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Fluxes of nitrogen oxides over a temperate deciduous forest

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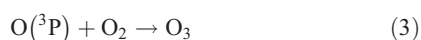
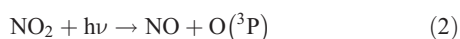
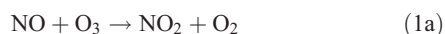
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[1] Eddy covariance flux measurements of NO, NO₂, and O₃ were obtained above the mixed deciduous canopy at Harvard Forest in central Massachusetts from April to November 2000. Net deposition of NO_x was observed throughout the measurement period, with average velocity of ~ 0.2 cm s⁻¹. At night, NO₂ is deposited at a rate that depends nonlinearly on NO₂ concentration and cannot be explained by N₂O₅ hydrolysis, suggesting HONO formation by heterogeneous disproportionation of NO₂. During the day, photochemically driven NO_x fluxes conform to the predicted behavior based on gradients of light and eddy diffusivity through the canopy, with residual net flux attributable to both stomatal and nonstomatal processes. The results were consistent with a compensation point for NO₂ near 1.5 nmol mol⁻¹. These results were confirmed by independent evidence from NO, NO₂, and O₃ profiles acquired at the site over several years. If the rate of NO_x deposition observed at this site in April through November continues during the winter, it would have a larger potential impact on tropospheric chemistry because mixing depths are shallow and chemical NO_x oxidation is slow during winter; the impact at night is important for the same reasons. The results contradict widely used parameterizations of NO₂ deposition that both overestimate stomatal uptake and do not allow for surface uptake when stomates are closed. *INDEX TERMS*: 0315 Atmospheric Composition and Structure: Biosphere/atmosphere interactions; 0322 Atmospheric Composition and Structure: Constituent sources and sinks; 0345 Atmospheric Composition and Structure: Pollution—urban and regional (0305); 0365 Atmospheric Composition and Structure: Troposphere—composition and chemistry; *KEYWORDS*: nitrogen oxides, fluxes, troposphere

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1. Introduction

[2] Tropospheric NO₂ is the critical nitrogen oxide radical in the photochemical formation of O₃. The NO_x species, NO and NO₂, rapidly interconvert in fast photochemical reactions (1)–(3). The cycle of reactions (1a), (2), and (3) has no net effect on O₃, but the sequence of reactions (1b), (2), and (3) produces O₃.



Production of O₃ is limited by removal of NO₂, either by chemical reactions to make the nonradical species HNO₃, or by deposition to the surface. Chemical removal during the day is mainly due to reaction of NO₂ with OH radicals, while at night removal proceeds via reaction with O₃ to make the nitrate radical (NO₃), which reacts with NO₂ to form N₂O₅ that can hydrolyze to HNO₃. These chain-terminating steps are slow compared to the chain propagating reactions (1)–(3), allowing one molecule of NO_x to be cycled many times before removal. Up to 100 molecules of O₃ may be generated for each NO_x molecule entering the remote troposphere, with ~ 5 O₃ thought to be typically produced per unit NO_x in the rural eastern United States [Liu *et al.*, 1987; Hirsch *et al.*, 1996].

[3] Deposition of NO₂ to surfaces bypasses atmospheric oxidation to HNO₃ and reduces the amount of NO_x exported to the remote troposphere, thus potentially reducing tropospheric O₃ production even if NO₂ deposition constitutes a small fraction of the total flux of fixed N to the surface [Jacob *et al.*, 1993; Liang *et al.*, 1998]. Chamber studies using NO₂ concentrations between 0.1

and 100 nmol mol⁻¹ have reported significant rates for uptake of NO₂ by vegetation through stomates [Rondón *et al.*, 1993; Thoene *et al.*, 1996; Gefßler *et al.*, 2000; Sparks *et al.*, 2001], adsorption on foliar surfaces [Gefßler *et al.*, 2002], and deposition to forest floors and soils [Hanson and Lindberg, 1991; Eugster and Hesterberg, 1996]. The NO₂ compensation point, the ambient concentration below which NO₂ is emitted from stomates, has been estimated by linear regression from higher concentrations [Thoene *et al.*, 1996] and observed directly [Rondón *et al.*, 1993; Gefßler *et al.*, 2000; Sparks *et al.*, 2001; Gefßler *et al.*, 2002] at values between 0.5 and 3 nmol mol⁻¹. Release of NO₂ by vegetation could significantly elevate global O₃ production rates because NO_x concentrations are lower than the inferred compensation point over much of the globe [Lerdau *et al.*, 2000].

[4] NO does not readily combine with atmospheric radicals to make stable nonradicals, and it is not expected to deposit to surfaces in significant amounts, though there is very little experimental confirmation [Hanson and Lindberg, 1991; Wesely and Hicks, 2000]. NO is emitted from soils, typically at low rates in temperate forests [Williams *et al.*, 1992; Munger *et al.*, 1996]. Nevertheless, rapid photochemical exchange between NO, O₃ and NO₂ make simultaneous quantification of fluxes for these species a requirement for elucidating NO_x deposition rates [Fitzjarrald and Lenschow, 1983].

[5] We present here NO and NO₂ eddy covariance fluxes at the Harvard Forest Environmental Measurement Site for spring throughfall, 2000, augmented by 12 years of data at the site for vertical profiles of NO₂ and NO concentrations and eddy covariance fluxes of O₃ and NO_y. The measurements represent the first simultaneous data for fluxes of NO, NO₂, and O₃ above a tall forest canopy with chemically specific eddy covariance instrumentation. The results allow us to critically test current assumptions about deposition and emission fluxes of NO_x at the whole-canopy scale.

2. Methods

[6] We studied a 50- to 70-year-old mixed deciduous forest of red oak and red maple, with scattered hemlock, red and white pine at Harvard Forest in central Massachusetts (42.54N, 72.18W; elevation 340 m). The landscape is ~95% forested and moderately hilly; the distance to the closest paved road is >1.5 km and to the nearest small town >10 km. Two wind directions dominate: northwest flows bring relatively cool, dry, unpolluted air, and southwest winds bring warm, humid, significantly more polluted air [Moody *et al.*, 1998; Munger *et al.*, 1996]. We utilized two towers for this study: a permanent 30-m Rohn 25G tower, used since 1990 to measure eddy fluxes of CO₂, NO_y, and O₃, along with vertical profiles of NO, NO₂, O₃, and other species; and a temporary steel scaffolding tower, 23 m high (just above the top of the canopy), situated about 100 m to the SE and outfitted with a sonic anemometer and a tunable diode laser absorption spectrometer (TDLAS) configured to measure eddy covariance fluxes and concentrations of NO₂ and concentrations of HNO₃ [Horie, 2002]. Data from the two towers were matched and averaged into an hourly

data set, available at <http://www-as.harvard.edu/chemistry/index.html>.

2.1. NO_x Photolysis-Chemiluminescence (P-C) Measurements

[7] The P-C system measures NO and NO₂ concentration profiles at heights of 29, 24.1, 18.3, 12.7, 7.5, 4.5, 0.8, and 0.3 m on the permanent tower [Munger *et al.*, 1996]. From late August to mid October 2000, it was reconfigured to measure NO concentrations at 10 Hz at the 29 m sampling height in order to determine eddy covariance fluxes for NO. Calibrations were made using standard gas mixtures for NO and NO₂, traceable to NIST. The NO₂ and NO standards were cross calibrated by titrating the NO standard with O₃ and by analyzing both using the NO_y detector equipped with a hot gold catalyst for reducing NO_y to NO.

2.2. NO₂ TDLAS Measurements

[8] The TDLAS [Horie *et al.*, 1999; Horie, 2002] ran two diodes simultaneously from April through November 2000. One channel measured NO₂ concentrations at 1 Hz and the other measured concentrations of HNO₃. The transition line strengths and pressure broadening coefficients used to compute absorption cross sections were obtained from the HITRAN spectral database [Rothman *et al.*, 1998]. TDLAS calibration relied on independent laboratory determination of spectroscopic parameters (light path length, laser purity and frequency, laser tuning rate function, and laser line width), which were also checked in the field monthly using standard additions of HNO₃ and NO₂, ambient water vapor, and a Germanium etalon. Multiple levels of cross calibration were maintained throughout the experiment. Independent cross-checks were obtained by comparing concentrations of H₂O inferred using spectral lines in both the NO₂ and HNO₃ spectral ranges (1590 and 1720 cm⁻¹, respectively) with H₂O data from hygrometers on the main tower. One-second, integrated measurements of both zero air and a constant NO₂ permeation tube source show that the RMS noise over a wide range of [NO₂] was 80 pmol mol⁻¹. NO₂ precision does not scale with trace gas concentration in the TDLAS; instead, precision is limited by the background noise of the photovoltaic detectors, the output power of the laser diode, and the finite reflectivity of the multiple pass mirrors. Short- and long-term temperature and pressure instabilities within the instrument limit the improvement of precision obtained from longer integration times to a factor of 2 to 4. Concentrations of NO₂ obtained by TDLAS and P-C instruments were directly compared during 459 hours of concurrent measurements (4 April to 29 August 2000); the orthogonal distance regression of hourly data for the TDLAS versus P-C had a slope of 1.1 ± 0.2 (95% confidence interval, R² = 0.91) and an intercept indistinguishable from zero [Horie, 2002].

2.3. O₃ UV Absorbance and Chemiluminescence Measurements

[9] O₃ concentration profiles and eddy covariance flux measurements on the main tower employ UV absorbance (Dasibi 1003AH, sampling at the same heights as above) and C₂H₄-chemiluminescence (29 m sampling height), respectively [Munger *et al.*, 1996]. The Dasibi 1003AH was calibrated by comparison to a calibration unit or return

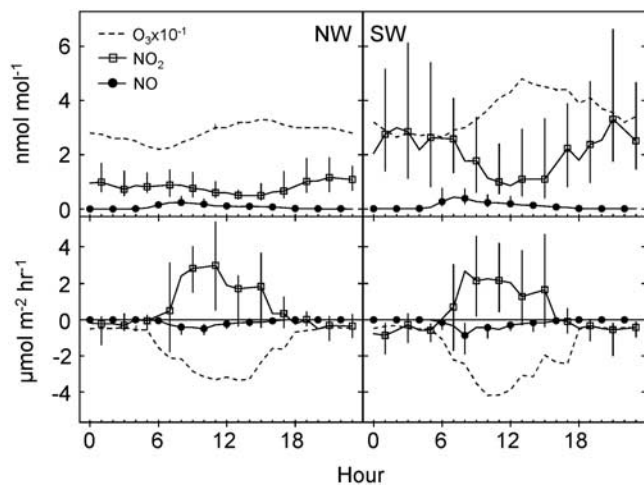


Figure 1. Diel cycles of (top) median concentrations and (bottom) fluxes for the (left) northwest (270°–45°) and (right) southwest (180°–270°) wind sectors at Harvard Forest, April–November 2000, for NO₂ at 22 m (squares), NO at 29 m (circles), and O₃ × 10⁻¹ at 29 m (dashed line, shown for reference). For NO and NO₂, symbols are plotted at even and odd hours, respectively (lines go through hourly points), and vertical bars indicate 25th and 75th percentiles.

to the factory, and the C₂H₄-chemiluminescence instrument calibration was checked by routine comparison to the Dasibi 1003AH.

2.4. Micrometeorology and Auxiliary Measurements

[10] Triaxial sonic anemometers were oriented westward at the eddy covariance sampling height on each tower; data for vertical and horizontal wind velocities and virtual temperatures were acquired at 10 Hz to compute eddy covariance fluxes of heat, momentum, NO, and NO₂, and at 5 Hz for O₃ [Munger *et al.*, 1996, 1998]. Photosynthetic photon flux density (PPFD, 400–700 nm) was measured continuously at 29 m on the main tower using a LI-COR quantum sensor.

2.5. Eddy Covariance Fluxes

[11] We computed 30-min mean fluxes of trace gases from the covariance of linearly detrended time series for vertical velocity (w') and trace gas concentrations (C'). To correct for relatively slow instrument response times, data for virtual temperature and w were processed using instantaneous measurements by the sonic anemometer, then repeated using a set of single-pole low-pass numerical filters matched with the time response of each gas sensor. The ratio of the “filtered” heat flux for each 30-min interval to the unfiltered heat flux provided an estimate of flux lost by instrumental smoothing of high-frequency fluctuations [Goulden *et al.*, 1996; Munger *et al.*, 1996, 1998]. Corrections were typically 20% or smaller for all chemical species.

3. Results and Discussion

[12] Figure 1 shows median values at each hour of the day for NO, NO₂, and O₃ concentrations (top panels) and fluxes (bottom panels) for April through November 2000, segregated by wind sector. Photochemical production of NO during the day and conversion to NO₂ at night are apparent

(top panels). Concentrations of O₃ are typically much greater than NO_x, thus chemical reactions do not significantly perturb the vertical O₃ gradients and fluxes set by mixing and surface deposition [Munger *et al.*, 1996].

3.1. Nighttime Fluxes

[13] We observed downward fluxes of NO₂ during all measurement seasons in 2000 (spring, summer, and fall) at night, for both clean (NW) and polluted (SW) wind sectors (Figure 1, bottom panels). Drainage flows at the surface in stable nighttime conditions make it difficult to quantify ecosystem exchange of respired CO₂ at night by eddy covariance [Staebler and Fitzjarrald, 2004]. However, the small turbulent fluxes of depositing species observed by eddy covariance are likely an accurate measure of their vertical exchange. When momentum is not being transported downward, neither are scalars.

[14] The downward nighttime NO₂ flux increases with NO₂ concentration (Figure 2). The polynomial regression function $F_{NO_2} = V_0[NO_2] + a[NO_2]^2$ applied to hourly nighttime data yields $R^2 = 0.60$, $V_0 = -0.08 \pm 0.03$ ($p = 0.01$), and $a = -0.013 \pm 0.001$ ($p < 0.0001$). The quadratic dependence in the regression is largely driven by the two nights during the sampling period when the median concentration of NO₂ exceeded 10 nmol mol⁻¹. Excluding these nights, neither the overall fit nor the coefficient a is statistically significant. However, even for concentrations below 10 nmol mol⁻¹, the coefficient V_0 is negative and significant ($V_0 = -0.16 \pm 0.06$, $p = 0.005$). The regression suggests that the nighttime deposition velocity of NO₂ ($V_d(NO_2) = -F_{NO_2}/[NO_2]$) increases from approximately

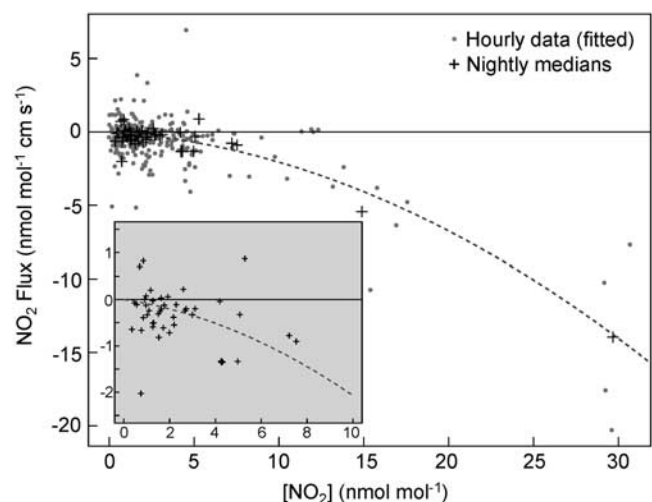


Figure 2. Nighttime hourly (dots) and median nightly (pluses) NO₂ flux versus concentration. The curve represents the function $F_{NO_2} = F_0 + V_0[NO_2] + a[NO_2]^2$, with coefficients derived from a least squares fit to the hourly data. The flux is expressed in units of concentration times velocity (nmol mol⁻¹ cm s⁻¹) in order to simplify the interpretation of the coefficients in the least squares fit. Pressure and temperature corrections have been taken into account in the conversion from density to mixing ratio flux units. The inset shows an expansion of both x and y axes with nightly medians only. See color version of this figure in the HTML.

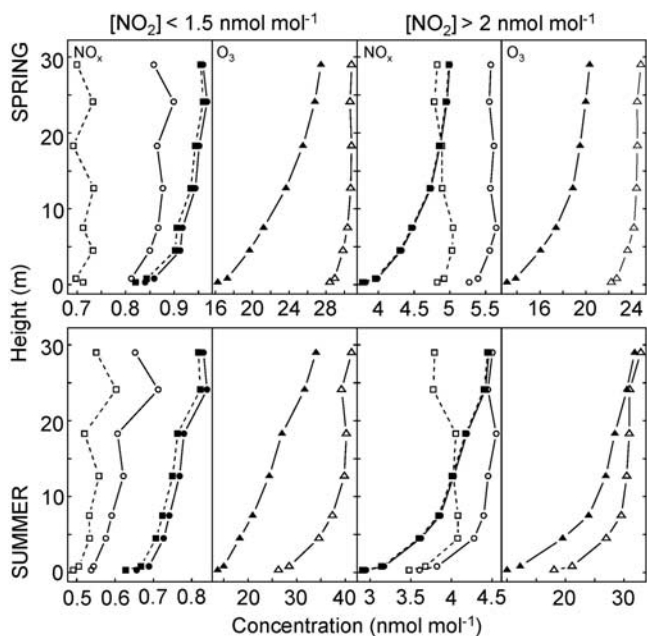


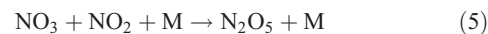
Figure 3. Mean vertical profiles of NO₂ (squares, dashed lines), NO_x = NO₂ + NO (circles, solid lines), and O₃ (triangles, solid lines) during the day (open symbols) and night (solid symbols) on the main Harvard Forest tower in 2000. (top) April and May data. (bottom) Data from June, July, and August. (September–November profiles are unavailable for this year.) (left) Limited to [NO₂] < 1.5 nmol mol⁻¹. (right) Limited to [NO₂] > 2 nmol mol⁻¹. Note the clear influence of forest canopy development from spring to summer on the NO₂ profile at higher [NO₂] and the masking of this effect at low [NO₂] by apparent canopy-level NO₂ emission. The year 2000 profiles are representative of those observed between 1990 and 2002.

0.2 cm s⁻¹ at [NO₂] = 1 nmol mol⁻¹ to 0.5 cm s⁻¹ at [NO₂] = 30 nmol mol⁻¹. This NO₂ deposition is equal to 6–20% of nighttime NO_y flux observed by eddy covariance at the site [Horii, 2002].

[15] The nighttime deposition of NO₂ in the canopy is confirmed by observations of NO, NO₂, and O₃ concentration profiles at the main tower (Figure 3). Concentrations of NO above the canopy at night are very small, often below the instrumental detection limit and indistinguishable from zero. Nighttime [NO_x] profiles exhibit a deposition signature similar to O₃, with dC/dz > 0, notwithstanding the small soil emissions of NO at the site of ≤ 0.9 μmol m⁻² h⁻¹ [Munger et al., 1996]. We compared the NO_x and O₃ fluxes by inferring average eddy diffusivity, K, derived from the flux-gradient relationship for each species: $F = -K \, dC/dz$, using the measured eddy covariance fluxes of NO₂ and O₃ above the canopy and the average concentration gradient between 5 m and 24 m. We found nighttime values of K ≈ 2400 cm² s⁻¹ with remarkable consistency: K_{NO_x}/K_{O₃} varied between 0.5 and 1.5 with a mean value ~1 independent of season (April–August) and NO_x concentration.

[16] Evidence for deposition to surfaces such as leaves, litter, bark, and soil has been reported previously by Eugster and Hesterberg [1996] and Hanson and Lindberg [1991, and references therein]. Hydrolysis of N₂O₅ on forest

surfaces below the sensor height is a possible mechanism (reactions (4)–(6)). The rate-limiting step is formation of NO₃ in reaction (4) [e.g., Jacob, 2000]:



We estimated an upper limit for NO₂ deposition due to this process by calculating production of NO₃ in reaction (4) from data for O₃ and NO₂ (Figure 3) and assuming 100% efficiency for reactions (5) and (6) and concluded that hydrolysis of N₂O₅ accounts for at most a minor fraction (<30%) of the observed nighttime NO₂ deposition at all observed concentrations.

[17] Nighttime deposition of NO₂ (and the possible non-linear dependence of flux on concentration) could also be explained by heterogeneous conversion to HONO and HNO₃, as observed on aerosols [Notholt et al., 1992]. The mechanisms and kinetics of the associated NO₂ reactions on hydrated surfaces are not fully understood. Heterogeneous NO₂ hydrolysis likely proceeds by the overall reaction



[Goodman et al., 1999]. Laboratory studies have shown first-order kinetics in NO₂ for reaction (7) at μmol mol⁻¹ (parts per million) concentrations, with NO₂ adsorption as the rate-limiting step, but the rate at low concentrations on forest surfaces is unknown. The atmospheric mechanism is likely to involve formation of N₂O₄ as an intermediate on the surface [Finlayson-Pitts et al., 2003]. As in the N₂O₅ hydrolysis mechanism, the aqueous phase HNO₃ produced in reaction (7) is not likely to be released to the gas phase, but HONO might be returned to the atmosphere.

[18] Harrison et al. [1996] observed upward HONO fluxes over vegetated surfaces at NO₂ concentrations above 10 nmol mol⁻¹, with a quadratic dependence of [HONO] on [NO₂]; likewise, HONO measurements by Thornberry et al. [2001] are consistent with heterogeneous production from NO₂. If the observed NO₂ flux at Harvard Forest is the result of reaction (7), then approximately half could return to the atmosphere as gas phase HONO. Since HONO is rapidly photolyzed to NO and OH upon sunrise, up to half of the nighttime NO₂ deposition would therefore not contribute to daily net loss of NO_x from the atmosphere. The possible heterogeneous HONO formation at the surface during the night is important to daytime HO_x chemistry, delivering a burst of OH and NO radicals to the surface layer at sunrise when HONO photolyzes.

3.2. Daytime Fluxes

[19] During daylight hours we observed large compensating fluxes, upward for NO₂ and downward for NO (Figure 1); these are largely due to the influence of the forest canopy on the vertical NO₂ photolysis gradient (reaction (3)) [Gao et al., 1993]. Higher irradiance above the canopy favors production of NO (reaction (2)); lower light below

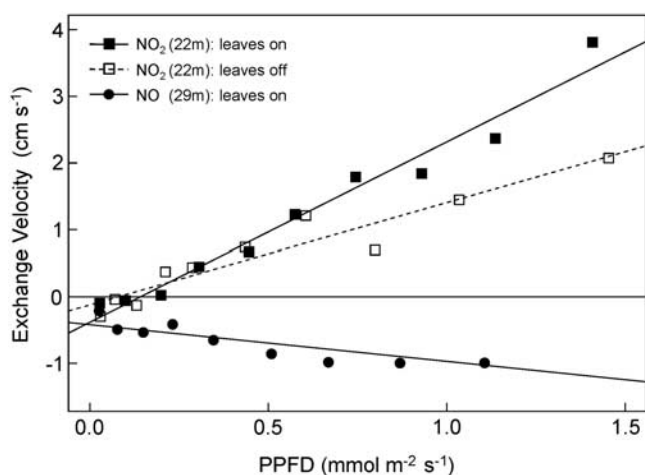


Figure 4. Daytime emission velocities of NO₂ (squares) and NO (circles) versus photosynthetic photon flux density (PPFD). Symbols are medians of hourly data binned by percentiles of above-canopy PPFD. NO₂ flux data are segregated into full-canopy conditions from early June to mid October (solid symbols), and leafless periods in April–May and October–November (open symbols). Linear fits to binned median points are $V_{NO_2}(\text{leaves on}) = (-0.4 \pm 0.1) + (2.7 \pm 0.2) \text{ PPFD}$, $R^2 = 0.97$; $V_{NO_2}(\text{leaves off}) = (0.1 \pm 0.1) + (1.5 \pm 0.2) \text{ PPFD}$, $R^2 = 0.90$; $V_{NO}(\text{leaves on}) = (-0.42 \pm 0.07) + (-0.5 \pm 0.1) \text{ PPFD}$, $R^2 = 0.78$. Midsummer leaf area index at Harvard Forest is ~ 3.4 , whereas the area index for bare stems and twigs is only 0.9 (Harvard Forest Online Data Archive, available at <http://www-as.harvard.edu/chemistry/index.html>).

the canopy shifts the balance toward NO₂ (reactions (1a) and (1b)). The magnitude of the daytime upward flux of NO₂ in Figure 1 is greater than the corresponding downward flux of NO, an artifact of the different sampling heights for NO (29 m on the main tower) and the TDLAS NO₂ (22 m on the auxiliary tower). Gao *et al.* [1993] showed in a modeling study that the fluxes of NO and NO₂ just above the canopy should be very sensitive to height.

[20] The eddy covariance fluxes of NO and NO₂ are correlated on an hourly basis during the daytime: An orthogonal distance fit yields $F_{NO_2} = -(4.0 \pm 0.1)F_{NO}$, $R^2 = 0.80$. The exchange velocities for NO and NO₂ follow light levels as measured by PPFD (Figure 4), as expected for vertical gradients associated with photolysis (Figure 3). There is a seasonal change in the dependence of daytime NO₂ flux on PPFD, correlated with canopy development, with a lower slope of the NO₂ exchange velocity versus PPFD for leafless periods. This result is also consistent with the expected effect of a smaller above- to below-canopy light gradient under leafless conditions, although it does not rule out a chemical or biological role in NO₂ exchange for the leaves. Since PPFD is a commonly measured and modeled quantity, we make use of this quantity as a proxy for photochemical activity in a simple parameterization of NO_x flux at our site.

3.3. Parameterization of NO_x Fluxes

[21] Global 2-D and 3-D models often parameterize NO₂ flux using canopy resistances assumed similar to O₃ [e.g.,

Wang *et al.*, 1998; Wesely and Hicks, 2000]. Although leaf-level measurements indicate a high degree of stomatal control over NO₂ flux similar to the stomatal uptake of O₃, the dearth of canopy-level NO₂ flux measurements calls this assumption into question. Recent results also raise doubts about the dominance of stomatal control of O₃ deposition to a forest canopy in the presence of biogenic hydrocarbons [Kurpius and Goldstein, 2003]. We therefore seek to assess the processes responsible for the observed NO_x surface fluxes independent of assumptions of similarity to O₃. In the following parameterization, we adopt the hypothesis that photochemical NO–NO₂ cycling via reactions (1)–(3) arising from canopy-induced light gradients has no effect on net deposition and represent it in our data set by a simple light-dependent term. We then use a multiple regression to determine the most likely contributions of other processes to the net flux.

[22] We represent fluxes of NO₂ by the sum of the following terms: $V_0 \cdot [\text{NO}_2]$, a constant exchange velocity term; $a \cdot [\text{NO}_2]^2$, a concentration-dependent exchange velocity term; $(b + f) \cdot \text{PPFD} \cdot [\text{NO}_2]$, a light-dependent photochemical cycling term allowing for differences between full-canopy and leafless conditions; and $\gamma \cdot g_c$, a term representing exchange processes under stomatal control. Stomatal conductance (g_c) (calculated using the measured water vapor flux and the Penman-Monteith equation after Shuttleworth *et al.* [1984]) correlates with PPFD, but the diel variations of the two differ in terms of morning/afternoon asymmetry and at dawn and dusk, making it possible to distinguish stomatally controlled from simple light-dependent processes, if both are strong enough. Thus we parameterize the flux of NO₂, 2 m above the canopy, day and night, by the function

$$F_{NO_2} = V_0 \cdot [\text{NO}_2] + a \cdot [\text{NO}_2]^2 + (b + f) \cdot \text{PPFD} \cdot [\text{NO}_2] + \gamma \cdot g_c \quad (8)$$

where f is added to the coefficient of PPFD to account for the observed difference between full-canopy and leaves-off conditions. The regression was run to determine the coefficients V_0 , a , b , and f using observed F_{NO_2} and $[\text{NO}_2]$ on the full data set as well as various subsets of data, including high-NO₂, low-NO₂, daytime, and nighttime conditions (Table 1). The values of the coefficients are consistent whether they are computed from the full data set or the selected subsets. However, for low-NO₂ conditions some coefficients have low statistical significance. The coefficient of the stomatal term, γ , is significant in most of the fits and follows the expected behavior for a stomatal process with a compensation point. The $\gamma \cdot g_c$ term appears to contribute a downward component of NO₂ flux when NO₂ is high ($>2 \text{ nmol mol}^{-1}$) and an upward flux when NO₂ is low ($<1.5 \text{ nmol mol}^{-1}$). Low-NO_x conditions dominate at the site [Munger *et al.*, 1996].

[23] Fluxes of NO from soils at our site are measurable but small. The major contributor to FNO during the day is the simple dependence on PPFD. We include a constant exchange velocity term ($V_{0_NO} \cdot [\text{NO}]$) and a light-dependent term ($b' \cdot \text{PPFD} \cdot [\text{NO}]$), giving the NO flux 7 m above the canopy as

$$F_{NO} = V_{0_NO} \cdot [\text{NO}] + b' \cdot \text{PPFD} \cdot [\text{NO}]. \quad (9)$$

Table 1. Multifactor Regressions of NO₂ and NO Flux Data Using All Available Hourly Observations of [NO₂], PPFD, FNO₂, and FNO and Calculated Stomatal Conductance, g_c, at Harvard Forest, April–November 2000^a

	All Data (R ² = 0.60, N = 928)	[NO ₂] < 1.5, nmol mol ⁻¹ (R ² = 0.15, N = 464)	[NO ₂] > 2, nmol mol ⁻¹ (R ² = 0.75, N = 306)	Day Only (R ² = 0.63, N = 412)	Night Only (R ² = 0.60, N = 316)
V ₀	-0.14 ± 0.03	-0.3 ± 0.4	-0.12 ± 0.04	-0.39 ± 0.09	-0.08 ± 0.03
a	-0.009 ± 0.002	0.2 ± 0.3	-0.010 ± 0.002	0.008 ± 0.009	-0.013 ± 0.001
b	2.23 ± 0.07	1.7 ± 0.4	2.4 ± 0.1	2.3 ± 0.1	NA
f	-1.32 ± 0.09	-1.2 ± 0.5	-1.4 ± 0.1	-1.3 ± 0.1	NA
γ	0.2 ± 0.2	0.8 ± 0.4	-1.6 ± 0.8	0.5 ± 0.3	NA

^aFor z = 22 m: FNO₂ = V₀ · [NO₂] + a · [NO₂]² + (b + f) · PPFD · [NO₂] + γ · g_c. Day-only and night-only regressions exclude dawn and dusk. Note the overall consistency of direction and magnitude of statistically significant terms, with the exception of the coefficient of g_c, which changes sign between high- and low-NO₂ conditions. Terms with p > 0.1 are italicized. For each term, units are as follows: V₀, cm s⁻¹; a, (nmol mol⁻¹)⁻¹ cm s⁻¹; b, cm s⁻¹ (μmol m⁻² s⁻¹)⁻¹; f, cm s⁻¹ (μmol m⁻² s⁻¹)⁻¹, where f = 0 for leaves-on conditions; and γ, nmol mol⁻¹.

The resulting parameterization indicates a small deposition velocity, with large fluxes due to photochemical cycling, using all available data (956 hours), R² = 0.92, V_{0-NO} = -0.25 ± 0.02 cm s⁻¹, and b' = -0.87 ± 0.03 cm s⁻¹ (μmol m⁻² s⁻¹)⁻¹.

3.4. Analysis of Net NO_x Flux

[24] Because net NO flux (soils and deposition) appears small, and the purely gas phase photochemical contributions to NO and NO₂ fluxes arising from canopy light gradients should cancel when considering net ecosystem flux, we estimate the net flux of NO_x from the FNO₂ parameterization (8) by omitting the photochemical cycling components:

$$\begin{aligned} \text{FNO}_x(\text{net}) &\approx \text{FNO}_2 - (b + f) \cdot \text{PPFD} \cdot [\text{NO}_2] \\ &= V_0 \cdot [\text{NO}_2] + a \cdot [\text{NO}_2]^2 + \gamma \cdot g_c. \end{aligned} \quad (10)$$

Figure 5 illustrates the relative contributions of each term to net flux and the resulting net NO₂ exchange velocity, for all conditions and for high-NO₂ periods, using the coefficients from Table 1, measured NO₂ concentrations, and calculated g_c.

[25] When NO₂ concentrations are low during the day, some NO₂ is emitted under stomatal control but is offset by NO₂ deposition to other surfaces. When NO₂ concentrations are high during the day, NO₂ is deposited under stomatal control, as much as doubling the rate of NO₂ deposition. For most conditions at Harvard Forest, nonstomatal deposition accounts for the largest fraction of the flux, during the night as well as the day. *Lerdau et al.* [2000] noted that leaf-level observations of NO₂ emission at low ambient concentrations were inconsistent with current understanding of NO_x chemistry and observations in the remote troposphere. The overwhelming impact of nonstomatal deposition at the canopy level appears to provide a resolution to this conundrum.

[26] At Harvard Forest the daytime net NO_x flux was ~2% of total daytime NO_y deposition during the summer [Horii, 2002]. The lifetime of NO_x associated with deposition, on the basis of an average net NO_x deposition velocity of 0.2 cm s⁻¹ and regionally representative mixed layer depths [Holzworth, 1967], varies between 10 days in summer and 4 days in winter, assuming surface deposition continues throughout the year as it does from early spring through late fall. This lifetime may decrease to less than 24 hours during shallow nighttime inversions. In comparison, the characteristic time for NO₂ oxidation by OH varies from less than 12 hours in summer to more than 5 days in

winter [Munger *et al.*, 1998]. Evidently, direct NO_x deposition to forest surfaces has the largest impact on tropospheric chemistry when mixing layer depths are shallow and when other oxidative processes are relatively slow.

[27] The daytime NO_x and O₃ profiles for April–August 2000 (Figure 3) are qualitatively consistent with the independent eddy covariance flux measurements and net NO_x flux analysis. Under low-NO_x conditions, profiles have a weak gradient, implying a net NO_x flux approaching zero, with nonstomatal deposition to the ground offset by apparent emission in the upper canopy; the persistent NO_x enhancement at 24 m may correspond to stomatal or other UV-induced NO₂ emission reported from vegetation at ambient concentrations below 1 nmol mol⁻¹ [Hari *et al.*,

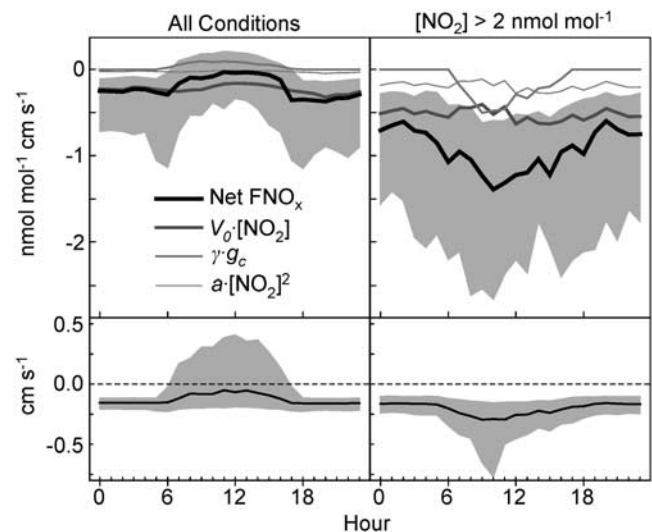


Figure 5. (top) Individual contributing terms (colored lines) and the net NO₂ flux (black line) based on the coefficients in Table 1, measured [NO₂], and calculated g_c. The shaded regions indicate reasonable bounds on the net flux: The upper bound is the 75th percentile of net FNO₂ calculated using the upper confidence limit of the coefficients; the lower bound is the 25th percentile calculated with the lower confidence limit of the coefficients. (bottom) Net NO_x deposition velocity with upper and lower bounds calculated as in the top panels. Panels on the left use all data and the corresponding coefficients from Table 1. Panels on the right use data hours when NO₂ > 2 nmol mol⁻¹ and high-NO₂ coefficients. See color version of this figure in the HTML.

2003]. The profiles indicate that net NO_x deposition is stronger at higher [NO_x] compared to lower [NO_x], and during the summer compared to the early spring. As discussed earlier, the nighttime profiles in Figure 3 also supported the results of the net NO_x flux analysis showing deposition during nondaylight hours.

4. Conclusion

[28] On the basis of both flux data and vertical profiles, we find that nighttime NO₂ flux at Harvard Forest is predominantly downward, including clean air conditions. The results contradict widely used parameterizations of NO₂ deposition (e.g., big-leaf, resistance-in-series models, following the Wesely [1989] scheme) that do not provide for surface uptake when stomates are closed. Because N₂O₅ hydrolysis is insufficient to account for the observed deposition flux (particularly at the highest NO₂ concentrations observed), HONO production may be important. Daytime fluxes of NO and NO₂ are strongly coupled and dominated by simple photochemical exchange that does not contribute to net NO_x flux. We conclude that NO₂ and NO fluxes can be parameterized in terms of known physical processes to derive a deposition velocity for NO₂ of ~0.2 cm s⁻¹. Stomatal influence is weaker than typically assumed in models, and it changes sign depending on NO₂ concentrations, with overall near zero deposition or emission in the daytime for NO_x less than ~1.5 nmol mol⁻¹, and significant stomatal uptake for high NO_x. These results identify a canopy scale deposition process that is not apparent in leaf-level measurements. They highlight the importance of including both nonstomatal deposition and a compensation point in NO₂ surface exchange parameterizations embedded in models of tropospheric chemistry and transport. Overall, we have found by direct measurements in field conditions that vegetation serves as a net sink for NO_x radicals, with the sink strength increasing as NO_x concentration rises. Contrary to previous assumptions, the sink persists at night and in the early spring and late fall dormant seasons. The persistent deposition due to nonstomatal processes provides a potential resolution to the problem that extensive NO₂ emission from foliage where NO₂ is below the compensation point would be inconsistent with our understanding of NO_x chemistry and with observations in the remote troposphere.

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