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Dry and wet deposition of inorganic nitrogen compounds to a tropical pasture site (Rondônia, Brazil)

I. Trebs¹, L. L. Lara², L. M. M. Zeri³, L. V. Gatti⁴, P. Artaxo⁵, R. Dlugi⁶, J. Slanina⁷, M. O. Andreae¹, and F. X. Meixner¹

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Abstract. The input of nitrogen (N) to ecosystems has increased dramatically over the past decades. While total (wet + dry) N deposition has been extensively determined in temperate regions, only very few data sets of N wet deposition exist for tropical ecosystems, and moreover, reliable experimental information about N dry deposition in tropical environments is lacking. In this study we estimate dry and wet deposition of inorganic N for a remote pasture site in the Amazon Basin based on in-situ measurements. The measurements covered the late dry (biomass burning) season, a transition period and the onset of the wet season (clean conditions) (12 September to 14 November 2002) and were a part of the LBA-SMOCC (Large-Scale Biosphere-Atmosphere Experiment in Amazonia – Smoke, Aerosols, Clouds, Rainfall, and Climate) 2002 campaign. Ammonia (NH₃), nitric acid (HNO₃), nitrous acid (HONO), nitrogen dioxide (NO₂), nitric oxide (NO), ozone (O₃), aerosol ammonium (NH₄⁺) and aerosol nitrate (NO₃) were measured in real-time, accompanied by simultaneous meteorological measurements. Dry deposition fluxes of NO2 and HNO3 are inferred using the "big leaf multiple resistance approach" and particle deposition fluxes are derived using an established empirical parameterization. Bi-directional surface-atmosphere exchange fluxes of NH₃ and HONO are estimated by applying a "canopy compensation point model". N dry and wet deposition is dominated by NH₃ and NH₄⁺, which is largely the consequence of biomass burning during the dry season. The

Correspondence to: I. Trebs (ivonne@mpch-mainz.mpg.de)

grass surface appeared to have a strong potential for daytime NH $_3$ emission, owing to high canopy compensation points, which are related to high surface temperatures and to direct NH $_3$ emissions from cattle excreta. NO $_2$ also significantly accounted for N dry deposition, whereas HNO $_3$, HONO and N-containing aerosol species were only minor contributors. Ignoring NH $_3$ emission from the vegetation surface, the annual net N deposition rate is estimated to be about $-11\,\mathrm{kgN}$ ha $^{-1}\,\mathrm{yr}^{-1}$. If on the other hand, surface-atmosphere exchange of NH $_3$ is considered to be bi-directional, the annual net N budget at the pasture site is estimated to range from -2.15 to $-4.25\,\mathrm{kgN}$ ha $^{-1}\,\mathrm{yr}^{-1}$.

1 Introduction

The supply of reactive nitrogen (N) to global terrestrial ecosystems has doubled since the 1960s as a consequence of human activities, such as fertilizer application, cultivation of N fixing legumes and production of nitrogen oxides by fossil-fuel burning (Galloway, 1998). The deposition of atmospheric N species constitutes a major nutrient input to the biosphere. On a long-term scale, the increase of N inputs into terrestrial ecosystems may result in (i.) intensified trace gas exchange (ii.) enhanced leaching of nitrate and soil nutrients (e.g., K⁺, Mg²⁺, Ca²⁺), (iii.) ecosystem eutrophication and acidification, (iv.) reduction in biodiversity, and (v.) increased carbon storage (Vitousek et al., 1997). Enhanced carbon storage due to N deposition has been shown to increase the terrestrial carbon sink in N-limited temperate

¹Max Planck Institute for Chemistry, Biogeochemistry Department, P.O. Box 3060, 55020, Mainz, Germany

²Centro de Energia Nuclear na Agricultura (CENA), Laboratorio de Ecologia Isotópica, Universidade de São Paulo (USP), Av. Centenario, 303 13400-970, Piracicaba, São Paulo, SP, Brazil

³Max Planck Institute for Biogeochemistry, Department Biogeochemical Processes, Hans-Knöll-Straße 10, 07745 Jena, Germany

⁴Instituto de Pesquisas Energéticas e Nucleares, CQMA, Atmospheric Chemistry Laboratory, Av. Prof. Lineu Prestes, 2242, Cidade Universitaria, CEP 055508-900, São Paulo, SP, Brazil

⁵Instituto de Física, Universidade de São Paulo (USP), Rua do Matão, Travessa R, 187, CEP 05508-900, São Paulo, SP, Brazil

⁶Working Group Atmospheric Processes (WAP), Gernotstrasse 11, 80804 Munich, Germany

⁷Peking University, College of Environmental Sciences, Beijing 100871, China

ecosystems, which may have substantial impacts on global CO₂ concentrations (e.g., Townsend et al., 1996).

N deposition is considered to be relevant in the tropics due to widespread biomass burning activity and increasing fertilizer application. It was suggested by e.g., Matson et al. (1999) and Asner et al. (2001) that in contrast to temperate ecosystems, nitrogen-rich/phosphorus (P)-limited tropical rainforest soils may have a reduced productivity following excess N deposition, resulting in a decreased C-Moreover, the humid tropical zone is a major source area for biogenic nitrous oxide (N2O) and nitric oxide (NO) emissions from soils (Reiners et al., 2002). Enhanced N inputs to tropical forests are likely to increase nitrification/denitrification rates and, hence, the emission of NO and N₂O to the atmosphere (Hall and Matson, 1999). The conversion of tropical rainforest into cultivated land and pasture may lead to a sustained disturbance of the natural N cycle. During clearing and burning of tropical rainforest, biomassassociated N is volatilized and a large fraction is emitted in form of gaseous NH₃ (Trebs et al., 2004), which may result in considerable N losses of tropical ecosystems (Kauffman et al., 1998; Kauffman et al., 1995). This is affirmed by the fact that, in contrast to old growth forests, plant growth in deforested areas is suggested to be limited by N rather than by P (Davidson et al., 2004; Oliveira et al., 2001).

The deposition of atmospheric N compounds occurs via dry and wet processes. Nitrogen dioxide (NO₂), ammonia (NH₃), nitric acid (HNO₃) and nitrous acid (HONO) are the most important contributors to N dry deposition. HNO₃ usually features a rapid downward (net deposition) flux to the surface (Huebert and Robert, 1985). By contrast, the exchange of NO, NH₃, HONO and NO₂ between surface and atmosphere may be bi-directional. The rates of production and consumption in vegetation elements and/or soils as well as the ambient concentration determine whether net emission or net deposition of these species takes place. Turbulent diffusion controls the transport of gases and particles from the surface layer to the Earth's surface. The uptake of trace gases by surfaces is considered to be dependent on physico-chemical and biological surface properties (Hicks et al., 1987), but also on the solubility and reactivity of the gaseous compound (Wesely, 1989). Hence, soil characteristics, plant stomatal activity and trace gas chemical properties largely determine the deposition velocity. The atmospheric dry removal of aerosol particles, which may contain N species such as ammonium (NH_4^+) and nitrate (NO_3^-) , is a function of the particle size (Nicholson, 1988) but also depends on the particle density (e.g., Seinfeld and Pandis, 1998). Dry deposition is enhanced for large particles (especially those larger than a few micrometers) due to the additional influence of gravitational settling.

N wet deposition is a result of in-cloud scavenging ("rain-out") and below-cloud scavenging ("washout") of atmospheric N constituents (Meixner, 1994). The total (wet + dry) N deposition ranges from -1 to $-2 \, \text{kgN ha}^{-1} \, \text{yr}^{-1}$ for

rural locations (e.g., North Canada) up to -30 to -70 kgN ha⁻¹ yr⁻¹ for regions that receive N from urban pollution and/or agricultural activities (e.g., North Sea, NW Europe and NE U.S.) (Howarth et al., 1996). In contrast to moderately fluctuating air pollution levels that prevail in Europe, the United States and Asia throughout the year, tropical environments such as the Amazon Basin experience every year a dramatic change from the "green ocean" clean background atmosphere to extremely polluted conditions during the biomass burning season. Only few studies exist were atmospheric N wet removal was determined experimentally in the tropics (Clark et al., 1998; Galloway et al., 1982; Likens et al., 1987; Srivastava and Ambasht, 1994). The chemical composition of precipitation in the Amazon region was determined in previous studies by Andreae et al. (1990), Stallard and Edmond (1981), Lesack and Melack (1991) and Williams et al. (1997). Galy-Lacaux et al. (2003) have estimated N wet + dry deposition for different (tropical) African ecosystems. However, their dry deposition measurements are based on the application of integrating filter methods which are prone to artifacts (cf. Slanina et al., 2001). Since 1999, N wet + dry deposition are monitored on the Caribbean Virgin islands (St. John Island; see http://www.epa.gov/castnet///sites/vii423.html). Global chemistry and transport models (CTMs) such as MOGUN-TIA have been applied to estimate total N deposition on a global scale (e.g., Holland et al., 1999). Model results suggest that net N deposition in the northern temperate latitudes exceeds contemporary tropical N deposition by almost a factor of two. However, reliable experimental information about N dry deposition in tropical environments, required to validate these model predictions, has been lacking.

Kirkman et al. (2002) determined the surface-atmosphere exchange of NO_x ($NO + NO_2$) and ozone (O_3) at a pasture site in the Amazon Basin (Rondônia, Brazil). In this paper, we complement their study by additionally estimating the surface-atmosphere exchange fluxes of NH_3 , HNO_3 , HONO, aerosol NO_3^- and NH_4^+ at the same pasture site. Our analysis is based on real-time measurements, supported by simultaneous measurements of meteorological quantities covering the late dry (biomass burning) season, the transition period, and the onset of the wet season (clean conditions). Fluxes of NO_2 , HNO_3 , NH_3 and HONO are estimated by inferential methods. N wet deposition was determined by collection of rainwater and subsequent analyses. We estimate the total (wet + dry) annual N deposition at this pasture site and the relative contribution of the individual N species.

2 Experimental

2.1 Field site

Measurements were performed during 12-23 September 2002 (dry season, biomass burning), 7-31 October 2002

(transition period) and 1-14 November 2002 (wet season, clean conditions) as part of the LBA-SMOCC (Large-Scale Biosphere-Atmosphere Experiment in Amazonia – Smoke, Aerosols, Clouds, Rainfall, and Climate) 2002 campaign (Andreae et al., 2004) at a pasture site in the state of Rondônia, Brazil (Fazenda Nossa Senhora Aparecida, FNS, 10°04.70′ S, 61°56.02′ W, 145 m a.s.l.). The site is located in the south-western part of the Amazon Basin. The location and a simplified sketch of the measurement site are shown in Fig. 1. The old growth rain forest at FNS was cleared by slash and burn activities in 1977. The vegetation at FNS is dominated by C4 grass species (*Brachiaria brizantha*) with small patches of Brachiaria humidicola and very few isolated palms and bushes, and the site is used as a cattle ranch (~200 "Blanco" cattle, Bos indicus hybrid). The stocking rate at FNS was about one animal per hectare when field measurements took place. The pasture does not receive any fertilizer and is not harvested. FNS is located within a strip of cleared land about 4 km wide and several tens of kilometers long (Culf et al., 1996). The towns Ouro Preto do Oeste (\sim 40 800 inhabitants) and Ji-Paraná (\sim 110 000 inhabitants) are situated approximately 8 km and 40 km to the ENE and ESE of the site, respectively.

The instrumentation for trace gas/aerosol sampling and online analyses was arranged in an air conditioned wooden house. Rain samples were collected nearby the house (see Fig. 1). An automatic weather station ($Met\ 1$) was located in a distance of $\sim\!20\,\mathrm{m}$ to the S, and a meteorological tower ($Met\ 2$) was situated $\sim\!200\,\mathrm{m}$ to the NE of the inlets for trace gas and aerosol measurements. While the sampling site provides a sufficient uniform fetch expanding for 1–2 km from the sampling location in each direction (Andreae et al., 2002), local flow distortions may be caused by the wooden house and some instrument shelters. A more detailed description of the measurement site is given in Andreae et al. (2002) and Kirkman et al. (2002).

2.2 Sampling and analysis

Table 1 summarizes the specifications of the instruments for the measurement of trace gases, aerosol species and meteorological quantities. Water-soluble N containing trace gases (NH₃, HNO₃ and HONO) and related aerosol species (NH₄⁺ and NO₂) were measured on-line. Air was taken from a height of 5.3 m above ground through a sophisticated inlet system, which was designed to reduce wall losses of soluble gases (especially HNO₃) and to minimize aerosol losses due to non-isokinetic sampling (see Trebs et al., 2004). Soluble gases were scavenged with a wet-annular denuder (WAD) (Wyers et al., 1993), which was combined with a Steam-Jet-Aerosol Collector (SJAC) (Khlystov et al., 1995) to collect particulate N species. For both gaseous and aerosol compounds, sample collection was followed by subsequent online analysis (ion chromatography (IC)) for anions and flowinjection analyses (FIA) for NH₄⁺. Cycle times were set to

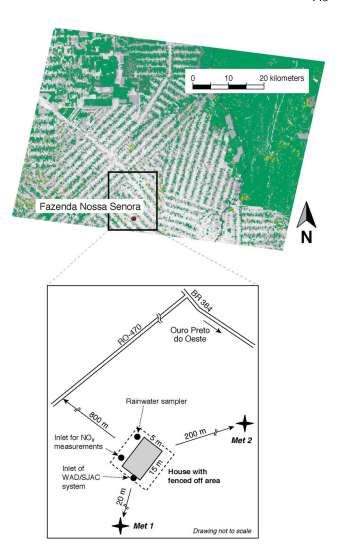


Fig. 1. Location of the LBA-SMOCC measurement site *Fazenda Nossa Senhora Aparecida* (FNS) in Rondônia, Brazil.

20 min (dry season), 40 min (transition period) and 60 min (wet season) (Trebs et al., 2004). Aerosol samples of either $PM_{2.5}$ ($D_p \le 2.5 \,\mu m$) or total suspended particulate matter (TSP) were collected. A detailed description and verification of the measurement method and of the inlet system can be found in Slanina et al. (2001) and Trebs et al. (2004).

The chemiluminescence NO/NO_x analyzer (Thermo Environment Instruments) (see Table 1) was equipped with a molybdenum converter to transform ambient NO_2 to NO. However, the converter basically responds to the sum of $NO_2 + HNO_3 + HONO + PAN + aerosol <math>NO_3^- + organic$ nitrates. Therefore, it is likely that NO_2 measurements might be biased by a positive artifact (Fehsenfeld et al., 1990). During our study, the inlet line for NO/NO_x measurements had a length of 25 m (inner diameter = 4.4 mm). Therefore, highly soluble and sticky species such as HNO_3 and HONO are assumed to be at least partly removed within the long inlet

Table 1. Specifications of the instrumentation for the measurement of trace gases, aerosol species and meteorological quantities at FNS during LBA-SMOCC 2002 (all heights are above ground).

Parameter/position/ height	Time resolution	Technique or sensor	Model, manufacturer	Detection limit/ precision	
NH ₃ , HNO ₃ , HONO (5.3 m)	20–60 min ⁽¹⁾	Wet-annular denuder (WAD), IC, FIA ⁽²⁾	ECN, Petten, Netherlands	\leq 0.015 ppb (3 σ) for acids, \leq 0.118 ppb (3 σ) for NH ₃	
Aerosol (NH $_4^+$, NO $_3^-$) (5.3 m)	20–60 min	Steam-Jet Aerosol Collector (SJAC), IC, FIA	ECN, Petten, Netherlands	\leq 0.015 ppb (3 σ) for anions, \leq 0.118 ppb (3 σ) for aerosol NH ₄ ⁺	
NO concentration (10 m)	5 min	Gas-phase chemilumi- nescense	Model 42C TL (trace level),Thermo Environment Instruments Inc., USA	0.05±0.025 ppb	
NO ₂ concentration (10 m)	5 min	Catalytic conversion of NO ₂ to NO by molybdenum converter (at 325°C), gas phase chemiluminescence	Model 42C TL (trace level), Thermo Environment Instru- ments Inc., USA	0.05±0.025 ppb	
O_3 concentration $(10 m)$	5 min	UV absorption	Model 49C Thermo Environment Instruments Inc., USA	1±0.5 ppb	
Air temperature (Met 1, 0.5 m and 5 m)	1 min	Pt-100 Ω resistance sensor	MP-103A-CG030-W4W Rotronic, Switzerland	±0.1 K	
Relative humidity (Met 1, 0.5 m and 5 m)	1 min	Capacitive sensor	MP-103A-CG030-W4W Rotronic, Switzerland	±1.5%	
Surface wetness (Met 1)	1 min	Surface wetness grids at soil surface	237 WSG, Campbell Scientific Ltd., UK	_	
Global radiation flux $(Met\ 1,\ 5\ m)$	1 min	Pyranometer sensor	LI200SZ (LI-COR, Lincoln, Nebraska, USA)	<±3%	
Eddy covariance; three dimensional wind and temperature fluctuations $(Met 2, 4m)$	10 Hz	3-D ultrasonic anemometer	Solent 1012R2, Gill Instruments, UK	±5–20%	
H_2O mixing ratio (Met 2, 4 m)	10 Hz	Infrared closed-path absorption	IRGA LI-COR 6262 (LI-COR, Lincoln, Nebraska, USA)	±1%	
Shortwave radiation in and out (Met 2, 8.5 m)	1 min	Pyranometer sensor	Kipp & Zonen Pyranometer CM 21	±2%	
Longwave radiation in and out (Met 2, 8.5 m)	1 min	Pyrgeometer sensor	Kipp & Zonen Pyrgeometer CG	±3%	

^{(1) 20} min: dry season (12–13 September), 40 min: transition period (7–31 October), 60 min: wet season (1–14 November)

tubing. In addition, mixing ratios of HNO₃ and HONO were usually below 0.5 ppb (Trebs et al., 2004), indicating that interferences would be marginal in case any of these gases would reach the chemiluminescence analyzer. However, elevated emissions of isoprene in the area may result in considerable concentrations of PAN such that we may not rule out interferences due to this compound. Aerosol NO₃⁻ was eliminated by the application of an inlet filter.

Also listed in Table 1 are those meteorological sensors that were used to measure the quantities involved in this study,

namely air temperature (*T*), relative humidity (*RH*), surface wetness, global radiation flux, momentum, latent and sensible heat flux and ingoing/outgoing short- and longwave radiation. Eddy covariance measurements were conducted using a Gill 3D-sonic anemometer. The H₂O mixing ratio was monitored by a fast LI-COR infrared gas analyzer, and its analog output was directly fed to the ultrasonic anemometer A/D converter. Post-processing of the eddy covariance data (EDDYWSC, software by Alterra, Wageningen University Research, Netherlands) resulted in 30 min averages

⁽²⁾ IC: ion chromatography, FIA: flow injection analysis

of sensible heat flux, latent heat flux, friction velocity and Monin-Obukov length. More details on the eddy covariance measurements and corresponding data evaluation/calibration procedures are given in Araujo et al. (2002).

Precipitation was sampled from 12 September to 14 November 2002 using a wet-only rainwater collector (Aerochem Metrics). A total of 23 rainstorm events were collected representing $\sim 100\%$ of the precipitation in this period. Rain samples were stored in the dark at 4°C using polyethylene bottles which were previously cleaned with deionized water and preserved with Thymol. In order to trace possible contaminations, the sample pH was measured directly after sampling and before analysis. Analyses of NH_4^+ , NO_3^- and NO_2^- were performed for all samples using a Dionex DX600 ion chromatograph at the Laboratório de Ecologia Isotópica, CENA/USP (São Paulo, Brazil). The detection limit was 0.05 μ M for all species. More details about sampling and analysis procedures are provided by Lara et al. (2001).

Moreover, a twin Differential Mobility Particle Sizer (DMPS) was employed to measure the dry aerosol particle size distribution in the diameter range from 3 to 850 nm (cf. Rissler et al., 2004). The size distribution of particles with aerodynamic diameters from 1 to $4 \mu m$ was measured with an Aerodynamic Particle Sizer (TSI APS 3310).

3 Theory: Estimation of N dry and wet deposition

3.1 Trace gas fluxes

Dry deposition fluxes of trace gases have been estimated using the inferential method, which is based on the "big leaf multiple resistance approach" (Wesely and Hicks, 1977; Hicks et al., 1987). The deposition flux (F) (μ g m⁻² s⁻¹) of a nonreactive trace gas for which the surface is a sink under all ambient conditions is defined by:

$$F = -V_d \cdot X(z_{\text{ref}}) = -\frac{X(z_{\text{ref}})}{R_a + R_b + R_c} \tag{1}$$

where $X(z_{\text{ref}})$ is the trace gas concentration (μ g m⁻³) at the reference height z_{ref} (m) and V_d denotes the dry deposition velocity (m s⁻¹), which is the reciprocal of the sum of the turbulent resistance (R_a) (s m⁻¹), the quasi-laminar or viscous boundary layer resistance (R_b) (s m⁻¹), and the surface resistance (R_c) (s m⁻¹). According to Hicks et al. (1987) R_a between the reference height (z_{ref}) and the roughness length z_0 (m) is given by:

$$R_a = \frac{1}{\kappa \times u_*} \left[\ln \left(\frac{z_{\text{ref}}}{z_0} \right) - \Psi_{H_{\left(\frac{z_{\text{ref}}}{L} \right)}} \right] \tag{2}$$

where κ denotes the von Karman constant (0.41) and L is the Monin-Obukov length (m), a measure of atmospheric stability that is derived from the sensible heat flux and the friction velocity u_* (Garratt, 1992). $\Psi_H(z_{\text{ref}}/L)$ is the stability correction function for heat and inert tracers in its integral

form (see Thom, 1975). The roughness length z_0 of the grass surface at the FNS site was taken as 0.11 m (cf. Kirkman et al., 2002) and $z_{\rm ref}$ was 5.3 m and 10 m for the WAD/SJAC and for the NO_x measurements, respectively (Table 1). To account for conditions when the reliability of micrometeorological techniques was low, data were rejected for $u_* \le 0.01$ m s⁻¹ and $z_{\rm ref}/L \ge 5$, i.e. when extremely low turbulence and/or very high thermal stability was prevailing. Also, data were rejected for $z_{\rm ref}/L \le -5$, which reflects cases of very high thermal turbulence production (when Monin-Obukov similarity is no longer valid (Ammann, 1999). Thus, about 10% of the dataset were not used for the flux calculations.

 R_b determines the exchange of gaseous matter by molecular-turbulent diffusion across the viscous laminar sublayer immediately above the vegetation elements and can be described by (Hicks et al., 1987):

$$R_b = \frac{2}{\kappa \times u_*} \left(\frac{Sc}{Pr}\right)^{\frac{2}{3}} \tag{3}$$

where Sc and Pr are the Schmidt and Prandtl number, respectively. Pr is 0.72 and Sc is a strong function of the molecular diffusivity of the trace gas. Values for Sc were taken from Hicks et al. (1987) and Erisman et al. (1994) for the different trace gas species. The surface resistances R_c could not be directly determined from our field measurements; hence values were adopted from the literature (see Sect. 4.4).

The inferential method is valid for trace gases whose mixing ratio just above the soil and/or vegetation elements is zero. The observation of a net NO_2 deposition flux to the FNS pasture by Kirkman et al. (2002) justifies the application of the inferential model for NO_2 in our study. This is also valid for HNO_3 , which typically features a rapid downward flux with negligible R_c and corresponding high V_d (Hanson and Lindberg, 1991).

By contrast, NO, HONO and NH₃ may be both deposited to and emitted from surfaces. Formally, this can be accounted for by a so-called canopy compensation point concentration $X_c(\mu g \text{ m}^{-3})$ that generally refers to the concentration of the compound just above the soil and/or vegetation elements (Nemitz et al., 2004a). X_c represents a concentration analogue of R_c and is the air concentration at which competing chemical and biological consumption and production processes balance each other (i.e., the net flux is zero) (see Sutton et al., 1995):

$$F = \frac{X_c - X(z_{\text{ref}})}{R_a + R_b} \tag{4}$$

The net NO emission from the FNS pasture site determined by Kirkman et al. (2002) was very low (0.65 ngN m $^{-2}$ s $^{-1}$ or 0.17 kgN ha $^{-1}$ yr $^{-1}$), thus we neglected any contribution of NO to the surface-atmosphere exchange of N species in our study. HONO is generally assumed to be formed by heterogeneous reaction of NO₂ with surface water (Harrison et al., 1996) and it may subsequently be emitted from

plant foliar cuticles or soil surfaces. Since there is no indication for any direct HONO emissions by plants (Schimang et al., 2006), the HONO compensation point concentration X_c (HONO) is expected to be a function of the NO₂ mixing ratio (see Sect. 4.4).

To predict the bi-directional surface-atmosphere exchange of NH₃ at the FNS site, we applied a dynamic resistance model proposed by Sutton et al. (1998). Besides uptake and emission of NH₃ via plant stomata, the dynamic model accounts for absorption of NH₃ by epicuticular water films under very humid conditions, and subsequent re-evaporation (capacitive leaf surface exchange). Since the FNS site is used as a cattle ranch and the NH3 flux directly from the soil is assumed to be negligible compared to that originating from cattle excreta, we considered a direct NH3 flux from cattle manure and urine $F(NH_3)_e$. The net NH₃ flux $F(NH_3)$ can be related directly to the NH₃ canopy compensation point concentration $X_c(NH_3)$ (Sutton et al., 1998) and is composed of its component fluxes through plant stomata, $F_s(NH_3)$, the flux in or out of the epicuticular water film (adsorption capacitor), $F_d(NH_3)$, and $F(NH_3)_e$:

$$F(NH_{3})_{t} = \underbrace{\frac{X_{s}(NH_{3}) - X_{c}(NH_{3})}{R_{s}(NH_{3})}}_{F_{s}(NH_{3})} + \underbrace{\frac{X_{d}(NH_{3})_{t} - X_{c}(NH_{3})}{R_{d}(NH_{3})}}_{F_{d}(NH_{3})} + F(NH_{3})_{e}$$

$$= \underbrace{\frac{X_{c}(NH_{3}) - X(NH_{3}, z_{ref})}{R_{d} + R_{b}}}$$
(5)

where $X_s(NH_3)$ denotes the NH₃ stomatal compensation point concentration ($\mu g \, m^{-3}$) and $R_s(NH_3)$ denotes the NH₃ stomatal resistance (s m⁻¹). $X_d(NH_3)_t$ is the NH₃ adsorption concentration ($\mu g \, m^{-3}$) associated with the "leaf surface capacitor" at time step t and $R_d(NH_3)$ is the charging resistance of the capacitor (s m⁻¹) (see Sutton et al., 1998). $X_c(NH_3)$ is then determined by Sutton et al. (1998):

$$X_{c}(\text{NH}_{3})_{t} = \frac{X(\text{NH}_{3}, z_{\text{ref}})/(R_{a} + R_{b}) + X_{s}(\text{NH}_{3})/R_{s}(\text{NH}_{3}) + X_{d}(\text{NH}_{3})_{t}/R_{d}(\text{NH}_{3}) + F(\text{NH}_{3})_{e}}{(R_{a} + R_{b})^{-1} + R_{s}(\text{NH}_{3})^{-1} + R_{d}(\text{NH}_{3})^{-1}}$$
(6)

 $X_s(NH_3)$ can be parameterized according to (Farquhar et al., 1980; Sutton et al., 1994):

$$X_s(\text{NH}_3) = \frac{161512}{T_S} \cdot 10^{(-4507.11/T_S)} \cdot \Gamma \cdot 17000 \tag{7}$$

where T_S is the surface temperature (K) which was derived from the outgoing longwave radiation by applying the Stefan-Boltzmann law. Γ is the ratio of apoplastic $[NH_4^+]/[H^+]$ and was adopted from the literature (Sect. 4.4). $R_s(NH_3)$ can be calculated from the measured latent heat flux for relatively dry daytime conditions and in the absence of precipitation according to Nemitz et al. (2004a) (see

Sect. 4.4). Furthermore, $X_d(NH_3)$ at time step t is a function of the adsorption charge and of the capacitance of the epicuticular water film (for details see Sutton et al., 1998). The values for the adsorption charge were adopted from Sutton et al. (1998) and the capacitance is a function of the epicuticular pH (see Sect. 4.4)

3.2 Aerosol fluxes

Up to date, no well established bulk resistance models exist for the dry deposition of particles. Significant discrepancies have been observed between experimental results and model predictions (Ruijgrok et al., 1995). The theoretical framework proposed by Slinn (1982) is widely used in modeling studies to predict particle deposition velocities. However, Wesely et al. (1985) derived an empirical parameterization for the dry deposition velocity V_p (m s⁻¹) of submicron sulfate aerosols (D_p =0.1–1.0 μ m) to grass surfaces:

$$V_p = u_* \cdot 0.002$$
, for $L \ge 0$ (8a)

$$V_p = u_* \cdot 0.002 \cdot \left[1 + \left(\frac{-a}{L} \right)^{\frac{2}{3}} \right], \text{ for } L < 0$$
 (8b)

where $a=300 \,\mathrm{m}$. This approach generally results in much higher V_p values for submicron particles than predicted by the Slinn model. Several other studies (e.g., Garland, 2001; Nemitz et al., 2004b; Vong et al., 2004) also showed that V_p to surfaces of low aerodynamic roughness may be much larger than predicted by the Slinn model, even for particles of different chemical composition than studied by Wesely et al. (1985). In this study either PM_{2.5} or TSP was sampled. NH_4^+ is known to be largely attributed to fine mode aerosols with $D_p \le 1.0 \,\mu\text{m}$ (Seinfeld and Pandis, 1998), which was also observed during the SMOCC measurement campaign (Fuzzi et al., 2005¹). By contrast, aerosol NO₃⁻ exhibited a bimodal size distribution (Falkovich et al., 2005). Since we did not find an empirical relationship in the literature to estimate V_p for aerosols with $D_p \ge 1.0 \,\mu\text{m}$ and the contribution of coarse mode aerosol NO₃⁻ relative to that of fine mode aerosol NH₄⁺ is presumably small (cf. Sect. 4.2, Table 2), the parameterization by Wesely et al. (1985) is considered as a reasonable approximation for both PM_{2.5} and TSP samples.

¹Fuzzi, S., Decesari, S., Facchini, M. C., Cavalli, F., Emblico, L., Mircea, M., Andreae, M. O. Trebs, I., Hoffer, A., Guyon, P., Artaxo, P., Rizzo, L. V., Lara, L. L., Pauliquevis, T., Maenhaut, W., Raes, N., Chi, X., Mayol-Bracero, O. L., Soto, L., Claeys, M., Kourtchev, I., Rissler, J., Swietlicki, E., Tagliavini, E., Schkolnik, G., Falkovich, A. H., Rudich, Y., Fisch, G., and Gatti, L. V.: Overview of the inorganic and organic composition of size-segregated aerosol in Rondônia, Brazil, from the biomass burning period to the onset of the wet season, J. Geophys. Res., submitted, 2005.

Table 2. Summary of trace gas and aerosol mixing ratios $^{(*)}$ during the dry season (12–13 September), the transition period (7–31 October) and the wet season (1–14 November) at FNS during LBA-SMOCC 2002 (conversion factors from ppb to μ g m⁻³ for standard conditions of 298.15 K and 1000 hPa: NO: 1.21, NO₂: 1.86, O₃: 1.94, NH₃: 0.69, HNO₃: 2.54, HONO: 1.89, aerosol NH₄⁺: 0.73, aerosol NO₃⁻: 2.77).

		Dry se	eason			Transitio	n period			Wet se	eason	
Species	m (ppb)	P 0.25 (ppb)	P 0.75 (ppb)	n (X)	m (ppb)	P 0.25 (ppb)	P 0.75 (ppb)	n (X)	m (ppb)	P 0.25 (ppb)	P 0.75 (ppb)	n (X)
NO	0.09	0.07	0.15	372	0.09	0.07	0.17	295	0.12	0.07	0.13	229
NO_2	4.54	2.85	6.46	630	1.78	1.23	2.56	697	1.07	0.82	1.76	268
O_3	24.3	11.76	34.87	826	25.59	15.91	32.95	781	14.74	10.77	19.39	316
NH_3	1.81	1.10	2.91	298	1.06	0.5	1.74	236	0.55	0.38	0.85	60
HNO_3	0.16	0.10	0.25	317	0.06	0.03	0.13	210	0.06	0.04	0.08	52
HONO	0.12	0.08	0.27	323	0.07	0.05	0.09	315	0.06	0.04	0.07	139
Aerosol NH ₄ ⁺	1.01	0.73	1.51	291	0.54	0.33	0.85	267	0.47	0.32	0.62	66
Aerosol $NO_3^{\frac{1}{2}}$	0.34	0.17	0.61	297	0.09	0.06	0.15	282	0.06	0.04	0.07	33

^(*) m: median, P0.25: 0.25 percentile, P0.75: 0.75 percentile, n: number of determined data points above the limit of detection (for aerosol species $PM_{2.5}$ and bulk measurements were included). NO_x/O_3 data were synchronized to the WAD/SJAC data.

3.3 Determination of characteristic time scales

The resistance-based approaches presented above to calculate surface-atmosphere exchange fluxes rely on the "constant flux layer assumption", which implies that the trace compounds are considered as chemically-non-reactive tracers, such that their flux within the atmospheric surface layer is constant. However, sufficiently accurate fluxes of compounds that undergo rapid chemical transformation can be estimated as long as characteristic chemical time scales are one order of magnitude larger than turbulent transport times (Damköhler ratio $D_r < 0.1$) (De Arellano and Duynkerke, 1992). Following De Arellano and Duynkerke (1992) the characteristic time of turbulent transport τ_{turb} (s) can be calculated as:

$$\tau_{\text{turb}} = \kappa \cdot (z_{\text{ref}} + z_0) \cdot \frac{u_*}{\sigma_w^2} \tag{9}$$

where σ_w denotes the standard deviation of the vertical wind component (m s⁻¹). Atmospheric stability strongly determines the time scale of turbulent transport, which typically ranges from a couple of seconds under unstable conditions up to 2.5 h under stable conditions (for a layer of 10 m height) (Dlugi, 1993).

For the characteristic chemical time scale of the NO-NO₂-O₃ triad, its photo-stationary equilibrium has to be considered. NO₂ is rapidly photolyzed to NO in the troposphere and it is re-formed by reaction of NO with O₃. The overall chemical time scale of the NO-NO₂-O₃ triad τ (NO-NO₂-O₃) is given by the combination of τ (NO)=(k₂×[O₃])⁻¹, τ (NO₂)= k_1^{-1} =j(NO₂)⁻¹ and τ (O₃)=(k_2 ×[NO])⁻¹ (Lenschow, 1982) (where k is the reaction rate constant) and was calculated in accordance to

Kirkman et al. (2002). Thereby, $j(NO_2)$ was estimated from global radiation data using a relation derived from simultaneous measurements of global radiation and $j(NO_2)$ in Amazonia during LBA-EUSTACH (cf. Kirkman et al., 2002).

HONO is rapidly photolyzed during daylight hours. The chemical time scale for daytime HONO photolysis is given by $\tau(\text{HONO})_{\text{photol.}} = j(\text{HONO})^{-1}$, whereby the parameterization provided by Kraus and Hofzumahaus (1998) was used to relate j(HONO) to $j(\text{NO}_2)$. The chemical time scale of heterogeneous HONO formation at nighttime $\tau(\text{HONO})_{\text{het.}}$ was derived by considering the HONO production rate P_{HONO} (ppb h⁻¹) determined directly from our measurements (see Sect. 4.3). Homogeneous daytime HONO formation may proceed via reaction of NO with OH radicals; however, this process is very slow and can be neglected compared to daytime HONO photolysis (Lammel and Cape, 1996).

Photochemical reactions involving NH₃ and HNO₃ are not considered to contribute to any flux divergence since their chemical time scale is much larger than the time scale of turbulent transport (Pandis et al., 1995). By contrast, time scales to achieve gas/aerosol equilibrium between gaseous NH₃, HNO₃ and particulate NH₄NO₃ may occur within the time frame of turbulent transport (few seconds for submicron particles) and have to be explicitly considered here (Dlugi, 1993; Meng and Seinfeld, 1996). As reported by Trebs et al. (2005), the measured concentration product of NH₃×HNO₃ was persistently below the theoretical equilibrium dissociation constant of the pure NH₃/HNO₃/NH₄NO₃ system during daytime (*RH*<90%), but approached the theoretical equilibrium value during nighttime (*RH*>90%). The desired equilibration time scale τ(NH₃-HNO₃-NH₄NO₃)

can be estimated according to (Wexler and Seinfeld, 1992):

$$\tau (NH_3 - HNO_3 - NH_4NO_3)^{-1}$$

$$= 3\overline{D} \times \int_0^\infty \frac{m(R_p)dR_p}{\left(1 + \frac{\lambda}{\alpha + R_p}\right) \times R_p^2 \times \rho_p}$$
(10)

whereby the aerosol mass size distribution $m(R_p)dR_p$ was related to the measured aerosol number size distribution. R_p is the particle radius (m), \overline{D} is the geometric mean of the diffusivity of semi-volatile gaseous species (m² s⁻¹), m is the water-soluble particle mass (kg m⁻³), λ is the mean free path of air $(6.51\times10^{-8}\,\mathrm{m}$ at 293.15 K), α denotes the accommodation coefficient $(0.001<\alpha<1)$ (Wexler and Seinfeld, 1990), and ρ_p is the particle density. A value of ρ_p =1.35±0.15 g cm⁻³ was used as determined by Reid and Hobbs (1998) for Amazonian biomass burning aerosols.

The comparison of τ_{turb} with $\tau(\text{NO-NO}_2\text{-O}_3)$, $\tau(\text{HONO})_{\text{photol.}}$, $\tau(\text{HONO})_{\text{het.}}$ and $\tau(\text{NH}_3\text{-HNO}_3\text{-NH}_4\text{NO}_3)$ provides a test whether reactive species can be treated as a passive tracer during their vertical transport within the surface layer ($D_r < 0.1$) and, consequently, if the application of the "big leaf multiple resistance approach" and the "canopy compensation point model" are justifiable.

3.4 N wet deposition

Storm size influences the chemical composition of the rainwater, in the sense that larger storms tend to be more dilute. This dependence demands the use of volume weighted means (VWM) for the calculation of monthly and annual concentration averages. For each rainwater solute, the concentrations measured in the sample were combined to create a VWM concentration for each collection date:

$$VWM_a = \frac{\sum_{i=1}^n c_{ai} \times v_i}{\sum v_i} \tag{11}$$

where c_{ai} is the concentration of species a in sample i, nis the number of samples and v_i is the volume of precipitation solution for sample i. The rain water concentrations measured during SMOCC were extrapolated to the whole year. Since storm size can vary from year to year we used long-term rainfall amounts to obtain the annual N deposition rate (cf. Lara et al., 2001). Rainfall amounts and air mass trajectories during the SMOCC period were comparable to conditions during the same period of previous years. Thus, these months can be considered as representative of the meteorological conditions during this period in this area. The N wet deposition rates for September, October and November were derived by summation of the VWM concentration determined for each precipitation event throughout the month, and subsequent multiplication by the mean of rainfall in the respective month obtained from historical time series (see http://www.aneel.gov.br). The annual N wet deposition was derived in a similar way; for instance, the VWM concentration obtained during September was considered as representative of the entire dry season and then multiplied by the historical mean rainfall for the dry season. The same procedure was performed for October and November representing transition period and wet season, respectively.

4 Results and Discussion

In the following, the general patterns of our results will be shown as diel courses of the medians of measured and inferred quantities. Diel courses will be presented either for part of the dry season (biomass burning, 12–23 September) or, where the investigated quantities are relatively independent of season, for the entire measurement campaign (12 September–14 November 2002). To derive surface-atmosphere exchange fluxes of N-containing compounds, meteorological data and NO_x/O_3 data were synchronized to the time resolution of the WAD/SJAC system. The convention of negative downward fluxes (net deposition) and positive upward fluxes (net emission) has been adopted.

4.1 Meteorological conditions

In the Amazon Basin, nighttime radiative cooling usually results in the formation of a shallow, decoupled nocturnal boundary layer of high thermodynamic stability characterized by very low wind speeds; while the development of a deepening, convectively mixed layer starts with the heating of the surface in the morning (Nobre et al., 1996; Fisch et al., 2004). Median nighttime wind speeds (at height $z=5.0 \,\mathrm{m}$) at FNS were low ($\leq 1 \text{ m s}^{-1}$). By contrast, wind speeds were substantially higher during daytime, ranging from 2 to 3 m s⁻¹. The 24-h average friction velocity (u_*) was $\sim 0.16 \,\mathrm{m \ s^{-1}}$ and reached maxima of $\sim 0.4 \,\mathrm{m \ s^{-1}}$ around noon. Differences of u_* between the dry season, the transition period, and the wet season were marginal. As a consequence of high global radiation fluxes (500–900 W m⁻², median) during daytime, ambient temperatures at the FNS site ($z=5.0 \,\mathrm{m}$) ranged between 30 and 36°C during sunlight hours, while nighttime temperatures were much lower (20– 25°C). High relative humidities (RHs) (z=5.0 m) were observed during nighttime (90–100%), while daytime RH values usually dropped to 40–50% during afternoon hours. Local meteorology changed only marginally from the dry season to the wet season. Strong rain events ($>30 \text{ mm h}^{-1}$) occurred in the afternoon hours during the transition period and the wet season. A more detailed description of diel meteorological conditions is presented by Trebs et al. (2005).

4.2 Mixing ratios

Median diel variations of mixing ratios of NO, NO₂, O₃, NH₃, HNO₃, HONO and the inorganic aerosol species NH $_4^+$ and NO $_3^-$ are shown in Fig. 2a–h for 12 to 23 September (dry

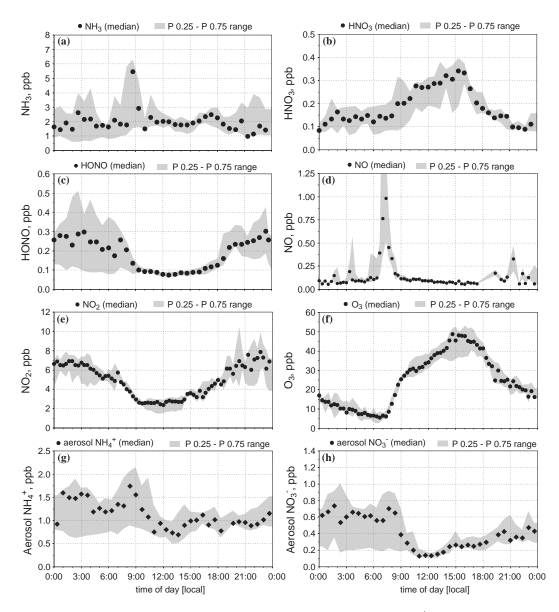


Fig. 2. Diel variations of (a) NH₃, (b) HNO₃, (c) HONO, (d) NO, (e) NO₂, (f) O₃, (g) aerosol NH₄⁺ (PM_{2.5}) and (h) aerosol NO₃⁻ (PM_{2.5}) measured during 12–23 September 2002 (dry season, biomass burning) at FNS during LBA-SMOCC 2002 (conversion factors from ppb to μ g m⁻³ for standard conditions of 298.15 K and 1000 hPa: NH₃: 0.69, HNO₃: 2.54, HONO: 1.89, NO: 1.21, NO₂: 1.86, O₃: 1.94, aerosol NH₄⁺: 0.73, aerosol NO₃⁻: 2.77). Symbols and grey shading represent medians and interquartile ranges (0.25 to 0.75 percentiles), respectively.

season, biomass burning). Table 2 summarizes ambient mixing ratios measured during the dry season (12–23 September), the transition period (7–31 October) and the wet season (1–14 November). A detailed discussion of seasonal and diel cycles observed for NH₃, HNO₃, HONO and aerosol NH₄⁺ and NO₃⁻ is given in Trebs et al. (2004, 2005).

Despite intensive biomass burning activity during the dry season, NO mixing ratios were very low (Fig. 2d). The sharp peak between 06:00 and 09:00 LT was most likely due to rapid photolysis of accumulated nighttime NO₂ shortly after sunrise, when O₃ mixing ratios were still too low to re-

oxidize significant amounts of NO (see Fig. 2e, f), but may also be partially attributed to rapid HONO photolysis. NO_2 was the most abundant N-containing trace gas during all three seasons and reached an average mixing ratio of \sim 5 ppb during the dry season. NO_2 featured a pronounced diel cycle with nighttime mixing ratios two times higher than during daytime (Fig. 2e). Apparently, NO_2 was accumulated in a shallow nocturnal boundary layer of high thermodynamic stability due to: (i.) the low water-solubility of NO_2 and consequently its low affinity to be taken up by epicuticular water films and (ii.) chemical production through reaction of

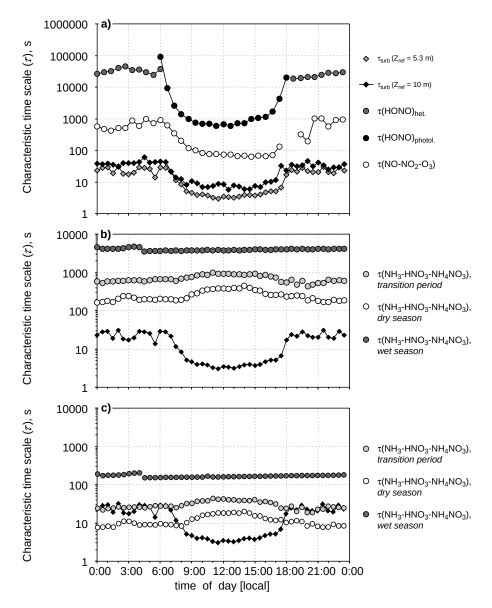


Fig. 3. Diel variation of characteristic turbulent time scale τ_{turb} for the reference heights z_{ref} =10 m and z_{ref} =5.3 m in comparison to (a) chemical time scale of the NO-NO₂-O₃ triad (τ (NO-NO₂-O₃)), daytime HONO photolysis (τ (HONO)_{photol.}) and heterogeneous HONO formation at night (τ (HONO)_{het.}, dry season only), (b) upper estimate of equilibration time scales for the NH₃/HNO₃/NH₄NO₃ system τ (NH₃-HNO₃-NH₄NO₃) (α =0.1, PM_{inorg.} ≤20%) and (c) lower estimate of τ (NH₃-HNO₃-NH₄NO₃) (α =1, PM=100%) at FNS during LBA-SMOCC 2002. Except for τ (HONO)_{het.}, data from all seasons were used.

NO with O_3 in the absence of NO_2 photolysis. Through the transition period until the wet season, NO_2 mixing ratios declined by a factor of four. O_3 mixing ratios exhibited a typical diel variation, which mirrors that of NO_2 (high values during daytime and lower values during the night; Fig. 2f). This was mainly caused by (i.) photochemical daytime production, (ii.) convective mixing within the boundary layer and from the free troposphere during daytime and (iii.) dry deposition and reaction with NO in a thermally stable stratified nocturnal boundary layer.

4.3 Characteristic timescales

Characteristic turbulent time scales (τ_{turb}) have been calculated according to Eq. (9) for z_{ref} =10 m (NO_x/O₃ measurements) and for z_{ref} =5.3 m (WAD/SJAC measurements) (cf. Table 1). To calculate the characteristic time scale for heterogeneous nighttime HONO buildup (τ (HONO)_{het.}) (Fig. 3a), only dry season nighttime HONO production rates (P_{HONO}) were considered. For the transition period and the wet season P_{HONO} could not be determined since the HONO diel variation was substantially reduced (cf. Trebs et

al., 2004). In contrast to other studies (e.g., Alicke et al., 2003; Harrison and Kitto, 1994; Lammel and Cape, 1996), our measurements revealed a relatively small average value of $P_{\rm HONO}$ =0.04 ppb h⁻¹. Figure 3a shows that $\tau_{\rm turb}$ was at least two orders of magnitude smaller than $\tau({\rm HONO})_{\rm photol.}$ and $\tau({\rm HONO})_{\rm het.}$, resulting in $D_r \ll 0.1$. Considering the chemical time scale of the NO-NO₂-O₃ triad $\tau({\rm NO-NO_2-O_3})$ (Fig. 3a), largest D_r values are found between 17:00 LT and 20:00 LT, exceeding a value of 0.2 ($z_{\rm ref}$ =10 m). However, during all other periods D_r for the NO-NO₂-O₃ triad ranged between 0.05 and 0.1. Therefore, we conclude that the application of Eqs. (1) and (4) to calculate surface-atmosphere exchange fluxes of NO₂ and HONO are justified, since chemical transformations are too slow to affect the vertical constancy of turbulent fluxes.

The estimation of the equilibration time scale $\tau(NH_3$ -HNO₃-NH₄NO₃) was performed by integrating over the measured particle size distribution according to Eq. (10), whereby two different cases were considered. Case 1 is an upper estimate (Fig. 3b), using an accommodation coefficient α =0.1 (Wexler and Seinfeld, 1992) and taking into account only the inorganic water-soluble aerosol fraction (≤20% of PM_{tot}) (cf. Trebs et al., 2005). Case 2, the lower estimate (Fig. 3c), was calculated using $\alpha=1$ and assuming that the entire aerosol mass (water-soluble and non-soluble species) are available to equilibrate. As shown in Fig. 3b, c τ (NH₃-HNO₃-NH₄NO₃) increased substantially from the dry season through the transition period to the wet season. This is obviously caused by much higher particle number concentrations measured during the dry season when biomass burning took place. For case 1 (upper estimate) (Fig. 3b), D_r for the NH₃-HNO₃-NH₄NO₃ triad during the dry season ranges from 0.1 to 0.17 at nighttime and is significantly smaller than 0.1 during the day. During the transition period and wet season, D_r for the NH₃-HNO₃-NH₄NO₃ triad is always significantly smaller than 0.1 (Fig. 3b). However, for case 2 (lower estimate) (Fig. 3c) nighttime D_r substantially exceeds a critical value of 0.1 during the dry season and the transition period.

To verify the theoretically derived values, results from a laboratory study will be discussed briefly. Condensation and evaporation of NH₃ and HNO₃ to/from particles have been investigated under controlled laboratory conditions. Particles were collected during field campaigns in 1991 (Brunnemann et al., 1996; Seidl et al., 1996) and 1993/1994 in the eastern part of Germany (Melpitz). The chemical aerosol composition was dominated by (NH₄)₂SO₄, NaCl and soot and is comparable to that described in Brunnemann et al. (1996) and Seidl et al. (1996). More than 90% of the NH₄NO₃ mass was found in the accumulation mode ($D_p \le 1 \mu m$). Aerosol samples were exposed to temperature step changes of 1 to 20 K above 278 K, which simulates a moderate diel cycle. RH was held constant for these cases. A second set of experiments was performed where RH was varied between 30 and 96% at constant temperature. In both cases non-volatile

Table 3. Ranges of daytime and nighttime surface resistances R_c for HNO₃ and NO₂ used to estimate flux scenarios at FNS during LBA-SMOCC 2002.

	Scenario	$R_c(\text{HNO}_3)$ [s m ⁻¹]	$R_c(NO_2)$ [s m ⁻¹]
Day	high flux	1	75
	low flux	50	550
Night	high flux	1	80
	low flux	15	435

SO₄²⁻, Na⁺ and K⁺ were considered as reference for the relative change of NO₃⁻ and NH₄⁺ while changing thermodynamic conditions. The mass loss of ammonium nitrate was recorded as a function of time for particles with a radius of r<0.8 μ m and r>1.6 μ m. Timescales τ (NH₃-HNO₃-NH₄NO₃) were calculated using the algorithm proposed by (i.) Kramm and Dlugi (1994) and (ii.) Meng and Seinfeld (1996), resulting in $\tau (NH_3-HNO_3-NH_4NO_3)=100-500$ s for fine mode particles and $\geq 880 \,\mathrm{s}$ for coarse mode particles. These values are comparable to characteristic times reported by Harrison et al. (1990) and Meng and Seinfeld (1996) and are in strong favor of our case 1 (upper estimate, Fig. 3b), which implies that the equilibration time scale was always much larger than that of turbulent transport. Nevertheless, it should be noted that the influence of the large soluble organic aerosol fraction typical for the Amazon Basin (cf. Trebs et al., 2005) on gas/aerosol partitioning processes is not exactly known. Taking into account the potential role of WSOC in enhancing aerosol water uptake and subsequently the uptake of gaseous species, equilibration time scales may be equal or even faster than turbulent transport (cf. Fig. 3c).

4.4 The inferential approach: selection of input parameters

Some surface parameters required for the inferential method to estimate of surface-atmosphere exchange fluxes for trace gases (Eqs. 1–7) could not be directly derived from the results of our field measurements. Thus, lower and upper scenarios were estimated (except for HONO; see below), comprising a certain range of surface-atmosphere exchange fluxes. These scenarios were obtained by varying surface parameters over a selected range based on results from studies in temperate latitudes. Note that for all parameters and quantities presented, values indicated as "low" and values indicated as "high" correspond to the estimated lower and upper fluxes, respectively.

Surface resistances R_c (HNO₃) and R_c (NO₂): R_c (HNO₃) was found to be zero in many studies (e.g., Dollard et al., 1987; Huebert and Robert, 1985). Very recently, Nemitz et al. (2004a) showed that non-zero, however, relatively small R_c (HNO₃) may exist (R_c =15–95 s m⁻¹). The ranges

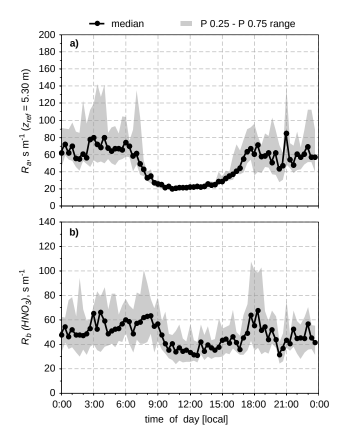


Fig. 4. Diel variations of (a) turbulent resistance (R_a) and (b) quasi-laminar boundary layer resistance (R_b) exemplary for HNO₃ at FNS during LBA-SMOCC 2002. Symbols and grey shading represent medians and interquartile ranges (0.25 to 0.75 percentiles), respectively. Data from all seasons were used.

of $R_c(\text{HNO}_3)$ used to estimate lower and upper surfaceatmosphere exchange fluxes are given in Table 3. The surface uptake of HNO_3 is considered to be enhanced by the presence of epicuticular water films under the humid conditions at the site (lower R_c values were chosen for nighttime). $R_c(\text{NO}_2)$ has been determined previously at the FNS site by Kirkman et al. (2002), hence the maximal and minimal diel variations of $R_c(\text{NO}_2)$ were taken directly from Kirkman et al. (2002) to estimate lower and upper surface-atmosphere exchange, respectively. Corresponding averages of maximal and minimal $R_c(\text{NO}_2)$ values are given in Table 3.

Stomatal resistance $R_s(NH_3)$ and charging resistance $R_d(NH_3)$: $R_s(NH_3)$ is not known for Brachiaria brizantha grass species, therefore the diffusion of H_2O through plant stomata is used for the calculation of upper and lower flux estimates. Thus, $R_s(NH_3)$ is estimated according to Nemitz et al. (2004a) using the measured latent heat flux LE. This approach is only valid for relatively dry daytime conditions (10:00–18:00 LT) and in the absence of precipitation. The transfer of H_2O through plant stomata during daytime represents an upper boundary for trace gas stomatal exchange.

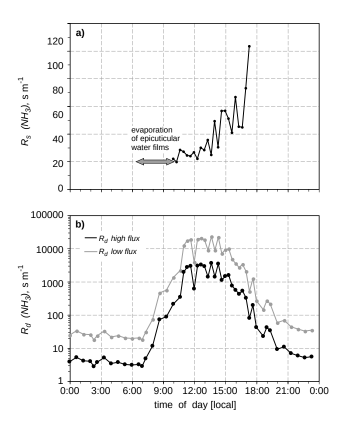


Fig. 5. Median diel variations of estimated (a) NH₃ stomatal resistance $R_s(\text{NH}_3)$ during daytime, using the measured latent heat flux LE ($R_s(\text{NH}_3)$) was set to $1000 \,\mathrm{m \ s^{-1}}$ for nighttime periods) and (b) lower (high flux) and upper (low flux) NH₃ charging resistance of the surface water layer ($R_d(\text{NH}_3)$) at FNS during LBA-SMOCC 2002. Data from all seasons were used.

Since stomata are thought to be closed during nighttime, $R_s(NH_3)$ is set to 1000 s m^{-1} for nighttime periods. The calculation of $R_d(NH_3)$ is performed in accordance with Sutton et al. (1998).

Compensation point concentration $X_c(HONO)$: Up to date, only two studies provide an estimate for $X_c(HONO)$ (Harrison and Kitto, 1994; Stutz et al., 2002). Here, the relationship $X_c(HONO)=0.03\cdot X(NO_2, z_{ref})$ was used, which was found for grassland in the recent study by Stutz et al. (2002). This corresponds on average to $X_c(HONO)=85$ ppt at the FNS site.

Apoplastic $[NH_4^+]/[H^+]$ ratio Γ and epicuticular pH: For Γ , we have chosen values of 100 (lower estimate) and 200 (upper estimate), which range at the lower end of data reported for grass in the literature (Loubet et al., 2002; Spindler et al., 2001; van Hove et al., 2002). This may be justified by the poor soil N status at FNS (Kirkman et al., 2002), because N absorbed by the root medium strongly affects the leaf tissue NH_4^+ concentration (Schjoerring et al., 1998a). The capacitance of the epicuticular water film is a function of the pH (see Sect. 3.1, Sutton et al., 1998), that is predominantly

acidic (Flechard et al., 1999) and is taken as 4.5 (lower estimate) and as 4.0 (upper estimate) (cf. Sutton et al., 1998).

 NH_3 flux from cattle excreta $F(NH_3)_e$: In order to estimate the contribution of cattle excreta to the net NH_3 flux, we considered results of Boddey et al. (2004), who investigated the cycling of N in *Brachiaria* pastures in the south of the Brazilian province Bahia. According to their results, *Bos indicus* cattle excreted 37 kgN animal⁻¹ yr⁻¹ in manure and 49 kgN animal⁻¹ yr⁻¹ in urine when the pastures were stocked with two animals per hectare. The stocking rate at FNS was one animal per hectare. About 8% of the excreted N may be released as NH_3 (A. F. Bouwman, personal communication, 2004). Hence, the average $F(NH_3)_e$ is estimated to be ~ 10 ngN m⁻² s⁻¹, which is applied for the entire measurement period (Eqs. 5, 6).

4.5 Resistances, NH₃ canopy compensation point, transferand deposition velocities

The turbulent resistance R_a and the quasi-laminar boundary layer resistance R_b (Fig. 4a, b): These resistances feature a typical diel variation with lowest values during daytime (R_a (5.3 m)=20 s m⁻¹, R_b =30 s m⁻¹), because of strong turbulent mixing within the surface layer, and higher values during nighttime (R_a (5.3 m)=70 s m⁻¹ and R_b =50 s m⁻¹).

The stomatal resistance $R_s(NH_3)$ and the charging resistance $R_d(NH_3)$ (Fig. 5a, b): The median diel cycle of $R_s(NH_3)$ estimated from LE during daytime (Fig. 5a) reaches very low values, falling below 50 s m⁻¹ between 10:00 and 14:00 LT. In the afternoon $R_s(NH_3)$ increases and attains values > 100 s m⁻¹ after 17:00 LT. $R_s(NH_3)$ is set to a value of 1000 s m⁻¹ for nighttime periods (see above; not shown). The simulated $R_d(NH_3)$ (Fig. 5b) is very high during daytime (>1000 s m⁻¹) and remains below 100 s m⁻¹ at night. This is obviously due to the fact that grass leaves became dry during sunlight hours (see below, Fig. 6), resulting in a very low capacity of the foliar cuticle to adsorb NH₃.

The NH₃ canopy compensation point concentration $X_c(NH_3)$ (Fig. 6): The $X_c(NH_3)$ scenario for the dry season calculated according to Eq. (6) lies well within the range of values determined for grassland in other studies (Hesterberg et al., 1996; Meixner et al., 1996; Spindler et al., 2001; Sutton et al., 2001) and is a strong function of surface temperature (cf. Eq. 6, 7). $X_c(NH_3)$ is predicted to be particularly high at daytime, although low values of Γ were used to run the model. This is caused by prevailing high surface temperatures at the FNS pasture site (35–40°C at daytime and 20– 25°C at night during the dry season). NH₃ deposits and dissolves in epicuticular water films at high RHs during nighttime. After sunrise the increasing surface temperature causes the NH₃ partial pressure above the epicuticular solution to increase in accordance to Henry's law. Hence, $X_d(NH_3)$ increases, which in turn results in higher $X_c(NH_3)$ values. This is visible in Fig. 6 as a distinctive peak of $X_c(NH_3)$ after sunrise between 08:00 and 09:00 LT. Once most of the

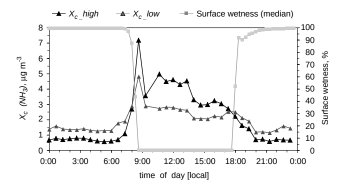


Fig. 6. Median diel variation of the simulated lower (low flux) and upper (high flux) NH₃ canopy compensation point $X_c(NH_3)$, and the median diel variation of the measured surface wetness, shown exemplarily for a period during the dry season (12–23 September) at FNS during LBA-SMOCC 2002. For details about low and high estimates, see text.

epicuticular NH₄⁺ has evaporated, X_c (NH₃) subsequently decreases (after 09:00 LT). Lower surface temperatures during the transition period and the wet season have caused lower estimates of X_c (NH₃) (0.3–1 μ g m⁻³ during nighttime and 1–4 μ g m⁻³ during daytime).

Transfer- and deposition velocities $V_{\rm tr.}(NH_3)$, $V_{\rm tr.}(HONO)$, $V_d(HNO_3)$, $V_d(NO_2)$ (Fig. 7a–d) and $V_p(aerosol)$ (Fig. 8): For compounds featuring a bidirectional surface exchange the concept of a dry deposition velocity $V_d = (R_a + R_b + R_c)^{-1}$ (see Eq. 1) is no longer useful (e.g., Kramm and Dlugi, 1994). Thus, for bi-directional NH₃ and HONO surface-atmosphere exchange the term transfer velocity $V_{\rm tr.} = -{\rm sgn}(X_c - X(z_{\rm ref}))/(R_a + R_b)$ (see Eqs. 4, 5) will be used henceforth.

The estimated median diel variation of V_{tr} (NH₃) (Fig. 7a) indicates net deposition (= downward flux) at nighttime $(V_{tr.}(NH_3)>0)$ and net emission (= upward flux) $(V_{\rm tr.}({\rm NH_3})<0)$ during the day. Typically, $V_{\rm tr.}({\rm NH_3})$ is predicted to range between $-2.6 \,\mathrm{cm} \,\mathrm{s}^{-1}$ during the day and 1.5 cm s⁻¹ at nighttime. For cases of NH₃ deposition, $V_{\rm tr.}({\rm NH_3})$ is in good agreement with values of 0.1-2 cm s⁻¹ observed by Erisman and Wyers (1993), Hesterberg et al. (1996), Nemitz et al. (2004a), Phillips et al. (2004) and Rattray and Sievering (2001). For cases of NH₃ emission, $V_{tr.}(NH_3)$ agrees well with values reported by e.g., Nemitz et al. (2004a). The application of the dynamic resistance model from Sutton et al. (1998), which uses nonzero $X_c(NH_3)$ values and takes into account that deposited NH₃ may (re-)evaporate from surfaces, represents a rather conservative N deposition estimate (lower boundary). Since $X_c(NH_3)$ was assumed to be zero in several studies (Goulding et al., 1998; Russell et al., 2003; Singh et al., 2001; Tarnay et al., 2001), which implies that NH₃ is persistently net deposited to the surface (see Eq. 1), we have complemented the upper and lower bi-directional flux scenarios by

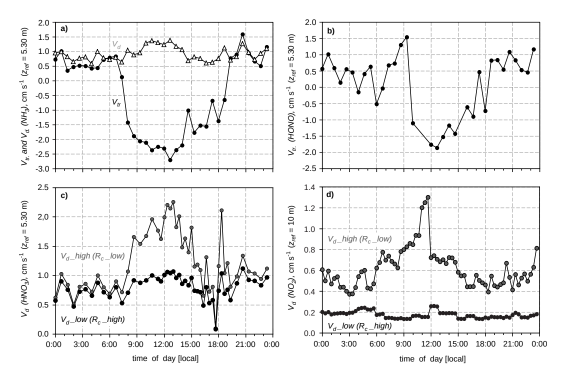


Fig. 7. Median diel variation of estimated transfer- $(V_{tr.})$ and deposition velocities (V_d) for (a) NH₃ $(z_{ref}=5.3 \, \text{m})$ (b) HONO $(z_{ref}=5.3 \, \text{m})$, (c) HNO₃ $(z_{ref}=5.3 \, \text{m})$ and (d) NO₂ $(z_{ref}=10 \, \text{m})$ at FNS during LBA-SMOCC 2002. The bi-directional NH₃ flux was complemented by a "deposition only scenario" (cf. V_d in a). Data from all seasons were used, except for $V_{tr.}$ (HONO), which only represents results from the dry season, since negligible small upward HONO fluxes were predicted during transition period and wet season.

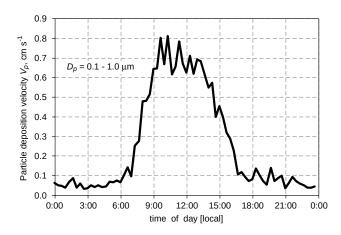


Fig. 8. Median diel course of the particle deposition velocity V_p ($z_{\rm ref}$ =5.3 m) using the empirical parameterization derived by Wesely et al. (1985) for aerosol particles with D_p =0.1–1 μ m at FNS during LBA-SMOCC 2002.

additionally calculating a "deposition only" scenario according to Eq. (1) (V_d in Fig. 7a), where $R_c({\rm NH_3})$ was calculated from the parallel resistances $R_s({\rm NH_3})$ and $R_d({\rm NH_3})$ (see Fig. 5a, b). Assuming NH₃ to be net deposited to the pasture site results in V_d =0.5–1.5 cm s⁻¹ (Fig. 7a).

The estimated median diel variation of $V_{\rm tr.}$ (HONO) (Fig. 7b, dry season only) similarly shows net deposition at night and net emission during the day, with $V_{\rm tr.}$ (HONO) ranging from -1.7 cm s⁻¹ to 1.5 cm s⁻¹. For cases of HONO deposition, these values are comparable to $V_{\rm tr.}$ (HONO) of 0–1.7 cm s⁻¹ reported by Harrison and Kitto (1994) and Stutz et al. (2002).

The estimates of $V_d(\mathrm{HNO_3})$ shown in Fig. 7c, reveal maximal median values of 2.3 cm s⁻¹ at around 13:00 LT when $R_c(HNO_3)=1$ s m⁻¹ is applied (see Table 3, upper flux estimate), coinciding with the period of highest turbulence. The lower estimate, when $R_c(HNO_3)$ is taken as 50 s m⁻¹ during daytime and as 15 s m^{-1} during nighttime (see Table 3), results in median $V_d(HNO_3)$ values nearly equal during day and night ($\leq 1.1 \text{ cm s}^{-1}$), which is attributed to the compensating effect of lower surface uptake and low turbulent exchange during nighttime. Although $V_d(HNO_3)$ was found to be higher $(3-4 \text{ cm s}^{-1})$ in other studies (e.g., Erisman et al., 1988), our values are still in reasonable agreement with deposition velocities determined by Dollard et al. (1987), Müller et al. (1993), Nemitz et al. (2004a) and Rattray and Sievering (2001) for vegetative canopies similar to that at FNS. It should be noted here that for a compound that is exchanged with a very small R_c (as in the case for HNO₃), V_d is mainly dependent on $R_a + R_b$. Thus, this comparison reflects largely the differences in turbulence and surface roughness.

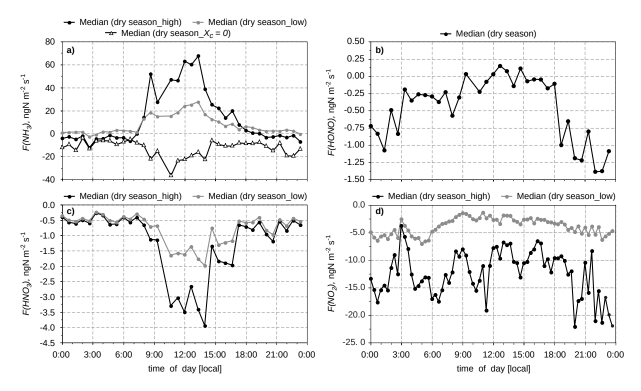


Fig. 9. Median diel courses of estimated lower (low) and upper (high) surface-atmosphere exchange fluxes for: (a) NH₃ (z_{ref} =5.3 m) (b) HONO (z_{ref} =5.3 m), (c) HNO₃ (z_{ref} =5.3 m) and (d) NO₂ (z_{ref} =10 m). Bi-directional NH₃ flux scenarios were complemented by a "deposition only" scenario (X_c (NH₃)=0). Only one scenario could be calculated for HONO due to the restricted availability of input parameters (cf. Sect. 4.4). Median diel courses are presented exemplarily for a period during the dry season (12–23 September) at FNS during LBA-SMOCC 2002.

The median diel course of $V_d({\rm NO_2})$ shown in Fig. 7d exhibits maxima of $\sim 1.3~{\rm cm~s^{-1}}$ during daytime, while minimal values are below $0.2~{\rm cm~s^{-1}}$. These values compare relatively well with ranges of $V_d({\rm NO_2})$ reported by Kirkman et al. (2002). However, the dry deposition of ${\rm NO_2}$ is thought to be mainly featured by uptake through plant stomata, which would imply that much higher dry deposition velocities (and hence much lower R_c) occur during daytime. As stated by Kirkman et al. (2002), measured day- and nighttime R_c values for ${\rm NO_2}$ at FNS were similar (see also Table 3), which was assumed to be mainly a result of stomatal closure due to high water vapor pressure deficit at lower RH's during daytime.

Deposition velocities V_p ($z_{\rm ref}$ =5.3 m) predicted for particles (Fig. 8) are estimated to be highest during daytime (\leq 0.8 cm s⁻¹) in accordance with higher values of u_* . During nighttime, V_p generally remains below 0.1 cm s⁻¹. As previously indicated, these values are much larger than V_p predicted by Slinn (1982) and are therefore considered as an upper estimate. It should be pointed out that aerosol water uptake at high RHs and the resulting particle growth during the deposition process may enhance deposition velocities (Khlystov, 1998; Gallagher et al., 1997). In our study, this would be particularly relevant for nighttime periods when

RH usually exceeded 90%. However, a quantification of this effect would go beyond the scope of this work.

4.6 Surface-atmosphere exchange fluxes

In this section, the measured mixing ratios (Sect. 4.2), the input parameters for the inferential approach (discussed in Sect. 4.4) and the results presented in section 4.5 will be used to estimate and discuss surface-atmosphere exchange fluxes of N-containing trace gases and aerosol particles.

Bi-directional fluxes of gaseous NH₃, F(NH₃) (Fig. 9a): Fluxes found at locations in temperate regions with vegetative surfaces similar to the grass surface at FNS are comparable to our upper bi-directional estimate of F(NH₃) for the dry season (biomass burning) (Fig. 9a) (Erisman and Wyers, 1993; Flechard et al., 1999; Meixner et al., 1996; Spindler et al., 2001). The net emission peak between 08:00 and 09:30 LT in the upper estimate corresponds to the peak of the predicted NH₃ canopy compensation point (cf. Fig. 6) and the highest observed NH₃ mixing ratios (see Fig. 2a). As already mentioned above, this feature may be related to desorption of NH₃ from epicuticular water films due to an increase of surface temperature after sunrise (see also Trebs et al., 2005). Relatively high median NH₃ net emission fluxes (5–70 ngN m⁻² s⁻¹, upper estimate) during daytime

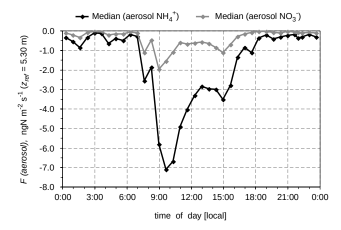


Fig. 10. Median diel courses of inferred deposition fluxes of aerosol NH₄⁺ and NO₃⁻ shown exemplarily for a period during the dry season (12–23 September) at FNS during LBA-SMOCC 2002.

(08:00–17:30 LT) are the result of (i.) relatively high NH₃ mixing ratios (Fig. 2a), (ii.) direct NH₃ emission from cattle excreta, (iii.) high surface temperatures and (iv.) corresponding high values of simulated $X_c(NH_3)$ (see Fig. 6). The simulated nighttime NH₃ net deposition (upper estimate) (19:00–06:30 LT) is on average -2 to -13 ngN m^{-2} s⁻¹. The lower NH₃ flux estimate shown in Fig. 9a suggests that the net exchange of NH₃ may be significantly smaller when a higher epicuticular pH (4.5) (Sect. 4.4), a lower apoplastic $[NH_4^+]/[H^+]$ ratio Γ (Sect. 4.4) and higher $R_d(NH_3)$ (Fig. 5b) are applied. The estimated NH₃ net deposition for this case during nighttime is negligible and the predicted daytime NH3 net emission varies between 1 and $28 \text{ ngN m}^{-2} \text{ s}^{-1}$. During the transition period and the wet season, however, the lower NH₃ flux estimate (not shown) features emission during day and night. The "NH3 deposition only" scenario ($X_c(NH_3)=0$; Fig. 9a) exhibits highest fluxes during daytime (08:00-15:00 LT), with values ranging between -20 and -40 ngN m⁻² s⁻¹. At nighttime, net deposition fluxes between -5 and $-20 \,\mathrm{ngN} \;\mathrm{m}^{-2} \;\mathrm{s}^{-1}$ were estimated.

The predicted bi-directional NH $_3$ flux scenarios are most sensitive to the pH of the epicuticular water film and the H $^+$ and NH $_4^+$ concentrations in the apoplastic fluid (Γ). If the pH of the epicuticular water film were >4.5 and Γ constant (100–200), the upper NH $_3$ flux estimate would show net emission also during nighttime. On the other hand, increasing Γ beyond a value of 200 would result in daytime NH $_3$ net emissions significantly higher than observed in urban regions in Europe or North America.

Bi-directional fluxes of gaseous HONO, F(HONO) (Fig. 9b): The estimated median diel HONO flux during the dry season (12 to 23 September), shows a small net emission during the afternoon but net deposition of up to -1.3 ngN m⁻² s⁻¹ from 17:00–08:40 LT. We like to note, that the HONO flux scenario presented here probably reflects the

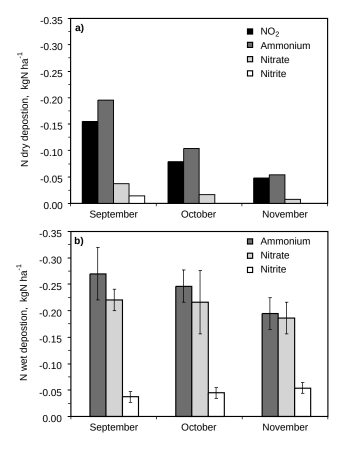


Fig. 11. Monthly estimates of (a) N dry deposition of NO_2 , total ammonium (aerosol $NH_4^+ + NH_3$), total nitrate (aerosol $NO_3^- + HNO_3$) and nitrite (HONO) (averages of lower and upper flux estimates are shown, except for NH_3 which represents an average of all three estimates (cf. Fig. 9a) and (b) N wet deposition of ammonium, nitrate and nitrite (error bars represent measurement uncertainties) for September (dry season, biomass burning), October (transition period) and November (wet season, clean conditions) at FNS during LBA-SMOCC 2002.

largest uncertainty of all flux estimates since presently neither $X_c(HONO)$, nor the exact HONO formation mechanism are well known.

Fluxes of gaseous HNO₃, $F(HNO_3)$ (Fig. 9c): The estimated median diel HNO₃ flux scenario during the dry season is characterized by highest net deposition values from 09:00–16:30 LT (\sim -0.5 to -4 ngN m⁻² s⁻¹, upper estimate), coinciding with highest estimated deposition velocities (cf. Fig. 7c) and the maximal observed HNO₃ mixing ratios (cf. Fig. 2b). Predicted nighttime HNO₃ deposition fluxes are lower in accordance with lower turbulence (lower V_d) and lower HNO₃ mixing ratios. The higher values of $R_c(\text{HNO}_3)$ applied for the lower flux estimate (Table 3) result in daytime net deposition fluxes that are a factor of two lower than for the upper flux estimate.

Fluxes of gaseous NO_2 , $F(NO_2)$ (Fig. 9d): The diel course of the estimated NO_2 flux scenario (dry season) is always

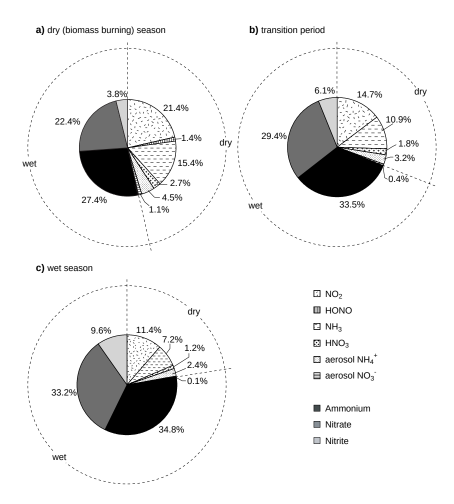


Fig. 12. Contribution of individual inorganic N compounds to the estimated total N deposition for (a) September (dry season, biomass burning), (b) October (transition period) and (c) November (wet season, clean conditions) at FNS during LBA-SMOCC 2002. Dry deposition of gaseous species is presented using averages of lower and upper flux estimates.

characterized by downward fluxes (net deposition). Maximal net deposition values range from -4 to -22 ngN m⁻² s⁻¹, whereas the lower scenario is characterized by values that are at least a factor of two smaller. Median nighttime fluxes (18:00–08:00 LT) are usually higher than during the day, which is mainly due to higher NO₂ mixing ratios measured during nighttime (Fig. 2e).

Fluxes of aerosol NH_4^+ and NO_3^- , F(aerosol) (Fig. 10): The estimated median diel flux of aerosol NH_4^+ for the dry season exhibits a net deposition pattern characterized by highest values just after 09:00 LT (\sim -7 ngN m⁻² s⁻¹), which is consistent with maximal aerosol NH_4^+ mixing ratios observed during this time (see Fig. 2g) and increased turbulent mixing (increasing u_*) after sunrise. Although aerosol NH_4^+ mixing ratios were generally lower during daytime, the dry deposition of aerosol NH_4^+ is estimated to be highest because of high V_p predicted during sunlight hours (see Fig. 8). The median diel course of the aerosol NO_3^- flux shown in Fig. 10 for the dry season resembles that estimated for aerosol NH_4^+ , although the net deposition flux was at least

three times lower than for aerosol NH_4^+ , which is a consequence of lower aerosol NO_3^- mixing ratios (see Fig. 2g, h). The highest aerosol NO_3^- net deposition is predicted at around 09:00 LT (\sim -2 ngN m⁻² s⁻¹).

4.7 Seasonal cycles of N dry and wet deposition

Figure 11a, b illustrates the estimated monthly N dry deposition (downward flux only) for NO₂, total ammonium, total nitrate and total nitrite. The dry deposition rates of NO₂ and total ammonium (NH₃ + aerosol NH₄⁺) are estimated to be highest and decline by about 70% from September (late dry season, biomass burning) to November (onset of the wet season, clean conditions) (Fig. 11a). This indicates the influence of biomass burning during the dry season, which significantly enhances the dry deposition fluxes of NO₂ and NH₃. The dry deposition of total nitrate (HNO₃ + aerosol NO₃⁻) also decreases by about 70% from September to November, but was on average 4 times lower than that of NO₂ and total ammonium. This is in accordance with lower mixing

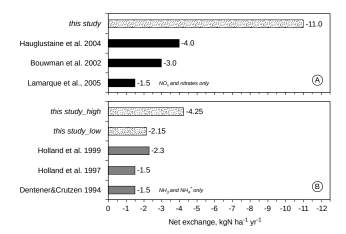


Fig. 13. Comparison of the annual N (wet + dry) deposition predicted by global chemistry and transport models for the Amazon region with the net annual N (wet + dry) exchange estimated in this study. Panel (**A**) represents the case when NH_3 is considered to be deposited only and in panel (**B**) the lower ("this study_low") and upper ("this study_high") bi-directional flux estimates are shown.

ratios observed for nitrate in gas and aerosol phase (Table 2) (cf. Trebs et al., 2004). The contribution of nitrite (HONO) to the dry deposition in September is smallest, and is negligible in October and November.

The estimated N wet deposition (Fig. 11b) is considerably higher than N dry deposition at the FNS site. The obtained wet deposition rate of nitrate is consistent with the results of Lara et al. (2006). N wet removal is dominated by ammonium and nitrate, and wet deposition of nitrite is at least 3 times smaller. Interestingly, the decrease of wet deposition of ammonium and nitrate from September to November is only $\sim\!20\%$, hence, much less pronounced than that found for N dry deposition.

During the late dry season (biomass burning) in September, on average 46.4% of the total N deposition can be attributed to dry deposition and ~53.6% to wet deposition (Fig. 12a). With increasing rainfall amounts, the contribution of dry deposition drops to only 31% and 22.4% to the total N deposition during October (transition period) and November (onset of the wet season, clean conditions), respectively (Fig. 12b, c).

N dry deposition at FNS was dominated by NO_2 and NH_3 . During September (dry season, Fig. 12a) the contribution of NO_2 and NH_3 to the total N deposition is estimated to be 21.4% and 15.4%, respectively. By contrast, the N dry deposition of HNO₃ and HONO is estimated to be small, contributing only 2.7% and 1.4% to the total N deposition during the dry season, respectively (Fig. 12a). Aerosol NH_4^+ and aerosol NO_3^- show comparably small contributions to the total N deposition during the dry season (4.5% and 1.1%, respectively).

Through the transition period (October) until the wet season (November), the contribution of NO_2 dry deposition to the total deposition declines to about 11% (Fig. 12b, c). The relative contribution of NH_3 to the total estimated N deposition decreases to 11% during the transition period and to about 7% during the wet season. The N dry removal associated with HNO_3 , HONO and aerosol NH_4^+ and NO_3^- is estimated to be negligibly small during the transition period and the wet season (<6% in total).

4.8 Annual budget of surface-atmosphere exchange

The annual budget of surface-atmosphere exchange of inorganic N species at the FNS pasture site has been determined making the following assumptions: (i.) September is representative for the entire dry season (May to September) $(PM_{10}=100-200 \,\mu g \,m^{-3})$, (ii.) October is representative for the two transition periods (April and October) (PM₁₀=50- $100 \,\mu \text{g m}^{-3}$), and (iii.) November is representative for the entire wet season (November to March) ($PM_{10} \le 25 \mu g \text{ m}^{-3}$) (Artaxo et al., 2002). If NH₃ is considered to be deposited only to the pasture site (i.e., no bi-directional exchange, cf. Fig. 9a), the annual N budget (i.e., the annual net N exchange at the pasture site) would be $-11 \text{ kgN ha}^{-1} \text{ yr}^{-1}$. Using the bi-directional scenarios, the annual N (wet + dry) deposition is estimated to range between -6.83 and -8.94 kgN $ha^{-1} yr^{-1}$. Thereof, -1.57 to $-3.68 kgN ha^{-1} yr^{-1} can be$ attributed to dry deposition and \sim -5.26 kgN ha⁻¹ yr⁻¹to wet deposition. Our estimates suggest that 2.58 to 6.79 kgN ha⁻¹ yr⁻¹ may be emitted in form of gaseous NH₃, HONO and NO from the pasture site (the annual estimate for NO emission was taken from Kirkman et al., 2002), whereof at least 90% are expected to be in the form of NH₃. Thus, considering the bi-directional scenarios, the annual N budget may range from -2.15 to -4.25 kgN ha⁻¹ yr⁻¹ (cf. Fig. 13).

Regarding soil emission of nitrous oxide (N₂O), not included in this study, Garcia-Montiel et al. (2003) reported that the emission of N₂O from Rondônian pastures is even smaller than that of NO. Kirkman et al. (2002) estimated that the FNS site constitutes a NO₂ sink of $-0.73 \, \mathrm{kgN} \, \mathrm{ha}^{-1} \, \mathrm{yr}^{-1}$. They assumed that their measurements from 24 September to 27 October 1999 (dry-wet transition period) were representative for the wet and dry season and that these seasons are about equal in length. However, based on our measurements and using the minimal and maximal $R_c(\mathrm{NO}_2)$ values of Kirkman et al. (2002), our annual NO₂ deposition estimate ranges from -0.76 to $-2.4 \, \mathrm{kgN} \, \mathrm{ha}^{-1} \, \mathrm{yr}^{-1}$, which is considerably higher, mainly due to higher NO₂ mixing ratios observed in our study.

Moreover, it should be noted that water-soluble organic nitrogen (WSON) constitutes a significant part of the total N fraction of Amazonian aerosols. It was found by Mace et al. (2003), that organic N may represent 45% and 43% of the total N in wet and dry season aerosol samples, respectively. Obviously, the total deposition of atmospheric N would be

significantly higher if these organic species were also taken into account.

Finally, Fig. 13 shows a comparison of our estimated annual net N exchange with predictions for annual N (wet +dry) deposition by global chemistry and transport models (CTMs) (Dentener and Crutzen, 1994; Holland et al., 1997, 1999; Bouwman et al., 2002; Hauglustaine et al., 2004; Lamarque et al., 2005). The values from Dentener and Crutzen (1994) and Holland et al. (1997) are averages of N deposition predicted for the Amazon region (MOGUNTIA model). The estimates from Dentener and Crutzen (1994) exclude dry deposition of NO_x and nitrates. Holland et al. (1999) (MOGUNTIA) present predictions for tropical grasslands, considering all components of N exchange. Bouwman et al. (2002) (STOCHEM model), Hauglustaine et al. (2004) (LMDz-INCA model) and Lamarque et al. (2005) provide the N deposition for South America, but do not take into account bi-directional exchange of NH₃. The most recent study by Lamarque et al. (2005) (average of six different CTMs) presents N deposition resulting from NO_x and nitrates, excluding deposition of NH₃. In the case where NH₃ is considered to be deposited only (Fig. 13, panel A) our estimates exceed those of global CTMs by more than a factor of two. In case of bi-directional exchange (Fig. 13, panel B), the result of our lower scenario agrees relatively well with predictions by global CTMs, whereas the result of the upper scenario is two times higher than CTM estimates.

It is noteworthy that the FNS site is situated in one of the most disturbed regions of the Brazilian Amazon. During the dry season, the site is characterized by high pollution levels compared to other Amazonian ecosystems. In remote Central Amazonia (smaller fire frequency/density), wet deposition rates of nitrate and ammonium were previously estimated by Stallard and Edmond (1981), Andreae et al. (1990) and Williams et al. (1997) to be $-1.7 \,\text{kgN ha}^{-1} \,\text{yr}^{-1}$, $-2.1 \,\text{kgN}$ $ha^{-1} yr^{-1}$ and $-2.8 kgN ha^{-1} yr^{-1}$, respectively. The corresponding value from our study $(-4.7 \text{ kgN ha}^{-1} \text{ yr}^{-1}; \text{ wet}$ deposition of ammonium + nitrate, nitrite excluded) is about twice as high as these results. However, the estimates of total N (dry + wet) deposition (see Fig. 13), calculated by global CTMs (which consider remote as well as polluted conditions), are equal to the deposition (wet only) rates for remote Amazonian sites given by Stallard and Edmond (1981), Andreae et al. (1990) and Williams et al. (1997).

5 Summary and conclusions

We have estimated dry and wet deposition of inorganic N at a tropical pasture site (Rondonia, Brazil) based on real-time measurements of inorganic N containing gases and aerosol species during the late dry (biomass burning) season, the transition period and the onset of the wet season (clean conditions) (LBA-SMOCC 2002). HNO₃, NO₂, aerosol NH₄⁺ and NO₃⁻ were considered to be net deposited to the pasture site

under all ambient conditions, while surface-atmosphere exchange of HONO and NH₃ is considered to be bi-directional. All fluxes have been determined by inferential techniques, making use of aerodynamic and surface related resistances that have been deduced from measurements or were taken from recent literature. For the least-well defined quantities, lower and upper cases have been considered. Bi-directional NH₃ fluxes were predicted using a dynamic resistance model from Sutton et al. (1998). NO₂ and NH₃ are the most important contributors to N dry deposition and their fluxes largely control the net N exchange pattern at the pasture site. Mixing ratios of NO₂ and water-soluble N species in gas and aerosol phase are significantly enhanced when widespread biomass burning takes place during the dry season, resulting in high N deposition rates. The contribution of N dry deposition to the total N (dry + wet) deposition decreased substantially from the dry (biomass burning) season to the wet season. In the case where bi-directional NH₃ exchange is considered, our results suggest that the annual N budget at the pasture site may range from -2.15 to -4.25 kgN ha⁻¹ yr⁻¹. When NH₃ is assumed to be deposited only, the overall net exchange is $-11 \text{ kgN ha}^{-1} \text{ yr}^{-1}$.

The largest uncertainties in our estimates are related to the bi-directional NH₃ exchange, namely (i.) the surface water (epicuticular) pH, (ii.) the ratio of NH₄⁺ and H⁺ concentration in the apoplastic fluid (Γ) and (iii.) direct NH₃ emissions from cattle excreta. These quantities strongly control the NH₃ canopy compensation point concentration. In this study, low values of (i.) the epicuticular pH (4-4.5) and (ii.) the ratio of NH₄⁺ and H⁺ concentration in the apoplastic fluid (Γ =100–200) were adopted. The latter can be justified by the low soil N status of the pasture. Choosing this parameterization, the characteristic diel pattern of the NH₃ exchange found in temperate latitudes could be reproduced (daytime net emission, nighttime net deposition). A comparison of our results to those of global CTMs suggests that net N deposition rates to Amazonian pastures may be higher than currently predicted by global CTMs.

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