Origin of tropospheric NO\textsubscript{x} over subarctic eastern Canada in summer

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Abstract. The origin of NO\textsubscript{x} in the summertime troposphere over subarctic eastern Canada is investigated by photochemical modeling of aircraft and ground-based measurements from the Arctic Boundary Layer Expedition (ABLE 3B). It is found that decomposition of peroxyacetyl nitrate (PAN) can account for most of the NO\textsubscript{x} observed between the surface and 6.2 km altitude (aircraft ceiling). Forest fires represent the principal source of PAN in the region, implying the same origin for NO\textsubscript{x}. There is, however, evidence for an unidentified source of NO\textsubscript{x} in occasional air masses subsiding from the upper troposphere. Isoprene emissions from boreal forests maintain high NO\textsubscript{x} concentrations in the continental boundary layer over eastern Canada by scavenging OH and NO\textsubscript{2}, thus slowing down conversion of NO\textsubscript{2} to HNO\textsubscript{3}, both in the daytime and at night. This effect is partly compensated by the production of CH\textsubscript{3}CO\textsubscript{3} radicals during isoprene oxidation, which slows down the decomposition of PAN subsiding from the free troposphere. The peroxy radical concentrations estimated from concurrent measurements of NO and NO\textsubscript{2} concentrations during ABLE 3B are consistent with values computed from our photochemical model below 4 km, but model values are low at higher altitudes. The discrepancy may reflect either a missing radical source in the model or interferences in the NO\textsubscript{2} measurement.

1. Introduction

Nitrogen oxides (NO\textsubscript{x} = NO + NO\textsubscript{2}) control the tropospheric production of O\textsubscript{3} and OH. Knowledge of the processes that govern tropospheric NO\textsubscript{x} concentrations is essential to understanding the present, past and future oxidizing capacity of Earth's atmosphere [Isaksen and Hov, 1987; Crutzen and Zimmermann, 1991; Thompson, 1992]. Distributions of NO\textsubscript{x} in the global troposphere have been simulated using two or three-dimensional chemistry, transport, and deposition models and estimated inventories of NO\textsubscript{x} emissions [Levy and Moxim, 1987; Penner et al., 1991; Ehhalt et al., 1992; Kasibhatla et al., 1993]. However, the origin of NO\textsubscript{x} in the remote troposphere remains uncertain. Transport of primary NO\textsubscript{x} from source regions is limited, because NO\textsubscript{x} is oxidized to HNO\textsubscript{3} in a matter of a few days. It appears that diffuse chemical sources must be responsible for maintaining the NO\textsubscript{x} levels in the remote troposphere.

Peroxyacetyl nitrate (PAN, CH\textsubscript{3}C(O)OONO\textsubscript{2}), which is produced in the oxidation of hydrocarbons, represents an ubiquitous reservoir of NO\textsubscript{x} in the remote troposphere [Singh et al., 1986, 1990, 1992, 1994a]. Since PAN is stable at the low temperatures of the upper troposphere, it may provide a vehicle for the long-range transport of NO\textsubscript{x} from source regions [Crutzen, 1979; Singh and Hanst, 1981]. The importance of PAN decomposition as a source of NO\textsubscript{x} in the remote troposphere has been evaluated recently in photochemical model analyses of observations taken over western Alaska in summer (Arctic Boundary Layer Expedition (ABLE) 3A) and at Mauna Loa, Hawaii (Mauna Loa Observatory Photochemistry Experiment (MLOPEX)). Jacob et al. [1992] found that PAN decomposition in ABLE 3A could account for most of the NO\textsubscript{x} observed below 4 km but not above. Liu et al. [1992] and Walega et al. [1992] found that PAN decomposition could provide only a minor source of NO\textsubscript{x} in MLOPEX. The origin of NO\textsubscript{x} in MLOPEX remains unclear (R. B. Chatfield, The anomalous HNO\textsubscript{3}/NO\textsubscript{x} ratio of remote tropospheric air: Is there conversion of nitric acid to formic acid and NO\textsubscript{y}, submitted to Geophysical Research Letters, 1994).

We present here a photochemical investigation of the NO\textsubscript{x} budget in the troposphere over eastern Canada in summer, using data from the ABLE 3B expedition conducted in July-August, 1990 [Harriss et al., 1994]. This expedition offered a comprehensive documentation of air chemistry over the region including mixing ratios of O\textsubscript{3}, NO, NO\textsubscript{2}, PAN, NO\textsubscript{y}, CO, and hydrocarbons measured from aircraft up to 6.2 km altitude and biosphere-atmosphere exchange fluxes measured from an instrumented tower at a boreal woodland site near Schefferville, Quebec (see special section "The Northern Wetlands Study and the Arctic Boundary Layer Expedition 3B: An International and Interdisciplinary Field Campaign" in Journal of Geophysical Research, 99 (D1), 1421-1953, 1994). We analyze the origin of NO\textsubscript{x} in ABLE 3B by using a combination of zero-dimensional (0-D) and one-dimensional (1-D) models constrained with the observations (details presented in section 2). The 0-d model calculations are used for the free tro-
posphere, yielding an ensemble of snapshots of local photochemistry along the aircraft flight tracks from which regional statistics for the NO\textsubscript{2} budget can be obtained. The 1-D model calculations are used for the continental boundary layer (CBL) over the boreal woodland and account for diel variations in vertical mixing and biogenic isoprene emissions. The woodlands of eastern Canada are large sources of isoprene [Blake et al., 1994]. The ABLE 3B data provide a rare chance to study the chemistry of isoprene under the low NO\textsubscript{2} conditions which are characteristic of the CBL over remote regions.

The NO\textsubscript{2} budgets in the free troposphere and in the continental boundary layer are presented in sections 3 and 4, respectively. Conclusions are in section 5. The appendix provides discussion on the feasibility of using the concurrent measurements of NO and NO\textsubscript{2} in ABLE 3B to calculate the concentration of peroxy radicals and test the accuracy of photochemical models.

2. Methods

A map of the ABLE 3B region is shown in Figure 1. Air masses of various chemical compositions were encountered by the aircraft, reflecting influences from forest and tundra fires, industrial and urban pollution, stratospheric intrusions, and tropical outflow (Table 1). The ABLE 3B mission design placed particular emphasis on sampling biomass fire plumes. Air masses influenced by biomass fire emissions, as diagnosed by CO concentrations greater than 120 ppb [Talbot et al., 1994], accounted for about 30% of the aircraft observations in the free troposphere. Lidar measurements of aerosol concentrations in the 2 to 6 km column during the expedition suggest that air influenced by biomass burning occupied, on average, 13% of the free troposphere, background air, 42%, stratospherically influenced air, 35%, and other types, 12% [Browell et al., 1994].

We use a 0-D model to calculate the concentrations of radicals and other secondary species at chemical steady state in the free troposphere. The calculations are constrained with the ensemble of measurements taken aboard the aircraft including temperature, dew point, pressure, UV radiation fluxes (zenith and nadir), and concentrations of NO, PAN, HNO\textsubscript{3}, O\textsubscript{3}, CO, acetone, C\textsubscript{1-7} alkanes, C\textsubscript{2-3} alkenes, benzene, and toluene. Measurements of NO are more reliable than those of NO\textsubscript{2} owing to possible interference in the NO\textsubscript{2} measurement (see appendix). The calculations are conducted for 3-min averaging intervals in the aircraft observations, representing the time resolution of the NO\textsubscript{2} measurement. There are 165 intervals in the free troposphere (2.5-6.2 km) where concurrent measurements are available for all the above input variables except acetone. The data are sparse for acetone; missing data are filled based on the correlation between acetone and CO (Figure 2) (see also Singh et al. [1994b]).

The 0-D model is inadequate in the continental boundary layer (CBL) (below 2.5 km), where the concentrations of radicals are sensitive to the abundance of biogenic isoprene and its oxidation products [Jacob and Wofsy, 1990]. The lifetime of isoprene is only a few hours, while the carbonyls produced successively in the isoprene oxidation chain have lifetimes of a few hours to a few days. One cannot assume that the isoprene oxidation products are in local chemical steady state with the isoprene concentrations measured aboard the aircraft. We use therefore a time-dependent, 1-D model for the CBL, following Trainer et al. [1987, 1991] and Jacob and Wofsy [1988, 1990], and apply this model to simulate observations over the Schefferville tower site on August 7 when detailed measurements from the tower and from an aircraft spiral over the tower are available. The tower site is 0.5 km above sea level. The CBL in our 1-D model extends to 2.25 km above ground level, representing the afternoon maximum of mixed layer depth as measured locally from rawinsondes [Fitzjarrald and Moore, 1994]. Photochemical calculations are conducted at seven grid points (0.01, 0.1, 0.2, 0.5, 1.0, 1.5, and 2.0 km above the canopy). Vertical transport is simulated with an eddy diffusion parameterization based on local measurements of the mixed layer depth z\textsubscript{L} and of the fluxes of momentum and sensible heat [Lamb et al., 1975]. The eddy diffusion coefficient between z\textsubscript{L} and 2.25 km is adjusted to reproduce the observed vertical profiles of O\textsubscript{3} mixing ratio, resulting in a ventilation lifetime of 4.5 days for the CBL.

![Figure 1. Map of the Arctic Boundary Layer Experiment (ABLE) 3B study region. The hatched areas indicate regions of intensive aircraft flight. The tower site was located near Schefferville, Quebec.](image-url)
measurements and hourly ground measurements. The deposition relative humidity are specified based on interpolation of aircraft 3.0 km above sea level and the mean concentrations of peroxides concentrations of O₃, NO, NO₂, and PAN measured between 2.5-3.5 km above sea level. The diurnal variations of temperature and peroxides have the same deposition velocity as O₃. Soil emission spirals below 2 km. The did cycle of isoprene emission is specified according to Jacob et al. [1993], assuming a leaf area index of 2 and with the midday emission rate adjusted to match the isoprene concentrations measured at the tower, typically 0.33 cm s⁻¹ during the day and 0.03 cm s⁻¹ at night (J. W. Munger et al., manuscript in preparation, 1994). We assume that PAN and carbonyls calculated in the 0-D model under background con-

The CBL model uses as upper boundary conditions the mean concentrations of O₃, NO, NO₂, and PAN measured between 2.5-3.0 km above sea level and the mean concentrations of peroxides and carbonyls calculated in the 0-D model under background conditions. Vertical profiles of HNO₃, CO, and hydrocarbons other than isoprene are specified using observations from the aircraft spirals below 2 km. The diurnal variations of temperature and relative humidity are specified based on interpolation of aircraft measurements and hourly ground measurements. The deposition flux of O₃ to the surface is calculated on the basis of the hourly mean O₃ deposition velocities measured at the tower, typically 0.33 cm s⁻¹ during the day and 0.03 cm s⁻¹ at night (J. W. Munger et al., manuscript in preparation, 1994). We assume that PAN and peroxides have the same deposition velocity as O₃. Soil emission of NO and dry deposition of NO₂ were negligibly small [Bakwin et al., 1994]. The diel cycle of isoprene emission is specified according to Jacob et al. [1993], assuming a leaf area index of 2 and with the midday emission rate adjusted to match the isoprene concentrations measured in the CBL during the aircraft spiral. The resulting 24-hour average emission flux of isoprene is 6.1x10¹⁰ molecules cm⁻² s⁻¹. In comparison, Klinger et al. [1994] estimated a mean isoprene emission rate of 5.5x10¹⁰ molecules cm⁻² s⁻¹ along a successional gradient in the Hudson Bay low-
lands.

The photochemical model includes HO₂-NO₂-hydrocarbon chemistry, based on recent compilations of kinetic and product data [Atkinson, 1990; Atkinson et al., 1992; DeMore et al., 1992]. The mechanism for photochemical decomposition of isoprene follows Tuazon and Atkinson [1989, 1990a, b]. Nighttime isoprene chemistry follows Paulson and Seinfeld [1992]. Cross sections for photolysis of PAN and the rate for PAN oxidation by OH are from recent measurements (R. K. Talukdar et al., manuscript in prepara-

Table 1. Mean Characteristics of Air Masses Observed in the Free Troposphere Over Eastern Canada in Summer 1990

| Type 1, Altitude, km | Type 2, Altitude, km | Type 3, Altitude, km | Type 4, Altitude, km |
|---------------------|---------------------|---------------------|---------------------|
| 2.5-3.5             | 3.5-4.5             | 4.5-6.2             | 2.5-3.5             |
| 3.5-4.5             | 4.5-6.2             | 2.5-3.5             |                     |
| 4.5-6.2             | 2.5-3.5             |                     |                     |
| n                   | 24                  | 17                  | 3                   |
| T, °C               | -3                  | -6                  | 2                   |
| DPT, °C             | -8                  | -17                 | -3                  |
| NO                  | 6                   | 8                   | 13                  |
| NO₂                 | 20                  | 29                  | 39                  |
| PAN                 | 190                 | 200                 | 480                 |
| HNO₃                | 41                  | 57                  | 200                 |
| NO₄                 | 260                 | 510                 | 900                 |
| O₃                  | 46                  | 54                  | 58                  |
| CO                  | 100                 | 99                  | 150                 |
| Ethyne              | 80                  | 82                  | 240                 |
| Ethene              | 24                  | 24                  | 130                 |
| Ethane              | 730                 | 780                 | 1100                |
| Propane             | 59                  | 79                  | 200                 |
| Butane              | 10                  | 15                  | 74                  |
| Benzene             | 35                  | 34                  | 99                  |
| Acetone             | 1200                | 1100                | 2100                |
| ΔNOₓ, %             | 2                   | 40                  | 18                  |

Air mass types are defined following Talbot et al. [1994] as 1, Regional background; 2, biomass burning influence; 3, tropical outflow; and 4, stratospheric influence. The mean characteristics are computed from aircraft observations for the 165, 3-min intervals used in our photochemical modeling calculations. Volume mixing ratios are in parts per trillion (ppt) except for CO, O₃ in parts per billion. Unlisted species were generally near or below their detection limits; 10 ppt propene, 5 ppt toluene, and 2 ppt for >C₄ alkanes. Abbreviations are n, number of observations; T, temperature; DPT, dew point; PAN, peroxyacetyl nitrate.

ΔNOₓ is the percent of NOₓ not accounted for by observations of NOₓ, PAN, and HNO₃; negative values indicate that the sum of concentrations of these species exceeded the observed concentration of NOₓ. For further details on the NOₓ mass balance, see Sandholm et al. [1994].

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In the 0-D model the NO₂ photolysis rate coefficient J_NO₂ is obtained from the UV measurements by zenith and nadir Eppler radiometers [Madronich, 1987; Chameides et al., 1990]. Photolysis rate coefficients for other species are calculated using a six-stream radiative transfer model for the clear sky, Rayleigh-scattering atmosphere [Logan et al., 1981] and are scaled by the ratio of J_NO₂ calculated with the model to J_NO₂ derived from the UV measurements. In these calculations, surface albedo is fixed at 0.06, a value obtained from extrapolation of Eppler measurements to ground level. The overhead ozone column is taken from daily satellite measurements at a resolution of 1° latitude by 1° longitude (total ozone mapping spectrometer) and ranges from 290 to 360 Dobson units. The ratio of J_NO₂ obtained from Eppler mea-

Air masses are defined using observations of HNO₃, CO, and the ratio of CO₂:NO found in the background air [Heikes and Thompson, 1983; Dentener and Crutzen, 1993]. However, the NH₄²SO₄ equivalent ratio in ABLE 3B averaged 1.4 for background air [Gorzelak et al., 1994], implying that sulfate aerosol would be present as solid (NH₄)₂SO₄ [Tang et al., 1978]. Conversion of N₂O₅ to HNO₃ on solid aerosols is negligi-

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The CBL model uses as upper boundary conditions the mean concentrations of O₃, NO, NO₂, and PAN measured between 2.5-3.0 km above sea level and the mean concentrations of peroxides and carbonyls calculated in the 0-D model under background conditions. Vertical profiles of HNO₃, CO, and hydrocarbons other than isoprene are specified using observations from the aircraft spirals below 2 km. The diurnal variations of temperature and relative humidity are specified based on interpolation of aircraft measurements and hourly ground measurements. The deposition flux of O₃ to the surface is calculated on the basis of the hourly mean O₃ deposition velocities measured at the tower, typically 0.33 cm s⁻¹ during the day and 0.03 cm s⁻¹ at night (J. W. Munger et al., manuscript in preparation, 1994). We assume that PAN and peroxides have the same deposition velocity as O₃. Soil emission of NO and dry deposition of NO₂ were negligibly small [Bakwin et al., 1994]. The diel cycle of isoprene emission is specified according to Jacob et al. [1993], assuming a leaf area index of 2 and with the midday emission rate adjusted to match the isoprene concentrations measured in the CBL during the aircraft spiral. The resulting 24-hour average emission flux of isoprene is 6.1x10¹⁰ molecules cm⁻² s⁻¹. In comparison, Klinger et al. [1994] estimated a mean isoprene emission rate of 5.5x10¹⁰ molecules cm⁻² s⁻¹ along a successional gradient in the Hudson Bay low-
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Figure 2. Relationship of acetone with CO. The line represents a least squares fit through the data. Mixing ratios of CO were averaged to measurement intervals of acetone. Data include all acetone measurements over eastern Canada during ABLE 3B.

Figure 3. Sources and sinks of NOx in the free troposphere (2.5-6.2 km) during ABLE 3B. Values are daytime means obtained by averaging the rates (parts per trillion per hour) computed for individual 3-min intervals, first according to time of day (2-hour bins, with morning/afternoon folding) and then over 12 hours from 0600 to 1800 LT. The data were segregated by air mass type (Table 1). The number of data points used in the computation are, respectively, 165 for all data, 82 for background air, 33 for stratospheric influence, 16 for tropical outflow, and 34 for biomass fire influence. The top abscissa scale is for the case of biomass burning influence, and the bottom abscissa scale is for all other cases.

3. NOx in the Free Troposphere

Mean production and loss rates of NOx in the free troposphere during daytime are shown in Figure 3 for individual air mass types. The reaction rates calculated for each 3-min interval were first averaged according to time of day (2-hour bins, with morning/afternoon folding to overcome a lack of measurements in the early morning) and then averaged over daytime hours (0600-1800 LT). Nighttime chemistry is assumed negligible in the free troposphere for reasons discussed above. We find that sources and sinks of NOx are in close balance in background air (Table 1, type 1). Decomposition of PAN represents the largest source for NOx, and therefore can explain most of the observed NOx during ABLE 3B. Photoreduction of HNO3 to NO is negligibly slow.

The effect of uncertainties in measured concentrations and rate constants must be considered in our NOx mass balance. The rates for PAN formation and thermal decomposition are estimated to have uncertainties of 20% at 298 K [Atkinson et al., 1992]. The precision and accuracy of the PAN measurement are estimated to be 10% and 25%, respectively, of PAN mixing ratios [Singh et al., 1994a]. The principal sources of the CH3CO2 precursor to PAN in the model are photolysis of acetone, oxidation of acetaldehyde, and decomposition of PAN. The error caused by inferring acetone concentrations from observations of CO can be estimated to be 0.25 ppb (Figure 2), which would cause a mean error of 0.2 parts per trillion per hour (hereafter, ppt h⁻¹) in the computation of PAN production. Although the above uncertainties are substantial, they are not so large as to affect our central conclusion that PAN decomposition can account for most of the NOx observed during ABLE 3B.

Forest and tundra fires were found to be the major source for PAN, acetone, and other nonmethane hydrocarbons during ABLE 3B [Singh et al., 1994b; Talbot et al., 1994; Wofsy et al., 1994]. Long-range transport of midlatitude pollution was found to be of secondary importance. Nitrogen oxides emitted from biomass burning are efficiently converted to PAN in the fire plumes, because of the abundance of reactive hydrocarbons [Jacob et al., 1992], and additional PAN is formed on the regional scale following photolysis of pyrogenic acetone.

Air masses sensibly influenced by biomass burning in ABLE 3B (Table 1, type 2) showed indication of rapid decomposition of
PAN (Figure 3). The resulting net source for NOx in Figure 3 is 10 ppt h$^{-1}$, but the mean concentration of NOx in these air masses was only 40-50 parts per trillion (hereafter, ppt). Reconciling the model NOx budgets with the observed NOx concentrations would require rapid dilution of the air masses with the regional background. However, an underestimate of PAN formation is also possible, because many primary and secondary species, particularly oxygenated hydrocarbons, may be present in type 2 air masses but not included in the model. Such species could decompose to yield CH$_3$CO$_2$ radicals and promote PAN formation.

Tropical air masses originating from the Pacific Ocean (Table 1, type 3) showed a near balance between chemical sources and sinks of NOx (Figure 3). However, the total NOx was exceeded by the sum of individual NOy species in these air masses. Reasons for the abnormal NOx composition are unclear; measurement errors could not be ruled out, though it is unlikely that possible errors can account for all of missing NOx [Sandholm et al., 1994].

Air masses subsiding from the upper troposphere (Table 1, type 4) represent the only case where decomposition of PAN was insufficient to account for the observed NOx. One possible explanation for the missing NOx source is decomposition of unidentified nitrogen compounds. Nearly 50% of total NOx in type 4 air masses was unaccounted for by measurements of NOx, PAN, and HNO3 (Table 1). Figure 4 shows the relation between the net loss of NOx computed in the model, equivalent to a missing source of NOx, and the NOx deficit in air of type 4. The missing source of NOx appears to increase with the NOx deficit. The NOx budget would be balanced if the missing species representing the NOx deficit were converted to NOx in the free troposphere with chemical lifetime of a few weeks. A remarkable feature of air masses of type 4 was the low mixing ratio of HNO3 (42 ppt on average). Such a concentration would be obtained in about 3 days from oxidation of NOx (Figure 3). Reduction of HNO3 to NOx on a timescale of 3 days would balance the NOx budget (Figure 3), but the chemical lifetime of HNO3 against photolysis and reaction with OH is about 2 weeks.

One way to explain the NOx and HNO3 budgets as well as the missing NOx species in air masses of type 4 would be by reaction of CH$_2$O with HNO3 on concentrated sulfuric acid aerosols, producing methyleneglycol nitrates (HOCH$_2$ONO$_2$, CH$_2$(ONO$_2$)$_2$) [Travaglini, 1938]. Methyleneglycol nitrates have low solubility and would volatilize from the aerosols to the gas phase. They are likely photodissociated to release NOx, by analogy with methyl nitrate; they would have a lifetime on the order of a few weeks. Sandholm et al. [1994] found the observed NOx deficit species to increase with O3 concentration (or altitude) and to increase with the photochemical age of the air mass as measured by the CO/C$_2$H$_4$ ratio. These observations seem consistent with possible formation of HOCH$_2$ONO$_2$ and CH$_2$(ONO$_2$)$_2$ in the upper troposphere.

A comparison can be made between the NOx budgets in ABLE 3B and those in ABLE 3A. Free troposphere concentrations of NOx, PAN, and O$_3$ were similar in both expeditions [Sandholm et al., 1992, 1994; Singh et al., 1992, 1994a] temperatures were also similar. Therefore we expect PAN decomposition to also have accounted for most of the NOx in ABLE 3A. In their modeling of ABLE 3A data, Jacob et al. [1992] argued that PAN decomposition could account for the NOx below about 4 km but not at higher altitudes. They used a PAN decomposition rate constant taken from Lurmann et al. [1986] that is about 30% lower than used here [Atkinson et al., 1992].

4. NOx in the Continental Boundary Layer

We now turn to an analysis of the origin of NOx in the continental boundary layer (CBL) over the boreal woodland at Schefferville using the 1-D model for August 7 described in section 2. We address the following questions. (1) Does decomposition of PAN subsiding from the free troposphere account for NOx in the CBL? (2) How does vegetation emission of isoprene affect NOx and PAN? (3) How sensitive is the NOx budget in the CBL to the RO$_2$ + HO$_2$ reactions? The last question is motivated by the lack of kinetic data for the reactions of organic peroxy radicals (RO$_2$) arising from photochemical oxidation of isoprene. Reactions of these peroxy radicals with each other are probably slow [Madrornich and Calvert, 1990], but reactions with NO could be rapid [Atkinson, 1990]. The reaction products are assumed to be organic peroxides (ROOH) which may photolyze, react with OH, or be removed by deposition. The latter two sinks would represent real loss of radicals from the atmosphere.

The model NOy deficit is the difference between the measured NOx and the sum of NOx, peroxyacetyl nitrate (PAN), and HNO3 measured simultaneously. The line represents a least squares fit (slope = 0.00154).
Figure 5 shows aircraft measurements of ambient temperature, absolute humidity, and mixing ratios of CO over the tower site on the afternoon of August 7. The air mass below 2 km originated from the Hudson Bay region 5 days prior to aircraft measurements and was not modified by rain or combustion emissions during the transit period [Shipham et al., 1994]. The composition of that air mass is typical of the regional background (Table 1). Tropical influence from the Pacific is apparent above 2 km and is manifested in Figure 5 by the low mixing ratios of CO (<80 ppb). This tropical influence was transitory [Shipham et al., 1994]; therefore we assume that the boundary layer had been in contact with a free troposphere of background composition in the few days before being overridden by the tropical air. This assumption dictates our choice of upper boundary conditions (section 2).

Figure 6 shows the comparisons of model mixing ratios of isoprene, O$_3$, NO, NO$_2$, and PAN with observations. The model simulates NO, NO$_2$, and PAN within the measurement uncertainties. The vertical distribution of isoprene is controlled by turbulent mixing and by OH oxidation in the boundary layer (the isoprene emission flux was adjusted to match the observed isoprene concentrations.) Concentrations of O$_3$ are controlled mainly by transport from the free troposphere and deposition. The net photochemical production of O$_3$ in the CBL is small, about 20% of the flux from the free troposphere.

The sensitivity simulation without isoprene emission yields O$_3$, NO, and NO$_2$ within the range of measurements. However, the PAN mixing ratio in the CBL falls below the measurement by a factor of more than 2. In the standard simulation, oxidation of isoprene yields high concentrations of the CH$_3$CO$_2$ radical. As a result, PAN decomposition is compensated by rapid PAN formation. Our simulation with no isoprene predicts concentrations of NO$_2$ and PAN close to observed in the CBL over Alaskan tundra during ABLE 3A [Bakwin et al., 1992; Sandholm et al., 1992; Singh et al., 1992]. Isoprene is not emitted by the major tundra plants (lichens and mosses).

The sensitivity simulation without RO$_2$ + HO$_2$ reactions yields mixing ratios of NO a factor of 2 lower than the standard simulation and significantly lower than the measurements. The afternoon mixing ratio of total peroxy radicals ($\Sigma$RO$_2$) in this simulation is higher than 100 ppt (Figure 7).

Figure 8 shows the simulated NO$_2$ and PAN budgets in the boundary layer for the standard simulation. The formation of HNO$_3$ is nearly balanced by decomposition of PAN subsiding from the free troposphere; net exchange of NO$_2$ between the CBL and the free troposphere is small. The production of HNO$_3$ occurs mostly in the daytime. At night, isoprene reacts rapidly with NO$_3$, producing isoprene nitrate radicals [Atkinson et al., 1988; Dlugokencky and Howard, 1989] which release NO$_2$ upon further reactions after sunrise [Paulson and Seinfeld, 1992]; this effectively prevents the nighttime formation of N$_2$O$_5$ and hence the loss of NO$_2$ to HNO$_3$ via N$_2$O$_5$ hydrolysis. Thus isoprene suppresses formation of HNO$_3$ by depleting OH during daytime (Figure 7) and by reacting with NO$_3$ at night. As a result, the lifetime of NO$_2$ in the CBL is considerably longer with than without isoprene, 2.9 versus 1.2 days. Loss of NO$_2$ in the former case could be more rapid than computed here if isoprene nitrate radicals react on aerosols to yield HNO$_3$.

5. Summary

The origin of NO$_2$ in the summertime subarctic troposphere over eastern Canada was studied by modeling aircraft and ground observations from the ABLE 3B expedition. It is found that decomposition of PAN can account fully for the observed NO$_2$ concentrations in the free troposphere below 6 km except in occasional air masses subsiding from the upper troposphere. There is evidence that other organic nitrates are present in these air masses, and their decomposition may provide significant sources for NO$_2$. We speculate that HNO$_3$ may react with CH$_2$O in concentrated sulfuric acid aerosols to produce HOC$_2$HONO$_2$ and CH$_2$(ONO$_2$)$_2$ and that these nitrates would photolyze on a timescale of weeks to release NO$_2$.

Decomposition of PAN subsiding from aloft appears to provide the primary source of NO$_2$ in the continental boundary layer (CBL) over eastern Canada woodlands. The NO$_2$ budget in the CBL is strongly influenced by isoprene emission from vegetation. On the one hand, isoprene increases the lifetime of NO$_2$ in the CBL by scavenging OH in the daytime and NO$_3$ at night (the isoprene nitrate radicals produced at night are assumed to return
NO$_3$ upon sunrise). On the other hand, the production of CH$_3$CO$_3$ radicals from isoprene oxidation slows down PAN decomposition and hence the source of NO$_3$. For the conditions in ABLE 3B, the net effect of isoprene is to increase NO$_3$ concentrations in the boundary layer, providing thus a small boost for O$_3$ production. Forest fires appeared to be the most important source for PAN during ABLE 3B. Formation of PAN occurs in fresh biomass burning plumes and also in the regional atmosphere following dispersal of pyrogenic acetone and other hydrocarbons. Subsequent transport and decomposition of PAN can then maintain a uniform distribution of NO$_3$ in the troposphere.

### Appendix: Constraints on Peroxy Radicals

A critical variable in predictions of the photochemical activity of the atmosphere is the total concentration of peroxy radicals (ΣRO$_2$). Concurrent measurements of NO, NO$_2$, and O$_3$ concentrations, together with $J_{\text{NO}_2}$, allow, in principle, a direct computation of ΣRO$_2$ concentrations [Ritter et al., 1979; Kelly et al., 1980; Parrish et al., 1986; Völtz et al., 1988; Chameides et al., 1990; Ridley et al., 1992; Davis et al., 1993]. Rapid cycling of NO and NO$_2$ occurs in the daytime through the following reactions:

![Diagram](image)

**Figure 7.** Concentrations of (a) OH and (b) ΣRO$_2$ (sum of peroxy radicals, values in parts per trillion) at 1.0 km above ground for the simulations described in Figure 6.
where $k_{2,i}$ is the rate constant of reaction (2) for the $i$'th peroxy radical. Photochemical steady state between NO and NO$_2$ is established on a timescale of a few minutes. The steady state relation is given by

$$J_{SO} [NO_2] = (k_1 [O_3] + \sum_i k_{2,i} [R_i O_2]) [NO]$$

(4)

The values of $k_{2,i}$ for CH$_3$O$_2$, CH$_3$CO$_3$, and other organic peroxy radicals are within 20% of that for HO$_2$ for the range of observed temperatures (-30°C to 20°C) [Atkinson et al., 1992]. We choose the rate constant of the HO$_2$ + NO reaction, simply denoted $k_2$, as an approximation of $k_{2,i}$ values. We represent the sum of peroxy radicals by

$$[\Sigma R_i O_2] = \frac{[NO_2]}{k_2 [NO]} \frac{k_1 [O_3]}{k_2}$$

(5)

which may be evaluated from measurements of ambient temperature, $J_{SO_2}$, and concentrations of O$_3$, NO, and NO$_2$. We call this quantity "implied" $\Sigma R_2$.

Errors for this implied $\Sigma R_2$ may be estimated from measurement errors for the individual variables,

$$\Delta [\Sigma R_2] = \left[ \frac{\Delta [NO_2]}{J_{SO_2}} + \frac{\Delta [NO]}{[NO]} + \frac{\Delta [O_3]}{[NO_2]} \right] [\Sigma R_2 + \frac{k_1 [O_3]}{k_2}]$$

(6)

Measurement errors for O$_3$ are small and therefore neglected. The uncertainties on $k_1$ and $k_2$ are respectively 20% and 25% [Atkinson et al., 1992] but are neglected as they also occur in model calculations. The values of $k_1/k_2$[O$_3$] averaged 65 ppt in the free troposphere during ABLE 3B. Stated measurement precisions for 3-min averaging intervals were about 20% for NO at 10 ppt and 20% for NO$_2$ at 30 ppt [Sandholm et al., 1994]. Measurement uncertainties on $J_{SO_2}$ are about 20% [Madronich, 1987; Shetter et al., 1992]. Hence a typical measurement error for the $\Sigma R_2$ concentration would be at least 60%. Much larger errors for $\Sigma R_2$ are expected when measured mixing ratios of NO fall below 10 ppt and NO$_2$ below 30 ppt, respectively. Therefore it is not instructive to compare model $\Sigma R_2$ with the implied $\Sigma R_2$ at 3-min averaging intervals. We reduce the uncertainty in the implied $\Sigma R_2$ by averaging over a large number of intervals.

Figure A1 shows the implied $\Sigma R_2$ mixing ratios as a function of NO averaged over all intervals for which data are available for constraining the model. Values of $\Sigma R_2$ increase with decreasing NO and exceed 200 ppt for NO less than 5 ppt. Concentrations of $\Sigma R_2$ of a few 100 ppt would result in rapid O$_3$ production (of the order of 1 ppb h$^{-1}$), which seems inconsistent with the concentrations of O$_3$ typically observed in the free troposphere. Further, this high level of $\Sigma R_2$ could not be maintained by known sources of odd hydrogen radicals. On the other hand, possible unknown errors in the NO measurements have been estimated to be at or below 3.5 ppt [Sandholm et al., 1994], so the implied $\Sigma R_2$ at a few ppt NO may not be reliable. Figure A2 compares simulated and implied mixing ratios of $\Sigma R_2$ for the subset of data with NO above 10 ppt (i.e., x5 measurement noise). The implied $\Sigma R_2$ mixing ratios show large variances and appear to increase with altitude. The model underpredicts the implied $\Sigma R_2$ by a factor of 2-3 above 4 km; the discrepancy is less at lower altitudes.

The discrepancy could conceivably reflect a large missing source of odd hydrogen radicals in the model at high altitudes.
Alternatively, it is possible that interferences in NO2 measurements, increasing with altitude, may be responsible. In particular, the agreement between simulated and implied mixing ratios of PAN also increased with altitude and were many times larger than those of NO2 [Singh et al., 1994a]. If a few percent of PAN decomposed in the sampling tubing, the interference on the NO2 measurement would be significant. For the instrument configuration used in ABLE 3B the wall reaction efficiency would need to be approximately \(1 \times 10^{-4}\) for a 50% conversion efficiency of HNO4 \(\rightarrow\) NO2 + HO2 and \(1 \times 10^{-5}\) for a 5% conversion efficiency of PAN.

Measurements of NO are more reliable and were therefore chosen as constraint in the 0-D model calculations. Since the disagreement between implied and modeled \(\Sigma R O_2\) concentrations is less at lower altitudes, we also use the NO2 measurements as model constraints in the boundary layer calculations.

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