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Ground-based measurements of NO\textsubscript{x} and total reactive oxidized nitrogen (NO\textsubscript{y}) at Sable Island, Nova Scotia, during the NARE 1993 summer intensive

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Abstract. Measurements of NO, NO\textsubscript{2}, and total reactive oxidized nitrogen (NO\textsubscript{y}) were added to ongoing measurements of aerosols, CO, and O\textsubscript{3} at Sable Island (43º55'N, 60º01'W), Nova Scotia, during the North Atlantic Regional Experiment (NARE) 1993 summer intensive. Ambient levels of NO\textsubscript{x} and NO\textsubscript{y} were found to be highly variable, and elevated levels can be attributed to the transport of polluted continental air or presumably to relatively fresh emissions from sources upwind (e.g., ship traffic). The median values for NO\textsubscript{x} and NO\textsubscript{y} are 98 and 266 parts per trillion by volume (pptv), respectively. A midday pollution episode occurred during which elevated NO\textsubscript{x} and NO\textsubscript{y} were observed with enhanced levels of O\textsubscript{3}, CO, and condensation nuclei. Air masses of recent tropical marine origin characterized by low and constant levels of O\textsubscript{3} and CO were sampled after Hurricane Emily. The correlation between ozone and CO is reasonably good, although the relation is driven by the single pollution episode observed during the study. The correlation of O\textsubscript{3} with NO\textsubscript{x} and with NO\textsubscript{y}-NO\textsubscript{x} is complicated by the presumed NO\textsubscript{y} removal processes in the marine boundary layer. Examination of the radiosonde data and comparisons of the surface data with those obtained on the overflying aircraft provide clear indications of vertical stratification above the site.

1. Introduction

Active and reservoir reactive nitrogen compounds play an important role in tropospheric chemistry. Oxides of nitrogen (NO\textsubscript{x} = NO + NO\textsubscript{2}) are critical in determining atmospheric oxidation rates by influencing the photochemical production of ozone and, consequently, hydroxyl radical concentrations in the troposphere [e.g., Crutzen, 1974, 1979; Chamedies, 1978; Fishman et al., 1979; Fishman, 1985; Liu et al., 1980, 1983, 1987; Logan et al., 1981; Logan, 1983, 1985]. Recent studies indicate that background levels of NO\textsubscript{x} in rural areas of both the northern and the southern hemispheres may have increased by as much as a factor of 2 during the past 100 years and that the tropospheric ozone budget is now strongly influenced by photochemical production resulting from increased levels of NO\textsubscript{x} [e.g., Volz and Kley, 1988; Sandroni et al., 1992]. Additionally, evidence from satellite measurements indicates that O\textsubscript{3} exported from the source regions may contribute to the abundance of tropospheric O\textsubscript{3} in remote marine environments [e.g., Fishman et al., 1986, 1990; Fishman and Larsen, 1987; Parrish et al., 1993a; Jacob et al., 1993].

Measurements of total reactive oxidized nitrogen (NO\textsubscript{y}), especially in conjunction with measurements of speciated reactive nitrogen, have proven to be valuable in assessing the photochemical processing that has occurred in an air parcel [e.g., Fahey et al., 1986; Doddridge et al., 1991; Jaffe et al., 1991; Ridley, 1991; Parrish et al., 1991, 1993b; Atlas et al., 1992; Hübeler et al., 1992; Sandholm et al., 1992, 1994]. In this paper, total reactive oxidized nitrogen is defined as NO\textsubscript{y} = NO + NO\textsubscript{2} + NO\textsubscript{3} + 2N\textsubscript{2}O\textsubscript{5} + HONO + HO\textsubscript{2}NO\textsubscript{2} + PAN + HNO\textsubscript{3} + aerosol nitrate + other organic nitrates (where N is in the > +II oxidation state). Since the concentration of total reactive oxidized nitrogen is a more conserved quantity than the concentrations of the individual NO\textsubscript{x} species, the observed relationship among NO\textsubscript{x} and other tracers such as CO, C\textsubscript{2}Cl\textsubscript{4}, and O\textsubscript{3} can aid in testing our current understanding of the sources of these compounds [e.g., Hübeler et al., 1992]. In addition, the distribution and chemistry of NO\textsubscript{y} is of primary interest in establishing inflow/outflow regional budgets for nitrogen with regard to the tropospheric acid transport/deposition problem [Fehsenfeld et al., 1987].

Emissions of ozone precursors along the eastern coast of North America can be transported to the temperate North Atlantic Ocean. A recent study by Parrish et al. [1993a] based on the measurements of CO and O\textsubscript{3} conducted at three island sites in the Canadian Maritime Provinces (including Sable Island, the site of this present study) shows that elevated O\textsubscript{3} is often correlated with high levels of CO in the summer months.
Parrish et al. [1993a] concluded that the observed linear relationship between ozone and CO indicates summertime photochemical production of ozone from anthropogenic precursors emitted from the source region on the North American continent. In the summer of 1993 (July-September), the North Atlantic Regional Experiment (NARE) 1993 summer intensive was conducted to study the influence of anthropogenic emissions (mainly from the North American continent) on the distribution of ozone and its precursors over the North Atlantic. A detailed description of the NARE program and the summer intensive is presented by Fehsenfeld et al. [this issue, (a & b)]. During this campaign, measurements of NO, NO2, and NOy (made by the University of Michigan) were added to ongoing measurements of O3, CO (D. D. Parrish and J. S. Holloway, NOAA Aeronomy Laboratory, 1993), and aerosol scattering coefficients, number concentration, and composition (J. Ogren, NOAA Climate Monitoring and Diagnostics Laboratory, 1993) at Sable Island, Nova Scotia. In addition, solar UV radiation and water vapor content were measured, and grab samples were collected in canisters and were later analyzed for nonmethane hydrocarbons (D. Blake and N. Blake, the University of California at Irvine, 1993). This paper focuses on the abundance and variability of reactive nitrogen compounds and their relationship with O3 and CO.

2. Experiment

2.1. Site Description

The measurements were made at Sable Island (43°55'N; 60°01'W), Nova Scotia, Canada (see Figure 1). The island is located approximately 180 km off the coast of Nova Scotia and is about 300 km east of Halifax. Physically, the island is 40 km long and approximately 1 km wide. It is composed of small sand dunes (< 30 m in height) covered with grass and other low vegetation. It is a refuge for seals, wild horses, and migratory birds. Public access is restricted, and the only permanent residents are the staff of the Canadian Atmospheric Environment Service weather station. The measurement site was at the weather station which is approximately 8 km from the western end of the island, 100 m from the north beach, and 800 m from the south beach. Ongoing measurements of O3 and CO were carried out in the main operation building (see Parrish et al., [1993a] for details), whereas measurements of reactive nitrogen, aerosols, and wind speed and direction were made in a building located approximately 100 m southwest of the main operation building. A 10-m scaffold was erected next to the instrument building to mount the NOx and NOy inlet systems.

The only major source of pollution on the island was a set of diesel generators for the weather station located approximately 180 m to the east of the sampling site, downwind of prevailing southwesterly flow. Other island sources included the oil-fired water heaters of the residences located to the northeast and infrequent local vehicle traffic. An oil production platform is located approximately 40 km to the southwest of the island, and two ships are permanently stationed there. Ship traffic in the vicinity of the island is erratic. Depending on weather conditions, transatlantic shipping routes occasionally pass 20 to 40 km north and south of the island. Emissions from the oil production facility (flaring and power generation) as well as from ships may have influenced the observed levels of chemical species at Sable Island.

Figure 1. A map showing the relation of Sable Island with the eastern coast of North America.

2.2. Instrumental Methods

Nitric oxide was measured with a chemiluminescence detector utilizing the reaction of NO with O3, and ambient NO2 was measured by photolyzing NO2 to NO upstream with subsequent detection of NO [e.g., Ridley et al., 1987, 1988, 1989; Carroll et al., 1990a,b, 1992]. One detector alternated between measuring NO and NO2. NOx species were converted to NO on a gold catalyst with CO as a reducing reagent [Bollinger et al., 1983; Fahey et al., 1985, 1986], followed by chemiluminescence detection of NO using a second detector. Only the features relevant to this field study will be described in detail.

Figure 2 shows a schematic of NOx and NOy inlet systems. Ambient air was sampled at a height of approximately 9 m on a 10-m scaffold. For NOx detection, ambient air was sampled through the photolysis cell (Pyrex glass; ID, 6.4 cm; length, 38 cm) at a flow rate of 2 standard liters per minute (slpm). The residence time of sampled air in the sampling system (3/8 inch PFA sample line and photolysis cell) was less than 7 s (~5.7 s in the photolysis cell). The pressure within the cell was maintained at 125 torr and the temperature of the cell was maintained at ~13°C. Operation of the photolysis cell at reduced pressure was required due to the use of a large cell. By lowering the pressure, concentrations of ozone and water vapor in the sampled air were lowered, which prevented condensation and reduced interfering reactions between ambient ozone and NO within the photolytic cell. A 300 W, high-pressure, Xe arc lamp was used to photolyze NO2 molecules. A
For NO\textsubscript{x} measurements, ambient air was drawn at a rate of 1 slpm through a gold-plated stainless steel inlet provided by D. D. Parrish (ID, 0.19 inches; length, 3.5 inches; heated to 100°C) into the gold catalytic converter maintained at 300°C. Three standard cubic centimeters per minute (sccm) of carbon monoxide from an aluminum cylinder (99.99%, Scott Specialty Gases) was added to the sample stream. The gold-plated inlet was made as short as possible and heat was added to minimize the loss of HNO\textsubscript{3} in the inlet.

Since mass flowmeters can be sensitive to temperature changes, both of the mass flow controllers in the NO\textsubscript{x} and NO\textsubscript{y} inlet boxes were held at 35°C. To avoid any contamination of sampled air by the inlet boxes, both enclosures were equipped with a 6-inch fan, and exhaust was vented through a dryer hose extending 15 m away from the sampling platform. The two chemiluminescence detectors, along with the ozone sources, calibration module and data acquisition and control systems, were located inside the same building housing the instruments for measuring aerosols and the data acquisition system for recording wind speed and wind direction measurements. Pump exhaust and dumped calibration standards were also vented through the dryer hose.

Calibrations were conducted roughly every 3.5 hours by adding a small flow of standard gas to the ambient airstream. Calibration standards were added at levels of ~3 parts per billion by volume (ppbv). The NO standard mixture (in N\textsubscript{2}, Scott Marrin Inc.) was compared with the National Institute of Standards and Technology (NIST) Standard Reference Material, and it agreed (difference < 3%). A flowing NO\textsubscript{2} standard was generated via titration of the NO standard with O\textsubscript{3} generated by photolyzing O\textsubscript{2} with a mercury vapor lamp. The NO\textsubscript{2} standard was used to determine the conversion efficiencies of both the photolytic and the catalytic converters. In addition, gaseous HNO\textsubscript{3} obtained from a permeation tube was used to check the catalytic converter efficiency in the field and in the laboratory after the field experiment. Since the level of NO\textsubscript{2} (extent of titration) was found to vary with time, it was determined indirectly by quantifying the NO remaining after titration during each calibration period.

Throughout the experiment, the sensitivities of both detectors with respect to NO were very stable. Average instrument sensitivity was 4.87 (standard deviation, ±0.13) counts per second per parts per trillion by volume (cps/pptv) for the NO\textsubscript{x} detector and 3.80 (standard deviation, ±0.09) cps/ppmv for the NO\textsubscript{y} detector. The efficiency of the NO\textsubscript{2} photolytic converter ranged from ~16% at the beginning of the experiment (as a result of imperfect optics adjustment) to ~24% throughout the rest of the experiment. The Xe lamp was not replaced during the experiment (23 days) since there was no indication of a decreasing NO\textsubscript{2} conversion efficiency. In the NO\textsubscript{y} detection system the NO\textsubscript{2} conversion efficiency in the gold converter never dropped below 95% (in both ambient air and synthetic air). During the field study, the HNO\textsubscript{3} conversion efficiency was evaluated in zero air (HNO\textsubscript{3} was added at the position of 2 inches from the front end of the sample inlet tube, Figure 2). However, the signal of the NO\textsubscript{x} detector was not stable and exhibited sinusoidal behavior for a 3-hour period, which may be attributed to unstable delivery of the HNO\textsubscript{3} standard. After the field study, a similar test was conducted in our laboratory on the two converters used in the field, without reconditioning following their most recent use in hopes of maintaining them in the same condition as in the field. The results show that the HNO\textsubscript{3} in zero air was converted at an efficiency of ~95% and ~100% on the two converters, respectively. The efficiency was determined by comparing the signal of the NO\textsubscript{x} detector when the Au catalyst was maintained at 300°C to that at 500°C, assuming that nitric acid was converted completely at 500°C. The high conversion efficiency of HNO\textsubscript{3} obtained in the lab was taken as an indication that HNO\textsubscript{3} was converted at
high efficiency in the field as well because (1) both converters were routinely reconditioned, as described below, every second day throughout the study period; and (2) field tests conducted by the NOAA Aeronomy Lab group at a similar North Atlantic site (Chebogue Point, Nova Scotia) showed that high conversion efficiency (> 84%) of HNO$_3$ in ambient air was achieved and maintained through a similar but even less frequent reconditioning practice. The designs of the converter and the inlet used by the UM group are identical to those used by AL group.

Artifact tests, cleaning of the photolysis cell, and reconditioning of the gold catalyst were typically carried out every other day. During artifact tests, ultrapure zero air (Scott Marrin Inc.) was used in place of ambient air. During the experiment, 23 such tests were conducted. Average artifact signals were 1.3 ± 1.0 (standard deviation) pptv for NO and 7.4 ± 3.3 pptv for NO$_2$. The NO$_3$ system showed artifact signals that decreased with time, with a value of ~ 100 pptv observed at the beginning of the experiment dropping to ~ 20 pptv toward the end of the study, excepting a brief period when a signal as high as 200 pptv was observed immediately after the HNO$_3$ conversion efficiency test. The mean NO$_3$ artifact signal was 53 pptv with a standard deviation of 39 pptv. During data reduction, artifact signals in NO, NO$_2$, and NO$_y$ modes were treated differently. Since it is possible that interferents in the zero air tank may also have contributed to the artifact signal obtained during the artifact tests, only half of the NO and NO$_2$ artifact signals were attributed to real instrument bias/contamination and were subtracted from the data (but included in estimates of total uncertainty). It should be stressed that NO and NO$_2$ artifact signals were relatively small compared with typical ambient levels at Sable Island (median measured values: daytime NO, ~ 20 pptv; NO$_2$, ~ 70 pptv). NO$_3$ artifact signals, however, were subtracted in their entirety from the data as contamination by metal carbonyls from a dirty CO source at the beginning of the campaign was indicated.

For maintenance of the instrument the photolysis cell was first washed with a solution of 5% NaOH and then rinsed with deionized water (Baxter Scientific Products). The gold catalytic converter was heated to 550°C with a flow of zero air of 1 slpm for about 5 hours, followed by three additional hours of flow during the catalyst cooling-down period [see Hübler et al., 1992]. The gold-plated inlet was also heated to 150°C with a flow of zero air of 0.5 slpm. These relatively simple procedures seemed to be effective in keeping the photolysis cell clean and the efficiency of the catalytic converter high with respect to NO$_2$.

Detection limits (signal to noise ratio = 2) for NO, NO$_2$, and NO$_3$ were calculated to be 2 pptv, 9 pptv and 3 pptv, respectively, for 2-min integrations. The precision for levels well above detection limits was ±10%, ±17%, and ±2% for NO, NO$_2$, and NO$_3$, respectively. The precision estimate is based on photon-counting statistics and is calculated according to the following ambient levels: NO ~ 20 pptv; NO$_2$ ~ 55 pptv and NO$_3$ ~ 290 pptv. Overview uncertainties in the measurements were estimated to be (−35%, +28%) for NO, (−39%, +33%) for NO$_2$, and (−41%, +36%) for NO$_3$ (calculated also for above ambient levels). They included the precision and the uncertainties in artifact, the NO calibration standard (2%), the NO$_3$ conversion efficiency (15%) [Hübler et al., 1992], and mass flowmeter calibrations. In this study, the change in the precampaign and postcampaign sample mass flowmeters calibrations (up to 18%) greatly exceeded those previously observed. Because we were not able to determine whether this was a step or a gradual change, interpolated values were used to determine sample flows in the data reduction.

It should be emphasized that accurate determination of total uncertainty in NO$_y$ measurements is difficult because of the complex issues involved in the measurements, such as converter efficiency, converter poisoning, artifact, and interference from non-NO$_y$, nitrogen-containing compounds [Crosley, 1994]. Sea salt may cause problems for NO$_y$ measurements: it may poison the Au catalyst causing drops in conversion efficiencies; it may remove HNO$_3$ within the inlet if sea-salt deposition occurs in the inlet. Such removal of HNO$_3$ depends on ambient humidity. In the present study, although we could not be absolutely sure, there seemed no indication of converter poisoning by sea salt at Sable Island (NO$_2$ conversion efficiencies were > 95% and HNO$_3$ conversion efficiencies were also believed to be high, as discussed previously). Concerning the removal of HNO$_3$ within the inlet, we hoped to reduce this problem by heating the inlet to 100°C. However, the effectiveness of this approach was not evaluated in the field. Postcampaign tests on HNO$_3$ were conducted using dry cylinder air. It should be noted that this measurement site is not in the immediate sea spray zone where the sea-salt problems were discovered.

Artifacts found in this study were quite high and variable, which were mainly attributed to carbonyl compounds in the CO cylinder and to outgassing from the converter and/or sampling line. Although each zero air test could determine the artifact signal accurately at a particular time, linearly interpolated values between the two adjacent artifacts (24-48 hours apart) may introduce errors because (1) two adjacent artifact tests may be not close enough in time for linear interpolation to represent the artifacts between the two tests, especially when the two individual artifact tests differed significantly (e.g., after the HNO$_3$ test), and (2) change of artifact with time may not follow a straight line. In this uncertainty estimate we included an uncertainty of 5% from artifact determination (relative to ambient NO level of 290 pptv), or 15 pptv. This quantity may have been underestimated for the period immediately after the HNO$_3$ test in the field (early morning on September 1) but seemed reasonable for the rest of the study period.

The instruments, controlled by a computer, were repetitively switched among zero (instrument background), measurement, and calibration modes. Instrument backgrounds were determined every 30 min. The NO$_2$ and NO$_3$ instruments were configured to allow maximum overlap of measurement modes. In the NO$_3$ instrument, NO and NO$_2$ measurement modes (each lasting 138 s) alternated between zero modes (each lasting 138 s). The NO$_2$ instrument had a single measurement mode between zeros. After each transition between modes, 18 s of data were discarded to allow for signal equilibration. The remaining data were averaged over 1-min intervals, and data presented in this paper are 5-min averages.

The following procedures were employed to finalize the data: (1) The data were corrected for artifacts, as described earlier, and for zero-volume efficiency which arises from incomplete conversion of NO in the zero volume by O$_3$. Zero-volume efficiency was determined during each artifact test. It should be mentioned that this zero-volume efficiency is very close to unity for the two detectors (0.979 and 0.997, respectively). (2) Interference with the NO$_2$ measurement, resulting
from the reaction \( \text{NO} + \text{O}_3 \rightarrow \text{NO}_2 + \text{O}_2 \) in the photolysis cell, was evaluated during each calibration sequence by determining the detector response when the photolysis cell was bypassed. The subsequent correction lowers \( \text{NO}_2 \) mixing ratios. However, the magnitude of change of the data resulting from this correction never exceeded 10% and was typically less than 5%. (3) Data associated with periods of instrument malfunction were removed (< 1% of data). (4) Data associated with local pollution were removed (e.g., exhaust from the power generator and passing vehicles). About 11% of the data were excluded for this reason. (5) A wind speed filter was applied and data were removed during periods when the wind speed was less than 1 m/s (~ 2% of the data). This was done to minimize the contamination of the air samples by possible outgassing of the building. (6) Residual signals from the calibration standards were removed. In the \( \text{NO}_x \) instrument, a residual signal often existed at a level of ~ 4 pptv after each calibration. Thus \( \text{NO} \) and \( \text{NO}_2 \) data obtained during the first 10 min following calibrations were removed. Approximately 7% of \( \text{NO} \) and 5.3% of \( \text{NO}_2 \) were excluded. Since ambient \( \text{NO}_x \) levels at Sable Island were much higher than the calibration residual, such that no residual was clearly detected, this correction was not applied to the \( \text{NO}_2 \) mixing ratios.

Prior to the Sable Island experiment, an informal intercomparison of \( \text{NO}_x \) and \( \text{NO}_y \) measurements was conducted at Chebogue Point, Nova Scotia. Groups from the NOAA Aeronomy Lab, the University of Maryland, and the University of Michigan participated. The data from the Aeronomy Lab (AL) and the University of Michigan (UM) groups were compared. The primary conclusions from the \( \text{NO}_x \) intercomparison are (1) the two measurement sets are very well correlated for both \( \text{NO} \) and \( \text{NO}_2 \) over the observed concentration range; (2) UM values are about 20% higher than AL values for both species; and (3) the measurements show no significant offsets for either species. The primary conclusions for \( \text{NO}_y \) are (1) the Chebogue Point site was particularly poor for intercomparisons due to the interferents (presumably ammonia) from the adjacent dairy farm and many spikes in \( \text{NO}_x \) and \( \text{NO}_y \); (2) the two data sets are fairly well correlated at least above 2 ppbv; (3) the UM data are about 11% higher than the AL data; and (4) the UM data may be positively offset relative to the AL data below 2 ppbv. Overall, the agreement between the UM and the AL data was found to be good; the systematic differences between the two instrument sets can be attributed to inconsistencies between the UM and the AL calibration standards and to uncertainties associated with UM's mass flowmeter calibrations.

3. Results and Discussion

Measurements of reactive nitrogen (\( \text{NO}, \text{NO}_2 \), and \( \text{NO}_x \)) started on the evening of August 14 and ended on the early morning of September 6. Daytime temperatures during the study period were mild with an average value of 18°C and a diurnal difference of about 2°C. Relative humidity was typically high, averaging 80% during the day and near 100% at night. Southwesterly winds slightly predominated; wind speed averaged around 3.7 m/s with no strong diurnal variation. Figure 3 shows the time series of \( \text{NO}_x \), \( \text{NO}_y \), \( \text{NO}/\text{NO}_x \), oxidation products of \( \text{NO}_x \) (\( \text{NO}_y/\text{NO}_x \)), \( \text{CO}, \text{O}_3 \), aerosol number concentration, scattering coefficient, temperature, relative humidity, wind speed, and wind direction. Aerosol number concentrations were determined for particles greater than 15 nm in diameter (TSI, model 3760 CNC). Scattering coefficients (at wavelength of 550 nm) were measured at relative humidity of less than 30% and for particles less than 10 μm in diameter. Three large time gaps in the reactive nitrogen time series indicate the periods when air sampled at the site was contaminated by exhaust from the island generator, during which the instruments were switched from ambient air to zero air. These three periods are August 16-18, August 20-21, and September 2-3.

In general, levels of \( \text{NO}_x \), \( \text{NO}_y \), and to a lesser extent condensation nuclei exhibited variations on a relatively short timescale, whereas \( \text{CO}, \text{O}_3 \), and aerosol scattering coefficient varied on a much longer timescale (an example can be seen on August 22-23). Additionally, the magnitude of changes relative to mixing ratios of \( \text{NO}_x \) and \( \text{NO}_y \) are much more pronounced than for the other species (here, logarithm scales are adopted for the \( \text{NO}_x \) and \( \text{NO}_y \) axes). Quite often, increases in \( \text{NO}_x \) and \( \text{NO}_y \) mixing ratios were observed without concurrent enhancements in CO levels, and often aerosol concentrations showed enhancements during these \( \text{NO}_x \) and \( \text{NO}_y \) peaks. An example of these events can be seen around noon on August 21. Typically, these peaks in \( \text{NO}_x \) levels lasted from a half hour to several hours and such periods occurred throughout this study. The ratio of \( \text{NO}_x \) to \( \text{NO}_y \) is found to increase during these \( \text{NO}_x \) and \( \text{NO}_y \) peaks, which suggests that reactive nitrogen observed during these periods may have come from relatively fresh sources in the area. Exhaust from the island generators could contaminate air samples measured at the site in two ways: (1) an air parcel carrying pollutants to the site sampled shortly after contamination and (2) an air parcel carrying pollutants that had been transported away from the site for some time and returned to the site due to a change in the flow pattern. However, the chemical and surface wind data associated with these peak events were examined, and no indication that the air sampled during these periods had been contaminated by the island generators was found. Instead, sources upwind of Sable Island (e.g., ship exhaust and emissions from the oil production facility) may have contributed to these observed peaks. This speculation is supported by the fact that diesel generators at the oil production platform and diesel engines that are widely used on ships tend to burn fuel efficiently, discharging exhaust with high \( \text{NO}_x \) to \( \text{CO} \) ratios and large amount of particles. Additionally, a least one radio contact was made by the Sable Island Weather Station during an \( \text{NO}_x \) peak event, and there was indeed a ship passing the island upwind of the measurement site. Surface wind, reactive nitrogen, and aerosol concentration data were examined in an attempt to find particular wind direction window(s) in which these sources appear dominant. These sources appeared to exist in all wind directions except for easterly and southeasterly flows (data associated with northeasterly flows were contaminated by the generators).

Although the levels of reactive nitrogen sometimes varied independently of \( \text{CO} \) and ozone, there were periods when reactive nitrogen, \( \text{CO} \), and ozone were correlated. For example, on August 24-26, \( \text{NO}_x \), \( \text{CO} \), and \( \text{O}_3 \) levels all varied quite smoothly, exhibiting three maxima for a 36-hour period. It is also noteworthy that during this period, \( \text{NO}_x/\text{NO}_y \) remained low and less variable (0.2-0.4).

A major pollution episode occurred from midnight on August 26 to midnight on August 29. During this period, \( \text{NO}_x \), \( \text{NO}_y \), \( \text{CO} \), \( \text{O}_3 \), scattering coefficient, and condensation nuclei
Figure 3. Twelve-day time series of NO\textsubscript{x}, NO\textsubscript{y}, CO, O\textsubscript{3}, NO\textsubscript{x}/NO\textsubscript{y}, NO\textsubscript{x}-NO\textsubscript{y}, aerosol concentration (for particles > 15 nm in diameter), scattering coefficient (550 nm, RH < 30%), temperature, relative humidity, wind speed and wind direction. Note that NO\textsubscript{x} and NO\textsubscript{y} are in log scale. Tick marks on the time axes indicate starting of a day (0000), and periods for "clean" and "polluted" events (see text in the later section for definitions) are marked in CO time series.

(a) August 14 - August 26

as well as nonmethane hydrocarbon species (not shown) showed significant enhancements. For example, ozone reached at least 85 ppbv and CO at least 250 ppbv on the morning of August 28; NO\textsubscript{x} mixing ratios were as high as 1700 pptv (median value for this study, 266 pptv). The ratio of NO\textsubscript{x} to NO\textsubscript{y} was less than 0.1 in the afternoon of that day, suggesting that the air parcel was well aged in terms of the degree of photochemical processing. This event is referred to in this paper as the "polluted" event. Four-day isentropic back trajectories calculated by J. Moody suggest that the air mass traveling over Sable Island during the pollution episode may have come from the highly industrialized northeastern and/or midwestern United States. During the same period, surface winds at Sable Island were also from southwest. It should be mentioned here that these back trajectories were calculated for altitudes usually much higher than the sampling height and as a result vertical stratification may limit their application to this data set (see Angevine et al., [this issue], for details). Indications of stratification will be discussed below in this paper. Another interesting feature during the pollution episode is worth mentioning: NO\textsubscript{x} (as well as NO\textsubscript{x}-NO\textsubscript{y}) and condensation nuclei reached their peak levels around 0300 on August 28 and then showed a rapid decrease, whereas CO and O\textsubscript{3} remained rather constant and then showed a slower decrease. We attribute this behavior to the removal of relatively soluble species such HNO\textsubscript{3}, aerosol nitrate, and condensation nuclei along the air mass trajectory. Carbon monoxide and O\textsubscript{3} are less affected by this type of removal process. This phenomenon is discussed below and is also addressed by Roberts et al. [this issue].

In contrast to the pollution episode, air masses with different characteristics were sampled for a 36-hour period on September 4-5 following the passage of Hurricane Emily. The air was characterized by low and stable levels of CO (~ 80
ppbv) and ozone (~15 ppbv). Mixing ratios of reactive nitrogen, however, were quite variable. Again the existence of relatively fresh sources of reactive nitrogen can be seen. In this paper we refer to this period as the "clean" event, which will be examined in greater detail below. Local surface winds were south-southwesterly and back trajectory calculations suggest that air traveling over Sable Island during this "clean" event originated in the tropical Atlantic Ocean.

If the "polluted" and "clean" episodes are excluded, the remaining data may represent "regional" background conditions. This category accounts for the majority of the data collected in this study.

Vertical stratification in the marine boundary layer is an issue. There is evidence that vertical mixing at Sable Island was limited. Each day, at 0815 and 2015 (Atlantic standard time), radiosondes (VIZ Canada, model: VIZ RAP) were launched by Atmospheric Environment Service (AES) staff as part of their normal operation. The radiosondes showed vertical profiles that primarily indicated stable or neutrally stable conditions within the marine boundary layer during launching times throughout the study. Figure 4 shows typical profiles of potential temperature and water vapor mixing ratio. Although no sounding information is available at noon or in the early afternoon when surface heating-induced vertical mixing tends to be maximum, surface temperature data suggest that surface heating by solar radiation was small (average and maximum afternoon temperature is ~18°C and 20°C, respectively; and diurnal difference is only ~2°C). Further evidence for the lack of vertical mixing comes from a direct comparison of chemical measurements made by the overflying aircraft with surface measurements. These results are presented in a later section. In short, surface levels of O₃, CO, and NOₓ were quite different from those at altitudes as low as 300 m, indicating the existence of vertical stratification. Additional discussion of the vertical sounding data is presented by M. A. Carroll et al. (manuscript in preparation, 1996).

Frequency distributions of O₃, CO, NOₓ, and NOᵧ are shown in Figure 5. For each species the "polluted" event (August 25, 2146 AST – August 29, 1752 AST), the "clean" event (September 4, 1602 AST – September 5, 1752 AST), and the "rest" of the data are shown separately. From the CO and O₃ distributions, one can see that during the "clean" event both
ozone and CO levels were low and the median mixing ratios were 19 and 78 ppbv, respectively. The mean values for O$_3$ and CO mixing ratios in this subdata set are 19 ppbv and 79 ppbv, respectively. Their histograms are approximately represented by a Gaussian distribution. In the "polluted" case, however, both O$_3$ and CO levels are much higher than in the "clean" case. Median mixing ratios are 38 ppbv and 147 ppbv for O$_3$ and CO, respectively. The majority of data collected during the study falls in the "rest" category which has a median of 27 ppbv and 108 ppbv for O$_3$ and CO, respectively. These distributions lie between those of the "clean" and "polluted" cases.

For reactive nitrogen the above grouping approach is not so obvious as it is for ozone and CO. For example, even in the "clean" event, both NOx and NOy were highly variable in their respective levels. Although the median mixing ratio of NOy is 189 pptv, the mean value is 321 pptv and 4.6% of the data in this category exceed 1000 pptv. NO$_x$ showed a similar distribution with a median of 98 pptv and a mean of 153 pptv. This suggests the existence of sources characteristic of high NO$_x$ to CO ratios in the area that contribute to the reactive nitrogen levels observed at Sable Island. Evidence of these sources can be seen in all three subdata sets. The pollution episode was responsible for the extremely high concentrations of O$_3$ and CO, but this does not hold true in the case of reactive nitrogen.

Therefore the general picture is that at Sable Island the variability of CO and O$_3$ seems to be determined by the long-range transport of air masses with different histories, whereas the levels of NO$_x$ and NOy are primarily determined by the sources within relatively short distances. This may be a feature characteristic of the western North Atlantic, and the impact of these sources on the regional ozone budget needs to be further assessed.

Diurnal distributions of NO, NO$_2$, NO$_x$, NOy, NO$_x$/NOy,
Figure 5. (continued)
NO$_x$, NO$_y$, CO, and O$_3$ are presented in Figure 6. Shown are hourly averaged values, medians, central 67%, central 95%, and range. The increase in the difference between means and medians in the case of ozone and CO is primarily due to the fact that during the pollution episode, CO and O$_3$ peaked between 0800 am and 1200 AST on the morning of August 28. Thus extremely high concentrations in that single incident resulted in skewed distributions. This is also true for reactive nitrogen, NO$_x$ and NO$_y$. However, the largest difference between the means and the medians of NO$_x$ and NO$_y$ occurs at earlier morning hours than those associated with O$_3$ and CO, as a result of the fact that both NO$_x$ and NO$_y$ peaked around 0600 AST, whereas CO and O$_3$ data peaked around 1000 (there is no data record of ozone and CO between 0300 to 1000 on August 28). Thus in a discussion of diurnal behavior our focus is on the median and central 67% of the data since they are less affected by the small number of extreme values.

Carbon monoxide levels did not have an obvious diurnal cycle and the median mixing ratio remained ~ 110 ppbv throughout a diurnal period. Ozone showed some diurnal behavior, with a broad maximum around local noon (1100 AST) and a minimum in the morning around 0600 AST. However, the degree of enhancement at noon and depression during the early morning in O$_3$ levels was much less than that observed at the surface sites in the continental boundary layer [e.g., Parrish et al., 1993b]. The smaller early morning decrease as compared with that at continental surface sites may be attributed to the combination of the following processes in this marine boundary layer site: (1) less vigorous development of nocturnal inversion which allows some transport of ozone aloft; (2) advection that transports the ozone over the ocean at night. Here, horizontal transport of ozone at night is supported by the surface wind observations which suggests that wind speed often remained constant from daytime to
nighttime. The noon/afternoon maximum is also smaller at this site, which may be attributed to weaker local photochemical production of ozone, and/or less vigorous transport of ozone from ozone rich regions aloft during daytime.

A diurnal variation of median NO\textsubscript{x} may exist, as daytime values (~270 pptv) were typically larger than those at night (~220 pptv). The median NO\textsubscript{y} showed the opposite diurnal trend, with daytime values generally lower than nighttime values. The combination of these two opposite diurnal trends resulted in a distinct diurnal pattern in the NO\textsubscript{x}/NO\textsubscript{y} ratio whose median was around 0.3 during the daytime and about 0.5 at night. The NO\textsubscript{x} diurnal trend can be explained by one or a combination of the following two processes: (1) stronger transport during daytime, presumably through some degrees of downward mixing of NO\textsubscript{x}, rich air masses; and (2) stronger removal of NO\textsubscript{x} components such as HNO\textsubscript{3} and aerosol nitrate at night. Photochemical conversion of NO\textsubscript{x} to HNO\textsubscript{3} may contribute to the daytime decrease of NO\textsubscript{x} levels. The diurnal cycles of NO and NO\textsubscript{y}-NO\textsubscript{x} are as expected, with maxima occurring during the daytime and minima at night.

The relationship of reactive nitrogen species with CO and O\textsubscript{3} can provide valuable information on source origin, transport, photochemical transformation and physical removal processes involving these species. As discussed by Feisenfeld et al. [this issue (a)], a previous study conducted by Parrish et al. [1993a] found that the ozone correlated well with CO, an anthropogenic tracer, at three surface sites near the Atlantic Coast of Canada (Sable Island was one of the sites) and concluded that elevated ozone is produced from precursors of anthropogenic origin, i.e., NO\textsubscript{x} and volatile organic compounds (VOCs). Figure 7 shows the correlation plot between ozone and CO obtained during the present study. Here, the relation between these two species was found to be similar to that observed in the previous study. For example, the slope ($\Delta$[O\textsubscript{3}]/$\Delta$[CO]) is 0.41 in this study, versus 0.30 obtained previously; the correlation coefficient is 0.62 versus 0.68. However, it is noted that the observed correlation in this study is driven primarily by a single pollution episode resulting from the transport of pollutants from the North American continent, whereas in the previous study, multiple pollution events were observed.

If elevated levels of O\textsubscript{3} are a result of photochemistry involving anthropogenic precursors and if both ozone and reactive nitrogen undergo similar physical (dilution and removal) processes, a positive correlation between ozone and NO\textsubscript{y}-NO\textsubscript{x} would be expected. Such a relation has been observed in many places over the North American continent [e.g., Trainer et al., 1993; Olszyna et al., 1994; Kleinman et al., 1994]. However, as shown in Figure 8, a simple correlation between ozone and NO\textsubscript{y}-NO\textsubscript{x} obtained from the present study is not obvious. Both daytime and nighttime data are included in this figure since there is no strong diurnal cycle of boundary layer development such as impacts nighttime ozone levels at continental sites. It is worth mentioning that the data points associated with ozone mixing ratios higher than 50 ppbv are from the peak period of the pollution episode on August 28. Here, we see indications of the removal of reactive nitrogen from the air parcels arriving at the site. For example, at about 0740 AST, mixing ratios of NO\textsubscript{y}-NO\textsubscript{x}, O\textsubscript{3}, and NO\textsubscript{x} were 1440 pptv, 77 ppbv, and 232 pptv (NO\textsubscript{x}/NO\textsubscript{y} = 0.14), respectively. At about 1021, their respective levels became 660 pptv, 86 ppbv, and 72 pptv (NO\textsubscript{x}/NO\textsubscript{y} = 0.10). NO\textsubscript{y}-NO\textsubscript{x} dropped by more than a factor of 2 and NO\textsubscript{x} levels decreased by a factor of 3, while O\textsubscript{3} levels rose slightly. The decrease of total reactive oxidized nitrogen (NO\textsubscript{y}) in the second air parcel must have resulted from its removal, if one assumes that the two air parcels had similar initial loading of anthropogenic emissions. This assumption is partially supported by the observation of similar levels of CO (~250 ppbv) in the two air parcels and by the fact that fog was present at Sable Island during the peak episode up until noon on August 29. An indication of NO\textsubscript{y} removal can also be seen in the correlation plot of NO\textsubscript{y} with O\textsubscript{3}. If one forces a linear fit (two sided) to the data, the obtained slope ($\Delta$[O\textsubscript{3}]/$\Delta$[NO\textsubscript{y}-NO\textsubscript{x}], ppbv/ppbv) is 86 (r\textsuperscript{2} = 0.41), which is much higher than the slopes from regression analyses applied to observations obtained in continental...
to the speculation that air might have become unstable in the afternoon. The G-1 flew over the site at an altitude of about 160 m. Mixing ratios of surface $O_3$, $CO$, and $NO_x$ are comparable to their respective values at 160 m altitude. However, according to the aircraft data, both ozone and $CO$ concentrations increase with altitude: ozone increasing from 24 to 43 ppbv and CO from 85 to 143 ppbv, with altitude changes from 300 to 1500 m. These observations may indicate that local mixing only extended throughout the lowest few hundred meters or that vertical stratification resulted in stable layers, including one just above the site.

Case 2. The G-1 flew over Sable Island on August 25 around 0930 AST at an altitude of ~ 340 m. The daytime temperature was approximately 19°C and it was sunny in the morning and became cloudy in the afternoon. Surface ozone data agreed with the aircraft data although there appeared to be some finer structure in the ozone vertical profile between the surface and the 340-m level. There are large differences in the CO and $NO_x$ mixing ratios between the surface and the 340-m level; upper level values are much higher than those at the surface.

Case 3. The King Air flew near Sable Island around 1134-1154 AST on August 28 at an altitude of about 400 m. The daytime maximum temperature was 19-20°C, and it became cloudy in the afternoon. Vertical sounding (Figure 4) indicated stable condition around launching time. Again surface levels of $O_3$, CO and $NO_x$ are much lower than those at ~ 400 m.

In summary, there are clear indications of vertical stratification above the site in all three cases examined.

### 4. Conclusions

$NO_x$, $NO_2$, and $NO_y$ were measured at Sable Island from August 14 to September 6 during the NARE 1993 summer intensive. $NO_x$ and $NO_2$ levels were found to be highly variable, whereas CO and $O_3$ showed much less variability. The medians for $NO_2$ and $NO_y$ were 98 and 266 pptv, respectively.

### Table 1. Airborne Versus Ground-Based Measurements

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<tr>
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<th>$NO_x$, pptv</th>
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<tr>
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<td>75 - 125</td>
<td>300 - 450</td>
</tr>
<tr>
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<tr>
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<td>21 - 21</td>
<td>98 - 103</td>
<td>333 - 356</td>
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<tr>
<td></td>
<td>(21)</td>
<td>(100)</td>
<td>(335)</td>
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<tr>
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<td>41 - 62</td>
<td>120 - 180</td>
<td>1200 - 1800</td>
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<tr>
<td></td>
<td>(42)</td>
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<td>(128)</td>
<td>(300)</td>
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<tr>
<td>Ground</td>
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</tr>
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* Numbers in parentheses are mixing ratios at the time approximately when aircraft flew over the island.
Elevated levels of reactive nitrogen can be attributed to the transport of polluted continental air or presumably to relatively fresh emissions from sources upwind (e.g., ship exhaust). A midday pollution episode occurred on August 26-29, and air masses of recent tropical marine origin characterized by low and constant levels of CO and O₃ were sampled on September 4-5.

Ozone correlated reasonably well with CO, although the correlation is primarily driven by the single pollution episode. The correlation between ozone and NOₓ(NO₂) is complicated, and [O₃]/[NOₓ(NO₂)] is much higher than those values obtained from continental studies. This result is primarily attributed to the removal of NOₓ along the air mass trajectory. An indication of the removal of NOₓ was seen on the morning of August 28, when NOₓ levels dropped by more than a factor of 2 while CO and O₃ remained at elevated levels.

By comparing results from the surface and overflying aircraft measurements, we found clear indications of vertical stratification. Studies under way include the estimation of peroxy radical levels and photochemical production and loss of ozone (K. A. Duderstadt et al., manuscript in preparation, 1996), further characterization of air mass origin focusing on transport within the marine boundary layer and correlation analyses for origin categories and specific air parcels (M. A. Carroll et al., manuscript in preparation, 1996), and removal mechanisms for reactive nitrogen in the marine environment.

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