONO, HNO_3 , and NH_4NO_3 aerosols as well as $14 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ as wet deposition, total nitrogen deposition results in approx. $21.7 \text{ kg N ha}^{-1} \text{ yr}^{-1}$, thereby more than four times exceeding the ecosystem-specific critical load of $5 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ (cf. Bobbink et al., 2010; UNECE, 2004).

4 Conclusions

Eddy-covariance flux measurements of ammonia using a QCL in combination with an ‘inertial inlet box’ were conducted at a peatland site in an agricultural landscape. This methodology has high potential for (1) an establishment in long-term observation networks with the aim to improve nitrogen budgets and transfer calculations at local and regional scale as well as (2) providing deeper insight into the mechanisms of ammonia transfer and the ecosystems’ responses to ammonia loads in the atmosphere by offering continuous flux observations at unprecedentedly high temporal resolution. In the present study, we interpret changing diurnal patterns of ammonia concentration and fluxes as well as a tipping point followed by decreasing deposition velocities and increasing canopy resistance, as a sign of non-stomatal leaf surface NH_3 saturation in response to elevated NH_3 from agricultural activities (i.e. manure spreading), but also by delayed plant physiological effects after the onset of the growing season. Temperature, radiation, and surface wetness were identified to partially regulate ammonia exchange at the site; however, the seasonal concentration pattern was clearly dominated by emissions from agricultural practices in the surrounding area. Overestimations of non-stomatal resistances due to low acid to ammonia ratios were assumed to be responsible for deviations between modeled and measured flux estimates. On a total cumulative campaign basis the estimated net NH_3 exchange differed only by 6 % between the model and independent flux measurements. Lower QCL-based deposition values than those of using denuder systems were likely due to a better representation of the emission component in the net signal of eddy-covariance fluxes as well as better adapted site-specific model parameters, particularly R_w . Further research is needed on long-term stability of the QCL system and on avoidance of unintended reactions of ammonia within the inlet and sample tube.

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Tables

Table 1:

Characterization for four subperiods of the measurement campaign (I to IV) with different NH_3 concentration (c_{NH_3} , mean, standard deviation, maximum and minimum) and weather regimes (mean air temperature $\bar{T}_a$ sum of precipitation P and net radiation R_n).

5

Period	Time	n	$\bar{c}_{\text{NH}_3}$ (ppb)	SD of c_{NH_3} (ppb)	Min c_{NH_3} (ppb)	Max c_{NH_3} (ppb)	$\bar{T}_a$ (°C)	P (mm)	R_n (MJ m ⁻² d ⁻¹)
I	Feb 18 – Mar 04	643	11.3	5.6	1.2	34.7	4.5	6.4	1.71
II	Mar 05 – Mar 15	528	22.2	12.1	3.4	87.8	6.3	3.7	4.86
III	Mar 16 – Apr 10	1296	16.5	11.7	2.2	87.4	8.6	35.5	5.19
IV	Apr 29 – May 07	450	8.0	4.2	1.7	21.0	10.3	15.5	8.14

Table 2: Data classification and results of Kruskal-Wallis test on the NH₃ flux; the null hypothesis of identical population is rejected, when the p-value is below the significance level of $\alpha = 0.05$, the Post-hoc-test confirms if the distributions in all groups are significantly different, if not, the equal groups are listed (see Section 3.2 for further details and Tab. S1 for tests on other variables).

5

Meteorological variable	Groups			p-value	Post-Hoc
	1	2	3		
Air temperature	<5°C	5 – 10°C	>10°C	<0.001	All differ
Precipitation	0 mm	>0 mm		<0.001	All differ
Days after last rain	1 – 2 d	2 – 5 d	>5 d	<0.001	1=2
Net radiation	<0 W m ⁻²	0 – 150 W m ⁻²	>150 W m ⁻²	<0.001	All differ

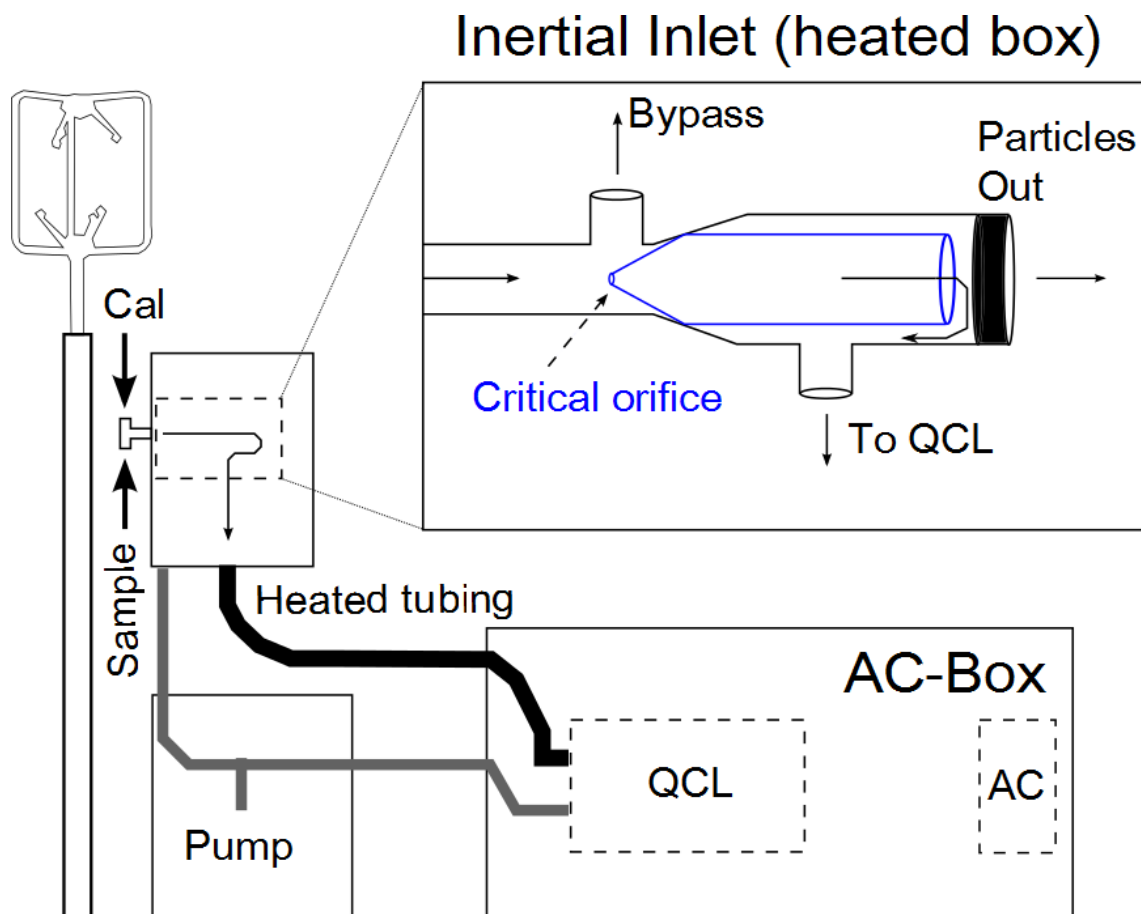


Figure 1: Schematic overview of the measurement setup. An ultrasonic anemometer (height 2.5 m) is mounted closely to the heated 'inertial inlet' box (39 cm × 34 cm × 21 cm) containing a critical glass orifice to reduce the pressure regime inside the sample line (total length 3.5 m) to 5.3 to 8 kPa. After passing the critical orifice, a sharp turn of the flow path leads to a reduction of particles (>300 nm) of approx. 50 %, thereby reducing unintentional chemical interactions. The heated tubing (black tube) leads the sample air (flow rate = approx. 17 l min⁻¹) to the Quantum Cascade Laser (QCL), which is housed in an air-conditioned (AC) box. The bypass air and the particles out flow are combined in the inertial inlet box and together sucked to the pump (left part of the grey tube).

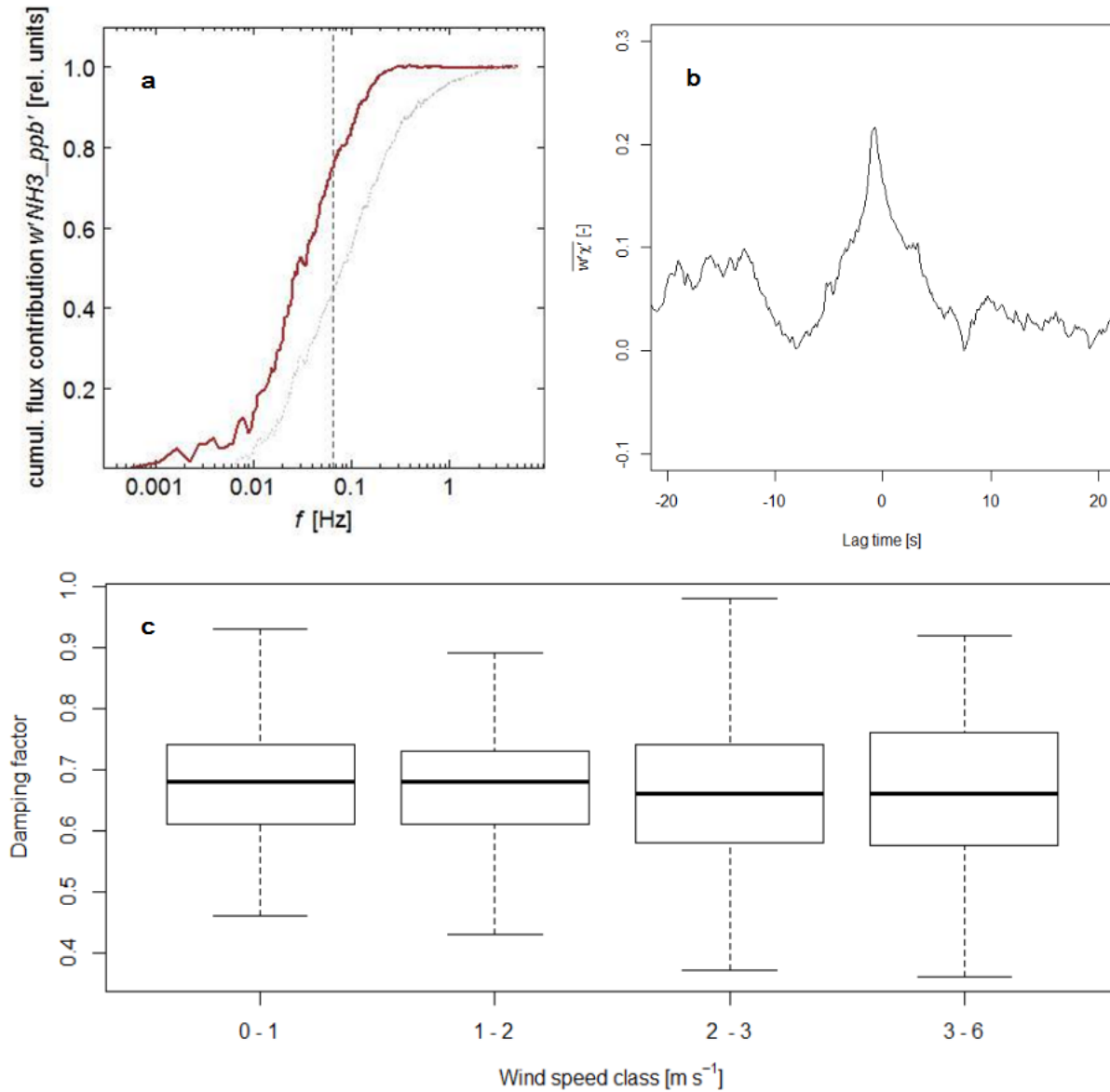
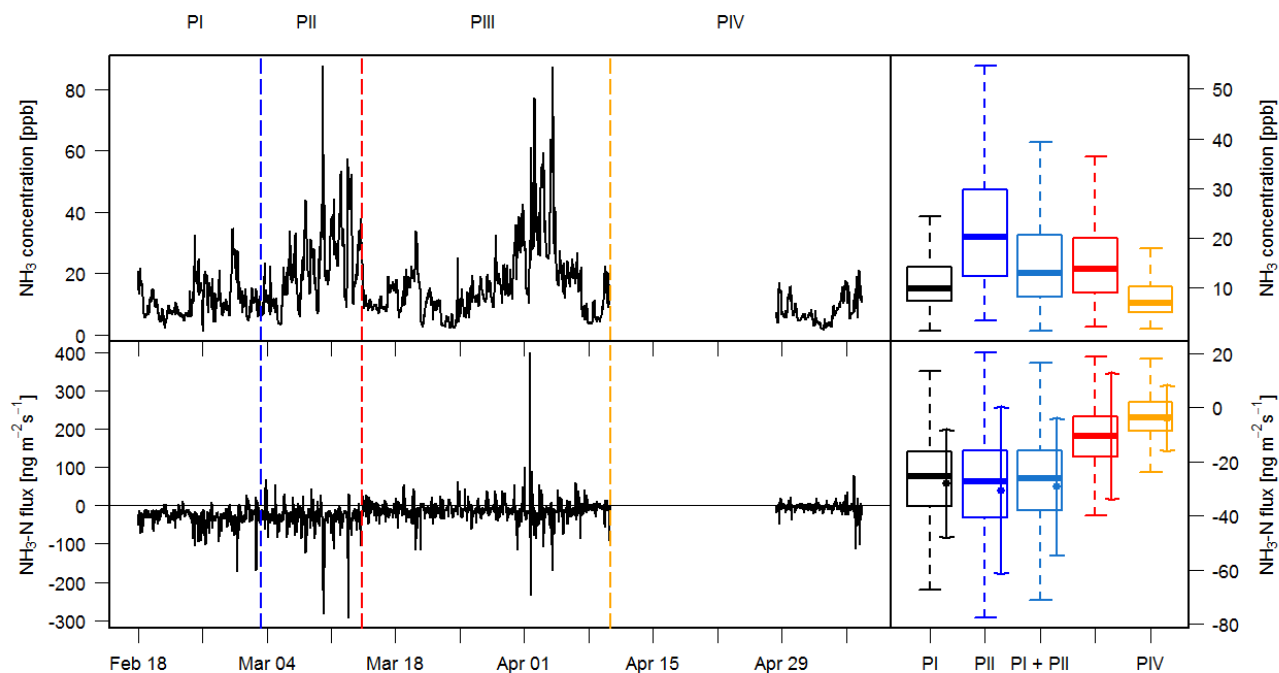


Figure 2: a) Normalized flux ogives, i.e. cumulative cospectra, of sensible heat (grey) and NH_3 flux (red). Data were recorded on 18 February 2014, 03:30 – 04:00 pm. The vertical dotted line indicates the frequency at which the two ogives are compared to determine the empirical high-frequency damping factor for the NH_3 flux. b) Example of a covariance function of vertical wind speed and ammonia concentration (19 Feb 2014 02:30 pm), after correction, peaking at a time lag of 0 s. c) Variation of the

damping factor separated by wind speed classes. Bold horizontal lines in boxplots show the median, fine horizontal lines indicate lower and upper quartile values, whiskers represent 1.5 times the interquartile range.



5 **Figure 3: Half-hourly measured ammonia concentrations and fluxes. Vertical lines indicate beginning of periods listed in Table 1. Each period is represented by a boxplot with bold horizontal lines showing the median, fine horizontal lines indicating lower and upper quartile values, whiskers representing 1.5 times the interquartile range and dots with arrows indicating the mean and standard deviation.**

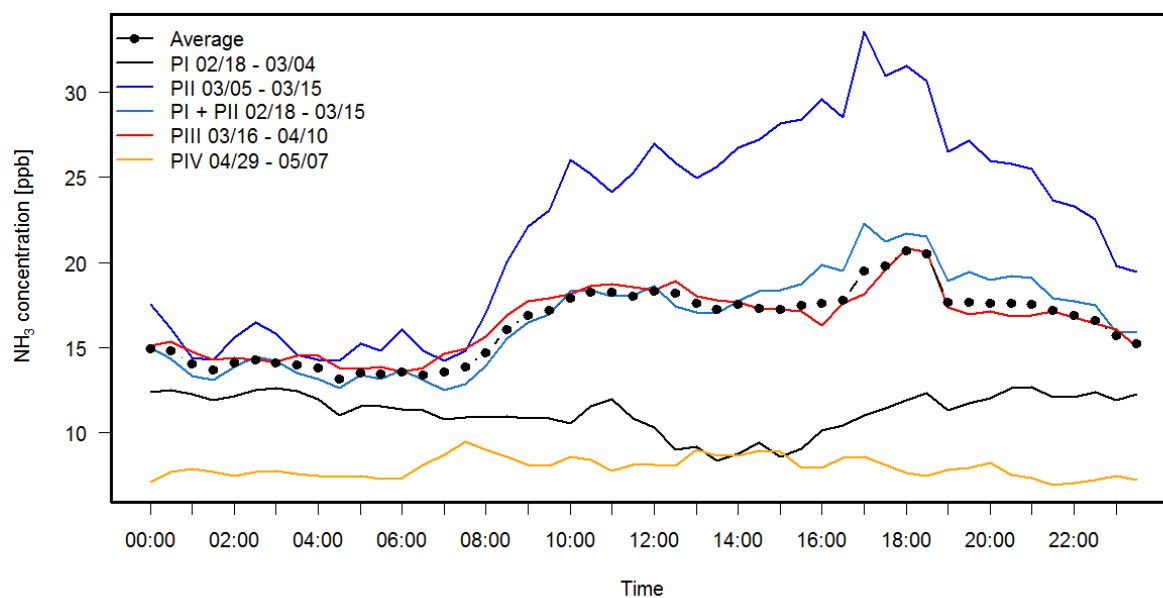


Figure 4: Mean diurnal variation of ammonia concentrations separated into different periods.

5

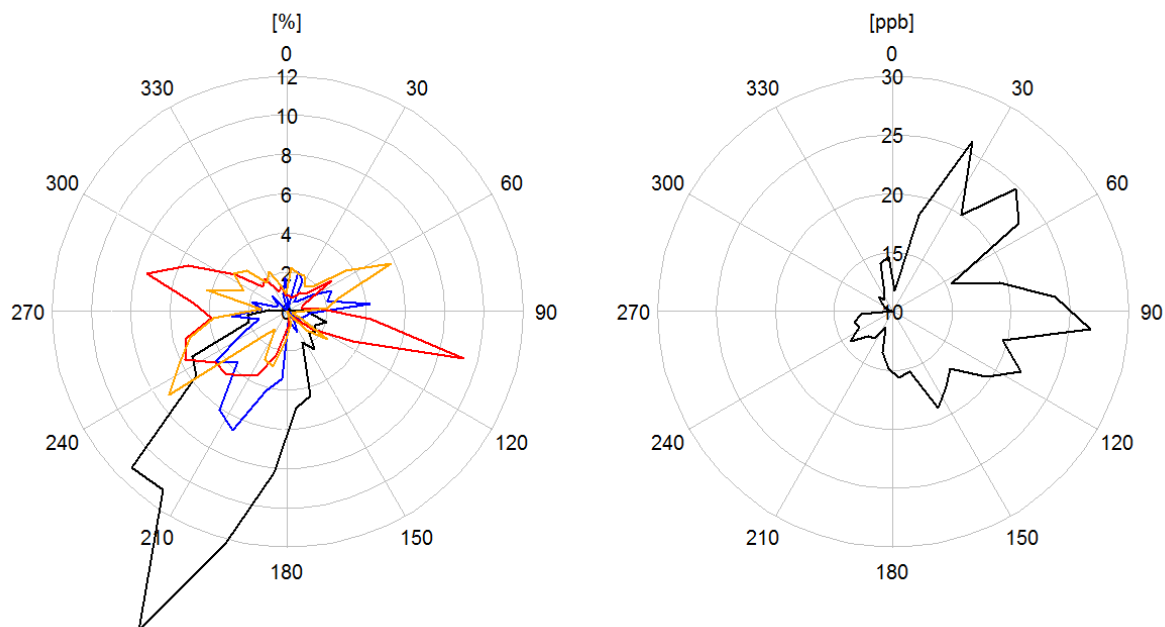
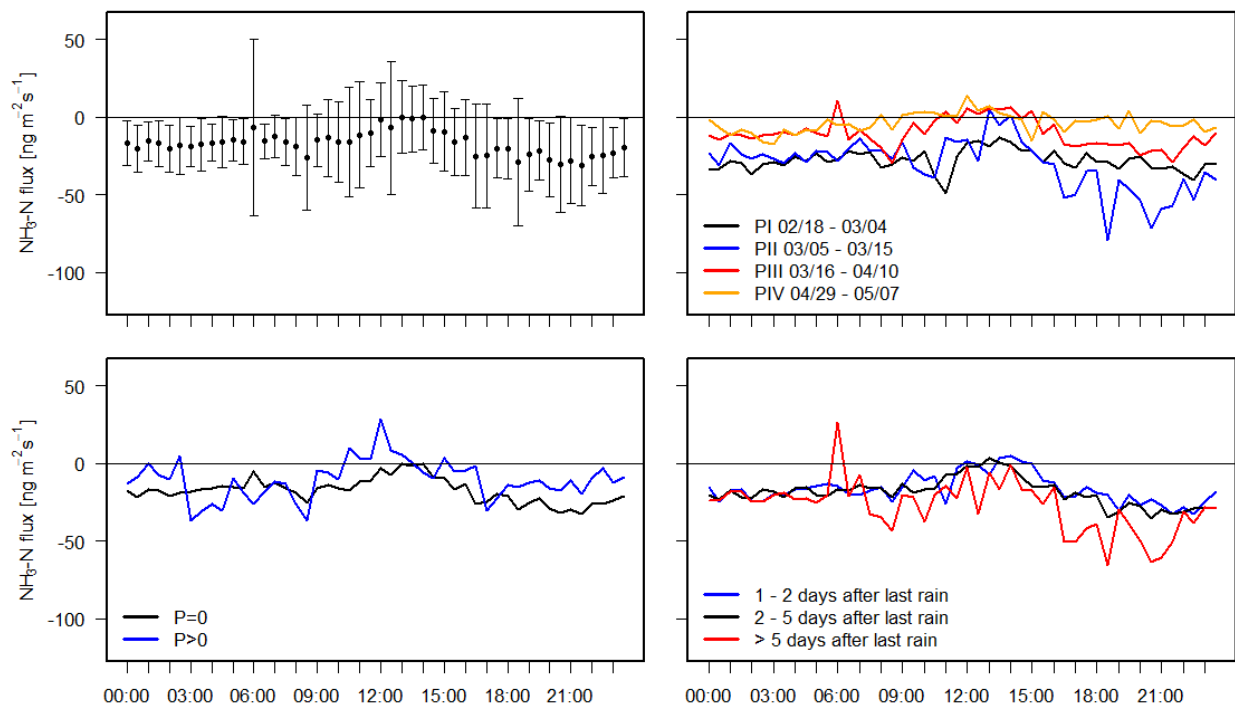


Figure 5: Frequency distribution of wind direction for each period (left panel, PI (02/18 – 03/04) black, PII (03/05 – 03/15) blue, PIII (03/16 – 04/10) red, PIV (04/29 – 05/07) orange) and ammonia concentration (right panel) for each 10° wind sector.



5 **Figure 6: Mean diurnal variation of ammonia fluxes with standard deviation (upper left panel), separated by periods (upper right panel), by precipitation (lower left panel), and by days after last rain (lower right panel).**

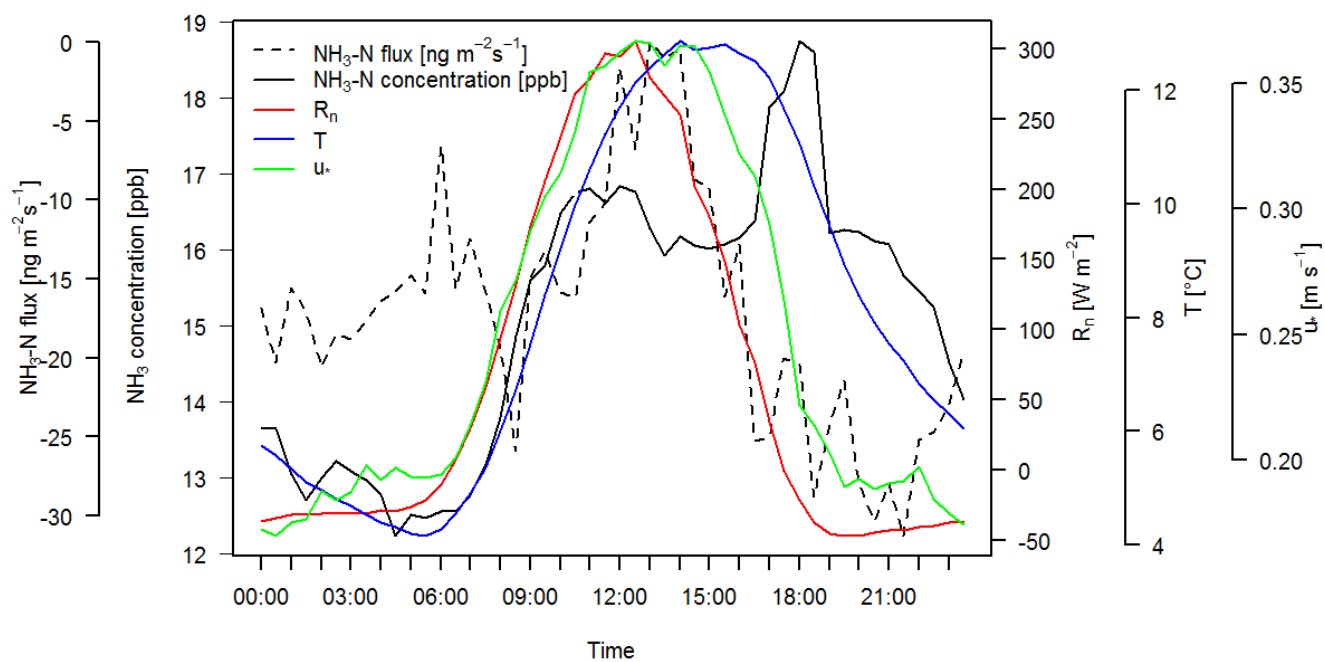


Figure 7: Mean diurnal cycles of ammonia concentration, ammonia flux, net radiation, air temperature, and friction velocity.

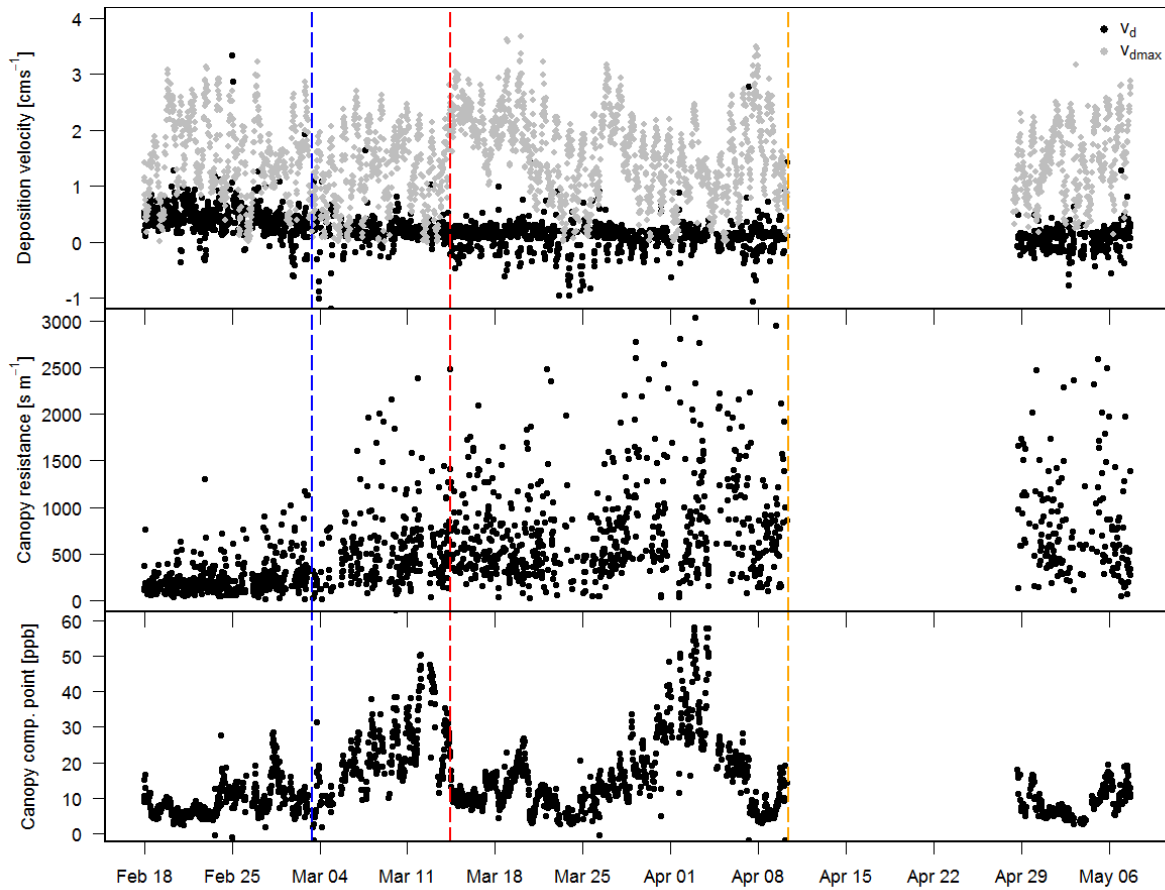


Fig. 8: Upper panel: Deposition velocity, negative values indicate emission, and maximum deposition velocity, middle panel: canopy resistance, negative values are not shown, as they are not defined in the resistance framework and mostly correspond to phases of emission, lower panel: canopy compensation point derived from measured data. Note that the upper and middle panels are based on a one-layer deposition-only and the lower panel to a one-layer canopy compensation point framework (cf. Sutton et al. 1993).

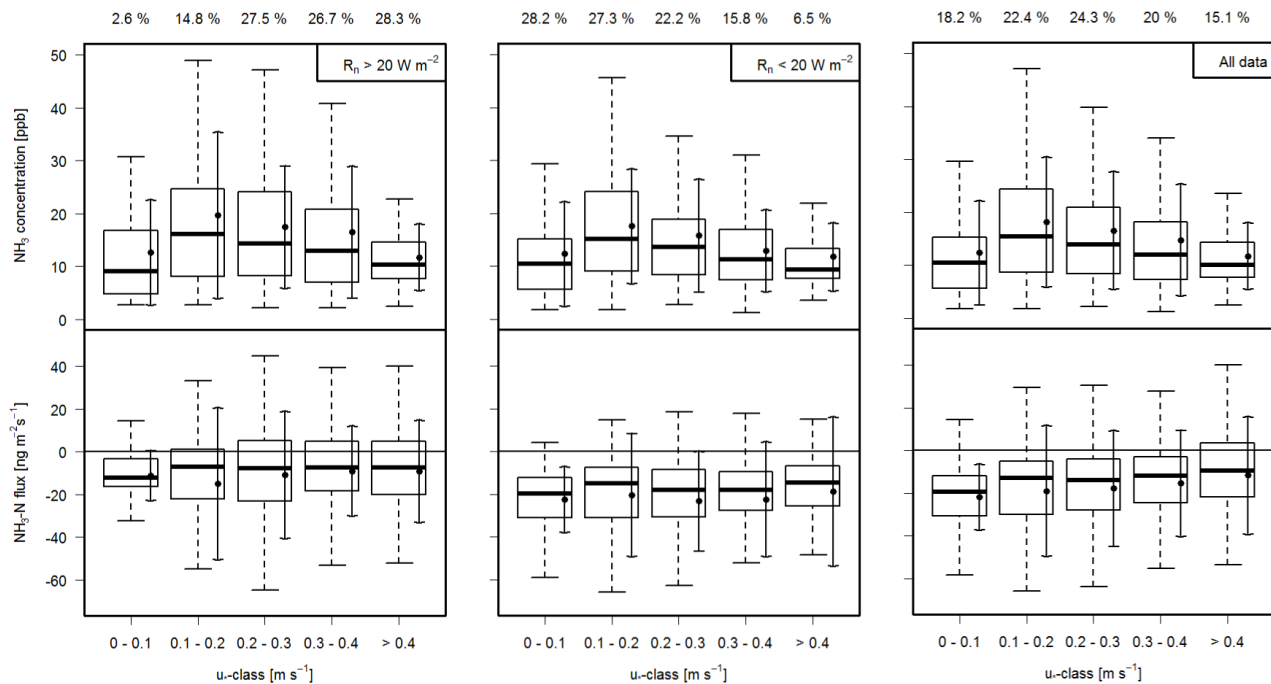
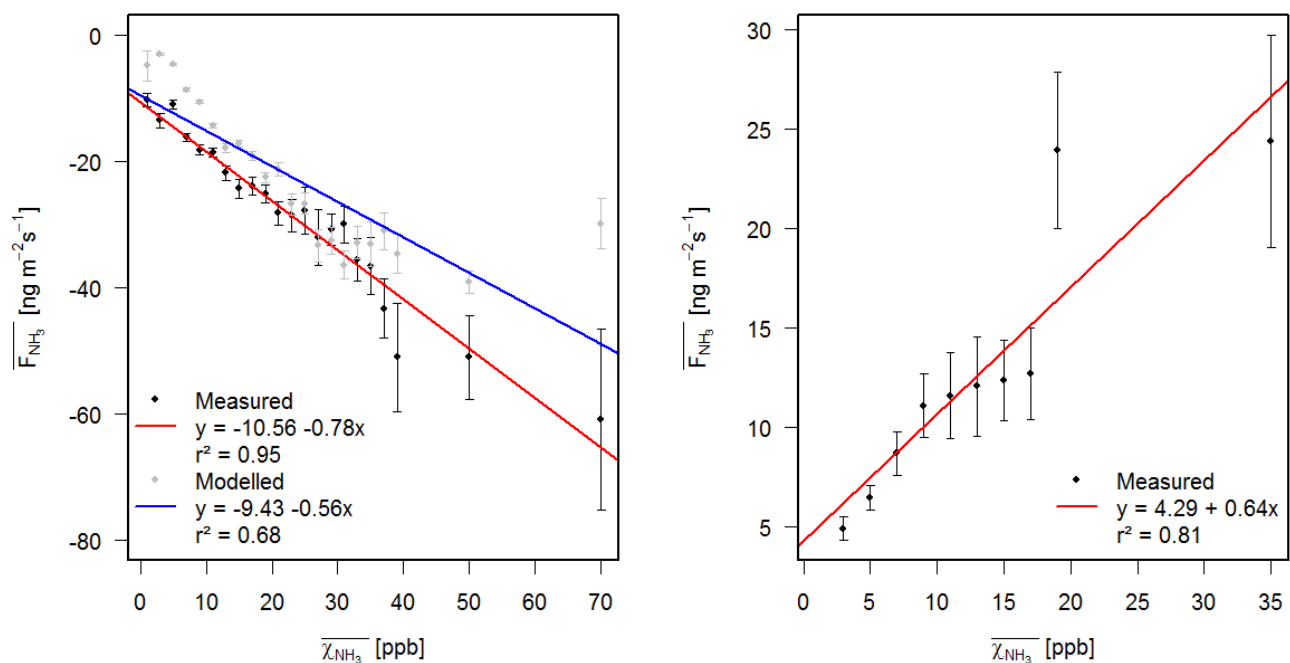


Figure 9: Dependency of ammonia concentrations and fluxes on u_* , with panels from left to right showing daytime ($R_n > 20 \text{ W m}^{-2}$), nighttime ($R_n < 20 \text{ W m}^{-2}$), and all data, respectively. Bold horizontal lines in boxplots show the median, fine horizontal lines indicate lower and upper quartile values, whiskers represent 1.5 times the interquartile range and dots with arrows indicate the mean and standard deviation. Values in upper row specify percentages of data in the respective u_* category given below.



5 **Figure 10: Dependency of NH_3 fluxes, measured and modelled, on bin-averaged NH_3 concentrations (bin sizes are 2 ppb for concentrations < 40 ppb and < 20 ppb, in the left and right panel, respectively; for concentrations > 40 ppb bin sizes are 20 ppb) during deposition (100% of the modelled and 84 % of the measured data, left panel) and emission periods (16% of the measured data, right panel), error bars indicate standard error.**

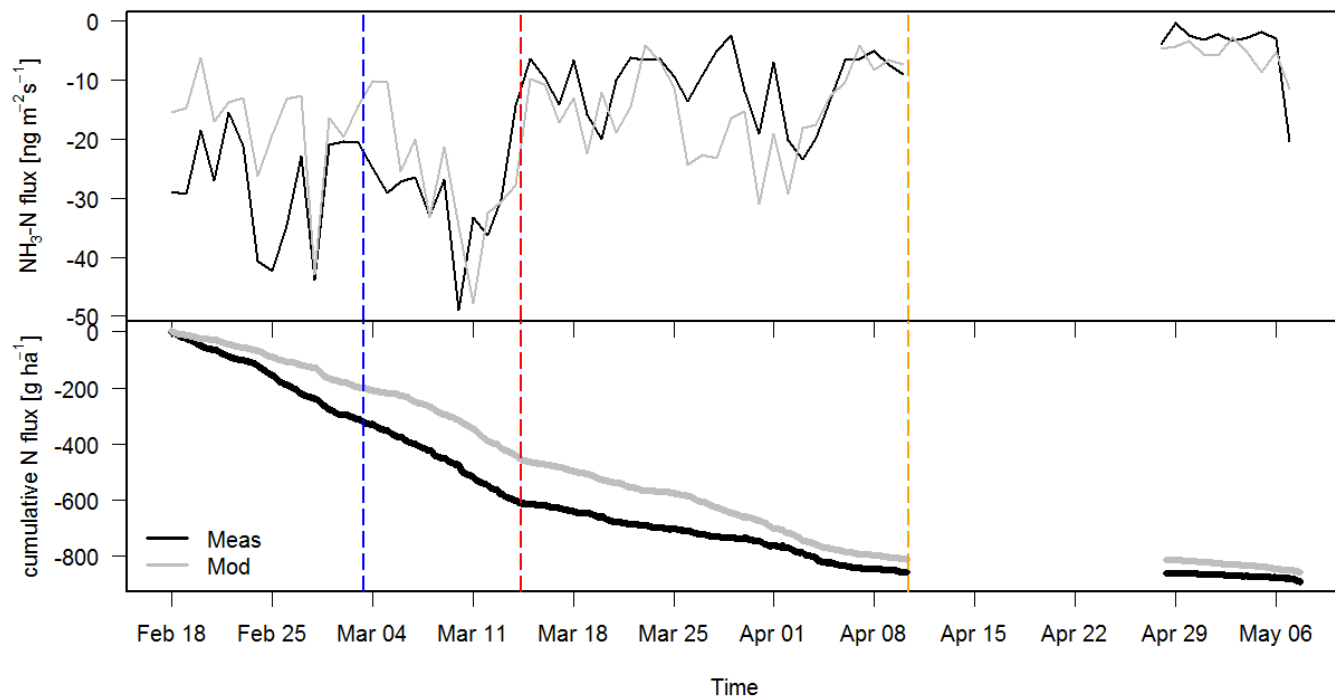


Figure 11: Comparison of measured and modeled daily mean NH_3 fluxes (upper panel) and cumulative NH_3 flux (lower panel) based on half-hourly data during the measurement campaign. Vertical lines indicate the start of Periods II, III, and IV as listed in Table 1.

Supplementary material

Parameters of the Weseley model

We used the original formulation of the Wesely (1989) model with a minimum R_s for H_2O of 200 s m^{-1} , scaled by the ratio of the molecular diffusivities of H_2O and NH_3 :

$$R_s = 200 \cdot \frac{D_{H_2O}}{D_{NH_3}} \cdot \left(1 + \left(\frac{200}{S_t + 0.1} \right)^2 \right) \frac{400}{T \cdot (T - 40)}$$

where S_t is global radiation in $W\text{ m}^{-2}$, D_x is the molecular diffusivity of H_2O and NH_3 in air, respectively, in $m^2\text{ s}^{-1}$, and T is the surface temperature in $^{\circ}C$. Note that we were not able to optimize these parameters due to a lack of data in the dry range, where cuticular deposition is restricted. A $\pm 100\%$ change in the minimum stomatal resistance leads to a change in total cumulative ammonia flux between -7% and $+19\%$ (for 300 s m^{-1} and 100 s m^{-1} , respectively).

Table S1: Data classification and results of Kruskal-Wallis test on the deposition velocity, the canopy compensation point, the emission potential and canopy resistance; the null hypothesis of identical population is rejected, when the p-value is below the significance level of $\alpha = 0.05$, the Post-hoc-test confirms if the distributions in all groups are significantly different, if not, the equal groups are listed (see Section 3.2 for further details).

Deposition velocity	Groups			p-value	Post-Hoc
	1	2	3		
Air temperature	<5°C	5 – 10°C	>10°C	<0.001	All differ
Precipitation	0 mm	>0 mm		0.811	All equal
Days after last rain	1 – 2 d	2 – 5 d	>5 d	0.115	All equal
Net radiation	<0 $W\text{ m}^{-2}$	0 – 150 $W\text{ m}^{-2}$	>150 $W\text{ m}^{-2}$	<0.001	All differ
Canopy resistance	Groups			p-value	Post-Hoc
	1	2	3		
Air temperature	<5°C	5 – 10°C	>10°C	0.149	All equal
Precipitation	0 mm	>0 mm		0.005	All differ
Days after last rain	1 – 2 d	2 – 5 d	>5 d	<0.001	1=2
Net radiation	<0 $W\text{ m}^{-2}$	0 – 150 $W\text{ m}^{-2}$	>150 $W\text{ m}^{-2}$	<0.001	All differ
Canopy compensation	Groups			p-value	Post-Hoc

point	1	2	3		
Air temperature	<5°C	5 – 10°C	>10°C	<0.001	All differ
Precipitation	0 mm	>0 mm		<0.001	All differ
Days after last rain	1 – 2 d	2 – 5 d	>5 d	<0.001	All differ
Net radiation	<0 W m ⁻²	0 – 150 W m ⁻²	>150 W m ⁻²	<0.001	All differ
Emission potential	Groups			p-value	Post-Hoc
	1	2	3		
Air temperature	<5°C	5 – 10°C	>10°C	<0.001	All differ
Precipitation	0 mm	>0 mm		<0.001	All differ
Days after last rain	1 – 2 d	2 – 5 d	>5 d	<0.001	1=2
Net radiation	<0 W m ⁻²	0 – 150 W m ⁻²	>150 W m ⁻²	<0.001	All differ

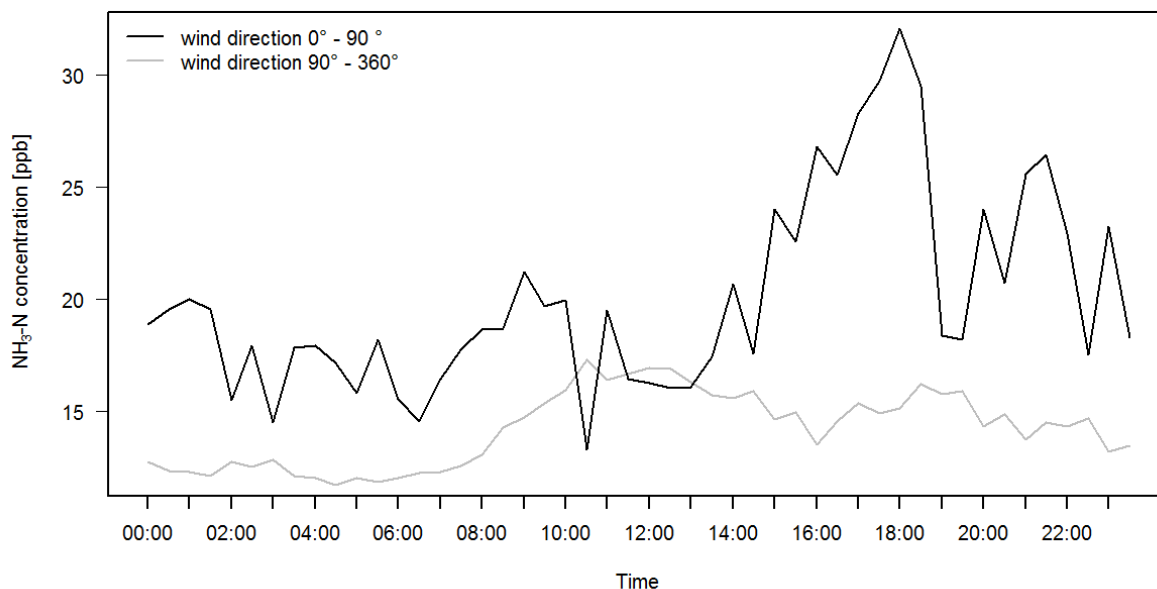


Fig. S1: Mean diurnal variation of ammonia concentrations separated by wind direction.

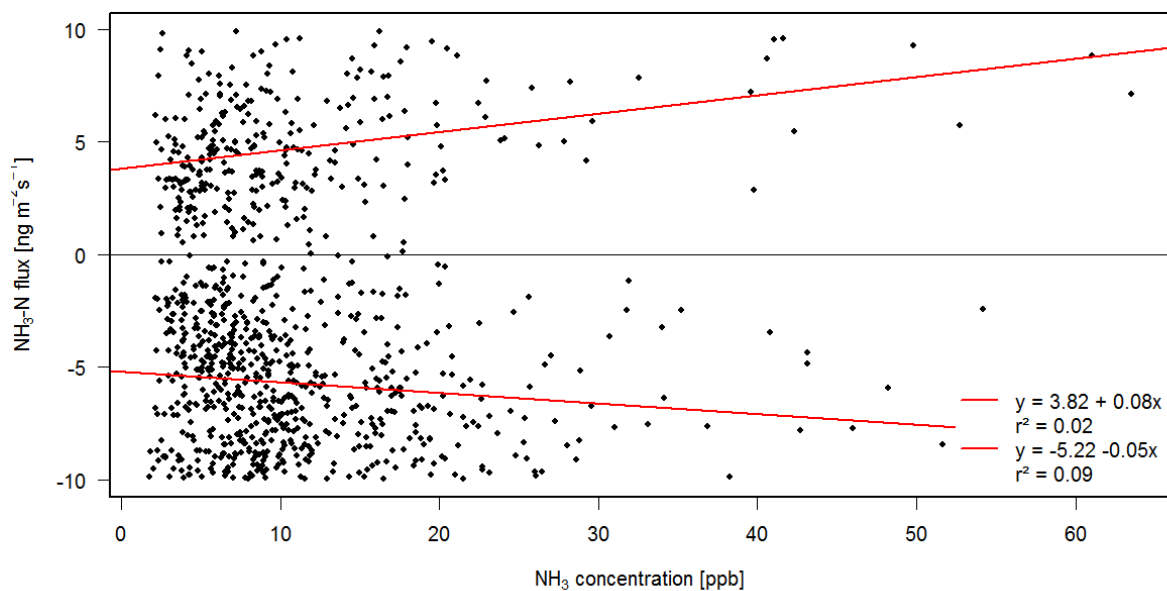


Fig. S2: Half-hourly scatter plot showing the dependency of NH_3 fluxes (only in a range of -10 to 10 ng N m⁻²s⁻¹) on NH_3 concentration, red line: linear regression above for emission, below for deposition, for coefficients and r^2 see legend

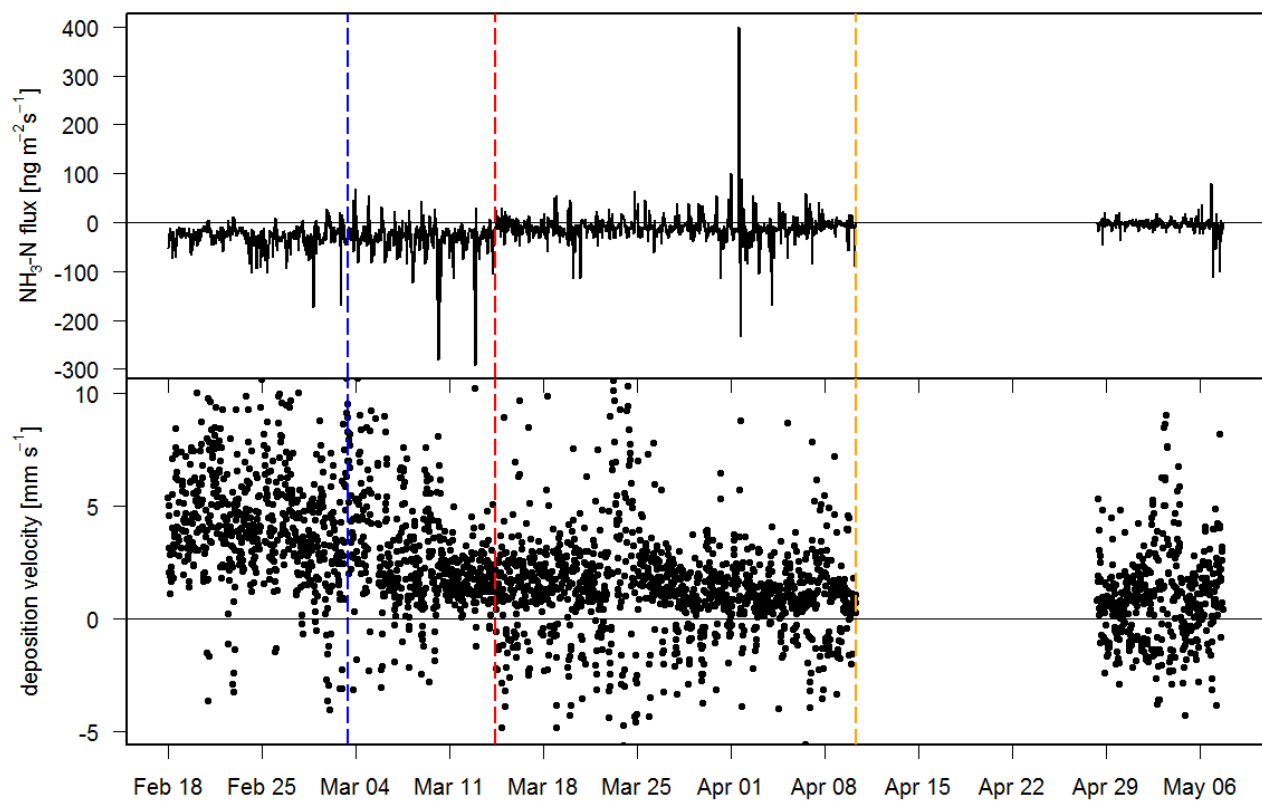


Fig. S3: Half-hourly ammonia fluxes (upper panel) and half-hourly ammonia deposition velocities (lower panel) during the whole campaign.

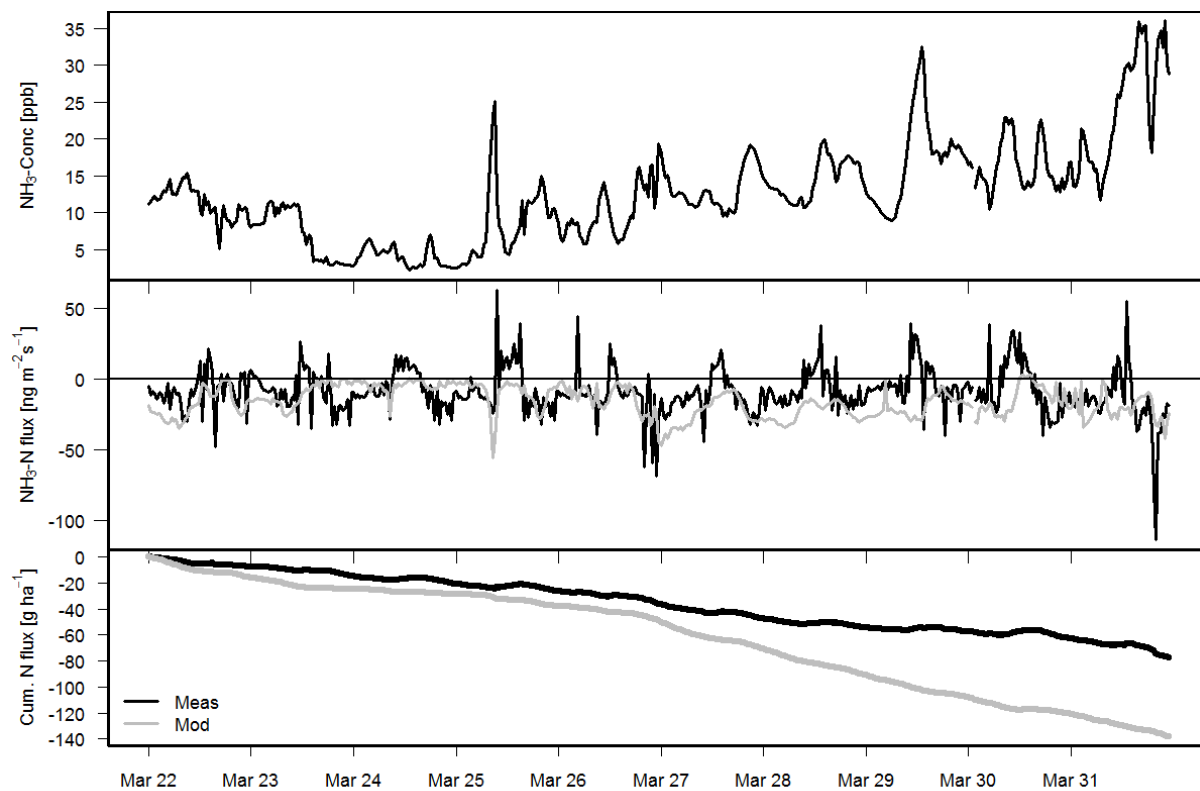


Fig. S4: Measured ammonia concentrations (upper panel), comparison of measured and modeled half-hourly ammonia fluxes (middle panel) and cumulative ammonia flux (lower panel) based on half-hourly data during one week of the measurement campaign.