Biomass burning related ozone damage on vegetation over the Amazon forest: A model sensitivity study.

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Abstract

The HadGEM2 Earth System climate model was used to assess the impact of biomass burning on surface ozone concentrations over the Amazon forest and its impact on vegetation, under present-day climate conditions. Here we consider biomass burning emissions from wildfires, deforestation fires, agricultural forest burning, residential and commercial combustion. Simulated surface ozone concentration is evaluated against observations taken at two sites in the Brazilian Amazon forest for years 2010 to 2012. The model is able to reproduce the observed diurnal cycle of surface ozone mixing ratio at the two sites, but overestimates the magnitude of the monthly averaged hourly measurements by 5-15 ppb for each available month at one of the sites. We vary biomass burning emissions over South America by +/-20, 40, 60, 80 and 100% to quantify the modelled impact of biomass burning on surface ozone concentrations and ozone damage on vegetation productivity over the Amazon forest. We used the ozone damage scheme in the “high” sensitivity mode to give an upper limit for this effect. Decreasing South American biomass burning emissions by 100% (i.e. to zero) reduces surface ozone concentrations (by about 15ppb during the biomass burning season) and suggests a 15% increase in monthly mean net primary productivity averaged over the Amazon forest, with local increases up to 60%. The simulated impact of ozone damage from present-day biomass burning on vegetation productivity is about 230 TgC/yr. Taking into account that
uncertainty in these estimates is substantial, this ozone damage impact over the Amazon forest is of the same order of magnitude as the release of carbon dioxide due to fire in South America; in effect to potentially double the impact of biomass burning on the carbon cycle.
Introduction

Biomass burning is a global source of aerosol and trace gases, including ozone (O₃) precursors, and can lead to local and regional O₃ pollution. Tropospheric O₃ is a greenhouse gas and, above background concentrations, an air pollutant: it is harmful to human health (e.g. Lippmann 1993; Burnett et al., 1997) and it damages plants (e.g. Rich et al., 1964; Fiscus et al., 2005; Felzer et al., 2007; Ainsworth et al., 2012). Tropospheric O₃ is a product of photochemical reactions whose main precursors are nitrogen oxides (NOₓ), carbon monoxide (CO), methane (CH₄) and volatile organic compounds (VOCs) (Seinfeld and Pandis, 1998). VOCs are particularly important in Amazonia because of the large natural biogenic and biomass burning emissions (Karl et al., 2007).

In the Amazon forest, biomass burning is mostly anthropogenic, and mainly occurs during the dry season (August to October). Biomass burning emissions drastically change the composition of the atmosphere, e.g. diurnal maximum mixing ratios of tropospheric O₃ varies from 12 parts per billion (ppb), during the wet season, to values as high as 100 ppb in the biomass burning affected dry season (Kirkman et al., 2002, Sigler et al., 2002, Artaxo et al., 2002, 2005, Rummel et al., 2007).

Surface O₃ mixing ratios over 40 ppb are known to produce visible leaf injury and damage to plants, reducing crop productivity and posing a threat to food security; nonetheless different climatic conditions (e.g. soil moisture and water stress) also play a role in determining leaf stomatal closure and hence there will be variable impacts of the same O₃ concentrations (Ashmore, 2005), e.g. tropical rainforest vegetation may be particularly sensitive to surface O₃, even at concentrations below 40ppb (a threshold associated with extra-tropical vegetation), due to high stomatal conductances. Moreover, tropical vegetation evolved in low background O₃ concentrations and could be more sensitive to O₃. In leaves, cellular damage caused by O₃ not only reduces photosynthetic rates but also requires increased resource allocation to detoxify and repair leaves (Ainsworth et al., 2012). Ozone damage to vegetation reduces plant productivity, decreasing the amount of carbon absorbed by plants, hence has an impact on climate via and indirect radiative forcing (Sitch et al., 2007).

Tropical rain forests play an important role in the global carbon budget, as they cover 12% of the Earth’s land surface and contain around 40% of the terrestrial biosphere’s carbon (Ometto et al., 2005, Taylor & Lloyd, 1992). It has been estimated that they may account for as much as 50% of
the global net primary productivity (Grace et al., 2001). Depending on age, land use and large scale meteorological conditions, tropical forest ecosystems can act as net carbon sources, sinks, or they can be in approximate balance (Lloyd et al., 2007, Gatti et al., 2013), but it is uncertain if global environmental changes are forcing these ecosystems outside their range of natural variation (Sierra et al., 2007). However, biomass burning may further reduce natural sinks in the neighbouring intact forest, via air pollution and O₃ damage on vegetation, and thus current estimates of the effects of biomass burning on the carbon cycle (Le Quéré et al., 2009) may be underestimated. Biomass burning is also an important aerosol source: regional levels of particulate matter are very high in the dry season in Amazonia (Artaxo et al., 2013), and the increase in diffuse radiation due to changes in aerosol loadings can increase net ecosystem exchange (NEE) quite significantly (Oliveira et al., 2007, Cirino et al., 2013). After a certain level of aerosol optical depth, the decrease in radiation fluxes can reduce significantly NEE over Amazonia (Cirino et al., 2013). This study does not consider the effects of the changes in diffuse radiation due to biomass burning on photosynthesis, or the impact of aerosols on O₃ chemistry via changing photolysis rate. That will be the focus of a separate study. Our specific aim is to estimate the effect of O₃-induced changes on vegetation productivity due to biomass burning.

Importantly, Sitch et al. (2007) performed their assessment of the potential impact of O₃ on vegetation using an offline simulation where monthly mean O₃ concentrations derived with a global chemistry climate model were used in determining the impacts of O₃ damage. Here we use an online flux-gradient approach to quantify the impact of biomass burning on surface O₃ concentration and O₃ damage on vegetation over the Amazon forest (see model description). The HadGEM2 (Hadley Centre Global Environment Model 2; Collins et al., 2011; Martin et al., 2011) Earth System climate model is used to study these interactions. We show results of the evaluation of surface O₃ simulated with HadGEM2 against observations in the Amazon forest and model experiments quantifying the impact of biomass burning on plant productivity.

**Methods**

We used HadGEM2 to simulate surface O₃ concentrations and O₃ damage on vegetation for present-day (2001-2009) climate conditions. Our version of HadGEM2 includes the O₃ damage scheme developed by Sitch et al. (2007). We evaluated simulated surface O₃ against observations taken at two sites in the Amazon forest: Porto Velho (Brazil; 8.69∞S; 63.87∞W), a site heavily impacted by
biomass burning emissions, and site ZF2 in the Cuieiras forest reserve in Central Amazonia (Brazil; 2.59°S; 60.21°W). A description of the sites can be found in Artaxo et al. (2013). In a sensitivity study we varied biomass burning emissions over South America by +/-20, 40, 60, 80, 100% to quantify the potential impact of biomass burning on surface O_3 concentrations and O_3 damage over the Amazon forest.

**Model description**

HadGEM2 is a fully coupled Earth-system model (Collins et al., 2011). It is built around the HadGEM2 atmosphere-ocean general circulation model and includes a number of earth system components: the ocean biosphere model diat-HadOCC (Diatom-Hadley Centre Ocean Carbon Cycle, a development of the HadOCC model of Palmer and Totterdell, 2001), the Top-down Representation of Interactive Foliage and Flora Including Dynamics (TRIFFID) dynamic global vegetation model (Cox, 2001), the land-surface and carbon cycle model MOSES2 (Met Office Surface Exchange Scheme; Cox et al. 1998, 1999; Essery et al. 2003), the interactive Biogenic Volatile Organic Compounds (iBVOC) emission model (Pacifico et al., 2012), the United Kingdom Chemistry and Aerosol (UKCA) model (O’Connor et al., 2014) and an interactive scheme of O_3 damage on vegetation (Sitch et al., 2007; Clark et al., 2011).

The configuration used here is a version of HadGEM2-UKCA with extended tropospheric chemistry (N96L38), the resolution is 1.25° latitude x 1.875° longitude (~200 x 140 km) at the equator with 38 vertical levels extending up to 39 km altitude. The land-based anthropogenic, biomass burning, and shipping emissions are taken from Lamarque et al. (2010), and represent a decadal (1997-2006) mean centered on the year 2000. The use of an emission pattern from 1997-2006 can lead to an overestimation of O_3 concentrations by the model, since the emissions vary on a year to year basis and are expected to be lower in recent years due to the reduction in Amazonian deforestation via burning, consequently reducing the amount of O_3 precursors. HadGEM2 runs at a 30-minute time step with the exception of global radiation, which is updated every 3 hours and provides radiative fluxes between those time steps via interpolation. This configuration is described and evaluated in O’Connor et al. (2014) with the exception of the Extended Tropospheric Chemistry (ExtTC) that has been applied in this work. The ExtTC mechanism has been designed to represent the key species and reactions in the troposphere in as much detail as is necessary to simulate atmospheric composition-climate couplings and feedbacks while retaining the capability to
conduct decade-long climate simulations. UKCA-ExtTC simulates the spatial distribution and
evolution in time of 89 chemical species, 63 of which are model tracers. The model includes
emissions from anthropogenic, biogenic, soil, and wildfire sources for 17 species: nitrogen oxides
(NOx = NO +NO2), CH4, carbon monoxide (CO), hydrogen (H2), methanol, formaldehyde,
acetaldehyde and higher aldehydes, acetone, methyl ethyl ketone, ethane (C2H6), propane (C3H8),
butanes and higher alkanes, ethene (C2H4), propene (C3H6), isoprene, (mono)terpenes, and a lumped
species representing aromatics (toluene + xylene) from anthropogenic sources.

Emissions of biogenic species (isoprene, terpenes, methanol, acetone) are computed by iBVOC and
provided to UKCA at every time step. The isoprene emission scheme is that of Pacifico et al.
(2011). Terpenes, methanol, and acetone emissions are simulated with the model described in
Guenther et al. (1995). Anthropogenic and wildfire emissions are prescribed from monthly mean
emission data sets prepared for CMIP5 using the historic scenario (Lamarque et al., 2010). Given
the difficulty in prescribing a diurnal cycle for fire emissions, these monthly mean emissions are
kept constant during the day. Wetland methane emissions are prescribed from data from Gedney et
al. (2004). Soil-biogenic NOx emissions are prescribed using the monthly distributions provided by
the Global Emissions Inventory Activity (http://www.geiacenter.org/inventories/present.html),
which are based on the global empirical model of soil-biogenic NOx emissions of Yienger and Levy
(1995). NOx emissions from global lightning activity are parameterized based on the convective
cloud top height following Price and Rind (1992, 1994) and are thus sensitive to the model climate.
UKCA also includes a dry deposition scheme based on the resistance in-series approach as outlined
in Wesely (1989). Physical removal of soluble species is parameterized as a first-order loss process
based on convective and stratiform rainfall rates (Collins et al., 2011).

The TRIFFID vegetation module of HadGEM2 simulates the dynamics of five plant functional
types (PFTs): broadleaf trees, needleleaf trees, shrubs, and C3 and C4 grass (i.e., grasses using the
C3 and C4 photosynthetic pathway, respectively). Changes in the extent of croplands over time are
not simulated but are prescribed from land use maps prepared for the Coupled Model
Intercomparison Project 5 (CMIP5, Taylor et al., 2012). Here we use the historic (1850–2000; Hurtt
et al., 2009) data sets, as described in Jones et al. (2011). A further four surface types (urban, inland
water, bare soil, and ice) are used in the land-surface scheme for the calculation of water and energy
exchanges between the land and the atmosphere. Each model grid box can include varying
proportions of several vegetation and/or surface types. The model does not include interactive
deforestation due to fire.
The parameterization of O₃ damage on vegetation is that of Sitch et al., (2007). This scheme uses a flux-gradient approach to model O₃ damage, rather than empirical approaches based on the accumulated O₃ exposure above 40 ppb (e.g. Felzer, et al. 2005). The Sitch et al. (2007) parameterization assumes a suppression of net leaf photosynthesis by O₃ that varies proportionally to the O₃ flux through stomata above a specified critical O₃ deposition flux. The critical deposition flux depends on O₃ concentration near the leaves, but also on stomatal conductance. This scheme also includes a relationship between stomatal conductance and photosynthesis, determining a reduction in stomatal conductance through O₃ deposition. As the O₃ flux itself depends on the stomatal conductance, which in turn depends upon the net rate of photosynthesis, the model requires a consistent solution for the net photosynthesis, stomatal conductance and the O₃ deposition flux. This approach to modelling O₃ effects on photosynthesis accounts for the complex interaction between CO₂ and O₃ effects, and can be used to study future climate impacts. This scheme includes a ‘high’ and ‘low’ parameterization for each PFT to represent species more sensitive and less sensitive to O₃ effects; in our analysis we use the ‘high’ sensitivity mode to establish the maximum response. The model was calibrated with data from temperate and boreal vegetation. Calibration data for other ecosystems, including tropical vegetation, are currently unavailable.

Description of the model experiments

All simulations use HadGEM2 in its atmosphere-only configuration, i.e., with all implemented couplings between atmosphere and land surface (including carbon cycle) active but without the atmosphere-ocean coupling. HadGEM2 was initialized with equilibrium concentrations of the major chemical components (O₃, CO, H₂, total reactive nitrogen (NO₃), BVOCs) taken from the CMIP5 simulation (see description of the simulations in Jones et al., 2011). Methane mixing ratios were prescribed as specified by CMIP5, with values of 1750 ppb for present-day. The decade-mean CO₂ atmospheric mixing ratio was 368 ppm.

Monthly means of sea surface temperature and sea ice cover were prescribed using climatologies derived from the appropriate decade of the Hadley Centre CMIP5 transient climate run Jones et al., (2011). The vegetation distribution for each of our simulations was prescribed using the simulated vegetation averaged for the same decade from this transient climate run, on which we superimposed crop area as given in the CMIP5 historic and future land use maps (Hurtt et al., 2009; Riahi et al., 2007).
We performed a 9-year (2001-2009) control simulation for present-day climate conditions initialized from a centennial transient climate simulation with ocean couplings (Jones et al., 2011). We analysed the last 8 years of the simulation, as the first year of simulation was used as spin-up. A single year is considered sufficient for spin-up because one year is around five times longer than the lifetime of the longest lived atmospheric species (with the exclusion of methane) involved in O3 chemistry. The control simulation was driven by anthropogenic and wildfire emissions of trace gases and aerosols via historical scenarios (Global Fire Emissions Database GFEDv2; Lamarque et al., 2010; Van der Werf et al., 2006) of anthropogenic and wildfire emissions.

HadGEM2 is able to reproduce the main spatial distribution of surface temperature (Figure S1) and precipitation (Figure S2). Surface temperature simulated with HadGEM2 exhibits a bias in the region of up to 2°C colder than in the observations over the Amazon forest. Simulated precipitation rate is in reasonable agreement with observations. The model is able to reproduce the main features of the seasonal cycle of precipitation, but tends to simulate less precipitation in September and November than the observations (Figure S03).

Simulated HadGEM2 NPP is compared against a meta-analysis of field data from the Ecosystem Model Data Model Intercomparison project (EMDI) (Olson et al., 2001). Measurements from the 81 ‘class A’ ("well documented and intensively studied") sites, representative of all major global biomes, are compared against our simulations. Traditionally, global vegetation models underestimate NPP in tropical ecosystems, and tend towards an asymptote of ~1000 g C m⁻² (Prentice et al., 2007). HadGEM2 is able to reproduce the main geographical variations of NPP globally (Figure S4), especially in the Northern Hemisphere, where more plentiful observations are available. In addition HadGEM2 is able to better simulate higher tropical NPP.

Ozone concentration simulated with HadGEM2-UKCA-ExtTC agrees better with observations at higher altitudes and higher latitudes (Figure S5). The model performs more poorly than the ACCENT mean over tropical areas, especially closer to the surface. Comparison with a selection of observed profiles of O₃ concentration shows the model overestimates O₃ for some locations but is in extremely good agreement for others. Over the tropics the agreement is better in the few continental profiles than the marine environment (Figure S6). Some differences may be expected given that the observations are from campaigns with specific meteorological conditions, while the model simulations represent a multi-year mean from the model. Comparison with a selection of surface O₃...
observations (Figure S7) confirms again how the model shows a better agreement with observations taken at higher latitudes.

We also perform 10 experiments that differ from the control simulation in terms of assumed biomass burning emissions, i.e. biomass burning emissions over South America are either increased or decreased by +/-20, 40, 60, 80, 100%, while emissions over the rest of the world are kept unchanged. The vegetation distribution was not adjusted for loss of vegetation due to fire. We define biomass burning emissions as those from wildfires, deforestation fires, agricultural forest burning, residential and commercial combustion, including fuel wood burning, charcoal production and biofuel combustion for cooking and heating (Lamarque et al., 2010). The dominant fire types in South America are from deforestation and degradation fires in an arc around Amazonia, with some regional hotspots of agricultural burning (see Figure 13 in Van der Werf et al., 2010). Between 2001 and 2009 the percentage contribution to annual fire emissions from fire types (deforestation and degradation, grassland and savanna, woodland, forest, agriculture) are (59%, 22%, 10%, 8%, 2%) over Southern Hemisphere South America (Figure 13 van der Werf et al., 2010), with minor differences in this region between this dataset (Global Fire Emissions Database GFEDv3) and the earlier GFEDv2 used in this study (see Fig. 16 in Van der Werf et al., 2010). The residential and commercial combustion contribution accounts for 1 and 8% of the total annual biomass burning emissions of CO and NOx respectively.

This set of experiments allows us to simulate the impact of biomass burning on surface O3 and vegetation productivity. The control simulation was also used to evaluate surface O3 mixing ratios against measurements over the Amazon forest.

**Model site-level Evaluation**

Over the data-sparse Amazonian region, comprehensive spatial data sets of surface O3 concentration are extremely limited. We evaluated simulated surface O3 against observations from two sites that have full annual analyses of O3 concentration: Porto Velho (Brazil; 8.69°S; 63.87°W) and site ZF2 in the Cuieiras forest reserve (Brazil; 2.59°S; 60.21°W). O3 mixing ratios were measured with a UV absorption analyser (Thermo 49i, USA). Observations from both sites have an estimated 4% uncertainty, considering zero noise, zero and span drifts reported in the instrument manual, and the frequency of zero and span checks performed along the experiments.
The Porto Velho sampling site is located in a forest reserve about 5 km NE (generally upwind) from the city of Porto Velho. Large land use change and regional biomass burning makes its atmospheric conditions characteristic of those of the Amazon forest with significant human interference (Brito et al., 2014). The whole region of Porto Velho has been subject to land use change since the 1980s. In Porto Velho, the dry season is from June to October and the wet season from November to May. Measurements of surface O\(_3\) mixing ratios were taken from November 2011 to October 2012 in a forest clearance, at 5 m a.g.l.

The Cuieiras forest reserve in Central Amazonia encloses 380 km\(^2\) of pristine tropical rainforest. The reserve is located in the central Amazon Basin, 60 km NNW of downtown Manaus and 40 km from the metropolis margins. This site is relatively undisturbed, as no biomass burning occurs in the forest reserve. Here rain showers are frequent with a short dry season from July to October. Measurements were taken at 39 m a.g.l. at the TT34 tower. The forest canopy height near the tower varied between 30 and 35 m, and the site is described in Martin et al. (2010), Rizzo et al. (2013) and Artaxo et al. (2013). Most of the time, the prevailing trade winds blow over 2000 km of the intact tropical forest before reaching the measurement tower. However, the site was also affected by regional transport of pollutants, either from biomass burning or urban sources (Rizzo et al., 2013). Measurements of surface O\(_3\) mixing ratios were taken from April 2010 to June 2012, with the exclusion of a few months due to instrument maintenance.

We compared simulated (averaged over 8 years of simulations) against observed average O\(_3\) diurnal cycles at each site for each available month. The model overestimates observed monthly averaged hourly O\(_3\) mixing ratios at the surface by about 5-15 ppb for all months at the Porto Velho site, but it reproduces the diurnal and seasonal cycle, including those months affected by biomass burning, i.e. August and September, at the Porto Velho site (Figure 1). The model is able to reproduce the diurnal cycle, including magnitude, at the ZF2 site for about 8 months out of 24. The model overestimates surface monthly averaged hourly O\(_3\) mixing ratios by about 5-10 ppb for the rest of months, which are also the months with lower surface O\(_3\) mixing ratios (Figure 2).

**Results**
Our analysis is focused on the region enclosed in the red rectangle in figure 3, this is a highly vegetated region with homogeneous topography, and it includes the two sites used for the model evaluation (Porto Velho and ZF2 in the Cuieiras forest reserve). This region of analysis is covered by two PFTs in HadGEM2: broadleaf trees, which is the predominant, and C3 grass (Figure 3).

Surface O3 mixing ratios simulated with HadGEM2 are higher during the months of August, September and October over the Amazon forest, and in particular over our region of analysis, because of the higher biomass burning emissions in the model during these months. Monthly average surface O3 mixing ratios in our control simulation peaks at 55-60 ppb in this region (Figure 4), while the average over the region of analysis is peaked at about 30 ppb in August and September, less in October (Figure 5a, black line).

Monthly total Net Primary Productivity (NPP) in our control simulation reaches its minimum during the months of August and September (Figure 5b, black line), at about 300 TgC/month, corresponding to the end of the dry season.

Decreasing biomass burning emissions over South America by -20%, -40%, -60%, -80%, -100% decreases surface O3 mixing ratios and increases net productivity. Vice versa, increasing biomass burning emissions over South America by +20%, +40%, +60%, +80%, +100% increases surface O3 mixing ratios over the region of analysis and subsequently reduces net productivity because of O3 damage on vegetation (Figure 5c).

These sensitivity tests suggest that decreasing biomass burning emission by 100% over South America brings monthly mean surface O3 mixing ratios averaged over the region of analysis to about the observed 15 ppb for each month (Figure 5a, dark blue line), even during the dry season, with no values over 35 ppb for any grid-cell (Figure 6). Increasing biomass burning emissions by 100% suggests that monthly mean mixing ratios of surface O3 averaged over the region of analysis reach 40 ppb in August (Figure 5a), with peaks of about 65-70 ppb in some grid-cells (Figure 6a).

For both increases and decreases of between 20 and 80% in South American biomass burning the model simulates almost linear changes in surface O3 mixing ratios (Figure 6, the figure shows increases and reductions by 40, 60 and 100%).

Suppressing biomass burning emissions (i.e. decreasing biomass burning emission by 100%) over South America increases total NPP over the region of analysis by about 15%, to about 350-370 TgC/month, with peak increases of 60% for a few grid-cells, in August and September (Figure 6b):
this quantifies the impact of present-day biomass burning on vegetation productivity. When increasing biomass burning emissions over South America by 100%, monthly total NPP over the region of analysis is reduced by about 10%, i.e. to about 250 TgC/month, in August and September (Figure 5b), with peak values of 50-60% reductions for few grid-cells (Figure 6c). For reductions by 20 to 80% in South American biomass burning the model varies NPP almost linearly (Figure 5c). However, the increase in South American biomass burning by 20 to 80% determine a very similar decrease in NPP, e.g. between 7 and 10% decrease in August (Figure 5c). Both increasing and reducing South American biomass burning from 20 to 80% increases the number of grid-cells where a significant variation of NPP takes place (Figure 6b). The percentages given above are significant against inter-annual variability in the control simulation, i.e. we only take into account of the variations above one standard deviation in the control simulation. We also exclude from our analysis the grid-cells with low productivity, i.e. where NPP in the control simulation is below 50 gC/m²/month (i.e. we focus on high productivity regions, e.g. forests).

Discussion and Conclusions

The HadGEM2 model overestimates the magnitude of the O₃ diurnal cycle at the two sites used in the evaluation. Overestimation of simulated O₃ in the Amazonian boundary layer has been observed in other modelling studies, especially in clean air conditions (Bela et al., 2014). Nonetheless, our model reproduces the main features of the diurnal and seasonal cycle. In particular, the model is able to reproduce the increase in surface O₃ during the biomass burning season.

As stated earlier in the model description section, biomass burning emissions are prescribed as monthly mean and kept constant during the day, and this can have an impact on the hourly and day-to-day variation of surface O₃. For example, O₃ production will respond differently if biomass burning emissions occur during the day or at night, affecting simulated surface O₃ mixing ratios. These issues can be improved by modelling fire and biomass burning emissions interactively. The inclusion of an interactive fire model in HadGEM is currently under development.

The model overestimates surface O₃ mixing ratios by 5-15 ppb for several months at the ZF2 site in the Cuieiras forest reserve and for all available months at the Porto Velho site. The reasons for these systematic biases in surface O₃ mixing ratio are likely manifold. In a complex, highly coupled
system such as the HadGEM2 Earth System Model (ESM) it is not always easy to disentangle all processes and attribute model biases to specific components.

We attribute the systematic biases in the surface $O_3$ mixing ratio to the following, most likely reasons:

1. Model resolution in both the horizontal and the vertical dimension
2. Uncertainties in emissions, both magnitude, seasonality and location
3. Uncertainties in the $O_3$ dry deposition at the surface

Other factors such as uncertainties in the chemical mechanism, the photolysis rates, lightning NOx production over the area and transport of $O_3$ and precursors will certainly contribute. We will briefly discuss the three most important (in our opinion) factors that contribute to the systematic biases.

The relatively coarse resolution of a global ESM simulates mixing ratios of trace species (both trace gases and aerosols) that represent averages over large areas. This issue has been discussed previously in the literature, mostly in relation to air quality modelling (see, e.g., Valari and Menut, 2008; Tie et al., 2010; Appel et al. 2011; Thompson and Selin, 2012). In our case one grid box equals approximately 30,000 km$^2$ (i.e., 200x150 km$^2$ in longitude and latitude). The implicit averaging pertains both to emission and concentration fields; the predominant consequence is a dilution in each grid-cell. Depending on the chemical regime, this can lead to reduced or enhanced net $O_3$ production. Additionally, HadGEM2-ES has a relatively coarse vertical resolution. HadGEM2-ES has a lowest model layer depth of 40m (global average) and the vertical profile of $O_3$ will undoubtedly show a gradient as the loss mechanism for $O_3$ is dominated by the surface (e.g. Colbeck and Harrison, 1967). The measurement level may explain part of the model overestimation, since it is well known that $O_3$ mixing ratios strongly decrease with height due to deposition within the canopy. The lowest layer of the model has a midpoint height 20 metres above the displacement height for the particular gridbox (generally approximated as 2/3 of the average height of the obstacle, in this case the canopy), while measurements were taken at 5 m and 39 m a.g.l., respectively, at Porto Velho and ZF2 which are located either in or just above canopy level. Rummel et al. (2007) reports a 5-15 ppb $O_3$ decrease from 52 to 11 m a.g.l. in a forest site in Amazonia. This steep gradient near the surface is due to surface deposition but also due to in-canopy chemical processing (c.f., e.g., Stroud et al., 2005; Gordon et al., 2014). The latter is not represented in HadGEM2-ES.
The remote environment of the Amazon forest is dominated by relatively high concentrations of VOC, particularly of biogenic origin, and low concentrations of nitrogen oxides, NOx. It is a NOx-limited environment. In such an environment O₃ is destroyed by reactions with BVOC (mainly isoprene and (mono-)terpenes). This destruction is more pronounced the higher the BVOC concentration becomes. Consequently, conditions in the global model are likely to differ from that of a measurement at a specific point such as those we compare to in Figures 1 and 2. It is a known problem in model evaluation.

Another issue related to model resolution, when comparing global models to point-like observations, is the uncertainty in global emission inventories, both with respect to magnitude and location. In particular the latter will result in discrepancies between modelled concentrations of O₃ and its precursors and point-like observations. But the uncertainties in emission magnitude are also substantial and can reach a factor of two or more in case of biogenic VOC (e.g., Guenther et al., 2006; Arneth et al., 2008, 2011; Pacifico et al., 2011, 2012).

Thirdly, and again related to model resolution, is the representation of O₃ dry deposition at the surface. Its magnitude and diurnal cycle will depend on boundary layer turbulence, surface roughness, land surface type, vegetation type, soil moisture, photosynthetic activity, and more. In a recent sensitivity study by Folberth et. al (in preparation) O₃ surface concentrations showed the largest sensitivity to perturbations in O₃ surface dry deposition fluxes. Underestimating O₃ surface dry deposition, in particular during the night preventing a complete flush of the PBL with respect to O₃, will lead to systematic biases.

A comparison with Rummel et al. (2007) indicates that ozone dry deposition velocities on average compare favourably with observations. Rummel at al. (2007) reported day-time velocities of up to 2 cm/s and night-time velocities of typically around 0.5 cm/s during the wet season and velocities between 0.3 cm/s and 1.0 cm/s during day-time and 0.3 cm/s and 0.8 cm/s during the dry season for one site in the Amazon region. HadGEM2-ES predicts annual mean O₃ deposition velocities of 0.5 to 0.6 cm/s (see Figure S8) in fair agreement with the observations. Furthermore, the model is able to capture well the variability between the wet season and the dry season. However, more data are needed to conduct a robust evaluation, but, this admittedly crude comparison is sufficient to demonstrate a basic capability of HadGEM2-ES to reproduce observed ozone deposition velocities in the Amazon region to a reasonable degree.
Interestingly, however, the latter process may also represent a redeeming feature of the model. According to our model of O$_3$ plant damage it is the total O$_3$ flux into the plant that determines the amount of damage caused to the photosynthetic activity and, hence, carbon assimilation. However, the total O$_3$ flux (or dose) is a function of both O$_3$ surface concentrations and dry deposition, i.e. for plants there is a compensation effect when concentrations are overestimated while deposition velocities are underestimated. Underestimating the O$_3$ dry deposition flux implies reduced O$_3$ plant uptake, and consequently an underestimation of the plant damage and productivity losses. However, it also leads to higher O$_3$ concentrations, which subsequently act to increase plant O$_3$ uptake and damage, compensating for the initial effects on productivity. Still, a detailed assessment and quantification of this interdependence of O$_3$ concentration and dry deposition fluxes is beyond the scope of this study and must be referred to future research.

August, September and October are the months when biomass burning and surface O$_3$ concentrations are higher over the Amazon forest, but also the months when plant productivity is at its lowest which will tend to suppress the impact of O$_3$ damage on plant productivity. This is because stomatal conductance is reduced due to water limitations (also accounted for in the model) during the dry season, thus reducing the flux of both carbon dioxide and O$_3$ into the leaves, and consequently reducing O$_3$ plant damage.

Ashmore (2005) noted how O$_3$ exposure is poorly correlated with flux into leaves and also the potential for damagingsly high O$_3$ fluxes in leaves at concentrations significantly below 40 ppb at maximum stomatal conductance. Consequently, global vegetation models as used in this study have adopted flux-based parameterizations to represent O$_3$ impacts on vegetation, moving away from application of the earlier exposure based metrics, e.g. accumulated O$_3$ exposure above a threshold of 40 ppb, AOT40.

The parameterization of O$_3$ damage used in this study is calibrated for high-latitude vegetation. Unfortunately data for calibrating this O$_3$ damage scheme for tropical vegetation are currently not available and observations of O$_3$ damage in the Amazon forest are very limited. Observations of O$_3$ damage on tropical forests are urgently needed, including observations at moderate (e.g. 20-30 ppb) and high surface O$_3$ mixing ratios.

The simulated impact of present-day biomass burning on vegetation productivity over our area of analysis is about 230 TgC/yr (i.e. the difference between the dark blue line and the black line in Fig.
5b) using the “high” sensitivity mode in the O₃ damage scheme. Taking into account that the uncertainty in these estimates is substantial, this O₃ damage impact over the Amazon forest is of the same order of magnitude as the release of CO₂ due to land fire in South America, as quantified in van der Werf et al., (2010; 293 TgC/yr from table 7 of that paper); in effect to potentially double the impact of biomass burning on the CO₂ fluxes. This highlights the urgent need for more tropical data on plant O₃ damage to better constrain estimates.

Despite overestimating surface O₃ mixing ratios our model simulates only a moderate reduction in NPP associated with elevated O₃ due to biomass burning emissions. Given that our model systematically overestimates O₃ mixing ratio, assuming accurate dry deposition, and that we use our model in the high sensitivity mode, our simulations where we increase biomass burning emissions by 100% suggest a maximum 10% average reduction in monthly plant productivity, and peak reductions of 50-60% reductions in few grid-cells. This is because, despite the increase in biomass burning, monthly average surface O₃ mixing ratios do not exceed a moderate 40 ppb. Moreover, our model does not include deforestation due to fire, which would reduce vegetation cover when increasing biomass burning emissions in our sensitivity experiments, reducing NPP, and BVOC emissions, further. However, local and daily/hourly impact of O₃ damage on plant productivity can be higher.

Estimates of the magnitude of the reduction in plant productivity due to O₃ damage can be improved with additional field studies and improving the representation of tropospheric O₃ in ESMs (sources, chemistry and sinks). Nevertheless, considering these processes in a coupled system can provide an improvement in robustness of conclusions, as e.g. it can treat processes with a specific diurnal cycle, like photosynthesis and surface O₃, interactively on a short time scale (e.g. half an hour in our model).
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Figures

1. Comparison of measured (dots) and simulated (stars) monthly averaged diurnal cycle of surface O$_3$ mixing ratios at the Porto Velho site, including measured day-to-day variability (grey lines) and standard deviation (dashed lines) for the model results. The measurements have an uncertainty of 4%.

2. Comparison of measured (dots) and simulated (stars) monthly averaged diurnal cycle of surface O$_3$ mixing ratios at the ZF2 site in the Cuieiras forest reserve, including measured day-to-day variability (grey lines) and standard deviation (dashed lines) for the model results. The measurements have an uncertainty of 4%. We show one of the two available years of observations. Legend as in Figure 1.

3. Vegetation cover in HadGEM2 for the month of September. The red rectangle is our region of analysis. The two sites used in the model evaluation (the sites of Porto Velho and ZF2 site in the Cuieiras forest reserve) are also marked.

4. Monthly average surface O$_3$ mixing ratio simulated with HadGEM2 for the month of September (average over 8 years of simulations).

5. Clockwise from the top-left: (a) Simulated monthly surface O$_3$ mixing ratios; (b) Simulated monthly total NPP; (c) Simulated monthly variation in total NPP. The plots show the results for the control simulation (i.e. using the decadal mean biomass burning emissions from Lamarque et al. (2010) centered on year 2000; 2000 BB emissions) and the various experiments with increased (+) or decreased (-) biomass burning emissions over South America by 20, 40, 60, 80 and 100%. All data are averaged over the region of analysis for 8 years of simulations.

6. From the left: simulated variation in surface O$_3$ mixing ratios and NPP over the region of analysis for the months of August, September and October.

7. Probability density function (histogram) of the variation in NPP for the same months. The plots show the variation between the experiments with South American biomass burning increased/decreased by 40, 60 and 100% and the control simulation.
Figure 1
Figure 2

ZF2 Cuieras forest (2.59°S, 60.21°W)
Figure 3
Figure 5
Figure 7