**Constraints on the sources of tropospheric ozone from 210Pb-7Be-O3 correlations**

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Constraints on the sources of tropospheric ozone from $^{210}$Pb-$^7$Be-O$_3$ correlations

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[1] The $^{210}$Pb-$^7$Be-O$_3$ relationships observed in three aircraft missions over the western Pacific (PEM-West A and B, TRACE-P) are simulated with a global three-dimensional chemical tracer model (GEOS-CHEM) driven by assimilated meteorological observations. Results are interpreted in terms of the constraints that they offer on sources of tropospheric ozone (O$_3$). Aircraft observations of fresh Asian outflow show strong $^{210}$Pb-O$_3$ correlations in September–October, but such correlations are only seen at low latitudes in February–March. Observations further downwind over the Pacific show stronger $^{210}$Pb-O$_3$ correlations in February–March than in September–October. The model reproduces these results and attributes the seasonal contrast to strong O$_3$ production and vertical mixing over east Asia in September–October, seasonal shift of convection from China in September–October to Southeast Asia in February–March, and slow but sustained net O$_3$ production in Asian outflow over the western Pacific in February–March. Seasonal biomass burning over Southeast Asia in February–March is responsible for the positive $^{210}$Pb-O$_3$ correlations observed at low latitudes. The model reproduces the observed absence of $^7$Be-O$_3$ correlations over the western Pacific during September–October, implying strong convective and weak stratospheric influence on O$_3$. Comparison of observed and simulated $^7$Be-O$_3$ correlations indicates that the stratosphere contributes less than 20–30% of O$_3$ in the middle troposphere at northern midlatitudes even during spring.

INDEX TERMS: 0368 Atmospheric Composition and Structure: Troposphere—constituent transport and chemistry; 0365 Atmospheric Composition and Structure: Troposphere—composition and chemistry; 0345 Atmospheric Composition and Structure: Pollution—urban and regional (0305); KEYWORDS: lead-210 and beryllium-7, Asian outflow of ozone, biomass burning, stratosphere-troposphere exchange


1. Introduction

[2] Understanding the factors controlling tropospheric ozone (O$_3$) is a central issue in tropospheric chemistry. Tropospheric O$_3$ is a greenhouse gas, a surface air pollutant, and a key player in controlling the oxidizing power of the atmosphere [Thompson, 1992]. Industrialization of the Asian continent is expected to be the principal driver of global change in atmospheric composition over the next decades [Ehhalt and Prather, 2001]. Better understanding of the Asian source of O$_3$ and its global impact is essential. We recently quantified the sources contributing to tropospheric O$_3$ over the Asian Pacific Rim in different seasons through a global three-dimensional (3-D) model analysis of Hong Kong and Japanese ozonesonde observations [Liu et al., 2002]. In this paper, we apply the same model to examine the constraints offered by aircraft observations of $^{210}$Pb-$^7$Be-O$_3$ correlations in Asian outflow over the western Pacific.

[3] Lead-210 (half-life 22 years) is the decay daughter of $^{222}$Rn (half-life 3.8 days) which is emitted from soils by decay of $^{226}$Ra [Turekian et al., 1977]. Beryllium-7 (half-life 53.3 days) is produced by cosmic ray spallation reactions with nitrogen and oxygen in the stratosphere and
upper troposphere. Both $^{210}\text{Pb}$ and $^7\text{Be}$ attach to aerosols following production, and are principally removed by precipitation scavenging. Lead-210 is a tracer of continental influence [Balkanski et al., 1993], while $^7\text{Be}$ is a tracer of stratospheric influence [Viezee and Singh, 1980; Dibb et al., 1992, 1994] and subsidence [Feely et al., 1989; Koch et al., 1996]. One may therefore expect $^{210}\text{Pb}$-$^7\text{Be}$-$\text{O}_3$ correlations to provide information on the origin of $\text{O}_3$. Strong correlations of $\text{O}_3$ with $^7\text{Be}$ observed at surface and mountain sites have been interpreted in a number of studies as evidence for a stratospheric or upper tropospheric source of $\text{O}_3$ [Johnson and Viezee, 1981; Reiter et al., 1983; Prospero et al., 1995; Moody et al., 1995; Tsutsumi et al., 1998]. A recent global 3-D model analysis suggests that there are confounding factors in this interpretation [Li et al., 2002a].

[4] Aircraft observations of $^{210}\text{Pb}$ and $^7\text{Be}$ up to 12-km altitude have been made by J. E. Dibb on a number of atmospheric chemistry field campaigns over the past decade. Detection limits for 10–30 min samples are 0.5 femtocurie per standard cubic meter ($\text{fCi SCM}^{-1}$) for $^{210}\text{Pb}$, and 10 fCi SCM$^{-1}$ for $^7\text{Be}$, sufficiently low to provide valuable data throughout the troposphere. We focus our attention on three campaigns (PEM-West A and B, TRACE-P) for which the data sets are particularly extensive and aimed at characterization of Asian outflow (Figure 1). The data reveal perplexing $^{210}\text{Pb}$-$^7\text{Be}$-$\text{O}_3$ correlations [Dibb et al., 1996, 1997, 2003] that provide as we will see important constraints on the sources of tropospheric $\text{O}_3$ and the role of Asian outflow.

[5] Our analysis will be based on the GEOS-CHEM global 3-D model of tropospheric chemistry [Bey et al., 2001a] driven by assimilated meteorological observations from the Goddard Earth Observing System data assimilation system (GEOS DAS) at the NASA Global Modeling and Assimilation Office (GMAO). The GEOS-CHEM model has been used extensively for simulations of tropospheric $\text{O}_3$, as reviewed in the next section, and has also been used for global simulation of $^{210}\text{Pb}$ and $^7\text{Be}$ [Liu et al., 2001]. It was recently applied to interpret observed $^{210}\text{Pb}$-$^7\text{Be}$-$\text{O}_3$ correlations at two North Atlantic surface sites [Li et al., 2002a].

2. Model Simulations

[6] Global simulations were conducted for the years 1991, 1994, and 2001 using GEOS-CHEM version 4.6 for 1991 and 1994, and version 4.26 for 2001 (see http://www-as.harvard.edu/chemistry/trop/geos). The GEOS data are from the GEOS1 assimilation for 1991 and 1994 ($2^\circ \times 2.5^\circ$ horizontal resolution, 20 sigma vertical levels), and GEOS3 for 2001 ($1^\circ \times 1^\circ$, 48 levels). The horizontal resolution is degraded here to $4^\circ \times 5^\circ$ for computational expediency. The model uses the advection scheme of Lin and Rood [1996] and the moist convective mixing scheme of Allen et al. [1996]. We assume complete vertical mixing within the GEOS-diagnosed mixed layer. Chemical tracer evaluations of convective transport in the GEOS fields have been presented by Allen et al. [1996], Liu et al. [2001], and Bell et al. [2002].

[7] Liu et al. [2001] presented a detailed description of the simulation capability for $^{210}\text{Pb}$ and $^7\text{Be}$ in GEOS-CHEM. Simulation of aerosol wet deposition includes scavenging in wet convective updrafts (with midlevel entrainment and detrainment), and first-order rainout and washout from both convective and large-scale precipitation. Return to the atmosphere following evaporation of precipitation is allowed. There is also a small dry deposition sink represented with a resistance-in-series model [Wesely and Hicks, 1977]. Cirrus precipitation not accounted for in the
assimilated meteorological data \cite{Lawrence and Crutzen, 1998} improves some aspects of the $^{210}$Pb-$^7$Be simulation and degrades others \cite{Liu et al., 2001}. It is not included in the present simulation.

The $^{210}$Pb source in the model assumes a uniform $^{222}$Rn emission of 1.0 atom cm$^{-2}$ s$^{-1}$ from land under nonfreezing conditions. Following Jacob and Prather \cite{1990}, we reduce the flux by a factor of 3 under freezing conditions. Zero emission is assumed from oceans and ice-covered surfaces. The $^7$Be source is taken from Lal and Peters \cite{1967} as a function of altitude and latitude. About two thirds of the $^7$Be is generated in the stratosphere and one third in the troposphere. The dependence of $^7$Be production on season or longitude is small and is neglected here.

Our initial simulation of $^7$Be given by Liu et al. \cite{2001} showed that the $^7$Be cross-tropopause flux in the GEOS1 meteorological fields is too fast by a factor of 3–4, consistent with the excessive $O_3$ flux from the stratosphere found by Bey et al. \cite{2001a}. We have since found the same problem in the GEOS3 fields. In the stratosphere, $^7$Be concentrations are determined by a local balance between production and radioactive decay. We correct the excessive $^7$Be flux from the stratosphere in our tropospheric simulations by scaling down globally the stratospheric $^7$Be source \cite{Liu et al., 2001}. Bey et al. \cite{2001a} found that although the magnitude of the cross-tropopause $O_3$ flux is too large, the latitudinal and seasonal variations of that flux are consistent with current knowledge \cite{Stohl et al., 2003}. Thus a uniform reduction in the $^7$Be cross-tropopause flux should not induce large errors in the latitudinal or seasonal distribution of this flux. The reader is referred to Liu et al. \cite{2001} for more discussion of this issue. For $O_3$ we circumvent the cross-tropopause flux problem by using the Synoz flux boundary condition \cite{McLinden et al., 2000}, which imposes a cross-tropopause $O_3$ flux of 475 Tg yr$^{-1}$.

After the correction for the cross-tropopause transport, we find that the model gives a good simulation of observed $^{210}$Pb and $^7$Be concentrations and deposition fluxes at surface sites worldwide with no systematic global bias, and that the observed latitudinal and seasonal distributions are successfully reproduced \cite{Liu et al., 2001}. PEM-West A and TRACE-P results were not presented by Liu et al. \cite{2001} and therefore we will present some discussion here.

The simulation of tropospheric $O_3$ in GEOS-CHEM uses a detailed representation of tropospheric $O_3$-NO$_x$-hydrocarbon chemistry \cite{Bey et al., 2001a}. The model solves the chemical evolution of 80 chemical species with 53 reactions and 67 species. The GEOS3 fields are presented by Bey et al. \cite{2001a}. The model reproduces the climatological monthly mean $O_3$ concentrations in the ozonesonde record to within usually 10 parts per billion by volume (ppbv), and captures the phase of the seasonal cycle to within 1–2 months, although it underestimates the seasonal amplitude at northern midlatitudes. More specific evaluations of model results with $O_3$ observations in different regions of the world have been conducted for the western Pacific \cite{Bey et al., 2001b}, the Asian Pacific Rim \cite{Liu et al., 2002}, the Middle East \cite{Li et al., 2001}, the United States \cite{Fiore et al., 2002a, 2002b, 2003a, 2003b}, the North Atlantic \cite{Li et al., 2002a, 2002b}, and the tropics \cite{Chandra et al., 2002, Martin et al., 2002}.

\cite{13} Our $O_3$ simulations presented here focus on the periods of the aircraft missions but include an 18-month initialization for each period. To investigate source regions of $O_3$ in the model, we decompose $O_3$ (actually odd oxygen or $O_x$) into tagged tracers where tagging indicates different source regions \cite{Wang et al., 1998; Fiore et al., 2002a; Li et al., 2002a, 2002b; Liu et al., 2002}. In the tagged $O_3$ tracer simulation, we use daily mean 3-D fields of $O_3$ production rates and loss frequencies archived from the standard full-chemistry simulation and transport five separate $O_3$ tracers originating from different regions, i.e., the stratosphere, the upper troposphere (400 hPa-tropopause), the middle troposphere (700–400 hPa), the continental lower troposphere (surface-700 hPa), and the marine lower troposphere. The decomposition is linear so that summing the concentrations of all five tracers reproduces the results from the standard simulation.

\cite{14} The radionuclide simulations presented here include a multi-year initialization starting with low concentrations of $^{222}$Rn, $^{210}$Pb and $^7$Be. This long initialization is necessary to bring $^{210}$Pb into steady state in the stratosphere.

3. Analysis of $^{210}$Pb-$^7$Be-$O_3$ Relationships

\cite{15} In this section we present a comparison of model results for $^{210}$Pb, $^7$Be, and $O_3$ with observations up to 12-km altitude from the three aircraft missions of Figure 1, and examine the constraints offered by $^{210}$Pb-$^7$Be-$O_3$ relationships on the sources of $O_3$. The $O_3$ data are time averages over the 10–30 min sampling interval for the radionuclides. Model output is sampled every 3 hours along the flight tracks. Concentrations of $^7$Be were below the detection limit (0.5 fCi SCM$^{-1}$) for 24%, 14%, and 33% of the samples from PEM-West A, PEM-West B, and TRACE-P, respectively. In those cases we assume a concentration of half the detection limit. Observations of $^{210}$Pb were never below the detection limit (0.5 fCi SCM$^{-1}$).

\cite{16} To describe the $^{210}$Pb-$^7$Be-$O_3$ relationships, we calculate the lines of best fit using the reduced-major-axis (RMA) method \cite{Hirsch and Gilroy, 1984}. Standard errors for the intercept and the slope are computed as described by Miller and Kahn \cite{1962}. We also tried a chi-square fitting (or weighted least-squares fitting) technique \cite{Press et al., 1992} and find that it yields results generally consistent with those obtained from the RMA method.

3.1. PEM-West A

\cite{17} This mission (September–October 1991) took place during the tail end of the summer monsoon season. We compare simulated and observed vertical distributions of $^{210}$Pb, $^7$Be, and $O_3$ concentrations over three regions (Figure 1) where intensive flights were flown. Following
Dibb et al. [1996], we call these regions “remote Pacific” (RP) and “near Asia” (NA1 and NA2). NA2 sampled northern midlatitudes outflow while NA1 and RP are subtropical or tropical. The comparisons are shown in Figure 2. The $^{210}$Pb observations near Asia show much higher values than those over the remote Pacific, as would be expected. The vertical gradients of $^{210}$Pb concentrations are generally weak and many samples in the free troposphere show higher concentrations than in the boundary layer, reflecting the role of convective transport during that season in lifting $^{222}$Rn to the free troposphere. Observed concentrations for $^{7}$Be are low relative to the typical 400–1000 fCi SCM$^{-1}$ range observed in other regions [Dibb et al., 1996], reflecting the general upwelling conditions and frequent rainfall of the summer monsoon as well as a seasonal minimum in transport from the stratosphere [Stohl et al., 2003]. Simulated concentrations of $^{210}$Pb and $^{7}$Be are consistent with these patterns.

Figure 2. Comparison between simulated (open circles) and observed (crosses) vertical distributions of $^{210}$Pb, $^{7}$Be, and O$_3$ concentrations during PEM-West A for the three regions in Figure 1: near Asia (NA1 and NA2) and remote Pacific (RP). Model results are 3-hour instantaneous concentrations sampled along the flight tracks for the specific flight days and times.

[18] The model reproduces the O$_3$ observations in PEM-West A with no systematic bias (Figure 2). Observed O$_3$ concentrations in the free troposphere near Asia are substantially (18 ppbv on average) higher than over the remote Pacific region, which may be largely explained by the lower latitude of the latter region where photochemical destruction exceeds production [Davis et al., 1996].

[19] Figure 3 shows the simulated and observed relationships of O$_3$ with $^{210}$Pb and $^{7}$Be during PEM-West A for three altitude bins (0–3, 3–8, 8–12 km) near Asia and over the remote Pacific. The observations show a positive covariation of $^{210}$Pb and O$_3$ at all altitudes near Asia [Dibb et al., 1996] with the slope $\Delta$O$_3$/\Delta$^{210}$Pb ranging from 2.8 ± 0.3 to 4.3 ± 0.9 ppbv/fCi SCM$^{-1}$. Strong O$_3$-$^{210}$Pb correlation is also observed over the remote Pacific in the upper but not in the lower troposphere. The model reproduces these observed $^{210}$Pb-O$_3$ relationships but tends to overestimate...
(underestimate) the observed correlations in the upper troposphere (middle troposphere). It reproduces well the observed \( \text{C}_1 \text{O}_3 / \text{C}_2 \text{Pb} \) values in the upper troposphere in both regions, implying a good simulation of deep convective outflow of \( \text{O}_3 \) from the Asian continent, as discussed below.

[20] The PEM-West A data show little correlation between the \( \text{Be} \) and \( \text{O}_3 \) observations anywhere (Figure 3b) and the model overestimates the correlations in the upper troposphere. Even in the model, we find from the tagged tracer analysis that the stratosphere contributes less than 20% to upper tropospheric \( \text{O}_3 \) over the PEM-West A domain. We previously found that stratospheric \( \text{O}_3 \) accounts for only about 6–8 ppbv \( \text{O}_3 \) in the upper troposphere over Hong Kong during this season [Liu et al., 2002].

[21] Dibb et al. [1996] hypothesized that frequent wet convection was the mechanism responsible for the \( \text{Pb}-\text{O}_3 \) relationships observed in the free troposphere during PEM-West A. This hypothesis agrees with what is seen in our model. In late summer-early fall, we find that convection is the principal process transporting surface pollution (\( \text{O}_3 \) and its precursors) to the middle and upper troposphere, and this is followed by transport eastward to the western Pacific [Liu et al., 2002]. During PEM-West A, the “near Asia” regions (Figure 1) are directly influenced by this convective outflow (Figure 4). Part of this outflow circulates around the western Pacific subtropical high, leading to positive \( \text{Pb}-\text{O}_3 \) correlations in the upper troposphere over the remote Pacific.

3.2. PEM-West B

[22] We compare simulated and observed vertical distributions of \( \text{Pb}, \text{Be}, \) and \( \text{O}_3 \) concentrations in Figure 5,
for similar regions (Figure 1) as PEM-West A. The $^{210}$Pb-$^{7}$Be-O$_3$ relationships are presented in Figure 6. The $^{210}$Pb and $^{7}$Be results were previously discussed by Liu et al. [2001]. While the PEM-West A $^{210}$Pb concentrations show relatively weak vertical gradients, the PEM-West B $^{210}$Pb observations indicate a decreasing trend with altitude near Asia, as the continental outflow is largely confined to the lower troposphere. The high $^{7}$Be concentrations in some of the samples indicate stronger seasonal stratospheric influence than during PEM-West A. The model captures this influence and the contrasting behavior between the two missions. The model overestimates the $^{7}$Be concentrations in the middle troposphere over the remote Pacific, but this appears to be due to insufficient aerosol scavenging rather than excessive stratospheric input [Liu et al., 2001].

[23] Bey et al. [2001b] previously examined the Asian outflow of O$_3$ to the western Pacific during PEM-West B using an earlier version of GEOS-CHEM. Near Asia the substantial enhancement of O$_3$ during PEM-West B relative to PEM-West A reflects the stronger continental outflow and intrusion of stratospheric air into the upper troposphere in spring. Over the remote Pacific, O$_3$ values in the free troposphere are about twice as much of those in fall due to efficient net production of O$_3$ [Crawford et al., 1997a]. The model reproduces these general patterns in the O$_3$ observations and shows no significant bias.

[24] O$_3$ and $^{210}$Pb concentrations only weakly correlate near Asia either in the model or in the observations (Figure 6a). The observed $^{210}$Pb-O$_3$ relationship in the upper troposphere near Asia shows two branches because of the latitudinal variation of O$_3$; $^{210}$Pb and O$_3$ concentrations appear positively correlated at low latitudes but negatively correlated at midlatitudes (not shown). By contrast, $^{210}$Pb and O$_3$ are correlated at all altitudes over the remote Pacific, with $\Delta$O$_3$/$\Delta$/$^{210}$Pb ranging from 2.3 ± 0.6 to 9.2 ± 0.7 ppbv/fCi SCM$^{-1}$ and increasing with altitude (Figure 6a). This reflects the slow but sustained rate of O$_3$ production in Asian outflow in that season [Crawford et al., 1997b], as shown here in Figure 7. Strong production of O$_3$ in Southeast Asian boundary layer is due to seasonal biomass burning (Figure 7c). The remote Pacific domain, situated at lower latitudes than the near Asia domain, sampled a greater influence of biomass burning outflow exported to the Pacific in convection and warm conveyor belts (WCBs) [Liu et al., 2003].

[25] The model shows strong O$_3$-$^{7}$Be correlations in the free troposphere, reflecting subsidence associated with the dry air stream of midlatitude cyclones [Cooper et al., 2002]...
and the western Pacific subtropical High, but these correlations are not apparent in the observations perhaps because of complications from aerosol scavenging (Figure 6b). Tagged tracer simulations indicate that about 20% of model O₃ at 500 hPa originates from the stratosphere.

3.3. TRACE-P

Most flights during TRACE-P were near Asia (Figure 1). We compare in Figure 8 the simulated and observed vertical distributions of ²¹⁰Pb, ⁷Be and O₃ concentrations over the TRACE-P period for two regions separated at 30°N (see Figure 1). The ²¹⁰Pb-⁷Be-O₃ relationships are presented in Figure 9. As during PEM-West B, the TRACE-P ²¹⁰Pb observations indicate a generally decreasing trend with altitude. The higher extreme values of ²¹⁰Pb in the boundary layer during TRACE-P are consistent with a stronger continental outflow than during PEM-West B [Liu et al., 2003]. The ⁷Be observations show larger variability, in particular in the middle and lower troposphere, than those for PEM-West B. The lack of vertical trend is partly due to the occurrence of high ⁷Be values in the boundary layer, which we cannot reproduce in the model. The latter do not correlate with other indicators of subsidence and instead are mostly associated with Asian outflow (200–450 ppbv CO, 20–70 fCi SCM⁻¹ ²¹⁰Pb); their source is unclear (J. E. Dibb, personal communication, 2003). They were not observed in PEM-West B (Figure 5).

[27] The observed O₃ concentrations generally do not correlate with ²¹⁰Pb concentrations north of 30°N (NA2 in Figure 9a). They are however correlated at all altitudes south of 30°N, with ΔO₃/Δ²¹⁰Pb increasing with altitude similarly to the remote Pacific data during PEM-West B (Figure 6a). Although TRACE-P was shifted ~3 weeks later in spring than PEM-West B and thus featured stronger photochemical production of O₃ [Davis et al., 2003], the overall patterns for O₃ production and transport are similar (compare Figure 7 and Figure 10). As discussed earlier for PEM-West B, the WCB and convective export of effluents from Southeast Asian biomass burning dominates O₃ production and generates the positive ²¹⁰Pb-O₃ correlation observed and simulated in the low-latitude middle and upper troposphere. Boundary layer outflow behind cold fronts is associated with O₃ production (positive ²¹⁰Pb-O₃ correlation) south of 30°N but not further north. Similar to our results for PEM-West B, the model shows strong O₃-⁷Be correlations that are not seen in the observations (Figure 9b) but are associated with only moderate stratospheric influence (30% of O₃ at 500 hPa north of 30°N).

4. Summary and Conclusions

[28] We have used a global 3-D model of tropospheric chemistry driven by assimilated meteorological data to analyze the observed ²¹⁰Pb-⁷Be-O₃ relationships during three aircraft campaigns over the western Pacific: PEM-West A (September–October 1991), PEM-West B (February–March 1994), and TRACE-P (February–April 2001). The objective of this study was to examine the constraints offered by ²¹⁰Pb-⁷Be-O₃ relationships on the sources of tropospheric O₃, with focus on Asian outflow to the Pacific.

[29] During PEM-West A, O₃ and ²¹⁰Pb concentrations are correlated at all altitudes near Asia, and in the upper troposphere only over the remote Pacific. In contrast, during PEM-West B and TRACE-P, the ²¹⁰Pb-O₃ correlations are strong in the low-latitude regions and over the remote Pacific, but are generally absent at midlatitudes near Asia. Our model analyses show that frequent wet convection in September–October (PEM-West A) and the shift of convection from China in late summer-early fall to Southeast Asia in spring (PEM-West B and TRACE-P) are responsible for this contrast. Boundary layer outflow behind cold fronts...
and sustained slow net $O_3$ production over the western Pacific during spring eventually leads to a positive $^{210}$Pb-$O_3$ correlation in air masses sampled in the boundary layer over the remote Pacific far downwind of the Asian continent. Seasonal biomass burning over Southeast Asia in spring is responsible for the positive $^{210}$Pb-$O_3$ correlations in the middle and upper troposphere at low latitudes.

[30] During PEM-West A, no $^7$Be-$O_3$ correlations are found either near Asia or over the remote Pacific owing to weak stratospheric influence. During PEM-West B and TRACE-P, the model shows much stronger $^7$Be-$O_3$ correlations than the observations, even though the stratospheric influence on $O_3$ in the model is rather modest (20–30% of total $O_3$ at northern midlatitudes at 500 hPa). This result implies that the model at least does not underestimate the stratospheric contribution to the observed springtime maximum in $O_3$ at northern midlatitudes. We have argued previously that this maximum is mostly the result of anthropogenic pollution [Wang et al., 1998; Li et al., 2002a].

[31] Overall, our results show that a global 3-D model interpretation of observed $^{210}$Pb-$^7$Be-$O_3$ correlations provides an important test of our understanding of the factors controlling tropospheric ozone. Our simulated $^{210}$Pb-$O_3$ and $^7$Be-$O_3$ correlations tend to be stronger than observed, presumably because of fewer factors of variability in the model. The stronger $^7$Be-$O_3$ correlations in the model could also be due to overestimate of subsidence. Fine-resolution models with more complete and detailed physical and chemical processes should produce larger variability and have potential to improve the simulation. Better knowledge of the $^{222}$Rn source distribution (assumed here to be

![Figure 5. Comparison between simulated (open circles) and observed (crosses) vertical distributions of $^{210}$Pb, $^7$Be, and $O_3$ concentrations during PEM-West B for the three regions in Figure 1: near Asia (NA1 and NA2) and the remote Pacific (RP). Model results are 3-hour instantaneous concentrations sampled along the flight track for the specific flight days and time. Stratospheric data above 8 km ($^7$Be > 1500 fCi SCM$^{-1}$ and $O_3$ > 150 ppbv) are not shown.](image-url)
Figure 6. Scatterplots of O\textsubscript{3} against (a) \textsuperscript{210}Pb and (b) \textsuperscript{7}Be during PEM-West B in three altitude bins (0–3, 3–8, 8–12 km) over near Asia (NA1 and NA2, left column) and over the remote Pacific (RP, right column), respectively. Model results (open circles and dashed lines) are compared with the observations (crosses and solid lines). Upper tropospheric samples with \textsuperscript{7}Be > 1500 fCi SCM\textsuperscript{-1} or O\textsubscript{3} > 100 ppbv are not used. The lines of best fit are calculated by the RMA method if the Pearson correlation coefficient R is greater than 0.3. See caption of Figure 3 for details.
(b) $^7$Be–O$_3$ relationships during PEM-West B

Figure 6. (continued)
PEM-West B, Feb-Mar 1994

(a) $^{222}$Rn concentrations (300 hPa)

(b) Horizontal fluxes and net production rate of $O_3$ (300 hPa)

(c) Horizontal fluxes and net production rate of $O_3$ (950 hPa)

Figure 7. (a) Simulated mean $^{222}$Rn concentrations (pCi SCM$^{-1}$) at 300 hPa for February–March 1994, indicative of frequent convection over Southeast Asia. (b) Simulated mean horizontal fluxes (arrows, moles cm$^{-2}$ s$^{-1}$) and net production rate (contours, ppbv day$^{-1}$) of $O_3$ at 300 hPa for February–March 1994. (c) Same as Figure 7b except for 950 hPa. Contour levels for $^{222}$Rn concentrations are 2, 4, 6, 8, 10. Contour levels for net production rate of $O_3$ are $-3$, $-1$, 0, 1, 3, 4, 8, 12, 16, 20.

TRACE-P, Feb-Apr 2001

Figure 8. Comparison between simulated (open circles) and observed (crosses) vertical distributions of $^{210}$Pb, $^7$Be and $O_3$ concentrations during TRACE-P for the two regions in Figure 1: NA1 and NA2. Model results are 3-hour instantaneous concentrations sampled along the flight track for the specific flight days and time. Stratospheric data ($^7$Be $> 1500$ fCi SCM$^{-1}$ and $O_3 > 150$ ppbv) are not shown.
Figure 9. Scatterplots of O$_3$ against (a) $^{210}$Pb and (b) $^{7}$Be during TRACE-P in three altitude bins (0–3, 3–8, 8–12 km) over near Asia (NA1, left column; and NA2, right column). See Figure 1 for the definitions of NA1 and NA2. Model results (open circles and dashed lines) are compared with the observations (crosses and solid lines). Samples with $^{7}$Be $>$ 1500 fCi SCM$^{-1}$ or O$_3$ $>$ 100 ppbv are not used, except one highly polluted air mass sample (106.6 ppbv O$_3$, 34 fCi/SCM $^{1210}$Pb, and below-detection-limit $^{7}$Be) in the 0–3 km altitude bin (NA1). The lines of best fit are calculated by the RMA method if the Pearson correlation coefficient R is greater than 0.3. See caption of Figure 3 for details.
(b) $^{7}$Be-O$_3$ relationships during TRACE-P

Figure 9. (continued)
uniform over land) would strengthen the constraints from the observed $^{210}$Pb–O$_3$ correlations.

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