Annual cycle of horizontal in-mixing into the lower tropical stratosphere

Paul Konopka,1 Jens-Uwe Groß,1 Felix Plöger,1 and Rolf Müller1

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[1] Based on the HALOE and SHADOZ observations of ozone (O3) and on a simple conceptual model of transport and photochemistry, the seasonality of O3 within the stratospheric part of the tropical tropopause layer (TTL) extending between 360 and 420 K potential temperature is discussed. We show that the seasonality of O3 diagnosed on pressure (p) surfaces has a significantly larger annual cycle compared with the same kind of analysis on surfaces with constant potential temperature (θ), in particular around p = 80 hPa, where the strongest annual variation in tropical temperature occurs. Thus by using θ instead of p as the vertical coordinate, the (seasonal) adiabatic variability is removed, and consequently, a much smaller seasonal cycle of O3 remains, which can be understood as a consequence of chemistry, cross-isentropic transport (upwelling), and horizontal, i.e., quasi-isentropic, transport (in-mixing). Furthermore, we show that the observed, θ-related seasonality of O3, with highest values during boreal summer, cannot be understood only by photolytical O3 production in slowly rising air masses, which are well isolated from the extratropics. By using the SHADOZ climatology at θ = 360 K and quantifying the photochemical production of O3 in ascending air above θ = 360 K, we determine the residual variability between the observations (SHADOZ, HALOE) and the calculated O3 values and, consequently, interpret this residuum as horizontal in-mixing from the extratropical stratosphere. We find that between 380 and 420 K, in-mixing contributes to about 40% of the observed O3 mixing ratios during boreal summer.


1. Introduction

[2] The tropical tropopause layer (TTL) has been described as a transition layer coupling the convectively dominated free troposphere with the Brewer-Dobson circulation in the stratosphere. Whereas the upward transport in the free troposphere is locally generated (convection, turbulence), the extratropical wave drag drives the meridional transport and the tropical upwelling within the Brewer-Dobson circulation (the so-called “downward control” principle [Haynes et al., 1991; Holton et al., 1995; Callaghan and Salby, 2001; Fueglistaler et al., 2009]).

[3] As recently deduced from satellite and in situ data, the composition of air within the TTL, in particular at the tropical tropopause (i.e., around p ≈ 100 hPa or θ ≈ 380 K; p, pressure; θ, potential temperature) shows a clear seasonal cycle. In particular, high water vapor (H2O), high ozone O3, low carbon monoxide (CO) during summer (seasons are relative to the northern hemisphere) alternate with low H2O, low O3 and high CO during winter [Folkins et al., 2006; Schoeberl et al., 2006; Randel et al., 2007]. Furthermore, Randel et al. [2007] also inferred that a large annual cycle of O3 (and CO) can be diagnosed on constant p levels within a narrow vertical layer extending around 80 hPa and that this annual cycle is approximately in-phase with the well-known variation in tropical temperature above the tropical tropopause [Reed and Vleck, 1969] that also has the largest amplitude around 80 hPa.

[4] The composition of air within the TTL, in particular the question of how well the tropics are isolated from the extratropics, has been intensively discussed in the past. Although the existence of a mixing barrier between the tropics and extratropics can be diagnosed from the meteorological winds [Chen, 1995; Plumb, 1996; Haynes and Shuckburgh, 2000], evidence of horizontal in-mixing, i.e., of meridional, quasi-isentropic transport from the extratropics into the TTL, was frequently observed in measurements of tracer species [Volk et al., 1996; Avallone and Prather, 1997; Folkins et al., 1999].

[5] Using ozone sondes and aircraft observations of N2O/O3 correlations, signatures of stratospheric contributions were found within the TTL above 14 km and well below the tropopause, identifying this region as a transition zone separating the troposphere from the stratosphere [Tuck et al., 1997]. Recently, Marcy et al. [2007] showed that more than 60% of their airborne in situ HCl observations within the TTL (up to ~100 pptv below θ ≈ 390 K) have stratospheric sources and that this HCl is well correlated with O3.

1ICG-1: Stratosphere, Forschungszentrum Jülich, Jülich, Germany.

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The main purpose of this study is to discuss how the horizontal in-mixing does contribute to the seasonality of O$_3$ within the stratospheric part of the TTL, roughly confined by potential temperature values $q = 360$ K (zero clear sky radiative heating) and 420 K.

2. Annual Cycle of Tropical O$_3$

To analyze the annual cycle of tropical O$_3$ we use the satellite data collected in the 10-year climatology of the Halogen Occultation Experiment (HALOE [Groß and Russell, 2005]) and the in situ observations of the Southern Hemisphere ADDitional OZonesondes (SHADOZ) network [Thompson et al., 2007]. In Figure 1 (top left and right), HALOE O$_3$ climatology based on observations from 1991 to 2002 is shown, averaged over the latitude range $\pm 10^\circ$N, with $p$ and $\theta$ as the vertical coordinate, respectively. Even though HALOE samples the tropics only intermittently in a single year, the 10-year climatology contains enough data for an average annual cycle of observations on a monthly basis. Furthermore, because the HALOE instrument did not measure temperature, the $\theta$ coordinate in the original HALOE data set was derived from the UKMO temperature [Groß and Russell, 2005].

Whereas a clear maximum in summer can be seen at each $p$ level above $\approx 120$ hPa, the use of $\theta$ as the vertical coordinate significantly reduces the seasonal variability of O$_3$ (see O$_3$ along the white isentropes plotted in the top left panel). An example of this effect can be seen at $p = 68/100$ hPa and $q = 420/380$ K (middle and bottom panels), where both the HALOE (pink) and the SHADOZ (green) time series are shown. The SHADOZ time series on $p$ levels are harmonic fits reported by Randel et al. [2007]. Using the observed temperature profiles, the $p$-related observations were transformed to $\theta$ levels (right middle and bottom panels). Each green circle represents the average value over 7 stations (from 11 available SHADOZ stations, the 7 stations closest to the equator were chosen), the error bars denote the station-to-station variance. The error bars of the HALOE climatology denote the standard deviation of all HALOE observations within the considered month and $\pm 10^\circ$N latitude range.

Figure 1. Seasonal cycle of O$_3$ within the TTL averaged within the latitude range $\pm 10^\circ$N using the (left) pressure $p$ and (right) potential temperature $\theta$ as the vertical coordinate as derived from the 10-year HALOE climatology [Groß and Russell, 2005]. In the middle and bottom panels, the corresponding time series are compared with the SHADOZ observations. The black bars estimate the amplitude of the seasonal cycle of O$_3$ on $p$ levels resulting from the seasonal adiabatic variability, i.e., obtained by assuming that $\partial O_3/\partial \theta|_{\theta\text{-const}} = 0$ (see text for details).
[9] It should be noted that the horizontal and vertical resolution of the HALOE instrument is of the order of 400 m and 2 km, respectively [Bruhl et al., 1996], whereas the vertical resolution of the ozone sondes of the SHADOZ network is much higher, i.e., between 150 and 200 m [Thompson et al., 2007]. On the other hand, the HALOE observations sampled over the 1991–2002 period cover the considered ±10°N latitude range almost uniformly. In contrast, the SHADOZ climatology is biased by the geographic positions of the 7 stations considered of which 5 are located in the southern hemisphere. Above the tropical tropopause, the accuracy of both instruments is better than 10% and 5% for the HALOE, and SHADOZ data, respectively [Bhatt et al., 1999; Thompson et al., 2007]. Between 100 and 140 hPa, HALOE accuracy decreases to 25% and the climatological average is biased by the fact that cloud-free conditions are necessary for the HALOE retrieval algorithm. Further, although at p = 100 hPa, the SHADOZ and HALOE data agree reasonably well, there is an offset of about 20 ppbv at θ = 380 K mainly due to the difference between the temperatures measured by the SHADOZ sondes and the UKMO-based HALOE temperature.

[10] To estimate how the choice of the vertical coordinate influences the seasonality of O₃, let us compare the time variation of O₃ on θ and p = const. surface, i.e.:

\[
\frac{\partial O_3}{\partial t}|_\theta = \frac{\partial O_3}{\partial t}|_p + \frac{\partial O_3}{\partial p} \frac{dp}{dt}|_\theta
\]  

(1)

Assuming that ozone variation at θ = const. is negligible, one obtains

\[
\frac{\partial O_3}{\partial t}|_p = -\frac{\partial O_3}{\partial p} \frac{1}{\kappa} \frac{dT}{dt}|_\theta
\]  

(2)

where \(dp/dt\) was derived from the definition of the potential temperature, \(\theta = T(p)/\rho\). In log-pressure coordinates with \(z = -H \ln(p/p_0)\) (\(H = 7\) km), equation (2) can be rewritten as

\[
\frac{\partial O_3}{\partial t}|_z = -\frac{\partial O_3}{\partial z} \frac{1}{\kappa} \frac{dT}{dt}|_\theta
\]  

(3)

Finally, assuming a sinusoidal variation of O₃ and T of the form \(f = f + \Delta f \sin(\omega t + \phi)\) and \(\Delta T/T \approx \Delta T/T\), we obtain

\[
\frac{\Delta O_3}{O_3}|_z = \frac{d \ln O_3}{dz} \frac{H}{\kappa} \frac{\Delta T}{T}|_\theta
\]  

(4)

Thus using the numerical values of \(d \ln O_3/dz\) and \(\Delta T/T\) given by Randel et al. [2007] (Figures 3 and 5), one obtains \(\Delta O_3 = 90, 80, 6\) ppbv at \(p = 68, 83, 100\) hPa where \(O_3\) was set to 400, 200 and 70 ppbv, respectively. The resulting variability (2ΔO₃) is shown in Figure 1 as black vertical lines. Note that this adiabatic variability (defined as \(\partial O_3/\partial t = 0\) at \(\theta = \text{const}\)) is only slightly smaller than the full variability on \(p\) surfaces of the harmonic fits of the SHADOZ time series (green).

[11] We conclude that the seasonal variability of O₃, and in particular its annual cycle, is much larger if pressure is used as the vertical coordinate instead of the potential temperature. This effect dominates around \(p = 80\) hPa (i.e., where \(\Delta O_3/O_3\) has a maximum) where also the strongest annual variation in tropical temperature occurs. Consequently, the large annual cycle of O₃ at \(p\) levels as recently discussed by Randel et al. [2007] contains a significant adiabatic part that can be removed if the annual cycle is diagnosed on \(\theta\) levels. Thus the analysis on \(\theta\) surfaces allows us to separate more precisely the contribution of the cross-isentropic (diabatic) transport from the horizontal in-mixing on the seasonality of O₃. We pursue this idea in the next section.

3. O₃ Budget Within the TTL: A Simple Model

[12] O₃ in the TTL is determined, first, by the tropospheric O₃ mixing ratios which define the entry level, second, by photolytical ozone production during ascent in the TTL and, third, by horizontal in-mixing of O₃-rich stratospheric air. In the following, we discuss the influence of all these processes on the distribution of O₃ within the upper part of the TTL (i.e., above \(\theta = 360\) K and between ±10°N).

[13] Here we construct a simple conceptual model describing these three processes that influence ozone. In particular, we use two-dimensional trajectories (i.e., in \(p \times \theta\) space) to describe vertical transport and calculate photolytical ozone production along such trajectories. The change of the \(\theta\) coordinate of the air parcels is driven by \(\theta\) depicted in Figure 2 (top) whereas the chemical production of O₃ along each trajectory is derived from the ozone production rate \(dO_3/dt\) shown in Figure 2 (bottom).
the latter values result in too strong tropical upwelling. Furthermore, mean vertical velocity derived from the upward propagation of the H$_2$O tape recorder signal have a minimum around 60 hPa of the order of 0.5 K/d ($\approx$0.3 mm/s [Mote et al., 1998; Niwano et al., 2003; Schoeberl et al., 2008], thick green line in Figure 3).

To take these uncertainties into account, we shift $\dot{\theta}$ above $\theta = 350$ K toward lower values using a time-independent offset and replace $\dot{\theta}$ with $[\dot{\theta}(1 - \gamma(\theta - 350))]$ with $\gamma = 0.035$ and 0.55. The thick red (default) and thin gray lines in Figure 3 show the corresponding annual means. Whereas the first scenario reproduces the upward vertical velocity of the H$_2$O tape recorder (at $\theta = 450$ K, the red line crosses the green one), the second case approximates the lower bound of the tropical vertical velocities recently discussed by Yang et al. [2008] (Figure 5) where radiative heating rate calculations with the SHADOZ climatology of O$_3$ and temperature as well as the H$_2$O profiles measured with the a frost-point hygrometer were used. With this scenario we make sure that slowest reasonable upwelling is taken into account even though the coverage of the SHADOZ ($-15^\circ$S $-10^\circ$N) data is more representative for the southern hemisphere and, consequently, the vertical velocities discussed by Yang et al. [2008] probably underestimate the upwelling in the $+10^\circ$N range discussed here.

In the following, we use these scenarios of a reduced $\dot{\theta}$ to estimate the upwelling in the lower tropical stratosphere. Both chosen corrections define scenarios that result in longer residence times than most estimates [Krüger et al., 2008; Plüger et al., 2009]. In particular, the mean residence times between $\theta = 360$ and 380 K vary between 45 and 60 days, for the red and gray curves, respectively, with longest summer residence times given by 71 and 95 days. In this way, we obtain an upper bound of the possible photolytical production of O$_3$ in the tropical lower stratosphere. In addition, to avoid some spurious transport due to the assimilation procedure [Schoeberl et al., 2003], we also use $\dot{\theta}$ calculated without the assimilation increments (thin blue line).

Figure 2 (bottom) illustrates the seasonality of the ozone production rate, $dO_3/dt$ at the equator (O$_3$ production through oxygen photolysis). Using $p$ as the vertical coordinate, $dO_3/dt$ was determined with a photolytical model described by Becker et al. [2000]. The absolute values of $dO_3/dt$ agree fairly well with published results [Osterman et al., 1997] and show an exact semiannual cycle with maxima during the equinox indicating that only the zenith angle and the overhead column of O$_3$ (overhead O$_3$ column is defined in $p$ space and not in $\theta$ space) determine the value of $dO_3/dt$ (not shown). By transferring these values from $p$ to $\theta$ coordinate, the annual cycle of the tropical tropopause pressure slightly modulates this cycle. For example, $p$ varies between 100 and 90 hPa at $\theta = 380$ K in summer and winter, respectively, with the highest $dO_3/dt$ in winter.

Generally, the use of a mean vertical velocity in the tropospheric part of the TTL, i.e., roughly below $\theta = 360$ K, strongly underestimates the real variability of transport dominated by rapid upward convective mass transport and slow large-scale subsidence. Furthermore, the modeling of the tropospheric ozone production, driven by hydrogen and nitrogen radicals [Wennberg et al., 1998], requires very
4. Model Results

Omissions and the mixing ratios of the photochemically formed SHADOZ climatology discussed in the previous section. Bottom, respectively). We overcome these difficulties by confining our two-dimensional information on the atmospheric composition. We written with the SHADOZ climatology). The air parcels need approximately 4–5 months to overcome the distance between 370 and 450 K in rough agreement with the slope of the H2O + 2CH4 tape recorder signal (the red line is derived from the position of the corresponding maxima in the HALOE climatology). The fastest diabatic ascent occurs in winter as a consequence of the seasonality of the tropical temperature.

[24] The accumulated, photochemically produced O3 along the trajectories, P(O3), is depicted in Figure 4 (bottom). Similar to the ozone production rates (Figure 2, bottom), the seasonality of P(O3) shows a strong semiannual cycle for 380 < θ < 450 K. In addition, this semiannual variability is overlaid by a weak annual cycle with a maximum in summer.

[25] This weak annual cycle of P(O3) is driven by the seasonal variation of the temperature and, following our analysis, can be understood as a superposition of two opposite annual cycles: a smaller cycle of dO3/dt with a maximum in winter (see Figure 2, bottom) and a slightly stronger cycle of upwelling with the longest residence time (and strongest contribution to P(O3)) in summer. Generally, longer residence times can lead to a photochemical summer maximum in P(O3) but our sensitivity studies indicate that this effect cannot completely explain the observed summer maximum, in particular within the θ range between 380 and 420 K (Figure 1, top right). Here the values of P(O3) are comparable with the observations only for unrealistically long residence times within the stratospheric part of the TTL (of the order of one year). The reason for this small contribution to P(O3) is low values of dO3/dt at those altitudes.

[26] Finally, we determine the residuum, ΔO3, between the HALOE or SHADOZ climatology and the photochemically produced P(O3) shown in Figure 4 (bottom). We interpret the corresponding fraction, i.e., ΔO3/O3 in %, as a result of horizontal in-mixing from the sub- and extratropics into the considered ±10°N band whereas both advection driven by the horizontal winds and mixing driven by breaking planetary waves contribute to such transport. The percentages of ΔO3/O3, inferred both from the HALOE and SHADOZ climatology, are shown in Figure 5 (top and bottom, respectively).

[27] Thus there is a large annual cycle of in-mixing of O3 into the tropical stratosphere, with the highest contributions (up to 50%) between 380 and 420 K during boreal summer and with slightly higher values derived from the SHADOZ climatology. Furthermore, the SHADOZ-based results show a more pronounced semiannual cycle if compared with the HALOE-based analysis. The pattern deduced from the HALOE- and SHADOZ-based analysis becomes more similar if the HALOE climatology between −15°N and the equator (instead of ±10°N) is used (not shown). This is consistent with the fact that most (5 out of 7) of the SHADOZ stations are in the southern hemisphere and, consequently, do not uniformly cover the latitude range between ±10°N. The differences in the absolute values probably result from the lower accuracy and coarser vertical resolution of the HALOE data as well as from the different temperatures used to transform p-related data on the θ surfaces (see discussion related to Figure 1).

[28] The pattern of in-mixing shows only a weak dependence on the assumed scenarios of upwelling (see Figure 3) although the absolute values vary for summer conditions between 30 and 60% for the slowest (gray line in Figure 3) and fastest ascent scenario (ERA-Interim with no assimila-

Figure 4. (top) Two-dimensional trajectories derived from the mean values of θ shown in Figure 2 (top) and shifted toward lower values as described in the previous section (thick red line in Figure 3). The trajectories were started from all the points of the considered time-θ plane (Δt = 10 day, Δθ = 5 K). Not all trajectories are shown. Within the gray region, i.e., below 360 K, the O3 mixing ratios are overwritten with the SHADOZ climatology. The thick red line denotes the position of maxima of the H2O tape recorder (HALOE climatology, H2O + 2CH4). (bottom) Photochemically formed and accumulated O3, P(O3), calculated along the trajectories shown above. Here dO3/dt plotted in Figure 2 (bottom) was used and the SHADOZ climatology was applied for the initial value at θ = 360 K.

detailed information on the atmospheric composition. We overcome these difficulties by confining our two-dimensional model to the region above θ = 360 K. For the lower boundary at θ = 360 K (and for levels below) we apply the SHADOZ climatology discussed in the previous section.

4. Model Results

[22] The resulting idealized trajectories of ascending air masses and the mixing ratios of the photochemically formed O3 along these trajectories are shown in Figure 4 (top and bottom, respectively).

[23] In particular, the slope of the trajectories measures the residence time at a given θ level. Here only trajectories above 360 K contribute to our calculations (i.e., within, the gray region in Figure 4 (top) O3 mixing ratios are overwritten with the SHADOZ climatology). The air parcels need approximately 4–5 months to overcome the distance between 370 and 450 K in rough agreement with the slope of the H2O + 2CH4 tape recorder signal (the red line is
production of O implicates more photochemically produced O (i.e., diabatic upwelling), quasi-isentropic transport (i.e., in- 

[2009]. Furthermore, our pressure levels is much more pronounced compared with 

quantify the contributions of the cross isentropic transport becomes more likely. Above 

the trajectories, i.e., longer residence time differences result from different contributions of the photo-

chemistry along the trajectories, i.e., longer residence time or from the northern extratropics) cannot be 

determined either, so we now discuss some additional arguments known from the literature.

There are also some other limitations on our simple estimate. First, our analysis does not allow us to say whether the isentropic in-mixing is a purely advective or diffusive process. Furthermore, the direction of in-mixing (from the southern or from the northern extratropics) cannot be determined either, so we now discuss some additional arguments known from the literature.

A weak and permeable subtropical jet in the summer northern hemisphere [Chen, 1995; Haynes and Shuckburgh, 2000] indicates that this in-mixing can be understood as enhanced meridional transport from the northern extratropics into the TTL. In addition, the most pronounced hemispheric asymmetry of the climatological flow pattern in the vicinity of the tropopause originates from the Asian summer monsoon that manifests as a strong anticyclone in the upper troposphere [Dethof et al., 1999; Randel and Park, 2006; Park et al., 2007]. Thus enhanced in-mixing of O3 in summer is likely driven by the Asian summer monsoon extending well into the lower stratosphere (up to about 20 km or θ = 420 K [Randel and Park, 2006]).

The strong hemispheric differences in isentropic transport were also discussed by Tuck et al. [1997] using backward trajectories (Figure 18) and by Chen [1995] using a semi-Lagrangian transport model. Chen [1995] showed that enhanced equatorward transport on θ surfaces between 340 and 380 K occurs both during summer and during winter (semiannual cycle) and that this transport is most

Figure 5. Fraction of in-mixed O3, i.e., ΔO3/O3 in %, derived from the difference between the climatology (HALOE, top; SHADOZ, bottom) and the accumulated P(O3). ΔO3 results from the horizontal in-mixing of O3-rich mid- and high-latitude stratospheric air into the lower tropical stratosphere.

5. Discussion and Conclusions

The presented study shows that the annual cycle of O3 within the lower tropical stratosphere diagnosed on pressure levels is much more pronounced compared with a similar analysis on levels with θ = const. Whereas a significant part of the p-related variability can be understood as an adiabatic effect on a seasonal scale, i.e., caused by the annual cycle of the tropical temperature (and, consequently, by the annual cycle of θ on p = const levels), our analysis on θ levels shows a much smaller amplitude of O3, in agreement with similar results discussed by Fueglistaler et al. [2009]. Furthermore, our θ-related analysis allows us to quantify the contributions of the cross isentropic transport (i.e., diabatic upwelling), quasi-isentropic transport (i.e., in-

mixing) and photochemistry to the observed seasonality of O3 in the lower tropical stratosphere.

The contribution of photolytical production strongly increases with altitude and dominates the fraction of O3 above ≈440 K in agreement with previous studies [see, e.g., Avallone and Prather, 1997]. In particular, the photochemistry shows a pronounced semiannual cycle where the highest photolytical production occurs twice per year during the equinox. Below this level, in particular below θ = 420 K, transport rather than photochemistry influences a significant fraction of O3.

Our analysis of the residuum between the observed O3 and the calculated O3 (i.e., tropospheric contribution + photolytical production) shows that this difference, interpreted here as in-mixing, is significant even if very conservative assumptions are made on the vertical ascent (slower than H2O tape recorder) and on the ozone production (equator values). Between June and August, the highest values around 40% were found in the θ range between 380 and 420 K.

Our analysis also shows that the effect of reduced upwelling in summer and, consequently, enhanced O3 production due to longer residence times in the lower tropical stratosphere is too weak to explain the observed summer maximum in O3. On the other hand, during winter, faster upward transport and better dynamical isolation of the tropics reduce the impact of in-mixing to values of around 20%. This seasonality of transport was also suggested by Yang et al. [2008] who found that during boreal summer the upward mass flux has a near-zero minimum around 70 hPa and, consequently, the mass flux below this level is decoupled from that above. Note that because O3 does not behave as a passive tracer, the mentioned percentages of in-mixing are related only to O3 and estimate the lower bound of the horizontal transport into the lower tropical stratosphere.

There are also some other limitations on our simple estimate. First, our analysis does not allow us to say whether the isentropic in-mixing is a purely advective or diffusive process. Furthermore, the direction of in-mixing (from the southern or from the northern extratropics) cannot be determined either, so we now discuss some additional arguments known from the literature.

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The strong hemispheric differences in isentropic transport were also discussed by Tuck et al. [1997] using backward trajectories (Figure 18) and by Chen [1995] using a semi-Lagrangian transport model. Chen [1995] showed that enhanced equatorward transport on θ surfaces between 340 and 380 K occurs both during summer and during winter (semiannual cycle) and that this transport is most
intensive from the northern summer hemisphere (annual cycle). This supports also our interpretation of the differences between the HALOE and SHADOZ-based results. SHADOZ observations sample the southern hemisphere more effectively and, consequently, enhance the relative contribution of the semiannual in comparison to the annual cycle.

[36] In contrast to the absolute values, the annual cycle of in-mixing is a very robust feature of our analysis. Finally, our study also shows that the picture of the TTL where “O₃-poor air of tropospheric origin enters and rises slowly in near isolation” [Avalone and Prather, 1997] should be revised, in particular between 380 and 440 K, by a picture of the TTL where the annual cycle of upwelling and in-mixing modulates the composition of air within the TTL.

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