What's the real role of iron-oxides in the optical properties of dust aerosols?

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Abstract:
Iron oxides compounds constitute an important component of mineral dust aerosol. Several previous studies have shown that these minerals are strong absorbers at visible wavelengths and thus that they play a critical role in the overall climate perturbation caused by dust aerosol. When compiling a database of complex refractive indices of possible mineral species of iron-oxides to study their optical properties, we found that uniformly continuous optical constants for a single type of iron-oxides in the wavelength range between 0.2 μm and 50 μm is very scarce and that the use of hematite to represent all molecular or mineral iron-oxides types is a popular hypothesis. However, the crucial problem is that three continuous datasets for complex refractive indices of hematite are employed in climate models, but there are significant differences between them. Thus, the real role of iron-oxides in the optical properties of dust aerosols becomes a key scientific question, and we address this problem by considering different refractive indices, size distributions, and more logical weight fractions and mixing states of hematite. Based on the microscopic observations, a semi-external mixture that employs an external mixture between Fe-aggregates and other minerals and partly internal mixing between iron-oxides and aluminosilicate particles is advised as the optimal approximation. The simulations demonstrate that hematite with a spectral refractive indices from Longtin et al. (1988) shows approximately equal absorbing capacity to the mineral illite over the whole wavelength region from 0.55 μm to 2.5 μm, and only enhances the optical absorption of aerosol mixture at λ<0.55 um. Using the dataset from Querry (1985) may overestimate the optical absorption of hematite at both visible
and near-infrared wavelengths. More laboratory measurements of the refractive index of iron-oxides, especially for hematite and goethite in the visible spectrum, should therefore be taken into account when assessing the effect of mineral dust on climate forcing.

1. Introduction

Iron oxides in dusts are now identified as being important component for a number of climatic, environmental and biological processes. Over the past decade, iron-oxide minerals have been shown to be able to strongly absorb solar radiation (Tegen et al., 1997; Sokolik and Toon, 1999; Lafon et al., 2004; Qin and Mitchell, 2009; Redmond et al., 2010), and thus have a direct impact on the Earth’s radiation balance (Balkanski et al., 2007; Smith and Grainger, 2014; Scanza et al., 2015). Based on the ability to absorb acidic gases and water vapor (Baltrusaitis et al., 2007; Lafon et al., 2004; Qin and Mitchell, 2009; Redmond et al., 2010), iron oxides also contribute to heterogeneous reactions and cloud processes (Shi et al., 2011; Dupart et al., 2012), further influencing the radiation balance. In particular, layers of dust on snow and ice cover accelerate the melting of snow and ice by diminishing the surface albedo (Painter et al., 2010; Ginot et al., 2014) and the heat-absorbing properties of iron oxides in these dust layers can add to this effect (Kaspari et al., 2013; Reynolds et al., 2013; Dang and Hegg, 2014). Moreover, the deposited iron-bearing dust aerosols provide critical nutrients to marine and terrestrial ecosystems, which associated with consequential important drawdown of atmospheric carbon dioxide (Jickells et al., 2005; Shao et al., 2011; Nickovic et al., 2013). However, these effects can lead to either positive or negative net radiative forcing depending mostly on the underlying surface albedo, vertical profile (optical depth and height of dust layer), particle size distribution and mineralogy (Liao and Seinfeld, 1998; Claquin et al., 1999). This large uncertainty results from our limited knowledge of the physical, chemical and optical properties of atmospheric iron oxides on various space and time scales (Tegen et al., 1997; Sokolik et al., 2001; Formenti et al., 2011).

The element iron can be found among numerous mineralogical species, such as feldspars, clays (e.g. illite, smectite, chlorite and biotite), iron-oxides, iron-hydroxides and so on. A useful mineralogical classification frequently used in soil science distinguishes two categories of iron: (i) “structural iron” (in either the Fe(II) or Fe(III) oxidation states), trapped in the crystal lattice of aluminosilicate minerals; and (ii) iron (in the Fe(III) oxidation state), in the form of discrete oxide
or hydroxide particles (Lafon et al., 2004). According to the classical terminology of soil scientists (Sumner, 1963; Anderson and Jenne, 1970; Angel and Vincent, 1978), the latter kind of iron will be referred to as free-iron and its corresponding oxides and hydroxides as iron-oxides. Ten of 16 known iron oxides, hydroxides and oxide-hydroxides are known to occur in nature, with goethite, hematite and magnetite being the most abundant as rock-forming minerals; ferrihydrite, maghemite and lepidocrocite being intermediate in abundance in many locations; and wüstite, akaganéite, ferroxyhyte, and bernalite being the least abundant (Cornell and Schwertmann, 2006; Guo and Barnard, 2013). Dust aerosols from arid and semi-arid regions typically contain goethite, hematite, ferrihydrite and magnetite, and based on the mass contribution, hematite and goethite are the major components of free-iron in the atmospheric dust aerosols (Schroth et al., 2009; Shi et al., 2012; Takahashi et al., 2013). Hematite (Fe₂O₃) is very common in hot, dry soils and imparts a red color to its sediments. Goethite (α-FeOOH) is a common weathered product in soils and loesses. It occurs in moist, acidic soils (Schwertmann, 1993), and is brown to yellow in color.

Sokolik and Toon (1999) found that hematite is an especially strong absorber at ultra-violet (UV) and visible wavelengths, and it can also enhance the absorption of clay minerals and quartz through the formation of aggregates. Derimian et al. (2008) mentioned that iron oxides (primarily hematite and goethite) only affect the optical absorbing ability of aeolian dust at short wavelengths (the blue spectral region). Since the importance of hematite relative to other dust mineral components was discussed more fully in the study of Sokolik and Toon (1999), most subsequent modeling studies have assumed the iron-oxides in dust aerosols to be in the form of hematite. An opposite viewpoint was put forward, however, by Balkanski et al. (2007) who have argued that the optical absorption of dust with hematite at visible wavelengths might be lower than previous thought. The main reason is that internal mixing rule calculations with a hematite content of 1.5 volume % (or 2.8 weight %) was supposed to be representative for median dust absorption and was, as well, consistent with the AERONET measurements.

However, recent measurements on dust samples from east Asia, northern Africa and western Africa (Lafon et al., 2006; Formenti et al., 2008; Reynolds et al., 2014) indicated that goethite was presented in higher concentration than hematite (i.e. the ratio of goethite to hematite is about 7:3, a result that will be detail introduced in Section 3.2.2). For dust in snow, goethite is also the dominant ferric oxide that detected by reflectance spectroscopy and thus appears to be the main
iron-oxide control on absorption of solar radiation (Reynolds et al., 2013). More recently hematite and goethite has been taken into account interactively in global climate simulations due to the availability of global mineralogical distribution maps (Nickovic et al., 2012; Journet et al., 2014).

Furthermore, Müller et al. (2009) have measured the spectral imaginary refractive indices of hematite over the wavelengths from 590 nm to 790 nm during laboratory experiments, but these derived values are much lower than the data that employed in Sokolik and Toon (1999). The theoretical simulations of optical scattering of hematite and goethite at 470, 550, and 660 nm by Meland et al. (2011) also found that differences are apparent for hematite in both the phase function and polarization results at 660 nm where the imaginary indices from different references differ. Additionally, two studies that refer to optical properties of hematite in dust samples have also argued that the imaginary values of hematite refractive index in Sokolik and Toon (1999) are more than a factor of 2 larger than those reported by Bedidi and Cervelle (1993) and Longtin et al. (1988) at wavelengths below 600 nm (Moosmüller et al., 2012; Wagner et al., 2012).

Thus, what is the real role of iron-oxides in determining the overall impact of the optical properties of dust aerosols? This study will focus on investigating this important scientific question by considering heterogeneous optical refractive indices, mixing states and more logical abundance of iron-oxides.

2. Method and simulation

2.1 Complex refractive index

The complex refractive index (optical constant) is the most basic and significant parameter for calculating the optical properties of aerosols, but values for the optical constants of hematite in the wavelength range 0.2 to 50 μm are scarce in published references. Table 1 compiles information about the complex refractive indices of the major constituents of free-iron that we have found in the published literature. Sokolik and Toon (1999) employed the refractive indices of hematite from Querry et al. (1978), but Querry et al. (1978) mainly studied the optical constants of limestone and, as far as we can discover, do not contain any work on hematite at all. Since the publication of Sokolik and Toon (1999), studies (Höller et al., 2003; Alfaro et al., 2004; Mishra and Tripathi, 2008; Otto et al., 2009; Munoz et al., 2006; Meland et al., 2011; Wagner et al., 2012) have misquoted the data as Querry et al. (1978) or Querry (1987) when modeling the optical
properties of hematite. Actually, Querry (1987) is entitled “Optical constants of minerals and other materials from the millimeter to the UV” and gives optical constants of 29 materials, but still without referring to hematite. After careful searching, we have found that Querry (1985) has tabulated values for the refractive index \((m = n + ki)\) of hematite. Moreover, Longtin et al. (1988) also reported spectral refractive indices for hematite from earlier measurements by Steyer (1974), Onari et al. (1977), Galuza et al. (1979) and Kerker et al. (1979). This dataset also has been used in modeling the contribution of hematite to the optical properties of atmospheric dust aerosols in recent studies (Klaver et al., 2011a; Köhler et al., 2011; Hansell et al., 2011).

Recently, the third unpublished continuous refractive indices of hematite (named TA2005 in Table 1) from the Aerosol Refractive Index Archive (ARIA) of Oxford University have been employed in the Community Atmosphere Model (Scanza et al., 2015). Because this work has not been peer-reviewed and because TA2005 and QE1985 show a similar wavelength dependency, we pay special attention to the differences of refractive indices between QE1985 and LG1988 in this paper.

Any errors due to uncertainties in the optical constants would be directly reflected in the calculated results referred to above.

**Table 1.** Summary of the published complex refractive indices for major constituents of free-iron at different wavelengths (with their references).

| Iron-oxide species | Wavelength (μm) | Reference source | Abbreviation |
|--------------------|----------------|------------------|--------------|
| Hematite           | 8.3-50         | Popova et al. (1973) | PV1973       |
| Hematite           | 1.0-333        | Onari et al. (1977) | OA1977       |
| Hematite           | 0.25-0.7       | Shettle and Fenn, (1979) | SF1979   |
| Hematite           | 0.35-0.65      | Hsu and Matijevic, (1985) | HM1985     |
| Hematite           | 0.21-90        | Querry (1985) | QE1985       |
| Hematite           | 0.3-300        | Longtin et al. (1988) | LG1988     |
| Hematite           | 0.2-0.7        | Gillespie and Lindberg, (1992) | GL1992   |
| Hematite           | 0.2-4.5        | Krekov (1992) | KE1992       |
| Hematite           | 0.35-0.75      | Bedidi and Cerville (1993) | BC1993   |
| Hematite           | 0.1-1000       | Triaud (2005)* | TA2005      |
| Hematite           | 5-50           | Marra et al. (2005) | MR2005     |
| Hematite           | 5-2000         | Glotch and Rossman, (2009) | GR2009   |
| Hematite           | 0.59-0.79      | Müller et al. (2009) | ML2009     |
| Magnetite          | 0.21-55        | Querry (1985) | QE1985       |
| Magnetite          | 5-50           | Mukai (1989) | MK1989       |
| Magnetite          | 0.25-0.7       | Gillespie and Lindberg, (1992) | GL1992 |
| Magnetite          | 0.1-1000       | Amaury et al. (2002) | AU2002     |
| Magnetite          | 5-100          | Glotch and Rossman, (2009) | GR2009     |
Goethite  0.45-0.75  Bedidi and Cerville, (1993)  BC1993
Goethite  8-50  Glotch and Roman, (2009)  GR2009
Wüstite  0.2-500  Henning (1995)  HN1995
Wüstite  10-500  Henning and Mutschke, (1997)  HN1997

* The data was taken from http://www.atm.ox.ac.uk/project/RI/hematite.html

Bedidi and Cerville (1993) also presented refractive indices for hematite and goethite derived from reflectance measurements at wavelengths of 350-750 nm. Glotch and Rogers (2007) reported the optical constants of hematite, goethite and magnetite over the infrared (IR) wavelengths, and Hsu and Matijević (1985) also measured the refractive indices of hematite in the wavelengths of 350-650 nm.

Hematite is a uniaxial crystal which crystallizes in the trigonal system, whose optic axis corresponds to the crystallographic c-axis. Perpendicular to the c-axis are two radial a-axes. The dielectric constants of bulk hematite must therefore be measured for two principal polarizations of the incident light, namely one with the electric vector in any direction perpendicular to the c-axis (the so-called ordinary ray or –O ray) and the other with the electric vector along the crystalline c-axis (the extraordinary ray or –E ray). In this work we have calculated the average refractive indices for anisotropic hematite from all references in Table 1. The formula used is adopted from Longtin et al. (1988) and is:

\[ n_{avg} = \frac{(2n_{E-ray} / 3 + n_{O-ray} / 3) + (2k_{E-ray} / 3 + k_{O-ray} / 3)i}{3} \]

The reported values for the complex refractive index from the sources in Table 1 are markedly different, especially the imaginary part which controls the optical absorption. In order to visually demonstrate the variation of optical constants from different references, values of the real (n) and logarithmic values of imaginary parts (k) are shown in Figure 1.
Fig. 1. Spectral distribution of the imaginary and real parts of the complex refractive index for different iron oxides from Table 1 at wavelengths of 0 to 40 μm (Panels (a) and (c)) and expanded for wavelengths of 0 to 1.5 μm (Panels (b) and (d)).

For the real part of the refractive index for iron-oxides, there is a reasonable agreement between the hematite and magnetite datasets from the different references (Figure 1c). Because the real refractive index of hematite shows large fluctuations at wavelengths longer than 18 μm due to anisotropic refraction, the agreement between the different datasets decreases at these wavelengths. For goethite we are aware of only two sets of optical constants: one at visible wavelengths from Bedidi and Cervelle (1993) and the other at IR wavelengths from Glotch and Rogers (2007), but the wavelength gap between these two datasets hampers continuity. Unfortunately, Meland et al. (2011) have checked the former dataset for goethite using simulations according to Mie and T-Matrix theories and show that it may be in error. Nevertheless, we can see that goethite has optical constants similar to hematite. The real refractive index of hematite is larger than that of magnetite at wavelengths less than 2μm, but is smaller between 2 and 33 μm (Figures 1c&d).

For the imaginary part of the refractive index of iron-oxides, hematite and goethite have different optical properties at short wavelengths, both in terms of magnitude and spectral dependence (Bedidi and Cervelle, 1993). Between 460 and 700 nm the imaginary part of the complex refractive index (representing absorption) of goethite is up to 3 times smaller than that of hematite. As a consequence, the proportions of hematite and goethite in mineral dust can...
potentially change the magnitude and the spectral dependence of shortwave absorption of mineral
dust. However, the limited and discontinuous refractive indices of goethite have constrained the
evaluation of the effects of specific compositions of goethite and hematite to dust optical
properties and solar radiation balance over broader wavelength ranges.

From Figure 1a, we clearly see that the $k$ values for hematite from QE1985 and from
LG1988 show significant differences for wavelengths between 650 nm and 15 $\mu$m. These
differences are present at visible wavelengths and disappear at ultraviolet wavelengths, but the two
datasets have similar trends at UV and visible wavelengths (Figure 1b). Note that the hematite
optical constants vary dramatically across the visible wavelengths. In particular, the imaginary
part of the index for hematite shows a sharp decrease with increasing wavelength in the red. As a
consequence there is a large variability in the imaginary refractive index values for hematite taken
from different published references, particularly at 2 $\mu$m where the values from different sources
differ by a factor of 8600. Thus, this study firstly focuses on what will be the result if these
heterogeneous optical constants of hematite are used as input for the calculation of radiation
transfer models.

2.2 Particle size distribution

Size distribution is another important factor that affects the optical properties of particles.
Because Sokolik and Toon (1999) has employed the refractive index dataset for hematite from
QE1985 to calculate the radiative properties, we adopt here the same particle size distribution but
with the refractive index dataset for hematite from LG1988 to compare our results with Sokolik
and Toon (1999). The lognormal number size distribution is applied to dust aerosols:

$$
\frac{dn}{d \ln r} = \frac{N_0}{\sqrt{2\pi} \ln\sigma} \exp\left[-\frac{1}{2} \left(\frac{\ln r - \ln r_0}{\ln\sigma}\right)^2\right],
$$

where $r_0$ is the median radius, $\sigma$ is the geometric standard deviation, and $N_0$ is the total
particle number density of the component in particles per cubic centimeter.

In order to compare with the results of Sokolik and Toon (1999), the optical properties of
minerals are calculated on the assumption that they have one size mode but varying median radius.
The particle size modes are selected as $r_0 = 0.5$ and 0.7 $\mu$m, and $\sigma = 2.0$. The size mode with
median radius $r_0 = 0.5$ μm is believed to be representative of the particle size distribution of the long-lived, long-distance-transport mode of airborne dust (Patterson and Gillette, 1977; Arimoto et al., 1997). The larger $r_0$ is representative of a particle size mode which occurs near the dust source (Gomes and Gillette, 1993). In reality, the size distribution of dust aerosols can have one or several modes, characterized by a specific composition (Mahowald et al., 2013).

2.3 Theoretical simulations

Images from scanning electron microscopy (SEM) reveal non-spherical, irregular and compact shapes of the dust particles (Figure 3), but Otto et al. (2009) and Klaver et al. (2011b) have shown that spherical/non-spherical differences only influence the single scattering albedo by less than 1%. Meland et al. (2011) have also shown that moderate departures from spherical shape are relatively unimportant in determining the scattering matrix for particles with high refractive index values, such as hematite. Therefore, we expect the aerosol asphericity to have a negligible impact on our calculated results of optical properties and subsequent calculations using the Mie theory (which assumes a spherical morphology for the dust particles).

There are several different computer codes that can be used to compute optical properties for a lognormal particle size distribution. The theoretical light scattering simulations in this paper have used the MieTab software. MieTab uses a FORTRAN code with continued fraction modification produced by W. J. Lentz from the Mie code originally developed by Dave and Center (1968). This modified code can be obtained from http://diogenes.iwt.uni-bremen.de/vt/laser/codes/ddave.zip. In order to validate the accuracy of MieTab, we firstly compared it with a double precision Lorenz-Mie scattering code and a double precision T-Matrix code for a lognormal particle size distribution from Mishchenko et al. (2002). The double precision Lorenz-Mie and T-Matrix codes are available from http://www.giss.nasa.gov/staff/mmishchenko/t_matrix.html.

In addition to the wavelength dependent optical constants and the size distribution, the T-Matrix theory also requires assumptions about the particle shape. In this work we use an aspect ratio of 1.000001 to represent a spherical particle shape, because use of an aspect ratio exactly equal to 1 causes computational overflow in some cases. The calculated results from the three codes at different wavelengths and complex refractive indices for the same size distribution are
listed in Table 2. The good agreement of the results from the three codes demonstrates that the possibility of computational error affecting the interpretation of the calculated optical properties of iron-oxides can be neglected.

### Table 2. Comparison of simulated optical properties between MieTab, Lorenz-Mie and T-matrix methods.

| Wavelength | Code type | MieTab | T-matrix | Lorenz-Mie | MieTab | T-matrix | Lorenz-Mie | MieTab | T-matrix | Lorenz-Mie |
|------------|-----------|--------|----------|------------|--------|----------|------------|--------|----------|------------|
| m=n+ki     |           |        |          |            |        |          |            |        |          |            |
| n=3.102, k=0.0925 | Q_{ext} | 2.0950 | 2.1332 | 2.1657 | 2.1440 | 2.1917 | 2.2585 | 2.3250 | 2.5042 | 2.0226 |
| n=3.007, k=0.00974 | Q_{sca} | 1.2640 | 1.3284 | 1.3233 | 1.8500 | 1.9048 | 1.9587 | 2.3240 | 2.5033 | 2.0216 |
| n=2.66, k=0.00003 | Q_{abs} | 0.8310 | 0.8048 | 0.8404 | 0.2940 | 0.2870 | 0.2998 | 0.0010 | 0.0009 | 0.0010 |
|             | \omega_{0} | 0.6033 | 0.6227 | 0.6116 | 0.8629 | 0.8691 | 0.8673 | 0.9996 | 0.9996 | 0.9995 |

### 3. Results and discussion

#### 3.1 Basic optical properties

We focus here on modeling the spectral optical properties of iron-oxides which are needed for climate modeling: the volume extinction coefficient $\beta_{ext}$ (which is the sum of the scattering coefficient $\beta_{sca}$ and the absorption coefficient $\beta_{abs}$), the single scattering albedo $\omega_0$, and the asymmetry parameter $g$ (a cosine weighted integral of the scattering phase function). This set of parameters allows the calculation of radiation forcing in most climate models. Figure 2 shows calculated optical parameters for hematite (with complex refractive indices from QE1985 and LG1988) and illite with varying median radius at solar and infrared wavelengths. The volume total extinction coefficients $\beta_{ext}$ have been normalized as $\beta_{ext}^*$ for particle number concentration $N = 1 \text{ cm}^{-3}$. 

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Fig. 2. The calculated spectral optical properties for hematite, goethite and illite with different complex refractive index and size distribution. (a) Volume cross section of extinction, (b) Volume cross section of absorption, (c) Single scattering albedo and (d) Asymmetry factor.

Figure 2a and Figure 2b demonstrate how the normalized spectral extinction coefficient and the normalized spectral absorption coefficient vary due to the differences in the refractive indices and median radius of the minerals. As shown in Figure 2a, $\beta_{ext}$ for hematite has a spectrum which is clearly distinguishable from that for illite at UV, visible and IR wavelengths. One point should be noted: hematite has a lower normalized spectral extinction coefficient than illite at wavelengths less than 1.3 $\mu$m, which means that hematite has a weaker optical extinction capacity than illite at these wavelengths. In the IR region, the spectral features of hematite in $\beta_{ext}$ show large differences in volatility, and mimic the features in the refractive index of hematite. The magnitude of $\beta_{ext}$ depends on the parameters of the particle size distribution.

Figure 2b shows the equivalent normalized spectral absorption coefficient for hematite from QE1985 and LG1988 at wavelengths less than 5 $\mu$m. The normalized absorption coefficient of hematite from QE1985 is about 100 times larger than that for illite at both visible and near-IR wavelengths, but the normalized absorption coefficient for hematite from LG1988 has larger values than those for illite at wavelengths less than 1 $\mu$m and about 100 times smaller values than...
those for illite at wavelengths between 1 μm and 5 μm. If we adopt the complex refractive indices of QE1985, the calculated absorption coefficient of hematite indicates that hematite is an especially strong absorber at UV and visible wavelengths. Conversely, the calculated absorption coefficient of hematite using LG1988 data suggests that hematite is an important aerosol component only for short-wavelength absorption. Considering the whole region from 0.2 μm to 5 μm, hematite with complex refractive indices from LG1988 has an approximately equal absorbing capacity to that of illite.

Figure 2c illustrates the single scattering albedo of hematite and illite at wavelengths between 0.2 μm and 5 μm for different particle size distributions. The single scattering albedo of hematite is about 0.6 for wavelengths λ < 0.55 μm and varies little from \( r_0 = 0.7 \) μm to \( r_0 = 0.5 \) μm. In contrast, illite has \( \omega_0 \) in the range from about 0.9 to 1 for \( \lambda < 0.55 \) μm, showing strong spectral dependence at short wavelengths. For \( \lambda > 0.55 \) μm, illite has \( \omega_0 \) of about 0.99 for 0.55 < \( \lambda < 2.0 \) μm and this gradually reduces to about 0.5 for 2.0 < \( \lambda < 5.0 \) μm, while hematite shows large differences of \( \omega_0 \) depending on the source of the refractive indices. The single scattering albedo decreases to about 0.35 at UV wavelengths for hematite nanoparticles (which are always observed as aggregates with other clay mineral particles) with \( r_0 = 0.01 \) μm and \( \sigma = 2.0 \), but it rapidly increases to nearly 1 at wavelengths \( \lambda > 0.7 \) μm.

Figure 2d compares the asymmetry parameter of hematite with refractive indices from QE1985 and LG1988 at UV and visible wavelengths with size modes of \( r_0 = 0.7, 0.5, \) and 0.01 μm. For \( r_0 = 0.5 \) and 0.7 μm, hematite from QE1985 has \( g = 0.3 - 0.99 \), \( g \) decreasing as \( \lambda \) increases. The magnitudes of \( g \) from LG1988 are in the range from 0.2 to 0.99 with a few fluctuations. For \( r_0 = 0.01 \) μm, both datasets put \( g \) in the range from about 0.15 to 0.38. Thus, the magnitude of \( g \) depends significantly on the particle size distribution.

3.2 Physical and mineralogical properties

3.2.1 Size and Morphology

Many electron microscopy observations (Greeland et al., 1968; Tipping, 1981; Postma and Brockenhuus-Schack, 1987; Poulton and Canfield, 2005; Raiswell and Anderson, 2005; Shi et al., 2009; Deboudt et al., 2012; Wagner et al., 2012; Guo and Barnard, 2013) have shown that poorly
ordered iron-oxides commonly occur as spheroidal to ellipsoidal nanoparticles that may be single or aggregated, and may be unattached or attached to quartz or clay minerals (Figure 3). The reasons for the aggregation and the attachment are not well understood but are likely to be related to interactions of surface charge characteristics between iron-oxides and quartz or clay minerals (Poulton and Canfield, 2005). According to Hinds (1982), the binding mechanisms that hold separate aerosols together in an agglomerate formed in the air include the van der Waals force, the electrostatic force and the surface tension of adsorbed liquid films. As mentioned above, the dispersed nanoparticles of iron-oxides which are attracted to larger dust particles have more prominent optical absorption than aggregated iron-oxides, but the use of size distributions for nanoparticles (such as $r_0 = 0.01\ \mu m$ and $\sigma =2.0$) will overestimate the optical absorption of iron-oxides in natural dust aerosol samples.

Fig. 3. Representative morphology of iron-oxide aggregates in dust samples observed by SEM.

3.2.2 Abundance of total iron and iron-oxides

By employing the optical parameters of hematite and clay minerals calculated above, we can model the optical effects of hematite in dust aerosols. Problems associated with this are the actual variation of iron-oxides content and the state of the mixture with other minerals and these should be accounted for when modeling the optical properties of dust aerosols.

The total iron content in dust aerosol bulk samples is always measured in terms of the Fe$_2$O$_3$
mass percentage of the total oxide mass by elemental analysis (i.e. XRF, PIXE or ICP). Ganor and Foner (1996) gave a median Fe content of 2.9% for dust storms in Israel. The observed Fe percentage for the Dunhuang site in China during ACE-Asia is 4.0 ± 0.9% (Zhang et al., 2003). A value of 4.45 ± 0.49% (Guieu et al., 2002) has been proposed as characterizing Saharan dust. The elemental analysis by XRF yielded total iron oxide contents between 2.0 and 5.0 weight % for four Saharan mineral dust samples of different color and origin (Linke et al., 2006b). Moreover, Lafon et al. (2004) and Lafon et al. (2006) reported that the total iron content (the Fe₂O₃ percentage) varies from 6.2% to 8.7% in six atmospheric samples and three wind tunnel generated samples. The total iron content ranges from 1.82% to 11.8% (with an especially high value of 30.0% in the sample collected from Bamako, Mali) in entrained PM₁₀ from ten soil samples representing the Arabian Peninsula, the Sahara and Sahel regions and samples from northeast Africa and south-central Asia (Moosmüller et al., 2012). Furthermore, percentage values of iron content from several datasets reported by previous studies vary mainly between 4 and 11% (e.g., Gomes and Gillette, 1993; Chiapello et al., 1997; Gao et al., 2001; Journet et al., 2014).

One additional aspect should be discussed here for the proper interpretation of the obtained data: how representative is the total iron concentration of the free iron oxide content of dust aerosols? Free-iron is present as a major aerosol component affecting the short-wavelength absorption of mineral dust. However, iron oxide represents only part of the total iron, which may also exist in the crystal lattice of numerous other dust minerals. The iron oxide-to-total iron ratio in natural and soil-derived aerosols has been characterized by applying an adapted reductive extraction method as commonly used in soil science (Lafon et al., 2004; Lafon et al., 2006). This method provides no structural information about the extractable iron and therefore cannot distinguish between the presence of goethite or hematite in the samples. Fortunately, this method can provide an upper limited to the free-iron content for the optical modeling.

Lafon et al. (2004) and Lafon et al. (2006) reported considerable variability in the iron oxide-to-total iron ratio for various regions and sampling conditions and that there is no clear relationship between the oxide-to-total iron ratio and dust origin or aging. Based on all the values for the oxide-to-total iron ratio reported in the published literature (Lafon et al., 2004; Lafon et al., 2006; Alfaro et al., 2004; Formenti et al., 2008; Klaver et al., 2011a), we have calculated an average of 0.52. Formenti et al. (2014a) reported that iron oxides account, by mass, for 0.38 to
0.72 of the total elemental iron based on X-ray absorption analysis of samples of mineral dust emitted from or transported to western Africa. Reynolds et al. (2013) reported that the percent iron in goethite and hematite relative to iron in all iron-bearing phases ranges from 0.2 to 0.52 for dust samples in Australia as determined from Mössbauer spectra. Based on an average compiled from the literature, Kandler et al. (2009) and Kandler et al. (2011) have assumed that only 20% of the total iron content is hematite when determining the complex refractive index of dust aerosols. Alfaro et al. (2004) found in their dust samples comparable total iron contents in the range of 3.0 to 6.5 weight %, and they assigned a significant amount of 2.8 to 5.8 % of this iron as present in iron oxide mineral phases. This result is consistent with the reported 2.8 - 5.0% of free-iron in aerosol samples collected from three different locations over the world (Lafon et al., 2004; Lafon et al., 2006). Takahashi et al. (2011) indicated that the content of iron-oxides is less than 5 weight % in Asian dust. Moreover, Klaver et al. (2011a) reported that the iron oxides-to-total iron ratio for the analysed samples varied between 0.4 and 0.61, accounting for between 1% and 3% of the total gravimetric mass, and Formenti et al. (2008) also illustrated that iron oxides (speciation hematite and goethite) represented 2.4% and 4.5% of the total mineral dust mass. As mentioned above, only about half of the total iron content is represented by free-iron.

XRD analytical technology has also been applied to identify the content of hematite and goethite in some cases and less than 2% iron oxides was detected (Shi et al., 2005; Linke et al., 2006b; Kandler et al., 2009; Lawrence et al., 2010; Klaver et al., 2011a; Wagner et al., 2012; Formenti et al., 2014b). Depending on the crystal phase of interest, this method has a detection limit of 0.1 to 0.5 weight % for iron oxides (Balsam et al., 2014). Discrepancies between the quantified free-iron content detected by the method of Lafon et al. (2006) and the hematite or goethite contents determined by XRD could be due to difficulties of the Rietveld method associated with poor crystallographic ordering of iron oxides in mineral dusts.

Single particle analysis has also been conducted for detecting the free iron oxides. Fe-rich particles (iron oxides) represented no more than 5% of the particle number in aerosol samples and hematite or goethite were found more often in the fine fraction (Chou et al., 2008; Kandler et al., 2009; Schladitz et al., 2009; Kang et al., 2009; Scheuvens et al., 2011; Malek et al., 2011; Wagner et al., 2012; Menéndez et al., 2014).

Table 3. Summary of global reported ratios of hematite to goethite (Hm/Gt) in dust aerosols.
| Location                | Type          | Method | Average value of Hm/Gt | Reference               |
|-------------------------|---------------|--------|------------------------|-------------------------|
| Niger (1)               | Aerosol*      | DRS    | 0.5625                 | Lafon et al. (2006)     |
| Tunisia (1)             | Aerosol*      | DRS    | 0.4085                 | Lafon et al. (2006)     |
| China-Zhenbeitai (1)    | Aerosol       | DRS    | 0.3514                 | Lafon et al. (2006)     |
| Niger (99)              | Aerosol       | DRS    | 0.4286                 | Formenti et al. (2008)  |
| Niger (12)              | Aerosol       | XAS    | 0.5771                 | Formenti et al. (2014)  |
| Gran Canaria (19)       | Aerosol       | DRS    | 0.9048                 | Lázaro et al. (2008)    |
| North Atlantic (9)      | Aerosol       | DRS    | 0.9276                 | Arimoto et al. (2002)   |
| Muztagata (7)           | Aerosol       | DRS    | 0.6918                 | Xu et al. (2014)*       |
| Golmod (29)             | Aerosol       | DRS    | 0.7262                 | Yang et al., (2014)     |
| Tazhong (6)             | Aerosol       | DRS    | 0.9157                 | Lu et al. (2011)        |
| Dunhuang (29)           | Aerosol       | DRS    | 0.8762                 | Shen et al. (2006)      |
| Yulin (32)              | Aerosol       | DRS    | 0.7158                 | Shen et al. (2006)      |
| Horqin (22)             | Aerosol       | DRS    | 0.7448                 | Shen et al. (2006)      |
| Australia (6)           | Aerosol       | MS     | 0.4571                 | Reynolds et al. (2014)  |

* Dust aerosol produced by wind tunnel; + Unpublished paper of the fourth author, private communication.

The technology of diffuse reflectance spectroscopy (DRS) has normally been to quantify the ratio of hematite to goethite in a particular dust sample (Lafon et al., 2006; Shen et al., 2006; Lázaro et al., 2008; Formenti et al., 2008). The accurate quantification of goethite and ferrihydrite in dust is extremely difficult owing to similarities in structure and associated absorption spectra of these two minerals (Scheinost et al., 1998; Torrent and Barrón, 2002; Schroth et al., 2009). This has the critical implication that the content of goethite measured by absorption spectroscopy is actually the sum of goethite and ferrihydrite. This does not, however, affect the optical calculations due to their optical similarity. Table 3 summarizes the measured ratios of hematite to goethite in global dust aerosol samples and shows higher ratios of Hm/Gt in Asian dust samples compared to African samples. Over the whole world, it is concluded that goethite predominates over hematite with a relative abundance of 50% - 75% of iron oxides in dust aerosols.

Based on the above reported results, we conclude that the iron-oxides account for approximately half of the mass of elemental Fe and for between 2 and 5% of the dust mass. Most of them are composed of goethite, representing between 50 and 75% of the iron oxide mass.
3.2.3 Mixing states

As free-iron particles are always mixed with other kinds of particle, the condition of the mixture could be important for their ability to scatter and absorb radiation. The 3D structure of iron-oxide particles obtained by tomography reveals that these Fe-rich inclusions are often found at the surface of aluminosilicate particles but that some are also included inside particles (Deboudt et al., 2012). Inversions calculated assuming external mixing are better able to explain the wavelength dependence of dust absorption by varying only hematite concentration than inversions using internal mixing (Koven and Fung, 2006; Formenti et al., 2014a). Thus, a semi-external mixing assumption is clearly an optimal approximation for iron-oxides mixed with aluminosilicate particles. Moreover, this assumption has the advantages of simplicity of calculation, interpretation, and the possibility of comparing with model results.

3.3 Further simulation and verification

Sokolik and Toon (1999) also suggested that the radiative properties of a mixture would strongly depend on the relative abundance of individual minerals due to the large variations in the optical properties of individual minerals. Therefore, we model the optical properties for a range of possible weight fractions of hematite in the clay-size mode while the remaining mass is illite which represents the clay minerals. As mentioned above, we adopt 0% hematite as the lower limit for the aerosol samples with no free-iron particles, 2.5% hematite for the transported dust aerosol samples, 5% hematite for the locally emitted dust samples and 7.5% hematite for the upper limit. Due to the limited and discontinuous refractive indices of goethite, this setting may underestimate the actual optical effects of goethite in dust aerosol. Using the density of hematite (5.3 g/cm³) and illite (2.75 g/cm³), volumetric hematite fraction was converted from the mass hematite fraction to calculate the effective complex refractive indices for dust.

Dust mineralogical composition is often expressed as a weighted fraction of individual components in the total dust sample. Because the relationship of number concentration and mass concentration can be expressed as:

\[ M_j = M_0 \cdot W_j = N_i \cdot \rho_j \cdot \frac{4}{3} \pi (r_o)^3 \cdot \exp \left[ \frac{9}{2} \left( \frac{1}{\sigma} \right)^2 \right], \]
where $M_i$, $N_i$, $W_i$ and $\rho_i$ are the particle mass concentration, number concentration, weight fraction and density of the $i$-th mineral in the mixture respectively and $M_0$ is the total particle mass concentration of the mixture sample, the optical properties of external mixtures of minerals can be modeled by:

$$K_{ext}^{mix} = \sum (K_{ext(i)}^{*} \cdot N_i) = \sum \left( K_{ext(i)}^{*} \cdot \frac{M_0 \cdot W_i}{\rho_i \cdot \frac{4}{3} \pi (r_0)^3 \cdot \exp \left[ \frac{9}{2} \ln \sigma \right]^2 \right),$$

Figure 4a shows the differences between the single scattering albedo calculated for the mixture of illite and hematite with complex refractive indices from LG1988 at visible and infrared wavelengths. Compared to $\omega_0$ for illite with no hematite, it is not hard to see that $\omega_0$ for mixtures with different amount of hematite show significant differences in four wavelength ranges, namely, 0.2 - 0.7 µm, 2.5 - 8.7 µm, 8.7 - 12.5 µm and 15.0 - 17.5 µm. For $0.2 < \lambda < 0.7$ µm and $15.0 < \lambda < 17.5$ µm, $\omega_0$ for the mixtures is smaller than $\omega_0$ for pure illite, which means the presence of hematite enhances the optical absorption of the mixtures. For $2.5 < \lambda < 8.7$ µm, $\omega_0$ for the mixtures is larger than for pure illite, which means the presence of hematite enhances the optical scattering of the mixtures. But for $8.7 < \lambda < 12.5$ µm, $\omega_0$ shows more complicated fluctuations.
Fig. 4. Spectral single scattering albedo (SSA) for mixtures of illite and hematite with varying percentages and refractive indices at wavelengths of 0-20 μm (a) and 0-3 μm (b) with comparison to field observed results at 550 nm.

Figure 4b shows the differences between \( \omega_0 \) for illite mixed with different amounts of hematite with refractive indices from QE1985 at solar wavelengths. The magnitude of \( \omega_0 \) for a mixture increases when the median radius \( r_0 \) increases. In order to compare the effects for hematite with refractive indices from two different sources for the same mixture state, we also plot \( \omega_0 \) for illite mixed with 7.5\% hematite with refractive indices from LG1988 in Figure 4b.
Evidently, for $0.55 < \lambda < 2.5 \mu m$, the dataset of QE1985 will lead to higher optical absorption, although the two datasets have the same optical scattering and absorption for $\lambda < 0.55 \mu m$.

Another coming question is what is the competition between theoretical calculated values and field observed results? In order to compare our calculated SSAs with measured values, we review all reported SSAs during different dust campaigns or inferred from AERONET measurements, and listed them in Table 4. The measured results for dust mixed with BC were excluded during our review progress, such as the results from the campaign of AMMA. The measurements from DABEX are comparable but on the lower edge of previous measurements performed at 550 nm during the TARFOX, SHADE, GERBILS, SAMUM, NAMMA and Fennec 2011 field campaigns. Absorption from the mineral dust as measured using the corrected nephelometer and Particle Soot Absorption Photometer (PSAP) combination suggests that single scattering albedos at 550 nm (SSA550) range from 0.91 to 0.97 (with a mean of 0.97) for iron oxide mass fractions between 1.3 and 3.5% (Klaver et al., 2011a). The SSAs at 532 nm were reported as 0.99±0.001 and 0.98 ± 0.002 for the Cairo 2 and Morocco dust samples with Fe2O3 mass fractions of 4.5 and 3.63%, respectively (Linke et al., 2006). Johnson and Osborne (2011) revealed that the use of the mineral dust refractive indices from Balkanski et al. (2007) assuming 1.5% hematite gave reasonable agreement with the measured single scattering albedo, consistent with the findings of Klaver et al. (2011a). Haywood et al. (2011) shown that mineral dust is relatively non-absorbing at 550 nm due to the relatively small fraction of iron oxides present (1–3%). Balkanski et al. (2007) addressed this difference and argued that dust absorption at visible wavelengths might be lower than previously thought because mixing rule calculations with hematite content of 1.5% by volume, supposedly representative of median dust absorption, showed a very good agreement with the AERONET measurements. The same question is presented in Figure 7a of Formenti et al. (2014a): that the calculated SSA using a higher imaginary index of iron-oxides also overestimated the absorbing ability of Saharan dust in comparison with field observation.

We compare our calculated SSA curves with the measured values in Figure 4b. It is shown that the observed SSAs were mostly ranged in 0.94-0.99 during different dust campaigns, but much lower (0.944-0.95) for the AERONET which cannot exclude the presence of black carbon with higher absorbing. Our calculated result could consistent with the higher part (0.97-0.99) of
measured SSAs, but higher than the lower part (0.95-0.97) due to the effect of coarse particles during different dust campaigns. Thus, the iron oxide content alone cannot explain the variability of the single scattering albedo. This suggests that more complete knowledge of the dust mineralogical composition and size distribution with varied multi-modes is needed as input to more rigorous modeling.

Table 4. Review of measured dust single scattering albedos during dust campaigns and AERONET observations.

| References                | SSA_550nm (Mean) | S.D. | Observations |
|---------------------------|------------------|------|--------------|
| Haywood et al. (2001)     | 0.97             | 0.02 | TARFOX       |
| Haywood et al. (2003)     | 0.97             | 0.02 | SHADE        |
| Johnson and Osborne (2011)| 0.97             | 0.02 | GERBILS      |
| Müller et al. (2011)      | 0.96             | 0.03 | SAMUM_2      |
| Petzold et al. (2011)     | 0.975            | 0.15 | SAMUM        |
| Jeong et al. (2008)       | 0.96             | 0.01 | NAMMA        |
| Osborne et al. (2008)     | 0.99             | 0.02 | DABEX        |
| Ryder et al. (2013)       | 0.965            | 0.015| Fennec 2011  |
| Lack et al. (2009)        | 0.95             | 0.01 | TexAQS/GoMACCS |
| Linke et al. (2006)       | 0.985            | 0.006| Laboratory   |
| Zhu et al. (2007)         | 0.95             | 0.01 | AERONET      |
| Kim et al. (2011)         | 0.944            | 0.005| AERONET      |

In order to check the further effects of the mixing state of hematite on the single scattering albedo, we use two sets of assumptions that bracket the actual state of hematite mixing: internal mixing (Int), in which individual dust particles are a combination of all components present; and external mixing (Ext), in which different components exist as separate particles.

For the case of an external mixture of particles, the average optical properties are calculated by summing over the optical properties of the individual species. Approximations have to be made to calculate the optical properties of internal mixed particles. Three common approximations for the calculation of these latter optical properties are the volume mixing method, the Bruggeman approximation and the Maxwell-Garnett approximation (Chyýlek et al., 1988; Bohren and Huffman, 1998). Detail information about the three methods is given by Sokolik and Toon (1999).

We have calculated the single scattering albedo (SSA) of illite-hematite mixtures with different hematite contents using internal mixing according to the above three internal approximations and also using external mixing.

The calculated SSA values for illite-hematite mixtures using internal and external mixture

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assumptions as a function of wavelength and hematite mass fraction are illustrated in Figure 5. For the case of external mixing, the SSAs at 405 nm show good agreement for refractive indices from QE1985 and LG1988, but the calculated SSAs at 870 nm for hematite with refractive indices from QE1985 are much smaller than those using LG1988. This is explained by Figure 4b where the two datasets have the same optical scattering and absorbing properties for $\lambda < 0.55 \mu m$ but the dataset of QE1985 leads to higher optical absorption for $\lambda > 0.55 \mu m$. The calculated SSAs with the three different internal mixing methods are all much smaller than those for external mixing both at 405 nm and 870 nm since the assumption of an external mixture results in less absorption and less wavelength dependence of absorption than does the assumption of an internal mixture for small amounts of hematite. The basic reason for this is due to the extremely high imaginary refractive index for hematite at short wavelengths. For the case of internal mixing, the SSAs from the volume mixing method are smaller than for the other methods. This is due to the averaged imaginary refractive index being larger than for the other two approximations. On the basis of the study of Peterson (1968), only the effective refractive index of the non-metallic part of the dust can be calculated using the volume mixing method. Thus, adopting the volume mixing method to calculate the optical properties of aerosol samples will lead to a smaller SSA (Levoni et al., 1997; Sokolik and Toon, 1999; Shi et al., 2005; Höller et al., 2003; Ebert et al., 2004; Kandler et al., 2007; Kandler et al., 2009; Petzold et al., 2009; Otto et al., 2009; Wagner et al., 2012).

The calculated SSAs using the Bruggeman approximation are consistent with those from the Maxwell-Garnet approximation for low hematite contents at both 405 nm and 870 nm but differ from them for hematite content larger than 10%. Both the Maxwell-Garnet and Bruggeman approximations are derived from the same integral equation for the propagation of electromagnetic waves in an inhomogeneous medium but under a different set of approximations (Chyžleš et al., 1988; Bohren and Huffman, 1998). In previous studies, the Bruggeman approximation (Sokolik and Toon, 1999; Lafon et al., 2006; Koven and Fung, 2006; Mishra and Tripathi, 2008; Thomas and Gautier, 2009; McConnell et al., 2010; Klaver et al., 2011a; Wagner et al., 2012; Mishra et al., 2012) has been more often used for calculating the complex refractive index of silicate-hematite mixtures than the Maxwell-Garnet approximations (Balkanski et al., 2007; Hansell Jr et al., 2011).

The Bruggeman approximation allows for the calculation of an effective dielectric constant of multicomponent mixtures without distinguishing between matrix and inclusions. If we do not
know which is the main body for the silicate and the hematite in an aerosol, it is better to choose
the Bruggeman approximation. For the Maxwell-Garnet approximation a decision must be made
as to which component is the matrix and which is an inclusion. The Maxwell-Garnett
approximation is designed for small inclusions inside a host matrix and thus it is not suitable for
hematite >50% in the mixture, so the calculated SSAs show abnormal fluctuation for hematite >50%
in Figure 5. This phenomenon disappears if we consider the inverse Maxwell-Garnet
approximation which makes hematite the host matrix. Actually, the variation of Fe$_2$O$_3$ has been
constrained within the range 0–10%, so we advise the use of the Maxwell-Garnet approximation
in which the inclusions should be identical in composition but may be different in volume, shape
and orientation.

The comparison between the laboratories measured SSAs with known abundance of
iron-oxides with our theoretically calculated SSAs by different mixing rules, would give us further
insights into the actual mixing states of iron oxides and accuracy of different refractive indices.
Laboratory measured SSAs by extinction and photoacoustic absorption measurements at different
wavelengths have been reported in Linke et al. (2006a) and Moosmüller et al. (2012). Moosmüller
et al (2012) has demonstrated that SSAs are much smaller at 405 nm than at 870 nm and that
SSAs at both wavelengths are dominated by and linearly correlated with the iron content. These
measured results are also shown in Figure 5 for comparison with our theoretically calculated SSAs.
The measured SSAs at 405 nm show good agreement with our external mixing SSA values
calculated from both LG1988 and QE1985 for hematite less than 10%, while they are much larger
than our internal mixing SSA values at 405 nm (Figure 5a), potentially indicating that the dust
samples are mainly external mixing and are accompanied by a very small degree of internal
mixing.

In contrast, the measured SSA values at 870 nm are much larger than our external and
internal mixing SSA values calculated from refractive indices from QE1985 but show good
agreement with our external mixing SSA values calculated using LG1988 values (Figure 5b). This
illustrates the fact that the complex refractive indices of hematite from QE1985 have greatly
overestimated absorption at 870 nm.
Fig. 5. The single scattering albedo (SSA) as a function of varying hematite content at wavelengths of (a) 405 and (b) 870 nm, with different mixing states (Internal and External mixing), different complex refractive index sources (QE1985 and LG1988) and mixing approximations (Volume, Maxwell-Garnet and Bruggeman). For comparison, the measured values using a photoacoustic instrument with integrating reciprocal nephelometer and linear fits from Moosmüller et al. (2012) are also shown.

For purposes of quantitatively illustrating the optical effects of hematite in the mixtures, we calculate $\Delta \omega_0$ as the difference between the single scattering albedo of mixtures with given abundances of hematite and pure illite. Thus, we have

$$\Delta \omega_0 = \frac{\omega_{0,\text{mixture}} - \omega_{0,\text{illite}}}{\omega_{0,\text{illite}}} \times 100\%.$$ 

Here, we assume $M_0 = 100 \mu g/cm^3$ and a lognormal size distribution with $r_0 = 0.5 \mu m$, $\sigma =$
2.0. The single scattering albedo of pure illite represents the case when the dust contains no hematite. If the calculated values of $\Delta \omega_0$ are negative, this means that hematite has high absorption at the corresponding wavelengths. Figure 6 shows $\Delta \omega_0$ calculated for hematite with refractive indices from QE1985 and LG1988 at $\lambda < 20 \mu m$. It demonstrates that hematite with refractive indices from LG1988 only enhances the optical absorption of dust mixtures for $\lambda < 0.55 \mu m$, but hematite with refractive indices from QE1985 enhances the absorption for $\lambda < 2.5 \mu m$. The magnitude of $\Delta \omega_0$ for 5% hematite over these wavelengths is approximately about 1%. Therefore, the use of refractive indices for hematite from QE1985 in climate models would lead to overestimation of the optical absorption at both visible and near-IR wavelengths. Another apparent difference is the positive magnitude of $\Delta \omega_0$ for hematite with refractive indices from the two datasets at wavelengths of $2.5 < \lambda < 8.7 \mu m$. 
Fig. 6. The difference $\Delta \omega_0$ between the single scattering albedo (SSA) of pure illite and the single scattering albedo of mixtures with different contents and refractive indices of hematite. (a) Complex refractive index of hematite from Qurrey (1985), (b) Complex refractive index of hematite from Longtin (1988).

Given all that, the complex refractive index of iron-oxides is therefore a key parameter in effects of dust aerosols on the radiation balance, and the optical constants of hematite from different sources become a major source of uncertainty in radiative forcing calculation. Further work is needed to provide experimental measurements of the refractive index of iron-oxides, especially for hematite and goethite in the visible region of the spectrum. More complete knowledge of the dust mineralogy, morphology and size distribution with varied multi-modes is
needed as input to more rigorous modeling.

4. Summary

In this paper we have investigated the spectral optical properties of iron-oxides with considering different refractive indices, size distributions, and more logical weight fractions and mixing states of iron-oxides. The iron-oxides account for approximately half of the mass of elemental Fe and for between 2 and 5 % of the dust mass. Most of them are composed of goethite, representing between 50 and 75 % of the iron oxide mass. The iron-oxides commonly occur as spheroidal to ellipsoidal nanoparticles that may be single or aggregated, and may be unattached or attached to quartz or clay minerals, which could be expressed as semi-external mixing state. Moreover, the spectral SSA values determined in the present study show a strong wavelength-dependence with a steep decrease from the visible to the near-UV. There are still problems that need to be solved in order to accurately study the real role of iron-oxides in determining the overall impact of dust aerosols on climate perturbation, as follows:

1. Although there have been many published investigations of the complex refractive index of different iron-oxides, uniformly continuous optical constants for a single type of iron-oxides from 0.2 um to 50 μm are very scarce. Some of them are inconsistent and careful checking of their accuracy is therefore essential.

2. The abundance of specific iron-oxide types (such as goethite and magnetite) remains unknown. Although many studies have measured the mass ratio of goethite to hematite as about 7:3, the absence of goethite optical constants at 0.75-8.5 μm restricts the usefulness of this ratio. Thus, using hematite to represent all types of iron-oxides is a popular hypothesis.

3. Microscopic observations and optical simulations have shown that semi-external mixtures employing both external mixtures of Fe-aggregates and other minerals and partly internal mixing between iron-oxides and aluminosilicate particles is the optimal mixing approximation.

4. For hematite, there are two datasets of complex refractive indices that differ significantly. Compared with LG1988, the complex refractive indices of QE1985 greatly overestimate the optical absorption at both visible and near-IR wavelengths. Comprehensive laboratory measurements of the refractive indices of iron-oxides, especially of hematite and goethite in the visible spectrum, should therefore be made in order to accurately assess the effect of mineral dust
on climate perturbation.

Theoretically calculated SSA values are comparable to values observed in recent laboratory and field studies in the range of 0.97-0.99. The iron oxide content alone cannot explain the variability of the single scattering albedo, and the lower SSAs could be explained as the presence of coarse dust particles and high-absorbing black carbon in natural transported dust aerosol. More complete knowledge of the dust mineralogical composition and size distribution with dynamic varied multi-modes is needed as input to more rigorous modeling. Furthermore, field observations focus on the mean and standard deviations of the modes for various sources, surface wind speeds and transport meteorology should also be conducted.

Acknowledgement

We are grateful to Dr. Yahui Yue at ITPCAS and Dr. Yongliang Li at BNU for completing the SEM–EDX analysis and Dr. Michael Mishchenko at NASA GISS for offering the T-matrix and Lorenz-Mie codes. We are also grateful for the helpful comments made by the editor of Prof. David Covert, the reviewer of Prof. Yves Balkanski and another anonymous reviewer. This work was supported by the National Natural Science Foundation of China (Grant No. 41205108, 41571063 and 41271074).

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