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Ammonia emissions from deciduous forest after leaf fall

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Abstract. The understanding of biochemical feedback mechanisms in the climate system is lacking knowledge in relation to bi-directional ammonia (NH₃) exchange between natural ecosystems and the atmosphere. We therefore study the atmospheric NH₃ fluxes during a 25-day period during autumn 2010 (21 October to 15 November) for the Danish beech forest Lille Bogeskov to address the hypothesis that NH₃ emissions occur from deciduous forests in relation to leaf fall. This is accomplished by using observations of vegetation status, NH₃ fluxes and model calculations. Vegetation status was observed using plant area index (PAI) and leaf area index (LAI). NH₃ fluxes were measured using the relaxed eddy accumulation (REA) method. The REA-based NH₃ concentrations were compared to NH₃ denuder measurements. Model calculations of the atmospheric NH₃ concentration were obtained with the Danish Ammonia Modelling System (DAMOS). The relative contribution from the forest components to the atmospheric NH₃ flux was assessed using a simple two-layer bi-directional canopy compensation point model. A total of 57.7% of the fluxes measured showed emission and 19.5% showed deposition. A clear tendency of the flux going from deposition of −0.25 ± 0.30 µg NH₃-N m⁻² s⁻¹ to emission of up to 0.67 ± 0.28 µg NH₃-N m⁻² s⁻¹ throughout the measurement period was found. In the leaf fall period (23 October to 8 November), an increase in the atmospheric NH₃ concentrations was related to the increasing forest NH₃ flux. Following leaf fall, the magnitude and temporal structure of the measured NH₃ emission fluxes could be adequately reproduced with the bi-directional resistance model; it suggested the forest ground layer (soil and litter) to be the main contributing component to the NH₃ emissions. The modelled concentration from DAMOS fits well the measured concentrations before leaf fall, but during and after leaf fall, the modelled concentrations are too low. The results indicate that the missing contribution to atmospheric NH₃ concentration from vegetative surfaces related to leaf fall are of a relatively large magnitude. We therefore conclude that emissions from deciduous forests are important to include in model calculations of atmospheric NH₃ for forest ecosystems. Finally, diurnal variations in the measured NH₃ concentrations were related to meteorological conditions, forest phenology and the spatial distribution of local anthropogenic NH₃ sources. This suggests that an accurate description of ammonia fluxes over forest ecosystems requires a dynamic description of atmospheric and vegetation processes.

1 Introduction

Atmospheric ammonia (NH₃) plays an important role in air quality and critical load studies of natural and semi-natural ecosystems. NH₃ is a reactive nitrogen compound (Nᵢ), which contributes to the formation of ammonium aerosols (NH₄⁺) through atmospheric chemical reactions (Hertel et al., 2012) and is leading to deposition to terrestrial and marine ecosystems (de Leeuw et al., 2003; Duce et al., 2008; Massad et al., 2010; Zhang et al., 2010). An enhanced load of Nᵢ in terrestrial ecosystems can increase the rate of acidification and eutrophication processes and thereby reduce biodiversity.
and increase ecosystem vulnerability to extreme weather and insect attacks (Bobbink et al., 2010; Erisman et al., 2007; Stevens et al., 2004; Sutton et al., 2011; Xiankai et al., 2008). In nutrient-limited ecosystems, nitrogen deposition can also work to increase the C-sequestration and growth of vegetation (de Vries et al., 2009).

Budgets of atmospheric NH$_3$ for water and land areas are being carried out using atmospheric models (e.g. Bartnicki et al., 2011; de Leeuw et al., 2003; Geels et al., 2012a; Hertel et al., 2003; Langner et al., 2009). Model calculations indicate that particular forest ecosystems are exposed to critical load exceedances of nitrogen (Geels et al., 2012b; Hertel et al., 2013). Generally the understanding of major biochemical feedback mechanisms in the terrestrial climate system suffers from large uncertainties (Arneth et al., 2010) and lacking knowledge of the bi-directional (two-way) NH$_3$ exchange between the land/water surface and the atmosphere (Massad et al., 2010). Because natural NH$_3$ emissions have been assessed to a rather limited extent (Massad et al., 2010; Nemitz et al., 2001), they are not yet included in operational air pollution models (Menut and Bessagnet, 2010).

Limited field studies have shown that the deposition velocity of NH$_3$ for forests is relatively high and variable (Andersen et al., 1999; Duyzer et al., 1994; Wyers et al., 1992) and thereby complicates the understanding of forest NH$_3$ exchange processes and makes model validation difficult. High deposition velocities for forests were simulated by four inferential models used across European sites, but large differences (up to factor 3) were found between model results (Flechard et al., 2011).

Sources of atmospheric NH$_3$ are conceptually considered to be anthropogenic (e.g. Gyldenkerne et al., 2005; Hertel et al., 2012) and primarily related to agricultural activities (e.g. Bouwman et al., 1997; Reis et al., 2009). In Europe, agricultural emissions arise from farm buildings (34–43 %), manure (22–26 %), fertilisers (17–26 %) and grazing animals (6–10 %) (Skjøth and Geels, 2013). Open water areas (e.g. Barrett, 1998; Sørensen et al., 2003) and natural land areas (e.g. Duyzer et al., 1994) have also been shown to emit NH$_3$. NH$_3$ emissions from forests are generally not included in official emission inventories (Reis et al., 2009) or the more detailed national inventories (Gyldenkerne et al., 2005; Velthof et al., 2012). However, NH$_3$ flux studies of forests indicate bi-directional flux patterns for NH$_3$ (Andersen et al., 1999; Erisman and Wyers, 1993; Sutton et al., 1997; Wyers and Erisman, 1998). Emissions of NH$_3$ from ecosystems are found to take place when the atmospheric NH$_3$ concentration is lower than the stomatal NH$_3$ compensation point (Wichink Kruit et al., 2007; Mattsson et al., 2009; Schjoerring et al., 1998), from decomposing leaf litter (David et al., 2009; Nemitz et al., 2000a), and by cuticular desorption (Pryor et al., 2001). Furthermore, Wang et al. (2011) discovered a seasonal dependence of the NH$_3$ compensation points of beech leaves and found largest emission potential in relation to the late senescent leaves. The effects of leaf NH$_3$ emissions in relation to leaf fall still remain to be quantified, particularly at canopy scale.

The main objective of this paper was to assess the NH$_3$ flux $F_{\text{NH}_3}$ for a Danish deciduous forest in the leaf senescence period using high-resolution atmospheric measurements and local-scale concentration-deposition modelling. We investigate the hypothesis that NH$_3$ emissions occur from deciduous forests in relation to leaf fall by correlating this with NH$_3$ emissions and explore the importance of including such emissions in models. To do this, half-hourly measurements of the NH$_3$ concentration and flux were conducted using the relaxed eddy accumulation (REA) technique for a Danish beech forest site in the leaf fall period 21 October to 15 November 2010 (Fig. 1). The measured concentrations were compared to model calculations using the Danish Ammonia MOdelling System (DAMOS), and a simple two-layer bi-directional canopy compensation point model was used to interpret the measured fluxes.

2 Methods

2.1 Field site

The field station (Lille Bøgeskov) is located in the central part of Zealand (55°29′13″ N, 11°38′45″ E) with a surrounding landscape characterised primarily by agricultural activities. Lille Bøgeskov covers approximately 2.5 km$^2$ with the field station located in the centre of the forest. The field station includes a flux tower (57 m) and a scaffolding tower (24 m) (Fig. 2).
The forest consists predominantly of 82 yr-old beech trees (Fagus sylvatica) with an average canopy height of 26 m. Scattered stands of conifers constitute about 20 % of the forest area. The mean summer peak of LAI has been measured to be 4.6 since year 2000 with maximum LAI just above 5 (Pilegaard et al., 2011). In 2010, defoliation was observed to begin on 23 October and leaf fall to end on 8 November.

The soils are brown and consist of Alfisols and Mollisols. Dead plant material consisting mainly of leaves and twigs from the beech trees constitutes the top 0–3 cm. Below is a 10–40 cm deep organic layer. In the upper organic soil layers, the C/N ratio is about 20 and the pH is low (4–5) (Østergård, 2000).

2.2 Leaf area index

In order to relate the atmospheric data to forest canopy development, the plant area index (PAI) was measured during the growing season (May–November) every 14–30 days using the LAI-2000 Plant Canopy Analyzer with a 270° view cap (LAI-2000 PCA). PAI was estimated using one above-canopy reading and 10 below-canopy readings conducted along an 18 m transect. The above-canopy readings were performed outside the forest edge. Measurements were made during uniformly overcast sky conditions, as recommended by the manufacturer. In order to assess the leaf area index (LAI), observations of leaf defoliation were used to adjust the PAI data by linear interpolation to zero LAI at the time when there were no more green leaves present in the canopy. The end of the defoliation and leaf fall periods were determined from daily digital photos of the canopy using a camera mounted on top of the tower. The uncertainty of measurements was calculated as the standard deviation of the 10 below-canopy readings.

2.3 Local meteorological measurements

The wind components in x-, y- and z-directions were measured at 10 Hz sampling using an ultra-sonic anemometer (Metek-uSonic-3 Scientific) installed above the forest canopy at 34 m height (Fig. 2). Half-hourly averaged values of wind velocity, wind direction, friction velocity, temperature, and Monin–Obukhov length were calculated from the 10 Hz sampling. Precipitation, relative humidity, soil temperature, soil water content, and global radiation were obtained from the European Fluxes Database Cluster (www.europe-fluxdata.eu) as described in Pilegaard et al. (2011).

2.4 Ammonia flux measurements

2.4.1 Relaxed eddy accumulation (REA)

The vertical turbulence-driven flux of NH₃ (F_{NH₃}) was estimated using the REA technique (Businger and Oncley, 1990). REA simplifies the eddy accumulation methods (Hicks and Mcmillen, 1984), where the sampling speed must be proportional to the vertical wind velocity, by relaxing the sampling at a constant flow rate (Businger and Oncley, 1990). REA combines measurements of the vertical momentum flux and the difference between the average trace gas concentration of upward and downward moving eddies.

A system to measure canopy-scale F_{NH₃} (µg NH₃-N m⁻² s⁻¹) based on the REA technique was installed in the flux tower at the forest field station (Fig. 2) in the period from 21 October to 15 November 2010. Two short breaks occurred due to technical work on the instruments. The system consisted of three parts: (1) a sonic anemometer measuring vertical wind speed, (2) an inlet system, and (3) an analytical detection system (Fig. 3) to detect the concentration signals. The sonic anemometer was located in the mast at a height of 34 m to control the conditional sampling of atmospheric NH₃ in the up- and downdrafts respectively. The inlet system, comprised by two wet effluent diffusion denuders (WEDDs), was located just below the sonic anemometer at 33 m. The WEDDs collected atmospheric NH₃ from upward and downward eddies separately by diffusion into a water film (Hensen et al., 2009). The aqueous NH₃ solution from the two WEDDs was pumped with a constant flow directly to the analytical detection system which was located at the top floor of a scaffolding tower. A fluorescent compound was produced by mixing o-phthalaldehyde (OPA), sulphite, and the aqueous NH₃ solution (Sorensen et al., 1994). The liquid was heated to 60 °C to enhance the formation of the fluorescent compound before injection into the fluorescence detector. Detailed information of the WEDD and the analytical
system can be found in Sorensen et al. (1994). To prevent freezing of the fluent when air temperature was near freezing point, the water was mixed with ethanol. The analytical system was calibrated using standard calibration fluids of 0, 10 and 25 ppb NH$_3$. Half-hourly estimates of $F_{NH_3}$ were calculated from Eq. (1):

$$F = \beta \sigma_w (\chi \uparrow - \chi \downarrow),$$

where $\chi \uparrow$ and $\chi \downarrow$ are the average NH$_3$ concentration in the up- and downdrafts respectively, $\sigma_w$ is the standard deviation of the vertical wind velocity $w'$, and $\beta$ is a coefficient to be determined by the probability distribution of $w$. The $\beta$ coefficient is well defined for an ideal Gaussian joint frequency distribution of $w$ and $\chi$. However, turbulent transport, especially over very rough surfaces, often violates the underlying assumption of a linear relationship between $w$ and $\chi$ (Ruppert et al., 2006); thus the use of a $\beta_0$ coefficient determined from a proxy scalar (such as the sensible heat flux) better reflects the correct $\beta$ coefficient for a certain measurement period. A dynamic deadband was introduced as a threshold for partitioning $\chi \uparrow$ and $\chi \downarrow$ (Businger and Oncley, 1990) where sampling only took place when the vertical wind velocity exceeded a predefined deadband velocity $w_0$. The $\beta$-coefficient has to be corrected for the choice of deadband velocity:

$$\beta = \beta_0 \exp \frac{-0.75 \cdot w_0}{\sigma_w},$$

where $w_0$ is the dynamic deadband (set to 0.5$\sigma_w$ m s$^{-1}$ in this study; Hensen et al., 2009), $\beta_0$ is the coefficient when $w_0 = 0$, and $\beta_0$ was calculated based on the sensible heat flux:

$$\beta_0 = \frac{w' \cdot T}{\sigma_w \cdot (T \uparrow - T \downarrow)},$$

where $w' \cdot T$ is the sensible heat flux, and $T \uparrow$ and $T \downarrow$ represent temperatures when the fluctuating component of $w$ is directed upward and downward respectively. According to the theoretical principles of the REA method, $\beta_0$ is $\sim 0.6$ for a smooth surface. However, Gao (1995) found $\beta_0$ to decrease when measuring close to tall canopies, and Ren et al. (2011) experimentally determined $\beta_0$ to $0.42 \pm 0.02$. Therefore, if the calculated $\beta_0$ is less that 0.2 or larger than 0.6, then $\beta_0 = 0.4$ was used following the value found by Ren et al. (2011).

The measurement uncertainty of NH$_3$ concentration was estimated from the relative uncertainty, based on the mean value of three calibration campaigns conducted during the measurement period. For each calibration, concentration liquids of 0, 10 and 25 ppb were used, and the detection limit was estimated. A few measurements on 25 October were found to be lower than the estimated detection limit and excluded from further analysis. The reliability of the $F_{NH_3}$ measurements was assessed by comparing the atmospheric NH$_3$ concentration measurements from the REA system ($c_{RNH_3}$) with concentration measurements performed using diffusion denuders ($c_{DNH_3}$).

### 2.5 Flux partitioning

The contribution to the atmospheric NH$_3$ flux from the stomatal, cuticular and ground in the forest was assessed by simulating the forest component fluxes $F_s$, $F_w$ and $F_g$ (µg NH$_3$-N m$^{-2}$ s$^{-1}$) using a simple two-layer bi-directional canopy compensation point biosphere–atmosphere modelling approach (Nemitz et al., 2001). The model includes a NH$_3$ stomatal compensation point and allows NH$_3$ emissions from the ground layer caused by e.g. soil emissions or litter decomposition. The NH$_3$ canopy compensation point $\chi_c$ and the component fluxes are calculated as (Nemitz et al., 2001)

$$\chi_c = \frac{\chi_c - \chi_s}{R_s},$$

$$F_s = \frac{\chi_c - \chi_s}{R_s},$$

$$F_w = \frac{\chi_c - \chi_s}{R_w},$$

$$F_g = \frac{\chi_c - \chi_s}{R_g},$$

where $\chi_s$ and $\chi_g$ are the stomatal and ground compensation points (µg NH$_3$-N m$^{-2}$ s$^{-1}$), $R_s$, $R_b$, $R_c$, $R_w$ and $R_g$ are the aerodynamic, boundary layer, stomatal, cuticular, and ground resistances (s m$^{-1}$), respectively. The total forest flux $F_t$ (µg NH$_3$-N m$^{-2}$ s$^{-1}$) is calculated as the sum of the three component fluxes. The leaf physiological parameters such as apoplastic pH and NH$_3^+$ concentration are normally used to calculate the stomatal emission potential $\Gamma_s$ [NH$_3^+$/H$^+$] and the stomatal compensation point $\chi_s$, and for this purpose a value for $\Gamma_s$ of 250 found by Wang et al. (2011) in the late senescent period for the forest was used. The ground layer emission potential $\Gamma_g$ [NH$_3^+$/H$^+$] was fitted to $F_g$ (Eq. 7). It was found that, after leaf fall, a constant $\Gamma_g$ value of 80 000 reproduces the measured net flux reasonably well. It should be noted however that $R_g$ is calculated as the sum of an in-canopy resistance $R_{ac}$ and a ground boundary layer resistance $R_{bg}$ that are parameterised for agricultural crops and free-water surfaces, respectively. Uncertainties regarding these parameterisations therefore exist when applied to a forest ecosystem.

### 2.6 Denuder measurement

Atmospheric NH$_3$ concentration $c_{DNH_3}$ (µg NH$_3$-N m$^{-3}$) at 29.8 m height (Fig. 2) was sampled using seven 15 cm-long glass diffusion denuders (Fern, 1979) for the period 26 October to 11 November. The denuders were coated with a solution of citric acid in methanol which effectively captures NH$_3$. Air was pumped through the denuders at 300 mL min$^{-1}$. Chemical analyses of denuders were performed using ion chromatography. The uncertainty of measurements was estimated as the standard deviation of the
three denuders, and the detection limit was calculated as 3 times the standard deviation of the unexposed denuders (blank) to be 0.003 µg NH₃·N m⁻³, following the method used by Andersen et al. (1999).

2.7 Model calculations using DAMOS

The atmospheric NH₃ deposition and concentration (c NH₃) was modelled using the DAMOS system (Geels et al., 2012b). DAMOS is a combination of the regional-scale DEHM model (Brandt et al., 2012; Christensen, 1997) and the local-scale Gaussian dispersion/deposition model OML-DEP (Sommer et al., 2009). DEHM includes a detailed NH₃ emission model (Gyldenkærne et al., 2005; Skjøth et al., 2004, 2011) and provided in this study the wet NH₃ emission model (Gyldenkærne et al., 2005; Skjøth et al., 2004, 2011) and provided in this study the wet NH₃ deposition and upwind boundary conditions for the local-scale OML-DEP model. OML-DEP was used to simulate atmospheric NH₃ concentrations and deposition rates using a one-way dry deposition scheme (Simpson et al., 2003), a high spatial resolution (100 m) land cover (Nielsen et al., 2000) and local field NH₃ emissions (Gyldenkærne et al., 2005; Skjøth et al., 2004). Estimates of field emissions considered the annual animal production in each agricultural building, and the location of buildings and storage facilities were represented by their geographical coordinates (Fig. 4). Temporal changes of emissions due to variable meteorological conditions, particularly air temperature and length of local growing season, were also taken into account (Skjøth et al., 2011, 2004). OML-DEP provided hourly simulations throughout the measurement period for 40 × 40 receptor points in a 16 km × 16 km grid. The flux tower was located in the centre of the modelling domain. A full description of DAMOS can be found in Geels et al. (2012b) and Hertel et al. (2013).

3 Results

3.1 Meteorological and soil observations

The meteorological and soil conditions in the flux measurement period (21 October to 15 November 2010) are presented in Fig. 5. A series of frontal passages characterised the environmental conditions which caused winds to primarily originate from the west and southwest although interrupted by shorter periods of northerly winds between the 1/2 November and 8/9 November (Fig. 5a). The friction velocity varied mainly between 0 and 1 m s⁻¹; however, during the period 3–14 November, three episodes with u* larger than 1 m s⁻¹ were observed with wind speeds of up to 8 m s⁻¹ (Fig. 5b). The period with westerly winds was characterised by near-neutral situations, while the shorter periods with more northerly winds had episodes with either stable or unstable conditions (Fig. 5f). Observed temperatures remained in the range 5–14 °C until 7 November, after which temperatures decreased and stabilised at 2–5 °C during the rest of the period (Fig. 5d). The same pattern was seen in soil temperature, which decreased from 9 to 5 °C during the period (Fig. 5e). The period 21 October to 7 November had episodes with a few mm of rain almost every day (Fig. 5g), which led to high soil water content and high relative air humidity (Fig. 5i and h). It did not rain from 8 to 14 November.

3.2 High-resolution ammonia fluxes

Figure 6b shows the measured NH₃ fluxes. F NH₃ varied from −0.25 ± 0.30 µg NH₃·N m⁻² s⁻¹ in the beginning of the measurement period to 0.67 ± 0.28 µg NH₃·N m⁻² s⁻¹ in the end of the period. The mean flux was 0.06 ± 0.17 µg NH₃·N m⁻² s⁻¹ (Table 1). A clear tendency of F NH₃ going from negative (deposition) to positive (emission) fluxes was seen throughout the measurement period. Even though estimates were related to large uncertainties (0.006–0.344 µg NH₃·N m⁻² s⁻¹), the occurrence of emissions was convincing. Comparing LAI and F NH₃ during leaf fall showed a gradual increase of F NH₃ from negative to positive fluxes following defoliation (Fig. 6a and b). When defoliation ended (LAI reached zero) on 23 October, the deposition decreased towards zero (non-existent flux). In the leaf fall period (23 October to 8 November), F NH₃ gradually turned to positive fluxes, and a peak NH₃ emission of 0.50 µg NH₃·N m⁻² s⁻¹ was reached on 3 November. Following leaf fall, F NH₃ stayed positive and three emission peaks (8/9, 10 and 12 November) of up to 0.67 ± 0.28 µg NH₃·N m⁻² s⁻¹ were observed.
Fig. 4. NH$_3$ emissions (kg NH$_3$-N yr$^{-1}$) from point sources in 2008 (data are described in Sect. 2.6). The flux tower is marked with a white star in Lille Bøgeskov (the green scratched area). The concentration roses show measured concentration using REA (upper) and modelled concentration using DAMOS (lower) related to measured and simulated wind direction, respectively. Red colours indicate the highest assessed concentrations and blue indicate lowest concentrations.

Table 1. Statistics of the atmospheric ammonia (NH$_3$) flux measured $F_{NH_3}$ above Lille Bøgeskov using relaxed eddy accumulation in 33 m and simulated using the two-layer bi-directional model $F_t$ in the period 21 October to 15 November.

|        | Mean        | Min         | Max         | Stdev       |
|--------|-------------|-------------|-------------|-------------|
| $F_{NH_3}$ | 0.06        | -0.25      | 0.67        | 0.15        |
| $F_t$   | 0.07        | -0.03      | 0.47        | 0.11        |

3.3 Stomatal, cuticular and ground NH$_3$ fluxes

In Fig. 7 the forest component fluxes $F_s$, $F_w$, $F_g$ and the total flux $F_t$ simulated using the bi-directional NH$_3$ compensation point model are illustrated together with the measured flux $F_{NH_3}$. The dimensionless ratio $\Gamma_g$ [NH$_4^+$/H$^+$] was fitted to $F_g$ and set to 300 before 30 October and to 80 000 in the following period. A ratio of 80 000 for $F_g$ is extremely high, but still in the range of what has previously been found for fertilised grassland (Sutton et al., 2009). The bi-directional model was able to reproduce the temporal variations in the NH$_3$ emissions following 30 October but underestimated the deposition fluxes by on average 75 % (minimum = 6.27 % and maximum = 99.89 %) before 30 October. Following 30 October, the simulated ground flux comprises most of the total flux (142 %), whereas stomatal and cuticular fluxes represent only 0.9 % and 41 % respectively.

3.4 Observed ammonia concentrations

Atmospheric NH$_3$ concentrations measured by the REA system ($c_{RNH_3}$) varied from less than 0.03 to 2 µg NH$_3$-N m$^{-3}$ in the measurement period (Fig. 6b). The mean $c_{RNH_3}$ was 0.56 ± 0.35 µg NH$_3$-N m$^{-3}$, and the detection
limit was found to be 0.03 µg NH₃-N m⁻³. From 26 October to 11 November, the mean cRNH₃ was 0.68 ± 0.35 µg NH₃-N m⁻³, and the mean concentration obtained from denuder measurements (cRNH₃) was 0.67 ± 0.05 µg NH₃-N m⁻³ and above the detection limit (Table 2). A clear diurnal pattern was seen in cRNH₃ particularly in the beginning of the measurement period, which indicated increasing cRNH₃ at night and decreasing cRNH₃ during daytime (Fig. 6b). The concentration roses (Fig. 4) indicate that the dominating contribution of atmospheric NH₃ originated from the southwest, where many smaller point sources were located, but the highest concentrations were related to northwesterly wind directions, where two NH₃ point sources were located at the forest edge. Relatively low concentrations were seen when wind directions were from the northeast, where only few NH₃ emission sources were located (Fig. 4).

3.5 Simulated ammonia concentration using DAMOS

Mean three-hourly modelled atmospheric concentrations by DAMOS (cRNH₃) for the entire period (Fig. 6b) varied between 0.03 ± 0.015 and 2.51 ± 1.255 µg NH₃-N m⁻³, and mean cRNH₃ was 0.50 ± 0.25 µg NH₃-N m⁻³ (Table 2). It should be noted that the emission signal from the Danish area includes only point sources (stables and storage) as diffuse sources from agricultural areas are inactive in the period 1 October to 1 February due to Danish legislations on fertilisation management. This legislation on actual farming practice is dynamically incorporated in the NH₃ emission model. Apart from five to six peaks exceeding 1 µg NH₃-N m⁻³, the simulated level of cRNH₃ remained between 0 and 1 µg NH₃-N m⁻³, and no decreasing or increasing trends were observed for the period. The cRNH₃ peaks exceeding 1 µg NH₃-N m⁻³ (Fig. 6b) were related mainly to contributions from local point sources located in the southwestern sector (Fig. 4) and to difficulties of the model in handling low friction velocities and changes in atmospheric stability.

4 Discussion

Despite the different measurement heights of REA (33 m) and the denuders (29.8 m), the mean atmospheric NH₃ concentration measured for the period 26 October to 11 November 2010 are in good agreement, i.e. 0.68 ± 0.35 (REA) and 0.67 ± 0.05 (Denuder), and the simulated concentrations are also in a comparable range, i.e. 0.56 ± 0.28 µg NH₃-N (DAMOS) (Table 2). Even though we compare atmospheric measurements of NH₃ concentration from two different measurement systems installed at two different heights separated by 3.2 m (cRNH₃ in 33 m and cRNH₃ in 29.8 m), we expect that the measured concentrations are comparable due to the relatively high standard deviation on the concentrations measured by the REA system. Furthermore, the concentrations assessed are comparable with measurements for forests in other Danish and US studies (Andersen et al., 1999; Pryor et al., 2001). Many studies have reported that forests generally act as efficient sinks taking up the atmospheric NH₃ (i.e. Erisman and Wyers, 1993; Duyzer et al., 1994). In this study where leaf absorption of NH₃ is almost non-existent due to leaf fall, 57.7% of the measured fluxes indicated emissions and 19.5% indicated depositions. The atmospheric ammonia flux measurements show a temporal correlation with the canopy’s vegetation development (Fig. 6a and b), suggesting that FNH₃ decreases as LAI decreases, and that NH₃ emissions occur in the leaf fall period. Apart from decomposition of litter, also natural emissions of NH₃ linked to the leaf stomatal compensation point may cause NH₃ emissions to increase in the senescent period (Wang et al., 2011, 2013). However such processes are not yet taken into account in the DAMOS system, and thus impacts of forest NH₃ emissions are not represented in the cRNH₃ model calculations. The relations between the NH₃ fluxes and the vegetation status are discussed in the following along with the model performance.

4.1 LAI and NH₃ fluxes in the leaf fall period

Before 30 October, the measured cRNH₃ was less than 0.5 µg NH₃-N m⁻³, and during the leaf fall period
(23 October to 8 November) a clear increase in the $c_{RNH_3}$ was seen (Fig. 6b). After 30 October, the $c_{RNH_3}$ level was typically higher than the mean ($0.56 \pm 0.35 \mu g\text{ NH}_3\text{-N m}^{-3}$) for the measurement period. This increase was found to be related to the increasing forest NH$_3$ flux that was observed after 2 November (Fig. 6a). The modelled concentration $c_{mNH_3}$ did not show such an increase in the same period, indicating that a natural source, the forest, was causing the increase in $c_{RNH_3}$. As defoliation ended, $F_{NH_3}$ turned positive and changed from showing net deposition to net emissions of NH$_3$ following the leaf fall period. The increased emissions could be due to a combination of increased litter emissions and decreased leaf absorption, but also to changes in the turbulent transfer above the canopy related to the leaf fall. During and after the leaf all period, four evident NH$_3$ emission peaks (on 2–4, 8/9, 10/11 and 12–14 November) occurred (Fig. 6a), and clear relations to the friction velocity (Fig. 5b) are seen for all four peak emissions, indicating that the turbulent flow above the forest canopy controls a large part of the emission flux. Other than the friction velocity, no clear control pattern of other environmental or climatic conditions is seen that can explain the emission events. However, as precipitation occurred during most of the period (Fig. 5g), the emissions can also be caused by volatilisation of NH$_3$ from moist soil and wet leaves, as suggested by Pryor et al. (2001). The emissions found on 2–4 November and 12–14 November were related to precipitation events followed by dry periods with relatively high air temperatures. Such conditions enhance the decomposition process of plant material and may explain the emissions on these occasions. Such conditions were not present on 8/9 November and 10/11 November, when peak emissions were also measured. Studies based on dynamic chamber techniques and within-canopy profile measurements of the NH$_3$ flux have reported NH$_3$ emissions from intensively managed ecosystems, and suggest re-emission from senescent leaves and decomposition of leaf litter to be a strong source of NH$_3$ emissions, particularly in humid conditions (David et al., 2009; Nemitz et al., 2000a). The emission potential for senescent leaves or leaf litter of grassland was studied in the GRAMINAE Integrated Experiment (Sutton et al., 2009) and reported by David et al. (2009), who found that the emission potential was a hundred times larger than that of green leaves, and that emissions were larger when the litter was moisturer than when dry. Soil conditions such as temperature, moisture, pH, and nitrogen content have also been found to be controlling factors for the NH$_3$ emissions (Riedo et al., 2002; Roelle and Aneja, 2002; Walker et al., 2013), however, not as crucial as the leaf litter. Despite the low pH of the soils in Lille Bøgeskov, the conditions for decomposition are relatively good, and volatilisation of NH$_3$ due to the microbiological breakdown of organic material could occur. However, we are aware that the observed NH$_3$ emissions are measured during relatively low air and soil temperatures (below 10°C) that slow down the decomposition processes.

The forest component fluxes from the bi-directional NH$_3$ compensation point model indicated high emission fluxes from the ground layer following 30 October correlated to the atmospheric NH$_3$ emissions. The decreased deposition in the senescence period could also be caused by reduced leaf uptake of NH$_3$ through stomata, decreased cuticular

|                          | Mean (µg NH$_3$-N m$^{-3}$) | Min (µg NH$_3$-N m$^{-3}$) | Max (µg NH$_3$-N m$^{-3}$) | Stdev (µg NH$_3$-N m$^{-3}$) |
|--------------------------|-------------------------------|-----------------------------|-----------------------------|-------------------------------|
| $c_{RNH_3}$ (REA)        | 0.68                          | 0                           | 1.98                        | 0.40                          |
| $c_{dNH_3}$ (Denuder)    | 0.67                          | –                           | –                           | –                             |
| $c_{mNH_3}$ (DAMOS)      | 0.56                          | 0.03                         | 2.51                        | 0.45                          |

Table 2. Statistics of the atmospheric ammonia (NH$_3$) concentration conducted using the relaxed eddy accumulation (REA) measurement technique (33 m) ($c_{RNH_3}$), denuder measurements (29.8 m) ($c_{dNH_3}$) and the DAMOS model ($c_{mNH_3}$) for Lille Bøgeskov in the period 26 October to 11 November 2010.
desorption and larger NH$_3$ emission potential of the senescent leaves related to remobilisation of nitrogen during leaf senescence (Wang et al., 2011, 2013). The bi-directional model showed a slightly decreased cuticular flux $F_{\text{cu}}$ (Fig. 7) during senescence (21/30 October), however, no significant effect of stomatal control of the NH$_3$ flux. The model underestimated the deposition flux by on average 75% before 30 October and suggested cuticular deposition to be the controlling process, while the stomatal flux was nearly non-existent (emissions up to 0.003 µg NH$_3$·N·m$^{-2}$·s$^{-1}$). Assuming that the maximum possible flux permitted by turbulent transport can be calculated as $F_{\text{Max}} = -c_{\text{NH}_3}/R_u$, the measured deposition flux in the period 21–30 October is much larger than $F_{\text{Max}}$. However, the $F_{\text{Max}}$ parameterisation is based on the assumptions of horizontal and vertical homogeneity, and no chemical reactions must occur within the gradient. It has earlier been found that these assumptions were violated for NH$_3$ exchange between surfaces and the atmosphere (Sorensen et al., 2003; Duyzer et al., 1994). Furthermore, due to the uncertainty of the measurements in the period 21–30 October ($\pm$0.14 µg NH$_3$·N·m$^{-2}$·s$^{-1}$), the deposition flux measured could just as well be less than $F_{\text{Max}}$.

Following leaf fall, the magnitude and temporal structure of the measured NH$_3$ emission fluxes could be adequately reproduced with the bi-directional resistance model. The magnitude was achieved by using a $\Gamma_g$ value of 80 000 following leaf fall, which is in the range of values found for senescing plant material (e.g. Sutton et al., 2009), but larger than measurements of the litter emission potential established by Wang et al. (2011) during the previous autumn. In the absence of in-canopy turbulence measurements, the parameterisation of the in-canopy transport resistance is poorly constrained. Any scaling error in this resistance would result in a similar scaling error of $\Gamma_g$. Because a constant value of $\Gamma_g$ is used after leaf fall, the temporal variability in the model result is dominated by the variability in $u_*$, on which the parameterisation of in-canopy resistance is based. The results demonstrate that the emissions are consistent with a source inside the canopy, the connection of which to the atmosphere is regulated by turbulence. A minor fraction of the NH$_3$ emitted from the leaf litter at ground level is predicted to be re-captured by the surfaces of the tree canopy ($F_{\text{w}}$), and this fraction decreases as the PAI decreases. The model underestimates the net flux on 10 November and overestimates the flux on 13/14 November. The first period is at the end of a drying period and during very low turbulence when convective processes contribute to in-canopy transport that would be underestimated by a $u_*$-based parameterisation. Alternatively, the re-establishment of good agreement on 12 November follows a precipitation event, and this might indicate that a secondary effect of moisture on the mineralisation and thus emission rate further modulates the emission of the ground surface.

4.2 OML-DEP model results

The simulated NH$_3$ concentration level in the senescent period fitted well the measured concentrations, but the modelled concentrations were too low following leaf fall (Fig. 6b). On 10 November the measured $c_{\text{NH}_3}$ increased rapidly up to 1.98 µg NH$_3$·N·m$^{-3}$, caused by the forest NH$_3$ emission, while the modelled $c_{\text{NH}_3}$ showed two narrow peaks that were related to changes in the atmospheric stability from the meteorological input to the model. $c_{\text{NH}_3}$ was twice the magnitude of $c_{\text{NH}_3}$ in this period, indicating that the missing contribution to $c_{\text{NH}_3}$ from the forest is of a relatively large magnitude and therefore important to study and also include in model calculations.

Inadequate descriptions of surface properties such as aerodynamic roughness, stomatal resistance and processes related to the bi-directionality of atmospheric NH$_3$ fluxes represent uncertainties in current biosphere–atmosphere exchange modelling of NH$_3$ (Simpson et al., 2011). The complex mechanisms controlling these exchange processes cause current model parameterisations of biological and chemical pathways and processes to be empirical and based on few existing datasets (Flechard et al., 2011; Menut and Bessagnet, 2010; Pouliot et al., 2012). In DAMOS, the deposition is calculated for various land use categories and the surface resistance is divided into stomatal and non-stomatal components. For NH$_3$ the non-stomatal component includes, among other things, an acidity ratio between SO$_2$ and NH$_3$ (Emberston et al., 2000; Simpson et al., 2003), while a stomatal compensation point is not included. Many on-going studies explore methods to include these processes in models to calculate the NH$_3$ exchange between vegetative surfaces and the atmosphere (Wichink Kruit et al., 2012; Massad et al., 2010). Riedo et al. (2002) coupled a two-layer resistance model to a NH$_3$ exchange model to include ecosystem N dynamics for an intensively managed grassland. The model, PaSim, was able to simulate effects of cutting and fertilisation, but postulated that leaf litter was the reason for underestimation of NH$_3$ emission peaks. Other modelling concepts simulated leaf NH$_3$ emissions by including the stomatal compensation point and a litter layer with the emission potential, $\Gamma$, being dependent on the relative air humidity (Nemitz et al., 2000b). Bi-directional approaches for simulating ammonia emissions are being developed for several chemistry transport models (CTMs) such as the CMAQ, DEHM, EMEP, CHIMERE, and LOTUS-EUROS models (Aas et al., 2012; Skjøth et al., 2011; Cooter et al., 2012; Hamauou-Laguel et al., 2012). Here, calculations of the ammonia emissions as a function of ambient conditions are expected to improve calculations and understanding considerably (e.g. Hendriks et al., 2013). But to our knowledge, these methods do not include ammonia emissions from litter fall from forest. Further experiments conducting ground layer and canopy layer information of the stomatal and ground layer emission potential,
\( \Gamma \), are necessary to develop new parameterisations to forest–atmosphere exchange of \( \text{NH}_3 \).

Despite these efforts, we are still lacking knowledge of most biosphere–atmosphere exchange processes of N compounds (including potential feedback mechanisms) (Arneth et al., 2010). Flux studies in combination with model calculations are therefore needed in order to highlight the knowledge gaps and target future model improvements.

### 4.3 Diurnal dynamics in atmospheric ammonia concentration

Measured \( c_{\text{RNH}_3} \) indicated a clear diurnal pattern in the beginning of the measurement period with increased concentrations during night and decreased concentrations during day, showing that \( c_{\text{NH}_3} \) varied significantly over shorter timescales (Fig. 6b). Stable atmospheric conditions during nighttime suppress the mixing of air and reduce the atmospheric boundary layer height, thereby leading to higher \( \text{NH}_3 \) concentrations in the lower atmosphere. The model only captured these diurnal variations to a small extent (24–26 October) before defoliation ended, and it even showed anti-correlations for the three days 21–24 October. It is known that Gaussian models including OML have problems when meteorological conditions change from stable to unstable and low friction velocities prevail (Olesen et al., 2007). One of the main problems is that the Gaussian formulation assumes stationary conditions during the calculation period, which is not appropriate under certain conditions such as low wind speeds (see full discussion in Olesen et al., 2007). In the current study, the comparison with high-resolution measurements showed that the observed diurnal variation \( c_{\text{RNH}_3} \) is hard to capture by the model. Some of the conceptual limitations can be improved with more advanced parameterisations such as updating the description of horizontal dispersion in OML. Other limitations require the Gaussian OML to be replaced with more advanced models (Olesen et al., 2007). The lower performance of OML during these few and special meteorological conditions does not affect the main conclusions: that there is a need to represent \( \text{NH}_3 \) releases from natural ecosystems, such as forests during leaf fall, for accurate high spatial and high-temporal (diurnal) atmospheric simulation of \( c_{\text{NH}_3} \). Therefore, more data on \( \text{NH}_3 \) fluxes are necessary to improve our understanding of controlling parameters in the biosphere–atmosphere \( \text{NH}_3 \) exchange processes.

### 5 Conclusions

Measurements in a deciduous beech forest showed that \( F_{\text{NH}_3} \) changed from negative (deposition) to positive (emission) fluxes during the leaf fall period 21 October to 15 November 2010, causing increased atmospheric \( \text{NH}_3 \) concentration. This change was temporally correlated to the vegetation status of the forest. The observations of \( \text{LAI} \) and \( \text{PAI} \) showed that the period with increased \( c_{\text{RNH}_3} \) was in the leaf fall period, and peak emissions up to 0.67 \( \pm \) 0.28 \( \mu \text{g} \, \text{NH}_3 \cdot \text{N} \, \text{m}^{-2} \cdot \text{s}^{-1} \) were observed. The simple two-layer bi-directional canopy compensation model was able to adequately reproduce the magnitude and temporal structure of the measured \( \text{NH}_3 \) emission fluxes following leaf fall and indicated that the forest ground layer (soil and litter) acted as the main contributing component to the \( \text{NH}_3 \) emissions. The mean \( \text{NH}_3 \) concentrations were well simulated using DAMOS before leaf fall, but were underestimated following leaf fall. This points to the need for representing forest leaf fall and associated \( \text{NH}_3 \) emissions in chemical transport models when simulating nitrogen deposition to forests. Besides influence on the atmosphere–forest exchange of \( \text{NH}_3 \) from the forest’s phenology, variations in meteorological and soil conditions, and the canopy turbulence, our observations support the hypothesis that \( \text{NH}_3 \) emissions occur from deciduous forests in relation to leaf fall possibly due to increased litter emissions or decreased leaf absorption or a combination of these. Additionally, diurnal variations of \( c_{\text{RNH}_3} \) related to meteorological conditions (i.e. radiation control on stomatal resistance), forest phenology (i.e. LAI), and the spatial distribution of local anthropogenic \( \text{NH}_3 \) sources were found. This suggests that dedicated process studies including manipulation of ecosystems would be very valuable for improved understanding of \( \text{NH}_3 \) fluxes from natural vegetation. Despite large uncertainties associated with the results obtained in this study due to the assessment techniques and the limited dataset, the good agreement between \( c_{\text{RNH}_3} \), \( c_{\text{RNH}_3} \), and \( c_{\text{RNH}_3} \) gives confidence in the data. The results must be considered as a good contribution to improve our understanding of the processes related to natural \( \text{NH}_3 \) emissions. This knowledge can be used in direct ecosystem manipulation studies or model studies in order to quantify the \( \text{NH}_3 \) emission flux from ecosystems and the total biosphere–atmosphere net flux of reactive nitrogen.

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