Dear Fiona,

Thank you for the positive feedback and comments to our manuscript. We have made additional clarifications in the methodology section that we hope address any remaining issues and misunderstandings.

Specifically, we have we have added "offline" when describing the OsloCTM3 (first paragraph section 2.1) and specified "fixed meterological data" in section 2.3. We have also modified the beginning of section 2.4, hich now reads: "We calculate the instantaneous top-of-the atmosphere radiative forcing of anthropogenic aerosols due to aerosol-radiation interactions (RFari) [*Myhre et al.*, 2013b]). The radiative transfer calculations are performed offline with a multi-stream model using the discrete ordinate method [*Stamnes et al.*, 1988]."

Thank you again for considering our study for publication in Geoscientific Model Development.

Sincerely, Marianne T. Lund CICERO, Center for International Climate Research

1 2 3	Concentrations and radiative forcing of anthropogenic aerosols from 1750-2014 simulated with the OsloCTM3 and CEDS emission inventory
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Abstract

We document the ability of the new generation Oslo chemistry-transport model, OsloCTM3, to accurately simulate present-day aerosol distributions. The model is then used with the new Community Emission Data System (CEDS) historical emission inventory to provide updated time series of anthropogenic aerosol concentrations and consequent direct radiative forcing (RFari)

from 1750 to 2014.

Overall, the OsloCTM3 performs well compared with measurements of surface concentrations and remotely sensed aerosol optical depth. Concentrations are underestimated in Asia, but the higher emissions in CEDS than previous inventories result in improvements compared to observations. The treatment of black carbon (BC) scavenging in OsloCTM3 gives better agreement with observed vertical BC profiles relative to the predecessor OsloCTM2. However, Arctic wintertime BC concentrations remain underestimated, and a range of sensitivity tests indicate that better physical understanding of processes associated with atmospheric BC processing is required to simultaneously reproduce both the observed features. Uncertainties in model input data, resolution and scavenging affects the distribution of all aerosols species, especially at high latitudes and altitudes. However, we find no evidence of consistently better model performance across all

observables and regions in the sensitivity tests than in the baseline configuration.

Using CEDS, we estimate a net RFari in 2014 relative to 1750 of -0.17 W m⁻², significantly weaker than the IPCC AR5 2011-1750 estimate. Differences are attributable to several factors, including

stronger absorption by organic aerosol, updated parameterization of BC absorption, and reduced sulfate cooling. The trend towards a weaker RFari over recent years is more pronounced than in

the IPCC AR5, illustrating the importance of capturing recent regional emission changes.

- 62 1 Introduction
- 63

Changes in anthropogenic emissions over the industrial period have significantly altered the 64 abundance, composition and properties of atmospheric aerosols, causing a change in the radiative 65 energy balance. The net energy balance change is determined by a complex interplay of different 66 types of aerosols and their interactions with radiation and clouds, causing both positive (warming) 67 and negative (cooling) radiative impacts. Global aerosols were estimated by the Intergovernmental 68 Panel on Climate Change fifth assessment report (IPCC AR5) to have caused an effective radiative 69 forcing (ERF) of -0.9 W m⁻² over the industrial era from 1750 to 2011, but with considerable 70 uncertainty (-1.9 to -0.1 W m⁻²) [Boucher et al., 2013]. This large uncertainty range arises from a 71 number of factors, including uncertainties in emissions and the simulated spatiotemporal 72 distribution of aerosols, their chemical composition and properties. 73 74 Historical emission estimates for anthropogenic aerosol and precursor compounds are key data needed for climate and atmospheric chemistry transport models in order to examine how these 75 drivers have contributed to climate change. The Community Emissions Data System (CEDS) 76 77 recently published a new time series of emissions from 1750 to 2014, which will be used in the 78 upcoming CMIP6 [Hoesly et al., 2018]. CEDS includes several improvements, including annual 79 temporal resolution with seasonal cycles, consistent methodology between different species, and 80 extending the time series to more recent years, compared to previous inventories and assessments

- [e.g., Lamarque et al., 2010; Taylor et al., 2012]. During the period from 2000 to 2014, global 81 emissions of black carbon (BC) and organic carbon (OC) have increased, while nitrogen oxide 82 83 (NOx) emissions have been relatively constant after 2008, and sulfur dioxide (SO₂) emissions were back at 2000 levels in 2014, after a temporary increase [Hoesly et al., 2018]. Furthermore, both 84 CEDS and other recent emission inventories report considerably higher estimates of global BC 85 and OC emissions in recent years than earlier inventories [Granier et al., 2011; Klimont et al., 86 87 2017; Lamarque et al., 2010; Wang et al., 2014]. The global trend in emissions is driven by a strong increase in emissions from Asia and Africa, and a decline in North America and Europe. 88 Capturing such geographical differences is essential, as the distribution, lifetime and radiative 89
- 90 forcing of aerosols depend on their location.

91 After emission or formation, particles undergo transport, mixing, chemical aging and removal by dry and wet deposition, resulting in a short atmospheric residence time, and a highly heterogeneous 92 93 distribution in space and time. Consequently, accurate representation of observed aerosols remains 94 challenging, and previous studies have shown that considerable diversity in the abundance and 95 distribution of aerosols exist between global models. Bian et al. [2017] found that the atmospheric 96 burden of nitrate aerosols differ by a factor of 13 between the models in AeroCom Phase III, caused by differences in both chemical and deposition processes. A smaller, but still considerable, model 97 spread in the simulated burden of organic aerosols (OA) from 0.6-3.8 Tg was found by Tsigaridis 98 99 et al. [2014]. It was also shown that OA concentrations on average were underestimated. There has been particular focus on BC aerosols over recent years. Multi-model studies have shown 100

variations in global BC burden and lifetime up to a factor of 4-5 [Lee et al., 2013; Samset et al., 101 2014]. Previous comparisons of modeled BC distributions with observations have also pointed to 102 two distinct features common to many models: an overestimation of high altitude concentrations 103 at low- to mid-latitudes and discrepancies in the magnitude and seasonal cycle of high-latitude 104 105 surface concentrations (e.g., [Eckhardt et al., 2015; Lee et al., 2013; Samset et al., 2014; Schwarz et al., 2013]. As accurate representation of the observed aerosol distributions in global models is 106 crucial for confidence in estimates of radiative forcing (RF), these issues emphasize the need for 107 broad and up-to-date evaluation of model performance. 108

109 The diversity of simulated aerosol distributions, and discrepancies between models and measurements, stem from uncertainties in the model representation aerosol processing. Knowledge 110 of the factors that control the atmospheric distributions is therefore needed to identify potential 111 model improvements and need for further observational data, and to assess how remaining 112 uncertainties affect the modeled aerosol abundances and, in turn, estimates of RF and climate 113 impact. A number of recent studies have investigated the impact of changes in aging and 114 scavenging processes on the BC distribution, focusing on aging and wet scavenging processes (e.g., 115 [Bourgeois and Bey, 2011; Browse et al., 2012; Fan et al., 2012; Hodnebrog et al., 2014; Kipling 116 et al., 2013; Lund et al., 2017; Mahmood et al., 2016]), resulting in notable improvements, at least 117 118 for specific regions or observational data sets. However, with some notable exceptions [e.g., Kipling et al., 2016], few studies have focused on impacts of scavenging and other processes on a 119 broader set of aerosol species or the combined impact in terms of total aerosol optical depth (AOD). 120

Here we use the CEDS historical emission inventory as input to the chemistry-transport model 121 122 OsloCTM3 to quantify the change in atmospheric concentrations over the period of 1750 to 2014. The OsloCTM3 is an update of the OsloCTM2, and includes several key changes compared to its 123 predecessor. The significant existing model spread and sensitivity to process parameterizations 124 underlines the need for careful and updated documentation of new model versions, and the 125 increasing amount of available measurement data allows for improved evaluation. Before the 126 model is used to quantify historical time series, we therefore evaluate the simulated present-day 127 aerosol concentrations and optical depth against a range of observations. To get a first-order 128 overview of how uncertainties in key processes and parameters affect the atmospheric abundance 129 130 and distribution of aerosols in the OsloCTM3, we perform a range of sensitivity simulations. In addition to changes in the scavenging (solubility) assumptions, runs are performed with different 131 emission inventories, horizontal resolution, and meteorological data. The impact on individual 132 species and total AOD, as well as on the model performance compared with observations, is 133 investigated. Finally, we present updated estimates of the historical evolution of radiative forcing 134 due to aerosol-radiation interactions from pre-industrial to present, taking into account recent 135 literature on aerosol optical properties. Section 2 describes the model and methods, while results 136 are presented in Sect. 3 and discussed in Sect. 4. The conclusions are given in Sect. 5. 137

- 138
- 139 2 Methods
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141 2.1 OsloCTM3

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143 The OsloCTM3 is an offline global 3-dimensional chemistry-transport model driven by 3-hourly meteorological forecast data [Søvde et al., 2012]. The OsloCTM3 has evolved from its predecessor 144 145 OsloCTM2 and includes several updates to the convection, advection, photodissociation and 146 scavenging schemes. Compared with OsloCTM2, the OsloCTM3 has a faster transport scheme, an 147 improved wet scavenging scheme for large scale precipitation, updated photolysis rates and a new lightning parameterization. The main updates and subsequent effects on gas-phase chemistry were 148 described in detail in *Søvde et al.* [2012]. Here we document the aerosols in OsloCTM3, including 149 BC, primary and secondary organic aerosols (POA, SOA), sulfate, nitrate, dust and sea salt. The 150 aerosol modules in OsloCTM3 are generally inherited and updated from OsloCTM2. The 151 following paragraph briefly describes the parameterizations, including updates new to this work. 152

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The carbonaceous aerosol module was first introduced by Berntsen et al. [2006] and has later been 154 155 updated with snow deposition diagnostics [Skeie et al., 2011]. The module is a bulk scheme, with aerosols characterized by total mass and aging represented by transfer from hydrophobic to 156 hydrophilic mode at a constant rate. In the early model versions, this constant rate was given by a 157 global exponential decay of 1.15 days. More recently, latitudinal and seasonal variation in transfer 158 159 rates based on simulations with the microphysical aerosol parameterization M7 were included [Lund and Berntsen, 2012; Skeie et al., 2011]. Previous to this study, additional M7 simulations 160 have been performed to include a finer spatial and temporal resolution in these transfer rates. 161 Specifically, the latitudinal transfer rates have been established based on experiments with 10 162 instead of four emission source regions and with monthly, not seasonal resolution. In OsloCTM3 163 164 the carbonaceous aerosols from fossil fuel and biofuel combustion are treated separately, allowing 165 us to capture differences in optical properties in subsequent radiative transfer calculations (Sect. 2.4). In contrast to the OsloCTM2, OsloCTM3 treats POA instead of OC. If emissions are given 166 167 as OC, a factor of 1.6 for anthropogenic emissions and 2.6 for biomass burning sources is used for the OC-to-POA conversion, following suggestions from observational studies [Aiken et al., 2008; 168 Turpin and Lim, 2001]. Upon emission, 20% of BC is assumed to be hydrophilic and 80% 169 hydrophobic, while a 50/50 split is assumed for POA [*Cooke et al.*, 1999]. An additional update 170 in this work is the inclusion of marine primary organic aerosols following the methodology by 171 172 Gantt et al. [2015], where emissions are determined by production of sea spray aerosols and 173 oceanic chlorophyll A. Monthly concentrations of chlorophyll A from the same year as the meteorological data is taken from the Moderate Resolution Imaging Spectroradiometer (MODIS; 174 available from https://modis.gsfc.nasa.gov/data/dataprod/chlor_a.php), while sea spray aerosols 175 176 are simulated by the OsloCTM3 sea salt module. The climatological annual mean total emission of marine POA is scaled to 6.3 Tg based on *Gantt et al.* [2015]. The scaling factor depends on the

178 chosen sea salt production scheme (see below) and to some degree on the resolution; here we have

used a factor of 0.5.

The formation, transport and deposition of SOA are parameterized as described by Hoyle et al. 180 [2007]. A two product model (Hoffmann et al., 1997) is used to represent the oxidation products 181 of the precursor hydrocarbons and their aerosol forming properties. Precursor hydrocarbons which 182 are oxidized to form condensable species include both biogenic species such as terpenes and 183 isoprene, as well as species emitted predominantly by anthropogenic activities (toluene, m-xylene, 184 methylbenzene and other aromatics). The gas/aerosol partitioning of semi-volatile inorganic 185 aerosols is treated with a thermodynamic model [Myhre et al., 2006]. The chemical equilibrium 186 between inorganic species (ammonium, sodium, sulfate, nitrate and chlorine) is simulated with the 187 Equilibrium Simplified Aerosol model (EQSAM) [Metzger et al., 2002a; Metzger et al., 2002b]. 188 The aerosols are assumed to be metastable, internally mixed and obey thermodynamic gas/aerosol 189 equilibrium. Nitrate and ammonium aerosols are represented by a fine mode, associated with sulfur, 190 and a coarse mode associated with sea salt, and it is assumed that sulfate and sea salt do not interact 191 through chemical equilibrium [Myhre et al., 2006]. The sulfur cycle chemistry scheme and 192 aqueous-phase oxidation is described by Berglen et al. [2004]. 193

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The sea salt module originally introduced by Grini et al. [2002] has been updated with a new 195 production parameterization following recommendations by Witek et al. [2016]. Using satellite 196 retrievals, Witek et al. (2016) evaluated different sea spray aerosol emission parametrizations and 197 198 found the best agreement with the emission function from Sofiev et al. [2011] including the sea surface temperature adjustment from Jaeglé et al. [2011]. Compared to the previous scheme, the 199 global production of sea salt is reduced, while there is an increase in the tropics. This will have an 200 impact on the uptake of nitric acid in sea salt particles, consequently affecting NOx, hydroxide 201 202 (OH) and ozone levels. However, here we limit the scope to aerosols. The Dust Entrainment and 203 Deposition (DEAD) model v1.3 [Zender et al., 2003] was implemented into OsloCTM2 by Grini et al. [2005] and is also used in OsloCTM3. As a minor update, radiative flux calculations, required 204 for determination of boundary layer properties in the dust mobilization parameterization [Zender 205 206 et al., 2003], now uses radiative surface properties and soil moisture from the meteorological fields. 207

208 Aerosol removal includes dry deposition and washout by convective and large-scale rain. Rainfall is calculated based on European Center for Medium-Range Weather Forecast (ECMWF) data for 209 convective activity, cloud fraction and rain fall. The efficiency with which aerosols are scavenged 210 211 by the precipitation in a grid box is determined by a fixed fraction representing the fraction of this 212 box that is available for removal, while the rest is assumed to be hydrophobic. The parameterization distinguishes between large-scale precipitation in the ice and liquid phase, and 213 the OsloCTM3 has a more complex cloud model than OsloCTM2 that accounts for overlapping 214 215 clouds and rain based on Neu and Prather [2012]. When rain containing species falls into a grid box with drier air it will experience reversible evaporation. Ice scavenging, on the other hand, can 216

217 be either reversible or irreversible. For further details about large-scale removal we refer the reader to Neu and Prather [2012]. Convective scavenging is based on the Tiedtke mass flux scheme 218 (Tiedtke 1989) and is unchanged from the OsloCTM2. The solubility of aerosols is given by 219 constant fractions, given for each species and type of precipitation (i.e., large-scale rain, large-220 221 scale ice, and convective) (Table 2). Dry deposition rates are unchanged from OsloCTM2, but the OsloCTM3 includes a more detailed land use dataset (18 land surface categories at 1°x1° 222 horizontal resolution compared to 5 categories at T42 resolution), which affects the weighting of 223 deposition rates for different vegetation categories. Re-suspension of dry deposited aerosols is not 224 225 treated.

- 226
- 227 2.2 Emissions
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The baseline and historical simulations use the CEDS anthropogenic [Hoesly et al., 2018; Smith et 229 al., 2015] and biomass burning (BB4CMIP) [van Marle et al., 2017] emissions. The CEDS 230 231 inventory provide monthly gridded emissions of climate-relevant greenhouse gases, aerosols and precursor species from 1750 to 2014 using a consistent methodology over time. Anthropogenic 232 CEDS emissions are comparable to, but generally higher than, other existing inventories [Hoesly 233 234 et al., 2018]. Biogenic emissions are from the inventory developed with the Model of Emissions 235 of Gases and Aerosols from Nature under the Monitoring Atmospheric Composition and Climate project (MEGAN-MACC) [Sindelarova et al., 2014] and are held constant at the year 2010 level. 236 Here we use the CEDS version released in 2016 (hereafter CEDSv16). In May 2017, after 237 completion of our historical simulations, an updated version of the CEDS emission inventory was 238 239 released after users reported year-to-year inconsistencies in the country/sector level gridded data. The emission totals were not affected, but there were occasional shifts in the distribution within 240 countries (http://www.globalchange.umd.edu/ceds/ceds-cmip6-data/). The potential implications 241 for our simulations are discussed below. 242

- 243 Two other emission inventories are also used. The ECLIPSEv5 emission dataset was created with
- the Greenhouse Gas Air Pollution Interactions and Synergies (GAINS) model [Amann et al.,
- 245 2011] and provides emissions in 5 year intervals from 1990 to 2015, as well as projections to 2050
- [Klimont et al., 2017]. The 1990-2015 emission series was recently used to simulate changes in
- 247 aerosols and ozone and their RF [*Myhre et al.*, 2017]. Here we only use emissions for 2010 in the
- 248 sensitivity simulation.
- The Representative Concentration Pathways (RCPs) [*van Vuuren et al.*, 2011] were developed as a basis for near- and long-term climate modeling and were used in CMIP5 and Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP) experiments. While the four RCPs span a large range in year 2100 RF, emissions of most species have not diverged significantly in 2010 and we select the RCP4.5 for use here [*Thomson et al.*, 2011]. Table S1 summarized total global emissions of BC, OC, NO_x and SO₂ in 2010 in each of the three scenarios.
- In the simulations with the ECLIPSEv5 and RCP4.5 inventories, biomass burning emissions are from the Global Fire Emission Database Version 4 (GFED4) [*Randerson et al.*, 2017]. The

BB4CMIP emissions are constructed with GFED4 1997-2015 emissions as a basis [*van Marle et al.*, 2017] and emissions in 2010 are similar in both datasets. Hence, any difference between the

- sensitivity simulations stems from differences in the anthropogenic inventory.
- 260
- 261
- 262 2.3 Simulations
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Time slice simulations with CEDSv16 emissions for 1750, 1850 and from 1900 to 2014 are performed (every ten years from 1900-1980, thereafter every five years), one year with six months spin-up. The model is run with <u>fixed</u> year 2010 meteorological data and a horizontal resolution of 2.25x2.25 degrees (denoted 2x2), with 60 vertical layers. While *Søvde et al.* [2012] used meteorological data from the ECMWF IFS model cycle 36r1, we apply here meteorology from the ECMWF OpenIFS cycle 38r1 (<u>https://software.ecmwf.int/wiki/display/OIFS/</u>).

270

Additional model runs are performed to investigate the importance of differences in key processes 271 272 for the aerosol distributions and model performance (Table 1). In addition to the CEDSv16 emissions, the model is run with ECLIPSEv5 and RCP4.5 emission inventories for anthropogenic 273 emissions and GFEDv4 biomass burning emissions. Additionally, we perform simulations with 274 1.125x1.125 degrees (denoted 1x1) horizontal resolution. To investigate the importance of 275 276 meteorology, the simulation with CEDSv16 emissions is repeated with meteorological data for 277 year 2000 instead of 2010. Year 2000 is selected due to its opposite El Niño-Southern Oscillation (ENSO) index compared to 2010. Finally, three model runs are performed with increased and 278 decreased aerosol removal by large-scale ice clouds and decreased aerosol scavenging by liquid 279 (large-scale and convective) precipitation. To modify the scavenging, we tune the fixed fractions 280 that control aerosol removal efficiency in the model (see Sect. 2.1). Table 2 summarizes fractions 281 used in the baseline configuration and the three sensitivity tests. A decrease and increase in 282 283 efficiency of 0.2 is adopted for scavenging of all aerosols by liquid clouds (except hydrophobic BC and POA) and ice clouds, respectively. Note that there is no test with increased removal by 284 liquid clouds, as, with the exception of hydrophobic BC, POA and SOA, 100% efficiency is 285 already assumed. For ice clouds we also reduce the efficiency to a fraction of 0.1, or 0.001 if the 286 value is 0.1 in the baseline configuration. We note that these changes do not represent realistic 287 uncertainty ranges based on experimental or observational evidence, as there are limited 288 constraints in the literature, but are chosen to explore the impact of a spread in the efficiency with 289 which aerosols act as ice and cloud condensation nuclei. 290

291

292 2.4 Radiative transfer

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We calculate the <u>instantaneous top-of-the atmosphere</u> radiative forcing of anthropogenic aerosols due to aerosol-radiation interactions (RFari) [*Myhre et al.*, 2013b]). The radiative transfer 296 calculations are performed offline with a multi-stream model using the discrete ordinate method 297 [Stamnes et al., 1988]. The model includes gas absorption, Rayleigh scattering, absorption and scattering by aerosols, and scattering by clouds. The RFari of individual aerosols is obtained by 298 separate simulations, where the concentration of the respective species is set to the pre-industrial 299 300 level. The aerosol optical properties have been updated from earlier calculations using this radiative transfer model [Myhre et al., 2007; Myhre et al., 2009], in particular those associated 301 with aerosol absorption. The Bond and Bergstrom [2006] recommendation of a mass absorption 302 coefficient (MAC) for BC of around 7.5 m² g⁻¹ for freshly emitted BC and an enhancement factor 303 of 1.5 for aged BC was used previously. In the present analysis, we apply a parametrization of 304 MAC from observations over Europe by Zanatta et al. [2016], where MAC depends on the ratio 305 of non-BC to BC abundance. The mean MAC of BC from these observations around 10 m² g⁻¹ at 306 630 nm [Zanatta et al., 2016]. The measurements in Zanatta et al. [2016] represent continental 307 European levels. For very low concentrations of BC, the formula given in Zanatta et al. [2016] 308 309 provides very high MAC values. We have therefore set a minimum level of BC of 1.0e-10 g m⁻³ for using this parameterization, and for lower concentrations we use Bond and Bergstrom [2006]. 310 In addition, we have set a maximum value of MAC of 15 $m^2 g^{-1}$ (637 nm) to avoid unrealistic high 311 values of MAC compared to observed values. Organic matter has a large variation in the degree 312 313 of absorption [e.g., Kirchstetter et al., 2004; Xie et al., 2017], from almost no absorption to a strong absorption in the ultraviolet region. Here, we have implemented absorbing organic matter 314 according to refractive indices from Kirchstetter et al. [2004]. The degree of absorption varies by 315 source and region and is at present inadequate quantified: Here we assume 1/3 of the biofuel 316 organic matter and ¹/₂ of the SOA from anthropogenic volatile organic carbon (VOC) precursors. 317 318 The remaining fractions of biofuel, fossil fuel and marine POA and SOA (anthropogenic and all natural VOCs) are assumed to be purely scattering organic matter. As these fractions are not 319 sufficiently constrained by observational data and associated with significant uncertainty, we also 320 perform calculations with no absorption by organic matter for comparison. 321 322

323 2.5 Observations

324

A range of observational datasets are used to evaluate the model performance in the baseline simulation. Note that we use the term "black carbon" in a qualitative manner throughout the manuscript to refer to light-absorbing carbonaceous aerosols. However, when comparing with measurements, we use either elemental carbon (EC) or refractive BC (rBC), depending on whether the data is derived from methods specific to the carbon content of carbonaceous aerosols or incandescence methods, in line with recommendations from *Petzold et al.* [2013].

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332 Measured surface concentrations of EC, OC, sulfate and nitrate are obtained from various networks.

333 For the US, measurements from IMPROVE (Interagency Monitoring of Protected Visual

Environments) and CASTNET (Clean Air Status and Trends Network) are used. For Europe, data

from EMEP (European Monitoring and Evaluation Programme) [Tørseth et al., 2012] and

ACTRIS (Aerosols, Clouds and Trace gases Research InfraStructure) [Cavalli et al., 2016; Putaud 336 et al., 2010] is used. EMEP and ACTRIS sites are all regional background sites, representative for 337 a larger area. To broaden the geographical coverage we also compare the model output against 338 additional observations from the CMA Atmospheric Watch Network (CAWNET) in China [Zhang 339 et al., 2012] and those reported in the literature from India (see Kumar et al. [2015] for more 340 341 details). CASTNET, IMPROVE, EMEP and ACTRIS data is from year 2010, while CAWNET observations were sampled in 2006-2007 and the observational data base from India compiled by 342 Kumar et al. [2015] cover a range of years. IMPROVE provides mass of aerosols using filter 343 analysis of measurements of particulate matter with diameter of less than 2.5 micrometers (PM_{2.5}), 344 while CASTNET uses an open-face filter pack system with no size restriction to measure 345 concentrations of atmospheric sulfur and nitrogen species [Lavery et al., 2009]. Mass of individual 346 species from the CAWNET network is obtained from aerosol chemical composition analysis 347 performed on PM₁₀ samples [Zhang et al., 2012]. EMEP and ACTRIS measurements of EC and 348 349 OC are in the $PM_{2.5}$ range, whereas nitrate and sulfate measurements are filter-based with no size 350 cutoff limit. Data resulting from EMEP and ACTRIS are archived in the EBAS data base ((http://ebas.nilu.no) at NILU - Norwegian Institute for Air Research, and are openly available (see 351 also Data availability). 352

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354 Modeled AOD is evaluated against the Aerosol Robotics Network (AERONET). AERONET is a global network of stations measuring radiance at a range of wavelengths with ground-based sun-355 photometers, from which aerosol column burden and optical properties can be retrieved [Dubovik 356 and King, 2000; Holben et al., 1998]. The comparison with AERONET data was done using the 357 358 validation tools available from the AeroCom data base hosted by Met Norway (http://aerocom.met.no/data.html). We also compare against AOD retrievals from MODIS-Aqua 359 and Terra (level 3 atmosphere products, AOD550 combined dark target and deep blue, product 360 version 6) [MOD08, 2018] and the Multi-angle Imaging SpectroRadiometer (MISR) (level 2 361 362 aerosol product, product version 4) [MISR, 2018].

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Figure S1 depicts the locations of all the stations. For comparison with surface concentrations and 364 AERONET AOD, the model data is linearly interpolated to the location of each station using 365 annual mean, monthly mean (concentrations) or 3-hourly output (AOD), depending on the 366 367 resolution of the observations. In the case of AERONET, high mountain stations (here defined as 368 having an elevation higher than 1000 meter above sea level) are excluded following Kinne et al. [2013]. For comparison with observed OC surface concentrations, modeled OA is converted to 369 OC using factor of 1.6 for POA and 1.8 for SOA. Unless measurements are restricted to the PM2.5 370 size range, the comparison includes both fine and coarse mode modeled nitrate (Sect. 2.1). Several 371 statistical metrics are used to assess the model skill, including correlation coefficient (R), root 372 mean square error (RMSE), variance and normalized mean bias (NMB). 373

The modeled vertical distribution of BC is compared with aircraft measurements of refractory BC (rBC) from the HIAPER Pole-to-Pole Observations (HIPPO) campaign [*Wofsy et al.*, 2011].

Vertical profiles of BC from OsloCTM2 have been evaluated in several previous studies (e.g., *Samset et al.* [2014]) and a more thorough comparison of OsloCTM3 results against a broader set
of campaigns is provided by *Lund et al.* [2018]. In the present analysis we focus on data from the
third phase (HIPPO3) flights, the only phase that was conducted in 2010, i.e., the same year as our

sensitivity simulations. Model data is extracted along the flight track using an online flight
 simulator. The data is separated into five latitude regions and vertical profiles constructed by
 averaging observations and model output in 13 altitude bins.

383

384 3 Results

385

We first document the aerosol distributions simulated in the baseline model configuration, focusing on the anthropogenic contribution, and compare with observations, multi-model studies and results from the sensitivity tests. With the present-day model performance evaluated, we then present the updated historical development of RFari of anthropogenic aerosols.

390

391 3.1 Evaluation of present-day aerosol distributions

392

The global mean aerosol burdens and atmospheric residence times (ratio of burden to total wet 393 plus dry deposition) in the baseline simulation are summarized in Table 3 (top row), with spatial 394 395 distribution shown in Fig. S2. Compared to results from the AeroCom III experiment, the OsloCTM3 sulfate burden of 5.4 mg m⁻² estimated here is about 50% higher than the multi-model 396 mean of 3.5 mg m⁻² and 35% higher than OsloCTM2 [Bian et al., 2017]. While the total SO₂ 397 emission is only 5% higher in the present study than in the OsloCTM2 AeroCom III simulation, 398 399 the atmospheric residence time of sulfate is 50% longer, suggesting that the burden difference is mainly attributable to changes in the parameterization of dry and large-scale wet deposition in 400 OsloCTM3 (Sect. 2.1). The nitrate burden is nearly a factor three higher than both the AeroCom 401 multi-model mean and OsloCTM2 burden, and higher than all nine models contributing in 402 AeroCom III [Bian et al., 2017]. This is mainly due to a higher burden of coarse mode nitrate 403 404 aerosols, associated with less efficient scavenging of sea salt in OsloCTM3 than OsloCTM2. The global budgets of OA simulated by the AeroCom II models was analyzed by Tsigaridis et al. 405 [2014]. The burden of OA in the OsloCTM3 of 3.4 mg m⁻² is close to their multi-model mean of 406 3.1 mg m⁻² and 25% higher than the OsloCTM2. The OsloCTM3 estimate includes the contribution 407 408 from marine OA emissions (Sect. 2.1), which may explain part of the difference as marine OA was included in some of the AeroCom II models, but not OsloCTM2. However, the marine POA only 409 contributes around 3% to the total OA. Additionally, the residence time of OA of 5.3 days is longer 410 than in the OsloCTM2 AeroCom II experiment. The global BC burden of 0.23 mg m⁻² is also close 411 to the mean of the AeroCom II models of 0.25 mg m⁻² [Samset et al., 2014]. We note that different 412 emission inventories were used in the AeroCom experiments and the present analysis, however, 413

the comparison shows that the aerosol burdens simulated by OsloCTM3 fall within the range ofexisting estimates from global models.

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417 Figure 1 shows results from the baseline OsloCTM3 simulation against annual mean measured 418 surface concentrations of EC, OC, sulfate and nitrate in Europe, North America and Asia. Overall, the OsloCTM3 shows a high correlation of 0.8-0.9 with measured surface concentrations. There 419 is a general tendency of underestimation by the model, with the lowest NMB and RMSE for BC 420 and nitrate (-23%) and the highest for sulfate (-52%). There are, however, notable differences in 421 model performance between data sets in different regions, as seen from Table S2. For all species, 422 the NMB and RMSE are highest for measurements in China. For instance, excluding the 423 CAWNET measurements, reduces the NMB for sulfate in Fig. 1 from -52% to -31% (not shown). 424 In contrast, the correlation with CAWNET observations is generally similar to, or higher than, 425 other regions/networks. In the case of BC and nitrate, the model slightly overestimates 426 427 concentrations in Europe and North America, but underestimates Asian measurements. The best 428 overall agreement is generally with IMPROVE observations in North America. Differences in instrumentation between different networks can affect the evaluation. Lavery et al. [2009] found 429 that measurements from CASTNET typically gave higher nitrate surface concentrations than 430 values obtained from co-located IMPROVE stations, which could partly explain the NMB of 431 opposite sign in these two networks in Table S2. For BC, we also include measurements from 432 across India compiled by Kumar et al. [2015]. This is a region where emissions have increased 433 strongly, but where evaluation of the model performance so far has been limited due to availability 434 of observations. The model underestimates concentrations with a NMB of -43%, however, the 435 436 correlation of 0.60 is similar to the comparison with data from China and higher than the other regions. An examination of the monthly concentrations (Fig. S3) shows that the largest 437 discrepancies occur during winter, with the largest bias found for measurements in North East 438 India. One possible reason could be missing or underestimated emission sources. This finding is 439 440 similar to the comparison of measurements against WRF-chem by Kumar et al. [2015]. The seasonality of BC concentrations has also been an issue at high northern latitudes, where earlier 441 versions of the OsloCTM strongly underestimated winter and springtime surface concentrations at 442 Arctic stations [Lund et al., 2017; Skeie et al., 2011], similar to many other models [Eckhardt et 443 444 al., 2015]. This Arctic underestimation persists in the current version of the model. Seasonal differences exist also in other regions, but not consistently across measurement networks. 445 Compared with EC measurements from EMEP/ACTRIS the correlation is poorer during winter 446 and spring, and the model underestimate concentrations in contrast to a positive NMB in summer 447 448 and fall. However, due to the relatively low number of stations, these values are sensitive to a few stations with larger measurement-model discrepancies. For both IMPROVE and EMEP/ACTRIS, 449 the model underestimation of sulfate is larger during summer and fall, but with opposite seasonal 450 differences in correlation. In general, the number of stations and evaluation of data from only one 451 452 year limits the analysis of seasonal variations.

We do not evaluate ammonium concentrations in the present analysis, as that requires a detailed 453 discussion of the nitrate and sulfate budgets, which has been covered by the recent multi-model 454 evaluation by Bian et al. [2017] based on an AeroCom Phase III experiment, in which the 455 OsloCTM3 participated. Results showed that most models tend to underestimate ammonium 456 457 concentrations compared to observations in North America, Europe and East Asia, with a multimodel mean bias and correlation of 0.886 and 0.47, respectively. The OsloCTM3 shows good 458 agreement with ammonium measurements in North America, but has a bias and correlation close 459 the model average in the other two regions. 460

461 In May 2017, after completion of our historical simulations, an updated version of the CEDS emission inventory was released after an error in the code was reported (see Sect. 2.2). This 462 resulted in occasional shifts in the spatial distribution of emissions within countries with large 463 spatial extent (e.g., USA and China). Since the emission totals were not affected, the impact on 464 our RFari estimates is likely to be small, but shifts in the emission distribution could influence the 465 model evaluation, in particular for surface concentrations. While repeating all simulations would 466 require more resources, we have repeated the year 2010 and 1750 runs. Figure S4 shows the 467 comparison of modeled concentrations against IMPROVE measurements with the two emission 468 inventory versions, CEDSv16 and CEDSv17. In the case of BC, the comparison shows a 5% higher 469 470 correlation and 15% lower RMSE with the CEDSv17 than CEDSv16. A similar improvement is found for nitrate, with 26% higher correlation and 12% lower RMSE, while in the case of OC and 471 sulfate, the difference is small (< 5%). Smaller differences of between 2-10% are also found in the 472 comparison against measurements in Europe and Asia (not shown). Hence, using the updated 473 474 version of the emission inventory has an effect on the model performance in terms of surface concentrations, but without changing the overall features or conclusions. The net RFari in 2010 475 relative to 1750 is 2% stronger with the CEDSv17 inventory, a combined effect of slightly higher 476 global BC burden and lower burdens of sulfate and OA. 477

As shown in Table S2, the model overestimate surface concentrations in some regions and 478 underestimate them in others. Compensating biases can influence the evaluation of total AOD. 479 Moreover, the biases differ in magnitude between different species. Moving one step further, we 480 therefore examine the average aerosol composition in the three regions where this is possible with 481 482 our available measurements. Figure 2 shows the relative contribution from different aerosols 483 species to the total mass in the IMPROVE, EMEP, ACTRIS and CAWNET measurements and the corresponding model results. The number of available aerosol species varies between the 484 measurement networks and we include sea salt from IMPROVE and ammonium from CAWNET. 485 486 Additionally, the number of stations where simultaneous measurements of all species were available also differ substantially, with 16 for CAWNET, 5 for EMEP/ACTRIS and 172 for 487 IMPROVE. Overall, the relative composition is well represented by the model. The agreement is 488 particularly good for the IMPROVE network. Compared to measurements from CAWNET, the 489 490 model has a lower relative contribution from OC and more sulfate. In the case of Europe, nitrate 491 aerosols also constitute a significantly larger fraction in the model than in the observations. The

evaluation of nitrate is complicated by possible differences in the detection range of 492 instrumentation compared to the size of the two nitrate modes in the model (Sect. 2.1). The 493 comparison against EMEP nitrate data includes both coarse and fine mode modeled nitrate. 494 Excluding the coarse mode, the fraction of total mass attributable to nitrate decreases from 43% to 495 496 28%, which is much closer to the observed 30% contribution. However, this affects the comparison in Figure 1, resulting in a negative NMB of -34%, compared to -23% when including both coarse 497 and fine mode. This suggest that part, but not all, of the nitrate represented as a coarse mode in the 498 model is measured by the instrument, pointing to a need for a more sophisticated size distribution 499 in the model to make better use of available observations. The low number of available stations 500 501 from EMEP/ACTRIS could also an important factor.

Next, we examine total AOD. Figure 3 shows modeled AOD and aerosol absorption optical depth 502 (AAOD), AOD retrieved from MODIS-Aqua and comparison of modeled AOD with AERONET 503 504 observations. Modeled global, annual mean AOD and AAOD is 0.13 (Fig. 3a) and 0.0051 (Fig. 3b), respectively. The overall spatial pattern of modeled AOD agrees well with MODIS (Fig. 3c), 505 however, the latter gives a higher global mean of 0.16 and clearly higher values in North India and 506 parts of China, as well as Central Africa. These peak values are similar to MODIS-Terra, but less 507 pronounced in the AOD retrieved from MISR (Fig. S5), illustrating important differences between 508 509 different remote sensing products. Nevertheless, an underestimation of modeled AOD in Asia is consistent with results from the evaluation of surface concentrations and can also be seen in the 510 comparison against AERONET, as discussed below. The OsloCTM3 shows a good agreement 511 with measured AOD from the AERONET network, with an overall correlation of 0.82 and RMSE 512 513 of 0.11, when using monthly mean data from 266 stations (Fig. 3d). Note that the modeled global mean AOD is 0.13, but the model mean at the AERONET stations is 0.175 (Fig 3d) and has only 514 a NMB of -11.8%. Many of the AERONET stations tend not to be regional background sites, but 515 can be influenced by local pollution (e.g., *Wang et al.* [2018]) 516

However, there are notable regional differences in model performance. Fig. S6 compares modeled 517 AOD against AERONET stations in Europe, North America, India and China separately. The best 518 agreement is found for Europe and North America, with NMB of -0.4% and -13%, respectively, 519 and RMSE of approx. 0.05. The correlation is higher for North America (0.71) than Europe (0.63). 520 521 A relatively high correlation of 0.71 is also found for stations in China. However, the NMB and 522 RMSE is higher (-34.5% and 0.25). There are significantly fewer observations for comparison with modeled AOD over India, but the ones available give NMB and RMSE on the same order of 523 magnitude as for China, but a lower correlation (0.45). 524

- 525 Ground-based measurements can also provide information about column absorption aerosol
- 526 optical depth (AAOD). Such information has been used to constrain the absorption of BC and
- provide top-down estimate of the direct BC RF (e.g., [*Bond et al.*, 2013]). However, retrieval and
- application of AERONET AAOD is associated with a number of challenges and uncertainties (e.g.,
- 529 [*Samset et al.*, 2018]), hence such an evaluation is not performed here.

530 Recent literature has pointed to important representativeness errors arising when constraining models using observations due to the coarse spatial and temporal scales of global models compared 531 with the heterogeneity of observations. Schutgens et al. [2016a] found differences in RMSE of up 532 to 100% for aerosol optical thickness when aggregating high resolution model output over grid 533 534 boxes representative of the resolution of current global models compared to small areas corresponding to satellite pixels. Smaller, but notable, differences of up to 20% were found when 535 monthly mean model data was used, as in the present analysis. However, that did not account for 536 issues related to temporal collocation, which can also introduce considerable errors [Schutgens et 537 al., 2016b]. In a recent study, Wang et al. [2018] found a spatial representativeness error of 30% 538 539 when constraining AAOD modeled at a 2°x2° horizontal resolution against AERONET retrievals. However, further work is needed to investigate whether similar biases exist for AOD. 540

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542 3.2 Sensitivity of aerosols distributions to model input and process parameterization

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As shown in the section above, the OsloCTM3 performs well compared against observed AOD. Still, a number of factors influence the simulated distributions of individual aerosol species. To assess the importance of key uncertainties for modeled distributions and model performance, we perform a range of sensitivity simulations (Table 1) to examine the importance of emission inventory, scavenging assumptions (Table 2), meteorological data and resolution for the modeled aerosol distributions and model performance.

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551 Global aerosol burdens and AOD in each sensitivity run are summarized in Table 3 (corresponding 552 atmospheric residence times are given in Table S3). The BC burden is particularly sensitive to reduced scavenging by large-scale ice clouds (LSIDEC), resulting in a 40% higher burden 553 compared to the baseline. In contrast, an equal increase in the scavenging efficiency (LSIINC) 554 result in a decrease in burden of only 9%, while decreased scavenging by liquid precipitation 555 (SOLDEC) gives a 13% higher burden. The lower BC emissions in the ECLv5 and CMIP5 556 557 inventories give a global BC burden that is 9 and 22% lower, respectively. For sulfate, ammonium 558 and OA, we also find the largest burden changes in the LSIDEC case, followed by SOLDEC. The change in the LSIDEC is particularly large for OA and is driven by changes in SOA. For SOA, the 559 560 changes are determined not only by modifying the scavenging, but also by changes in POA 561 concentrations, which gas-phase secondary organics can partition onto. Increasing the horizontal resolution results in a slightly higher burden for all species, except sea salt. 562

563 While sensitivity tests may give similar changes in the total global burdens, the spatial distribution 564 of changes can differ substantially. Figure 4 shows the ratio of AOD and total burden by species 565 and altitude in each sensitivity simulation to the baseline. As expected, varying the emission 566 inventories results in changes that are largely confined to the main source regions (Figs.4a,b). 567 Using the CMIP5 inventory results in considerably lower concentrations over Asia, the Middle 568 East and North Africa, reflecting the higher emissions in the more recent inventory. Over central 569 North America the AOD is higher, mainly due to more ammonium nitrate, whereas the higher AOD over Eastern Europe and part of Russia is a result of higher sulfate concentrations. Similar 570 characteristics are found when using ECLv5, but the relative differences are smaller. Reducing or 571 increasing the large-scale ice cloud scavenging gives the largest relative changes in AOD at high 572 573 latitudes, while changes in the solubility assumption for liquid precipitation affects AOD most over Asia, where aerosol burdens are high, and around the equator where convective activity is 574 strong. In general, the burden of BC, OA and dust is significantly affected by changes in the 575 scavenging assumptions, while nitrate responds more strongly to different emission inventories, 576 likely due to the complicated dependence on emissions of several precursors and competition with 577 ammonium-sulfate. We also note that at higher altitudes the absolute differences in the burden of 578 nitrate are small. Changes in AOD resulting from using different meteorological input data are 579 more heterogeneous and are most notable in regions where effects of choosing data from years 580 581 with opposite ENSO phase are expected, e.g., west coast of South America and South East Asia. 582 There is also a notable change in the Atlantic Ocean, where mineral dust is a dominating species. The meteorological data can affect production, deposition and transport of dust directly, as well as 583 indirectly through ENSO-induced teleconnections as suggested by e.g., Parhi et al. [2016]. 584

For BC, OA and dust, the largest impact relative to the baseline are seen above 600 hPa in the 585 586 LSIDEC case. Change in LSIDEC are also important in the case of sulfate and sea salt, but occur at lower altitudes. In contrast to the other aerosol species, differences in emission inventories are 587 most important for nitrate. In a recent study, Kipling et al. [2016] investigated factors controlling 588 the vertical distribution of aerosols in the HadGEM3-UKCA. It was found that in-cloud 589 590 scavenging was very important in controlling the vertical mass concentration of all species, except dust. For dust, it was also found that dry deposition and sub-cloud processes played key roles, 591 processes not examined in the present analysis. Moreover, Kipling et al. [2016] performed 592 sensitivity simulations by switching transport and scavenging on and off to get the full effect of a 593 594 given process, while we perform smaller perturbations to investigate uncertainties. Here we find significant impacts of changes in ice-cloud removal efficiency (Table 2) on the vertical distribution 595 of BC, OA and dust, while large-scale liquid and convective precipitation is more important for 596 sea salt and nitrate 597

598 Our sensitivity tests show that changes in input data, resolution or scavenging can lead to notable changes in the aerosol distributions. The next question is then how these changes affect model 599 performance compared to observations. Figure 5a compares modeled and measured surface 600 concentrations of BC, OC, sulfate and nitrate in each simulation using all observations in Fig. 1. 601 602 For BC, the sensitivity tests have little or no impact on correlation, but there is a markedly better 603 agreement in terms of standard deviation (i.e., model becomes closer to observations) for CEDSv16/CMIP6 compared to RCP/CMIP5, reflecting the higher emissions in the former. Similar, 604 but smaller, effects are also found for the other species. The improvement from RCP/CMIP5 to 605 606 CEDSv16/CMIP6 is especially seen for measurements in Asia. A higher resolution is also found 607 to reduce the bias, in particular for BC. Figure 5b shows the comparison against AERONET AOD

in each sensitivity simulation. Again, there is a higher correlation and lower bias in the 1x1RES
run than in the baseline, while the opposite is found in the RCP/CMIP5 and ECLv5 cases. For both
observables, the improvement in the 1x1RES simulation may result from a better sampling at a
finer resolution, improved spatial distribution or a combination. The most pronounced changes
results from using meteorological data from year 2000, in which case the correlation is reduced
from around 0.8 to 0.7.

For both observables, the difference in model performance between the baseline and scavenging 614 sensitivity tests is small. This may partly be an effect of the geographical coverage of stations; the 615 616 majority of measurements are from stations in more urban regions, whereas simulated burden changes occur in remote regions, particularly at high latitudes and altitudes (Fig. 4). We therefore 617 also perform evaluations against AOD from regional sub-sets of AERONET stations. Ten of the 618 AERONET stations used in the present analysis are located north of 65°N (Fig. S1). A comparison 619 of monthly mean simulated AOD in each of the sensitivity runs against observations in this region 620 shows the best agreement with the baseline simulation and with the ECLv5 emission inventory, 621 with a considerably higher bias when scavenging parameters are modified (Fig. S7a). This is 622 particularly the case in the LSIDEC run, where concentrations of all species increase at high 623 latitudes compared to the baseline (Fig. 4). In contrast, the reduced concentrations in LSIINC, 624 625 results in a negative bias. We note that most of these stations have missing values in the winter months, which is when the model underestimate BC concentrations in the Arctic, hence limiting 626 the evaluation. Decreased scavenging efficiency also leads to a higher bias than in the baseline for 627 observations in Europe and North America (not shown). In Asia, where the model already 628 629 underestimates aerosols in the baseline configuration, the bias is reduced since concentrations increase. However, differences are smaller than north of 65°N. Moreover, given the notable 630 exacerbation in model performance in other regions, it is likely that other sources of uncertainty 631 (e.g., emissions) are more important for the model-measurement discrepancies in Asia. A similar 632 633 comparison is performed for 15 AERONET stations located in North Africa and the Middle East 634 (Fig. S7b), where the dust influence is strong. Changing the meteorological year and reducing scavenging results in higher dust burdens (Table 3). Again, the agreement is better in the baseline 635 run than in these sensitivity runs. In particular, the METDATA run result in a higher bias and a 636 637 lower correlation, which is not surprising as dust production depends also on meteorological conditions. The changes compared to the baseline CEDSv16/CMIP6 simulation cannot be entirely 638 attributed to differences in dust concentrations, as seen from the RCP/CMIP5 and ECLv5 639 640 simulations where the dust production is equal to the baseline. Several studies have pointed to the importance of spatial resolution for improved model performance compared to observations (e.g., 641 642 [Sato et al., 2016; Schutgens et al., 2017; Schutgens et al., 2016a; Wang et al., 2016]). Wang et al. 643 [2016] found significant reductions in NMB of BC AAOD relative to AERONET when using a high resolution (10 km) emission data and model output. In our analysis, moving from 2°x2° to 644 1°x1° horizontal resolution also results in a slightly higher correlation and reduced bias and errors 645 when compared to all AERONET stations (Fig. 5b). The impact is largest for AOD in China and 646 India, the NMB is reduced (from -34% and -24% (Fig. S6) to -20% and -10%, respectively). 647

However, the opposite effect is found for AERONET stations in Europe and North America. Of course, the $1^{\circ}x1^{\circ}$ resolution is still very coarse compared to the grid sizes used in the abovementioned studies.

651 Changes away from near-source areas are also evaluated in terms of BC concentrations by a comparison with observed vertical distribution from the HIPPO3 campaign, where remote, marine 652 air over the Pacific was sampled across all latitudes (Sect. 2.5). To limit the number of model runs, 653 we focus on only one phase of the HIPPO campaign here, but a more comprehensive evaluation 654 of OsloCTM3 vertical BC distribution against aircraft measurements was performed by Lund et 655 al. [2018]. Figure 6 shows observed average vertical BC concentration profiles against model 656 657 results from each sensitivity test. The OsloCTM3 reproduces the vertical distribution well in low and mid-latitudes over the Pacific in its baseline configuration, although near-surface 658 concentrations in the tropics are underestimated. This is a significant improvement over the 659 OsloCTM2, where high-altitude concentrations in these regions typically were overestimated. The 660 baseline configuration of OsloCTM3 includes updates to the scavenging assumptions based on 661 previous studies investigating reasons for the high-altitude discrepancies (e.g., [Hodnebrog et al., 662 2014; Lund et al., 2017]. At high northern and southern latitudes, the model underestimates the 663 observed vertical profiles in the baseline. Increasing the model resolution does not have any 664 665 notable impact on the vertical profiles. There is a notable increase in high-latitude concentrations when large-scale ice cloud scavenging is decreased. However, there is a simultaneous exacerbation 666 of model performance in the other latitude bands, pointing to potential tradeoffs when tuning 667 global parameters, as also illustrated by Lund et al. [2017]. Due to the significant altitude 668 669 dependence of the radiative effect of BC (e.g., [Samset et al., 2013]), high altitude overestimations will contribute to uncertainties in BC RFari. We also note that HIPPO3 was conducted in 670 671 March/April: Comparison with aircraft measurements from other seasons show a smaller underestimation at high latitudes [Lund et al., 2018]. 672

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674 3.3 Pre-industrial to present-day aerosols

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With confidence in the model ability to reasonably represent current aerosol distributions 676 established, we next present an updated historical evolution of anthropogenic aerosols from pre-677 industrial to present-day, and the consequent direct radiative effect (RFari) (Sect. 2.4). Figure 7 678 679 shows the net change in total aerosol load from 1750 to 2014. Full times series by species are given in Table S4. To keep in line with the terminology used in the IPCC AR5, we now separate out 680 biomass burning BC and POA in a separate species "biomass". We also note that only the fine 681 mode fraction of nitrate contributes to the RFari and is inlcuded in Fig. 7. To illustrate the 682 contributions from additional emissions during the past 14 years, we also include the 2000-1750 683 684 difference. The values from the present study are also compared with results from the AeroCom II models [Myhre et al., 2013a], where emissions over the period 1850 to 2000 from Lamarque et 685 686 al. [2010] were used.

The most notable difference compared to the AeroCom II results is seen for biomass aerosols. 688 Biomass burning emissions have high interannual variability and this affects the analysis. While 689 the 1750-2014 difference is 0.23 mg m^{-2} , taking the difference between year 1750 and 2000 (black 690 asterisk) results in a net change of only 0.03 mg m⁻². There is also a much larger change in the 691 burden of biomass aerosols in the AeroCom experiments, reflecting a more than 100% higher 692 emissions in 2000 compared to 1850 Lamarque et al. [2010] inventory. However, biomass aerosols 693 comprises both scattering OA and absorbing BC and, as seen below, these nearly cancel in terms 694 of RFari. Changes in sulfate and OA from pre-industrial to 2000 are slightly higher in the present 695 696 analysis than in AeroCom II, and the influence of additional emissions since 2000 is seen. The OsloCTM3 is well below the AeroCom multi-model mean for nitrate. The OsloCTM2 was found 697 to be in the low range, but the multi-model was also influenced by some models giving high 698 699 estimates [Myhre et al., 2013a]

Using the CEDSv16 emissions, we estimate a net RFari from all anthropogenic aerosols in 2014 relative to 1750 of -0.17 W m⁻². The RFari from sulfate is -0.30 W m⁻², while the contributions from OA (combined fossil fuel plus biofuel POA and SOA), nitrate and biomass aerosols are smaller in magnitude of -0.09, -0.02 and -0.0004 W m⁻², respectively. The RFari due to fossil fuel and biofuel BC over the period is 0.31 W m⁻².

Figure 8a shows the time series of RFari by component, as well as the net, in the present analysis 705 (solid lines), and corresponding results reported in the IPCC AR5 (dashed lines). The net RFari 706 over time is mainly determined by the relative importance of compensating BC and sulfate RFari. 707 708 The most rapid increase in BC RFari is seen between 1950 and 1990, as emissions in Asia started to grow, outweighing reductions in North America and Europe [Hoesly et al., 2018]. After a period 709 of little change between 1990 and 2000, the rate of change increases again over the past two 710 decades, following strong emission increases in Asia and South Africa. Similarly, cooling 711 712 contribution from sulfate aerosols strenghtened from around mid-century. However, in contrast to BC, the evolution is fairly flat after 1990. The last 20 years has seen a continuous reduction in 713 sulfur dioxide (SO₂) emissions in Europe, from around 30 to 5 Tg yr⁻¹ in CEDSv16, with a similar 714 trend in North America. While emissions in China continue to increase well into the 2000s, a 715 716 stabilization is seen after 2010, following introduction of stricter emission limits as part of a program to desulfurize power plants [Klimont et al., 2013]. During the same period, emissions in 717 India have risen. However, the net global SO_2 emission trend over the past few years is a slight 718 decline [*Hoesly et al.*, 2018]. This development is reflected in the net RFari, which reaches its peak 719 720 (i.e., strongest negative value) around 1990 and gradually becomes weaker thereafter. This trend 721 is more pronounced in the present analysis that in the IPCC AR5 estimates, where the forcing due to sulfate is more flat in recent decades, suggesting that projected emission estimates 722 underestimated recent decreases in SO₂. The minimum net RFari value is also reached later in the 723 724 latter. Moreover, a recent study suggests that current inventories underestimate the decline in 725 Chinese SO₂ emissions and estimate a 75% reduction since 2007 [Li et al., 2017]. In this case, the

weakening trend could be even stronger than estimated here. The insert in Fig. 8a focuses on recent estimates of total RFari over the period 1990-2015. Using the ECLv5 emission inventory, *Myhre et al.* [2017] found a global mean RFari due to changes in aerosol abundances over the period 1990-2015 of 0.05 (\pm 0.04) W m⁻². Our results using CEDSv16 emissions are in close agreement with these findings.

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732 Over the past decades, there has been shift in emissions, from North America and Europe to South and East Asia. This is also reflected in the zonally averaged net RFari over time in Fig. 8b. RFari 733 declined in magnitude north of 40°N after 1980, with particularly large year-to-year decreases 734 735 between 1990 and 1995, and from 2005 to 2010, and strengthened in magnitude between 10°-30°N. The RFari also strengthened in the Southern Hemisphere subtropical region, reflecting 736 incressing emission in Africa and South America after 1970. However, the peak net RFari is 737 738 considerably weaker in 2014 than the peak in 1980. This mainly is due to fact that simultaneously 739 with the southwards shift, the sulfate burden has declined, while the BC burden has increased steadily at the same latitudes, resulting in a weaker net RF. The past decade, the net RFari has 740 switched from negative to positive north of 70°N, due to a combination of stronger positive RF of 741 742 BC and from biomass burning aerosols.

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Table S5 shows changes in burden, AOD, AAOD, RFari, and normalized RF over the period 1750-

745 2010 for individual aerosol components and the net RFari. Compared to earlier versions of

OsloCTM [Myhre et al., 2009; Myhre et al., 2013a] the normalized RF with respect to AOD is

747 lower because of short lifetime of BC resulting in smaller abundance of BC above clouds, whereas

normalized RF with burden is comparable to earlier estimates because of higher MAC compensate

749 for short lifetime of BC. Weaker normalized RF of OA (POA and SOA) than earlier OsloCTM

versions is due to the inclusion of absorbing OA.

In the present study we have used an updated parameterization of BC absorption based on *Zanatta et al.* [2016] (Sect. 2.4), which takes into account the ratio of non-BC-to-BC material and results in a MAC of 12.5 m² g⁻¹ at 550 nm. This is 26% higher than the 9.94 m² g⁻¹ using the approach from *Bond and Bergstrom* [2006]. Using the latter, we estimate a BC RFari in 2014 relative to 1750 of 0.23 W m⁻², 25% lower than the 0.31 W m⁻² calculated based on *Zanatta et al.* [2016]. These results emphasize the importance of assumptions and uncertainties related to the BC absorption.

The magnitude of RFari by scattering aerosols is sensitive to assumptions about absorption by organic aerosols, so-called brown carbon (BrC). Observational studies have provided evidence for the existence of such particles, and modeling studies suggest they could be responsible for a substantial fraction of total aerosol absorption, although the spread in estimates is wide (e.g., *Feng et al.* [2013] and reference therein). In the present study we assume a considerable fraction of absorption by OA (Sect. 2.4). Assuming purely scattering aerosols, the RFari from OA is -0.13W m⁻²; acounting for BrC absorption this is weakened to -0.09 W m⁻². Splitting total OA RFari into 765 contributions from primary and secondary aerosols, we find that purely scattering POA gives a RFari of -0.07 W m⁻² compared to -0.06 Wm⁻² with absorption. The corresponding numbers for 766 SOA are -0.06 and -0.03 W m⁻². This indicates that in OsloCTM3, the absorbing properties of SOA 767 768 are relatively more important than for POA. This is likely due to the generally higher altitude of 769 SOA than POA (Fig. S8) in combination with the increasing radiative efficiency of absorbing aerosols with altitude [Samset et al., 2013]. However, due to the weaker overall contributions from 770 771 OA, our results indicate that differences in parameterization of BC absorption can be more important than uncertainties in absorption by BrC for the net RFari. 772

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774 4 Discussion

776 Our estimate of total net RFari in 2014 relative to 1750 is weaker in magnitude than the best estimate for the 1750-2010 period reported by IPCC AR5. The difference is due to a combination 777 778 of factors, including weaker contributions from both cooling aerosols and BC. Despite 779 considerably higher BC emissions in the CEDSv16 inventory compared to older inventories, we 780 calculate a weaker BC RFari than reported in AR5, hence going in the opposite direction of explaining the difference to IPCC AR5 total RFari. The IPCC AR5 best estimate for fossil fuel 781 and biofuel BC of 0.4 (0.05 to 0.8) W m⁻² [Boucher et al., 2013] was based mainly on the two 782 studies by Myhre et al. [2013a] and Bond et al. [2013], who derived estimates of BC RFari of 0.23 783 $(0.06 \text{ to } 0.48) \text{ W m}^{-2}$ and $0.51 (0.06 \text{ to } 0.91) \text{ W m}^{-2}$, respectively. The spread between the two is 784 largely attributed to methodological differences: Bond et al. [2013] used an observationally 785 weighted scaling of results to match those based on AERONET AAOD, which was not adopted 786 by *Myhre et al.* [2013a]. Such ad-hoc adjustments typically result in higher estimates (*Wang et al.* 787 788 [2018] and references therein). Moreover, a recent study by Wang et al. [2018] suggest that 789 representativeness error arising when constraining coarse resolution models with AERONET AAOD could result in a 30% overestimation of BC RFari, which explains some of the differences 790 between bottom-up and observationally constrained numbers. The BC RFari estimate from the 791 792 present study is around 20% higher than the AeroCom multi-model mean from Myhre et al. [2013a] when calculated over the same period 1850-2000. This reflects the higher emissions in the 793 794 CEDSv16 emission inventory than in *Lamarque et al.* [2010], as well as a higher MAC.

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A significant range from -0.85 to +0.15 W m⁻² surrounds the central RFari estimate of -0.35 W m⁻ 796 ² from IPCC AR5 [Boucher et al., 2013], caused by the large spread in underlying simulated 797 aerosol distributions. Deficiencies in the ability of global models to reproduce atmospheric aerosol 798 799 concentrations can propagate to uncertainties in RF estimates. As shown in Sect. 3, the OsloCTM3 800 generally lies close to or above the multi-model mean of anthropogenic aerosol burdens from recent studies and is found to perform reasonably well compared with observations and other 801 global models, with improvements over the predecessor OsloCTM2. In particular, recent progress 802 towards constraining the vertical distribution of BC concentrations has resulted in improved 803 agreement between modeled and observed vertical BC profiles over the Pacific Ocean with less of 804

the high-altitude overestimation seen in earlier studies. However, as shown by *Lund et al.* [2018], there are discrepancies compared to recent aircraft measurements over the Atlantic Ocean. A remaining challenge is the model underestimation of Arctic BC concentrations. However, this is seen mainly during winter and early spring, when the direct aerosol effect is small due to lack of sunlight. In contrast, the higher emissions in the CEDSv16 inventory also results in an improved agreement with BC surface concentrations over Asia.

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812 In general, we find lower surface sulfate concentrations in the model compared with measurements. This could contribute to an underestimation of the sulfate RFari, which is weaker in the present 813 study than in IPCC AR5. An underestimation of observed AOD in Asia is also found, however, 814 the implication of this bias on RF is not straightforward to assess, as it is complicated by the mix 815 of absorbing and scattering aerosols. We note that the global mean sulfate burden is higher in the 816 817 OsloCTM3 than in most of the global models participating in the AeroCom III experiment (Sect. 818 3.1, Bian et al. [2017]), and that the OsloCTM3 performs similarly to or better than other AeroCom Phase III models in terms of nitrate and sulfate surface concentrations, at least for measurements 819 from CASTNET [Bian et al., 2017]. Nevertheless, the model diversity in simulated nitrate and 820 821 sulfate remains large and, although all models capture the main observed features in concentrations, further work is needed to resolve the differences and improve model performance for these species. 822

823

While a comprehensive quantitive uncertainty analysis of the updated RFari estimate is not 824 possible within the scope of this study, we explore the order of magnitude uncertainties due to 825 "internal" factors such as scavenging parameterizations and model resolution by performing 826 827 sensitivity tests. Changes in global burden on the order of 10-20%, and up to 65%, were found (Sect. 3.2). However, compared to observations of surface concentrations in near-source regions, 828 total AOD and vertical distribution of BC concentrations, we saw that the model generally 829 performed the best in its baseline configuration. Furthermore, the largest changes in the simulated 830 831 AOD and aerosol distributions were found in high-latitude regions, whereas changes over land where the concentrations, and hence subsequent RF is localized, were smaller. For certain regions 832 and observables, there were notable differences between the baseline and sensitivity simulations. 833 For instance, an improvemet in the baseline compared to using the CMIP5 emission inventory was 834 835 seen for BC surface concentrations, in particular in Asia, while the NMB of AOD compared to AERONET stations in the same region was reduced in the simulation with higher spatial resolution. 836 The importance of using the correct meteorological year was also seen. Such uncertainties will 837 translate to the RFari estimates, along with uncertainties in optical properties such as absorption 838 839 by organic aerosols and parameterization of BC absorption (Sect. 3.3).

840

841 Estimates of radiative impacts depend critically on the confidence in the emission inventories. A

detailed discussion of uncertainties in the CEDS inventory is provided by *Hoesly et al.* [2018]. On

a global level, the uncertainty in SO_2 emissions tend to be relatively low, although there is an

indication of missing SO₂ sources in particular in the Persian Gulf [*McLinden et al.*, 2016], whereas

emission factors for BC, OC, NOx, CO and VOCs have higher uncertainties. Uncertainties in

country-specific emissions can also be much larger, which is particularly true for carbonaceous
aerosols. In future CEDS versions, a quantitative uncertainty analysis is planned [*Hoesly et al.*,
2018], which will provide valuble input to modeling studies.

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850 Our study does not include anthropogenic dust, i.e., wind-blown dust from soils disturbed by human activities such as land use practices, deforestation and agriculture, and fugitive combustion 851 and industrial dust from urban sources. These sources could contribute an important fraction of 852 emissions and ambient PM_{2.5} concentrations in some regions [Paul et al., 2012; Sajeev et al., 2017], 853 but are missing from most models today. For instance, a recent study found a 2–16 mg m⁻³ increase 854 in PM2.5 concentrations in East and South Asia from anthropogenic fugitive, combustion, and 855 industrial dust emissions. However, the transport processes and optical properties, and hence, 856 radiative impact, is poorly known. We also do not include the effect of aerosol-cloud interactions, 857 858 which are crucial for the net climate impact of aerosols. For instance, recent studies suggest that 859 the impact of BC on global temperature response is small due to largely compensating direct and rapid adjustment effects [Samset and Myhre, 2015; Stjern et al., 2017]. The composition and 860 distribution of aerosols and oxidants in the pre-industrial atmosphere is uncertain and poorly 861 constrained by observations. However, while this is an important source of uncertainty in estimates 862 of RF due to aerosol-induced cloud albedo changes, it is less important for RFari because the 863 forcing scales quite linearly with aerosol burden [Carslaw et al., 2017]. 864

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867 5 Conclusions

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869 In this study, we have documented the third generation of the Oslo chemical transport model 870 (OsloCTM3) and evaluated the simulated distributions of aerosols, including results from a range of simulations to investigate the sensitivity to uncertainties in scavenging processes, input of 871 emissions and meteorological data and resolution. We have then used the new historical CEDS 872 873 emission inventory (version 2016; CEDSv16), which will also be used in the upcoming CMIP6, to simulate the temporal evolution of atmospheric concentrations of anthropogenic aerosols, and 874 quantified the temporal evolution of the subsequent radiative forcing due to aerosol-radiation 875 interactions (RFari). 876

877

878 The total AOD from the OsloCTM3 is in good agreement with observations from the AERONET network with a correlation of 0.82 and a normalized mean bias (NMB) of -11.8%. Regionally, the 879 880 underestimation of observed AOD is higher for stations in China and India than in Europe and North America, as also reflected from the comparison against measured aerosol surface 881 882 concentrations. High correlations 0.80-0.90 are also found for surface concentrations of BC, OC, sulfate and nitrate aerosols compared with all measurements across Europe, North America and 883 Asia. The corresponding NMB range from -23% for BC and nitrate to -46% and -52% for OC and 884 sulfate, respectively. The OsloCTM3 performs notably better than its predecessor OsloCTM2 in 885

terms of high-altitude BC distribution as compared with observed BC concentration profiles over
the Pacific Ocean from the HIPPO3 campaign. In constrast, the model continues to underestimate
observed surface levels of BC during winter and spring. Compared with other recent estimates of
aerosol burdens, the OsloCTM3 generally lies close to or above the mean of other global models.

Increasing or reducing the scavenging efficiency, moving to a finer resolution, and using the wrong 890 meteorological year or a different emission inventory results in changes in the global mean aerosol 891 burdens of up to 65%. The burdens of BC, OC and sulfate are particularly sensitive to a reduced 892 efficiency of removal by large-scale ice clouds; a 10 percentage point reduction increases the 893 894 global burden by 40%, 65% and 20%, respectively. A corresponding increase in the efficiency gives around 10% lower burdens. A significantly better agreement with BC surface concentrations 895 is found when using the CEDSv16 emission inventory compared with the RCP4.5. Furthermore, 896 a notable reduction in the bias of AOD compared to AERONET observations in Asia is found 897 when increasing the horizontal resolution, while the correlation is reduced when using the wrong 898 meteorological year. However, we find no clear evidence of consistently better model performance 899 across all observables and regions in the sensitivity tests than in the baseline configuration. This 900 may in part be influenced by the geographical coverage of observations, as the largest differences 901 in concentrations and AOD from the baseline is found at high altitudes and latitudes where the 902 903 availability of constraining measurements is limited.

Using the CEDSv16 historical emission inventory we estimate a total net RFari from all 904 anthropogenic aerosols, relative to 1750, of -0.17 W m⁻². This is significantly weaker than the best 905 estimate reported in the IPCC AR5, due to a combination of factors resulting in weaker 906 907 contributions from both cooling aerosols and BC in our simulations. Our updated RFari estimate is based on a single global model. As shown by previous studies, there is a large spread estimates 908 of RFari due to the spread in modeled aerosol distributions. The present analysis shows that 909 uncertainties in emissions, scavenging and optical properties of aerosols can have important 910 911 impacts on the simulated AOD and subsequent forcing estimates within one model. Additional studies to place our estimates in the context of multi-model spread and provide a comprehensive 912 uncertainty analysis are needed ahead of the IPCC Sixth Assessment Report. 913

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- 916 Data availability

The CEDS anthropogenic emissions data is published within the ESGF system <u>https://esgf-</u> node.llnl.gov/search/input4mips/. Surface observations used in this study are collected from the following publicly available databases: the EBAS database (http://ebas.nilu.no/) hosted by NILU – Norwegian Institute for Air Research. The US national Clean Air Status and Trends monitoring network (CASTNET), available at <u>http://www.epa.gov/castnet</u>. The Interagency Monitoring of Protected Visual Environments (IMPROVE), a collaborative association of state, tribal, and

923 federal agencies, and international partners, with the US EPA as the primary funding source and

924 support from the National Park Service. Data available from http://vista.cira.colostate.edu/Improve/. MODIS and MISR AOD retrievals are downloaded from 925 https://giovanni.gsfc.nasa.gov/giovanni/. Aircraft measurements from the HIPPO3 flights 926 available from https://www.eol.ucar.edu/node/524. Model output available upon request from 927 928 Marianne T. Lund (m.t.lund@cicero.oslo.no).

- 929
- 930 Code availability
- The OsloCTM3 is stored in a SVN repository at the University of Oslo central subversion
- system and is available upon request. Please contact m.t.lund@cicero.oslo.no. In this paper, we
- use the official version 1.0, OsloCTM3v1.0.
- 934
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- 949 Competing interests
- 950 The authors declare that they have no conflict of interest.
- 951 952 953 954 References 955 956 957 Aiken, A. C., P. F. DeCarlo, J. H. Kroll, D. R. Worsnop, J. A. Huffman, K. S. Docherty, I. M. Ulbrich, C. Mohr, 958 J. R. Kimmel, D. Sueper, Y. Sun, Q. Zhang, A. Trimborn, M. Northway, P. J. Ziemann, M. R. Canagaratna, T. 959 B. Onasch, M. R. Alfarra, A. S. H. Prevot, J. Dommen, J. Duplissy, A. Metzger, U. Baltensperger, and J. L. 960 Jimenez: O/C and OM/OC Ratios of Primary, Secondary, and Ambient Organic Aerosols with High-Resolution Time-of-Flight Aerosol Mass Spectrometry, Environmental Science & Technology, 42(12), 961
- 962 4478-4485, doi:10.1021/es703009q, **2008**

963 964 965 966 967	Amann, M., I. Bertok, J. Borken-Kleefeld, J. Cofala, C. Heyes, L. Höglund-Isaksson, Z. Klimont, B. Nguyen, M. Posch, P. Rafaj, R. Sandler, W. Schöpp, F. Wagner, and W. Winiwarter: Cost-effective control of air quality and greenhouse gases in Europe: Modeling and policy applications, Environmental Modelling & Software, 26(12), 1489-1501, doi: <u>http://dx.doi.org/10.1016/j.envsoft.2011.07.012</u> , 2011
968 969 970 971	Berglen, T. F., T. K. Berntsen, I. S. A. Isaksen, and J. K. Sundet: A global model of the coupled sulfur/oxidant chemistry in the troposphere: The sulfur cycle, Journal of Geophysical Research-Atmospheres, 109(D19), doi:10.1029/2003jd003948, 2004
972 973 974	Berntsen, T., J. Fuglestvedt, G. Myhre, F. Stordal, and T. F. Berglen: Abatement of greenhouse gases: Does location matter?, Climatic Change, 74(4), 377-411, doi:10.1007/s10584-006-0433-4, 2006
975 976 977 978 979	Bian, H., M. Chin, D. A. Hauglustaine, M. Schulz, G. Myhre, S. E. Bauer, M. T. Lund, V. A. Karydis, T. L. Kucsera, X. Pan, A. Pozzer, R. B. Skeie, S. D. Steenrod, K. Sudo, K. Tsigaridis, A. P. Tsimpidi, and S. G. Tsyro: Investigation of global particulate nitrate from the AeroCom Phase III experiment, Atmos. Chem. Phys., 2017(17), 12911-12940, doi: <u>https://doi.org/10.5194/acp-17-12911-2017</u> , 2017
980 981 982	Bond, T. C., and R. W. Bergstrom: Light absorption by carbonaceous particles: An investigative review, Aerosol Science and Technology, 40(1), 27-67, doi:10.1080/02786820500421521, 2006
983 984 985 986 987 988 989	Bond, T. C., S. J. Doherty, D. W. Fahey, P. M. Forster, T. Berntsen, B. J. DeAngelo, M. G. Flanner, S. Ghan, B. Kärcher, D. Koch, S. Kinne, Y. Kondo, P. K. Quinn, M. C. Sarofim, M. G. Schultz, M. Schulz, C. Venkataraman, H. Zhang, S. Zhang, N. Bellouin, S. K. Guttikunda, P. K. Hopke, M. Z. Jacobson, J. W. Kaiser, Z. Klimont, U. Lohmann, J. P. Schwarz, D. Shindell, T. Storelvmo, S. G. Warren, and C. S. Zender: Bounding the role of black carbon in the climate system: A scientific assessment, Journal of Geophysical Research: Atmospheres, 118(11), 5380-5552, doi:10.1002/jgrd.50171, 2013
990 991 992 993 994 995 996	Boucher, O., D. Randall, P. Artaxo, C. Bretherton, G. Feingold, P. Forster, K. VM., Y. Kondo, H. Liao, U. Lohmann, P. Rasch, S. K. Satheesh, S. Sherwood, B. Stevens, and X. Y. Zhang Clouds and Aerosols: In: Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change [Stocker, T.F., D. Qin, GK. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex and P.M. Midgley (eds.)]. Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA., 2013
997 998 999 1000	Bourgeois, Q., and I. Bey: Pollution transport efficiency toward the Arctic: Sensitivity to aerosol scavenging and source regions, Journal of Geophysical Research: Atmospheres, 116(D8), n/a-n/a, doi:10.1029/2010JD015096, 2011
1001 1002 1003	Browse, J., K. S. Carslaw, S. R. Arnold, K. Pringle, and O. Boucher: The scavenging processes controlling the seasonal cycle in Arctic sulphate and black carbon aerosol, Atmospheric Chemistry and Physics,

1004 12(15), 6775-6798, doi:10.5194/acp-12-6775-2012, **2012**

1005 1006 Carslaw, K. S., H. Gordon, D. S. Hamilton, J. S. Johnson, L. A. Regayre, M. Yoshioka, and K. J. Pringle: 1007 Aerosols in the Pre-industrial Atmosphere, Current Climate Change Reports, 3(1), 1-15, 1008 doi:10.1007/s40641-017-0061-2, 2017 1009 1010 Cavalli, F., A. Alastuey, H. Areskoug, D. Ceburnis, J. Čech, J. Genberg, R. M. Harrison, J. L. Jaffrezo, G. Kiss, 1011 P. Laj, N. Mihalopoulos, N. Perez, P. Quincey, J. Schwarz, K. Sellegri, G. Spindler, E. Swietlicki, C. 1012 Theodosi, K. E. Yttri, W. Aas, and J. P. Putaud: A European aerosol phenomenology -4: Harmonized 1013 concentrations of carbonaceous aerosol at 10 regional background sites across Europe, Atmospheric 1014 Environment, 144, 133-145, doi:https://doi.org/10.1016/j.atmosenv.2016.07.050, 2016 1015 1016 Cooke, W. F., C. Liousse, H. Cachier, and J. Feichter: Construction of a 1 degrees x 1 degrees fossil fuel 1017 emission data set for carbonaceous aerosol and implementation and radiative impact in the ECHAM4 1018 model, Journal of Geophysical Research-Atmospheres, 104(D18), 22137-22162, 1019 doi:10.1029/1999jd900187, 1999 1020 1021 Dubovik, O., and M. D. King: A flexible inversion algorithm for retrieval of aerosol optical properties from 1022 Sun and sky radiance measurements, Journal of Geophysical Research: Atmospheres, 105(D16), 20673-1023 20696, doi:10.1029/2000JD900282, 2000 1024 1025 Eckhardt, S., B. Quennehen, D. J. L. Olivié, T. K. Berntsen, R. Cherian, J. H. Christensen, W. Collins, S. 1026 Crepinsek, N. Daskalakis, M. Flanner, A. Herber, C. Heyes, Ø. Hodnebrog, L. Huang, M. Kanakidou, Z. 1027 Klimont, J. Langner, K. S. Law, M. T. Lund, R. Mahmood, A. Massling, S. Myriokefalitakis, I. E. Nielsen, J. K. 1028 Nøjgaard, J. Quaas, P. K. Quinn, J. C. Raut, S. T. Rumbold, M. Schulz, S. Sharma, R. B. Skeie, H. Skov, T. Uttal, K. von Salzen, and A. Stohl: Current model capabilities for simulating black carbon and sulfate 1029 1030 concentrations in the Arctic atmosphere: a multi-model evaluation using a comprehensive measurement 1031 data set, Atmos. Chem. Phys., 15(16), 9413-9433, doi:10.5194/acp-15-9413-2015, 2015 1032 1033 Fan, S. M., J. P. Schwarz, J. Liu, D. W. Fahey, P. Ginoux, L. W. Horowitz, H. Levy, Y. Ming, and J. R. 1034 Spackman: Inferring ice formation processes from global-scale black carbon profiles observed in the 1035 remote atmosphere and model simulations, Journal of Geophysical Research: Atmospheres, 117(D23), 1036 D23205, doi:10.1029/2012JD018126, 2012 1037 1038 Feng, Y., V. Ramanathan, and V. R. Kotamarthi: Brown carbon: a significant atmospheric absorber of 1039 solar radiation?, Atmos. Chem. Phys., 13(17), 8607-8621, doi:10.5194/acp-13-8607-2013, 2013 1040 1041 Gantt, B., M. S. Johnson, M. Crippa, A. S. H. Prévôt, and N. Meskhidze: Implementing marine organic 1042 aerosols into the GEOS-Chem model, Geosci. Model Dev., 8(3), 619-629, doi:10.5194/gmd-8-619-2015, 1043 2015 1044 1045 Granier, C., B. Bessagnet, T. Bond, A. D'Angiola, H. Denier van der Gon, G. J. Frost, A. Heil, J. W. Kaiser, S. 1046 Kinne, Z. Klimont, S. Kloster, J.-F. Lamarque, C. Liousse, T. Masui, F. Meleux, A. Mieville, T. Ohara, J.-C. 1047 Raut, K. Riahi, M. G. Schultz, S. J. Smith, A. Thompson, J. van Aardenne, G. R. van der Werf, and D. P. van

1048 1049 1050	Vuuren: Evolution of anthropogenic and biomass burning emissions of air pollutants at global and regional scales during the 1980–2010 period, Climatic Change, 109(1), 163, doi:10.1007/s10584-011-0154-1, 2011
1051 1052 1053 1054	Grini, A., G. Myhre, J. K. Sundet, and I. S. A. Isaksen: Modeling the annual cycle of sea salt in the global 3D model Oslo CTM2: Concentrations, fluxes, and radiative impact, Journal of Climate, 15(13), 1717-1730, doi:10.1175/1520-0442(2002)015<1717:mtacos>2.0.co;2, 2002
1055 1056 1057 1058	Grini, A., G. Myhre, C. S. Zender, and I. S. A. Isaksen: Model simulations of dust sources and transport in the global atmosphere: Effects of soil erodibility and wind speed variability, Journal of Geophysical Research-Atmospheres, 110(D2), doi:10.1029/2004jd005037, 2005
1059 1060 1061	Hodnebrog, Ø., G. Myhre, and B. H. Samset: How shorter black carbon lifetime alters its climate effect, Nat Commun, 5, 5065, doi:10.1038/ncomms6065, 2014
1062 1063 1064 1065 1066 1067	Hoesly, R. M., S. J. Smith, L. Feng, Z. Klimont, G. Janssens-Maenhout, T. Pitkanen, J. J. Seibert, L. Vu, R. J. Andres, R. M. Bolt, T. C. Bond, L. Dawidowski, N. Kholod, J. I. Kurokawa, M. Li, L. Liu, Z. Lu, M. C. P. Moura, P. R. O'Rourke, and Q. Zhang: Historical (1750–2014) anthropogenic emissions of reactive gases and aerosols from the Community Emission Data System (CEDS), Geosci. Model Dev., 2018(11), 369-408, doi: <u>https://doi.org/10.5194/gmd-11-369-2018</u> , 2018
1068 1069 1070 1071 1072	Holben, B. N., T. F. Eck, I. Slutsker, D. Tanré, J. P. Buis, A. Setzer, E. Vermote, J. A. Reagan, Y. J. Kaufman, T. Nakajima, F. Lavenu, I. Jankowiak, and A. Smirnov: AERONET—A Federated Instrument Network and Data Archive for Aerosol Characterization, Remote Sensing of Environment, 66(1), 1-16, doi: <u>https://doi.org/10.1016/S0034-4257(98)00031-5</u> , 1998
1073 1074 1075	Hoyle, C. R., T. Berntsen, G. Myhre, and I. S. A. Isaksen: Secondary organic aerosol in the global aerosol- chemical transport model Oslo CTM2, Atmos. Chem. Phys., 7(5675-5694), 2007
1076 1077 1078 1079	Jaeglé, L., P. K. Quinn, T. S. Bates, B. Alexander, and J. T. Lin: Global distribution of sea salt aerosols: new constraints from in situ and remote sensing observations, Atmos. Chem. Phys., 11(7), 3137-3157, doi:10.5194/acp-11-3137-2011, 2011
1080 1081 1082 1083	Kinne, S., D. O'Donnell, P. Stier, S. Kloster, K. Zhang , H. Schmidt, S. Rast, M. Giorgetta, T. F. Eck, and B. Stevens: MAC-v1: A new global aerosol climatology for climate studies, J. Adv. Model. Earth Syst., 5, 704-740, doi:10.1002/jame.20035, 2013
1084 1085 1086 1087	Kipling, Z., P. Stier, C. E. Johnson, G. W. Mann, N. Bellouin, S. E. Bauer, T. Bergman, M. Chin, T. Diehl, S. J. Ghan, T. Iversen, A. Kirkevåg, H. Kokkola, X. Liu, G. Luo, T. van Noije, K. J. Pringle, K. von Salzen, M. Schulz, Ø. Seland, R. B. Skeie, T. Takemura, K. Tsigaridis, and K. Zhang: What controls the vertical

1088 distribution of aerosol? Relationships between process sensitivity in HadGEM3–UKCA and inter-model

1089 1090	variation from AeroCom Phase II, Atmos. Chem. Phys., 16(4), 2221-2241, doi:10.5194/acp-16-2221- 2016, 2016
1091 1092 1093 1094 1095	Kipling, Z., P. Stier, J. P. Schwarz, A. E. Perring, J. R. Spackman, G. W. Mann, C. E. Johnson, and P. J. Telford: Constraints on aerosol processes in climate models from vertically-resolved aircraft observations of black carbon, Atmos. Chem. Phys., 13(12), 5969-5986, doi:10.5194/acp-13-5969-2013, 2013
1096 1097 1098 1099	Kirchstetter, T. W., T. Novakov, and P. V. Hobbs: Evidence that the spectral dependence of light absorption by aerosols is affected by organic carbon, Journal of Geophysical Research: Atmospheres, 109(D21), n/a-n/a, doi:10.1029/2004JD004999, 2004
1100 1101 1102 1103	Klimont, Z., K. Kupiainen, C. Heyes, P. Purohit, J. Cofala, P. Rafaj, J. Borken-Kleefeld, and W. Schöpp: Global anthropogenic emissions of particulate matter including black carbon, Atmos. Chem. Phys., 17(14), 8681-8723, doi:10.5194/acp-17-8681-2017, 2017
1104 1105 1106	Klimont, Z., S. J. Smith, and J. Cofala: The last decade of global anthropogenic sulfur dioxide: 2000–2011 emissions, Environmental Research Letters, 8(1), 014003, 2013
1107 1108 1109 1110	Kumar, R., M. C. Barth, G. G. Pfister, V. S. Nair, S. D. Ghude, and N. Ojha: What controls the seasonal cycle of black carbon aerosols in India?, Journal of Geophysical Research: Atmospheres, 120(15), 7788-7812, doi:10.1002/2015JD023298, 2015
1111 1112 1113 1114 1115 1116	Lamarque, J. F., T. C. Bond, V. Eyring, C. Granier, A. Heil, Z. Klimont, D. Lee, C. Liousse, A. Mieville, B. Owen, M. G. Schultz, D. Shindell, S. J. Smith, E. Stehfest, J. Van Aardenne, O. R. Cooper, M. Kainuma, N. Mahowald, J. R. McConnell, V. Naik, K. Riahi, and D. P. van Vuuren: Historical (1850–2000) gridded anthropogenic and biomass burning emissions of reactive gases and aerosols: methodology and application, Atmos. Chem. Phys., 10(15), 7017-7039, doi:10.5194/acp-10-7017-2010, 2010
1117 1118 1119 1120 1121	Lavery, T. F., C. M. Rogers, R. Baumgardner, and K. P. Mishoe: Intercomparison of Clean Air Status and Trends Network Nitrate and Nitric Acid Measurements with Data from Other Monitoring Programs, Journal of the Air & Waste Management Association, 59(2), 214-226, doi:10.3155/1047-3289.59.2.214, 2009
1122 1123 1124 1125 1126 1127 1128	Lee, Y. H., J. F. Lamarque, M. G. Flanner, C. Jiao, D. T. Shindell, T. Berntsen, M. M. Bisiaux, J. Cao, W. J. Collins, M. Curran, R. Edwards, G. Faluvegi, S. Ghan, L. W. Horowitz, J. R. McConnell, J. Ming, G. Myhre, T. Nagashima, V. Naik, S. T. Rumbold, R. B. Skeie, K. Sudo, T. Takemura, F. Thevenon, B. Xu, and J. H. Yoon: Evaluation of preindustrial to present-day black carbon and its albedo forcing from Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP), Atmospheric Chemistry and Physics, 13(5), 2607-2634, doi:10.5194/acp-13-2607-2013, 2013
1129	

1130 1131 1132	Li, C., C. McLinden, V. Fioletov, N. Krotkov, S. Carn, J. Joiner, D. Streets, H. He, X. Ren, Z. Li, and R. R. Dickerson: India Is Overtaking China as the World's Largest Emitter of Anthropogenic Sulfur Dioxide, Scientific Reports, 7(1), 14304, doi:10.1038/s41598-017-14639-8, 2017
1133 1134 1135 1136	Lund, M. T., and T. Berntsen: Parameterization of black carbon aging in the OsloCTM2 and implications for regional transport to the Arctic, Atmos. Chem. Phys., 12(15), 6999-7014, doi:10.5194/acp-12-6999-2012, 2012
1137 1138 1139 1140	Lund, M. T., T. K. Berntsen, and B. H. Samset: Sensitivity of black carbon concentrations and climate impact to aging and scavenging in OsloCTM2–M7, Atmos. Chem. Phys., 17(9), 6003-6022, doi:10.5194/acp-17-6003-2017, 2017
1141 1142 1143 1144	Lund, M. T., B. H. Samset, R. B. Skeie, D. Watson-Parris, J. M. Katich, J. P. Schwarz, and B. Weinzierl: Short Black Carbon lifetime inferred from a global set of aircraft observations. Accepted manuscript. , npj Climate and Atmospheric Science 1, 31, doi:10.1038/s41612-018-0040-x, 2018
1145 1146 1147 1148	Mahmood, R., K. von Salzen, M. Flanner, M. Sand, J. Langner, H. Wang, and L. Huang: Seasonality of global and Arctic black carbon processes in the Arctic Monitoring and Assessment Programme models, Journal of Geophysical Research: Atmospheres, 121(12), 7100-7116, doi:10.1002/2016JD024849, 2016
1149 1150 1151 1152	McLinden, C. A., V. Fioletov, M. W. Shephard, N. Krotkov, C. Li, R. V. Martin, M. D. Moran, and J. Joiner: Space-based detection of missing sulfur dioxide sources of global air pollution, Nature Geoscience, 9, 496, doi:10.1038/ngeo2724
1153	https://www.nature.com/articles/ngeo2724#supplementary-information, 2016
1154 1155 1156 1157	Metzger, S., F. Dentener, M. Krol, A. Jeuken, and J. Lelieveld: Gas/aerosol partitioning - 2. Global modeling results, Journal of Geophysical Research-Atmospheres, 107(D16), doi:10.1029/2001jd001103, 2002a
1158 1159 1160 1161	Metzger, S., F. Dentener, S. Pandis, and J. Lelieveld: Gas/aerosol partitioning: 1. A computationally efficient model, Journal of Geophysical Research-Atmospheres, 107(D16), doi:10.1029/2001jd001102, 2002b
1162 1163 1164 1165 1166 1167 1168	MISR (2018), Data product specification for the MISR level 2 aerosol product, Garay, M.J., et al., <u>https://eosweb.larc.nasa.gov/project/misr/DPS_AEROSOL_V023.20180125.pdf</u> (accessed 04/26/2018), edited. MOD08: MODIS Level 3 Atmosphere Products (MOD 08), Data Products Handbook Volume 2. <u>https://modis.gsfc.nasa.gov/data/dataprod/dataproducts.php?MOD_NUMBER=08</u> (accessed 04/26/2018), 2018
1169 1170 1171	Myhre, G., W. Aas, R. Cherian, W. Collins, G. Faluvegi, M. Flanner, P. Forster, Ø. Hodnebrog, Z. Klimont, M. T. Lund, J. Mülmenstädt, C. Lund Myhre, D. Olivié, M. Prather, J. Quaas, B. H. Samset, J. L. Schnell, M.

- 1172 Schulz, D. Shindell, R. B. Skeie, T. Takemura, and S. Tsyro: Multi-model simulations of aerosol and ozone
- 1173 radiative forcing due to anthropogenic emission changes during the period 1990–2015, Atmos. Chem.
- 1174 Phys., 17(4), 2709-2720, doi:10.5194/acp-17-2709-2017, **2017**
- 1175
- 1176 Myhre, G., N. Bellouin, T. F. Berglen, T. K. Berntsen, O. Boucher, A. L. F. Grini, I. S. A. Isaksen, M.
- 1177 Johnsrud, M. I. Mishchenko, F. Stordal, and D. TanrÉ: Comparison of the radiative properties and direct 1178 radiative effect of aerosols from a global aerosol model and remote sensing data over ocean, Tellus B,
- 1179 59(1), 115-129, doi:10.1111/j.1600-0889.2006.00226.x, **2007**
 - 1180
 - 1181 Myhre, G., T. F. Berglen, M. Johnsrud, C. R. Hoyle, T. K. Berntsen, S. A. Christopher, D. W. Fahey, I. S. A. 1182 Isaksen, T. A. Jones, R. A. Kahn, N. Loeb, P. Quinn, L. Remer, J. P. Schwarz, and K. E. Yttri: Modelled
 - 1183 radiative forcing of the direct aerosol effect with multi-observation evaluation, Atmospheric Chemistry
 - 1184 and Physics, 9(4), 1365-1392, doi:10.5194/acp-9-1365-2009, **2009**
 - 1185
 - 1186 Myhre, G., A. Grini, and S. Metzger: Modelling of nitrate and ammonium-containing aerosols in presence 1187 of sea salt, Atmos. Chem. Phys., 6, 4809-4821, **2006**
 - 1188
 - 1189 Myhre, G., B. H. Samset, M. Schulz, Y. Balkanski, S. Bauer, T. K. Berntsen, H. Bian, N. Bellouin, M. Chin, T.
 - 1190 Diehl, R. C. Easter, J. Feichter, S. J. Ghan, D. Hauglustaine, T. Iversen, S. Kinne, A. Kirkevåg, J. F.
 - 1191 Lamarque, G. Lin, X. Liu, M. T. Lund, G. Luo, X. Ma, T. van Noije, J. E. Penner, P. J. Rasch, A. Ruiz, Ø.
 - 1192 Seland, R. B. Skeie, P. Stier, T. Takemura, K. Tsigaridis, P. Wang, Z. Wang, L. Xu, H. Yu, F. Yu, J. H. Yoon, K.
 - 1193 Zhang, H. Zhang, and C. Zhou: Radiative forcing of the direct aerosol effect from AeroCom Phase II
 - simulations, Atmos. Chem. Phys., 13(4), 1853-1877, doi:10.5194/acp-13-1853-2013, **2013a**
 - 1195
 - 1196 Myhre, G., D. Shindell, F.-M. Brèon, W. Collins, J. Fuglestvedt, J. Huang, D. Koch, J.-F. Lamarque, D. Lee,
 - 1197 B. Mendoza, T. Nakajima, A. Robock, G. Stephens, T. Takemura, and H. Zhang: Anthropogenic and
 - 1198 natural radiative forcing. In: Climate Change 2013: The Physical Science Basis. Contribution of Working
 - 1199 Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change [Stocker, T.F.,
 - 1200 D., Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex and P.M. Midgley (eds).
- 1201 Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA **2013b**
- 1202
- Neu, J. L., and M. J. Prather: Toward a more physical representation of precipitation scavenging in global
 chemistry models: cloud overlap and ice physics and their impact on tropospheric ozone, Atmos. Chem.
 Phys., 12(7), 3289-3310, doi:10.5194/acp-12-3289-2012, **2012**
- 1206
- Parhi, P., A. Giannini, P. Gentine, and U. Lall: Resolving Contrasting Regional Rainfall Responses to El
 Niño over Tropical Africa, Journal of Climate, 29(4), 1461-1476, doi:10.1175/jcli-d-15-0071.1, 2016

- Paul, G., P. J. M., G. T. E., H. N. Christina, and Z. Ming: Global scale attribution of anthropogenic and
 natural dust sources and their emission rates based on MODIS Deep Blue aerosol products, Reviews of
 Geophysics, 50(3), doi:doi:10.1029/2012RG000388, **2012**
- 1213

- 1214 Petzold, A., J. A. Ogren, M. Fiebig, P. Laj, S. M. Li, U. Baltensperger, T. Holzer-Popp, S. Kinne, G.
- 1215 Pappalardo, N. Sugimoto, C. Wehrli, A. Wiedensohler, and X. Y. Zhang: Recommendations for reporting
- 1216 "black carbon" measurements, Atmos. Chem. Phys., 13(16), 8365-8379, doi:10.5194/acp-13-8365-2013,
- 1217 **2013**
- 1218
- 1219 Putaud, J. P., R. Van Dingenen, A. Alastuey, H. Bauer, W. Birmili, J. Cyrys, H. Flentje, S. Fuzzi, R. Gehrig, H.
- 1220 C. Hansson, R. M. Harrison, H. Herrmann, R. Hitzenberger, C. Hüglin, A. M. Jones, A. Kasper-Giebl, G.
- 1221 Kiss, A. Kousa, T. A. J. Kuhlbusch, G. Löschau, W. Maenhaut, A. Molnar, T. Moreno, J. Pekkanen, C.
- 1222 Perrino, M. Pitz, H. Puxbaum, X. Querol, S. Rodriguez, I. Salma, J. Schwarz, J. Smolik, J. Schneider, G.
- 1223 Spindler, H. ten Brink, J. Tursic, M. Viana, A. Wiedensohler, and F. Raes: A European aerosol
- 1224 phenomenology 3: Physical and chemical characteristics of particulate matter from 60 rural, urban,
- and kerbside sites across Europe, Atmospheric Environment, 44(10), 1308-1320,
- 1226 doi:<u>https://doi.org/10.1016/j.atmosenv.2009.12.011</u>, **2010**
- 1227
- 1228 Randerson, J. T., G. R. van der Werf, L. Giglio, G. J. Collatz, and P. S. Kasibhatla: Global Fire Emissions
- 1229 Database, Version 4.1 (GFEDv4). ORNL DAAC, Oak Ridge, Tennessee, USA.,
- 1230 doi:<u>https://doi.org/10.3334/ORNLDAAC/1293</u>, **2017**
- 1231
- 1232 Sajeev, P., V. M. Randall, S. Graydon, L. W. Crystal, D. Aaron van, B. Michael, K. H. Daven, K. Zbigniew, V.
- 1233 Chandra, K. G. Sarath, and Z. Qiang: Anthropogenic fugitive, combustion and industrial dust is a
- 1234 significant, underrepresented fine particulate matter source in global atmospheric models,
- 1235 Environmental Research Letters, 12(4), 044018, 2017
- 1236
- Samset, B. H., and G. Myhre: Climate response to externally mixed black carbon as a function of altitude,
 Journal of Geophysical Research: Atmospheres, 120(7), 2913-2927, doi:10.1002/2014JD022849, 2015
- 1239
- Samset, B. H., G. Myhre, A. Herber, Y. Kondo, S. M. Li, N. Moteki, M. Koike, N. Oshima, J. P. Schwarz, Y.
 Balkanski, S. E. Bauer, N. Bellouin, T. K. Berntsen, H. Bian, M. Chin, T. Diehl, R. C. Easter, S. J. Ghan, T.
 Iversen, A. Kirkevåg, J. F. Lamarque, G. Lin, X. Liu, J. E. Penner, M. Schulz, Ø. Seland, R. B. Skeie, P. Stier,
 T. Takemura, K. Tsigaridis, and K. Zhang: Modelled black carbon radiative forcing and atmospheric
 lifetime in AeroCom Phase II constrained by aircraft observations, Atmos. Chem. Phys., 14(22), 12465-
- 1245 12477, doi:10.5194/acp-14-12465-2014, **2014**
- 1246
- Samset, B. H., G. Myhre, M. Schulz, Y. Balkanski, S. Bauer, T. K. Berntsen, H. Bian, N. Bellouin, T. Diehl, R.
 C. Easter, S. J. Ghan, T. Iversen, S. Kinne, A. Kirkevåg, J. F. Lamarque, G. Lin, X. Liu, J. E. Penner, Ø. Seland,
 R. B. Skeie, P. Stier, T. Takemura, K. Tsigaridis, and K. Zhang: Black carbon vertical profiles strongly affect
 its radiative forcing uncertainty, Atmos. Chem. Phys., 13(5), 2423-2434, doi:10.5194/acp-13-2423-2013,
 2013
- 1252
- Samset, B. H., C. W. Stjern, E. Andrews, R. A. Kahn, G. Myhre, M. Schulz, and G. L. Schuster: Aerosol
 Absorption: Progress Towards Global and Regional Constraints, Current Climate Change Reports,
- 1255 doi:10.1007/s40641-018-0091-4, **2018**
- 1256

1258 1259	Sato, Y., H. Miura, H. Yashiro, D. Goto, T. Takemura, H. Tomita, and T. Nakajima: Unrealistically pristine air in the Arctic produced by current global scale models, Scientific Reports, 6, 26561, doi:10.1038/srep26561
1260	http://www.nature.com/articles/srep26561#supplementary-information, 2016
1261 1262 1263 1264	Schutgens, N., S. Tsyro, E. Gryspeerdt, D. Goto, N. Weigum, M. Schulz, and P. Stier: On the spatio- temporal representativeness of observations, Atmos. Chem. Phys., 17(16), 9761-9780, doi:10.5194/acp- 17-9761-2017, 2017
1265 1266 1267 1268	Schutgens, N. A. J., E. Gryspeerdt, N. Weigum, S. Tsyro, D. Goto, M. Schulz, and P. Stier: Will a perfect model agree with perfect observations? The impact of spatial sampling, Atmos. Chem. Phys., 16(10), 6335-6353, doi:10.5194/acp-16-6335-2016, 2016a
1269 1270 1271 1272	Schutgens, N. A. J., D. G. Partridge, and P. Stier: The importance of temporal collocation for the evaluation of aerosol models with observations, Atmos. Chem. Phys., 16(2), 1065-1079, doi:10.5194/acp-16-1065-2016, 2016b
1273 1274 1275 1276	Schwarz, J. P., B. H. Samset, A. E. Perring, J. R. Spackman, R. S. Gao, P. Stier, M. Schulz, F. L. Moore, E. A. Ray, and D. W. Fahey: Global-scale seasonally resolved black carbon vertical profiles over the Pacific, Geophysical Research Letters, 40(20), 2013GL057775, doi:10.1002/2013GL057775, 2013
1277 1278 1279 1280	Sindelarova, K., C. Granier, I. Bouarar, A. Guenther, S. Tilmes, T. Stavrakou, J. F. Müller, U. Kuhn, P. Stefani, and W. Knorr: Global data set of biogenic VOC emissions calculated by the MEGAN model over the last 30 years, Atmos. Chem. Phys., 14(17), 9317-9341, doi:10.5194/acp-14-9317-2014, 2014
1281 1282 1283 1284	Skeie, R. B., T. Berntsen, G. Myhre, C. A. Pedersen, J. Ström, S. Gerland, and J. A. Ogren: Black carbon in the atmosphere and snow, from pre-industrial times until present, Atmospheric Chemistry and Physics, 11(14), 6809-6836, doi:10.5194/acp-11-6809-2011, 2011
1285 1286 1287	Smith, S. J., Y. Zhou, P. Kyle, H. Wang, and H. Yu: A Community Emissions Data System (CEDS): Emissions for CMIP6 and Beyond. , 2015
1288 1289 1290 1291	Sofiev, M., J. Soares, M. Prank, G. de Leeuw, and J. Kukkonen: A regional-to-global model of emission and transport of sea salt particles in the atmosphere, Journal of Geophysical Research: Atmospheres, 116(D21), n/a-n/a, doi:10.1029/2010JD014713, 2011
1292 1293 1294 1295	Stamnes, K., S. C. Tsay, W. Wiscombe, and K. Jayaweera: Numerically stable algorithm for discrete- ordinate-method radiative transfer in multiple scattering and emitting layered media, Appl. Opt., 27(12), 2502-2509, doi:10.1364/AO.27.002502, 1988

1297 Stjern, C. W., B. H. Samset, G. Myhre, P. M. Forster, Ø. Hodnebrog, T. Andrews, O. Boucher, G. Faluvegi, 1298 T. Iversen, M. Kasoar, V. Kharin, A. Kirkevåg, J.-F. Lamarque, D. Olivié, T. Richardson, D. Shawki, D. 1299 Shindell, C. J. Smith, T. Takemura, and A. Voulgarakis: Rapid Adjustments Cause Weak Surface 1300 Temperature Response to Increased Black Carbon Concentrations, Journal of Geophysical Research: 1301 Atmospheres, 122(21), 11,462-411,481, doi:10.1002/2017JD027326, 2017 1302 1303 Søvde, O. A., M. J. Prather, I. S. A. Isaksen, T. K. Berntsen, F. Stordal, X. Zhu, C. D. Holmes, and J. Hsu: The 1304 chemical transport model Oslo CTM3, Geosci. Model Dev., 5(6), 1441-1469, doi:10.5194/gmd-5-1441-1305 2012, 2012 1306 1307 Taylor, K. E., R. J. Stouffer, and G. A. Meehl: An Overview of CMIP5 and the Experiment Design, Bulletin 1308 of the American Meteorological Society, 93(4), 485-498, doi:10.1175/bams-d-11-00094.1, 2012 1309 1310 Thomson, A. M., K. V. Calvin, S. J. Smith, G. P. Kyle, A. Volke, P. Patel, S. Delgado-Arias, B. Bond-1311 Lamberty, M. A. Wise, L. E. Clarke, and J. A. Edmonds: RCP4.5: a pathway for stabilization of radiative 1312 forcing by 2100, Climatic Change, 109(1), 77, doi:10.1007/s10584-011-0151-4, 2011 1313 1314 Tsigaridis, K., N. Daskalakis, M. Kanakidou, P. J. Adams, P. Artaxo, R. Bahadur, Y. Balkanski, S. E. Bauer, N. 1315 Bellouin, A. Benedetti, T. Bergman, T. K. Berntsen, J. P. Beukes, H. Bian, K. S. Carslaw, M. Chin, G. Curci, 1316 T. Diehl, R. C. Easter, S. J. Ghan, S. L. Gong, A. Hodzic, C. R. Hoyle, T. Iversen, S. Jathar, J. L. Jimenez, J. W. 1317 Kaiser, A. Kirkevåg, D. Koch, H. Kokkola, Y. H. Lee, G. Lin, X. Liu, G. Luo, X. Ma, G. W. Mann, N. 1318 Mihalopoulos, J. J. Morcrette, J. F. Müller, G. Myhre, S. Myriokefalitakis, N. L. Ng, D. O'Donnell, J. E. 1319 Penner, L. Pozzoli, K. J. Pringle, L. M. Russell, M. Schulz, J. Sciare, Ø. Seland, D. T. Shindell, S. Sillman, R. B. 1320 Skeie, D. Spracklen, T. Stavrakou, S. D. Steenrod, T. Takemura, P. Tiitta, S. Tilmes, H. Tost, T. van Noije, P. G. van Zyl, K. von Salzen, F. Yu, Z. Wang, Z. Wang, R. A. Zaveri, H. Zhang, K. Zhang, Q. Zhang, and X. 1321 1322 Zhang: The AeroCom evaluation and intercomparison of organic aerosol in global models, Atmos. Chem. 1323 Phys., 14(19), 10845-10895, doi:10.5194/acp-14-10845-2014, 2014 1324 1325 Turpin, B. J., and H.-J. Lim: Species Contributions to PM2.5 Mass Concentrations: Revisiting Common 1326 Assumptions for Estimating Organic Mass, Aerosol Science and Technology, 35(1), 602-610, 1327 doi:10.1080/02786820119445, 2001 1328 1329 Tørseth, K., W. Aas, K. Breivik, A. M. Fjæraa, M. Fiebig, A. G. Hjellbrekke, C. Lund Myhre, S. Solberg, and 1330 K. E. Yttri: Introduction to the European Monitoring and Evaluation Programme (EMEP) and observed 1331 atmospheric composition change during 1972 - 2009, Atmos. Chem. Phys., 12(12), 5447-5481, 1332 doi:10.5194/acp-12-5447-2012, 2012 1333 1334 van Marle, M. J. E., S. Kloster, B. I. Magi, J. R. Marlon, A. L. Daniau, R. D. Field, A. Arneth, M. Forrest, S. 1335 Hantson, N. M. Kehrwald, W. Knorr, G. Lasslop, F. Li, S. Mangeon, C. Yue, J. W. Kaiser, and G. R. van der 1336 Werf: Historic global biomass burning emissions for CMIP6 (BB4CMIP) based on merging satellite

- 1337 observations with proxies and fire models (1750–2015), Geosci. Model Dev., 10(9), 3329-3357,
- 1338 doi:10.5194/gmd-10-3329-2017, **2017**

1340 van Vuuren, D. P., J. Edmonds, M. Kainuma, K. Riahi, A. Thomson, K. Hibbard, G. C. Hurtt, T. Kram, V. 1341 Krey, J.-F. Lamarque, T. Masui, M. Meinshausen, N. Nakicenovic, S. J. Smith, and S. K. Rose: The 1342 representative concentration pathways: an overview, Climatic Change, 109(1), 5, doi:10.1007/s10584-1343 011-0148-z, **2011** 1344 1345 Wang, R., E. Andrews, Y. Balkanski, O. Boucher, G. Myhre, B. H. Samset, M. Schulz, G. L. Schuster, M. 1346 Valari, and S. Tao: Spatial Representativeness Error in the Ground-Level Observation Networks for Black 1347 Carbon Radiation Absorption, Geophysical Research Letters, 45, 2106-2114, 1348 doi:10.1002/2017GL076817, 2018 1349 1350 Wang, R., Y. Balkanski, O. Boucher, P. Ciais, G. L. Schuster, F. Chevallier, B. H. Samset, J. Liu, S. Piao, M. 1351 Valari, and S. Tao: Estimation of global black carbon direct radiative forcing and its uncertainty 1352 constrained by observations, Journal of Geophysical Research: Atmospheres, 121(10), 5948-5971, 1353 doi:10.1002/2015JD024326, 2016 1354 1355 Wang, R., S. Tao, Y. Balkanski, P. Ciais, O. Boucher, J. Liu, S. Piao, H. Shen, M. R. Vuolo, M. Valari, H. 1356 Chen, Y. Chen, A. Cozic, Y. Huang, B. Li, W. Li, G. Shen, B. Wang, and Y. Zhang: Exposure to ambient black 1357 carbon derived from a unique inventory and high-resolution model, Proceedings of the National 1358 Academy of Sciences, 111(7), 2459-2463, doi:10.1073/pnas.1318763111, 2014 1359 1360 Witek, M. L., D. J. Diner, and M. J. Garay: Satellite assessment of sea spray aerosol productivity: 1361 Southern Ocean case study, Journal of Geophysical Research: Atmospheres, 121(2), 872-894, 1362 doi:10.1002/2015JD023726, 2016 1363 1364 Wofsy, S. C., H. S. Team, T. Cooperating Modellers, and T. Satellite: HIAPER Pole-to-Pole Observations 1365 (HIPPO): fine-grained, global-scale measurements of climatically important atmospheric gases and 1366 aerosols, Philosophical Transactions of the Royal Society a-Mathematical Physical and Engineering 1367 Sciences, 369(1943), 2073-2086, doi:10.1098/rsta.2010.0313, 2011 1368 1369 Xie, M., M. D. Hays, and A. L. Holder: Light-absorbing organic carbon from prescribed and laboratory 1370 biomass burning and gasoline vehicle emissions, Scientific Reports, 7(1), 7318, doi:10.1038/s41598-017-1371 06981-8, 2017 1372 1373 Zanatta, M., M. Gysel, N. Bukowiecki, T. Müller, E. Weingartner, H. Areskoug, M. Fiebig, K. E. Yttri, N. Mihalopoulos, G. Kouvarakis, D. Beddows, R. M. Harrison, F. Cavalli, J. P. Putaud, G. Spindler, A. 1374 1375 Wiedensohler, A. Alastuey, M. Pandolfi, K. Sellegri, E. Swietlicki, J. L. Jaffrezo, U. Baltensperger, and P. 1376 Laj: A European aerosol phenomenology-5: Climatology of black carbon optical properties at 9 regional 1377 background sites across Europe, Atmospheric Environment, 145, 346-364, 1378 doi:https://doi.org/10.1016/j.atmosenv.2016.09.035, 2016 1379 1380 Zender, C. S., H. Bian, and D. Newman: Mineral Dust Entrainment and Deposition (DEAD) model: 1381 Description and 1990s dust climatology, Journal of Geophysical Research: Atmospheres, 108(D14),

1382 doi:doi:10.1029/2002JD002775, 2003

1383 1384 1385	Zhang, X. Y., Y. Q. Wang, T. Niu, X. C. Zhang, S. L. Gong, Y. M. Zhang , and J. Y. Sun: Atmospheric aerosol compositions in China: spatial/temporal variability, chemical signature, regional haze distribution and
1386	comparison with global models. , Atmos. Chem. Phys., 12, 779-799, 2012
1387	
1388	
1389	
1390	
1391	
1392	
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Table 1: Summary and description of simulations in this study

Name	Athropogenic	Year	Res	Description
	emissions			
CEDSv16/CMIP6	CEDS, version	2010	2x2	Baseline simulation, 2.25x2.25 degree resolution
	released in 2016			
ECLv5	ECLIPSEv5	2010	2x2	As baseline, but with ECLIPSEv5 emissions
RCP/CMIP5	RCP4.5	2010	2x2	As baseline, but RCP4.5/CMIP5 emissions
LSIDEC	CEDS	2010	2x2	Reduced scavenging of all aerosols by large-scale ice clouds
LSIINC	CEDS	2010	2x2	Increased scavenging of all aerosols by large-scale ice clouds
SOLDEC	CEDS	2010	2x2	Decreased scavenging of all aerosols by convective and large-scale liquid preciptation
1x1RES	Ix1RES CEDS		1x1	Same as baseline, but on 1.125x1.125 degree resolution
METDTA	CEDS	2010	2x2	Year 2010 emissions, but 2000 meteorology
Historical	CEDS/	1750-	2x2	Time-slice simulations for year 1750, 1850, 1900, 1910,
		2014		1920, 1930, 1940, 1950, 1960, 1970, 1980, 1985, 1990,
				1995, 2000, 2005, 2010, 2014

Table 2: Fraction of aerosol mass available for wet scavenging by convective, large-scale liquid
and large-scale ice precipitation in baseline setup and in the three sensitivity tests.
Phil=hydrophilic, phob=hydrophobic.

Simulation	Precipitation	Sulfate	OM	OM	BC	BC	Nitrate	SOA	Sea	Dust
	type		phil	phob	phil	Phob			salt	
CEDSv16/	Convective	1	1	1	1	1	1	0.8	1	1
CMIP6	LS-liquid	1	1	0	1	0	1	0.8	1	1
	LS-ice	0.1	0.1	0.2	0.1	0.2	0.1	0.16	0.1	0.5
LSIINC	LS-ice	0.3	0.3	0.4	0.3	0.4	0.3	0.32	0.3	0.7
LSIDEC	LS-ice	0.001	0.001	0.1	0.001	0.1	0.001	0.001	0.001	0.1
SOLDEC	Convective	0.8	0.8	0.8	0.8	0.8	0.8	0.6	0.8	0.8
	LS-liquid	0.8	0.8	0	0.8	0	0.8	0.6	0.8	0.8

1414 Table 3: Global, annual mean aerosol burdens $[mg m^{-2}]$ and total AOD in the baseline and

1415 sensitivity simulations. Parentheses in the top row give the atmospheric residence time (ratio of

1416 burden to total wet plus dry scavenging) [days]. Corresponding values for the sensitivity

simulations are given in Table S3.

				NH4	Nitrate	Nitrate				
Simulation	BC	OA	Sulfate	(fine+coarse)	(fine)	(coarse)	Sea salt	Dust	AOD	
CEDS/CMIP6	0.23	3.4 [§]	5.4	0.68	0.17	3.9	12	39		
	(4.4)	(5.3)	(5.4)	(3.5)	(4.2)	(5.2)	(0.46)	(3.4)	0.13	
ECLv5	0.21	3.1	5.1	0.65	0.15	3.7	12	39	0.13	
RCP/CMIP5	0.18	3.2	5.3	0.63	0.13	3.7	12	39	0.13	
LSIINC	0.21	2.8	4.9	0.63	0.17	3.4	11	39	0.12	
LSIDEC	0.32	5.3	6.5	0.79	0.16	4.7	14	43	0.16	
SOLDEC	0.26	3.6	6.1	0.78	0.16	5.2	15	42	0.15	
1x1RES	0.24	3.4	5.6	0.71	0.19	3.6	12	38	0.14	
METDTA	0.22	3.0	5.5	0.69	0.16	3.8	12	42	0.13	
[§] SOA: 1.1 mg m ⁻² [5.8 days] and POA: 2.3 mg m ⁻² [5.1 days]										





Figures



Figure 1: Annual mean modeled versus measured aerosol surface concentrations of a) EC, b)

1448 OC, c) sulfate and d) nitrate from the IMPROVE, EMEP, ACTRIS, CASTNET and CAWNET
1449 measurements networks.



- 1452 Figure 2: Aerosol composition (fraction of total aerosol mass) derived from the IMPROVE, EMEP,
- 1453 ACTRIS and CAWNET networks (left column) and corresponding OsloCTM3 results (right column).



Figure 3: Annual mean (year 2010) modeled a) AOD and b) AAOD, c) MODIS-Aqua AOD
retrieval and d) scatter density plot of comparison of simulated AOD against monthly mean
AERONET observations.



Figure 4: Ratio of each sensitivity simulation relative to the baseline for AOD (columns 1 and 3)
and total burden by species in each model layer (columns 2 and 4).



1471 Figure 5: Taylor diagram of modeled and measured aerosol surface concentrations in the baseline
1472 simulation and sensitivity tests using all observations in Fig. 1.



1478 Figure 6: Modeled vertical BC profiles against rBC aircraft measurements in five different

latitudes bands over the Pacific Ocean from the HIPPO3 flight campaign. Model data is

extracted along the flight track using an online flight simulator. Black lines: mean of

observations (solid), mean + plus 1 standard deviation (dashed). Colored lines: OsloCTM3

baseline (CEDSv16/CMIP6) (solid), sensitivity simulations (dashed).



Figure 7: Change in anthropogenic aerosol load over the period 1750 to 2014 using CEDSv16
emissions. Black symbols show the 1750 to 2000 difference and red symbols show multi-model
mean and OsloCTM2 results from the AeroCom II experiments [Myhre et al., 2013a].



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Figure 8: a) Time evolution of RFari. Solid lines show OsloCTM3 results from the current study, 1514

while dashed lines show results from IPCC AR5[Myhre et al., 2013b]. The inset shows the change 1515

1516 in total RFari between 1990 and 2015 in the current study compared with IPCC AR5 and multimodel mean and OsloCTM2 results from Myhre et al. [2017] using ECLv5 emissions. b) zonal 1517

1518 mean RFari 1750-2014.