Aerosol absorption and radiative forcing

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Abstract

We present a comprehensive examination of aerosol absorption with a focus on evaluating the sensitivity of the global distribution of aerosol absorption to key uncertainties in the process representation. For this purpose we extended the comprehensive aerosol-climate model ECHAM5-HAM by effective medium approximations for the calculation of aerosol effective refractive indices, updated black carbon refractive indices, new cloud radiative properties considering the effect of aerosol inclusions, as well as by modules for the calculation of long-wave aerosol radiative properties and instantaneous aerosol forcing. The evaluation of the simulated aerosol absorption optical depth with the AERONET sun-photometer network shows a good agreement in the large scale global patterns. On a regional basis it becomes evident that the update of the BC refractive indices to Bond and Bergstrom (2006) significantly improves the previous underestimation of the aerosol absorption optical depth. In the global annual-mean, absorption acts to reduce the short-wave anthropogenic aerosol top-of-atmosphere (TOA) radiative forcing clear-sky from −0.79 to −0.53 W m\(^{-2}\) (33 \%) and all-sky from −0.47 to −0.13 W m\(^{-2}\) (72 \%). Our results confirm that basic assumptions about the BC refractive index play a key role for aerosol absorption and radiative forcing. The effect of the usage of more accurate effective medium approximations is comparably small. We demonstrate that the diversity in the AeroCom land-surface albedo fields contributes to the uncertainty in the simulated anthropogenic aerosol radiative forcings: the usage of an upper versus lower bound of the AeroCom land albedos introduces a global annual-mean TOA forcing range of 0.19 W m\(^{-2}\) (36 \%) clear-sky and of 0.12 W m\(^{-2}\) (92 \%) all-sky. The consideration of black carbon inclusions on cloud radiative properties results in a small global annual-mean all-sky absorption of 0.05 W m\(^{-2}\) and a positive TOA forcing perturbation of 0.02 W m\(^{-2}\). The long-wave aerosol radiative effects are small for anthropogenic aerosols but become of relevance for the larger natural dust and sea-salt aerosols.
1 Introduction

Atmospheric aerosols play an important role in the global climate system through modifications of the global radiation budget: directly, by scattering and absorption of radiation (e.g. Ångström, 1962; McCormic and Ludwig, 1967; Schulz et al., 2006); indirectly, by the modification of cloud properties and abundance (e.g. Twomey, 1974, 1977; Albrecht, 1989; Lohmann and Feichter, 2005); and semi-directly, by the effect of the direct and indirect aerosol effects on cloud properties and abundance via the modification of the thermal structure of the atmosphere and the surface energy budget (Hansen et al., 1997; Ackerman et al., 2000; Lohmann and Feichter, 2005). Aerosol absorption is of particular importance for both direct and semi-direct aerosol effects; however, the magnitude of aerosol absorption on the global scale and its contribution to global warming is subject to considerable uncertainty (Cooke and Wilson, 1996; Cooke et al., 1999; Hansen et al., 2002; Jacobson, 2002; Menon et al., 2002; Chung and Seinfeld, 2002, 2005; Bond and Sun, 2005; Kinne et al., 2006; Roeckner et al., 2006; Stier et al., 2006b; Schulz et al., 2006).

A number of factors complicate the understanding of aerosol absorption, even at the local scale. The radiative properties of an individual particle are determined by the particle composition and the associated component refractive indices as well as by its size and shape (e.g. Bohren and Huffman, 1998). It follows from Mie theory that, to a first order, aerosol extinction per unit aerosol mass is largest when the aerosol radius is comparable to the wavelength of the radiation (e.g. Bohren and Huffman, 1998). This has important implications for natural and anthropogenic aerosol radiative effects. Anthropogenic aerosols are formed predominantly by combustion and gas-to-particle conversion, resulting in average particle sizes at which most of the anthropogenic aerosol mass resides in the sub-micron accumulation mode size-range. By contrast, the predominant sources of natural particles are sea salt and mineral dust, both emitted by wind-driven processes as primary particles with a large contribution in the super-micron coarse mode size ranges (e.g. Seinfeld and Pandis, 2006).
fore, it can be expected that the long-wave aerosol radiative effects will be of primary importance for the larger natural aerosols.

Black carbon (BC), also referred to as “elemental carbon” or “light absorbing carbon”, emitted from incomplete combustion processes, is the dominant absorber of solar radiation under present day conditions, followed by mineral dust (DU) that has typically an imaginary part of the refractive index that is about two orders of magnitude smaller than for BC (e.g. Kinne et al., 2003). While present day BC is predominantly of anthropogenic origin (Ito and Penner, 2005; Stier et al., 2006b), the contribution of anthropogenically disturbed soils to the DU emissions is uncertain (Tegen and Fung, 1995; Mahowald and Luo, 2003; Tegen et al., 2004; Mahowald et al., 2004). The refractive index of representative BC and DU particles is subject to considerable uncertainty (Bond and Bergstrom, 2006; Balkanski et al., 2007), resulting in diverse assumptions in aerosol models and measurement retrievals.

As BC is the predominant anthropogenic aerosol absorber, it is a focal point of this study. Observations show that black carbon particles have complex non-spherical morphologies upon emission, depending on the source type and combustion temperature (Mikhailov et al., 2006; Andreae and Gelenčsér, 2006) affecting their radiative properties (Martins et al., 1998a; van Poppel et al., 2005). Microphysical ageing processes and subsequent mixing with other aerosol components tend to transform these structures to more spherical morphologies, supporting the spherical approximation and application of Mie theory in large-scale modelling (Martins et al., 1998b; Mikhailov et al., 2006) that will also be used throughout this study.

The radiative properties of an aerosol population depend further on the aerosol mixing state, that is the degree to which the chemical components occur as independent particles (external mixing) as compared to a component mixture in each individual particle (internal mixing). Within the framework of Mie theory, internal mixtures are generally represented by effective medium approximations that quantify the macroscopic radiative properties of the mixture as an average over the microscopic arrangement of the components. Practically this is approached through mixing rules that calculate an
effective complex refractive index from the refractive indices of the individual components. A wide range of mixing rules have been developed of which the simple volume-weighted averaging of the component complex refractive indices is most widely used in atmospheric applications. The internal mixing of a population of black carbon with other aerosol components enhances the co-single scattering albedo of the population, and therefore the absorption optical depth, i.e. the column integrated extinction owing to absorption (Ackerman and Toon, 1981; Chýlek et al., 1995; Jacobson, 2000; Stier et al., 2006c). However, in recent work we showed that microphysical ageing processes leading to the internal mixing of initially hydrophobic BC with hydrophilic aerosols, such as sulfate, decrease the atmospheric lifetime and abundance of BC and compensate for the absorption enhancement by internal mixing (Stier et al., 2006c).

The actual atmospheric aerosol absorption over the atmospheric column with given aerosol radiative properties depends on its global spatio-temporal distribution. In particular, the vertical distribution relative to clouds and the surface albedo play a role, as overlying clouds reduce the incoming solar radiation and the surface albedo and underlying clouds regulate the upwelling solar radiation available for absorption. The simulated or retrieved aerosol absorption depends also on the radiative transfer scheme and its spectral resolution, as well as on the assumptions about aerosol effects within clouds.

Both interstitial aerosols and aerosols embedded in cloud droplets and ice crystals affect cloud radiative properties. It has been estimated that absorption from embedded aerosols could be a non-negligible contributor to short-wave cloud absorption, potentially reaching up to 1–3 W m$^{-2}$ (Chýlek et al., 1996) in the global annual-mean. In their lower estimate Chýlek et al. (1996) have found relatively small global impact, in agreement with other studies (Chuang et al., 2002; Liu et al., 2002); however, as pointed out in Chýlek et al. (1996), the underlying assumptions about the BC abundance in their work and the work of Liu et al. (2002) might not be applicable for highly polluted areas.

The effect of atmospheric absorption on the top-of-atmosphere (TOA) aerosol radiative forcing, i.e. net change in the radiative fluxes at TOA introduced by the aerosols,
depends crucially on the effective albedo of the underlying surface. The aerosol single scattering albedo (SSA), i.e. the fraction of radiation extinguished by scattering of the total radiation extinguished by both scattering and absorption, decreases with increasing aerosol absorption. For a certain threshold SSA, the critical single scattering albedo (Haywood and Shine, 1995), the combined aerosol-surface system reflects less solar radiation back to space than the surface of a specific surface albedo alone. In this case the SW TOA forcing becomes positive and the aerosols warm the earth system. It is important to note that optically deep clouds under an aerosol layer also serve effectively as a high albedo surface. Thus, absorbing aerosols above a cloud layer have the potential to introduce positive TOA forcings as evident in the spatial distribution of the all-sky forcing estimates in Schulz et al. (2006).

This complex dependency of aerosol absorption on the aerosol microphysical state, i.e. the composition, size distribution, and mixing state, complicates its simulation in global aerosol models as well as the retrieval from remote sensing instruments.

Early global circulation model based estimates of climatic effects of aerosol absorption focused on specific events such as nuclear winter (Pittock et al., 1986; Turco et al., 1990) or the Kuwait oil fires after 1990 (Bakan et al., 1991). The effects of black carbon on the atmospheric radiation balance and the potential for positive TOA forcing have been conceptually explored by Haywood and Shine (1995). Subsequently, a number of studies investigated the general radiative effects of absorbing carbonaceous aerosols, employing bulk aerosol schemes in global circulation models (Cooke and Wilson, 1996; Liousse et al., 1996; Cooke et al., 1999; Chung and Seinfeld, 2002, 2005). By now, most multicomponent global aerosol models include a bulk representation of the predominant absorbers, black carbon and mineral dust, with prescribed size distributions and assumptions about the aerosol mixing state. A few studies prescribe ageing times from externally-mixed hydrophobic to hydrophilic BC that in some studies is then treated as an internal mixture with sulfate for the radiation calculations. See Textor et al. (2006) for an overview of current aerosol models. A few global aerosol models include a prognostic treatment of the aerosol mixing that allows simulation of...
variations in the BC mixing state under different aerosol emission scenarios (Jacobson, 2001; Stier et al., 2005).

There exist a number of studies in which the global distribution of aerosol absorption has been computed, generally as part of an overall analysis of aerosol radiative forcing. As noted, the absorption portion is likely the most uncertain component of direct aerosol radiative forcing. The objective of the present study is to carry out a comprehensive examination of aerosol absorption. A focus is on evaluating the sensitivity of the global distribution of aerosol absorption to key uncertainties in the process representation. These include aerosol absorbing component refractive indices, surface albedo, refractive indices mixing rules, and the absorption of aerosol inclusions in cloud droplets and ice crystals. Predictions of present day aerosol absorption optical depths are evaluated with observations from the global AERONET sun-photometer network (http://aeronet.gsfc.nasa.gov; Holben et al., 1998; Dubovik and King, 2000; Holben et al., 2001). Another key uncertainty in model based estimates of aerosol absorption are the emissions of carbonaceous aerosols that have been discussed in much detail (Cooke et al., 1996; Lioussse et al., 1996; Cooke, 1997; Cooke et al., 1999; van der Werf et al., 2003; Bond et al., 2004; Schaap et al., 2004; Rao et al., 2005; Ito and Penner, 2005) and will not be considered in this study.

To facilitate this investigation we further extend and evaluate the comprehensive aerosol climate model ECHAM5-HAM (Stier et al., 2005), concentrating on the representation of aerosol absorption. The ECHAM5-HAM model is described in Sect. 2 with a brief summary of the base configuration as in Stier et al. (2005) in Sect. 2.1 and its extension for this study in Sect. 2.2. The base and sensitivity simulations are described in Sect. 3. Results and discussion are presented in Sect. 4 followed by conclusions in Sect. 5.
2 Model description

In this study we employ the aerosol-climate model ECHAM5-HAM with a prognostic representation of the composition, size distribution, and mixing state of the major global aerosol components: sulfate, black carbon, particulate organic matter, sea salt, and mineral dust (Stier et al., 2005). A brief description of ECHAM5-HAM, employed in its version echam_5.3.01.ham.1.51, is presented in the first part of this section. More details and an extensive evaluation of this base model can be found in Stier et al. (2005). The extensions of the representation of aerosol absorption in ECHAM5-HAM are given in the second part of the section.

2.1 ECHAM5-HAM

2.1.1 The atmospheric general circulation model ECHAM5

The atmospheric general circulation model (GCM) ECHAM5 (Roeckner et al., 2003) is the fifth-generation climate model developed at the Max Planck Institute for Meteorology. ECHAM5 solves prognostic equations for vorticity, divergence, surface pressure, and temperature, expressed in terms of spherical harmonics with a triangular truncation. Non linear processes and the physical parameterisations are solved on a corresponding Gaussian grid. Water vapour, cloud liquid water, cloud ice, and trace components are transported in grid-point space with a flux form semi-Lagrangian transport scheme (Lin and Rood, 1996). ECHAM5 contains a microphysical cloud scheme (Lohmann and Roeckner, 1996) with prognostic equations for cloud liquid water and ice. Cloud cover is predicted with a prognostic-statistical scheme solving equations for the distribution moments of total water (Tompkins, 2002). Convective clouds and convective transport are based on the mass-flux scheme of Tiedtke (1989) with modifications by Nordeng (1994). The short-wave radiation scheme (Fouquart and Bonnel, 1980) has currently 4 spectral bands, on for the visible and ultra-violet, and three for the near-infrared. The long-wave radiation scheme (Mlawer et al., 1997; Morcrette et al., 1998)
1998) has 16 spectral bands.

2.1.2 The aerosol module HAM

The microphysical aerosol module HAM (Stier et al., 2005) predicts the evolution of an ensemble of seven interacting internally- and externally-mixed log-normal aerosol modes. In the current setup, the components comprise: sulfate, black carbon, particulate organic matter, sea salt, and mineral dust. The aerosol mixing state is prognosed within the possible mixing state configurations illustrated in Table 1. The modes are composed either of hydrophobic compounds or of an internal mixture of hydrophobic and hydrophilic compounds, henceforth denoted as hydrophilic modes. The main components of HAM are the microphysical core M7, a sulfur chemistry scheme, a module to calculate the aerosol radiative properties, a deposition module, and an emission module.

The microphysical core M7 (Vignati et al., 2004) calculates coagulation among the modes, condensation of gas-phase sulfuric acid on the aerosol surface, binary nucleation of sulfate, and water uptake.

The sulfur cycle model (Feichter et al., 1996) treats the prognostic variables dimethyl sulfide (DMS), sulfur dioxide, sulfate and their gas- and aqueous-phase reaction pathways, using pre-calculated monthly mean oxidant fields.

Aerosol radiative properties, as well as the sink processes dry deposition, sedimentation, and wet deposition, are parameterised in dependence on the prognostic aerosol size distribution, composition, and mixing state and coupled to the ECHAM5 meteorology.

Aerosol radiative properties are calculated in the framework of Mie theory. For each aerosol mode, effective refractive indices are calculated by volume-averaging the refractive indices of all components, including aerosol water, which is parameterised in terms of ambient relative humidity. The effective complex refractive indices and the Mie size-parameters for each mode serve as input to look-up tables for the aerosol radiative properties, providing extinction cross-section, single scattering albedo, and asymmetry.
parameter to the ECHAM5 radiation scheme. To allow the investigation of a wide range of refractive indices we extended the upper boundary of the range of the imaginary part of the refractive index in the look-up tables from 0.5 \( i \) to 1.0 \( i \) for this study.

Aerosol wet deposition is parameterised in terms of the aerosol size distribution and mixing state via mode-specific scavenging ratios, specifying embedded and interstitial aerosol fractions in the cloudy part of a grid box and in convective updrafts. The actual wet deposition is calculated from the resulting embedded aerosol content based on the precipitation formation and re-evaporation calculated by the ECHAM5 cloud scheme. Aerosol and gas dry deposition velocities are calculated based on a serial resistance approach (Stier et al., 2005; Ganzeveld et al., 2006).

Emissions of mineral dust are calculated online depending on the ECHAM5 10-m wind speed, soil moisture, and snow cover (Tegen et al., 2002, 2004). Freshly emitted BC and mineral dust are assumed hydrophobic. Sea salt emissions are parameterised in terms of the simulated 10-meter wind speed and sea-ice cover (Schulz et al., 2004). Emissions of DMS are calculated interactively from DMS seawater concentrations of Kettle and Andreae (2000) depending on the simulated 10-meter wind speed and sea surface temperature, applying the sea-air exchange formulation of Nightingale et al. (2000). Terrestrial biogenic DMS emissions are prescribed (Pham et al., 1995). For all other compounds, emission strength, distribution, and height are based on the AeroCom aerosol model inter-comparison (http://nansen.ipsl.jussieu.fr/AEROCOM/) emission inventory representative for the years 2000 and 1750 (Dentener et al., 2006). The emissions for the years 1750 and 2000 are summarised in Table 2.

### 2.2 Extensions of ECHAM5-HAM

#### 2.2.1 Mixing rules for refractive indices

The prognostic treatment of the aerosol mixing state requires the interactive computation of the aerosol radiative properties from the composition of each mode. In standard HAM, the effective refractive index of each mode is obtained by volume-weighted aver-
aging of the refractive index of its components. However, as discussed in Chýlek et al. (1995) and Lesins et al. (2002), the volume-weighted mixing of the refractive indices for strongly absorbing inclusions, such as BC, in a less absorbing medium, such as sulfate, results in an overestimate of the effective imaginary part of the refractive index as compared to an average of explicit calculations over all possible positions of an inclusion in the host medium. A more accurate approximation for effective refractive indices can be obtained from mixing rules derived from effective medium approaches, quantifying the macroscopic radiative properties as an average over the microscopic arrangement of the components.

A general formulation for mixing rules of *n* components can be expressed as (Aspnes, 1982):

\[
\frac{\epsilon_{\text{eff}} - \epsilon_0}{\epsilon_{\text{eff}} + 2\epsilon_0} = \sum_{k=1}^{n} f_k \frac{\epsilon_k - \epsilon_0}{\epsilon_k + 2\epsilon_0}
\]  

(1)

where \(\epsilon_{\text{eff}}\) is the complex effective dielectric constant of the mixture, \(\epsilon_0\) is the complex dielectric constant of the host medium, and \(\epsilon_k\) and \(f_k\) are the complex dielectric constant and volume fraction of component *k*, respectively.

The appropriate mixing rule for the inclusion of an insoluble core in a liquid host medium is the Maxwell-Garnet mixing rule (Garnett, 1904, 1906), see Chýlek et al. (2000) and Lesins et al. (2002). However, it requires the explicit choice of a host and an inclusion medium. As the volume fractions of the components in the internally-mixed aerosol modes of HAM are highly variable and the modes can be composed of mixtures of either dry or solid components, the application of the Maxwell-Garnet mixing rule is not straightforward. The mixing rule of Bruggeman (1935), best suited for internal mixtures of solid compounds (see Chýlek et al., 2000; Lesins et al., 2002), does not require the explicit definition of host and inclusion media. The resulting real and imaginary parts of the refractive indices obtained from the Bruggeman mixing rule lie in-between the results of the two possible choices of host and inclusion media in the Maxwell-Garnet mixing rule (Chýlek et al., 2000).
In the Maxwell-Garnet mixing rule one of the mode components \((k=1)\) is chosen as the host medium, i.e. \(\varepsilon_1=\varepsilon_0\), so that the first term of the sum in Eq. (1) vanishes and the system reduces to:

\[
\frac{\varepsilon_{\text{eff}} - \varepsilon_1}{\varepsilon_{\text{eff}} + 2\varepsilon_1} = \sum_{k=2}^{n} f_k \frac{\varepsilon_k - \varepsilon_1}{\varepsilon_k + 2\varepsilon_1}
\]

so that

\[
\varepsilon_{\text{eff}} = \varepsilon_1 \frac{1 + 2 \sum_{k=2}^{n} f_k \frac{\varepsilon_k - \varepsilon_1}{\varepsilon_k + 2\varepsilon_1}}{1 - \sum_{k=2}^{n} f_k \frac{\varepsilon_k - \varepsilon_1}{\varepsilon_k + 2\varepsilon_1}}
\]

For the Bruggeman mixing rule no choice of a host medium is required as it is assumed that the components are embedded in a host medium of the effective dielectric constant, i.e. \(\varepsilon_0=\varepsilon_{\text{eff}}\), so that the left-hand side in Eq. (1) vanishes and the system is reduced to the implicit equation for \(\varepsilon_{\text{eff}}\):

\[
f(\varepsilon_{\text{eff}}) = \sum_{k=1}^{n} f_k \frac{\varepsilon_k - \varepsilon_{\text{eff}}}{\varepsilon_k + 2\varepsilon_{\text{eff}}} = 0
\]

2.2.2 Calculation of effective aerosol refractive indices employing the Bruggeman mixing rule

As the component volume fractions for the modes of ECHAM5-HAM are highly variable, we employ in this study the Bruggeman mixing rule as an alternative method for the calculation of the mode effective complex refractive indices \((n_{\text{eff}}=\sqrt{\varepsilon_{\text{eff}}})\) in ECHAM5-HAM. (The Maxwell-Garnet mixing rule has also been implemented but requires somewhat arbitrary choices of host and inclusion media.) Using

\[
f'(\varepsilon_{\text{eff}}) = \sum_{k=1}^{n} f_k \frac{-3\varepsilon_k}{(\varepsilon_k + 2\varepsilon_{\text{eff}})^2}
\]
we solve Eq. (4) with a Newton iteration over the index $\nu$:

$$e^{\nu+1}_{\text{eff}} = e^{\nu}_{\text{eff}} - \frac{f(e^{\nu}_{\text{eff}})}{f'(e^{\nu}_{\text{eff}})}$$

The number of required iterations for convergence was determined not to exceed six iterations at any grid point in a one-year test simulation, so that a fixed number of seven iterations is applied globally.

2.2.3 Parameterisation of the effect of aerosol inclusions on cloud radiative properties employing the Maxwell-Garnett mixing rule

To facilitate the investigation of aerosol inclusions on the cloud radiative properties, we recompute the standard ECHAM5 cloud radiative properties (Roeckner et al., 2003; Rockel et al., 1991) including aerosol effects. Analogous to the parameterisation of the aerosol radiative properties, we tabulate the cloud radiative properties in a look-up table as a function of the effective refractive index of the cloud droplets with aerosol inclusions.

In standard ECHAM5, short-wave cloud radiative properties are parametrised from idealised size-distributions (Rockel et al., 1991) under the assumption of spherical cloud droplets and ice crystals. From a series of Mie calculations, fits are derived to express the resulting radiative properties of cloud droplets and ice crystals in terms of the effective radius (Roeckner et al., 2003) that is empirically related to the radius of average volume by Johnson (1993) for water droplets and by Moss (personal communication as in Roeckner et al., 2003) for ice crystals.

For the extended parameterisation we repeated the Mie calculations of the cloud radiative properties for varying cloud droplet or ice crystal refractive indices. For each complex pair of the refractive indices, we use Mie calculations (Toon and Ackerman, 1981) to compute, as in ECHAM5, radiative properties for cloud droplets and ice crystals as an average over idealised gamma size-distributions following Hansen and Travis.
\[
 n(r) = C r^{1-3\nu_{\text{eff}}} e^{-r/\nu_{\text{eff}}} 
\]

(7)

where \( C \) is a constant and \( \nu_{\text{eff}} \) is the effective variance of the distribution, with the specifications given in Table 3 based on Rockel et al. (1991).

The computed radiative properties for cloud droplet and ice crystal extinction cross section, single scattering albedo, and asymmetry factor are provided in a look-up table with three dimensions: Mie effective size-parameter \( (\chi_{\text{eff}} = 2\pi r_{\text{eff}}/\lambda) \), as well as real and imaginary parts \( (n_r \text{ and } n_i) \) of the effective refractive index of the cloud water – aerosol mixture.

The standard ECHAM5 cloud droplet concentration \( N_l \) is prescribed within the boundary layer as \( 220 \times 10^6 \text{ m}^{-3} \) over land and as \( 80 \times 10^6 \text{ m}^{-3} \) over sea. Above the boundary layer, \( N_l \) decreases exponentially to \( 50 \text{ m}^{-3} \) in the upper troposphere over both land and ocean. The cloud ice crystal number concentration is calculated from the ice water mixing ratio via an empirical parameterisation of the effective radius (Moss, 1996, personal communication as in Roeckner et al., 2003). In an extended version of ECHAM5-HAM, the standard bulk stratiform cloud scheme of ECHAM5 (Lohmann and Roeckner, 1996) has been replaced by a microphysical cloud parameterisation with a prognostic treatment of the cloud droplet number concentration and ice crystal number concentration (Lohmann et al., 2007).

Contrary to standard ECHAM5-HAM, in which all aerosol within the cloudy fraction of a grid box is assumed interstitial with identical radiative properties as for clear-sky, the in-cloud aerosol content is split into interstitial aerosol and cloud droplet/crystal inclusions that are considered for the cloud radiative properties. For the calculation of the cloud droplet and ice crystal effective refractive indices, the aerosol content associated with water droplet / ice crystal inclusions is calculated consistently from the prescribed ECHAM5-HAM scavenging ratios (Stier et al., 2005), defined as the fraction of aerosol embedded in cloud droplets or ice crystals within the cloudy fraction of a grid box. The embedded aerosol content is assumed uniformly distributed over all cloud droplets and

7184
ice crystals. The volume fraction of the aerosol inclusions can be considered small for cloud droplets and ice crystals so that the Maxwell-Garnett mixing rule (Eq. 3) is employed for the calculation of the effective dielectric constants, from which the complex refractive index is derived by taking the complex root $n_{\text{eff}} = \sqrt{\varepsilon_{\text{eff}}}$. For the case of pure water clouds, as applied in the CL-INT sensitivity study (see Sect. 3), the aerosol volume fractions in Eq. (3) are zero and the effective refractive index reduces to the host medium value. While all aerosol components could be considered for the calculation of the cloud droplet/crystal refractive indices via Eq. (3), here we activate only the effect of BC on the cloud radiative properties, as the orders of magnitude larger BC imaginary part of the refractive index will dominate the total aerosol effect.

2.2.4 Long-wave aerosol radiative properties

The radiation module of ECHAM5-HAM has been extended to also cover the long-wave radiative properties of aerosol, in analogy to the original short-wave radiation module (Stier et al., 2005). The long-wave aerosol radiative properties are pre-computed from Mie theory for the 16 long-wave spectral bands of the Mlawer et al. (1997); Morcrette et al. (1998) radiation scheme of ECHAM5 and supplied in look-up tables with the three dimensions: Mie median size-parameter ($X_g = 2\pi r_g/\lambda$), and real and imaginary parts of the refractive index. Hereby, $r$ is the number median radius of the log-normal mode and $\lambda$ is the wavelength. Effective refractive indices of internally-mixed modes are calculated by volume-weighted averaging of the component refractive indices (default) or by the Bruggeman mixing rule as described in 2.2.1. References for the implemented long-wave refractive indices are given in Table 4, and the spectral dependence of the refractive indices is depicted in Fig. 1.

2.2.5 Instantaneous forcing calculation

We have extended ECHAM5-HAM by a module for the calculation of instantaneous direct aerosol radiative forcing (as already applied in Schulz et al., 2006). Due to the
internal mixing of mode components and the resulting nonlinear effects of individual components on the aerosol microphysical state and radiative properties (Stier et al., 2006c), it is not possible to directly calculate radiative forcing for individual components. Instead, we calculate total aerosol radiative forcing by a double call of the ECHAM5 radiation scheme, without and with the radiative properties of the total aerosols. The instantaneous direct aerosol radiative forcing is then computed as the difference in the separate clear-sky and all-sky, short-wave and long-wave net radiative fluxes, without and with aerosols. No adjustment of stratospheric temperatures is performed in the forcing calculations.

3 Simulation setup

To investigate aerosol absorption and its radiative effects we perform a series of sensitivity studies with the ECHAM5-HAM model in which we vary key uncertainties in model parameterisations and compare the results to observations as well as the Stier et al. (2005) base simulation.

For each simulation we run ECHAM5-HAM from October 1999 to December 2000 and constrain the large-scale meteorology to the year 2000 by nudging (Jeuken et al., 1996) the model to the ECMWF ERA40 reanalysis data (Simmons and Gibson, 2000). Only the year 2000 data are analysed. We employ a horizontal resolution of T63 in spectral space with a corresponding resolution of 1.8° × 1.8° on a Gaussian grid. The vertical resolution is set to 31 levels, extending from the surface up to 10 hPa. Aerosol and aerosol precursor emissions are identical in all simulations, except for small variations in the interactive sources owing to small variations in the local scale meteorology. To isolate the effect of anthropogenic emissions, each study is performed twice with emissions from the AeroCom emission inventory (Dentener et al., 2006) representative for the years 2000 and 1750. An overview of the simulations performed is given in Table 5.

The first series of simulations (BASE, BB-M, BB-H) is aimed at exploring the sensi-
tivity of the aerosol absorption and radiative forcing to uncertainties in the BC refractive indices. The BASE simulation is physically identical to that of Stier et al. (2005), however, has been performed with an updated model version, as described in Sect. 2.

In the BB-M and BB-H simulations we employ the medium and highly absorbing BC refractive indices from Bond and Bergstrom (2006), respectively, as given in Table 6. As Bond and Bergstrom (2006) report the BC refractive indices only at a wavelength of 550 nm, we use the same spectral dependence as in BASE (Hess et al., 1998) to spectrally extrapolate BC refractive indices for BB-M and BB-H.

Variations in the aerosol absorption have been attributed partly to the calculation of the effective refractive indices for internally mixed aerosols, in particular for internal mixtures of BC (Chýlek et al., 1995; Lesins et al., 2002; Jacobson, 2000). To quantify the effect of mixing rules on global atmospheric absorption and radiative forcing we replace in the BRUG simulation the most commonly applied volume-weighted averaging applied in all other simulations with the more elaborate Bruggeman (1935) mixing rule.

Variations in the TOA forcing of absorbing aerosols are often partly attributed to differences in the assumed surface albedos (e.g. Schulz et al., 2006). To quantify this effect, we derive upper and lower estimates of the global surface albedo distribution, as applied in current global aerosol models, and apply those fields in ECHAM5-HAM. We derive these upper and lower bounds from the surface albedo fields submitted by the GISS (Koch et al., 2006), ECHAM5-HAM (Stier et al., 2005), LOA (Reddy et al., 2005), LSCE (see Textor et al., 2006), UIO-GCM (Kirkevåg and Iversen, 2002; Kirkevåg et al., 2005; Iversen and Seland, 2002, 2003), UMI (Liu et al., 2005), and SPRINTARS (Take-mura et al., 2005) global aerosol models to the AeroCom forcing experiment (Schulz

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1 For SPRINTARS, the AeroCom database contained the visible instead of the broad-band short-wave surface albedo. This field was used in the calculation of the upper and lower bound of the broad-band surface albedos. However, a reanalysis of the derived upper and lower bounds without SPRINTARS resulted in almost indistinguishable lower and upper bounds with global annual means of 0.19 and 0.35, respectively. Therefore, we decided not to repeat the sensitivity studies with the corrected fields.
et al., 2006). At each grid point of ECHAM5-HAM we select the highest and lowest value of the surface albedo provided by any of the AeroCom models. The resulting upper and lower bounds of the AeroCom surface albedo distributions are depicted in Fig. 2. Please note that the protocol of the AeroCom forcing experiment specified albedo broadly as “surface solar broadband albedo (0.2–4.0 μm)”. Therefore, some of the submissions provide the prescribed background surface albedo while others provide the applied effective surface albedo, including the contributions from interactively computed snow and ice covers. This might cause a high bias of the upper bound of the surface albedo estimate at high latitudes. For the sensitivity studies with ECHAM5-HAM, we use the the derived lower (ALB-L) and upper (ALB-H) bounds to replace the prescribed background ECHAM5 land surface albedo (Hagemann, 2002) used in all other simulations. The background albedo is modified during the simulation based on the simulated snow cover. We do not modify the original sea and lake surface albedo of ECHAM5, which is set to a constant value of 0.07 (Roeckner et al., 2003).

To quantify the contribution of aerosol absorption to aerosol radiative forcing, we perform a study with transparent aerosols (TRANS), i.e. with the imaginary parts of all aerosol refractive indices set to $n_i=10^{-9}$.

The effect of aerosol inclusions on short-wave cloud absorption and the resulting radiative effects are investigated in the CL-INC, CL-INT, and CL-EXT simulations. All employ the new cloud radiative properties described in Sect. 2.2.3. CL-INC is set up as BB-M but with in-cloud aerosol split in interstitial aerosol and cloud droplet/crystal inclusions that are considered in the calculation of the cloud radiative properties via the Maxwell-Garnett mixing rule. CL-INT is identical to CL-INC but considers only the interstitial aerosol and not the effect of cloud droplet/crystal inclusions on the new cloud radiative properties. In CL-EXT, all in-cloud aerosol is considered interstitial, i.e. externally mixed with the cloud droplets.
4 Results

In this section we will systematically evaluate the sensitivity of aerosol absorption and radiative forcing to key uncertainties in aerosol modelling.

The general setup of our simulations is identical to that in Stier et al. (2005), and the sensitivity studies performed differ only in the representation of the aerosol and surface radiative properties, so that the extensive evaluation of ECHAM5-HAM in Stier et al. (2005) does also apply for this study. As the present study focuses on the representation of aerosol absorption, the changes in all simulations affect only minimally the global aerosol optical depth as compared to Stier et al. (2005), with a global annual mean value of 0.14.

4.1 Total aerosol absorption and its evaluation

The simulated global distribution of clear-sky total aerosol absorption optical depth (AAOD), its seasonal evaluation with AERONET sun-photometer measurements, and the simulated clear-sky total aerosol absorption for the BASE, BB-M, BB-H, and BRUG scenarios are depicted in Figs. 3–6, respectively. For the AAOD evaluation at each AERONET site, the simulated daily mean AAOD values at the model grid-box containing the site have been sampled at the available measurement dates. For the BB-M sensitivity study, all included daily-mean AAODs are additionally given as seasonally colour-coded scatterplots for each AERONET site in Appendix A, Fig. A1. The same plots for the BASE, BB-H, and BRUG study can be found in the supplementary online material (http://www.atmos-chem-phys-discuss.net/7/7171/2007/acpd-7-7171-2007-supplement.pdf), SOM-Figs. 1–3. Aerosol radiative properties and forcings for all simulations are given in Table 7.

The global distribution of the AAOD and aerosol absorption shows maxima in the biomass burning regions of Central Africa and South America, in the Saharan dust plume, and in the regions with high anthropogenic activities over India and South East Asia, as well as secondary maxima over the Eastern U.S. and Central to Eastern Eu-
rope. For all simulations, the AERONET evaluation of AAOD shows generally a good agreement in these large-scale global patterns. Also the seasonality is well captured at many AERONET sites, such as the July to September onset of the Southern Hemispheric biomass burning season in South America and Africa.

On a regional basis, general and seasonal biases as well as differences among the simulations become more clearly evident. For example, AAODs at the urban Mexico City site (99.2 W, 19.3 N) as well as over the central European site Lille (3.1 E, 50.6 N) are generally underestimated, and the AAOD increase during the Asian spring dust-season at the Mongolian Dalanzadgad site (104.4 E, 43.6 N) is not captured. The BASE simulation has a tendency to generally underestimate AAOD, in particular over the European and US source regions. The increase of the imaginary part of the BC refractive index from BASE to BB-M and BB-H (Table 6), generally improves the agreement with the AERONET retrieved AAOD. The incomparability of different stations and the consideration of a highly variable number of measurements at each site (Fig. 5) do not seem to allow deriving a global measure of the overall model performance. However, from comparison of Figs. 3–6 and SOM-Figs. 1–3 we conclude that the BB-M (and the very similar BRUG) case seems to provide the best representation of the AERONET retrieved AAOD. Thus, the BB-M simulation will be used as a reference for comparison with the other sensitivity studies.

4.2 Anthropogenic aerosol absorption and radiative forcing

4.2.1 BASE case

The simulated short-wave anthropogenic aerosol absorption and TOA radiative forcings are depicted in Fig. 7; corresponding global annual-mean total and anthropogenic aerosol radiative properties and forcings are given in Table 7.

For the BASE case, as submitted to the AeroCom forcing experiment (Schulz et al., 2006), the global annual mean short-wave clear- and all-sky anthropogenic aerosol absorption are 0.81 W m$^{-2}$ and 0.74 W m$^{-2}$, these are 46% and 45% of the total
Aerosol absorption generally reduces the amount of solar radiation reflected back to space. Over high surface albedo areas the combined aerosol-surface system reflects less solar radiation back to space than the high albedo surface alone, resulting in a positive TOA forcing and a warming effect. This can be observed in the clear-sky TOA aerosol radiative forcing over North Africa and West Australia. In the all-sky case, clouds below absorbing aerosols serve as an effective high albedo surface so that large regions over South America and regions with persistent fields of stratocumulus clouds of the west coast of Southern Africa show positive TOA aerosol radiative forcings. The global annual-mean short-wave clear-sky and all-sky anthropogenic TOA aerosol radiative forcings are $-0.65 \text{ W m}^{-2}$ and $-0.27 \text{ W m}^{-2}$, respectively.

4.2.2 Sensitivity to BC refractive indices

To investigate the sensitivity of aerosol absorption and radiative forcing to the BC refractive index we increase the imaginary part of the BC refractive index from 0.44 in BASE to the medium and high recommendations by Bond and Bergstrom (2006) of 0.71 in BB-M and 0.79 in BB-H (at $\lambda = 550 \text{ nm}$). As evident in Fig. 7, these changes increase the anthropogenic short-wave aerosol absorption from 0.81 to 1.20 and 1.31 $\text{ W m}^{-2}$ in the clear-sky and from 0.74 to 1.11 and 1.21 $\text{ W m}^{-2}$ in the all-sky case, respectively. It is interesting to note that while this increase of absorption reduces the TOA forcing from $-0.65$ to $-0.53$ ($-19\%$) and $-0.50 \text{ W m}^{-2}$ ($-23\%$) for the clear-sky case, the all-sky forcing decreases from $-0.27$ to $-0.13$ ($-52\%$) and $-0.09 \text{ W m}^{-2}$ ($-67\%$). This highlights the strong sensitivity of the all-sky TOA aerosol radiative forcing to aerosol absorption and the crucial role of the relative location of absorbing aerosols and clouds. For the BB-M study, which shows the best agreement of AAOD with the AERONET sunphotometer network, the TOA aerosol radiative forcing becomes positive over large regions of Africa, South America, Asia and Australia.
4.2.3 Sensitivity to refractive index mixing rules

In the BRUG study, we replace the simple volume-weighted averaging calculation of the effective refractive indices for internally mixed modes in BB-M by the Bruggeman (1935) mixing rule. As shown in Table 7, this change decreases the total aerosol clear-sky and all-sky absorption from 2.35 to 2.12 W m$^{-2}$ (–10%) and from 2.15 to 1.95 W m$^{-2}$ (–9%), respectively. This relatively small change of the aerosol absorption, as compared to the change associated with the variations of the BC refractive indices from BASE to BB-H, does not allow one to identify significant improvements in the evaluation of AAOD with the AERONET sun-photometer network between BB-M (Fig. 4) and BRUG (Fig. 6). The anthropogenic aerosol clear-sky and all-sky TOA radiative forcings intensify from BB-M to BRUG from –0.53 to –0.58 W m$^{-2}$ (+9%) and from –0.13 to –0.18 W m$^{-2}$ (+39%), respectively.

4.2.4 Sensitivity to surface albedo

The large gradients in the TOA forcing at discontinuities of the surface albedo fields, such as the coasts of Africa, readily suggest a strong dependence of aerosol radiative forcing on the surface albedo. To quantify the effect of common assumptions for the surface albedo on the aerosol absorption and radiative forcing, we replace the land surface albedo field applied in BB-M by the upper (ALB-H) and lower (ALB-L) bound of the reported surface albedos in the AeroCom forcing experiment, as described in Sect. 3. The resulting annual-mean difference in the broad-band land surface albedo between the simulations ALB-H and ALB-L is shown in Fig. 8. It is clearly evident that significant surface albedo difference exist between the two simulations, reaching values of 0.5. Interestingly, the differences are not limited to high latitude areas where different assumptions in snow covers and albedos could play a role in the AeroCom upper bound of the surface albedo. Instead, differences of up to about 0.4 occur over all continents with the largest absolute differences over the high albedo desert areas. The global annual-mean difference is 0.04 and the difference averaged over the land
areas is 0.13.

The resulting modification of the upwelling surface short-wave radiative fluxes cause small changes in the clear- and all-sky anthropogenic aerosol absorption of less than 0.05 W m\(^{-2}\) (Table 7). However, the corresponding anthropogenic TOA aerosol radiative forcing changes from −0.53 in BB-M to −0.61 (+15 %) in ALB-L and −0.42 W m\(^{-2}\) (−21 %) in ALB-H for the clear-sky case and the all-sky forcing change from −0.13 in BB-M to −0.18 (+39 %) in ALB-L and −0.06 W m\(^{-2}\) (−54 %) in ALB-H (Table 7 and Fig. 7). The difference of clear-sky and all-sky anthropogenic direct aerosol radiative forcings between the ALB-H and ALB-L simulations is shown in Fig. 9. Over large regions, such as East Asia, the differences of the clear-sky TOA anthropogenic forcing due to the use of different surface albedo datasets reach up to 2–5 W m\(^{-2}\), locally up to 10 W m\(^{-2}\). The global annual-mean difference in the anthropogenic TOA aerosol radiative forcing between the cases with the upper and the lower bounds of the AeroCom land-surface albedo is 0.20 W m\(^{-2}\) (clear-sky) and 0.12 W m\(^{-2}\) (all-sky), respectively.

4.3 Contribution of aerosol absorption to radiative forcing

The difference between the BB-M and TRANS simulations allows one to isolate the effect of aerosol absorption on radiative forcing. Unlike studies with absorbing aerosols omitted, this method does not alter the aerosol microphysical state by changing the number of emitted primary particles that can act as seeds for secondary particle growth (Stier et al., 2006a; Pierce and Adams, 2007).

The comparison of the global patterns of short-wave anthropogenic aerosol radiative forcing in Fig. 7 shows the very similar distribution of negative aerosol radiative perturbations in the main anthropogenic source regions. However, without aerosol absorption the short-wave TOA forcings are universally negative and the positive forcings over extended regions in BB-M change their sign in TRANS. In the global annual mean, absorption acts to reduce the negative TOA short-wave anthropogenic aerosol forcing of BB-M by 0.26 W m\(^{-2}\) for the clear-sky and 0.34 W m\(^{-2}\) for the all-sky case, as indicated by the differences between the TRANS and BB-M cases in Table 7.
4.4 Long-wave aerosol radiative effects

The simulated instantaneous year 2000 mean long-wave total aerosol radiative forcings from the BASE simulation are given in Fig. 10 and Table 7. The largest long-wave atmospheric aerosol net absorption occurs in the high-latitude storm track regions, owing to large sea salt and associated aerosol water burdens, and in the Saharan dust plume. The wind driven sea salt emissions are spatio-temporally correlated with storms and the associated clouds and water vapour. As the long-wave radiation in these regions is predominantly attenuated by the water vapour and cloud water, the all-sky long-wave aerosol absorption in the storm tracks becomes smaller compared to long-wave absorption in the predominantly clear-sky dust plume. Dust storms frequently lift mineral dust to high altitudes (e.g. Colarco et al., 2003), resulting in a higher mass fraction of DU at high altitudes than for sea salt (see Fig. 11 in Textor et al., 2006). Owing to the colder temperatures at high altitudes, it follows from Stefan-Boltzmann’s law that less of the absorbed long-wave radiation is re-emitted than for low-altitude absorption that effectively re-emits at higher temperatures, similar to the underlying surface. Consequently, the long-wave aerosol radiative forcing at the TOA shows a maximum in the Saharan dust plume, reaching about 2–5 W m\(^{-2}\) in the annual mean. On a monthly mean basis, clear-sky long-wave TOA aerosol radiative forcings in the Saharan dust source region reach about 10–20 W m\(^{-2}\) (not shown), in good agreement with the satellite-derived estimate by Zhang and Christopher (2003). It needs to be emphasised that the present setup of ECHAM5-HAM does not include the super-coarse modes of mineral dust and sea salt (see Stier et al., 2005) so that the long-wave aerosol radiative effect of natural aerosols could be underestimated. However, owing to the short lifetime of super-coarse particles this effect is probably limited to the source regions.

As expected from theory, the simulated annual-mean long-wave anthropogenic aerosol absorption (0.03 W m\(^{-2}\)) and TOA radiative forcing (0.02 W m\(^{-2}\)) are small (Fig. 11, note the different scales) compared to the short-wave absorption and radia-
tive forcings. Maxima of anthropogenic absorption and TOA forcing reach about 0.2 and 1 W m$^{-2}$ in the main anthropogenic source regions, respectively.

4.5 Cloud radiative properties considering the effects of aerosol inclusions

Simulated aerosol and aerosol-cloud short-wave absorption optical depths from the simulation with the consideration of interstitial aerosols and black carbon inclusions on the cloud radiative properties CL-INC, the simulation with interstitial in-cloud aerosols but without the black carbon inclusion effect CL-INT, and the simulation with all in-cloud aerosol considered interstitial CL-EXT are given in Fig. 12. While the cloud absorption optical depth, i.e. the absorption optical depth of embedded aerosols, for pure water clouds in CL-INT and CL-EXT is negligible, the consideration of black carbon inclusions for the cloud radiative properties in CL-INC results in an inclusion AAOD of $0.6 \times 10^{-3}$. The consideration of all in-cloud aerosol as interstitial increases the interstitial AAOD in CL-EXT to $0.3 \times 10^{-3}$ as compared to $0.2 \times 10^{-3}$ in CL-INT and CL-INC.

From the simulated short-wave total aerosol atmospheric absorption differences between CL-INC and CL-INT given in Fig. 13, it is evident that the absorption by BC inclusions results in an all-sky absorption of 0.05 W m$^{-2}$, with maxima in regions with high inclusion AAODs. However, the absorption enhancement over the fully externally mixed in-cloud aerosol population through comparison with the the difference of CL-EXT – CL-INT is very small.

The resulting short-wave TOA total aerosol forcing perturbation owing to BC inclusions, shown in Fig. 13 as the difference between CL-INC and CL-INT, is small with a global annual-mean of 0.02 W m$^{-2}$. The TOA forcing perturbation owing to the consideration of all aerosols as an external mixture in CL-EXT as opposed to the case of only the interstitial residual in CL-INT, given as the difference between CL-EXT and CL-INC, shows small positive values in the regions with high AAODs and small negative values over regions with high hydrophilic aerosol burdens. This can be explained by the fact that the effect of non BC aerosol components, such as sea salt, on the cloud radiative properties is generally considered small and was thus neglected in CL-INT.
while their effect is included in the fully externally mixed interstitial aerosol in CL-EXT. Thus, the enhanced scattering due to interstitial sea salt in CL-EXT results in the small negative forcing perturbation over the oceans.

5 Conclusions

We have presented a comprehensive examination of aerosol absorption with a focus on evaluating the sensitivity of the global distribution of aerosol absorption to key uncertainties in the process representation.

For this purpose we employed the aerosol-climate model ECHAM5-HAM with prognostic representation of the aerosol composition, size-distribution, and mixing-state (Stier et al., 2005) and extended it by:

- Alternative refractive indices for black carbon (Bond and Bergstrom, 2006) and extended look-up tables of the short-wave aerosol radiative properties covering a wider refractive index range.

- Alternative methods for the calculation of effective refractive indices of internally mixed modes based on the Bruggeman and Maxwell-Garnett effective medium approaches.

- A new look-up table approach for the ECHAM5 cloud radiative properties, considering the effect of aerosol inclusions in cloud droplets and ice crystals employing the Maxwell-Garnett effective medium approach.

- A long-wave aerosol radiation module in analogy to the short-wave look-up table approach.

- A module for the calculation of instantaneous aerosol forcing.

- Alternative lower- and upper-bound land surface-albedo datasets derived from the AeroCom aerosol model inter-comparison forcing experiment submissions.
For our investigation of aerosol absorption we performed pairs of present day (2000) and pre-industrial (1750) simulations for BASE: as in Stier et al. (2005), BB-M: updated medium absorbing refractive indices from Bond and Bergstrom (2006), BB-H: updated highly absorbing refractive indices from Bond and Bergstrom (2006), BRUG: as BB-M but with Bruggeman (1935) mixing rule for effective refractive indices, ALB-L: as BB-M but employing a lower bound of currently used land-surface albedos derived from AeroCom, ALB-H: as BB-M but employing a higher bound of currently used land-surface albedos derived from AeroCom, CL-INC: as BB-M but with consideration of aerosol inclusions for cloud radiative properties.

Evaluation of the simulated aerosol absorption optical depth with retrievals from the AERONET sun-photometer network shows for all simulations a good agreement in the large scale global patterns with maxima in the biomass burning regions of Central Africa and South America, in the Saharan dust plume, and in regions with high anthropogenic activities over India and South East Asia. On a regional basis, general and seasonal biases, such as an underestimation of the AAOD at the urban Mexico City and central European sites, and differences among the simulations become evident. The spatio-temporally very inhomogeneous measurements do not seem to allow one to derive a global measure of the model performance. However, it becomes clear from the seasonal evaluation at 38 AERONET sites that the update of the BC refractive indices from Hess et al. (1998) to Bond and Bergstrom (2006) significantly improves the previous underestimation of AAOD in the BASE simulation. From this evaluation we conclude that the BB-M and the similar BRUG simulation seem to provide the best representation of the AERONET retrieved AAOD – for the black carbon emissions used in all simulations (Bond et al., 2004).

From comparison of the BB-M simulation with that simulation in which all aerosols are artificially made completely non-absorbing, we calculate that absorption acts to reduce the global annual-mean negative short-wave anthropogenic aerosol forcing of BB-M by 0.26 W m\(^{-2}\) (33 %) for the clear-sky and 0.34 W m\(^{-2}\) (72 %) for the all-sky case.
Our results confirm that the basic assumptions about the BC refractive index play a key role for aerosol absorption and radiative forcing; changing it from the widely used values $n=1.75+0.44i$ in BASE (Hess et al., 1998) to the medium absorbing values of $n=1.85+0.71i$ recommended by Bond and Bergstrom (2006) (values at $\lambda=550\text{ nm}$) in BB-M alters the annual-mean short-wave anthropogenic aerosol forcing clear-sky from $-0.65\text{ W m}^{-2}$ to $-0.53\text{ W m}^{-2}$ and all-sky from $-0.27$ to $-0.13\text{ W m}^{-2}$. By comparison, replacing the commonly used volume-weighted mixing for the calculation of the effective refractive indices for the internally mixed modes by the more accurate Bruggeman (1935) mixing rule changes the BB-M forcings only by $-0.06\text{ W m}^{-2}$ for clear skies and by $-0.05\text{ W m}^{-2}$ for all skies.

The effect of the enhanced aerosol absorption in BB-M, which shows better agreement with the AERONET retrieved aerosol absorption optical depth than BASE, to further reduce the negative aerosol radiative forcing is in contradiction to some recent, more negative, satellite-based estimates of anthropogenic TOA aerosol radiative forcings: Bellouin et al. (2005) report clear-sky $-1.9\pm0.3\text{ W m}^{-2}$ and all-sky $-0.8\pm0.1\text{ W m}^{-2}$; Chung et al. (2005) report clear-sky $-1.08\text{ W m}^{-2}$ and all-sky $-0.35\pm0.25\text{ W m}^{-2}$; Yu et al. (2006) report clear-sky $-1.33\pm0.82\text{ W m}^{-2}$. It has to be pointed out that the used passive satellite sensors cannot retrieve aerosols in the presence of clouds, adding to the uncertainty of the all-sky satellite estimates.

However, not only uncertainties in the aerosol system itself affect estimates of aerosol radiative forcing. Other factors in its calculation also contribute to the large diversity in current aerosol forcing estimates. In particular, the surface albedo plays a crucial role as its value determines the sign of the top-of-atmosphere aerosol radiative forcing for an aerosol column with given aerosol single scattering albedo. To quantify the effect of common assumptions for the surface albedo on aerosol absorption and radiative forcing, we replaced the land surface albedo field applied in BB-M by fields of the the upper and lower bound of the reported surface albedos in the AeroCom forcing experiment. Higher surface albedos result in a positive contribution of aerosol absorption to the TOA forcing so that it is reduced in the global annual-mean from $-0.53$...
to –0.42 W m\(^{-2}\) (clear-sky) and from –0.13 to –0.06 W m\(^{-2}\) (all-sky) using the upper-bound surface albedo and enhanced from –0.53 to –0.61 W m\(^{-2}\) (clear-sky) and from –0.13 to –0.18 W m\(^{-2}\) (all-sky) using the lower-bound of the surface albedo fields. It needs to be emphasised that also the ocean surface albedo and its implementation in current aerosol forcing calculations is subject to considerable uncertainty, as indicated by a large diversity in the AeroCom submissions. However, owing to ambiguities in the diagnostics of the AeroCom submissions we refrained to include the ocean albedos in our sensitivity study so that the simulated global mean forcing diversity is likely to be a lower estimate.

The anthropogenic long-wave aerosol radiative effects predict a global annual-mean absorption of 0.06 W m\(^{-2}\) (clear-sky) and 0.03 W m\(^{-2}\) (all-sky) and a TOA forcing of 0.03 W m\(^{-2}\) (clear-sky) and 0.02 W m\(^{-2}\) (all-sky), negligible, as expected from the typical sub-micron size of anthropogenic aerosols. For the larger natural dust and sea-salt aerosols, long-wave radiative effects become of importance so that the long-wave total aerosol absorption is 1.33 W m\(^{-2}\) (clear-sky) and 0.72 W m\(^{-2}\) (all-sky) while the TOA forcing is 0.37 W m\(^{-2}\) (clear-sky) and 0.21 W m\(^{-2}\) (all-sky). However, the present setup of ECHAM5-HAM does not include the super-coarse modes of mineral dust and sea salt so that the long-wave aerosol radiative effect of natural aerosols close to the source regions could be underestimated. A version of ECHAM5-HAM including super-coarse aerosols is in development and will allow one to quantify these effects in the near future.

As previous studies indicated, the simulated effect of absorption by aerosol inclusions embedded in cloud droplets and ice crystals on the global radiation budget is small. The global annual-mean total aerosol short-wave AAOD of BC inclusions in clouds is 0.6×10\(^{-3}\), resulting in an absorption of 0.05 W m\(^{-2}\) and a positive TOA forcing perturbation of 0.02 W m\(^{-2}\). In published literature AODs and AAODs are generally reported as “clear-sky” without a precise definition. Our results indicate that a more detailed diagnostics of these quantities, separate for clear-sky and cloudy-sky cases, might improve the understanding about the diversity of aerosol radiative effects in current models and remote sensing products.
We intentionally did not address two key uncertainties in estimates of aerosol absorption and radiative forcing in this study: the uncertainty associated with emission inventories, which has been extensively discussed in the respective literature and the radiative effects of mixing of BC by micro-physical interaction with other aerosol components, that we have scrutinised in previous work. Uncertainties of the mass of present day BC emissions are estimated globally to be at least a factor of two with fossil fuels emissions ranging from 2.8 Tg(C) yr\(^{-1}\) to 8.0 (see Bond et al., 2004; Ito and Penner, 2005, and references therein) and wildfire emissions ranging from 1.8 Tg(C) yr\(^{-1}\) (year 2000 estimate, Hoelzemann et al., 2004) to 7.1 Tg(C) yr\(^{-1}\) (year 1997 estimate, Generoso et al., 2003). Robust estimates of emission size distributions representative for global modelling studies are generally not available, resulting in a significantly larger uncertainty in the emission number flux of primary BC particles. These uncertainties directly propagate into simulations of the aerosol absorption and radiative effects. However, without tighter constraints on atmospheric aerosol radiative properties, ambiguities will not allow one to use absorption measurements to improve the emission inventories by forward or inverse modelling. The radiative implications of mixing of BC with other aerosol components are more complex than is frequently portrayed. The mixing with non-absorbing, hydrophilic aerosols, such as sulfate, affects atmospheric absorption in opposing ways: increasing absorption through enhancement of the BC absorption efficiency by internal mixing, but also decreasing absorption through ageing initially hydrophobic BC to a hydrophilic state, enhancing its removal by wet scavenging, and consequently decreasing BC lifetime and abundance (Stier et al., 2006c).

Our analysis of the effect of the use of different land-surface albedo in current modelling studies highlights that uncertainties in the host model parameters, independent of the uncertainty associated with the aerosol simulations, can significantly contribute to the large diversity in current estimates of aerosol radiative effects. The estimated uncertainties would be even larger if the differing assumptions in the sea surface albedo or variations in the cloud fields could have been taken into account.

The demonstrated uncertainties related to assumptions in current aerosol models
and satellite forcing retrievals could be quantified in an inter-comparison study with prescribed aerosol radiative properties.

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Ginoux, P., Gong, S., Grini, A., Hendricks, J., Horowitz, L., Huang, P., Isaksen, I., Iversen,
T., Kirkevåg, A., Kristjansson, J. E., Kloster, S., Koch, D., Kroll, M., Lauer, A., Lamarque,
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Table 1. Modal structure of the aerosol module HAM where SU (sulfate), BC (black carbon), POM (particulate organic matter), SS (sea salt), DU (mineral dust) denote the components in the respective modes.

<table>
<thead>
<tr>
<th>Modes</th>
<th>Hydrophilic</th>
<th>Hydrophobic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nucleation</td>
<td>SU</td>
<td></td>
</tr>
<tr>
<td>Aitken</td>
<td>SU, BC, POM</td>
<td>BC, POM</td>
</tr>
<tr>
<td>Accumulation</td>
<td>SU, BC, POM, SS, DU</td>
<td>DU</td>
</tr>
<tr>
<td>Coarse</td>
<td>SU, BC, POM, SS, DU</td>
<td>DU</td>
</tr>
</tbody>
</table>
Table 2. Global annual aerosol and aerosol-precursor emissions used in the simulations in Tg yr\(^{-1}\) and Tg(S) yr\(^{-1}\) for sulfur species.

<table>
<thead>
<tr>
<th>Species</th>
<th>Source</th>
<th>Year 1750</th>
<th>Year 2000</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMS</td>
<td>Marine Biosphere</td>
<td>23.5</td>
<td>23.5</td>
</tr>
<tr>
<td></td>
<td>Terrestrial Biosphere</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
<td>SO(_2)</td>
<td>Volcanoes</td>
<td>14.6</td>
<td>14.6</td>
</tr>
<tr>
<td></td>
<td>Vegetation Fires</td>
<td>0.7</td>
<td>2.1</td>
</tr>
<tr>
<td></td>
<td>Industry, Fossil-Fuel,</td>
<td>0.8</td>
<td>54.2</td>
</tr>
<tr>
<td></td>
<td>Bio-Fuels</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total sulfur</td>
<td></td>
<td>39.9</td>
<td>94.6</td>
</tr>
<tr>
<td>BC</td>
<td>Vegetation Fires</td>
<td>1.0</td>
<td>3.0</td>
</tr>
<tr>
<td></td>
<td>Fossil-Fuel</td>
<td>0.0</td>
<td>3.0</td>
</tr>
<tr>
<td></td>
<td>Bio-Fuels</td>
<td>0.4</td>
<td>1.6</td>
</tr>
<tr>
<td>Total BC</td>
<td></td>
<td>1.4</td>
<td>7.7</td>
</tr>
<tr>
<td>POM</td>
<td>Biogenic</td>
<td>19.1</td>
<td>19.1</td>
</tr>
<tr>
<td></td>
<td>Vegetation Fires</td>
<td>12.7</td>
<td>34.7</td>
</tr>
<tr>
<td></td>
<td>Fossil-Fuel</td>
<td>0.0</td>
<td>3.4</td>
</tr>
<tr>
<td></td>
<td>Bio-Fuels</td>
<td>2.0</td>
<td>9.1</td>
</tr>
<tr>
<td>Total POM</td>
<td></td>
<td>33.8</td>
<td>66.3</td>
</tr>
<tr>
<td>SS</td>
<td>Wind driven</td>
<td>5050±5</td>
<td>5050±5</td>
</tr>
<tr>
<td>DU</td>
<td>Wind driven</td>
<td>820±10</td>
<td>820±10</td>
</tr>
</tbody>
</table>
Table 3. Size-distribution parameters $r_{\text{eff}}$ [µm] and $\nu_{\text{eff}}$ of the 32 water droplet and 40 ice crystal distributions (Rockel et al., 1991).

<table>
<thead>
<tr>
<th>Type</th>
<th>$r_{\text{eff}}$</th>
<th>$\nu_{\text{eff}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>0.188, 0.375, 0.75, 1.5, 3, 6, 12, 24</td>
<td>0.01, 0.1, 0.175, 0.25</td>
</tr>
<tr>
<td>Ice</td>
<td>0.375, 0.75, 1.5, 3, 6, 12, 24, 40, 60, 80</td>
<td>0.001, 0.1, 0.175, 0.25</td>
</tr>
</tbody>
</table>
Table 4. References for the long-wave aerosol refractive indices shown in Fig. 1.

<table>
<thead>
<tr>
<th>Species</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SU</td>
<td>Toon et al. (1976)</td>
</tr>
<tr>
<td>BC</td>
<td>Bond and Bergstrom (2006) spectrally</td>
</tr>
<tr>
<td></td>
<td>extrapolated with Hess et al. (1998)</td>
</tr>
<tr>
<td>POM</td>
<td>Hess et al. (1998)</td>
</tr>
<tr>
<td>SS</td>
<td>Shettle and Fenn (1979); Nilsson (1979)</td>
</tr>
<tr>
<td>DU</td>
<td>Irina Sokolik (pers. comm., 2006)</td>
</tr>
<tr>
<td>WAT</td>
<td>Downing and Williams (1975)</td>
</tr>
</tbody>
</table>
Table 5. Studies to explore the sensitivity of the simulated absorption to uncertain model parameters. Each scenario is simulated with AeroCom year 2000 and 1750 emissions.

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BASE</td>
<td>Base run as in Stier et al. (2005)</td>
</tr>
<tr>
<td>BB-M</td>
<td>Updated medium absorbing refractive indices from Bond and Bergstrom (2006)</td>
</tr>
<tr>
<td>BB-H</td>
<td>Updated highly absorbing refractive indices from Bond and Bergstrom (2006)</td>
</tr>
<tr>
<td>BRUG</td>
<td>As BB-M but with Bruggeman (1935) mixing rule for refractive indices</td>
</tr>
<tr>
<td>ALB-L</td>
<td>As BB-M but with low AeroCom surface albedo estimate</td>
</tr>
<tr>
<td>ALB-H</td>
<td>As BB-M but with high AeroCom surface albedo estimate</td>
</tr>
<tr>
<td>TRANS</td>
<td>As BB-M but with imaginary part of all aerosol refractive indices set to $n_i = 10^{-9}$</td>
</tr>
<tr>
<td>CL-INC</td>
<td>As BB-M but with new cloud radiative properties and in-cloud aerosol split in interstitial and cloud droplet/crystal inclusions that are considered via cloud radiative properties</td>
</tr>
<tr>
<td>CL-INT</td>
<td>As CL-INC with new cloud radiative properties and interstitial aerosol but without consideration of cloud droplet/crystal inclusions via cloud radiative properties</td>
</tr>
<tr>
<td>CL-EXT</td>
<td>As CL-INC with new cloud radiative properties but all in-cloud aerosol considered as interstitial, i.e. externally mixed with cloud droplets</td>
</tr>
</tbody>
</table>
**Table 6.** Complex BC refractive indices at $\lambda$=550 nm used in the sensitivity studies.

<table>
<thead>
<tr>
<th>Name</th>
<th>Refractive Index</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>BASE</td>
<td>1.75+0.44 $i$</td>
<td>Hess et al. (1998)</td>
</tr>
<tr>
<td>BB-M</td>
<td>1.85+0.71 $i$</td>
<td>Bond and Bergstrom (2006)</td>
</tr>
<tr>
<td>BB-H</td>
<td>1.95+0.79 $i$</td>
<td>Bond and Bergstrom (2006)</td>
</tr>
<tr>
<td>TRANS</td>
<td>$1.85+10^{-9} i$</td>
<td>Sensitivity study</td>
</tr>
</tbody>
</table>
Table 7. ECHAM5-HAM simulated short- and long-wave aerosol radiative properties and forcings for the different sensitivity studies. Aerosol optical depths are given at wavelength $\lambda = 550$ nm.

<table>
<thead>
<tr>
<th>Study</th>
<th>AAOD fClear-Sky Total [10^{-3}]</th>
<th>AAOD Clear-Sky Antrop. [10^{-3}]</th>
<th>Absorption Clear-Sky Total [W m^{-2}]</th>
<th>Absorption All-Sky Total [W m^{-2}]</th>
<th>Absorption Clear-Sky Anthrop. [W m^{-2}]</th>
<th>Absorption All-Sky Anthrop. [W m^{-2}]</th>
<th>Forcing TOA Clear-Sky Total Aerosol [W m^{-2}]</th>
<th>Forcing TOA All-Sky Total Aerosol [W m^{-2}]</th>
<th>Forcing TOA Clear-Sky Anthrop. [W m^{-2}]</th>
<th>Forcing TOA All-Sky Anthrop. [W m^{-2}]</th>
</tr>
</thead>
<tbody>
<tr>
<td>SW</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BASE</td>
<td>2.1</td>
<td>1.3</td>
<td>1.78</td>
<td>1.63</td>
<td>0.81</td>
<td>0.74</td>
<td>-4.02</td>
<td>-2.51</td>
<td>-0.65</td>
<td>-0.27</td>
</tr>
<tr>
<td>BB-M</td>
<td>3.0</td>
<td>2.0</td>
<td>2.35</td>
<td>2.15</td>
<td>1.20</td>
<td>1.11</td>
<td>-3.84</td>
<td>-2.30</td>
<td>-0.53</td>
<td>-0.13</td>
</tr>
<tr>
<td>BB-H</td>
<td>3.2</td>
<td>2.2</td>
<td>2.50</td>
<td>2.30</td>
<td>1.31</td>
<td>1.21</td>
<td>-3.80</td>
<td>-2.24</td>
<td>-0.50</td>
<td>-0.09</td>
</tr>
<tr>
<td>BRUG</td>
<td>2.6</td>
<td>1.7</td>
<td>2.12</td>
<td>1.95</td>
<td>1.05</td>
<td>0.97</td>
<td>-3.90</td>
<td>-2.37</td>
<td>-0.58</td>
<td>-0.18</td>
</tr>
<tr>
<td>ALB-L</td>
<td>3.1</td>
<td>2.0</td>
<td>2.42</td>
<td>2.22</td>
<td>1.18</td>
<td>1.08</td>
<td>-4.29</td>
<td>-2.62</td>
<td>-0.61</td>
<td>-0.18</td>
</tr>
<tr>
<td>ALB-H</td>
<td>3.0</td>
<td>2.0</td>
<td>2.38</td>
<td>2.18</td>
<td>1.24</td>
<td>1.14</td>
<td>-3.55</td>
<td>-2.09</td>
<td>-0.42</td>
<td>-0.06</td>
</tr>
<tr>
<td>TRANS</td>
<td>0.0</td>
<td>0.0</td>
<td>0.08</td>
<td>0.11</td>
<td>0.01</td>
<td>0.02</td>
<td>-3.96</td>
<td>-2.47</td>
<td>-0.79</td>
<td>-0.47</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Study</th>
<th>SW</th>
<th>LW</th>
</tr>
</thead>
<tbody>
<tr>
<td>BASE</td>
<td>1.33</td>
<td>0.72</td>
</tr>
<tr>
<td>BB-M</td>
<td>0.66</td>
<td>0.06</td>
</tr>
<tr>
<td>BB-H</td>
<td>0.14</td>
<td>0.03</td>
</tr>
<tr>
<td>BRUG</td>
<td>0.37</td>
<td>0.21</td>
</tr>
<tr>
<td>ALB-L</td>
<td>0.03</td>
<td>0.02</td>
</tr>
</tbody>
</table>
Fig. 1. Long-wave component refractive indices as given in Table 4.
Fig. 2. Global distribution and mean of the upper and lower bound of broadband short-wave (0.2–4.0 μm) land surface albedo fields sampled from submissions the AeroCom forcing experiment.
Fig. 3. ECHAM5-HAM simulated BASE year 2000 mean clear-sky aerosol absorption optical depth [1] at wavelength $\lambda = 550$ nm, comparison of simulated (red) and AERONET sun-photometer retrieved (black) clear-sky aerosol absorption optical depth [1] as seasonal distribution (clockwise, starting at 12 o’clock) at respective AERONET locations, simulated clear-sky short-wave aerosol absorption [W m$^{-2}$]. Global annual-mean values are given in the titles.
**Fig. 4.** ECHAM5-HAM simulated BB-M year 2000 mean clear-sky aerosol absorption optical depth [1] at wavelength $\lambda = 550$ nm, comparison of simulated (red) and AERONET sunphotometer retrieved (black) clear-sky aerosol absorption optical depth [1] as seasonal distribution (clockwise, starting at 12 o’clock) at respective AERONET locations, simulated clear-sky short-wave aerosol absorption [W m$^{-2}$]. Global annual-mean values are given in the titles.
Fig. 5. ECHAM5-HAM simulated BB-H year 2000 mean clear-sky aerosol absorption optical depth [1] at wavelength $\lambda = 550$ nm, comparison of simulated (red) and AERONET sun-photometer retrieved (black) clear-sky aerosol absorption optical depth [1] as seasonal distribution (clockwise, starting at 12 o-clock) at respective AERONET locations, simulated clear-sky short-wave aerosol absorption [W m$^{-2}$]. Global annual-mean values are given in the titles.
Fig. 6. ECHAM5-HAM simulated **BRUG** year 2000 mean clear-sky aerosol absorption optical depth [1] at wavelength $\lambda = 550$ nm, comparison of simulated (red) and AERONET sunphotometer retrieved (black) clear-sky aerosol absorption optical depth [1] as seasonal distribution (clockwise, starting at 12 o’clock) at respective AERONET locations, simulated clear-sky short-wave aerosol absorption [W m$^{-2}$]. Global annual-mean values are given in the titles.
Fig. 7. Simulated year 2000 anthropogenic aerosol short-wave clear-sky atmospheric absorption, all-sky atmospheric absorption, clear-sky TOA forcing, and all-sky TOA forcing [W m\(^{-2}\)] for the BASE, BB-M, BB-H, BRUG, ALB-L, ALB-H, and TRANS scenarios. Global annual-mean values are given in the titles.
Fig. 8. Annual-mean difference in the effective ECHAM5-HAM broad-band surface albedo between the simulations with the upper (ALB-H) and lower (ALB-L) bound of the AeroCom land surface albedos. The global annual mean value in the title includes unchanged ocean areas to allow for a comparison with the global forcings in Fig. 9.
Fig. 9. Annual-mean difference of clear-sky and all-sky anthropogenic direct aerosol radiative forcings between simulations applying the upper (ALB-H) and lower (ALB-L) bound of the AeroCom land surface albedos given in Fig. 8.
Fig. 10. Year 2000 instantaneous long-wave total aerosol TOA direct radiative forcing and atmospheric absorption from the BASE simulation.
Fig. 11. Year 2000 instantaneous long-wave anthropogenic aerosol TOA direct radiative forcing and atmospheric absorption from the BASE simulation.
Fig. 12. Annual-mean in-cloud broad-band visible (0.25–0.69 µm) aerosol absorption optical depths from the CL-INC, CL-INT, and CL-EXT simulations, integrated over the cloudy layers of the atmospheric column. Inclusion AAOD: inclusion aerosol AAOD, i.e. cloud absorption optical depth with consideration of black carbon inclusions; Interstitial: interstitial AAOD.
Fig. 13. Simulated year 2000 total aerosol all-sky short-wave atmospheric absorption and TOA radiative forcing differences between CL-INC and CL-INT and between CL-EXT and CL-INT.
Appendix A

Scatterplots of daily-mean simulated and AERONET retrieved absorption optical depth for the BB-M sensitivity study in Fig. 4

Fig. A1.
Fig. A1. Continued. Scatterplots of year 2000 daily-mean ECHAM5-HAM simulated and AERONET retrieved absorption optical depth at wavelength \( \lambda = 550 \text{ nm} \) for the BB-M simulation as used in Fig. 4b). The ECHAM5-HAM output is sampled for each measurement date at the model grid box containing the measurement station. The seasonality is colour coded.