Radiative Forcing of the Direct Aerosol Effect from AeroCom Phase II Simulations

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Abstract. We report on the AeroCom Phase II direct aerosol effect (DAE) experiment where 16 detailed global aerosol models have been used to simulate the changes in the aerosol distribution over the industrial era. All 16 models have estimated the radiative forcing (RF) of the anthropogenic DAE, and have taken into account anthropogenic sulphate, black carbon (BC) and organic aerosols (OA) from fossil fuel, biofuel, and biomass burning emissions. In addition several models have simulated the DAE of anthropogenic nitrate and anthropogenic influenced secondary organic aerosols (SOA). The model simulated all-sky RF of the DAE from total anthropogenic aerosols has a range from −0.58 to −0.02 Wm⁻², with a mean of −0.27 Wm⁻² for the 16 models. Several models did not include nitrate or SOA and modifying the estimate by accounting for this with information from the other AeroCom models reduces the range and

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slightly strengthens the mean. Modifying the model estimates for missing aerosol components and for the time period 1750 to 2010 results in a mean RF for the DAE of $-0.35 \text{ Wm}^{-2}$. Compared to AeroCom Phase I (Schulz et al., 2006) we find very similar spreads in both total DAE and aerosol component RF. However, the RF of the total DAE is stronger negative and RF from BC from fossil fuel and biofuel emissions are stronger positive in the present study than in the previous AeroCom study. We find a tendency for models having a strong (positive) BC RF to also have strong (negative) sulphate or OA RF. This relationship leads to smaller uncertainty in the total RF of the DAE compared to the RF of the sum of the individual aerosol components. The spread in results for the individual aerosol components is substantial, and can be divided into diversities in burden, mass extinction coefficient (MEC), and normalized RF with respect to AOD. We find that these three factors give similar contributions to the spread in results.

1 Introduction

Global estimates of the direct aerosol effect (DAE) have improved tremendously over the last two decades, due to improvements in processes taken into account and handling of complexities in the aerosol schemes (Bauer et al., 2010; Chin et al., 2009; Seland et al., 2008; Stier et al., 2005; Takemura et al., 2002). The first study of radiative forcing (RF) of the DAE with a global aerosol model included only sulphate aerosols (Charlson et al., 1991). Now multi-model studies with multi aerosol components and including some complex aerosol microphysical schemes have been performed (Schulz et al., 2006).

In the estimate of the RF from the direct aerosol effect from AeroCom Phase I models (Schulz et al., 2006) anthropogenic sulphate, black carbon and organic carbon were considered. In the interim increased attention has been given to the direct aerosol effect of anthropogenic nitrate and secondary organic aerosols (SOA). The formation of ammonium nitrate is complex and is dependent on sufficient ammonia and thus competition with sulphate is important (Adams et al., 2001; Metzger et al., 2002). With the expected reduction in SO$_2$ emission in the future, nitrate may become more important as a climate forcer (Bauer et al., 2007; Bellouin et al., 2011). Over the last few years increasing consideration has also been paid to secondary organic aerosols (SOA), with advanced understanding of their anthropogenic influence (Hoyle et al., 2011; Robinson et al., 2007). Anthropogenic influence on SOA is now included with varying degrees of complexity in some global aerosol models (Hoyle et al., 2009; Liao and Seinfeld, 2005; Spracklen et al., 2011; Tsigeridis et al., 2006).

The mean RF of the total direct aerosol effect in Schulz et al. (2006) from 9 global aerosol models was $-0.22 \text{ Wm}^{-2}$. This is much weaker than several observational based estimates (Bellouin et al., 2005, 2008; Chung et al., 2005; Quaas et al., 2008). Observational based estimates have been made possible through the advancement of remote sensing from ground and space (Holben et al., 1998; Remer et al., 2008). Myhre (2009) showed that estimates of the direct aerosol effect must rely on global aerosol model estimates, but consistency between models and observational based methods could be reached by accounting for anthropogenic changes in the aerosol optical properties in the latter method.

Various factors lie behind the large range in the global estimates of the DAE, which in AeroCom Phase I was from $-0.41$ to $+0.04 \text{ Wm}^{-2}$ (Schulz et al., 2006). Schulz et al. (2006) showed that the variations in RF could be separated into diversities in aerosol residence time, aerosol mass extinction coefficients, and normalized forcing (RF divided by aerosol optical depth). Normalized forcing was found to be the single largest contributor to the variability. Differences in the aerosol vertical profile, known to be significant between the models, is among the factors that contribute to differences in the normalized RF (Heald et al., 2011; Schwarz et al., 2010; Textor et al., 2006; Zarzycki and Bond, 2010).

In this paper RF of the DAE from 16 recent state-of-the-art global aerosol models in the AeroCom Phase II are presented. Two of the 16 global aerosol models are from Asia, six from Europe and eight from North America. Additional models compared to Schulz et al. (2006) are included in Phase II, and the models have been further developed over the 7–8 yr since AeroCom Phase I. Results for anthropogenic nitrate and SOA are included in the present study. Evaluation of the global aerosol models is an ongoing activity within AeroCom, with comparisons of various measurements being performed (Huneeus et al., 2011; Koch et al., 2009; Koffi et al., 2012). Activities to understand the model differences are also an important part of AeroCom (Randles et al., 2012; Samset et al., 2012; Stier et al., 2012; Textor et al., 2007).

2 Methods

The RF of the total anthropogenic DAE is calculated as the difference between the reflected solar radiation at TOA for simulations with present (2000, or 2006 for a few models) and pre-industrial (1850) emissions of aerosols and their precursors, denoted respectively as CTRL and PRE simulations, but the same cloud and surface conditions. The RF has been simulated by 16 global aerosol models with various complexities with regard to number of aerosol species included, aerosol microphysical treatment and their aerosol optical properties. The host model is also of great importance since aerosol transport and removal is dependent on the transport scheme and distribution of precipitation, respectively. In the radiative transfer calculations cloud distribution and properties and surface albedo, among other factors, are important.
Table 1. Model description, general information. If meteorology was nudged or driven by reanalysis fields, the year 2006 meteorology was used.

<table>
<thead>
<tr>
<th>Model</th>
<th>Type</th>
<th>Resolution</th>
<th>Levels</th>
<th>Meteorology</th>
<th>Responsible</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCC</td>
<td>GCM</td>
<td>2.8°×2.8°</td>
<td>26</td>
<td>NCEP/NCAR reanalysis</td>
<td>Hua Zhang, Zhili Wang</td>
</tr>
<tr>
<td>CAM4-Oslo</td>
<td>GCM</td>
<td>2.5°×1.8°</td>
<td>26</td>
<td>Produced by CAM4 atmospheric physics with CAM4-Oslo cloud tuning and boundary data from the data ocean and sea-ice models of CCSM4.</td>
<td>Alf Kirkevåg, Trond Iversen, Øyvind Seland, X. Liu, R. C. Easter, Steve Ghan, P. J. Rasch, J.-H. Yoon, Fangqun Yu, Gan Luo, Xiaoyan Ma, Susanne Bauer, Kostas Tsigaridis, Susanne Bauer, Huisheng Bian, Hongbin Yu, Thomas Diehl, Mian Chin, Guangxing Lin, Joyce Penner, Li Xu, Cheng Zhou, Yves Balkanski, Michael Schulz, Didier Hauglustaine</td>
</tr>
<tr>
<td>CAM5.1</td>
<td>GCM</td>
<td>2.5°×1.8°</td>
<td>30</td>
<td>CAM5.1</td>
<td>X. Liu, R. C. Easter, Steve Ghan, P. J. Rasch, J.-H. Yoon</td>
</tr>
<tr>
<td>GEOS-Chem</td>
<td>CTM</td>
<td>5.0°×4.0°</td>
<td>47</td>
<td>GEOS-5, reanalysis, nudged</td>
<td>Fangqun Yu, Gan Luo, Xiaoyan Ma, Susanne Bauer, Kostas Tsigaridis, Susanne Bauer, Huisheng Bian, Hongbin Yu, Thomas Diehl, Mian Chin, Guangxing Lin, Joyce Penner, Li Xu, Cheng Zhou, Yves Balkanski, Michael Schulz, Didier Hauglustaine</td>
</tr>
<tr>
<td>GISS-MATRIX</td>
<td>GCM</td>
<td>2.5°×2.0°</td>
<td>40</td>
<td>Nudged to NCEP winds</td>
<td>Susanne Bauer, Kostas Tsigaridis</td>
</tr>
<tr>
<td>GISS-modelE</td>
<td>GCM</td>
<td>2.5°×2.0°</td>
<td>40</td>
<td>Nudged to NCEP winds</td>
<td>Susanne Bauer, Kostas Tsigaridis</td>
</tr>
<tr>
<td>GMI</td>
<td>CTM</td>
<td>2.5°×2.0°</td>
<td>72</td>
<td>GEOS-4 MERRA reanalysis for 2006, nudged</td>
<td>Huisheng Bian, Hongbin Yu, Thomas Diehl, Mian Chin, Guangxing Lin, Joyce Penner, Li Xu, Cheng Zhou, Yves Balkanski, Michael Schulz, Didier Hauglustaine</td>
</tr>
<tr>
<td>GOCART</td>
<td>CTM</td>
<td>2.5°×2.0°</td>
<td>30</td>
<td>GEOS-4 DAS (Goddard Earth Observing System version 4 Data Assimilation System), reanalysis for year 2006</td>
<td>Susanne Bauer, Kostas Tsigaridis, Susanne Bauer, Huisheng Bian, Hongbin Yu, Thomas Diehl, Mian Chin, Guangxing Lin, Joyce Penner, Li Xu, Cheng Zhou, Yves Balkanski, Michael Schulz, Didier Hauglustaine</td>
</tr>
<tr>
<td>IMPACT</td>
<td>CTM</td>
<td>5.0°×4.0°</td>
<td>46</td>
<td>DAO assimilation fields for 1997, reanalysis.</td>
<td>Susanne Bauer, Kostas Tsigaridis, Susanne Bauer, Huisheng Bian, Hongbin Yu, Thomas Diehl, Mian Chin, Guangxing Lin, Joyce Penner, Li Xu, Cheng Zhou, Yves Balkanski, Michael Schulz, Didier Hauglustaine</td>
</tr>
<tr>
<td>INCA</td>
<td>GCM</td>
<td>3.8°×1.9°</td>
<td>19</td>
<td>ECMWF reanalysis from the Integrated Forecast System (IFS) model for year 2006.</td>
<td>Jean-Francois Lamarque, Didier Hauglustaine</td>
</tr>
<tr>
<td>HadGEM2</td>
<td>GCM</td>
<td>1.8°×1.2°</td>
<td>38</td>
<td>ERA Interim data for 2006, nudged</td>
<td>Nicolas Bellouin, Kai Zhang, Philip Stier, Johann Feichter, Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
<tr>
<td>ECHAM5-HAM</td>
<td>GCM</td>
<td>1.8°×1.8°</td>
<td>31</td>
<td>Model nudged with ECMWF analysis for the year 2006</td>
<td>Nicolas Bellouin, Kai Zhang, Philip Stier, Johann Feichter, Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
<tr>
<td>NCAR-CAM3.5</td>
<td>GCM</td>
<td>2.5°×1.9°</td>
<td>26</td>
<td>GCM-generated</td>
<td>Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
<tr>
<td>OsloCTM2</td>
<td>CTM</td>
<td>2.8°×2.8°</td>
<td>60</td>
<td>ECMWF reanalysis from the Integrated Forecast System (IFS) model for year 2006</td>
<td>Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
<tr>
<td>SPRINTARS</td>
<td>GCM</td>
<td>1.1°×1.1°</td>
<td>56</td>
<td>NCEP/NCAR reanalysis (temperature and horizontal wind), nudged.</td>
<td>Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
<tr>
<td>TM5</td>
<td>CTM</td>
<td>3.0°×2.0°</td>
<td>34</td>
<td>ECMWF ERA-Interim reanalysis for year 2006</td>
<td>Jean-Francois Lamarque, Gunnar Myhre, Ragnhild B. Skeie, Terje Berntsen, Toshihiko Takamura, Ping Wang, Ana Ruiz</td>
</tr>
</tbody>
</table>

Tables 1 and 2 show a list of the models included in the AeroCom Phase II DAE experiment, together with the resolutions used, aerosols components treated and comments on their microphysics scheme. Meteorological data for year 2006 have been used for all the models, except for IMPACT with meteorological data for 1997 and for CAM4-Oslo, NCAR-CAM3.5, and CAM5.1 where nudging possibilities were not available. A protocol describing the data submitted from the global aerosol models are available from the AeroCom web site (http://aerocom.met.no/aerocomhome.html). Most of the simulations for this DAE experiment are performed with identical cloud cover and cloud optical properties in PRE and CTRL, i.e. no indirect aerosol effects are included, but one (CAM5.1) estimated the DAE from the CTRL-PRE difference (Δ) in the direct radiative forcing by all aerosols in each simulation (Ghan et al., 2012): Δ (S - S_clean), where S is the net downward solar at TOA calculated.
Table 2. Model description, aerosol information.

<table>
<thead>
<tr>
<th>Model</th>
<th>S</th>
<th>BC</th>
<th>OC</th>
<th>BB</th>
<th>SOA</th>
<th>NO3</th>
<th>Aerosol microphysics</th>
<th>References for aerosol module</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCC</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>–</td>
<td>12 bin sizes for each aerosol with radii between 0.005–0.01, 0.01–0.02, 0.02–0.04, 0.04–0.08, 0.08–0.16, 0.16–0.32, 0.32–0.64, 0.64–1.28, 1.28–2.56, 2.56–5.12, 5.12–10.24, and 10.24–20.48 µm.</td>
<td>Zhang et al. (2012a)</td>
</tr>
<tr>
<td>CAM4-Oslo</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>–</td>
<td>Mass conc. of SO$_4$, BC, OM, sea-salt and dust in four size-classes are tagged according to production mechanism. Based on 44 sectional size bins and lognormal distributions at the point of emission, look-up tables yield physical properties of the processed aerosols.</td>
<td>Kirkevåg et al. (2012)</td>
</tr>
<tr>
<td>CAM5.1</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>3 internally-mixed log-normal modes</td>
<td>Liu et al. (2012)</td>
</tr>
<tr>
<td>GISS-MATRIX</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Aerosol microphysical scheme</td>
<td>Bauer et al. (2008, 2010)</td>
</tr>
<tr>
<td>GISS-modelE</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Aerosol mass based scheme</td>
<td>Koch et al. (2007, 2006); Bauer et al. (2007); Tsigaridis et al. (2013)</td>
</tr>
<tr>
<td>GMI</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Y</td>
<td>5 bin sizes for dust, 4 bin sizes for sea-salt, 3 bin size for nitrate and sulfate, all aerosols with log-normal size distributions.</td>
<td>Bian et al. (2009)</td>
</tr>
<tr>
<td>GOCART</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Parameterized with prescribed dry particle sizes: 8 bins for dust, 4 bins for sea salt, 1 bin for sulfate, BC, and OA, with log-normal distributions, particle growth parameterized as a function of RH</td>
<td>Chin et al. (2000, 2002, 2009); Ginoux et al. (2001)</td>
</tr>
<tr>
<td>IMPACT</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>*</td>
<td>4 bin sizes for sea-salt and mineral dust, pure sulfate treated using 2 modes with predicted size and coagulation and condensation of SO$_4$ with other aerosols explicitly resolved.</td>
<td>Wang and Penner (2009); Lin et al. (2012); Xu and Penner (2012)</td>
</tr>
<tr>
<td>INCA</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Y</td>
<td>Soluble and insoluble aerosol treated separately, modal assumptions with lognormal size distributions. We distinguish between accumulation, coarse and super coarse modes.</td>
<td>Balkanski et al. (2004); Schulz (2007); Balkanski (2011); Szopa et al. (2012)</td>
</tr>
<tr>
<td>HadGEM2</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Y</td>
<td>Component aerosol mass transported in Aitken, accumulation, coarse, and dissolved modes. Size distributions assumed lognormal for interaction with radiation.</td>
<td>Bellouin et al. (2011)</td>
</tr>
<tr>
<td>ECHAM5-HAM</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Modal method, log-normal size distributions, 7 modes (4 soluble, 3 insoluble). Internal mixing is assumed for aerosol compositions considered in each mode, while external mixing is assumed among different aerosol modes.</td>
<td>Vignati et al. (2004); Stier et al. (2005); Zhang et al. (2012b)</td>
</tr>
</tbody>
</table>
Table 2. Continued.

<table>
<thead>
<tr>
<th>Model</th>
<th>S</th>
<th>BC</th>
<th>OC</th>
<th>BB</th>
<th>SOA</th>
<th>NO$_3$</th>
<th>Aerosol microphysics</th>
<th>References for aerosol module</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCAR-CAM3.5</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>Y</td>
<td>Bulk-aerosol model, except 4-bins for sea-salt and mineral dust</td>
<td>Lamarque et al. (2012)</td>
</tr>
<tr>
<td>OsloCTM2</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>8 bin sizes for sea-salt and mineral dust, aerosol mass scheme for other aerosols</td>
<td>Myhre et al. (2007, 2009);</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>with log-normal size distributions in calculations of optical properties</td>
<td>Skeie et al. (2011)</td>
</tr>
<tr>
<td>SPRINTARS</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>–</td>
<td>–</td>
<td>6 bins for dust, 4 bins for sea salt, 1 bin for sulfate, BC, and OA, with log-normal</td>
<td>Takemura et al. (2005, 2009)</td>
</tr>
<tr>
<td>TM5</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Modal method, log-normal size distributions, 7 modes (4 soluble, 3 insoluble).</td>
<td>Vignati et al. (2004);</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Internal mixing is assumed for aerosol compositions considered in each mode, while</td>
<td>Aan de Brugh et al. (2011)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>external mixing is assumed among different aerosol modes.</td>
<td></td>
</tr>
</tbody>
</table>

Note: GOCART only includes SOA from biogenic sources (terpene oxidation)

* The NO$_3$ values for forcing from this model were simulated using the same model as the IMPACT model described here, but did not include the chemistry of formation of SOA and used the simplified NO$_x$ chemistry described in Feng and Penner (2007). The resolution was 2.5 x 2.0.

with all aerosols present, and $S_{\text{clean}}$ is the solar calculated neglecting scattering and absorption by all aerosols.

Aerosol optical properties are treated differently in the global aerosol models but in all models the optical properties are derived using Mie theory; see further description in model references. All models include treatment of sulphate, black carbon, primary organic carbon, sea salt, and mineral dust. Internal mixing between BC and scattering aerosols enhances the absorption from BC (Fuller et al., 1999; Haywood and Shine, 1995), and half of the 16 models include some degree of internal mixture for BC. Most of the models have used emissions of aerosols and their precursors from Lamarque et al. (2010). CAM4-Oslo and SPRINTARS have used these IPCC AR5 emissions for PRE (year 1850), but for the CTRL simulations emissions for year 2006 from the AeroCom Phase II dataset has been used (HCA0 v1 or v2 by T. Diehl, see http://aerocom.met.no/emissions.html). This dataset also includes emissions estimates of BC, SO$_2$ and POM from aviation. BCC used year 2000 emissions, but preindustrial emissions for 1750. GEOS-Chem uses emissions for 2006 which is close to AeroCom emission for 2000 and employs pre-industry emission corresponding to 1750 (Dentener et al., 2006). TM5 used 1850 emissions from Lamarque et al. (2010) and for 2006 a linear interpolation between 2005 and 2010 for the representative concentration pathway RCP4.5 (Thomson et al., 2011).

If the diagnostics for the different components were difficult to establish, separate model simulations were performed to allow for an attribution to eg biomass burning (BB) or fossil fuel (FF) and bio fuel (BF) sources. For some of the models with internally-mixed aerosols the separation of AOD and RF by aerosol component is challenging. Several groups have solved this by additional radiation calls. Even with this method a few models have been unable to properly extract the component AOD. Where relevant, these results have been removed from the analysis. All simulations were performed such that the cloud fields were identical, e.g. by neglecting the influence of aerosols on clouds or keeping the influence constant.

Only anthropogenic aerosol effects will be treated in the present study, and reference to the DAE apply to the anthropogenic change in the aerosol distribution and its influence on the scattering and absorption of solar radiation. Results will be presented for sulphate, nitrate, BC from fossil fuel and biofuel, OA from fossil fuel and biofuel, BC and OA combined from biomass burning, and SOA.

3 Results

All results are for anthropogenic aerosols only, unless otherwise stated. Anthropogenic aerosols are defined as the difference between the CTRL and PRE simulations described above.

3.1 Total direct aerosol effect

The underlying albedo is crucial for RF of the direct aerosol effect (Haywood and Shine, 1995). Scattering aerosols are efficient at producing a RF over dark surfaces and absorbing aerosols are efficient over bright surfaces. Figure 1a shows the latitudinal variation of the top of the atmosphere (TOA) albedo of the 16 models and the satellite retrieval from
CERES (Smith et al., 2011; Wielicki et al., 1996). Note that the albedo at the TOA consists of reflection at the surface and scattering in the atmosphere from clouds, aerosols, and Rayleigh scattering, and thus the scattering may occur above the anthropogenic aerosols. The higher albedo at high latitudes is mainly due to high surface albedo from snow and ice, but larger cloud fraction also contributes. For most of the models the albedo compares well to the CERES data, except between 30° N and 30° S where almost all overestimate the albedo. By inspection of the geographical distribution (not shown) of the albedo the overestimation is largest in regions with low cloud fraction, indicating problems with too large cloud fractions in these regions for the models. The GOCART model has a very low albedo compared to the rest of the models and the CERES data, which is identified to be caused by the meteorological data. The agreement between the simulated latitudinal variation in effective broadband surface albedo (computed from surface level down and up welling short wave radiation fluxes as diagnosed in the models) is in general excellent (see Fig. 1b), except at high latitudes due to differences in snow cover or albedo for snow and for the Northern Hemisphere sea ice coverage and albedo for sea ice.

Figure 2 shows the latitudinal variation in the RF of the total DAE for all sky (Fig. 2a) and clear sky (Fig. 2b) conditions. The negative RF is at maximum around 30° N for most of the models, since the anthropogenic change in aerosol burden (see Fig. 3) has its maximum in this region combined with a relatively high solar radiation flux. GISS-Matrix and GMI have the strongest all-sky RF in the Northern Hemisphere, but for clear-sky the IMPACT model shows the strongest RF in most of the Northern Hemisphere. The RF in the Southern Hemisphere is generally weak. How clouds affect the RF of the direct aerosol effect depends on a variety
of factors. Among them are the degree of aerosol absorption, vertical position of aerosols in relation to clouds, and the cloud distribution in general. The all-sky RF is around half in magnitude of the clear sky RF, except for the models with close to zero all sky RF. All models turn over into a positive net RF at high northern latitudes. This is caused by the higher surface and cloud albedo at high latitudes, as can be seen in Fig. 1. The surface and cloud albedo is the main cause of the positive RF since it will be later shown that aerosol optical properties are not a cause for the positive RF (see Fig. 3). CAM4-Oslo has the strongest positive all-sky RF at high latitudes. This model has more anthropogenic absorbing aerosols at high latitudes than the other models (see Fig. 3). This is partly due to the different CTRL emission data sets, which are for year 2006 and have more biomass burning from forest fires than the 2000 emissions. Assumed emission height of biomass burning (up to 6 km) may be another reason. The treatment of convective mixing of aerosols and aerosol precursors probably also leads to somewhat overestimated aerosol burdens in CAM4-Oslo (see Kirkevåg et al., 2012).

Figure 3 shows the latitudinal variation in the anthropogenic AOD (550 nm) used in the radiative transfer calculations, anthropogenic absorbing AOD (AAOD) (550 nm), anthropogenic single scattering albedo (550 nm), and the change in single scattering albedo (550 nm) between pre-industrial (PRE) and current (CTRL). The anthropogenic AOD in Fig. 3a shows a strong hemispheric difference which is a consistent model characteristic. CAM4-Oslo has the highest and CAM5.1 and BCC the lowest anthropogenic AOD almost at all latitudes. The pattern for the anthropogenic AAOD (Fig. 3b) is rather similar in the models, with two peaks, one around the Equator where biomass burning emissions of BC are high, and the second around 30° N, where fossil fuel and biofuel emissions of BC are high. The range in the anthropogenic single scattering albedo (defined as the ratio of anthropogenic AOD-AAOD and AOD) (Fig. 3c) is large, with most values lying between 0.9 and 1.0 but even lower values than 0.9 seen at high latitudes for a few
models. The lowest anthropogenic single scattering albedo is simulated near the Equator and can be seen in Fig. 3b to be caused by strong anthropogenic absorption AOD that is not associated with a similar strong maximum in the anthropogenic AOD in Fig. 3a. This pattern is caused by large BB emissions of BC and OA in this region. Most models simulate a reduction in the single scattering albedo during the industrial era at all latitudes, except GISS-MATRIX, GO-CART, TM5 and IMPACT. The reduction in the single scattering albedo in the majority of the models is due to stronger growth in absorbing aerosol (BC) than scattering aerosols relative to pre-industrial distribution of absorbing and scattering aerosols. GO-CART has an increase in the single scattering albedo in the NH due to low single scattering albedo in the PRE simulation compared to most of the other models. IMPACT shows a weaker increase in single scattering albedo than GO-CART, but still positive in the NH. As will be shown later this model has a much larger anthropogenic influenced SOA than the other models, which is likely the cause of the increase in single scattering albedo.

Table 3 summarizes the global and annual mean values for RF, anthropogenic AOD, anthropogenic absorption AOD (AAOD), atmospheric absorption, anthropogenic single scattering albedo, and cloud fraction. The anthropogenic AOD has a mean of 0.030, which is slightly higher than the Schulz et al. (2006) value of 0.029. A main reason for this small increase is likely the inclusion of additional aerosol components in this experiment. The anthropogenic AAOD of 0.0015 is lower than the value of 0.0019 for AAOD attributed to BC in Schulz et al. (2006). However, here it should be mentioned that some of the models in this experiment include absorption by OA which is non-negligible at 550 nm and thus the anthropogenic AAOD is not solely due to BC. A main reason for the lower AAOD compared to Schulz et al. (2006) is likely due to the much smaller anthropogenic BB emissions of BC in a majority of the models using the Lamarque et al. (2010) data. Figure 4 shows the model simulated global and annual mean total RF and the RF for the six aerosol components. The simulated mean RF for the DAE is $-0.27\,\text{Wm}^{-2}$ which is stronger negative than the mean of $-0.22\,\text{Wm}^{-2}$ found in Schulz et al. (2006). A stronger negative RF of the DAE in this experiment than in Schulz et al. (2006) is in accordance with the stronger AOD and weaker AAOD as noted above. It can also be noted that the models in this experiment have lower global and annual mean cloud cover (mean of 57 % for the 13 models reporting cloud cover) than in the models in Schulz et al. (2006) (mean of 63 %). As shown in many previous studies of the DAE the total RF consists of aerosol components causing negative RF and positive RF. Figure 4 shows clearly that this balance varies substantially between the models. Further, it shows that some models have weak RF for the aerosol components (CAM5 and BCC) whereas other models have stronger aerosol component RF (CAM4-Oslo, OsloCTM2 and GMI).

![Fig. 4. Radiative forcing from the six components, overlain with the (unmodified) model total forcing (yellow bars).](image)

Not all models have all six aerosol components included. Figure 5 shows the total RF where modifications for missing aerosol components have been taken into account. We calculate the mean of each aerosol component from all models that treat it, then add that mean to the total RF for each component that an individual model doesn’t treat. Since the missing aerosol components in the models are predominantly scattering aerosols (nitrate, SOA, and in some few cases BB aerosols), the modification leads to stronger negative RF for all models with modification. The mean RF of the DAE changes from $-0.27\,\text{Wm}^{-2}$ to $-0.32\,\text{Wm}^{-2}$ with the aerosol component modification.

Table 3 also shows the normalized clear sky RF (NRF, clear sky RF divided by AOD), which ranges from $-17\,\text{Wm}^{-2}$ to $-76\,\text{Wm}^{-2}$ with a mean of $-27\,\text{Wm}^{-2}$. The NRF depends on a variety of factors such as spatial distribution, optical properties, and the vertical profile. The weakening of the DAE RF due to clouds (Schulz et al., 2006) depends on vertical positioning of the aerosols and the clouds (Chand et al., 2009; Liao and Seinfeld, 1998; Samset and Myhre, 2011) as well as the degree of absorption of the aerosols (single scattering albedo) (Haywood and Shine, 1995).

Atmospheric absorption by the aerosols is calculated as the difference between the RF at TOA and radiative effect of aerosols at the surface. Absorbing aerosols such as BC are the main cause for the atmospheric absorption by the aerosols, but scattering aerosols enhance the absorption by gases in the atmosphere and make a small contribution (Randles et al., 2012; Stier et al., 2012). Figure 6 shows
the atmospheric absorption by anthropogenic aerosols as a function of absorbing AOD. The mean atmospheric absorption (see Table 3) is 0.75 W m\(^{-2}\), which is weaker than the 0.82 W m\(^{-2}\) reported in Schulz et al. (2006). The normalized values (atmospheric absorption divided by absorbing AOD) are shown on the figure. Normalized values range from 360 W m\(^{-2}\) to 935 W m\(^{-2}\) with a mean of 525 W m\(^{-2}\) and the median 10% lower. Some of the models with lowest anthropogenic atmospheric absorption have the largest deviation from the mean of the normalized values. The correlation between anthropogenic absorption AOD and atmospheric absorption is 0.73 for the 12 models available for this part of the study. IMPACT has high atmospheric absorption mainly, as discussed above, due to a large fraction of SOA that absorbs

### Table 3. Global mean anthropogenic value for all-sky and clear sky RF, normalized RF (NRF) with respect to AOD for clear sky, atmospheric absorption, atmospheric absorption divided by AAOD, AOD, anthropogenic fraction of AOD, AAOD, single scattering albedo (SSA), combined natural and anthropogenic change in SSA from PRE simulation to CTRL simulation, and present day cloud fraction (CLT). For NCAR-CAM3 the sum of component forcings is used.

<table>
<thead>
<tr>
<th>Model</th>
<th>Load [mg m(^{-2})]</th>
<th>MEC [m(^{2}) g(^{-1})]</th>
<th>AOD [W m(^{-2})]</th>
<th>RF [W m(^{-2})]</th>
<th>NRFB [W g(^{-1})]</th>
<th>NRFA [W m(^{-2})]</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCC</td>
<td>1.29</td>
<td>5.4</td>
<td>0.0069</td>
<td>-0.14</td>
<td>-108</td>
<td>-20.0</td>
</tr>
<tr>
<td>CAM4-Oslo</td>
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<td>12.3</td>
<td>0.0342</td>
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<td>-173</td>
<td>-14.0</td>
</tr>
<tr>
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<td>10.1</td>
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</tr>
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<td>SPRINTARs</td>
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<td>0.0220</td>
<td>-0.37</td>
<td>-172</td>
<td>-16.6</td>
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</tbody>
</table>

| Mean            | 1.78                 | 12.7                        | 0.0213              | -0.32               | -185                | -16.0               |
| Median          | 1.69                 | 10.3                        | 0.0212              | -0.32               | -180                | -16.5               |
| Stddev          | 0.47                 | 8.6                         | 0.0096              | 0.11                | 71                  | 3.7                 |
Fig. 5. Model total RFs. Black bars show the bare modelled forcing, the colored bars show the forcing modified for untreated components (see text for details). The yellow bar shows the AeroCom mean of the total RF of DAE. Solid lines inside the boxes show the model mean, dashed lines show the median. The boxes indicate one standard deviation, while the whiskers indicate the max and min of the distribution. The yellow shaded bar shows the AeroCom mean when aerosol component adjustment is made for missing aerosol components.

Fig. 6. Correlation between anthropogenic absorption AOD and atmospheric absorption. Numbers show ratio AtmAbs/AAOD, the lines indicate the mean and one standard deviation of this ratio. \( R^2 = 0.73 \)

Fig. 7. Component and total RF. Total RF has been modified for missing components in individual models. Solid lines inside the boxes show the model mean, dashed lines show the median. The boxes indicate one standard deviation, while the whiskers indicate the max and min of the distribution.

at short wavelengths, while the absorption is much weaker at 550 nm which is the reported wavelength for absorption AOD. This leads to the strongest normalized atmospheric absorption of 935 W m\(^{-2}\) for IMPACT. The mean of the anthropogenic fraction of AOD (Table 3) is 24 %, slightly smaller than in Schulz et al. (2006), where a different preindustrial year (here 1850 instead of 1750) is one contributor to this reduction. Half of the models are close to the mean anthropogenic fraction, whereas two of the models have quite low anthropogenic fractions of 12–14 % and one has 35 %.

Figure 7 summarizes the component and aerosol component modified total RFs and their intermodel variability. Model means are shown as solid lines in the middle of the boxes, median values as dashed lines. The boxes indicate one standard deviation, and the bars show the maximum and minimum single values.

In the following sections we discuss the individual aerosol components in detail, highlighting model results that deviate significantly from the mean.

3.2 RF of sulphate

The RF of the direct aerosol effect of sulphate is shown in Fig. 4 for all models. In addition Figure 7 shows the mean, median, standard deviation and the range for sulphate aerosols and for other aerosol components. Table 4 summarizes the aerosol burden, mass extinction coefficient (MEC) (550 nm), AOD (550 nm), RF, and NRF with respect to both burden and AOD. The mean RF of sulphate from the AeroCom Phase II models of \(-0.32 \text{ W m}^{-2}\) is weaker than the mean of AeroCom Phase I models of \(-0.35 \text{ W m}^{-2}\) in Schulz et al. (2006). The range in RF is very similar in the two AeroCom DAE experiments. The global mean MEC is calculated as the ratio of global mean AOD to global mean burden. The
Fig. 8. Zonal mean SO$_4$ RF (a), burden (b), AOD at 550 nm (c), normalized RF with respect to AOD (NRF(A)) (d).

Table 5. Same as Table 4 for BC from FF and BF emissions.

<table>
<thead>
<tr>
<th>Model</th>
<th>Load [mg m$^{-2}$]</th>
<th>MEC [m$^2$ g$^{-1}$]</th>
<th>AOD [1]</th>
<th>RF [W m$^{-2}$]</th>
<th>NRFB [W g$^{-1}$]</th>
<th>NRFA [W m$^{-2}$]</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Stddev</td>
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<td>3.9</td>
<td>0.0005</td>
<td>0.07</td>
<td>630</td>
<td>37.1</td>
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</tbody>
</table>
MEC depends on various factors, where growth by water uptake for sulphate aerosols is a major cause of the difference between the models. To illustrate the importance of water uptake, the MEC in OsloCTM2 is 4.0 m$^2$ g$^{-1}$ for low relative humidity (below 30%) and is 10.1 m$^2$ g$^{-1}$ as a global mean with ambient relative humidity. The mean of the MEC is 12.7 m$^2$ g$^{-1}$, but also without the two models with very high MEC (NCAR-CAM3.5 and GISS modelE) the MEC is 9.7 m$^2$ g$^{-1}$ and 7% higher than in Schulz et al. (2006).

The weaker normalized RF with respect to AOD and stronger with respect to burden than in Schulz et al. (2006) could be a result of sulphate at generally lower altitudes, but other factors could contribute.

Figure 8 shows the latitudinal variation in RF, burden, AOD, NRF (with respect to AOD) for sulphate. All models have a maximum in the burden at around 30° N with a large difference in the magnitude. The difference in the magnitude of the burden is particularly large at high northern latitudes and is likely linked to transport and scavenging problems in the models (Rasch et al., 2000). The difference in RF of sulphate at high latitudes is not as evident as for the burden (and AOD), since NRF at high latitudes is weak due to the high surface albedo, and some of the models with strongest burden (and AOD) have weak NRF. The RF of sulphate is very weak in the Southern Hemisphere.

### 3.3 RF of BC

In Fig. 9 the latitudinal variation is shown for RF, burden, AOD, NRF (with respect to AOD) for BC from FF and BF emissions. Global numbers are shown in Table 5. The importance of surface and cloud albedo is evident in the NRF for BC, especially if it is compared to NRF for sulphate in Fig. 8d, which weakens (for a majority of the models) rather than strengthens over the poles as for BC. The much stronger NRF at high latitudes for BC than sulphate and the relatively large range in burden at high latitudes are also clearly evident for the RF. Schulz et al. (2006) showed the latitudinal variation in burden of BC (total and not only FF and BF as in Fig. 9b). There is no indication of a reduced range in the BC burden among the AeroCom models from Phase I to Phase II. For the maximum in RF at around 30° N there is almost an order of magnitude difference between the model with strongest RF (CAM4-Oslo) and the model with the weakest RF (BCC). The model differences in the NRF are larger for BC than for sulphate, and main reason for this is likely a strong dependence on altitude of BC and
variations in absorbing properties of BC (Samset and Myhre, 2011; Zarzycki and Bond, 2010). The model mean mass extinction coefficient is 10.5 m$^2$ g$^{-1}$. Unfortunately the model mass absorption coefficient for BC from FF and BF emissions was not diagnosed. Based on observations, Bond et al. (2006) recommend a mass absorption coefficient (MAC) of around 7.5 m$^2$ g$^{-1}$ for fresh BC particles and 50% higher for aged BC. Jacobson (2012) and Chung et al. (2012) use higher absorption enhancement factors. As the single scattering albedo for BC often is in the range of 0.2 to 0.3 (Bond and Bergstrom, 2006), the BC MEC should be around 25% higher than the MAC, implying the models underestimate the MEC and MAC. The low MEC for a few of the models is the cause of the generally low mean MEC; but half of the models have MEC according to the recommendations. Internal mixing of BC with other aerosol components increases the absorption, so simulating sufficiently high MEC requires representing internal mixing of BC and its effects on absorption. The difference in MEC for models using internal mixture (12.1 m$^2$ g$^{-1}$) is on average 34% higher than for models assuming external mixture (9.1 m$^2$ g$^{-1}$). However, recent work has questioned the importance of the enhancement of MEC from internal mixing, at least in certain regions (Cappa et al., 2012).

The model mean global average burden of BC is 0.14 mg m$^{-2}$ and a RF of 0.18 Wm$^{-2}$. The RF estimate is similar to the IPCC AR4 estimate, but much lower than Ramanathan and Carmichael (2008). However, the estimates here and in IPCC AR4 (Forster et al., 2007) are solely from FF and BF, whereas Ramanathan and Carmichael (2008) included also BC from BB emissions and estimate forcing for all BC, not just anthropogenic. The spread in NRF with respect to burden is larger than in NRF with respect to AOD due to variations in MEC included in the former. For the two NRF measures the model difference is shown to arise from differences in the vertical profile of BC (Samset et al., 2012), host model difference (Stier et al., 2012) and likely differences in optical properties, especially the single scattering albedo of BC.

### 3.4 RF of OA

The RF of primary OA from FF and BF is weak and has a mean of $-0.03$ Wm$^{-2}$. The RF together with burden, MEC, AOD and NRF are shown in Table 6. Primary OA from biomass burning and SOA are described in separate sections below. The burden of OA is on average about 20% of the sulphate burden; however, this fraction varies between the models. RF from primary OA from FF and BF is around 10% of the sulphate RF, and thus the NRF with respect to burden for OA is much weaker than for sulphate. One main reason for this is the much smaller water uptake for OA. This is also reflected in the MEC. Due to the smaller water uptake for OA compared to sulphate the particles are smaller, leading to a larger Ångström exponent and a stronger NRF with respect to AOD. Compared to BC from FF, OA has a lower burden in the Arctic (see Fig. 10). Combined with the weak NRF in the Arctic for scattering aerosols, this causes the quite weak RF from OA in this region seen in Fig. 10.

<table>
<thead>
<tr>
<th>Model</th>
<th>Load [mg m$^{-2}$]</th>
<th>MEC [m$^2$ g$^{-1}$]</th>
<th>AOD [1]</th>
<th>RF [W m$^{-2}$]</th>
<th>NRFB [W g$^{-1}$]</th>
<th>NRFA [W m$^{-2}$]</th>
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<tr>
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<td>0.0017</td>
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<td></td>
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<td>0.0020</td>
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<td>-113</td>
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<tr>
<td>Median</td>
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<td>0.0017</td>
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<td>Stddev</td>
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<td>3.0</td>
<td>0.0010</td>
<td>0.01</td>
<td>41</td>
<td>6.9</td>
</tr>
</tbody>
</table>
Another cause for the differences in NRF between the models as well as between sulphate and OA are differences in the single scattering albedo for OA. Several models have single scattering albedo for OA at 550 nm at 0.96 with increasing values for higher relative humidity. Some OA components have absorption at short solar wavelength with a reduction in the absorption with wavelength stronger than for BC (Jacobson, 2012; Kanakidou et al., 2005). Uncertainties in the refractive indexes and thus absorption for OA are large.

However, some of the major components of OA do not absorb (Kanakidou et al., 2005). OsloCTM2 has among the strongest RF for OA from FF and BF emissions and uses pure scattering OA aerosols. One additional simulation was performed with OsloCTM2 using refractive indexes for OA as in CAM4-Oslo, resulting in a 25% weakening in the RF of OA from FF and BF emissions. In the INCA model a larger effect was found when applying pure scattering OA instead.
of their standard simulation, resulting in a 72% enhancement in the RF of OA from FF and BF emissions.

3.5 RF SOA

Five models have explicitly performed RF simulations for SOA; the two GISS-models have SOA included in the total DAE simulations and some information is available for GISS-modelE. The uncertainties in the modelling of SOA are large (Spracklen et al., 2011), and thus it is not surprising that we find a huge range in the burden of anthropogenic SOA from 0.09 to 0.97 mg m\(^{-2}\) (see Table 7). The burden in IMPACT-Umich is substantially higher than the other models. Figure 11 shows that burden of SOA has a larger fraction around the Equator than for the other aerosol

![Fig. 11. Zonal mean SOA RF (a), burden (b), AOD at 550 nm (c), NRF(A) (d).](image-url)
Table 9. Same as Table 4 for combined OA and BC from BB emissions.

<table>
<thead>
<tr>
<th>Model</th>
<th>Load [mg m(^{-2})]</th>
<th>MEC [m(^2) g(^{-1})]</th>
<th>AOD [1]</th>
<th>RF [W m(^{-2})]</th>
<th>NRF [W g(^{-1})]</th>
<th>NRFA [W m(^{-2})]</th>
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<tr>
<td>BCC</td>
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<td>GOCART</td>
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<tr>
<td>HadGEM2</td>
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<td>0.0038</td>
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<td>0.0056</td>
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<td>NCAR-CAM3.5</td>
<td>0.21</td>
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<td>0.02</td>
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</tr>
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<td>0.0028</td>
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</table>

Mean 0.94 7.7 0.0054 -0.00 7 -2.1
Median 0.48 7.0 0.0038 -0.00 24 4.5

Atmos. Chem. Phys., 13, 1853–1877, 2013

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3.6 RF nitrate

Eight models have simulated RF of nitrate aerosols, explicitly. The resulting range of nitrate burdens has a larger span among the aerosol components than for sulphate and primary BC and OA, ranging from 0.14 to 0.90 mg m\(^{-2}\) (see Table 8 and Fig. 12). The mean RF is -0.08 Wm\(^{-2}\) but, like the burden, the RF also varies widely among the AeroCom Phase II models with a range from -0.12 to -0.02 Wm\(^{-2}\). The pattern of the zonal mean burden of nitrate resembles the sulphate pattern. There is a tendency for models with weak sulphate burden to have a strong nitrate burden and vice versa and to some extent this can be expected since ammonium nitrate can only be formed if there is an excess of ammonia and thus it competes with sulphate (Adams et al., 2001). However, there are some exceptions to this relationship with some models having both low burden for sulphate and nitrate. The hygroscopic growth for nitrate is generally stronger than for sulphate (Fitzgerald, 1975) so one might expect stronger or at least similar MEC for nitrate compared to sulphate, but this is not the case for GMI and NCAR-CAM3.5. However, regional variation in the aerosols and thus water uptake may influence the results. The NRF with respect to burden and with respect to AOD are rather similar for nitrate and sulphate.

The mean RF of the DAE of nitrate is substantial and with its potentially larger role in the future it is important to investigate the model differences and compare with observations. We encourage studies comparing nitrate surface concentrations and in the atmosphere from aircraft measurements, which over Europe indicate a large role of nitrate (Crosier et al., 2007; Putaud et al., 2010), as well as over ocean where observations show small fractions fine mode nitrate (Quinn and Bates, 2005).

3.7 RF BB

The RF for simulated BB aerosol varies between positive and negative RF, highly dependent on SSA and likely also the altitude of the BB aerosols. The mean RF from the models is close to zero. The BB RF is weak in magnitude, but it is important to note that the RF of BB consists of a positive RF from BC and a negative RF from OA of much larger magnitude than for the total RF from BB aerosols. The MECs for BB aerosols given in Table 9 are in the upper range of what is measured during aerosol campaigns in Africa (Haywood et al., 2003; Johnson et al., 2008). In Schulz et al. (2006) the burden of primary carbonaceous was given separately for BC and OA, but not differentiated by sources. Since RF of OAFF and BCFF are at least not weaker and the total primary carbonaceous burden is lower than in Schulz et al. (2006) this
Fig. 12. Zonal mean NO$_3$ RF (a), burden (b), AOD at 550 nm (c), NRF(A) (d).

Fig. 13. Model mean RF (left) and standard deviation (right).

indicates lower BB burden in Phase II compared to Phase I. The model range in burden of BB is large as illustrated by the large difference between the mean and the median values. Zonal mean RF, burden, AOD and NRF with respect to AOD is shown for BB aerosols in Fig. S1.

4 Discussion

The geographical distributions of the RF of the total DAE and its uncertainty (given as one standard deviation) are shown in Fig. 13. As shown in earlier studies the RF of the DAE is strongly negative in industrialized regions of Asia, Europe and Northern America and reaches positive RF at high latitudes and in other regions with high albedo (either high surface albedo or high cloud cover).

The model mean RF and standard deviation of DAE for sulphate, BC, and OA are shown in Fig. S2. Maximum strength in RF for sulphate, BC, and OA is in Southeast Asia. For sulphate a quite strong RF is also simulated in Europe and Eastern US. Over regions with high surface albedo such as Sahara and Arctic the RF for sulphate and OA are quite weak, but RF from BC is relatively strong. The standard deviations are highest in the regions of high RF.
The geographical distribution of RF by nitrate and SOA (Fig. S3) show similar pattern as for the other aerosol components (Fig. S2). For BB aerosols the geographical pattern of RF is very different from the other aerosol components. In some areas (southeast U. S.) this has been caused by reduction in BB emissions, but in other regions differing signs may be caused by the underlying albedo such as southern Africa (Chand et al., 2009; Keil and Haywood, 2003), where low albedo over land causes negative RF and large coverage of low lying clouds with BB aerosols above cause positive RF over ocean to the west of the coast. The high surface albedo at high latitudes is the major cause of the positive RF at high latitudes.

Schulz et al. (2006) introduced a way to investigate the contribution to the uncertainty from various key components in estimates of the RF of the direct aerosol effect. The same approach is shown in Fig. 14 for burden, MEC, NRF (with respect to AOD) and RF. The figure shows for each of burden, MEC, NRF and RF the contribution for the individual models to the diversity in the RF. For example one model may have a burden that contributes to a strong RF, but a MEC of opposite contribution. For sulphate the uncertainties in RF are slightly stronger than for the burden, MEC, NRF. However, the range for burden, MEC, NRF seems quite similar. CAM5, IMPACT, and BCC have the weakest RF. Weak MEC is shown to be a main reason for this for CAM5 and BCC, whereas for IMPACT burden and NRF are the main factors. CAM4-Oslo has the strongest sulphate RF. Here the burden is the main reason.

Whereas Schulz et al. (2006) found that diversity in NRF clearly dominated the diversity in RF of BC, Fig. 14 shows that diversities in burden and MEC are at least equally important in this study. Illustrative of how differences in burden, MEC, and NRF may compensate each other, the HadGEM2 model has the highest BC burden, but a RF of BC close to the mean, which can be seen to be caused by a rather low MEC. For OA the dominant cause for the spread in RF is differences in MEC, overwhelming the importance of differences in burden and NRF. ECHAM5-HAM has the weakest MEC of 1.6 m² g⁻¹ and IMPACT the strongest of 14.1 m² g⁻¹. The weak MEC for ECHAM5-HAM explains why it shows the weakest RF among the global aerosol models, since its burden and NRF are close to mean values.

The reasons for the differences in burden, MEC and NRF are numerous. For differences in NRF Randles et al. (2012) show that the radiative transfer scheme itself contributes to the diversity and Stier et al. (2012) further explores that host model differences such as surface albedo and clouds cause a substantial part of the diversity for the RF of the DAE. The differences in the aerosols vertical profile (Koffi et al., 2012; Schwarz et al., 2010) contribute to differences in NRF, especially for BC. Samset et al. (2012) show that 20–50 % of the diversity in RF of BC is caused by the differences in the BC vertical profile. Further studies within AeroCom will explore other causes of the diversity in the results. Since most of the models in this study apply similar data sets for anthropogenic emissions of aerosols and their precursors, the uncertainties in this study do not include the full uncertainty
in the RF of the DAE. The difference in the background aerosol concentration (pre-industrial) between the models is of small importance for the RF of the DAE since RF is relatively linearly depending on the anthropogenically perturbed concentrations at current levels of the aerosol species.

Global aerosol models with strong RF for sulphate have a tendency to also have strong RF for BC and similar for OA in relation to BC. The RF sulphate and RF BC have an anti-correlation of 0.62. Figure 15 shows the relationship between burden, RF and NRF for sulphate, BC and OA. Aerosol burden is dependent on dry and wet deposition and the transport schemes in the models, and the simulated burdens of various aerosol compounds are therefore correlated. Similarly, there is a tendency also for relationship between the NRF (with respect to burden) for various aerosol compounds. The total DAE which is the combined effect of all aerosol components with both positive RF (BC, and in some cases OA) and negative RF (rest of components) thus also consists of RF contributions that are correlated. Altogether, this leads to smaller range in the total DAE than combining the range through error propagation rules for the individual aerosol components. This is illustrated in Fig. 16, which shows the PDFs for the individual aerosol components and their sum, and the modelled total RF. A narrower PDF for the model total DAE than aerosol component sum is evident. Further, Figure 16 illustrates the smaller model diversity when the aerosol missing component modification is performed compared to the total DAE directly from the various models.

The emission data used in most of the simulations (Lamarque et al., 2010) is generated for the period 1850 to 2000. Anthropogenic emissions of aerosols and their precursors had started in 1850 and emission changes have occurred after 2000. Skeie et al. (2011) performed RF simulations for OsloCTM2 for the different aerosol species for several time intervals for the period 1750 to 2000 based on Lamarque et al. (2010), as well as for 2010 based on the RCP4.5 scenario. GISS model simulations in Shindell et al. (2012) were also performed under the RCP4.5 scenario for years 2000 and 2010. Figure 17 shows a scaling of the AeroCom RF simulation for 1850-2000 to the extended time period 1750–2010.
To achieve this, we first calculate the aerosol component specific ratios of OsloCTM2 RF between 2010 and 1750 (Skeie et al., 2011) to RF between 2000 and 1850 (present work). Similar ratios are extracted using the GISS values from 2000 to 2010. The ratios from these two models are combined, and applied to each of the aerosol components individually. To scale the total RF, we then take the multi-model mean from the present work and add the sum of the component scalings. BB is excluded from this analysis, due to large uncertainties in BB activity between 1750 and 1850 as well as the inhomogeneous RF (see Fig. S3). The strengthening in the RF for 1750–2010 relative to 1850–2000 is particularly strong for OA from FF and BF and nitrate (40–45%), quite important for BC from FF and BF and SOA (25–30%), but rather small for sulphate (5%). For OA from FF and BF, nitrate and SOA the largest strengthening occurs for the period 1750–1850 in comparison to the 2000–2010 period. For BC and sulphate the strengthening is rather similar in the 1750–1850 and 2000–2010 periods. The increase in BC emission from 2000 to 2010 is of similar magnitude in other emission data sets as we have applied in for the scaling (Granier et al., 2011). In Skeie et al. (2011) the increase in RF from BC (from FF and BF emissions) for the time period 2000 to 2010 was slightly larger than the associated increase in emissions due to also changes in the spatial pattern of the emissions. The increase in the RF of the primary carbonaceous aerosols from 2000 to 2010 were very similar in OsloCTM2 and GISS, but somewhat weaker for the secondary components in the GISS model compared to OsloCTM2. This indicates that uncertainties in the scaling are largest for nitrate and SOA which had a quite large change from 1750 to 1850. Taking into account the changes between 1750–1850 and 2000–2010 for the aerosol components changes the total DAE from $-0.32 \text{Wm}^{-2}$ (modified for missing aerosol components) for 1850–2000 to $-0.35 \text{Wm}^{-2}$ for the 1750–2010 period.

5 Conclusions

We have documented the RF of the DAE for anthropogenic aerosol emissions from 16 global aerosol models, combined and for sulphate, BC, OA, BB, nitrate and SOA separately. This AeroCom Phase II DAE experiment shows many similarities with the Phase I (Schulz et al., 2006), even though more models are included here and model development has taken place over the intervening years. We summarize here the most important findings:

- The total RF of DAE is more strongly negative in the current simulations than in AeroCom Phase I. The main reason for this is addition of new species such as nitrate and SOA.
- Sulphate and BC from FF and BF emissions are the two aerosol components with strongest absolute RF. The RF of the DAE of sulphate is very similar in the simulations presented here and in Schulz et al. (2006). On the other hand BC RF from FF and BF emissions is almost twice as strong compared to in Schulz et al. (2006).
- The global aerosol models have undergone substantial development over the last 7–8 yr, and while there are some changes relative to AeroCom Phase I (Schulz et al., 2006), the present analysis shows that the main estimates across AeroCom Phase I and II are quite robust. This is both with respect to mean estimates and uncertainty range, with the main exception that the BC RF from FF and BF is substantial stronger in Phase II compared to Phase I.
- The mean from the 16 models of the RF of the total DAE is $-0.27 \text{Wm}^{-2}$. Several of the AeroCom models have not included either nitrate aerosols or SOA or both. Modifying these models with information from AeroCom models altogether results in a mean RF of $-0.32 \text{Wm}^{-2}$. The model simulations are to a large extent performed for the period 1850–2000 and using available information to scale to 1750–2010 results in a mean RF of the total DAE of $-0.35 \text{Wm}^{-2}$.
- The AeroCom Phase II simulated 1850–2000 (four models use slightly different time period) mean RF of the DAE for sulphate is $-0.32 \text{Wm}^{-2}$, BC from FF and BF emissions is $+0.18 \text{Wm}^{-2}$, OA from FF and BF emissions is $-0.03 \text{Wm}^{-2}$, BB is $-0.00 \text{Wm}^{-2}$, nitrate is $-0.08 \text{Wm}^{-2}$ and SOA is $-0.06 \text{Wm}^{-2}$. When these numbers were scaled to 1750–2010 numbers the RF for sulphate is $-0.34 \text{Wm}^{-2}$, BC from FF and BF emissions is $+0.23 \text{Wm}^{-2}$, OA from FF and BF emissions is $-0.05 \text{Wm}^{-2}$, BB is $-0.00 \text{Wm}^{-2}$ (no change included), nitrate is $-0.11 \text{Wm}^{-2}$ and SOA is $-0.08 \text{Wm}^{-2}$.
- The differences between models are large for the DAE of the aerosol components, as illustrated by the relative

![Fig. 17. Model mean RF per component (internal bars), modified from the study timeperiod of 1850–2000 to 1750–2010, based on numbers from Skeie et al. (2011). Details to be given in text.](image-url)
standard deviation of more than 40% for RF of BC and a range from 0.05 to 0.37 W m\(^{-2}\). A further understanding of the causes of the uncertainties is attempted by decomposing the results into diversities due to burden, MEC, and NRF. In general these three factors contribute equally to the uncertainty in the DAE.

- We find a correlation in the magnitude of RF of BC (positive) and RF of sulphate or OA (negative) among the models, due to transport, deposition, and optical properties being treated similarly in each of the models. We therefore find a smaller uncertainty in the total DAE than the sum of the DAE by aerosol components.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys.net/13/1853/2013/acp-13-1853-2013-supplement.pdf.

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