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Ozone production and transport over the Amazon Basin during the dry-to-wet and wet-to-dry transition seasons

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Abstract. The Regional Carbon Balance in Amazonia (BARCA) campaign provided the first Amazon Basin-wide aircraft measurements of ozone (O₃) during both the dry-towet (November and December 2008) and wet-to-dry (May 2009) transition seasons. Extremely low background values (<20 ppb) were observed to the west and north of Manaus in both seasons and in all regions during the wet-to-dry transition. On the other hand, elevated O₃ levels (40–60 ppb) were seen during the dry-to-wet transition to the east and south of Manaus, where biomass burning emissions of O₃ precursors were present. Chemistry simulations with the CCATT-BRAMS and WRF-Chem models are within the error bars of the observed O_3 profiles in the boundary layer (0–3 km a.s.l.) in polluted conditions. However, the models overestimate O₃ in the boundary layer in clean conditions, despite lacking the predominant NO source from soil. In addition, O₃ simulated by the models was either within the error bars or lower than BARCA observations in mid-levels (3-5 km a.s.l.), and lower than total tropospheric O₃ retrieved from the OMI/MLS instruments, which is primarily comprised of middle troposphere O₃ and thus reflects long-range transport processes. Therefore, the models do a relatively poor job of representing the free troposphere-boundary layer gradient in O3 compared with aircraft and satellite observations, which could be due to missing long-range and convective transport of O₃ at mid-levels. Additional simulations with WRF-Chem showed that the model O_3 production is very sensitive to both the O_3 deposition velocities and the NO_x emissions, which were both about one-half of observed values. These results indicate the necessity of more realistic model representations of emissions, deposition, and convective processes for accurate monitoring and prediction of increases in O_3 production in the Amazon Basin as the regional population grows.

1 Introduction

In the Amazon Basin, trace gases from biomass-burning, urban, and biogenic emissions are important sources of ozone precursors, which are efficiently transported by intense convective activity to the upper troposphere, where they can be dispersed over long distances by regional and global circulations. Additionally, convective overshooting may inject heat, moisture, and trace gases into the tropical tropopause layer, impacting stratospheric ozone and other aspects of global climate (Fueglistaler et al., 2009). In the dry-to-wet transition season, regional smoke and haze plumes from biomass burning are observed (Longo et al., 2009). On the other hand, in the wet-to-dry transition season, biogenic emission of volatile organic compounds (VOCs), particularly from the Amazon rainforest, may maintain the atmospheric oxidative capacity for generating ozone and other photochemical pollutants (Lelieveld et al., 2008).

The Amazon Basin continues to rapidly urbanize, and anthropogenic emissions of O₃ precursors are also expected to grow. Emissions from cities in the tropics may have a larger impact on the upper troposphere due to high solar radiation levels and intense convective transport (Gallardo et al., 2010). In the upper troposphere, O₃ acts as a greenhouse gas, increasing surface radiative forcing (Smithson, 2002). Inhalation of elevated levels of ozone can irritate the lungs, aggravate asthma, and cause emphysema, bronchitis, and premature death (Schwela, 2000). High ozone concentrations can also decrease photosynthesis in plants and damage leaf tissue, harming wild ecosystems and reducing crop productivity (Reich and Amundson, 1985). Thus, an improved understanding/quantification of O3 temporal and spatial variability in the tropical rainforest environment is important for projecting future impacts of land use and climate change in the Amazon Basin and other tropical rainforest regions worldwide on their expanding human populations and significant biodiversity.

Analyses of satellite, aircraft, and ground-based observations of O_3 over Amazonia since the 1980s have demonstrated the influence of long-range transport of African biomass burning and Northern Hemisphere inputs, local fire sources, NO soil and biogenic VOC emissions, and convective transport on spatial and seasonal variability in O_3 . In particular, data from the ABLE-2B aircraft and ground campaign during the 1987 wet-to-dry transition season and the BARCA observations offer the opportunity to compare the regional O_3 distribution across decades.

Previous analyses of satellite ozone data have noted early year O₃ maximums in the tropical Southern Hemisphere primarily associated with cross-Atlantic transport of biomass burning emissions from Africa (Fishman and Larson, 1987; Thompson et al., 1996), Northern Hemisphere fires, and lightning NO_x (Edwards et al., 2003). In the Amazon region, ground-based and aircraft campaigns (e.g., Crutzen et al., 1985; Kirchhoff et al., 1990; Browell et al., 1996; Kaufman et al., 1998; Longo et al., 1999; Andreae et al., 2001, 2002, 2004; Zhou et al., 2002; Cordova et al., 2003; Rummel et al., 2007; Kuhn et al., 2010; Martin et al., 2010; Toon et al., 2010) have observed daytime background O₃ levels of 10–20 ppb, decreasing to very low values (circa 5 ppb) at night due to O₃ deposition to the forest. However, nighttime values can be increased up to 30 ppb by convective downdrafts (Betts et al., 2002; Cordova et al., 2003). Elevated levels of 60-80 ppb are found due to production from regional fire emissions and recirculated urban pollution from SE Brazil, as well as deep convective transport of boundary layer air to the middle and upper troposphere.

Thus, satellite observations enable the attribution of tropical O_3 maxima to biomass burning and lightning NO_x sources, while ground-based measurements allow the identification of key surface processes in the Amazon Basin affecting O₃ amounts. These processes include O₃ production from soil NO_x emissions and removal via dry deposition to the forest canopy. Aircraft campaigns complete the suite of observations, allowing the examination of convective lofting of surface emissions, with biomass burning emissions being of particular importance on the regional scale. In situ data on cloud properties and chemical species, as well as observations of land use changes, boundary layer dynamics, and larger-scale cloud-aerosol interactions, are scant in this region. Therefore, models are essential tools for understanding the interactions of physical and chemical processes that contribute to O₃ formation, as well as monitoring and predicting atmospheric chemistry composition, weather, and climate at local, regional, and global scales. In turn, the observations help constrain uncertainties in the model representations of parameterized convection, turbulence, land surface, and other subgrid scale processes that affect the simulated transport and chemical transformation of the atmospheric composition (Beck et al., 2013).

Motivated by the impact of O_3 in the Amazon Basin on human and ecosystem health and global climate, we collected aircraft observations of O_3 during BARCA and conducted regional chemistry simulations in order to answer the following scientific questions: how does O_3 vary spatially, seasonally, and across decades over the Amazon Basin? What are the sources and sinks of O_3 in this region? How well can stateof-the-art regional chemistry models reproduce O_3 distributions over the Amazon Basin?

The structure of this paper is as follows. In Sect. 2, the measurements taken during the BARCA aircraft campaign are presented, followed by the meteorological conditions and emission regimes during the two phases of the campaign. The ABLE-2 campaigns from the 1980s are also described in this section. Sections 3.1-3.3 detail the aircraft observations, the setups of the CCATT-BRAMS and WRF-Chem simulations, and the ground-based and remote sensing observations used in the analysis. In Sect. 4.1, the O₃ aircraft observations are presented, followed by the analysis of observed and modeled transition season meteorology in Sect. 4.2, and the findings from the O₃ simulations and process studies in Sect. 4.3. Final discussions and conclusions are found in Sect. 5.

2 BARCA aircraft campaigns

The Regional Carbon Balance in Amazonia (BARCA) Large-Scale Biosphere-Atmosphere (LBA) experiment was an aircraft campaign was based in Manaus and conducted during the dry-to-wet (November and December 2008) and wet-to-dry (May 2009) transition seasons. BARCA was the first flight campaign to sample O_3 and other trace gases on a regional scale in both transition seasons. It offers a unique opportunity, alongside satellite observations and modeling studies, to understand the regional ozone distribution in the Amazon under different meteorological and emission regimes.

The BARCA flights were conducted with the EMB 110 Bandeirante aircraft of the Brazilian National Institute for Space Research (INPE). In situ measurements were made of carbon dioxide (CO₂), carbon monoxide (CO), methane (CH_4) , ozone (O_3) , and aerosol number concentration and optical properties. Flask samples were collected to determine CO₂, CH₄, sulfur hexafluoride (SF₆), CO, nitrous oxide (N_2O) , hydrogen, and the oxygen : nitrogen ratio (O_2 / N_2) . The flights consisted of quasi-Lagrangian measurements, which attempt to sample an air parcel at multiple locations along its path in order to constrain regional and basin-wide fluxes of these species. The aircraft had a ceiling of 4500 m, and flights usually consisted of ascending and descending vertical profiles separated by short (5-30 min) horizontal legs. A detailed description of the aircraft measurements can be found in Andreae et al. (2012). Fig. 1 shows a map of the flight tracks from BARCA A and B. Both experiment periods included flights to the north, south, and east of Manaus, as well as local flights near Manaus. Only BARCA A included flights to the west of Manaus, because intense convective activity in that region during BARCA B precluded flying. During BARCA B, fire activity was low throughout the Amazon region due to heavy precipitation, while during BARCA A, intense fire activity occurred on the northern coast of Brazil and scattered fires were present throughout the southeastern Amazon.

Andreae et al. (2012) summarized the BARCA campaign, meteorological background, carbon monoxide and aerosol observations, and CO results from several regional transport and chemistry models. These included versions of the CCATT-BRAMS and WRF-Chem simulations analyzed in greater detail in this paper. Meteorological analysis showed that during BARCA A, when the Inter-Tropical Convergence Zone (ITCZ) was to the north of the Amazon Basin, inflow to the Amazon was primarily from the Southern Hemisphere. During BARCA B, the ITCZ extended to 20° S and air at low levels was of Northern Hemisphere origin, including some smoke from West African fires. On the other hand, the midtropospheric air was of mixed origin.

The highest CO levels were observed on the flights on 25–27 November in the southeastern Amazon, influenced by regional biomass burning, since maximum values were observed from 1–3 km. These are typical of injection heights of smoke plumes from savanna fires (Freitas et al., 2007). The excess CO from biomass burning was between about 30 and 200 ppb, increasing from north to south across the Basin. According to analysis of tracer simulations, during BARCA A, biomass burning contributed on average about 56 ppb (31 %) to the total CO of around 180 ppb, while the background was 110 ppb (61 %). Biomass burning influence was indicated by CO mixing ratios of up to 300 ppb, condensation nuclei (CN) approaching 10 000 cm⁻³, and a low CN to CO ratio (Δ CN / Δ CO) signifying aged smoke. This influence



Figure 1. Flight tracks during BARCA.

was highest in the southern Amazon from 1 to 3 km. Manaus back trajectories at 500 and 4000 m came from eastern Amazon fires rather than the intense African fires occurring at the same time. During BARCA B, little biomass burning influence was observed. CN counts were $300-500 \text{ cm}^{-3}$ and a CO enhancement of circa 10 ppb above the mixing ratios in air entering the basin from the Atlantic was seen. Small boundary layer enhancements were attributed to a source from the oxidation of biogenic VOCs (Andreae et al., 2012).

Andreae et al. (2012) also showed simulated vertical CO profiles from CCATT-BRAMS and WRF-Chem simulations, as well as the Stochastic Time Inverted Lagrangian Transport (STILT) model with two different meteorological field inputs and the WRF greenhouse gas module (WRF-GHG). The simulated CO profiles matched mean observed values, but were overly vertical (too low near the surface and too high above 3 km). This suggested that the models had too much convective transport or vertical mixing from the planetary boundary layer (PBL) schemes. However, the probability densities were consistent with observations in the boundary layer, indicating that horizontal dispersion was reasonable. Beck et al. (2013) evaluated different CH₄ wetland emissions schemes and maps using WRF-GHG. They found the best agreement with BARCA CH₄ data for days when convective transport, as evaluated by comparison of upstream Tropical Rainfall Monitoring Mission (TRMM) and WRF precipitation amounts, was well represented in the model. This indicates that proper representation of convective transport in models is essential for prediction of vertical distributions of pollutants in the Amazon Basin.

It is interesting to compare BARCA data to observations from the NASA Amazon Boundary Layer Experiments ABLE campaigns (ABLE-2A and -2B), which took place during the dry season of 1985 and wet-to-dry transition of 1987. During the dry season (July–August 1985), the Amazon Boundary Layer Experiment (ABLE-2A) integrated aircraft, ground-based, and satellite observations to study the processes affecting the chemical composition in mixed layers over Amazonia (Harriss et al., 1988). Jacob and Wofsy (1988) used a photochemical model of the Amazonian boundary layer to study the diurnal cycle of isoprene, NO_v and O₃ during ABLE-2A. They found that photochemical production spurred by NO emissions from soils increased daytime O₃ to about 20 ppb. However, at night, dry deposition to the forest caused O₃ to drop below 5 ppb. Model results were consistent with the NO values of 25-60 ppt observed in the lower boundary layer over central Amazonia (Torres and Buchan, 1988). Isoprene emissions were found to have little effect on O₃ levels, as the oxidation of CO would produce sufficient hydrogen oxides (HO_x) to generate 20 ppb of O₃. However, O₃ production in the model was highly sensitive to NO_x emissions, and downward transport from the free troposphere became the dominant source of O3 in the PBL when NO emissions were decreased below the average value of $44 \pm 14 \,\mu g \,\text{Nm}^{-2} \,\text{h}^{-1}$ NO measured by Kaplan et al. (1988). Lidar observations during ABLE-2A showed highly variable O3 levels, with some small regions with up to 30-40 ppb, attributed to variable NO flux from the canopy (Browell et al., 1988). ABLE-2B was conducted during the wet-to-dry transition season (April-May 1987) (Harriss et al., 1990). Periodic inputs from the Northern Hemisphere were found to be a pollution source over Amazonia, and dry deposition in the region provided a significant sink in the global O₃ budget. As part of ABLE-2, near-continuous O_3 surface measurements (1.5 m above the soil surface) showed daytime maximums of 3.7 ppb inside a forest and 5.7 ppb in a clearing (typical standard deviations of 0.3 ppb). Additionally, tower measurements at the clearing site showed higher O₃ values of 6.7 ppb at 7 m above the soil surface and 6.9 ppb at 15 m above the soil surface (Kirchhoff et al., 1990). Furthermore, 20 ozonesondes launched in the clearing showed typical mixing ratios of 40 ppb from 500 to 300 hPa, with values about 10 ppb lower in the wet than dry season. Andreae et al. (2012) showed that CO mixing ratios were about 10 ppb higher during ABLE-2B than in BARCA B everywhere except the southern region, reflecting the global trend towards decreasing CO emissions since the 1980s, particularly in the Northern Hemisphere. The CO comparison also showed a similar enhancement of 10-20 ppb in the lowest 1 km above the surface, attributed to diffuse biogenic sources, and also indicated that the much higher enhancements during the dry season in BARCA A must be due to anthropogenic or biomass burning inputs. The O₃ comparison is expected to yield information about long-term trends in O₃ production in the Amazon Basin, as well as the relative importance of biogenic, urban, and fire sources.

3 Data and Methods

3.1 BARCA aircraft measurements

During the BARCA campaign, in situ measurements of O_3 were conducted aboard the EMB 110 Bandeirante INPE aircraft using a dual-cell, UV Photometric analyzer (Ozone Analyzer, Model 49i, Thermo Fisher Scientific, United States). During BARCA A, 1 min averages of the original 1 s data were taken, while during BARCA B 1 s data were stored. The detection limit for both campaigns was 1 ppb. The intake for O_3 was forward facing, located 185 mm from the fuselage on the lower fuselage in front of the propellers to minimize effects of turbulence. The inlet lines consisted of stainless steel tubes with a bend radius of 100 mm and an inner diameter of 11.5 mm. The sample air was not heated or dried before measurement, so reported values are molar mixing rations, nmol mol⁻¹, abbreviated "ppb", with respect to ambient humid air (Andreae et al., 2012).

3.2 Model description and simulation setup

Simulations of BARCA A and B were conducted with the Chemistry Coupled Aerosol-Tracer Transport model to the Brazilian developments on the Regional Atmospheric Modeling System (CCATT-BRAMS; Freitas et al., 2009; Longo et al., 2013) and Weather Research and Forecasting with Chemistry (WRF-Chem; Grell et al., 2005) Version 3.4.1 coupled chemistry and meteorology models. The model physics and chemistry options that were used are listed in Table 1. Both models used a two-way nested grid configuration, with a 140 km grid covering Africa and South America (southwest corner: 60° S, 100° W, northeast corner: 20° N, 50° W), to encompass the cross-Atlantic transport of biomass burning emissions from Africa, and a 35 km resolution grid covering most of South America (southwest corner: 35° S 85° W, northeast corner: 15° N, 30° W), as depicted in Fig. 3.

The simulations were initialized on 1 October 2008 00:00 UTC and 1 April 2009 00:00 UTC for BARCA A and B, respectively. Boundary conditions and analysis nudging on the outer domain were given by the NCEP GFS analysis (http://rda.ucar.edu/datasets/ds083.2/) with a 6-hourly time resolution and $1^{\circ} \times 1^{\circ}$ spatial resolution. Chemistry initial and boundary conditions were provided by 6-hourly analyses from the Model of Atmospheric Chemistry at Large Scale (Modélisation de la Chimie Atmosphérique Grande Echelle, MOCAGE) global model (Josse et al., 2004; Teyssèdre et al., 2007) with a T42 (circa 2.8°) spatial resolution. Sea surface temperature was provided by the NOAA Optimum Interpolation (OI) Sea Surface Temperature (SST) V2 (available at http://www.esrl.noaa.gov/psd/data/gridded/ data.noaa.oisst.v2.html) with $1^{\circ} \times 1^{\circ}$ spatial resolution. Soil moisture was initialized with the TRMM-based soil moisture operational product (GPNR) developed by Gevaerd and Freitas (2006).



Figure 2. O₃ observations during (a) BARCA A, clean conditions (west, north, and around Manaus regions), (b) BARCA A, polluted conditions (east and south regions), and (c) BARCA B. The central mark is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers, and outliers are plotted individually as red plus signs. Values are drawn as outliers if their values exceed $q_3 + w(q_3 - q_1)$ or are less than $q_1 - w(q_3 - q_1)$, where q_1 and q_3 are the 25th and 75th percentiles, respectively, and w is the maximum whisker length with the default value of 1.5. For normally distributed data, the whiskers encompass approximately the 2.7 to 99.3 percentiles.

Table 1. CCATT-BRAMS and WRF-Chem physics and chemistry options for the BARCA simulations.

	CCATT-BRAMS	WRF-Chem
Short/longwave radiation	Based on CARMA	RRTMG
Cloud microphysics	Single moment bulk	WSM-5
Deep/shallow convection	Grell and Dévényi (GD)	Grell 3-D
Trace gas chemistry	RACM	RACM
Photolysis	F-TUV	F-TUV
Aerosol scheme	Smoke aerosol	GOCART
Wet deposition	Convective and grid scales	Convective scale only

The PBL parameterization in CCATT-BRAMS is based on Mellor and Yamada (1982), while in WRF-Chem the Mellor-Yamada Nakanishi and Niino 2.5 level PBL scheme (MYNN; Nakanishi and Niino, 2004) was used. In CCATT-BRAMS, shallow and deep convection are parameterized based on the mass-flux approach of Grell and Dévényi (2002). CCATT-BRAMS also uses the turbulent kinetic energy (TKE) from the planetary boundary layer (PBL) scheme to determine if convection will be triggered within a grid cell. In WRF-Chem the Grell 3-D (G3) scheme was used, which includes shallow convection and subsidence spreading of convective



Figure 3. Land surface albedo (fraction) and locations of the coarse (140 km resolution) and nested (35 km resolution) domains for WRF-Chem simulations.

outflow into neighboring grid cells. The Noah land surface model (Koren et al., 1999) was used in WRF-Chem and the Land Ecosystem–Atmosphere Feedback model v.2 (LEAF-2; Walko et al., 2000) was utilized in CCATT-BRAMS. Land use was provided by the United States Geological Survey (USGS) global 1 km vegetation data set, updated with a land cover map for the Brazilian legal Amazon region for use in meteorological models (PROVEG) (Sestini et al., 2003). PROVEG is based on the Landsat Thematic Mapper (TM) images with spatial resolution of 90 m×90 m from the year 2000 and deforestation data from the Amazon Deforestation Monitoring Program (PRODES) for the year 1997. For WRF-Chem, albedo and greenness fraction were calculated offline using the updated vegetation data set, Moderate Resolution Imaging Spectroradiometer (MODIS) Normalized Difference Vegetation Index (NDVI) data from the years 2001–2002, and vegetation parameters from the LEAF-2 land surface model as implemented in CCATT-BRAMS.

Emissions were generated with PREP-CHEM-SRC (Freitas et al., 2011) Version 1.2. Fire emissions were estimated from GOES, AVHRR, and MODIS fire count data, using the Brazilian Biomass Burning Emission Model (3BEM; Longo et al., 2009). Anthropogenic emissions were estimated from the RETRO, GOCART, and EDGAR v4.0 global databases updated with South American inventories (Alonso et al., 2010). Emissions are obtained from RETRO if available for that species (CO, NO_x, chlorinated hydrocarbons, acids, esters, alcohols, ethers, benzene, ketones, methanal, other alkanals, other aromatics, C₂H₂, C₂H₄, C₂H₆, C₃H₆, C₃H₈, C₄H₁₀, C₅H₁₂, C₆H₁₄ plus higher alkanes, other VOCs, toluene, trimethylbenzenes, xylene), then from EDGAR v4.0 (NMVOC, SO₄, CO₂, SF₆, N₂O), otherwise from GOCART (BC, OC, SO₂, DMS) in order to use the most consistent emissions inventory possible. Biogenic emissions were provided by a monthly climatology for the year 2000 produced with the Model of Emissions of Gases and Aerosols from Nature (MEGAN; Guenther et al., 2006). The MEGAN 2000 climatology includes numerous biogenic species (acetaldehyde, formaldehyde, other ketones, acetone, isoprene, propane, methane, propene, ethane, methanol, sesquiterpenes, ethene, monoterpenes, and toluene), but not soil NO emissions. In WRF-Chem, the same Gaussian diurnal cycle with peak at 15:00 UTC (11:00 LT) is applied to both anthropogenic and biogenic emissions, while in CCATT-BRAMS the diurnal cycle of biogenic emissions follows the solar radiation cycle. In both models, the biomass burning daily cycle peaks at 18:00 UTC (15:00 LT).

In both CCATT-BRAMS and WRF-Chem, the Regional Atmospheric Chemistry Mechanism (RACM) was used (Stockwell et al., 1997). In WRF-Chem, the Goddard Chemistry Aerosol Radiation and Transport (GOCART; Chin et al., 2002) aerosol scheme was used with aerosol direct radiative effects. CCATT-BRAMS has a smoke aerosol scheme with intensive optical properties (extinction efficiency, single scattering albedo, and asymmetry parameter) calculated in an offline Mie code based on observations of climatological size distribution and complex refractive index from AERONET sites in the southern Amazon (Rosário, 2011; Rosário et al., 2013).

CCATT-BRAMS includes scavenging of soluble species in the convective scheme following Berge (1993), as described in Freitas et al. (2005), where the wet removal rates are a function of the precipitation rate, liquid water content, and precipitable water. In the cloud microphysics scheme the wet deposition follows Barth et al. (2001), whereby low solubility species partition into the liquid phase according to Henry's Law and high solubility species by diffusionlimited mass transfer. In WRF-Chem, at the convectiveparameterizing scale, a constant fraction of gas and aerosol species in convective updrafts are removed (complete removal for sulfur dioxide (SO₂), sulfate (H₂SO₄), ammonium (NH₃), nitric acid (HNO₃), and sea salt; no removal for hydrophobic organic (OC) and black carbon (BC) and dimethyl sulfide (DMS); and 50% removal for all other aerosol species). On the other hand, no wet scavenging is included for cloud water and precipitation resolved by the microphysics scheme, because this option is not currently available in WRF-Chem for the RACM chemical mechanism. O₃ production in the upper troposphere is affected by the net convective transport of soluble HO_x precursors (including hydrogen peroxide (H2O2), methyl hydroperoxide (CH₃OOH), and formaldehyde (CH₂O)). However, uncertainties remain about the scavenging efficiencies of these and other soluble species by deep convective storms. Simulations of an idealized thunderstorm by several cloud-resolving models yielded varying results for CH₂O, H₂O₂, and HNO₃ in convective outflow due to differing microphysics and assumptions about retention of chemical species during cloud drop freezing (Barth et al., 2007b).

The CCATT-BRAMS simulations employ a lightning NO_x parameterization based on convective cloud top height (Stockwell et al., 1999). In WRF-Chem, lightning production of NO_x was not included, because these parameterizations have not yet been evaluated for the Amazon region. In the tropics and over continents, lightning production is comparable to other sources of NO_x, including biomass burning and soil release, and it is the primary source over oceans (Bond et al., 2002). Since lightning NO_x production peaks in the upper troposphere, it could be an important contributor to ozone production. The impact of wet deposition and lightning NO_x production on O₃ distributions will be more closely examined in future modeling studies of tropospheric chemistry in the Amazon.

For model results evaluation, the mean vertical O_3 profiles for observations, CCATT-BRAMS, and WRF-Chem were calculated for the regions to the west, north, south, east, and around Manaus. Horizontal flight legs were excluded from the analysis to eliminate the influence of plumes in the boundary layer. As the model output has a much coarser spatial and temporal resolution than the aircraft measurements, the model value is interpolated to the observation time and location. To calculate the mean simulated profiles, the four grid points closest in latitude and longitude to each observation were determined at the 2 model hours that bracket the observations. At each of these grid points and hours, vertical profiles were extracted from the model output and then



Figure 4. Monthly mean precipitation $(mm h^{-1})$ on the 35 km resolution domain (dark gray line) for November 2008 from (a) TRMM 3B43, (b) CCATT-BRAMS, and (c) WRF-Chem and for May 2009 from (c) TRMM 3B43, (d) CCATT-BRAMS and (f) WRF-Chem. The subregions for the precipitation comparison are indicated by black lines.



Figure 5. Mean daily cycle of surface incident shortwave radiation from the Manaus AERONET site (solid line, dotted line denotes 1 standard deviation), WRF-Chem (crosses), and CCATT-BRAMS (circles) for the BARCA A period (October–November 2008).

linearly interpolated to the observed GPS height. The four points from each time were averaged, weighting by the inverse distance to the observed longitude and latitude. Finally, the prior and posterior hour values were averaged with appropriate weights. Thus, 16 model points were used with spatial and temporal weights to obtain each model value for comparison to observations. The observed and model time series were then separated into five regions to the west, north, east, and south of Manaus, and in the region of Manaus itself. The mean value and standard deviation were calculated for each region and 500 m vertical bin.

To facilitate comparison of other models with the data presented in Fig. 2, mean profiles from the large regions corresponding to clean (west, north, and around Manaus regions) and polluted (east and south regions) regions during BARCA A and all regions during BARCA B are presented in Fig. 16. From the models, all horizontal grid points falling within the corresponding region's longitude and latitude bounds for each flight day (Table 6) and the closest model output times (12:00–18:00 UTC / 08:00–14:00 LT) were averaged into 500 m vertical bins.

3.3 Satellite and ground-based O₃ and meteorological data

In addition to the in situ O_3 data, the model results were compared with OMI/MLS monthly mean tropospheric ozone mixing ratios and total column ozone (http://acd-ext.gsfc. nasa.gov/Data_services/cloud_slice/#pub) (Ziemke et al., 2006). In this product, the tropospheric values are estimated by subtracting the stratospheric contribution from total column measurements. A cloud-slicing method is used to detect O₃ inside optically thick clouds. This method was able to detect elevated O₃ levels of around 50 ppb in the upper parts of convective clouds over South America and Africa, comparable to background cloud-free levels in the tropics (Ziemke et al., 2009). In this study, the model total tropospheric O_3 column and mean tropospheric O₃ mixing ratio were calculated by summing O₃ mixing ratios, weighted by the model level air density, from the first model level to the level below the tropopause. The tropopause level was determined by the

Table 2. Monthly mean precipitation $(mm h^{-1})$ for TRMM 3B43, CCATT-BRAMS and WRF-Chem models for three regions: the Amazon $(15^{\circ} \text{ S}-10^{\circ} \text{ N}, 50-80^{\circ} \text{ W})$, northeast Brazil $(15^{\circ} \text{ S}-0^{\circ} \text{ N}, 35-50^{\circ} \text{ W})$, and southeast South America $(15-35^{\circ} \text{ S}, 35-65^{\circ} \text{ W})$.

		Nov 2008	May 2009			
Region	TRMM 3B43	CCATT-BRAMS	WRF-Chem	TRMM 3B43	CCATT-BRAMS	WRF-Chem
Amazon	0.24	0.22	0.51	0.20	0.15	0.40
Northeast	0.12	0.07	0.08	0.37	0.23	0.49
Southeast	0.19	0.11	0.24	0.10	0.06	0.07

Table 3. Values of RMSE and bias for CCATT-BRAMS and WRF-Chem simulations for 26 METAR and 52 SYNOP stations in the Amazon Basin for BARCA A (October–November 2008) and BARCA B (April–May 2009).

		Oct-N	ov 2008		Apr–M		
			CCATT- BRAMS	WRF-Chem		CCATT- BRAMS	WRF-Chem
T (K)	Mean Obs.	295.97			293.89		
	RMSE		2.30	2.81		1.70	2.44
	Bias		1.04	-2.42		-0.06	-2.28
T _d (K)	Mean Obs.	289.26			288.49		
	RMSE		2.68	1.72		1.76	1.67
	Bias		-1.92	-0.81		-0.99	-0.83
Wind Spd.	Mean Obs.	3.00			2.59		
$(m s^{-1})$	RMSE		1.41	1.33		1.15	1.00
	Bias		-0.60	0.16		-0.51	0.07
Sfc. Press.	Mean Obs.	1013.17			1016.09		
(hPa)	RMSE		2.16	1.43		1.09	1.34
	Bias		-2.01	-1.02		-0.79	-1.17
Precip. TRMM	Mean Obs.	0.49			0.62		
$(mm h^{-1})$	RMSE		2.42	4.50		3.03	7.12
	Bias		0.28	3.47		0.25	5.84

World Meteorological Organization (WMO) definition of a temperature lapse rate less than 2 K km^{-1} (Logan, 1999).

The models were also compared with soundings measuring O₃, temperature, and relative humidity conducted at sites in Paramaribo, Suriname $(5.8^{\circ} \text{ N}, 55.2^{\circ} \text{ W})$ and Natal, Brazil $(5.4^{\circ} \text{ S}, 5.4^{\circ} \text{ W})$ during the BARCA periods as part of the Southern Hemisphere ADditional OZonesondes (SHADOZ) network (http://croc.gsfc.nasa.gov/shadoz/) (Thompson et al., 2003a, b, 2007).

Monthly mean precipitation over the Amazon region was obtained from the 3B43 TRMM and Other Data Precipitation Product at a spatial resolution of $0.25^{\circ} \times 0.25^{\circ}$ (obtained from http://trmm.gsfc.nasa.gov/) (Kummerow et al., 1998; Kawanishi et al., 2000). TRMM 3B43 is derived from retrievals of 3-hourly precipitation amount from the precipitation radar (PR), TRMM microwave imager (TMI), and visible and infrared scanner (VIRS) aboard the TRMM satellite, merged with rain gauge data from the Climate Anomaly Monitoring System (CAMS) and the Global Precipitation Climatology Project (GPCP). Satellite estimates of precipitation are used for model evaluation due to their more complete spatial and temporal coverage compared to rain gauge data. Buarque et al. (2011) found that mean annual rainfall from Brazilian rain gauge and TRMM 3B42 3-hourly data at 488 sites in the Amazon Basin for the years 2003-2005 agreed within 5%. Other characteristics of the rainfall distribution, such as the number of days with rainfall, differed more substantially. Mean precipitation during the dry-to-wet (November 2008) and wet-to-dry (May 2009) transition seasons was calculated for the TRMM 3B43 data and the CCATT-BRAMS and WRF-Chem models for three regions: the Amazon (15° S–10° N, 50–80° W), northeast Brazil (15° S–0° N, 35-50° W), and southeast South America (15-35° S, 35-65° W). The values are listed in Table 2. The mean precipitation on the 35 km resolution domain for the 2 months is shown in Fig. 4, as well as the delineations of the subregion boxes.



Figure 6. Mean daily cycles of surface (**a**) latent (LE) and sensible (H) heat and (**c**) incident shortwave (S_{in}) and incoming (L_{in}) and outgoing (L_{out}) longwave radiation fluxes for a forest site and (**b**) heat and (**d**) radiation fluxes for a pasture site, comparing observations (solid lines) from von Randow et al. (2004) for the dry-to-wet transition season (July–September 1999–2000) and from WRF-Chem (crosses) and CCATT-BRAMS (circles) for the BARCA A period (October–November 2008).



Figure 7. Mean daily cycles of surface (**a**) latent (LE) and sensible (H) heat and (**c**) incident shortwave (S_{in}) and incoming (L_{in}) and outgoing (L_{out}) longwave radiation fluxes for a forest site and (**b**) heat and (**d**) radiation fluxes for a pasture site, comparing observations (solid lines) from von Randow et al. (2004) for the wet-to-dry transition season (February–March 1999, January–March 2000) and from WRF-Chem (crosses) and CCATT-BRAMS (circles) for the BARCA B period (April–May 2009).

				Pasture	;			
PBL Height (km)		Method	TKE	Theta	WRF MYNN	TKE	Theta	WRF MYNN
BARCA A (Nov 2008)	CCATT-BRAMS	Mean SD	1.103 0.621	1.610 0.646		1.143 0.581	1.636 0.640	-
	WRF-Chem	Mean SD	1.211 0.655	1.131 0.390	0.983 0.423	1.258 0.665	1.087 0.470	0.991 0.455
BARCA B (May 2009)	CCATT-BRAMS	Mean SD	0.628 0.515	1.067 0.554		0.669 0.527	1.049 0.564	-
	WRF-Chem	Mean SD	0.828 0.443	0.922 0.336	0.815 0.288	0.845 0.432	0.933 0.282	0.766 0.272

Table 4. PBL height at 21:00 UTC (17:00 LT) estimated from CCATT-BRAMS and WRF-Chem using methods based on turbulent kinetic energy (TKE) and theta (θ) and the diagnostic from the WRF MYNN PBL scheme.

Surface downward shortwave radiation (Level 1.5) data obtained with a Kipp and Zonen CM-21 pyranometer (305–2800 nm) were obtained from the Solar Radiation Network (SolRad-Net) site at Manaus (2.56° S, 60.04° W, 93 m a.s.l.) (http://aeronet.gsfc.nasa.gov/cgi-bin/bamgomas_interactive).

Mean daily cycles of fluxes of sensible and latent heat and radiation were obtained from flux tower measurements for the wet (February–March 1999, January–March 2000) and dry (July–September 1999–2000) seasons at forest (Rebio Jarú, 10.08° S, 61.93° W, 145 m a.s.l.) and pasture (Fazenda Nossa Senhora, 10.75° S, 62.37° W, 293 m a.s.l) tower sites (von Randow et al., 2004).

Surface meteorological station data was obtained for the BARCA region for October–November 2008 and April–May 2009 from 52 SYNOP (INMET) and 26 METAR (airport) stations, the locations of which are depicted in Fig. 10. The models were also evaluated against TRMM 3B42 3-hourly precipitation rates at the 78 surface station locations in the Amazon. Values of root mean squared error (RMSE) and bias for the CCATT-BRAMS and WRF-Chem simulations are shown in Table 3.

Meteorological soundings from the Manaus airport (3.15° S, 59.98° W) were conducted at 00:00 UTC (12 in October–November 2008, 60 in April–May 2009) and 12:00 UTC (49 in October–November 2008, 60 in April–May 2009). During BARCA A, 13 additional soundings were conducted at 18:00 UTC from 18 November–1 December 2008.

Fisch et al. (2004) found that in the dry season (14– 25 August 1994), higher sensible heat fluxes over pasture increase the maximum height at 21:00 UTC (17:00 LT) of the convective boundary layer (CBL) from around 1100 m for forest (Rebio Jarú) to 1650 m over pasture (FNS). On the other hand, during the wet season (January–February 1999) the height of the CBL was similar for both land types, around 1000 m. The simulated height of the PBL at 21:00 UTC above the forest and pasture sites (Table 4) was analyzed from model output using two different methods: TKE, the first level above the surface where the turbulent kinetic energy (TKE) from the PBL schemes dropped below $0.01 \text{ m}^2 \text{ s}^{-1}$ and theta, the first level above the surface where theta exceeded theta of the level below by 0.6 K. In addition, WRF MYNN is the diagnostic from the WRF PBL scheme which takes into account TKE as well as stability.

3.4 Results and discussion

3.5 BARCA O₃ observations

The vertical distributions of O₃ measured by the aircraft during BARCA A and B are depicted in Fig. 2. Observations during the dry-to-wet transition (BARCA A) are plotted separately for clean (west, north, and around Manaus regions) and fire-influenced polluted (east and south regions) conditions. The longitude and latitude bounds and flight dates included in each geographic region from BARCA A and BARCA B are listed in Table 6. The O₃ distributions are similar during BARCA A in the clean regions and BARCA B, with median values ranging from 10 to 25 ppb. However, there is more variability, as measured by the difference between the 25th and 75th percentiles, in the BARCA A data. This may be due to downward mixing of O3 transported longrange from fires in Africa or recirculated from the polluted southeast Brazil region. In the fire-influenced regions during BARCA A, medians range from 25 to 45 ppb, peaking at a typical plume injection height for savanna fires of 2-3 km. The highest variability is seen in polluted conditions during BARCA A, particularly at 2-3 km, indicating the influence of small-scale fire plumes. This variability of O₃ in the PBL presents a challenge to the regional models, since the effects of small-scale processes such as plume rise and convection are parameterized and averaged across the grid cell.



Figure 8. Mean vertical profiles at Manaus from radiosoundings (black, gray line denotes 1 standard deviation), CCATT-BRAMS (blue) and WRF-Chem (green) for October–November 2008 at (a) 00:00, (b) 12:00, and (c) 18:00 UTC.

3.6 Observed and simulated meteorology

Tropospheric O_3 distributions are driven by both chemical processes, including chemistry and emissions of O_3 precursors, and meteorological ones, such as solar radiation, tracer transport, and removal. During the dry-to-wet transition season, increased actinic fluxes stimulate the production of hydroxyl (OH) radicals from O_3 photolysis that can lead to net O_3 production (Seinfeld and Pandis, 2006). In November 2008, a band of elevated precipitation extended in TRMM 3B43 observations from the northwest Amazon to southeast Brazil but the northern Amazon between Manaus and Belém was relatively dry (Fig. 4a).

On the other hand, lower levels of O_3 in the wet-to-dry transition season are largely associated with increased presence of convective clouds and precipitation. Decreased surface temperatures and incident solar radiation due to cloudiness suppress emissions of biogenic VOCs such as isoprene (Fall and Wildermuth, 1998). In addition, higher surface humidity and precipitation decrease the occurrence of fires (Morton et al., 2013; Chen et al., 2013) that emit NO_x and VOCs (Freitas et al., 2007). O₃ precursors are further decreased by wet removal within the storms (Barth et al., 2007a). In May 2009, intense precipitation as observed by TRMM 3B43 stretched from the northwest Amazon to the northeast coast of Brazil (Fig. 4b). In radiosoundings at Manaus, a more pronounced decrease in dew point temperature is observed in May 2009 (Fig. 9) than in November 2008 (Fig. 8) in upper levels (300–400 hPa) from 00:00 to 12:00 UTC, likely due to precipitation.

Land cover also impacts surface heat and moisture exchange and can thus affect convective triggering. In both



Figure 9. Mean vertical profiles at Manaus from radiosoundings (black, gray line denotes 1 standard deviation), CCATT-BRAMS (blue) and WRF-Chem (green) for April–May 2009 at (**a**) 0 and (**b**) 12:00 UTC.



Figure 10. Locations of surface meteorological stations for model evaluation.

transition seasons, surface sensible heat fluxes are higher and latent heat fluxes are lower at the pasture compared to forest sites (Figs. 6a–b and 7a–b). However, incident solar radiation and thereby peak sensible heat flux (Fig. 7) are lower in the wet-to-dry than dry-to-wet transitions (Fig. 6) for both forest and pasture sites.

Now we summarize the key findings of the model-data meteorological comparison and their implications for the chemistry simulations. The models capture the seasonal spatial distribution of precipitation over northern South America (Fig. 4), and the signs of NE-SE differences are correctly modeled by both models during both seasons, i.e., the NE is drier than the SE during November and vice-versa during May. For the Amazon, CCATT-BRAMS slightly underestimates the precipitation rates in both seasons, but the rate in WRF-Chem is about twice that of TRMM 3B43 (Table 2). This may lead to errors in the strength and vertical distribution of convective transport and the amount of convective wet removal.

Peak shortwave radiation during the dry-to-wet transition at Manaus is within the error bars of the observations for both models (Fig. 5). However, for the southern Amazon forest and pasture sites, peak shortwave radiation may be overestimated (underestimated) by $50-100 \text{ Wm}^{-2}$ by CCATT-BRAMS (WRF-Chem) (Figs. 6–7), suggesting that there is insufficient (excessive) cloudiness in the models. This will increase (decrease) surface temperature and evaporation, and therefore increase (decrease) O₃ production from photolysis.

In the dry-to-wet transition season (Fig. 6), the peak observed Bowen ratio (sensible/latent heat flux) is lower at the forest site than the pasture site (0.23-0.38 vs. 0.8). However, in WRF-Chem, the Bowen ratio at 13:00 LT shows a smaller contrast between the forest and pasture sites (0.40 vs. 0.51), due to underestimated sensible heat flux at the pasture site. In the wet-to-dry transition season (Fig. 7), the peak Bowen ratio is lower at both forest and pasture sites for this season (0.18-0.39 vs. 0.33-0.59). On the other hand, in WRF-



Figure 11. O₃ as observed (black circles) and simulated with CCATT-BRAMS (blue stars) and WRF-Chem (base case: green diamonds, 2DEPVEL: cyan circles, and 0.5ENOx: yellow squares) from BARCA flights from (a) Manaus to Belém on 18 November 2008, (b) Belém to Manaus on 19 November 2008, (c) Manaus to Belém on 21 May 2009, and (d) Belém to Manaus on 23 November 2009.

Chem, the peak latent and sensible heat flux and thus the Bowen ratio are nearly constant at the forest and pasture sites (0.39 vs. 0.38). This indicates that the Noah land surface model is not properly representing the impact of conversion of forest to pasture and the resulting increase in sensible heat flux.

At the surface stations (Table 3), both models overestimate precipitation on average. Dew point temperature is underestimated by 1–2 K and temperature is underestimated in all cases by 0.1–2.4 K except by CCATT-BRAMS during BARCA A, which overestimated temperature by about 1 K. All of these biases will decrease photochemical O₃ production at the surface. The models generally show good agreement with soundings at Manaus (Figs. 8–9), but excess moisture (positive dewpoint bias of 10 K) in CCATT-BRAMS above 500 hPa may lead to increased photochemical production of O₃ at mid-levels.

Next we compare the CBL heights for wet and dry seasons reported by Fisch et al. (2004) with the simulated PBL heights in the dry-to-wet and wet-to-dry transitions (Table 4). The models represent the pattern of lower PBL heights in the wet-to-dry than dry-to-wet transitions, and similar PBL heights at the forest and pasture sites. However, for the dryto-wet transition, the PBL heights are indistinguishable between forest and pasture sites for both models, and generally closer to the observed forest (1.1 km) than pasture (1.65 km) values. Additionally, for the wet-to-dry transition, the mean PBL height for all models and diagnostics except theta for CCATT-BRAMS are lower than observed (1 km). Overall, the models underestimate the PBL depth, which may contribute to an overestimate of O₃ near the ground. Despite these limitations, the models are able to capture the meteorological contrast between the dry-to-wet and wet-to-dry transition seasons.

3.7 Observed and simulated chemistry

3.7.1 Mean O₃ profiles

An example of observed and simulated O_3 during the flight legs between Manaus and Belém in BARCA A and B is shown in Fig. 11. While the models capture the pattern of increasing O_3 values with height, the models underestimate elevated O_3 values from 2.5 to 4.5 km and overestimate low values near the surface (1–1.5 km). The models also do not reproduce the variability in the high values, likely due to the aircraft intersection of biomass burning plumes. This is expected given the coarse horizontal grid resolution. Thus, mean profiles are analyzed in order to study differences among the regions and seasons and to assess the models' abilities to capture the impacts of such small-scale processes on regional O_3 distributions.

The mean vertical O_3 profiles for observations, CCATT-BRAMS and WRF-Chem for the regions to the west, north, south, east, and around Manaus are shown for BARCA A and B in Figs. 12 and 14, respectively, and NO profiles corresponding to the aircraft tracks are depicted in Figs. 13 and 15, respectively. Mean profiles from longitudinal surveys over Amazonia of O_3 during ABLE-2A (Browell et al., 1988) and ABLE-2B (Harriss et al., 1990) and NO during ABLE-2A (Torres and Buchan, 1988) are included for comparison. In BARCA B, O_3 values were at or near background values in all regions, ranging from 8–15 ppb at the surface to



Figure 12. Mean vertical O_3 profiles for BARCA A flights for observations (black, gray line denotes 1 standard deviation), CCATT-BRAMS (blue), and WRF-Chem (base case: green, 2DEPVEL: cyan, and 0.5ENOx: yellow) simulations by region: (a) north, (b) east, (c) west, (d) south, and (e) around Manaus. ABLE-2A observations (gray) from the same regions are included for comparison.

2-15 ppb at 4-4.5 km, and the models are generally within 5-10 ppb of the observations. During BARCA A, while the W region still had low O₃ values (5 ppb at the surface to 20 ppb at 4–4.5 km), the N, S, and M regions ranged from 15-20 ppb at the surface to 30-35 ppb at 4-4.5 km, and the E region presented the most elevated values, from 25-55 ppb. ABLE-2A O₃ profiles are similar in all regions, ranging from 15-20 ppb near the surface to 30-40 ppb from 4-6 km, so that the BARCA values are higher in the fire-influenced east and south regions, lower in the north and west regions, and very similar around Manaus. The profiles from ABLE-2B are within 1 standard deviation of the BARCA B measurements, except for the north region, where they are lower (5-15 ppb). These results suggest an increasing influence of fire emissions to the east and south of Manaus, but that O₃ in clean regions has not changed much.



Figure 13. Mean vertical NO profiles corresponding to BARCA A flights for CCATT-BRAMS (blue) and WRF-Chem (base case: green, 2DEPVEL: cyan, and 0.5ENOx: yellow) simulations by region: (a) north, (b) east, (c) west, (d) south, and (e) around Manaus. ABLE-2A observations (gray) from the same regions are included for comparison.

A similar model behavior is seen in the broad regional mean profiles (Fig. 16). All simulations over-estimate O_3 throughout the PBL and lower troposphere during clean conditions in BARCA A, but under-estimate O_3 in polluted conditions. This is especially true from 2 to 4 km where biomass burning plumes detrain O_3 precursors. During BARCA B all simulations show good agreement.

In order to understand the possible sources of model error, we now individually examine the contributions of different chemical sources and sinks, including surface emissions and deposition, boundary inflow, and chemistry within the PBL.

3.7.2 Emissions

The sensitivities of O_3 production to NO_x or BVOC emissions depend upon the relative amounts of VOCs and NO_x present. Under clean conditions with a high VOC : NO_x ratio, O_3 production is NO_x sensitive; increases in NO_x will lead

to more O₃ while increased VOCs will have little impact. On the other hand, in polluted areas with a high NO_x: VOC ratio, the system is VOC sensitive; that is, increased VOCs contribute to O₃ production but higher NO_x actually depletes O₃. Emissions of BVOCs can increase O₃ production by the following mechanism. Oxidation of BVOCs can lead to formation of hydroperoxyl (HO₂) and organic peroxy (RO_2 / radicals, which react with NO to form NO₂. NO₂ in turn photolyzes to form $O({}^{3}P)$, which reacts with O_{2} to form O_{3} (National Research Council, 1991). We expect the polluted east/south regions during BARCA A to be VOC sensitive and the clean west, north, and around Manaus regions during BARCA A and all regions in BARCA B to be NO_x sensitive. Kuhn et al. (2010) determined via aircraft transects in the Manaus urban plume that most of the VOC reactivity was provided by isoprene emissions from the surrounding rainforest, and NO_x emissions suppressed O₃ production close to urban sources, but stimulated it downwind.

For BARCA, the simulated mean monthly emission rates for two O₃ precursors, NO_x (anthropogenic and biomass burning) and isoprene (biogenic) are shown in Fig. 17. In November 2008, elevated NO_x emission rates of up to $5 \times$ 10^{-5} kg m⁻² d⁻¹ are seen from an area of intense biomass burning in the northeast Amazon, as well as from more scattered fires in the southeast Amazon. These are both regions that were overflown by the aircraft (Fig. 1). In May 2009, the Amazon region is largely free of fire. Because biogenic NO emissions (e.g., from soil) were not included in the MEGAN climatology used in this study, background NO emissions (in absence of fire) are likely too low. Typical model anthropogenic NO_x emissions values over the Amazon, primarily from biofuel sources, were $0.008-13 \,\mu g \,\mathrm{N \,m^{-2} \,hr^{-1} \,N}$. These NO_x emissions included in the models were less than one-third of the mean values of $44 \pm 14 \,\mu g \,\mathrm{N \,m^{-2} \,h^{-1}}$ NO measured by Kaplan et al. (1988) during ABLE-2A. This is considered a threshold value for NO_x-driven O₃ production to be the dominant O₃ source in the PBL. The model emissions were also much lower than the mean emissions from forest of $35.8 \,\mu$ g N m⁻² h⁻¹ NO measured in the 1998 dry season (Garcia-Montiel et al., 2003). Wetting the forest soil resulted in small pulses of NO and therefore the mean emissions are expected to be higher in the wet season than dry season.

Isoprene emissions are highest in the western and southern Amazon, reaching 15×10^{-5} kg m⁻² d⁻¹ in November 2008 and $5-10\times 10^{-5}$ kg m⁻² d⁻¹ in the aircraft sampling region. Due to decreased surface temperature and incident solar radiation in the rainy season, isoprene emissions in the Amazon Basin are much lower during BARCA B, $3-5\times 10^{-5}$ kg m⁻² d⁻¹. The MEGAN emissions are consistent with isoprene emission measurements above the Amazonian canopy: a normalized flux of 5.76×10^{-5} kg m⁻² d⁻¹ in July 2000 at the end of the rainy season (Rinne et al., 2002) and an average noontime flux of $18.7 \pm 5.5 \times 10^{-5}$ kg m⁻² d⁻¹ in September 2004 during the dry season (Karl et al., 2007).



Figure 14. Mean vertical O_3 profiles for BARCA B flights for observations (black, gray line denotes 1 standard deviation), CCATT-BRAMS (blue), and WRF-Chem (base case: green, 2DEPVEL: cyan, and 0.5ENO_x: yellow) simulations by region: (a) north, (b) east, (c) west, (d) south, and (e) around Manaus. ABLE-2A observations (gray) from the same regions are included for comparison.

3.7.3 Deposition

Figures 18 and 19 show the average O_3 dry deposition fluxes and median daytime deposition velocities, respectively, as simulated on the 35 km resolution domain by the CCATT-BRAMS and WRF-Chem models for November 2008 and May 2009. In the Amazon Basin, O_3 deposition fluxes are higher in the dry-to-wet transition season, with values reaching 3.5 nmol m⁻² s⁻¹ for CCATT-BRAMS and 6 nmol m⁻² s⁻¹ for WRF-Chem in the northeast Amazon, near the region of concentrated biomass burning. These values are also seen along the northern Andes and southeast Brazil, due to recirculation of O_3 -rich air. In the wet-to-dry transition season, O_3 deposition is at a minimum in the western Amazon, with values of 0.5–1 nmol m⁻² s⁻¹ for CCATT-BRAMS and 2 nmol m⁻² s⁻¹ for WRF-Chem. For both models, deposition velocities are higher over the rainforest than

Table 5. Observed average O_3 dry deposition flux (nmol m⁻² s⁻¹) and daytime (11:00–21:00 UTC) median deposition velocity (cm s⁻¹) in the dry and wet seasons (Rummel et al., 2007), and from WRF-Chem and CCATT-BRAMS simulations from November 2008 (dry-to-wet transition) and May 2009 (wet-to-dry transition) for Reserva Biológica Jarú (RBJ), Fazenda Nossa Senhora (RNS), and Reserva Ducke (RD).

			Dry Seaso	Wet Season			
Site		Observed	CCATT- BRAMS	WRF-Chem	Observed	CCATT- BRAMS	WRF-Chem
RBJ (forest)	Flux v _d	-5.69 0.6	-2.43 0.3	-3.25 0.5	-2.93 1.2	-1.59 0.4	-2.39 0.8
FNS (pasture)	Flux v _d	-4.68 0.6	$-3.06 \\ 0.4$	-2.49 0.4	-2.04 0.7	-2.07 0.4	-2.04 0.7
RD (forest)	Flux v _d				-1.82 1.6	$-1.63 \\ 0.4$	-2.68 0.6

Table 6. Longitude and latitude bounds and dates for each region of the BARCA A and B campaigns.

BARCA A (Nov 2008)						BARCA B (May 2009)				
Region	Longitude		Latit	ude	Days	Longitude		Latitude		Days
West	-60.06	-54.24	-12.00	-3.03	29, 30	-61.16	-59.46	-3.71	-2.39	28
North	-62.00	-59.11	-3.04	2.89	23	-61.81	-60.00	-3.04	3.71	19
Around Manaus	-61.52	-58.50	-4.39	1.00	16, 22	-62.14	-60.00	-4.07	-2.16	15, 17
East	-108.73	-48.45	-3.04	-1.33	18, 19	-60.34	-44.82	-4.39	0.14	21, 22, 23, 26
South	-67.69	-60.01	-3.40	0.12	25, 26	-63.93	-60.01	-8.77	-3.04	27

in the savanna to the east or south of the Amazon Basin, and higher in the wet-to-dry transition than in the dry-to-wet transition. These patterns are also seen in the tower observations in Table 5.

O₃ surface fluxes and dry deposition velocities predicted by the models were compared with observations from several field campaigns (Table 5). These include during the dry (May 1999) and wet (September-October 1999) seasons at Reserva Biológica Jarú (RBJ, forest) and Fazenda Nossa Senhora (FNS, pasture) from LBA-EUSTACH (Rummel et al., 2009; Kirkman et al., 2002) and during the wet season at Reserva Ducke (RD, forest tower near Manaus, 2.95° S, 59.95° W) from ABLE 2B (April-May 1987) (Fan et al., 1990) and at FNS from LBA-TRMM (January-February 1999) (Sigler et al., 2002). For the observations, the means of the hourly (WRF-Chem) and 3-hourly (CCATT-BRAMS) O₃ dry deposition fluxes (nmol $m^{-2} s^{-1}$) and the medians of the daytime (11:00-21:00 UTC) hourly mean deposition velocities (cm s^{-1}) are shown. The values were extracted from the grid points closest to the tower locations. In the observations, O₃ fluxes are larger in the dry season, due to higher O₃ mixing ratios. However, deposition velocities are higher in the wet season, and O₃ deposition to the Amazon forest constitutes a globally significant O₃ sink (Rummel et al., 2009). Both models capture these patterns, but the models underestimate the deposition velocities by 15–75 %, which may be partially

responsible for the low O_3 fluxes at the Jarú forest site in both seasons and the pasture site in the dry season.

3.7.4 Boundary conditions

The mean tropospheric and total tropospheric column O₃ from OMI/MLS, CCATT-BRAMS, and WRF-Chem for November 2008 and May 2009 are shown in Figs. 20 and 21, respectively. The models significantly underestimate the total columns from satellite and middle altitudes from BARCA. For both BARCA A and B, the models represent the pattern of lower O₃ over the Amazon and higher values over northeast Brazil (BARCA A only) and at 30° S, although the values are strongly underestimated. In November 2008 (Fig. 14), OMI/MLS mean tropospheric O₃ concentrations show an inflow of elevated O₃ (mean ca. 55 ppb, total 40-45 DU) on the northeast Brazilian coast due to cross-Atlantic transport from biomass burning in southern and sub-Saharan Africa. Additionally, a band of elevated O₃ (mean 55–60 ppb, total 35-40 DU) passes over the South American continent at around 30° S, also from cross-Atlantic transport. During this month, Northern Hemisphere O₃ levels to the north of South America are relatively low (mean 35-40 ppb, total 25-30 DU). On the other hand, the tropospheric ozone distribution in May 2009 (Fig. 15) is characterized by a band of low ozone extending over the Amazon Basin and northeast Brazil between 10° S and 10° N (mean 25-35 ppb, total



Figure 15. Mean vertical NO profiles corresponding to BARCA B flights for CCATT-BRAMS (blue) and WRF-Chem (base case: green, 2DEPVEL: cyan, and 0.5ENOx: yellow) simulations by region: (**a**) north, (**b**) east, (**c**) west, (**d**) south, and (**e**) around Manaus.

20-25 DU). In addition, slightly elevated values are found at around 30° S, primarily over the oceans (40–55 ppb, 30– 35 DU), and higher ozone is seen in the Northern Hemisphere (mean 50–55 ppb, total 35–40 DU to the north of 10° N). Both models capture the overall distribution (inflow in NE Brazil in November 2008, lower values over the Amazon Basin, elevated at 30° S) but values are underestimated relative to OMI/MLS. In general the models overestimate O₃ in the PBL compared to aircraft measurements, but underestimate the total column values relative to the OMI/MLS satellite product. This suggests that the total column values in Amazonia are dominated by global pollution from Africa, rather than local O₃ production from biomass burning. A typical OMI averaging kernel (cloud-free ocean conditions) shows maximum sensitivity from 594-416 hPa (Zhang et al., 2010). Therefore, OMI may not be detecting O_3 in the PBL from local sources, but rather primarily seeing global pollution from Africa.

Above the boundary layer, from 3 to 4 km a.g.l., chemical inflow at the eastern boundary of South America may con-



Figure 16. O_3 as observed (black circles) and simulated with CCATT-BRAMS (blue stars) and WRF-Chem (base case: green diamonds, 2DEPVEL: cyan circles, and 0.5ENOx: yellow squares) during (a) BARCA A, clean conditions (west, north, and around Manaus regions), (b) BARCA A, polluted conditions (east and south regions), and (c) BARCA B.

tribute to O₃ elevated above background. In order to evaluate the model representation of this inflow, vertical profiles from SHADOZ soundings on the northeast coast of South America during the BARCA A and B periods were compared with CCATT-BRAMS and WRF-Chem (Fig. 22). In addition, 120 h back trajectories from the sounding locations at heights of 1500, 6000, and 9000 m above ground level (gal) were calculated with the HYSPLIT model (http://ready.arl. noaa.gov/hypub-bin/trajtype.pl?runtype=archive) using meteorological data from the NCEP/NCAR global reanalysis. Inflow at Paramaribo originated either in the Caribbean or the tropical Atlantic, while at Natal, air parcels came from anticyclonic recirculation from southeastern Brazil or the tropical Atlantic. Both models generally represent the SHADOZ O₃ profiles up to 600 hPa, but do not capture layers of elevated O_3 above 500 hPa. These are likely to be either from pollution recirculated from southeast Brazil or possibly from African biomass burning. The models also do not reproduce thinner layers of high O₃ below 600 hPa. For example, at Natal on 7 November 2008 (Fig. 22c, air of African origin at circa 850 and circa 470 hPa) and 19 November 2008 (Fig. 22d, air from the central African coast at circa 850 hPa and recirculation from southeastern Brazil at circa 470 and circa 310 hPa) and at Paramaribo on 11 May 2009 (Fig. 22f, air of tropical Atlantic origin at all three levels), both models underestimate O₃ above 500 hPa by 40-60 ppb (model values of 30-50 ppb vs. observations maximum values of 80-100 ppb). A previous analysis of ozone soundings and air-



Figure 17. Mean emission rates $(10^{-5} \text{ kg m}^{-2} \text{ d}^{-1})$ from PREP-CHEM-SRC for the 35 km domain (dark gray outline) for NO_x for (a) BARCA A (November 2008) and (b) BARCA B (May 2009) and isoprene for (c) BARCA A and (d) BARCA B periods.



Figure 18. Average O_3 dry deposition flux (nmol m⁻² s⁻¹) as simulated on the 35 km resolution domain (dark gray outline) by the CCATT-BRAMS model for (a) November 2008 and (b) May 2009 and by the WRF-Chem model for (c) November 2008 and (d) May 2009.



Figure 19. Same as Fig. 12, but daytime (11:00–21:00 UTC) median deposition velocity (cm s⁻¹).



Figure 20. Mean tropospheric O₃ (ppb) on the 35 km domain from (a) OMI/MLS, (b) CCATT-BRAMS and (c) WRF-Chem and total tropospheric column O₃ (Dobson units) from (d) OMI/MLS, (e) CCATT-BRAMS and (f) WRF-Chem for November 2008.



Figure 21. Same as Fig. 14, but for May 2009.

craft measurements at Natal suggested that increases in tropospheric ozone in the southern Hemisphere springtime may be due to stratospheric intrusion (Logan, 1985). This is consistent with the November 2008 profiles at Natal; the models may not be capturing the intrusion of stratospheric air masses seen in the observations, indicated by upper tropospheric (> 500 hPa) layers with elevated O₃ and very low relative humidity (< 20 %). On the other hand, at Paramaribo on 6 and 25 November 2008 and at Paramaribo on 4 May 2009, when air masses at all levels were of Northern Hemisphere origin, the models reproduced the nearly constant with altitude O₃ values of 30–40 ppb.

3.7.5 Chemistry

The excess O₃ in the PBL in the models could be due to either a low deposition sink, as O₃ dry deposition velocities in the models are about half of observed values, or excessive model sensitivity to NO_x emissions, or both. Two additional simulations were conducted with WRF-Chem to evaluate the model sensitivity to these processes: (1) doubling the calculated deposition velocity for O₃ only (2DEPVEL) and (2) halving the NO_x surface emission rates (0.5ENOx). The O3 profiles corresponding to BARCA flights for these two simulations are also included in Figs. 12 and 14. The corresponding NO profiles from all model simulations as well as a mean profile over Amazonia from ABLE-2A are depicted in Figs. 13 and 14. The 0.5ENOx simulation reduces O₃ more than 2DEPVEL throughout the entire profile. In the dry-towet transition, 2DEPVEL reduces O₃ in the lower PBL by about 25%, while 0.5ENOx decreases O₃ by around 40%, and in the wet-to-dry-transition the reductions are about 10 and 30%, respectively. In general the 0.5ENOx O₃ profiles are lower than observed in the first 500 m above the surface, but they provide the best representation of the data for the north and west regions in the dry-to-wet transition. They also provide a similarly good fit as 2DEPVEL for the east, Manaus, and south regions, while in the wet-to-dry transition 0.5ENOx is closer to the observed value from 0-500 m in all regions except the north. During BARCA A, NO in all WRF-Chem simulations in the north, west, and Manaus regions is 10–15 ppt from 0–500 m above the surface, increasing to a maximum of 20-50 ppt at 2 km a.g.l., and is generally lower than the ABLE-2A observations in the PBL. In the east and south, where biomass burning influence was seen, NO in 0-500 m a.g.l. increased from 20-50 ppt in the base simulation to 35-60 ppt in 2DEPVEL due to decreased O₃ and conversion of NO to NO₂, and was generally within 1 standard deviation of the ABLE-2A measurements in the PBL. In BARCA B, NO simulated by WRF-Chem is very low, 5-10 ppt in the entire profile, except for the west region, where a mean NO of 30 ppt is seen from 0-500 m a.g.l. This is again due to very low O₃, and for the Manaus region, where anthropogenic NO_x sources may have contributed to NO values of 20 ppt. These results suggest that adjustment of dry deposition parameterizations are needed to increase O₃ deposition velocities by about a factor of two in agreement with ground observations. Future research will compare simulated NO_x fields with observations from more recent field campaigns, as the results of these simulations also suggest that O_3 in WRF-Chem is very sensitive to NO_x emissions.

In summary, chemistry simulations of the BARCA periods with CCATT-BRAMS and WRF-Chem overestimated O_3 in the PBL by 5–10 ppb in the wet-to-dry transition (BARCA B), with background levels observed (10–20 ppb) in all regions. In the dry-to-wet transition (BARCA A), the models



Figure 22.

generally reproduced elevated O₃ levels in the northeast and southeast Amazon, where biomass burning emissions of precursors led to significant enhancements of ambient O₃. However, the models overestimate O_3 in the PBL by 5–10 ppb, whereas from 2 to 4 km the modeled values are generally lower than observations. These discrepancies of models with observations may result from an overly mixed (constant with altitude) profile due to overly active vertical mixing from the PBL scheme from 1 to 2 km or too much downward convective transport of O_3 from 2 km to the surface, as observed by Betts et al. (2002). In addition, the models may be missing sources of O₃ and/or precursors at 3-4.5 km in the model inflow boundary conditions. In the lower boundary layer, the surface sink of O₃ (dry deposition) may be too low, or overestimation of NO_x sources may produce too much O₃. Additional simulations with WRF-Chem showed that O₃ in the lower boundary layer was about twice as sensitive to increases in O₃ deposition velocity as reductions in NO_x emissions, but both simulations achieved better agreement with observations than the base-case simulation. Although NO emissions over the forest were less than half of observed values, likely due to the lack of inclusion of soil emissions, sufficient O_3 production occurred to match or exceed aircraft observations, suggesting that the model chemistry is overly NO_x -sensitive.

4 Conclusions

The BARCA campaign offered the first regional aircraft survey of O_3 in the Amazon Basin in both the dry-to-wet and wet-to-dry transition seasons. In both seasons, extremely low background O_3 values (< 20 ppb) were observed to the west and north of Manaus, and in the wet-to-dry transition low O_3 was also measured to the east and south and in the region around Manaus. These background values are the lowest observed on Earth, due to a combination of isolation from



Figure 22. Vertical profiles of potential temperature, relative humidity, and O_3 from SHADOZ soundings (black), CCATT-BRAMS (blue), and WRF-Chem (green) and HYSPLIT back trajectories at 13:00 UTC at 1500 m (circa 850 hPa, red), 6000 m (circa 470 hPa, blue), and 9000 m (circa 310 hPa, green) for: Paramaribo on (a) 6 November and (b) 25 November 2008, Natal on (c) 7 November and (d) 19 November 2008, and Paramaribo on (e) 4 May and (f) 11 May 2009.

anthropogenic and biomass burning NOx sources and O3 deposition to the forest canopy, and the ecosystem and atmospheric chemistry is adjusted to these very low values. According to models, the chemistry in the Amazon is very sensitive to NO_x emissions from soils, so that even a small overestimate of NO_x emissions generates too much O_3 in the PBL. However, it is likely that the model chemistry is incorrect in the PBL, because the models have about the right amount of NO_x but far too much O_3 in the PBL. Further simulations with WRF-Chem showed that the model O3 production is very sensitive to both the O₃ deposition velocities and the NO_x emissions, which were both about one-half of observed values. In polluted, VOC-sensitive conditions, approximately the correct net amount of O₃ is generated in the PBL. This suggests there is insufficient VOC reactivity in the models, since the correct amounts of O₃ deposition velocities and NO_x emissions would both decrease O₃ production. Additionally, in clean, NO_x-sensitive conditions, proportionally more O₃ is produced per unit NO_x emissions and the O₃ deposition velocities are still too low, resulting in an overestimate. Therefore, we conclude that the current model chemistry produces much more O₃ per unit NO_x than the atmosphere at very low NO_x, but may be about right in polluted conditions. In addition, simulated O₃ was lower than both the OMI/MLS total tropospheric O₃ and the BARCA observations in mid-levels, indicating that the models are missing sources at mid-levels from long-range and convective transport.

As the regional population grows in the Amazon basin, leading to increases in both urban and fire NO_x sources, this is indeed a big concern because PBL O_3 is lower in clean areas than the models predict, so that the change to

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polluted conditions is larger, and that the chemistry to define the path to higher NO_x conditions is poorly represented. Future modeling studies can include more complete organic chemistry and biogenic emissions, including NO emissions from soil, as well as improved representation of lightning NO_x production, dry deposition, convective transport, and wet scavenging processes, to address this NO_x sensitivity. Additionally, future field campaigns in the Amazon that include aircraft observations of nitrogen oxides and hydrocarbons and ground-based measurements of NO flux from the forest canopy may allow better constraints on the Amazonian O_3 budget.

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