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## Internal structures of dust particles

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# TEM analysis of the internal structures and mineralogy of Asian dust particles and the implications for optical modeling

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## Abstract

Mineral dust interacts with incoming/outgoing electromagnetic radiation in the atmosphere. This interaction depends on the microphysical properties of the dust particles, including size, mineral composition, external morphology, and internal structure. Ideally all these properties should be accounted for in dust remote sensing, the modeling of single-scattering properties, and radiative effect assessment. There have been many reports on the microphysical characterizations of mineral dust, but no investigations of the internal structures or mineral composition of individual dust particles. We explored the interiors of Asian dust particles using the combined application of focused ion beam thin-slice preparation and high-resolution transmission electron microscopy. The results showed that individual dust particles consisted of numerous mineral grains, which were organized into several types of internal structure: single and polycrystalline cores of quartz, feldspars, calcite, and amphibole often with oriented clay coatings; individual clay agglomerates of nano-thin clay platelets showing preferred to random orientations commonly with coarser mineral inclusions; and platy coarse phyllosilicates (muscovite, biotite, and chlorite). Micron to submicron pores were scattered throughout the interior of particles. Clays in the coatings and agglomerates were dominated by nano-thin platelets of the clay minerals of illite-smectite series including illite, smectite, and their mixed layers with subordinate kaolinite and clay-size chlorite. Submicron iron oxide grains, dominantly goethite, were distributed throughout the clay agglomerates and coatings. Unlike the common assumptions and simplifications, we found that the analyzed dust particles were irregularly shaped with birefringent, polycrystalline, and polymineralic heterogeneous compositions. Accounting for this structural and mineralogical makeup may improve the remote sensing retrieval of dust and the evaluation of radiation effects, but will also require sophisticated single-scattering modeling. In particular, the observed internal structures of dust particles such as clay coatings, preferred orientation, embedded grains in clays, and pores, likely have a great impact on the light scattering of dust particles. The distribution and size of structural components

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Extensive microphysical characterizations have been performed for single dust particles to determine their chemical composition (Okada et al., 1990; Anderson et al., 1996; Ro et al., 2005; Gao et al., 2007; Kandler et al., 2007), mineralogical composition (Jeong, 2008; Jeong et al., 2014), and particle size distributions (Reid et al., 2003). However, the microphysical properties of individual particles have not been fully resolved, despite the abundant reports. Microphysical properties are determined by the laboratory analyses of dust samples on a filter or the real-time analysis of particles through means such as optical particle counting and time-of-flight mass spectrometry, to determine size, morphology, and chemical or mineralogical types (Kulkarni et al., 2011 and references therein). However, a “single” particle is rarely a single crystal or mineral, but commonly polycrystalline and polymineralic (Falkovich et al., 2001; Jeong, 2008; Jeong et al., 2014). Thus, the microphysical data obtained from “single” particles are often the result of the numerous mineral grains, composed of different mineral species. Nevertheless, dust particles have usually been grouped into several chemical and mineralogical types based on the resultant properties (Anderson et al., 1996; Gao et al., 2007; Jeong et al., 2014). The chemistry, mineralogy, external morphology, and size distribution of “single” particles are helpful, but insufficient, in modeling the true single-scattering properties of mineral dust.

The geometric characteristics of the internal structure of single dust particles and the varying dielectric properties of the structural components are largely unknown key factors in the evaluation of dust single-scattering properties. For example, Vilaplana et al. (2006) found that the linear polarization of scattered radiation has a fundamentally different size dependence and characteristics in terms of absorbing dust particles with or without internal structure. Likewise, Nousiainen et al. (2011a) observed that the single-scattering properties of spheroids with empty cavities could not be mimicked by solid spheroids of varying sizes, shapes and compositions, suggesting fundamentally different single-scattering properties as a result of particle porosity. Similarly a modeling study by Nousiainen et al. (2011b) found that, for irregular ice crystals, the internal structure had the greatest potential of all the parameters considered to change



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single-scattering properties. Nousiainen et al. (2003) and Muinonen et al. (2009) found that internal structure, assumed in their studies to be random structure in the absence of observational data, is potentially quite significant for the single-scattering properties of dust particles much larger than the wavelength. In contrast to considerable efforts expended to investigate how the single-scattering properties of dust particles depend on particle shape (e.g., Nousiainen, 2009; Nousiainen and Kandler, 2014), relatively little attention has been paid to effects arising from internal structures. Undoubtedly the main reasons for this are the general lack of information regarding the microphysical characteristics of the particle interiors and the limitations imposed by the computational methods needed to solve single-scattering properties.

Iron oxides of complex refractive indices with large real and imaginary parts contribute greatly to the single-scattering properties of mineral dust (Sokolik and Toon, 1999; Lafon et al., 2006; Koven and Fung, 2006; Balkanski et al., 2007; Derimian et al., 2008; Moosmüller et al., 2012). However, calculations have shown that their contributions vary greatly depending on assumptions about the mixing state and mineralogy of iron oxides (Sokolik and Toon, 1999; Lafon et al., 2006). However, direct analyses of mixing state and mineralogy were not performed for individual particles.

To summarize, almost no investigations dedicated to the analysis of the internal structures and mineralogical makeup of dust particles have been published so far. Consequently, the impact of these factors on dust single-scattering properties and radiative effects, as well as on the interpretation of remote sensing data, remains largely uninvestigated, or has otherwise been based on hypothetical models of internal structure (Nousiainen, 2009 and references therein). Yet, these are clearly important factors to consider in radiation-related applications. In this study, we explored the interiors of individual Asian dust particles using high-resolution transmission electron microscopy (TEM). Electron-transparent thin slices were prepared for TEM analysis using a focused ion beam (FIB) technique. We report the structural and mineralogical details of the Asian dust particles, and discuss the implications of our findings for single-scattering properties and, consequently, for remote sensing and radiative effects.

## 2 Samples and method

Asian dust particles were collected on a borosilicate glass-fiber filter using a Thermo Scientific high-volume sampler around Seoul on 31 March 2012 and in Andong, Korea, on 17 March 2009. Meteorological, mineralogical, and physical properties of the 2012 dust were previously reported by Jeong et al. (2014). Satellite dust-index images (National Meteorological Satellite Center, 2013) showed that the source of both the 2012 and 2009 dusts was the Gobi desert, situated around northern China and southern Mongolia.

The individual dust particles were preliminarily characterized using a TESCAN LMU VEGA scanning electron microscope (SEM), equipped with an IXRF energy dispersive X-ray spectrometer. The dust particles on the filter were transferred onto a carbon adhesive tape attached onto a SEM stub by lightly contacting tape to the filter surface. After platinum coating for 60 s, the predominant mineralogy of the dust particles was analyzed using energy dispersive X-ray spectrometry (EDXS). High-resolution SEM images were acquired with a JEOL JSM 6700F field emission gun (FEG) SEM.

Dust particles for FIB sample preparation were selected on the basis of the predominant particle mineralogy and morphology determined by EDXS and FEG SEM. The SEM stub was placed on a SMI3050TB FIB instrument for preparing thin slices of approximately  $(5\text{--}12) \times (5\text{--}6) \mu\text{m}^2$  area and about  $\sim 100$  nm in thickness. Carbon was first deposited on the target particle in a thickness of  $\sim 1 \mu\text{m}$  for welding the loose agglomerates of fine particles, and then a gallium ion beam was sputtered to cut one thin slice from each individual particle. Altogether 35 slices were prepared and analyzed by TEM. Of these, 26 slices had a good flatness and a wide area sufficient for the TEM analysis. However, only a limited analysis of a small area was possible in nine slices, which had broken due to the cleavage of minerals or loose agglomeration during the transfer from the FIB instrument to the TEM chamber. Some of the slices were not suitable for lattice fringe imaging due to their thickness. The FIB slices were imaged using three microscopes: a JEOL JEM 2100F FEG STEM at 200 kV and a JEOL JEM 3010 at 300 kV

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is one of the most important advantages of FIB milling of geological samples compared to the traditional Ar ion milling. Our TEM images discussed in later Sect. 3.1 preserve the large and small pores of delicate shapes, all of which are reasonably interpreted on the basis of grain agglomeration and mineral growth/dissolution

Mineral identification was based on lattice fringes and EDXS chemical compositions. The identification of non-phylosilicate and relatively coarse phyllosilicate minerals (muscovite, biotite, and chlorite) was straightforward, but that of nano-thin phyllosilicates (clay minerals) was delicate. The lattice fringes of clay minerals parallel to basal plane were recorded normally above the magnification of  $\times 200000$ . The identification of clay minerals was based on the spacings of lattice fringes corresponding to the spacing of repeat units of the crystal structures: 1.0 nm for illite,  $\sim 1.0$  nm for smectite and vermiculite,  $\sim 7.0$  nm for kaolinite, and  $\sim 1.4$  nm for chlorite (Fig. 1). Kaolinite and chlorite were directly identified from their EDXS and lattice fringes (Fig. 1). However, illite, smectite, vermiculite, and illite-smectite (vermiculite) mixed layers could not be positively distinguished from each other because smectite (1.4–1.6 nm unit layers in hydrated state) was dehydrated and contracted under the high vacuum of the TEM chamber, showing  $\sim 1.0$  nm lattice fringes similar to those of illite (1.0 nm lattice fringe) (Fig. 1). EDXS can be used for identifying illite and smectite with interlayer cations K and Ca, respectively. However, although illite could form a thick plate and be positively identified from the clear 1.0 nm lattice fringe and high K and Al contents, illite and smectite platelets are normally very thin, consisting of only a few repeat units. They cannot be separately analyzed using EDXS, even when using an electron microbeam that is as small as possible. Additionally, mixed layering of illite and smectite is common in the soil and geological environments (Weaver, 1989, Środoń, 1999). Therefore, in practice, we cannot distinguish between nano-thin illite and smectite. To avoid over-interpretation, nano-thin platelets of clay minerals showing  $\sim 1.0$  nm lattice fringes with varying K and Ca contents were grouped into illite-smectite series clay minerals (IS-CMs). ISCMs are likely nano-scale mixtures of nano-thin platelets of illite, smectite, and illite-smectite mixed-layers.

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### 3 Results and discussion

In what follows, we will first present the results of our TEM analyses of dust particles. Schematic models for the common structural types observed will then be proposed. Finally, we will discuss the possible implications of the structural features discovered on dust optical modeling and on climate and remote sensing applications.

#### 3.1 TEM observations

##### 3.1.1 Clay-rich particles

TEM data for three clay-rich particles showing different internal structures are presented here because they are the most abundant particle type in Asian dust (Jeong, 2008; Jeong et al., 2014). SEM images of the clay-rich dust particle #1 show a rough surface composed of submicron clay grains (SEM images in Fig. 2a and b). Low-magnification TEM images of the FIB slice prepared from the particle in Fig. 2a show an agglomerate of randomly oriented platelets of clay minerals, which are tightly interlocked with each other (Fig. 2c). EDXS analyses suggest that the clays are mainly ISCM with some chlorite and kaolinite. Minor quantities of submicron iron and titanium oxide grains are also randomly distributed in the clay matrix (Fig. 2c). Confirmation of the specific mineral species of the iron and titanium oxides was not possible because the quality of lattice fringes were poor due to the large slice thickness. The areal fractions of these oxides in Fig. 2c are approximately 0.9 and 0.3 %, respectively. High-magnification TEM images of the clay show loose, disrupted, and nano-thin clay platelets (Fig. 2d–f). Lattice fringes confirm ISCMs (Fig. 2e–f) intermixed with coarser platelets of chlorite (Fig. 2e) and kaolinite (Fig. 2f). There are many pores (total 2.2 %) of approximately 1  $\mu\text{m}$  in diameter (Fig. 2c). Pores may have been formed through the dehydration of expandable clay minerals such as smectite in the high vacuum TEM chamber. However, some of the pores (arrow in Fig. 2c) are unlikely to have formed in this way by dehydration.

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the slice prepared from the particle (Fig. 5c) reveals a quartz grain wrapped with thin coatings ( $\sim 200$  nm thick), which are composed of oriented submicron platelets of clay minerals (Fig. 5d and e). The clay minerals are mostly ISCMs, based on the  $\sim 1.0$  nm lattice fringe and EDXS (Fig. 5f). Submicron goethite grains identified from the lattice fringe and EDXS are scattered within the clay coatings (Fig. 5g).

Another quartz-rich particle shown in the SEM images (Fig. 6a and b) also has an irregular morphology and a surface coated with micron to submicron clay platelets. A TEM image of the slice prepared from the particle reveals a thick clay coating ( $\sim 1$   $\mu$ m thick) with a quartz core (Fig. 6c). The coating is primarily composed of nano-thin platelets of clay minerals showing a high degree of preferred orientation (Fig. 6c and d). Submicron quartz particles were also found in the clay coatings (Fig. 6d). The lattice fringe shows that the clay minerals are mostly ISCMs (Fig. 6e) with some chlorite (Fig. 6f). The ISCMs were tightly adhered to the surface of the quartz core (Fig. 6e). Fine iron oxides were also found within the clay coating, but were not common (Fig. 6d). Magnified images show subparallel, chaotic arrangement of nano-thin ISCM platelets (1–10 nm thick; Fig. 6d–f).

### 3.1.3 Plagioclase-rich particle

A plagioclase-rich particle shown in the SEM images (Fig. 7a and b) has an irregular shape with surface clay coatings. The slice prepared from the particle reveals clay coatings of  $\sim 0.2$  to  $1$   $\mu$ m thick (Fig. 7c). Magnified images of the coating show nano-thin platelets of clay minerals oriented along the plagioclase surface (Fig. 7d and e). A submicron quartz grain is embedded in the clay coating (Fig. 7g) and pores are found around the quartz grain. The lattice fringe images show that most of the clay platelets are ISCMs with some chlorite ( $\sim 1.4$  nm) and kaolinite (0.7 nm) (Fig. 7e, f, and h). In the cavity inside the plagioclase, short tubes of halloysite are found attached to the cavity wall (Fig. 7c, and i). Halloysite is a kaolin group clay mineral, typically forming during the chemical weathering of plagioclase (Jeong and Kim, 1993). The cavity in the plagioclase core was formed by dissolution during weathering in the source soils.



### 3.1.4 Calcite-rich particle

The particle in the SEM images (Fig. 8a and b) is calcite-rich with a clay surface coating. The slice made from the particle reveals a polycrystal consisting of micron to submicron-sized calcite crystals (Fig. 8c–e). The calcite polycrystal was coated with thin (< 200 nm) ISCM clay layers (Fig. 8d and e).

### 3.1.5 Amphibole-rich particle

An amphibole-rich particle is encrusted with submicron clay platelets (Fig. 9a and b). A TEM image of a slice prepared from the particle reveals that the surface of the amphibole is coated with subparallel stacks of ISCM platelets subordinately with chlorite, quartz, calcite, and titanium oxide (Fig. 9c and d). Submicron pores were distributed within the coatings, particularly around the larger grains such as chlorite and calcite (Fig. 9d). Lattice fringes show that the dominant clay minerals are ISCMs (Fig. 9e).

### 3.1.6 Biotite-rich particle

SEM images (Fig. 10a and b) reveal a biotite flake coated with submicron clay plates. A slice perpendicular to the plate shows long lenticular iron oxide grains formed along the biotite layers (Fig. 10c and d). The iron oxide was identified as goethite (5.6 % area fraction) using EDXS and a 0.42 nm lattice fringe (Fig. 10e). EDXS analyses showed that biotite was depleted of interlayer K, indicating K loss during weathering in the source soils. Goethite intergrowth in the weathered biotite is the result of oxidative weathering of biotite, where iron ions are released from the biotite lattice following oxidation (Jeong et al., 2006).

### 3.1.7 Chlorite-rich particle

The SEM images (Fig. 11a and b) show a chlorite-rich flaky particle covered with clay minerals. The slices prepared perpendicular to the flake show that half of the flake

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consists of subparallel ISCM clay minerals (Fig. 11c). The other half is chlorite with long lenticular voids (6.2%) that were propped by submicron crystals of goethite (4.3%) (Fig. 11c and d) as identified by EDXS and a 0.41 nm lattice fringe (Fig. 11e). The EDXS analysis of chlorite showed a slight increase in silicon (Si) content compared with that of fresh chlorite, indicating oxidative weathering of chlorite followed by the formation of goethite crystals that consumed the iron released from chlorite lattices to maintain a charge balance. The growth of goethite crystals is responsible for the formation of lenticular voids.

### 3.1.8 Iron-oxide-rich particle

An iron-rich agglomerate particle is shown in Fig. 12a. An overall TEM image of a slice prepared from the particle shows the association of an irregular magnetite crystal and clusters of iron-oxide nanograins (Fig. 12b and c), which were identified as goethite by electron diffraction (inset in Fig. 12c). The micron pores and goethite were likely formed by the dissolution of magnetite during the weathering in the source soils.

### 3.1.9 Polycrystalline/polymineralic rock fragment

Minerals form compact, heterogeneous solids such as igneous, metamorphic, and sedimentary rocks in Earth's lithosphere. Some rocks consist of coarse mineral grains up to several centimeters in size, while other rocks consist of fine mineral grains, the sizes of which are as small as a few micrometers in size. The grain sizes of minerals in rocks commonly exceed the particle size of long-range transported dust. Therefore, original rock fragments composed of several interlocked mineral grains are relatively rare in long-range transported dust. The TEM image of a slice (Fig. 13b–c) prepared from the dust particle in Fig. 13a shows a rock-fragment particle composed of quartz, plagioclase (albite), biotite and chlorite grains of a few micrometer sizes identified by EDXS and electron diffractions (Fig. 13d–f). The mineral grains are not a loose agglomerate of fine soil grains, but compactly interlocked with each other to form heterogeneous

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solids. The rock fragments have some pores (Fig. 13b) and are coated with ISCM clay layers of  $\sim 0.5\ \mu\text{m}$  thick scattered with goethite grains (Fig. 13c). The particle in Fig. 13c is a polycrystalline/polymineralic rock fragment, whereas the particle in Fig. 8 is an example of a polycrystalline/monomineralic rock fragment.

### 3.2 Structural models

To facilitate the optical modeling of mineral dust particles, the internal structures observed in Asian dust particles were grouped into idealized classes. We identified three major types of internal structure: Type I, coarse non-phyllsilicate minerals; Type II, nanocrystalline clay agglomerates; and Type III, coarse phyllsilicate plates.

Type-I dust particles have core grains of non-phyllsilicate minerals including quartz, plagioclase, calcite, K-feldspar, and amphibole in the order of abundance (Fig. 14). They exist as either monomineralic crystals (Figs. 5–9) or polymineralic rock fragments (Fig. 13). The monomineralic particles can also be further described as monocrystalline (Figs. 5, 6, 7 and 9) or polycrystalline (Fig. 8). Although some surfaces of the coarser core crystals are directly exposed, almost all the surfaces are covered with nanocrystalline ISCM clay coatings (ca.  $0.2\text{--}1\ \mu\text{m}$ ). Therefore, there are six subtypes, as presented in Fig. 14. In Asian dust sources, all silt-size mineral grains have been observed to be coated with clay minerals according to an electron microscopic analysis of the silty soils as shown in Fig. 4 of Jeong (2008). The clay coatings are features acquired in the source soils via repeated wet-drying cycles, and do not form in the atmosphere through, for example, cloud processing.

Type-II dust particles are clay agglomerates composed mainly of nanocrystalline clay minerals (Fig. 15). ISCM is the most abundant mineral group in Asian dust (Jeong et al., 2014). As shown in the Figs. 2d and 4d, the orientations of the nano-thin ISCM platelets are always subparallel in the nano scale. However, at larger scales, fabrics of clay agglomerates are diverse, ranging from complete lamination (Fig. 2c) to random (Fig. 4c). The clay agglomerates often have micron-scale pores of lenticular or irregular shapes (Figs. 2c, 3c, and 4c). Agglomerates of pure clays are rare. Many clay-rich

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agglomerates include larger non-phyllsilicate grains (quartz, plagioclase, K-feldspar, and calcite) and coarser phyllosilicates (muscovite, biotite, and chlorite). Therefore, clay agglomerates could be further classified into eight subtypes (Fig. 15).

Type-III particles are coarse phyllosilicates of muscovite, biotite, and/or chlorite (Fig. 16). The platy morphologies are regulated by the well-developed cleavages along the (001) basal planes. They are commonly coated with ISCM clays. Another feature of the internal structures is the occurrence of goethite (iron-oxyhydroxide) along the cleavages in the weathered biotite and chlorite, which is a feature acquired in the source soils. The lenticular voids occur in the weathered biotite and chlorite. Although we have not presented the data of muscovite, it is reportedly highly resistant to oxidative weathering due to the absence of iron. Thus, goethite microinclusions and lenticular voids are not expected in the internal structures of muscovite. Coarse phyllosilicates could be further classified into four subtypes, as presented in Fig. 16.

### 3.3 Implications for the optical modeling of dust particles

Individual dust particles are often composed of several mineral species. Their mineral grains and pores are arranged to form several types of internal structures, some of which could contribute significantly to the single-scattering properties of mineral dust. Our results imply that the presence of internal structures in natural dust particles is a rule rather than an exception. To quantify their effects on single-scattering properties, sophisticated simulations should be carried out. While this is clearly beyond the scope of the present study, we can nevertheless offer our first impressions and speculate on the possible impacts.

#### 3.3.1 Internal structures

In regards to the geometrical characteristics of the internal structure, the most important factor is the size scale of the structure compared to the wavelength of radiation. This impacts not only the effectiveness of the structure in influencing single-scattering

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properties (sub-wavelength structures interact with radiation only weakly) but also impacts how it should be accounted for in modeling. If the structures are small compared to the wavelength, and sufficiently randomly located, effective medium approximations (e.g., Chylek et al., 2000) may be used, after which the particle can be treated as a homogeneous material with mean dielectric properties. However, as the example in Sect. 3.3.2 shows, the mixing of dielectrically very different materials can lead to strong effects, and to large errors if all the assumptions are not satisfied. In general, the geometrical characteristics should be explicitly accounted for in the most accurate light-scattering simulations (e.g., Kocifaj and Videen, 2008; Kocifaj et al., 2008). For example, effective medium approximations are unlikely to be accurate for the coated structure present in Fig. 6, while for the embedded crystals in an ISCM matrix (Fig. 4) those approximations may work well. For particles with large pores inside, which present large structures with high dielectric contrasts, the effective medium approximations are likely to also fail (Nousiainen et al., 2011a). Therefore, the pores with fractional areas extending up to 16.3% are clearly significant and should be accounted for, preferably explicitly.

Another interesting aspect is that in many particles studied, the internal structure is far from random. Instead, we often see varying types of ordered structure. From the single-scattering point of view, the most important is whether these structures are also preferentially oriented. For example, the embedded constituent crystals may have preferred orientation within the particle (e.g., Figs. 4c–e and 15). Such a structure may act to make the whole particle seemingly birefringent, even if composed of isotropic materials. Whether this is significant for the particles' single-scattering properties depends on the strength of this structural birefringence and the overall shape of the particle. From Nousiainen et al. (2009) we know that polarization quantities in particles are sensitive to birefringence; whereas, Dabrowska et al. (2012) reports that the effect increases with increasing particle aspect ratio. For example, the preferred orientation of the platelets in clay layers (e.g., Figs. 14 and 16) may give rise to structural birefringence, especially if the particle is elongated, because then there will be more platelets oriented parallel

to the longer particle axis than perpendicular to it. Likewise, ordered layered structures, such as those seen in Figs. 10c, d, and 11c, d, may give rise to structural birefringence. Again, this depends on how preferential the orientations of such structures are in the particle.

### 3.3.2 Mineralogy

Mineral dust particles can be composed of numerous grains of several mineral species with different refractive indices and sizes. The single-scattering properties of the particles will depend on the internal mixing state, size, and distribution of the constituent grains. For the effect to be substantial, however, the refractive indices of the different grains must vary considerably. Iron oxides are considered to be the most important minerals in this respect, because they are relatively common and the real and imaginary parts of their refractive indices are considerably higher than those for most other mineral species typically encountered in atmospheric dust. There are many reports that even small amounts of iron oxides can be significant for the single-scattering properties of dust. For example, Sokolik and Toon (1999) found that even 1 % hematite mixed with kaolinite was sufficient to decrease the dust particles' modeled single-scattering albedo by  $\sim 10\%$  when assuming an internal mixture treated with effective medium approximation instead of an external mixture. Similarly, when Balkanski et al. (2007) attempted to constrain dust refractive indices by varying the hematite content in the internal mixture to fit AERONET data, they found that a subtle variation in hematite contents and their mixing state were critical in explaining the observed refractive indices by AERONET, and in evaluating the global net radiative effect. Lindqvist et al. (2013) also found that a few volume percent of hematite was sufficient to impact the simulated single-scattering properties of dust particles.

Confirmations of the specific properties possessed by iron oxides are rare, despite their great importance and many related assumptions. The identification of iron oxide minerals (hematite or goethite) has not been attempted other than by the diffuse reflectance spectroscopic analysis of Lafon et al. (2006). Goethite has different

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wavelength-dependent refractive indices from hematite (Bedidi and Cervelle, 1993). In addition, the actual impact of iron oxides depends on the grain size and distribution of iron oxides within a particle. For example, widely distributed small iron oxide grains will lead to stronger absorption than a few larger grains of identical total mass.

5 The results of our study indicate that goethite dominates over hematite, and is evenly distributed in the clay agglomerates, normally as submicron-size grains. Lafon et al. (2006), using diffuse reflectance spectroscopy, also showed that goethite was a major iron oxide in dust samples collected near the desert margin of China (38°17' N, 109°43' E). The estimated areal fractions of goethite range from 0.6% to 5.6% for those particles where it was present. The even distribution of submicron goethite particles suggests potentially considerable impacts on the particles' single-scattering properties.

10 Biotite, chlorite, and their weathered equivalents have also been found above trace quantities in Asian dust (Jeong et al., 2014). Their dark color and iron-rich chemical compositions suggest clearly higher imaginary parts of the complex refractive indices than those of colorless minerals such as quartz, feldspars, muscovite, calcite, or IS-CMs. However, their complex refractive indices have not been experimentally measured over a wide range of wavelengths (Mooney and Knacke, 1985). In addition, the grain sizes and spatial distribution of titanium oxides (possibly rutile and anatase) within dust particles are similar to those of iron oxides. Their high refractive indices (Cardona and Harbeke, 1965) may also significantly contribute to the single-scattering properties of dust, which deserve further investigation.

### 3.4 Implications for climate and remote sensing

25 The radiative impacts of mineral dust are ultimately derived from their single-scattering properties averaged over all sizes and shapes present. In principle, it is possible that the internal structures observed cause systematic effects on single-scattering dust properties that persist through the averaging and can therefore be important in

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radiation-related applications. The most important of such applications in atmospheric sciences are climate modeling and remote sensing.

For the internal structure of particles to substantially impact climate through radiation, two prerequisites must be satisfied: (1) the dust single-scattering properties need to be affected sufficiently, and (2) the dust optical depth needs to be sufficiently large. Räsänen et al. (2013) recently investigated the impact of dust-particle non-sphericity on climate and found the effect to be negligible on a global scale, but possibly important locally. We can expect the same to hold for the impact of internal structure. There exists the potential for large and fairly systematic impacts to single-scattering properties similar to those due to particle non-sphericity. However, the globally averaged dust optical depth is too small to allow for significant global impacts. Local and regional effects are possible, but depend on the impact on the single-scattering properties, which are yet to be quantified. We also note that atmospheric dust loadings during the last glacial maximum were an order-of-magnitude larger than today, thereby inducing much stronger global effects (Harrison et al., 2001).

In terms of remote sensing, the potential for important implications is much greater. Remote observations are usually directional and therefore dependent on the differential single-scattering properties that can be quite sensitive to the physical properties of particles. Again, the impacts depend on how the single-scattering properties are affected, which is yet to be quantified; however, it is safe to assume that different types of remote sensing measurements will be affected differently. For example, polarization quantities are more likely to be affected than the intensity (Nousiainen et al., 2009, 2012). Further, we speculate that lidar observations, looking at the exact backscattering angle, may be particularly sensitive to internal structure. We also emphasize that the impact may extend to atmospheric remote sensing beyond aerosol measurements, because the radiative impact of aerosols often needs to be accounted for and corrected even when measuring other atmospheric constituents, or when measuring through the atmosphere.

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Even though the present investigation considered a few dust particles, the major types of internal structure obtained by TEM analysis can be integrated with statistical data obtained by SEM single particle analysis as done by Jeong et al. (2014) to estimate the proportions of the structural types, and followed by the simulation of the optical properties of bulk dust. However, evidently there are long steps toward the optical simulation of bulk dust. Optical property of each structural type can be modeled first, and then we may progress to model bulk dust considering the proportions of the structural types.

### 4 Summary and conclusion

Optical models for the interaction between dust and electromagnetic radiation are important in the evaluation of net radiative effects, and in the processing of remote sensing data. All microphysical properties, including size distributions, particle morphology, and composition should be known and accounted for to allow for realistic optical single-scattering treatment. Of the many uncertainties in bulk microphysical properties, the most uncertain are the properties of individual particles that ultimately govern the radiative effects. That is to say, when computing bulk properties, averaging should in principle be performed for single-scattering properties rather than for microphysical properties; what is averaged matters, because the microphysical properties and the resulting single-scattering properties are not linearly proportional. For the most accurate radiative treatments, single-particle microphysical properties are thus needed. In the past, optical models have been based on many assumptions and simplifications of the mineralogical and structural properties of individual particles such as species, size, mixing state, and arrangement of constituent minerals. This study directly explored the interior of individual Asian dust particles, revealing many novel microphysical details of the constituent mineralogy and internal structures.

Individual dust particles are composed of several mineral species of varying grain sizes. Iron oxides, known as the most important minerals with large real and imaginary



parts of the refractive index, were scattered as submicron-sized grains throughout clay agglomerates. Goethite was the dominant iron oxide. In addition, we suggest that submicron titanium oxides, chlorite, and biotite are worth considering as optically significant minerals. The internal structures of individual dust particles were formed by the patterned arrangement of nano-to-micron-sized mineral grains and pores. Internal structures could be grouped into three major types: coarse cores of quartz, feldspars, calcite, and amphibole with oriented clay coatings; clay agglomerates of nano-thin clay platelets; and coarse platy phyllosilicates of muscovite, biotite, and chlorite. Nano-thin platelets of clay minerals were dominated by the illite-smectite series clay minerals with subordinate kaolinite and clay-sized chlorite.

The observed internal structures and mineralogy are potentially important factors for the single-scattering properties of Asian dust particles. For example, the contrasting dielectric properties of pores and constituent minerals may greatly impact light scattering by dust particles, while structural birefringence by the preferred alignment of nano-thin clay-mineral platelets or micron-size phyllosilicate plates may also produce significant effects. Directional remote sensing, for example lidar examining the exact backscattering angle, is strongly dependent on differential single-scattering properties, and may be particularly sensitive to a particles' internal structure. In addition, local and regional net radiative effects due to dust may depend on the structural and compositional properties of dust particles. The microphysical parameters of individual dust particles considered in this study can be explicitly accounted for in single-scattering modeling if sophisticated methods, such as a discrete-dipole approximation by Draine and Flatau (1994), are used. Such modeling studies can illustrate the means by, and degree to, which microphysical parameters influence dust particle single-scattering properties, and will allow for further investigation of the dust radiative effect and remote-sensing implications. In the future, we plan to both carry out such simulations and to measure the internal structures for more dust particles and from different sources.

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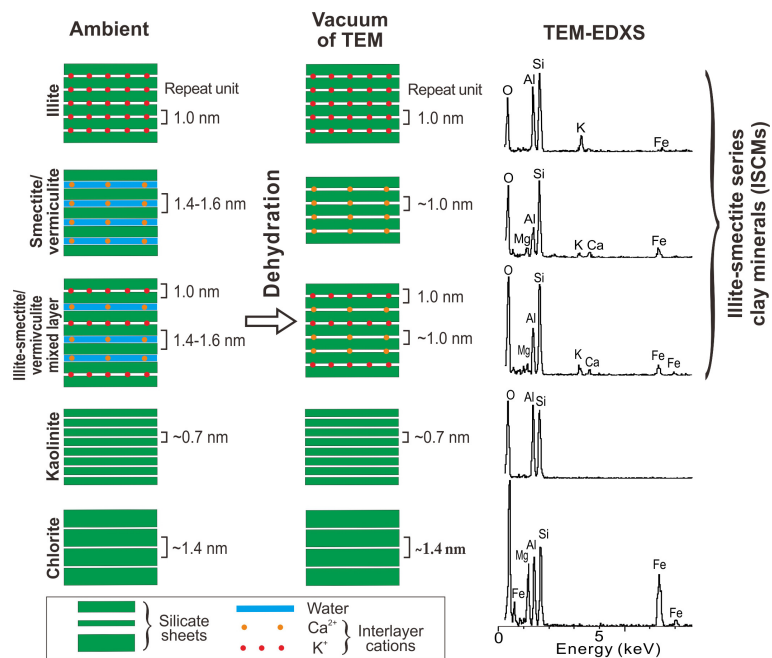


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**Fig. 1.** Identification criteria of clay minerals based on the spacing of repeat units measured from a TEM lattice fringe image (left two columns) and EDX spectra (right column). Repeated stacking of silicate sheets in clay minerals viewed along the crystallographic a–b plane.

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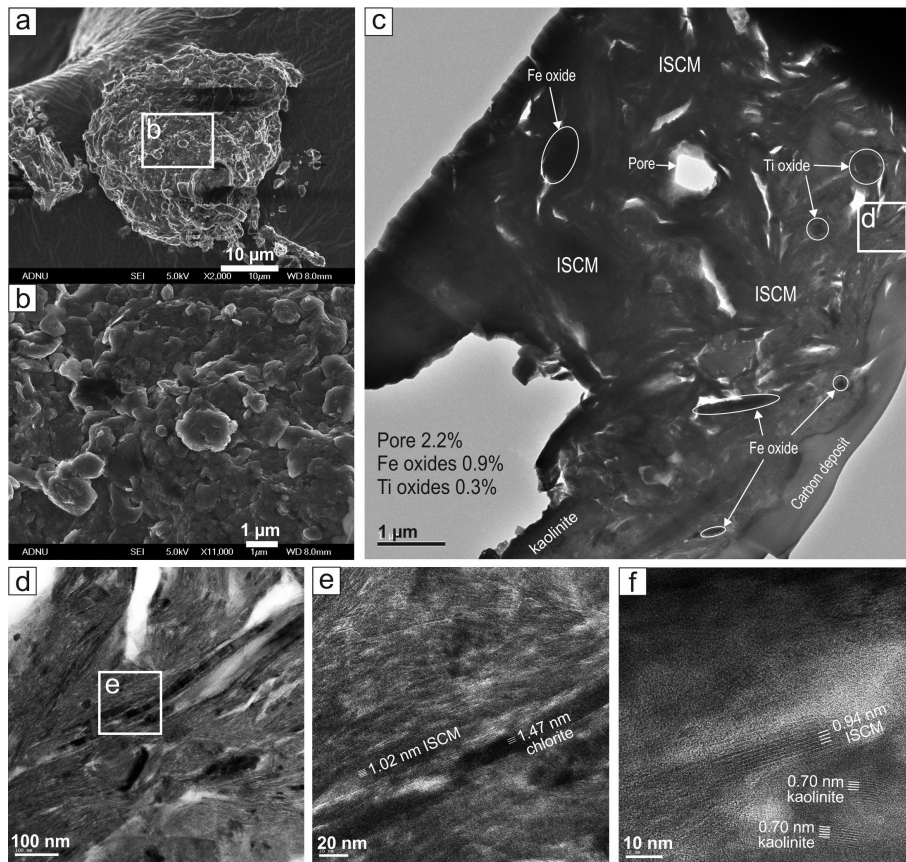
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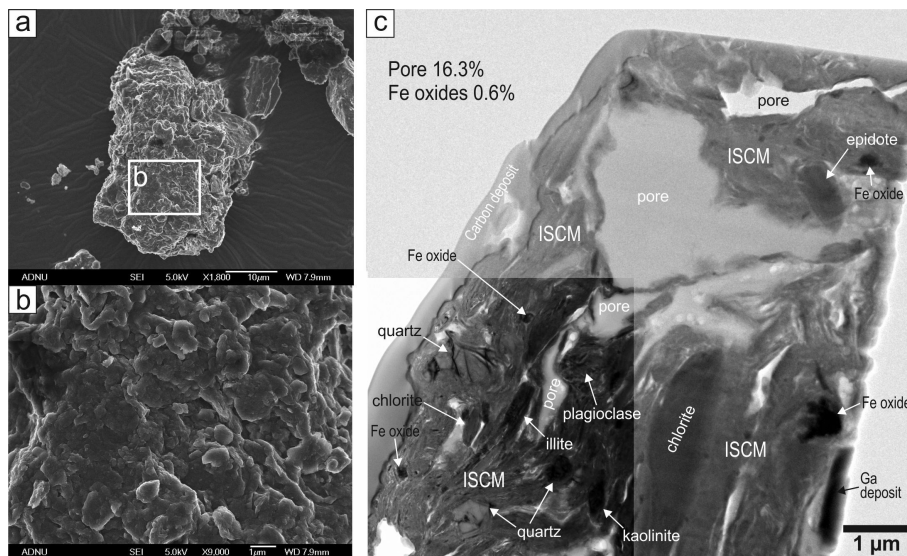






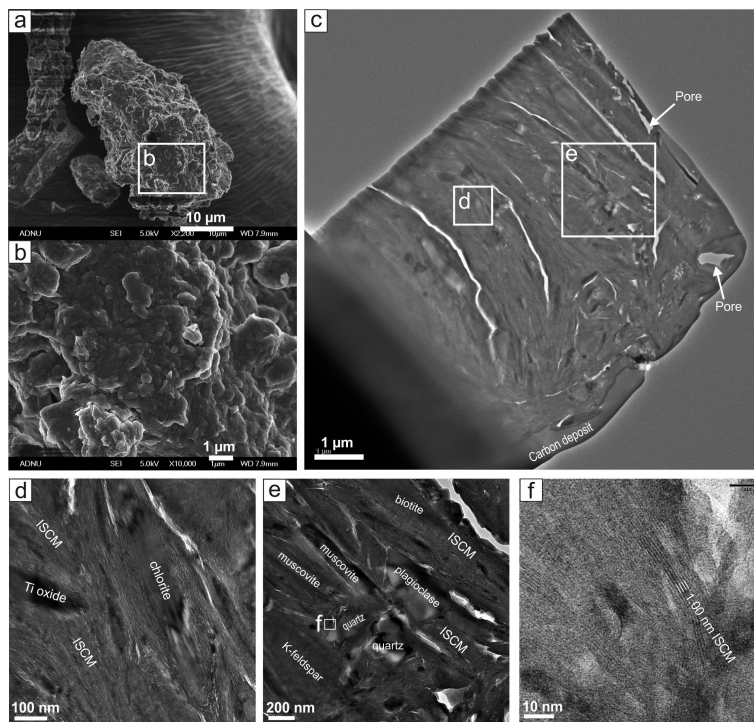
**Fig. 2.** Clay-rich dust particle #1 from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of FIB slice prepared from the particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of ISCM and chlorite. **(f)** TEM lattice fringe image of ISCM and kaolinite.

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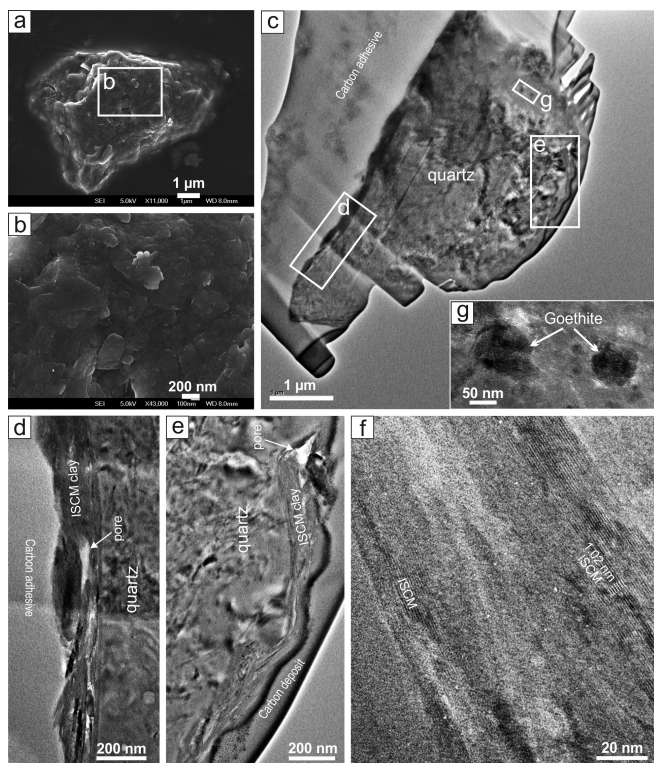
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**Fig. 3.** Clay-rich dust particle #2 from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. Pores are filled with carbon deposited prior to slicing.

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**Fig. 4.** Clay-rich dust particle #3 from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d, e)** TEM images magnified from the boxes in **(c)**. **(f)** TEM lattice fringe image of ISCMs.



**Fig. 5.** Quartz-rich dust particle #1 from 2009 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d, e)** TEM images magnified from the box in **(c)**. **(f)** TEM lattice fringe image of ISCMs. **(g)** Goethite spheres magnified from the box in **(c)**. **(e)** TEM lattice 767 fringe image of ISCMs. **(f)** TEM lattice fringe image of ISCM and chlorite.

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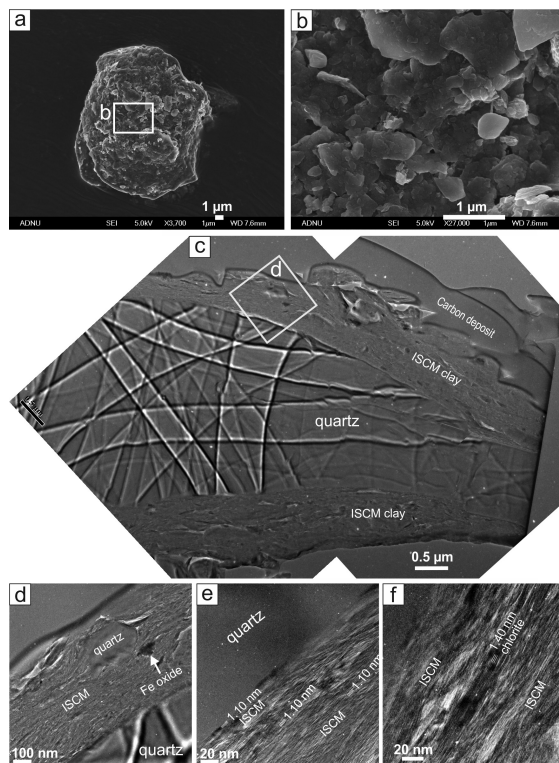
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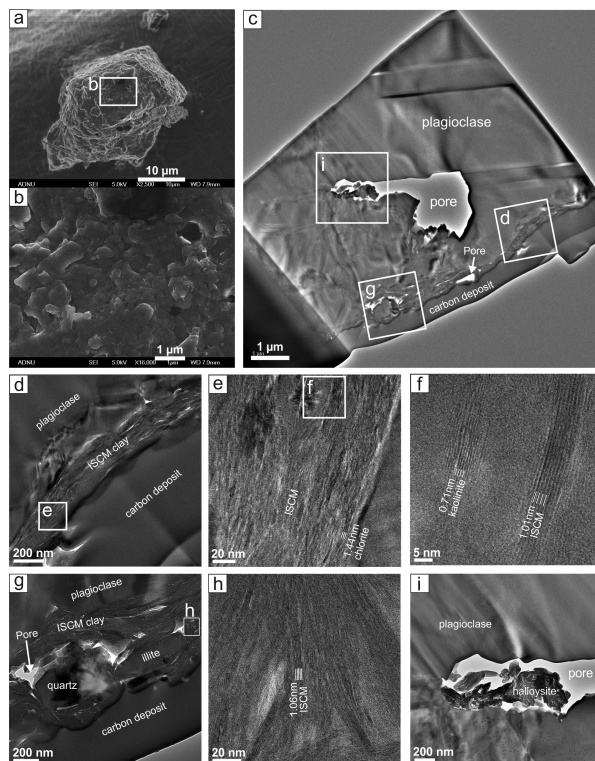






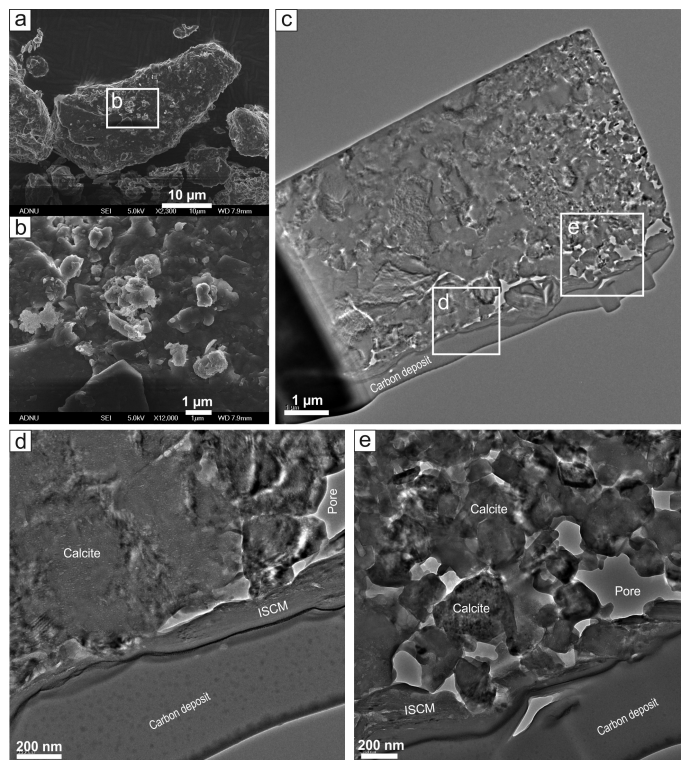
**Fig. 6.** Quartz-rich dust particle #2 from 2009 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of ISCMs. **(f)** TEM lattice fringe image of ISCM and chlorite.

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**Fig. 7.** Plagioclase-rich dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of ISCMs and chlorite magnified from the box in **(d)**. **(f)** TEM lattice fringe image of ISCM and kaolinite magnified from the box in **(e)**. **(g)** TEM image magnified from the box in **(c)**. **(h)** TEM lattice fringe image of ISCMs magnified from the box in **(g)**. **(i)** TEM image of internal pore and halloysite magnified from the box in **(c)**.

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**Fig. 8.** Calcite-rich dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle, **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d, e)** TEM images magnified from the box in **(c)**.

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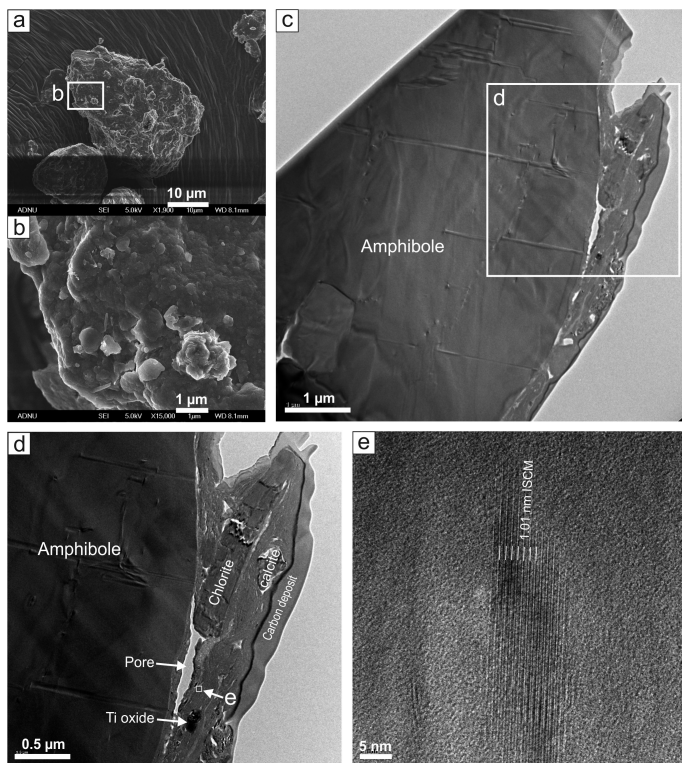
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**Fig. 9.** Amphibole-rich dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle, **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of ISCMs magnified from the box in **(d)**.

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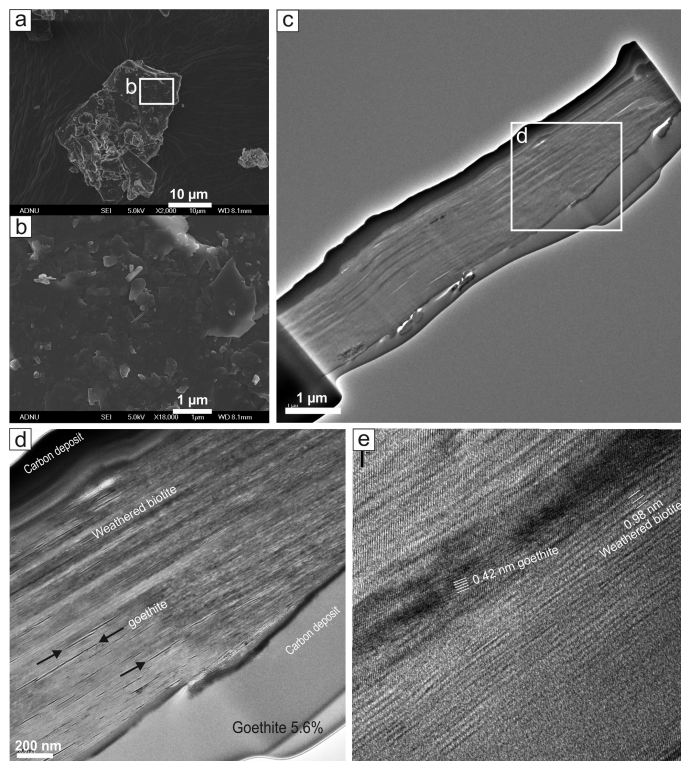
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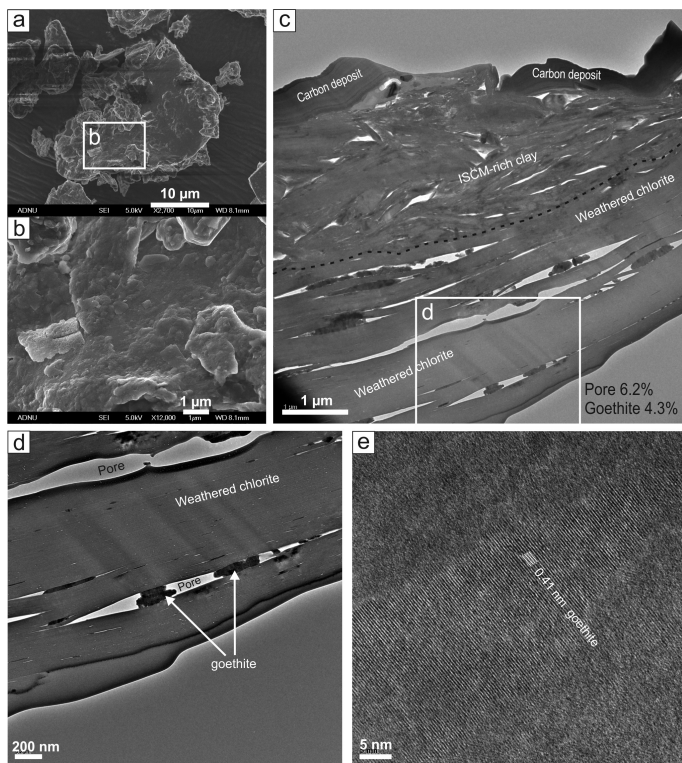




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**Fig. 10.** Biotite dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the biotite particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of goethite in weathered biotite.



**Fig. 11.** Chlorite-rich dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** SEM image of the particle surface magnified from the box in **(a)**. **(c)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(d)** TEM image magnified from the box in **(c)**. **(e)** TEM lattice fringe image of goethite in the pore of weathered chlorite.

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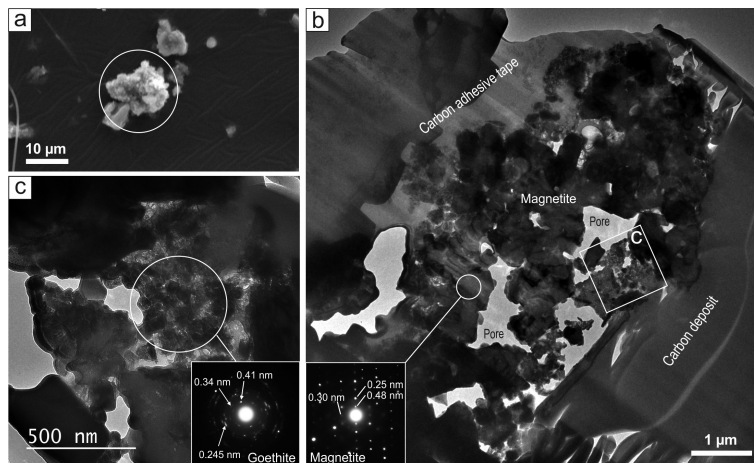
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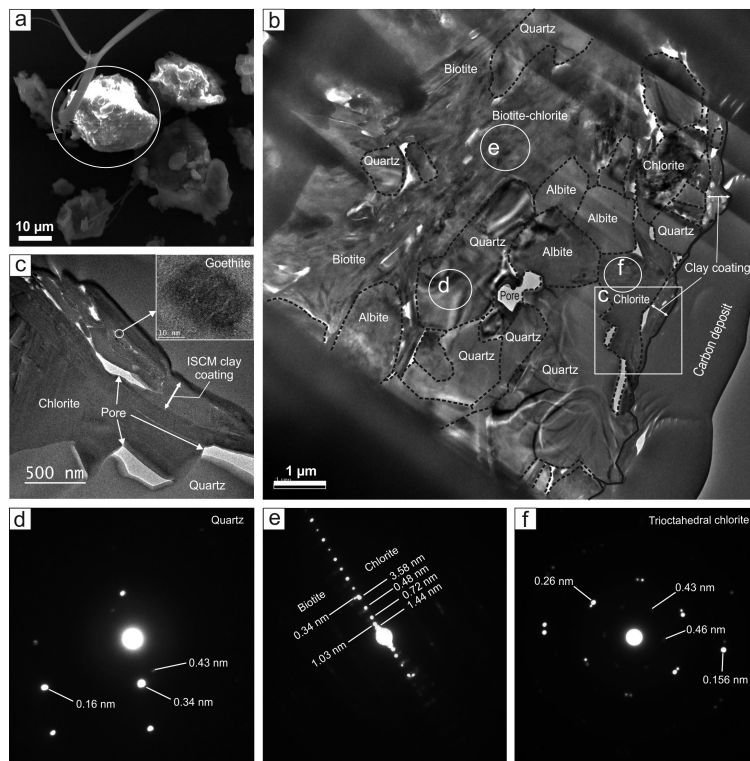
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**Fig. 12.** Iron-oxide-rich dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** Overall TEM image of the FIB slice prepared from the particle in **(a)**. **(c)** TEM image magnified from the box in **(b)**. Electron diffraction patterns of the circled areas in **(b)** and **(c)** indicate goethite and magnetite, respectively.

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**Fig. 13.** Rock fragment dust particle from 2012 Asian dust. **(a)** SEM image of the dust particle. **(b)** Overall TEM image of the FIB slice prepared from the rock fragment particle in **(a)**. **(c)** TEM image magnified from the box in **(b)** showing the coatings of oriented nano-thin clay minerals with pore and goethite nanograins. **(d–f)** Electron diffraction patterns of the areas marked with circles.

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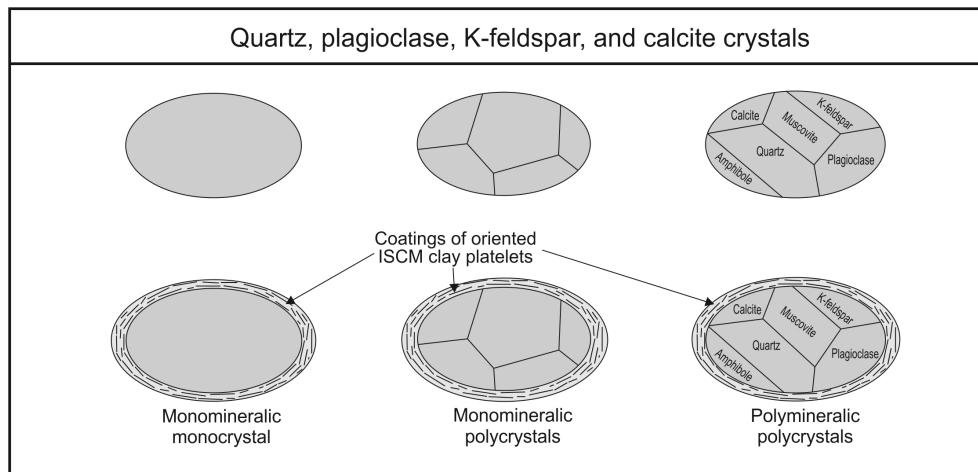
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**Fig. 14.** Structural models for single and polycrystals of quartz, plagioclase, K-feldspar, and calcite with clear or clay-coated surfaces. Clay minerals are dominated by ISCMs.

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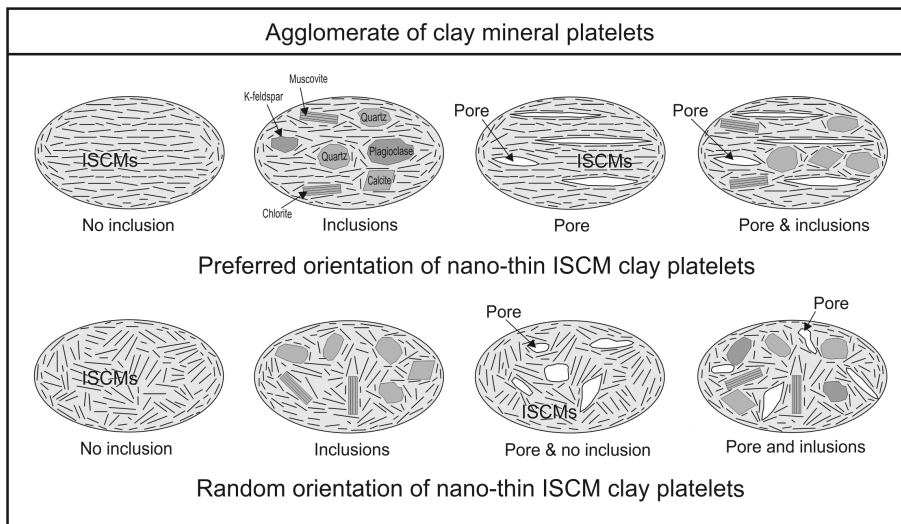
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**Fig. 15.** Structural models for ISCM-clay-rich particles with preferentially or randomly oriented nano-thin clay platelets, and with pores and inclusions of nonphyllosilicates, micas (muscovite and biotite), and chlorite.

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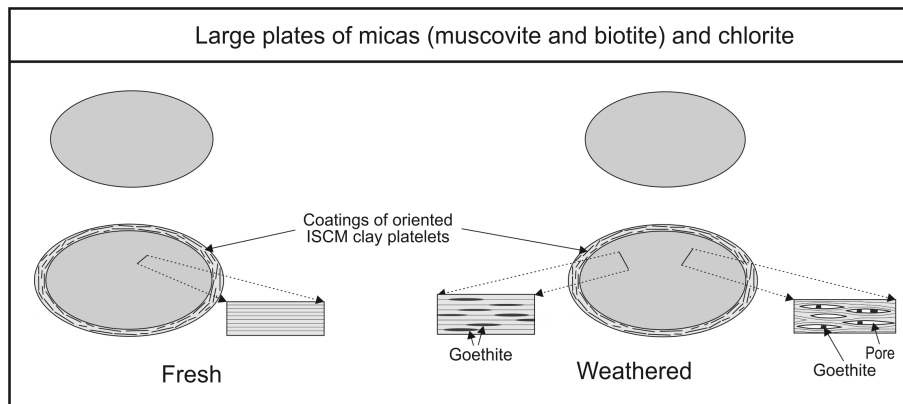
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**Fig. 16.** Structural models for plates of micas (muscovite and biotite) and chlorite, which are either fresh or weathered. Weathered biotite and chlorite contain goethite crystals and lenticular pores.

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