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An estimate of the stratospheric contribution to springtime tropospheric ozone maxima using TOPSE measurements and beryllium-7 simulations

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[1] Measurements of tropospheric ozone (O_3) between $30^\circ N$ and $70^\circ N$ show springtime maxima at remote locations. The contribution of seasonal changes in stratosphere–troposphere exchange (STE) to these maxima was investigated using measurements from the Tropospheric Ozone Production about the Spring Equinox Experiment (TOPSE) campaign and the beryllium-7 (7Be) distribution from a calculation driven by fields from the Goddard Earth Observing System Data Assimilation System (GEOS DAS). Comparison with TOPSE measurements revealed that upper tropospheric model-calculated 7Be mixing ratios were reasonable (a change from previous calculations) but that lower tropospheric mixing ratios were too low most likely due to an overestimation of scavenging. Temporal fluctuations were well captured although their amplitudes were often underestimated. Analysis of O_3 measurements indicated that O_3 mixing ratios increased by 5–10% month⁻¹ for $\theta < 300$ K (the underworld) and by 10–15% month⁻¹ for $\theta > 300$ K (the tropospheric middleworld). 7Be mixing ratios decreased with time for $\theta < 290$ K and increased with time for $\theta > 300$ K. Model-calculated middleworld increases of 7Be were a factor of 2 less than measured increases. 7Be with a stratospheric source (strat- 7Be) increased by 4.6–8.8% month⁻¹ along TOPSE flight paths within the tropospheric middleworld. Increases in strat- 7Be were not seen along TOPSE flight paths in the underworld. Assuming changes in tropospheric O_3 with a stratospheric source are the same as changes in strat- 7Be and that 50% of O_3 in the region of interest is produced in the stratosphere, changes in STE explain 20–60% of O_3 increases in the tropospheric middleworld and less than 33% of O_3 increases in the underworld. *INDEX TERMS:* 0368 Atmospheric Composition and Structure: Troposphere—constituent transport and chemistry; 3362 Meteorology and Atmospheric Dynamics: Stratosphere/troposphere interactions; 3337 Meteorology and Atmospheric Dynamics: Numerical modeling and data assimilation

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1. Introduction

[2] Measurements of tropospheric ozone (O_3) between $30^\circ N$ and $70^\circ N$ show broad summer time maxima at sites affected by pollution and springtime maxima at more remote locations [Oltmans, 1981; Logan, 1985]. The summertime maxima are attributed to rapid photochemical production of O_3 during a period when ultraviolet radiation is at its peak. The springtime maxima are usually attributed to seasonal changes in the cross-tropopause transport of O_3

[Danielsen and Mohnen, 1977] and/or in the photochemical processing and abundance of O_3 precursors [Penkett and Brice, 1986]. The relative importance of each of these factors is still unclear. Monks [2000] summarizes observations of springtime maxima and hypotheses for their cause(s) in a recent review article.

[3] An increased understanding of the processes contributing to the springtime maxima was a major goal of the Tropospheric Ozone Production about the Spring Equinox Experiment (TOPSE) (E. Atlas et al., The TOPSE experiment: Introduction, submitted to *Journal of Geophysical Research*, 2002). The TOPSE field experiment consisted of seven deployments and 38 science flights (TOPSE flights 5–42, flights 1–4 were test flights) between 4 February and 23 May 2000. Each deployment began and ended at National Center for Atmospheric Research's (NCAR's) Research Aviation Facility in Broomfield, CO ($39^\circ 55' N$, $105^\circ 07' W$). Each deployment included flights to Winnipeg ($49^\circ 53' N$, $97^\circ 09' W$) and Churchill, Manitoba ($58^\circ 47' N$,

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94°12'W). Deployments 2 and 4–7 also included flights to Thule, Greenland (76°32'N, 68°45'W). Trace species and atmospheric parameters were measured on each of the NCAR C-130 flights at altitudes of up to 8 km.

[4] In this paper, seasonal changes in the cross-tropopause transport of O₃ are investigated by analyzing TOPSE measurements, Ertel's potential vorticity (PV), and output from a beryllium-7 (⁷Be) simulation. The focus of this study is the TOPSE region (defined here to be 100°–60°W, 40°–80°N) and period (defined here to be 1 February to 31 May 2000). Past studies of seasonal variations in cross-tropopause exchange are discussed in section 2. The data and model output used in this study are described in section 3. Model-calculated and measured ⁷Be mixing ratios are compared in section 4. Seasonal trends in ⁷Be and their implications for O₃ are discussed in section 5. Results are summarized in section 6.

2. Seasonal Variations in Cross-Tropopause Exchange

[5] The cross-tropopause net upward flux (F) of O₃ during an exchange event can be calculated by subtracting the downward flux of O₃ during the event from the upward flux of O₃ during the event.

$$F_Q = M_u Q_1 - M_d Q_2, \quad (1)$$

where M_u and M_d are the upward and downward components of the mass flux during the event and Q_1 and Q_2 are the O₃ mixing ratios in the upper troposphere and lower stratosphere, respectively. The net effect of several stratosphere–troposphere exchange (STE) events on O₃ is an increase in tropospheric O₃ because stratospheric O₃ mixing ratios exceed tropospheric mixing ratios by roughly an order of magnitude. Equation (1) can be rewritten as [e.g., *Gettelman and Sobel, 2000*]:

$$F_Q = M Q_2 + M_u (Q_1 - Q_2), \quad (2)$$

where the flux is now comprised of one part equal to the net mass flux (M) multiplied by the O₃ mixing ratio in the lower stratosphere and another part that is equal to the upward mass flux multiplied by the difference between upper tropospheric and lower stratospheric O₃ mixing ratios. Assuming M_u is zero, a defensible assumption in the middle stratosphere, the cross-tropopause flux of O₃ can be computed as a residual of the O₃ budget in the region between the tropopause and the middle stratosphere [e.g., *Gettelman and Sobel, 2000*].

[6] The net mass flux across the tropopause is determined by the frequency of STE events and the strength of the wave-driven and turbulence-driven Brewer–Dobson circulation [Brewer, 1949; Dobson, 1956]. This diabatic circulation consists of upward motion in the tropics and descending motion at higher latitudes (Figure 1). Its connection with STE can be understood by dividing the atmosphere into three parts, the underworld, middleworld, and overworld [Hoskins, 1991; Holton et al., 1995]. The underworld and overworld are the regions of the atmosphere nearest and farthest from the Earth, respectively. Isentropes within these regions do not cross the tropopause.

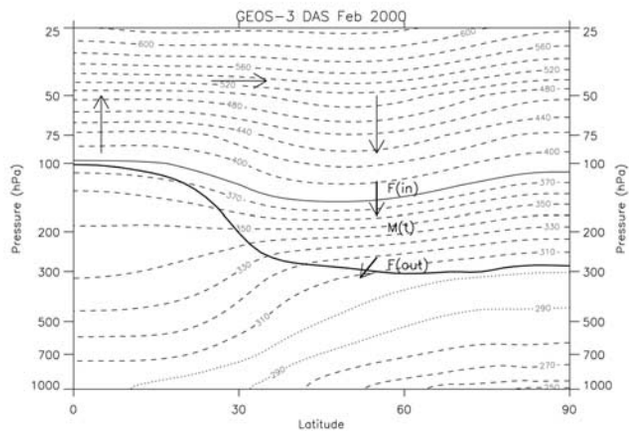


Figure 1. Zonally averaged mean (February 2000) θ (thin lines) and tropopause pressure (thick solid line) from the GEOS-3 DAS. Contour interval is 10 (20) K for $\theta < (>)$ 380 K. The 380 K isentrope is shown with a thin solid line. The 290 and 300 K isentropes are shown with dotted lines. $M(t)$ denotes the mass of the lowermost stratosphere (upper middleworld), the region between the tropopause and the 380 K isentrope. $F(\text{in})$ denotes the flux into the upper middleworld from the overworld. $F(\text{out})$ denotes the flux from the upper to lower middleworlds (i.e., the net stratosphere-to-troposphere exchange). Notation and concept from the study of *Appenzeller et al.* [1996].

In the absence of diabatic processes, air cannot move between the underworld and overworld. The remaining portion of the atmosphere is called the middleworld. Isentropes in this region, which has an upper bound at ~ 380 K, intersect the tropopause. Therefore, it includes portions of the upper troposphere and extratropical lower stratosphere. Seasonal variations in the strength of the Brewer–Dobson circulation lead to seasonal variations in the downward transport of mass between the stratospheric overworld and middleworld [Holton, 1990]. This downward transport is important because mass in the stratospheric middleworld can be transported isentropically into the troposphere.

2.1. Estimates of STE From Meteorological Data

[7] *Appenzeller et al.* [1996] studied seasonal variations in transport between the stratospheric overworld and middleworld and in STE using United Kingdom Meteorological Office (UKMO) stratospheric analyses. They found that transport between the stratospheric overworld and middleworld has a maximum in early winter and a minimum in late spring or early summer. *Gettelman and Sobel* [2000] found a similar seasonal cycle using fields from version 1 of the Goddard Earth Observation System Data Assimilation System (GEOS-1 DAS). By continuity, the net flux (F_{out}) across the tropopause can be calculated by subtracting the change in the mass of the stratospheric middleworld with time (dM/dt) from the net flux into the stratospheric middleworld from above ($F_{\text{in}}(t)$) where $F_{\text{in}}(t)$ and $F_{\text{out}}(t)$ are defined to be positive for downward directed mass fluxes (see Figure 1) [Appenzeller et al., 1996].

$$F_{\text{out}}(t) = F_{\text{in}}(t) - dM/dt \quad (3)$$

The seasonal cycle of STE is not completely determined by the seasonal cycle in the residual circulation because seasonal changes in the position of the tropopause lead to changes in the mass of the stratospheric middleworld. The tropopause moves upward and poleward during the transition from winter to summer and downward and equatorward during the transition from summer to winter [Staley, 1982]. The result is a net transport of mass from the stratospheric to the tropospheric middleworld during the spring and a net transport of mass from the tropospheric to the stratospheric middleworld during the fall. Appenzeller *et al.* [1996] found that the net downward transport across the tropopause is large and increasing slowly (by 10–20%) between December and June. A broad peak exists because springtime decreases in inflow from the overworld are offset by springtime decreases in the mass of the stratospheric middleworld. Gettelman and Sobel [2000] found an earlier (late winter) and more defined peak in downward transport across the tropopause when they calculated STE using the Wei method [Wei, 1987].

[8] Cross-tropopause transport is of the most interest when it results in the irreversible transport of material between stratosphere and troposphere. Irreversible STE in the extratropics occurs primarily by small-scale mixing within tropopause folds [Danielsen, 1968], cutoff lows [Bamber *et al.*, 1984], and planetary-scale anticyclones [Chen, 1995]. Seasonal cycles in the frequency and/or intensity of any of these features can lead to seasonal changes in STE. Folding events are more than twice as common as cutoff lows [Ebel *et al.*, 1996]. They also show clear seasonal cycle with a minimum in the summer and a broad peak between midautumn and midspring [Ebel *et al.*, 1996; Elbern *et al.*, 1998]. Cutoff low events show a slight preference for the summer [Price and Vaughan, 1992].

[9] Wernli and Bourqui [2002] quantified seasonal and latitudinal variations in extratropical STE by examining European Center for Medium Range Weather Forecasting (ECMWF)-based air trajectories that cross the 2.0×10^{-6} K m² kg⁻¹ s⁻¹ PV (2.0 PVU) surface, the “traditional” dynamical tropopause [Holton *et al.*, 1995]. They found that net exchange was downward throughout the year in the midlatitudes. The amplitude of the downward exchange was largest between January and April and smallest between August and October. At higher latitudes (65°–90°N), the net exchange was slightly upward and did not vary significantly with season. In the vertical, STE was largest in a 150 hPa layer below the climatological tropopause. This layer corresponds to approximately 285–320 K in the winter and 310–340 K in the summer. Seasonal variations in tropopause height are at least partially responsible for differing seasonal cycles between the lower and upper portions of the middleworld. Analyses of ECMWF fields show vigorous exchange throughout the year in the lower middleworld ($\theta < 330$ K) [Chen, 1995] and a summertime peak in exchange in the upper middleworld [Chen, 1995; Postel and Hitchman, 1999].

2.2. Estimates Derived From Model-Calculated and Measured Radionuclide Distributions

[10] Seasonal changes in the cross-tropopause flux of O₃ can also be estimated through the analysis of radionuclide distributions. Radionuclides that can act as stratospheric

tracers include strontium-90 (⁹⁰Sr) and the isotopes ⁷Be and ¹⁰Be [Husain *et al.*, 1977; Dutkiewicz and Husain, 1985]. Typical stratospheric mixing ratios of these species exceed tropospheric mixing ratios by over an order of magnitude. ⁹⁰Sr is a species with a 39.9-year lifetime (mean life with respect to radioactive decay) that is produced by nuclear detonations. ⁷Be is a species with a 76 day lifetime that is produced by cosmic radiation, while ¹⁰Be is a species with a 3.6 million year lifetime that is produced by cosmic radiation. Danielsen and Mohnen [1977] studied seasonal variations in STE using seasonal variations in the ⁹⁰Sr deposition rate and observed relationships between ⁹⁰Sr, PV, and O₃. Their analysis indicated that the net transport of mass between stratosphere and troposphere peaked during mid-May. Fitting a cosine to the seasonal variation, they estimated that exchange rates of mass (O₃) during mid-May exceeded exchange rates during mid-November by a factor of 3.0 (4.8) while exchange rates of mass (O₃) during mid-May exceeded exchange rates during mid-January by a factor of 2.0 (2.8).

[11] The interpretation of seasonal peaks in radionuclide data is complicated because a balance between four processes (STE, vertical and horizontal mixing within the troposphere, and scavenging) controls seasonal variations in tropospheric radionuclide mixing ratios [Feely *et al.*, 1989]. The relative importance of each of these processes depends on location and season. For example, analysis of ⁷Be, ¹⁰Be, and ²¹⁰Pb (a species with a 32.3-year lifetime that is produced predominantly in the troposphere by the decay of Radon-222) aerosol samples taken at Alert, Northwest Territories, Canada indicates that seasonal variations in vertical mixing within the troposphere are the main factor responsible for springtime peaks in ⁷Be and ¹⁰Be at that location [Dibb *et al.*, 1994].

[12] Chemistry and transport model (CTM) calculations aid in the interpretation of radionuclide data. Rehfeld and Heimann [1995] studied seasonal changes in the intensity of the meridional circulation and STE by comparing model-calculated and observed mixing ratios of ⁹⁰Sr, ⁷Be, and ¹⁰Be. They found that seasonal changes in scavenging and STE contribute to seasonal changes in ⁷Be. However, seasonal changes in the ¹⁰Be/⁷Be ratio were dominated by seasonal changes in STE. The ¹⁰Be/⁷Be ratio in the upper troposphere peaked during February and March indicating that STE peaked during this period. The ratio in the lower troposphere peaked during March and April indicating that it takes about a month to mix air from the upper to lower troposphere. Synoptic conditions following STE are often conducive to large-scale subsidence that transports tracers from the upper to the lower troposphere [Moody *et al.*, 1996; Cooper *et al.*, 1998].

[13] Koch and Rind [1998] evaluated the intensity of the meridional circulation in the lower stratosphere using output from simulations of ⁷Be and ¹⁰Be with the Goddard Institute for Space Studies (GISS) GCM. They found that the net transport is poleward and vigorous during the winter and spring and equatorward and weak during the summer. Koch *et al.* [1996] evaluated processes affecting the vertical mixing of tracers in the troposphere using output from ⁷Be and ²¹⁰Pb simulations. They found that seasonal changes in middle and upper tropospheric mixing are affected by seasonal changes in the strength of the meri-

dional circulation. However, transport to the surface is more closely related to seasonal changes in convective mixing.

2.3. Estimates Derived From Model-Calculated and Measured O₃ Distributions

[14] Seasonal changes in the cross-tropopause flux of O₃ have also been estimated through the analysis of meteorological parameters, O₃ measurements and/or model output. M. A. Olsen et al. (Estimating downward cross-tropopause ozone flux using column ozone and potential vorticity, submitted to *Journal of Geophysical Research*, 2002) estimated the net downward flux of O₃ across the Northern Hemisphere midlatitude tropopause using the relationship between total column O₃ and PV. They found that exchange rates in mid-April exceeded exchange rates in mid-September by a factor of 2. Exchange rates increased by ~25% between mid-January and mid-April. They attributed the seasonal cycle to seasonal changes in the frequency and strength of baroclinic systems.

[15] *Roelofs and Lelieveld* [1997] studied seasonal changes in Northern Hemisphere STE of O₃ using a GCM, the European Center Hamburg Model, version 4 (ECHAM4), coupled to a tropospheric chemistry model. They found that transport of stratospheric O₃ into the troposphere increased by nearly 20% between winter and spring. However, the winter to spring increase due to transport was less than the winter to spring increase in photochemically produced O₃. *Wang et al.* [1998] studied the causes of springtime O₃ maxima using a CTM driven by fields from the GISS GCM. They attributed springtime maxima to the “superimposed” contributions of O₃ transported from the stratosphere that peaked in the early spring, and O₃ produced in the troposphere that peaked in the late spring.

2.4. Summary

[16] Seasonal variations in the cross-tropopause flux of O₃ have been studied by a variety of independent methods. With few exceptions, a springtime peak is seen in STE. The contribution of this springtime peak in STE to the springtime O₃ maxima is the focus of this study.

3. Data and Model Description

[17] The contribution of STE to the springtime O₃ maxima will be investigated using meteorological parameters derived from the GEOS-3 DAS [*Schubert et al.*, 1993; *Bloom et al.*, 1996], measurements from the TOPSE campaign, and the ⁷Be distribution from a CTM calculation. These tools are discussed in the following sections.

3.1. University of Maryland (UMD) CTM/GEOS-3 DAS

[18] The distribution of ⁷Be was simulated using the UMD-CTM, a version of the Goddard CTM [*Allen et al.*, 1996a, 1996b] that includes wet scavenging and dry deposition. The calculations were driven by data from the GEOS-3 DAS. The mixing ratio change due to advection is calculated using a multidimensional and semi-Lagrangian extension of the piecewise parabolic method (PPM) [*Lin and Rood*, 1996]. Convective mixing in the UMD-CTM [*Allen et al.*, 1996b] is calculated using cumulus mass flux and detrainment output from the Relaxed Arakawa–Schu-

bert (RAS) algorithm [*Arakawa and Schubert*, 1974; *Moorathi and Suarez*, 1992] that is used to parameterize convection in the GEOS GCM. Three-hour-averaged planetary boundary layer depths are available from the GEOS-3 DAS. Turbulent mixing within the boundary layer is parameterized using a fractional mixing scheme [*Allen et al.*, 1996a]. Dry deposition is parameterized based on surface type [e.g., *Brasseur et al.*, 1998] with deposition velocities from the study of *Sander and Crutzen* [1996]. Wet removal of soluble species is accomplished through a wet scavenging algorithm described by *Chin et al.* [2000] using precipitation rates from the GEOS-3 DAS. Scavenging by cirrus precipitation [*Lawrence and Crutzen*, 1998; *Liu et al.*, 2001] is not included.

[19] Data from the GEOS-3 DAS are archived on a 48 sigma layer 1° in latitude by 1° in longitude (1° × 1°) grid. The UMD-CTM was run at a 2° × 2.5° resolution with 35 sigma–pressure layers. The model tops of the GEOS-3 DAS and the UMD-CTM are 0.01 hPa. Pressures at the tops and bottoms of the first 25 GEOS-3 DAS and UMD-CTM layers (~1000 to ~37.6 hPa) are identical. The uppermost 23 GEOS-3 DAS layers are mapped onto 10 UMD-CTM layers. The resulting fields are mapped onto a 2° × 2.5° grid. The UMD-CTM grid includes 8 layers between the surface and 850 hPa and 13 layers between 850 and 100 hPa with a vertical resolution in the expected TOPSE tropopause region of 1.4–1.8 km. The sigma–pressure interface of the UMD-CTM is ~242 hPa. Six-hour-averaged *u* and *v* components of the wind are taken from the GEOS-3 DAS. The winds are interpolated in time to the transport time. The vertical velocity is calculated kinematically each transport time step (15 min) by assuming the vertical velocity is zero at the top of the model and integrating downward.

[20] Ertel’s PV was calculated on the UMD CTM grid using fields from the GEOS-3 DAS. The formula for PV can be expressed in the form [e.g., *Nash et al.*, 1996]:

$$PV = -g(\zeta_{\sigma} + f)\partial\theta/\partial p, \quad (4)$$

where *g* is the acceleration due to gravity, ζ_{σ} is the vertical component of the relative vorticity on a sigma surface, *f* is the Coriolis parameter, θ is the potential temperature, and *p* is the pressure. The ζ_{σ} is given by

$$\zeta_{\sigma} = \partial v/\partial x - \partial u/\partial y, \quad (5)$$

where *u* and *v* are the zonal and meridional wind components along the *x* and *y* Cartesian coordinates. Wind components are available at 00, 06, 12, and 18 UT. PV was calculated for every 6-hour period between 1 February and 31 May 2000.

[21] The tropopause pressure was also calculated as a function of latitude and longitude on the UMD-CTM grid. It was defined to be the largest pressure (pressure < 600 hPa) at which Ertel’s PV equaled 2.5 PVU. However, if the resulting tropopause pressure was less than the pressure at the 380 K surface, the pressure at the 380 K surface was used as the tropopause pressure. Estimates of PV at the midlatitude tropopause vary from 1.5 [*WMO*, 1996] to 3.5 [*Hoerling et al.*, 1991] PVU. A value of 2.5 PVU was chosen as a compromise between these values. The impact on strato-

sphere-to-troposphere and troposphere-to-stratosphere exchange of varying the tropopause definition (from 1 to 4.5 PVU) is discussed by *Morgenstern and Carver* [2001].

3.2. ⁷Be and O₃ Data

[22] *Dibb et al.* [2002] describe the procedure used to determine ⁷Be activities during the TOPSE mission. Atmospheric aerosol samples were collected on filters during 12–24 min constant altitude sampling periods. The filters were shipped to a laboratory and analyzed for ⁷Be activity using gamma spectroscopy. Activities were adjusted for loss by decay during the time period between sampling and analysis.

[23] Two hundred and ninety eight aerosol samples were collected on 31 different days during TOPSE flights 5–42. Three of the samples were taken in the stratosphere (PV > 2.5; mean sampling pressure < tropopause pressure). Of the remaining 295, eight had mean sampling pressures within 10 hPa of the tropopause pressure. These eight samples included four samples from deployment 5. ⁷Be mixing ratios (see appendix) from 239 samples exceeded detection limits (0.8–4.5 milli-Becquerels (mBq) SCM⁻¹), which varied with sampling pressure and time. Mixing ratios less than the detection limit were set to the detection limit. This step introduced a positive bias of 0.0–0.4 mBq SCM⁻¹ (0.0–1.8%) in the mean ⁷Be mixing ratio (The mean ⁷Be mixing ratio decreased from 19.9 to 19.5 mBq SCM⁻¹ when the mixing ratios at these points were set to zero). The median and maximum mixing ratios during the TOPSE mission were 11.4 and 175 mBq SCM⁻¹, respectively. The standard deviation was 26.7 mBq SCM⁻¹.

[24] O₃ measurements were collected using the NCAR Atmospheric Chemistry Division (ACD) fast chemiluminescence instrument [*Ridley et al.*, 2002]. HNO₃ measurements were taken using a mist chamber sampling technique and ion chromatographic analysis [*Talbot et al.*, 1999] (*E. Scheuer et al.*, Seasonal distributions of fine aerosol sulfate in the North American Arctic basin during TOPSE, submitted to *Journal of Geophysical Research*, 2002). Five-minute average O₃ and HNO₃ mixing ratios from these measurements are available as part of the 300-s TOPSE data merges that are available through the NCAR ACD. Mean O₃ and HNO₃ mixing ratios for the 12–24 min aerosol sampling periods were calculated by averaging data from the 5-min merges over the aerosol sampling periods.

[25] The effect of seasonal trends in STE on seasonal trends in O₃ and ⁷Be in the troposphere are evaluated in section 5. Aerosol samples collected below 1 km or above the tropopause are not included in the trend analysis. Surface layer samples are excluded in order to remove “trends” associated with bromine chemistry over ice-covered surfaces [*Barrie et al.*, 1988]. Twenty-one samples (three, one, three, six, seven, zero, and one from deployments 1–7, respectively) were collected below 1 km. Three stratospheric samples (two from deployment 3 and one from deployment 5) were also removed. The total number of samples used for the correlation and trend analysis in section 5 is 274.

3.3. ⁷Be Calculations

[26] ⁷Be is produced when neutrons created by collisions between primary cosmic rays and atoms collide with nitrogen and oxygen atoms. Since cosmic rays follow the Earth’s magnetic field, ⁷Be production is largest at latitudes pole-

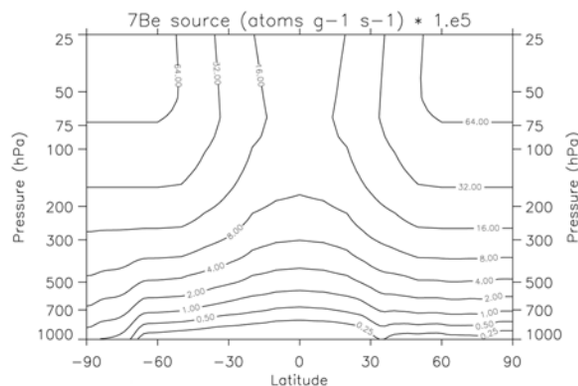


Figure 2. ⁷Be source (atoms g⁻¹ s⁻¹ * 1.e-5) as a function of latitude and pressure.

ward of 60°. In the vertical, production is largest in the lower stratosphere. ⁷Be production also varies with the solar cycle. It is smallest when the solar cycle is near its maximum and largest when the cycle is near its minimum. The solar cycle was near its maximum in 2000. The ⁷Be source used in this study is given by *Lal and Peters* [1967]. The magnitude of the source, 0.080 atoms ⁷Be m⁻² s⁻¹, is appropriate for a solar maximum year. The ⁷Be source used in this study is shown as a function of latitude and pressure in Figure 2.

[27] It is important to note that ⁷Be has a significant tropospheric source. Therefore, in order to study STE using ⁷Be, it is desirable to solve for the distribution of ⁷Be with a tropospheric source (trop-⁷Be) separately from ⁷Be with a stratospheric source (strat-⁷Be) [e.g., *Liu et al.*, 2001]. This was achieved by dividing the ⁷Be source into stratospheric and tropospheric components using the tropopause pressure. Separate calculations for tropospheric and stratospheric ⁷Be began 1 January 2000 following a 6-month spin up from an initial condition of ⁷Be = 0 using GEOS-3 DAS fields for July–December 2000. The spin up did not use data from 1999 because the GEOS-3 assimilation did not begin until November 1999.

[28] Time-averaged (1 February to 31 May 2000) budgets for trop-⁷Be and strat-⁷Be are shown for the globe and the TOPSE region (100°–60°W, 40°–80°N) in Table 1. Overall, seventy-one percent of model-calculated ⁷Be is produced in the stratosphere. This value agrees remarkably well with the value (70%) given by *Lal* [1963] and *Lal and Peters* [1967]. The extremely good agreement is partially fortuitous given uncertainties in the source distribution and the tropopause location. Radioactive decay is the dominant loss mechanism for ⁷Be with a stratospheric source (strat-⁷Be), while wet scavenging is the dominant loss mechanism for ⁷Be with a tropospheric source (trop-⁷Be). Dry deposition plays a relatively minor role. Within the TOPSE region, because of the relatively low tropopause, strat-⁷Be accounts for 84% of the total ⁷Be production. The relative importance of radioactive decay is also enhanced due to low scavenging rates over the northern portion of the TOPSE region.

[29] The zonally averaged model-calculated ⁷Be distribution in mBq SCM⁻¹ (see appendix) and its stratospheric component (strat-⁷Be/total-⁷Be) are shown for the TOPSE

Table 1. Monthly Average Global and TOPSE Region (100°–60°W, 40°–80°N) ⁷Be Budget for Trop-⁷Be and Strat-⁷Be^a

	Trop- ⁷ Be		Strat- ⁷ Be	
	Mass	Loss (%)	Mass	Loss (%)
<i>Global Budget</i>				
Source	3.61		8.69	
Decay	0.90	24.6	6.75	81.6
Wet dep	2.63	72.3	1.45	17.6
Dry dep	0.11	3.1	0.07	0.9
Total	3.64	100.0	8.27	100.0
<i>TOPSE Region Budget</i>				
Source	0.065		0.355	
Decay	0.018	35.4	0.251	90.9
Wet dep	0.029	58.4	0.023	8.5
Dry dep	0.003	6.2	0.002	0.6
Total	0.050	100.0	0.276	100.0

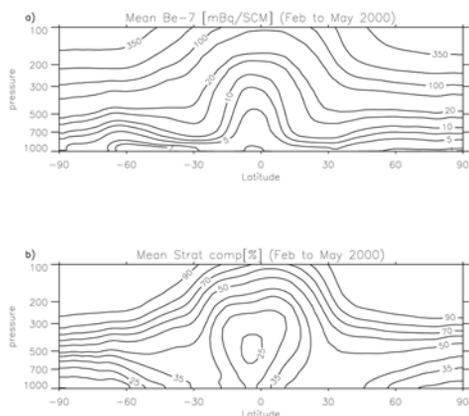
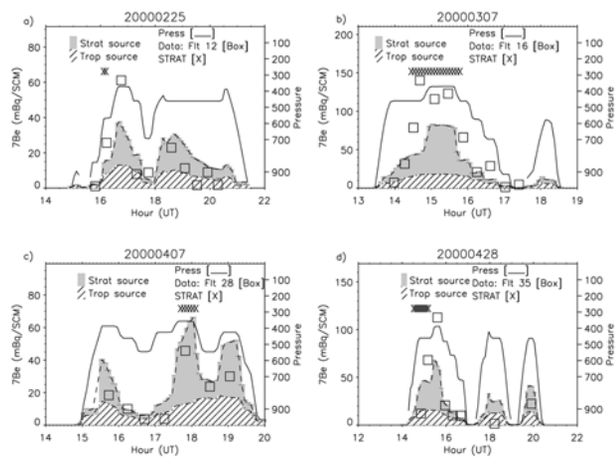
^aThe magnitude of the ⁷Be source and its sinks is given in the mass column (g month⁻¹). The percent each loss term contributes to the total loss is shown in the loss column.

period in Figures 3a and 3b. As expected, stratospheric mixing ratios exceed tropospheric mixing ratios by an order of magnitude. At a given pressure within the troposphere, mixing ratios are largest in the descending branches of the Hadley circulation and lowest at latitudes where scavenging is large. In general, model-calculated mixing ratios and stratospheric components are very similar to values shown by Liu *et al.* [2001] for a ⁷Be simulation without cirrus scavenging. The stratospheric component in the troposphere varies from 15% to 80%. As expected, low values are seen in regions of ascent such as the tropics while high values are seen in regions of subsidence such as the subtropics.

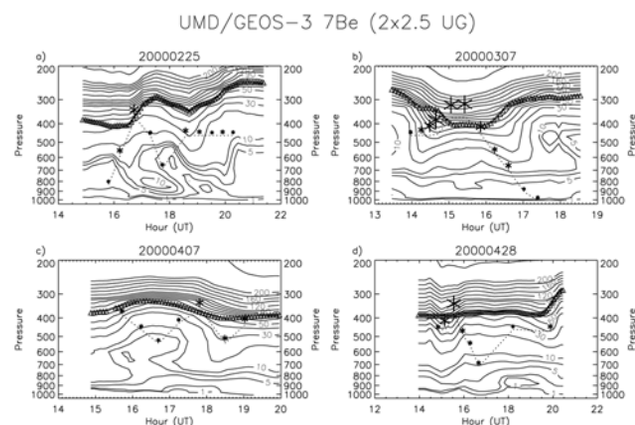
4. Comparison of Model-Calculated and Observed ⁷Be

4.1. Individual Flights

[30] Model-calculated and observed ⁷Be mixing ratios during flights 12 (25 February), 16 (7 March), 28 (7 April),


Figure 3. Zonally averaged mean (1 February to 31 May 2000) model-calculated ⁷Be as a function of latitude and pressure. (a) Mixing ratio in units of 1, 2, 5, 7.5, 10, 15, 20, 50, 100, 200, and 350 mBq SCM⁻¹ and (b) Percent with a stratospheric source. Contour intervals of 20%, 25%, 30%, 35%, 40%, 50%, 60%, 70%, 80%, and 90%.

Figure 4. Model-calculated versus measured ⁷Be on (a) 25 February, (b) 7 March, (c) 7 April, and (d) 28 April. Measured mixing ratios are shown with boxes. The hatched (shaded) region shows the contribution of model-calculated ⁷Be with a tropospheric (stratospheric) source. A solid line shows the mean sampling pressure. Time periods when stratospheric influence is believed to be large are shown with an X.

and 35 (28 April) are compared in Figures 4a–4d and 5a–5d. These flights were chosen because more than changes in pressure controlled ⁷Be variations during these flights. The 25 February, 7 April, and 28 April flights originated in Thule and ended in Churchill. The 7 March flight began and ended in Churchill. In general, model-calculated and observed ⁷Be mixing ratios track well. Within each flight, mixing ratios show a strong dependence on pressure; although considerable variation is also seen during several constant altitude segments (e.g., 18–21 UT, 25 February and 14–15 UT, 7 March). The model does a reasonable job


Figure 5. Curtain plots comparing model-calculated ⁷Be mixing ratios above and below the flight paths. Measured mixing ratios are proportional to asterisk size and range from 0.9 to 113.0 mBq SCM⁻¹. The lowest points on each of the asterisks give sampling pressures. Dates shown are (a) 25 February, (b) 7 March, (c) 7 April, and (d) 28 April. The location of the tropopause is shown with triangles.

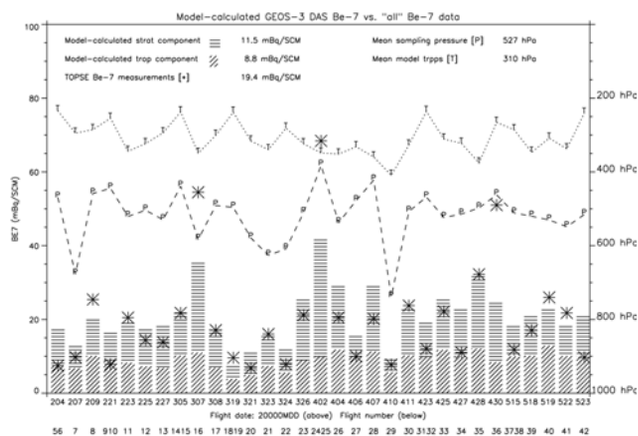


Figure 6. Model-calculated versus measured ⁷Be mixing ratio as a function of flight date. The component of model-calculated ⁷Be with a tropospheric (stratospheric) source is shown with diagonal (horizontal) lines. The mean sampling pressure of the aircraft is shown with “P’s.” The mean tropopause pressure is shown with “T’s.”

of capturing this constant altitude variation. Periods when analysis of temporal fluctuations in ⁷Be, O₃, and HNO₃ data indicated that stratospheric influence was large are shown with X’s [Dibb *et al.*, 2002]. Both model-calculated and observed mixing ratios are enhanced during these periods; although, model-calculated mixing ratios are usually too low. The underestimation of variability is not surprising. Model-calculated mixing ratios were obtained by interpolating 2° × 2.5° model output at 00, 06, 12, and 18 UT to the measurement location and time. Sampling periods were only 12–24 min, which translates to 75–150 km (1/2–1 grid box) at typical C-130 speeds.

[31] Curtain plots for each of the four flights are shown in Figures 5a–5d. The location of the GEOS-3 DAS tropopause is shown by triangles. Lower stratospheric mixing ratios exceed upper tropospheric mixing ratios by approximately an order of magnitude. Clearly, the model is able to maintain large gradients across the tropopause. Comparison of ⁷Be measurements and model output indicates that the location of tropopause folds is reasonably well simulated by the model.

4.2. Mean Distributions

[32] Mean measured and model-calculated ⁷Be mixing ratios, tropopause pressures, and sampling pressures are shown as a function of flight date in Figure 6. Mean values for each date were calculated by weighting individual measurements and model values at the sampling locations by sampling lengths. Overall, mean model-calculated and measured mixing ratios differ by only 5%.

[33] Observed daily average ⁷Be mixing ratios exceed 30 mBq SCM⁻¹ on 7 March, 2 April, 28 April, and 30 April. Mean mixing ratios are high on these days because several of the individual measurements were taken at relatively low pressures and/or within tropopause folds (e.g., 7 March (see Figure 5b), 2 April (not shown), and 28 April (see Figure 5d)). Model-calculated mean mixing ratios on these days are also much larger than normal. In general, model-calculated ⁷Be

mixing ratios are too low within tropopause folds. The low bias may be caused by the relatively coarse resolution of the model or by errors in the source specification. ⁷Be mixing ratios increase rapidly with height in the lower stratosphere. Therefore, small errors in the source distribution could lead to significant errors in mixing ratios. The ability of the CTM to capture the location and time of high ⁷Be mixing ratios is encouraging. It suggests that the location and time of major STE events are reasonably well simulated by the UMD-CTM when driven by GEOS-3 DAS fields.

[34] Mean model-calculated and observed ⁷Be profiles in the southern (latitudes < 60°N) and northern (latitudes > 60°) portions of the TOPSE region are shown in Figures 7a and 7b, respectively. In general, model-calculated mixing ratios are too low in the lower troposphere and slightly high in the upper troposphere. The low bias is most noticeable in the southern portion of the TOPSE region where mixing ratios in the lowest troposphere are too low by almost a factor of 2. An upper tropospheric high bias of 10–30% is evident in the northern portion of the TOPSE region. Model-calculated upper tropospheric mixing ratios in the southern portion of the TOPSE region appear to be reasonable given the large variability of observed mixing ratios in the upper troposphere.

[35] The low bias in the lower troposphere may be caused by an overestimation of wet scavenging within stratiform clouds. Scavenging rates within stratiform clouds are proportional to the precipitation rate and inversely proportional to the liquid water content [Giorgi and Chameides, 1986]. Monthly average GEOS-3 DAS precipitation rates for the 100°–60°W, 30°–60°N region during the TOPSE period differ from satellite-gauge precipitation estimates from the Global Precipitation Climatology Project Version 2 Combined Precipitation Data Set [Huffman *et al.*, 1995; Susskind *et al.*, 1997] by less than 20%. In this study, a cloud condensed water content of 0.5 × 10⁻³ kg m⁻³ [Giorgi and Chameides, 1986] was used when calculating the fraction of a grid volume experiencing large-scale convection. This value is appropriate for highly soluble gases and aerosols but appears to be a factor of 3 too small for ⁷Be [Brost *et al.*, 1991; Liu *et al.*, 2001].

[36] The reasonable agreement in the upper troposphere is significant because previous calculations driven by GEOS

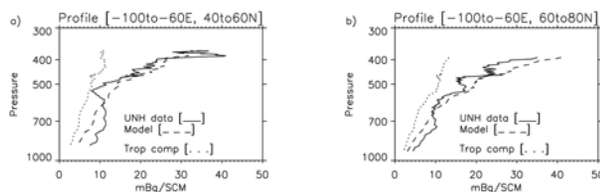


Figure 7. Vertical profiles of ⁷Be mixing ratio for the southern (sampling latitudes ≤ 60°N) (a) and northern (sampling latitudes > 60°N) (b) portions of the TOPSE region. The mean profiles were obtained by applying a 25-point smoother to the raw fields after weighting them by sampling time and sorting them by pressure. The mean profile calculated using measurements (model output) is shown with a solid (dashed) line. The mean profile of trop-⁷Be is shown with a dotted line.

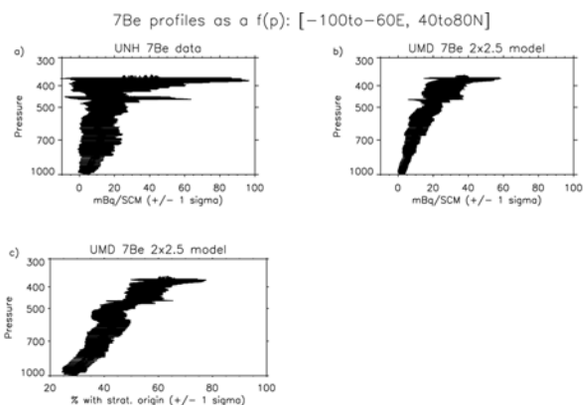


Figure 8. Vertical profiles of observed ⁷Be (a), model-calculated ⁷Be (b), and percent of model-calculated ⁷Be with a stratospheric source (c). The profiles were obtained by applying a 25-point smoother to the raw fields after sorting them by pressure. The standard deviation was calculated for each 25-point subset, and the mean $\pm 1\sigma$ is shown.

DAS fields have overestimated net STE by a factor of ~ 3 [Bey et al., 2001; Liu et al., 2001]. The reasonable agreement is consistent with results obtained by Gettelman and Sobel [2000]. They found that Northern Hemisphere extratropical net winter cross-tropopause fluxes derived using GEOS-1 DAS fields differ from UKMO-based estimates by only 15%. The main difference between this calculation and the calculations of Liu et al. and Bey et al. is the vertical grid. A sigma–pressure coordinate system was used for this simulation while a sigma coordinate system was used for the previous calculations. CTM-calculated STE may be extremely sensitive to the vertical coordinate.

[37] The variability of observed and model-calculated mixing ratios as a function of pressure is shown in Figures 8a and 8b, respectively. Model-calculated mixing ratios are less variable than observed, especially in the upper troposphere. Both the measurements and model output show a peak in variability in the upper troposphere. The variation of the model-calculated stratospheric component (the percent of total ⁷Be with a stratospheric source) with pressure at the location of TOPSE measurements is shown in Figure 8c. It equals 25–30% near the surface and increases steadily with height reaching 40–45% by 650 hPa. The stratospheric component then remains steady to ~ 475 hPa before increasing rapidly and reaching 64% by 365 hPa.

5. Evaluation of Seasonal Changes in O₃ During the TOPSE Period

[38] The evaluation of seasonal changes in O₃ mixing ratios during the TOPSE period will be divided into four sections. In the first section, correlations between O₃ and four stratospheric tracers (PV, model-calculated ⁷Be, measured ⁷Be, and HNO₃) will be assessed using measurements and model output along TOPSE flight paths. In the second section, seasonal trends in O₃, ⁷Be, and PV will be estimated using measurements and model output along TOPSE flight paths. In the third section, the contributions of changes in mass convergence, scavenging, and net radio-

active production to seasonal changes in ⁷Be will be evaluated for the southern and northern portions of the TOPSE region. Implications for seasonal changes in O₃ will be discussed in section 4.

5.1. Tracer–Tracer Correlations During TOPSE

[39] Correlations between O₃ and PV, O₃ and measured ⁷Be, O₃ and model-calculated tot-⁷Be, O₃ and model-calculated strat-⁷Be, O₃ and HNO₃, and model-calculated ⁷Be and PV are shown as a function of θ for sampling latitudes less than and greater than 60°N in Figures 9a and 9b, respectively. Only free troposphere data points are included. O₃ is most highly correlated with ⁷Be. Correlations range from 0.7 to 0.9 in the upper troposphere to 0.5–0.7 in the lower troposphere. The moderate correlations in the lower troposphere indicate that ⁷Be retains some value as a tracer of stratospheric air even in the lower troposphere. Correlations between O₃ and model-calculated ⁷Be (both tot-⁷Be and strat-⁷Be) differ by only a small amount from correlations between O₃ and measured ⁷Be. Correlations between PV and ⁷Be are slightly higher than correlations between PV and O₃ throughout the troposphere. In both cases, correlations with PV decrease rapidly below ~ 290 K. Correlations between O₃ and HNO₃ also fall off rapidly with decreasing θ . Clearly, the value of PV and HNO₃ as stratospheric tracers decreases rapidly with decreasing θ . The θ range for which PV is a useful tracer of stratospheric air can be seen in Figure 10. This figure shows the correlation between PV and model-calculated ⁷Be within the TOPSE region (100°–60°W) as a function of θ and latitude. Correlation coefficients in the lower stratosphere and upper troposphere exceed 0.85 within 15 K of the tropopause. Correlations decrease rapidly in the midtropo-

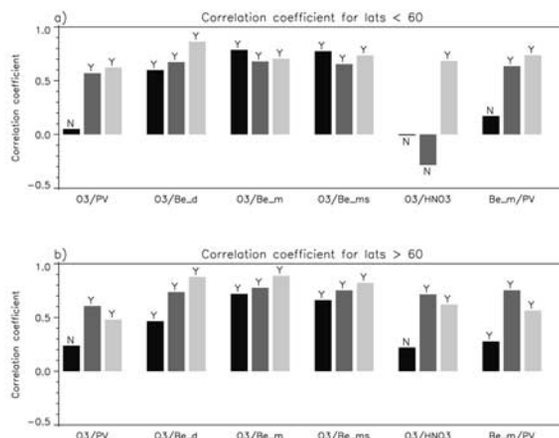


Figure 9. Correlation coefficient between O₃ and PV, O₃ and measured ⁷Be, O₃ and model-calculated tot-⁷Be, O₃ and model-calculated strat-⁷Be, O₃ and HNO₃, and model-calculated ⁷Be and PV. Correlations are shown for $\theta < 290$ K (darkest bar), $290 < \theta < 300$ K (dark bar), and $\theta > 300$ K (lightest bar). (a) Sampling latitudes $\leq 60^\circ$ N and (b) sampling latitudes $> 60^\circ$ N. The correlations in the southern (northern) portion of the TOPSE region were calculated using 34, 40, and 72 (63, 37, and 28) data points. Statistically significant (insignificant) correlations at a 95% confidence level are indicated with a Y(N).

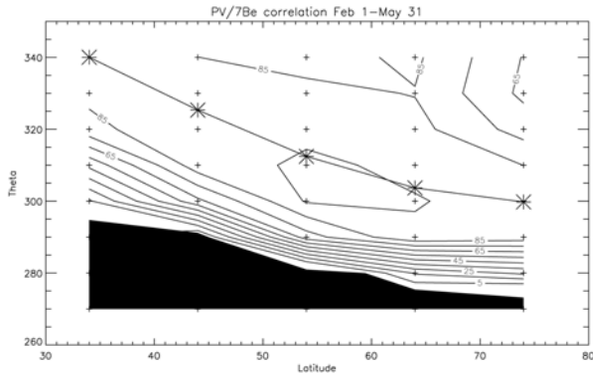


Figure 10. Mean PV/⁷Be correlation for 1 February 2001 to 31 May 2001. The correlations were determined after mapping PV and ⁷Be distributions at 0, 6, 12, and 18 UT onto eight θ surfaces between 270 and 340 K. Area-weighted mean correlations for 100°–60°W within five 10° latitude bins between 30°N and 80°N are shown as a function of θ . Regions where correlations are statistically insignificant at a 95% confidence level and/or negative are shaded. The mean location of the tropopause within each latitude bin is shown with a large asterisk. Data point locations are shown with small “+”s.

sphere reaching ~ 0 for isentropic surfaces more than 40 K below the tropopause. PV, because it varies little in the lower troposphere, is a poor tracer of stratospheric air in the lower troposphere.

5.2. Analysis of Observed Trends in O₃ and ⁷Be

[40] Before assessing trends, the 274 free troposphere data points (see section 3.2) were placed into six bins using three θ ($\theta < 290$ K, 290 K $< \theta < 300$ K, and $\theta > 300$ K) and two PV (PV < 0.75 PVU, PV > 0.75 PVU) categories. Binning tracers by potential temperature (θ) reduces meteorological variability associated with day-to-day changes in air mass, as air parcels remain on the same isentrope during the adiabatic ascent and descent associated with the passage of synoptic-scale features. Diabatic processes will change θ ; however, outside of regions of active convection, changes in θ following air parcels are typically less than 1 K d⁻¹ [Holton, 1979]. Deployment-averaged values of O₃, PV, measured ⁷Be, and model-calculated ⁷Be are shown for each of the bins in Figures 11, 12, and 13. Mean trends of O₃ and PV are shown in Table 2. With one exception, O₃ mixing ratios (see Figures 11a and 11c, 12a and 12c, and 13a and 13c) show a statistically significant increase with time during the TOPSE period. Increases of 4–7 ppbv month⁻¹ are typical. O₃ mixing ratios in the ~ 295 K/ >0.75 PVU bin do not show an increase with time as extensive sampling within the “tropospheric portion” of a 7 March fold caused PV values in this bin to decrease by 0.2 PVU month⁻¹ (Figure 12c and Table 2). This sampling bias caused PV to decrease with time and O₃ to remain steady during the TOPSE period.

[41] Trends in measured ⁷Be, model-calculated tot-⁷Be, and model-calculated strat-⁷Be are shown in Figures 11b and 11d, 12b and 12d, and 13b and 13d and in Table 3. In most cases, agreement is reasonable in well-sampled bins where biases between model-calculated and measured mix-

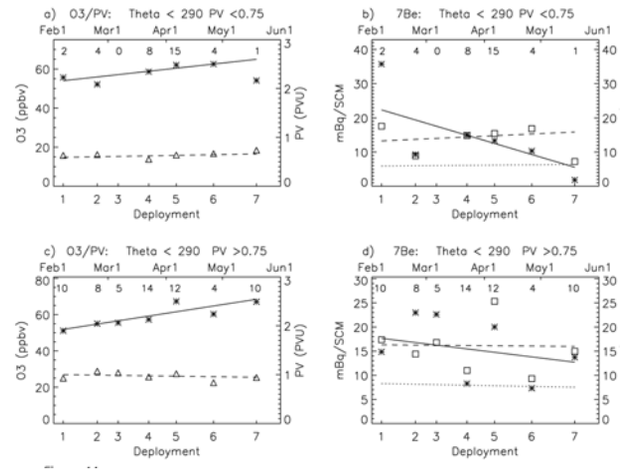


Figure 11. Mean O₃, PV, and measured and model-calculated ⁷Be for each deployment. Mean values during each deployment were calculated by weighting individual measurements during each deployment by the sampling length for ⁷Be. A weighted least squares line was fit to the deployment averages. The line was fit after weighting each deployment average by the number of samples during the deployment (given along the top of each plot). Only sampling points with $\theta < 290$ K are included. (a) O₃ (*) and PV (Δ) for sampling points with PV < 0.75 PVU. (b) measured ⁷Be (*) and model-calculated ⁷Be (boxes) for sampling points with PV < 0.75 PVU. (c) O₃ (*) and PV (Δ) for sampling points with PV > 0.75 PVU. (d) measured (*) and model-calculated (boxes)-⁷Be for sampling points with PV > 0.75 PVU. Linear fits to deployment averages are shown with solid (measured O₃ and ⁷Be) and dashed (PV and model-calculated ⁷Be) lines. A linear fit to deployment average strat-⁷Be is shown with a dotted line.

ing ratios are relatively small and poor in undersampled bins. For example, measurements show a large negative trend (-34.9% month⁻¹) in the <290 K/ <0.75 PVU bin (Figure 11b) and a large positive trend (20.7% month⁻¹) in the 295 K/ <0.75 PVU bin (Figure 12b). Model-calculated

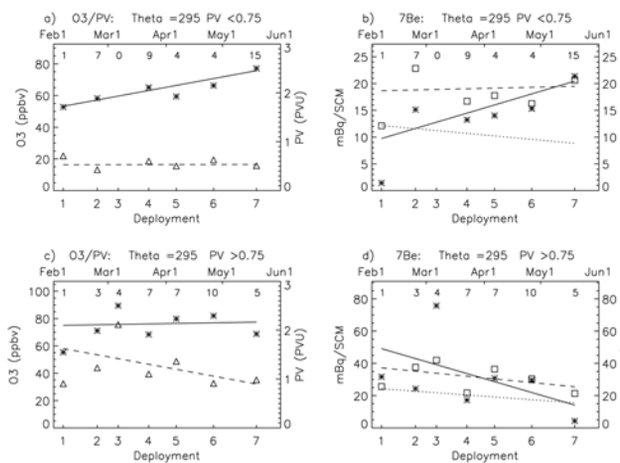


Figure 12. Same as Figure 11, but for sampling points with $290 < \theta < 300$ K.

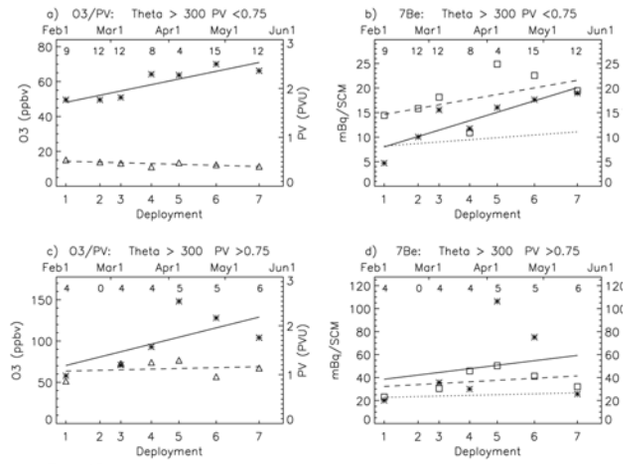


Figure 13. Same as Figure 11, but for sampling points with $\theta > 300$ K.

trends in these bins are much smaller in magnitude. Inspection of Figures 11b and 12b indicates that these large “trends” are dependent upon the representativeness of one or two aerosol samples. For example, the large decrease in the <290 K/ <0.75 PVU bin is driven by two high mixing ratios during deployment 1 and one low mixing ratio during deployment 7 (Figure 11b). O₃ and ⁷Be measurements in the 295 K/ <0.75 PVU bin show a similar pattern with an increase with time between deployments 1 and 7. However, the increase observed in ⁷Be may be overestimated as it is partially caused by one very low mixing ratio during deployment 1. Model-calculated mixing ratios may be more reliable than measured mixing ratios during “undersampled” deployments because model-calculated ⁷Be mixing ratios are less variable than measured mixing ratios (see Figures 8a–8c). Differences between model-calculated and measured trends are significantly smaller in other bins. Discounting the undersampled bins, ⁷Be mixing ratios increase with time in the upper troposphere ($\theta > 300$ K) and change little with time in the lower and middle troposphere.

[42] Deployment-averaged O₃ and ⁷Be mixing ratios were especially large for $\theta > 300$ K during flight 24 of deployment 5 (Figures 13c and 13d). This flight began in Broom-

Table 2. Trends in O₃ and PV for Six Different θ /PV Bins^a

θ /PV	O ₃ (%)		PV	Npts
<290 / <0.75	3.2	5.4	(Y)	34
~ 295 / <0.75	6.5	10.1	(Y)	40
>300 / <0.75	6.6	11.3	(Y)	72
<290 / >0.75	4.8	8.1	(Y)	63
~ 295 / >0.75	0.7	0.9	(N)	37
>300 / >0.75	17.1	17.0	(Y)	28

^aThe θ /PV bins ($\theta < 290$ K, 290 K $< \theta < 300$ K, $\theta > 300$ K) and PV (PV < 0.75 PVU, PV > 0.75 PVU) are shown in column 1. O₃ trends expressed in ppbv month⁻¹ and % month⁻¹ are shown in columns 2 and 3. Statistically significant (insignificant) O₃ trends at a 95% confidence level are identified in column 4 with a Y(N). The significance tests assume a Student’s *t* distribution with 5 or 6 degrees of freedom depending on the number of deployments with at least one sample in the bin. The PV trend in PVU month⁻¹ is shown in column 5. The number of data points in each bin (Npts) is shown in column 6.

Table 3. Trends in Measured-⁷Be, Model-Calculated Tot-⁷Be, and Model-Calculated Strat-⁷Be For Six Different θ /PV Bins^a

θ /PV	⁷ Be (Data)		Tot- ⁷ Be (Model)		Strat- ⁷ Be (Model)	
<290 / <0.75	-4.9	-34.9% (Y)	0.8	5.2% (N)	0.1	2.2% (N)
~ 295 / <0.75	3.1	20.7% (Y)	0.2	1.3% (N)	-1.0	-9.2% (Y)
>300 / <0.75	3.5	25.8% (Y)	2.0	11.3% (Y)	0.8	8.8% (Y)
<290 / >0.75	-1.4	-9.4% (N)	-0.1	-0.6% (N)	-0.2	-2.8% (N)
~ 295 / >0.75	-10.2	-31.0% (Y)	-3.4	-10.7% (Y)	-2.5	-12.6% (Y)
>300 / >0.75	6.0	12.2% (N)	2.7	7.2% (N)	1.2	4.6% (N)

^aThe θ /PV bins are shown in column 1. Trends in mBq SCM⁻¹ month⁻¹ are shown in columns 2, 5, and 8. Trends in % month⁻¹ are shown in columns 3, 6, and 9. Statistically significant (insignificant) trends at a 95% confidence level are identified in columns 4, 7, and 10 with a Y(N).

field, CO and ended in Winnipeg, Manitoba. A curtain plot (not shown) for this flight indicated that several of the samples were taken within a tropopause fold. Only one of the samples was classified as stratospheric using the criteria described in section 3.2; however, three additional samples with elevated O₃ and ⁷Be had mean sampling pressures within 10 hPa of the 6-hour average tropopause pressure. These samples could easily have been classified as stratospheric with a slightly more relaxed criteria. The sensitivity of trends to the tropopause definition is a concern and indicates that trends for PV < 0.75 PVU are more reliable than trends for PV > 0.75 PVU.

[43] For a given pressure level in the troposphere, mean values of θ decrease with increasing latitude (e.g., Figure 1). Therefore, upper tropospheric sampling bins contain more low latitude (latitudes $< 60^\circ$ N) sampling points than lower tropospheric bins. The percent of samples collected at low latitudes is 25% for $\theta < 290$ K, 33% for $\theta \sim 295$ K, and 83% for $\theta > 300$ K. Since ⁷Be increases with time for $\theta > 300$ K, it also increases with time in the southern portion of the TOPSE region. Changes in the northern portion of the TOPSE region are small (see section 5.3.1).

5.3. Analysis of Model-Calculated ⁷Be Trends in the Troposphere

5.3.1. Troposphere-Average Trends

[44] In general, ⁷Be mixing ratios along TOPSE flight paths increased with time in the upper troposphere and decreased or remained constant with time in the middle and lower troposphere. What changes were seen for the troposphere as a whole during the TOPSE period? Changes with time of troposphere-average model-calculated ⁷Be mixing ratios are shown for the southern (100°–60°W, 40°–60°N) and northern (100°–60°W, 60°–80°N) portions of the TOPSE region in Figures 14a and 14b, respectively. Trends in trop-⁷Be, strat-⁷Be, and tot-⁷Be are also shown in Table 4. The troposphere averages were obtained by weighting ⁷Be mixing ratios at tropospheric grid volumes by the mass of each grid volume. Grid volumes are considered to be tropospheric if they reside entirely within the troposphere (i.e., if the pressure at the top of the volume is greater than the GEOS-3 DAS tropopause pressure at that grid point). The contributions of ⁷Be with a tropospheric (trop-⁷Be) and a stratospheric (strat-⁷Be) source to the troposphere-average mixing ratios are also shown.

[45] Troposphere-average mixing ratios increased by 3.0% month⁻¹ in the southern portion of the region and by 1.3% month⁻¹ in the northern portion of the region.

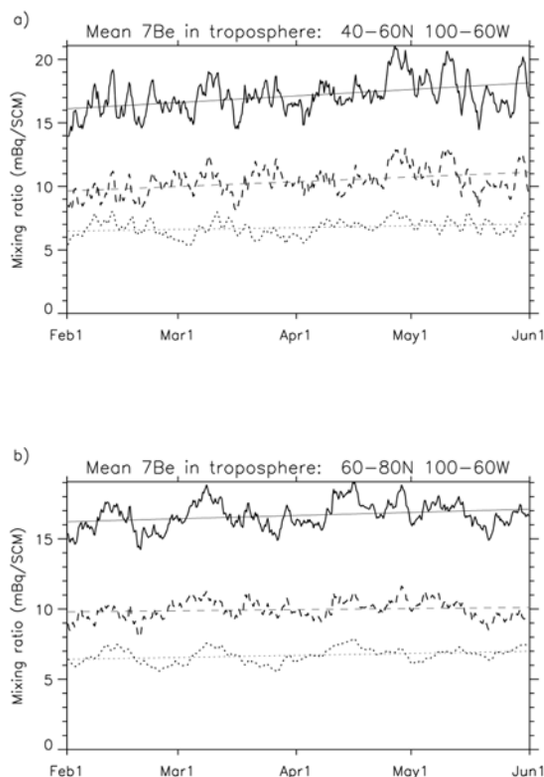


Figure 14. Time series of mean troposphere-average ⁷Be mixing ratios for 100°–60°W, 40°–60°N (a) and 100°60°W, 60°80°N (b). The solid lines show total ⁷Be mixing ratios. The dashed and dotted lines show the contribution from strat-⁷Be and trop-⁷Be, respectively.

Both trop-⁷Be and strat-⁷Be increased with time; although, the relative increase of each component varied between the southern and northern portion of the TOPSE region. Trop-⁷Be mixing ratios increased by 2.1% in the southern portion of the region and by 2.2% in the northern portion of the region. Strat-⁷Be mixing ratios increased by 3.5% in the southern region but by only 0.8% in the northern region.

5.3.2. Budget Analysis

[46] Insight into the processes responsible for the changes with time of troposphere-average ⁷Be mixing ratios in the southern and northern portions of the TOPSE region can be obtained by analyzing the ⁷Be budget in each region. Changes in troposphere-average ⁷Be mixing ratios due to mass convergence (advection and/or STE), imbalances between radioactive production and decay, and scavenging (both wet and dry) are shown for the southern and northern

Table 4. Trends (% Month⁻¹) in Model-Calculated Tot-⁷Be, Trop-⁷Be, and Strat-⁷Be for the Southern and Northern Portions of the TOPSE Region for 1 February to 31 May and 1 February to 30 April

Field	February–May		February–April	
	South	North	South	North
Tot- ⁷ Be	3.0	1.3	4.6	3.2
Trop- ⁷ Be	2.1	2.2	2.3	3.2
Strat- ⁷ Be	3.5	0.8	6.1	3.3

portions of the TOPSE region in Figures 15a–15c and 16a–16c, respectively.

[47] In the southern portion of the region, day-to-day fluctuations in troposphere-average ⁷Be mixing ratios are dominated by changes in mass convergence during February and March and by a combination of changes in mass convergence and scavenging during April and May (Figure 15a). During February, trop-⁷Be decreases due to increases in mass divergence (due to net advection out of the region) and scavenging. Strat-⁷Be mixing ratios remain steady as increases in STE overcome losses by scavenging and advection. Troposphere-average total ⁷Be mixing ratios decrease slightly during this period due to the increase in scavenging. During March, advection continues to be a net sink of trop-⁷Be and presumably strat-⁷Be while STE continues to be a net source of strat-⁷Be. However, neither process changes in strength during March and changes in ⁷Be due to these processes are small. During April, net loss of trop-⁷Be via advection decreases rapidly; although, increases in trop-⁷Be are small because scavenging also increases rapidly during this period (see Figure 15b). Strat-⁷Be increases significantly during April. The increases

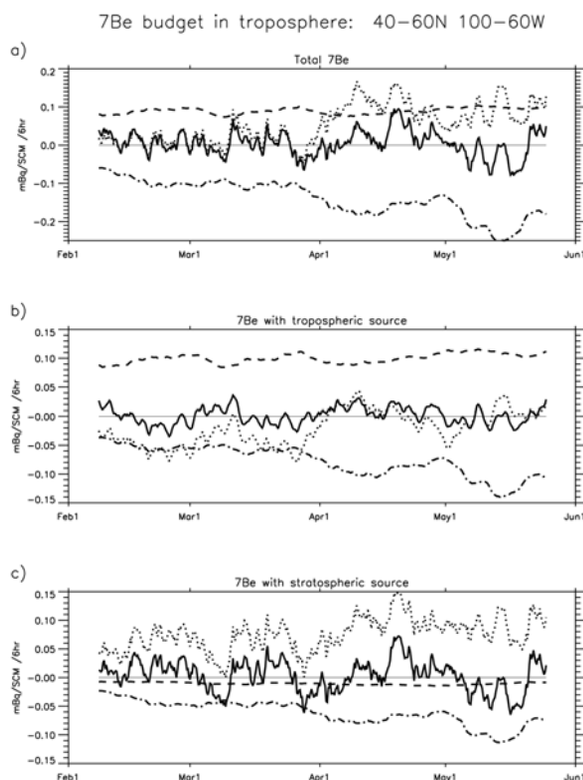


Figure 15. ⁷Be budget within the troposphere for the southern portion of the TOPSE region. The thick solid line shows the change in mixing ratio during each 6-hour period due to all processes. The thick dotted, dashed, and dot-dashed lines show the changes due to mass convergence, imbalances between radioactive production and decay, and scavenging. The zero line is shown with a thin solid line. In order to reduce noise, a boxcar smoother of width 57 (14 days) was applied to the time series before plotting. Because of the smoothing, only fields for 8 February to 24 May are shown.

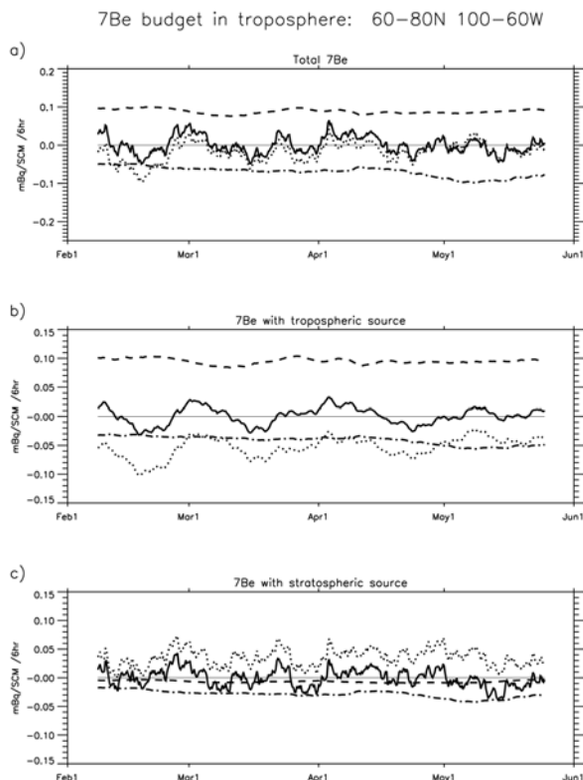


Figure 16. Same as Figure 15, but for northern portion of TOPSE region.

are most likely caused by increases in STE; although, increases in mass convergence due to horizontal advection also play a role. Mixing ratios of both trop-⁷Be and strat-⁷Be decrease during May as changes due to advection and/or STE level off while scavenging continues to increase.

[48] In the northern portion of the region, fluctuations in troposphere-average ⁷Be mixing ratios are smaller and controlled by changes in mass convergence due to advection and/or STE. Seasonal trends are also smaller (see Table 4). In general, mass convergence leads to an increase in strat-⁷Be due to STE (Figure 16c) and a decrease in trop-⁷Be via advection out of the region or troposphere-to-stratosphere exchange (Figure 16b). Contributions to strat-⁷Be from STE (the dotted line in Figure 16c) increase during February remain constant during March and April and decrease slowly throughout much of May. Overall increases during the February–May period are small. Increases in trop-⁷Be exceed increases in strat-⁷Be (see Table 4) because the increase due to STE does not change significantly between mid-February and late May, while the net transport of trop-⁷Be out of the high latitude region decreases from February to May.

[49] Changes in troposphere-average ⁷Be mixing ratios due to seasonal changes in the difference between radioactive production and decay are also shown in Figures 15 and 16. Radioactive production was held fixed throughout the simulation. However, the partitioning of the ⁷Be source between the troposphere and stratosphere changes with time as the mean tropopause moves upward and poleward during the spring. This movement leads to an increase in the

tropospheric component (Figure 15b) and a decrease in the stratospheric component (Figure 15c). With the ⁷Be source held fixed, seasonal increases in scavenging also lead to an imbalance between radioactive production and decay. Decay is suppressed and the net change in production–decay during the TOPSE period is positive (Figure 15c). However, the overall contribution of imbalances between radioactive production and decay to the ⁷Be trend is small.

5.4. Implications for Seasonal Changes in O₃

[50] The differing temporal trends in ⁷Be between the southern/high θ portion of the TOPSE region and the northern/low θ portion of the TOPSE region are consistent with the belief that changes in STE impact the tropospheric middleworld more than the underworld [Hoskins, 1991]. Potential temperature surfaces of less than ~ 300 K are in the underworld throughout the TOPSE period (see Figure 1). It is also consistent with studies that show that total stratosphere-to-troposphere exchange falls off dramatically north of 50°N [Wernli and Bourqui, 2002]. The impact of seasonal increases in STE on the underworld is indirect and difficult to quantify with ⁷Be since many different processes will affect its distribution before it reaches the underworld.

[51] O₃ trends are positive throughout the troposphere but are larger in the tropospheric middleworld than in the underworld. The underworld trends may be partially caused by seasonal changes in STE; however, seasonal changes in photochemical processing and/or in the abundance of O₃ precursors are much more likely explanations. Browell *et al.* [2002] estimated that seasonal changes in STE could explain less than 20% of the observed increase in O₃ at latitudes poleward of 60°N . Seasonal changes in STE are at least partially responsible for the springtime maxima in the tropospheric middleworld. The contribution is likely to decrease with time during the late spring after the springtime peak in STE. This decrease would be consistent with the decrease seen in tropospheric middleworld O₃ and ⁷Be mixing ratios between the sixth and seventh deployments (Figures 13a and 13b).

[52] The percent changes in strat-⁷Be shown in Tables 3 and 4 can be used to estimate the contribution of seasonal changes in STE to the springtime O₃ maxima. This estimation assumes that O₃ with a stratospheric source increases by the same percent as ⁷Be with a stratospheric source (strat-⁷Be). Assuming that this is the case, the increase in O₃ with time ($d\text{O}_3/dt$) can be calculated by multiplying the increase in strat-⁷Be with time [$d(\text{strat-}^7\text{Be})/dt$] by the fraction of total tropospheric O₃ in the region of interest with a stratospheric source [$f_{\text{strat}}(\text{O}_3)$]:

$$d\text{O}_3/dt = d(\text{strat-}^7\text{Be})/dt * f_{\text{strat}}(\text{O}_3) \quad (6)$$

For $\theta > 300$ K (the tropospheric middleworld), changes with time of strat-⁷Be along TOPSE flight paths equals 4.6% (month^{-1}) for $\text{PV} < 0.75$ PVU and 8.8% for $\text{PV} > 0.75$ PVU (see Table 3). Assuming $f_{\text{strat}}(\text{O}_3)$ equals 0.5, these changes would result in a 2.3% (lower bound) to 4.4% increases in tropospheric middleworld O₃. However, model-calculated trends of tot-⁷Be within the tropospheric middleworld (7.2% and 11.3%) are nearly a factor of 2 less than measured trends (12.2% and 25.8%) (see Table 3). If measured trends are accurate, increases in tropospheric middleworld O₃

could be as large as 8.8% (upper bound). Nearly 50% (72 of 146) of samples taken in the southern portion of the TOPSE region had values of θ exceeding 300 K. Changes in strat-⁷Be with time in this region equaled 3.5% for 1 February to 31 May and 6.1% for 1 February to 30 April (see Table 4). These changes could lead to 1.7–6.1% increases in O₃ within this region. The upper bound of 6.1% assumes that $f_{\text{strat}}(\text{O}_3)$ equals 0.5 for this region and that model-calculated trends are too low by a factor of 2.

[53] For $\theta < 300$ K (the underworld), changes with time of strat-⁷Be along TOPSE flight paths are negative in bins where PV decreases with time and only slightly positive in bins where PV is constant or increasing slightly with time (see Table 3). The largest increase in strat-⁷Be with time is 2.2% ($\theta < 290$ K; PV < 0.75 PVU). This change could lead to O₃ increases of $\sim 1\%$. Measured ⁷Be trends are not helpful in determining if 1% is a reasonable upper bound. They range from -35% to $+20\%$. Nearly 80% (100 of 128) of samples taken in the southern portion of the TOPSE region had values of θ less than 300 K. Changes in strat-⁷Be with time in this region equaled 0.8% for 1 February to 31 May and 3.3% for 1 February to 30 April (see Table 4). These changes could lead to 0.4–1.7% (an upper bound) increases in O₃ within this region.

[54] These estimates are highly uncertain for several reasons including the following: (1) Seasonal increases in scavenging affect ⁷Be but not O₃. Therefore, percent increases in strat-⁷Be may be less than percent increases in O₃ with a stratospheric source. (2) Poleward and upward movement of the tropopause during spring decreases the source of strat-⁷Be. This effect is minor (see section 5.3.2) but does lead to a slight underestimation of the importance of seasonal increases in STE. (3) The fraction of O₃ in the region of interest with a stratospheric source [$f_{\text{strat}}(\text{O}_3)$] varies with time and location.

[55] This analysis indicates that seasonal changes in STE could lead to 2–9% increases in O₃ within the tropospheric portion of the middleworld. These increases can be compared to measured increases of 10–15% (see Table 2). Therefore seasonal changes in STE explain 20–60% of observed O₃ increases within the tropospheric portion of the middleworld. Increases in STE result in underworld O₃ increases of less than 1.7%. These increases can be compared to observed O₃ increases in the underworld of 5–10%. Therefore, seasonal changes in STE account for less than a third of observed increases in underworld O₃.

6. Summary

[56] Measurements of tropospheric O₃ between 30°N and 70°N show springtime maxima at remote locations. The contribution of seasonal changes in STE to the springtime O₃ maxima was investigated using meteorological parameters derived from the GEOS-3 DAS, measurements from the TOPSE campaign, and the ⁷Be distribution from a calculation with a 35 sigma–pressure layer version of the UMD-CTM driven by GEOS-3 DAS fields.

[57] Separate calculations were performed for ⁷Be with a tropospheric (trop-⁷Be) and a stratospheric (strat-⁷Be) source. Comparison of model-calculated ⁷Be mixing ratios with measurements revealed that upper tropospheric mixing ratios were reasonable (albeit a bit high, especially in the

northern part of the TOPSE region). The good agreement is significant because previous CTM calculations with a sigma vertical coordinate overestimated upper tropospheric mixing ratios by a factor of 3. Lower tropospheric mixing ratios were too low, especially in the southern part of the TOPSE region, most likely due to an overestimation of scavenging. In general, temporal fluctuations in ⁷Be mixing ratios were well captured even during constant altitude portions of TOPSE flights, although the magnitudes of the fluctuations were usually underestimated.

[58] Analysis of TOPSE O₃ measurements as a function of θ indicated that O₃ mixing ratios increased by 5–10% month⁻¹ for $\theta < 300$ K (the underworld) and by 10–15% month⁻¹ for $\theta > 300$ K (the tropospheric middleworld) between the first (5 February 2000) and last (18 May 2000) deployment.

[59] Model-calculated and measured trends in ⁷Be, a tracer of stratospheric air, were more variable during this period. In general, ⁷Be mixing ratios decreased with time for $\theta < 290$ K and increased with time for $\theta > 300$ K. Model-calculated increases in the middleworld were approximately a factor of 2 less than measured increases. ⁷Be and O₃ trends also varied with latitude since nearly 50% of low latitude ($< 60^\circ\text{N}$) samples were collected in the middleworld while nearly 80% of high latitude ($> 60^\circ\text{N}$) samples were collected in the underworld. Model-calculated mean tropospheric ⁷Be mixing ratios increased by 3.0% month⁻¹ in the southern portion of the TOPSE region and by 1.3% month⁻¹ in the northern of the TOPSE region. The differing temporal trends in ⁷Be between the southern/high θ portion of the TOPSE region and the northern/low θ portion of the TOPSE region are consistent with studies that indicate that changes in STE impact the tropospheric middleworld more than the underworld.

[60] Seasonal changes in advection and STE contributed to a 4.6–8.8% increase in model-calculated ⁷Be with a stratospheric source (strat-⁷Be) along TOPSE flight paths within the tropospheric middleworld. Increases in strat-⁷Be in the southern portion of the TOPSE region totaled 3.5%. Changes in strat-⁷Be along TOPSE flight paths in the underworld were often negative and no larger than 2.2%. Increases in strat-⁷Be within the northern portion of the TOPSE region totaled only 0.8%. Assuming changes in O₃ with a stratospheric source are the same as changes in ⁷Be with a stratospheric source and that 50% of O₃ in the region of interest was produced in the stratosphere, changes in STE explain 20–60% of O₃ increases in the tropospheric middleworld and less than 33% of O₃ increases in the underworld.

Appendix

[61] The ⁷Be source used in this study was expressed in atoms ⁷Be g⁻¹ s⁻¹; however, ⁷Be amounts are usually expressed in mBq or femto-Curies (fCi) per cubic meter at standard temperature and pressure (SCM). Milli-Becquerels and femto-Curies are measures of radioactive activity. One mBq SCM⁻¹ equals 27 fCi SCM⁻¹. The activity of a radioactive sample can be defined in terms of Becquerels where a Becquerel equals one disintegration per second. The activity of ⁷Be is proportional to its mixing ratio. It can be calculated by dividing the mixing ratio of ⁷Be (atoms ⁷Be g⁻¹) by its lifetime with respect to radioactive decay (76.88

* 86,400 s). The mixing ratio of ⁷Be in terms of Bq SCM⁻¹ can be obtained by dividing its activity by the density at STP (1.293 kg m⁻³). One Bq SCM⁻¹ equals 1000 mBq SCM⁻¹. Therefore, 1 mBq SCM⁻¹ equals 0.1947 atoms ⁷Be g⁻¹. Since mBq SCM⁻¹ are proportional to mixing ratio, ⁷Be activities when expressed in terms of mBq SCM⁻¹ will be referred to as mixing ratios.

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References

- Allen, D. J., R. B. Rood, A. M. Thompson, and R. D. Hudson, Three-dimensional radon-222 calculations using assimilated meteorological data and a convective mixing algorithm, *J. Geophys. Res.*, **101**, 6871–6881, 1996a.
- Allen, D. J., P. Kasibhatla, A. M. Thompson, R. B. Rood, B. G. Doddridge, K. E. Pickering, R. D. Hudson, and S.-J. Lin, Transport-induced interannual variability of carbon monoxide determined using a chemistry and transport model, *J. Geophys. Res.*, **101**, 28,655–28,669, 1996b.
- Appenzeller, C., J. R. Holton, and K. H. Rosenlof, Seasonal variation of mass transport across the tropopause, *J. Geophys. Res.*, **101**, 15,071–15,078, 1996.
- Arakawa, A., and W. H. Schubert, Interaction of a cumulus cloud ensemble with the large-scale environment, part 1, *J. Atmos. Sci.*, **31**, 674–701, 1974.
- Bamber, D. J., P. G. W. Healey, B. M. R. Jones, S. A. Penkett, A. F. Tuck, and G. Vaughan, Vertical profiles of tropospheric gases: Chemical consequences of stratospheric intrusions, *Atmos. Environ.*, **18**, 1759–1766, 1984.
- Barrie, L. A., J. W. Bottenheim, R. C. Schnell, P. J. Crutzen, and R. A. Rasmussen, Ozone destruction and photochemical reactions at polar sunrise in the lower Arctic atmosphere, *Nature*, **334**, 138–141, 1988.
- Bey, I., D. J. Jacob, R. M. Yantosca, J. A. Logan, B. D. Field, A. M. Fiore, Q. Li, H. Y. Liu, L. J. Mickley, and M. G. Schultz, Global modeling of tropospheric chemistry with assimilated meteorology: Model description and evaluation, *J. Geophys. Res.*, **106**, 23,073–23,095, 2001.
- Bloom, S. C., L. L. Takacs, A. M. DaSilva, and D. Ledvine, Data assimilation using incremental analysis updates, *Mon. Weather Rev.*, **124**, 1256–1271, 1996.
- Brasseur, G. P., D. A. Hauglustaine, S. Walters, P. J. Rasch, J.-F. Müller, L. K. Emmons, and M. A. Carroll, MOZART, a global chemical transport model for ozone and related chemical tracers, 1, Model description, *J. Geophys. Res.*, **103**, 28,265–28,289, 1998.
- Brewer, A. W., Evidence for a world circulation provided by the measurement of helium and water vapor distribution in the stratosphere, *Q. J. R. Meteorol. Soc.*, **75**, 351–363, 1949.
- Brost, R. A., J. Feichter, and M. Heimann, Three-dimensional simulation of Be-7 in a global climate model, *J. Geophys. Res.*, **96**, 22,423–22,445, 1991.
- Browell, E. W., et al., Ozone, aerosol, potential vorticity, and trace gas trends observed at high latitudes over North America from February to May 2000, *J. Geophys. Res.*, **108**, doi:10.1029/2001JD001390, in press, 2002.
- Chen, P., Isentropic cross-tropopause mass exchange in the extratropics, *J. Geophys. Res.*, **100**, 16,661–16,673, 1995.
- Chin, M., R. B. Rood, S.-J. Lin, J.-F. Müller, and A. M. Thompson, Atmospheric sulfur cycle simulated in the global model GOCART: Model description and global properties, *J. Geophys. Res.*, **105**, 24,671–24,687, 2000.
- Cooper, O. R., J. L. Moody, J. C. Davenport, S. J. Oltmans, B. J. Johnson, X. Chen, P. B. Shepson, and J. T. Merrill, Influence of springtime weather systems on vertical ozone distributions over three North American sites, *J. Geophys. Res.*, **103**, 22,001–22,013, 1998.
- Danielsen, E. F., Stratospheric–tropospheric exchange based upon radioactivity, ozone, and potential vorticity, *J. Atmos. Sci.*, **25**, 502–518, 1968.
- Danielsen, E. F., and V. A. Mohnen, Project Dustorm Report: Ozone transport, in situ measurements, and meteorological analyses of tropopause folding, *J. Geophys. Res.*, **82**, 5867–5877, 1977.
- Dibb, J. E., L. D. Meeker, R. C. Finkel, J. R. Southon, M. W. Caffee, and L. A. Barrie, Estimation of stratospheric input to the Arctic troposphere: ⁷Be and ¹⁰Be in aerosols, at Alert, Canada, *J. Geophys. Res.*, **99**, 12,855–12,864, 1994.
- Dibb, J. E., R. W. Talbot, E. M. Scheuer, G. L. Seid, DeBell, B. A. Lefler, and B. Ridley, Stratospheric influence on the northern North American free troposphere during TOPSE: Be-7 as a stratospheric tracer, *J. Geophys. Res.*, **107**, doi:10.1029/2001JD001347, in press, 2002.
- Dobson, G. M. B., Origin and distribution of polyatomic molecules in the atmosphere, *Proc. R. Soc. London, Ser. A*, **236**, 187–193, 1956.
- Dutkiewicz, V. A., and L. Husain, Stratospheric and tropospheric components of ⁷Be in surface air, *J. Geophys. Res.*, **90**, 5783–5788, 1985.
- Ebel, A., H. Elbern, J. Hendricks, and R. Meyer, Stratosphere–troposphere exchange and its impact on the structure of the lower atmosphere, *J. Geomagn. Geoelectr.*, **48**, 135–144, 1996.
- Elbern, H., J. Hendricks, and A. Ebel, A climatology of tropopause folds by global analyses, *Theor. Appl. Climatol.*, **59**, 181–200, 1998.
- Feely, H. W., R. J. Larsen, and C. G. Sanderson, Factors that cause seasonal variations in beryllium-7 concentrations in surface air, *J. Environ. Radioact.*, **9**, 223–249, 1989.
- Gettelman, A., and A. H. Sobel, Direct diagnoses of stratosphere–troposphere exchange, *J. Atmos. Sci.*, **57**, 3–16, 2000.
- Giorgi, F., and W. L. Chameides, Rainout lifetimes of highly soluble aerosols and gases as inferred from simulations with a general circulation model, *J. Geophys. Res.*, **91**, 14,367–14,376, 1986.
- Hoerling, M. P., T. K. Schaack, and A. J. Lenzen, Global objective tropopause analysis, *Mon. Weather Rev.*, **119**, 1816–1831, 1991.
- Holton, J. R., *An Introduction to Dynamic Meteorology*, Academic, San Diego, Calif., 1979.
- Holton, J. R., On the global exchange of mass between the stratosphere and troposphere, *J. Atmos. Sci.*, **47**, 392–395, 1990.
- Holton, J. R., P. H. Haynes, M. E. McIntyre, A. R. Douglass, R. B. Rood, and L. Pfister, Stratosphere–troposphere exchange, *Rev. Geophys.*, **33**, 403–439, 1995.
- Hoskins, B. J., Towards a PV– θ view of the general circulation, *Tellus*, **43A**, 27–35, 1991.
- Huffman, G. J., R. F. Adler, B. Rudolf, U. Schneider, and P. R. Keehn, Global precipitation estimates based on a technique for combining satellite-based estimates, rain gauge analysis, and NWP model precipitation information, *J. Clim.*, **8**, 1284–1295, 1995.
- Husain, L., P. E. Coffey, R. E. Meyers, and R. T. Cederwall, Ozone transport from stratosphere to troposphere, *Geophys. Res. Lett.*, **4**, 363–365, 1977.
- Koch, D., and D. Rind, Beryllium 10/beryllium 7 as a tracer of stratospheric transport, *J. Geophys. Res.*, **103**, 3907–3917, 1998.
- Koch, D., D. J. Jacob, and W. C. Graustein, Vertical transport of tropospheric aerosols as indicated by ⁷Be and ²¹⁰Pb in a chemical tracer model, *J. Geophys. Res.*, **101**, 18,651–18,666, 1996.
- Lal, D., On the investigations of geophysical processes using cosmic ray produced radioactivity, in *Earth Science and Meteoritics*, compiled by J. Geiss and E. D. Goldberg, 312 pp., North-Holland, New York, 1963.
- Lal, D., and B. Peters, Cosmic ray produced radioactivity on the Earth, in *Handbuch der Physik*, **46/2**, edited by K. Sitte, pp. 551–612, Springer-Verlag, New York, 1967.
- Lawrence, M. G., and P. J. Crutzen, The impact of cloud particle gravitational settling on soluble trace gas distributions, *Tellus*, **50B**, 263–289, 1998.
- Lin, S. J., and R. B. Rood, Multidimensional flux form semi-Lagrangian transport schemes, *Mon. Weather Rev.*, **124**, 2046–2070, 1996.
- Liu, H., D. J. Jacob, I. Bey, and R. M. Yantosca, Constraints from ²¹⁰Pb and ⁷Be on wet deposition and transport in a global three-dimensional chemical tracer model driven by assimilated meteorological fields, *J. Geophys. Res.*, **106**, 12,109–12,128, 2001.
- Logan, J. A., Tropospheric ozone: Seasonal behavior, trends, and anthropogenic influence, *J. Geophys. Res.*, **90**, 10,463–10,482, 1985.
- Monks, P. A., A review of the observations and origins of the spring ozone maximum, *Atmos. Environ.*, **34**, 3545–3561, 2000.
- Moody, J. L., J. C. Davenport, J. T. Merrill, S. J. Oltmans, D. D. Parrish, J. S. Holloway, H. Levy II, G. L. Forbes, M. Trainer, and M. Buhr, Meteorological mechanisms for transporting O₃ over the western North Atlantic Ocean: A case study for August 24–29, 1993, *J. Geophys. Res.*, **101**, 29,213–29,227, 1996.
- Moorthi, S., and M. J. Suarez, Relaxed Arakawa–Schubert: A parameterization of moist convection for general circulation models, *Mon. Weather Rev.*, **120**, 978–1002, 1992.
- Morgenstern, O., and G. D. Carver, Comparison of cross-tropopause transport and ozone in the upper troposphere and lower stratosphere region, *J. Geophys. Res.*, **106**, 10,205–10,221, 2001.
- Nash, E. R., P. A. Newman, J. E. Rosenfeld, and M. R. Schoeberl, An objective determination of the polar vortex using Ertel's potential vorticity, *J. Geophys. Res.*, **101**, 9471–9478, 1996.

- Oltmans, S. J., Surface ozone measurements in clean air, *J. Geophys. Res.*, *86*, 1174–1180, 1981.
- Penkett, S. A., and K. A. Brice, The spring maximum in photo-oxidants in the Northern Hemisphere troposphere and lower stratosphere, *Nature*, *319*, 655–675, 1986.
- Postel, G. A., and M. H. Hitchman, A climatology of Rossby wave breaking along the subtropical tropopause, *J. Atmos. Sci.*, *56*, 359–373, 1999.
- Price, J. D., and G. Vaughan, Statistical studies of cut-off low systems, *Ann. Geophys.*, *10*, 96–102, 1992.
- Rehfeld, S., and M. Heimann, Three dimensional atmospheric transport simulation of the radioactive tracers ²¹⁰Pb, ⁷Be, ¹⁰Be, and ⁹⁰Sr, *J. Geophys. Res.*, *100*, 26,141–26,161, 1995.
- Ridley, B., et al., Ozone depletion events observed in the high latitude surface layer during the TOPSE aircraft program, *J. Geophys. Res.*, *107*, doi:10.1029/2001JD001507, in press, 2002.
- Roelofs, G.-J., and J. Lelieveld, Model study of the influence of cross-tropopause O₃ transports on tropospheric O₃ levels, *Tellus*, *49B*, 38–55, 1997.
- Sander, R., and P. J. Crutzen, Model study indicating halogen activation and ozone destruction in polluted air masses transported to the sea, *J. Geophys. Res.*, *101*, 9121–9138, 1996.
- Schubert, S. D., R. B. Rood, and J. Pfaendtner, An assimilated data set for earth science applications, *Bull. Am. Meteorol. Soc.*, *74*, 2331–2342, 1993.
- Staley, D. O., Strontium-90 in surface air and the stratosphere: Some interpretations of the 1963–75 data, *J. Atmos. Sci.*, *39*, 1571–1590, 1982.
- Susskind, J., P. Piraino, L. Rokke, L. Iredell, and A. Mehta, Characteristics of the TOVS Pathfinder Path A dataset, *Bull. Am. Meteorol. Soc.*, *78*, 1449–1472, 1997.
- Talbot, R. W., et al., Reactive nitrogen budget during the NASA SONEX mission, *Geophys. Res. Lett.*, *26*, 3057–3060, 1999.
- Wang, Y., D. J. Jacob, and J. A. Logan, Global simulation of tropospheric O₃–NO_x–hydrocarbon chemistry, 3, Origin of tropospheric ozone and effects of nonmethane hydrocarbons, *J. Geophys. Res.*, *103*, 10,757–10,767, 1998.
- Wei, M. Y., A new formulation of the exchange of mass and trace constituents between the stratosphere and the troposphere, *J. Atmos. Sci.*, *44*, 3079–3086, 1987.
- Wernli, H., and M. Bourqui, A Lagrangian “1-year climatology” of (deep) cross-tropopause exchange in the extratropical Northern Hemisphere, *J. Geophys. Res.*, *107*, doi:10.1029/2001JD000812, 2002.
- World Meteorological Organization, Atmospheric ozone 1985: Global Ozone Research and Monitoring Project, Rep. 16, 1094 pp., WMO, Geneva, Switzerland, 1996.

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