Space weathering of silicates simulated by successive laser irradiation: In situ reflectance measurements of Fo\textsubscript{90}, Fo\textsubscript{99+}, and SiO\textsubscript{2}

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(Received 16 June 2015; revision accepted 19 October 2015)

Abstract—Pulsed-laser irradiation causes the visible-near-infrared spectral slope of olivine (Fo\textsubscript{90} and Fo\textsubscript{99+}) and SiO\textsubscript{2} to increase (redden), while the olivine samples darken and the SiO\textsubscript{2} samples brighten slightly. XPS analysis shows that irradiation of Fo\textsubscript{90} produces metallic Fe. Analytical SEM and TEM measurements confirm that reddening in the Fo\textsubscript{90} olivine samples correlates with the production of “nanophase” metallic Fe (npFe\textsuperscript{0}) grains, 20–50 nm in size. The reddening observed in the SiO\textsubscript{2} sample is consistent with the formation of SiO or other SiO\textsubscript{x} species that absorb in the visible. The weak spectral brightening induced by laser irradiation of SiO\textsubscript{2} is consistent with a change in surface topography of the sample. The darkening observed in the olivine samples is likely caused by the formation of larger npFe\textsuperscript{0} particles, such as the 100–400 nm diameter npFe\textsuperscript{0} identified during our TEM analysis of Fo\textsubscript{90} samples. The Fo\textsubscript{90} reflectance spectra are qualitatively similar to those in previous experiments suggesting that in all cases formation of npFe\textsuperscript{0} is causing the spectral alteration. Finally, we find that the accumulation of successive laser pulses cause continued sample darkening in the Vis-NIR, which suggests that repeated surface impacts are an efficient way to darken airless body surfaces.

INTRODUCTION

Airless planetary bodies are constantly being bombarded with multiple types of radiation and hypervelocity meteorite impacts, such that their surface regoliths undergo a diversity of physical, chemical, and optical changes known as space weathering (Hapke 2001; Clark et al. 2002; Chapman 2004). Identifying the “markers” that indicate a space weathered regolith is important not only for understanding how the surface evolves over time but also for accurately determining the composition of the intrinsic surface by remote sensing techniques. One of the most commonly utilized techniques for characterizing surfaces of airless bodies is optical reflectance spectroscopy. Ideally the remotely sensed reflectance spectrum of a surface can be compared with a relevant laboratory spectrum of extraterrestrial or analog material to yield information on the surface’s composition, mineralogy, and origin.

For the case of planetary surfaces with an abundance of mafic minerals, such as S-type asteroids and the Moon, there are key indicators for surface alteration via radiation or meteorite impact. Compared to unweathered minerals of assumed similar composition, the near-infrared (\textasciitilde0.6–2.5 \mu m) reflectance spectra is usually more sloped (redder), lower in reflectance (darker), and contains weaker absorption features (Gaffey 2010).

Laboratory studies have been generally successful in reproducing the space weathering characteristics seen in lunar and S-type asteroid spectra by focusing on the changes caused by energetic ions and micrometeorite hypervelocity impacts. Using keV protons as a solar wind analog, early studies (Nash 1967; Hapke 1973) showed that irradiation caused the reflectance spectrum of the analog and returned lunar samples to evolve toward the in situ lunar soil spectra, becoming darker, redder, and having attenuated absorptions. Through more detailed studies (see Hapke [2001] for a review),
Hapke was able to show that to create spectral changes consistent with space weathering, the sample needed to contain Fe. The effect was enhanced when using a fine-grained, powder sample (Pieters et al. 1993; Hapke 2001). The chemical reduction of Fe found in both ion irradiated and natural lunar soils (Yin et al. 1975; Housley and Grant 1977) seemed to support the assertion that solar wind chemical reduction of Fe could be causing spectral results of weathering. Subsequent analytical transmission electron microscopy (TEM) observations supported the hypothesis that the Fe was chemically reduced and found that most of it was in the form of 5–100 nm diameter metallic Fe spherules residing in altered rims on lunar regolith grains, as well as in the glassy portions of glass-welded aggregates known as agglutinates (Keller and McKay 1993, 1994, 1997). For the solar wind specifically, laboratory studies have since shown that direct keV ion irradiation can chemically reduce Fe on the surfaces of Fe-bearing minerals such as olivine (Dukes et al. 1999; Davoisine et al. 2008; Loeffler et al. 2009), and that this reduction correlates with the observed spectral reddening (Loeffler et al. 2009). In addition, Noble et al. (2007) showed that simply adding fine-grained (<10 nm) metallic Fe to an nonabsorbing silicate gel caused reddening of the sample spectrum.

Early simulations of meteorite impacts focused on vitrification of regolith grains, because glass formation appeared to alter the reflectance spectra of various types of samples (Nash and Conel 1973; Hapke et al. 1975). However, there is still some debate on why darkening and attenuation of absorption bands were found in some experiments (Adams and McCord 1971; Nash and Conel 1973) and not in others (Hapke et al. 1975); see Hapke (2001) for more discussion. Other experiments that investigated the synergistic effects of heating and comminution during impacts found little change in reflectance with comminution alone, but qualitative reddening and darkening with repeated comminution-melting cycles (Clark and Fanale 1992). However, these results have also been questioned (Hapke 2001).

More recent attempts to simulate the conditions in micrometeorite impacts have used pulsed-laser beams with instantaneous power densities (W cm$^{-2}$) similar to those of typical impacts by micron-sized meteorites (Moroz et al. 1996; Sasaki et al. 2001; Brunetto et al. 2006a; Loeffler et al. 2008; Noble et al. 2011). These studies have shown laser irradiation, like ion irradiation, can cause the near-infrared reflectance spectra of mafic minerals to redden, and may cause significant darkening as well. These spectral changes are consistent with those in weathered lunar samples, which are known to contain npFe$^0$. However, although the SEM results of Moroz et al. (1996) showed that all laser-irradiated samples contained glassy deposits and that the olivine samples contained Fe-rich regions, the spatial resolution of their microprobe analysis was not sufficient for them to determine whether any of these regions contained reduced “nanophase” Fe (npFe$^0$). Later, Sasaki et al. (2001, 2003) used transmission electron microscopy and electron spin resonance (ESR) to show that the Fe-rich regions formed in laser-irradiated olivine and pyroxene samples contained reduced npFe$^0$, supporting the hypothesis that the observed optical changes are driven by the presence of reduced npFe$^0$ particles. Most recently, Brunetto et al. (2006a) suggested that, besides formation of npFe$^0$, structural modification induced by the pulsed laser may be important, as the rapid melting of the sample may not only change the phase of the silicates but could also induce irregular structures that may change the scattering properties of the minerals.

These previous laser irradiation studies have clearly shown that the laser can generate both melt and vaporized material on mineral and rock surfaces creating a variety of structures which, given the presence of Fe in the target material, can contain npFe$^0$ particles. However, there have been no experiments explicitly showing that the observed spectral changes correlate directly with the formation of metallic Fe, because none of the previous studies employed a microanalytical technique that could monitor metallic Fe content during laser irradiation. In addition, although previous investigations studied different mafic minerals, none varied the Fe content within the same mineralogical family, a strategy which could clarify the role structural modification plays in the alteration of reflectance spectra. In this work, we have investigated the role that Fe content plays when a key silicate, olivine, is irradiated with a pulsed laser operated to simulate conditions during micrometeorite impact. Besides being directly applicable to previous laser irradiation studies mentioned above, olivine is a major component in chondritic meteorites and has been identified in numerous asteroids (Sanchez et al. 2014), and is also a significant lunar mineral (Gaffey et al. 1993; Hapke 2001; Clark et al. 2002). This article is a companion work to our previous studies on the spectral and chemical effects of vapor redeposition and ion irradiation (Loeffler et al. 2008, 2009). The unique aspect of our approach has been that, in addition to measuring the reflectance spectra of our sample, we can also monitor the chemical evolution of our sample using X-ray photoelectron spectroscopy (XPS). This has allowed us to show the correlation between spectral reddening and chemical reduction of Fe in ion irradiated Fo$_{90}$ San Carlos olivine (Loeffler et al. 2009).
In the current experiments, we used a 193 nm ns-pulsed laser to irradiate, under ultrahigh vacuum, flat-sectioned surfaces of single-crystals and powders of Fo$_{90}$ San Carlos olivine, as well as powders of nearly pure synthetic forsterite and SiO$_2$. During pulsed-laser irradiation, we monitored both the near-infrared reflectance spectra and the surface chemical composition of our samples in situ, e.g., without exposing the samples to the atmosphere. In addition to the in situ measurements, we also performed ex situ measurements using ultraviolet-visible reflectance spectroscopy, and scanning and transmission electron microscopies on select samples to measure changes in surface structure, including Fe nano-particle production.

**EXPERIMENTAL METHODS**

**Sample Preparation and In Situ Analysis**

The majority of the experiments reported here were on powders sieved to a 45–125 μm size fraction and pressed into an aluminum ring to make a ~10 mm diameter by ~2 mm thick pellet disk. The slightly larger grain size in our pellets, compared to our previous work (<45 μm; Loeffler et al. 2008, 2009), was needed to prevent the sample from disintegrating during laser irradiation. The pellets were prepared from natural San Carlos olivine (Fo$_{90}$; Mg$_{1.8}$Fe$_{0.2}$SiO$_4$; Penn Minerals), synthetic SiO$_2$ (99.99%; Alfa Aesar) and synthetic forsterite (Fo$_{99+}$; Reade Advanced Materials). The synthetic forsterite is listed as containing 0.02 wt% Fe$_2$O$_3$ as an impurity (≥Fo$_{99.9}$), along with Al$_2$O$_3$ (0.03), TiO$_2$ (0.09), Na$_2$O (0.11), CaO (0.22), and K$_2$O (0.12). In addition to the pressed pellets, we also performed experiments on flat unpolished sections of San Carlos olivine single-crystals several mm in size (Arizona Gems and Crystals).

After preparation, the samples were mounted on a copper holder and transferred into an oil-free ion-pumped ultrahigh vacuum (UHV) chamber with a base pressure of 5–10 $\times$ 10$^{-10}$ Torr (see Loeffler et al. 2009). Surface analyses of the samples were performed before, during, and after irradiation using X-ray photoelectron spectroscopy (XPS) and near-infrared reflectance (NIR) spectroscopy. More details on sample preparation and our in situ analysis methods are given in Loeffler et al. (2009).

**Laser Irradiation**

A typical approach to simulating micrometeorite impacts employs high-energy pulses from a nanosecond laser (Yamada et al. 1999; Sasaki et al. 2001; Brunetto et al. 2006a). This technique has the advantage, compared to the more traditional approach of accelerating dust particles from a dust accelerator (Cintala 1992), of uniform irradiation over macroscopic areas with a relatively high-energy flux. Analysis of laser-irradiated samples shows that this method successfully produces spectral changes consistent with micrometeorite impact (Sasaki et al. 2001). Besides the experimental evidence, the justification for using ns-pulsed lasers to mimic dust impacts has also been made by comparing the composition of thermoionic plasma emitted from samples that were impacted by accelerated dust and laser beams (Kissel and Krueger 1987), normalized to the deposited power density (W cm$^{-2}$). Kissel and Krueger (1987) showed that, above the laser ablation threshold, similar ejected ion mass spectra and absolute yields are obtained when the power density of the laser beam is ~100 times lower than that for dust impact, as a result of the different efficiency of conversion of impact energy to temperature (Sugita et al. 2003). With this in mind, we estimate that the UV laser pulses used in our experiments (~10$^8$ W cm$^{-2}$) are roughly equivalent to a 1–20 μm particle (2 g cm$^{-2}$) traveling ~10–15 km s$^{-1}$. On impact, this is equivalent to a power density of 10$^{10}$ W cm$^{-2}$ assuming that all the energy is deposited over an area given by the cross section of the dust particle, and a depth of 3 particle diameters. We note that even micrometeorite impacts as slow as 0.05 km s$^{-1}$ can produce a thermionic plasma and, hence, vaporization (Smith and Adams 1973).

In the experiments reported here, we irradiated our samples at normal incidence with a GAM ArF excimer laser that emits 10 ns pulses of 193 nm light at 1 Hz. During irradiation, the normal of the sample surface was horizontal. The laser beam entered the analysis chamber through a quartz-fused silica window and was imaged to a 0.5 mm$^2$ spot on the sample surface; each pulse had an energy fluence of 1 J cm$^{-2}$. The beam was rastered uniformly across the sample with mirrors actuated by a motion control system synchronized to the laser. During each raster sequence, the sample was monitored to ensure that only the pellet material (not the Al ring) was irradiated. Finally, we note that in the text when we describe one laser pulse, we mean that the laser has been rastered across the sample one time, so that each region of the analyzed area has been irradiated once.

**Ex Situ Analysis**

After irradiation and in situ analysis, selected samples were subsequently analyzed by additional methods after removal from the UHV irradiation chamber. These additional ex situ analyses included: ultraviolet-visible spectroscopy, analytical field-emission scanning electron microscopy (FE-SEM) and analytical field-emission scanning transmission electron...
microscopy (FE-STEM). For the ultraviolet-visible measurements, we used a FilmTek fiber optic spectrometer over the wavelength range of 250–600 nm, at a resolution of 0.2 nm. Light was incident normal to the sample’s surface and reflected light was collected at 45 degrees. The reflectance spectrum was obtained by dividing the collected light from the sample by the reference, after subtraction of the dark spectrum, to yield the reflectance value: 
\[ R = \frac{I_{\text{sample}} - I_{\text{Dark}}}{I_{\text{reference}} - I_{\text{Dark}}} \]

The reference sample was a Spectralon™ reflectance standard obtained from Labsphere. However, as the absolute reflectance values are not provided at the lower wavelengths, we do not consider our processed reflectance spectrum to be an absolute one. Given the minor difference between the phase angle in the UV and near-infrared reflectance measurements, lack of absolute standard in the UV etc., we are not surprised by the slight reflectance variation in the data we can see in SiO2 and to a lesser extent, in the other two samples (compare Figs. 1, 2, and 5).

Following ex situ optical spectroscopy, field-emission scanning electron microscopy characterization was performed on selected samples utilizing a JEOL 7600F field-emission SEM, which can acquire secondary/backscattered electron images (SEI/BSE) and compositional element maps with a Thermo® energy-dispersive X-ray (EDX) spectrometer system. Regions of particular microstructural or compositional interest were then prepared for field-emission scanning transmission electron microscopy characterization by focused ion beam (FIB) lift-out sectioning using an FEI Quanta dual electron/ion beam milling instrument. FE-STEM characterization utilized a JEOL 2500SE FE-STEM with compositional element mapping capabilities provided by a Thermo® EDX compositional spectrum imaging system. X-ray counts for individual EDX spectra and element maps were converted to wt% element concentrations using the Cliff-Lorimer quantification technique, which is based on empirically determined k-factors (Cliff and Lorimer 1975).

RESULTS

Fluence Dependence of Near-Infrared Reflectance

Figure 1 shows the near-infrared reflectance (unpolarized) of the Fo90-pressed pellet sample, measured in situ as a function of the accumulated number of individual 1 J cm⁻² rastered laser scans. The Fo90 absolute reflectance spectrum (Fig. 1 top) shows the absorption features characteristic of polycrystalline olivine, due to electronic excitation of the outer 3d-electrons of Fe²⁺ (Burns 1993). The spectrum of the unirradiated sample is similar to spectra reported in previous studies on San Carlos olivine (Yamada et al. 1999; Sasaki et al. 2001; Brunetto et al. 2006a). The data in Fig. 1 show clear evidence that consecutive laser irradiations cause systematic changes in the Fo90 pellet sample: the spectral slope increases, the overall reflectance decreases, and 1 µm absorption band attenuates. To highlight the changes in the sample solely due to irradiation, we show the ratio of the spectra of the irradiated olivine sample to that of the unirradiated olivine sample (Fig. 1 bottom). Similar changes in spectral shape have been reported previously for various types of laser-processed olivine samples, including pulsed-laser deposits formed on powdered olivine substrates using olivine irradiation targets (Loeffler et al. 2008), direct laser ablation of olivine (Brunetto et al. 2006a), olivine impacted by high-energy (60–400 keV) ions (Strazzulla et al. 2005; Brunetto et al. 2006b), and low energy (4 keV) ions (Loeffler et al. 2009). As in our previous ion irradiation experiments (Loeffler et al. 2009), we find that there is no clear shift in the band center during irradiation.

To determine if pulsed-laser irradiation can alter the reflectance and composition of a non-Fe-bearing
mineral, we irradiated pressed pellets of both Fo99+ and SiO2 powders. Figure 2 (top) shows the absolute reflectance spectrum of SiO2 powder. The spectral changes induced by the laser are much less than what was observed in the Fo90, yet the overall trend is clearly greater than our estimated experimental error (<1%). Figure 2 (middle) shows the absolute reflectance spectra of the Fo99+ powder during irradiation, while Fig. 2 (bottom) shows the ratio of the spectrum of the irradiated sample to that of the unirradiated sample; the spectra (from top to bottom at 700 nm) correspond to 7, 25, and 49 laser pulses.

Although the synthetic forsterite has less than a tenth of the Fe as compared to our San Carlos sample, its spectral slope increases and, after the first laser pulse, overall reflectance decreases with increasing irradiation. These results are different from what we observed when we vapor deposited Fo99+ onto silicate powders, where only a low amount of reddening was observed (Loeffler et al. 2008).

To quantify the spectral changes observed in our three types of samples, we compare the changes in the spectral slope (reddening) in Fig. 3 (top). As previously (Loeffler et al. 2009), we define reddening as $r = (R_{2.0 \mu m} - R_{0.7 \mu m})/1.3 \mu m$, where 1.3 $\mu m$ is the wavelength interval. For the Fo90 pellet sample, only one laser pulse is needed to bring it halfway to its most reddened value, achieved after 49 pulses. Fo99+ and SiO2 redden in a more gradual fashion with laser fluence, and do not reach steady state within the time frame of our experiment. We note that although the laser causes the Fo99+ sample to redden less than the Fo90 sample initially, the Fo99+ sample continues to redden with laser irradiation and after 49 pulses the two samples are nearly equally reddened. The SiO2 is actually slightly “blue” before irradiation, but reddens moderately as it is irradiated. By the time the fluence reaches 99 pulses, the reddening is still more than a factor of three less than what we observed for the other samples and does not appear to have reached a steady state.

In addition to reddening, data for sample darkening are shown in Fig. 3, with darkening parameterized as $1 - R/R_0$ at 2 $\mu m$, where $R_0$ is reflectance of the unirradiated sample. The Fo90 pellet sample darkens...
during irradiation and appears to reach a steady-state value of \( \sim 15\% \) after 99 laser pulses. The \( \text{Fo}_{99}+ \) and \( \text{SiO}_2 \) samples show a different behavior. After only one pulse, both samples appear to brighten slightly. The \( \text{SiO}_2 \) continues to slowly brighten throughout the irradiation up to a value of \( \sim 6\% \). However, after the initial brightening, additional irradiation does not change the \( \text{Fo}_{99}+ \) sample’s reflectance until after 13 pulses, where the sample darkens linearly up to the last set of pulses (49 pulses) to a value of 5%.

For the \( \text{Fo}_{90} \) sample, we quantified the area of the 1 \( \mu \)m absorption band by converting the spectrum to optical depth units, taking the natural logarithm of the reflectance, and subtracting a nonlinear baseline (Fig. 3 bottom). This method is the same as that used in our previous ion irradiation study on \( \text{Fo}_{90} \) (Loeffler et al. 2009). Figure 3 shows that, as the sample is irradiated, the area of the 1 \( \mu \)m absorption band decreases by 20% after 99 laser pulses. The slight decrease in the absorption band depth also is evident in the olivine ratio spectrum (Fig. 1 bottom, 99 pulse spectrum).

**X-Ray Photoelectron Spectroscopy**

During laser irradiation, changes in the surface chemical composition of the \( \text{Fo}_{90} \) and \( \text{Fo}_{99}+ \) samples were monitored with XPS. We acquired initial survey spectra and also high-resolution spectra for chemical analysis of the main elements (O, C, Fe, Mg, and Si). The spectra were compared as follows. First, the spectra were all shifted to compensate for residual electrostatic charging, so that the peak position of Si-2p oxide was aligned; this shift was <0.5 eV. Next, the absolute binding energy scale was determined from the position of the C-1s peak for adventitious carbon in the initial spectra at 284.8 eV. The binding energies measurements for our samples are in good agreement with previously published values of olivine (Dukes et al. 1999), forsterite (Hochella and Brown 1988), fayalite (Schott and Berner 1983), and \( \text{Mg}_2\text{Fe}_{0.8}\text{SiO}_4 \) (Yin et al. 1971). The \( \text{Fo}_{99}+ \) sample was found to contain detectable Fe, but the amount was so low that the chemical changes could not be easily followed.

For the \( \text{Fo}_{90} \) and \( \text{Fo}_{99}+ \) samples, we did not observe significant changes in the O-1s (531.4 eV), Mg-2p (50.1 eV), or Si-2p (101.8 eV) photoelectron peaks resulting from laser irradiation. In particular, no formation of metallic Mg or elemental Si was detected, although the Mg-2p and Si-2p peaks did broaden slightly suggesting that some chemical reduction occurred. As the pellet samples were not prepared under vacuum conditions, they all showed initially some atmospheric contamination, evidenced by the presence of the C-1s adventitious carbon peak at 284.8 eV. After 1–2 laser pulses, the surface carbon signal decreased to the noise level of the instrument. In addition to the adventitious carbon peak, there appeared to be a second, minor, C-1s peak present at 283.8 eV that remained throughout the experiment indicating carbon intrinsic to the samples.

Besides the removal of carbon surface contaminants at low fluences, the only other major change observed was chemical reduction, apparent in the Fe-2p transition of the \( \text{Fo}_{90} \) sample (Fig. 4). The initial peak position of the Fe-2p band indicates that Fe is not in the form of Fe\(^{2+}\) (709.1 eV) expected for the bulk olivine sample, but rather as Fe\(^{3+}\) (711.4 eV) on the surface. These results are in agreement with previous experiments (Schott and Berner 1983; Dukes et al. 1999). Continued irradiation of the \( \text{Fo}_{90} \) olivine powder increasingly reduces the Fe on the surface to the metallic state (Fe\(^0\), 706.7 eV), attaining a steady-state level of near 0.7 Fe\(^0\)/Fe represented by the 49 pulse spectrum. This value is similar to the steady-state value observed as a result of ion irradiation (Loeffler et al. 2009), but slightly lower than that of a laser-irradiated flat olivine section. We will return to this topic in the discussion section.

After laser irradiation and in situ analysis, we exposed the laser-irradiated \( \text{Fo}_{90} \) powder sample to atmosphere for \( \sim 10 \) min, and then reacquired the XPS and NIR spectra. The Fe on the surface of the olivine is nearly completely re-oxidized (top spectrum in Fig. 4), as determined by XPS, while the reflectance spectra is virtually unchanged (data not shown here), consistent
with our previous work (Loeffler et al. 2009). This implies that the reoxidized layer, while thicker than the XPS sampling depth (~5–8 nm for our sample) (Seah and Dench 1979) is much thinner than the layer created by the laser.

Ex Situ UV Reflectance

We also measured the ultraviolet-visible reflectance of some of our samples before and after irradiation (Fig. 5). The left panel shows the sample reflectance before irradiation; it is clear that there is significant UV absorption in the reflectance spectrum of the Fo90 and Fo99+ samples, while the spectrum of SiO2 is spectrally flat within error. The SiO2 spectrum is consistent with previous studies (Philipp 1966). The absorption observed for the Fo90 sample is also expected, as it is derived from the relatively strong Fe oxide charge transfer band at 260 nm, an allowed transition for Fe2+-O2-/C0 in olivine (Hapke 2001). The absorption present in the Fo99+ spectrum is presumably due to the Fe impurities present in the sample, which can exist as Fe2+ or Fe3+, as the closest reported transition in the region for a completely iron-free sample is due to the band gap transition near ~150 nm (Nitsan and Shankland 1976). We note that previous studies have shown that even very low Fe/Mg ratios in synthetic forsterite can show observable absorption bands as both Fe cations absorb in this region (Shankland 1968; Nitsan and Shankland 1976; Balberg and Pinch 1978).

After irradiation with 99 laser pulses, all samples showed a decrease in reflectance in the ultraviolet. Surprisingly, the SiO2, with no known absorption bands above ~120 nm and no metallic Fe present, shows a definite decrease in UV reflectance, which extends up to ~500 nm. The shape of the reflectance curves for the Fo90 and Fo99+ samples look fairly similar after irradiation. Both spectra lose their curvature below 400 nm and become linear, dropping in reflectance with decreasing wavelength. Examination of the slope for these two samples shows that between 300 and 400 nm the slope decreases by a factor of two for Fo99+ and a factor of 3.5 for Fo90. This decrease in slope with irradiation is consistent with the observation by Hendrix and Vilas (2006), who found that the spectral slope in this region is also lower for weathered asteroids and lunar soil.

Analytical Field-Emission SEM and STEM

After 99 laser pulses, the Fo90 pressed-pellet powder sample was analyzed by FE-SEM and FE-STEM.
methods. Figure 6 shows a representative region of the sample surface imaged by FE-SEM backscattered electron (BSE) imaging. Figure 6a shows areas (indicated by arrows) with locally high concentrations of submicrometer, spherule-shaped grains that were confirmed to be metallic Fe (npFe$^0$) based on subsequent FE-STEM characterization (see Fig. 7). The npFe$^0$ is hosted in laser-altered material concentrated within narrow gaps between grains and on the sloping sides of grains that border larger surface depressions. Averaged over the area of Fig. 6a, the npFe$^0$ modal percentage is approximately 0.5% based on the corresponding threshold-processed image in Fig. 6b. The average grain size of npFe$^0$ visible at the resolution of most BSE images is approximately 100 nm, but a broad size distribution is typical as shown by the higher magnification BSE image in Fig. 6c.

A FIB cross section for FE-STEM characterization was prepared from the area shown by the white line in Fig. 6a. The cross section encompasses a continuous layer of laser-altered material covering a single olivine grain that borders a surface depression (see Fig. 6a). FE-STEM bright-field (BF) imaging shows that the altered layer is similar in thickness to what has been observed in laser-irradiated plagioclase grains (Moroz et al. 2014). Specifically the altered layer is thickest (0.8–1.0 µm) on the side of the grain that slopes down into the depression, then progressively tapers to a thickness of <0.5 µm over the top or “crest” of the grain away from depression. The images also show that in the thickest part of the altered region, there are three microstructurally distinct layers. EDX shows that each of these layers are composed of silicate glass with varying modal fractions of npFe$^0$ spherules and/or nanocrystalline silicate material (Figs. 7a and 7b). In addition, clusters of larger npFe$^0$ spherules 100–300 nm diameter occur locally within the altered layer, consistent with our SEM observations (Fig. 6). The first sublayer, which forms the immediate surface of the altered grain, is 50–300 nm wide and composed of approximately 40–50 modal% npFe$^0$ spherules 5–20 nm in size (Figs. 7a and 7b).

Underneath sublayer 1 is a glass layer (sublayer 2; Fig. 7), which has a much lower modal concentration of npFe$^0$ (<0.5%) relative to sublayer 1 and contains scattered nanoscale regions with short range crystalline order identifiable in high-resolution TEM images. The third sublayer in the sequence, at the bottom of the altered region, is between 200 and 400 nm thick and is composed predominantly of more irregularly shaped nanocrystalline grains that are 10–50 nm in size (Fig. 7a). In portions of this layer, the grains are intergrown along shared grain boundaries with little or no surrounding glass, while in other areas they are isolated within a glassy matrix. The d-spacings of both individual reflections and ring patterns in selected-area electron diffraction patterns of the aggregates suggest most of these grains are olivine, with a subordinate amount of npFe$^0$.

In quantified EDX element maps, the elemental wt% of Fe in the topmost sublayer (sublayer 1) was enhanced by a factor of 2.5 higher relative to the wt% Fe in the underlying olivine grain, while Mg and Si were depleted by 20% and 10% respectively. Sublayers 2 and 3 both show trends in Fe and Mg that are opposite to sublayer 1: Fe is depleted by 80–90% and Mg is enriched by 20–30% relative to the underlying
olivine. Si is, however, depleted in sublayers 2 and 3 in much the same manner as in sublayer 1.

DISCUSSION

Brief Overview of Laser Ablation of Insulators

It is well known that the surfaces of solid materials can be heavily modified by high-power pulsed-laser irradiation. The process by which surface material is removed (ablated) is highly dependent on the laser fluence (J cm\(^{-2}\)): at low fluences the material can leave through simple evaporation from the surface, while at higher fluences a plasma can be created in the ejected plume. Predicting the degree to which a laser will ablate any material is complicated, as the interaction will depend on a number of factors, such as the laser fluence, pulse width, wavelength, and absorption coefficient of the sample. For wide band-gap materials, such as the insulators studied here, although the absorptivity of the sample is low at the 193 nm wavelength used, the sample can still absorb light. This occurs through multiphoton absorption (Tsai et al. 1988; Shen et al. 1989), via defects initially present at the surface or subsequently created during irradiation (Ihlemann et al. 1992; Pedraza 1998). In multiphoton absorption, the electrons are excited into the conduction band of the solid by absorbing more than one photon, whereby they can absorb light at any wavelength, subsequently transferring energy in the solid. Defects are present in all materials to some degree and break the periodicity of the lattice structure, creating absorption states in the band gap that can absorb light. Regardless of how the light is absorbed, the damage can produce more defects and changes in surface structure that can further enhance a sample's efficiency to absorb light. Furthermore, the absorption of high intensity light and ablation efficiency of transparent material has also been shown to increase with the surface roughness (Ihlemann et al. 1992), potentially as a result of avalanche ionization (Bloembergen 1974), where the increased electric fields on the rough surfaces produce dielectric breakdown, creating a plasma that renders the material opaque. Thus, although the materials studied here have band gaps >8 eV (Nitsan and Shankland 1976), larger than the laser photon energy in our experiments, we expected that both olivine and SiO\(_2\) would be ablated at some level by our laser.

Factors in Comparing Laser Irradiation Effects in Different Minerals

For the current set of measurements, as in previous space weathering experiments (Loeffler et al. 2008, 2009), we studied samples that were similar in composition, Fo\(_{90}\) and Fo\(_{99}\), to investigate how irradiation effects varied in samples that had different intrinsic Fe contents. However, comparing sample-to-sample effects in the ion irradiation experiments on the basis of fluence is subject to fewer uncertainties than these laser experiments, where the amount of light absorbed during each pulse and how it is absorbed may vary with mineral composition, as the material optical properties change.

Reddening and Darkening in SiO\(_2\)

Of the three samples, the spectral changes resulting from laser irradiation of the SiO\(_2\) sample are the least prominent (Fig. 3). After 49 pulses, the reddening is ~10
times less for SiO$_2$ than olivine samples and the change in reflectance at 2 $\mu$m is more than five times lower and in the other direction (the sample brightens).

Spectral reddening in the SiO$_2$ sample is not evident until after the seventh laser pulse (Fig. 3), when the spectral slope begins to increase slowly with fluence. This threshold is similar to what has been reported for ablation studies on crystalline and amorphous SiO$_2$ (Ihlemann 1992; Pedraza 1998), which show similar fluence thresholds for the onset of ablation. This behavior has been attributed to the buildup of defects in the sample, which increase the sample’s absorptivity; the absorption for SiO$_2$ has been modeled to be extremely low at 193 nm (Awazu 2004). Once the threshold has been reached, the sample ablates at a nearly linear rate (Ihlemann et al. 1992; Pedraza 1998). We note that the threshold we observe is slightly lower than in previous studies. However, we speculate this can still explain why little to no spectral change is observed for a low number of laser pulses, as the threshold appears to be highly dependent on a number of factors, such as sample surface roughness (Ihlemann et al. 1992), the phase of the SiO$_2$ (Pedraza 1998), and laser wavelength (Awazu 2004). Once the ablation threshold is reached in our samples the spectral slope increases linearly with increased irradiation until the end of the experiment. The reddening observed can be explained by the formation of SiO$_x$, where x < 2 (Philipp 1971), which absorb strongly in the UV. We note that the formation of SiO and SiO$_{1.5}$ in laser-irradiated samples of SiO$_2$ has been reported previously, using XPS (Awazu 2004) and is consistent with the broadening of the Si-2p peak in our Fo$_{90}$ and Fo$_{99+}$ samples that were measured with XPS.

In addition to reddening, the SiO$_2$ brightens at 2 $\mu$m during irradiation. Half of the brightening (~3%) occurs after the first laser pulse, which is likely due to the removal of the adventitious carbon layer on the surface grains or simply a minor change in the macroscopic structure of the pressed pellet, i.e., some of the weakly bound grains may have fallen off of the sample. Similar to the case of reddening, no spectral changes occur until after seven pulses, whereby there is a slow increase in the reflectance at 2 $\mu$m. By the end of the experiment, the reflectance at 2 $\mu$m is about 3% higher than it was after seven laser pulses. We suspect that this small change is due to changes in sample topography, such as surface melting/smoothing of the surface. We note that previous experiments showed that ablating Fo$_{99+}$ onto unirradiated Fo$_{99+}$ powders caused significant topography changes (see fig. 9 in Loeffler et al. 2008) and brightened the powders by ~4%. A similar conclusion was made by Moroz et al. (2014), who observed brightening in plagioclase pellets that had been laser irradiated. We point out that brightening has also been reported for dark minerals that have been ion irradiated, although the process may be different (Vernazza et al. 2013), as the initial reflectance is much lower.

**Reddening and Darkening in Fo$_{90}$ and Fo$_{99+}$**

Previous laboratory measurements have shown that the absorption of Fo$_{90}$ and Fo$_{99+}$ at 193 nm are comparable (Nitsan and Shankland 1976), and thus in the following discussion we consider that the absorbed laser fluence for these two olivine samples is similar. This is consistent with our earlier studies that showed the ablation rate for these two minerals was similar (Loeffler et al. 2008).

Spectral reddening and darkening occur during laser irradiation for both Fo$_{90}$ and Fo$_{99+}$ pellet samples. For the Fo$_{90}$ sample, the reddening occurs quickly and irradiation beyond ~13 pulses does not additionally increase the spectral slope of the olivine, yet the fraction of metallic Fe continues to increase (Fig. 8). The lack of correlation between the metallic fraction and spectral reddening is different from our previous ion irradiation experiments, plotted for comparison in Fig. 8, which showed that these two parameters were correlated (Loeffler et al. 2009). We suspect that the difference observed here is that, while repeated laser pulses increase the Fe$^0$/Fe on the surface, they also heat the sample, allowing the metallic Fe grains to undergo microstructural coarsening (e.g., Ostwald ripening), increasing the size of the iron nanoparticles. The larger particles do not contribute to the reddening but are more effective at darkening the sample (Noble et al. 2007). This hypothesis is also supported by our FE-SEM and FE-STEM observations, which show a small number of larger (100–200 nm) npFe$^0$ grains that are likely growing at the expense of the smaller (~50 nm) grains.

During laser irradiation, the spectral changes observed in the Fo$_{99+}$ pellet sample are initially less than in the Fo$_{90}$ sample, which is consistent with the hypothesis that reddening and darkening are being caused by the production of reduced Fe nanoparticles from the precipitation of Fe present in low concentration in the sample. Given that spectral reddening has been correlated with small npFe$^0$ (Noble et al. 2007), the observation that the Fo$_{99+}$ sample continues to redden up to the point where it is similar in magnitude to the Fo$_{90}$ (Fig. 3) sample suggests that after a large number of laser pulses the small npFe$^0$ present in each sample may be similar. Consequently, the number of large Fe precipitates would have to be much smaller in the Fo$_{99+}$ sample, given the significantly lower concentration of Fe, which is reasonable considering that the Fo$_{99+}$ sample does not

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begin to darken until after 13 laser pulses. It is at this point that the rate of spectral reddening decreases, suggesting that some of the smaller grains precipitated into larger npFe$_0$ particles. Future studies with the electron microscope could confirm this hypothesis.

It is interesting to note that the different weathering rates for our two olivine samples are also consistent with recent laboratory studies (Moroz et al. 2014) on the laser irradiation of natural plagioclase (An$_{47-52}$), which contained an intermediate Fe concentration (0.72 wt% FeO). In those experiments, one laser pulse caused the spectral slope to increase two times more than it increased after a single shot in our Fo$_{99+}$ sample and about two times less than it increased after a single shot in our Fo$_{90}$ sample. Subsequent irradiation of Moroz et al.’s plagioclase sample further increased the spectral slope, suggesting that similar trends as those quantified in our experiments (Fig. 3) would be observed in those experiments with successive irradiation. More detailed comparison between these studies, as well as others, is desired, but is not currently possible given the differences in the experimental conditions, such as laser wavelength and fluence, as well as the likelihood that the mineral composition and structure also affect its susceptibility to weathering (Yamada et al. 1999; Brunetto et al. 2006a; Moroz et al. 2014).

**Comparison to Other Experiments on Fo$_{90}$**

In Fig. 9, we compare these laser irradiation experiments with other irradiation experiments on Fo$_{90}$. To focus on the spectral changes observed in the experiments, we divided the spectrum of the irradiated olivine sample by that of the unirradiated olivine sample. We note that this is a qualitative comparison as the scattering properties of the sample depend on both the grain size and phase angle, and these are slightly different for the different studies. Figure 9 (top) compares experiments where Fo$_{90}$ was irradiated with a ns-pulsed laser (this study), irradiated with 4 keV He$^+$ ions (Loeffler et al. 2009), and coated with vapor produced from irradiating another olivine sample (Loeffler et al. 2008), while Fig. 9 (bottom) compares the reflectance spectra of Fo$_{90}$ taken after a single laser pulse. It is clear from these experiments that regardless of the irradiation approach, the shape of the weathered spectra are similar, which suggests that in all cases these changes are likely caused by the chemical reduction of Fe through the formation of npFe$_0$. Furthermore, judging from Fig. 9 (bottom), the spectral change was the largest when the sample was irradiated at the highest laser fluence.

**Metallic Fe Formation in Fo$_{90}$**

Much of work investigating how the metallic Fe is created during irradiation has been performed previously by Hapke and coworkers (see Hapke [2001] and references therein). The main conclusion from these studies is that the metallic Fe is only formed when material is sputtered from the surface and redeposited, as opposed to being directly produced by irradiation. Later experiments using the laser to evaporate Fo$_{90}$ confirmed with chemical analysis that the evaporated deposits do indeed contain metallic Fe (Loeffler et al. 2008). However, using similar in situ techniques it was also found that metallic Fe was formed when sectioned rocks of olivine, where redeposition is assumed to be negligible, were irradiated with He ions (Dukes et al. 1999; Davoisine et al. 2008; Loeffler et al. 2009), suggesting that direct irradiation can also produce metallic Fe. This debate has also been extended to the spectral changes occurring as a result of laser irradiation (e.g., Sasaki et al. 2001). Thus, we also irradiated multiple flat Fo$_{90}$ sections. In all cases, XPS spectra showed that the Fe in the rock reduced to Fe$_0$ at a faster rate than in the powdered sample, and in some cases the metallic Fe on the surface reached equilibrium after a single laser pulse. The difference between the rock and the powder could be due to a significant percentage of the redeposited material being oxidized, as was observed previously (Loeffler et al. 2008). Yet, given the discussion above regarding how the laser ablation of insulators are strongly affected by the sample roughness (Ihlemann et al. 1992), this conclusion is less definitive than in the case of ion irradiation.
Implications for Natural Space Weathering

The implications of the current experimental results for natural space weathering depend to varying degrees on how well the experiments simulate the processes and products of natural small-scale (e.g., micrometeorite) hypervelocity impacts. Previous studies have used energetic comparisons between laser irradiations and micrometeorite impacts to estimate regolith impact-exposure ages based on laser irradiation experiments (Sasaki et al. 2001; Loeffler et al. 2009). Although the current results could conceivably refine such estimates, the energy comparison approach remains hampered by fundamental unknowns, such as the efficiency at which impact kinetic energy is converted into thermal energy or the relative importance that the momentum transferred from the impactor may play in altering the surface. The problem is compounded by additional lack of knowledge concerning how impact-generated thermal energy itself is converted into a given amount of shock melt and npFe\(^0\), especially when mechanical shock adds to the process. Once these aspects of the hypervelocity impact process become well quantified through theory and experiment, a better basis of extrapolation between laser irradiation results and natural hypervelocity impacts should be possible.

The optical reflectance results, at least for the Fo\(_{90}\) olivine pellet sample, suggest the samples are good spectral analogs for natural space weathered regoliths of similar mineralogical composition, such as ordinary chondrite parent materials on S-type asteroids. Finding naturally space weathered samples of similar mineralogical composition to the Fo\(_{90}\) olivine pellet for microscopic analysis, such as those in olivine-rich meteorite/asteroidal regolith breccias or the Hayabusa samples, has been difficult (Noble et al. 2010; Keller et al. 2013; Noguchi et al. 2014; Thompson et al. 2014). Lunar soils, which have been studied extensively by microscopic analysis, are typically dominated by minerals other than olivine, and thus are harder to compare. However, we point out that the laser-irradiated Fo\(_{90}\) samples studied here appear to be similar in their npFe\(^0\) microstructure to impact vapor and agglutinitic melt deposits in mature lunar soils (e.g., Keller and Clemett 2001; Keller and McKay 1997). For instance, microscope analysis shows that sublayers 1–3 are amorphous, indicating that these regions have either been melted or evaporated and redeposited during laser irradiation. Sublayer one is likely a vapor deposit, as the enrichment of Fe and high modal concentration of npFe\(^0\) <20 nm size present is consistent for what has been found in vapor deposits on lunar regolith grains (Keller and Mckay 1997; Keller and Clemett 2001). Sublayer 2 and 3 are likely melt deposits, as they are chemically similar to the bulk grain but lack the fine-grained npFe\(^0\) found in sublayer 1 characteristic of vapor deposits. The nanocrystalline olivine found in sublayer 3, adjacent to the olivine substrate, also likely crystallized from the melt formed by the laser, possibly promoted by local thermal gradients or heterogeneous nucleation on the underlying olivine.

Assuming there is sufficient basis to use the current Fo\(_{90}\) and Fo\(_{99+}\) sample results as an analog for impact-generated asteroid space weathering, a few implications can be drawn. The step-wise experimental approach of this study, in which optical reflectance was tracked as a function of increasing laser fluence, has produced data supporting the concept that as asteroid regoliths accumulate melt and vapor deposits from micrometeorite impact, their optical properties should reach a steady state or “saturation.” This saturation would necessarily be tied to the amount of npFe\(^0\) produced in the deposits.
as a function of accumulated micrometeorite impacts over time, an amount that may be much smaller compared to lunar samples (Noble et al. 2010; Keller et al. 2013; Noguchi et al. 2014; Thompson et al. 2014). Alternatively, however, other optically active nanoscale grains such as Fe-sulfides could substitute, in their optical effects, for the npFe$^0$ (Keller et al. 2013). In either case, the current results suggest that optical reddening, which is correlated with npFe$^0$ grains less than ~10–20 nm in size (Noble et al. 2007), could be expected to reach saturation at an earlier stage in regolith evolution compared to darkening (Fig. 3). This saturation would be achieved when the volume fraction of sub-20 nm npFe$^0$ in the regolith became similar to that of the Fo$_{90}$ pellet sample after 10 laser pulses (this volume fraction was not measured by TEM or other methods in this study). Darkening, on the other hand, would not reach steady state until the volume fraction of larger (>40 nm) npFe$^0$ grains became similar to the value of approximately 0.5 vol% we estimated for the 99 pulse Fo$_{90}$ pellet sample. Assuming a steady-state micrometeorite flux, with no resurfacing from major impacts, the time needed to reach steady state in the case of either reddening or darkening would necessarily take longer if the olivine had less Fe than the Fo$_{90}$ composition. Whether it would be accelerated for compositions more Fe-rich than Fo$_{90}$ cannot be determined based on our experiments, but has been discussed by Moroz et al. (2014).

CONCLUSIONS

Pulsed-laser irradiation, used to simulate micrometeorite impacts, can alter the reflectance spectrum of Fe-bearing and non-Fe-bearing minerals. For Fo$_{90}$ and Fo$_{99}$+ olivine samples, the irradiation causes the spectral slope to redden and the overall reflectance to decrease. In addition, irradiation causes the band area of the 1 μm absorption band, observable in the Fo$_{90}$ samples, to decrease slightly during irradiation, with no shift in peak position. Laser irradiation of SiO$_2$ results in a reddened spectral slope similar to that of the olivine sample and brightening after an incubation dose. The spectral reddening observed from the ultraviolet to the near-infrared is consistent with the formation of SiO or other SiO$_x$ species, which absorb light in this region. The low degree of brightening is consistent with a change in the surface topography of the sample, which has been observed previously. The observation that the near-infrared spectra of non-Fe-bearing minerals can also be altered via irradiation suggests that other minerals may also contribute to spectral change due to space weathering observed via remote sensing.

In situ surface chemical analysis of the Fo$_{90}$ samples shows that the majority of Fe present on the mineral surface reduces to a metallic state. However, this reduction does not correlate with spectral reddening, as it had in previous ion irradiation studies, as the sample stops reddening while npFe$^0$ continues to increase with further laser irradiation. However, the chemical reduction of Fe does correlate fairly well with the spectral changes if we include both darkening and reddening. This difference, as compared to our previous ion irradiation studies, is reasonable as while successive laser pulses create new small npFe$^0$ particles, they also catalyze the aggregation of small npFe$^0$ into larger npFe$^0$, involved in darkening. This hypothesis is supported by the observation that the sample continues to darken after the reddening reaches equilibrium and by our electron microscope studies, which show the presence of both small (20–50 nm) and large (100–400 nm) npFe$^0$ in the irradiated sample.

Extrapolating our results to airless bodies, the sequence of spectral alteration observed in our Fo$_{90}$ samples, in which end-state or “saturated” optical reddening precedes end-state darkening as laser deposits accumulate, suggests a similar relative sequence may occur on asteroid bodies with regoliths dominated by olivine. The sequence would be tied to differences in formation efficiency, optical absorption efficiency or both, for smaller npFe$^0$ (<40 nm) associated with vapor-dominated deposition that promote reddening, as opposed to larger npFe$^0$ (>40 nm) responsible for darkening.

Acknowledgments—This work was supported by the NASA Cosmochemistry, Lunar Advanced Science and Exploration Research, and Solar System Workings programs, as well as NSF’s Astronomy and Astrophysics Research Grants program (C.D. and R.B.). The authors would like to thank Lindsay Keller for his comments on the manuscript.

Editorial Handling—Dr. Carle Pieters

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