1 Sensitivity of tropospheric loads and lifetimes of short

2 lived pollutants to fire emissions

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8 Abstract

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The capability of global Chemistry and Transport Models (CTMs) to simulate atmospheric composition and its spatial and temporal changes highly relies on the input data used by the models, in particular the emission inventories. Biomass burning emissions show large spatial, diurnal, seasonal and year-to-year variability. In the present study, we applied a global 3D CTM to evaluate uncertainties in the computed atmospheric composition associated with the use of different biomass burning emissions and identify areas where observational data can help to reduce these uncertainties. We find the emission inventory choice to lead to regional differences in the calculated load of aerosols up to a factor of 4. Assumptions on the injection height of the biomass burning emissions are found to produce regionally up to 30% differences in the calculated tropospheric lifetimes of pollutants. Computed changes in lifetimes point to a strong chemical feedback mechanism between emissions from biomass burning and isoprene emissions from vegetation that are linked via NO_x-driven oxidant chemistry, NO_x-dependent changes in isoprene oxidation products, aerosol emissions and atmospheric transport. These interactions reduce isoprene load in the presence of biomass burning emissions by 15%, calculated for the same amount of isoprene emitted into the troposphere. Thus, isoprene load and lifetime are inversely related to the quantities of pollutants emitted by biomass burning. This feedback is shown to be able to increase the global annual secondary aerosol yield from isoprene emissions, defined as the ratio of tropospheric loads of secondary aerosol from isoprene oxidation to isoprene emissions, by up to 18%.

1 Introduction

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2 Atmospheric composition is affected by emissions of reactive gases and aerosols to the atmosphere by several natural (e.g. soils, vegetation, oceans, volcanoes, wild fires) and 3 anthropogenic sources (e.g. industrial and residential activities, transport, and shipping). 4 5 Among these sources biomass burning plays a central role for atmospheric chemistry via 6 changes in the atmospheric composition but also impacting on the ecosystem functioning 7 through atmospheric deposition of nutrients and the lifecycle of vegetation (Keywood et al., 8 2013). Biomass burning is positioned between the natural (wild fires) and human-induced 9 (intentional burning) sources of atmospheric pollutants since a fraction of open fires is 10 induced by humans for agricultural and city expansion purposes (Levine et al., 1995) or for 11 protection against fire itself (Mutch, 1994). Biomass burning is an important source of trace 12 constituents to the atmosphere including radiatively and chemically reactive gases and aerosols (Andreae and Merlet, 2001; Akagi et al., 2011). It is the largest source of primary 13 14 carbonaceous aerosols (Bond et al., 2004) and the second largest source of volatile organic 15 compounds (VOC) in the atmosphere after the emissions from vegetation (Guenther et al., 2012) and of carbon monoxide (CO) after anthropogenic emissions (Kanakidou and Crutzen, 16 1999; Pfister et al., 2005). 17 18 Emissions from biomass burning and their transformation in the atmosphere affect air quality 19 (Lelieveld et al., 2004), interact with radiation (Reid et al., 2005) and the atmospheric water 20 cycle and thus affect climate (Rosenfeld, 1999). In turn climate change is seen to impact on 21 wild fire occurrence and intensity. For instance the exceptionally intensive 1997/1998 22 Indonesia fires have been attributed to the combined strength of the El Niño and the Indian 23 Ocean Dipole (Field et al., 2009). 24 Significant changes in the trends of atmospheric concentrations of CH₄ and CO have been 25 attributed to the changes in the biomass burning emissions (Simmonds et al., 2005). Most of 26 these emissions occur in the tropics that are subject to intensive photochemistry in the 27 presence of high humidity conditions and significant convective activities (Chatfield and 28 Delany, 1990; Crutzen, 1994). During summer in the high latitudes boreal forest fires 29 contribute about 12% to the global biomass burning emissions (Lavoué et al., 2000) and can 30 be so intensive and convective that their emissions reach the high troposphere and low 31 stratosphere (Fromm et al., 2000).

1 Tropical photochemistry is controlling the lifetime of most atmospheric pollutants (Crutzen, 2 1994; Keywood et al., 2013), including reactive greenhouse gases like methane (CH₄) and 3 ozone (O_3) , and thus their persistence in the atmosphere to impact on radiation and climate. 4 Up to about 25% of the net global photochemical production of tropospheric ozone has been 5 attributed to biomass burning emissions and chemistry in the atmosphere (Crutzen and 6 Andreae, 1990; Jaffe and Wigder, 2012). Long range transport of biomass burning aerosols 7 has been seen to happen fast within one or two weeks both downwind tropical (Edwards et al., 8 2006; Dirksen et al., 2009) and high latitude sources (Jaffe et al., 2004). Thus this source is 9 affecting atmospheric pollutant levels in remote environments. For instance, chemical ageing 10 of fire plumes has been identified as contributor to the high ozone over the Atlantic ocean 11 (Lelieveld et al., 2004). Therefore it is important to simulate the impact of biomass burning 12 emissions on tropospheric composition and pollutant lifetimes and to evaluate the 13 uncertainties in such simulations. 14 Several biomass burning emission inventories have been constructed based on burned area, active fire detections, and plant productivity from satellite observations (van der Werf et al., 15 16 2010) or on assimilated Fire Radiative Power derived from satellite observations (Kaiser et 17 al., 2012) and experimentally determined pollutant emission factors (Andreae and Merlet, 18 2001) and assumptions on the state of burning of the biomass (smoldering or flaming, van der 19 Werf et al. (2006)). All these factors introduce uncertainties in the emissions (Granier et al., 20 2011; Wiedinmyer et al., 2011). In particular, the size of small fires can be overestimated and 21 the number of fires can be underestimated when seen by satellites (Wiedinmyer et al., 2011). The injection height of fire emissions (Dentener et al., 2006; Freitas et al., 2007; Sofiev et al., 22 23 2012) is an additional cause of discrepancies in the model estimates of the impact of these 24 fires on tropospheric composition. The height distribution proposed by Dentener et al. (2006) 25 (used in this work) is based on wildfire location and type, where the distribution described in 26 Sofiev et al. (2012) is based on the fire characteristics (fire intensity, temperature of plume, 27 type of source) as well as the meteorological conditions (atmospheric boundary layer height, 28 free troposphere). These two approaches show similarities in emission heights over North 29 America and Oceania, but over Eurasia, Australia and South America the two methods show 30 significant differences (Sofiev et al., 2013). A plume height climatology over North America has been also derived by analysis of 5-year satellite observations by MISR (Val Martin et al., 31 32 2010) which compared to the Dentener et al (2006) vertical distribution of fires there (2000-

1 6000 meters) shows lower mean injection heights (500-1500 meters) for boreal fires but is in 2 agreement for temperate and tropical fires. Plume rise models evaluated against that 3 climatology have been shown to underestimate the observed plume heights (Val Martin et al., 4 2012). Guan et al. (2008) using the NCAR CAM3.1 model found that the calculated CO 5 concentrations downwind biomass burning emission areas, can increase by up to 150 ppb 6 depending on the assumptions in the injection height of the emissions. Boreal forest fire 7 emissions occurring high in the troposphere have been detected by Colarco et al. (2004) to be 8 transported from Canada to Washington D.C. in the U.S.A. where they have been mixed with 9 boundary layer air. Long range transport of biomass burning pollutants has been followed by 10 lidar and satellite observations and the simulations have been shown to be sensitive to the 11 injection height of the emissions as well as to the entrainment of air into the boundary layer 12 over U.S.A. Note that boreal fires plumes can reach the upper troposphere where their impact 13 is different from that in the boundary layer due to the non-linearities in the atmospheric 14 chemistry (Chatfield and Delany, 1990) and the different photochemical conditions there. 15 Leung et al. (2007) global modeling study of the impact of boreal fire emissions on air 16 pollutants levels, found a much larger enhancement in ozone when about half the emissions 17 were released above the boundary layer than when all emissions were occurring in the boundary layer. They attributed these differences to the role of peroxyacetyl nitrate (PAN) as 18 19 carrier of NO_x downwind burning areas. Jaffe et al. (2004) found that the intensive Siberian 20 fires in 2003 enhanced the background ozone over the Pacific Norhwest U.S.A., resulting to 21 exceedance of ozone air quality standard. Hodzic et al. (2006) studying AOT over Europe 22 during the 2003 Portuguese fires identified high altitude transport of smoke particles from 23 Portugal to The Netherlands, that has been both observed by POLDER-2 and simulated by the CHIMERE model. Williams et al. (2012) simulated the African fires in 2005 using the TM4 24 25 model and three different biomass burning emission inventories, two global and one regional. 26 They calculated differences in the ozone global burden resulting from the use of different 27 biomass burning inventories that range between +1.7% and +4.6% compared to the simulation using GFEDv3 biomass burning emission inventory. 28 29 The present study aims to evaluate uncertainties in model estimates of biomass burning 30 impacts on atmospheric composition that are associated with the use of different emission inventories in the same model. The study also aims to identify locations where additional 31 32 observations can provide constrains for biomass burning emission estimates. For this purpose

- a global 3D Chemistry and Transport Model (CTM) is applied to evaluate uncertainties in the
- 2 atmospheric composition and major pollutants lifetimes computed using recently updated and
- 3 commonly used biomass burning emissions. Based on the computed model sensitivity to
- 4 biomass burning emissions, we also identify areas where observational data can help to
- 5 reduce these uncertainties.

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2 Model Description

- 8 The model used for this study is the global 3-D CTM TM4-ECPL (Kanakidou et al., 2012).
- 9 The model accounts for gas and multiphase chemistry to describe tropospheric ozone
- 10 chemistry and all major aerosol components (primary and secondary). It contains explicit
- 11 chemistry of C₁ to C₅ volatile organic compounds (VOCs) and a highly simplified
- representation of a-pinene and β -pinene chemistry. The model calculates secondary organic
- 13 aerosol (SOA) formation by VOC oxidation and subsequent gas-to-particle partitioning of
- semivolatile products (Tsigaridis and Kanakidou (2007) as updated by Myriokefalitakis et al.
- 15 (2010)). Chemical aging of organic aerosol (OA) is also taken into account. For primary
- organic aerosol (POA) and black carbon (BC) chemical ageing is considered to occur by
- oxidation of organic material that coats the particles and is driven by O₃ (Tsigaridis and
- 18 Kanakidou, 2003); while for SOA chemical ageing to non-volatile SOA (Tsigaridis and
- 19 Kanakidou, 2003) is considered to occur by reaction with OH at the rate of 4.10⁻¹² molec
- 20 ¹cm³s⁻¹, very close to that of the H-abstraction reaction of pinonic acid with OH (Praplan et
- 21 al., 2012). BC emissions are by 20% soluble while terrestrial POA emissions are by 50%
- soluble. For both BC and POA the insoluble fraction is converted to soluble during aging.
- 23 Multiphase chemical production of SOA is parameterized as described in Myriokefalitakis et
- 24 al. (2011). Gas-to-particle partitioning of inorganic components is solved using the
- 25 ISORROPIA II aerosol thermodynamic model that also calculates the aerosol-water (Nenes et
- al., 1998; Fountoukis and Nenes, 2007). For this study the TM4-ECPL model uses a 3°x2°
- 27 longitude-latitude grid and 34 hybrid levels up to 0.1 hPa (with the first 4 model vertical
- 28 layers between surface and 900 hPa) and is driven by the European Centre for Medium-range
- Weather Forecasts (ECMWF) ERA-Interim meteorological data (Dee et al., 2011) for the year
- 30 2008 for all the sensitivity simulations.

2.1 Natural emissions

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- 2 Isoprene, terpenes and biogenic volatile organic compounds (BVOC) emissions in the TM4-
- 3 ECPL model are taken from the MEGAN-MACC inventory (Sindelarova et al., 2014) for the
- 4 year 2008, which is a product of the MEGANv2.1 model (Guenther et al., 2012). Dust
- 5 emissions are from AeroCom (Aerosol Comparisons between Observations and Models;
- 6 (Dentener et al., 2006) calculated for the year 2008 by E. Vignati (personal communication,
- 7 2011). Marine emissions of sea-salt aerosols and organic gases and aerosols are calculated
- 8 online driven by meteorology and sea water productivity as described by Myriokefalitakis et
- 9 al. (2010) and Vignati et al. (2010).

2.2 Anthropogenic emissions

- Anthropogenic emissions used for this experiment are the ECLIPSE (Evaluating the CLimate
- and Air Quality ImPacts of Short-livEd Pollutants) version 4.0 emissions (Klimont et al.,
- 13 2013), available in 0.5°x0.5° spatial resolution. The ECLIPSE anthropogenic inventory was
- 14 initially provided as sectoral including the agricultural waste burning sector (AWB). Since
- 15 AWB is either included in the anthropogenic emissions or in the biomass burning emissions,
- 16 caution was taken to avoid double counting of the emissions. For this, the AWB emissions
- 17 (Table 3) are considered separately for the simulations that have been performed for this study
- 18 (Table 4). The AWB in the ECLIPSE database amounts to 4.5% of the total anthropogenic
- 19 pollutants emissions (approximately 34.5 Tg a⁻¹) for the year 2008 (see Table 1 for more
- 20 information). Anthropogenic emissions of all basic pollutants are used (CO, nitrogen oxides
- 21 (NO_x), black carbon aerosol (BC), particulate organic carbon (OC), sulfur dioxide and sulfates
- (SO_x) as well as speciated non methane volatile organic compounds (NMVOCs; for a list of
- 23 the NMVOCs used in the model see supplementary material S1).

2.3 Biomass burning emissions

- 25 For the present study a number of sensitivity simulations have been performed (Table 4) using
- different biomass burning emissions (Table 2) and AWB emissions (Table 3), all for the year
- 27 2008. For the base simulation (S0.0), the biomass burning emissions from the Global Fire
- 28 Emission Database v 3.1 (GFEDv3; van der Werf et al. (2010)) are used, excluding the AWB
- sector (Table 3), hereafter called GFEDv3-ECLIPSE biomass burning emissions (S0.X),
- 30 while AWB emissions are taken from the ECLIPSE anthropogenic emissions developed in the

framework of the ECLIPSE project. Additional simulations have been performed (Table 4) using both biomass burning and AWB emissions from the GFEDv3 (van der Werf et al., 2010) (S1.X), as well as AWB from ECLIPSE and biomass burning emissions from the Atmospheric Chemistry and Climate Model Intercomparison Project's (ACCMIP; Lamarque et al. (2013); http://ecaad.sedoo.fr) (S2.X) or from the Fire INventory from NCAR (FINN; Wiedinmyer et al. (2011) http://bai.acd.ucar.edu/Data/fire/) (S3.X) and finally a simulation where no biomass burning emissions were taken into account (S4.0). Since the injection height of these emissions contributes to the uncertainty of the model results, biomass burning emissions are considered in the model either to be injected at heights following Dentener et al. (2006), or to be emitted solely in the lowest model layer (see list of simulations in Table 4). The temporal variability of theses biomass burning inventories per emitted species for 2008 is shown in Fig. 1. This figure depicts the differences between the inventories in their seasonality and amplitude (also annual totals in Table 2); while Fig S2 in the supplementary material shows spatial difference in the annual BC emissions between the inventories. The ACCMIP inventory shows the largest magnitude in the temporal variation of these emissions. All inventories show a July-Sept. primary maximum while they differ in the secondary maximum between Jan and April. The AWB emissions that are not included in the GFEDv3-ECLIPSE biomass burning inventory significantly contribute to NMVOC and NH₃ emissions during spring and summer.

3 Experiment setup

The impact of the use of different biomass burning emission inventories to the calculated tropospheric loads and lifetimes of the main pollutants and the sensitivity of the model results to the wild fire emissions have been evaluated based on nine different simulations. For all simulations the model setup was exactly the same, except for the biomass burning emissions inventory used and its vertical distribution application. A summary of the simulations here performed is provided in Table 4. The GFEDv3-ECLIPSE inventory and height distribution for biomass burning emissions have been used as the base case scenario (S0.0). All scenarios named SX.0 assume the same fractional height distribution of the emissions according to Dentener et al. (2006) where all the scenarios named SX.1 assume all open biomass burning emissions to occur at surface. For scenario S4.0, open biomass burning emissions are set to zero. Note that we have chosen to account for monthly mean emissions since not all

- 1 inventories have higher temporal resolution. This is the reason we have also chosen to
- 2 validate the model results comparing to monthly mean observations.

3 4 Results

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- 4 To evaluate the ability of the model to reproduce the observations, the computed
- 5 concentrations are compared with measurements. The differences in the fields computed by
- 6 the various emission inventories provide a measure for the robustness of the model results
- 7 with regard to the biomass burning impacts. Comparison of the simulated tropospheric
- 8 concentrations of pollutants between the various scenarios reveals the spatial and temporal
- 9 differences due to the different inventories and could indicate which inventory is performing
- 10 the best. Ultimately these differences will point to areas where additional observations can
- 11 contribute to reduce uncertainties of the emission inventories as will be further discussed.
- Finally, tropospheric lifetimes are calculated to provide information on how the location and
- strength of the emissions affect the persistence of the pollutants in the atmosphere.

4.1 Comparison with ground measurements

- 15 Surface observations of Ozone from the European Monitoring and Evaluation Programme
- 16 (EMEP) monitoring network (Europe), Ozone and CO observations from the World Data
- 17 Centre for Greenhouse Gases (WDCGG) database (Global) and particulate Organic Carbon
- 18 (OC) observations from the Aerosol Comparisons between Observations and Models
- 19 (AeroCom) phase II database (Global) (Tsigaridis et al., 2014) have been used for the model
- evaluation. The locations of measurements are shown in Fig. S1 in the supplement. While all
- 21 available data have been used for model evaluation, only comparisons at stations that have
- been selected to make evident differences between the simulations using different biomass
- burning emission inventories are shown for OC (Fig. 2), CO (Fig. 3) and O₃ (Fig. 4).
- 24 Concentration fields of primary pollutants emitted by biomass burning are more strongly
- 25 affected by the different emission inventories and injection heights. Thus, OC computed
- 26 concentrations (Fig. 2) and BC concentrations (not shown) present the largest diversity,
- 27 between simulations followed by CO (Fig. 3), which is emitted by fires, but has also
- secondary sources.
- 29 The simulated OC for the various scenarios and their differences from the observations in the
- 30 tropics, the subtropics and high latitudes at locations affected by biomass burning emissions

1 are shown in Fig. 2. Due to limited observational data from the tropics where most of the 2 biomass burning occurs, for the following comparisons all available data have been used 3 independent of the year. Modeled differences for OC due to emission inventory choice can 4 exceed a factor of three at Alta Floresta (Fig. 2c) and eight at Rondonia (Fig. 2d) during the 5 biomass burning months. Using the ACCMIP inventory the largest OC levels are computed at 6 the tropical station of Alta Floresta in August and September, whereas the GFEDv3-ECLIPSE 7 and GFEDv3 inventories include large amounts of OC injections at the subtropical stations of 8 California in June, July and August (Fig. 2b and g). Different emission inventories 9 significantly affect the model performance over and downwind locations where wildfires 10 occur. Unfortunately, current observational sites do not provide sufficient constraint for the 11 emission databases evaluation. 12 Tsigaridis et al. (2014) OC global model intercomparison exercise has indicated that among 13 the thirty-one models contributing to that study, some models emit all biomass burning 14 aerosols at the surface, while most models distribute them to a number of layers above the 15 surface, typically within the boundary layer. Most models are using GFEDv3 and ACCMIP 16 inventories and all models appear to have similar seasonality in primary OC emissions with 17 increased emissions during Northern Hemisphere summer due to the enhanced contribution of 18 Northern Hemisphere biomass burning emissions from temperate and boreal forests to the 19 total OC fluxes. Kaiser et al. (2012) found systematic model underestimation of smoke 20 aerosol optical depth (AOD) observed by MODIS that can be as high as a factor of 3 on the 21 global scale when emissions from bottom-up inventories like GFED are used. Petrenko et al. 22 (2012) have demonstrated that such underestimate strongly varies by region. 23 Similar to OC results are obtained for CO, as seen in Fig. 3, where during the biomass 24 burning season different quantities of CO are calculated depending on the inventory used. At 25 Yonagunijima (Fig. 3a) CO concentration differences computed using the different 26 inventories maximize in spring and models are underestimating measurements by 25%. Such 27 differences between inventories are large at the East Trout Lake station in Canada, where in 28 June and July model results differ by up to 150 ppb (a factor of 2.5). These results reflect the 29 extremely high emissions in the GFEDv3-ECLIPSE and GFEDv3 inventories for this region that are not seen in the measurements (Fig. 3b). The assumption that all emissions occur near 30 31 the surface leads to about 60% higher CO surface concentrations than when emissions are 32 distributed vertically. At the areas where biomass burning occurs and downwind of them,

- these emissions contribute between 10 and 75% to the total CO levels during the burning
- 2 season.

- 3 Comparisons of O₃ simulations with surface measurements (Fig. 4) show noticeable
- 4 difference between the simulation that neglects wildfire emissions (S4.0) and all other
- 5 simulations, at stations like Mt. Kenya (Fig. 4f), La Quiaca observatory (Fig. 4g) and Hok
- 6 Tsui (Fig. 4d), which are located in the vicinity or outflow of tropical biomass burning. These
- 7 are areas where O₃ levels are the most sensitive to the different biomass burning emission
- 8 scenarios. For instance, at La Quiana observatory (Fig. 4g), differences as high as 10 ppb of
- 9 O₃ (i.e. ~25%) are computed for October when using the different emission scenarios. The
- 10 FINN inventory results in the highest computed O₃ levels, while omitting biomass burning
- reduces O_3 levels by ~35%. However, very small sensitivity is seen between the scenarios
- with wildfire emissions for the other locations in Fig. 4. Thus, evaluating these inventories
- 13 requires densifying air quality monitoring close to the major biomass burning sources in the
- tropics, which are virtually absent. Furthermore, we have calculated the ratio of the standard
- deviation to the mean of all model simulations to identify locations where biomass burning
- emission inventories produce the largest model divergence. In Fig. 5 these ratios are shown
- 17 for OC and indicate that systematic observations over boreal regions, Alaska, South Asia and
- 18 Indonesia can help constrain the used biomass burning emission inventories.

4.2 Comparison with ozonesondes and satellite observations

- 20 Because the impact of biomass burning is not restricted to the surface concentrations of
- 21 pollutants but also extends in the free troposphere, we have also compared model results with
- 22 ozonsondes as well as with O₃ and CO mid tropospheric columns as observed by
- 23 Tropospheric Emission Spectrometer (TES) satellite instrument. In addition, simulated O₃
- 24 profiles have been compared with available ozonesondes data from WDCGG after
- 25 interpolating into layers of 50 hPa from surface to the top of the atmosphere as described in
- detail by Myriokefalikakis et al. (2015 in preparation). Figure S6 in the supplement shows
- that there is no statistical difference in the performance of the different scenarios with regard
- 27 that there is no statistical difference in the performance of the different sections with regar
- 28 to ozonesonde observations.
- 29 Similar results are obtained from the comparison of model results to the TES global survey
- data version 4 with focus on the relatively sensitive in the middle/lower free troposphere,
- 31 using data from 7 TES pressure levels between 800 and 400 hPa. The TES products are

- provided in 67 levels in vertical with a varying layer thickness (Beer et al., 2001). In order to
- 2 compare TM4-ECPL model results with the TES observations, the methods presented in
- 3 (Voulgarakis et al., 2011) have been used. Thus, the 3 hourly model outputs are sampled at
- 4 the times and locations of the TES measurements, then they are interpolate onto the 67 TES
- 5 pressure levels in vertical, and finally the TES a priori profiles and averaging kernels are
- 6 applied. The processed observational and model data are regridded to original 3°x2° in
- 7 longitude by latitude horizontal resolution in order to smooth out gaps in the observations.
- 8 More details are provided in Myriokefalitakis et al. (in preparation, 2015) where a detailed
- 9 model evaluation is presented including comparison with satellite observations.
- 10 Point-by-point comparisons of the results for the different simulations performed for the
- present study against available TES observations for all model grids on daily mean basis are
- shown in Figures S7 in the supplement. No simulation and thus no emission database stands
- out for its performance in reproducing the observations.

4.3 Tropospheric loads

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- 15 The global annual mean tropospheric loads for selected gases and aerosol components as
- 16 computed for the base case scenario (S0.0) are shown in Fig. 6 for OC, CO, NOx, O₃, OH,
- and isoprene. Fig. S3 (in the supplement) shows similar results for BC, SO₄²⁻, NO₃⁻, HNO₃
- and NH₄⁺. Although changes in the wildfire emissions do not significantly impact the global
- 19 tropospheric load of most pollutants as shown in Table 5, regionally significant differences
- are computed (e.g. for BC, the difference can reach a factor of 7, Fig. S4b) as will be further
- 21 discussed. The choice of wildfire emission inventory impacts on the calculated tropospheric
- 22 load of tracers. The most sensitive pollutants to wildfire emissions are found to be OC and
- 23 BC, while O₃ shows small sensitivity.

24 **4.3.1** Contribution of wildfires emissions on tropospheric loads.

- 25 The contribution of wildfires to the tropospheric load of pollutants can be calculated by
- 26 comparison of S0.0 (base case) with S4.0 that neglects the emissions. Wildfires increase the
- 27 tropospheric loads of: OC by ~30%, BC by ~35%, CO by about 13%, NH₄⁺ by 10%, HNO₃
- 28 by 8%, NO_x by 5%, and SO_4^{2-} and O_3 by 3% (Table 5).
- 29 Previous studies for CO with the NOAA GFDL GCTM have shown biomass burning to
- 30 contribute from 15 to 30% to the total CO background (Galanter et al., 2000). This is in

1 agreement with the measurements by Crounse et al. (2009) in central Mexico which attributed 2 21-31% of CO load to biomass burning emissions. This impact presents large temporal and 3 spatial variability since it occurs during the burning season that lasts only a few months per 4 year and is marked by tropical and boreal forest fires. Ziemke et al. (2009) modeling study with the Global Modeling Initiative (GMI) chemical transport model shows a global increase 5 6 in CO between 21% and 53% due to biomass burning. The tropospheric O₃ load has been 7 shown to correlate with that of CO during biomass burning events with a slope of O₃/CO of 8 about 1 (Honrath et al., 2004). However, other studies have shown only small changes in the 9 tropospheric ozone on global scale (4-5% increase computed by Ziemke et al. (2009)), where 10 regionally different impacts are computed, ranging for 10%-40% increase depending on 11 region and season (Galanter et al., 2000). Aircraft observations in Boreal Canada showed no 12 distinguishable within the smoke plume and in clean air (Parrington et al. (2013), while 13 substantial O₃ enhancement has been measured in air masses downwind fire locations (Palmer 14 et al., 2013). 15 The spatial variability of the annual mean impact of wildfire emissions on the tropospheric loads of OC, CO, NO_x, O₃, OH and isoprene is depicted in Fig. 7a-f and on BC, SO₄²⁻, NO₃-, 16 17 HNO₃ and NH₄⁺ in Fig. S5a-e (supplement). The most affected pollutants are OC (Fig. 7a) 18 and BC (Fig. S5a) with computed local reduction due to the omission of wildfires by almost 19 100%, in agreement with previous studies where a reduction of 50 % has been measured in 20 Beijing (Duan et al., 2004), and up to 66% in Central Mexico (Crounse et al., 2009). Our 21 results also show that annual mean local impacts on O₃ and CO, pollutants that have strong secondary sources, maximize at 20-30% in the tropics. As expected, the NO_x tropospheric 22 23 load is mostly affected by biomass burning both in the extra-tropics since fires contribute by 24 50% to the NO_x load at the outflow of boreal fires and in the tropical regions of south 25 America, Africa and N. Australia where burning is significant (Fig. 7c) in agreement with 26 previous studies that show up to 75% reduction near equatorial Africa (Galanter et al., 2000). 27 As a consequence of the NO_x and O_3 reductions when fire emissions are omitted, the 28 computed hydroxyl radical (OH) load (Fig. 7e) is significantly reduced (5-10%) over the same 29 regions; while larger percent reductions are computed at high northern latitudes where OH 30 loads are generally very low due to the very weak photochemistry there.

4.3.2 Impact of injection height

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2 The effect of height distribution of wildfire emissions on the computed tropospheric loads has 3 been studied by comparing the simulations SX.0 with the respective simulations SX.1. Fig. 8 4 presents such comparisons for BC. Both OC and BC are strongly affected by the injection height parameterization, since emitting aerosols above the boundary layer reduces aerosols 5 6 available near the surface for loss via dry deposition. The largest differences are computed for 7 the high latitudes over N. America and China where emission height distribution assumptions 8 can result in differences of about 25% (Fig. 8). Previous studies conducted with the GEOS-9 Chem model over the south eastern Asia during 2001, show a decrease of 20-40% of BC 10 surface concentrations when injected at height (Jian and Fu, 2014). In the same study it is 11 shown that biomass burning injection height has much larger impact on BC than CO (50%-12 150% more BC calculated at 700hPa, than when emitted in the boundary layer). Differences are positive over source areas (since more is emitted near the surface in SX.1) and negative 13 14 downwind (since less is transported away from source regions due to the increased deposition 15 flux at the source regions). Additional comparisons are presented in the supplementary 16 material (Fig. S6a-f). Assumptions in the biomass burning emissions injection height 17 marginally affect CO and O₃, with computed differences in the global annual mean 18 tropospheric load smaller than 2.5%.

4.3.3 Chemical feedbacks between biomass burning and vegetation

emissions

- 21 It is interesting to examine the impact of wildfire emissions on isoprene tropospheric load.
- 22 Isoprene is the single most important biogenic volatile organic compound (BVOC) emitted by
- vegetation (more than 50% of total annual BVOC emissions). The changes in OH described
- in section 4.3.1 (Fig. 7e), the main tropospheric oxidant that consumes isoprene, led to
- 25 opposite in sign changes of isoprene (Fig. 7f). Such results indicate a strong chemical
- 26 feedback between biomass burning and species emitted by vegetation.
- 27 This feedback is linking isoprene destruction and aerosol formation via the oxidants
- 28 (hydroxyl-OH- and nitrate-NO₃- radicals and ozone) that consume isoprene and produce
- 29 semi-volatile organics but also via primary biomass burning aerosols that provide surface for
- 30 organics to condense on. In the presence of fires, for the same isoprene emissions from
- vegetation (Fig. 7e) more nitrogen oxides (NO_x) (Fig. 7c) are emitted leading to higher OH

radicals in the extended biomass burning region (up to 20% regionally) and slightly lower over northern hemisphere regions with intensive anthropogenic NO_x emissions and their outflow. Thus, isoprene ambient levels are reduced with the highest reduction over and downwind tropical forested areas. Isoprene global tropospheric column is calculated to be lower by 15% in S0.0 than in S4.0 (fig. 7f). However, due to the NO_x-dependence of the semivolatile organic compounds formation from isoprene oxidation the total isoprene_SOA concentrations change little (2%). This implies an overall 13% reduction in semi-volatile organic compounds formation yield from isoprene oxidation that comes to compensate for the increased isoprene oxidation. In addition, the primary organic aerosols (POA) emitted by biomass burning provide surface for partitioning of semi-volatile compounds, thus significantly increasing the partitioning of organic vapors to the aerosol phase that in turn also stimulate further partitioning to the aerosol phase. Thus, the isoprene-SOA partitioning to the aerosol phase increases by 16% in depletion of the gas phase isoprene-SOA precursors. This enhancement is consistent with, although much lower than derived from results by Kanakidou et al. (2000) on the enhancement of SOA formation from biogenic VOC due to partitioning on POA from pollution sources. That earlier study was using higher aerosol yields from BVOC than here and did not account for the later studied NO_x-dependence of these yields; it also presented changes due to both combustion and fossil fuel POA. It has also shown that the use of different parameters in the two product yield representation of SOA formation from BVOC can lead to up to 70% of differences in the computed SOA tropospheric burden depending on atmospheric conditions. Tsigaridis et al. (2006) have evaluated the importance of the consideration of NO_x-dependent SOA formation by calculating changes in the SOA burden and characteristics and found that in the current troposphere about 72% of the total SOA mass is formed under NO_x-driven chemistry while in the past this fraction was lower (48%). Note however that large uncertainties and gaps in knowledge exist in the kinetics of isopreneaerosol formation. Rollins et al. (2009) studying the NO₃ radical-driven chemistry of isoprene-SOA formation, have demonstrated the complexity of isoprene chemistry with respect to SOA formation with a drastic increase in aerosol yield when both double bounds of isoprene are oxidized, thus documenting the aerosol yield dependence on the level of oxidation of the precursors. Ervens et al. (2008) investigations have shown that isoprene aqueous phase chemistry is more efficient (about 40% aerosol yield) than gas phase chemistry (about 3% of aerosol yield) in forming SOA and depends on the water content in the

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atmosphere and the pH. Carlton et al. (2009) review of laboratory measurements, field experiments and modeling studies concerning SOA formation from isoprene, documented differences in SOA yield parameterizations that most rely on a single set of chamber experiments, while aerosol yields are known to depend on various factors including the relative importance of NO_x versus peroxide chemistry, temperature (that affects aerosol components volatility based on their enthalpy of vaporization) and pre-existing aerosol loading. They have calculated differences in SOA load induced by the NO_x dependence parameterizations that are up to 30% of the total simulated OA over Eastern USA.

This feedback in the presence of biomass burning emissions increases by about 18% the global mean aerosol yield from isoprene emissions that is defined as the ratio of the tropospheric load of secondary organic aerosol from isoprene oxidation to the emissions of isoprene, while locally this difference can exceed 40%. Impacts on the tropospheric loads of the first generation gaseous products of isoprene are smaller, i.e. about 10%. The supplementary figure S11 shows the spatial distribution of the percent changes in the aerosol yield from isoprene emissions as computed comparing simulations S4.0 and S0.0. This figure points to the areas where the impact of biomass burning emissions (in percent) on the apparent SOA yield from isoprene is calculated by our model to be significant. These areas are the high latitude zone of North America and Asia, the tropical regions over land as well as the outflow from biomass burning regions. Note however that most isoprene_SOA formation occurs over land.

Our results demonstrate the strong coupling between tropospheric chemistry, biomass burning and vegetation emitted species. They show that it is critical for the evaluation of the impact of these emissions on tropospheric chemistry to consistently account for BVOC emissions from vegetation and the co-location/co-occurrence of biomass burning emissions in the area. Co-location of vegetation and biomass burning emissions is linked to the model grid size since co-location area increases with lowering the horizontal resolution of the model. In this respect, to further investigate the impact of the feedback strength to the model resolution, a lower resolution set of simulations has been also performed. These low resolution simulations give results similar to the higher resolution with regard to the feedback strength (relative changes between S0.0 and S4.0). Thus, the percent increases do not seem to be affected by the resolution of the model, while the computed tropospheric loads of isoprene and secondary

- 1 organic aerosol differ between the high and low resolution simulations with low resolution
- 2 simulation computing about 10% lower SOA and 4% lower isoprene loads.

4.4 Tropospheric lifetimes

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4 The lifetimes of pollutants provide a measure of pollutant persistence in the atmosphere. They 5 are here computed as the ratio of the tropospheric load to the loss rate (sum of chemical loss 6 and deposition fluxes) for each model column (first 22 vertical layers of the model). Global 7 mean tropospheric lifetimes are derived from the computed global burdens and losses. 8 Changes in chemistry as discussed above, as well as changes in deposition of pollutants due to 9 the modification of their spatial distribution, affect the lifetime of these compounds in the troposphere. Thus, isoprene's lifetime is increased in S4.0, as previously explained, by almost 10 20% compared to S0.0. The global tropospheric lifetimes of all other species are less 11 12 impacted by the choice of the emission inventory, with a maximum of about 12% for OC. 13 This is in agreement with previously calculated differences reported in literature. For instance, 14 such differences resulting from the use of 3 different biomass burning inventories (two global 15 and one regional) in the TM4 model coupled with the CBM4 chemical mechanism do not 16 exceed 5% for the African domain (Williams et al. (2012). Table 6 shows the calculated 17 global tropospheric lifetimes of pollutants for each scenario. The maximum percentage 18 differences from the base case scenario (S0.0) are computed for the S4.0 simulation that 19 neglects all wildfire emissions. 20 The lifetimes of pollutants, computed as the ratio of the tropospheric load to the loss rate (sum 21 of chemical loss and deposition fluxes) for each model column, show sensitivity to both the 22 height distribution of the emissions and the different emission inventories. The sensitivity of the BC lifetime to the height of injection of the biomass burning emissions is depicted in Fig. 23 24 9, where the difference in calculated tropospheric lifetimes of OC attributed to emission injection height alone can reach 30% (right panels). The differences produced by injection 25 26 height for other species are provided in Fig. S7 (supplement). The use of different biomass 27 burning emission inventories led to up to almost 90% local differences for OC as seen in Fig. 28 9g. The maximum differences are computed in the tropics and over the boreal forests in 29 Canada and eastern Russia using the ACCMIP and FINN inventories (Fig. 9e,g). The overall 30 impact of biomass burning emissions (simulations S4.0 versus S0.0) on the regional lifetimes 31 of tracers is shown in Fig. 10, where significant increases in O₃ (up to about 25%) and CO (up

1 to about a factor of 2) lifetimes are calculated when wild fire emissions are neglected. 2 Biomass burning is reducing O₃ lifetime in the burning regions of the tropics and the boreal 3 forests. This is mainly due to the reaction of O₃ with NO emissions and subsequent HNO₃ 4 formation. The impact of fire emissions on chemistry can be seen through the increases in the 5 regional lifetime of CO and isoprene in S4.0 (Fig. 10a,d), where local differences can reach 6 160%. OC and BC lifetimes are highly affected with local computed differences up to almost 90% (OC) and 150% (BC) (Fig. 10e-f). Similar results are produced for SO_4^{2-} lifetimes where 7 the local differences in calculated tropospheric lifetimes range from about -25% to 25% near 8 9 the tropics (Fig. 10g) and above the boreal forests of Russia and Canada where most open 10 biomass burning events occur. Note that aerosols species like OC and BC have significant 11 primary emissions from biomass burning and are removed from the atmosphere by dry and 12 wet deposition, while carbon monoxide, isoprene and O₃ loads and lifetimes are driven by strong chemical production and loss terms. Thus, aerosol species behave differently than these 13 14 short lived chemically reactive gases. 15 The tropospheric NO_v lifetime (NO_v=sum of NO_x, HNO₃, peroxyacetyl nitrate and organic 16 nitrates) due to depositional losses strongly responds to the wild fire emissions used in the 17 model, with differences between about -40% and 70% when taking into account all 18 simulations that have been performed for this study. Focusing on central Canada and north 19 eastern Asia, the S2.0 simulation results in a large increase in NO_v lifetime compared to S0.0 20 (Fig. 11b) that is weaker for the S1.0 (Fig. 11a). These differences are mainly attributed to the 21 spatial distribution of the emissions favoring different chemistry pathways and resulting in 22 different fractional composition of NO_y and thus different dry and wet removal fluxes. Note 23 that high differences in NO_v lifetime due to deposition are calculated over the tropical regions 24 with both positive and negative differences over and off-shore tropical Africa. There, both 25 NO_v burden and deposition losses are increasing due to biomass burning emissions and the corresponding NO_v lifetime in S0.0 varies roughly between 4 to 50 days (Figure S12a). When 26 27 wild fire emissions are omitted in the model, the absolute differences in NO_v lifetime between simulations S4.0 and S0.0 in the tropics vary from -6.5 to +5 days (Figure S12b). Thus, the 28 29 NO_v lifetime is increased by up to 67% locally (Fig. 11d), although on global scale a small 30 lifetime change (about -2%) is computed (Table 6).

5 Conclusions

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2 The CTM sensitivity simulations performed here show that the choice of wildfire emission 3 inventory has a significant impact on the simulated tropospheric concentrations of both primary emitted and secondary produced species, and as a result on the tropospheric lifetimes 4 5 of gaseous and aerosol pollutants. 6 The differences introduced by the choice of biomass burning emissions are usually between -7 30% and 30% above and downwind of biomass burning hotspots (near the tropics, boreal 8 forests of Russia and Canada) and can reach up to a factor of about 7 (e.g. for BC Fig. S4). 9 These impacts maximize for primary pollutants over source areas and for secondary pollutants 10 downwind. They are either due to the spatial and temporal differences in the emitted amounts 11 of primary pollutants, or to the resulting changes in the levels of oxidants and thus to the 12 impact of the primary pollutants on the concentrations of the chemically produced or 13 destroyed tracers. The injection height of the wildfire emissions is found to affect both the 14 tropospheric load and the lifetimes of the pollutants. Regionally up to 30% differences are 15 computed in the calculated tropospheric lifetimes of pollutants. Tropospheric column of OC is 16 mostly affected by different emission injection height with regional differences ranging from -17 20% to 25% and those attributed to the different emission inventories ranging from -70% to 18 450% (Fig. S8b). 19 Interestingly, isoprene, mainly emitted by vegetation, shows sensitivity to the biomass 20 burning emissions, with increasing tropospheric concentrations (and lifetime) when fire 21 emissions decrease mainly due to the reduction in OH radical concentrations. This leads to an 22 increase of the global mean aerosol yield from isoprene, defined as the ratio of tropospheric 23 loads of secondary aerosol from isoprene oxidation to isoprene emissions, by about 18% 24 when biomass burning emissions are taken into account. This fractional increase shows no sensitivity to the model resolution. 25 26 Finally, comparison of model results to observations shows the limitations of current 27 observations in evaluating the biomass burning emission inventories. Such evaluation requires 28 densifying air quality monitoring close to and downwind the major biomass burning sources 29 in the tropics, as well as over boreal regions, Alaska, South Asia and Indonesia where our simulations using different biomass burning emission inventories show the larger diversity. 30

6 Acknowledgements

- 2 This work has been supported by the EU-FP7 project ECLIPSE (FP7-ENV-2011-282688).
- 3 ND acknowledges support from the EU-FP7 project PEGASOS (FP7-ENV-2010-265148).
- 4 The authors would like to thank the reviewers for their pertinent comments that helped
- 5 improving this work.

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Table 1 Anthropogenic emissions (Tg a⁻¹) used in this study and fraction of emissions that corresponds to the AWB sector included in the ECLIPSE anthropogenic emissions inventory. Both absolute quantities and percentage of the total anthropogenic emissions from (Klimont et al., 2013) are presented.

	BC	CO	NOx	OC	SOx	NMVOC
ECLPSE (with AWB)	5.38	527.1	43.97	11.56	45.95	140.47
AWB on ECLIPSE	0.333	27.46	0.296	1.281	0.173	4.255
% contribution of AWB	6.19	5.21	0.67	11.08	0.38	3.03
to total anthropogenic						

Table 2 Total annual amounts of pollutants emitted by wild fires according to the different inventories used, for 2008 in Tg a^{-1} . NO_x is reported as NO. (*)GFEDv3.1 without the AWB is here called GFEDv3.1-ECLIPSE

	BC	CO	NO_x	OC	SO ₂	NMVOC	NH ₃	Spatial resolution	Temporal resolution
GFEDv3.1- ECLIPSE*	1.695	264.205	3.751	15.197	0.940	44.414.	3.320	0.5°x0.5°	Monthly
FINN	1.939	338.576	5.998	20.202	1.102	63.476	5.410	1°x1°	Monthly
ACCMIP	2.620	460.419	5.479	23.309	1.929	80.869	9.203	0.5°x0.5°	Monthly

Table 3 Agricultural Waste Burning sector as provided for different emission inventories in Tg a⁻¹ for the year 2008. NOx is reported as NO

	ВС	co	NOx	ос	SOx	NMVOC
ECLIPSE	0.333	27.46	0.296	1.281	0.173	4.255
GFEDv3.1	0.064	12.57	0.143	0.497	0.027	1.296
ACCMIP	0.162	21.22	0.444	0.775	0.220	2.857

Table 4 Summary of simulations performed for this work.

Height	inventory	Varying	Surface	AWB	
S0.0	GFEDv3 ECLIPSE	X		ECLIPSE	
S0.1	GFEDVS ECLIFSE		X	ECLIISE	
S1.0	GFEDv3.1	X		GFEDv3.1	
S1.1	GFEDV3.1		X	Gredv3.1	
S2.0	ACCMIP	X		ECLIPSE	
S2.1	ACCMIP		X	ECLIPSE	
S3.0	TETNINI	X		ECI IDCE	
S3.1	FINN		X	ECLIPSE	
S4.0	zero			ECLIPSE	

Table 5 Total annual mean tropospheric load of pollutants for all simulations in Tg a⁻¹.

	S0.0	S0.1	S1.0	S1.1	S2.0	S2.1	S3.0	S3.1	S4.0
СО	319.12	318.37	317.26	316.20	341.47	339.63	331.58	330.37	283.88
O_3	416.17	415.52	415.35	414.82	422.17	421.29	423.04	422.03	405.25
NO_x	1.299	1.293	1.286	1.282	1.330	1.323	1.390	1.378	1.200
$\mathrm{SO_4}^{2\text{-}}$	1.914	1.908	1.913	1.906	1.933	1.923	1.911	1.905	1.868
HNO ₃	2.196	2.188	2.181	2.181	2.235	2.228	2.229	2.219	2.048
$\mathbf{NH_4}^+$	0.498	0.487	0.514	0.496	0.516	0.496	0.507	0.492	0.460
Isoprene	0.266	0.267	0.267	0.268	0.247	0.248	0.253	0.254	0.315
OC	0.111	0.110	0.110	0.109	0.121	0.120	0.117	0.116	0.072
BC	0.136	0.135	0.131	0.131	0.146	0.146	0.133	0.133	0.088

Table 6 Calculated annual mean tropospheric lifetimes of pollutants for all the simulations performed.

	S0.0	S0.1	S1.0	S1.1	S2.0	S2.1	S3.0	S3.1	S4.0
CO (days)	41.48	41.44	41.43	41.35	41.82	41.67	41.45	41.40	41.67
O_3 (days)	24.58	24.62	24.59	24.63	24.39	24.43	24.33	24.39	25.19
NO _y (days)	7.342	7.300	7.293	7.255	7.358	7.297	7.628	7.541	7.184
SO ₄ ²⁻ (days)	4.446	4.442	4.448	4.444	4.427	4.423	4.421	4.419	4.426
HNO ₃ (days)	2.804	2.805	2.793	2.800	2.792	2.796	2.774	2.775	2.776
NH ₄ ⁺ (days)	4.979	4.932	5.032	4.962	4.961	4.905	4.928	4.894	4.862
Isoprene (hours)	4.457	4.475	4.466	4.482	4.137	4.152	4.236	4.250	5.270
OC (days)	6.031	5.998	6.046	6.012	5.925	5.894	5.839	5.819	5.302
BC (days)	6.927	6.908	6.962	6.941	6.889	6.871	6.583	6.572	6.261

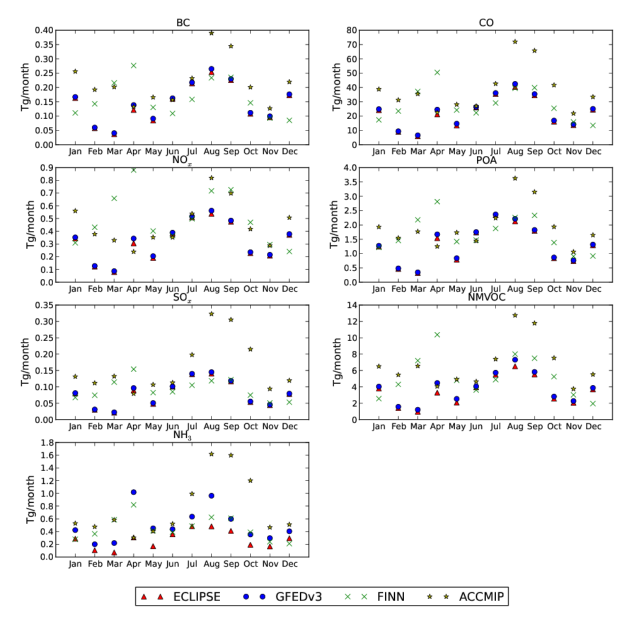


Fig. 1 Monthly variation and differences of biomass burning emission inventories for the year 2008 for all species used in the model. For simplicity, NMVOC are summed up. NOx are presented in NO, SOx in SO_2 and NMVOCs in total mass.

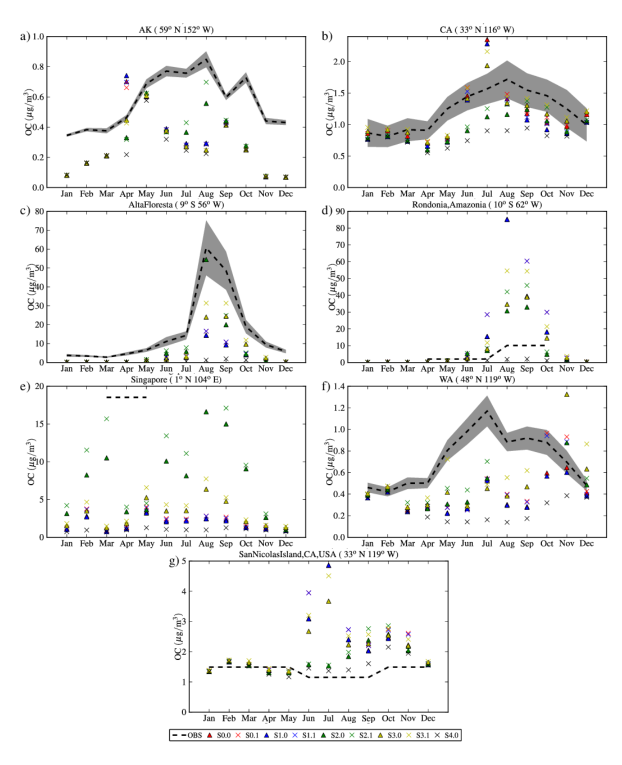


Fig. 2 Comparison of monthly mean model results with observations of organic carbon (OC) at southern Alaska (a), California State, USA (b), Alta Floresta, Brazil (c), Rondonia, Amazonia (d), Singapore (e), Washington State, USA (f) and San Nicolas Island, California, USA (g). The dashed line with the gray shaded area shows the monthly mean value of observations with the standard deviation based on their interannual variability, while the colored symbols show the calculated values for the specific station. Triangles are for simulations assuming a vertical distribution of wildfire emissions, while the x symbols show the simulations assuming that all open biomass burning emissions occur near the surface. Details on the simulations are given in Table 4.



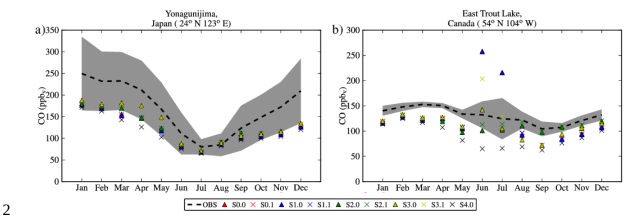


Fig. 3 Comparison of monthly mean model results with CO surface observations at Yonagunijima, Japan (a) and at East Trout Lake, Canada (b). Lines and symbols as in Fig. 2 but for CO.

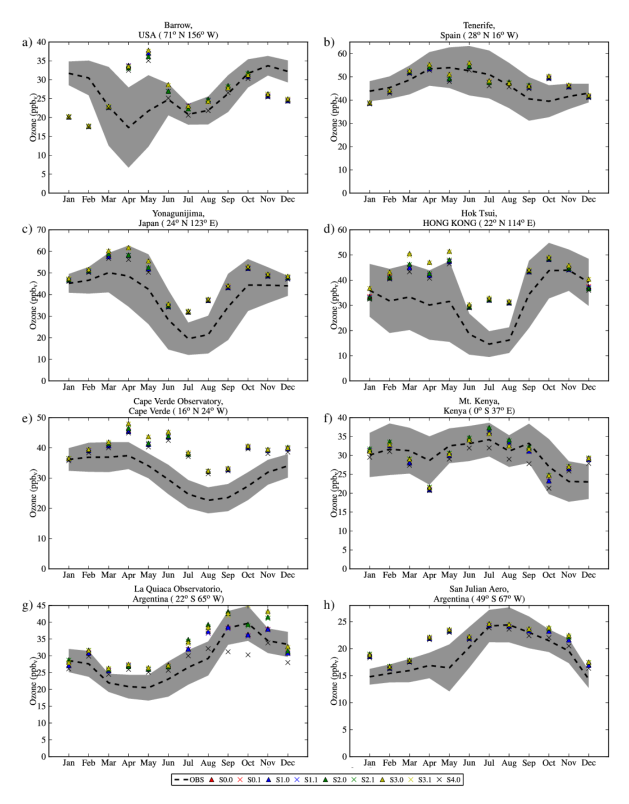


Fig. 4 Comparison of monthly mean surface ozone measurements with model results at Barrow, USA (a), Tenerife, Spain (b), Yonagunijima, Japan (c), Hok Tsui, Hong Kong (d), Cape Verde Observatory, Cape Verde (e), Mount Kenya, Kenya (f), La Quiaca Observatory, Argentina (g) and San Julian Aero, Argentina (h). Lines and symbols as in Fig. 2 but for O₃.

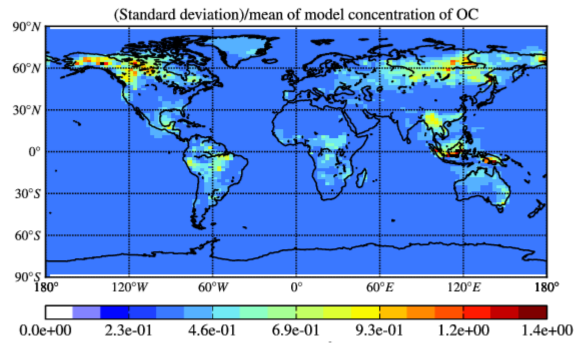


Fig. 5 Spatial distribution of the ratio of the standard deviation to the mean of all model simulations, based on annual mean of the computed surface OC concentrations.

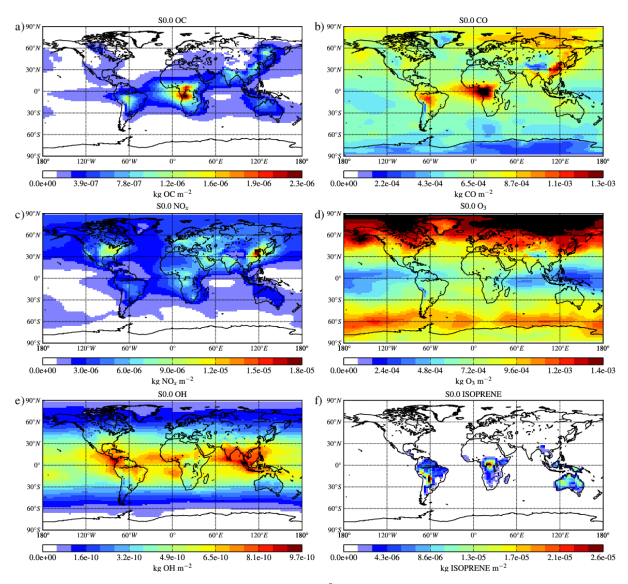


Fig. 6 Calculated annual mean tropospheric load in (Kg m⁻²) of selected species for the base case scenario (S0.0). Areas with black exceed the maximum value of the colorbar.

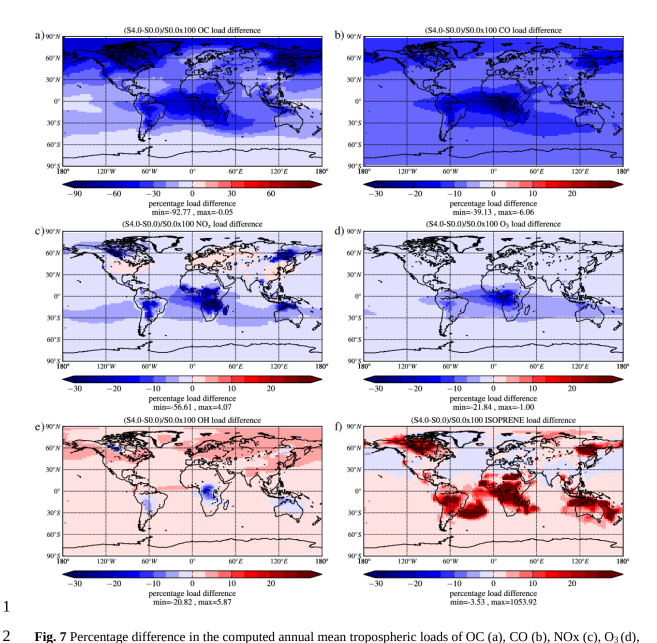


Fig. 7 Percentage difference in the computed annual mean tropospheric loads of OC (a), CO (b), NOx (c), O_3 (d), OH (e), isoprene (f) – attributed to wildfire emissions calculated as (column_S4.0 – column_S0.0)/(column S0.0)x100. The scale is from -30% to 30% (-90% to 90% for OC); the minimum and maximum differences are printed under each panel.

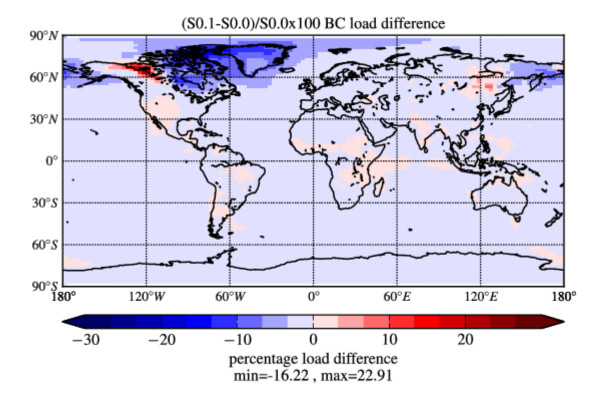


Fig. 8 Percentage difference of annual mean computed tropospheric load of BC attributed to wildfire emission injection height calculated as $(load_S0.1 - load_S0.0)/(load_S0.0)x100$. The scale is from -30% to 30%; the minimum and maximum percent differences are printed under each panel.

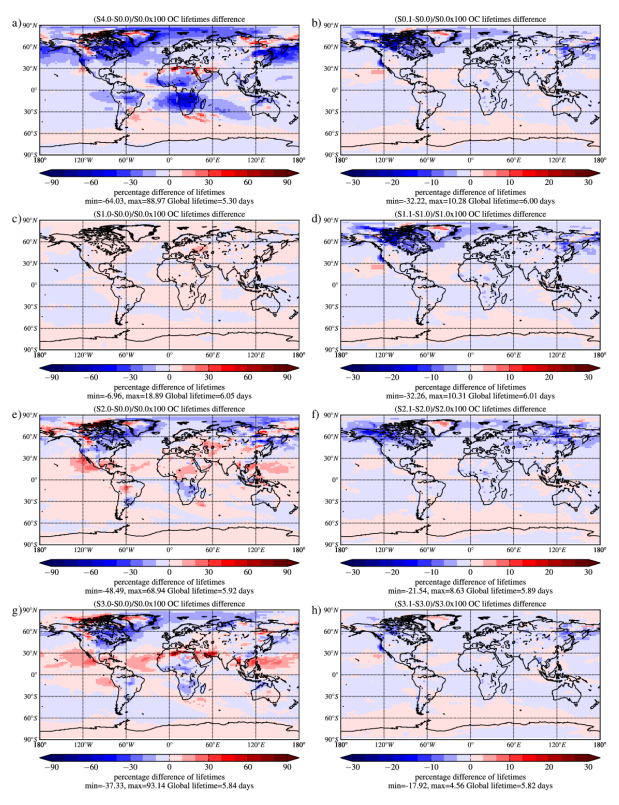


Fig. 9. Percent impact on the computed annual mean tropospheric lifetime of OC of: (left panels) the different emission inventories calculated as the percent difference between simulations SX.0 and simulation S0.0; and of (right panels) height distribution calculated as the percent difference between simulations SX.1 and simulations SX.0.. The colorbar ranges from -90% to 90% for the surface differences and -30% to 30% for the differences induced by height distribution. The minimum and maximum local lifetimes percent changes as well as the global lifetime are printed under each panel.

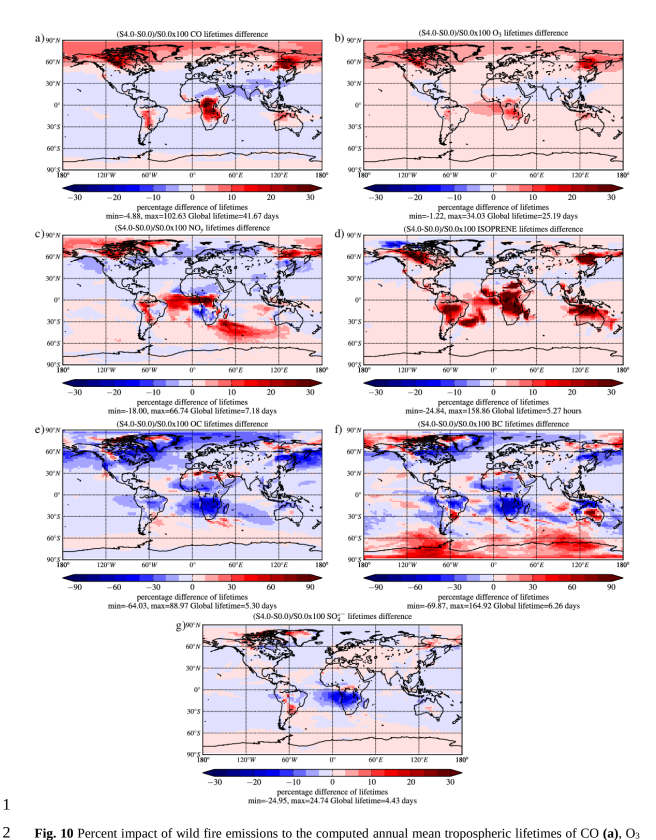


Fig. 10 Percent impact of wild fire emissions to the computed annual mean tropospheric lifetimes of CO (a), O₃ (b), NO_y (c), isoprene (d), OC (e), BC (f) and SO₄²⁻ (g) depicted as the percentage difference of S4.0 and S0.0. The colorbar ranges from -30% to 30% (-90% to 90% for OC and BC). The minimum and maximum local lifetimes percent changes as well as the global lifetime are printed under each panel.

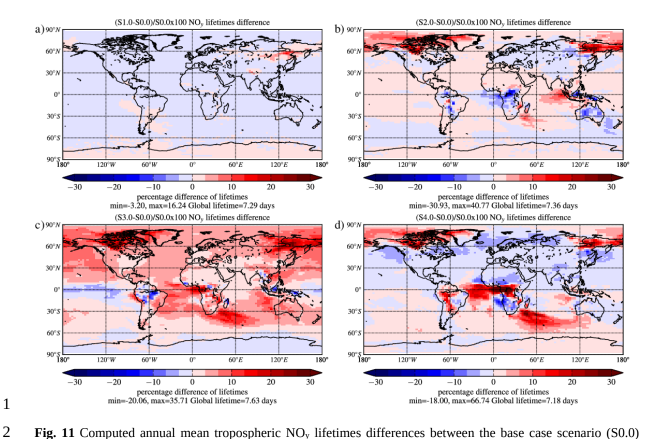


Fig. 11 Computed annual mean tropospheric NO_y lifetimes differences between the base case scenario (S0.0) and S1.0 (a), S2.0 (b), S3.0 (c) and S4.0 (d), computed by reference to S0.0. The colorbar ranges from -30% to 30%. The minimum and maximum local lifetimes percent changes as well as the global lifetime are printed under each panel.