Springtime photochemical ozone production observed in the upper troposphere over east Asia

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[1] Aircraft observations of ozone and its precursors (NO, NOx, CO, and nonmethane hydrocarbons) were made near Japan (26°–44°N) between 21 and 24 April 1998 to investigate the effect of transport and chemistry on the tropospheric ozone over east Asia in spring. The average mixing ratios of ozone and its precursors in the upper troposphere were higher than those observed during February–March 1994. Significantly higher values of CO indicate that the influence of surface sources mediated by convection was large throughout the troposphere during the period. Highly polluted air masses were observed in the upper troposphere at 8–11 km over the Japan Sea on 24 April. These air masses were influenced by cumulus convection associated with a cold front over northeast China about 1 day prior to the observation. However, the majority of observed air masses in the upper troposphere were not directly affected by the recent emissions of the ozone precursors from east Asia. Instead, convection over other regions in the northern midlatitude, followed by long-range transport, affected the abundances of ozone precursors in the upper troposphere. In these air masses, ozone was positively correlated with NOx and C3H8, indicating the effect of photochemical ozone production. The diurnal-average column-integrated rate of ozone production, estimated by a photochemical box model, was larger than the NH average and local stratospheric flux at NH midlatitudes in spring by a factor of 3–20. The net production rates in the majority of the air masses were estimated to be 0.5–4.4 ppbv d–1 in the upper troposphere. These results indicate the important role photochemistry plays in controlling the upper tropospheric ozone abundance in spring.

INDEX TERMS: 0368 Atmospheric Composition and Structure: Troposphere—constituent transport and chemistry; 0345 Atmospheric Composition and Structure: Pollution—urban and regional (0305); 0365 Atmospheric Composition and Structure: Troposphere—composition and chemistry; KEYWORDS: Asian outflow, ozone, ozone precursors, convective transport, western Pacific

1. Introduction

[2] East Asia is the region where rapidly growing industrial activities are causing large increases in emissions of photochemically active pollutants, such as CO, NOx (= NO + NO2), and nonmethane hydrocarbons (NMHCs). For example, emissions of NOx in East Asia are anticipated to increase by a factor of 5 from 1990 to 2020 [van Aardenne et al., 1999]. Emissions of these active pollutants from this continental region have potential impacts on the tropospheric ozone budget over the North Pacific region due to the eastward transport of the polluted air masses in the free troposphere by strong westerlies at midlatitudes. Knowing the impact of surface pollutants in the present-day atmosphere will enable us to anticipate how ozone will respond to future changes.

[3] The transport of pollutants from east Asia to the North Pacific is generally most effective in spring, due to a combination of active convection over the continent and strong westerlies [Merrill, 1989]. Aircraft observations showed high levels of total reactive nitrogen (NOx) in the free troposphere over the Hawaiian region in spring, probably due to export from northern latitude continental surface regions [Ridley et al., 1997]. Transport of Asian pollution, such as CO and NMHCs, to the northwestern contiguous United States during spring was observed by ground-based...
measurements [Jaffe et al., 1999]. In addition to the transport of pollution, solar ultraviolet radiation intensifies as spring advances because of changes in solar zenith angle and total ozone [Yienger et al., 1999]. Active photochemistry combined with transport can contribute to the springtime increase of ozone in the free troposphere over east Asia, as shown by the recent analysis of ozonesonde data obtained over Japan [Logan, 1999].

[4] In situ and remote measurements of an extensive suite of trace gases were made over the western Pacific during the Pacific Exploratory Mission-West (PEM-W)-A in September–October 1991 and PEM-W-B in February–March 1994 to investigate chemical processes and long–range transport of trace species over the northwestern Pacific and to estimate the impact of human activities on these species over this region [Hoell et al., 1996, 1997]. Comparisons of these data indicate that the air masses sampled in early spring (PEM-W-B) were more polluted than those sampled in fall (PEM-W-A). A first indicator for the effects of surface pollutants is the hydrocarbon data. Below 5 km, the mixing ratios of hydrocarbons, which originated from the continental surface, were much larger during PEM-W-B than those during PEM-W-A due to the strong outflow from the Asian continent [Blake et al., 1997; Gregory et al., 1997; Kondo et al., 1997a; Talbot et al., 1997]. However, in the upper troposphere, mixing ratios of pollutants were smaller than those below 5 km during PEM-W-B because of the lack of deep convection, which transports pollutants from the lower troposphere, as is typical for winter/early spring [e.g., Newell et al., 1997]. An implicit assumption in this discussion is that air parcels below/above 5 km sampled by the aircraft came from below/above 5 km upwind. Thus the trace gas abundance measured by the aircraft over the Pacific could be used as a proxy for that over the Asian continent.

[5] A second indicator of the effects of Asian outflow is the column ozone budget. Photochemical model calculations indicate that net ozone production occurred between 0–12 km at 20°–50°N during PEM-W-B, in contrast to PEM-W-A where net ozone loss at lower altitudes balanced net production at higher altitudes at 18°–42°N [Crawford et al., 1997].

[6] Comprehensive observations in April, when stronger convective activities are expected to transport air from below 5 km to the upper troposphere, would better quantify the role of sources on springtime ozone increases in the upper troposphere over east Asia. As a part of the Biomass Burning and Lightning Experiment (BIBLE) campaigns, three aircraft observation flights were made near Japan in April 1998, called BIBLE T. Major objectives of this study were (1) to study the influence of continental outflow on the distributions of trace gases up to the tropopause over east Asia in April, and (2) to study the relative contribution of photochemical production to the spring increase of tropospheric ozone over these regions. In this paper, effects of convective transport and photochemical ozone production in the upper troposphere are discussed based on the observed data.

2. Aircraft Observations

[7] During the BIBLE T experiment, atmospheric sampling was conducted aboard the Gulfstream (G-II) aircraft at 26°–44°N near Japan. The base of operations was Nagoya International Airport. The horizontal flight tracks are shown in Figure 1. The intensive aircraft observations were carried out on three flights on 21, 22, and 24 April after two test flights. The G-II aircraft flew from Nagoya south to 26°N over the Pacific on April 21 (flight 3). On 22 April (flight 4), the aircraft flew east and then went north, over the western Pacific. Measurements were made up to 44°N over the Japan Sea on 24 April (flight 5). The G-II aircraft flew mostly in the upper troposphere above 7 km and the maximum flight level was 13.2 km. Altitude profiles of constituents were also obtained at two locations as shown in Figure 1.

[8] During BIBLE T, ozone, CO, NO, NOy, H2O, NMHCs, and meteorological parameters were measured. The instruments aboard the aircraft and their characteristics are listed in Table 1. Ozone was measured by a UV absorption photometer with a detection limit of 3 ppbv at 13 km and less than 3 ppbv at lower altitudes [Kita et al., 2002]. The CO mixing ratio was measured by an automated gas chromatograph (GC) with a reduction gas detector (RGD) [Kita et al., 2002]. It was measured with a time interval of 20 s using three independent GC-RGD units. Because of the uncertainty of the CO standard gases used for the GC-RGD during this experiment, the accuracy of the 20-s data was 10% at 200–300 ppbv, 20% at 100–200 ppbv, and 30% at 0–100 ppbv. NO and NOy measurements were made using a chemiluminescence technique with a 1-s time resolution. The precision of the 10-s NO and NOy measurements at 10 km was 3 and 5 pptv for NO and NOy values of 100 and 200 pptv, respectively [Kondo et al., 1997b; Koike et al., 2000]. The actinic flux was measured using UV radiometers to calculate the photolysis rate
coefficient of NO2 (J(NO2)) [Junkerman et al., 1989]. The J(NO2) filter radiometer was absolutely calibrated by Meteorologie Consult (Glashütten, Germany) and the precision was $2 \times 10^{-4}$ s$^{-1}$ for 10-s data. The photostationary state of NO$_2$/NO ratios above 8 km were calculated using the measured values of ozone, temperature, pressure, and J(NO2). The ratios were calculated for solar zenith angles lower than 67°. The NOx mixing ratios were calculated from these NO$_2$/NO ratios and the observed NO mixing ratios. The dew point was measured with two different types of dew point hygrometers (model 1011A provided by General Eastern and model CR-2 provided by Buck Research). The lower detection limits are 75°C for the 1011A and 95°C for the CR-2. The H$_2$O mixing ratios were derived from these measurements over the ranges of 1–100,000 ppmv (1011A) and 0.01–27,000 ppmv (CR-2). Whole air samples were collected into 2-L stainless steel sampling canisters and more than sixty samples were obtained per flight. The samples were analyzed for a suite of NMHCs in the laboratory at the University of California, Irvine [Blake et al., 1996].

3. Results and Discussion

3.1. Altitude Profiles

Figures 2a–2e show the altitude distributions of ozone, CO, C$_3$H$_8$, NO, and NO$_y$ observed during 21–24 April (flights 3–5). Ozone, CO, NO, and NO$_y$ values averaged for 10-s were used. The median values are also shown in the right panel with 1-km resolution. Data obtained during takeoff and landing in Nagoya below 2 km were excluded because these data showed a large variability due to local pollution. Data below 2 km obtained over the ocean during fights 3 and 5 were used. The bars shown in the median profiles indicate the range spanned by 67% of the data around the median values. Because the number of data below 2 km is limited, the median values below 2 km may not be representative for the BIBLE T period. During these flights, the measurements were made below the tropopause, which was defined by the temperature obtained by the routine radiosonde observations at several sites in Japan by the Japan Meteorological Agency (JMA). Figures 2a–2e also show altitude profiles of the median values observed during PEM-West A (September–October 1991) and PEM-West B for comparison. The PEM-W-A data obtained at the same latitudes (26°–44°N) as for BIBLE T were used. The data used in this comparison were obtained during flights 5–13 (17 September to 6 October 1991). The PEM-W-B data shown in Figures 2a–2e were limited to 20°–30°N because the tropopause height north of 30°N was as low as 8–9 km [Koike et al., 1997], and the data obtained above 8 km north of 30°N were from the lowermost stratosphere. In order to compare BIBLE T data with those at 0–12 km in the troposphere, the PEM-W-B data obtained north of 30°N

| Table 1. Parameters and Measurement Techniques for the BIBLE T |
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| **Species/Items** | **Techniques** | **Sampling Time Interval** | **Precision (10 s)** | **Accuracy** |
| O$_3$ | UV absorption | 1 s | <3 ppbv | 5% |
| CO | GC/HgO-reduction | 20 s | 2 ppbv | 5% |
| NO, NO$_y$ | Chemiluminescence | 1 s | 3 pptv (NO), 5 pptv (NO$_y$) | 8% (NO), 17% (NO$_y$) |
| Actinic Flux | UV radiometer | 1 s | $2 \times 10^{-4}$ s$^{-1}$ | 8% |
| H$_2$O | Dew point hygrometer | 10 s | 0.2–1.0°C | <5% |
| NMHCs | Whole air sampling/GC | 5 min | <3 pptv | 2–20% |

Figure 2a. Altitude profiles of the observed ozone mixing ratios during BIBLE T. The 10-s averaged data was used. The median values shown in the right panel are compared with those observed during PEM-West A (September–October 1991) and PEM-West B (February–March 1994). The bars indicate the central 67% values. Highly polluted air masses observed during flight 5 are also shown by crosses.

The parameters and measurement techniques for the BIBLE T are shown in Table 1. The measurement techniques include UV absorption, GC/HgO-reduction, Chemiluminescence, UV radiometer, Dew point hygrometer, and Whole air sampling/GC.
were excluded. The PEM-W-B data used were obtained during flights 7, 8, and 12–16 (13 February to 7 March 1994). Both of the PEM-W-A and B flights used here were made at the same longitude (120°–150°E) as BIBLE T. During both PEM–W-A and PEM-W-B, NO and NOy measurements were made by Nagoya University (NU) using a chemiluminescence technique [Kondo et al., 1996, 1997b] and by the Georgia Institute of Technology using a laser-induced fluorescence technique [Sandholm et al., 1997]. Comparisons of NOy measured by NU and the sum of reactive nitrogen species showed good agreement over the Pacific [Kondo et al., 1997a] and the North Atlantic (slope >0.9 and $r^2 \approx 0.9$) [Talbot et al., 1999]. In this paper, the NO and NOy values measured by NU during PEM-W-A and PEM-W-B were used. The median ozone mixing ratios during BIBLE T were 71–84 ppbv at 7–13 km and 55–62 ppbv below 7 km. The ozone concentrations during BIBLE T were higher than those observed during PEM-W-A and PEM-W-B, by 11–35 ppbv at all altitudes. Although all the data were obtained below the tropopause, stratospheric intrusion likely influenced the median ozone value of 170 ppbv around 13 km.

[10] Figure 3a compares the median ozone mixing ratios during BIBLE T with ozonesonde data obtained in Japan in April for the years 1993–1997. The ozonesonde observations were routinely conducted at four stations in Japan: Sapporo (43.1°N, 141.3°E), Tateno (36.1°N, 140.1°E), Kagoshima (31.6°N, 130.6°E), and Naha (26.2°N, 127.7°E), about once a week by JMA. Data obtained above the tropopause were excluded. The BIBLE T data may not be representative of conditions in spring because the data were obtained over a period of three consecutive days, so

Figure 2b. Same as Figure 2a but for CO.

Figure 2c. Same as Figure 2a but for C3H8. The 5-min data were used.
the results might well have been influenced by synoptic scale events. However, comparison with ozonesonde data shows that the aircraft ozone median values agree well with the median values of the 5-year ozonesonde data between 2–12 km (50–85 ppbv), within a variability of ±15 ppbv. The median ozone mixing ratios during PEM-W-A and PEM-W-B also agree with those of the 5-year ozonesonde data (30–65 ppbv) within a variability of approximately ±20 ppbv for September–October (Figure 3b) and for February–March (Figure 3c), respectively. Therefore the difference in the ozone levels during BIBLE T and PEM-W-B should be mainly due to the lack of convective activities during PEM-W-B compared to BIBLE T, which is a reflection of the seasonal variation of synoptic scale meteorology.

[11] During BIBLE T, the CO mixing ratios at 5–12 km were fairly constant at about 200 ppbv, which were higher than those observed during PEM-W-A and PEM-W-B by a factor of 1.5–2, as shown in Figure 2b. At 2–5 km, the CO mixing ratios during BIBLE T were similar to those during PEM-W-B, but higher than those during PEM-W-A by a factor of 1.5, as anticipated from the seasonal variation at the surface: a maximum in late winter/spring and a minimum in summer/early autumn at remote sites in the Northern Hemisphere [Novelli et al., 1994, 1998]. In addition to the seasonal cycle of CO at the surface driven by the reaction with OH, increased CO emissions from biomass burning in late winter-spring in the Northern Hemisphere [e.g., Galanter et al., 2000] could enhance the whole CO column as observed during BIBLE T. It

Figure 2d. Same as Figure 2a but for NO. Several spikes due to aircraft emissions larger than 800 pptv were excluded.

Figure 2e. Same as Figure 2a, but for NO$_y$. Several spikes larger than 2.5 ppbv were excluded.
should be noted that the vertical gradient of the CO mixing ratios at 4–8 km during BIBLE T and PEM-W-A was much smaller than that during PEM-W-B. The small vertical gradient was also seen for C3H8 (Figure 2c) during BIBLE T. The lifetimes of CO and C3H8 are roughly 2 months and 10±6 days, respectively (depending on the OH concentration). Since both CO and C3H8 are mainly emitted from the surface, these altitude distributions indicate that major convective activity occurred at least once within 10 days prior to the flights and throughout the troposphere at northern midlatitudes. Profiles for shorter-lived hydrocarbons indicate that 15% of the samples experienced convective activity within the last 2 days, as discussed in section 3.2.

The median NO and NOy mixing ratios at 2–7 km were 10–20 pptv and 240–480 pptv, respectively (Figures 2d and 2e). They were similar to those observed during PEM-W-A and PEM-W-B. The median NO and NOy mixing ratios at 7–13 km were 40–350 pptv and 440–1230 pptv, respectively. The NO values above 9 km during BIBLE T were much higher than those during PEM-W-A, although they were more or less similar to those during PEM-W-B. The increase in the NO mixing ratios with altitude has also been previously observed \[\text{Liu et al.}, 1980; \text{Kley et al.}, 1981; \text{Drummond et al.}, 1988; \text{Ridley et al.}, 1994\]. This increase is due to the longer chemical lifetime of NOx and additional sources of NOx in the upper troposphere, such as lightning, aircraft emissions, and stratospheric intrusions. The BIBLE T NOx mixing ratios were similar but slightly higher than those during PEM-W-A and PEM-W-B above 7 km. There is no simple explanation for the lower concentrations of NO at 2–7 km during BIBLE T in contrast to the higher concentrations of CO and C3H8, because of the shorter lifetime of NO compared to those of CO and C3H8, together with the fewer number of data samplings below 7 km than above.

3.2. Identification of Air Masses Influenced by Pollutants From East Asia

The origins of air masses sampled in the upper troposphere were investigated using 10-day isentropic back trajectories. The trajectories were calculated with the trajectory model developed by \text{Matuzono et al.} [1998], using the European Centre for Medium-range Weather Forecasts (ECMWF) 2.5° resolution data. Figure 4 gives a summary of trajectories for air masses sampled above 7 km. Almost all the air masses were transported over the Asian continent by the dominant westerlies in the upper troposphere at northern midlatitudes. Above 10 km, the air masses were transported from over the Atlantic Ocean and even the U.S. continent within 10 days. Although the range of latitude that the trajectories passed varied from flight to flight, the flow of the air masses were mostly confined to the 35°–45°N latitude band. These trajectories suggest that the observed air masses were not significantly affected by high-latitude or tropical air. The air masses identified to be strongly influenced by east Asian pollution are discussed below in this section.

The profiles of C3H8 indicate that convective activities in April occurred with sufficient frequency that they maintained high upper tropospheric concentrations of gases with atmospheric lifetimes of 10 days. The C2H2/CO ratio can be used as an indicator of the relative degree that air masses have been photochemically processed on a timescale of 1–3 weeks, which is the photochemical lifetime of C2H2 \[\text{e.g., Smyth et al.}, 1996\]. In the study of remote western Pacific air during PEM-W-A, C2H2/CO values of 1 pptv/ppbv or lower were identified as photochemically well-aged
Although the ratio contains uncertainty due to the complexity of sources and atmospheric mixing, the median C\textsubscript{2}H\textsubscript{2}/CO value of 1.2 pptv/ppbv above 7 km indicates that a majority of the air masses were photochemically aged. Convective transport over the Asian, European, and North American continents all contribute to the observed high upper tropospheric concentrations. The effects of emissions and vertical transport could not be separated from each continent due to the complexity of source distributions and transport, given lifetimes of 10 days during which the air masses were transported over very long distances in the free troposphere. However, the origin of some air masses observed in the upper troposphere on 24 April has been identified to be the boundary layer over east Asia as detailed below.

In order to identify the location of the convection, 5-day backward trajectories, combined with the cloud image data obtained by the Japanese Geostationary Meteorological Satellite-5 (GMS-5), were used. The resolution of the cloud images was 0.25° in latitude and longitude for the infrared (IR)-1 (λ = 10.5–11.5 μm) and IR-2 (λ = 11.5–12.5 μm) channels. The difference in the brightness temperature (DBT) between the 2 channels depends sensitively on the optical thickness of the clouds. In this analysis, optically thick clouds were defined as those with DBT values smaller than 1 K [Inoue, 1985, 1989]. The air temperature at a position of an air mass along its back trajectory was compared with the brightness temperature at the position for each 1-hour period. The air mass was considered to have been affected by the outflow of convective clouds if the brightness temperatures were lower than the air temperatures at the time when the horizontal locations of the air masses overlapped with those of the optically thick clouds.

Table 2. Observed Values of Chemical Species Between 8 and 11 km at 43°–44°N, 137°E Over the Japan Sea on 24 April\textsuperscript{a}

| Chemical Species | CO, ppbv | O\textsubscript{3}, ppbv | C\textsubscript{2}H\textsubscript{2}, pptv | NO, pptv | NO\textsubscript{x}, pptv |
|------------------|---------|----------------|-----------|--------|------------------|
| Japan Sea (24 April) | 250–317 | 83–96 | 260–442 | 160–680 | 1080–2030 |
| Median values during BIBLE T | 210 | 78 | 121 | 130 | 541 |

\textsuperscript{a}Median values at 8–11 km during BIBLE T are also shown for reference.
convective transport associated with a cold front over east Asia can transport a significant amount of the ozone precursor species from the boundary layer to the upper troposphere in spring. A global 3-D model study of Bey et al. [2001] has suggested the importance of frontal lifting as a mechanism of the export of CO from China and formation of an Asian pollution plume in the middle troposphere over the Pacific during PEM-W-B.

3.3. Correlations Analysis

We now examine the correlations of different species above 7 km. In this analysis, we excluded data strongly impacted by the recent convection over the northeastern part of China as discussed in 3.2. The meteorological analysis using the same method indicates that the rest of the air masses were not impacted by recent (within 2 days) convection over east Asia, consistent with the low C2H2/CO ratios in the majority of the sampled air masses, as discussed in 3.2.

Also as described in 3.1, some influence of stratospheric intrusions was seen around 13 km, just below the tropopause. The CO mixing ratios at these altitudes were 110–160 ppbv, which was lower than those below 13 km. The low H2O mixing ratios of 15–60 ppmv (not shown) were similar to those observed in stratospheric air masses at midlatitudes [Kondo et al., 1997a; Kotamarthi et al., 1997]. The ozone and NOx mixing ratios were mostly higher than 100 ppbv and 500 pptv, respectively, as seen in Figures 2a and 2e. These low values of CO and H2O, together with high values of ozone and NOx, indicate that the air masses obtained around 13 km were affected by stratospheric input. These stratospherically influenced data were also excluded from the following analysis.

Correlations among ozone and its precursors can be used to investigate contributions of emissions of precursors from continents. The NOx mixing ratios derived from the measured NO and the NOy mixing ratios in these air masses are plotted against the C3H8 mixing ratios in Figure 7. The NOx mixing ratios are well correlated with the C3H8 mixing ratios (r2 = 0.67). On the contrary, NOy showed no significant correlations with CO (not shown). The lifetime of NOy, after emission from the surface, is controlled by heterogeneous processes. These results show that effective loss of NOy occurred with a time constant similar to the lifetime of C3H8 (about 10 days), which is determined by the reaction with OH. Because the lifetime of CO is about 2 months, NOy is lost much more rapidly than CO, reducing the NOy-CO correlation in moderately aged air masses. Mixing of fresh pollution or moderately aged air with background air (well-aged air) will further weaken the NOy-CO correlation because NOy is more depleted than CO in background air.

The NOx-C3H8 correlation is somewhat more scattered (r2 = 0.48) than the NOy-C3H8 correlation. Conversion of NOx into HNO3 and PAN mainly controls the lifetime of NOx after emission from the surface and transport to the upper troposphere. The lifetime of NOx is shorter than NOy and C3H8, leading to the NOx-C3H8 correlation more scattered than the NOy-C3H8 correlation.

These results suggest that NOx and NOy in these air masses were supplied dominantly by the upward transport of polluted air near the surface within the photochemical lifetimes of C3H8. However, during upward transport by deep convection, lightning can add a significant amount of NO to the polluted air masses rich in NOx, CO, and NMHCs [e.g., Liu et al., 1999]. Therefore these correlations cannot,
Figure 6. Temporal variation of cloud distributions during a time interval of about 12 hours before the observation. Marks indicate the positions of air masses in which enhancements of ozone precursors were observed at 8–11 km. They were estimated by the trajectory calculations.
by themselves, unambiguously separate the contribution of lightning from the production by anthropogenic activities near the surface.

Ozone was correlated with NOy ($r^2 = 0.66$) and C$_3$H$_8$ ($r^2 = 0.56$) in these air masses independent of the flights, as shown in Figures 8 and 9. Generally, a tight, positive correlation between ozone and NOy indicates air masses that have been photochemically processed in the troposphere [Buhr et al., 1996; Daum et al., 1996] and/or influenced by downward transport from the stratosphere [Murphy et al., 1993; Koike et al., 1997] (Figure 8). In addition to this, the positive correlations of NOx and NOy with C$_3$H$_8$, as well as the high mixing ratios of CO and C$_3$H$_8$, excludes significant effects of stratospheric air. Mixing of air high in NOy, C$_3$H$_8$, and O$_3$ with clean air can produce the correlations discussed above. However, high values of

Figure 7. Scatterplots of C$_3$H$_8$ versus (top) NOx and versus (bottom) NOy. All the data above 7 km were used after excluding the data from highly polluted air masses and stratospheric air masses (see text).

Figure 8. Scatterplot of ozone versus NOy. Data used are the same as those in Figure 5. The dashed line represents a typical slope of regressions from the middle to northern latitude lower stratosphere [Murphy et al., 1993; Buhr et al., 1996].

Figure 9. Scatterplot of ozone versus C$_3$H$_8$. Data used are the same as those in Figure 5.
ozone, with simultaneously high NOx and C3H8, are still required to produce the observed correlations, indicating that photochemical production of ozone had occurred in sampled air masses.

[24] Ranges of the ozone and NOx values during BIBLE T were similar to those in the observed air masses in the free troposphere over Hawaii in April and May [Ridley et al., 1997]. While trajectories arriving over Hawaii frequently passed over the Asian continent, our results indicate that these air masses were not necessarily or solely affected by the surface sources of the Asian continent, but may also have been affected by other continents in the northern midlatitudes.

3.4. In Situ Production of Ozone in the Troposphere

[25] Although the correlations between ozone and its precursors demonstrate the effects of photochemical ozone production, the altitude region where the ozone production mainly occurred cannot be identified. As already discussed, the concentrations of trace species with lifetimes longer than 10 days were strongly influenced by convection in April. Ozone in the upper troposphere may not necessarily have been produced at the altitudes of sampling, but may have been transported from lower altitudes where ozone can be produced, depending on the levels of the ozone precursors and the length of time during which the air masses stayed there. The contribution of the in situ ozone production at the time of the observation was assessed by calculating the diurnal average of the net ozone production rates. The Atmospheric and Environmental Research, Inc. (AER) photochemical box model [Kotamarthi et al., 1997; Ko et al., 2000] was used for the calculations. The model calculates diurnal steady state concentrations of radicals and chemical species by constraining the observed values of ozone, NO, CO, H2O, NMHCs, and J(NO2) with the measurements along the flight tracks. It should be noted that the model results are from a local steady state model that does not account for the upwind or downwind photochemical conditions.

[26] Production (P) and loss (L) rates of ozone were calculated using the 10-s merged data for all altitudes but excluding take-off and landing, the polluted air masses from flight 5, and air masses influenced by stratospheric intrusions. The median values for each 1-km altitude step were integrated to derive column rates. A summary of the ozone budget for the 0–13-km column is given in Table 3. Figure 10 also shows median altitude profiles of the calculated diurnal averages of P, L, and the local net ozone production rates (P-L). The tropospheric (0–13 km) column-integrated rate of ozone production is $7.4 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$. The average stratospheric fluxes in NH and at midlatitudes in spring have been estimated to be $3.3-4 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$ and $10-20 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$, respectively, by global chemical transport models (GCTM) [Hauglustaine et al., 1998; Wang et al., 1998a, 1998b]. The integrated ozone production rate is 3–20 times larger than the stratospheric fluxes, suggesting that photochemical production is a dominant factor for the springtime increase of ozone. The integrated value of P during BIBLE T is comparable to that of 58 molecules cm$^{-2}$ s$^{-1}$ estimated for 20°–30°N during PEM-W-B [Crawford et al., 1997]. Net production of $8 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$ at 0–2 km is larger than that estimated for 20°–30°N during PEM-W-B by a factor of 2. In contrast, the result showed net destruction of ozone at 2–7 km ($7 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$) during BIBLE T, while net production of ozone prevailed at 2–8 km ($9 \times 10^{10}$ molecules cm$^{-2}$ s$^{-1}$) during PEM-W-B. However, it should also be noted that the above estimate is not statistically representative of the season because of the limited amount of data below 7 km (Table 3). Especially, the amount of data at 0–2 km is only 7% of the total.

**Table 3. Budget of Tropospheric Ozone During BIBLE T**

| Altitude, km | Number of model points | Column rates, $10^{10}$ molecules cm$^{-2}$ s$^{-1}$ |
|-------------|------------------------|---------------------------------------------------|
| 0–2         | 32                     | 29 ($12/23$)                                      |
| 2–7         | 114                    | 22 ($23/16$)                                      |
| 7–13        | 301                    | 23 ($3/21$)                                      |
| 0–13        | 447                    | 74 ($32/51$)                                      |
| Production (P) O3 | 25 ($9/5$)            | 25 ($9/5$)                                       |
| Loss (L) O3  | 30 ($10/20$)           | 6 ($1/2$)                                        |
| Net (P-L)   | 8 ($20/16$)            | 9 ($8/12$)                                       |

*Median diurnal average rates and central ±67% values calculated by the model. Because entries are medians, Net (P-L) and P(O3)-L(O3) are not identical.

Figure 10. Median altitude profiles of the calculated diurnal-average rates of ozone production (P), destruction (L), and the net ozone production (P-L) for each 1-km altitude step. The values are calculated from 1-min data at solar zenith angles less than 60°. Data points used for the calculation are the same as those in Figure 5. The bars indicate the central 67% values.
data. The large variability in $P$ at 0–2 km, combined with the large contribution (40%) to the column-integrated production rate, leads to reduced statistical reliability in the estimate of the column-integrated production rate.

The median values at 7–13 km show that the P-L values are positive, which correspond to the range of 0.5–4.4 ppbv d$^{-1}$ (0.7–5.1% d$^{-1}$). Figure 11 shows a scatterplot of the P-L values versus NO at 7–13 km. Critical NO values are defined as the levels of NO at which ozone formation and destruction are balanced [Crutzen, 1979]. The critical NO values ranged between 9–13 pptv at 7–13 km during BIBLE T. The median measured NO value at this altitude region was about 40 pptv, which was higher than the critical NO values by a factor of 3–4, indicating the importance of photochemical ozone production on the ozone budget in the upper troposphere in spring. The net production rate in the upper troposphere was also shown to be positive during PEM-W-A and B [Davis et al., 1996; Crawford et al., 1997]. The median net ozone production rate of 1.4 ppbv d$^{-1}$ for 26°–44°N at 7–13 km during BIBLE T is comparable to that of 1.7 ppbv d$^{-1}$ estimated for 20°–30°N at 8–12 km during PEM-W-B [Crawford et al., 1997]. Net ozone production has been shown to prevail in the upper troposphere north of Japan during April 1998 [Yienger et al., 1999]. The year-round net ozone production in the upper troposphere is compensated for by mixing with lower tropospheric air, where ozone is destroyed in summer and fall. An increase in the total column ozone occurs in winter and spring when net ozone production also occurs in the lower troposphere [Yienger et al., 1999]. Considering the longer photochemical lifetime of ozone in the upper troposphere (2–6 months) [Wang et al., 1998b] than that in the lower troposphere, continuous ozone production in the upper troposphere from winter to spring contributed to the higher ozone values during BIBLE T than those during PEM-W-B. However, net ozone production in the middle and lower troposphere in spring followed by vertical mixing also contributes to the springtime ozone amounts in the upper troposphere. More extensive observations both in the lower and upper troposphere in spring are needed to separate these two effects.

The P-L values in the highly polluted air masses discussed in 3.2 were also estimated to be 3.2–8.5 ppbv d$^{-1}$ (3.4–9.5% d$^{-1}$), which correspond to the high NO mixing ratios of 160–680 pptv (Table 2). The net diurnal-averaged production rate in these air masses was significantly larger than those in other air masses during BIBLE T and those at 20°–30°N during PEM-W-B, by a factor of 2–4. This result suggests that convective activity over east Asia has a large impact on the ozone levels over the whole Pacific region through eastward transport of ozone precursors.

4. Summary and Conclusions

Aircraft measurements of ozone, NO, NO$_2$, CO, and NMHCs were made mainly in the upper troposphere at 26°–44°N near Japan during April 1998. The observed ozone mixing ratios were about 80 ppbv at 7–13 km and 60 ppbv at 2–7 km. At all altitude levels, the ozone concentrations during BIBLE T were higher by 10–30 ppbv than those observed during PEM-West B at 20°–30°N. The average ozone values during BIBLE T were very similar to the climatological values obtained by ozone sonde measurements over Japan in April during 1993–1997.

The CO and C$_3$H$_8$ mixing ratios below 3 km during BIBLE T were lower than those during PEM-West B, as expected by their seasonal variations. The vertical gradients of these species above 5 km were small during BIBLE T because of active convection in April. In fact, highly polluted air masses were observed in the upper troposphere at 8–11 km over the Japan Sea on 24 April. They were transported from the boundary layer to the upper troposphere by cumulus convection associated with a cold front over northeast China during the 1–2 days prior to sampling.

In air masses not influenced by recent convection over east Asia, NO$_x$ and NO$_3$ were positively correlated with C$_3$H$_8$, suggesting significant contributions of anthropogenic activities on the levels of reactive nitrogen. However, other sources of reactive nitrogen, including the effect of NO production by lightning, need to be assessed to quantitatively determine the contribution of the surface sources. Ozone was well correlated with NO$_x$ and C$_3$H$_8$. The slope of the O$_3$–NO$_x$ correlation was 3 times smaller than typically observed in the lowermost stratosphere. The diurnal-average column-integrated gross production rate of tropospheric ozone, estimated by a photochemical box model, was larger than the NH average and local stratospheric flux at NH mid-latitudes in spring by a factor of 3–20. These results demonstrate the dominance of tropospheric photochemistry as a source of ozone over stratospheric input. Although the number of data is limited, the results imply that photochemical production is a dominant factor of the springtime increase of ozone.
The net ozone production rate in these air masses was estimated to be 1–4 ppbv d⁻¹ at 7–13 km. The ozone production rate is comparable to that calculated for 20°–30°N in the upper troposphere during PEM-West B. Continuous ozone production in the upper troposphere from winter to spring contributes to the enhancement of the ozone values in April, as observed during BIBLE T relative to those of PEM-West B. In addition, ozone produced in the lower troposphere, where net ozone production prevailed during PEM-West B, can also be transported to the upper troposphere by convection in April. Conversely, ozone produced in the upper troposphere can be transported downward by mixing associated with convection. The relative importance of in situ ozone production in the upper troposphere as compared with transport of ozone from below still needs to be assessed by further measurements.

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