

Quantification of the factors controlling tropical tropospheric ozone and the South Atlantic maximum

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[1] We quantify the processes controlling the tropical tropospheric ozone burden with particular attention to the tropical Atlantic, using a global chemical transport model (GEOS-Chem) constrained by satellite and in situ observations of O₃, NO₂, and HCHO. Lightning is the dominant contributor to tropical tropospheric O₃, accounting for more than 37% of the O₃ burden over the Atlantic on annual average. The contributions from biomass burning, soils, and fossil fuels are 4 to 6 times smaller, despite comparable source strengths. This discrepancy can be explained by the tropical ozone production efficiency of lightning (32 mol/mol), soils (14 mol/mol), biomass burning (10 mol/mol), and fossil fuel (13 mol/mol) sources, as calculated using sensitivity simulations with a 1% perturbation. The role of volatile organic compound emissions on the tropical Atlantic ozone burden is negligible (<2.5%). Stratosphere-troposphere exchange accounts for less than 5% of the regional O₃ burden. The tropical Atlantic O₃ burden is more strongly influenced by nitrogen oxides from Africa (>30%) than from South America (>18%) or the eastern tropics (>11%). Lightning is responsible for more than 39% of the atmospheric oxidation capacity, higher than other sources. The dominant sources of uncertainty in the tropical oxidation rate are the lightning magnitude and the cloud convective parameterization.

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1. Introduction

[2] The tropical troposphere largely controls the oxidation capacity and radiation budget of the global atmosphere. Ozone (O₃) plays a key role in the global oxidizing power of the atmosphere, as the main producer of OH, the primary atmospheric oxidant [Thompson, 1992]. Ozone production is driven by the supply of nitrogen oxides from lightning, biomass burning, fossil fuel combustion, and soils [Jacob *et al.*, 1996]. Dynamics also strongly influence photochemistry, forcing convergence in the lower troposphere, and vertical ascent of air masses through deep convection into the upper troposphere where chemical species have a longer lifetime, and can be redistributed globally. We use a global chemical transport model to disentangle these processes by interpreting satellite retrievals of tropical tropospheric O₃.

[3] Tropical tropospheric O₃ columns have been inferred from a number of instruments including the GOME [Burrows *et al.*, 1999], and the OMI [Levelt *et al.*, 2006], and MLS instruments [Waters *et al.*, 2007]. The GOME instrument

onboard the European Remote Sensing-2 satellite provided continuous global monitoring of O₃ atmospheric columns through observation of solar backscatter over 1995–2003. GOME observes the atmosphere in the nadir view with a 40 km along track by 320 km across track horizontal resolution. Daily tropospheric O₃ retrievals have been performed by Liu *et al.* [2005]. The OMI and MLS instruments were launched in July 2004 onboard the Aura spacecraft. OMI is a nadir-scanning instrument that detects backscattered solar radiance to measure the O₃ column with near global daily coverage at a resolution of up to 13 km × 24 km at nadir. The MLS instrument is a thermal-emission microwave limb sounder that measures vertical profiles of O₃ in the mesosphere, the stratosphere, and the upper troposphere from limb scans [Froidevaux *et al.*, 2007]. Daily tropospheric O₃ columns have been retrieved by Ziemke *et al.* [2006] by combining measurements from both MLS and OMI. The measurement uncertainty for both retrievals is 5–8 Dobson units (DU).

[4] The middle and upper troposphere of the tropical Atlantic exhibits a persistent ozone maximum as part of the well-known zonal wave-one [Fishman *et al.*, 1990; Logan and Kirchhoff, 1986; Thompson *et al.*, 2003]. At this altitude, ozone is critical for the global warming earth system, with maximum positive radiative effect [de F. Forster and Shine, 1997]. Numerous studies have attempted to understand the origin of that ozone maximum, giving evidence of various influence by the different regional sources, such as biomass burning [Pickering *et al.*, 1996; Edwards *et al.*, 2003; Jenkins

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and Ryu, 2004b; Sauvage et al., 2006], biogenic sources [Meyer-Arnek et al., 2005], lightning [Thompson et al., 2000; Martin et al., 2002; Jenkins and Ryu, 2004a], dynamics [Krishnamurti et al., 1993, 1996], long-range transport [Chatfield et al., 2004], as well as stratospheric-tropospheric exchange [Weller et al., 1996]. Moxim and Levy [2000] through model analysis quantified the tropospheric mass of NO_x transported into that region during September. More recently Wang et al. [2006] have suggested that both the Hadley and Walker circulation can contribute to the zonal wave-one pattern.

[5] Despite these accomplishments, many critically important gaps remain in our knowledge of the tropical tropospheric ozone, the Atlantic ozone maximum, and atmospheric oxidation. For instance, there has been no quantitative estimation of the different sources contributing to the annual O₃ maximum. Indeed, such analysis implies the use of a thoroughly evaluated global chemical transport model, with an accurate emission inventory of O₃ precursors, such as NO_x. Moreover, no study to date has addressed the contribution of each source to OH concentrations. The short OH lifetime (around 1 s) results in a chemical rather than transport control dependence [Bloss et al., 2006]. Manning et al. [2005] suggest a recent 20% decrease in global OH concentrations. Labrador et al. [2004] find that global mean OH is highly sensitive to the NO_x from lightning. A decadal rise in OH simulated with the MOZART-2 chemical transport stems from an increase in lightning NO_x [Fiore et al., 2006]. Determining processes controlling the abundance of OH are then critical to understand how the oxidation capacity of the troposphere is changing.

[6] To address these outstanding questions, we use a three-dimensional global chemical transport model (GEOS-Chem), with emissions of NO_x (lightning, biomass burning, soils) and volatile organic compounds (biomass burning) constrained by in situ and satellite observations [Sauvage et al., 2007]. The emission inventory in that simulation has been validated with in situ measurements from aircraft (MOZAIC), and ozonesondes (SHADOZ) as well as GOME satellite measurements over the tropics. The resultant simulation provides a useful tool to fully understand the ozone maximum over the Atlantic. A summary of that simulation is given in section 2.

[7] The purpose of this paper is (1) to conduct sensitivity studies on NO_x emissions with a global chemical transport model to understand the processes controlling tropical tropospheric O₃ and the maximum over the South Atlantic, and (2) to quantify the influence of different sources and regions on the South Atlantic. We will address these specific points in section 3, after providing in section 2 an overview of the chemical transport model and developments by Sauvage et al. [2007]. Finally, we estimate errors in the factors controlling tropical tropospheric O₃ in section 4.

2. Description of the Simulation and Methodology

[8] We use the global chemical transport model (GEOS-Chem, Bey et al. [2001], <http://www-as.harvard.edu/chemistry/trop/geos/index.html>) driven with GEOS-4 meteorological fields for the year 2000. The GEOS-Chem model includes a detailed simulation of tropospheric O₃-NO_x-hydrocarbon chemistry as well as of aerosols and their precursors. The

model presently includes sulfate, nitrate, ammonium, black and organic carbon, mineral dust and sea salt [Park et al., 2004; Park et al., 2005; Alexander et al., 2005; Fairlie et al., 2007]. The aerosol and gaseous simulations are coupled through formation of sulfate and nitrate, HNO₃(g)/NO₃⁻ partitioning of total inorganic nitrate, heterogeneous chemistry on aerosols [Jacob et al., 1999; Evans and Jacob, 2005], and aerosol effects on photolysis rates [Martin et al., 2003]. The stratosphere to troposphere flux of O₃ in the model of 500 Tg/yr is prescribed by the synthetic ozone method [McLinden et al., 2000].

[9] The model version used here is based on version 7-02-04 with substantial improvements in emissions and heterogeneous chemistry by Sauvage et al. [2007] as summarized below. The spatial distribution of lightning is scaled to reproduce seasonal mean lightning flash rates from the lightning imaging sensor and optical transient detector satellite instruments [Christian et al., 2003] to improve the spatial distribution of lightning in the model. Emissions of lightning NO_x remain linked to deep convection following the parameterization of Price et al. [1997] with vertical profiles from Pickering et al. [1998] as implemented by Wang et al. [1998]. The magnitude of the global lightning source is 6.0 Tg N/yr of which 3.3 Tg N/yr is within 24°S–24°N. The evaluation of ozone vertical profiles conducted by Sauvage et al. [2007] at all tropical sites as part of the Measurement of Ozone and Water Vapor by Airbus In-service Aircraft (MOZAIC) and Southern Hemisphere Additional Ozonesondes (SHADOZ) programs has shown that the local redistribution of lightning emissions reduces the original model bias by 5–20 ppbv, versus the in situ O₃ measurements in the upper troposphere.

[10] Sauvage et al. [2007] also applied tropospheric NO₂ and HCHO columns from GOME as top-down constraints on emission inventories of NO_x from soils and biomass burning, and VOCs emitted from biomass burning. Biomass burning emissions include 7 Tg N/yr as NO, 4 Tg C/yr as HCHO, and 27 Tg C/yr as lumped alkenes. The soil source of 9 Tg N/yr is from Jaeglé et al. [2005]. The top-down constraints on emissions reduced model biases in the lower troposphere by 5–20 ppbv versus in situ observations from MOZAIC program and SHADOZ network.

[11] Uptake of HO₂ on biomass burning aerosol was disabled based on recent laboratory [Thornton and Abbott, 2005] and field measurements [Yamasoe et al., 2000]. Uptake of HNO₃ was implemented with a reaction probability of 0.1 as measured by Hanisch and Crowley [2003]. These two updates to the heterogeneous chemistry improved the lower tropospheric O₃ simulation by 5 ppbv versus MOZAIC observations in biomass burning and desert regions [Sauvage et al., 2007].

[12] The evaluation by Sauvage et al. [2007] of the tropical simulation compared to MOZAIC and SHADOZ in situ data demonstrated the capability to represent the main tropospheric ozone features. The simulation generally reproduces MOZAIC and SHADOZ tropical O₃ profiles to within 5 to 10 ppbv. GOME observations are well simulated for NO₂ ($r^2 = 0.86$, bias = 2×10^{14} molecules cm⁻²) and HCHO ($r^2 > 0.6$, bias = 6×10^{14} molecules cm⁻²).

[13] We use this improved simulation to assess the budget and origin of tropical (24°S–24°N) O₃ with particular attention to the South Atlantic (24°S–0°/35°W–10°E). Sensitivity studies are conducted by perturbing the main

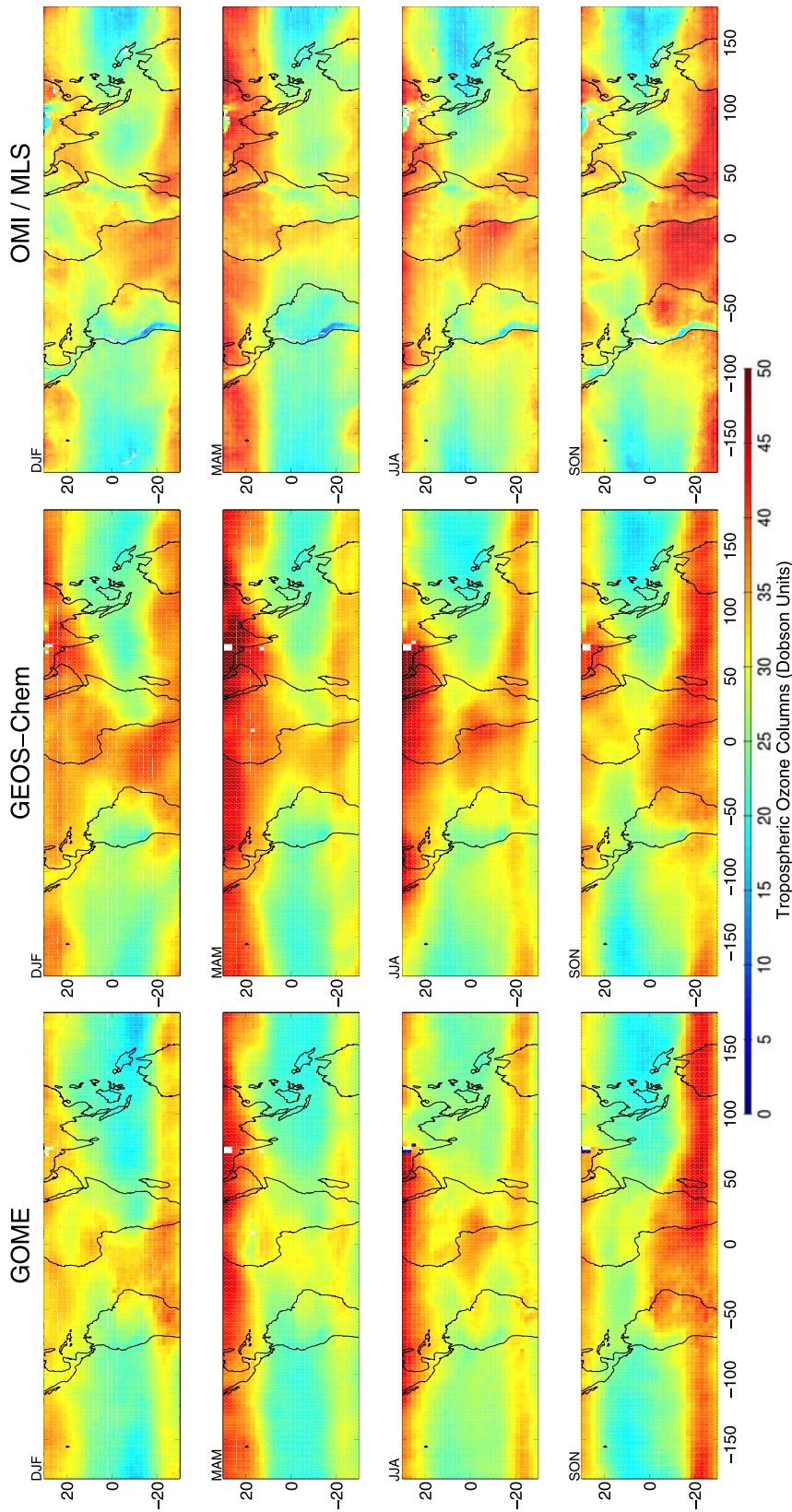


Figure 1. Seasonal tropospheric ozone columns (DU), as seen by GOME for the year 2000 (left), GEOS-Chem for the year 2000 (middle), and OMI/MLS for 2004–2005 (right).

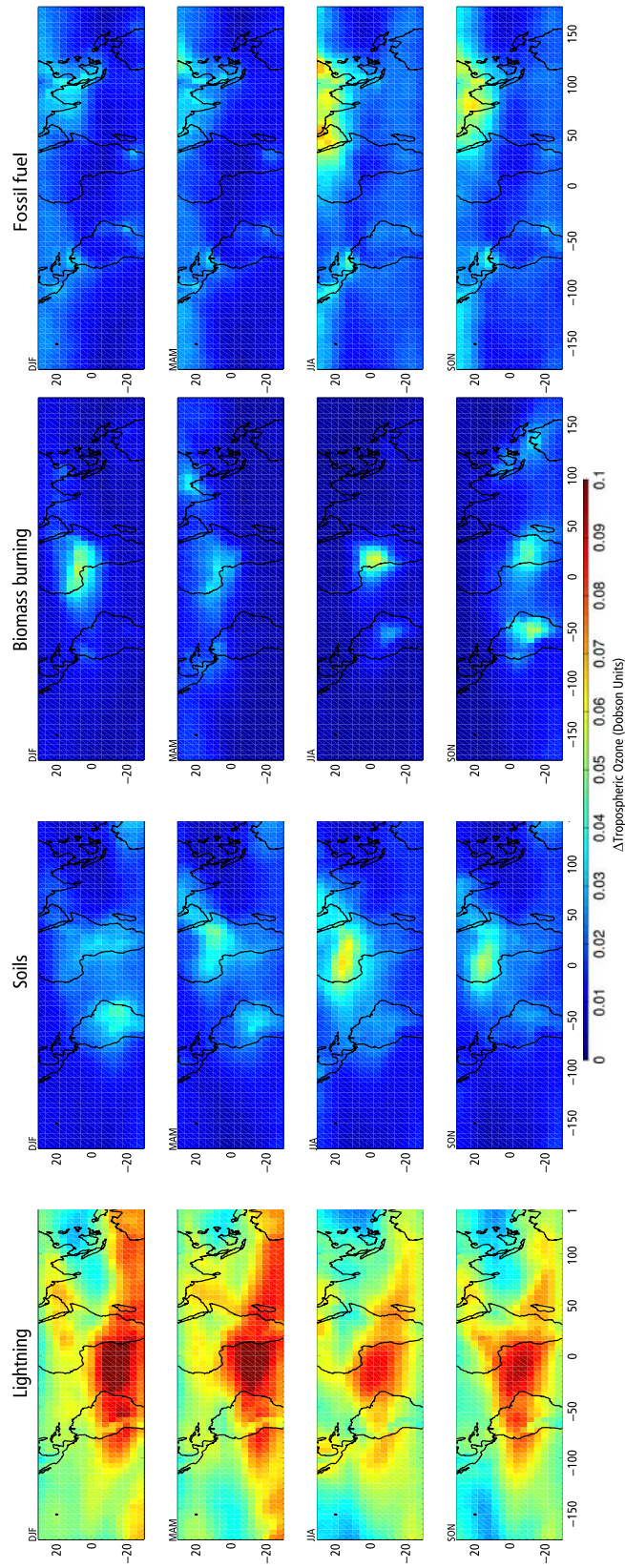


Figure 2. Calculated seasonal sensitivity of the tropospheric ozone column to the NO_x source, as determined by decreasing each source by 1%. The tropospheric ozone column differences (DU) between standard and sensitivity simulations are represented for lightning, soils, biomass burning, and fossil fuel combustion.

Table 1. Source Contribution to the South Atlantic Ozone Burden (Tg O₃)^a

Season	Total ^b	Light. NO _x	Soil NO _x	Biomass burning NO _x	Fossil fuel NO _x	STE O ₃	Background ^c	Afr. NO _x	S. Am. NO _x	East NO _x
DJF	8.87	3.48 ^d (0.025) ^e	0.67 (0.007)	0.81 (0.004)	0.38 (0.004)	0.39	2.26	3.00 (0.017)	1.66 (0.013)	0.79 (0.006)
MAM	7.73	3.57 (0.022)	0.67 (0.006)	0.53 (0.008)	0.31 (0.004)	0.22	1.65	2.83 (0.012)	1.26 (0.009)	0.66 (0.005)
JJA	8.93	2.74 (0.017)	0.69 (0.005)	0.78 (0.005)	0.80 (0.003)	0.47	3.00	2.67 (0.008)	1.28 (0.008)	1.40 (0.010)
SON	9.32	2.90 (0.019)	0.57 (0.003)	1.05 (0.008)	0.70 (0.003)	0.58	3.06	2.63 (0.010)	2.12 (0.011)	1.10 (0.008)
Annual	8.71	3.17 (0.021)	0.62 (0.006)	0.79 (0.005)	0.60 (0.003)	0.42	2.50	2.80 (0.012)	1.60 (0.010)	0.98 (0.007)

^aSources and regions do not sum to 100% due to chemical nonlinearity.

^bThe South Atlantic is defined over 35°W–10°E/24°S–0°.

^cThe ozone background is calculated here using a simulation that excludes all tropical NO_x emissions.

^dValues represent total contribution determined by difference with sensitivity simulation that excludes that source.

^eValues in parenthesis indicate the sensitivity to a 1% perturbation.

NO_x sources: lightning, biomass burning, soils, and fossil fuel combustion to assess their relative contribution. In order to investigate the stratospheric contribution on tropical tropospheric ozone, we use a tagged ozone tracer to follow the stratospheric ozone flux across the tropopause as described by Fiore *et al.* [2002]. The tagged simulation subjects ozone produced in different regions of the atmosphere to archived three-dimensional fields of production and loss frequencies, allowing tropospheric ozone to be deconstructed into components from stratosphere and troposphere. When discussing the dynamical influence (section 3.2), we also comment on the regime south of 24°S. We conduct simulations for the year 2000 following a spin-up of 6 months.

3. Tropical Tropospheric Ozone

[14] In the following section, we decrease NO_x emissions by 1% to examine the sensitivity of O₃ to perturbations that may result from anthropogenic and natural activity or an altered climate. This small perturbation is designed to account for nonlinear O₃ chemistry especially in the lower troposphere. *Kunhikrishnan and Lawrence* [2004] find that nonlinearity associated with changes in photochemical regime becomes important for perturbations larger than 20–30%. For comparison and to approximate the role of each source, we also perform sensitivity studies by completely excluding a specific NO_x source (–100%). Since ozone production increases with NO_x more rapidly at low concentrations of NO_x [Liu *et al.*, 1987], both approaches will underestimate the true source contribution. Values will be preceded by “>” to emphasize this nonlinearity. For ease of comparison to the tropospheric ozone column (TOC) and the O₃ burden, we present the absolute difference between the standard simulation and the simulation with 100% reduction, followed in parenthesis by the results from the 1% perturbation. We also determine an O₃ background to estimate the enhancement for each source. It is quite challenging to estimate an O₃ background due to ambiguity in definition. As our study is based on O₃ sensitivity to NO_x emissions, we determine the O₃ background by excluding all NO_x sources over the tropics.

3.1. The Tropics and the South Atlantic

[15] Figure 1 shows seasonal tropospheric O₃ columns from two satellite instruments (GOME: left panels; OMI: right panels) that we interpret here with GEOS-Chem simulations (middle panels). The three data sets exhibit similar features despite OMI/MLS data being for a different

year. The GEOS-Chem simulation well represents GOME ($r^2 > 0.9$, bias = 1.5 DU) and OMI/MLS ($r^2 > 0.8$, bias = 0.6 DU) on annual average over the tropics. The O₃ minimum over the Pacific is related to frequent convection of depleted O₃ associated with convection [Folkens *et al.*, 2002] and the Walker circulation [Krishnamurti *et al.*, 1993]. Persistent enhanced TOC are observed over the Atlantic in 2000 by GOME and in 2004 by OMI/MLS. The simulated ozone burden over the Atlantic is 10 Tg on annual average, with a maximum of 10.7 Tg in September–November (SON), during the maximum intensity of the zonal wave-one.

3.1.1. Sensitivity to Sources

[16] Figure 2 shows the seasonal sensitivity of tropospheric O₃ columns to NO_x emissions from either lightning, biomass burning, soils or fossil fuel combustion as determined from the difference between the standard simulation and a sensitivity simulation with 1% reduction. Table 1 contains the influence of the different NO_x sources on the O₃ burden calculated over the South Atlantic region. Lightning has the largest influence on tropical tropospheric ozone, accounting for >7.7 DU (0.056 DU) on annual average over the tropics and accounting for >37% of the annual average O₃ burden over the South Atlantic. Lightning has a large influence over a broad area [Lamarque *et al.*, 1996], affecting O₃ production far downwind of emissions [Jacob *et al.*, 1996]. As a result the spatial variation of the lightning influence on O₃ is significantly correlated with the tropical TOC ($r^2 = 0.93$). For comparison this correlation is much lower globally ($r^2 < 0.45$). The similarity is also apparent between tropical TOC (Figure 1) and the lightning influence on tropical TOC (Figure 2). The maximum lightning influence is found largely in the Southern Hemisphere (Figure 2) where the persistent zonal wave-one pattern is apparent in the three data sets (Figure 1). The minimum influence is found over the northern Pacific Ocean, where the TOC is less than 25 DU. Lightning emissions are clearly the dominant source driving the zonal wave-one. For instance lightning accounts for 12.8 DU of the South Atlantic maximum on annual average, similar to the study of *Martin et al.* [2002]. The seasonal influence of lightning is maximum in DJF–MAM with >8.3 DU (0.06 DU), reflecting higher lightning activity and weaker biomass burning emissions. Lightning explains >39% of the South Atlantic O₃ burden in DJF, versus >30% in JJA.

[17] The contributions of biomass burning, fossil fuel combustion, and soil NO_x emissions to the TOC are, respectively, 6, 5, and 4 times smaller (with the 1% sensitivity approach) than that from lightning. The contribution from

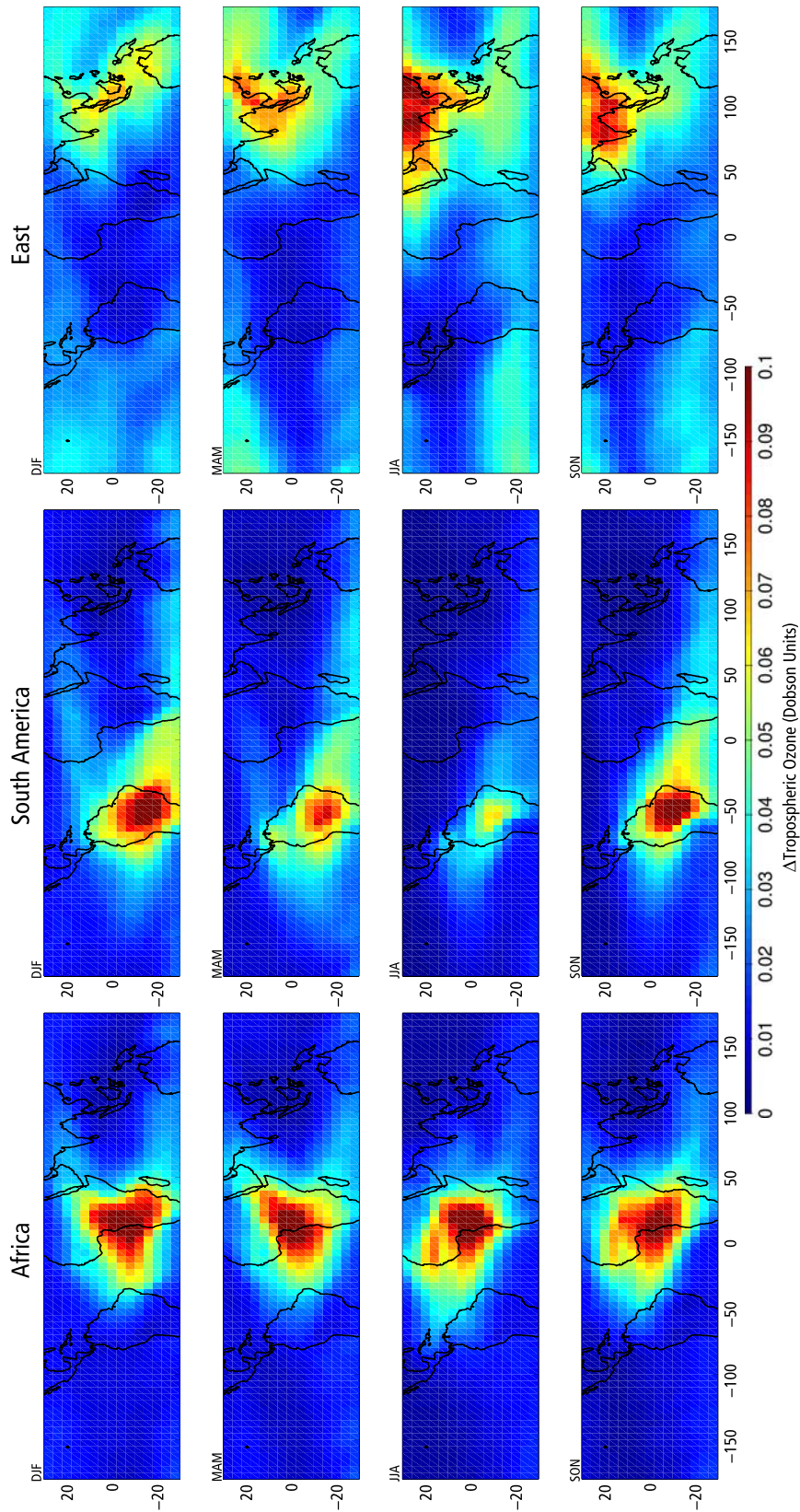


Figure 3. Seasonal sensitivity of the tropospheric ozone column to NO_x source, by decreasing each source by 1%. The tropospheric ozone column difference (DU) between the standard and sensitivity simulation is represented for (left) African sources, (middle) South American sources, and (right) eastern sources. Eastern sources are for India, South East Asia, and Oceania.

any of these sources does not exceed 2 DU (0.015 DU). A similarly weak influence is observed for the South Atlantic O₃ burden, with source sensitivity smaller than for lightning by factors of 4 for biomass burning and for soils, and 7 for fossil fuel (with 1% sensitivity approach). The weak effect of fossil fuel emissions reflects few sources in the surrounding area. The influence of surface sources is primarily close to emissions, and therefore has the largest influence on Africa.

[18] Soils exert considerable influence on TOC over Africa, with maximum effect during JJA through rain-induced emissions during the monsoon, as expected from the analysis of GOME NO₂ columns by Jaeglé *et al.* [2005]. Soil emissions in June are responsible for >8–10 DU (0.06–0.1 DU) over west equatorial Africa (17°W–35°E; 5°–10°N). Some influence of soil emissions on TOC is also found during rainy season over Brazil, during DJF with >2 DU (0.019 DU). Soil NO_x emissions have little seasonal variation in ozone over the South Atlantic.

[19] The contribution of biomass burning emissions to TOC is closely related to satellite observations of fires from the Along-Track Scanning Radiometer [Arino *et al.*, 2005] ($r^2 > 0.7$) reflecting the local effect of this source [Valks *et al.*, 2003]. The influence on O₃ is primarily in the burning hemisphere, in the Northern Hemisphere in DJF–MAM with >2.1 DU (0.010–0.015 DU), and the southern one in JJA–SON. This influence is predominantly on lower tropospheric O₃ [Savauge *et al.*, 2005]. Over the South Atlantic, biomass-burning emissions have a maximum influence of 11% during SON, and minimum of 7% during MAM. Biomass burning exhibits a pronounced nonlinearity due to its concentrated emissions, with a weaker influence than soils in case of a small perturbation.

[20] Fossil fuel combustion primarily affects the eastern tropics. Weaker sources are present over Africa and South America, except for particular sites (Rio de Janeiro, Sao Paulo, Gauteng Province).

[21] The influence of stratospheric O₃ on tropical TOC, as calculated from the ozone flux across the tropopause in the tagged ozone simulation, is 1 DU on annual average only half the magnitude of either soils or biomass burning. During DJF and MAM, most of the stratospheric source is over the north Atlantic and Africa, as noted by Cammas *et al.* [1998], in relation with the seasonal variability of the subtropical jet. In JJA most of the stratospheric contribution is in the Southern Hemisphere. Over the South Atlantic, the stratospheric source is negligible, contributing to 5% of the ozone burden on annual average.

[22] In the absence of all tropical NO_x sources, an O₃ background of 28% remains, reflecting transport and chemical production from extratropical sources. The sum of O₃ produced from all NO_x sources, stratosphere-troposphere exchange and the O₃ background explains only 90% of the ozone burden because of ozone production nonlinearity. However, the seasonal influence and strength of the different sources is mostly independent of the O₃ chemistry nonlinearity (except for surface sources).

3.1.2. Sensitivity to Regions

[23] Numerous studies have investigated the influence on ozone over the South Atlantic from regional emissions from Africa [Jonquière *et al.*, 1998; Krishnamurti *et al.*, 1996; Meyer-Arnek *et al.*, 2005; Edwards *et al.*, 2003], from South America [Pickering *et al.*, 1996; Jonquière and Marengo,

1998; Andreae *et al.*, 2001], from India [Lelieveld *et al.*, 2001] and how they can be redistributed over the Atlantic [Singh *et al.*, 1996; Thompson *et al.*, 1996; Chatfield *et al.*, 2004]. In the following we investigate in a consistent fashion the role of regional emissions and transport on ozone in the tropics and particularly the Atlantic.

[24] Figure 3 shows seasonal influences on tropical TOC from total African, South American, and Eastern (India, South-East Asia, Australia) emissions. These regions account, respectively, for 7.3 Tg N/yr (42%), 4.4 Tg N/yr (25%), and 5.5 Tg N/yr (33%) of tropical NO_x sources. All the regions and associated sources exhibit primarily local influence, close to the region where they are emitted. As a result, the South Atlantic O₃ maximum is influenced more by adjacent continents than by long range transport.

[25] Eastern sources exert the largest influence on tropical TOC, with >7 DU (0.025 DU) annual average, reflecting the divergent circulation. The maximum local effect is found during JJA and SON (>6.8 DU), but also on a broader area than sources situated over Africa and South America. Indeed, there is influence throughout the Pacific and Indian Oceans, in JJA and SON. Eastern emissions generally have little influence on the African continent. However, in Figure 3 a particularly large influence upon North Africa, close to Egypt, occurs through long-range transport in JJA, with >6 DU (0.037 DU). The upper level anticyclone associated with the Indian monsoon [Hoskins and Rodwell, 1995; Rodwell and Hoskins, 2001] merged with the Tropical Easterly Jet, has maximum amplitude during JJA [Hastenrath, 1985] and allows advection of Eastern source emissions to North Africa and the Atlantic. Eastern sources account for >11% of the South Atlantic ozone burden, with a maximum influence in JJA. Convective outflow is advected to the northern part of South Atlantic, where Hadley cells can ensure southernward and slow redistribution.

[26] Sources situated over Africa and South America exhibit maximum influence on the regional TOC. African sources account for >5 DU (0.020 DU) and South America for >4 DU (0.018 DU) of the entire tropical troposphere, on annual average, driven by the higher emissions over the African continent. Both regions have high influence in DJF when there is substantial lightning activity over South America, as well as lightning and biomass burning [Marufu *et al.*, 2000] over Africa. Africa, north of the equator, is the region most influenced by African sources, with >10 DU (0.061 DU) on annual average, versus >8 DU (0.057 DU) for Southern Africa. These regional sources account for most of the South Atlantic ozone burden. African emissions are responsible for >30% of the South Atlantic ozone burden, and South American emissions for >18% on annual average. These regions have more similar influence on the South Atlantic ozone burden with the 1% sensitivity simulation. African emissions have a greater effect in DJF with a contribution of more than 33%, largely due to lightning rather than north African biomass burning emissions. Similar conclusions can be made for the maximum influence of South America in SON of >22%, due to substantial lightning activity.

[27] The seasonal maximum of the zonal wave-one in SON reflects the confluence of both South American and African sources, with >22 and >28%, respectively, as well as long-range transport from eastern sources contributing >11% of the burden. In contrast, during the MAM minimum

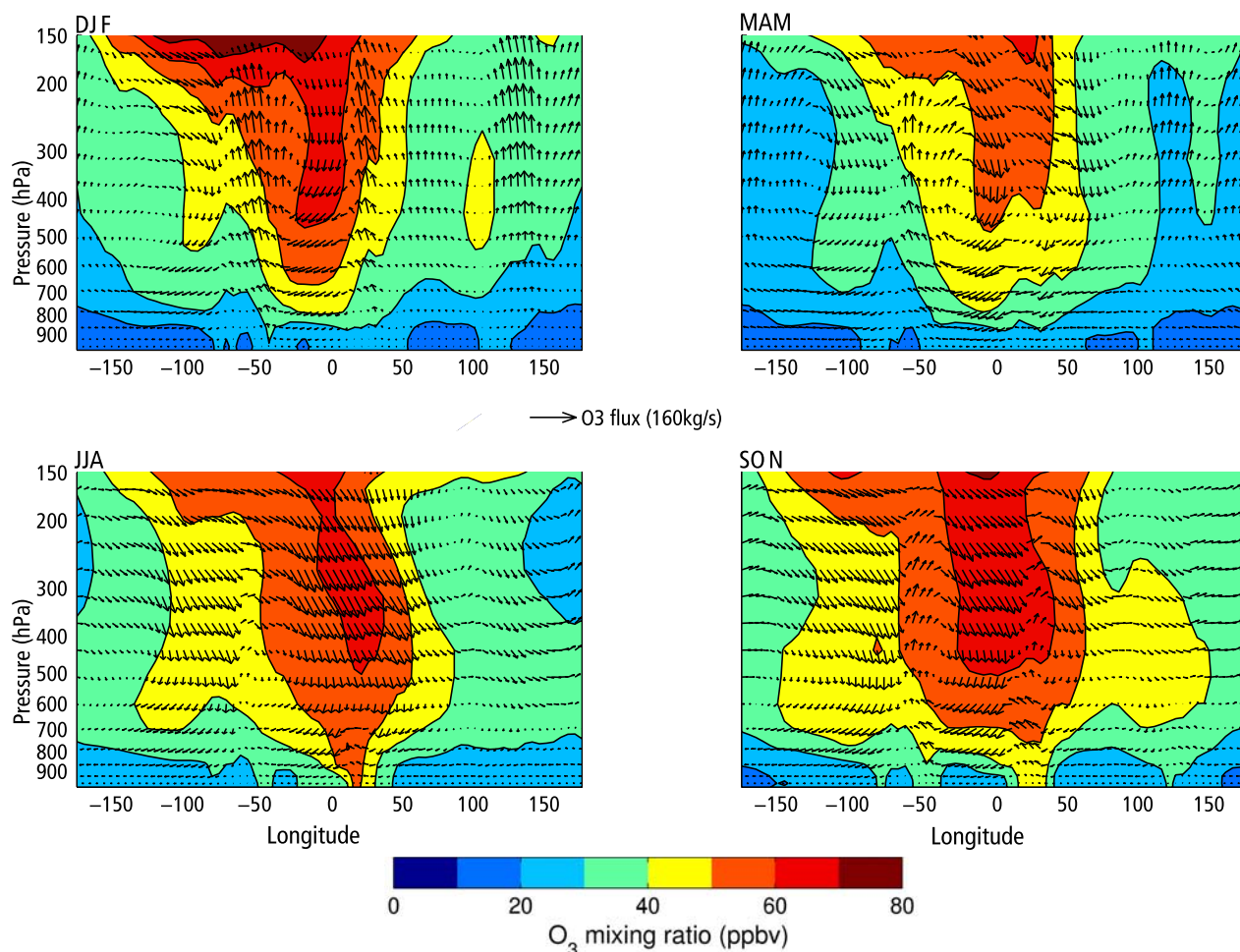


Figure 4. Vertical cross-section of the seasonal O₃ mixing ratios (ppbv) averaged over latitude, between 24°S and 0°. Arrows represent the zonal and vertical components of the O₃ flux in the cross section. The ozone flux is calculated for each grid point using the vertical and horizontal wind components. The vertical component has been scaled by a factor of 8 for better visibility.

of O₃ over the South Atlantic, Africa has a maximum influence, whereas South America and the East have minimum influence.

3.2. Dynamical Processes Affecting the Atlantic Ozone Maximum

[28] Lightning is the dominant “fuel” of the zonal wave-one with a maximum influence on O₃ over the South Atlantic as discussed in section 3.1, and apparent from Figures 1 and 2. It is noteworthy that lightning is absent over this region of the South Atlantic Anticyclone. This region exhibits persistent convergence and subsidence, allowing O₃ production and accumulation during downward transport [Newell, 1979; Krishnamurti et al., 1993; Jacob et al., 1996; Moxim and Levy, 2000; Wang et al., 2006]. The subtropical South Atlantic high, a quasi stationary anticyclone [Hastenrath, 1985], and energy balance drive the radiative subsidence over that region, through convergent flow at higher altitude, representing the “engine” of O₃ buildup over the South Atlantic. However, different flow regimes occur close to the equator and near the subtropics. For that reason in this section we also discuss the extratropical southern Atlantic (32°S–24°S).

[29] Figure 4 provides insight into the processes affecting the southern tropics and the South Atlantic O₃ maximum in particular. It shows the simulated zonal cross-section of the seasonal O₃ mixing ratio between 0° and 24°S. This simulation reproduces the year-round zonal wave-one pattern centered near 40°E–60°W, with maximum amplitude in SON and minimum in MAM, as inferred from in situ measurements [Thompson et al., 2003; Sauvage et al., 2006, 2007] and SAGE II data [Wang et al., 2006]. The extension of the wave one into the lower troposphere resulted from the improved African emissions calculated from satellite observations as implemented by Sauvage et al. [2007]. In contrast, the southern subtropics (not shown) exhibit a very weak wave one pattern.

[30] The ozone fluxes in Figure 4 clearly show the strong subsident feature of the O₃ flux for all seasons over the South Atlantic, allowing downwelling of air masses loaded with enhanced O₃ and its precursors. It supports the previous discussion of a preponderance of regional processes in controlling the O₃ column enhancement. The O₃ maximum is bounded by deep convection as illustrated by O₃ fluxes above South America (70°–35°W) and above Africa (20°–40°E). Over those regions the O₃ concentration is

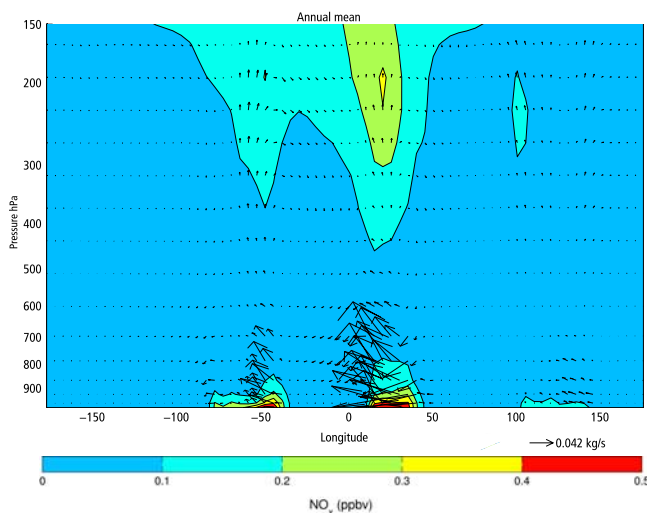


Figure 5. Vertical cross-section of the annual NO_x mixing ratios (ppbv) averaged over latitude, between 24°S and 0°. Arrows represent the zonal and vertical components of the NO_x flux in the cross section. The vertical component has been scaled by a factor of 8 for clarity.

lower than in the maximum of the zonal wave-one, and with strong vertical O₃ gradient. In the southern subtropics, subsidence is weak over the South Atlantic as this region is on the southern edge of the convergence area of the South Atlantic high. Our simulation of background O₃ exhibits a wave one pattern, in agreement with the passive tracer experiment of *Krishnamurti et al.* [1996]. However, the wave one pattern depicted with the simulation of background O₃ exhibits considerably weaker O₃ burden, indicating that the photochemical production is responsible for its magnitude as discussed below.

[31] Figure 5 shows annual mean NO_x fluxes. As described by *Jacob et al.* [1996], deep convection over continents injects NO_x from surface sources and lightning into the upper troposphere, driving O₃ production. The uplift of surface sources also exists in the subtropics (not shown), with weaker lightning influence compared to the tropics. O₃ increases during upper tropospheric zonal advection and descent over oceans. Destruction in the lower troposphere occurs with

high humidity and low NO_x concentrations. Figures 4 and 5 confirm this scheme, highlighting the regional convection over continents, the zonal advection as part of Walker circulation, and the radiative subsidence over the South Atlantic anticyclone area, as major contributors to the O₃ enhancement.

[32] Table 2 quantifies the regional O₃ budget over the tropical South Atlantic. Chemical production and loss are the dominant terms, reflecting the convective injection of NO_x into the free troposphere.

[33] Table 3 contains the seasonal contribution of each source to the NO_x burden over the tropical South Atlantic. Lightning is responsible for more NO_x in the tropical South Atlantic when the ITCZ is further south, during DJF-MAM, with 35 pptv influence. Lightning supplies 59% of the NO_x in the tropical South Atlantic, nearly three times more than all surface sources combined. For comparison, *Moxim and Levy* [2000] found 49% of tropospheric NO_x transported into that region comes from lightning. Our higher lightning ratios, (6 Tg N/yr versus 4 Tg N/yr), is part of that difference. It is interesting that in the subtropical part of the Atlantic where a very weak wave one pattern is found, lightning and total surface sources make similar contributions to NO_x (28 and 23 pptv, respectively).

[34] Figure 6 shows the annual (top) and seasonal (bottom, SON) tropical O₃ mixing ratio attributed to lightning, and to all surface sources. Lightning has the largest influence on the southern tropics. A particularly large lightning contribution to the O₃ mixing ratio appears at the top of the ascending branches (near 150–200 hPa) and in the middle and upper troposphere of the tropical Atlantic downwind of lightning emissions. The maximum influence of surface sources on O₃ appears in the lower troposphere above continents, mostly when biomass burning is active. The role of surface sources in the O₃ enhancement is much less than lightning and exhibits more seasonal variation. In the southern subtropics lightning still has the largest influence, but is somewhat weaker.

[35] Figure 7 highlights zonal transport through a meridional-vertical cross-section averaged over 35°W–10°E where the overturning of the Hadley cells is clearly visible. Strong convection occurs over northern Africa (5°N–10°N) injecting NO_x into the upper troposphere. This is followed by slow

Table 2. Tropospheric Ozone Budget of the South Atlantic^a

O ₃ budget parameters	Values, Tg O ₃ yr ⁻¹
Chemical Production ^b	151
Chemical Loss ^b	160
Deposition	6
Upward Influx	-15
Net Zonal Flux ^c	-24
Net Meridional Flux ^d	9

^aThe tropical South Atlantic is defined over 35°W–10°E/24°S–0°, from the surface to the tropopause.

^bThe chemical production and loss are calculated for odd oxygen family with $O_x = O_3 + O + NO_2 + 2NO_3 + PANs + HNO_3 + HNO_4$, to avoid accounting for rapid cycling of ozone with short-lived species that have little implication in the budget.

^cThe net zonal flux is calculated as the difference of the integrated flux for the eastern and the western edges of the South Atlantic.

^dThe net meridional flux is calculated as the difference of the integrated flux for the northern and the southern edges of the South Atlantic.

Table 3. Tropical Tropospheric OH and Tropical South Atlantic Tropospheric NO_x, and the Contribution From the Different Sources

Season	NO _x (pptv)		
	NO _x	NO _{x,light}	NO _{x,surf}
DJF	58	36	9
MAM	50	34	6
JJA	52	27	13
SON	56	30	13
ANNUAL	54	32	11
Season	OH (10 ⁶ molec cm ⁻³ s ⁻¹)		
	OH	OH _{light}	OH _{surf}
DJF	1.52	0.62	0.36
MAM	1.54	0.63	0.32
JJA	1.46	0.57	0.41
SON	1.55	0.55	0.50
ANNUAL	1.51	0.59	0.40

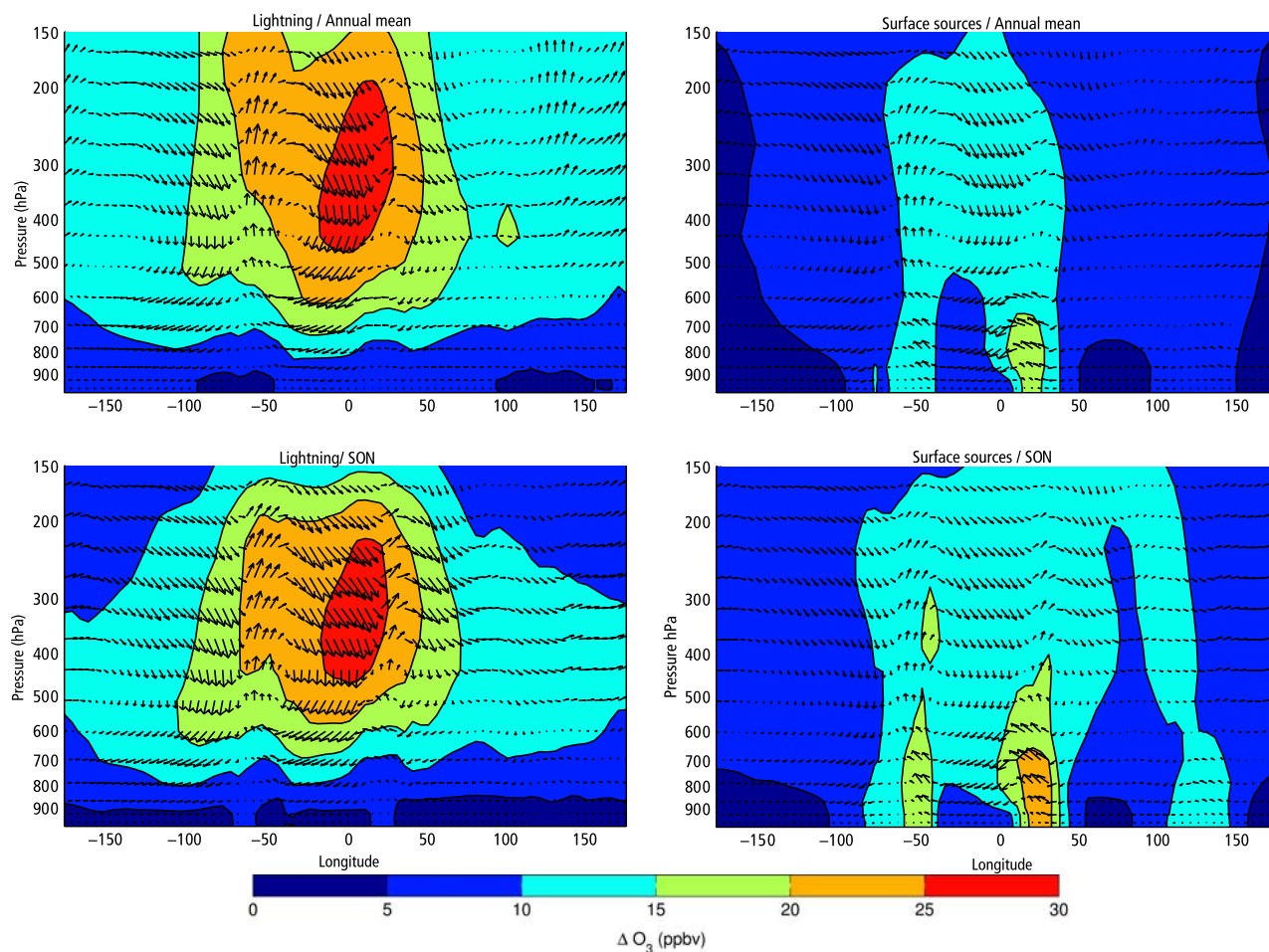


Figure 6. Vertical cross-section of the contribution of lightning (left), and surface sources (right), to the O₃ mixing ratios (ppbv) averaged over latitude, between 24°S and 0°. Arrows represent the meridional and vertical components of the source contribution to the O₃ flux. Top panels are annual mean and bottom panels are for SON.

poleward transport and strong subsidence over the South Atlantic. In contrast, to the North Atlantic, the South Atlantic can be supplied by surface sources in DJF (not shown) through a similar circulation depicted in Figure 7, confirming conclusions of *Chatfield et al.* [2004] and *Sauvage et al.* [2006] for the remaining O₃ paradox over the Atlantic. During DJF and SON the position of the ITCZ facilitates efficient redistribution of fire products into the southern middle troposphere.

[36] To summarize the South Atlantic O₃ maximum is driven by persistent radiative subsidence over the South Atlantic anticyclone area, sustained by injection of NO_x by deep convection over South America and Africa. O₃ is produced during transport converging into that area in both meridional and zonal transport pathways. If various sources play the role of “fuel” for this dynamical and photochemical “engine”, lightning emissions are the most important source.

[37] In the next section we examine why lightning has the most influence despite a source strength similar to biomass burning or soils.

3.3. Ozone Production Efficiency Over the Tropics

[38] The ozone production efficiency (OPE), the number of O₃ molecules produced per NO_x molecule consumed [*Liu*

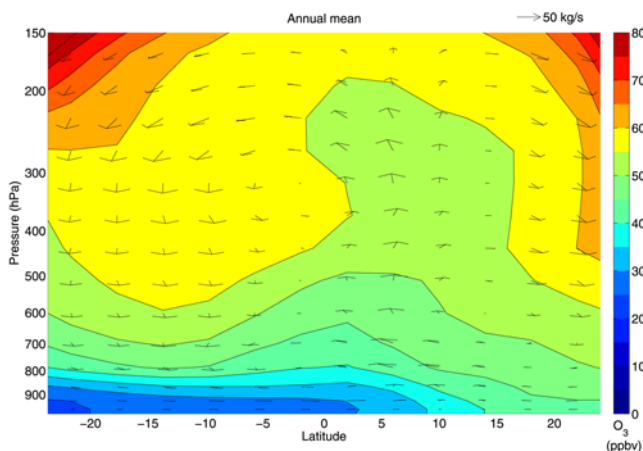


Figure 7. Vertical cross-section of the annual O₃ mixing ratios (ppbv) averaged over longitude, between 35°W and 10°E. Arrows represents the meridional and vertical components of the O₃ flux in the cross section. The vertical component has been scaled by a factor of 8 for visibility.

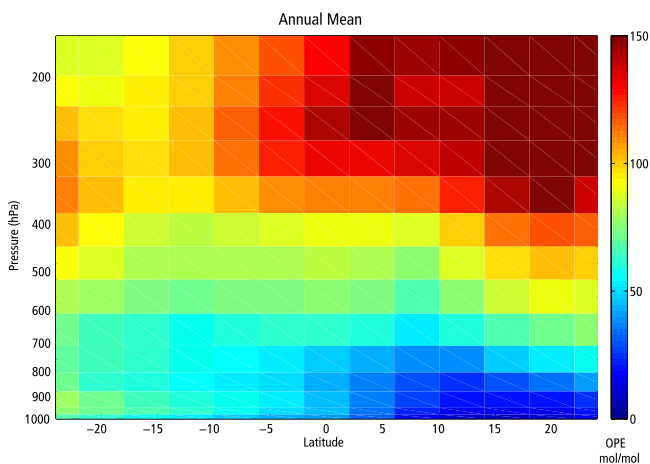


Figure 8. Annual mean Ozone Production Efficiency (OPE) represented in meridional (24°S–24°N) cross-section over the South Atlantic O₃ maximum between 35°W and 10°E.

et al., 1987], provides a useful metric to understand tropical tropospheric O₃. We calculate OPE in the model using HNO₃ production as a proxy for NO_x loss. The OPE provides a measure of the nonlinear dependence of net O₃ production on NO_x.

[39] Figure 8 shows the OPE along vertical cross-sections annually averaged over the tropical South Atlantic O₃ maximum meridional band (35°W–10°E). OPE generally increases with altitude due to intense radiation, low humidity, and relatively low NO_x concentrations. In the middle and upper troposphere the OPE is maximum, from 100 to more than 150 mol/mol. The upper tropospheric OPE minimum follows the ITCZ where humidity and NO_x concentrations are higher. At these altitudes the seasonal OPE (not shown) is much higher in the opposite hemisphere of convection. However, there is persistent high OPE in the tropical South Atlantic region with 98 mol/mol on annual average. As a result O₃ chemical production dominates the O₃ source of the South Atlantic (Table 2). Particularly low OPE appears in the lower troposphere north of 5°N during DJF and MAM because of the high NO_x concentrations from biomass burning over West Equatorial Africa.

[40] We assess the OPE of the different sources by archiving the production of O_x (PO_x) and HNO₃ (PHNO₃) in sensitivity simulations with perturbations to each source:

$$\text{OPE}_{\text{source}} = \frac{(\text{PO}_x)_{\text{standard}} - (\text{PO}_x)_{\Delta_{\text{source}}}}{(\text{PHNO}_3)_{\text{standard}} - (\text{PHNO}_3)_{\Delta_{\text{source}}}}$$

Results are depicted in Table 4. Due to chemical non-linearity, the results from this approach can not be directly related to the OPE of 98 mol/mol presented above. Therefore we calculate an OPE for all sources with a sensitivity simulation in which all NO_x sources are reduced by 1%. Then resulting OPE of 12 mol/mol can be directly compared to the individual sources that follow.

[41] Lightning exhibits the highest OPE, 32 mol/mol on annual average, versus 14 mol/mol for biogenic soil emissions, 10 mol/mol for biomass burning and 13 mol/mol for fossil fuel emissions. Soils and fossil fuels have nearly

constant OPE. In contrast, biomass burning has stronger seasonal variation of OPE. The diffuse nature of lightning NO_x emissions and direct emission into the middle and upper troposphere explains its high OPE.

3.4. Oxidizing Capacity of the Tropical Troposphere

[42] Table 3 contains the seasonal mean of tropical tropospheric OH concentrations, and the component of OH concentrations attributable to each NO_x source, as determined by difference between the standard simulation and simulations without a specific source. Our simulated tropical mean OH is 6% higher than that inferred from measurements of related species by *Spivakovsky et al.* [2000]. OH is highly sensitive to lightning NO_x, as noted by *Labrador et al.* [2004] in a global analysis. Indeed, this source contributes to >39% of the 1.5×10^6 molec cm⁻³ simulated over the tropics on annual average. Other NO_x sources account for substantially less OH, >8% for soils, >7% for biomass burning, and >11% for fossil fuel. The sensitivity simulations with a 1% perturbation reveal that the lightning contribution is three times higher than that from soils or fossil fuel, and five times higher than that from biomass burning.

[43] Lightning is the primary NO_x source determining the oxidation capacity of the troposphere. Future change in the lightning source would have considerable implications for atmospheric oxidation. Indeed, some studies predict an increase in lightning intensity in an altered climate [*Price and Rind*, 1994; *Hauglustaine et al.*, 2005]. Lightning has similar influence on both tropical tropospheric O₃ and OH concentrations. In the upper troposphere high NO and O₃ concentrations related to lightning emission appear, allowing greater production of OH through photolysis of O₃ and, as noted by *Labrador et al.* [2004], through reaction of NO with HO₂.

4. Error Estimates: Sensitivity Studies

[44] In the following we estimate the error of our O₃ budget analysis over the tropics and the South Atlantic, by focusing on the largest uncertainties.

4.1. Lightning Emissions

[45] Recent comparison of GEOS-Chem simulations with in situ and satellite observations yields upper and lower bounds for the lightning source of 8 and 4 Tg N/yr [*Martin et al.*, 2007; *Sauvage et al.*, 2007]. The resultant uncertainty in both the simulation of the South Atlantic O₃ burden and OH is 10%. Even for the lower limit of 4 Tg N/yr, lightning remains the main NO_x source driving the tropical O₃ burden, the South Atlantic maximum, and atmospheric oxidation.

Table 4. Average Tropical Tropospheric OPE (mol mol⁻¹) and for Different Sources (OPE_{Δ_{source}})

Ozone Production Efficiency	Value, mol mol ⁻¹
OPE	98
OPE _{Δ_{all}} ^a	12
OPE _{Δ_{light}}	32
OPE _{Δ_{soils}}	14
OPE _{Δ_{bb}}	10
OPE _{Δ_{fossil fuel}}	13

^aOPE_{Δ_{all}} is calculated by perturbing all NO_x sources by 1%.

[46] Another uncertainty arises from the vertical distribution of lightning emissions. Recent studies provide evidence that the NO_x production efficiency of intracloud (IC) flashes could be the same order of magnitude as cloud to ground (CG) flashes [DeCaria *et al.*, 2000; Fehr *et al.*, 2004]. A sensitivity study using an IC/CG ratio of 0.7 instead of 0.1 increases the ozone burden by 3.5% (0.32 Tg) on annual average. The influence on OH concentrations is negligible, around 2%.

4.2. Dynamics

[47] We also examined the sensitivity of our results to a different dynamical scheme (GEOS-3). The resultant annual mean O₃ burden over the South Atlantic decreases by 4%, with a corresponding decrease in tropical mean OH of 14%.

[48] There are three primary differences between GEOS-3 and GEOS-4. The cloud optical depth is higher in GEOS-3 than in GEOS-4 [Liu *et al.*, 2006; Wu *et al.*, 2007], reducing photolysis frequencies in GEOS-3. The GEOS-3 convective parameterization has a weaker deep outflow mode [Folkins *et al.*, 2006] and a higher altitude of convective detrainment. These differences yield O₃ variation of 10–20% in the middle and upper troposphere [Sauvage *et al.*, 2007] and a 10% variation on the production of O_x [Wu *et al.*, 2007]. Cloud radiative properties and convective transport are the major sources of uncertainty in the oxidizing capacity of the tropical troposphere.

4.3. Surface Sources

[49] Here we assess how uncertainties in surface sources and heterogeneous processes affect the O₃ budget over the Atlantic. As discussed in section 2, Sauvage *et al.* [2007] applied GOME observations of tropospheric NO₂ columns to provide top-down constraints on soils and biomass burning NO_x emissions, increasing global NO_x emissions by 2.6 Tg N/yr for soils and 1.1 Tg N/yr for biomass burning. As a result the South Atlantic O₃ burden increased by only 3%. According to Holland *et al.* [1999] NO_x sources are within 3–13 Tg N/yr for biomass burning and 1–21 Tg N/yr for soils. Even for emissions at the upper limit for both surface sources, and using the lower limit for lightning, lightning remains the major contributor to the South Atlantic O₃ burden and OH.

[50] We conduct a sensitivity simulation with the MEGAN inventory [Guenther *et al.*, 2006] to assess the error estimated on the contribution of biogenic volatile organic compounds. Isoprene emissions increase by 50%, to 600 Tg C/yr. However, the influence on the ozone burden of the South Atlantic maximum is negligible ($\pm 1.5\%$). Varying VOC emissions from biomass burning by a factor of 8 is also negligible, introducing a $\pm 2\%$ change in the South Atlantic O₃ burden.

4.4. Heterogeneous Chemistry

[51] Uncertainty in heterogeneous processes also has little effect on the tropical Atlantic O₃ burden. The two most uncertain processes examined here are uptake of HO₂ and HNO₃. Removing these processes changes the South Atlantic ozone burden by $\pm 2.5\%$.

4.5. Stratospheric Flux

[52] Upper and lower limits on the stratospheric O₃ flux are 680 Tg O₃/yr to 400 Tg O₃/yr [Olsen *et al.*, 2001;

Hauglustaine *et al.*, 1998]. This implies difference in the O₃ burden over the tropical South Atlantic $< \pm 6\%$.

5. Conclusions and Perspective

[53] In this study we used a state of the art global chemical transport model, using an emission inventory that was constrained with satellite observations and validated over the tropics [Sauvage *et al.*, 2007], to quantify the O₃ budget of the tropics and the South Atlantic ozone maximum. Sensitivity studies were conducted with perturbations on NO_x sources and regions, in which NO_x emissions were decreased by 1 and 100%. This approach allows assessment of ozone production nonlinearity on our conclusions.

[54] Lightning is the dominant contributor to tropical tropospheric ozone, accounting for more than 28% of the tropical tropospheric O₃ burden on annual average, as determined from a sensitivity simulation that excludes this source. The contributions from surface sources are smaller, with $>7\%$ for soils, $>7\%$ for biomass burning, and $>8\%$ for anthropogenic sources on annual average, despite similar source strengths. These values represent lower bounds due to O₃ production nonlinearity. However, the relative contribution of each source is largely independent of chemical nonlinearity, except for competition between surface sources. Stratosphere-troposphere exchange accounts for 5% of tropical tropospheric O₃. The O₃ background accounts for 30% of the tropical tropospheric O₃ burden in the absence of tropical NO_x sources, reflecting photochemical production and transport of NO_x and O₃ from outside the region.

[55] The greater contribution of lightning compared to other sources, despite similar strengths, is explained by the high ozone production efficiency (OPE) of the tropical upper troposphere. Lightning is emitted directly into the middle and upper troposphere, where low OH abundance and high NO/NO₂ ratios enhance OPE. The OPE of tropical lightning emissions (32 mol/mol) is twice that of any surface source.

[56] Lightning is the major contributor of the oxidizing capacity of the tropical troposphere, accounting for $>39\%$ of OH concentrations, much higher than all surface emissions combined ($>26\%$). Future evolution of the oxidizing capacity of the atmosphere will be closely coupled to lightning intensity.

[57] Over the South Atlantic O₃ maximum, lightning NO_x accounts for $>37\%$ of the O₃ burden in annual average, against $>7\%$ for soil NO_x, $>9\%$ for biomass burning NO_x, and $>5\%$ for fossil fuel NO_x sources. Stratospheric O₃ has little role, 5%. An O₃ background of 28% remains in the absence of all tropical NO_x emissions.

[58] The South Atlantic maximum is driven by dynamical processes, with permanent radiative subsidence over the quasi-stationary South Atlantic anticyclone area. This area acts as a convergence zone of meridional and zonal fluxes. Chemical production and loss are the dominant terms.

[59] The main geographic contribution appears to be regional with the two bounded continents. Convection in the ITCZ over these adjacent continents, Africa and South America (with respective NO_x sources representing 30 and 18% of the South Atlantic ozone burden), allows uplifting of “fuel”, i.e., O₃ precursors emitted by surface sources and most of all of lightning NO_x emissions. In the middle and

upper troposphere with low humidity rates and high UV radiation, O₃ can buildup and be pumped down by radiative subsidence, giving rise to the zonal wave-one pattern. Long-range transport has influence mostly in JJA, when the tropical easterly jet is maximum, accounting for 15% of the O₃ burden, against 11% on annual average.

[60] The maximum O₃ intensity over the South Atlantic in SON is explained by a combination of lightning and soil NO_x sources, but also a maximum intensity of fires over Africa and South America, as well as a maximum in long range transport.

[61] Despite nonlinearity in O₃ production, the previous statements are similar for 1 or 100% reduction of sources, in terms of seasonality and intensity. The primary difference comes in the relative contribution of soils versus biomass burning or fossil fuels, which is shifted depending of the configuration, and reflected in the regional contributions for South America and Africa.

[62] Finally, we assess uncertainties of our study by conducting sensitivity simulations of the lightning source strength, lightning vertical distribution, dynamical scheme, surface emissions, heterogeneous chemistry, and stratospheric flux. Uncertainty in the ozone burden over the Tropical Atlantic is driven by lightning emissions and dynamical fields (convective parameterization and clouds). Lightning and dynamical processes account for most of uncertainty in the oxidizing power of the tropical troposphere. These uncertainties do not change our primary conclusions, that lightning is the dominant NO_x source affecting the tropical O₃ burden.

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