INORGANIC NITROGEN OVER THE WESTERN NORTH ATLANTIC OCEAN

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Abstract. The concentrations of the reactive nitrogen species NO$_2$, NO$_x$ (= NO + NO$_2$), NO$_y$ (the sum of all compounds of nitrogen and oxygen with the exception of N$_2$O), particulate NO$_2$, and volatile NO$_2$ were measured from ship and aircraft platforms over the western North Atlantic Ocean as part of the GCE/CASE/WATOX experiment. Air masses sampled were divided into continentally influenced and typical marine on the basis of trajectories, and radon and black carbon measurements. From the NO$_y$ measurements on size separated aerosol and the altitude variations of volatile NO$_2$ and particulate NO$_2$, a significant interaction between volatile NO$_2$ and sea salt aerosol was indicated. The average marine concentrations measured were: 18 nmol m$^{-3}$ for NO$_2$, 29 nmol m$^{-3}$ for NO$_x$, 46 nmol m$^{-3}$ for NO$_y$, and 10 nmol m$^{-3}$ for total inorganic NO. The reactive nitrogen species were present at concentrations some 40 times those encountered in the remote Pacific Ocean, whereas the inorganic NO$_2$ was only 3 times higher.

INTRODUCTION

The North Atlantic Ocean is surrounded by continental areas which contain a number of the world's largest pollutant emitters. Industry in North America and Europe, and slash and burn agriculture in Central Africa emit huge amounts of C, S, and N compounds into the atmosphere, which prevailing winds regularly transport over the ocean. The anthropogenically enhanced concentrations of these elements may have a direct impact on the biogeochemical cycling of these elements in the marine and atmospheric environments. In addition, the presence of elevated concentrations of odd nitrogen would likely perturb the natural oxidative capacity of the atmosphere over the ocean and have an indirect impact on the biogeochemical cycling of O, C, S, and N itself over the North Atlantic Ocean region. Of all the world's oceans, the North Atlantic is the most influenced by anthropogenic inputs, so it is the first area in which the impact of anthropogenic activity on the marine atmospheric and surface water environments would be discernible.

Since the pollutant sources are almost exclusively on the continents, it is possible to examine a polluted air mass as it moves off the continent over the ocean without the complication of additional inputs. The North Atlantic Ocean is therefore a somewhat simplified system on which to examine the fate of pollution in the environment. While many of the processes may be specific to the particular region under study, the results from such a simplified system may assist in the overall understanding of processes that occur as the pollutants are transported from their source regions.

A number of studies, some completed but most ongoing, have been made on the atmosphere over the North Atlantic Ocean. The Western Atlantic Ocean Experiment (WATOX) has focused on the magnitude and fate of selected nitrogen,
sulphur, organic, and trace metal compounds advected from the continent of North America to the western Atlantic Ocean [Galloway et al., 1987; Galloway and Whelpdale, 1987]. The Atmosphere/Ocean Chemistry Experiment (AEROCE) has established a number of long-term stations to study the chemical cycling of a variety of trace species that play an important role in the atmospheric and marine biogeochemical cycles over and in the Atlantic Ocean [Levy, 1988]. The STRATOZ III program has performed a number of flights around the perimeter of the Atlantic Ocean [Rudolph et al., 1987; Drummond et al., 1988]. Several ship-based measurement programs have also been conducted [e.g., Church et al., 1990]. These programs have all shown the continents to have a significant impact on the composition of the atmosphere over the North Atlantic Ocean.

In the summer of 1988 the National Oceanic and Atmospheric Administration (NOAA) ship Mt. Mitchell conducted a cruise in the North Atlantic Ocean as part of the GCE/CASE/WATOX (Global Change Expedition/Coordinated Air-Sea Experiment/WATOX) program. There were atmospheric and oceanic components to the cruise, with the aim of the atmospheric component being to obtain data on the dynamics of chemical cycling over the North Atlantic Ocean. An overview of the cruise and the experiments is given by Pszenny et al. [1990a].

The current paper uses the nitrogen species data obtained on the first leg of the cruise, from Virginia to Bermuda, to address some aspects of nitrogen chemistry over the ocean. Of particular importance are the speciation and concentrations of the nitrogen compounds over the ocean and the degree of continental influence on these concentrations.

EXPERIMENT

During the GCE/CASE/WATOX cruise the Mt. Mitchell encircled a large area of the North Atlantic Ocean (Figure 1). To augment the measurements from the ship platform, aircraft overflights were conducted close to the North American east coast and in the vicinity of Bermuda to obtain vertical profiles of species concentrations. This paper focusses on leg 1 of the cruise, which commenced in Norfolk, Virginia, on July 15, 1988 (Julian day 197), and ended at Bermuda on July 29, 1988 (Julian day 211).

The nitrogen oxides are conveniently divided into two groups on the basis of their reactivity. There are the reactive species, such as NO, NO₂ (or their sum, NOₓ), and peroxyacetyl nitrate (PAN) which, since they have lifetimes of the order of minutes to hours, must be measured in situ, preferably with fast response instruments. The more stable inorganic nitrates, such as HNO₃ and particulate NO₃, can

Fig. 1. Map of the North Atlantic Ocean indicating the track of the NOAA vessel Mt. Mitchell for the 1988 Global Change Expedition.
be collected on filters and returned to the laboratory for analysis. NOy, which includes all compounds of nitrogen and oxygen with the exception of N2O, must also be measured in situ. There were in situ and filter systems on both the aircraft and the ship.

The aircraft system for reactive nitrogen measurements, and the results obtained, are described by Ray et al. [1990] and will not be discussed here. On the ship, NO2, NOx, and NOy were measured with a commercial luminol detector, a Unisearch LMA-3 with LNC-3 converter unit operated with the manufacturer's solutions [Drummond et al., 1989]. The ozone scrubber supplied was found to absorb NO2 and so was removed from the inlet line. NO2 was measured directly by the instrument. NOx was measured by passing the air through a chromium trioxide converter to transform the NO into NO2 prior to analysis for NO2. NOy was measured using a gold converter with carbon monoxide as the reducing agent, which converted the nitrogen species to NO [Fahey et al., 1985], which was then oxidized by the chromium trioxide and measured as NO2.

This reactive nitrogen instrumentation was housed behind the bridge. Separate inlet systems were installed on booms extending 3 m outboard off the bridge wings on each side of the ship and operated such that samples were always collected from the windward side. Teflon filters (1 μm) were placed at the ends of all lines to protect against sea salt contamination. Since it will be suggested that sea salt scavenges volatile NO3 from atmospheric samples, it is possible that some of the NOy concentrations reported here are underestimated. To minimize losses of polar NOy species to the walls of the inlet system prior to conversion, gold converters were placed at the end of each inlet line. A single NO to NO2 converter was used for both inlet systems. A schematic of the sampling system is given in Figure 2. The instrument performed a zero check, by passing the air sample through a ferrous sulphate reducer,
for 6 min at the top of each hour. The remainder of the hour was broken into 18-min cycles which in turn comprised 6 min of each of NO\(_2\), NO\(_x\), and NO\(_y\) measurements. Calibrations were performed daily in the NO\(_2\) mode using an NO standard prepared by precisely diluting a certified NO standard (Scott Specialty Gases) to approximately 5 parts per billion by volume (ppbv). The NO\(_2\) concentration from the instrument was corrected for instrument nonlinearity and ozone interference [Drummond et al., 1989]. The errors introduced by these corrections became significant below 100 part per trillion (pptv) and were of the same order as the measurements at 50 pptv; thus even though Drummond et al. [1989] claim such a configuration will have a detection limit of 5 pptv for NO\(_2\), for this data set the detection limit was 50 pptv. There were no PAN measurements on this leg of the cruise so it was not possible to correct for the known PAN interference. Measurements over this area show PAN concentrations below 20 pptv [Botteghi and Gallant, 1987; Rudolph et al., 1987]. Using 0.2 as the instrument sensitivity to PAN [Drummond et al., 1989] suggests that neglecting this interference would overestimate NO\(_2\) and NO\(_x\) by less than 4 pptv.

Three institutions measured particulate NO\(_2\) and volatile NO\(_x\) (largely HNO\(_3\) but possibly including contributions from organic nitrates and N\(_2\)O\(_5\)) using filter techniques on leg 1. The three groups operated a total of four samplers on the ship. The Atmospheric Environment Service (AES) triple filter pack [Anlauf et al., 1985] trapped particles on the front Teflon filter and volatile NO\(_3\) on the second nylon filter of a three-filter system. The NOAA Atlantic Ocean Marine Laboratory (AOML) had two systems, a high volume (hi-vol) bulk aerosol system and a filter pack. Daily hi-vol samples were collected on Whatman-41 filters under conditions very similar to those used historically for most marine boundary layer sampling [e.g., Savoie et al., 1990]. Semi daily filter pack samples were collected in parallel with the hi-vol samples. In the filter pack the particulate NO\(_3\) was collected on a quartz fiber filter and the volatile NO\(_x\) on a K\(_2\)CO\(_3\)-glycerol impregnated rayon filter.

Additional details about these systems are given by Pszenny et al. [1990b] and Keene et al. [1990]. The University of Virginia (UVA) and NOAA Environmental Research Laboratories (ERL) ran a single filter pack on the ship measuring particulate NO\(_3\) on a Teflon filter and volatile NO\(_3\), also on an impregnated rayon filter [Bardwell et al., 1990]. Intercomparison of the sea salt concentrations between the various systems suggested that approximately two thirds of the sea salt, by mass, was excluded by the AOML inlet and approximately 20% was excluded by the UVA sampler [Boatman et al., 1990]. The AOML analyzed some of their filters at sea, but most of the filters were stabilized, stored and returned to the laboratory for quantitative NO\(_3\) analysis using ion chromatography. Detailed procedures are given by Anlauf et al. [1985], Pszenny et al. [1990b], Keene et al. [1990] and Bardwell et al. [1990]. Two UVA/ERL filter pack systems were deployed on the aircraft, and the samples were analyzed identically to the ship samples. For one of the aircraft systems a cyclone was used to exclude particles of radius greater than 0.4 \(\mu\)m.

RESULTS AND DISCUSSION

Distinguishing Between Continental and Marine Air Masses

One of the objectives of this experiment was to determine the concentrations of reactive nitrogen species in the Atlantic marine atmosphere. This requires that air which has recently passed over a continent be identified.

For the first leg of the cruise the meteorology over the western Atlantic Ocean was dominated by a large high-pressure system which was centered in the vicinity of Bermuda and covered most of the North Atlantic Ocean (the so-called Bermuda High). As a result the air sampled at the ship would likely have come from over the ocean to the east in moving clockwise around this high. This conclusion is supported by both the isobaric and isotropic trajectories calculated for the ship's position [Stunder et al., 1990]. However, when the ship was close to the North American coast, the air actually passed over the continent before being sampled at the ship. Thus on the early part of the cruise the shipboard instruments were sampling air heavily impacted by the continent, but as the ship travelled eastward the sampled air no longer contacted the continent. Trajectories in these latter cases indicated a residence time over the ocean of more than 5 days, and so the composition of the air was typical of that over large areas of the western North Atlantic Ocean. The date when the sampled air was no longer impacted by the continent is difficult to ascertain precisely from the trajectories but appears to be around July 20 (Julian day 202).

Two tracer methods were used to give a better determination of the time of this transition: radon and black carbon. Radon 222 is only emitted from land surfaces and has a radioactive decay half-life of 3.8 days. Thus, it is a very good indicator of an air mass that has recently been in contact with land. Radon 222 was measured on this mission; and the data are discussed in detail by Hansen et al. [1990]. Figure 3 shows the radon measurements for the Virginia to Bermuda leg. As expected, the radon concentration close to shore was over 10 pCi m\(^{-3}\), showing the air mass being sampled to be of recent continental origin. The radon concentration dropped rapidly as the distance from the coast increased and became approximately constant with an average of 1.9 pCi m\(^{-3}\) after July 19 (Julian day 201). Consequently, 0000 GMT on July 20 was used as a conservative estimate of the transition from continentally influenced air to marine air. The black carbon data [Hansen et al., 1990] support this assignment. Black carbon can come only from combustion, and as the bulk of this occurs over land, these data can also be used as an indicator of continental influence. The average black carbon concentration was 307 ng m\(^{-3}\) prior to July 20 but only 14.8 ng m\(^{-3}\) afterward, supporting the classification assigned using the radon 222 data.

In this paper, data prior to July 20 (Julian day 202) are classified as continentally influenced and data collected after this date are classified as representative of the marine air mass over the western North Atlantic Ocean.
Volatile Nitrate and Particulate Nitrate Measurements and the Evidence for Sea Salt Interaction

There is a substantial body of literature indicating that a number of acidic gas phase species can interact with the alkaline sea salt aerosol released from the ocean's surface. Finlayson-Pitts [1983] and Finlayson-Pitts et al. [1988, 1989] have shown in laboratory experiments that NO₂, ClONO₂, and N₂O₅ react with sea salt to yield nitrosyl halides (XNO X= Cl, Br), Cl₂, and ClONO₂ respectively. Brimblecombe and Clegg [1988] used thermodynamic considerations to show that HNO₃ would be expected to displace HCl when dissolved in a very low pH (1-3) sea salt aerosol droplet. Although there is not yet any evidence that the reactions found by Finlayson-Pitts et al. play a significant part in the marine boundary layer chemistry and the aerosol is generally not sufficiently acidic for the simple displacement of HCl [see Keene et al., 1990] marine aerosol measurements do show a strong interaction between volatile NO₃ and the marine aerosol. Savoie and Prospero [1982] and Church et al. [1990] are two of a number of studies that have found that a large proportion of the particulate NO₃ resides in the coarse aerosol fraction, namely, the sea salt fraction.

Any interaction between volatile NO₃ and sea salt would have a significant impact on the nitrogen budget in the marine boundary layer, as it would remove the gas phase NO₃ and sequester it on particles with short atmospheric lifetimes, 2 days or so [Kritz and Rancher, 1980]. Examination of the data obtained in this study provides support for at least an interaction between the sea salt aerosol and volatile NO₃.

The UVA/ERL group gathered particulate and volatile NO₃ data from both the ship and from the aircraft and additionally with particle size segregation from the aircraft. Only the UVA/ERL data set is considered here, as it is the only one that has altitude information. There is little gain in precision in supplementing the shipboard measurements with those of the other groups. Table 1 summarizes the UVA/ERL inorganic NO₃ measurements. The data are grouped by sampling height and by sampling system, and within each of these classifications into continentally influenced and marine air. The aircraft measurements out of Bermuda are listed as a special case. The means with their standard deviations and the medians are plotted in Figure 4.

Concentration measurements are available from three altitudes, at 16 m above the sea surface as measured from the ship, and from 152 and 2591 m as measured from the...
TABLE 1. UVA/ERL Volatile and Particulate Nitrate Measurements

<table>
<thead>
<tr>
<th>Altitude m</th>
<th>Air Mass</th>
<th>Size Fraction</th>
<th>Particulate Nitrate, nmol m⁻³</th>
<th>Gas Phase Nitrate, nmol m⁻³</th>
<th>Gas Phase to Total Nitrate Ratio nmol m⁻³</th>
<th>No. of Points</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mean (s.d.) Median</td>
<td>Mean (s.d.) Median</td>
<td>Mean (s.d.) Median</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>Continental</td>
<td>Total</td>
<td>15.6 (6.1) 14.1</td>
<td>17.3 (22.7) 2.2</td>
<td>0.31 (0.26) 0.15</td>
<td>6</td>
</tr>
<tr>
<td>16</td>
<td>Marine</td>
<td>Total</td>
<td>6.9 (2.4) 6.6</td>
<td>2.2 (1.6) 1.5</td>
<td>0.24 (0.13) 0.22</td>
<td>14</td>
</tr>
<tr>
<td>152</td>
<td>Continental</td>
<td>Fine</td>
<td>4.4 (1.7) 4.8</td>
<td>12.5 (13.6) 7.1</td>
<td>0.57 (0.30) 0.64</td>
<td>4</td>
</tr>
<tr>
<td>152</td>
<td>Continental</td>
<td>Total</td>
<td>8.7 (4.7) 7.7</td>
<td>15.1 (12.1) 10.9</td>
<td>0.57 (0.18) 0.46</td>
<td>6</td>
</tr>
<tr>
<td>152</td>
<td>Marine</td>
<td>Fine</td>
<td>2.1 (0.2) 2.1</td>
<td>1.2 (0.2) 1.2</td>
<td>0.36 (0.08) 0.36</td>
<td>2</td>
</tr>
<tr>
<td>152</td>
<td>Marine</td>
<td>Total</td>
<td>4.3 (0.1) 4.3</td>
<td>2.0 (0.4) 2.0</td>
<td>0.32 (0.04) 0.32</td>
<td>2</td>
</tr>
<tr>
<td>152</td>
<td>Marine (BDA)</td>
<td>Fine</td>
<td>0.60 (0.4) 0.41</td>
<td>1.8 (2.8) 0.58</td>
<td>0.56 (0.25) 0.62</td>
<td>8</td>
</tr>
<tr>
<td>152</td>
<td>Marine (BDA)</td>
<td>Total</td>
<td>1.0 (0.3) 0.92</td>
<td>2.2 (0.4) 2.4</td>
<td>0.68 (0.07) 0.68</td>
<td>8</td>
</tr>
<tr>
<td>2591</td>
<td>Continental</td>
<td>Total</td>
<td>1.6 (0.3) 1.7</td>
<td>12.0 (4.2) 11.2</td>
<td>0.88 (0.88) 0.90</td>
<td>3</td>
</tr>
<tr>
<td>2591</td>
<td>Marine</td>
<td>Total</td>
<td>1.3 (N/A) 1.3</td>
<td>1.7 (N/A) 1.7</td>
<td>0.56 (N/A) 0.56</td>
<td>1</td>
</tr>
<tr>
<td>2591</td>
<td>Marine (BDA)</td>
<td>Total</td>
<td>0.64 (0.1) 0.58</td>
<td>2.5 (1.1) 2.7</td>
<td>0.75 (0.16) 0.82</td>
<td>4</td>
</tr>
</tbody>
</table>

BDA, Bermuda

Aircraft on its constant-altitude legs. Data from the cyclone-separated samples on the aircraft are only available for the lower aircraft sampling altitude (152 m) and are shown in Figure 4a. The degree to which the upper level air mass is related to the lower two is not that clear. Rawinsondes flown from the ship [Stunder et al., 1990] showed there to be a strong and persistent inversion at about 700 mb (~3000 m) with weak and erratic inversions in the 500 m to 2 km altitude range. The two lowest sampling altitudes are clearly within the marine boundary layer, but it may well be the case that some of the highest-altitude measurements may also be on air from within the marine boundary layer, although this is not supported by the aerosol measurements where the measured sea salt was always below detection limits.

Comparison of the aircraft data from the sample restricted to the fine aerosol fraction (Table 1 and Figure 4a) and the total aerosol (Table 1 and Figure 4c) show the volatile NO₃ to be lower in the fractionated sample than in the unperturbed sample. As the volatile NO₃ measured must be gas phase to pass through the first filter, these results suggest a small loss of volatile NO₃, probably on the walls of the cyclone itself [see Bardwell et al., 1990].

Much more noticeable are the differences in the particulate NO₃ concentrations. In all three cases there is approximately half the particulate NO₃ in the fine fraction aerosol compared with that in the total sample. This points to the measured particulate NO₃ being more abundant on the larger particles and is consistent with the observations in the literature [e.g., Savoie and Prospero, 1982; Church et al., 1990].

Further information comes from the variation in sodium, particulate NO₃, and volatile NO₃ with altitude. As the altitude increases, the concentrations of sodium decrease markedly. This is due largely to the sea surface source and large size of the sea salt particles and their high settling velocity, although there may also be a decrease in the collection efficiency of the large particles. The concentrations of the particulate NO₃ also decrease markedly, whereas those of the volatile NO₃ remain almost unchanged. This again supports the contention that a large fraction of the particulate NO₃ is associated with the sea salt. This also means that the distribution of the inorganic NO₃ between the gas and particle phases is not the same at all altitudes within the marine boundary layer. Figure 5 shows, in both the mean and median values, that there is an increase of a factor of 2 in the gas phase NO₃ to total NO₃ ratio between the ship data and the 152 m aircraft data and a further 50% in going to 2591 m. This strongly supports an interaction between the volatile NO₃ and the sea salt aerosol. It is not possible to tell from these data alone whether this interaction is a simple adsorption or a true volatile NO₃ to particulate NO₃ conversion.

This observation also has implications for the measurement of volatile and particulate nitrate in the marine environment. It is possible that once sea salt aerosol is present on a filter it can remove volatile NO₃ from the air sample, and thus inorganic NO₃ that was present in the air as volatile NO₃ may be measured as particulate NO₃ [see Keene et al. [1990]]. This effect is in the opposite direction to the well-known production of volatile nitrate from the volatilization of ammonium nitrate aerosol which is unlikely to be important in this case as the ammonium nitrate concentration is expected to be low. The extent of this problem will depend on the homogeneity of the environment over the sampling period. If the volatile NO₃ is in equilibrium with the sea salt prior to sampling and if the air mass is homogeneous in both inorganic nitrate and sea salt, then...
there will be no further interaction on the filter. A change in the concentrations or any conditions that govern the equilibrium processes could lead to errors in the measured partitioning between volatile and particulate nitrate. In environments where there is a large amount of volatile NO\textsubscript{3} or low sea salt concentrations, or for samples with low loadings, the bias should be small. Care must therefore be taken, in a sea-salt-rich environment, in using filter pack data to quantitatively differentiate between gas and particulate NO\textsubscript{3}. The measured particulate NO\textsubscript{3} should be considered an upper limit and the volatile NO\textsubscript{3} a lower limit.

Concentrations of the Inorganic Nitrogen Species

Reactive nitrogen. Figure 6 shows the concentrations of NO\textsubscript{2}, NO\textsubscript{x}, and NO\textsubscript{y} as measured from the ship on leg 1.
The instrument was not operated for the first day out of port to prevent contamination by the very polluted air of the Norfolk port area. The highest concentrations were encountered on the first few days of the cruise when the continentally influenced air was sampled. The concentrations decreased as the distance from the continental source increased owing to the combination of dispersion, transformation, and deposition processes.

The data after July 20 (day 202) are considered to be characteristic of North Atlantic marine background and should yield an estimate of the marine reactive nitrogen concentrations over the Western North Atlantic Ocean. However, before the discussion of these concentrations it should be noted that while the concentrations are generally uniform, there are two features that require comment.

The NO₃ concentrations go through a minimum on Julian day 200 rising to a much higher concentrations on Julian days 201 and 202 (July 20-21) then decreasing for the following 7 days, whereas the changes in NO₂ and NO₂ over this period were much smaller. By all previous arguments, the air after July 20 was free of continental influence and would be expected to show similar NO₃ concentrations over this period. Intercomparisons with the NO₂ monitor on the aircraft were performed on July 19 and
Fig. 6. Reactive nitrogen measurements from the first leg of the cruise.

TABLE 2. Marine Nitrogen Concentrations at Sea Level

<table>
<thead>
<tr>
<th>Source</th>
<th>NO₂</th>
<th>NOₓ</th>
<th>NO₃</th>
<th>Inorganic NO₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carroll et al. [1990]</td>
<td>0.46</td>
<td>0.62</td>
<td></td>
<td></td>
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<tr>
<td>(Pacific Ocean)</td>
<td></td>
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<td></td>
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<tr>
<td>Hastie et al. [1988]</td>
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<tr>
<td>(Bermuda data)</td>
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<tr>
<td>Wolff et al. [1986]</td>
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<td></td>
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<tr>
<td>(Bermuda winter)</td>
<td></td>
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<td></td>
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<tr>
<td>(Bermuda summer)</td>
<td></td>
<td></td>
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<tr>
<td>Misanchuk et al. [1987]</td>
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<td></td>
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<tr>
<td>(Atlantic Ocean)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Luke and Dickerson [1987]</td>
<td>55.3</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>(Atlantic Ocean)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Whelpdale et al. [1987]</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Atlantic Ocean)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>This work</td>
<td>18(8.5)</td>
<td>29(9.4)</td>
<td>46(17)</td>
<td></td>
</tr>
<tr>
<td>(AES)</td>
<td></td>
<td></td>
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<td>(UVA)</td>
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<tr>
<td>(AOML)</td>
<td></td>
<td></td>
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<tr>
<td>Savoie et al. [1990]</td>
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<td></td>
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<tr>
<td>(North Pacific)</td>
<td></td>
<td></td>
<td></td>
<td>5.2⁺</td>
</tr>
<tr>
<td>(South Pacific)</td>
<td></td>
<td></td>
<td></td>
<td>3.2⁺</td>
</tr>
</tbody>
</table>

Standard deviations, where available, are given in parentheses.

* Obtained by summing the particulate and volatile nitrate concentrations.

⁺ Particulate measurement only assuming gas phase scavenged by the collected particulate.
mass, or a change in local meteorology with an increase in wind speed, although the trajectories and weather maps give no such indication.

To obtain average marine reactive nitrogen concentrations over the western North Atlantic Ocean, all data after Julian day 202 have been averaged. These are presented in Table 2 along with some previous marine measurements for comparison. The concentrations are lower than those previously measured in marine air over the western Atlantic Ocean [Misanchuk et al., 1987; Luke and Dickerson, 1987], and at Bermuda [Hastie et al., 1988 and Wolff et al., 1986]. This is likely because these previous measurements were made in the winter, when there was a strong westerly flow carrying material from the continent over the ocean. The measurements of Misanchuk et al. [1987] were made within 50 km of the North American coast and so were even more strongly impacted by the transport of continental nitrogen emissions.

The concentrations reported here are much higher than those measured over the Pacific Ocean [Carroll et al., 1990; McFarland et al., 1979]. On the NASA Chemical Instrument Test and Evaluation (CITE) 2 program the average NOx concentration was 0.014 ppbv (0.62 nmol m\(^{-3}\)) in the marine boundary layer [Carroll et al., 1990]. Even these levels are thought to be above the natural Pacific Ocean marine background, due to air circulating from the continent to the aircraft several hundred miles offshore. McFarland et al. [1979] measured NO concentrations of 4 pptv from a ship in the mid-Pacific. Thus it is apparent that the reactive nitrogen encountered over the western Atlantic Ocean in this study is 40 to 50 times higher than that found in the marine boundary layer over the Pacific Ocean. As the ocean is not a significant source of oxidized nitrogen [Liu et al., 1983], this indicates that the surrounding continents are a major source of reactive nitrogen over the Atlantic Ocean.

The marine reactive nitrogen concentrations measured here are for summer and for air that has been over the ocean for more than 5 days. The measurements of Luke and Dickerson [1987] are for the winter when there was a strong flow off the continent, so the time that the air had spent over the ocean was only 1-2 days. There is, however, less than a factor of 2 difference in the reactive nitrogen concentration between these two regimes, and the reactive nitrogen concentration is still more than an order of magnitude above the probable background concentration [Carroll et al., 1989]. This difference cannot be explained by the small continental seasonal variation in the nitrogen sources and, as there are believed to be no oceanic sources, points toward large continental sources and a several-day lifetime for the reactive nitrogen species.

**Inorganic nitrate.** Figure 7 shows the concentrations of inorganic (gas and particulate) NO\(_2\) as measured by each of the three groups, operating NO\(_2\) systems on the first leg of the cruise. On the first day of measurements the UVA/ERL group measured much higher NO than those of the other groups, but by July 18 and for the remainder of leg 1 the measurements of total NO\(_2\) agreed well (see Figure 7). For the marine samples the measured concentrations were very consistent from sample to sample and between groups. The averaged marine total NO\(_2\) concentration over all three groups are given in Table 2 along with other reported values. The values reported here are similar to those reported over the Atlantic Ocean by Whelpdale et al. [1987] and for Bermuda by Hastie et al. [1988] and Wolff et al. [1986]. Savoie et al. [1990] report annually averaged total
NO$_3^-$ concentrations throughout the Pacific Ocean. These data were collected on a single particulate filter which was assumed to collect all the particulate nitrate and scavenge any of the small amount of volatile nitrate thought to be present. If there was a significant amount of volatile nitrate in the sampled air, then this assumption is not valid and the data yield only a lower limit for the total nitrate. In these environments, volatile nitrates are thought to be much lower in concentration than the particulate nitrates, so the data are then directly comparable with those obtained in this study. Savoie et al. [1990] report nitrate concentrations in the remote North Pacific approximately a factor of 2 lower than those reported here for the western Atlantic Ocean and remote South Pacific concentrations approximately a factor of 3 lower. As with the reactive nitrogen species, this suggests that the air over the western Atlantic Ocean is much more heavily impacted by the emissions from the surrounding continents than is the Pacific Ocean. While this data set is small and limited in spatial and temporal coverage compared with the extensive data of Savoie et al. [1990], the indications are that there is only a small difference in total inorganic NO$_3^-$ between the oceans when compared to the difference in the reactive nitrogen precursors, namely, a factor of 3 compared to a factor of 40. This difference may be explained by a slow conversion of reactive nitrogen into NO$_3^-$ and a much faster process removing this NO$_3^-$ from the marine boundary layer.

CONCLUSIONS

The concentration of nitrogen species over the western North Atlantic Ocean is elevated compared with remote marine areas, likely because of the advection of emitted materials from the continents. Both the reactive species, such as NO$_2$ and NO$_x$, and the less reactive but more readily deposited inorganic nitrates are found in higher amounts. The reactive species are present at approximately 40 times the concentration found over the Pacific and the total NO$_2$ at a factor of 1.5 to 3 higher, suggesting that the rate of NO$_2$ loss over the ocean is fast compared to the production of the NO$_3^-$ from the reactive precursors.

The distribution of inorganic NO$_3^-$ between particulate and volatile phases appears to have a strong dependence on altitude. A much higher fraction of the NO$_3^-$ is in the particulate phase at lower altitudes where the sea salt concentration is higher. This suggests a strong interaction of volatile NO$_3^-$ with the sea salt aerosol. Aircraft sampling at a greater number of altitudes would give greater insight into the processes responsible. This interaction also has implications for the measurement of these species using filter pack systems. The volatile and particulate NO$_3^-$ can both be quantitatively determined in the following cases: if the concentrations are homogeneous throughout the sampling period, if the volatile NO$_3^-$ concentration is high or the sea salt concentration low, or where the filter loading is low. In other cases there is the possibility of the filter packs giving an erroneous partitioning so that only the total inorganic NO$_3^-$ can be determined.

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