Reflectance spectroscopy (200–2500 nm) of highly-reduced phases under oxygen- and water-free conditions

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1. Introduction

Some common extraterrestrial minerals are extremely rare or absent in Earth materials. Important planetary materials, including oldhamite and nanophase iron, oxidize and hydrolyze rapidly in contact with Earth’s atmosphere and hydrosphere (e.g., Cooper et al., 2012; Haberle et al., 2013; Okada et al., 1981). Oldhamite is particularly susceptible to terrestrial weathering, and is rapidly altered to sulfates (bassanite, gypsum), oxhydroxides (portlandite), and carbonates (vaterite, calcite) as described by Okada et al. (1981). Nitrides and graphite are less reactive, but may still be affected by adsorbed atmospheric water or gases. We have conducted a spectroscopic study of synthetic versions of CaS (oldhamite), TiN (osbornite), CrN (carlsbergite), nanophase graphite, and iron powders (10 μm and 50 nm nominal particle sizes) under an oxygen-free anhydrous N2 atmosphere. In the subsequent discussion we use the mineralogic terms for these synthetic equivalents unless otherwise indicated. Given the considerable uncertainty concerning the extent to which known meteorites are representative of the asteroids, and the lack of any known meteorites that sample of the surface of Mercury, spectral studies of diverse highly-reduced materials are useful for understanding the properties of planetary surfaces using telescopic, orbiter and lander data sets. Objects which may contain highly-reduced mineral assemblages include the parent asteroids of enstatite chondrites, aubrites, iron meteorites, pallasites, mesosiderites (or similar materials not present in terrestrial meteorite collections), and the surface of Mercury.

2. Methods

The samples used in this study are described in Table 1. All of the samples are air-sensitive. Special procedures were implemented to preserve their integrity. Spectra were collected in a Plas-Labs 818 GBB glovebox with interior dimensions of 152.4W × 96.5D × 79H cm (Plas-labs Inc., Lansing, MI) under a dry N2 atmosphere, where dry nitrogen was passed through a series of Drierite filters. The glovebox also contained Drierite and sieves of Drierite filters. The glovebox also contained Drierite and sieves of Drierite filters. The glovebox also contained Drierite and sieves of Drierite filters. The glovebox also contained Drierite and sieves of Drierite filters.

Note

Spectra of highly-reduced mineral phases from 200 to 2500 nm provide new laboratory constraints on the surfaces of asteroids and other extremely reduced solid assemblages. Synthetic oldhamite (CaS) is distinguished by high ultraviolet reflectance (which decreases toward shorter wavelengths). Oldhamite and osbornite spectra show absorption features at ~401 nm and ~436 nm, respectively. Chemically pure synthetic oldhamite is spectrally distinct from naturally-occurring oldhamite from the Norton County aubrite, possibly due to differences in minor and trace element compositions, presence or absence of inclusions, or differences in oxidation/hydration (terrestrial weathering). Iron powders at 50 nm and 10 μm nominal particle sizes, nanophase graphite, and carlsbergite (CrN) all have very low reflectivity over the 200–2500 nm wavelength range. Carlsbergite has a slight blue spectral slope in the visible and near-infrared (400–2500 nm), while the iron powders and nanophase graphite show slight red slopes over this wavelength range.

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2.3. 250–2500 nm measurements (near UV, VIS, and near IR regions)

The 250–2500 nm reflectance spectra were measured with an Analytical Spectral Devices FieldSpec Pro HR spectrometer which acquires data from 350 to 2500 nm in 1.4 nm steps, with a spectral resolution of between 2 and 7 nm. The data are internally resampled by the instrument to output data ultimately at 1 nm intervals. Data below approximately 400 nm may be affected by low signal levels and hence data in the 350–400 nm region, particularly the abrupt changes in slope, are suspect. Unless otherwise indicated, spectra were measured at a viewing geometry of $i = 30^\circ$ and $e = 0^\circ$. Incident light was provided by an in-house 100 W quartz–tungsten–halogen collimated light source. Sample spectra were measured relative to a Spectralon (Labsphere, North Sutton, NH) standard and corrected for minor (less than ~2%) irregularities in its absolute reflectance. In each case, 200 spectra of the dark current, standard, and sample were acquired and averaged, to provide sufficient signal-to-noise for subsequent interpretation. Locations of reflectance minima in oldhamite and osborneite were determined by fitting a 3rd-order polynomial to the measured spectra.

2.2. 200–400 nm measurements (UV region)

The ultraviolet spectroscopy methods used in this study were initially described by Cloutis et al. (2008) and are briefly summarized here. The 200–400 nm spectra were measured with an Ocean Optics S-2000 spectrometer equipped with a grating that provides coverage from 200 to 859 nm with spectral resolution between 0.36 nm (at 200 nm) and 0.28 nm (at 859 nm). Illumination was provided by an Analytical Instrument Systems Inc. Mini-DTA light source with output from a 30 W deuterium lamp directed through a bifurcated fiber optic bundle consisting of six illumination fibers surrounding a central pick-up fiber feeding into the detector array. This assembly consisted of 400-μm diameter solarization-resistant (XSR) fibers, with transmission efficiencies between 23% and 40% across the 200–400 nm range. The fiber optic bundle was used in normal incidence and we used an integration time of 1000 ms and averaged 100 individual spectra. Measurements for each sample were made by first acquiring a dark current spectrum (with the input to the spectrometer blocked), a reference spectrum of our BaSO$_4$ standard (Alfa Aesar Puratronic grade: 99.998% metals basis; CAS #25583-20-4), a reference spectrum of our sample being imaged. The most appropriate definition of the instrument configuration is a biconical arrangement with $i$ and $e$ centered on ~0° and detector and illumination fields of view of 25.4°. The spectra were corrected for irregularities in the BaSO$_4$ standard using a calibrated deep UV mirror as described in Cloutis et al. (2008).

2.3. Note on terminology for fine iron particles

Various terms have been used in the planetary science literature for very small iron particles thought to influence the spectral properties of airless bodies, including nanophase iron (npFe) (Keller and McKay, 1993; Pieters et al., 2000), submicroscopic metallic iron (SMFe) (Hapke, 2001), and Brit-Pieters particles or microphase iron (Lucey and Riner, 2011; Riner and Lucey, 2012). In the present study, we have used commercially available iron powders of 50 nm and 10 μm particle sizes. The 50 nm powder consists of nanophase iron, applying the widely accepted definition adopted by the American Society for Testing and Materials (ASTM) “nanoparticle: sub-classification of ultrafine particle with lengths in two or three dimensions greater than 0.001 μm (1 nm) and smaller than about 0.1 μm (100 nm) and which may or may not exhibit a size-related intensive property” (ASTM, 2012). The 10 μm powder would be classified as a coarse particle according to the ASTM definition “a particle smaller than about 10 μm and larger than about 2.5 μm in size” (ASTM, 2012). For the 50 nm powder, we have used

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Albedo (560 nm) (%)</th>
<th>Reflectance ratios (wavelength positions given in nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>2400/560</td>
</tr>
<tr>
<td>CA5001 (oldhamite)</td>
<td>62.35</td>
<td>1.12</td>
</tr>
<tr>
<td>TIN001 (osborneite)</td>
<td>10.59</td>
<td>2.95</td>
</tr>
<tr>
<td>CAR101 (carlsbergite)</td>
<td>5.33</td>
<td>0.67</td>
</tr>
<tr>
<td>GRP200 (np graphite)</td>
<td>0.97</td>
<td>1.23</td>
</tr>
<tr>
<td>IRO103 (np iron, 10 μm)</td>
<td>2.81</td>
<td>1.48</td>
</tr>
<tr>
<td>IRO102 (np iron, 50 nm)</td>
<td>2.60</td>
<td>1.23</td>
</tr>
</tbody>
</table>
the term nanophase iron. For the 10 \( \mu \)m powder, we have used the term Britt-Pieters particles following (Lucey and Riner, 2011; Riner and Lucey, 2012). Where both iron particle sizes are referred to simultaneously, we have used the term iron powder.

3. Results

3.1. 350–2500 nm observations (near UV, VIS, and near IR)

Spectral slope is here defined as the ratio of reflectance at 2500 nm to that at 560 nm. Albedo and spectral slope parameters are summarized in Table 2, following (Cloutis et al., 2010). Oldhamite has a high albedo (62.35% at 560 nm) and a strong absorption feature with a minimum near 401 nm and shoulders centered near 546 nm and 763 nm, and possible weak features at \( \sim 1150 \) nm and \( \sim 1380 \) nm. Osbornite has an absorption feature with a minimum near \( \sim 436 \) nm and 10.59% reflectance at 560 nm (Fig. 1A and B). Carlsbergite is dark (5.33% reflectance at 560 nm) and is spectrally featureless with a blue slope (Fig. 1A and B). Neither of these phases has been spectrally characterized before.

Iron powders, at both 10 \( \mu \)m and 50 nm nominal particle sizes, have dark (2.81% and 2.60% reflectance at 560 nm respectively) and featureless spectra with nearly flat to slightly red spectral slopes (Fig. 1A and B; cf. Fig. 5d of Noble et al. (2007), for 50 nm particles). The similar spectral properties of these two iron particle sizes are somewhat surprising as the finer-grained sample is expected to be more red-sloped than the larger size (Lucey and Riner, 2011). It is likely that the similar spectral slopes are due to the fact that we are measuring pure powders as opposed to grains dispersed in a more transparent matrix (Noble et al., 2007). It is possible that the 50 nm particles are aggregating to create much larger effective particle sizes, as Rayleigh scattering would be expected to dominate for 50 nm particles, as the size parameter \( \pi D / \lambda \) is less than unity for the entire wavelength range studied here (Lucey and Riner, 2011). Nanophase graphite has extremely low albedo (0.97% at 560 nm), is spectrally featureless, and has a slight red slope (Fig. 1A and B). Its spectral properties are similar to other graphites that have been spectrally characterized, all of which show red-sloped spectra with a minimum in the 0.7 \( \mu \)m region (Cloutis et al., 1990, 1994). The lack of a minimum near 0.7 \( \mu \)m and the overall

Fig. 1. Absolute reflectance spectra of CaS (oldhamite, CAS001), TiN (osbornite, TIN001), CrN (carlsbergite, CAR101), nanophase graphite (carbon black, GRP200) and iron powders: 10 \( \mu \)m Britt-Pieters particles (IRO103), and 50 nm nanophase iron (IRO102). (A) Near-UV, visible and near infrared spectra collected with \( \iota = 30^\circ \) and \( \epsilon = 0^\circ \), with the reflectance of each spectrum at 400 nm scaled to match the reflectance measured by the ASD spectrometer at 400 nm. The narrow peaks in the UV spectra (positions denoted with small symbols below) are attributable the deuterium light source (Cloutis et al., 2008).
shape of graphite spectra are very sensitive to small differences in composition and structure (e.g., Blanco et al., 1988; Bussoletti et al., 1978; Draine, 1985; Rouleau and Martin, 1991).

3.2. 200–400 nm observations (UV)

Osbornite, carlsbergite, nanophase graphite and both iron powders have monotonically increasing reflectance with increasing wavelength in the ultraviolet. Osbornite, carlsbergite, nanophase graphite and the iron powders all have low reflectance from 200 to 400 nm. Osbornite is distinctive and has a high reflectance in the near ultraviolet, increasing from ~4.5% at 275 nm to ~34% at 400 nm. The high ultraviolet reflectance of oldhamite could facilitate remote detection. Spectra of pure oldhamite in the 350–2500 nm range in the present study are markedly different from those of Burbine et al. (2002). Similar to the results by Helbert et al. (2013), our spectra do not show the absorptions at ~490 and ~950 nm, and instead show a single strong absorption centered near 401 nm with shoulders centered near 360 nm and 375 nm and possible weak features at ~1150 nm and ~1380 nm (cf. Fig. 1 of this study with Fig. 3 of Burbine et al., 2002). The oldhamite investigated in this study is synthetic and close to chemically pure (99.9%, 500 ppm Sr; see Burbine et al., 2002). The oldhamite investigated in this study is synthetically pure and carlsbergite has not previously been measured over the wavelength range studied here. Osbornite (TIN) shows a visible absorption feature centered near 436 nm, probably due to charge transfer between the valance and conduction bands (N 2pₓ → Ti 3dᵧ) (Kuznetsov and Serpone, 2009; Patsalas and Logothetidis, 2001).

Observables spectral changes other than the appearance of absorption features can be caused by the presence of opaque materials like those studied here. For instance, finely-dispersed opaque phases can greatly darken a material and reduce the contrast of spectra features and can induce large changes in spectral slope (e.g., Clark, 1983; Cloutis et al., 2011; Milliken and Mustard, 2007).

4.1. Potential for detection on planetary surfaces

Oldhamite is the most likely candidate for remote detection based on its spectral characteristics including high UV reflectance and detectable spectral features in the visible and near-IR. Oldhamite-rich lithologies (commonly occurring as cliffs or veins) are known from several enstatite meteorites including the Abeel EH4 impact melt breccia (Rubin and Keil, 1983), the EL6 chondrite Jahl deh Kot Lalu (Rubin et al., 1997), the Norton County aubrite (Wheelock et al., 1994), and the Bustee aubrite (McCoY, 1998). Oldhamite is likely the product of impact melts on enstatite meteorite parent bodies (e.g., Rubin and Keil, 1983; Rubin et al., 1997). Oldhamite may be more abundant on some E-type asteroids, for instance, an oldhamite abundance of up to 42% has been suggested by Nedelcu et al. (2007) for Asteroid 2867 Steins. Clark et al. (2004) concluded from spectroscopic observations and compositional modeling that several E-type asteroids contain abundant sulfides, possibly including oldhamite, and that several E-types may not correspond with known enstatite-rich meteorites (E chondrites or aubrites). Oldhamite-rich lithologies may be widespread on the surface of Mercury. Direct measurements of the S concentration of the Hermean crust by the MErcury Surface, Space Environment, Geochemistry, and Ranging (MESSENGER) gamma ray and X-ray spectrometers showed high S abundances (estimated concentrations of ~4 wt.% and correlations of Mg and Ca with S suggestive of sulfide-rich lithologies (Evans et al., 2012; Nittler et al., 2011; Weider et al., 2011; Zolotov et al., 2013). The MESSENGER findings confirmed previous suggestions based primarily on radar observations for S-rich materials on Mercury’s surface (e.g., Sprague et al., 1995). Separation of CaS from sulfide melts by liquid immiscibility has been documented in both synthetic and natural systems (e.g., Floss et al., 1998; Skinner and Luce, 1971; Wheelock et al., 1994). Sulfide liquid immiscibility could therefore produce local enrichments of oldhamite in a variety of settings including impact melts (as melt sheets, ponds, and veins) and endogenic igneous rocks (e.g., Floss et al., 1998; Helbert et al., 2013; Wheelock et al., 1994). The thermal breakdown of sulfate minerals at subsolvus temperatures may play a role in the formation of ‘hollows’ on the surface of Mercury (Blewett et al., 2012; Helbert et al., 2013).

Osborneite is a trace constituent of many enstatite chondrites (e.g., Rubin, 1997), and has been reported in an oldhamite-rich lithology from the Bustee aubrite (McCoy, 1998). Osborneite, likely formed as an early Solar System condensate, has also been reported from comet Wild-2 (Chi et al., 2009). Carlsbergite is a fine-grained trace constituent of many iron meteorites (e.g., Axon et al., 1981; Buchwald and Scott, 1971).

4.2. Stellar envelopes and disks

Another, more speculative setting in which to search for spectral signatures for highly-reduced minerals is in extrasolar, astrophysical settings such as stellar disks and supernova outflows. Presolar diamonds, carbides and nitrides have been observed in primitive meteorites (e.g., Meibom et al., 2007; Nittler et al., 1995) and from comets (Chi et al., 2009). Recently, presolar oldhamite was identified as an inclusion in an SiC grain originating from
an Asymptotic Giant Branch (AGB) star (Hynes et al., 2011). The probable low concentrations of sulfides and nitriles would make detection difficult, however, both crystalline and amorphous silicates have been identified in stellar outflows (e.g., Gall et al., 2009), thus it is not inconceivable that other solid phases including some of those investigated here could also be detected under favorable circumstances. Therefore, while the spectra reported here are not directly comparable to those of dust in stellar envelopes, they may still provide some constraints on the presence of highly-reduced phases in astrophysical settings.

5. Conclusions

The highly-reduced phases studied here are characterized by a range of reflectance values, with oldhamite and osbornite having higher reflectance than the other highly-reduced phases. The presence of absorption features in these two materials opens up the possibility of their being spectrally detectable using these features. The differences in spectral properties between synthetic and meteoritic oldhamite suggests that it is possible to determine compositional variations in this material from spectral variations. It is also possible that differences in spectral properties between different subgroups of E-class asteroids (Clark et al., 2004) may be attributable to variations in oldhamite and other sulfides on these bodies.

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References

Clark, R.E. et al., 2004, E-type asteroid spectroscopy and compositional modeling. J. Geophys. Res. 105, 1–11.
Noble, S.K. et al., 2007. An experimental approach to understanding the optical effects of space weathering on Icarus 192, 629–642.
Skinner, R.J., Luce, F.D., 1971. Solid solutions of the type (Ca,Mg,Mn,Fe)3 and their use as geothermometers for the enstatite chondrites. Am. Miner. 56, 1269–1296.

