The oxidation state of sulfur in lunar apatite

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ABSTRACT

Lunar apatites contain hundreds to thousands of parts per million of sulfur. This is puzzling because lunar basalts are thought to form in low oxygen fugacity (fO2) conditions where sulfur can only exist in its reduced form (S0), a substitution not previously observed in natural apatite. We present measurements of the oxidation state of S in lunar apatites and associated mesostasis glass that show that lunar apatites and glass contain dominantly S6+, whereas natural apatites from Earth are only known to contain S4-. It is likely that many terrestrial and martian igneous rocks contain apatites with mixed sulfur oxidation states. The S4+/S6+ ratios of such apatites could be used to quantify the fO2 values at which they crystallized, given information on the portioning of S4+ and S6+ between apatite and melt and on the S4+/S6+ ratios of melts as functions of fO2 and melt composition. Such a well-calibrated oxybarometer based on this the oxidation state of S in apatite would have wide application.

Keywords: Moon, oxygen, apatite, sulfur

INTRODUCTION

Lunar apatites, melt inclusions, and glass beads contain concentrations of H, C, S, and Cl that suggest that at least some portions of the lunar crust, mantle, or both contain higher concentrations of these volatile elements than previously thought and perhaps even in concentrations similar to those observed for Earth (Saal et al. 2008; Boyce et al. 2010; Greenwood et al. 2011; Hauri et al. 2011; Chen et al. 2015; McCubbin et al. 2015; Wetzel et al. 2015). This information has led to significant recent interest in lunar volatiles and to efforts to explain these results in the context of models for the formation of the Moon. A puzzling aspect of the unexpectedly high volatile contents of lunar materials was the observation of hundreds to thousands of parts per million of S in apatite, similar to the levels observed in terrestrial igneous apatite (Boyce et al. 2010).

Terrestrial magmas typically formed at fO2 levels 4–5 orders of magnitude higher than the iron-wüstite oxygen buffer (referred to as IW+4 to IW+5). At these fO2 values, sulfur dissolved in the silicate liquids from which terrestrial apatites crystallize is present as sulfate (i.e., SO42- or S4+) and sulfide (i.e., S2-), but it has been generally believed that the sulfur in terrestrial apatites is present entirely as sulfate (Fleet 2005), with the sulfate anion in the apatite substituting for the phosphate anionic group, coupled with SiO44- or Na+ to maintain charge balance (Pan and Fleet 2002). However, lunar rocks reflect much lower fO2 conditions, including at the point of apatite saturation and crystallization. Lunar apatites crystallize in the interstices of lunar basalts from late-stage, highly differentiated liquids, since only in such liquids does phosphorus reach sufficient concentrations for the liquids to become saturated with respect to apatite. The fO2 of these late-stage liquids in lunar basalts are constrained from petrographic descriptions of the phases present in the interstices (e.g., coexistence of Fe-metal, ulvospinel, ilmenite, and sometimes fayalite and/or silica) and are as low as IW-1 (e.g., El Goresy 1976). This is more than 4 orders of magnitude lower than the fO2 values required to begin to stabilize sulfate in basaltic and andesitic melts (~IW+3.5; Botcharnikov et al. 2010; Jugo et al. 2010), and more than 3 orders of magnitude lower than needed in Fe-free or Fe-poor soda-lime, K2SiO3, albite, and hapol-trondhjemite liquids (~IW+2.5; Klimm et al. 2012). Thus, under the reducing conditions of lunar petrogenesis, it is anticipated that nearly all S dissolved in the interstitial silicate melts in these lunar magmas will be dissolved as sulfide (S2-), and that crystalline phases in equilibrium with the interstitial silicate melts likely contain nearly entirely sulfide. Since at the time of the discovery of hundreds to thousands of parts per million of S in lunar apatites sulfur was only known to be present in naturally occurring apatite as sulfate groups, it was not clear how to explain the incorporation of sulfur in lunar apatites (Boyce et al. 2010).

Boyce et al. (2010) speculated that the elevated S abundance of lunar apatite was due to sulfide substitution into the column anion site in apatite where F-, OH-, and Cl- anions normally sit. Although this hypothesized substitution has until now not been observed in natural apatites, in support of their speculation, Boyce et al. (2010) pointed out that fully S2- substituted apatites had been synthesized experimentally (Stuchl et al. 1986; Taitai and Lacout 1989; Henning et al. 2000), and since then, apatites that exhibit S-XANES spectral evidence for both sulfide and sulfate (Konecke et al. 2017a) have been synthesized.

The speculation that this occurs at the column anion site is also supported by ab initio calculations, which show that this substi-
tution of $S^2$ is most energetically favorable in chloride-bearing apatites (Kim et al. 2017). If natural apatite can simultaneously incorporate both oxidized and reduced sulfur ($S^{0}$ and $S^2$), it is possible that their proportions could be sensitive to the $S^0/S^2$ ratio of the apatite growth environment, and therefore could provide a proxy for the $f_O$ at which the apatites formed. This possibility is of interest because apatites are a common igneous phase in various rocks from Earth, Moon, and other planets and the determination of $f_O$ is a topic of considerable interest and importance (e.g., Haggerty and Meyer 1970; Taylor et al. 1972, 2004; Sato et al. 1973; Sato 1979; Carmichael 1991; Steele et al. 1992; Herd et al. 2002; Karner et al. 2006; Wadhwa 2008; Kelley and Cottrell 2009; Jugo et al. 2010; Wetzel et al. 2013; Brounce et al. 2017; Konecke et al. 2017a). In this paper, we report the results of X-ray absorption near edge structure spectroscopy (S-XANES, see SI Appendix) on S-bearing lunar apatite (from basalt samples 10044 and 12039) and terrestrial igneous apatite (from Durango) and demonstrate that $S^2$ dominates in the lunar apatites but is undetectable in the terrestrial apatite.

**SAMPLE DESCRIPTIONS**

Sample 12039 is a 3.2 ± 0.05 billion year old, low-TiO$_2$ basalt (Nyquist et al. 1979). Sample 10044 is a ~3.71–3.73 billion year old, high-TiO$_2$ basalt (Turner 1970; Guggisberg et al. 1979). Both rocks are inferred to have derived from lave flows (Klein 1972). They are slightly vesiculated (~1%) and dominated by compositionally zoned and skeletal pyroxene, plagioclase, and ilmenite (Klein 1972). For both samples, fractionation of the basaltic magma due to crystallization at low $f_O$ produced Fe-rich, late-stage liquids, which then separated into two immiscible liquids upon further crystallization—one very Si-rich, Fe-poor melt that occurs as quenched glass containing 78 wt% SiO$_2$, 0.5 wt% FeO$^+$, 3.2 wt% K$_2$O, 4.48 wt% CaO, and ~0.03 wt% MgO (vs. the bulk lava with ~47 wt% SiO$_2$, 20.39 wt% FeO$^+$, 0.11 wt% K$_2$O, 11.6 wt% CaO, and ~8.6 wt% MgO; Boyce et al. 2014), and another lower silica, Fe-rich melt (SiO$_2$ ~47.8 wt%, FeO$^+$ ~20 wt%, K$_2$O ~0.3 wt%, CaO ~11.2 wt%, MgO ~2.3 wt%; Roedder and Weiblen 1970; Pernet-Fisher et al. 2014). Apatite can be in contact with either glass in both samples (Pernet-Fisher et al. 2014), and recent experimental data indicate that the composition of apatites among conjugate liquids are indistinguishable with respect to major and minor elements in systems that undergo silicate-liquid immiscibility (McCubbin and Ustunisik 2018). Sulfide blebs are also associated with silicate glass in the mesostasis in both samples, suggesting that apatites analyzed in this study crystallized from and equilibrated with a sulfide-saturated silicate melt (Figs. 1–3). Features such as rounded silicate glass-sulfide contacts (e.g., rounded sulfide/K-glass contact in the upper right region of Figs. 1a and 1b) indicate that the sulfide was molten. The mesostasis in the thin section of sample 12039 studied here (thin section 4) also contains K,Si-rich glass, K,Ba-feldspar, troilite (distinguished from sulfide on the basis of the shape of their XANES spectra, Figs. 1 and 2), plagioclase, and pyroxenes (e.g., Fig. 1). Other studies of sample 12039 also report tranquillityite, native iron, cristobalite, and tridymite in the mesostasis (Bunch et al. 1972). The mesostasis in sample 10044 studied here (thin section 33) contains crystallized troilite, plagioclase, fayalite, SiO$_2$, and pyroxene (e.g., Fig. 3). Other studies of sample 10044 also report bauleyite, ulvöspinel, tranquillityite, K,Ba-feldspar in hand sample, as well as K,Si-rich glass and devitrified high-Fe glass, the last two of which are interpreted as being immiscible liquids (Beatty and Albee 1978).

Apatites in thin sections of both rocks have igneous textures as indicated by equant and skeletal grains with central cavities filled with glass, as well as compositionally zoned crystals (see Figs. 1 and 3; Piccoli and Candela 2002). These apatites contain hundreds to thousands of parts per million of S (determined via electron probe; Greenwood et al. 2011). Apatite grains in sample 12039 have been shown to be zoned with respect to F, Cl, SiO$_2$, and S, which was hypothesized to reflect variations in the composition of a residual melt during apatite crystallization (Greenwood et al. 2011).

Durango apatite is from volcanogenic deposits near Durango, Mexico, and is associated with magnetite in gas cavities and open breccias in sheeted flows and flow breccias from Cerro de Mercado. Halogen-rich gases are thought to have streamed through and further oxidized magnetite to hematite, setting the $f_O$ conditions under which Durango apatite formed at ~IW+6 (Lyons 1988). At these $f_O$ values, sulfur is expected in solution as $S^0$ (Botcharnikov et al. 2010; Jugo et al. 2010; Klimm et al. 2012).

**RESULTS**

The measured S$^0$/$S^2$ ratios vary between 0 and 45% in six lunar apatite grains in thin section 12039,4 (Figs. 1 and 2, Supplemental Fig. 2 and 3; Supplemental Table 1) and between 0 and 32% in two apatite grains in thin section 10044,33 (Fig. 3, Supplemental Fig. 4; Supplemental Table 1). The sulfur

![Figure 1](image_url) - **Figure 1.** (a and b) Backscatter electron and (c) S-P-Na (R-G-B) maps that show the locations of analyses in thin section 12039,4 area 4. Black circles in a mark the locations of cracks and pits that cross the analysis transect. Phases are labeled b (Ap = apatite; S = sulfide; Fa = fayalite; Plag = plagioclase; Pyx = pyroxene; K,Ba-feld = K,Ba-feldspar; K-glass = K-rich mesostasis glass). In c, sulfide grains/blebs appear as red, apatite grains appear as green, and mesostasis glass appears as bright blue. The size of the beam is smaller than the symbol size. Hollow diamonds are saved stage motor positions that were not analyzed. Filled white, yellow, and red diamonds are analyzed positions. (d) Calculated S$^0$/$S^2$ ratio from spectra at each analysis point from Figure 1. Locations of cracks and pits near analysis points (see a) are marked by gray dashed lines. Dark gray field marks S$^0$/$S^2$ > 3%. Analytical uncertainties are smaller than symbol size. (Color online.)
concentrations of the K, Si-rich mesostasis glass touching measured apatites in sample 12039 contain 69–107 ppm S, and two S-XANES analyses of the mesostasis glass demonstrate that the sulfur present in the mesostasis has the characteristic spectral features of dissolved S\(^{2-}\) in terrestrial basaltic liquids (e.g., Jugo et al. 2010; Brounce et al. 2017) but no spectral evidence for the presence of S\(^{6+}\) dissolved in the glass, resulting in a S\(^{6+}/\Sigma S\)~ 0% (Fig. 2). More than half of all measurements on apatite (28 analyses out of 48 total analyses) have S\(^{6+}/\Sigma S\) < 3%, which we consider to be indistinguishable from 0% (see SI Appendix). This is in contrast with our analyses of Durango apatite (the representative of terrestrial apatite considered in this study), which shows the characteristic spectral features of S\(^{6+}\), but no spectral evidence for the presence of S\(^{2-}\), resulting in a S\(^{6+}/\Sigma S\) = 100% (Supplemental1 Fig. 1).

Sulfur abundances of the same apatites and glasses were measured via electron microprobe (see SI Appendix) and vary from below the detection limit (~20 ppm based on measurements of nearby silicate minerals that are nominally free of S) up to 500 ppm in thin section 12039,4 (Supplemental1 Figs. 2, 3, and 5; Supplemental1 Table 1) and up to ~350 ppm in thin section 10044,33 (Supplemental1 Figs. 4 and 6; Supplemental1 Table 1). Individual apatite grains are heterogeneous with respect to S. Where analyzed, the S abundances of the mesostasis glass in 12039,4 range from below the detection limit to ~100 ppm S (Supplemental1 Fig. 5; Supplemental1 Table 1).

**FIGURE 2.** (a) Normalized S-XANES spectra for (top panel) gypsum (black curve), troilite (gray curve), and sulfide blebs in Apollo sample 12039,4 area 4 (red and yellow curves) and (bottom panel) epoxy in thin section 10044,33 (light green curve) and 12039,4 (dark green curve). (b) S-XANES spectra for analysis points on apatite (q, n), mesostasis glass (i), and sulfide bleb (j). In a and b, the position of absorption peaks traditionally assigned to S\(^{2-}\) (2466, 2470, and 2478 eV) and S\(^{6+}\) (2482 eV) are marked in vertical black dashed lines. The black curves are data; the green curves are synthetic spectra produced from linear combinations of spectra collected on gypsum and troilite to provide a best fit to the data (see Supplemental1 Materials). (Color online.)

**DISCUSSION**

In apatites from both 12039 and 10044, the locations of analyses that yield S\(^{6+}/\Sigma S\) > 3% correspond with the presence of fractures, pits, or both (Figs. 1–3; Supplemental1 Figs. 2–4). This suggests that the S\(^{6+}\) observed in these S-XANES spectra may not reflect primary sulfur incorporated into the apatite when it crystallized, but instead is either primary sulfide altered to sulfite or sulfur of secondary origin. Although it is possible that some portion of the S\(^{6+}\) signal is derived from the epoxies used in making the thin sections, the sharp absorption feature that appears at ~2474 eV in the epoxies, which does not appear in the mineral standards or in apatite measurements, and the low intensity of the S\(^{6+}\) peak in the epoxy (~2482 eV; Supplemental1 Fig. 1) in each thin section limits the proportion of epoxy that can contribute to the measured absorption spectra in apatite to <1% (Supplemental1 Fig. 1). There also could be secondary sulfate-bearing materials precipitated in the fractures and pits of the thin sections of these rocks, but, if so, we have no basis for evaluating whether these could be of terrestrial or lunar origin. Chlorine isotope measurements of the so-called “rusty” rock (Apollo sample 66095) suggest a lunar origin for oxidation- and hydration-induced, Cl-bearing minerals observed in Apollo 16 samples (Shearer et al. 2014), and it is thus possible that apatites in Apollo 11 and 12 samples studied here are susceptible to similar fumerolic alteration (e.g., Konecke et al. 2017b). On the other hand, measurements of the hydrogen isotope contents of lunar apatites near fractures reveal distinctly terrestrial surface water signatures (Greenwood et al. 2011), and this terrestrial water contamination may have been associated with deposition of micro-to-nano-scale S\(^{6+}\)-bearing phases. Finally, while all spectra that indicate S\(^{6+}/\Sigma S\) > 3% are near fractures and pits in the thin section, there are some spectra that are near fractures or pits and do not have significantly elevated S\(^{6+}/\Sigma S\) ratios (Figs. 1–3; Supplemental1 Figs. 2–4), suggesting that if the oxidized sulfur we have detected is from secondary sulfate-bearing materials in the fractures and pits of these thin sections, these sulfates are heterogeneously distributed. The key point is that analyses taken far from fractures and pits are uniformly lacking in S\(^{6+}\) absorption features (i.e., S\(^{6+}/\Sigma S\) < 3%; e.g., Fig. 3).

Lunar basalts are also variably affected by subsolidus reactions, for instance, as the result of interaction between their primary mineral assemblages and implanted solar wind hydrogen (e.g., sample 14053; Taylor et al. 2004). It is possible that some or all of the S\(^{2-}\) found in lunar apatites measured in this study was formed as the result of interaction between solar wind hydrogen and sulfur-bearing phases such as troilite (i.e., reducing Fe\(^{2+}\) to Fe\(^{0}\) and making S\(^{2-}\) available to diffuse into pre-existing, sulfide-poor or sulfide-free apatite), and thus do not record the f\(_{O_2}\) of the igneous system described by the bulk rock. However, unlikeapatite grains in sample 14053, where solar wind implantation may have lowered the hydrogen isotopic compositions of apatite grains in thin section from their primary values, the hydrogen isotopic compositions of apatite grains in samples 12039 and 10044 are higher (δD in apatites from 14053 vary from ~100 to ~200‰, from 12039 vary from ~400 to ~1000‰, from 10044 vary from +550 to +750‰). (Greenwood et al. 2011). Solar wind has low D/H ratios, so the D-enrichment of apatite in the rocks...
studies here limits the extent to which solar wind implantation influenced the incorporation of $S^{2-}$ into apatite. Apollo rocks are also variably influenced by impact-related metamorphism and metasomatism. For example, abundant apatite found in granulate 79215 was hypothesized to be condensed from a halogen-rich vapor (Treiman et al. 2014). However, apatite grains in the mare basalts studied here have distinctly igneous textures, including being embayed by and displaying resorption textures with the mesostasis melt (e.g., Figs. 1 and 3, Supplemental Figs. 2–4), indicating that apatite was present in the rock at the time the mesostasis was liquid. Our preferred interpretation of $S^{6+}$-only lunar apatites in this study is that this reflects the low $f_{O_2}$ conditions of the magma at the time of apatite crystallization.

The mesostasis glass in sample 12039 contains 70–110 ppm sulfur and that sulfur is entirely present as $S^{6+}$ (Supplemental Table 1, Supplemental Fig. 5). For Apollo 12 basalts, experiments and petrographic observations suggest that olivine and pigeonite are the liquidus phases at an $f_{O_2} \sim$ IW (Green et al. 1971; Rhodes et al. 1977). The presence of Fe-metal and troilite, and in some cases fayalite, Fe-metal, and silica, or of ulvöspinel, Fe-metal, and ilmenite in the mesostasis reflects $f_{O_2} \ll$ IW-1, consistent with differentiation (i.e., low-pressure crystallization and degassing) having led to a decrease of at least an order of magnitude in $f_{O_2}$. This could be driven by degassing of a C-H-O-S vapor, which has been shown to be an effective means of reducing the $f_{O_2}$ of Hawaiian magmas by ~1 order of magnitude (Moussallam et al. 2016; Helz et al. 2017; Brounce et al. 2017), and has been suggested for lunar basalts previously (Brett 1976).

Based on these previous results, the liquids from which lunar apatites in basalts crystallized clearly experienced $f_{O_2}$ values at which sulfur is expected in solution as $S^{2-}$, as confirmed by our S-XANES analyses of the mesostasis glass from sample 12039 and consistent with our S-XANES measurements on apatite from cracks and pits from both 12039 and 10044 (Figs. 1–4). These latter results contrast with our S-XANES results on the Durango apatite, which has $S^{6+}/2S = 100\%$ (Supplemental Fig. 1).

Paired glass-apatite measurements in sample 12039 enable an estimate of the partition coefficient of sulfide for apatite coexisting with K-Si-rich mesostasis melt. Taking the average sulfide concentration of the mesostasis glass in sample 12039 (88 ppm; Supplemental Table 1) and of the apatite measurements in the same thin section (176 ppm, assuming that $S^{6+}/2S$ of lunar apatite at the time of apatite crystallization is zero), then for conditions appropriate for apatite crystallization in the mesostasis of sample 12039, $D_{Ap/lq}^{S^{6+}} \sim 2 \pm 1$. Importantly, however, apatite-melt coefficients may vary as a function of temperature, pressure, melt composition, apatite composition, and/or oxygen fugacity, so the combined effects of these parameters must be investigated further before being applied broadly to determine sulfur abundances of coexisting melt from apatite $S^{2-}$ abundances.

**Figure 3.** (a and b) Maps as in Figure 1 that shows the locations of analyses in thin section 10044.33. (e) S-XANES spectra for analysis points on apatite. The position of absorption peaks traditionally assigned to $S^{2-}$ (2466, 2470, and 2478 eV) and $S^{6+}$ (2482 eV) are marked in vertical gray lines. All apatite measurements lack spectral evidence for $S^{4-}$ and thus have $S^{6+}/2S = 0$. (Color online.)

**Figure 4.** The relationship between $S^{2-}/2S$ ratios of igneous apatites and $f_{O_2}$ at the time of apatite crystallization from natural apatites (gray stars = this study) and experimentally grown apatites (white stars = Konecke et al. 2017a). Error bars on gray stars represent the full range of $S^{2-}/2S$ ratios observed in measurements of lunar apatites and Durango apatite (i.e., lunar apatite measurements yield $S^{2-}/2S$ ratios as high as 45%, but we hypothesize that the sulfate signal originates from alteration or secondary mineralization after sample collection; Durango apatite measurements show no spectral evidence of sulfide). Error bars on the experimentally grown apatite data point reflect the full range of $S^{2-}/2S$ of apatite reported for that experiment equilibrated at IW+3 (Konecke et al. 2017a). Also shown are the current accepted $f_{O_2}$ ranges of typical terrestrial and martian magmas (black horizontal lines with arrows). Larger dark star = Natural apatites, this study. Smaller open star = Experimentally grown, Konecke et al. (2017). terr = terrestrial.

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**IMPLICATIONS**

We have documented the occurrence of S\(^2\)-only apatite in nature (Figs. 1 and 3; Supplemental Figs. 3 and 4). Although not unexpected from the experimental synthesis of sulfosaapatites, ab initio calculations, and recent synthesis of mixed S\(^6\)/S\(^2\) apatites, this is the first documentation of its natural occurrence. This observation confirms speculation of Boyce et al. (2010) that the elevated S contents of lunar apatites crystallized from melts that record \(f_{\text{O}_2}\) values at which silicate melts are expected to have only S\(^2\) are due to the ability of S\(^2\) to partition into the apatite mineral structure. Finally, this study, in combination with the recent experimental work of Konecke et al. (2017a, 2017b), suggests that a S-in-apatite oxybarometer could be developed and applied to igneous rocks from various planetary bodies in our solar system.

The S\(^6\)/S\(^2\) of an individual silicate melt transitions from 0 to 100% over ~2 orders of magnitude in \(f_{\text{O}_2}\). The range over which this transition occurs for an individual melt depends on its major element composition, i.e., the midpoint of this transition for the melts for which this transition has been measured varies from QFM-0.5 (IW+3) to QFM+1 (IW+4.5) (Botcharnikov et al. 2010; Jugo et al. 2010; Klimm et al. 2012). Apatites coexisting with a given melt over the \(f_{\text{O}_2}\) range of this transition are anticipated to contain both S\(^6\) and S\(^2\) and thus could be used to quantify the \(f_{\text{O}_2}\) of the system at the time of apatite crystallization. In detail the \(f_{\text{O}_2}\) dependence of the transition in S\(^6\)/S\(^2\) for apatites coexisting with melt undergoing the same transition will depend on the individual partition coefficients of S\(^6\) and S\(^2\) between apatite and melt and the major element composition of the melt (including the oxidation states of other heterovalent elements). Thus, development of an oxybarometer based on the S\(^6\)/S\(^2\) of apatite would require experimental work to determine the individual partition coefficients of S\(^6\) and S\(^2\) between apatite and melt, and a quantification of the extent to which those partition coefficients vary as a function of pressure, temperature, oxygen fugacity, melt composition, and apatite composition. Although such experiments and measurements will probably not be simple, given the widespread occurrence of apatite in igneous rocks from Earth and other planets, and the interest in the \(f_{\text{O}_2}\) values at which they formed (and their variations), such an oxybarometer would likely be useful. In particular, given the \(f_{\text{O}_2}\) range over which this transition is expected to occur for terrestrial magmas based on their major element compositions (Botcharnikov et al. 2010; Jugo et al. 2010), the \(f_{\text{O}_2}\) ranges of terrestrial magmas (Carmichael 1991), and the capacity of XANES for measurements of sulfide/sulfate ratios in experimental and natural glasses (Brounce et al. 2017), S-XANES measurements can be expected to readily detect sulfide-dominated apatites, mixed sulfide-sulfate apatites, and sulfate-dominated apatites (as in the case for the Durango apatite we studied) in terrestrial igneous rocks and that these results can, with appropriate experimental calibration, be used quantitatively as an apatite oxybarometer for igneous rocks ranging in composition from basalt to rhyolite. In addition to Earth and the Moon, apatites are found in lavas from planetary bodies distributed throughout the solar system (McCubbin and Jones 2015) and an apatite-based oxybarometer could be useful in constraining the \(f_{\text{O}_2}\) of lavas from Mars and the howardite-eucrite-diogenite parent body (e.g., Herd et al. 2002; Shearer et al. 2006; Wadhwa 2008).

Finally, we note that S\(^2\)-likely partitions into the column anion site in apatite as a substitution for two F\(^-\), Cl\(^-\), and/or OH\(^-\) anions. Thus, the abundance of S\(^2\) in apatites may be useful in studies seeking to constrain the abundance of F\(^-\), Cl\(^-\), and H\(_2\)O in magmas based on their concentrations in apatite.

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