Detecting Sub-Micron Space Weathering Effects in Lunar Grains With Synchrotron Infrared Nanospectroscopy

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Abstract

Space weathering processes induce changes to the physical, chemical, and optical properties of space-exposed soil grains. For the Moon, space weathering causes reddening, darkening, and diminished contrast in reflectance spectra over visible and near-infrared wavelengths. The physical and chemical changes responsible for these optical effects occur on scales below the diffraction limit of traditional far-field spectroscopic techniques. Recently developed super-resolution spectroscopic techniques provide an opportunity to understand better the optical effects of space weathering on the sub-micrometer length scale. This paper uses synchrotron infrared nanospectroscopy to examine depth-profile samples from two mature lunar soils in the mid-infrared, 1,500 – 700 cm\(^{-1}\) (6.7–14.3 μm). Our findings are broadly consistent with prior bulk observations and theoretical models of space weathered spectra of lunar materials. These results provide a direct spatial link between the physical/chemical changes in space-exposed grain surfaces and spectral changes of space weathered bodies.

Plain Language Summary

The Moon's surface, unprotected from the space environment, is bombarded with solar wind ions and micrometeoroids. These interactions are part of a process known as space weathering, which changes the physical and optical properties of lunar soils and asteroid surfaces on a microscopic scale. Technological hurdles have hindered our understanding of the connection between the physical changes caused by space weathering and the optical properties thought to result from them. Using synchrotron infrared nanospectroscopy, we examined how various weathering processes affect the infrared spectral characteristics of lunar soil grains. With these insights, we can develop better space weathering models to predict how different surfaces may be affected. The data from this investigation can also be used to calibrate laboratory analog studies of space weathering and help interpret observations of bodies similar to the Moon.

1. Introduction

The Moon is subject to frequent micrometeoroid impacts and bombardment by energetic solar wind ions. The compositional and structural changes induced by these processes on the Moon and other airless bodies are collectively referred to as space weathering (Hapke, 2001; Pieters et al., 1993). In aggregate, these changes to the morphology, chemical composition, and crystal structure of individual regolith grains alter the optical properties of the bulk soil—relative to freshly exposed lunar regolith, reflectance spectra of space exposed soils have reddened, darkened continua, and weaker diagnostic absorption peaks in the visible to infrared (IR) wavelengths. These effects have also been observed in studies of S-type asteroidal surface soils (Noguchi et al., 2011, 2014) and simulated space weathering experiments (Kaluna et al., 2017; Lantz et al., 2017; Thompson et al., 2019).

The effects of space weathering occur on a spatial scale comparable to the wavelength of visible light, presenting a unique challenge to our understanding of how various weathering processes evolve and interact to produce optical changes. The physical changes induced by space weathering, including the production of nano-phase iron particles and damage to the soil's crystal structure (i.e., amorphization), have been found to occur predominantly within 100–200 nm of the grain surface (Noble et al., 2005; Pieters et al., 1993, 2000; Taylor et al., 2001). Hence, electron microscopy techniques are well-suited to characterize microstructural and micro-compositional changes. For instance, transmission electron microscopy
Journal of Geophysical Research: Planets

TEM of weathered lunar soils has demonstrated that many of the optical changes seen in weathered soil are associated with the presence of nano-phase iron (npFe\textsuperscript{0}) particles in amorphous rims coating mineral grains (Keller & McKay, 1993, 1997; Taylor et al., 2001, 2010), and micro-phase iron that occurs in agglutinates (Basu, 2005). In particular, npFe\textsuperscript{0} grains smaller than 40 nm in diameter cause spectral reddening and darkening, while larger iron particles cause only darkening (Lucey & Riner, 2011; Noble et al., 2007). Although the physical and chemical changes caused by space weathering can be detected via TEM, the localized optical effects of these changes cannot be directly interrogated using diffraction-limited spectroscopic techniques. Traditional diffraction-limited spectroscopic techniques cannot spatially resolve features much smaller than the wavelength of light used—most npFe\textsuperscript{0} is <40 nm in diameter. Until recently, computational modeling was required to determine the cumulative effects of space weathering on the optical properties of lunar soil (Hapke, 2001; Lucey & Noble, 2008; Lucey & Riner, 2011; Wohlfarth et al., 2019).

To bridge the gap between the optical effects of space weathering and the nano-scale physio-chemical phenomena that produce them, we used Synchrotron Infrared Nano Spectroscopy (SINS) to collect IR spectral data with sub-micrometer spatial resolution from depth-profile samples of space-exposed lunar soil. This technique is capable of ~20 nm spatial resolution, making it possible to assess the optical effects of weathering phenomena at a spatial resolution sufficient to resolve sub-micrometer products of lunar space weathering (Bechtel et al., 2014). Near-field infrared spectroscopy has previously been employed to analyze other extraterrestrial or planetary materials, including the Murchison meteorite (CM2; Kebukawa et al., 2010), a grain (Iris) from comet 81P/Wild 2 (Dominguez et al., 2014), and the Didim meteorite (H3-5; Yesiltas et al., 2020). This paper presents near-field infrared spectroscopic evidence of space weathering induced changes to mature lunar soils’ optical properties in the “fingerprint region” of the mid-infrared (1,500–700 cm\textsuperscript{-1}; 6.7–14.3 μm).

Figure 1. The geospatial context for lunar soil 79221 (samples 1–3). (a) NASA photograph AS17-142-21827 showing approximate in situ sample location (circled) as recorded at the time of collection. (b) 2 kV SE image of host grain for samples 1–3, extracted from the circled areas. The encircled white rectangles indicate the orientation of each sample. (c)–(e) 2 kV SE images of samples 1–3, respectively, on Si substrate after thinning and low-voltage polishing.
2. Materials and Methods

2.1. Sample Preparation

The examined samples were selected from fine-grained portions of, shown in Figures 1 and 2. The degree of surface exposure (maturity) of lunar soils is typically indicated by the ferromagnetic resonance (FMR) surface exposure index, given as FMR intensity divided by iron content ($I_f/FeO$) — see Morris (1976) for more detail. By this metric, both 79221 and 10084 are highly mature soils with $I_f/FeO$ values of 81 and 78, respectively (Rhodes & Blanchard, 1982; Taylor et al., 2001). To prepare the soils for sectioning, we secured a portion of each sample to an aluminum stub with carbon tape. We subsequently coated both sample and stub with $\sim 10$ nm of gold-palladium. The coated grains were imaged in secondary and back-scattered electrons using a Tescan Mira3 field-emission scanning electron microscope (FEG-SEM). An EDAX energy-dispersive X-ray (EDX) spectrometer on the SEM was used for elemental analyses and mineral identification.

As described in greater detail below, SINS utilizes an atomic force microscope (AFM) tip to enhance near-field resonances at a sample surface. To interrogate spectral response changes as a function of depth, we created depth-profile lift-outs from the grains identified via SEM-EDX. Qualitative markers of space exposure (e.g., surface blistering, micrometeoroid impact craters, and melt splash) were used to inform the site-selection for targeted liftout extraction. Lift-outs (initial thickness $\sim 1 \mu$m) were extracted from space-exposed regions of the target grains with an FEI Quanta 3D focused ion beam (FIB) equipped with a computer-con-
trolled Omniprobe micro-manipulator. These samples were transferred to an Omniprobe lift-out grid, upon which they were thinned to a thickness of 300–600 nm. Each sample was subsequently polished with a low-energy (5 kV, 48 pA) Ga⁺ beam for roughly two minutes per side to remove any surface damage created during the thinning procedure (Kato, 2004). The thinned lift-outs were then placed onto an ultra-flat (surface roughness <0.5 nm) Si chip.

Samples 1–3 were taken from a ~250 μm grain of 79221 with a composition consistent with anorthite-rich plagioclase (see Table 1). Sample 4 was extracted from a ~150 μm grain of 10084 with a composition consistent with Ti-, Al-rich augite, in agreement with prior studies of this sample and other Apollo 11 lunar rock samples (Ross et al., 1970). A fifth sample was taken from a terrestrial anorthite standard (Miyake Island, Japan). The studied samples and their characteristics are outlined in Table 2.

2.2. Experimental Methods

Near-field IR spectra were collected using SINS at Beamline 5.4 at the Advanced Light Source (Bechtel et al., 2014). This technique can be thought of as a combination of Fourier transform infrared spectroscopy (FTIR), scattering-type scanning optical microscopic (s-SNOM) techniques, and atomic force microscopy (AFM). Synchrotron IR light is coupled into an asymmetric Michelson interferometer consisting of a beam-
## Table 1

| Sample | Composition (atomic percent) | Description |
|--------|-----------------------------|-------------|
| 1–3    | O 60% Mg — Al 16% Si 16% Ca 8% — Ti 3% Fe — | Melt-splash coated Ti-, Al-rich Augite |
| 4      | O 57% Mg 6% Al 5% Si 17% Ca 4% Ti 3% Fe 7% | Mildly amorphized surface An-rich Plagioclase |

Abbreviations: EDX, energy-dispersive X-ray; SEM, scanning electron microscope.

The data presented in this work were collected over a broad range of mid-infrared wavenumbers, 5,000–700 cm$^{-1}$ (2.0–14.3 μm), with a spectral resolution of 8 cm$^{-1}$. At shorter wavelengths, however, the signal is dominated by noise caused by reduced tip-sample coupling. This high-frequency noise precludes the identification of any C-H or O-H stretch features in the sample spectra. As such, this work focuses primarily on the “fingerprint” region, 1,500–700 cm$^{-1}$ (6.7–14.3 μm). This range captures key features in the infrared spectra of both plagioclase and pyroxene while maximizing the signal-to-noise ratio. Of particular interest to lunar and remote-sensing applications, the explored spectral range encompasses the Christiansen feature—an important diagnostic feature in mid-infrared silicate spectra. Canonically, the CF is defined as an emissivity maximum associated with the frequency at which the real part of the effective dielectric constant (index of refraction) approaches unity (Christiansen, 1884). Since this condition occurs at wavelengths just short of the fundamental modes, the CF contains valuable information about silicate mineral composition (Conel, 1969; Salisbury et al., 2012).

To differentiate the near-field signal from the far-field (scattered) background, signals are detected at higher harmonics of the tip oscillation frequency, which arise from the non-linear near-field response. Here, we use the second harmonic response as a compromise between background suppression and signal-to-noise ratio. After demodulation, the interferometric signal is Fourier transformed to yield the complex near-field spectra. To first-order approximation, the spectral amplitude, $|A(\tilde{v})|$, is related to the real-valued component of the material’s complex dielectric function (i.e., the reflection coefficient) and the spectral phase, $\Phi(\tilde{v})$, is similarly related to the imaginary component of the dielectric function (i.e., the absorption coefficient; Govyadinov et al., 2014; Xu et al., 2012). However, this approximation may not be strictly valid due to the thickness of the samples studied, the presence of nanoscale heterogeneities therein, and variable oscillator strengths. Spectral features may therefore be shifted compared to conventional FTIR measurements (Mastel et al., 2015).

We collected a series of line scans oriented perpendicular to the space-exposed surface (vertically) to interrogate the effects of space weathering as a function of depth. Each line scan is composed of 20–60 evenly distributed points with an inter-point spacing of 20–100 nm. Three horizontally oriented (i.e., parallel to the grain surface) scans were collected to rule out systematic instrumental artifacts as the cause of observed depth-dependent changes. Background spectra were collected before and after each line scan. The experimental spectra were referenced to the average over relevant backgrounds. The location and orientation of each line scan are shown in Figure 4 and further details can be found in Table 3.

## 2.3. Data Analysis

The data presented in this work were pre-processed using a custom program that employs commercially available fast Fourier transform software packages (Wavemetrics Igor Pro). Background and instrumental responses were removed by referencing the amplitude and phase signals to a background spectrum collected on Si or Au before and after each sample scan. The referenced amplitude and phase signals were obtained via, respectively,

$$|A(\tilde{v})| = \frac{|A(\tilde{v})|_{\text{sample}}}{|A(\tilde{v})|_{\text{reference}}} \quad \text{and} \quad \Phi(\tilde{v}) = \Phi(\tilde{v})_{\text{sample}} - \Phi(\tilde{v})_{\text{reference}}. \quad (1)$$

| Sample | Soil | Composition | Description |
|--------|------|-------------|-------------|
| 1      | 79221| An-rich Plagioclase | Micrometeoroid impact crater |
| 2      | 79221| An-rich Plagioclase | Melt-splash coated |
| 3      | 79221| An-rich Plagioclase | Surface blistering |
| 4      | 10084| Ti-, Al-rich Augite | Mildly amorphized surface |
| —      | —    | Anorthite | Terrestrial mineral standard |
SINS amplitude spectra are generally more susceptible to topographical and instrumental artifacts than phase spectra. This susceptibility is partly because amplitude spectra typically present dispersive lineshapes, whereas phase spectra generally occur as Gaussian or Lorentzian profiles, potentially making weak features more difficult to see in amplitude data. Phase spectra have been shown to closely track the material’s local absorption coefficient (Stiegler et al., 2011; Taubner et al., 2004), which is crucial for the depth-profile studies in this work. As such, we will focus primarily on phase spectra to examine the relative changes of particular spectral features as a function of depth.

The collection depth for each spectrum was calculated relative to the bottom edge of the protective Pt cap. The interface of the space-exposed surface and the Pt cap was located by overlaying high-resolution SE images (in which the Pt was visually distinct from the sample) atop the AFM topographical images used for SINS target selection. Spectra collected from the Pt cap were not used for the analyses described below.

Figure 4. Close-up SE images (2 kV) of the samples shown in Figures 1c–1f and 2d, upon which each line scan site is superimposed. The direction of the scans oriented parallel to the space-exposed surface is indicated with an arrow. All vertically oriented scans start in the grain interior and end near the surface. (a) Sample 1 (79221), scans 1–3. (b) Sample 2 (79221), scans 1 and 2. The second scan on this sample is located below a melt-droplet (indicated by the red arrow) that was adhered to the surface. (c) Sample 3 (79221), scans 1–5. The first and second scans are located below an unusually thick vesiculated melt texture. Note that scan 4 ends roughly 2.4 μm from the surface. (d) Sample 4 (10084), scans 1–4. The first and third scans are oriented vertically, whereas scans 2 and 4 were collected parallel to the grain surface at different depths. (e) Scans 1 and 2 from the terrestrial anorthite standard, respectively oriented vertically and horizontally. Note that the discoloration (the dark square in the upper right) is a temporary charging effect caused by Ga+ ion beam use immediately prior to image capture. (f) An example demonstrating the alignment of the atomic force microscope (AFM) topographical image (overlaid at 50% opacity) over a reference 2 kV secondary electron (SE) image. We found general agreement in all AFM channels (i.e., tapping phase, amplitude, and topography), regarding the position of each scan and the Pt cap. However, the topographical image was used for navigation and is thus shown here.

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To assess the validity of the observed qualitative spectral changes over depth, we used a robust, iterative, non-linear least squares fitting (or peak deconvolution) algorithm. Peaks in phase spectra were fit to Lorentzians with a linear baseline, following the Lorentz model for dielectrics. The constraints and initial values used for the fitting procedure were similarly physically motivated. Together, these factors improved the efficiency and likelihood of convergence for our analyses. This approach is loosely similar to the Modified Gaussian Model (MGM) developed by Sunshine et al., 1990, with some fundamental changes implemented to account for the differences between SINS and far-field infrared spectroscopy. It should be noted that the MGM, however, is typically used with UV-visible or near infrared data. Similar fitting procedures have
been used previously to analyze a suite of extraterrestrial materials such as Martian meteorites (Sunshine et al., 1993), remote sensing data from Mars (Mustard & Sunshine, 1995), and lunar soils (including the two soils studied here; Noble et al., 2006). For a more thorough treatment of our quantitative model, please refer to the Supplementary Material.

2.4. Terrestrial Standard

For purposes of comparison, two scans were collected from a terrestrial anorthite standard. The phase spectra from these two scans are plotted in Figure 5. Spectra from the anorthite standard feature two strong absorption peaks likely corresponding to Si–O–Si asymmetric stretch Restrahl–en bands (Carmichael, 1988; Le Bras et al., 2003). The peak at approximately 1,150 cm$^{-1}$ (8.70 μm) is sharp and distinct (FWHM $\approx$ 60 cm$^{-1}$), whereas the peak at roughly 1,035 cm$^{-1}$ (9.66 μm) is substantially broader (FWHM $\approx$ 170 cm$^{-1}$) due to the presence of a shoulder roughly centered about 965 cm$^{-1}$ (10.36 μm). These features' positions and lineshapes closely match previously reported absorption spectra for anorthite-rich plagioclase (Estep et al., 1971; Williams & Jeanloz, 1989). Given this correspondence, the shallow trough at approximately 1,230 cm$^{-1}$ is likely related to the CF. Accordingly, the broad minimum centered at ~840 cm$^{-1}$ is likely associated with a low-phase feature between vibrational modes.

Table 3
Parameters for Each Line Scan performed

| Sample | Scan | Length (μm) | Points | Spacing (nm) | Depth(s)          |
|--------|------|-------------|--------|--------------|------------------|
| 1      | 1    | 1.70        | 18     | 95           | 0–1.70 μm        |
| 2      | 1    | 1.18        | 22     | 55           | 0–1.18 μm        |
| 3      | 3    | 0.89        | 14     | 65           | 0–0.89 μm        |
| 2      | 1    | 1.93        | 22     | 90           | 0–1.93 μm        |
| 2      | 2    | 1.61        | 21     | 75           | 0–1.61 μm        |
| 3      | 1    | 0.52        | 5      | 105          | 0–0.52 μm        |
| 2      | 2    | 1.28        | 18     | 70           | 0.10–1.38 μm     |
| 3      | 3    | 1.28        | 65     | 20           | 0.34–1.62 μm     |
| 4      | 4    | 2.45        | 50     | 50           | 2.34–4.79 μm     |
| 5      | 5    | 0.75        | 11     | 70           | 30–750 nm        |
|        | 4    | 0.77        | 17     | 45           | 0–770 nm         |
|        | 2    | 1.45        | 29     | 50           | 4.08–4.64 μm     |
|        | 3    | 0.80        | 17     | 45           | 0–800 nm         |
|        | 4    | 0.89        | 30     | 30           | 1.02 μm          |
|        | 2    | 1.84        | 24     | 75           | 0–1.84 μm        |
|        | 5    | 1.90        | 20     | 95           | 6.35 μm          |

Anorthite Standard

Note. Italic text denotes horizontally oriented scans (i.e., parallel to the space-exposed surface). Spectra were collected with 8 cm$^{-1}$ spectral resolution. At each point, a 500-scan measurement was collected over roughly six minutes. Backgrounds were collected before and after each line scan.

Figure 5. Line scan phase spectra collected on the terrestrial anorthite standard. Scan 1 (top panel, plotted in blue) is oriented perpendicular to the surface and starts in the grain interior. Scan 2 (bottom panel, plotted in green) is oriented parallel to the grain surface. Spectra collected at depth are plotted in darker colors, whereas the lighter colors indicate spectra collected from near the surface. The Christiansen feature (CF) and a low-phase feature (LPF) are indicated with arrows.
seen in the standard (peaks at roughly 1,040 and 1,150 cm\(^{-1}\)), but also contain some features not observed for the terrestrial standard. By contrast, sample 1 is characterized by spectra with relatively weak and broad features, making detailed interpretation challenging. Potential explanations for this divergence are explored further in the Discussion.

The depth-dependent spectral effects seen among samples from 79221 were not observed in spectra collected from the mineral standard, indicating that they are unlikely to have arisen due to instrumental effects (see Figure 5). Moreover, line scans collected from the terrestrial standard at a constant depth were not found to differ significantly from those collected at variable depths, offering supporting evidence that the observed variations result from space weathering-induced microstructural and chemical changes in the uppermost layers of lunar soil grains.

3. Results

3.1. Micrometeoroid Impact Crater (Sample 1)

SINS phase spectra, \(\Phi(\tilde{\nu})\), collected near the hypervelocity impact crater on sample 1 (see Figure 3) exhibit systematic variations between the grain interior (far from the crater bottom) and the grain surface (just below the crater). Primary among these variations is the loss of spectral contrast with increasing proximity to the surface, as shown in Figure 7 and described below.

Two peaks in the phase spectra at 830 cm\(^{-1}\) (12.05 \(\mu\)m) and 1,140 cm\(^{-1}\) (8.77 \(\mu\)m) broaden and display reduced spectral contrast with increasing proximity to the surface. These effects are most notable within 300 nm of the surface. Near the surface, the loss of spectral contrast causes the CF to become indistinguishable from the background.

3.2. Melt-Splash Coating (Sample 2)

Sample 2 was collected from a melt-splash coated region approximately 150 \(\mu\)m from the hypervelocity impact crater on sample 1 (see Figure 3). The spectra from this sample are less noisy and contain sharper peaks than those described in Subsection 3.1. As shown in Figure 8, these spectra evolve as a function of depth similar to those from sample 1. We observed depth-dependent loss of spectral contrast, particularly at longer wavelengths. This effect is most prominent among spectra collected from within 400 nm of the surface.

Spectra from the grain interior include several peaks in the range ~950–725 cm\(^{-1}\). At the surface, these features are difficult to distinguish from the background conclusively. The provenance of these features is explored further in the discussion. A “reddening” baseline accompanies this trend at wavelengths \(\geq 11 \mu\)m, wherein the apparent slope transitions from negative to slightly positive between depths of 890 and 400 nm. Two prominent peaks occur at 1,045 cm\(^{-1}\) (9.6 \(\mu\)m) and 1,165 cm\(^{-1}\) (8.6 \(\mu\)m) in spectra from all sampled depths. These features remain relatively stable over depth, with only some statistically insignificant broadening observed near the surface.

3.3. Surface Blistering (Sample 3)

The spectra from sample 3, characterized by its evidence of surface blistering (see Figure 3), are shown in Figure 9. We observed depth-dependent loss of spectral contrast, particularly at wavelengths of 8.5–12 \(\mu\)m. This effect is most prominent among spectra collected from within 250 nm of the surface.
Importantly, this sample's spectra contain features consistent with imperfect background subtraction. In particular, the peak at \( \approx 1,350 \text{ cm}^{-1} \) is consistent with some signal from the silicon substrate “bleeding through” the sample. In contrast to the sample's diagnostic features, this peak is stronger at the surface than in the grain interior.

3.4. Mildly Amorphized Pyroxene (Sample 4)

As shown in Figure 10, the spectra from sample 4 display few systematic variations over depth. Potential explanations for this are explored in the Discussion. In contrast with the spectra from samples 1–3 and the terrestrial anorthite standard, which contain several identifiable features, the spectra from this pyroxene sample are dominated by a roughly symmetric, prominent, broad peak at 1,040 \text{ cm}^{-1} (9.6 \mu m). Although this peak's intensity remains roughly constant at all depths, its width increases slightly at the surface. At depths greater than 550 nm, there appears to be a weak feature at \( \approx 750 \text{ cm}^{-1} \) (13.3 \mu m) that broadens near the surface. However, this feature should be interpreted with caution given the variability of these spectra at low wavenumbers (long wavelengths).

3.5. Depth-Dependent Spectral Effects

The collected data for samples 2 and 3 indicate that the total scattered intensity and spectral contrast are inversely correlated with distance from the space-exposed surface. Peaks present in SINS amplitude spectra from close to the surface of these two samples are significantly less distinguishable from the continuum than in spectra collected from the crystalline grain interior (see, e.g., Figure 8). Figure 11 illustrates this effect centered about a peak at 1,145 \text{ cm}^{-1} found in the phase spectra of sample 3. In scans from sample 2, peaks at higher wavenumbers were more effectively suppressed than those at lower wavenumbers. Whether this trend extends to samples 1 and 4 remains unclear.
The integrated amplitude response, analogous to total scattered intensity, was observed to evolve over depth (see Figure 12). This quantity was calculated for each collected spectrum by integrating the amplitude signal over the wavenumbers of interest \( \nu \in [700 \text{ cm}^{-1}, 2000 \text{ cm}^{-1}] \). In scans from samples 1–3, the total scattered intensity is strongly correlated with depth (i.e., spectra from close to the space-exposed surface are darker than those from within the grain interior). This darkening effect occurs in samples 1–3 over depths of 0–2,000 nm. In samples 2 and 3, darkening is most pronounced in the uppermost 500 nm. Interestingly, the darkening in spectra from near the impact crater (sample 1) occurs at a shallower slope than the other samples.

Data from sample 4 do not display a strong correlation between scattered intensity and depth.

4. Discussion

Many minerals have qualitatively different near-field and far-field IR spectra. Though the two spectra may share some features, there is generally no one-to-one correlation (Hermann et al., 2014; Huth et al., 2012; Pollard et al., 2015). This disparity means that we cannot definitively link spectral features observed in our samples to particular vibrational modes without further analysis. Despite this, many of the features we observed in our samples’ SINS spectra are consistent with the characteristic absorption features reported in the literature.

The \( \sim 830 \text{ cm}^{-1} \) feature observed with varying prominence in samples 1–3 appear to be related to the Si–O–Si or Al–O symmetric stretch features. Although prior work modeling the optical constants of labradorite has not found a similar peak at roughly 830 cm\(^{-1}\) (12.05 μm) in spectra of the complex coefficient (Ye et al., 2019), these peaks are present in a majority of the spectra from samples 1-3 and appear to follow a depth-dependent trend similar to other peaks. Similar features have additionally been reported in mid-in-
Figure 9. Computed fits of selected synchrotron infrared nano spectroscopy (SINS) phase spectra from scan 5 on sample 3, collected at the labeled distances from the blistered surface of the plagioclase grain shown in Figures 3c and 4c. Spectra are vertically offset (dashed lines) from one another for clarity. Shaded areas indicate 2σ confidence intervals. Diminished spectral contrast was observed at the surface for several diagnostic features. Key spectral features at 750, 1,015, and 1,145 cm$^{-1}$ are discussed in greater detail in the text.

Figure 10. Computed fits of selected synchrotron infrared nano spectroscopy (SINS) phase spectra from scan 1 on sample 4 at a range of depths from the blistered surface (see Figure 3d). Spectra are vertically offset (dashed lines) from one another for clarity. Shaded areas indicate 2σ confidence intervals. Spectral features were not observed to undergo substantial changes over depth, as discussed in greater detail within the text.
frared absorption spectra of anorthite and albite (Dorschner, 1971). That this feature is more pronounced for these samples result from the presence of the hypervelocity impact crater on the host grain for samples 1–3 and the high Al-content relative to other plagioclase minerals. Al-O tetrahedra are more susceptible to deformation under pressure than their Si-O counterparts (Johnson et al., 2003; Williams, 1998; Williams & Jeanloz, 1989). It has been speculated that (Si, Al)-O tetrahedra are susceptible to metastable “defects” under pressure that lead to, for example, Si–O–Si links between adjacent tetrahedra that may alter the stretch and bending vibrational modes (Santamaria-Perez et al., 2016). Alternatively, it is hypothetically possible that these peaks could be an indication of silanol (Si–O–H) that formed on the surface of the sample or on the substrate underneath the sample. Since the spectra were normalized to an average of standard spectra, which included scans collected on the Si-chip or nearby platinum cap, this scenario is highly unlikely.

Both soils 79221 and 10084 are classified as mature, with FMR maturity indices of L/FeO = 81 and 75, respectively (Morris, 1978). However, the maturity index is by definition a bulk property of soils. As such, the individual grains that comprise a mature soil are likely to have various exposure ages. Although sample 4 displays far less pronounced space weathering effects than samples 1–3, this is more likely to be a reflection of the different mineral chemistry.

Previous studies on experimentally shocked feldspars have shown that absorption bands weaken and broaden due to increasing glass content, particularly at shock pressures above ~20 GPa (Johnson et al., 2002, 2003; Nash et al., 1993). In contrast, pyroxenes are more resilient to increasing shock pressures. Studies show little change in spectral properties with shock pressures of 45 GPa and up to 65 GPa (Adams et al., 1979; Johnson et al., 2002).

Shock effects may also be responsible for the apparent dissimilarity between spectra from sample 1 and those from the anorthite standard. The region directly below the micrometeoroid impact crater (on sample 1) experienced much greater pressures than the material in samples 2 and 3. The most prominent feature at roughly 1,100 cm⁻¹ in phase spectra from sample 1 may result from a shock-induced spectral broadening of the 1,000 and 1,150 cm⁻¹ features seen for samples 2–3 and the terrestrial anorthite standard. These differences could alternatively be explained by the presence of compositional or structural inhomogeneities in the soil grain. Should this interpretation be correct, our observations serve to reinforce the value of SINS for spectroscopic investigation of micrometer-scale mineralogical variations. The signal produced by diffraction-limited techniques is an average over various mineral structures or compositions, making it unlikely that such minor deviations in chemical composition would be detectable.

In samples 2 and 3, we observed reduced total scattered intensity with increasing proximity to the space exposed surface (see Figure 12). This darkening may be associated with an increasing concentration of npFe⁰ near the surface (Lucey & Riner, 2011; Noble et al., 2007). Surface-correlated amorphization may be a more parsimonious explanation, however, since the host grain for samples 1–3 does not contain an appreciable amount of iron. Suppose that the observed darkening results from the amorphous surface layers produced by long-term exposure to the space environment. In that case, it is plausible that the hypervelocity impact (sample 1) vaporized or melted this layer; this could explain why no darkening was observed for sample 1 despite originating from the same soil grain as samples 2 and 3. This scenario is consistent with the widespread evidence of impact-induced shock and vitrification seen near the crater in sample 1 (see Figure 7). Similarly, we did not observe a robust correlation between total scattered intensity and depth for sample 4, suggesting that it contains limited concentrations of npFe⁰ or is otherwise more robust to the space environment over the studied wavelength range.
However, it is important to note that samples 2 and 3 were extracted parallel to one another and nearly perpendicular to sample 1. As such, the effect of crystallographic orientation cannot be ruled out when comparing results among the samples. The primary observations—namely, that there is depth-dependent spectral variation between the grain interior and the space-exposed surface—are not affected by this limitation. Where present, the darkening effect is most apparent within 500 nm of the grain surface (see Figure 12). This depth falls just outside of the observed range of thicknesses for amorphous rims in lunar soil (~10–350 nm; Burgess & Stroud, 2018; Christoffersen et al., 1996), but well within the range of thicknesses for glassy silicate layers (10–1,000 nm) thought to have been produced by micrometeoroid impacts (Noble et al., 2005). For comparison, the average implantation depth of solar wind-produced H and He has been estimated as ~20–100 nm (Christoffersen et al., 1996; Farrell et al., 2015; Tucker et al., 2019).

That we observed space weathering effects at depths greater than the penetration range of typical solar wind protons could suggest that the implanted hydrogen diffused into the grain. Although some diffusion undoubtedly occurs, it is unlikely to be the dominant cause of the observed effects given the relatively poor H-retention of lunar soil (Farrell et al., 2015). Alternatively, the effects observed could result from the occasional bombardment of the lunar surface by solar energetic particles (SEPs), which are substantially more energetic than solar wind ions. Hydrogen SEPs have kinetic energies of 2–10 MeV (Mewaldt et al., 2009), whereas typical solar wind H⁺ ions have kinetic energies of ~1 keV (Gosling et al., 1976). Although SEPs are likely to implant further into lunar soil than average solar wind ions, they occur far less frequently. Without
additional support from independent lines of evidence (e.g., observing tracks via TEM), these confounding factors preclude definitive conclusions about the role of SEPs in the weathering of our samples.

5. Conclusions

We used SINS to examine surface-correlated, mid-IR space weathering effects in lunar soil grains. In general, our results are consistent with the spectral changes previously hypothesized to be correlated with the microstructural and compositional changes measured by TEM. Crucially, however, our results demonstrate that SINS (and related techniques) can be used to investigate the spatial scales over which ion irradiation and micrometeoroid bombardment affect the soil’s optical properties. With a spatial resolution comparable to the scale of space-weathering induced microstructural and chemical changes, SINS can be used to establish a direct link to bulk space weathering effects. As such, we have shown that this technique fills the gap between TEM microstructural studies and far-field FTIR measurements.

The data presented above provide clear evidence supporting previous findings that space weathering effects result from highly localized features (on the order of tens of nanometers). We found that the effect size varies continuously (at the sampled spatial resolution) over a micrometer-scale range of depths. Our results additionally indicate that soil maturity indices should be used with caution when discussing micron-scale sub-samples of lunar soil. While the soil maturity index is a reliable predictor of large-scale weathering effects, our results reinforce the variability of exposure history among a soil’s constituent grains.

The techniques utilized in this study have been shown to produce results similar to those observed in bulk lunar soils. Taking advantage of techniques with spatial resolutions on the order of tens of nanometers, such as afforded by SINS, may prove useful for studying the relative contribution of each small-scale process (e.g., solar wind implantation, nano-phase iron production, and micrometeoroid impacts) to the overarching space weathering phenomenon. SINS data may also inform and refine the techniques used to simulate weathering phenomena in the laboratory. With a more detailed understanding of the spectral effects of charged-particle irradiation on mineral and soil grains, it may also be possible to draw parallels to, inter alia, silicate processing in the interstellar medium (Chiar & Tielens, 2006). Detailed studies of the association between specific molecular vibrational modes and the features present in SINS spectra of minerals may shed further light on various space weathering mechanisms. Information regarding the molecular bonds affected by space weathering, paired with precise chronometry and compositional measurements of weathered lunar soils, may help to constrain or validate current models of space weathering processes.

Data Availability Statement

The data used for this research is available at the Digital Research Material Repository at Washington University in St. Louis (Utt et al., 2020).

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