Authors' response

To gmd-2019-21 editor decision (04 Oct 2019): Thank you for your detailed comments regarding grammar and language. We appreciate these and changed our text accordingly. In the following, we will respond to the questions in detail.

- Page 1, Abstract, line 19: Change "associated to" to "associated with" and change "resistance the ocean" to "resistance
- over the ocean" Changed accordingly.
 - Page 2, Introduction, line 15: Replace "else" with "otherwise" Changed accordingly.
 - Page 2, Introduction, line 34: Replace "observation" with "observations" Changed accordingly.
 - Page 3, Introduction, line 2: Change "sides" to "sites" Changed accordingly.
 - Page 3, Introduction, line 15: Worth noting that those early observations were subject to interference by other species
 e.g. SO2 Thank you for pointing this out. We add this remark and write: *These early observations, however, were subject to interference by other species, e.g.*, SO₂.
 - Page 3, Introduction, line 27: Worth saying at this stage why the Luhar study gives lower estimates We add: In particular, Luhar et al. (2018) found that the average surface resistance of ozone over ocean ($r_c = 2200 \,\mathrm{sm}^{-1}$) is highly overestimated in most models. We add to sentence P3L14: [...] indicates much lower amounts [...] due to lower dry deposition to the oceans.
 - Page 4, Introduction, line 5: Change to "as the beginning and duration of the greening season" Changed accordingly.
 - Page 4, Section 2, line 15/16: Change to "publication focussing on this is planned" Changed accordingly.
 - Page 4, Section 2, line 17: Change "is given in a" to "is given at a" Changed accordingly.
 - Page 7, Section 2.1.3, line 2: Remove "," after both Changed accordingly.
- Page 7, Section 2.1.3, equation 12: I'm not clear what the index m is referring to can you explain or exclude if not necessary? The index m refers to molar. We made this more clear. We add molar to the sentence preceding the equation to [...] the leaf-level molar stomatal conductance in the mOSaic scheme [...]. And we move the last paragraph up and change it accordingly so that it follows the equation: The maximum molar stomatal conductance is given by g^k_{max,m} which is in units of mmols⁻¹ m⁻². A unit conversion to ms⁻¹ is necessary in our model: [...] To annotate the differing units, we use the index m in Eq. ([...]).
 - Page 7, Section 2.1.3, line 27: Change to "start of the greening" Changed accordingly.
 - Page 9, Section 2.1.3, line 3: Change "land surface types k" to "land surface type k" Changed accordingly.
 - Page 9, Section 2.1.3, line 4: Change "others" to "other" Changed accordingly.
 - Page 10, Section 2.2, line 28: Change "planed" to "planned" Changed accordingly.
 - Page 11, Section 2.2.1, line 10: Give web address Changed accordingly.
 - Page 12, Section 2.2.1, line 5: Change "planed" to "planned" Check for other examples Changed accordingly.
 - Page 12, Section 2.2.2, line 13: Change to "spectrally" Changed accordingly.
 - Page 13, Section 3, line 14: Give reference for CEDS here Changed accordingly.
 - Page 13, Section 3, line 17: Can you provide global annual total? NO_x emissions from soil and vegetation in the Oslo CTM3 are estimated to amount to $6.55 Tg(N)a^{-1}$. We add this accordingly.

10

15

30

35

5

- Page 14, Section 3.1, line 13: Change to "we show global distributions of the relative difference between mosaic and wesely_type for surface ozone, dry deposition velocity and total ozone dry deposition" Changed accordingly.
- Page 14, Section 3.1, line 14: Change "ozone burden" to "surface ozone" Changed accordingly.
- Page 14, Section 3.1, line 16: Change "then" to "than" Changed accordingly.
- Page 15, Section 3.2, line 5: Change to "the multi-model mean of the Task ... models" Changed accordingly.
 - Page 16, Section 3.2, line 5: Change to "available" Changed accordingly.

5

10

20

25

- Page 16, Section 3.2.1, line 7: Change "in consistency with" to "consistent with" Changed accordingly.
- Page 16, Section 3.2.1, line 10: Change to "hemispheres" Changed accordingly.
- Page 16, Section 3.2.1, line 11: Change to "focussing" Changed accordingly.
- Page 16, Section 3.2.1, line 28: Change to "as the new" Changed accordingly.
 - Page 16, Section 3.2.1, line 29: Change to "Arctic" Changed accordingly.
 - Page 16, Section 3.2.1, line 30: Change to "strong sensitivity to" Changed accordingly.
 - Page 17, Section 3.2.1, line 11: Change to "account" Changed accordingly.
 - Page 17, Section 3.2.1, line 11: Change to "but do not" Changed accordingly.
- 15 Page 17, Section 3.2.1, line 15: Change to "matches better" Changed accordingly.
 - Page 17, Section 3.2.1, line 21: Change to "as a reference" Also, in section 3.2.2 Changed accordingly.
 - Page 19, Section 3.2.2, line 12: Remove "both" Changed accordingly.
 - Page 19, Section 3.2.2, line 21: Change "regarding" to "for" Changed accordingly.
 - Page 19, Section 3.2.2, line 23: Change "In the case of" Changed accordingly.
 - Page 20, Section 3.2.2, line 19: Change to "in the Oslo" Changed accordingly.
 - Page 22, Section 3.2.2, line 6: Change "responses" to "responds" Changed accordingly.
 - Page 22, Section 3.3, line 16: It is unclear what you mean by "While on global average much more concise with the MACC..." We make our point more clear and change the sentences to: *Regarding the global average surface ozone concentration*, Wesely_type (Supplement S.8) is more consistent with the MACC-reanalysis than mOSaic, but shows the same tendency of enhanced ozone over the continents as the latter. This enhancement is apparent mostly in the deep tropics and over the north-western Indian subcontinent which coincides with regions of high intensity in incoming UV radiation.
 - Page 23, Section 3.4, line 8: Change to "At 4 of the 6 sites" and again, later on in the same section Changed accordingly.
 - Page 25, Section 4, line 4: Correct spelling i.e. "mosaic" Changed accordingly.
- Page 25, Section 4, line 6: Change to "(with the latter" Changed accordingly.
 - Page 25, Section 4, line 22: Change to "the operational" Changed accordingly.
 - Page 25, Section 4, line 30: Change to "In the case of" Changed accordingly.

- Page 25, Section 4, line 30: Change "is in the order of 1 magnitude" to "is an order of magnitude" Changed accordingly.
- Page 26, Section 4, line 3: Change to "breaking" Changed accordingly.
- Page 26, Section 4, line 6: Change to "Arctic" Changed accordingly.
- Page 26, Section 4, line 11: Remove "total" Changed accordingly.
- Page 26, Section 4, line 14: Change to "to changes of the order of" Changed accordingly.
 - Page 27: Section 4: Change all references of "arctic" to "Arctic" Changed accordingly.
 - Page 27, Section 4, line 3: Change to "affect stratospheric" Changed accordingly.
 - Page 27, Section 4, line 7/8: Change to "at higher" Changed accordingly.
 - Author contributions: Replace reference to EMEP scheme! Changed accordingly.

Update and evaluation of the ozone dry deposition in the Oslo CTM3 v1.0

Stefanie Falk¹ and Amund Søvde Haslerud^{2,a}

¹Department of Geosciences, University of Oslo, Oslo, Norway
²CICERO Center for International Climate Research, Oslo, Norway
^aKjeller Vindteknikk, Kjeller, Norway

Correspondence: Stefanie Falk (stefanie.falk@geo.uio.no)

Abstract. High concentrations of ozone in ambient air are hazardous not only to humans but to the ecosystem in general. The impact of ozone damage on vegetation and agricultural plants in combination with advancing climate change may affect food security in the future. While the future scenarios in themselves are uncertain, there are limiting factors constraining the accuracy of surface ozone modeling also at present: The distribution and amount of ozone precursors and ozone depleting substances, the

- 5 stratosphere-troposphere exchange as well as scavenging processes. Removal of any substance through gravitational settling or by uptake by plants and soil is referred to as dry deposition. The process of dry deposition is important for predicting surface ozone concentrations and understanding the observed amount and increase of tropospheric background ozone. The conceptual dry deposition velocities are calculated following a resistance-analogous approach wherein aerodynamic, quasi laminar, and canopy resistances are key components, but these are hard to measure explicitly. We present an update of the dry deposition
- 10 scheme implemented in the Oslo CTM3. We change from a purely empirical dry deposition parameterization to a more processbased one which is taking the state of the atmosphere and vegetation into account. We examine the sensitivity of the scheme to various parameters, e.g. the stomatal conductance-based description of the canopy resistance and the choice of ozone surface resistance, and evaluate the resulting modeled ozone dry deposition with respect to observations and multi-model studies. Individual dry deposition velocities are now available for each land surface type and agree generally well with observations.
- We also estimate the impact on the modeled ozone concentrations at the surface. We show that the global annual total ozone dry deposition decreases with respect to the previous model version (-37%), leading to an increase in surface ozone of more than 100% in some regions. While high sensitivity to changes in dry deposition to vegetation is found in the tropics and the northern hemisphere, the largest impact on global scales is associated to with the choice of prescribed ozone surface resistance over the ocean and deserts.

20 1 Introduction

Ozone is an important trace gas for all lifeforms on Earth. Depending on the place of its occurrence it has either a positive or negative connotation. In the stratosphere, ozone absorbs most of the ultraviolet (UV)-light from the sun within the range of 100–315 nm, thus shielding the Earth's surface from the most harmful UV-radiation. In addition, ozone is a potent greenhouse gas in both, stratosphere and troposphere. With a radiative forcing of $0.40 \pm 0.20 \text{ W m}^{-2}$, it is placed third, only surpassed by

CO₂ and CH₄ (IPCC - Intergovernmental Panel on Climate Change, 2013, Chapter 8).

In the troposphere and in particular in ambient air, ozone is considered as a highly toxic pollutant. Since the industrial revolution, tropospheric background ozone concentrations have been increasing in the northern hemisphere (IPCC - Intergovernmental Panel on Climate Change, 2013, Chapter 2). In recent years, the number of episodes of peak concentrations has been,

- 5 in general, decreasing in North America and Europe due to the implementation of air quality regulations (e.g., Fleming et al., 2018; Mills et al., 2018). At the same time, fast developing countries, like e.g., China or India, saw a significant increase in ozone related air pollution. Continuously high concentrations of ambient air ozone are hazardous to the whole ecosystem. It is estimated that ozone is cause to an increase in pre-mature deaths (WHO World Health Organization, 2008), an average global loss of yield in the four major crops (wheat, rice, maize, and soybean) of about 3 15% (Ainsworth, 2017) as well as 7% loss
- 10 in primary production in forestry (Wittig et al., 2009; Matyssek et al., 2012). The impact of ozone damage on vegetation and agricultural plants may affect food security in the future especially in Asia (Tang et al., 2013; Tai et al., 2014; Chuwah et al., 2015; Mills et al., 2018) and might be an important additional feedback to climate change (Sitch et al., 2007). Elevated ozone levels at a site may originate from both, the local production of ozone from its precursors, which are transported,

and from advection of ozone itself. Long-range ozone transport occurs regularly and might be most important in regions that

15 else otherwise lack precursors. Tropospheric ozone is produced in complex photochemical cycles involving precursor gases such as carbon monoxide (CO) or volatile organic substances (VOCs – also known as hydrocarbons) in the presents of nitrogen oxides (NO_x). A typical reaction mechanism for CO is sketched in the following. In a sequence of rapid reactions a peroxyl radical HO^o is formed through an initial reaction of CO with a hydroxyl radical •OH. Via a reaction between HO^o and NO, NO₂ is formed which is then photolyzed. The resulting atomic oxygen reacts then with O₂ (and also under the presence of available co-reactants) to form an ozone molecule. Such a cycle leads to a net production via:

$$\mathrm{CO} + 2\mathrm{O}_2 + h\nu \to \mathrm{CO}_2 + \mathrm{O}_3. \tag{R1}$$

Similar cycles involving VOCs exist (Monks et al., 2015). Another source of tropospheric ozone is downward transport from the stratosphere via stratosphere-troposphere exchange (STE) (WMO - Global Ozone Research and Monitoring Project, 2014). Based on observations, STE might only amount to 10% (550 ± 140 Tg a⁻¹) of the total global ozone budget in the troposphere, while ozone from chemical production is estimated to be 5000 Tg a⁻¹ (Monks et al., 2015). Ozone is removed from the atmosphere by photochemical reactions or scavenging processes. Major sinks are photolysis followed by a reaction with water vapor to from OH, reactions with HO₂, titration reactions, and dry deposition. We will come back to the latter later in this section and cover the implemented scheme in more detail in Section 2.1.

25

Since ozone is highly reactive, its global mean life-time in the troposphere is roughly 22 days but ranges between a few days in the tropical boundary layer to up to 1 year in the upper troposphere (Stevenson et al., 2005; Young et al., 2013). The abundance of tropospheric ozone therefore varies, e.g., with time of the day, season, altitude, location (Schnell et al., 2015), or weather conditions in general (Otero et al., 2018). Typical concentrations of surface ozone range from 10 ppb over the tropical Pacific to 100 ppb in the downwind areas of highly emitting sources (IPCC - Intergovernmental Panel on Climate Change, 2013, Chapter 8). This variability poses a challenge on both, trend analysis from observation observations as well as validation and intercomparison of models. From the observational side, the number of long-term observations (started before the 1950s) is limited and restricted to mainly European sidessites. Most of these indicate a doubling of tropospheric ozone since the 1950s (IPCC - Intergovernmental Panel on Climate Change, 2013, Chapter 2). But especially the very low pre-industrial ozone abundance cannot be reproduced by the likes of most models. These early observations, however, were subject to interference

- 5 by other species, e.g., SO₂, Among the participating models in the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP), there is a general tendency to underestimate tropospheric ozone burden (e.g., 10 – 20% negative bias at 250 hPa in the southern hemisphere (SH) tropical region) (IPCC - Intergovernmental Panel on Climate Change, 2013, Chapter 8). With respect to surface ozone, Schnell et al. (2015) conclude that all ACCMIP models, which reported hourly surface ozone, tend to overestimate surface ozone values in North America and Europe in comparison with available observations. A
- 10 key to fathom these slightly contradicting results may lie in the used dry deposition schemes. Removal of any substance from the atmosphere which is not involving rain, e.g., through gravitational settling or by uptake by plants, soil, and water, is referred to as dry deposition. The process of dry deposition is important for predicting surface ozone concentrations and understanding the observed amount and increase of tropospheric background ozone. It is estimated that $1000 \pm 200 \,\mathrm{Tg} \,\mathrm{a}^{-1}$ of ozone are removed from the atmosphere by dry deposition processes (Monks et al., 2015). A newer
- 15 study by Luhar et al. (2018), however, indicates much lower amounts $(722.8 \pm 87.3 \,\mathrm{Tg}\,\mathrm{a}^{-1})$ due to lower dry deposition to the oceans. Conceptually, dry deposition is a product between near-surface ozone concentration $[O_3](z_0)$ (e.g. the lowermost model level) and a dry deposition velocity $v_{DD}^{O_3}$. Species dependent dry deposition velocities v_{DD}^i , which are synonymously referred to as conductance G^i , for any gaseous species *i*, are typically calculated following a resistance-analogous approach

$$v_{\rm DD}^{i} = \frac{1}{R_a + R_b^i + R_c^i},$$
(1)

- 20 wherein aerodynamic R_a , quasi-laminar layer R_b^i , and canopy resistances R_c^i are key components (Wesely, 1989; Seinfeld and Pandis, 2006). For all gases, R_a is the same, while R_b^i and R_c^i vary from gas to gas and also depend on land surface types (e.g., ice/snow, water, urban, desert, agricultural land, deciduous forest, coniferous forest etc.). Originally, Wesely (1989) used fixed seasonal average dry deposition resistances for each land surface type. For all three types of resistances in this Wesely-type parameterization, more process-oriented formulations have been developed and validated over the years. Luhar et al. (2017)
- have validated ozone dry deposition to the ocean with respect to three different formulations of surface resistances. Based on the global atmospheric composition reanalysis performed in the ECWMF project Monitoring Atmospheric Composition and Climate (MACC) (MACC-II Consortium, 2011) and a more realistic process-based oceanic deposition scheme, Luhar et al. (2018) found that the ozone dry deposition to oceans amounts to $98.4 \pm 30.0 \,\mathrm{Tg} \,\mathrm{a}^{-1}$. In particular, Luhar et al. (2018) found that the average surface resistance of ozone over ocean ($r_c = 2200 \,\mathrm{sm}^{-1}$) is highly overestimated in most models. An update
- 30 on the ozone surface resistance over snow and ice covered surfaces has been provided from combined model and observation studies (Helmig et al., 2007, $v_{ice/snow}^{O_3} = 1/10000 \text{ m s}^{-1}$). Canopy conductance is parameterized at the single-leaf-level (stomatal conductance) for various plant function types (PFT) as well as for single plant species based on empirical studies (Jarvis, 1976; Ball et al., 1987; Simpson et al., 2012; Mills et al., 2017). But progress has also been made on process-oriented modeling of stomatal conductance (Anderson et al., 2000; Buckley, 2017). The variety of differing formulations and choices of parame-

ters leads to a wide spread of results in model intercomparisons (Hardacre et al., 2015; Derwent et al., 2018) and about 20% uncertainty on the resulting total dry deposition (Monks et al., 2015).

In Section 2, we will briefly describe the Oslo CTM3, give a detailed account of the new dry deposition scheme (Section 2.1) as well as present pre-processing of meteorological input data to compute necessary input to the dry deposition scheme such as the begin and duration of the greening season (GDAY, GLEN) and photosynthetic photon flux density (PPFD) (Section 2.2). In Section 3, we present sensitivity tests with respect to a manifold of parameters in the dry deposition scheme (Section 3.1) and validate our results with respect to results from the multi-model intercomparison of Hardacre et al. (2015) (Section 3.2), the MACC-reanalysis (Section 3.3), and to surface ozone observations (Section 3.4). In Section 4, we will summarize and discuss our results and draw conclusions for further development of the model.

our results and draw conclusions for further development of the m

2 Model description

The Oslo CTM3 is an offline, three dimensional, global chemistry transport model (CTM). The key components of the Oslo CTM3 have been described and evaluated by Søvde et al. (2012). A detailed account of the capabilities of the Oslo CTM3 in simulating anthropogenic aerosol forcing in the past and recent past using the Community Emission Data System (CEDS)

15 historical emission inventory (Hoesly et al., 2018) is given by Lund et al. (2018). The Oslo CTM3 can also be coupled to the Model of Emissions of Gases and Aerosols from Nature (MEGAN v2.10) (Guenther et al., 2006). A publication focusing focussing on this is planedplanned.

While the meteorological data driving the Oslo CTM3 is given in at a resolution of T159N80L60, with the highest model level at 0.02 hPa, it is very time and memory consuming to run the Oslo CTM3 with full chemistry at this resolution. Therefore,

20 we reduced the horizontal resolution to $2.25^{\circ} \times 2.25^{\circ}$ in our experiments. In the following, we will give a detailed account of the new dry deposition scheme and the equations that we use.

2.1 Ozone dry deposition scheme

In the original dry deposition scheme, the state of the atmosphere was not taken into account. Dry deposition velocities were rather parameterized following the work of Wesely (1989) with parameter updates from Hough (1991). This means that sea-

- 25 sonal day and night average deposition velocities for different land surface types (water, forest, grass, tundra/desert, and ice and snow) were in use. Day was distinguished from night by solar zenith angles below 90°. Winter was defined by temperatures below 273.15 K for gridboxes containing land masses. For ocean, winter and summer parameters are equal in this parameter-ization, therefore no distinctive treatment was needed for ocean gridboxes. In addition, a reduced uptake due to snow cover above 1 m for forest and 10 cm for grass/tundra, respectively, was taken into account. We will refer to this parameterization as
- 30 Wesely scheme.

Regarding the new dry deposition scheme, we mainly follow Simpson et al. (2012) in their description of dry deposition used in the European Monitoring and Evaluation Programme (EMEP) MSC-W model (see also, Emberson et al., 2000; Simpson et al., 2003; Tuovinen et al., 2004), which is used for air quality modeling implementing the Convention on Long-Range Transboundary Air Pollution (CLRTAP). We will refer to the new scheme as *mOSaic scheme* throughout the rest of the paper.

- 5 The mOSaic scheme is a more physical approach compared to the previously used Wesely scheme, because it takes state (e.g., pressure, temperature) of the atmosphere as well as dynamics (e.g., wind stress) of the boundary layer into account. To a certain degree, the global variety of plants and their variability throughout the seasons is also acknowledged. The mOSaic scheme is implemented for the gaseous species O₃, H₂O₂, NO₂, PAN, SO₂, NH₃, HCHO, and CH₃CHO. Since CO has a very small uptake and is not included in Simpson et al. (2003, 2012), the Wesely parameterization is kept. In addition to the gaseous
- 10 species, Simpson et al. (2012) also modify aerosol deposition velocities, namely black carbon (BC) and organic carbon (OC), sulfuric aerosols (SO₄, MSA) and secondary organic aerosols (SOA), but we have not updated our model with respect to these.

As displayed in Eq. (1), the dry deposition computation is subdivided into contributions from three different resistances. The main idea of a mosaic approach is to calculate these resistances separately for each land surface type k in each grid cell: $R_a^k, R_b^{i,k}$, and $R_c^{i,k}$. The grid cell average dry deposition velocity \overline{v}_{DD}^i is then defined by weighting each individual $v_{DD}^{i,k}$ by the corresponding land fraction factor f_k :

$$\overline{v}_{\rm DD}^i = \sum_k f_k v_{\rm DD}^{i,k} \tag{2}$$

2.1.1 Aerodynamic resistance

15

25

30

In general, the aerodynamic resistance describes the turbulent transport of any substance down to the surface. To derive R_a^k , we follow Simpson et al. (2003, 2012) and compute a local friction velocity at reference height z_{ref} (Eq. (52), Simpson et al., 20 2012)

$$u_*^k = \frac{\overline{u}(z_{\text{ref}}) \cdot \kappa}{\ln\left(\frac{z_{\text{ref}} - d_k}{z_0^k}\right) - \Psi_m\left(\frac{z_{\text{ref}} - d_k}{L}\right) + \Psi_m\left(\frac{z_0^k}{L}\right)},\tag{3}$$

with the average wind speed $\overline{u}(z_{ref})$ at reference height, the Kármán constant $\kappa = 0.40$, the integrated stability equation for momentum Ψ_m (e.g., Garratt, 1992), a grid average Obukhov length L, deplacement height d_k , and roughness length z_0^k $(d_k = 0.78 \cdot h_k(\text{lat}), z_0^k = 0.07 \cdot h_k(\text{lat})$ for forests, $d_k = 0.7 \cdot h_k(\text{lat}), z_0^k = 0.1 \cdot h_k(\text{lat})$ for vegetation other than forests). Taking the height of vegetation in to consideration, we have chosen the model level such that $\overline{z}_{ref} \approx 45 \text{ m}$. Using the derived u_*^k from Eq. (3), a local Obukhov length L_k can be obtained from (Eq. (8), Simpson et al., 2012):

$$L_k = -\frac{\rho c_p T_{2\mathrm{m}} u_*^k}{\kappa g H}.$$
(4)

Herein, H is the sensible heat flux, g is the standard gravitational acceleration, c_p the specific heat capacity, and T_{2m} the 2 m temperature. With these, we can compute the aerodynamical resistance for each land surface type (Eq. (8.8), Simpson et al., 2003)

$$R_a^k = \frac{1}{\kappa u_*^k} \left[\ln\left(\frac{z_{\text{ref}} - d_k}{z_0^k}\right) - \Psi_h\left(\frac{z_{\text{ref}} - d_k}{L_k}\right) + \Psi_h\left(\frac{z_0^k}{L_k}\right) \right],\tag{5}$$

with the integrated stability equation for heat Ψ_h (e.g., Garratt, 1992). Both integrated stability functions (Ψ_m , Ψ_h) and corresponding parameters are listed in supplement S.1.

2.1.2 Quasi-laminar layer resistance

The quasi-laminar layer resistance $R_b^{i,k}$ is species specific and differs over land and ocean surfaces. Over land, we use (Eq. (53), 5 Simpson et al., 2012)

$$R_b^{i,k} = \frac{2}{\kappa u_*} \cdot \left(\frac{\mathbf{S}\mathbf{c}_i}{\mathbf{P}\mathbf{r}}\right)^{\frac{2}{3}},\tag{6}$$

wherein Pr is the Prandtl number (0.72 for air and other gases) and Sc_i is the Schmidt number for a gas *i*. Eq. (6) differs from a similar formulation in Seinfeld and Pandis (2006) by a factor of roughly 1.25. From Sc_i = ν/D_i , with the kinematic viscosity of air ν , we derive a Schmidt number in water equivalent:

10
$$\operatorname{Sc}_{i} = \frac{D_{\operatorname{H_2O}}}{D_{i}} \cdot \operatorname{Sc}_{\operatorname{H_2O}},$$
 (7)

with the molecular diffusivity for any gas D_i , the Schmidt number of water (Sc_{H₂O} = 0.6) and its molecular diffusivity $(D_{H_2O} = 0.21 \cdot 10^{-4} \text{ m}^2 \text{ s}^{-1})$. The used ratios D_{H_2O}/D_i are taken from Simpson et al. (2012, Table S18).

Over ocean, we use (Eq. (54), Simpson et al., 2012)

15
$$R_b^i = \frac{1}{\kappa u_*} \cdot \ln\left(\frac{z_0}{D_i} \cdot \kappa u_*\right)$$
(8)

with an imposed lower threshold of 10 s m^{-1} and an upper limit of 1000 s m^{-1} . The computation of roughness length z_0 over ocean is divided into a *calm* and a *rough* sea case, with a threshold of 3 m s^{-1} . For calm sea, we apply the following upper limit (Hinze, 1975; Garratt, 1992, with a slightly higher coefficient of 0.135)

$$z_0^{\text{calm}} = \min\left\{2 \cdot 10^{-3}, 0.135 \cdot \frac{\nu}{u_*}\right\}.$$
(9)

20 The kinematic viscosity of air ν herein can be computed from

$$\nu = \frac{\mu}{\rho} = \frac{\mu(T)}{\frac{P_0}{T \cdot R_{\rm air}}}.$$
(10)

For the temperature dependent dynamic viscosity of air $\mu(T)$, we chose a linear fit to Sutherland's law through the origin within the temperature range $\{T \in \mathbb{R} | (243.15 < T < 313.15) \text{ K}\}$: $\mu(T) = 6.2 \cdot 10^{-8} \text{ kg m}^{-1} \text{ s}^{-1} \text{ K}^{-1} \cdot T$. But despite its rough accuracy, we found that the choice of $\mu(T)$ has no effect on $\overline{R}_b^{i,k}$ (Supplement S.2: Figs. S1–S2). In Eq. (10), ρ is substituted by

25 the air density using the ideal gas law. P_0 is the surface pressure, as T the 2 m temperature is chosen, and R_{air} is the universal gas constant for air. The rough sea case follows the method of Charnock (1955); Wu (1980):

$$z_0^{\text{rough}} = \min\left\{2 \cdot 10^{-3}, 0.018 \cdot \frac{u_*^2}{g}\right\}$$
(11)

with a gravitational acceleration $g = 9.836 \text{ m s}^{-2}$. The allowed maximum roughness length in both cases is set to 2 mm. Since the z_0 computed with this parameterization are rather small ($0 < z_0^{\text{calm}} < 1 \cdot 10^{-4} \text{ m}$, $0 < z_0^{\text{rough}} < 2 \cdot 10^{-3} \text{ m}$), R_b^i is set to its lower limit of 10 s m^{-1} in about 91% of all cases (see Supplement S.2: Fig. S3).

2.1.3 Surface resistance

5 The surface resistance consists of both , stomatal and non-stomatal resistances.

The stomatal conductance is a measure of the rate of CO_2 exchange and evapotranspiration through the stomata of a leaf. There are several environmental conditions affecting the opening and closing of the stomata and hence the capability of respiration (e.g., light, available water, etc.). Stomata sluggishness, a state in which the stomata can no longer fully close, has been reported as ozone induced damage (Hoshika et al., 2015), but is not taken into account in our formulation. To reflect part of the underland the loof level makes at exactly and level as a property of the stomata of the stomata conditions.

10 the underlying mechanism, the leaf-level molar stomatal conductance in the mOSaic scheme is computed using a common multiplicative ansatz (Ball et al., 1987; Mills et al., 2017):

$$g_{\text{sto, m}}^{k} = g_{\text{max, m}}^{k} \cdot f_{\text{light}}^{k} \cdot \max\left\{f_{\text{min}}^{k}, f_{T}^{k} \cdot f_{D}^{k} \cdot f_{\text{SW}}^{k}\right\}.$$
(12)

The factors herein are normalized and vary within the range 0-1. They account for leaf phenology (f_{phen}) , light (f_{light}) , temperature (f_T) , water vapor pressure deficit (f_D) , and soil water content (f_{SW}) . All factors differ with land use type k.

15 For clarity reasons, we drop this index in the following, as long as it is not necessary for the equation's completeness. The maximum molar stomatal conductance is given by $g_{\max,m}^k$ which is in units of mmol s⁻¹ m⁻². A unit conversion to m s⁻¹ is necessary in our model:

$$g_{\text{sto}}^{k} = g_{\text{sto, m}}^{k} \cdot R \cdot \frac{T_{0}}{P_{0}}.$$
(13)

Herein, R is the universal gas constant. To annotate the differing units, we use the index m in Eq. (12).

20

25

The temperature adjustment f_T is computed from

$$f_T = \frac{T_{2\mathrm{m}} - T_{\mathrm{min}}}{T_{\mathrm{opt}} - T_{\mathrm{min}}} \cdot \left(\frac{T_{\mathrm{max}} - T_{2\mathrm{m}}}{T_{\mathrm{max}} - T_{\mathrm{opt}}}\right)^{\beta},\tag{14}$$

with $\beta = \frac{T_{\text{max}} - T_{\text{opt}}}{T_{\text{opt}} - T_{\text{min}}}$. The parameters T_{min} , T_{max} and T_{opt} are tabulated for various plant functional types. All parameters are taken from Simpson et al. (2012, Tables S16, S19). Since f_T turns negative outside the range defined by T_{min} , T_{max} , we impose a lower limit of 0.01 for numerical reasons.

The water vapor deficit (VPD) is proportional to the saturation partial pressure of water $(P_{H_2O}^s)$ and relative humidity (RH)

$$VPD = P_{H_2O}^s \cdot (1 - RH/100).$$
(15)

Using tabulated values of f_{\min} , D_{\min} , D_{\max} , the water vapor pressure deficit penalty factor f_D can be computed:

$$f_D = f_{\min} + (1 - f_{\min}) \cdot \frac{D_{\min} - \text{VPD}}{D_{\min} - D_{\max}}.$$
 (16)

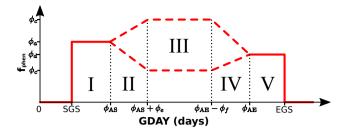


Figure 1. Sketch of the five different phases in plant phenology f_{phen} in accordance to Eq. (18).

The penalty factor with respect to available soil water (SW) f_{SW} is defined as

$$f_{\rm SW} = \begin{cases} 1 & \text{if } SW \ge 0.5, \\ 2 \cdot SW & \text{if } SW < 0.5. \end{cases}$$
(17)

SW is evaluated at a soil depths of $0.28 - 1 \,\mathrm{m}$, which corresponds to SWVL3 in OpenIFS.

The phenology of a plant typically describes its life-cycle throughout a year, e.g., at mid latitudes and for deciduous species, it starts with the emergence of leafs in spring and ends in fall. In the mOSaic scheme, phenology is parameterized with respect to the start of the greening season (SGS) and its end (EGS). Details about our treatment of these are given in Section 2.2.1. In summary, our adaption of the f_{phen} parameterization reads as follows:

$$f_{\text{phen}} = \begin{cases} if \quad \text{GLEN} \ge 365 & 1 \quad (explicitly \ excluding \ tropics) \\ if \quad \text{GDAY} = 0 & 0 \\ \\ else & \begin{cases} if \quad \text{GDAY} \le \phi_{\text{AS}} & \phi_a \\ if \quad \text{GDAY} \le \phi_{\text{AS}} + \phi_e & \phi_b + (\phi_c - \phi_b) \cdot (\text{GDAY} - \phi_{\text{AS}})/\phi_e & (18) \\ \\ if \quad \text{GDAY} \le \text{GLEN} - \phi_{\text{AE}} - \phi_f & \phi_c \\ \\ if \quad \text{GDAY} \le \text{GLEN} - \phi_{\text{AE}} & \phi_d + (\phi_c - \phi_d) \cdot (\text{GLEN} - \phi_{\text{AE}} - \text{GDAY})/\phi_f \\ \\ else & \phi_d \end{cases}$$

Herein, we use the SGS and EGS derived parameters day of greening season (GDAY), the time elapsed starting at the SGS, 10 and the total length of the greening season (GLEN), the time span between EGS and SGS. The parameters ϕ_a , ϕ_b , ϕ_c , and ϕ_d define start or end points in the five phases of phenology in the mOSaic scheme, while ϕ_e , ϕ_f , ϕ_{AS} , and ϕ_{AE} control the temporal timing (Fig. 1). If GLEN is zero we are, e.g., in Arctic regions and there is no vegetation anyway, therefore $f_{phen} = 0$. Before the start of the greening season (GDAY = 0) $f_{phen} = 0$. Since this phenology is tuned to northern hemisphere (NH) mid latitudes, it does not apply to the tropics. We therefore decided to set $f_{phen} = 1$ if GLEN is greater or equal to 365 which is the 15 case in the tropics.

Light in the wavelength band 400-700 nm to which the plant chlorophyll is sensitive is called photosynthetic active radiation (PAR). The integral of PAR over these wavelengths is the photosynthetic photon flux density (PPFD). The correction factor

 f_{light} in response to varying PPFD is:

 $f_{\text{light}} = 1 - \exp(-\alpha_{\text{light}} \cdot \text{PPFD}).$

Since $g_{\text{max, m}}$ in Eq. (12) is in units of , a unit conversion to is necessary in our model:

$$g_{\rm sto}^k = g_{\rm sto, \ m}^k \cdot R \cdot \frac{T_0}{P_0}.$$

5 Herein, R is the universal gas constant. In the mOSaic scheme, non-stomatal conductances are explicitly calculated for O_3 , SO_2 , HNO_3 , and NH_3 . For all other species, an interpolation between O_3 and SO_2 values is carried out. The non-stomatal conductance for O_3 consists of two terms, one depending on vegetation type and one depending on the soil/surface. For each land surface types type k, we can write

$$G_{\rm ns}^{\rm O_3,k} = \frac{{\rm SAI}_k}{r_{\rm ext}} + \frac{1}{R_{\rm inc}^k + R_{\rm gs}^{\rm O_3,k}}.$$
(20)

10 SAI_k is the surface area index for vegetation type k, which is LAI plus a value that represents cuticles and others other surfaces. The external leaf resistance is defined by

$$r_{\rm ext} = 2000 \, {\rm s} \, {\rm m}^{-1} \cdot F_T. \tag{21}$$

Herein F_T is a temperature correction factor for temperatures below $-1^{\circ}C$ and $\{F_T \in \mathbb{R} | (1 \le F_T \le 2)\}$

$$F_T = \exp\left(-0.2 \cdot (1 + \theta_{2m})\right). \tag{22}$$

15 θ_{2m} is the 2m temperature in °C. For most land surface types, SAI \equiv LAI. Some exceptions are:

$$SAI = \begin{cases} LAI + 1 & if forest / wetland, \\ LAI \cdot 5/3.5 & if cropland, 1st part of growing season, \\ LAI + 1.5 & if cropland, 2nd part of growing season, \\ 0 & if cropland, winter. \end{cases}$$
(23)

Extending the mOSaic scheme to the southern hemisphere, we use the growing season for crops defined in Table 1.

In this way, vegetation affects the conductance also by being there, not only by uptake through the stomata. The in-canopy resistance R_{inc} (Erisman et al., 1994) is then modified with respect to each (vegetated) land surface type in k

20
$$R_{\rm inc} = b \cdot {\rm SAI}_k \cdot \frac{h_k({\rm lat})}{u_*},$$
 (24)

where $h_k(\text{lat})$ is the latitude dependent vegetation height (see explanation at the end of this section) and $b = 14 \text{ m}^{-1}$ is an empirical constant. The canopy resistance described in Simpson et al. (2012) does not take temperature and snow into account and is zero for non-vegetated surfaces, but we will adopt the correction previously used in the Oslo CTM3 Wesely scheme.

Table 1. Definition of growing season for crops used in the Oslo CTM3 in northern hemisphere (NH) and southern hemisphere (SH).

	1st part (days)	2nd part (days)
NH	90–140	141–270
SH	272–322	323–452

As initially mentioned, the necessary depth of snow to cover a certain type of vegetation differs. Therefore, we calculate a snow cover fraction f_{snow} using the snow depth S_D , which is available in units of meter of water equivalent from the meteorological input data, scaled to 10% of the vegetation height. $R_{gs}^{O_3,k}$ is tabulated. We correct for temperature by F_T and for snow cover fraction:

5
$$\frac{1}{R_{gs}^{O_3,k}} = \frac{1 - f_{snow}^k}{\hat{R}_{gs}^{O_3,k}} + \frac{f_{snow}^k}{R_{snow}^{O_3,k}}.$$
 (25)

The bulk canopy conductance is then defined as:

$$G_c^k = \text{LAI} \cdot g_{\text{sto, m}}^k + G_{\text{ns}}^k, \tag{26}$$

wherein LAI is the one-sided leaf area index taken from ISLSCP2 FASIR, g_{sto} the leaf-level stomatal conductance, and G_{ns} the bulk non-stomatal conductance.

10

15

2.1.4 Latitude dependent vegetation height

The vegetation height $h_k(\text{lat})$ as described by Simpson et al. (2012) is linearly decreasing with latitude between 60° and 74°N . To adapt this to a global model, we made a few additional assumptions. The tabulated height for each vegetation type h_k in the mOSaic scheme is regarded as constant at mid latitudes ($40^\circ - 60^\circ$). Towards the poles, we decrease the height of each vegetation type using the same rate as described in Simpson et al. (2012). At a latitude of 74° a minimum height of $3/10 \cdot h_k$ is reached and kept constant. Towards the equator, we increase the height linearly so that at a latitude of 10° a maximum height of $2 \cdot h_k$ is reached which is then held constant. We also assume symmetry in both hemispheres. Presuming a typical tree height of 20 m at mid latitudes, this step-wise function yields a height of 8 m at high latitudes and 40 m in the tropics which is not unrealistic. For four example PFTs, results are shown in the Supplement (S.3, Fig. S4).

20 2.1.5 Mapping of land surface types

The Oslo CTM3 is configured to read land surface types from, either ISLSCP2 product from MODIS or Community Land Model (CLM) 2 categories, which have to be mapped to the land surface types used in the mOSaic scheme (Fig. 2). For both, MODIS and CLM 2 land surface categories, snow and ice cover is estimated from input meteorology, while f_L^{water} is defined as

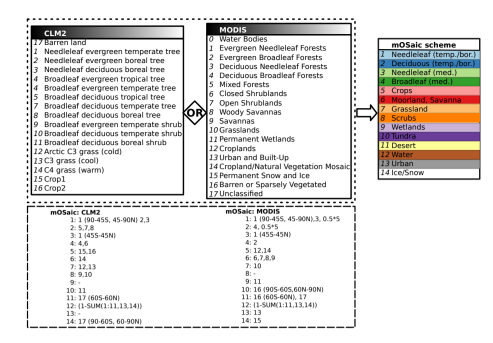


Figure 2. Mapping of land surface categories. Either land surface categories from ISLSCP2 product of MODIS or the Community Land Model (CLM) 2 can be chosen for mapping to the land surface types we use in the mOSaic scheme. Water bodies of MODIS are actually not mapped. For both, MODIS and CLM 2 land surface categories, snow and ice cover is estimated from input meteorology, while water is defined as $1 - \sum_k f_L^k$. From MODIS category *Barren or sparsely vegetated*, everything polward from 60° is defined as tundra, while everything equatorward is categorized as desert.

 $1 - \sum_{k} f_{L}^{k}$. From the MODIS category *Barren or sparsely vegetated*, everything polward from 60° is defined as tundra, while everything equatorward is categorized as desert. This mapping differs from the one used in the Wesely scheme.

2.2 Pre-processing

As mentioned in the previous section, there are two variables needed for computing the stomatal conductance which are not 5 directly available from the meteorological input data. The greening season, as the time of the year in the mid and high latitudes when it is most likely for plants to grow, and the photosynthetic photon flux density, as the amount of light that plants need to photosynthesize. In the following, we present the necessary pre-processing of the variables. It is <u>planed planned</u> to implement an online computation of these variables into the Oslo CTM3 later on.

2.2.1 Greening season

10 In Eqs. (20–23), Simpson et al. (2012) use prescribed start of growing season (SGS) and end of growing season (EGS) at 50 °N (d_{SGS} , d_{EGS}) together with lapse rates (∇d_{SGS} , and ∇d_{EGS}) to define phenology and dry deposition over agricultural areas. For the growing season of crops in the computation of non-stomatal conductances, we use also prescribed values (Table 1), while for

the stomatal conductances, as shown in Eq. (18), we use the SGS and EGS derived parameters: day of greening season (GDAY), the time elapsed starting at the SGS, and the total length of the greening season (GLEN), the time span between EGS and SGS. Since the parameterization of SGS and EGS in Simpson et al. (2012) is not applicable in a global model, another latitude dependent parameterization is needed. First, we used a parameterization which was already implemented in the Oslo CTM3

- 5 and which had been adopted from the Sparse Matrix Operational Kernel Emissions Biogenic Emission Inventory System (SMOKE-BEIS; https://www.epa.gov/air-emissions-modeling/biogenic-emission-inventory-system-beis). SMOKE-BEIS has fixed values for SGS and EGS for all regions but NH mid latitudes $(23^{\circ} < lat < 65^{\circ})$, where it uses lapse rates of $\nabla d_{SGS} = 4.5$ and $\nabla d_{EGS} = 3.3$. As this parameterization is optimized for North America, it does not work well in Europe, e.g., most of northern Scandinavia has no allocated vegetation period. This basically results in a suppression of canopy resistance in north-
- 10 ern Scandinavia.

20

In agriculture, there are different empirical rules to estimate the SGS and EGS. The simplest assumption is that greening starts after 5 consecutive days with a daily average temperature above 5° C and vice versa for EGS. Other estimates use growing degree days (Levis and Bonan, 2004; Fu et al., 2014a), include soil moisture (Fu et al., 2014b), or rely on satellite observations.

15 A comprehensive evaluation of different techniques is given by Anav et al. (2017). Another solution would be the usage of a proper land surface model, e.g. LPJ-GUESS, CLM, but the integration of such into the Oslo CTM3 is not planed_planned at the moment.

Based on the empirical rule (5 °C-days), we have pre-processed our meteorological input data offline. We added some additional criteria to prevent for *false spring*: If, within these 5 days, the average temperature drops below or rises above 5 °C, the counter is reset, respectively. First, we used the 5 °C-days criteria for $45^{\circ} < \text{lat} < 85^{\circ}$ in the NH, but extended it

also to $35^{\circ} < \text{lat} < 65^{\circ}$ in the SH. In all other cases and where the 5°C -days criteria fails, we still use the SMOKE-BEIS parameterization. The described algorithm written in python 2.7 has been included as Supplement S.4. An example map of the computed GLEN using the 5°C -days criteria in both hemispheres is shown in Fig. 3.

2.2.2 Photosynthetic photon flux density

- 25 From OpenIFS an accumulated surface PAR is available. It is integrated both, spactrally spectrally (presumably 400 700 nm) and temporally. For practical use in Eq. (19), we de-accumulate this field with respect to time and refer to the result as PPFD. The main obstacle is that PAR has been accumulated since model start, so that the first field kept from the original OpenIFS simulation (00 UTC) is 12 hours after model start (12 UTC on the previous day). In other words, the first time step of each day in the Oslo CTM3 has already accumulated PAR from 12 UTC on the previous day. De-accumulation of times 03 UTC to
- 30 21 UTC, simply means computing the difference

$$PPFD(t_i) = PAR(t_{i+1}) - PAR(t_i).$$
(27)

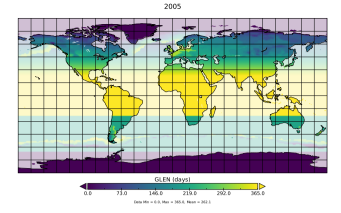


Figure 3. Pre-processing of greening season from meteorological surface temperature fields. Shown is the total length of the greening season (GLEN) for the year 2005. The 5 °C-days criteria has been used in both hemispheres mid–high latitudes. Ocean has been shaded to indicate that greening season will only affect land.

For de-accumulation of the remaining time step, the best choice is subtracting the difference between 21 UTC and 12 UTC of the previous day

$$PPFD(t = 00 \text{ UTC}) = PAR(t = 00 \text{ UTC}) - [PAR(t = 21 \text{ UTC} - 1 \text{ day}) - PAR(t = 12 \text{ UTC} - 1 \text{ day})]$$
(28)

and limit the result to positive values only. An example PAR de-accumulation for January 2nd 2005 is shown in Supplement S.5

5 (Figs. S5–S7). The resulting PPFD fields are still accumulated over a time period of 3 hours and should be divided by 3. A known issue in the OpenIFS (cycles \leq c41r2) causes surface PAR values to be about 30% below observations. To counter this, we decided to refrain from the division at this stage, but need to bear this in mind for later OpenIFS cycles.

3 Evaluation

In this section, we present results from a manifold of Oslo CTM3 model integrations testing different parameters of the mO-

- 10 Saic scheme. We focus on changes in ozone total dry deposition $\sum O_3^{DD}$, dry deposition velocities $v_{DD}^{O_3}$, concentrations in the lowermost model level $[O_3](p_0)$, and tropospheric burden $\sum_{trop} O_3$. We evaluate our results with respect to the multi-model comparison of ozone dry deposition by Hardacre et al. (2015) (Section 3.2), the MACC-reanalysis (Section 3.3), and observations (Section 3.4). The Oslo CTM3 is driven by meteorological input fields from ECMWF OpenIFS cy38r1. CEDS historical emission inventory (Hoesly et al., 2018) is used for anthropogenic emissions, while biomass burning is covered in daily reso-
- 15 lution by NASA's Global Fire Emissions Database, Version4 (GFEDv4). Biogenic emissions are taken from MEGAN-MACC output (Sindelarova et al., 2014), while emissions from soil and wetlands are computed by MEGAN. Resultant NO_x emissions are up-scaled to match Global Emissions InitiAtive (GEIA) inventory and are estimated to amount to $6.55 \text{ Tg}(\text{N}) \text{ a}^{-1}$. For oceanic emissions of CO, we use predefined global fields from POET (GEIA-ACCENT emission data portal, 2003). Emissions of CH₄ are taken from the EU project (EU GOCE 037048) Hydrogen, Methane and Nitrous oxide: Trend variability,

budgets and interactions with the biosphere (HYMN) for the year 2003 and scaled to oceanic amounts of CH_4 from NASA. In the following (Section 3.1), we will present the various model sensitivity studies.

3.1 Sensitivity studies

Due to significant differences between the mOSaic scheme and the previous Wesely scheme with respect to implementation,

- 5 it is not possible to fully disentangle and trace back every single difference in results to a respective change. Therefore, we conducted one reference simulation denoted as *mOSaic* and in total seven sensitivity studies to probe the parameter space for stomatal conductance (*mOSaic_offLight*, *mOSaic_offPhen*, and *mOSaic_SWVL1*), ozone surface resistance R^{O3} (*mOSaic_ice*, *mOSaic_desert*, and *mOSaic_hough*), and emissions (*mOSaic_emis2014*). A reference simulation featuring the Oslo CTM3 Wesely scheme has been conducted and will be referred to as *Wesely_type*, indicating that other implementations of the original
- 10 work by Wesely (1989) may exist in other models. All model experiments discussed in the following are summarized in Table 2. An *x* therein denotes that the model was run exactly in the configuration and with parameters as has been described in Section 2. For all model integrations, the meteorological reference year is 2005. This choice affects the direct comparison with data and studies that either show results based on decadal averages or differing years, because non-linearities in ozone formation and destruction make ozone concentrations sensitive to both, differences in local concentration of precursors and meteorological

15 conditions (Jin et al., 2013).

First, we have a closer look at the influence of certain parameters on the stomatal conductance. As indicated by the names, $mOSaic_offLight$ and $mOSaic_offPhen$ are rather extreme scenarios completely switching off the sensitivity to light and phenology in Eq. (12) by setting f_{light} and f_{phen} to a fixed value of 1, respectively. Because of the underlying research project's focus on aretie_Arctic_and alpine ecosystems, where water might only be available from upper soil layers, an experiment was

- 20 conducted using the uppermost soil water level (SWVL1) in the implementation of f_{SW} . After this, we want to confirm the importance of choice of R^{O_3} for different land surface types. We conducted three experiments looking at a $R_{ice/snow}^{O_3}$ update (Helmig et al., 2007) (*mOSaic_ice*), observed $R_{desert}^{O_3}$ (Güsten et al., 1996) (*mOSaic_desert*), and an approximation of R^{O_3} originally used in *Wesely_type* (Wesely, 1989; Hough, 1991). Finally, we run a simulation with emissions for the year 2014 instead of 2005 (*EMEP_emis2014*) to characterize the general influence of differing emissions on ozone.
- In Fig. 4, we show the average global distributions of the relative difference between *mOSaic* and *Wesely_type* on global maps by means of ozoneburden in the lowermost model level for surface ozone, dry deposition velocity, and total ozone dry deposition. The ozone burden surface ozone increases globally except for some regions covered by tropical forest. Especially in desert regions in Africa, America, and Asia, ozone burden the surface ozone increases by more then than 100%. Consistently, dry deposition velocities decrease globally by the same order of magnitude in these regions, while they increase over tropical
- 30 forest. With respect to total dry deposition, the picture is a bit less clear. We find a decrease of total dry deposition of ozone in desert regions and ocean covered areas and an increase in regions covered by tropical forest, while at mid and high latitudes in both hemispheres only small changes are visible. A possible explanation to this divergence especially in desert regions is the difference between the prescribed surface resistances R^{O_3} in the Wesely scheme in comparison to those used in *mOSaic*. We come back to this in the following sections.

Simulation	mOSaic scheme			Greening	season	R^{O_3}	Emissions	
	f_{phen}	$f_{ m light}$	fsw	SMOKE-BEIS	$5^{\circ}\mathrm{C}$ -days		(year)	
Wesely_type		n/a		х	n/a	Wesely (1989); Hough (1991)	2005	
mOSaic	х	х	х		х	Simpson et al. (2012)	2005	
mOSaic_offLight	х	1	х		х	Simpson et al. (2012)	2005	
mOSaic_offPhen	1	х	х		х	Simpson et al. (2012)	2005	
mOSaic_SWVL1	х	х	VL1		х	Simpson et al. (2012)	2005	
mOSaic_ice	х	х	х		х	Simpson et al. (2012); Helmig et al. $(2007)^{\dagger}$	2005	
mOSaic_desert	х	х	х		х	Simpson et al. (2012); Güsten et al. (1996)*	2005	
mOSaic_emis2014	х	х	х		х	Simpson et al. (2012)	2014	
mOSaic_hough	х	х	х		x	Wesely (1989); Hough (1991)*	2005	

Table 2. Summary of specifications of all simulations discussed in this section. For simplicity, only the tested parameters are listed. An x denotes that the model was run exactly in the configuration as has been described in Section 2.

[†] $R_{\text{ice/snow}}^{\text{O}_3} = 10000 \,\text{s} \,\text{m}^{-1}$; * $R_{\text{desert}}^{\text{O}_3} = 800 \,\text{s} \,\text{m}^{-1}$; * For adapted values see Supplement S.6

3.2 Comparison with modeling results

In the evaluation of our model, we closely follow suggestions by Hardacre et al. (2015). For the purpose of comparison with the multi-model mean of the therein participating Task Force on Hemispheric Transport of Air Pollution (TF HTAP) models, we also have re-gridded our data to a horizontal resolution of $3^{\circ} \times 3^{\circ}$. In Section 3.2.1, we look at zonal distributions of $[O_3](p_0)$, $u_{23}^{O_3}$ and $\sum ODD$ for all our constitution simplefiere and study accessed where a formula for a formula formu

5 $v_{\text{DD}}^{\text{O3}}$, and $\sum O_3^{\text{DD}}$ for all our sensitivity simulations and study seasonal cycles of hemispheric ozone as well as for nine land surface types (Section 3.2.2). From this, we estimate the total annual ozone dry deposition onto ocean, ice, and land surfaces and compare also with results from Luhar et al. (2017).

Dry deposition velocities are directly available only for the new model version. For *Wesely_type*, monthly averaged dry deposition velocities $v_{DD}^{O_3}$ had to be retrospectively estimated from the ratio between the total ozone dry deposition $\sum O_3^{DD}(p_0)$ and monthly averaged ozone amount in the lowermost model level $O_3(p_0)$

$$v_{\rm DD}^{\rm O_3} = \frac{\sum {\rm O}_3^{\rm DD}(p_0)}{{\rm O}_3(p_0)} \cdot c_{\rm month}.$$
(29)

Herein, $c_{\text{month}} = \frac{\Delta h_{\text{month}}}{s_{\text{month}}}$, with the monthly average height of the lowermost model level in each gridbox Δh_{month} and the respective number of seconds in a month s_{month} . In the case of *mOSaic*, resulting values for $v_{\text{DD}}^{\text{O}_3}$ from Eq. (29) are compatible with the values which are directly ailable available from model output.

15 3.2.1 Zonal distribution

10

The annual zonal average with respect to surface ozone concentration (Fig. 5a) displays on average, in consistency consistent with Fig. 4a, a global increase of surface ozone concentrations by 6 ppb comparing *mOSaic* to *Wesely_type*. This increase

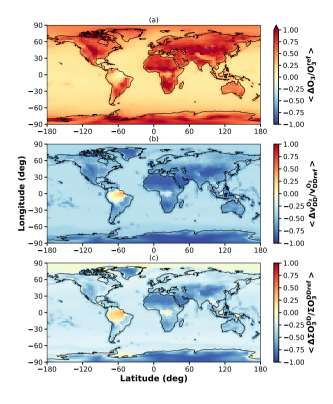


Figure 4. Relative difference between reference simulations *mOSaic* and the *Wesely_type* with respect to (a) average surface ozoneburden in the lowermost model level; (b) average ozone dry deposition velocity; (c) total amount of ozone removed from the atmosphere by dry deposition.

is largest in the zonal band (25-50) °N which contains the major deserts. In the deep tropics $(5 \circ S - 5 \circ N)$, the increase is smallest ($\mathcal{O}(5 \text{ppb})$). We find that the mOSaic scheme further intensifies the strong asymmetry between northern and southern hemisphere hemispheres as a consequence of the distribution of the continental land masses and vegetation thereon. Among the sensitivity studies focusing focusing on the stomatal conductance, there is only a low absolute variance. Neglecting the

- 5 dependence on light in the stomatal conductance formulation ($mOSaic_offLight$) or in other words allowing photosynthesis 24/7 – decreases the ozone concentration by 1 – 2 ppb in the tropics and NH mid latitudes, while choosing soil water at shallower depths ($mOSaic_SWVL1$) increases [O₃] insignificantly. Rather surprisingly, switching off the phenology completely ($mOSaic_offPhen$) amounts on average only to small difference (O(<1ppb)). Most remarkable, but expected due to the much smaller prescribed dry deposition velocity over ice and snow, $mOSaic_ice$ displays a doubling of surface ozone in the
- 10 high Arctics compared to Wesely_type ($\mathcal{O}(20 \text{ ppb})$) but affects ozone concentrations down to latitudes at about 50° in both hemispheres. Reducing $R_{\text{desert}}^{O_3}$ by 60% (mOSaic_desert) a reduction in the order of 1 ppb is found mainly limited to the NH. The largest impact on ozone concentrations ($\mathcal{O}(2-5 \text{ ppb})$) is found for the experiment mOSaic_hough which is closest to

Wesely_type, since we used on average the same R^{O_3} (see Supplement S.6). The scenario of differing emissions (2005 in comparison to 2014 or more specifically *mOSaic* compared to *mOSaic_emis2014*), yields higher ozone concentrations in the northern hemisphere in 2005 in accordance to a reduction in sulfur and NO_x emissions in south east Asia in later years. An opposite tendency is seen for latitudes south of 30 °N, where an increase in ozone precursors is seen in CEDS.

- The $v_{DD}^{O_3}$ are shown in Fig. 5b. The dry deposition velocities in the mOSaic scheme are well below the Wesely scheme and in remarkable agreement with the results shown by Hardacre et al. (2015). In the Arctics, except for *mOSaic_ice*, all model experiments are slightly above the multi-model-mean. This indicates, that with respect to the other models, the Helmig et al. (2007) surface resistance above ice and snow should be considered as the new standard for the Oslo CTM3. This may, however, lead to an overcompensation of the current arctic Arctic low-bias in surface ozone in the Oslo CTM3 and needs further
- 10 evaluation. The dry deposition velocities are of course independent of the emission scenario, but display a strong sensitivity on to f_{light} , f_{phen} , and especially on the choice of R^{O_3} . The shape of the normalized zonal average dry deposition velocities of the mOSaic scheme are more similar to the multi-model-mean than to *Wesely_type* (Supplement S.7: Fig. S8). The biggest exceptions are the zonal bands (50 - 70) °S (almost entirely covered by ocean), (12 - 30) °S (coinciding with the location of Australia and its desert regions), as well as its counterpart in the northern hemisphere (12 - 30) °N.
- The annual total ozone dry deposition is shown in Fig. 5c. In accordance to the previously described features, we observe a reduction of the global total ozone dry deposition in all sensitivity studies. In the most extreme case (NH subtropics and mid latitudes), the total ozone dry deposition drops to one-half of the amount given by *Wesely_type*. The occurrence of this reduction in the zonal bands, where the major deserts are located, points to a substantial difference in v_{desert}^{O3}. Consulting the parameter file used in the Wesely scheme, we indeed find v_{desert}^{O3} = v_{undra}^{O3} = 0.26 cm s⁻¹ (Hough, 1991), while in the mOSaic scheme ver higher in the Wesely scheme (v_{ice/snow}^{O3} = v_{water}^{O3} = 0.07 cm s⁻¹) than in the original parameter set (v_{ice/snow}^{O3} = v_{water}^{O3} = 0.05 cm s⁻¹, Simpson et al., 2012). These differences in surface resistances over huge parts of the unvegetated surface of the Earth accounts account for most of the qualitative difference between the Wesely and the mOSaic scheme, but does do not explain the quantitative difference (compare mOSaic_hough). We further elaborate on this in the following (Section 3.2.2).
- There seems to be a discrepancy between the Oslo CTM3 response and the multi-model-mean, since the Wesely scheme is similar to the multi-model-mean with respect to total annual ozone dry deposition, while the $v_{DD}^{O_3}$ of the mOSaic scheme matches matches better. This could be a sign of differences in photo-chemistry and transport (e.g. convective, advective, STE) between the Oslo CTM3 and the average TF HTAP model, but without comparing to the actual [O₃] of the TF HTAP models that participated in the model intercomparison, we cannot elaborate on this any further. This may also hint to issues in the Oslo CTM3
- 30 photo-chemistry, which may have a too high ozone production, or the actual removal of ozone from the atmosphere, which might have been adjusted to the less physical dry deposition velocities in the past, but this is subject to further investigations.

In Appendix Fig. A1a, the average zonal ozone dry deposition is shown separated by month. Where available, we have added the multi-model-mean given by Hardacre et al. (2015) as a reference. As for the global annual comparisons above, the mOSaic scheme matches the multi-model-mean values remarkably well with respect to dry deposition velocities, while

35 it strongly underestimates the total dry deposition. Qualitatively, there are two major phases apparent: NH and SH greening

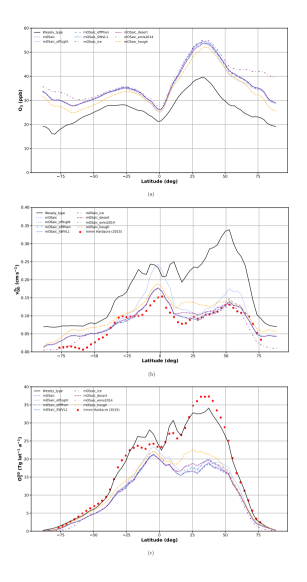


Figure 5. Comparison of the manifold of Oslo CTM3 integrations with respect to (a) Ozone concentrations in the lowermost model level, (b) Annual average ozone dry deposition velocity, (c) Total annual ozone dry deposition. The different colors indicate sets of simulation with similar baselines. The multi-model mean from the evaluation of TF HTAP models by Hardacre et al. (2015) is shown as a reference (where available).

season. Spring and summer in the NH is reflected in a pronounced peak of $v_{DD}^{O_3}$ in the northern mid latitudes, while it is absent in winter (SH summer). Spring and summer in the SH are marked by a southward shift of the tropical peak dry deposition velocity and a slight increase of $v_{DD}^{O_3}$ in the region (20 - 40) °S. In the Wesely scheme, NH mid latitude peak velocities appear in June compared to July in the mOSaic scheme, indicating that the seasonal cycles differ. The corresponding total monthly ozone dry deposition is shown in Appendix Fig. Ally In general, the seasonal patterns are guite similar in the Wesely scheme and

5 dry deposition is shown in Appendix Fig. A1b. In general, the seasonal patterns are quite similar in the Wesely scheme and

the mOSaic scheme, displaying a strong symmetry around 10°N in January/February and November/December, respectively. What differs most is the molding and intensity of the NH peak dry deposition. Both schemes reach the maximum in June/July but the peak is much more differentiated in March already in the Wesely scheme. Similarly, the SH tropical peak dry deposition is reached in August/September but sustained longer, into October, in the Wesely scheme. Since we have not conducted any

5 simulation with a meteorological year other than 2005, we cannot elaborate on whether this is a special feature of our chosen year or not.

3.2.2 Average seasonal cycles

To further disentangle the contributions of different regions to the global ozone budget, we will look at different projections of seasonal cycles.

- In Fig. 6, the total annual ozone dry deposition separated into mid and high latitudes in the northern hemisphere $(30^{\circ} 90^{\circ}N)$, the tropics and subtropics $(30^{\circ}S 30^{\circ}N)$, and the mid and high latitudes in the southern hemisphere $(30^{\circ} 90^{\circ}S)$ is shown. We have added the multi-model-mean by Hardacre et al. (2015) as a reference. While the total ozone dry deposition of *Wesely_type* agrees well with the multi-model-mean in any zonal band, the mOSaic scheme displays a much smaller total ozone dry deposition. This deviation appears to be almost the same for each zonal band (6 7%).
- As expected, the NH mid and high latitudes display a strongly pronounced seasonal cycle, while it is less pronounced in the tropics (due to the lack of seasons) and in the SH (due to the small percentage of vegetated surface). The highest ozone dry deposition is found in the tropics and amounts on average to the peak level of dry deposition in the NH for the multimodel-mean (Hardacre et al., 2015) and mOSaic scheme. In the Wesely scheme, the average tropical ozone dry deposition diverges by 5 Tg in comparison to its corresponding NH maximum. Compared to the multi-model-mean, the seasonal cycle in
- 20 the Oslo CTM3 NH appears to be shifted towards later in the year. The seasonal cycle in the tropics and subtropics only differs by magnitude otherwise the shapes are identical for both the mOSaic scheme, the Wesely scheme, and the multi-model-mean. The total amount of dry deposition of ozone differs strongly between the different model experiments, with *mOSaic_SWVL1* and *mOSaic_hough* displaying the lowest and highest amount, respectively. This indicates that surface ozone is much more sensitive to the choice of parameters ($\mathcal{O}(5 \text{ ppb})$ for *mOSaic_hough* in the tropics) than to slight changes in precursor emissions
- 25 ($\mathcal{O}(1 \text{ ppb})$ for *mOSaic_emis2014* in the tropics).

As suggested by Hardacre et al. (2015), we also look at ozone dry deposition velocities with respect to surface types separately. Since dry deposition velocities are not directly available for *Wesely_type*, we use Eq. (29) to estimate these. Based on a CLM 2 average dynamic land surface map, we generate masks for 9 different surface types (Fig. B1a) and use these to select gridboxes with a high percentage of these surface types, ranging from a meager 70% for cropland in the NH mid latitudes to

30 100% regarding for desert, ocean, snow and ice, and tropical forest. Thus, it is not possible to exclusively select gridboxes with 100% cover for each surface type. Since we have not performed a full unfolding on the data, the results should be treated with slight caution (e.g. over cropland). In the case of the mOSaic scheme, we have pre-selected the dry deposition velocities in accordance to the land surface type.

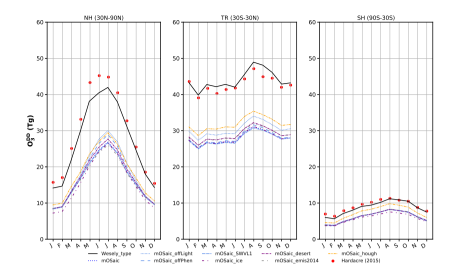


Figure 6. Seasonal cycle of total annual amount of ozone removed from the atmosphere through dry deposition separated into northern hemisphere (NH), tropics (TR), and southern hemisphere (SH). The multi-model mean from the evaluation of HTAP models by Hardacre et al. (2015) is shown as a reference.

In Fig. 7, the seasonal cycles of dry deposition velocities are shown for the nine surface categories. The patterns and absolute numbers differ substantially between the Wesely scheme and the mOSaic scheme and the multi-model-mean. The divergence of the average dry deposition velocities between Wesely_type and mOSaic in desert regions ($\Delta \overline{v}_{desert}^{O_3} = 0.20 \,\mathrm{cm \, s^{-1}}$) as well as grassland ($\Delta \overline{v}_{\text{grassland}}^{O_3} = 0.65 \text{ cm s}^{-1}$) is quite remarkable. The difference of *mOSaic* and *Wesely_type* to the multi-modelmean in tropical forest regions is $\Delta_{\text{mOSaic}} \overline{v}_{\text{tropical forest}}^{O_3} = 0.61 \,\mathrm{cm \, s^{-1}}$ and $\Delta_{\text{Wesely_type}} \overline{v}_{\text{tropical forest}}^{O_3} = 0.49 \,\mathrm{cm \, s^{-1}}$, respectively. 5 The multi-model-mean displays a rather pronounced seasonal cycle in desert regions $(0.10 \text{ cm s}^{-1} \le v_{\text{desert}}^{O_3} \le 0.15 \text{ cm s}^{-1})$, which cannot be reproduced with the mOSaic scheme. The dry deposition velocities over desert regions are consistent with the average values from the prescribed ozone surface resistances, which means that in the mOSaic scheme they are 1 order of magnitude lower than in the Wesely scheme. In the mOSaic scheme, dry deposition to deserts is dominated by contribution from R_b . From a limited number of ozone flux measurements in the Sahara desert, Güsten et al. (1996) deduced $v_{\text{desert. dav}}^{O_3} =$ 10 $0.1\,\mathrm{cm\,s^{-1}}, v_{\mathrm{desert, \, night}}^{\mathrm{O}_3} = 0.04\,\mathrm{cm\,s^{-1}}, \text{ and } \overline{v}_{\mathrm{desert}}^{\mathrm{O}_3} = 0.065\,\mathrm{cm\,s^{-1}}.$ This implies, that ozone dry deposition over desert regions is highly overestimated in the Wesely scheme as well as in TF HTAP models, while it may be underestimated in the mOSaic scheme. Similarly, the dry deposition velocities over water differ. From measurements during ship-campaigns a mean value of $\overline{v}_{water}^{O_3} = 0.019 \,\mathrm{cm \, s^{-1}}$ over the ocean has been deduced (Helmig et al., 2012). In a model study of different mechanisms of dry deposition to ocean waters by means of prescribed $\overline{v}_{water}^{O_3}$ and one- and two-layer gas exchange modeling, Luhar et al. 15 (2017) found $\overline{v}_{water}^{O_3}$ ranging between 0.018 cm s^{-1} (two-layer scheme) and 0.039 cm s^{-1} (prescribed). With $\overline{v}_{water}^{O_3} = (0.046 \pm 10^{-1} \text{ cm})^{-1}$ $(0.002) \,\mathrm{cm \, s^{-1}}$, the mOSaic scheme (Section 2.1.2) yields probably a too strong dry deposition to ocean, but is in line with the multi-model-mean. This implies that ozone concentrations might even become larger and dry deposition even lower in the model if a more advanced dry deposition scheme to the ocean would be implemented. With respect to vegetation, we might

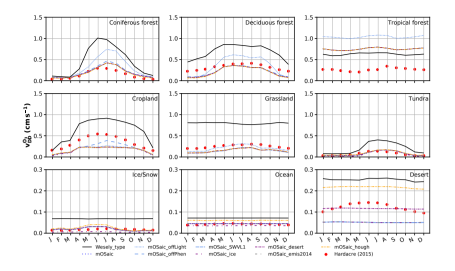


Figure 7. Average seasonal cycles of ozone dry deposition velocities separated by land use type. Results from (Hardacre et al., 2015) are shown as a reference. We refrain from showing the full extent of *EMEP_offLight* here, since it is an extreme scenario and has been discussed already.

be able to improve the model performance further by allowing more PFTs and phenologies, especially in regions covered by tropical forest (Anav et al., 2017) or in boreal regions.

Finally, we take a look at the different global as well as hemispheric dry deposition sinks for ozone (Table 3). Despite its vastness, the ocean amounts only to 35% of the global ozone sink due to dry deposition in the Oslo CTM3 with operative mOSaic scheme (*mOSaic*), while permanently ice and snow covered regions account for 1.2%. The remainder is deposited to land surfaces of which deserts might be the most neglected in process-modeling. The total annual dry deposition in the mOSaic scheme is one-third below the multi-model-mean result by Hardacre et al. (2015). But also the results of Luhar et al. (2017, 2018) yield a (19 - 27)% lower ozone dry deposition than the models participating in the model intercomparison, with deposition to ocean ranging between (12 - 21)% of the total annual ozone dry deposition. In particular, Luhar et al. (2018)

10 found that current model estimates of ozone dry deposition to the ocean may be three times too high compared to their analysis. This implies that the ozone dry deposition to the ocean in the Oslo CTM3 is too high as well.

Table 4 displays the average tropospheric ozone burden for all model experiments. Consistently with the previous findings, the mOSaic scheme increases the tropospheric ozone burden by 35 Tg. From various satellite ozone retrieval products, Gaudel et al. (2018, Tab. 5) deduce a lower limit estimate for global tropospheric ozone burden for the years 2010 - 2014 of 333 - 2010 - 2014 of 330 - 2014 of 30 - 2014 of

15 $345 \,\mathrm{Tg}$, but remark that this amount underestimates the actual tropospheric ozone burden, since it is only based on daytime retrievals. Nevertheless, the results of *mOSaic* lie 17% above that estimate and also well above the typical modeling range of $302 - 378 \,\mathrm{Tg}$ (Young et al., 2013). Despite the strong positive bias in ozone concentrations and accordingly low-bias in total dry deposition, the difference between *mOSaic* and *mOSaic* emis2014 (6 Tg), lies well within the range given by satellites for

Table 3. Total ozone dry deposition for the respective model experiment in $Tg a^{-1}$. The global ozone dry deposition has been weighted by ocean, ice and, land fraction in each gridbox, respectively. *Ice* herein refers to regions at high latitudes that are permanently covered by ice and snow.

Ocean		Ice			Land			Total			Δ^{\dagger}		
Experiment	NH	SH	Global	NH	SH	Global	NH	SH	Global	NH	SH	Global	
	$(Tg a^{-1})$		(Tga^{-1})		(Tga^{-1})			$(\mathrm{Tg}\mathrm{a}^{-1})$			(%)		
Wesely_type	160.5	147.7	308.2	7.0	6.4	13.4	417.3	190.8	608.2	613.4	344.9	958.3	36.7
mOSaic	108.1	105.3	213.4	4.3	3.1	7.4	236.2	130.3	366.4	368.3	238.6	606.9	0.0
mOSaic_offLight	110.5	106.3	216.8	4.3	3.0	7.4	263.1	145.3	408.4	399.9	254.7	654.6	7.3
mOSaic_offPhen	108.3	105.3	213.6	4.3	3.1	7.4	246.9	132.2	379.1	379.5	240.5	620.0	2.1
mOSaic_SWVL1	108.1	105.2	213.3	4.3	3.1	7.4	234.6	128.8	363.4	366.6	237.1	603.7	-0.5
mOSaic_ice	105.6	103.0	208.6	2.6	1.0	3.6	232.7	130.2	362.9	358.3	234.2	592.6	-2.4
mOSaic_desert	109.1	105.5	214.6	4.3	3.1	7.4	250.4	132.6	383.0	383.4	241.2	624.6	2.8
mOSaic_emis2014	108.9	107.0	215.9	4.3	3.1	7.3	238.3	133.8	372.1	370.8	243.9	614.7	1.3
mOSaic_hough	133.3	131.1	264.4	4.9	3.6	8.4	265.8	132.0	397.8	423.6	266.7	690.3	12.1

[†]: Relative change of global annual total in comparison to *mOSaic*.

the years 2005 and 2014 (Gaudel et al., 2018, Fig. 26). This indicates that the Oslo CTM3 responses responds well to given changes in global emissions.

Table 4. Annual mean tropospheric ozone burden for all experiments and 1σ standard deviation.

Experiment		op. O (Tg)	3
Wesely_type	364	±	23
mOSaic	399	±	31
mOSaic_offLight	395	±	30
mOSaic_offPhen	398	±	31
mOSaic_SWVL1	399	±	31
mOSaic_ice	401	±	31
mOSaic_desert	398	\pm	31
mOSaic_emis2014	405	±	32
mOSaic_hough	393	\pm	30

3.3 Comparison with MACC-reanalysis

In this section, we conclude the comparison of our results with respect to global ozone by looking at ECWMF's MACCreanalysis (MACC-II Consortium, 2011, data obtained from ECWMF's data center). In Fig, 8, we compare *mOSaic* ozone concentrations in the lowermost model level with surface concentrations deduced from the MACC-reanalysis for the year

- 5 2005. The MACC-reanalysis displays low ozone concentrations above all land masses except for the Greenland ice sheet. The lowest values are found in the deep tropics (e.g. northern South America and central Africa), while the highest values occur within (25-60)°N over the oceans. These low values over the oceans are relatively well reproduced by *mOSaic* (±20%). Over, e.g., South America, central and southern Africa, the Arabian peninsula, and the north-western Indian subcontinent, the ozone concentrations are elevated by up to 200%. While on global average much more concise Regarding the global average surface
- 10 <u>ozone concentration, Wesely_type</u> (Supplement S.8) is more consistent with the MACC-reanalysis , <u>Wesely_type</u> displays a similar tendency with respect to elevated ozone over continents. Enhanced ozone compared to MACC-reanalysis is found than *mOSaic*, but shows the same tendency of enhanced ozone over the continents as the latter. This enhancement is apparent mostly in the deep tropics and over the north-western Indian subcontinent (Supplement S.8) which coincides with regions of high intensity in incoming UV radiation. This may indicated an imbalance in the photo-chemical production and loss of O_3 in
- 15 the Oslo CTM3.

3.4 Comparison with ground-based observations

In this section, we compare our model results to observations at a selected number of sites which provide ozone flux measurements. For all comparisons, we use the original resolution of the Oslo CTM3 $(2.25^{\circ} \times 2.25^{\circ})$ instead of the re-gridded resolution $(3^{\circ} \times 3^{\circ})$.

In Fig. 9a, seasonal cycles of average ozone dry deposition fluxes for the six selected observation sites are shown. We have computed a model average for all sensitivity studies at the closest grid point and show the 1σ uncertainty band. The shaded area around the multi-model-mean indicates the broad range of model results but is not an actual uncertainty band since such is not given in Hardacre et al. (2015). At 4 of the 6 sites, the mOSaic scheme performs better than the Wesely scheme and similar to or better than the multi-model-mean. We use a χ^2 -test

25
$$\chi^2 = \sum_{i=1}^{12} \frac{\left(\overline{O}_{3\text{DD},i}^{\text{sim}} - \overline{O}_{3\text{DD},i}^{\text{obs}}\right)^2}{\sigma_i^2},$$
 (30)

with an estimated standard deviation of observation $\sigma_i = 1 \text{ mmol m}^{-2} \text{ s}^{-1}$ and divide it by the number of degrees of freedom (NDF) to assess this subjective analysis in a more objective way. The closer to 1 this test scores, the better does the simulation represent the observation. A score between 0 and 1 indicates that the estimated σ is too small. The results of the χ^2 -test are shown together with the divergences in Fig. 9b. The χ^2 -test reveals also that in 4 of the 6 cases the mOSaic scheme improves

30 the performance of the Oslo CTM3 with respect to observed ozone dry deposition fluxes, although a satisfying result is only achieved for two sites (Castel Porziano, Blodgett Forest). With only one full year of simulation, the model uncertainty regarding the seasonal cycle at observational sites cannot be properly quantified. Furthermore, the observational averages comprise at

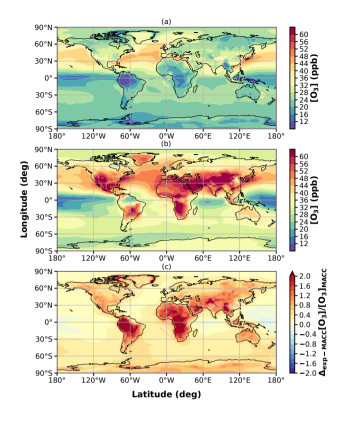


Figure 8. Mean ozone concentrations for the year 2005. (a) MACC-reanalysis (surface); (b) Oslo CTM3 *mOSaic* (lowermost model level); (c) Relative difference.

most 9 years worth of data. Statistically, these data may still be subject to interannual variability. Among other aspects, the horizontal as well as vertical resolution play an important role in the model performance. Although, we do not explicitly assess the impacts of differing resolutions in our model, we can assume that both high and low biases exist due to dilution of sources and sinks in coarse resolution models (Schaap et al., 2015). Good matches between observation and model are only to be expected if the station's location is representative for an area similar to the respective model gridbox and not substantially affected by differences in modeled and actual topography (e.g., major wind directions).

4 Summary and conclusions

5

10

We have presented an update of the dry deposition scheme in the Oslo CTM3 from purely prescribed dry deposition velocities (Wesely, 1989; Hough, 1991) to a more process-oriented parameterization taking the state of the atmosphere and vegetation into account. Based on the description of dry deposition in Simpson et al. (2003, 2012), we have implemented a moasic-mosaic approach to compute contributions to dry deposition by individual sub-grid land surface types. Aerodynamic, quasi-laminar,

24

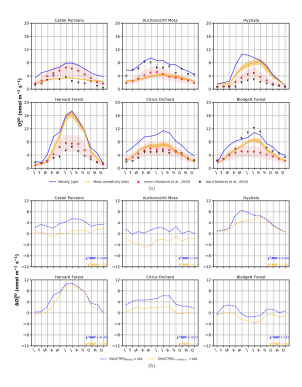


Figure 9. Ozone dry deposition fluxes at different observation sites. (a) Comparison between Wesely scheme and average result from all sensitivity studies with observational averages taken from Hardacre et al. (2015). The model uncertainty of the Oslo CTM3 is given as 1σ error band. The shaded area around the multi-model-mean indicates the broad range of different model results but is not an actual error band. (b) Model divergence from observation and χ^2 -test results.

and surface resistance (with the latter divided into stomatal and non-stomatal contributions) are calculated for each land surface category separately. Based on these, a land fraction-weighted mean is deduced. In addition, the various dry deposition velocities are now directly available from model output for diagnostics and further studies.

The new dry deposition scheme named *mOSaic* improves the modeled dry deposition velocities which are now compatible 5 with observation and model studies (e.g., Hardacre et al., 2015; Luhar et al., 2018). Dry deposition velocities are reduced by 6-60%. At the same time, the annual amount of ozone dry deposition decreases by more than 100% over all major desert areas and increases over tropical forest. Compared to results from a multi-model evaluation (Hardacre et al., 2015), the total annual ozone dry deposition in the Oslo CTM3 is $(38^{+1}_{-10})\%$ below average. However, there seems to be a tendency that newer TF HTAP models show a lower total annual dry deposition of ozone than older models, indicating that newer developments

10 lead to decreasing ozone dry deposition and increasing tropospheric ozone burden (e.g., Luhar et al., 2017, 2018; Hu et al., 2017).

We found the response of the Oslo CTM3 to the changes in dry deposition velocities from the old and the mOSaic scheme to be consistent. A decrease in $v_{\text{DD}}^{\text{O}_3}$ leads to a decrease in total ozone dry deposition and an increase in ozone concentration

 $[O_3]$. As the new scheme is quantitatively more similar to the multi-model-mean (Hardacre et al., 2015) with respect to dry deposition velocities, while the old scheme agrees better in terms of total dry deposition, there is an apparent discrepancy. By means of tropospheric ozone burden (Gaudel et al., 2018) and surface ozone concentrations deduced from the MACC-reanalysis (MACC-II Consortium, 2011), the Oslo CTM3 with the operational mOSaic scheme shows a pronounced high-bias

- 5 of tropospheric ozone. While the average bias is small or even reversed when using the old scheme, both display elevated ozone in comparison with the MACC-reanalysis in continental regions with high average incoming UV radiation (e.g., northern South America, central and southern Africa, the Himalayas). The reason behind this bias has not yet been resolved and may hint to, e.g., issues in photo-chemistry ([OH] related ozone production and loss) or previously introduced optimization of ozone removal with respect to the old, less physical dry deposition velocities.
- 10 Most of the qualitative change in ozone dry deposition in the Oslo CTM3 (-2 12%) can be attributed to changes in dry deposition over the ocean and deserts. This is mainly due to updates of the respective, prescribed ozone surface resistances R^{O_3} . In the case of desert and grasslands the difference between the old and new prescribed value is in the order of 1 an order of magnitude. Over the ocean, the absolute change in dry deposition is small, but it is accumulated over a large area which is especially amplified in the southern hemisphere. Small adjustments to the lower limits in our quasi-laminar layer
- 15 resistance formulation may help improve the Oslo CTM3 performance in this regard. With respect to available measurements of dry deposition velocities of ozone over desert (Güsten et al., 1996) and ocean (Helmig et al., 2012), the new Oslo CTM3 dry deposition scheme slightly underestimates ozone dry deposition velocities over the former and overestimate them over the latter. Regarding the vastness of the ocean and the ongoing desertification, it may be worthwhile to revise the dry deposition scheme for these regimes and add more process-oriented formulations, e.g., 2-layer gas exchange with ocean waters (Luhar
- 20 et al., 2017, 2018), wave braking breaking and spray (Pozzer et al., 2006).

Although dry deposition to ice and snow amounts to only 1% of the total global annual ozone dry deposition in *mOSaic*, a decrease in prescribed dry deposition velocity in accordance to combined measurements and model studies (Helmig et al., 2007) causes almost a doubling in the surface ozone concentrations in the high <u>Areties Arctic</u> and affects surface ozone concentrations down to latitudes at 50° in both hemispheres. Comparing with results from the multi-model evaluation (Hardacre

et al., 2015), we conclude that it is important to use this updated ozone dry deposition velocity to counter an Arctic surface ozone low-bias in models, however, this currently leads to an overcompensation (high-bias) in the Oslo CTM3.

We have studied the parameter space of the stomatal conductance parameterization and found that total surface ozone in the tropics and the northern hemisphere is most sensitive to changes therein. In the most extreme test case, the increase in global total dry deposition amounts to 7.3%, while the more realistic test cases, e.g. using differing years of emission amount

- 30 to changes in of the order of ±2%. This may indicate that future changes in vegetation cover and solar radiation at the surface due to changes in stratospheric ozone, cloud cover, or aerosols could also strongly influence the surface ozone burden in the tropics. Total column ozone in the tropics is predicted to decrease due to changes in the atmospheric circulation (e.g., WMO Global Ozone Research and Monitoring Project, 2014), while tropospheric and surface ozone increase. The combined effects of increasing emissions of ozone precursors and an increase in UV due to thinning of stratospheric ozone might permit more
- 35 UV light at ground and thus increase the ozone production.

An important factor in the global ozone budget are emissions of precursor substances. We cover this by using the same meteorology with different years of CEDS emissions. We chose the years 2005 and 2014 for our comparison. Ozone precursor emissions in 2014 are slightly lower in the NH while enhanced in the tropics and the SH. In 2014, surface ozone burden is higher in the southern hemisphere and in the tropics (5%) compared to 2005, while it is lower in the northern hemisphere (2%).

We also evaluated the model with respect to observed dry deposition fluxes at six sites in the northern hemisphere and found that the mOSaic scheme performs better than the old one, but is not able to reproduce the measurements at most sites quantitatively. This may be due to several reasons. The model resolution in both horizontal $(2.25^{\circ} \times 2.25^{\circ})$ and vertical (L60, $P_{\text{max}} = 0.02 \text{ hPa}$) does not capture all details in transport, thus affecting the distribution and transport (e.g., long-range, convection, and stratosphere–troposphere exchange) of ozone and its precursors. Depending on the location of the observation

- site and its respective representativeness for a larger area, ozone dry deposition and ozone concentrations are expected to be over- or underestimated in the model. Because of non-linearities in ozone formation and destruction, ozone concentrations are sensitive to both, differences in local concentration of precursors and meteorological conditions (Jin et al., 2013). In addition, a comparison of very few years of measurement to only one specific year of simulation may reflect the year to year variability
- 15 more than the actual model performance.

5

10

Future work on the Oslo CTM3 should resolve the ozone high-bias which may involve revising the photolysis- and chemical reaction computation as well as reaction rates. For a better modeling of ozone abundances, ocean emissions of very short-lived ozone depleting substances (VSLS) (Warwick et al., 2006; Ziska et al., 2013) which affect the stratospheric ozone (Hossaini et al., 2016; Falk et al., 2017) and a scheme covering aretic Arctic spring-time ozone depletion (e.g., Yang et al., 2010;
Toyota et al., 2011; Falk and Sinnhuber, 2018), could be worthwhile implementing. The general model performance could also be improved by allowing for more plant functional types and phenologies than currently used or implementing an actual photosynthesis-based modeling of plants. A more efficient parallelization of the code would enable computation on at higher horizontal resolutions.

Code and data availability. The Oslo CTM3 shall be publicly available on git-hub under a MIT license in the future. Until then, access can
be made granted under request. Model results can be made available under request.

Appendix A: Figures

30

Author contributions. Stefanie Falk has compiled the manuscript, finalized and revised the implementation of the stomatal conductance in the *mOSaic* dry deposition scheme of the Oslo CTM3, conducted the simulations, and analyzed and evaluated the results. Amund Søvde Haslerud has implemented the *mOSaic* dry deposition scheme and wrote the respective documentation. Both authors contributed to the writing and discussion of the paper.

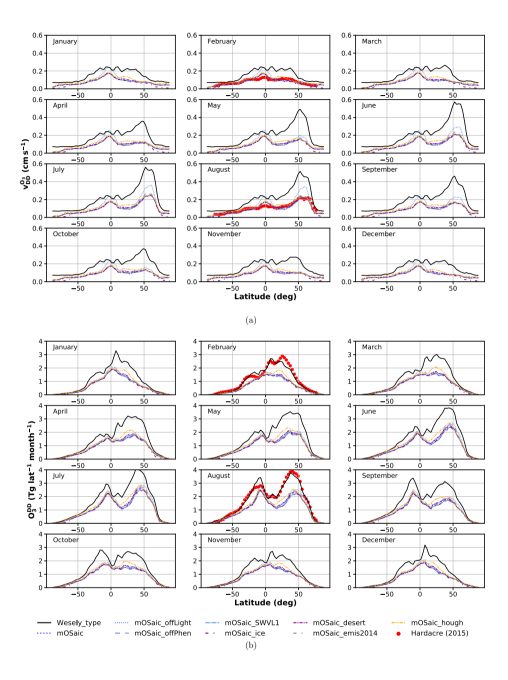


Figure A1. Comparison of the manifold of Oslo CTM3 integrations with respect to (a) Zonal average ozone dry deposition velocities; (b) Total annual amount of ozone removed from the atmosphere via dry deposition. The multi-model mean from the evaluation of TF HTAP models by Hardacre et al. (2015) is shown as a reference (where available).

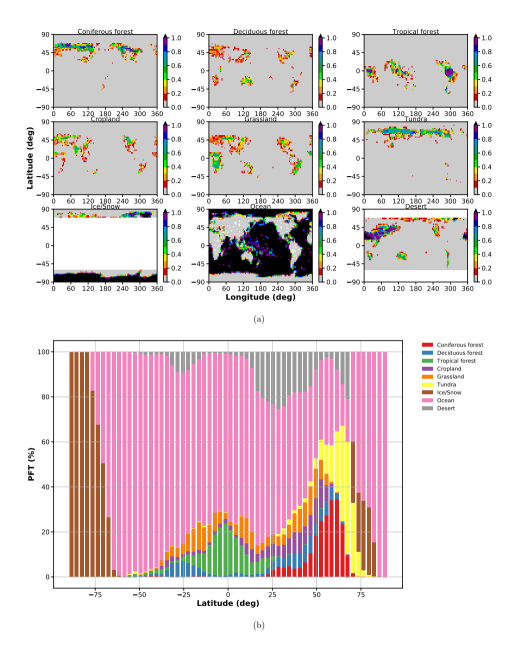


Figure B1. Partitioning of land surface types. (a) CLM 2 dynamic land surface types in $(0.5 \times 0.5)^{\circ}$ resolution; (b) Zonal distribution of land surface types.

5

Acknowledgements. This work was supported by the Norwegian Research Council (NRC) through the project The double punch: Ozone and climate stresses on vegetation (268073).

The simulations were performed on resources provided by UNINETT Sigma2 – the National Infrastructure for High Performance Computing and Data Storage in Norway (project nn2806k).

The used Leaf Area Index (LAI) and roughness length (*z*₀) are available online from Oak Ridge National Laboratory Distributed Active Archive Center, Oak Ridge, Tennessee, U.S.A. (https://doi.org/10.3334/ORNLDAAC/970). Community Emission Data System (CEDS) historical emission inventory is provided by the Joint Global Research Institute project (http://www.globalchange.umd.edu/ceds/.) Randerson, J.T., G.R. van der Werf, L. Giglio, G.J. Collatz, and P.S. Kasibhatla. 2018. Global Fire

10 Emissions Database, Version 4, (GFEDv4). ORNL DAAC, Oak Ridge, Tennessee, USA. https://doi.org/10.3334/ORNLDAAC/1293 We would like to thank Prof. Frode Stordal (Section for Meteorology and Oceanography, University of Oslo) for discussions regarding early drafts of the manuscript, Anne Fouilloux (scientific programmer at the same institute) for technical support as well as Olimpia Bruno (Karlsruhe Institute of Technology) and Franziska Hellmuth (University of Oslo) for valuable input regarding the aerodynamic resistance formulation. We would also like to thank the Center for International Climate Research (CICERO) for their support of this work.

References

5

15

Ainsworth, E. A.: Understanding and improving global crop response to ozone pollution, Plant J., 90, 886–897, https://doi.org/10.1111/tpj.13298, 2017.

Anav, A., Liu, Q., De Marco, A., Proietti, C., Savi, F., Paoletti, E., and Piao, S.: The role of plant phenology in stomatal ozone flux modeling, Glob. Change Biol., 24, 235–248. https://doi.org/10.1111/gcb.13823. https://onlinelibrary.wiley.com/doi/abs/10.1111/gcb.13823. 2017.

Anderson, M. C., Norman, J. M., Meyers, T. P., and Diak, G. R.: An analytical model for estimating canopy transpiration and carbon assimilation fluxes based on canopy light-use efficiency, Agr. Forest Meteorol., 101, 265–289, https://doi.org/10.1016/S0168-1923(99)00170-7, 2000.

Ball, J., Woodrow, I., and Berry, J.: Progress in Photosynthesis Research, chap. A Model Predicting Stomatal Conductance and its Contribu-

tion to the Control of Photosynthesis under Different Environmental Conditions, Springer, Dordrecht, 1987.
 Buckley, T. N.: Modeling Stomatal Conductance, Plant Physiol., 174, 572–582, https://doi.org/10.1104/pp.16.01772, 2017.
 Charnock, H.: Wind stress on a water surface, Q. J. Roy. Meteorol. Soc., 81, 639–640, https://doi.org/10.1002/qj.49708135027, https://rmets.onlinelibrary.wiley.com/doi/abs/10.1002/qj.49708135027, 1955.

Chuwah, C., van Noije, T., van Vuuren, D. P., Stehfest, E., and Hazeleger, W.: Global impacts of surface ozone changes on crop yields and land use, Atmos. Environ., 106, 11–23, https://doi.org/10.1016/j.atmosenv.2015.01.062, 2015.

- Derwent, R. G., Parrish, D. D., Galbally, I. E., Stevenson, D. S., Doherty, R. M., Naik, V., and Young, P. J.: Uncertainties in models of tropospheric ozone based on Monte Carlo analysis: Tropospheric ozone burdens, atmospheric lifetimes and surface distributions, Atmos. Environ., 180, 93–102, https://doi.org/10.1016/j.atmosenv.2018.02.047, 2018.
 - Emberson, L. D., for Monitoring, C. P., of the Long Range Transmission of Air Pollutants in Europe, E., meteorologiske institutt, N.,
- 20 and Meteorological Synthesizing Centre-West (Oslo, N.: Towards a Model of Ozone Deposition and Stomatal Uptake Over Europe, EMEP/MSC-W Note, Norwegian Meteorological Institute, https://books.google.no/books?id=hj_ntgAACAAJ, 2000.

Erisman, J. W., Van Pul, A., and Wyers, P.: Parametrization of Surface-Resistance for the Quantification of Atmospheric Deposition of Acidifying Pollutants and Ozone, Atmos. Environ., 28, 2595–2607, https://doi.org/10.1016/1352-2310(94)90433-2, 1994.

Falk, S. and Sinnhuber, B.-M.: Polar boundary layer bromine explosion and ozone depletion events in the chemistry-climate model EMAC

- v2.52: implementation and evaluation of AirSnow algorithm, Geosci. Model Dev., 11, 1115–1131, https://doi.org/10.5194/gmd-11-1115-2018, 2018.
 - Falk, S., Sinnhuber, B.-M., Krysztofiak, G., Jöckel, P., Graf, P., and Lennartz, S. T.: Brominated VSLS and their influence on ozone under a changing climate, Atmos. Chem. Phys., 17, 11 313–11 329, https://doi.org/10.5194/acp-17-11313-2017, https://www.atmos-chem-phys. net/17/11313/2017/, 2017.
- 30 Fleming, Z. L., Doherty, R. M., von Schneidemesser, E., Malley, C. S., Cooper, O. R., Pinto, J. P., Colette, A., Xu, X., Simpson, D., Schultz, M. G., Lefohn, A. S., Hamad, S., Moolla, R., Solberg, S., and Feng, Z.: Tropospheric Ozone Assessment Report: Present-day ozone distribution and trends relevant to human health, ELEMENTA-SCIENCE OF THE ANTHROPOCENE, 6, https://doi.org/10.1525/elementa.273, 2018.

Fu, Y., Zhang, H., Dong, W., and Yuan, W.: Comparison of Phenology Models for Predicting the Onset of Growing Season over the Northern

35 Hemisphere, Plos One, 9, https://doi.org/10.1371/journal.pone.0109544, 2014a.

Fu, Y. H., Piao, S., Zhao, H., Jeong, S.-J., Wang, X., Vitasse, Y., Ciais, P., and Janssens, I. A.: Unexpected role of winter precipitation in determining heat requirement for spring vegetation green-up at northern middle and high latitudes, Glob. Change Biol., 20, 3743–3755, https://doi.org/10.1111/gcb.12610, 2014b.

Garratt, J. R.: The Atmospheric Boundary Layer, University Press, Cambridge, 1992.

- 5 Gaudel, A., Cooper, O. R., Ancellet, G., Barret, B., Boynard, A., Burrows, J. P., Clerbaux, C., Coheur, P. F., Cuesta, J., Cuevas, E., Doniki, S., Dufour, G., Ebojie, F., Foret, G., Garcia, O., Granados-Munoz, M. J., Hannigan, J. W., Hase, F., Hassler, B., Huang, G., Hurtmans, D., Jaffe, D., Jones, N., Kalabokas, P., Kerridge, B., Kulawik, S., Latter, B., Leblanc, T., Le Flochmoen, E., Lin, W., Liu, J., Liu, X., Mahieu, E., McClure-Begley, A., Neu, J. L., Osman, M., Palm, M., Petetin, H., Petropavlovskikh, I., Querel, R., Rahpoe, N., Rozanov, A., Schultz, M. G., Schwab, J., Siddans, R., Smale, D., Steinbacher, M., Tanimoto, H., Tarasick, D. W., Thouret, V., Thompson, A. M.,
- 10 Trickl, T., Weatherhead, E., Wespes, C., Worden, H. M., Vigouroux, C., Xu, X., Zeng, G., and Ziemke, J.: Tropospheric Ozone Assessment Report: Present-day distribution and trends of tropospheric ozone relevant to climate and global atmospheric chemistry model evaluation, ELEMENTA-SCIENCE OF THE ANTHROPOCENE, 6, https://doi.org/10.1525/elementa.291, 2018.
 - GEIA-ACCENT emission data portal: POET, online, Global CO emissions (1990–2000), http://accent.aero.jussieu.fr/database_table_ inventories.php, 2003.
- 15 Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P. I., and Geron, C.: Estimates of global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from Nature), Atmos. Chem. Phys., 6, 3181–3210, https://doi.org/10.5194/acp-6-3181-2006, 2006.
 - Güsten, H., Heinrich, G., Mönnich, E., Sprung, D., Weppner, J., Ramadan, A. B., Ezz El-din, M. R. M., Ahmed, D. M., and Hassan, G. K. Y.: On-line measurements of ozone surface fluxes: Part II. Surface-level ozone fluxes onto the sahara desert, Atmos. Environ., 30, 911–918,
- 20 https://doi.org/10.1016/1352-2310(95)00270-7, 1996.
 - Hardacre, C., Wild, O., and Emberson, L.: An evaluation of ozone dry deposition in global scale chemistry climate models, Atmos. Chem. Phys., 15, 6419–6436, https://doi.org/10.5194/acp-15-6419-2015, 2015.
 - Helmig, D., Ganzeveld, L., Butler, T., and Oltmans, S. J.: The role of ozone atmosphere-snow gas exchange on polar, boundary-layer tropospheric ozone a review and sensitivity analysis, Atmos. Chem. Phys., 7, 2007.
- 25 Helmig, D., Lang, E. K., Bariteau, L., Boylan, P., Fairall, C. W., Ganzeveld, L., Hare, J. E., Hueber, J., and Pallandt, M.: Atmosphereocean ozone fluxes during the TexAQS 2006, STRATUS 2006, GOMECC 2007, GasEx 2008, and AMMA 2008 cruises, J. Geophys. Res.-Atmos., 117, https://doi.org/10.1029/2011JD015955, 2012.

Hinze, J. O.: Turbulence, p. 790, McGraw-Hill Series in Mechanical Engineering, McGraw-Hill, New York, 1975.

Hoesly, R. M., Smith, S. J., Feng, L., Klimont, Z., Janssens-Maenhout, G., Pitkanen, T., Seibert, J. J., V., L., Andres, R. J., Bolt, R. M., Bond,

- 30 T. C., Dawidowski, L., Kholod, N., Kurokawa, J., Li, M., Liu, L., Lu, Z., Moura, M. C. P., O'Rourke, P. R., and Zhang, Q.: Historical (1750-2014) anthropogenic emissions of reactive gases and aerosols from the Community Emissions Data System (CEDS), Geosci. Model Dev., 11, 369–408, https://doi.org/10.5194/gmd-11-369-2018, 2018.
 - Hoshika, Y., Katata, G., Deushi, M., Watanabe, M., Koike, T., and Paoletti, E.: Ozone-induced stomatal sluggishness changes carbon and water balance of temperate deciduous forests, Sci. Rep.-UK, 5, https://doi.org/10.1038/srep09871, 2015.
- 35 Hossaini, R., Patra, P. K., Leeson, A. A., Krysztofiak, G., Abraham, N. L., Andrews, S. J., Archibald, A. T., Aschmann, J., Atlas, E. L., Belikov, D. A., Bönisch, H., Carpenter, L. J., Dhomse, S., Dorf, M., Engel, A., Feng, W., Fuhlbrügge, S., Griffiths, P. T., Harris, N. R. P., Hommel, R., Keber, T., Krüger, K., Lennartz, S. T., Maksyutov, S., Mantle, H., Mills, G. P., Miller, B., Montzka, S. A., Moore, F., Navarro, M. A., Oram, D. E., Pfeilsticker, K., Pyle, J. A., Quack, B., Robinson, A. D., Saikawa, E., Saiz-Lopez, A., Sala, S., Sinnhuber,

B.-M., Taguchi, S., Tegtmeier, S., Lidster, R. T., Wilson, C., and Ziska, F.: A multi-model intercomparison of halogenated very short-lived substances (TransCom-VSLS): linking oceanic emissions and tropospheric transport for a reconciled estimate of the stratospheric source gas injection of bromine, Atmos. Chem. Phys., 16, 9163–9187, https://doi.org/10.5194/acp-16-9163-2016, http://www.atmos-chem-phys. net/16/9163/2016/, 2016.

- 5 Hough, A. M.: Development of a 2-Dimensional Global Tropospheric Model - Model Chemistry, J. Geophys. Res.-Atmos., 96, 7325-7362, https://doi.org/10.1029/90JD01327, 1991.
 - Hu, L., Jacob, D. J., Liu, X., Zhang, Y., Zhang, L., Kim, P. S., Sulprizio, M. P., and Yantosca, R. M.: Global budget of tropospheric ozone: Evaluating recent model advances with satellite (OMI), aircraft (IAGOS), and ozonesonde observations, Atmos, Environ, 167, 323–334. https://doi.org/10.1016/j.atmosenv.2017.08.036, 2017.
- 10 IPCC - Intergovernmental Panel on Climate Change: Climate Change 2013: The Physical Science Basis, 2013. Jarvis, P. G.: The interpretation of the variations in leaf water potential and stomatal conductance found in canopies in the field, Philos. T. Roy. Soc. B., 273, 593-610, https://doi.org/10.1098/rstb.1976.0035, http://rstb.royalsocietypublishing.org/content/273/927/593, 1976.
 - Jin, L., Loisy, A., and Brown, N.: Role of meteorological processes in ozone responses to emission controls in California's San Joaquin Valley, J. Geophys. Res.-Atmos., 118, 8010-8022, https://doi.org/10.1002/jgrd.50559, 2013.
- 15 Levis, S. and Bonan, G. B.: Simulating springtime temperature patterns in the community atmosphere model coupled to the community land model using prognostic leaf area, J. Climate, 17, 4531–4540, https://doi.org/10.1175/3218.1, 2004.
 - Luhar, A. K., Galbally, I. E., Woodhouse, M. T., and Thatcher, M.: An improved parameterisation of ozone dry deposition to the ocean and its impact in a global climate-chemistry model, Atmos. Chem. Phys., 17, 3749–3767, https://doi.org/10.5194/acp-17-3749-2017, 2017.
 - Luhar, A. K., Woodhouse, M. T., and Galbally, I. E.: A revised global ozone dry deposition estimate based on a new two-layer
- 20 parameterisation for air-sea exchange and the multi-year MACC composition reanalysis, Atmos. Chem. Phys., 18, 4329-4348, https://doi.org/10.5194/acp-18-4329-2018, 2018.
 - Lund, M. T., Myhre, G., Haslerud, A. S., Skeie, R. B., Griesfeller, J., Platt, S. M., Kumar, R., Myhre, C. L., and Schulz, M.: Concentrations and radiative forcing of anthropogenic aerosols from 1750 to 2014 simulated with the Oslo CTM3 and CEDS emission inventory, Geoscientific Model Development, 11, 4909–4931, https://doi.org/10.5194/gmd-11-4909-2018, https://www.geosci-model-dev.net/11/4909/2018/, 2018.
- 25

30

- MACC-II Consortium: MACC Reanalysis of Global Atmospheric Composition (2003-2012), https://atmosphere.copernicus.eu/catalogue#/, Copernicus Atmosphere Monitoring Service (CAMS), 2011.
- Matyssek, R., Wieser, G., Calfapietra, C., de Vries, W., Dizengremel, P., Ernst, D., Jolivet, Y., Mikkelsen, T. N., Mohren, G. M. J., Le Thiec, D., Tuovinen, J. P., Weatherall, A., and Paoletti, E.: Forests under climate change and air pollution: Gaps in understanding and future directions for research, Environ. Pollut., 160, 57-65, https://doi.org/10.1016/j.envpol.2011.07.007, 2012.
- Mills, G., Sharps, K., Simpson, D., Pleijel, H., Broberg, M., Uddling, J., Jaramillo, F., Davies, W. J., Dentener, F., Van den Berg, M., Agrawal, M., Agrawal, S. B., Ainsworth, E. A., Büker, P., Emberson, L., Feng, Z., Harmens, H., Hayes, F., Kobayashi, K., Paoletti, E., and Van Dingenen, R.: Ozone pollution will compromise efforts to increase global wheat production, Glob. Change Biol., 24, 3560-3574, https://doi.org/10.1111/gcb.14157, https://onlinelibrary.wiley.com/doi/abs/10.1111/gcb.14157, 2018.
- Mills, G., Harmens, H., Hayes, F., Pleijel, H., Büker, P., González-Fernandéz, I., Alonso, R., Bender, J., Bergmann, E. Bermejo, V., Braun, 35 S., Danielsson, H., Gerosa, G., Grünhage, L., Karlsson, P. E., Marzuoli, R., Schaub, M., and Simpson, D.: ICP Mapping Manual - Chapter III: Mapping Critical Levels for Vegetation, International Cooperative Programme on Effects of Air Pollution on Natural Vegetation and Crops, https://icpmapping.org/Latest_update_Mapping_Manual, minor edits OCT 2017, 2017.

- Mills, G., Pleijel, H., Malley, C. S., Sinha, B., Cooper, O. R., Schultz, M. G., Neufeld, H. S., Simpson, D., Sharps, K., Feng, Z., Gerosa, G., Harmens, H., Kobayashi, K., Saxena, P., Paoletti, E., Sinha, V., and Xu, X.: Tropospheric Ozone Assessment Report: Present-day tropospheric ozone distribution and trends relevant to vegetation, ELEMENTA-SCIENCE OF THE ANTHROPOCENE, 6, https://doi.org/10.1525/elementa.302, 2018.
- 5 Monks, P. S., Archibald, A. T., Colette, A., Cooper, O., Coyle, M., Derwent, R., Fowler, D., Granier, C., Law, K. S., Mills, G. E., Stevenson, D. S., Tarasova, O., Thouret, V., von Schneidemesser, E., Sommariva, R., Wild, O., and Williams, M. L.: Tropospheric ozone and its precursors from the urban to the global scale from air quality to short-lived climate forcer, Atmos. Chem. Phys., 15, 8889–8973, https://doi.org/10.5194/acp-15-8889-2015, 2015.
 - Otero, N., Sillmann, J., Mar, K. A., Rust, H. W., Solberg, S., Andersson, C., Engardt, M., Bergstrom, R., Bessagnet, B., Colette, A., Couvidat,
- 10 F., Cuvelier, C., Tsyro, S., Fagerli, H., Schaap, M., Manders, A., Mircea, M., Briganti, G., Cappelletti, A., Adani, M., D'Isidoro, M., Pay, M.-T., Theobald, M., Vivanco, M. G., Wind, P., Ojha, N., Raffort, V., and Butler, T.: A multi-model comparison of meteorological drivers of surface ozone over Europe, Atmos. Chem. Phys., 18, 12 269–12 288, https://doi.org/10.5194/acp-18-12269-2018, 2018.
 - Pozzer, A., Joeckel, P. J., Sander, R., Williams, J., Ganzeveld, L., and Lelieveld, J.: Technical note: the MESSy-submodel AIRSEA calculating the air-sea exchange of chemical species, Atmos. Chem. Phys., 6, 5435–5444, 2006.
- 15 Schaap, M., Cuvelier, C., Hendriks, C., Bessagnet, B., Baldasano, J. M., Colette, A., Thunis, P., Karam, D., Fagerli, H., Graff, A., Kranenburg, R., Nyiri, A., Pay, M. T., Rouil, L., Schulz, M., Simpson, D., Stern, R., Terrenoire, E., and Wind, P.: Performance of European chemistry transport models as function of horizontal resolution, Atmos. Environ., 112, 90–105, https://doi.org/10.1016/j.atmosenv.2015.04.003, 2015.
 - Schnell, J. L., Prather, M. J., Josse, B., Naik, V., Horowitz, L. W., Cameron-Smith, P., Bergmann, D., Zeng, G., Plummer, D. A., Sudo, K.,
- 20 Nagashima, T., Shindell, D. T., Faluvegi, G., and Strode, S. A.: Use of North American and European air quality networks to evaluate global chemistry–climate modeling of surface ozone, Atmospheric Chemistry and Physics, 15, 10581–10596, https://doi.org/10.5194/acp-15-10581-2015, https://www.atmos-chem-phys.net/15/10581/2015/, 2015.
 - Seinfeld, J. H. and Pandis, S. N.: Atmospheric Chemistry and Physics: From Air Pollution to Climate Change, chap. 19, John Wiley & Sons, New York, 2nd edn., 2006.
- 25 Simpson, D., Tuovinen, J. P., Emberson, L., and Ashmore, M. R.: Characteristics of an ozone deposition module II: Sensitivity analysis, Water Air Soil Poll., 143, 123–137, https://doi.org/10.1023/A:1022890603066, 2003.
 - Simpson, D., Benedictow, A., Berge, H., Bergstrom, R., Emberson, L. D., Fagerli, H., Flechard, C. R., Hayman, G. D., Gauss, M., Jonson, J. E., Jenkin, M. E., Nyiri, A., Richter, C., Semeena, V. S., Tsyro, S., Tuovinen, J.-P., Valdebenito, A., and Wind, P.: The EMEP MSC-W chemical transport model technical description, Atmos. Chem. Phys., 12, 7825–7865, https://doi.org/10.5194/acp-12-7825-2012, 2012.
- 30 Sindelarova, K., Granier, C., Bouarar, I., Guenther, A., Tilmes, S., Stavrakou, T., Muller, J. F., Kuhn, U., Stefani, P., and Knorr, W.: Global data set of biogenic VOC emissions calculated by the MEGAN model over the last 30 years, Atmos. Chem. Phys., 14, 9317–9341, https://doi.org/10.5194/acp-14-9317-2014, 2014.
 - Sitch, S., Cox, P. M., Collins, W. J., and Huntingford, C.: Indirect radiative forcing of climate change through ozone effects on the land-carbon sink, Nature, 448, 791–U4, https://doi.org/10.1038/nature06059, 2007.
- 35 Søvde, O. A., Prather, M. J., Isaksen, I. S. A., Berntsen, T. K., Stordal, F., Zhu, X., Holmes, C. D., and Hsu, J.: The chemical transport model Oslo CTM3, Geosci. Model Dev., 5, 1441–1469, https://doi.org/10.5194/gmd-5-1441-2012, 2012.
 - Stevenson, D. S., Dentener, F. J., Schultz, M. G., Ellingsen, K., van Noije, T. P. C., Wild, O., Zeng, G., Amann, M., Atherton, C. S., Bell, N., Bergmann, D. J., Bey, I., Butler, T., Cofala, J., Collins, W. J., Derwent, R. G., Doherty, R. M., Drevet, J., Eskes, H. J., Fiore, A. M., Gauss,

M., Hauglustaine, D. A., Horowitz, L. W., Isaksen, I. S. A., Krol, M. C., Lamarque, J.-F., Lawrence, M. G., Montanaro, V., Müller, J.-F., Pitari, G., Prather, M. J., Pyle, J. A., Rast, S., Rodriguez, J. M., Sanderson, M. G., Savage, N. H., Shindell, D. T., Strahan, S. E., Sudo, K., and Szopa, S.: Multimodel ensemble simulations of present-day and near-future tropospheric ozone, J. Geophys. Res.-Atmos., 111, 2005.

- Tai, A. P. K., Martin, M. V., and Heald, C. L.: Threat to future global food security from climate change and ozone air pollution, Nat. Clim.
 Change, 4, 817–821, https://doi.org/10.1038/NCLIMATE2317, 2014.
- Tang, H., Takigawa, M., Liu, G., Zhu, J., and Kobayashi, K.: A projection of ozone-induced wheat production loss in China and India for the years 2000 and 2020 with exposure-based and flux-based approaches, Glob. Change Biol., 19, 2739–2752, https://doi.org/10.1111/gcb.12252, 2013.
- Toyota, K., McConnell, J. C., Lupu, A., Neary, L., McLinden, C. A., Richter, A., Kwok, R., Semeniuk, K., Kaminski, J. W., Gong,
 S. L., Jarosz, J., Chipperfield, M. P., and Sioris, C. E.: Analysis of reactive bromine production and ozone depletion in the Arctic boundary layer using 3-D simulations with GEM-AQ: inference from synoptic-scale patterns, Atmos. Chem. Phys., 11, 3949–3979, https://doi.org/10.5194/acp-11-3949-2011, 2011.
 - Tuovinen, J. P., Ashmore, M. R., Emberson, L. D., and Simpson, D.: Testing and improving the EMEP ozone deposition module, Atmos. Environ., 38, 2373–2385, https://doi.org/10.1016/j.atmosenv.2004.01.026, Workshop on the Development and Application of New Methods
- 15 of Ozone Risk Assessment for Europe, Hindas, SWEDEN, NOV, 2002, 2004.
 - Warwick, N. J., Pyle, J. A., Carver, G. D., Yang, X., Savage, N. H., O'Connor, F. M., and Cox, R. A.: Global modeling of biogenic bromocarbons, J. Geophys. Res.-Atmos., 111, https://doi.org/10.1029/2006JD007264, 2006.

Wesely, M. L.: Parameterization Of Surface Resistances To Gaseous Dry Deposition In Regional-Scale Numerical-Models, Atmos. Environ., 23, 1293–1304, https://doi.org/10.1016/0004-6981(89)90153-4, 1989.

- 20 WHO World Health Organization: Health risks of ozone from long-range transboundary air pollution, 2008.
 - Wittig, V. E., Ainsworth, E. A., Naidu, S. L., Karnosky, D. F., and Long, S. P.: Quantifying the impact of current and future tropospheric ozone on tree biomass, growth, physiology and biochemistry: a quantitative meta-analysis, Glob. Change Biol., 15, 396–424, https://doi.org/10.1111/j.1365-2486.2008.01774.x, 2009.

WMO - Global Ozone Research and Monitoring Project: Scientific Assessment of Ozone Depletion: 2014, 2014.

- 25 Wu, J.: Wind-Stress Coefficients Over Sea-Surface Near Neutral Conditions A Revisit, J. Phys. Oceanogr., 10, 727–740, https://doi.org/10.1175/1520-0485(1980)010<0727:WSCOSS>2.0.CO;2, 1980.
 - Yang, X., Pyle, J. A., Cox, R. A., Theys, N., and Van Roozendael, M.: Snow-sourced bromine and its implications for polar tropospheric ozone, Atmos. Chem. Phys., 10, 7763–7773, https://doi.org/10.5194/acp-10-7763-2010, 2010.

Young, P. J., Archibald, A. T., Bowman, K. W., Lamarque, J.-F., Naik, V., Stevenson, D. S., Tilmes, S., Voulgarakis, A., Wild, O., Bergmann,

- 30 D., Cameron-Smith, P., Cionni, I., Collins, W. J., Dalsøren, S. B., Doherty, R. M., Eyring, V., Faluvegi, G., Horowitz, L. W., Josse, B., Lee, Y. H., MacKenzie, I. A., Nagashima, T., Plummer, D. A., Righi, M., Rumbold, S. T., Skeie, R. B., Shindell, D. T., Strode, S. A., Sudo, K., Szopa, S., and Zeng, G.: Pre-industrial to end 21st century projections of tropospheric ozone from the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP), Atmos. Chem. Phys., 13, 2063–2090, https://doi.org/10.5194/acp-13-2063-2013, https://www.atmos-chem-phys.net/13/2063/2013/, 2013.
- 35 Ziska, F., Quack, B., Abrahamsson, K., Archer, S. D., Atlas, E., Bell, T., Butler, J. H., Carpenter, L. J., Jones, C. E., Harris, N. R. P., Hepach, H., Heumann, K. G., Hughes, C., Kuss, J., Krueger, K., Liss, P., Moore, R. M., Orlikowska, A., Raimund, S., Reeves, C. E., Reifenhaeuser, W., Robinson, A. D., Schall, C., Tanhua, T., Tegtmeier, S., Turner, S., Wang, L., Wallace, D., Williams, J., Yamamoto, H., Yvon-Lewis,

S., and Yokouchi, Y.: Global sea-to-air flux climatology for bromoform, dibromomethane and methyl iodide, Atmos. Chem. Phys., 13, 8915–8934, https://doi.org/10.5194/acp-13-8915-2013, 2013.