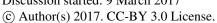
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Projected global tropospheric ozone impacts on vegetation under different 1

emission and climate scenarios 2

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Abstract

The impact of ground-level ozone (O₃) on vegetation is largely under-investigated at global 11 scale despite worldwide large areas are exposed to high surface O₃ levels and concentrations 12 are expected to increase in the next future. To explore future potential impacts of O₃ on 13 14 vegetation, we compared historical and projected O₃ concentrations simulated by six global atmospheric chemistry transport models on the basis of three representative concentration 15 pathways emission scenarios (i.e. RCP 2.6, 4.5, 8.5). To assess changes in the potential O₃ 16 threat to vegetation, we used the AOT40 metric. Results point out a significant overrun of 17 AOT40 in comparison with the recommendations of UNECE for the protection of vegetation. 18 In fact, many areas of the northern hemisphere show that AOT40-based critical levels will be 19 exceeded by a factor of at least 10 under RCP8.5. Changes in surface O₃ by 2100 range from 20 about + 4-5 ppb worldwide in RCP8.5 scenario to reductions of about 2-10 ppb in the RCP2.6 21 scenario. The risk of O₃ injury for vegetation decreased by 61% and 47% under RCP2.6 and 22 23 RCP4.5, respectively and increased by 70% under RCP8.5. Key biodiversity areas in South and North Asia, central Africa and Northern America were identified as being at risk from 24 high O₃ concentrations. To better evaluate the regional exposure of ecosystems to O₃ 25 pollution, we recommend the use of improved chemistry-climate modelling system, fully 26 27 coupled with dynamic vegetation models.

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Keywords: AOT40, Ozone, Representative Concentration Pathways, O3 injury on vegetation

2013; Proietti et al., 2016).

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Introduction

Tropospheric ozone (O₃) is a secondary air pollutant, i.e. it is not emitted as such in the air but 34 it is formed by reactions among precursors (e.g. CH₄, VOCs, NOx). Ozone is an important 35 greenhouse gas resulting in a direct radiative forcing of 0.35-0.37 W m⁻² on climate (Shindell 36 et al., 2009; Ainsworth et al., 2012). Despite significant control efforts and legislation to 37 reduce O₃ precursor emissions, tropospheric O₃ pollution is still a major air quality issue over 38 large regions of the Globe (Lefohn et al., 2010; Langner et al., 2012; Young et al., 2013; 39 40 Cooper et al., 2014; EEA, 2015; Sicard et al., 2016a,b). Long-range transport of O₃ and its precursors can elevate the local and regional O₃ background concentrations (Ellingsen et al., 41 2008; Wilson et al., 2012; Paoletti et al., 2014; Derwent et al., 2015; Xing et al., 2015; Sicard 42 et al., 2016a). Therefore, remote areas such as the Arctic region, can be affected (Langner et 43 al., 2012). The current tropospheric O₃ levels (35-50 ppb in the northern hemisphere, NH) are 44 high enough to damage both forests and crops by reducing growth rates and productivity 45 46 (Paoletti et al., 2009; Wittig et al., 2009; Anav et al., 2011; Mills et al., 2011; Ashworth et al.,

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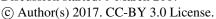
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Increasing atmospheric CO₂, nitrogen deposition and temperatures enhance plant growth, and increase primary production and greening of plants (Nemani et al., 2003; Zhu et al., 2016). At the global scale, a widespread increase of greening and net primary production (NPP) is observed over 25-50% of the vegetated area, while a decrease is observed over only 7% of the Globe (Nemani et al., 2003; Zhu et al., 2016). In contrast, a previous modeling study over Europe shows how O₃ reduces the mean annual gross primary production (GPP) by about 22% and the leaf area index by 15-20% (Anav et al., 2011). Similarly, Proietti et al (2016), using different in-situ measurements collected over 37 European forest sites, found a GPP decrease of 30% caused by O₃. At global scale, over the time period 1901-2100, GPP is projected to decrease by 14-23% (Sitch et al., 2007). As a consequence of reduced photosynthetic assimilation, the total biomass of trees is estimated to be decreased by 7% under the current O₃ mean concentrations (40 ppb) and by 17% under the O₃ mean concentrations expected in 2100 (97 ppb) compared to preindustrial O₃ levels (about 10 ppb, Wittig et al., 2009). Wittig et al. (2009) also reported that the total tree biomass of angiosperms was reduced by 23% at O₃ mean concentrations of 74 ppb, and by 7% at 92 ppb for gymnosperms. High surface O₃ levels, exceeding 40 ppb, do occur in many regions of the Globe with associated economic costs of several billion dollars per year (Wang and Mauzerall, 2004; Ashmore, 2005). Ashworth et al. (2013) reported an annual loss of 3.5% for

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wheat (very O₃-sensitive) and 1% for maize (more O₃-tolerant) for Europe in 2010 relative to 2000, while Holland et al. (2006) estimated a €4.5 billion loss in the production of 23 common crop species, due to surface O₃ exposure by 2020 relative to 2000.

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The international Tropospheric Ozone Assessment Report (TOAR) establishes a state-of-theart and an up-to-date scientific assessment of global O₃ metrics for climate change, human health and crop/ecosystem research (Lefohn et al. 2017). To assess the potential O₃ risk and protect vegetation from O₃, different metrics are used: the European and US standard (AOT40 and W126, respectively) are based on exposure-based metrics, while flux-based metrics have been introduced only recently (UNECE, 2010; Klingberg et al., 2014; EEA, 2015). Unlike the exposure-based metrics, which only rely on the surface O₃ concentration, the flux-based metrics were developed to quantify the accumulation of damaging O₃ taken up by vegetation through the stomata over a species-specific phenological time-window. These metrics also provide an information-rich tool in assessing the relative effectiveness of air pollution control strategies in lowering surface O₃ levels worldwide (Monks et al., 2015). By reducing plant photosynthesis and growth, high tropospheric O₃ levels will result in reduction in carbon storage by vegetation and, in fine an indirect radiative forcing as a consequence of the CO2 rising in the atmosphere (Sitch et al., 2007; Ainsworth et al., 2012). This CO₂ rising reduces stomatal conductance which decreases O₃ flux into plants leading to increased O₃ levels in the air of 3-4 ppb during the growing season over the NH by doubling of CO2 concentration (Fiscus et al., 2005; Sanderson et al., 2007).

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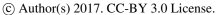
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Projected changes in tropospheric O₃ vary considerably among models (Stevenson et al., 2006; Wild, 2007) and emission scenarios. In earlier studies, the emissions of O₃ precursors were based on a high population growth, leading to very high projected surface O₃ concentrations by 2100 (Stevenson et al., 2000; Zeng and Pyle, 2003; Shindell et al., 2006). The last emission scenarios, i.e. the Representative Concentration Pathways (RCPs) were developed as part of the Fifth Assessment Report of the Intergovernmental Panel on Climate Change (Meinshausen et al., 2011; van Vuuren et al., 2011; Cubasch et al., 2013; Myhre et al., 2013). These scenarios include e.g. different assumptions on climate, energy access policies, and land cover and land use changes (Arneth et al., 2008; Kawase et al., 2011; Kirtman et al., 2013). Until now, studies on O₃ pollution impacts on terrestrial ecosystems are either limited to a single model or to particular regions (e.g. Clifton et al., 2014; Rieder et al., 2015) and only a few applications of global or regional models under the new RCPs scenarios

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were carried out (Kelly et al., 2012). In the framework of the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP), different simulations were performed by

Lamarque et al. (2013) and Young et al. (2013) from 16 global chemistry models.

A few issues about surface O_3 , such as a better understanding of spatial changes and a better assessment of O_3 impacts worldwide, are still challenging. To overcome these issues, the aim of this study is to quantify, for the first time, the spatial and temporal changes in the projected potential O_3 impacts on carbon assimilation of vegetation at global scale, by comparing the O_3 potential injury at present with that expected at the end of the 21^{st} century from different global chemistry models.

Materials and Methods

ACCMIP models and RCP scenarios

The global chemistry models used in this work have been developed under the ACCMIP project. A detailed description of the selected models and of the emission scenarios (i.e. RCPs) is included in Supplementary Information (SI). ACCMIP models have been widely validated and used to evaluate projected changes in atmospheric chemistry and air quality under different emission and climate assumptions (e.g. Lamarque et al., 2010; Fiore et al., 2012; Bowman et al., 2013; Lee et al., 2013; Voulgarakis et al., 2013). Lamarque et al. (2013) and Young et al. (2013) provided the main characteristics of 16 models and details for the ACCMIP simulations. Although within the ACCMIP project 16 models are available, due to the lack of hourly O₃ concentration here we only focus on 6 global chemistry models with different configurations (Table 1).

The length of historical and RCP simulations vary between models, but for all models the historical runs cover a period centered around 2000, while the time-slice of RCPs is centered around 2100 (Table 1). As for each model we compare the mean change between the historical and RCP simulations, a different length in the number of years used in the analysis does not affect the results.

Potential ozone injury on vegetation

The O₃ exposure-based index, i.e. AOT40 (ppb h), is a metric used to assess the potential O₃ risk to vegetation from local to global scales (Emberson et al., 2014). It is computed as sum

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137 of the hourly exceedances above 40 ppb, for daylight hours (8am-8pm) over species-specific growing seasons (UNECE, 2010). A recent study over Europe showed how computing 138 139 AOT40 only over the growing season (i.e. April-September) would lead to an underestimation of AOT40 up to 50% for conifer trees, while in case of deciduous trees the underestimation is 140 much smaller (< 5%, Anav et al., 2016). Besides, it should be noted that in Anav et al. (2016) 141 the AOT40 is computed year-round when the stomatal conductance is greater than 0. Here, 142 because of the lack of hourly meteorological data, we can only compute the AOT40 year-143 round and during the daylight hours. In case of risk assessment, this approach would lead to a 144 relevant overestimation of AOT40, mainly over polluted area of NH. Nevertheless, since the 145 aim of this study is to compare how O₃ stress to vegetation changes between historical period 146 and future, the overestimation of AOT40 does not affect our results. Therefore, we computed 147 AOT40 as follows: 148

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150 AOT 40=
$$\int_{0.1 \text{ ian}}^{3.1 \text{ dec}} \max(([O_3] - 40), 0) dt$$
 (1)

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where $[O_3]$ is hourly O_3 concentration (ppb) simulated by the models at the lower model layer and dt is time step (1h). The function "maximum" ensures that only values exceeding 40 ppb are taken into account. The O_3 concentration to be used in AOT40 calculation should be at the top of the canopy; however, most of models used here provide O_3 concentrations at 90-120 m. Nevertheless, even if the O_3 concentration is simulated at different elevations above the sea level, as for each model we compare the variation between present and future, the change is consistent because the elevation is the same. For the protection of forests, a critical level of 5,000 ppb.h (or 5 ppm.h) is recommended by UNECE (2010). Within the 2008/50/CE Directive, the critical level for agricultural crops (3 ppm.h) is adopted as the long-term objective value for the protection of vegetation by 2020.

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From the AOT40, a factor of risk for forests and crops can be computed (Anav et al. 2011; Proietti et al. 2016). Thus, the potential O₃ impact on photosynthetic assimilation (IO₃) is

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$$167 \quad IO3 = \alpha \times AOT40 \tag{2}$$

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where α is an empirically derived O_3 response coefficient representing the proportional change in photosynthesis per unit of ozone-uptake (Anav et al., 2011). The coefficient for coniferous trees $(0.7 \times 10^{-6} \text{ mm}^{-1} \text{ ppb}^{-1})$ and crops $(3.9 \times 10^{-6} \text{ mm}^{-1} \text{ ppb}^{-1})$ are based on the

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regressions of the ozone-uptake response curves (Reich, 1987), while the coefficient for deciduous trees and other vegetation types (2.6×10⁻⁶ mm⁻¹ ppb⁻¹) is based on Ollinger et al. (1997). From changes in the risk factor, we can highlight potential risk areas for vegetation.

Results and Discussion

Although differences in the simulated global O₃ spatial pattern were previously discussed and analyzed (e.g. Lamarque et al., 2013), we show the mean annual O₃ concentration at the lower model layer in Figure 1 because O₃ concentration explains AOT40 patterns. Then, in Figure 2 we show and discuss the AOT40 spatial and temporal distribution from the ACCMIP models for the historical and RCPs simulations, and finally in Figure 3 we show the percentage of variation of IO3, i.e. the change in the potential impact of O₃ on vegetation for the ACCMIP models computed comparing the RCPs simulations with historical runs. A detailed description of each figure, model by model, is included in Supplementary Information (SI).

Spatial pattern of historical ozone concentration and AOT40

The highest surface O_3 concentrations (Fig. 1) and potential O_3 injury (Fig. 2) are found in the NH, highlighting a hemispheric asymmetry. The multi-models O_3 mean concentration, averaged over the land points of the domain, is 37.9 ± 4.3 ppb in NH and 22.9 ± 3.8 ppb in SH (Table 3a). The NH extratropics (i.e. mid-latitudes beyond the tropics) has 65% more O_3 than the SH extratropics (data not shown). The highest AOT40 values are found in the NH, with an averaged AOT40 of 24.8 ± 10.1 ppm.h in NH and 2.5 ± 1.7 ppm.h in SH (Table 3a).

According to previous studies, the annual mean background O₃ concentrations at NH midlatitudes range between 35 and 50 ppb during the end of the 20th century (e.g. Cooper et al., 2012; IPCC, 2014; Lefohn et al. 2014). Similarly, we found historical surface O₃ mean concentrations ranging between 35 and 50 ppb and 35-50 ppm.h for AOT40 in the NH, with the highest values occurring over Greenland and in the latitude band 15-45°N, particularly around the Mediterranean basin, Near East, Northern America and over the Tibetan plateau (> 50 ppb and 70 ppm.h) while the lowest O₃ burden (15-30 ppb, < 20 ppm.h) was recorded in SH, particularly over Amazon, African and Indonesian rainforests. Tropospheric O₃ has a significant source from stratospheric O₃ (Parrish et al., 2012) and it can be transported by the large-scale Brewer-Dobson overturning circulation, i.e. an upward motion from the tropics and downward at higher latitudes, resulting in higher O₃ concentrations in the extratropics

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205 (Hudson et al., 2006; Seidel et al., 2008; Parrish et al., 2012). The six models are able to reproduce the spatial pattern of O₃ concentration and thus AOT40 worldwide. 206

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The highest historical O₃ mean concentrations are observed in GFDL-AM3 and the lowest are 208 found in MIROC-CHEM. In the early 2000s, the maximum global O₃ mean concentration (39 209 ppb) in GFDL-AM3 is associated to the lowest annual total NOx emissions (46.2 Tg, Table 210 2a) and low LNOx (4.4 Tg) while the minimum global O₃ mean concentration (28 ppb) in 211 212 MIROC-CHEM is related to the highest emissions of total NOx per year (57.3 Tg) and erroneously high LNOx (9.7 Tg per year, Lamarque et al., 2013). MIROC-CHEM simulates 213 58 gaseous species in the chemical scheme with constant present-day biogenic VOCs 214 emissions while GFDL-AM3 simulates 81 species (Stevenson et al., 2012; Lamarque et al., 215 2013). In GISS-E2-R, the hemispheric asymmetry in O₃ is more important with e.g. a mean 216 concentration of 22 ppb in SH and 42 ppb in NH. A stronger global AOT40 mean (26 ppm.h) 217 218 is observed in GISS-E2-R and the lowest (7 ppm.h) in MIROC-CHEM for historical simulations. Model-to-model differences are observed due to different natural emissions of O₃ 219 220 precursors (e.g. lightning NOx) and the used chemical schemes.

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Higher O₃ burdens (mean concentration > 50 ppb, AOT40 > 70 ppm.h) are simulated at highelevation areas, e.g. at Rocky and Appalachian Mountains and over the Tibetan plateau (Fig. 1, Fig. 2). At high-elevation, solar radiation, biogenic VOC emission, exchange between free troposphere and boundary layer, and stratospheric O₃ intrusion within the troposphere are more important that at the surface layer (Steinbacher et al. 2004; Kulkarni et al., 2011; Lefohn et al., 2012). Altitude reduces the O₃ destruction by deposition and NO (Chevalier et al., 2007). In addition, due to the high elevation, ambient air remains colder and dryer in summer, leading to lower summertime O₃ losses from photolysis (Helmig et al., 2007). The highelevation areas, characterized by higher O₃ burdens, are well simulated in GISS-E2-R and MOCAGE models.

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The Tibetan plateau, so-called "ozone valley", is the highest plateau in the world, with a mean height of 4000 m a.s.l. (Tian et al., 2008) with strong thermal and dynamic influences on 234 regional and global climate (Chen et al. 2011). High surface O₃ mean concentrations (40-60 ppb) were reported in previous studies (e.g. Zhang et al., 2004; Bian et al., 2011; Guo et al., 236 2015; Wang et al., 2015). Although this region is remote, road traffic, biofuel energy source, coalmines and trash burning are prevalent. These pollution sources contribute to significant

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amount of NOx, CO and VOCs (Wang et al., 2015). The high O₃ levels are attributed to the combined effects of high-elevation surface, thermal and dynamical forcing of the Tibetan plateau and *in-situ* photochemical production in the air trapped in the plateau by surrounding mountains (Guo et al., 2015; Wang et al., 2015). The dynamic effect, associated with the large-scale circulation, is more important than the chemical effect (Tian et al., 2008; Liu et al., 2010) and responsible for the high O₃ levels over the Tibetan plateau. The six models are able to well reproduce the high surface O₃ mean concentrations (> 50 ppb) over the Tibetan

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Higher O₃ mean concentrations (> 60 ppb) are also observed in Southwestern U.S., at the stations inland close to Los Angeles, in Northeastern U.S. and East Asia (e.g. Beijing) (Fig. 1). The American Southwest is an O₃ precursor hotspot where the industrial sources emit CH₄ and VOCs into the air (Jeričević et al., 2013) and the eastern and northern desert areas have 252 higher ambient O₃ than urban areas of southern California due to four factors: on-shore winds, gasoline reformulation, eastward population expansion and nighttime air chemistry (Arbaugh 254 and Bytnerowicz, 2003). The surface concentrations show higher O₃ levels in areas downwind of O₃ precursor sources, i.e. urban and well-industrialized areas, at distances of hundreds or even thousands of kilometers due to transport of O₃ and precursors, including "reservoir" 256 species such as PAN, lower O₃ titration by NO and higher biogenic VOC emission (Wilson et al., 2012; Paoletti et al., 2014; Monks et al., 2015; Sicard et al., 2016a). The higher O₃ levels in areas downwind of O₃ precursor sources are well simulated in GISS-E2-R and MOCAGE models.

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In the lower troposphere, O₃ can be removed by a large number of chemical reactions and by dry deposition (Sicard et al., 2016c). The O₃ dry deposition rates range from 0.01-0.05 cm s⁻¹ (oceans and snow) to 0.15-1.80 cm s⁻¹ for mixed wood forests (Wesely and Hicks, 2000; Zhang et al., 2003). The model performance is also related to the parameterization of the dry deposition rates.

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Over Greenland, mean O₃ concentrations during the historical runs, ranged from 40 to 55 ppb (Fig. 1) except in MIROC-CHEM (20-25 ppb). Similarly, Helmig et al. (2007) reported annual mean of surface O₃ concentrations of 47 ppb over Greenland between 2000 and 2005, particularly at the high-elevation Summit station (3200 m a.s.l.). Several investigations, about snow photochemical and oxidation processes over Greenland, concluded that photochemical

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273 O₃ production can be attributed to high levels of reactive compounds (e.g. oxidized nitrogen species) present in the surface layer during the sunlit periods due to local sources e.g. NOx 274 275 enhancement from snowpack emissions, Peroxyacetyl nitrate (PAN) decomposition, boreal 276 forest fires or ship emissions (Granier et al., 2006; Stohl et al., 2007; Legrand et al., 2009; Walker et al., 2012). PAN to NOx ratio increases with increasing altitude and latitude (Singh 277 et al., 1992). The PAN reservoir for NOx may be responsible for the increase in surface 278 O₃concentrations at high latitudes (Singh et al., 1992). Local O₃ production does not appear to 279 280 have an important contribution to the ambient high O_3 levels (Helmig et al., 2007), however the long-range O₃ transport can elevate the background concentrations measured at remote 281 sites, e.g. Greenland (Ellingsen et al., 2008; Derwent et al., 2010). Low dry deposition rates 282 for O₃, the downward transport of stratospheric O₃, the photochemical local production and 283 the large-scale transport (Legrand et al., 2009; Walker et al., 2012; Hess and Zbinden, 2013) 284 285 are known factors to explain higher O₃ pollution over Greenland.

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292 293 The surface O₃ concentrations (> 40 ppb) and AOT40 (> 60 ppm.h) are higher over deserts, downwind of O₃ precursor sources (e.g. Near East, Sierra Nevada, Colorado Desert), due to lower O₃ dry deposition fluxes, O₃ precursors long-range transport from urbanized areas and high insolation. Around the Mediterranean basin, elevated AOT40 values (> 60 ppm.h) are recorded, mainly due to the industrial development, road traffic increment, high insolation, sea/land breeze recirculation and O₃ transport (Sicard et al., 2013). All models, except MIROC-CHEM, are able to well reproduce the high surface O₃ mean concentrations over Greenland and over deserts.

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Projected changes in ozone concentration and AOT40

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Recent studies display a mean global increase in background O₃ concentration from a current level of 35-50 ppb (e.g. IPCC, 2014; Lefohn et al. 2014) to 55-65 ppb (e.g. Wittig et al., 2007) and up to 85 ppb at NH mid-latitudes by 2100 (IPCC, 2014). During the latter half of the 20th century surface O₃ concentrations have increased markedly at NH mid-latitudes (e.g. Oltmans et al., 2006; Parrish et al., 2012; Paoletti et al., 2014), mainly related to increasing anthropogenic precursor emissions related to economic growth of industrialized countries (e.g. Lamarque et al., 2005). Our results indicate that the future projections of the mean tropospheric O₃ concentrations and AOT40 vary considerably with the different scenarios and models (Fig. 1 and 2). The six models simulate a decrease of O₃ concentration by 2100 under the RCP2.6 and RCP4.5 scenarios, and an increase under the RCP8.5 scenario (Lamarque et

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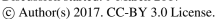


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al., 2011). In our study, the averaged relative changes in surface O₃ concentration means (and AOT40) for the different RCPs are: -21% (-75%) for RCP2.6, - 10% (-50%) for RCP4.5 and 309 310 + 14% (+69%) for RCP8.5 with a strong disparity between both hemispheres, e.g. - 8% in SH and - 25% in NH for RCP2.6 (Tables 3b-c). RCP8.5 is the only scenario to show an increase 311 in global background O₃ levels by 2100 (+ 23% in SH and + 11% in NH). 312 313 Under the RCP2.6 scenario, all models predict that tropospheric O₃ will strongly decrease 314 worldwide, except in Equatorial Africa where higher O₃ levels are observed in GFDL-AM3, 315 GISS-E2-R and MOCAGE. In CESM-CAM, GFDL-AM3 and MIROC-CHEM, a 316 homogeneous decrease in O₃ burden is simulated worldwide while in GISS-E2-R, MOCAGE 317 and UM-CAM, the strongest decrease in surface O₃ mean concentrations are found where 318 high historical O₃ concentrations were reported. Under RCP4.5 scenario, the surface O₃ mean 319 320 concentrations and AOT40 values are lower than historical runs worldwide for all models except in MOCAGE where deterioration is observed over Canada, Greenland and East Asia. 321 322 For all models, the surface O₃ levels and AOT40 are higher for RCP8.5 as compared to 323 historical runs and the highest increases occur in the Northwestern America, Greenland, Mediterranean basin, Near East and East Asia. The AOT40 values, exceeding 70 ppm.h, are 324 found over the Tibetan plateau and in Near East and over Greenland. For RCP8.5, GFDL-325 AM3 is the most pessimistic model and MIROC-CHEM the most optimistic. By the end of 326 the 21st century, similar patterns are evident for RCP4.5 compared to RCP2.6 and RCP4.5 327 simulation is intermediate between RCP2.6 and RCP8.5 ones. 328 329 For all models and RCPs, the O_3 hot-spots (mean concentrations > 50 ppb and AOT40 > 70 330 ppm.h) are over Greenland and South Asia, in particular over the Tibetan plateau. The highest 331 increases are observed in NH, in particular in Northwestern America, Greenland, Near East 332 333 and South Asia (> 65 ppb). For the three RCPs, no significant change in tropospheric O₃ is observed in SH and the SH extratropics makes a small contribution to the overall change. 334 335 A recent global study showed the geographical patterns of surface air temperature differences 336 for late 21st century relative to the historical run (1986-2005) in all RCP scenarios (Nazarenko 337 et al., 2015). The global warming in the RCP2.6 scenario is 2-3 times smaller than RCP4.5 338 scenario and 4-5 times smaller than RCP8.5 scenario (Nazarenko et al., 2015). For the three 339 RCPs, the greatest change is observed over the Arctic, above latitude 60°N, and in the latitude 340 band 15-45°N (IPCC, 2014; Nazarenko et al., 2015). The least warming is simulated over the 341

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large area of the Southern Ocean. For RCP8.5 scenario, the global pattern of surface O₃ levels and AOT40 (Fig. 1-2) is similar to surface air temperature increase distribution. For RCP8.5, significant increases in air temperature are simulated over latitude 60°N and over the Tibetan plateau (more than 5°C). An increase of 4-5°C over the Near East, East and South Asia, North and South Africa and Canada are simulated as well as + 1-3°C for the rest of the world (Nazarenko et al., 2015). The tropospheric warming is stronger in the latitude band 15-45°N (Seidel et al., 2008) and Hudson et al. (2006) have demonstrated that O₃ trends over a 24-year period in the NH are due to trends in the relative area of the tropics and mid-latitudes and Polar Regions. All models are able to reproduce the global pattern of air temperature changes distribution in agreement with surface O₃ concentrations changes.

The spread in precursor emissions (e.g. VOCs, NOx, CO) is due to the range of representation of biogenic emissions (NOx from soils and lightning, CO from oceans and vegetation) as well as the complexity of chemical schemes in particular for NMVOCs simulations (e.g. isoprene) from explicitly specified to fully interactive with climate. RCP2.6 scenario has the lowest O₃ precursor concentrations, and RCP8.5 has relatively low NOx, CO and VOCs emissions, but very high CH₄ (Table 2b). The global emissions of NOx (-44%), VOCs (-5%) CO (-40%) and CH₄ burden (-27%) decline, while LNOx increase by e.g. 7% under RCP2.6 (Table 2b). The CO (-32%) and NOx (-20%) emissions have decreased while LNOX (+33%), VOCS (+1%) and CH₄ burden have increased (+120%) under RCP8.5 scenario (Table 2b). The GISS-E2-R model shows a greater degree of variation than other models, with a stronger increase in CH₄ burden (+ 153%) and in VOCs emissions (+ 20%) for RCP8.5 (Table 2b).

Excluding CH₄ burden and VOCs emissions, all the RCPs include reductions and redistributions of O₃ precursor emissions throughout the 21st century, due to the air pollution control strategies worldwide. The changes in CH₄ burden are due to the different climate policies in model assumptions. In RCP2.6, CH₄ emissions decrease steadily throughout the century, in RCP4.5 it remain steady until 2050 and then decrease (Voulgarakis et al., 2013) and in RCP8.5 (no climate policy) it rapidly increase compared to 2000. Methane burdens are fixed in the models with no sources, except for the GISS-E2-R simulations in which surface CH₄ emissions are prescribed for future rather than concentrations (Shindell et al., 2012). The model chemical schemes vary greatly in their complexity, mainly due to the NMVOCs simulations (Young et al. 2013). Isoprene dominates the total NMVOCs emissions (Guenther et al., 1995). Inversely to other models with constant present-day isoprene emissions, the

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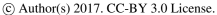
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376 GISS-ES2-R simulations incorporate climate-driven isoprene emissions, with greater BVOC emissions by 2100 and a positive change in total VOCs emissions across RCPs, related to the 377 378 positive correlation between air temperature and isoprene emission (e.g. Guenther et al., 2006; 379 Arneth et al., 2011; Young et al., 2013). 380 For RCP2.6 and RCP4.5 scenarios, there is a widespread decrease in O₃ in NH by 2100. The 381 overall decrease in O₃ concentration and AOT40 means for RCP4.5 are about half of that 382 383 between RCP2.6 and the historical simulation. For both scenarios, the changes are dominated by the decrease in O₃ precursor emissions in the NH extratropics compared to historical 384 simulations (Table 2b). In NOx saturated areas, annual mean O3 will slightly increase as a 385 result of a less efficient titration by NO, but the overall O₃ burden will decrease substantially 386 at hemispheric scale over time (Gao et al., 2013; Querol et al., 2014; Sicard et al., 2016a). In 387 RCP4.5, Gao et al. (2013) showed that the largest decrease in O₃ (4-10 ppb) occurs in summer 388 389 at mid-latitudes in the lower troposphere while the O₃ concentrations undergo an increase in winter. During the warm period, the photochemistry plays a major role in the O₃ production, 390 391 suggesting that the reduction in surface O₃ concentrations is in agreement with the large 392 reduction in anthropogenic O₃ precursor emissions (Sicard et al., 2016a) reducing the extent of regional photochemical O₃ formation (e.g. Derwent et al., 2013; Simpson et al., 2014). 393 394 Titration effect was also reported by Collette et al. (2012) over Europe by using six chemistry 395 transport models. 396 The O₃ increase can be also driven by the net impacts of climate change, i.e. increase in 397 stratospheric O₃ intrusion, changing LNOx and impacting reaction rates, through sea surface 398 temperatures and relative humidity changes (Lau et al., 2006; Voulgarakis et al., 2013; Young 399 et al., 2013). 400 401 402 Under the RCP8.5 scenario, the increase in surface O₃ concentrations, by 14% on average, can 403 be attributed to the higher CH₄ emissions coupled with a strong global warming, exceeding 2°C, and a weakened NO titration by reducing NOx emissions (Stevenson et al., 2013; Young 404 et al., 2013). The global CH₄ burden are 27% and 5% lower than 2000, for the RCP2.6 and 405 406 RCP4.5 scenarios respectively while for RCP8.5, the total CH₄ burden has more than doubled 407 compared to early 2000s and LNOx emissions increased by 33% (Table 2b). In addition, stronger increases are found over the high-elevation Himalayan Plateau reflecting increased 408 exchange with the free troposphere or stratosphere (Lefohn et al., 2012; Schnell et al., 2016). 409

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Several studies reported an increase in the stratospheric O₃ influx and higher stratospheric O₃ levels in response to a warming climate (e.g. Hegglin and Shepherd, 2009; Zeng et al., 2010). The downwards O₃ transport from the stratosphere is an important source of tropospheric O₃ (Hsu and Prather, 2009; Tang et al., 2011), therefore, stratospheric O₃ recovery also plays a partial role (e.g. + 11% for RCP8.5) in surface O₃ burden pattern. As an example, in MOCAGE, smaller reduction in global O₃ mean concentrations (-13%) and higher increase in stratospheric O₃ inputs (+20%) are observed for RCP2.6 (Table 3b). Similarly, for RCP8.5, the highest increase in O₃ mean concentrations (+23%) and stratospheric O₃ (+24%) are recorded in MOCAGE. In addition, lightning NOx emissions show significant upward trend from 2000 to 2100, in particular for the strongest warming scenario (RPC8.5) with greater convective and lightning activity (e.g. Williams, 2009; Lamarque et al., 2013). For RCP8.5, a reduction in surface O₃ concentrations is also simulated over the equatorial region, where the increased relative humidity, in a warmer climate, increases the O₃ loss rate (e.g. Johnson et al., 1999; Zeng and Pyle, 2003).

Risk areas for vegetation under RCP scenarios

Figure 3 shows the changes in the potential O_3 injury between present and future. It should be noted that a zero percentage of change (i.e. no change) for IO3, is simulated in sparsely vegetated regions (e.g. Gobi, Sahara, Near East, Western plateau and Greenland), while the change can be higher than 100% when the historical O_3 concentrations are lower than 40 ppb (i.e. AOT40 = 0 and IO3 = 0) and the O_3 concentrations exceed 40 ppb under RCPs (i.e. AOT40 > 0, IO3 > 0).

The potential O₃ impact for vegetation strongly decreases in NH for RCP2.6, except in MOCAGE where a slight increase in the risk factor (+ 15 %) is simulated at high latitudes and in South Asia. Conversely, the areas where the risk for vegetation increases (> 60 %) occur over Africa (+ 15% to + 60%) for all models, except in CESM-CAM where no change is observed across Africa. Under RCP4.5 scenario, the strongest increase in potential risk for vegetation (> + 60 %) is simulated by MOCAGE, markedly different from the other models, above the latitude 50°N. For all models, the potential O₃ impact for vegetation increases across Africa, from - 15% to + 60% while slight decreases or no change occur worldwide. Under RCP8.5 scenario, an increase of average O₃ over a significant part of the domain is simulated, therefore the exposure to O₃ pollution and impacts on vegetation will increase worldwide by 2100. An increase of the O₃ impacts on vegetation is simulated in Northern

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U.S., South America, Asia and Africa while a reduction in particular over Eastern U.S. and
 Southeastern China, and a slight increase (+ 15%) or decrease (- 15%) over Europe depending
 on the model, are simulated.

In summary, compared to the historical simulations, the averaged relative changes in the O₃ risk factor for the different RCPs are: - 61% for RCP2.6, - 47% for RCP4.5 and + 70% for RCP8.5 (Table 3d). We thus find a significant reduction in risk for vegetation for both RCP2.6 and RCP4.5 scenarios, except in South Africa and at high-latitudes in MOCAGE simulations, and a strong increase in global risk under RCP8.5. Under RCP2.6 and RCP4.5 scenarios, IO3 slightly increases in Africa and over North America and Asia (> latitude 60°N) in MOCAGE. The risk increases over the few areas where the O₃ concentrations increased between the historical period and 2100. Under both scenarios, the strongest reductions in risk are observed over Amazon, Central Africa and South Asia, i.e. where the O₃ concentrations have strongly declined between historical period and 2100. Under the RCP8.5, the areas where the highest projected O₃ mean concentrations are simulated (e.g. Greenland, deserts) are not associated to an increase in IO3 due to the absence of vegetation. Under RCP8.5, IO3 increases worldwide while a reduction is simulated over Southeast North America, northern Amazon, Central Africa and Southeast Asia, and a slighter reduction or a slight increase is simulated over Western Europe (depending on the model).

 The spatial pattern of IO3 is consistent with previous analyses on climate change and O₃ impacts on vegetation (e.g. Nemani et al., 2003; Zhu et al., 2016), i.e. the highest reduction in risk for vegetation, in particular under RCP8.5, occurs over areas where a strong increase in greening, LAI and NPP is observed due to global change and where a reduction in O₃ mean concentrations is found by 2100 (Fig. 1). The regions with the largest greening trends are in Southeast North America, northern Amazon, Europe, Central Africa and Southeast Asia with an average increase of the observed LAI exceeding 0.25 m² m⁻² per year (Zhu et al., 2016). The CO₂ fertilization effects (70%), nitrogen deposition (9%) and climate change (8%) explain the observed greening trend (Zhu et al., 2016). The changing climate alone produces persistent NPP increases and the regions with the highest increase in NPP, ranging from 1.0-1.5% per year, are in Southeast North America, northern Amazon, Western Europe, Central Africa and South Asia (Nemani et al., 2003). NPP increased by 6% globally between 1982 and 1999 and the highest increases are observed in tropical regions, with more than 1.5% per year over Amazon rainforest which accounts for 42% of the global NPP increase (Nemani et

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al., 2003). Amazon rainforest is one region where the effects are statistically significant. This is particularly important owing to the role of the Amazon rainforests in the global carbon cycle (Zhu et al., 2016). In these areas, the increasing effect of a warming climate on forests (e.g. increase of greening, LAI) is higher than the reduction in GPP due to O₃. Inversely, the risk for vegetation increases in particular in Africa, e.g. western Africa along the Gulf of Guinea, in South Brazil and over high-latitudes regions (> 60°N) in North America and Asia where a reduction or a slight increase in LAI (from - 0.05 to + 0.03 m² m⁻² per year) and strong decreases, by 1.0-1.5% per year, in NPP are simulated (Nemani et al., 2003; Zhu et al., 2016).

Our results are not in agreement with the high GPP reduction, due to O₃ effects, simulated by Sitch et al. (2007) between 1901 and 2100, with a projected GPP reduction exceeding 30% over Western Europe, eastern and western North America, Amazon, central Africa and East Asia where higher surface O₃ mean concentrations were projected. Previous studies reported that the reductions in GPP simulated by Sitch et al. (2007) are overestimated up to six times (Ren et al., 2011; Zak et al., 2011; Kvaleveg and Myhre 2013), mainly due to the lack of empirical data about the response of different species to O₃, the fact that a few experiments have shown no response, e.g. grasslands (Bassin et al., 2013), and the non-inclusion of the nitrogen limitation of growth (Kvalevag and Myhre, 2013).

The projected land covers widely vary under RCPs (Betts et al., 2015). In RCP2.6 scenario, the ground surface covered by croplands increases as a result of bio-energy production, with a more-or-less constant use of grassland. The RCP4.5 scenario focuses on global reforestation programs as part of global climate policy, as a result, the use of cropland and grassland decreases. Under RCP8.5, an increase in croplands and grasslands is applied mostly driven by an increasing global population (van Vuuren et al., 2011). Generally, the risk for vegetation strongly increases over shrublands (e.g. high-latitude region, Australia, South Africa) and savannas (e.g. South Brazil, Africa) and the risk decreases over forests, strongly over evergreen broadleaf forest and deciduous woodland over Africa and Amazon rainforests, and slighter over needleleaf forests in Northern America (Canada) and Northern Asia. The risk slightly decreases or slightly increases over grasslands (Central Asia and central Africa and U.S.). The largest decreases (50-80%) under RCP8.5 occur in Eastern U.S., Europe and Southeastern China, where the ground is mainly dominated by croplands, in all models except CESM-CAM.

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Conclusions

From six global atmospheric chemistry transport models, we illustrate the changes, i.e. differences for late 21^{st} century relative to the historical run, in ground-level O_3 concentrations and vegetation impact metric (AOT40). *In fine*, the potential O_3 impacts on vegetation worldwide are investigated to define potential risk areas for vegetation at global

scale by 2100.

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The six models are able to well reproduce the spatial pattern of historical O₃ concentration and AOT40 at global scale, in particular GISS-E2-R and MOCAGE are able to simulate the higher O₃ levels in areas downwind of precursor sources and at the high-elevation areas. The model outputs emphasize the strong asymmetry in the tropospheric O₃ distribution between NH and SH; substantially higher O₃ mean concentrations are observed in the NH (ca. 38 ppb), particularly in the latitude band 15-45°N, than in the SH (ca. 23 ppb). The natural emissions of O₃ precursors (e.g. lightning NOx, CO from oceans, isoprene) as well as the complexity of chemical schemes are significant sources of model-to-model differences.

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In this study, the projected mean tropospheric O₃ concentrations and AOT40 dependent on 528 global and regional emission pathways. Compared to early 2000s, the results suggest changes 529 in surface O_3 of - 9.5 ± 2.0 ppb (NH) and - 1.8 ± 2.1 ppb (SH) in the cleaner RCP2.6 scenario 530 and of $+4.4 \pm 2.8$ ppb (NH) and $+5.1 \pm 2.1$ ppb (SH) in RCP8.5 scenario. For RCP2.6 and 531 RCP4.5, absolute decreases are observed for the Mediterranean basin and the Western U.S. 532 due to less precursor emissions in the NH extratropics (e.g. reduction of 5-7 ppb over 533 Europe). Smaller reduction in surface O₃ levels in South and East Asia highlight the smaller 534 535 changes in O₃ precursor emissions due to the recent emission growth in this region (e.g. Zhang et al., 2009; Xing et al., 2015). For RCP8.5, all models show climate-driven increases 536 in ground-level O₃ in particular over the Western U.S, Greenland, South Asia and Northeast 537 538 China. The changes in surface O₃ over North America and Europe ranged from + 1-5 ppb 539 under RCP8.5. South Asia sees the greatest increase, up to more than 10 ppb for RCP 8.5. The O₃ increase can be attributed to substantial increase in CH₄ emissions coupled with a strong 540 global warming, exceeding 2°C, and a weakened NO titration and a greater stratospheric O₃ 541 influx (Kawase et al., 2011; Wild et al., 2012; Young et al., 2013). A decline in CH₄ 542 543 emissions will undoubtedly benefit future O₃ control.

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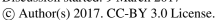
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545 The current surface O₃ levels (35-50 ppb in NH) are high enough to damage both forests and crops. About 50% of forests, grasslands and croplands might be exposed to high O₃ levels by 546 the end of the 21st century (Sitch et al., 2007; Wittig et al. 2009). Most important results from 547 the study are the significant overrun of exposure metric (AOT40) in comparison with the 548 AOT40-based critical level for the protection of forests (5 ppm.h) and crops (3 ppm.h). The 549 global models suggest that exposure-based critical levels will be exceeded over many areas of 550 the NH, and in parts of North America, East and South Asia they may be exceeded by a factor 551 exceeding 10 under RCP8.5. The critical level were defined for boreal and temperate 552 deciduous tree species, i.e. more consistent for regions in the latitude band 35-60°N. To 553 protect vegetation, the current AOT40 index appears inadequate for a realistic quantification 554 of O₃ impacts on vegetation (Paoletti and Manning, 2007; Mills et al., 2011; De Marco et al., 555 2015; Sicard et al., 2016b,c). As a result, in the last decade, the United Nations Convention on 556 Long-Range Transboundary Air Pollution (CLRTAP) has introduced the flux-based metric 557 for vegetation protection against effects of O₃, taking into account the modifying effects of 558 multiple climatic and phenological factors on O₃ uptake (Paoletti and Manning, 2007; Sicard 559 560 et al., 2016b,c). Ozone may be a major threat to biodiversity over large regions of the world (Sicard et al., 561 562 2016b), however the size of these areas remains uncertain. The potential O₃ impact on assimilation, IO3, provides a clear indicator of the potential risk to vegetation. The risk for 563 564 vegetation decreases by about 61% and 47% under RCP2.6 and RCP4.5, respectively and increases by 70% under RCP8.5, compared to early 2000s over the whole domain by 2100 565 and that the potential risk areas for vegetation vary worldwide according to the dominant 566 vegetation cover. The strongest increase of the O₃ impacts on vegetation is simulated in 567 Northern America and Asia and central Africa. The highest reduction in risk for vegetation 568 (i.e. Southeast North America, the northern Amazon, Central Africa and Southeast Asia) 569 occurs over areas where a strong increase in greening, LAI and NPP is observed and where a 570 571 reduction in O₃ mean concentrations is found by 2100. 572 Trees possess a defence capacity, e.g. through antioxidant activity and a capacity of repairing 573 injured tissues (Paoletti, 2007). The short-term response to O₃ is a reduction in productivity of 574 575 crops and forests and long-term changes in community composition could be observed due to species-specific O₃-sensitivity (Wittig et al., 2009). Generally, deciduous woodland are highly 576

O₃-sensitive risk areas, grasslands and needleleaf forests are moderately O₃-sensitive risk

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areas while the lower risk areas include evergreen broadleaf forests. However, crops are more sensitive to O_3 exposure than trees and deciduous trees are more sensitive than coniferous trees with lower stomatal conductance (Felzer et al., 2004; Ren et al., 2007; Wittig et al. 2009; Anav et al., 2011). To efficiently protect vegetation against O_3 pollution, suitable standards taking into account the detoxification processes (e.g. flux-based metric) are urgently needed.

As the vegetation atmosphere feedbacks are still under investigated, e.g. impacts of changes of vegetation on air chemistry, we recommend the use of improved chemistry-climate modelling system, fully coupled with dynamic vegetation models, to perform high resolution simulations and to better evaluate the regional exposure of ecosystems to air pollution.

 The risk reduction is possible through climate-change mitigation, e.g. reductions in air pollution, and adaptation actions. An efficient reduction in overall O₃ levels is expected over North America and Europe in all RCP scenarios and worldwide if CH₄ emissions are reduced (e.g. Kirtman et al., 2013; Pfister et al., 2014; Schnell et al., 2016). However, the increasing effect of a warming climate on surface O₃ concentrations is higher than the reduction achieved by the decline in O₃ precursor emissions (Revell et al., 2015; Hendriks et al., 2016), therefore, climate change and the measures and policies in e.g. Asia will need to be factored into future O₃ policies (Wilson et al., 2012; Lefohn and Cooper, 2015). Many ecosystems worldwide are unprotected from O₃ due to the lack of international efforts (Emberson et al., 2014). To be efficient, the mitigation actions for O₃ impacts on biodiversity must be as part of international emission reduction programmes.

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Figure 1: Surface ozone mean concentrations (in ppb) at the lower model layer for each ACCMIP model for the historical run and for RCP2.6, RCP4.5 and RCP8.5 simulations by 2100.

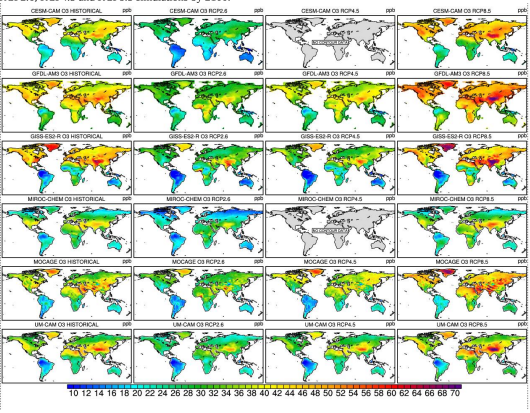






Figure 2: Surface AOT40 means (in ppm.h) at the lower model layer for each ACCMIP model for the historical run and for RCP2.6, RCP4.5 and RCP8.5 simulations by 2100.

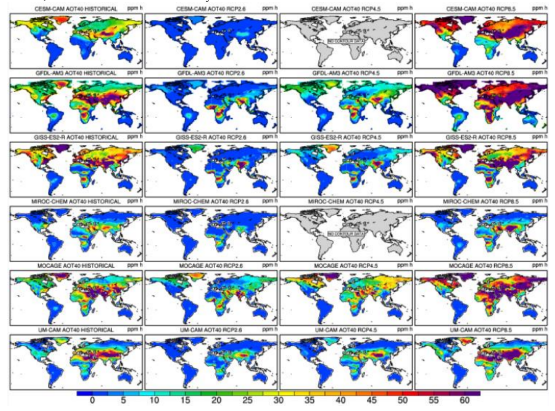






Figure 3: Simulated percentage changes (%) in the potential ozone impact on vegetation (IO3) for each ACCMIP model between RCP2.6, RCP4.5 and RCP8.5 simulations and the historical run.

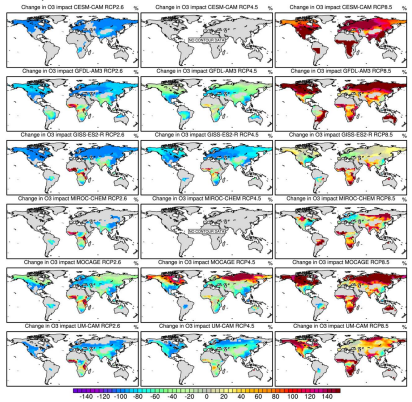






 Table 1: Characteristics of the models, including simulation time slice, spatial resolution, simulated gas species and associated bibliographic references (from Lamarque et al., 2013 and Young et al. 2013). Black carbon (BC), Organic carbon (OC), Secondary Organic Aerosols (SOA), Dimethylsulfide (DMS), Chemistry Climate Model (CCM), Chemistry Transport Model (CTM), Chemistry-General Circulation Model (CGCM).

Models	Туре	Simulation length	Resolution (lat/lon)	Number of vertical pressure levels & top level	Species simulated	References
CESM-CAM	ССМ	2000-2009 and 2100-2109	1.875/2.5	26 levels 3.5 hPa	16 gas species; constant present-day isoprene, soil NOx, DMS and volcanic sulfur, oceanic CO.	Lamarque et al., 2012
GFDL-AM3	ССМ	2001-2010 and 2101-2110	2.0/2.5	48 levels 0.017 hPa	81 gas species; SOx, BC, OC, SOA, NH ₃ , NO ₃ ; constant pre- industrial soil NOx; constant present-day soil and oceanic CO, and biogenic VOC; climate-sensitive dust, sea salt, and DMS.	Donner et al., 2011 Naik et al., 2012
GISS-E2-R	ССМ	2000-2004 and 2101-2105	2.0/2.5	40 levels 0.14 hPa	51 gas species; interactive sulfate, BC, OC, sea salt, dust, NO ₃ , SOA, alkenes; constant present-day soil NOx; climate-sensitive dust, sea salt, and DMS; climate-sensitive isoprene based on present-day vegetation.	Lee and Adams, 2011 Shindell et al., 2012
MIROC-CHEM	ССМ	2000-2010 and 2100-2104	2.8/2.8	80 levels 0.003 hPa	58 gas species ; SO ₄ , BC, OC; constant present-day VOCs, soil-NOx, oceanic-CO; climate-sensitive dust, sea salt and DMS.	Watanabe et al., 2011
MOCAGE	СТМ	2000-2003 and 2100-2103	2.0/2.0	47 levels 6.9 hPa	110 gas species; constant present-day isoprene, other VOCs, oceanic CO and soil NOx.	Josse et al., 2004 Krinner et al., 2005 Teyssèdre et al., 2007
UM-CAM	CGCM	2000-2005 and 2094-2099	2.50/3.75	19 levels 4.6 hPa	60 gas species; constant present-day biogenic isoprene, soil NOx, biogenic and oceanic CO.	Zeng et al., 2008, 2010





Table 2a: Annual total emissions of CO (Tg CO/year), NMVOCs (Tg C/year), NOx (Tg N/year, including lightning and soil NOx), total lightning NOx emissions (LNOx) and global atmospheric methane (CH₄) burden (Tg) for the historical simulations in each model (from Young et al., 2013 and * from Voulgarakis et al., 2013).

Models	Historical								
Models	CO	* CH ₄	NMVOCs	NOx	*LNOx				
CESM-CAM	1248	4902	429	50.0	4.2				
GFDL-AM3	1246	4809	830	46.2	4.4				
GISS-E2-R	1070	4793	830	48.6	7.7				
MIROC-	1064	4805	833	57.3	9.7				
CHEM									
MOCAGE	1168	4678	1059	47.9	5.2				
UM-CAM	1148	4879	535	49.2	5.1				

Table 2b: Simulated percentage (%) changes in total emissions of CO, NMVOCs, NOx (including lightning and soil NOx), total lightning NOx emissions (LNOx) and global atmospheric CH₄ burden for each model between 2100 and historical simulation for RCPs (from Young et al., 2013 and *Voulgarakis et al., 2013). The last row shows means and standard deviations (SD). Missing or not available data are identified (n.a).

Models	RCP2.6 scenario					RCP4.5 scenario				RCP8.5 scenario					
Models	CO	*CH ₄	VOCs	NOx	*LNOx	CO	*CH ₄	VOCs	NOx	*LNOx	CO	*CH ₄	VOCs	NOx	*LNOx
CESM-CAM	- 36.7	- 27.1	0	- 52.8	+ 7.1	n.a	n.a	n.a	n.a	n.a	- 30.1	+ 112.1	0	33.0	+ 29.7
GFDL-AM3	- 36.9	- 27.9	- 5.0	- 47.0	+ 12.6	- 47.4	- 9.3	- 3.6	- 41.5	+ 23.5	- 30.3	+ 116.1	- 1.9	22.4	+ 38.2
GISS-E2-R	- 42.8	- 21.0	+ 0.5	- 44.2	+ 3.8	- 54.9	+ 4.6	+ 6.9	- 39.2	+ 12.2	- 35.1	+ 152.7	+ 19.8	20.0	+ 26.2
MIROC-CHEM	- 43.1	- 28.2	- 7.1	- 36.0	+ 7.5	n.a	n.a	n.a	n.a	n.a	- 35.4	+ 116.0	- 3.4	- 6.9	+ 38.0
MOCAGE	- 39.4	- 28.8	- 6.5	- 45.7	+ 5.2	n.a	n.a	n.a	n.a	n.a	- 32.3	+ 113.4	- 2.8	22.9	+ 19.9
UM-CAM	- 39.0	- 27.9	- 11.3	- 40.6	+ 8.1	- 50.4	- 8.7	- 9.2	- 36.0	+ 17.5	- 32.0	+ 112.1	- 4.2	17.2	+ 43.6
Mean ± SD	- 39.7 ± 2.2	- 26.8 ± 3.7	- 4.9 ± 4.9	- 44.4 ± 4.3	+ 7.4 ± 2.0	- 50.9 ± 3.2	- 4.5 ± 9.4	- 2.0 ± 11.4	- 38.9 ± 2.3	+ 17.7 ± 3.7	- 32.5 ± 1.8	+ 120.4 ± 19.5	+ 1.3 ± 11.6	- 20.4 ± 7.0	+ 32.6 ± 10.8





Table 3a: Global and hemispheric (averaged over the domain) surface ozone mean concentrations (in ppb) and AOT40 means (in ppm.h) for the historical simulations in each model (North and South Hemisphere, i.e NH and SH). The last row shows means and standard deviations (SD).

Models	Ozone conc. global	Ozone conc. SH	Ozone conc. NH	AOT40 global	AOT40 SH	AOT40 NH
CESM-CAM	31.3	20.9	36.4	12.8	0.2	18.9
GFDL-AM3	38.6	30.6	42.9	21.8	4.7	30.8
GISS-E2-R	35.8	22.3	42.3	26.0	3.6	36.8
MIROC-CHEM	27.9	20.4	31.4	7.3	1.9	9.8
MOCAGE	32.9	21.5	38.3	22.9	3.5	31.8
UM-CAM	31.3	21.4	36.0	14.4	1.3	20.6
Mean ± SD	33.0 ± 3.8	22.9 ± 3.8	37.9 ± 4.3	17.5 ± 7.2	2.5 ± 1.7	24.8 ± 10.1

Table 3b: Simulated percentage (%) changes in global and hemispheric surface ozone mean concentrations and in global mean stratospheric ozone column (* from Voulgarakis et al., 2013) for each model between 2100 and historical simulation for RCPs (North and South Hemisphere, i.e NH and SH). The last row shows means and standard deviations (SD). Missing or not available data are identified (n.a).

	Surface ozone mean concentrations									* Stratospheric ozone			
Models	RCP2.6	RCP2.6	RCP2.6	RCP4.5	RCP4.5	RCP4.5	RCP8.5	RCP8.5	RCP8.5	RCP2.6	RCP4.5	RCP8.5	
	global	SH	NH	global	SH	NH	global	SH	NH	global	global	global	
CESM-CAM	- 29.1	- 20.6	- 31.3	n.a	n.a	n.a	+ 21.9	+ 22.5	+ 20.5	n.a	n.a	+ 5.3	
GFDL-AM3	- 20.5	- 10.8	- 24.5	- 11.7	- 6.9	- 13.5	+ 15.5	+ 18.6	+ 14.5	+ 3.3	+ 3.9	+ 8.4	
GISS-E2-R	- 23.5	- 5.8	- 27.9	- 20.4	- 6.3	- 23.9	+ 7.0	+ 19.3	+ 3.8	+ 8.0	+ 8.8	+ 15.1	
MIROC-CHEM	- 23.3	- 12.3	- 26.8	n.a	n.a	n.a	+ 3.9	+ 10.3	+ 2.2	+ 2.6	n.a	+ 4.2	
MOCAGE	- 12.8	+ 7.4	- 18.5	- 1.8	+ 17.7	- 7.0	+ 23.1	+ 40.4	+ 16.7	+ 19.9	n.a	+ 23.6	
UM-CAM	- 17.3	- 4.7	- 21.1	- 8.3	+ 0.9	- 10.8	+ 14.4	+ 24.3	+ 11.4	+ 6.7	+ 6.9	+ 7.4	
Mean ± SD	- 21.1 ± 5.6	- 7.8 ± 9.4	- 25.0 ± 4.7	- 10.5 ± 7.7	+ 1.4 ± 11.5	- 13.8 ± 7.2	+ 13.8 ± 7.1	+ 22.6 ± 10.0	$+ 11.5 \pm 7.3$	$+8.1 \pm 7.0$	$+6.5 \pm 2.5$	+ 10.7 ± 7.4	

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Table 3c: Simulated percentage (%) changes in global and hemispheric AOT40 means for each model between 2100 and historical simulation for RCPs (North and South Hemisphere, i.e NH and SH). Missing or not available data are identified (n.a).

		AOT40												
Models	RCP2.6	RCP2.6 SH	RCP2.6 NH	RCP4.5	RCP4.5	RCP4.5	RCP8.5	RCP8.5	RCP8.5 NH					
	global			global	SH	NH	global	SH						
CESM-CAM	- 96.9	- 99.9	- 96.8	n.a	n.a	n.a	+ 138.3	+ 150.0	+ 134.9					
GFDL-AM3	- 75.2	- 25.5	- 78.9	- 53.2	- 36.2	- 54.5	+ 96.3	+ 242.5	+ 85.1					
GISS-E2-R	- 78.1	- 13.9	- 81.2	- 75.0	- 27.8	- 77.2	+ 22.3	+ 83.3	+ 19.5					
MIROC-CHEM	- 74.0	- 10.5	- 80.6	n.a	n.a	n.a	+ 20.5	+ 78.9	+ 16.3					
MOCAGE	- 53.7	+ 68.6	- 59.7	- 17.5	+ 202.9	- 28.3	+ 85.1	+ 448.6	+ 67.0					
UM-CAM	- 73.6	+ 92.3	- 76.7	- 52.8	+7.7	- 54.8	+ 49.3	+ 176.9	+ 45.1					
Mean ± SD	- 75.2 ± 13.7	$+1.9 \pm 69.5$	- 79.0 ± 11.8	- 49.6 ± 23.8	+ 36.6 ± 112.4	- 53.7 ± 20.0	+ 68.6 ± 46.3	+ 196.7 ± 137.7	$+61.3 \pm 44.8$					

Table 3d: Simulated percentage (%) changes in potential O_3 impact on vegetation (IO3) for each model between 2100 and historical simulation for RCPs (North and South Hemisphere, i.e NH and SH). Missing or not available data are identified (n.a).

		Risk factor IO3												
Models	RCP2.6 global	RCP2.6 SH	RCP2.6 NH	RCP4.5 global	RCP4.5 SH	RCP4.5 NH	RCP8.5 global	RCP8.5 SH	RCP8.5 NH					
CESM-CAM	- 97.2	- 91.8	-97.5	n.a	n.a	n.a	+ 129.6	+146.8	+127.5					
GFDL-AM3	- 69.4	- 49.1	- 74.8	- 50.1	- 61.1	- 47.2	+ 91.9	+95.5	+90.4					
GISS-E2-R	- 66.1	- 20.7	- 74.3	- 71.7	- 53.3	- 74.6	+ 21.5	+56.6	+14.2					
MIROC-CHEM	- 41.4	- 18.9	- 51.9	n.a	n.a	n.a	+ 41.0	+103.8	+25.5					
MOCAGE	- 46.6	-22.8	- 51.4	- 7.0	- 38.0	- 1.0	+ 77.7	+68.2	+80.0					
UM-CAM	- 45.8	- 9.2	- 71.3	- 59.5	+ 2.0	- 69.0	+ 61.3	+84.2	+56.0					
Mean ± SD	- 61.1 ± 21.1	- 35.5 ± 30.7	- 70.2 ± 17.2	- 47.1 ± 28.1	- 37.6 ± 28.1	- 47.9 ± 33.4	+ 70.5 ± 38.4	$+92.5 \pm 31.7$	$+65.6 \pm 42.4$					