Impact of aircraft emissions on reactive nitrogen over the North Atlantic Flight Corridor region

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Abstract. The impact of aircraft emissions on reactive nitrogen in the upper troposphere (UT) and lowermost stratosphere (LS) was estimated using the NOy-O3 correlation obtained during the Subsonic Assessment (SASS) Ozone and Nitrogen Oxide Experiment (SONEX) carried out over the U.S. continent and North Atlantic Flight Corridor (NAFC) region in October and November 1997. To evaluate the large-scale impact, we made a reference NOy-O3 relationship in air masses, upon which aircraft emissions were considered to have little impact. For this purpose, the integrated input of NOx from aircraft into an air mass along a 10-day back trajectory (ANOx) was calculated based on the Abatement of Nuisance Caused by Air Traffic/European Commission (ANCAT/EC2) emission inventory. The excess NOy (dNOy) was calculated from the observed NOy and the reference NOy-O3 relationship. As a result, a weak positive correlation was found between the dNOy and ANOx, and dNOy and NOx/NOy values, while no positive correlation between the dNOy and CO values was found, suggesting that dNOy values can be used as a measure of the NOx input from aircraft emissions. The excess NOy values calculated from another NOy-O3 reference relationship made using in situ condensation nuclei data also agreed with these dNOy values, within the uncertainties. At the NAFC region (45°N–60°N) the median value of dNOy in the troposphere increased with altitude above 9 km and reached 70 parts per trillion by volume (pptv) (20% of NOy) at 11 km. The excess NOy was estimated to be about half of the dNOy values, corresponding to 30% of the observed NOy level. Higher dNOy values were generally found in air masses where O3 = 75–125 ppbv, suggesting a more pronounced effect around the tropopause. The median value of dNOx in the stratosphere at the NAFC region at 8.5–11.5 km was about 120 pptv. The higher dNOx values in the LS were probably due to the accumulated effect of aircraft emissions, given the long residence time of affected air in the LS. Similar dNOy values were also obtained in air masses sampled over the U.S. continent.

1. Introduction

Nitrogen oxide radicals (NOx = NO + NO2) play a critical role in the photochemistry of ozone (O3) and hydrogen oxide radicals (HOx = OH + peroxy radicals) in the upper troposphere (UT) and lowermost stratosphere (LS). The major sources of total reactive nitrogen (NOy = NO + NO2 + NO3 + HNO2 + HNO3 + HNO4 + 2N2O5 + peroxyacetyl nitrate (PAN) + other organic nitrates + aerosol nitrates) in the UT are transport from the stratosphere, NO production by lightning, aircraft NOx emissions, and convective transport of NOy-rich air from the continental surface where NOy is produced by fossil fuel combustion and other processes. According to three-dimensional (3-D) model calculations the contribution from each source is similar in magnitude at northern midlatitude UT during winter, while lightning becomes a major source in summer [Lamarque et al., 1996]. Because of a rapid growth in air traffic in recent years, greater attention has been paid to the impact of aircraft NOy emissions on the O3 chemistry in the UT and LS. Model calculations predict that aircraft emissions can have significant impact on large-scale NOx distributions in the UT and LS in the northern midlatitudes and high latitudes [Brasseur et al., 1996]. The emissions are especially extensive in the North Atlantic Flight Corridor (NAFC) region, where commercial air traffic is heavy at altitudes between 10 and 12 km [Baughcum et al., 1996; Gardner, 1998]. The maximum increase in NOx in the NAFC region was estimated to be between 50 and 90 parts per trillion by volume (pptv) at 200 hPa for July, based on several 3-D model calculations using a 1990 fleet of subsonic aircraft [Brasseur et al., 1996, 1998, and references therein]. Sharp increases in NOy concentrations within aircraft contrails were observed in the eastern NAFC region, and an aircraft source was successfully identified in some cases [Schlager et al., 1997; Klemm et al.,...
Figure 1. Flight tracks during the SONEX experiment. In this study, analyses were made in two study areas, the NAFC region at 45°N–60°N and 0°W–80°W (a part of flight tracks denoted by solid lines) and the region over the U.S. continent at 35°N–45°N and 70°W–120°W (dashed lines).

1998]. Although the large-scale effect (e.g., corridor effect) from aircraft emissions is more difficult to evaluate from the observation, a regional-scale enhancement in the NO/NOy ratio was identified in the LS over the east coast of the U.S. continent by comparing the observed ratios with those of trajectory model calculations [Witte et al., 1997].

The Subsonic Assessment (SASS) Ozone and Nitrogen Oxide Experiment (SONEX) was carried out using NASA's DC-8 aircraft in October and November 1997. Most of the aircraft measurements were made over the U.S. continent and the NAFC region (Figure 1). Four intensive flights were made from both Bangor, Maine (45°N, 68°W), and Shannon, Ireland (53°N, 9°W). A substantial amount of the data was obtained at altitudes between 8 and 11.5 km in both the UT and LS. Details of the experiment are described by Singh et al. [1999].

In this paper, the impact of aircraft emissions on the amount of reactive nitrogen was estimated using the NOy-O3 correlation. This is because a positive correlation between NOy and O3 was generally observed in the UT and LS [Murphy et al., 1993; Ridley et al., 1994; Kondo et al., 1996] and aircraft emissions can alter the correlation [Koike et al., 1997]. In Plate 1 a correlation plot between NOy and O3 using all of the 10-s SONEX data obtained above 8.5 km are shown. In this plate we excluded the data obtained within aircraft plumes, influenced by lightning, or affected by recent convective transport, as described in detail in the following section. As seen in this plate, the NOy values at 45°N–60°N and those over the U.S. continent are generally higher than those obtained over the west coast of the United States and at 30°N–45°N and 60°N–70°N, where air traffic density is lower. As described in the following sections, a reference NOy-O3 relationship will be derived using data from air masses, in which aircraft emissions are considered to have little impact, that is, “background air.” The excess NOy (dNOy) will be estimated using this reference relationship. The statistical analyses of these dNOy values will be made for the data obtained at 45°N–60°N over the Atlantic (corresponding to the NAFC) and the U.S. continent. The excess NOy (dNOy) will also be estimated using the reference NOy/NOx ratio in the background air.

2. Measurements and Data

2.1. Measurements

The NO and NOy instrument used for SONEX was very similar to that used during NASA's Pacific Exploratory Mission West-B (PEM-West B) conducted in 1994 [Kondo et al., 1997]. Both NO and NOy were measured using a chemiluminescence technique. NOy compounds were catalytically converted into NO on the surface of a heated gold tube with the addition of CO. The inlet tube for air sampling faced rearward discriminating against particles of diameter larger than about 1 μm. The data were recorded every 1-s; however, 10-s averaged data were used in this study. The precision of the 10-s NO and NOy measurements at 10 km estimated from the photon counts fluctuations (2σ) was 6 and 19 pptv for NO and NOy values of 100 and 800 pptv, respectively. The absolute accuracy was estimated to be 8 and 10% for these NO and NOy values.

During the SONEX experiment the same gold tube was used for all NOy measurements. Other than operating the instrument using synthetic air, no attempt was made to clean the tube. The conversion efficiency of NO2 in ambient air was 99 ± 1% during each flight. Possible interference in the NOy measurements from the conversion of HCN, NH3, and CH3CN was checked in the laboratory and during the flights. The conversion efficiency of HCN in dry synthetic air was between 0.5 and 4% on the ground before, during, and after the mission. It was also measured during the two test flights (at 6.4–11.9 km), using two independent NOy channels and by adding a known amount of HCN gas into one of the channels. For one flight the conversion efficiency was 2.1 ± 0.5% for the O3 and H2O
Plate 1. Correlation plot between NOy and O3. All SONEX 10-s data obtained above 8.5 km were used after excluding data obtained within aircraft plumes and data influenced by lightning or recent convective transport.

mixing ratios of 62–140 parts per billion by volume (ppbv) and 23–235 parts per million by volume (ppmv), respectively. For the other flight it was 1.6 ± 0.5%, although the O3 and H2O values were not measured. From these results, HCN was estimated to have contributed about 4 pptv to the SONEX NOy data assuming an average HCN mixing ratio of about 200 pptv [Rinsland et al., 1982; Mahieu et al., 1995]. The conversion efficiencies of NH3 and CH3CN were also measured in the laboratory at 1–3 and 0.5–2%, respectively. Assuming mixing ratios of NH3 and CH3CN in the UT of 1 pptv [Ziereis and Arnold, 1986] and 200 pptv [Hamm et al., 1989], the interference from these species is considered to be negligible.

During the SONEX experiment the 15 s averaged actinic flux was measured using grating monochromators to calculate photolysis rates of NO2 (J(NO2)) [Shetter and Müller, 1999]. The photostationary state NO2/NO ratios were calculated using the measured values of O3, temperature, pressure, and J(NO2) for periods when the solar zenith angles were lower than 87°. The NOx mixing ratios were calculated from these NO2/NO ratios and the observed NO mixing ratios. The calculated NO2/NO ratios and NOx values were compared with those calculated using a box model in which a more complete photochemistry was included [Jaegle et al., 1998]. At altitudes above 7 km they agreed within 30 and 10%, respectively. Simplified chemistry was used in this study to maximize the amount of NOx data.

In this study, 10-s averaged data of O3, N2O, and CO were also used. The O3 measurements were made using a chemiluminescence technique, and N2O and CO were measured using a tunable diode laser system. The concentration of condensation nuclei (CN) with a diameter larger than 15 nm (CN fine particle; TSI model 3760) [Anderson et al., 1999] was also used in this study. The unit of CN concentration was normalized to 0°C temperature and 1013 hPa (STP cm⁻³) to represent the
on the east coast. This area will be referred to as the "U.S. continent region."

When the measurements were made within the heavy air traffic region, sharp simultaneous increases in NO and NOy values were often seen [Anderson et al., 1999; T. Sugita et al., manuscript in preparation, 1999]. These air masses were most likely sampled within aircraft plumes. The increase in the 1-s NO and NOy data ranged from 100 pptv to a few ppbv. The time duration of the enhancement ranged from 1 to 60 s, with a median value of 4 s. Given the typical cruise speed of the NASA DC-8 aircraft of 200–250 m/s, the horizontal scales of the plumes were smaller than about 10 km with a typical size of about 1 km. On the basis of the Gaussian plume model, aircraft emission plumes are diluted to background concentrations within 5 to 10 hours for a typical wind shear condition with a corresponding horizontal size of 20 km [Schlager et al., 1997; Gerz et al., 1998]. Although the size of the NOy enhancement observed from the NASA DC-8 depends on the geometry of the plume crossing, the maximum size of the observed plume was comparable to these estimates. These estimates were also consistent with a plume age of 10 min to 10 hours estimated using peak NOx mixing ratios observed during SONEX [Anderson et al., 1999]. In this study, the data obtained within the plumes were excluded in the evaluation of the large-scale impact from the aircraft emissions. This was because the sampling frequency of the plumes depends on the location and time of the measurement (many plumes were observed when the measurements were made in air masses just after heavy air traffic) and an inclusion of these data could be a potential cause of a bias in overestimating the aircraft impact. The excluded data corresponded to 6% of the entire data set at the 8.5 to 11.5 km altitude range. Because they were relatively infrequent, the results of this study would not change significantly even if the plumes were included in the analyses.

A clear signature of NO production by lightning was seen during four flights on October 13 and 29, 1997, and November 3 and 9, 1997 [Allen et al., this issue; A. M. Thompson et al., unpublished manuscript, 1999]. In these cases the NOx and NOy mixing ratios increased more than 1 ppbv, and NOx/NOy ratios ranged between 0.4 and 1.0 suggesting that NO production by lightning had occurred within a few days prior to the measurements. These results were generally consistent with air mass trajectory analysis, convective activities using cloud images, and lightning activities detected by the U.S. ground-based lightning network [Allen et al., this issue; A. M. Thompson et al., unpublished manuscript, 1999]. Possible influences from lightning were also observed on October 20 and 28, 1997. In this study all of the data likely to have been influenced by NO production by lightning were excluded. It should be noted, however, that the influence of lightning cannot be identified in an air mass once the NOy in the air mass is diluted down near the background level. Consequently, we could not completely remove from our analysis the data which might have been affected by NO production from lightning. The contribution from lightning will be revisited in section 3.2.3.

The vertical profile of the CO in the troposphere is shown in Figure 2 using all of the data except for those obtained from aircraft plumes and clearly influenced by lightning. The median values and 67% ranges in the NAFC region are also shown. As seen in this figure, the CO mixing ratio generally decreased with altitude, and the median values were between 76 and 82 ppbv at 9 to 11 km. These values were systematically lower than the median CO values of 100 to 105 ppbv obtained in the
continental or maritime air masses in the UT over the middle to high latitude western Pacific in September and October [Kondo et al., 1996]. In Figure 3 a correlation plot between NOy and CO is shown using the data obtained at altitudes above 8.5 km. Different symbols are used for the UT and LS data. The NOy mixing ratio did not generally have a positive correlation with the CO mixing ratio in the UT. These results suggest that NOy values in most of air masses in the UT obtained during SONEX had not been influenced by recent convective transport of polluted continental surface air. To eliminate the possible influence from recent convection, the data with CO values higher than 100 ppbv were excluded in this study. In addition, data showing a clear positive correlation between CO and NOy were also excluded. The excluded data by these criteria corresponded to 19% of the entire UT data set obtained above 8.5 km.

The correlations of NOy with nonmethane hydrocarbons such as ethyne (C2H2), ethane (C2H6), and propane (C3H8), and halocarbons such as tetrachloroethylene (C2Cl4), methyl iodide (CH3I), and CHBr3 were also examined. No clear positive correlation was seen in the UT. In fact, NOy decreased with increasing mixing ratios of these hydrocarbons and halocarbons for NOy mixing ratios higher than 300 pptv (not shown). C2H2, C2H6, C3H8, and C2Cl4 are emitted in urban areas. CH3I and CHBr3 are considered to be good indicators of maritime air mass. The photochemical lifetimes of these species are comparable or shorter than that of CO. Consequently, the lack of a clear correlation of NOy with these species further confirms that the data selected in this study were generally free from the recent convective transport of air influenced by urban and other surface sources.

3. Results and Discussion

3.1. Reference NOy-O3 Relationship

To estimate the increase in the NOy mixing ratio due to aircraft emissions, we utilized a method that uses the NOy-O3 relationship. In this method a reference NOy-O3 relationship was estimated using data in air masses upon which aircraft emissions were believed to have had little impact, that is, "background air masses." To select the background air masses at 8.5–11.5 km, two independent approaches were taken. As described below, the agreement in the estimates on the excess NOy derived from these two approaches was evaluated, and the first approach described below was used for the further analyses.

For the first approach, 10-day back trajectories were calculated using a kinematic method for air masses sampled every 1 min on board the NASA DC-8. The European Centre for Medium-Range Weather Forecasts (ECMWF) data were used for this calculation. Then an integrated value of expected NOy input from aircraft emissions along each trajectory was calculated using the monthly mean values of the three-dimensional NOy emission distribution for October 1992, compiled in the Abatement of Nuisance Caused by Air Traffic/European Commission (ANCAT/EC2) emissions inventory [Gardner, 1998]. No chemical loss or diffusion process and no diurnal variation in the NOy emission rate were taken into account for this calculation, although some initial dilution effect was included because the emission rate was provided for each 1° x 1° in latitude and longitude and every 1 km in altitude. The typical emission rate in the corridor region was 2–5 pptv h⁻¹. The calculated value is denoted as \( \Delta NO_y \). For the present analysis, air masses in which the \( \Delta NO_y \) value was less than 40 pptv were used for the reference relationship irrespective of the location in which they were sampled. A large portion of the data in the \( O_3 = 100–200 \) ppbv range was obtained off the west coast of the United States (flight October 9, 1997). These air masses had been transported by westerly winds from locations over the Pacific Ocean where aircraft emissions are climatologically not very high. The data obtained at latitudes lower than 45°N and higher than 60°N were also selected as background for both the UT and LS. The criterion of 40 pptv within 10 days used in this study was admittedly arbitrary and subjective. A value of 40 pptv was chosen simply because enough data points in each \( O_3 \) range were selected by this criterion. However, the resultant estimates of aircraft effects were not very sensitive to the choice of this criterion as long as it was relatively small. If real background air masses were based on the criterion of \( \Delta NO_y = 0 \) pptv, the data selected under the present criterion would underestimate the aircraft impact. When 40 pptv was chosen as the criterion, 42 and 37% of the UT and LS air masses sampled above 8.5 km were classified as background air masses. In air masses when the \( \Delta NO_y \) values were lower or higher than 40 pptv, the median \( \Delta NO_y \) value was 17 or 107 pptv, respectively, in the UT, suggesting a large difference in the effect of aircraft on the two sets of air masses. The same feature was also seen in the LS where the median value of \( \Delta NO_y \) was 14 and 107 pptv. The median value of the NOy mixing ratio in the background air mass was then calculated for each 10 ppbv \( O_3 \) range, and a linear relationship was calculated using these median values in the UT and LS separately as shown in Figure 4. The relationships in the UT and LS were smoothly connected at the \( O_3 \) value of 100 ppbv. Using the reference NOy-O3 relationship (referred as the "low \( \Delta NO_y \) reference" hereafter), the expected NOy value in the background air mass was calculated using the \( O_3 \) value simultaneously observed. The difference between this expected NOy value and the observed NOy value was defined as an excess NOy (\( dNOy \)).

For the second approach the CN concentration was chosen as the criterion. This is because within the plume of the aircraft, enhancements of the CN concentration were often observed simultaneously with enhancements of the NO and NOy [Anderson et al., 1999]. Consequently, relatively higher CN
values can be expected when an air mass is influenced by aircraft emissions. As shown in Figure 5, the CN concentration increased with altitude both in the UT and LS. These vertical profiles could partly be due to aircraft emissions which are most intensive at altitudes between 10 and 12 km. In fact, the analysis made by Kondo et al. [1999] showed that the CN fine particle concentration positively correlated with the NOx mixing ratio and NOx/NOy ratio in the LS air masses obtained during SONEX. Moreover, the slope of the positive correlation between the CN and NOx values was generally close to the median value of the ratio between the enhanced levels of NO3 (δNO3) and CN (δCN) found in the aircraft plumes (δNO3/δCN = 0.11 pptv STP cm-3). They concluded that an increase in CN particles due to aircraft emissions constituted 29% of the observed CN fine particle concentration. These results suggest that air masses with low CN values can be used as background in the LS. A positive correlation between the CN concentration and NO3/NOy ratio was also found in the UT in air masses with CN < 5000 STP cm-3 (Figure 6). Although the overall correlation was partly due to the vertical profile of CN and NO3/NOy, a positive correlation was generally found even within each 1 km layer and within limited O3 ranges. These results suggest that aircraft emissions played some role as a source of CN in the UT air masses sampled during SONEX. Using the SONEX data, Anderson et al. [1999] estimated that 8% of the CN fine particles in the northern hemispheric UT originated from aircraft emissions, based on the emission index of CN obtained within the aircraft plumes. However, it should be noted that when a clear enhancement of NO and NOy due to lightning activity was seen during the above mentioned four flights, an enhancement of CN concentration up to 3 × 104 STP cm-3 was also seen. New particle formation in the cloud-scavenged air around cumulous clouds can elevate the CN concentration in the UT [e.g., Clarke et al., 1998]. Consequently, although low CN air masses were considered to be relatively unaffected by aircraft emissions, high CN air masses were not necessarily affected greatly. For the present analysis, air masses with a CN concentration lower than 1500 and 1000 STP cm-3 were used to make a reference NO3-CN relationship in the UT and LS, respectively, irrespective of the location in which they were sampled. A different criterion was chosen because the CN concentration was systematically higher in the UT than in the LS (Figure 5). These data corresponded to 28 and 47% of the UT and LS data. As for the “low ΔNOx reference,” a median NOx value was calculated for each 10 ppbv O3 range, and a linear relationship was calculated for the UT and LS data separately (Figure 4, denoted as the “low CN reference” hereafter). The dNOx values were calculated for every 10-s data using this relationship. Note that data obtained from background air masses were not excluded and were used in conjunction with other data for the analyses described below.

It should be noted that products from three-dimensional

Figure 4. Two reference NO3-O3 relationships made using the background air masses obtained at altitudes between 8.5 and 11.5 km. The “low ∆NOx” air masses received less than 40 pptv NOx input from aircraft emission along the 10-day back trajectory based on the ANCAT/EC2 emission inventory. In the “low CN” air masses, CN concentration was less than 1500 or 1000 STP cm-3 in the UT and LS, respectively. The bars indicate the central 67% range.

Figure 5. Vertical profile of the median values of the CN fine particle concentration in the NAFC region. The bars indicate the central 67% range. The O3 values are in units of ppbv.

Figure 6. Correlation plot between NO3/NOy and CN fine particle concentration at three altitude ranges and two O3 ranges for air masses in the NAFC region. The data within each range were further divided into four subsets so that each subset contained the same amount of data. The median values and central 67% ranges were calculated for these subsets.
Table 1. Median Value of $d\text{NO}_y$

| Reference 1a | Reference 2b |
|--------------|--------------|
| pptv | Percent | pptv | Percent |
| NAFC region | | | |
| Troposphere | | | |
| 8.5–9.5 km | 8 (-37–47) | 3 (-15–17) | 2 (-46–41) | 1 (-19–15) |
| 9.5–10.5 km | 43 (-11–110) | 16 (-4–34) | 37 (-21–103) | 13 (-7–32) |
| 10.5–11.5 km | 71 (-19–167) | 21 (-6–35) | 66 (-29–152) | 18 (-10–34) |
| 8.5–11.5 km | 28 (-26–122) | 11 (-10–29) | 22 (-36–108) | 8 (-13–28) |
| Stratosphere | | | |
| 8.5–11.5 km | 128 (-25–217) | 16 (-3–27) | 110 (-41–199) | 14 (-5–25) |
| U.S. region | | | |
| Troposphere | | | |
| 8.5–11.5 km | 32 (-64–108) | 11 (-35–39) | 25 (-70–102) | 9 (-39–40) |
| Stratosphere | | | |
| 8.5–11.5 km | 89 (3–294) | 13 (0–31) | 71 (-13–277) | 10 (-2–29) |

The central 67% ranges are shown in parentheses.

aEstimates using the low $\Delta\text{NO}_y$ (model input NO value along 10-day back trajectory) reference.
bEstimates using the low CN reference.

3.2. Validity and Uncertainty in the $d\text{NO}_y$ Estimate

3.2.1. Reference values. The median values of the $d\text{NO}_y$ calculated using the two references are shown in Table 1 for both the UT and LS data. The median $d\text{NO}_y$ values for each 1 km layer are also shown for the tropospheric data in the NAFC region (Figure 7). In this case, the $d\text{NO}_y$ values were calculated down to 6.5 km, although the reference was derived using data from 8.5–11.5 km. The two sets of $d\text{NO}_y$ values were generally in good agreement when relatively large uncertainties were considered. The agreement between the two sets of $d\text{NO}_y$ values confirmed the validity of the method adopted in this study. The $d\text{NO}_y$ values calculated using the low CN reference were slightly lower when compared to those calculated using the low $\Delta\text{NO}_y$ reference. The median of the differences in the $d\text{NO}_y$ values was 8 and 18 pptv in the UT and LS, respectively, suggesting the uncertainty in the $d\text{NO}_y$ estimation caused by selecting the background air mass. Because the difference in the two sets of the $d\text{NO}_y$ values was small and systematic, the values calculated from the low $\Delta\text{NO}_y$ reference are used in the following discussion.

Although it is not shown here, similar results were also obtained when the $d\text{NO}_y$ value was calculated using the $\text{NO}_y$-$\text{NO}_x$ relationship in the LS instead of using $\text{NO}_y$-$\text{O}_3$.

3.2.2. Correlation. In Figure 8 a correlation plot between $d\text{NO}_y$ and $\Delta\text{NO}_y$ is shown for the UT NAFC data (8.5–11.5 km). In this figure the median $d\text{NO}_y$ values for each 50 pptv range in the $\Delta\text{NO}_y$ values and a one-to-one line are also shown. A large variability was generally seen in the $d\text{NO}_y$ values presumably reflecting the wide variety of air mass histories. However, the median values of $d\text{NO}_y$ generally increased with the $\Delta\text{NO}_y$ values suggesting that the averaged $d\text{NO}_y$ values were reasonable estimates for the aircraft impact on reactive nitrogen. The $d\text{NO}_y$ values were generally lower than the $\Delta\text{NO}_y$ values, partly due to the fact that when the $\Delta\text{NO}_y$ value was higher, the NOx value observed tended to decrease more because of dilution. On the other hand, the $d\text{NO}_y$ value in the UT did not correlate with CO or C$_2$H$_4$ in the NAFC and U.S. continent regions (not shown), suggesting that the high $d\text{NO}_y$ values were generally not due to the convective transport of polluted air masses.

Figure 7. Vertical profile of the $d\text{NO}_y$ values in the troposphere in the NAFC region calculated from the two reference $\text{NO}_y$-$\text{O}_3$ relationships. The median values and central 67% ranges are shown for each 1 km layer. The median values of the $\Delta\text{NO}_y$ (integrated value of the model NO input along the 5- or 10-day back trajectory) are also plotted for the same air masses.
3.2.3. Influence from lightning. One of the largest error sources, which can result in overestimating the $d\text{NO}_y$ values in the UT, is influence from NO production by lightning. As described above, the data, which had a clear signature of NO production by lightning (i.e., distinct enhancement in the NO and NO$_v$ mixing ratios and the NO$_x$/NO$_y$ ratio), were excluded from the present analysis. However, these were the only cases in which data were removed showing the influence of recent NO production by lightning. Lightning activity generally has an effect similar to aircraft emissions: it increases the NO$_x$ and NO$_y$ mixing ratios, the NO$_x$/NO$_y$ ratio, and the CN concentration. In order to estimate the effect of aircraft on air masses which had been minimally affected by lightning, $d\text{NO}_y$ values were calculated using only air masses which had remained over the Atlantic Ocean more than 90% of time within 5 days prior to the measurements. This criterion was used because most lightning activities occur over land [e.g., Turman and Edgar, 1982]. In addition, air masses which had passed near the Florida Peninsula were also excluded because of the intensive lightning activity in that region. The selected data consisted of 27% of the UT data in the NAFC region. As a result, the median $d\text{NO}_y$ values in the 9.5–11.5 km range where the largest $d\text{NO}_y$ values were obtained decreased by 24%. Because these air masses generally came from lower latitudes where aircraft NO emissions were lighter, the reduction in $d\text{NO}_y$ values could also have been the result of the change in the effect of aircraft. However, this change in the $d\text{NO}_y$ value provides an estimate of the degree of possible overestimation of the effect of aircraft in this study.

3.2.4. Stagnated air mass within the NAFC region. A clear indication that aircraft emissions had caused an increase in $d\text{NO}_y$ was seen in air masses sampled during the flight made at the eastern NAFC region on October 23, 1997 (Figure 1). The correlation plot of NO$_x$-$d\text{NO}_y$ for this flight is shown in Figure 9. Most of the $d\text{NO}_y$ was found to be in the form of NO$_x$, suggesting that the NO$_x$ input had been made within the lifetime of NO$_x$ of about 5 days (UT at 40ºN–60ºN). The 10-day back trajectories of the air masses obtained during this flight are shown in Figure 10. As shown in this figure, these air masses remained over the ocean at least 10 days and stagnated in the NAFC region the last 5 days. The median values of the O$_3$ and CO mixing ratios in the UT (8.5–11.5 km) were 37 and 77 ppbv, and they changed little throughout the flight. Consequently, most of the change in NO$_x$ and NO$_y$ was quite likely due to aircraft emissions. The median and maximum values of the $\Delta\text{NO}_x$ (10 days) were 71 and about 300 pptv, respectively, at altitudes between 9.5 and 11.5 km. Almost all of the input from aircraft was made within the last 5 days in this calculation. The median and maximum values of the $d\text{NO}_y$ were 42 and about 200 pptv, respectively, in the 9.5–11.5 km altitude range. These data clearly indicate that aircraft emissions can raise the levels of NO$_x$ and NO$_y$ over a large region, and the effect was successfully detected by the method used in this study.

3.3. Vertical Profile of $d\text{NO}_y$

In the NAFC region of the troposphere, the $d\text{NO}_y$ values were close to zero at altitudes between 7 and 9 km, suggesting minor impact from aircraft emissions (Figure 7). At altitudes above 9 km, $d\text{NO}_y$ increased with altitude, reaching 43 and 71 pptv at 10 and 11 km, respectively. (37 and 66 pptv at these two altitudes when the low CN reference was used.) The medians
of the $\Delta NOy/NOy$ ratio at these two altitudes were 16 and 21%; that is, about 20% of the NOy in the UT likely originated from aircraft emissions. These results are also summarized in Table 1.

As described above, the model input NOy value along the air mass trajectory, $\Delta NOy_x$, was calculated for each data point. Consequently, a statistical analysis can be made on the $\Delta NOy$ value for the same data set from which the $dNOy$ values were calculated. In Figure 7 the median values of $\Delta NOy$ for the 5- and 10-day trajectories are shown. The median $\Delta NOy$ (10 days) was 25 pptv at 9 km and increased with altitude reaching 160 pptv at 11 km. The shape of the profile is consistent with that of the $dNOy$ value, although the $dNOy$ values were systematically lower than the $\Delta NOy$ values, a likely result of the dilution effect.

The median value of $dNOy$ in the UT over the U.S. continent region was calculated using all of the data obtained at 8.5–11.5 km range (Figure 11 and Table 1). This was because the amount of data was not sufficient to derive a vertical profile of the $dNOy$ values. The median value was 32 pptv, comparable to the 28 pptv in the NAFC region. Although the data over the U.S. continent were obtained from only two flights and the amount of data was smaller (4790 versus 1539 10-s data), the results here suggest a similar effect from aircraft emissions in the two regions.

The median value of $dNOy$ in the LS was calculated for the 8.5–11.5 km range (Figure 11 and Table 1). The median values of $dNOy$ and $dNOy/NOy$ in the NAFC region were 128 pptv and 16%, respectively (110 pptv and 14% when the low CN reference was used.) The median $dNOy$ value over the U.S. continent region was 89 pptv, again comparable to the NAFC value when the large central 67% ranges are considered. It should be noted that these $dNOy$ values were systematically higher than those in the UT both in the NAFC and U.S. continent regions, 28 pptv versus 89 pptv in the NAFC region. The contrast in the $dNOy$ values between the UT and LS was much higher than that in the $\Delta NOy$ (10 days) values, at least in the NAFC region (72 pptv versus 102 pptv). One possible explanation is the longer residence time of NOy in the LS than in the UT. In the UT the NOy level can be lowered by dry and wet deposition of HNO3 and aerosol nitrates through the vertical mixing of air. In the LS, NOy cannot be lost until the air mass is transported into the troposphere. Gettelman [1998] estimated the residence time was around 50 days for aircraft exhaust released in the lowermost stratosphere (LS) using the NASA 1992 zonal mean aircraft emission database. During SONEX all the stratospheric air masses were sampled in the LS as described above. Because the tropopause height was lower during late autumn, a larger amount of NOy could have been injected into the LS, and the LS air masses might have been influenced by the accumulated effect.

**3.4. Relationship of $dNOy$ With NOy/NOx, CN, and $O_3$**

In Figures 12a–12d the correlation plots between the $dNOy$ and NOy/NOx, CN and $O_3$ values are shown for the UT and LS using the data from both the NAFC and the U.S. continent regions. For these figures we used only the data in which the model input NOx values during the last 5 days were greater than 80% of the 10-day values (i.e., $\Delta NOy$ (5 days)/$\Delta NOy$ (10 days) > 0.8). This is because NOx molecules emitted from aircraft more than 5 days prior to the measurements could have already been converted to higher oxidized species, such as HNO3. In fact, this data selection generally improved the degree of correlation. The lifetime of NOx was estimated to be 4.8 and 3.2 days in the UT and LS ($O_3 = 100–200$ ppbv), respectively, at 40°N–60°N using the observed values of aerosol surface area, etc. In Figure 12a the $dNOy$ values were plotted against the NOy/NOx ratios obtained in the UT. Different symbols were used for the data where $O_3 = 0–50$ ppbv and 50–100 ppbv, respectively. The NOy/NOx ratio generally increased with $dNOy$ in both data sets. This is consistent with the fact that emissions of NOx from aircraft increases both the NOy/NOx ratio and $dNOy$ value. In other words, the change in the NOy/NOx ratio observed in the UT during SONEX was partly due to NOx from aircraft emissions. A positive correlation observed in the air masses where $O_3 = 0–50$ ppbv was mostly due to those sampled on October 23, 1997, which had been strongly affected by aircraft emissions as described above. The degree of the NOy/NOx increase was higher for air masses where $O_3 = 0–50$ ppbv than that for air masses where $O_3 = 50–100$ ppbv when data with the same $dNOy$ value were compared. This result is plausible because the NOy value was generally lower when $O_3$ was low (Plate 1) and the NOx input could cause a greater change in the NOy/NOx ratio.

In Figure 12b, $dNOy$ values are plotted against the CN concentration obtained in the UT. Kondo et al. [1999] and Anderson et al. [1999] calculated the ratio between enhanced levels of NOy ($\delta NOy$) and CN (6CN) observed within aircraft plumes. The $\delta NOy/6CN$ ratio ranged between 0.04 and 0.2 ppv STP cm$^{-3}$ with a median value of 0.11 ppv STP cm$^{-3}$. In Figure 12b, two lines having this slope and different intercepts are also plotted. As seen in this figure, the $dNOy$ value has a weak positive correlation with the CN concentration in both the air masses where $O_3 = 0–50$ ppbv and 50–100 ppbv. On the basis of the passive cavity aerosol spectrometer probe (PCASP) measurements the median value of the aerosol surface area for aerosol particles with diameters between 0.1 and $3.0 \mu m$ is 1.7 $\mu m^2$ cm$^{-3}$ in the UT in the NAFC region. Under this condition, CN is lost primarily by the Brownian coagulation of CN, for CN concentrations higher than about 1400 STP cm$^{-3}$. The half-lifetime of CN with a diameter of 15 nm at a CN concentration of 2500 STP cm$^{-3}$ is 1.3 days and decreases inversely with the CN mixing ratio [Pruppacher and Klett, 1977; B. Liley, personal communication 1999]. Because
the lifetime of CN was shorter than that of NO$_x$ and NO$_y$, the CN concentration should be generally lower than that expected from the slope in the aircraft plumes if aircraft emissions were the dominant source of CN. Consequently, the observed result was not inconsistent with the hypothesis that aircraft emissions contributed to a source of CN for air masses where O$_3$ = 50–100 ppbv. On the other hand, other CN sources were predominant for air masses where O$_3$ = 0–50 ppbv because much higher CN concentrations than those expected from aircraft emissions were observed in these air masses. In fact, no correlation between CN and NO$_x$/NO$_y$ was found for air masses where CN > 5000 STP cm$^{-3}$ as described above (Figure 6). However, it should be noted that a positive correlation having a similar slope to that within the aircraft plumes was seen in some air masses. As described above, air masses sampled on October 23, 1997, had probably been strongly affected by aircraft emissions (see subsection 3.2.4), and the O$_3$ values in these air masses were mostly below 50 ppbv. The data where NO$_x$ was higher than 60 pptv obtained on this day are shown using a different symbol in Figure 12b, and their CN-dNO$_y$ slope was found to be generally similar to that observed within the aircraft plumes. Consequently, aircraft emissions could have caused an increase in the CN concentration in the UT in some cases during SONEX, resulting in the positive correlation between CN and dNO$_y$, and CN and NO$_x$/NO$_y$, although the background level of the CN concentration was changed greatly by other CN sources.

In Figure 12c the correlation plot between dNO$_x$ and the NO$_x$/NO$_y$ ratio in the LS is shown. A positive correlation was generally found in the LS. Because the LS air mass was generally free from tropospheric reactive nitrogen sources, the results shown here strongly suggest that an NO$_x$ input from aircraft emissions primarily caused the NO$_x$ variation in the LS.

In Figure 12d the correlation plot between dNO$_y$ and CN in the LS is shown. A clear positive correlation between the two parameters was found suggesting that aircraft emissions were an important source of CN in the LS. The slope between dNO$_y$ and CN was generally smaller as compared to the median 3NO$_x$/6CN ratio of 0.11 pptv STP cm$^{-3}$ observed in aircraft plumes. This is reasonable because the lifetime of NO$_y$ is much longer than that of CN in the LS.
In Figures 13a and 13b the correlation plots between dNOy and O3, and NOx/NOy and O3, are shown using both the UT and LS data in the NAFC region. The median values of dNOy or NOx/NOy in seven O3 ranges are also shown. Systematically large dNOy values were generally found with O3 values between 75 and 125 ppbv. The dNOy values were generally distributed around zero for lower and higher O3 air masses. These features are quite similar to those of the correlation plot between NOx/NOy and O3, suggesting a relatively greater effect from aircraft emissions in the air masses with O3 values between 75 and 125 ppbv. These air masses likely remained around the tropopause altitude where NOx input from aircraft emissions is large. Air masses with a lower O3 mixing ratio in the UT could generally be more strongly influenced by lower-altitude air. This is consistent with the fact that the dNOy values were close to zero when the CO and C2H2 values were high. Similarly, air masses with O3 values higher than 125 ppbv are considered to be less affected by aircraft emissions. Further decreases in the NOx/NOy ratio with O3 values between 125 and 300 ppbv were likely because the loss rate of NOx via heterogeneous hydrolysis of N2O5 is faster for higher O3 values. Similar O3 dependences of dNOy and NOx/NOy values were also seen in the air masses obtained over the U.S. continent in the UT and LS. In this case, high NOx/NOy ratios of up to 0.3 were observed with O3 values as high as 160 ppbv.

3.5. Estimation of dNOx in the UT

The ratio between excess NOx (dNOx) and dNOy was estimated from the following expression:

\[
\frac{dNO_x}{dNO_y} = \left( \frac{NO_x}{NO_y} - \left( \frac{NO_x}{NO_y} \right)_{\text{ref}} \right) \times (1 - \gamma) \times \frac{1}{\gamma}
\]

where \(\gamma = \frac{dNO_y}{NO_y}\). For the reference NOx/NOy ratio we used the background air masses that were used to make the reference NOy,O3 relationship, that is, air masses with ANOy values (model NOx input along the 10-day back trajectory) lower than 40 pptv. In Figure 14 a vertical profile of the median NOx/NOy ratio in background air masses is shown with that in air masses in the NAFC region. The NOx/NOy ratios in the NAFC region were systematically higher than those in the background air masses. Using these NOx/NOy values and the dNOy/NOy values shown in Table 1 (low ANOy (10 days) reference), dNOx/dNOy was calculated for each 1 km altitude level. As a result, the dNOx/dNOy ratio was estimated to be 0.54 and 0.38 at altitudes of 10 and 11 km, respectively. Even if the median NOx/NOy values in the low CN air masses were used for the reference, similar results were obtained (Figure 14). Furthermore, even if only the data at 45°N–60°N were used for the reference to eliminate a possible effect from the change in the NOx/NOy ratio with latitude, similar results were obtained. These results suggest that about half of the excess NOx was likely to be in the form of NOx in the UT. Using an average dNOx/dNOy value of 0.46, we found the excess NOy to be 20 and 33 pptv at 10 and 11 km, respectively. The dNOx/NOy ratio can be also expressed using the NOx/NOy ratio and dNOx/NOy ratio by transforming (1). Using the same values shown above, we estimated that 30% of the NOy at 10 and 11 km was likely to have originated from aircraft emissions in the UT.

Figure 13. Correlation plot between (a) dNOy,O3 and (b) NOx/NOy,O3 for the data obtained in the NAFC region. The median values and central 67% ranges of the dNOy and NOx/NOy ratio are also shown for the seven O3 ranges.
4. Summary

The large-scale impact of aircraft emissions on reactive nitrogen in the UT and LS was estimated using the NO$_x$-O$_3$ correlation observed during SONEX, which was carried out over the U.S. continent and NAFC region in October and November 1997. For the present analysis we excluded data obtained within aircraft plumes, data clearly influenced by NO production from lightning, and data influenced by recent convective transport of air affected by surface sources. Two reference NO$_x$-O$_3$ relationships in the air masses, which were considered to be affected little by aircraft emissions, were made using the data obtained above 8.5 km. First, the integrated input of NO into an air mass along a 10-day back trajectory ($\Delta$NO$_x$) was calculated based on the ANCAT/EC2 aircraft emission inventory. When the $\Delta$NO$_x$ value was less than 40 pptv (lower 42 and 37% values in the UT and LS), that air mass was used for the reference NO$_x$-O$_3$ relationship, irrespective of the location from which it was sampled. Second, air masses with a CN concentration lower than 1500 and 1000 STP cm$^{-3}$ (lower 28 and 47% values) in the UT and LS were used to make the reference. Using these two reference NO$_x$-O$_3$ relationships, the two sets of the excess NO$_x$ (dNO$_x$) were calculated from observed values of NO$_x$ and O$_3$.

The two sets of dNO$_x$ values generally agreed within 8 and 18 pptv in the UT and LS, respectively, in spite of the very different nature of the criteria for the background air selection. The calculated dNO$_x$ values did not positively correlate with either the CO values or anthropogenically produced hydrocarbons such as C$_2$H$_2$, suggesting a negligible effect of the surface sources on the dNO$_x$ values. On the other hand, the weak positive correlation between the dNO$_x$ and NO$_x$/O$_3$ values indicated that the dNO$_x$ values generally reflected the effect of aircraft emissions. By calculating the dNO$_x$ values only for the air masses which remained over the ocean for 5 days prior to the measurement, the uncertainty in the dNO$_x$ values due to the influence of recent NO production by lightning was estimated to be 24%.

In the NAFC region (45°N–60°N) the median value of dNO$_x$ in the troposphere was close to zero between 7 and 9 km and increased with altitude above 9 km reaching 70 pptv (20% of NO$_x$) at 11 km. High values of dNO$_x$ and NO$_x$/O$_3$ ratio were generally found in air masses in which O$_3$ = 75–125 ppbv, suggesting a stronger effect around the tropopause. The median value of dNO$_x$ in the LS in the NAFC region at 8.5–11.5 km was about 120 pptv. The larger dNO$_x$ value observed in the LS was probably the result of the accumulated effect of aircraft emissions, given the long residence time of the affected air in the LS (50 days).

In general, the dNO$_x$ value positively correlated with the NO$_x$/O$_3$ ratio in both the UT and LS in the NAFC and the U.S. continent regions, suggesting that the observed large-scale changes in the NO$_x$/O$_3$ levels and NO$_x$ values in the UT and LS were partly caused by NO$_x$ from aircraft emissions. The slope of the positive correlation between the dNO$_x$ and CN values in the LS was generally consistent with that observed in the aircraft plumes, suggesting the importance of aircraft emissions as a source of CN in the LS. A positive correlation between the dNO$_x$ and CN values was also seen in a part of air masses in the UT, and its slope was not inconsistent with that observed in the aircraft plumes; however, the significance of aircraft emissions is still not clear.

The ratio between excess NO$_x$ (dNO$_x$) and dNO$_x$ was calculated using the reference NO$_x$/NO$_3$ ratio. In the UT (NAFC region) it was estimated that about half of dNO$_x$ was in the form of NO$_x$ (20 and 33 pptv at 10 and 11 km) and about 30% of the NO$_x$ had originated from aircraft emissions. These estimates were generally consistent with the results from the model calculations.

In this study, the degree of the large-scale impact of aircraft emissions on reactive nitrogen was estimated from the observed data. A more quantitative estimate will be made using sophisticated three-dimensional models when they can consistently reproduce most of the observed features of various species.

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