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Large-scale air mass characteristics observed over western Pacific during summertime

Permalink
https://escholarship.org/uc/item/6xq1j7xb

Journal
Journal of Geophysical Research: Atmospheres, 101(D1)

ISSN
0148-0227

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Publication Date
1996-01-20

DOI
10.1029/95jd02200

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Large-scale air mass characteristics observed over Western Pacific during summertime


Abstract. Remote and in situ measurements of gases and aerosols were made with airborne instrumentation to investigate the sources and sinks of tropospheric gases and aerosols over the western Pacific during the NASA Global Tropospheric Experiment (GTE)/Pacific Exploratory Mission-West A (PEM-West A) conducted in September-October 1991. This paper discusses the general characteristics of the air masses encountered during this experiment using an airborne lidar system for measurements of the large-scale variations in ozone (O₃) and aerosol distributions across the troposphere and airborne in situ instrumentation for comprehensive measurements of air mass composition. In low latitudes of the western Pacific the airflow was generally from the east, and under these conditions the air was observed to have low aerosol loading and low ozone levels throughout the troposphere. Ozone was found to be below 10 parts per billion volume (ppbv) near the surface to 40-50 ppbv in the middle to upper troposphere. In the middle and high latitudes the airflow was mostly westerly, and the background O₃ was generally less than 55 ppbv. On 60% of the PEM-West A flights, O₃ was observed to exceed these levels in regions that were determined to be associated with stratospheric intrusions. In convective outflows from typhoons, near-surface air with low ozone (<25 ppbv) was transported into the upper troposphere (>10 km). Several cases of continental plumes from Asia were observed over the Pacific during westerly flow conditions. These plumes were found in the lower troposphere with ozone levels in the 60-80 ppbv range and enhanced aerosol scattering. At low latitudes over the central Pacific the troposphere primarily contained air with background or low ozone levels; however, stratospherically influenced air with enhanced ozone (40-60 ppbv) was observed several times in the lower troposphere. The frequency of observation of the air masses and their average chemical composition are also discussed in this paper.

Introduction

The primary objectives of the NASA Global Tropospheric Experiment (GTE)/Pacific Exploratory Mission-West A (PEM-West A) field experiments were to investigate the atmospheric chemistry of ozone (O₃) and its precursors over the western Pacific, to examine the natural budgets of these species, and to assess the atmospheric impact of anthropogenic emissions [Davis et al., this issue (a); Hoell et al., this issue]. The first PEM-West mission focused on the late summer/early fall period (PEM-West A) when the climatological flow in the lower troposphere of the western Pacific is expected to be predominantly from the east [Merrill et al., 1985]. Under these flow conditions the air would have been over the remote Pacific for a long period of time. A second PEM-West mission was planned for the late winter to early spring period (PEM-West B) when a strong outflow from the Asian continent into the western Pacific is expected [Savoie and Prospero, 1989]. This is the season when the natural input of desert aerosols and the anthropogenic influence of Asia on the troposphere over the western Pacific is the largest. These two PEM-West missions represent the extreme conditions for determining the composition of the air and the tropospheric chemistry over the western Pacific.

The first instrumented aircraft study of the troposphere over the Pacific was Project GAMETAG (Global Atmospheric Measurements Experiment on Tropospheric Aerosol and Gases) conducted in August-September 1977 and April-May 1978 [Davis, 1980]. The GAMETAG program was designed to test models for short-lived photochemical species and provide survey data on a number of species over a latitude range from 70øN to 58øS. The latitudinal and vertical distribution of O₃ was studied, and anticorrelations were found between O₃ and water vapor (H₂O), suggesting a dynamical effect [Routhier and Davis, 1980]. These data were used in a modeling study which concluded that...
Plate 1. Aerosol (top) and O₃ (bottom) distributions obtained from nadir airborne DIAL measurements on September 18, 1991, over the western Pacific (east of Japan) on PEM-West A (PWA) survey flight from Anchorage, Alaska, to Yokota, Japan (PWA flight 5). The relative amount of atmospheric backscattering and O₃ mixing ratios in parts per billion by volume (ppbv) are defined by the color scales at the top of each display. Black represents values greater than the maximum level given on the color scale. Geometric altitudes are given in kilometers above sea level (ASL), and universal time (UT) is shown at the top of each display. Aircraft latitude and longitude information in degrees is given at the bottom of each display at each reference time.

Plate 2. Aerosol and O₃ distributions associated with a stratospheric intrusion observed in the upper troposphere southeast of Japan on September 22, 1991 (PWA flight 6).
O$_3$ in the equatorial Pacific was destroyed by a reaction between H$_2$O and O(1D) and that odd nitrogen is deposited as nitric acid (HNO$_3$) and particulate nitrate (NO$_3$) [Liu et al., 1983]. One of the key findings from Project GAMETAG was the unusually large O$_3$ mixing ratios found in the troposphere between San Francisco and Hawaii, which led to further understanding of stratospheric injections of O$_3$ into the troposphere [Danielsen, 1980; Danielsen and Hipskind, 1980].

The NASA GTE/Chemical Instrumentation Test and Evaluation (CITE 2) aircraft mission was conducted in part over the northwestern Pacific in August 1986 [Hoell et al., 1990]. The CITE 2 mission focused on intercomparison of techniques for measuring nitrogen dioxide (NO$_2$), nitric acid (HNO$_3$), and peroxyacetyl nitrate (PAN). In addition to establishing the measurement limitations of the various techniques, important new data were acquired on these and other species, including nitric oxide (NO), NO$_2$, O$_3$, carbon monoxide (CO), ethane (C$_2$H$_6$), and CFC-11 (CFCl$_3$) [Singh et al., 1990] in clean air regions of the Pacific.

Measurements of the composition and chemistry in the lower troposphere over the Pacific were made as part of the NOAA Mauna Loa Observatory Photochemistry Experiment (MLOPEX) research program during May-June 1988. Downslope flow conditions at the Mauna Loa Observatory permit the sampling of air from the lower free troposphere, and measurements made during these periods have led to further understanding of the distribution and photochemical interactions of reactive species in the remote marine troposphere [e.g., Ridley and Robinson, 1992].

The composition and chemistry of the marine boundary layer over the Pacific was studied as part of the Third Soviet-American Gas and Aerosol Experiment (SAGA 3) conducted in February-March 1990 over the western tropical Pacific aboard the former Soviet ship Akademik Korolev [Johnson et al., 1993], and aerosols were studied off the coast of China on survey cruises from 1985 to 1987 by Zhou et al. [1992]. These ship measurements have provided extensive data sets on trace gases and aerosols in the marine boundary layer over the Pacific.

To study the budget of O$_3$ across the troposphere, we must be able to examine the sources and sinks of O$_3$ as they are related to different air masses that are observed over the western Pacific. Contributions from stratosphere-troposphere exchange processes, natural photochemical processes, and anthropogenic photochemical processes must be examined across the entire troposphere to determine the large-scale magnitude of their impact on the tropospheric O$_3$ budget.

This paper reports the results of large-scale studies of the distribution of aerosols and O$_3$ over the western Pacific using primarily data from an airborne lidar system. These results, together with airborne in situ measurements of O$_3$ and aerosols [Gregory et al., this issue], NO$_x$ and NO$_y$ [Smyth et al., this issue], CO and methane (CH$_4$) [Gregory et al., this issue], carbon dioxide (CO$_2$) [Anderson et al., this issue], nitrous oxide (N$_2$O) [Collins et al., this issue], sulfur dioxide (SO$_2$) and sulfate (SO$_4$) [Thornton et al., this issue], PAN [Singh et al., this issue], non-methane hydrocarbons (NMHC) [Blake et al., this issue], hydrogen peroxide (H$_2$O$_2$) [Heikes et al., this issue], and other species [Hoell et al., this issue], and with the meteorological analyses of atmospheric transport [Bachmeier et al., this issue; Merrill et al., this issue; Newell et al., this issue] provide insights into factors contributing to the summertime tropospheric distribution of O$_3$ and aerosols over the western Pacific.

### Experimental Techniques

An airborne differential absorption lidar (DIAL) system was used to provide vertical profiles of O$_3$ and aerosols from near the surface to above the tropopause along the flight track of the NASA Ames Research Center (ARC) DC-8 aircraft. Simultaneous zenith and nadir lidar measurements of O$_3$ and aerosols were made from a range of about 750 m above and below the aircraft to above the tropopause in the zenith case and to about 300 m above the surface in the nadir case. The DIAL O$_3$ measurements were made using an on-line wavelength at 288.2 nm and the off-line wavelength at 299.5 nm, and the aerosol backscatter measurements were made at a laser wavelength of 1064 nm. An O$_3$ measurement accuracy of better than 10% or 2 ppbv (parts per billion by volume), whichever is larger, with a vertical resolution of 300 m in the nadir and 600 m

### Table 1. PEM-West A* Flight Information

<table>
<thead>
<tr>
<th>Flight Number</th>
<th>Mission</th>
<th>Latitude Range</th>
<th>Longitude Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Sept. 16 survey (NASA Ames to Anchorage, Alaska)</td>
<td>37°N-61°N</td>
<td>154°W-122°W</td>
</tr>
<tr>
<td>5</td>
<td>Sept. 17 survey (Anchorage to Yokota, Japan)</td>
<td>35°N-61°N</td>
<td>140°E-150°E</td>
</tr>
<tr>
<td>6</td>
<td>Sept. 22 marine wall (southeast of Japan)</td>
<td>26°N-36°N</td>
<td>139°E-149°E</td>
</tr>
<tr>
<td>7</td>
<td>Sept. 24 continental wall (east of Japan)</td>
<td>35°N-42°N</td>
<td>139°E-149°E</td>
</tr>
<tr>
<td>8</td>
<td>Sept. 25-26 marine photochemistry (southeast of Japan)</td>
<td>27°N-36°N</td>
<td>131°E-140°E</td>
</tr>
<tr>
<td>9</td>
<td>Sept. 27 typhoon (south of Japan)</td>
<td>18°N-36°N</td>
<td>121°E-142°E</td>
</tr>
<tr>
<td>10</td>
<td>Oct. 1 survey (Yokota to Okinawa)</td>
<td>20°N-26°N</td>
<td>114°E-128°E</td>
</tr>
<tr>
<td>11</td>
<td>Oct. 2 survey (Okinawa to Hong Kong)</td>
<td>22°N-26°N</td>
<td>116°E-124°E</td>
</tr>
<tr>
<td>12</td>
<td>Oct. 4 continental outflow (east of Taiwan)</td>
<td>22°N-30°N</td>
<td>114°E-128°E</td>
</tr>
<tr>
<td>13</td>
<td>Oct. 6 Japanese flyby (south of Japan)</td>
<td>13°N-22°N</td>
<td>114°E-145°E</td>
</tr>
<tr>
<td>14</td>
<td>Oct. 8 survey (Hong Kong to Guam)</td>
<td>1°SN-14°N</td>
<td>145°E-162°E</td>
</tr>
<tr>
<td>15</td>
<td>Oct. 10-11 ozone trough (southeast of Guam)</td>
<td>1°SN-14°N</td>
<td>145°E-162°E</td>
</tr>
<tr>
<td>16</td>
<td>Oct. 13 high-pressure ridge (southwest of Guam)</td>
<td>4°N-14°N</td>
<td>125°E-145°E</td>
</tr>
<tr>
<td>18</td>
<td>Oct. 17-18 survey (Guam to Wake Island)</td>
<td>1°SN-19°N</td>
<td>145°E-167°E</td>
</tr>
<tr>
<td>19</td>
<td>Oct. 18-19 survey (Wake Island to Hawaii)</td>
<td>15°N-21°N</td>
<td>167°E-158°W</td>
</tr>
<tr>
<td>20</td>
<td>Oct. 20 Mauna Loa flyby (Hawaii local)</td>
<td>18°N-21°N</td>
<td>158°W-154°W</td>
</tr>
<tr>
<td>21</td>
<td>Oct. 21-22 survey (Hawaii to Ames)</td>
<td>21°N-38°N</td>
<td>158°W-122°W</td>
</tr>
</tbody>
</table>

*Pacific Exploratory Mission-West A*
in the zenith and a horizontal resolution in both cases of about 36 km (assuming a 3-min averaging time and an aircraft speed of 200 m/s) was obtained with a precision of better than 5% or 1 ppbv [Browell, 1983; Browell et al., 1983, 1985a, b]. Comparisons between airborne DIAL and in situ O3 measurements were made throughout PEM-West A to constantly verify that their agreement was consistent with the above stated limits. The aerosol backscatter measurements were derived from the range-corrected lidar signal at the 1064-nm laser wavelength, and the resolution of these measurements was defined by the vertical averaging interval of 60 m and the horizontal averaging interval of 1.75 s or about 350 m. The accuracy and precision of the aerosol measurements were better than 1%. Detailed characteristics of the current airborne DIAL system and the O3 DIAL technique are given by Browell [1989].

In addition to the airborne DIAL system the DC-8 had instrumentation for in situ measurements of O3, aerosol size distribution, aerosol number density, NOx, NOy, CO, CH4, N2O, SO2, SO4, HNO3, PAN, NMHCs, CFCs, H2O2, 7Be, other trace species, and meteorological parameters such as temperature, dew point, and winds. A general description of these systems and their measurements is given by Hoell et al. [this issue], and more details are provided in companion papers in this issue.

**Data Results and Discussion**

Eighteen missions were conducted as part of the PEM-West A field experiment between September 16 and October 21, 1991. A list of the PEM-West A missions, the mission objectives, and the latitude and longitude ranges of the missions are given in Table 1, and the flight tracks for these missions are shown in Figure 1. During this field experiment, eight flights were survey missions between NASA ARC, Anchorage, Alaska, Yokota, Japan, Okinawa, Hong Kong, Guam, Wake Island, Hawaii, and NASA ARC; and ten flights were “local” missions for intensive measurements over the Pacific from the bases in Japan, Hong Kong, Guam, and Hawaii.

Ozone and aerosol distributions were measured remotely with the DIAL system on all flights during PEM-West A. These measurements provided nearly complete altitude coverage of O3 and aerosol distributions from near the surface to the tropopause region along the aircraft flight track.

The air masses observed during PEM-West A were broadly divided into six main categories: background air in the free troposphere, near-surface (mixed layer) air, stratospherically influenced air, convective outflows, plumes, and clean Pacific air. Examples are presented below for the O3 and aerosol distributions that were observed during PEM-West A (PWA). Characteristics are discussed for the different types of air masses, and an analysis is given of the extent to which the air masses were observed in various regions of the Pacific in different altitude ranges. The average O3 profile and chemical composition of major air mass types are also presented.

**Ozone Enhancement from Stratospheric Intrusions and Continental Outflow**

The survey flight from Anchorage, Alaska, to Yokota, Japan, on September 18, 1991 (PWA flight 5), revealed the presence of extensive stratospheric intrusions that extended down into the lower troposphere (<5 km) at midlatitudes over the western Pacific. Plate 1 shows the aerosol and O3 distribution observed east of Japan on that flight. An air mass with relatively low aerosol scattering (<200) and high O3 mixing ratios (>60 ppbv)
can be seen in the center of Plate 1. This air mass is bounded by air with enhanced aerosol scattering (>400) that sometimes exceeds the scattering in the clean regions by an order of magnitude (>2000) and is optically thick in some locations (note attenuation of lidar signals in those regions). There are also enhanced O_3 levels (>50 ppbv) in association with the enhanced aerosol regions below ~6 km. The upper level (~6 km) aerosol layers are clouds, and they are associated with intermediate levels of O_3 (30-50 ppbv).

The in situ measurements on the DC-8 indicated that we were flying in the lower stratosphere at 11 km along this flight track. The highest values of O_3 (262 ppbv) were obtained at 01:11 UT, and at that time the dew point (DPT) was -66°C; CO was 52 ppbv; CH_4 was 1670 ppbv; N_2O was 300 ppbv; SO_2 was 226 pptv (parts per trillion by volume); SO_4 was 208 pptv; NO was 30 pptv; NO_3 was 593 pptv; ^7Be was 2653 fCi/m^3, and PAN was 116 pptv at 01:08 UT. The levels of SO_2 and SO_4 were elevated due to the injection of gases and aerosols into the lower stratosphere from the Mount Pinatubo eruption that occurred in the Philippines in June 1991. The zenith DIAL measurements showed that O_3 continued to increase above the aircraft, further confirming that we were in the lower stratosphere. The air mass below us with the elevated O_3 and the relatively low aerosol scattering (compared to the tropospheric aerosol loading) appeared to be an extension of the air mass we were flying in at 11 km. A cross section of potential vorticity (PV) derived from the European Center for Medium-Range Weather Forecasting (ECMWF) meteorological analysis for a portion of our flight track and within 2 hours of our time is shown in Figure 2. The PV analysis confirmed the intrusion of air with enhanced PV levels (>4x10^-7 deg m^2 (kg s)^{-1}) extending down to ~3 km near 40°N and the sampling of the aircraft in the lower stratosphere at ~250 hPa (PV~20). The vertical and horizontal extent of the intrusion was clearly evident in the airborne DIAL data south of 50°N, and there was good correlation between the O_3 enhanced regions and the PV analysis. Large-scale stratospheric intrusions have been observed previously with an airborne lidar at midlatitudes in the spring [Browell et al., 1987] and at high latitudes in the summer [Browell et al., 1992, 1994]; however, this was the first observation of this large-scale transport process at midlatitudes in the summer.

An air mass backtrajectory was calculated to determine the origin of the air mass with enhanced aerosols and enhanced O_3 below ~6 km. The wind analysis along the flight track on September 18, 1991, indicated that the air mass with enhanced aerosols below ~6 km (Plate 1) was coming off the Asian continent, and the enhanced O_3 was thought to be due to photochemical O_3 production from precursor gases released over Asia. Air mass backtrajectories were computed for four altitudes (1.5, 4, 10, and 14 km) where different air mass characteristics were observed at 01:50 UT on flight 5 (Figure 3). The air in the lowest layer, which has the most aerosols, came south from western Russia before turning southeast over China and Japan. The 4-km layer came east from the middle of Russia, and the higher layers came directly east over the Asian continent. These results confirm that the air with enhanced aerosols and O_3 had a continental source.

A second example of a stratospheric intrusion observed during PEM-West A southeast of Japan is shown in Plate 2. In this case, the intrusion was observed to extend down from a tropopause height of about 14.5 km on the northern side of the intrusion (left side of plate) to below 9 km in the upper troposphere. The intru...
indicated by the O₃=100 ppbv level) decreases in altitude from troposphere; however, the O₃ levels were as high or higher than in the outflow region below about 3.5 km. Low O₃ (<20 ppbv) was observed in part of the light aerosol scattering region from the 3.5- to the 5.5-km region, and above ~5.5 km to the tropopause at ~16.5 km. The O₃ ranged from 40 to >100 ppbv in what appeared to be relatively clean air (low aerosol loading). The backtrajectory analysis done for this case showed that the low-altitude air (~3 km) traveled over portions of the ocean and land areas before arriving at our measurement location. The low O₃ component of the air resulted from the marine portion of the trajectory, and the enhanced aerosols and O₃ came from inputs from either Taiwan and/or from the coast of China. The clean air with elevated O₃ came from over China with the air at ~6 km descending from ~9 km during the trajectory. While the low aerosol loading and enhanced O₃ levels of these air masses are consistent with the characteristics of stratospheric intrusions discussed above, the PV analysis did not show evidence for a stratospheric intrusion that would explain the magnitude of the O₃ enhancement that was observed. Clouds associated with deep convection (>12 km) over China were observed from the aircraft during this flight, and in situ measurements of air mass composition at ~9 km confirmed that the air was not of stratospheric origin. Large aerosols can be washed out of the air during the deep convective events [Gatz, 1977], while insoluble gases will pass into the outflow region in the middle to upper troposphere to photochemically produce O₃ if the precursor gases are present [Pickering et al., 1989, 1991].

This discussion has presented examples of the major processes that were observed to increase O₃ in the troposphere over the western Pacific during PEM-West A. These processes include stratospheric intrusions which mix elevated concentrations of O₃ with the background tropospheric air to produce what we call stratospherically influenced air, and continental outflows which can increase O₃ if the precursor gases are present in the air that is directly advected over the Pacific in aerosol-laden plumes or in outflows from convection over the continent.

**Low Ozone Air and Convective Transport over Pacific**

The first observation of an extensive region of air containing low O₃ (<30 ppbv) was on October 1, 1991, on a survey flight that was originally planned to go from Yokota, Japan, to Hong Kong, but because of a problem with the weather radar on the DC-8 and the presence of Typhoon Nat over the South China Sea, was diverted to Okinawa. Plate 5 presents the aerosol and O₃ distributions observed east of the northern tip of the Philippines on this flight. Except for the aerosols contained predominantly in the marine boundary layer below ~1.5 km and some cloud activity seen on the right of the cross section, the atmosphere was clean of aerosols below ~11.5 km. The region of low O₃ (<30 ppbv) extended from the marine boundary layer to just below ~1.5 km, and there was a broad transition region of mixing between the stratosphere and the troposphere over the western Pacific during PEM-West A. These processes include stratospheric intrusions which mix elevated concentrations of O₃ with the background tropospheric air to produce what we call stratospherically influenced air, and continental outflows which can increase O₃ if the precursor gases are present in the air that is directly advected over the Pacific in aerosol-laden plumes or in outflows from convection over the continent.

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Plate 3. Aerosol (left side) and O₃ (right side) distributions in the continental outflow observed east of Japan on September 24, 1991 (PWA flight 7).
below the tropopause at ~16 km. There were also regions where \( O_3 \) was less than 20 ppbv over most of the depth of the troposphere. An extensive region of cirrus with a depth of ~4 km was observed just below the tropopause, and the \( O_3 \) associated with it was typical of the \( O_3 \) values near the surface. Figure 6 shows the backtrajectories for three representative altitudes in Plate 5. While the trajectories at the different altitudes are from different directions due to the variations in wind patterns with altitude around Typhoon Nat, they have the common characteristic of spending the last 10 days over the water at low latitudes. While the 800 and 475 hPa air parcels were probably in the inflow region of the typhoon, the 200-hPa level was in the outflow region of the typhoon as indicated by the presence of extensive cirrus clouds in the upper troposphere. For more discussion of typhoon dynamics and its implications for atmospheric chemistry, see Newell et al. [this issue].

On the flight southeast of Guam into the "ozone trough" on October 11, 1991 (PWA flight 15), the lowest \( O_3 \) levels were observed. The nadir and zenith \( O_3 \) distributions obtained near the equator are shown in Plate 6. In this case, \( O_3 \) was less than 10 ppbv below 3 km, less than 25 ppbv up to 13 km, and less than 35 ppbv up to 15 km. A region of even lower \( O_3 \) was observed between 6° and 10°N where \( O_3 \) was less than 15 ppbv from near the surface to above 11 km. The backtrajectory analysis for representative altitudes at 0240 UT is shown in Figure 7. At all altitudes the flow was from the east, and the air parcels had been over the Pacific for at least 10 days. Photochemical destruction of \( O_3 \) over the tropical Pacific due to hydroxyl (OH) chemistry is thought to be the cause of these very low \( O_3 \) levels [see Davis et al., this issue (b)].

**Tropospheric Occurrence of Different Air Masses**

To assess the relative impact of various air mass types on determining the composition of the troposphere over the Pacific, the nadir and zenith DIAL \( O_3 \) and aerosol data were used to categorize the air mass types observed on all flights during PEM-West A. Prior to establishing the criteria for the different air mass types, a reference or background \( O_3 \) profile had to be defined. The reference \( O_3 \) profile shown in Figure 8 is an approximation to the average of a series of \( O_3 \) profiles that were selected to be representative of air that was reasonably unaffected by recent sources/sinks of \( O_3 \) and aerosols. While the reference \( O_3 \) profile may be close to the average western Pacific midlatitude background \( O_3 \) profile, it is really only a model for use as a discriminator in this analysis. A better estimate of the average background \( O_3 \) profile will come from the air mass analysis to be discussed below.

Nine air mass types were identified during this field experiment, and they were categorized using the following criteria in each altitude range: (1) background or reference air (BKG/REF) in the free troposphere, \( O_3 \) within 80-120% of reference \( O_3 \) profile and low aerosol scattering (within factor of 2 of estimated scattering from aerosol-free region at same altitude); (2) near-surface air (NS), enhanced aerosol scattering (more than factor of 2 of estimated scattering from nearby aerosol-free region) below about 2 km and \( O_3 \) less than 120% of reference \( O_3 \) profile; (3) plumes with low \( O_3 \) in the free troposphere (LPLU), enhanced aerosol scattering and \( O_3 \) less than 80% of reference \( O_3 \) profile; (4) plumes with background \( O_3 \) in the free troposphere (BPLU): enhanced aerosol scattering and \( O_3 \) within 80-120% of reference \( O_3 \) profile; (5) plumes with high \( O_3 \) in the free troposphere (HPLU), enhanced aerosol scattering and \( O_3 \) greater than 120% of reference \( O_3 \) profile; (6) clean Pacific air (CP), \( O_3 \) less than 80% of reference \( O_3 \) profile and low aerosol scattering; (7) stratospherically influenced air (SINF), \( O_3 \) more than 120% of reference \( O_3 \) profile and aerosol scattering comparable to levels above the tropopause; (8) convective outflows from convection over the ocean (CO), \( O_3 \) less than 80% of reference \( O_3 \) profile.

![Figure 4. Backtrajectory analysis for 10 days starting from 41.7°N/148°E at 0450 UT on September 24, 1991 (PWA flight 7), at 150 hPa (~14 km) (solid line), 275 hPa (~10 km) (long-dashed line), 550 hPa (~5 km) (medium-dashed line), and 800 hPa (~2 km) (short-dashed line).](image)

![Figure 5. Backtrajectory analysis for 10 days starting from 24.5°N/121°E at 0220 UT on October 6, 1991 (PWA flight 13), at 375 hPa (~8 km) (solid line), 475 hPa (~6 km) (long-dashed line), and 700 hPa (~3 km) (medium-dashed line).](image)
Plate 4. Aerosol and O₃ distributions associated with convective outflow from the continent observed near Taiwan on October 6, 1991 (PWA flight 13).
Yokota to Okinawa

PEM West-A Flight 10 1 Oct 91

Aerosol Data

Relative Aerosol Scattering (IR)

Ozone Data

Ozone (ppbv)

Plate 5. Aerosol and O$_3$ distributions associated with clean Pacific air and convective outflow observed near Taiwan on October 1, 1991 (PWA flight 10).
Figure 6. Backtrajectory analysis for 10 days starting from 19°N/124°E at 0740 UT on October 1, 1991 (PWA flight 10), at 200 hPa (~12 km) (solid line), 475 hPa (~6 km) (long-dashed line), and 800 hPa (~2 km) (medium-dashed line).

and in vicinity of cirrus clouds; (9) convective outflows from convection over the continent (CO-C), O_3 more than 120% of reference O_3 profile, low aerosol scattering, and no evidence of stratospheric intrusions in PV analysis.

The results of the air mass analysis are presented in Plate 7. At high latitudes (40°-60°N) over the Pacific the average tropopause was at an altitude of ~8 km (>70% of SINF were above this altitude). This is also confirmed by the average O_3 profile for the SINF for this region (Figure 9) where the average O_3 at 8 km exceeded 80 ppbv. A significant percentage of the troposphere above 6 km (~20%) consisted of SINF and the balance of the troposphere was mainly composed of plumes with enhanced O_3 from continental outflows. The average O_3 profile for these plumes was 17-20 ppbv (~45%) greater than the reference O_3 profile (Figure 9). Less than 25% of the troposphere consisted of plumes with background O_3 levels. In this latitude range the main flow was from the Asian continent over the Pacific throughout the troposphere.

At midlatitudes (20°-40°N) over the western Pacific the average tropopause was at ~16 km, and the background air was the predominant air mass type throughout the free troposphere, representing about 40% of the troposphere from 3 to 16 km. This provides confirmation that the O_3 reference profile used in the air mass classifications does represent the average air mass type in this region. In addition, the average O_3 profile for this air mass type was very close to the reference O_3 profile chosen for the air mass discrimination (Figure 10). The SINF and convective outflows (marine) also contributed greatly to the composition in the upper troposphere. In the midtroposphere, many different air mass types were observed, and in the lower troposphere, plumes from continental outflows and convective outflows from the continent were the dominant air masses. Figure 10 shows the average O_3 profiles for these air masses in the midlatitude region. The high O_3 plumes had up to 100% higher O_3 than that found in the background air, and in the upper troposphere the O_3 in the SINF was ~40% higher than the background average. The O_3 levels in the convective outflows (marine) were comparable to O_3 levels 5 to 10 km lower in the troposphere.

The results for the western Pacific low-latitude region show that the tropopause was typically above 16 km and that outflows from convection over the ocean were present a large percentage of the time in the upper troposphere. The low O_3 associated with the clean Pacific air and convective outflows dominated the air masses in the southwestern Pacific. Figure 11 shows the average O_3 profiles for the these two dominant air mass types. The O_3 in the clean Pacific air near the surface was less than 9 ppbv, which is more than 50% lower than the O_3 reference profile. In the middle to upper troposphere, the low O_3 air in outflows was typically ≤33 ppbv, which is about 50% lower than the reference profile. Only a low percentage of plume cases were observed in this region, and they were mostly below 5 km. Background air mass conditions were found in a low percentage of observations (5-15%) across the troposphere.

In the central Pacific at low latitudes the tropopause was typically between 15 and 16 km, and background air was the main air mass type observed in the middle to upper troposphere. In spite of the high percentage of observations of background air in this region, the average O_3 profile was strongly influenced by the convective outflow (marine) air below the tropopause (Figure 12), and in the lower troposphere the average O_3 profile was a combination of clean Pacific, background, and SINF air mass types. Clean Pacific air was found on average of about 50% of the time below 9 km, and the amount of background air decreased with decreasing altitude in this region. The average O_3 profile for the clean Pacific air in the central Pacific was within 20% (~4 ppbv) of the same air mass in the western Pacific (Figure 11). A few plumes were observed below 4 km with enhanced O_3 in them. These plumes were advected into this region from biomass burning that was occurring on Borneo. The observation of SINF
in the lower troposphere came as a surprise. This air was transported from stratospheric intrusions at midlatitudes over the western Pacific to the low latitudes of the central Pacific by advection and descent around the climatological high-pressure system typically situated in the southwestern Pacific [Bachmeier et al., this issue]. This is the reason why no SINF was observed in the upper troposphere in this region. The O3 increased by as much as 110% in the lower troposphere due to the transported air from the stratosphere compared to the background O3 levels.

Tables 2-4 present summaries of the regional and altitude dependence of several major air mass types observed during PEM-West A. The distribution of SINF is shown in Table 2. The latitude dependence of the tropopause levels and the frequency of occurrence of enhanced O3 due to stratospheric intrusions are clearly seen in this table. A significant tropospheric extent of SINF at midlatitudes over the Pacific was observed with generally increasing frequency with altitude and covering an average of more than 39% of the troposphere above 9 km. The amount of SINF in the upper troposphere (>6 km) at high latitudes and in the lower troposphere (<5 km) at low latitudes over the central Pacific is given in this table. Table 3 gives the total percentage of the troposphere that contains all plume types. The amount of plumes below 8 km increased with increasing latitude over the western Pacific. This was predominantly the result of the advection of boundary layer continental air over the Pacific as a result of the climatological high-pressure system that is present in the southwestern Pacific during the summer. Some plumes in the lower troposphere were also observed in the central Pacific as a result of long-range transport in the 2- to 4-km range over thousands of kilometers into the central Pacific. The low O3 air, as represented by the clean Pacific air and convective outflow (marine) air (Table 4), consisted of 55–95% of the entire troposphere from 2-15 km at low latitudes in the western Pacific to 35–66% in the same altitude range in the central Pacific. At midlatitudes the extent of low O3 air was reduced to 18–38% in the 4- to 14-km range, and there was negligible evidence of low O3 air in the high-latitude region.

The average percentage of the troposphere that contains the different air mass types in the different locations is given in Table 5. In the low latitudes of the western Pacific the low-O3 air (clean Pacific and convective outflow marine air) represented 68% of the troposphere, while the background conditions (free troposphere and near-surface air) was 14%. In the central Pacific the low-O3 air was 37% of the troposphere and the background was 46%. At midlatitudes the enhanced-O3 air (enhanced-O3 plumes and SINF) and the background air were both 36% of the troposphere. At high latitudes the enhanced-O3 air comprised most of the troposphere with 64% in plumes and 27% in SINF.

Contribution of Different Air Masses to Ozone Budget

The fractional contribution of the different air mass types to the O3 budget in the different regions is shown in Plate 8. The fractional contribution is calculated by weighting the average O3 for each air mass type by its percentage of occurrence as observed during PEM-West A. At high latitudes the elevated O3 in SINF dominated the O3 budget (>80%) above 8 km, and continental
plumes (BPLU and HPLU) dominated the O₃ budget over the rest of the troposphere. At midlatitudes, SINF contributed over 30% of the O₃ above 6 km, while continental plumes contributed over 40% below that altitude. The background air contributed about 35% of the O₃ over the entire free troposphere with minor contributions (less than ~20%) from the other air mass types. At low latitudes in the western Pacific the average O₃ is less than 30 ppbv up to 13 km (Figure 11), and the O₃ budget is dominated above 7 km (>85%) by the clean Pacific and convective outflow (marine) air mass types. Below 7 km, the clean Pacific air provided about 40% of the average O₃, and CO-C and high-O₃ plumes contributed over 30% of the O₃. In the central Pacific the background air accounted for more than 50% of the O₃ above 7 km, while below 6 km, SINF and high-O₃ plumes were a significant source of the O₃ (40-60%).

Chemical Characteristics of Different Air Masses

A detailed chemical characterization of the various air mass types encountered during PEM-West A was made using the comprehensive in situ measurement capabilities on the DC-8 [Hoell et al., this issue]. Missions and time periods were identified that corresponded to cases where in situ sampling occurred within each air mass type characterized by the DIAL system (Table 6). These measurement periods primarily corresponded to level flight legs that had either been overflown or underflown in the same geographic region during wall-type missions. Table 6 also shows the altitudes and geographic locations of the measurements. Only one case of low-O₃ plumes (LPLU) and two cases of background-O₃ plumes (BPLU) and near surface air (NS) could be clearly identified as having been in situ sampled. While the DIAL classification scheme could readily identify these air mass types, the limited in situ sampling of these air masses must be recognized in the interpretation of their chemical characteristics. Table 7 gives a summary of the combined chemical signatures of each of the air mass types based on the in situ sampled cases. The average chemical composition derived for each air mass type represents the first attempt to relate the large-scale classification capabilities of the airborne DIAL system with the chemical signatures of these air masses.

The in situ measurements of background air (BKG) measurements were grouped primarily in the midtroposphere (5.2-8.6 km), while the plumes (BPLU, LPLU, and HPLU) were sampled over most of the lower troposphere (1.4-8.6 km). The clear cases of in situ measurements of stratospherically influenced air (SINF) were found in the mid to upper troposphere (8.6-11.0 km) at middle to high latitudes over the western Pacific and in the low to middle troposphere (3.4-4.6 km) at low latitudes over the central Pacific. The convective outflow-continental (CO-C) air was measured in the midtroposphere (7.8-8.6 km), and the clean Pacific (CP) air was measured in the low to midtroposphere (3.7-8.6 km). The convective outflow (CO) marine air was sampled in the upper troposphere (8.3-11.3 km). While the altitudes of these in situ measurements were generally consistent with the regions where the air masses were most often observed in the low to mid troposphere (<10 km), they cannot necessarily be considered to be representative of the same air mass type observed remotely by the DIAL system in the upper troposphere (~10-16 km). This is particularly true for SINF which has a chemical composition that is determined by significantly diluting air from the lower stratosphere with highly variable tropospheric air.

In general, many compounds follow the trends that might be expected for the different air masses. The chemical composition was similar for continental plumes with high O₃ levels (HPLU) and for continental convective outflow air (CO-C). Both had elevated average levels of O₃ (57-64 ppbv), CO (105-108 ppbv), CH₄ (1764-1777 ppbv), NOy (645-679 pptv), SO₂ (124-127 pptv), PAN (99-161 pptv), NOx (96-119 pptv), C₂H₆ (775-1122 pptv), C₂H₄ (189-230 pptv), Pb (11 fCi/m³ STP). These compounds are generally elevated in air from continental surface sources that have some pollution, and the backtrajectories also indicate that the observed air was from a continental source. The C₂H₂/CO and C₂H₄/C₂H₆ ratios in these air mass types were also elevated (1.72-2.10 and 0.097-0.167, respectively) indicating that these air mass types have had a short processing time from their source over the continent. While the insoluble gas composition was similar in these air mass types, the levels of H₂O₂ (348 pptv) and CH₃OOH (146 pptv) were greatly reduced in the continental convective outflow (CO-C) air (444 and 141 pptv, respectively).
Table 2. Percentage of Troposphere With Stratospherically Influenced Air

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<tr>
<th>Altitude, km</th>
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<td>20°-40°N Latitude</td>
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<td>100 (stratosphere)</td>
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*110°-150°E region except for high-latitude case, which is 150°E-160°W.
†150°E-160°W region.

Table 3. Percentage of Troposphere with Plumes

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<td>20°-40°N Latitude</td>
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*110°-150°E region except for high-latitude case, which is 150°E-160°W.
†150°E-160°W region.
compared to the high-\(O_3\) plumes (1030 and 435 pptv, respectively) due to the wet processing of the CO-C air during vertical convective transport over the land. The aerosols were also backscattering in the lidar returns, and this was also reflected in washed out of the CO-C air, as was evident in the low aerosol levels (13 pptv) in these air masses.

The chemical composition of the background \(O_3\) plumes (BPLU) reflects the continental pollution source of the air with elevated CO (126 ppbv), \(SO_2\) (163 pptv), \(NO_2\) (510 pptv), \(C_2H_6\) (874 pptv), and \(H_2O_2\) (4453 pptv). Due to the longer processing time, as indicated by a \(C_2H_6/CO\) ratio of -1, and the lower average altitude of the plumes, the \(O_3\) in these plumes was within the range of our background \(O_3\) criterion. The low \(O_3\) observed in some of the plumes (LPLU) was also associated with even longer processing times (\(C_2H_2/CO\) -0.84; \(C_3H_6/C_2H_6\)-0.076) and possible photochemical loss of \(O_3\) ; although there is some indication from the lower CO, \(SO_2\), \(NO_2\), and hydrocarbon levels that there was less pollution in these plumes.

The stratospherically influenced air (SINF) had the lowest average levels of CO (71 ppbv), \(N_2O\) (305.9 ppbv), \(C_2H_6\) (481 pptv), \(C_3H_6\) (51.0 pptv), \(H_2O_2\) (313 pptv), and nitrate (6.8 pptv) and the longest processing times as indicated by the low ratios of \(C_2H_2/CO\) (0.64) and \(C_2H_6/C_2H_4\) (0.052). All of these factors reflect the influence of the stratospheric air on the composition of SINF. The average SINF composition includes three cases of directly sampled stratospheric air (Flights 4 and 5) and four cases where the stratospheric air that had been highly diluted with background tropospheric air (Flights 6, 19, and 20). Thus the variability of many species in the average SINF composition is not unique with respect to the other air mass types; however, gases that are low in the stratosphere were generally lower in SINF than in all but the clean Pacific (CP) and near-surface (NS) air. The unexpectedly high levels of \(SO_2\) (214 pptv) and sulfate (390 pptv) in the average SINF composition is attributed to the presence of gas and aerosols in the stratospheric air from the Mount Pinatubo eruption in June 1991.

The clean Pacific and marine convective outflow (CO) air have similar chemical compositions for the insoluble species. In the case of marine convective outflow the levels of \(H_2O_2\) and \(CH_3OOH\) are less than half that for CP. This results from the same wet scavenging process that reduces the soluble gases associated with the convective outflow of continental air (CO-C). In addition, sulfate is lower in the marine convective outflow from the washout of aerosols. Long processing times for the clean Pacific air is indicated by the low values of \(C_2H_2/CO\) (0.85) and \(C_2H_6/C_2H_4\) (0.058) and the low levels of \(O_3\) (27 ppbv), PAN (16.4 pptv), \(NO_2\) (140 pptv), \(NO_3\) (27.4 pptv), and hydrocarbons. Additional discussion of the chemical characteristics of these air mass types is given by Liu et al. [this issue] and Smyth et al. [this issue].

**Conclusions**

The troposphere over the Pacific during the late summer/early fall was found to be composed of a complex combination of different air masses. The distribution of \(O_3\) and aerosols in these air masses has been used to differentiate between different air mass types. Backtrajectory analyses were used to identify the probable source (continental or marine) of the tropospheric air, and inference of stratosphere-troposphere exchange was made through PV analyses. The tropospheric composition was found to be greatly dependent on the region of the Pacific and the altitude range of the air masses. Nine types of large-scale air masses were characterized during PEM-West A. The spatial distribution, frequency of observation, and composition of these air masses were discussed above.

Significant enhancements in \(O_3\) levels (typically >40% greater than reference \(O_3\) profile) were found at high latitudes (40°-60°N) over the Pacific during the summer/fall due to photochemical \(O_3\) production in continental plumes in the low to midtroposphere and to air that has been previously involved in stratospheric intrusions (SINF). Enhanced PV levels were associated with the SINF, and backtrajectory analysis confirmed the origin of the plumes over the Asian continent. The high-\(O_3\) continental plumes and SINF represented 64% and 27%, respectively, of the troposphere in this high-latitude region.
Plate 7. Percentage of time different air masses were observed over the Pacific in the indicated geographic and altitude regions. The air masses were categorized as either background air (REF/BKG), plumes with background O₃ (BPLU), plumes with low O₃ (LPLU), plumes with high O₃ (HPLU), near-surface (NS) air, convective outflows (CO) marine air, stratospherically influenced (SINF) air, clean Pacific (CP) air, or convective outflows continental (CO-C). The number of independent air mass classifications (N) that went into the analysis at each altitude is also given.
Plate 8. Fractional contribution (percentage) of different air mass types to the O₃ budget over the Pacific in the indicated geographic and altitude regions. The air mass types are the same as in Plate 7.
### Table 4. Percentage of Troposphere With Clean Pacific or Convective Outflow (Marine) Air

<table>
<thead>
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<th>Central Pacific†</th>
<th>Western Pacific*</th>
<th>Central Pacific†</th>
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* 110°-150°E region except for high-latitude case, which is 150°E-160°W.
† 150°E-160°W region.

### Table 5. Average Percentage of Troposphere for Different Air Masses

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<th>Central Pacific†</th>
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* 110°-150°E region except for high-latitude case, which is 150°E-160°W.
† 150°E-160°W region.
Table 6. Air Mass In Situ Measurements

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BKG—background or reference air; NS—near surface air; BPLU—plumes with background O₃ in the free troposphere; LPLU—plumes with low O₃ in the free troposphere; HPLU—plumes with high O₃ in the free troposphere; SINF—stratospherically influenced air; CO-C—convective outflow from convection over the continent; CP—clean Pacific air; CO—convective outflow from convection over the ocean.

*Average values.

At midlatitudes (20°-40°N) over the western Pacific the troposphere was dominated by background O₃ and aerosol conditions (31%); however, O₃ was enhanced due to stratospheric influences in 15-51% of the troposphere between 8 and 16 km. Plumes were frequently observed (over 37% of the time) in the lower troposphere below 5 km, and as in the high-latitude results, the majority of plumes had enhanced levels of photochemically produced O₃. These were due to advection of air from urban sources or biomass-burning regions. Low O₃ was found in association with cirrus clouds in up to 30% of the cases in 8- to 15-km region. This air was transported from near the surface into the upper troposphere as a result of convective storm activity over the Pacific.

At low latitudes (0°-20°N) in the western Pacific, low-O₃ air (up to 50% lower than reference O₃ profile) was found in 68% of the troposphere. The extensive vertical distribution of the low-O₃ air was attributed to convective storm activity associated with the many typhoons in this region of the Pacific during the late summer. Photochemical destruction of O₃ in the moist tropical marine atmosphere was responsible for the low O₃ ≤30 ppbv throughout most of the troposphere and the very low O₃ levels (≤10 ppbv) below 2 km. In the central Pacific (150°E-160°W) at low latitudes the average O₃ levels were not so low as in the western Pacific. Background O₃ was observed over 46% of the troposphere, and clean Pacific air and convective outflows were found 37% of the time. Observations of enhanced O₃ in the lower troposphere (below 6 km) were unexpected. The low aerosol loading and enhanced PV values associated with this air confirmed that the enhanced O₃ was due to previous stratospheric intrusions. The SINF had probably resulted from stratospheric intrusions that occurred at midlatitudes over eastern Asia, and this air subsequently descended into the lower troposphere of the central Pacific around the anticyclonic circulation in the western South Pacific. Plumes with enhanced aerosols and enhanced O₃ (40-60 ppbv) were also observed in the lower troposphere over the central Pacific. These plumes were just above the marine boundary layer, and they were determined to have originated from distant fires in Borneo based on backtrajectory analyses and reports of extensive burning on the island.

The fractional contribution of the various air masses to the average O₃ profiles observed in the different regions was generally similar to the fractional occurrence of the air mass types.
Table 7. Air Mass Composition (Average Value (s. d.) / Median Value (Number of Cases) / Range of Values)

<table>
<thead>
<tr>
<th>Air Mass Type</th>
<th>(Number of Cases)</th>
<th>BKG (3)</th>
<th>NS (2)</th>
<th>BPLU (2)</th>
<th>LPLU (1)</th>
<th>HPLU (4)</th>
<th>SINF (7)</th>
<th>CO-C (5)</th>
<th>CP (8)</th>
<th>CO (4)</th>
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<tr>
<td>Altitude, km</td>
<td>7.0/6.5</td>
<td>0.35</td>
<td>1.6</td>
<td>3.8</td>
<td>4.8/5.2</td>
<td>8.4/9.1</td>
<td>8.7/9.0</td>
<td>6.1/6.4</td>
<td>10.8/11.7</td>
<td></td>
</tr>
<tr>
<td>Dew Pt., °C</td>
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<td>21.6</td>
<td>7.4</td>
<td>-0.4</td>
<td>-21.3/-17.2</td>
<td>-48.5/-44.2</td>
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<td>O₃, ppb</td>
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<td>12.8</td>
<td>15.9</td>
<td>23.1</td>
<td>64.4/57.0</td>
<td>124.0/60.2</td>
<td>57.4/58.7</td>
<td>27.0/26.6</td>
<td>33.5/27.9</td>
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<tr>
<td>CO, ppbv</td>
<td>76.3/74.4</td>
<td>92.5</td>
<td>126.3</td>
<td>88.6</td>
<td>105.5/117.8</td>
<td>71.2/67.9</td>
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<td>CH₄, ppbv</td>
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<td>1713.8</td>
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<td>1777.5/1769.8</td>
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<td>1729.6/1723.0</td>
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<td>N₂O, ppbv</td>
<td>308.99 (1)</td>
<td>309.34 (1)</td>
<td>308.77 (1)</td>
<td>309.25</td>
<td>309.20/309.47</td>
<td>305.93/309.13 (4)</td>
<td>309.53/309.54</td>
<td>309.05/309.04</td>
<td>309.08/309.17</td>
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<tr>
<td>SO₂, pptv</td>
<td>61.9/66.0 (2)</td>
<td>55.4</td>
<td>163</td>
<td>42.0</td>
<td>124/139 (3)</td>
<td>214/226 (6)</td>
<td>127.2/128.3</td>
<td>59.0/51.4</td>
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<tr>
<td>Cₛ₂, pptv</td>
<td>6.87/8.14 (2)</td>
<td>3.55</td>
<td>0.90</td>
<td>2.80</td>
<td>4.17/3.14 (3)</td>
<td>2.98/2.58 (5)</td>
<td>3.07/1.64</td>
<td>0.68/0.75</td>
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<td>PAN, pptv</td>
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<td>2.0</td>
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<td>161/177</td>
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<td>34.4/19.0</td>
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<td>CₛCl₄, pptv</td>
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<td>3.58</td>
<td>1.35</td>
<td>2.80</td>
<td>6.77/6.80</td>
<td>4.12/4.13</td>
<td>3.34/3.02</td>
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<td>NO₃, pptv</td>
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<td>65.2/91.5 (4)</td>
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<td>539</td>
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<td>1.03</td>
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<td>0.64/0.69 (6)</td>
<td>1.72/1.80</td>
<td>0.85/0.92</td>
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<td>CH₄/O₂</td>
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<td>0.042</td>
<td>0.091</td>
<td>0.076</td>
<td>0.167/0.177</td>
<td>0.052/0.045 (6)</td>
<td>0.097/0.090</td>
<td>0.058/0.060</td>
<td>0.072/0.074</td>
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*Number of cases averaged together and used for median (three or more cases only), and if fewer cases are used, the number is indicated in the parenthesis; *Georgia Institute of Technology; *Nagoya University; *ppbv/ppbv; *pptv/pptv.
in that region; however, the higher O_3 levels in the SINF and high-O_3 plume air masses resulted in these air mass types having an enhanced contribution to the average O_3 profile at the expense of air mass types that have low O_3 levels, such as the clean Pacific and marine convective outflow air. At high latitudes, air from the lower stratosphere dominated the O_3 profile above 8 km, and high O_3 associated with high aerosols in continental outflow plumes dominated the O_3 profile in the low to midtroposphere.

At midlatitudes in the western Pacific, SINF contributed over 35% of the O_3 above 6 km, and this is significant since the tropopause is above 16 km in this region. Many different types of air masses contributed to the average O_3 profile below 6 km; however, the high-O_3 plume air made the most significant (40-65%) influence on the O_3 level. With the predominant easterly flow at the low latitudes the average O_3 profile was low (typically <30 ppbv), and the clean Pacific and convective outflow air mass types determined the O_3 profile above 7 km, and in the lower troposphere the contribution of the clean Pacific air decreased with altitude to less than 35% at 2 km. Below 5 km, plumes from various continental sources (enhanced aerosols and O_3) combine to influence the average O_3 profile. A very surprising result was found in the central Pacific at low latitudes. While the background and clean Pacific air mass types dominated the average O_3 profile above 6 km, the air associated with SINF was found to have contributed 27 to 40% of the average O_3 below 6 km. This shows the unexpected importance of this enhanced O_3 source to the O_3 budget in this remote region of the Pacific.

In general, the average chemical composition of all the air mass types followed trends that either reflected anthropogenic influences from continental sources, stratospheric influences or long processing times over the tropical marine environment. Convective transport processes over the land and the ocean tended to decrease soluble gases and washout aerosols; however, the insoluble species remained to reflect the near-surface source of the air. Thus the continental convective outflow air resembled the insoluble composition of the continental plume air, and the marine convective outflow air resembled the insoluble composition of the clean Pacific air. With the exception of the stratospherically influenced air the average O_3 levels in the different air mass types were generally inversely related to the amount of processing of the air as defined by the C_2H_4/C_2O ratio. The amount of air mass processing (i.e., photochemistry plus mixing) significantly influenced the composition of the observed air masses [Liu et al., this issue; Smyth et al., this issue]. As was done for O_3, the contribution of various air mass types to the tropospheric budget of other compounds can be determined from the chemical composition and fractional occurrence information presented in this paper for each air mass type.

Acknowledgments. The authors express their appreciation to Bill McCabe, Jerry Williams, Nealé Mayo, and Byron Meadows for their support in operating the airborne DIAL system in the field for the measurement of O_3 and aerosol distributions. We also thank Syd Ismail, Greg Nowicki, Shane Mayor, and Susan Kooi for their assistance in the reduction of the DIAL data and Yong Zhu (MIT) for the PV figure preparation. We appreciate the cooperation of the NASA Ames Research Center's DC-8 flight crew in conducting this mission and the assistance of the European Center for Medium-Range Weather Forecasting for providing the potential vorticity analysis. This research was supported by the NASA Global Tropospheric Chemistry Program.

References


Browell, E. V., M. A. Fenn, C. F. Butler, and S. M. Beck. Source to the 03 budget in this remote region of the Pacific.


Smyth, S., et al., Comparison of free tropospheric Western Pacific air mass classification schemes for the PEM-West A experiment, _J. Geophys. Res._, this issue.


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(Received August 5, 1994; revised July 8, 1995; accepted July 14, 1995.)