Ferric saponite and serpentine in the nakhlite martian meteorites

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Abstract

Transmission electron microscopy and Fe-K X-ray absorption spectroscopy have been used to determine structure and ferric content of the secondary phase mineral assemblages in the nakhlite martian meteorites, NWA 998, Lafayette, Nakhl, GV, Y 000593, Y 000749, MIL 03346, NWA 817, and NWA 5790. The secondary phases are a rapidly cooled, metastable assemblage that has preserved Mg# and Ca fractionation related to distance from the fluid source, for most of the nakhlites, though one, NWA 5790, appears not to have experienced a fluid pathway. All nine nakhlite samples have also been analysed with scanning electron microscopy, electron probe micro analysis, Bright Field high-resolution transmission electron microscopy (HRTEM) and selected area electron diffraction. By measuring the energy position of the Fe-K XANES 1s → 3d pre-edge transition centroid we calculate the ferric content of the minerals within the nakhlite meteorites. The crystalline phyllosilicates and amorphous silicate of the hydrothermal deposits filling the olivine fractures are found to have variable Fe3+/ΣFe values ranging from 0.4 to 0.9. In Lafayette, the central silicate gel parts of the veins are more ferric than the phyllosilicates around it, showing that the fluid became increasingly oxidised. The mesostasis of Lafayette and NWA 817 also have phyllosilicate, which have a higher ferric content than the olivine fracture deposits, with Fe3+/ΣFe values of up to 1.0. Further study, via TEM analyses, reveal the Lafayette and NWA 817 olivine phyllosilicates to have 2:1 T–O–T lattice structure with a d001-spacing of 0.96 nm, whereas the Lafayette mesostasis phyllosilicates have 1:1 T–O structure with d001-spacings of 0.7 nm. Based on our analyses, the phyllosilicate found within the Lafayette olivine fractures is trioctahedral ferric saponite (Ca0.2K0.1P0.3(Mg2.6Fe2+1.3Fe3+1.7Mn0.1)P5.7[(Si6.7Al0.9Fe3+0.4)P8.0O20](OH)4)/C1nH2O, and that found in the mesostasis fractures is an Fe-serpentine (Ca0.1Mg0.7Fe3+1.0Al0.4)P2.2[Si2O5]OH4, with a ferric gel of similar composition in Lafayette and found as fracture fills throughout the other nakhlites.

1. INTRODUCTION

The eight nakhlites (plus six pairings) are a group of the SNC martian meteorites. Their unique combination of known formation depths (Mikouchi et al., 2003, 2012) and hydrothermal veining (Gooding et al., 1991; Changela and Bridges, 2010) makes them an important source of information about water-crust interaction on Mars (Bridges and Schwenzer, 2012).

The nakhlites are basaltic cumulate clinopyroxenite rocks, comprised mainly of sub-calcic augite with approximately 10% Fe-rich olivine, in all but one of the nakhlites (Miller-Range 03346). The cumulus grains are set in a mesostasis which includes plagioclase, K-feldspar, and Ti-magnetite (Treiman, 2005; Bridges and Warren, 2006; McCubbin et al., 2013). The nakhlites are thought to have formed in a thick basic–ultrabasic lava flow or shallow intrusion (Treiman et al., 1993). There are currently eight known distinct nakhlites, with several additional pairings.

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for the Yamato (Imae et al., 2002) and Miller-Range (Hallis and Taylor, 2011; Udry et al., 2012) meteorites.

Mikouchi et al. (2003, 2012) used likely igneous cooling rates to determine the formation depths of the nakhlites. Their study suggests MIL 09030/032/136 and Northwest Africa (NWA) 5790 crystallised 1–2 m from the surface, followed with increasing depth by NWA 817, Miller-Range (MIL) 03346, Yamato (Y) 000593 paired with Y 000749, Governador Valadares (GV) along with Nakhl, and to the bottom with Lafayette and NWA 998 at a burial depth of ~30 m.

Secondary minerals are present in veining within brittle fractures in the olivines and mesostasis (Bridges and Grady, 2000). Observations of truncation by fusion crust established the martian origin of the majority of the veining material in Nakhl (Gooding et al., 1991) and Lafayette (Treiman et al., 1993). Most of the vein deposits in Lafayette feature crystalline smectite phyllosilicates; the rest of the nakhlite samples feature brittle fractures dominated by poorly crystalline silicate gel (Changela and Bridges, 2010). Governador and Nakhl also contain siderite, and Nakhl has anhydrite and halite grains (Bridges and Grady, 2000).

The nakhlite hydrothermal assemblage resulted from an impact adjacent to the nakhlite parent rocks, as suggested by Changela and Bridges (2010) on the basis of saw-toothed, brittle fractures in the nakhlites. The fracture fills resulted from fluids circulating due to increased temperatures associated with impact and then rapid cooling. After the impact a CO₂-rich fluid at 150–200 °C, pH 6–8 with a water:rock ratio (W/R) < 300 precipitated Ca-rich siderite, predominantly within olivine. As the fluid cooled to 50 °C, at pH 9 and W/R of 6, Fe-rich phyllosilicate precipitated, followed in turn by the rapid formation of an amorphous gel (Bridges and Schwenzer, 2012). Changela and Bridges (2010) showed that the composition of gel and smectite varied with higher Mg# (Mg# = 100 Mg/(Mg + Fe)) in the base of the nakhlite pile and lower Mg# towards the top.

The nakhlites have a common crystallization age of 1.3 Ga, shown by Rb–Sr, Sm–Nd, U–Pb, and ⁴⁰Ar/³⁹Ar data (Nyquist et al., 2001). Cosmogenic nuclides ⁴He, ²¹Ne, and ⁴⁰Ar in the nakhlites suggest that they were all ejected from Mars in the same event 10–12 Ma (Korochantsyeva et al., 2011). Several methods have been implemented to measure the age of the hydrothermal assemblage, including Rb–Sr and Sm–Nd isotopic analyses (Shih et al., 1998; Misawa et al., 2003), and K–Ar (Swindle et al., 1997). The likely date of formation is taken as ≤670 Ma (Swindle et al., 2000).

TEM studies, by Changela and Bridges (2010), of the Lafayette veining material showed an Fe-rich smectite with lattice fringe d-spacings of 0.9–1.1 nm, sometimes sandwiched between 0.7 nm fringes, suggesting a mixture of smectite and serpentinite. The serpentine deposits were usually found amongst the Lafayette mesostasis, whereas the smectite formed mostly in the olivine fractures. The amorphous gel in the centre of veins is similar in composition to the smectite and serpentinite. In the current study, we have performed more TEM analysis in order to further characterise the crystalline and amorphous alteration phases in the nakhlites. In order to do this we have used High Resolution TEM and electron diffraction to measure lattice spacings and assess the extent of abundance of amorphous versus crystalline material of smectite composition. We have also used Fe-K X-ray Absorption Near-Edge Structure (XANES) and Extended X-ray Absorption Fine Structure (EXAFS), in order to determine the ferric-ferrous ratio semi-quantitatively. Estimating the oxidation state allows us to further constrain the identity of the secondary phase assemblage minerals found in the nakhlites, and provide a ground truth to the alteration minerals observed on the surface of Mars by remote sensing and lander missions (Bibring et al., 2006; Mustard et al., 2008; Michalski and Niles, 2010; Mangold et al., 2012).

2. METHODS AND SAMPLES

Nine nakhlites have been studied in polished sections and resin blocks: Lafayette (BM 1958, 775); Nakhl (BM 1911, 369); GV (BM 1975, M16); Y 000593 and Y 000749; MIL 03346; NWA 817; NWA 998; NWA 5790. The latter four samples are additional to the five nakhlites studied by Changela and Bridges (2010).

Veins were identified and characterised using Back Scattered Electron (BSE) imagery and Energy Dispersive X-ray (EDX) spectroscopy using a Phillips XL30 ESEM at the University of Leicester (UL) Advanced Microscopy Centre. EDX spectra were measured, with an accelerating voltage of 20 kV and beam current of ~1.0 nA. To assess the extent of the veining and the fraction percentage of the olivine grains in which they are found, the SEM images were observed using Adobe Photoshop to select the veining material and compare the pixel count to that of the olivine grains (including the veins) to calculate the fraction, from which an average is taken over several veins per nakhlite.

Following SEM analyses, Electron Probe Microanalysis (EPMA) data on some of the phyllosilicate and gel veins were taken using a Cameca SX100 at the Open University, UK. The measurements were made using TAP, LTAP, LLiF, LPET, and PET crystal spectrometers, a 10 µm focused beam, at an accelerating voltage of 20 kV, and beam current 20 nA.

Wafers measuring 5 µm x 15 µm and 50–100 nm thick, suitable for Transmission Electron Microscopy (TEM) analysis, were prepared using a FEI Quanta 200 3D Dual Focused Ion Beam (FIB-SEM) at UL. TEM analyses were performed with a JEOL 2100 TEM at UL. All HRTEM images were made with an accelerating voltage of 200 kV and beam current of ~110 µA. The d₀₀₀ basal spacings were measured from the HRTEM BF images using the Gatan Digital Micrograph Camera program, with an error of up to ±0.05 nm based on imaging pixel resolution. The measurements were typically taken from an image that had been filtered by “masking” the unwanted features in a FFT image, thus revealing the lattice structure with greater clarity. Further analyses in the TEM also utilised selected aperture electron diffraction (SAED), measured with an accelerating voltage of 200 kV and a camera length of 21.15 cm. The SAED imaging was calibrated in the Gatan Digital Micrograph Camera program, using an Agar
graphitized carbon standard with lattice spacings of 3.4 Å, which allows measurements to be a simple process of inverting the observed diffraction ring radius by 1 nm to find the lattice spacings of our samples.

The ferric-ferrous ratio (Fe\(^{3+}\)/2Fe) has been calculated from stoichiometry for reference materials: the Barwell L6 olivine (\((\text{MgFe})_2\text{SiO}_4\)\)); siderite (FeCO\(_3\)); magnetite (Fe\(_3\)O\(_4\)); two hematites (Fe\(_2\)O\(_3\)); and goethite (FeO(OH)), which were checked for purity with SEM-EDX. Fe\(^{3+}\)/2Fe values for the nakhlite Y 000749 olivine, nakhlite NWA 817 pyroxene, standard San Carlos olivine (Mg\# = 0.90), and a chromite were estimated using EPMA data, normalising to a total sum of 3 cations based on 4 oxygens for the olivines, 4 cations based on 6 oxygens for the pyroxene, and 24 cations based on 32 oxygens for chromite (Deer et al., 1992). Fe\(^{3+}\)/2Fe values for the nontronites (NG-1 and NAu-2), illite (IMt-1), and peridotite (JP-1) were calculated from online reference material chemical compositions (Geological Survey of Japan/AIST, 1995; The Clay Minerals Society, 2013).

In order to analyse the 1s → 3d transitions and the Fe bonding environment, X-ray Absorption Near-Edge Structure (XANES) and Extended X-ray Absorption Fine Structure (EXAFS) measurements were collected at the I-18 Beamline, of the Diamond Light Source synchrotron, Oxfordshire, UK. XRF mapping of the area of interest with a spatial resolution of 2 × 2.5 μm, observing elements Z > 20, was used, followed by Fe-K XANES and EXAFS measurements. All of the nakhlite samples and two standard reference materials (Barwell L6 olivine and a San Carlos olivine) were measured in fluorescence mode at 45° to the beam and the detector, with the remaining twelve reference materials (including Nontronite #142) in powdered form measured in transmission mode. All measurements were taken over an energy range 6900–7500 eV, with varying energy resolutions, focussing on a high resolution of 0.1 eV over the XANES region 7090–7145 eV, followed by varying energy steps of 0.2–0.4 eV over the EXAFS region up to 7500 eV.

The resultant data were processed and normalised through Athena, using the data reduction/fitting engine IFEFFIT. The 1s → 3d centroids were analysed using the normalised Fe-K XANES data. Based on a similar method used in Berry et al. (2003) and Wilke et al. (2001), a curved baseline is fitted under the 1s → 3d pre-edge peaks. This is achieved by firstly finding the boundaries of the 1s → 3d pre-edge region, by fitting a linear baseline across the peaks, and then increasing these boundary points by 1 eV to either side of the region and using these points to fit a curved baseline. This curved baseline is based on the gradient of the spectra at the boundary points before and after the 1s → 3d pre-edge region. Subtracting this baseline from the spectra, the energy position of the centroid is then calculated from the intensity-weighted average over the 1s → 3d pre-edge peaks.

For EXAFS analyses, the normalised data was analysed further using PySpline 1.2 in order to process the data suitable for analysis using EXCURV (Roy and Gurman, 2001). The results of this analysis reveal the bond lengths of neighbouring atoms, such as O and Si, to the Fe atoms with which the beam interacts. Error values in the bond lengths are given for each individual analysis, based on a 2-sigma fitting over the peaks in the Fourier Transform, and are calculated in EXCURV.

In addition to measuring Fe-K XANES and EXAFS at specifically chosen locations of 2 × 2.5 μm spot size, Fe-K XANES maps were made, visually showing the variation in ferrous/ferric proportions by measuring a mapping region at chosen energies, and repeating the maps in increments of increasing energy. Two maps were collected from Lafayette and Nakhl, of size 50 × 60 μm with spot size resolution 2 × 2.5 μm, over the energy ranges 7075–7215 eV i.e. over the Fe K absorption edge. The Fe-K XANES spectra produced for each pixel of the map were then normalised and analysed using MANTIS 1.10.

The various data presented here were measured from three separate sessions at the I-18 Beamline. To account for any variation in the exact position of peaks on the energy range from one session to another, the same standards, including Fe foil, olivine, and hematite, and specific locations on nakhlite samples, were measured at each session.

3. RESULTS

3.1. Alteration petrography

Martian hydrothermal minerals were found in all of the nine nakhlite samples, except NWA 5790. Between 5 and 22 veins were studied per nakhlite. The hydrothermal veins typically have a vein width of 5–10 μm, or can be up to ~26 μm in the case of Lafayette and NWA 817 (see Fig. 1). A similar pattern is seen in Fig. 1b, where Lafayette and NWA 817 both have veining fractions of ~10% of the olivine grains compared to the much lower fractions seen in the other nakhlites. The pre-terrestrial mineral assemblages are identified in our study on the basis of the close similarity between the different nakhlite samples, including the Nakhl and Lafayette veins which have been shown to be truncated by fusion crust and are thus of extra-terrestrial origin (Gooding et al., 1991; Treiman et al., 1993). However, the nakhlite meteorite finds sometimes contain terrestrial alteration. For instance, calcite was also observed in four of the nakhlite finds, NWA 998, NWA 5790, NWA 817 (Fig. 2c, Table 1), and Y 000749, for which evidence of an extra-terrestrial origin is lacking. Calcite veins are typical of terrestrial desert weathering. For those reasons, any fracture-filling deposits with a calcite composition, similar to that seen in Table 1, are assumed to be terrestrial in origin. The sulphate mineral jarosite has been identified cutting though fusion crust in Y 000749 and is thus of terrestrial origin (Changela and Bridges, 2010).

Most of the pre-terrestrial veins within nakhlites have a smooth, featureless appearance under BSE imaging, with no obvious crystalline structure. In contrast, the fractures within the olivine grains of Lafayette that contain crystalline material have a fibrous texture when imaged with BSE. The olivine fractures often contain carbonate blocks, which have a lighter BSE tone and flat but slightly rough texture and well-defined edges, determined from analysis.
as Ca-siderite (see Table 1), and also a typically central feature of amorphous (or poorly crystalline) silicate gel similar to that seen in the other nakhlites (Changela and Bridges, 2010). The crystallinity of the fracture fills is considered below on the basis of our transmission electron microscopy results. Within Lafayette crystalline phyllosilicates are also present in the mesostasis, but with lesser amounts of silicate gel compared to the olivine veins’ assemblage. Fl-Cl-apatite is also present in the mesostasis veining (see Table 1).

Brittle fractures in the Lafayette olivine, often with sawtooth edges to the vein, usually have at least minor deposits of siderite on the walls of the vein, followed by crystalline phyllosilicates which have replaced the siderite and grown inwards toward the centre of the vein, where the central amorphous silicate material is present (see Fig. 2a). In other places, the hydrothermal minerals appear to have exploited and filled a crack at the margins of two mineral grains, sometimes with pyroxene forming part of the cavity wall. In such sites, phyllosilicates are also observed to vein and replace Ca-siderite (Fig. 2c). The amorphous silicate gel is not usually present within these deposits.

A third, less common type is fracture-filled veins within the mesostasis (Fig. 2d), and Fig. 2b shows a rare occurrence of a mesostasis vein connecting with an olivine vein. Under HRTEM imaging the mesostasis alteration phases in Lafayette are dominated by crystalline phyllosilicates, and appear crystalline in BSE imaging. Amorphous gel has only occasionally been seen associated with serpentine veins, such as that seen Fig. 2b. HRTEM imaging of the mesostasis veining from NWA 817 was found to be predominantly amorphous with a very minor presence of weak crystallinity. Mesostasis veining has also been found in the nakhlites NWA 998, Y 000593, MIL 03346.

### 3.2. Phyllosilicate and gel compositions

We obtained at least one EPMA measurement of phyllosilicate or gel per nakhlite sample (except NWA 5790 due to the absence of any veining material in our sample). Representative single analyses and the average composition for the gel and phyllosilicate deposits of each nakhlite, measured by EPMA, are shown in Table 2. The FeO and Fe₂O₃ values for each nakhlite, presented in Table 2, have been determined based on the Fe-K XANES derived Fe³⁺/ΣFe values for each nakhlite, shown in Table 4 (an average Fe³⁺/ΣFe value was taken where a nakhlite has a range of values).

Plotting Fe versus Si (wt%) for the fracture filling deposits (Fig. 3a), reveals distinct groupings for each nakhlite and negative linear patterns. This trend has also been found in previous similar studies (Gooding et al., 1991; Treiman et al., 1993; Changela and Bridges, 2010). Fig. 3b shows Mg# = 100 Mg/(Mg + Fe) plotted against Fe/Si (wt%) ratio. This also shows the nakhlite cumulate pile stacking with a decreasing average Mg# up the pile from Lafayette and NWA 998 to MIL 03346 and Y 000593/Y 000749. These correlations do not extend to the primary mineral compositions, which do not show compositional fractionation trends between the nakhlites (e.g. Bridges and Warren, 2006). The higher Na+K contents in MIL 03346 (Fig. 3c) may also reflect its position near the furthest point from the fluid source, as the most soluble elements precipitated. The Lafayette Ca contents (Fig. 3d) are highest 0.5–2 wt% and the MIL 03346 lowest suggesting that the Ca precipitated early from the fluid.

Some of the veins, notably MIL 03346, contain relatively high S contents 0.5–4.5 wt%, equivalent to up to 10.7 wt% SO₃. The distribution of S is patchy with some veins (Table 2) having low contents, similar to the other nakhlites. It is known that Y 000749 (Changela and Bridges, 2010) and MIL 03346 (Hallis and Taylor, 2011) contain terrestrial sulphates and so it is likely that terrestrial alteration has exploited some of the martian veining in an analogous way to the calcite veining in the NWA meteorites (Fig. 3f).

Fig. 3g shows Al₂O₃ and MgO contents compared between the veining material in olivine and mesostasis fractures. The range of MgO contents of the olivine phyllosilicate and gel extend to higher values (13 wt%) than the mesostasis veins. However, the clearest difference is in Al₂O₃, with the mesostasis veining having higher contents than the olivine veining. The Al₂O₃ content of the Lafayette serpentine is 5.8% and for the saponite 4.3% (or 3.2% in the gel). A higher Al content in the serpentine, compared to the saponite, can also be seen in Fig. 3e. A similar relationship is also be seen in Fig. 3g for the mesostasis veins of three other nakhlites, Y 000593, MIL 03346, and NWA 817, with
Al$_2$O$_3$ content higher in the mesostasis fractures compared to very low (<1.0 wt%) Al$_2$O$_3$ contents in the olivine fractures. However, the MgO content shows little variation between the silicate deposits in the olivine fractures and the mesostasis fractures. Such relatively small variations in Al$_2$O$_3$ and MgO in the fracture filling deposits of olivine and mesostasis suggest large scale fluid circulation and mixing did not occur. The parental fluid of the nakhlite secondary minerals is considered in more detail in Bridges and Schwenzer (2012).

The Lafayette phyllosilicate (saponite and serpentine, $n = 25$) have an average of 12–13 wt% H$_2$O. (Table 2). The gel has average ranging from 12 wt% (Lafayette, $n = 24$), 14% (NWA 817, $n = 15$), 15% (Nakhla, $n = 6$), 15% (MIL 03346, $n = 23$) and 18 wt% H$_2$O (GV, $n = 3$).

3.3. Transmission electron microscopy

3.3.1. HRTEM

A total of thirteen TEM sections of hydrothermal deposits have been extracted using the FIB-SEM (Fig. 4), including four from Lafayette olivine fractures and two from the mesostasis fractures, one from a MIL 03346 olivine fracture, one from Nakhla, three from NWA 817 olivine fractures and another from a NWA 817 mesostasis fracture.
enhances the collapsed dehydrated state. The vacuum setup of the TEM also

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<tr>
<th>Lafayette</th>
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<th>NWA 817 terrestrial</th>
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<td>calcite</td>
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Table 1 Compositions of Lafayette apatite and siderite and NWA 817 terrestrial calcite.

All data measured using SEM, carbonate CO₂ contents calculated by difference of probe analysis from 100 wt% oxide. See Bridges and Grady (2000) and Bridges and Schwenzer (2012) for more nakhlite carbonate analyses.

HRTEM observations of the veining phyllosilicates found amongst the mesostasis of Lafayette (Fig. 5b) have a different lattice fringe structure to those of phyllosilicates surrounded by olivine. The d₀₀₁-spacings range 0.67–0.73 nm, averaging 0.70 nm. The lattice structure is also different when observed along [001] orientation, visible as repeating units of two layers. This two layer structure is a pairing of a single O cation plane with a single T plane, giving a 1:1 layered T–O phyllosilicate, characteristic of serpentine minerals (Zega et al., 2006).

The NWA 817 samples, and the central silicate gel of the Lafayette samples are predominantly non-crystalline, but some poorly developed crystallinity is present. Some crystalline lattice d₀₀₁-spacings have been observed under HRTEM, though only two or three lattice fringe spacings are visible. The minor lattice features found in NWA 817 olivine fracture samples have repeating units of three layers with d₀₀₁-spacings measurements averaging ~0.98 nm, similar to the T–O–T layers of the saponite features found in Lafayette, and also similar to the poorly crystalline material previously identified in NWA 817 by Gillet et al. (2002).

The general lack of crystalline lattice features observed in the central silicate gel regions of Lafayette olivine veins, as well as the veining deposits of the nakhlites NWA 817, MIL 03346, and Nakhla, suggest a dominantly amorphous nature. In addition to the weak crystallinity we have identified, some lattice spacings were found in Nakhla and Y 000749 by Gooding et al. (1991), Noguchi et al. (2009), Lee et al. (2013).

3.3.2. Selected area electron diffraction

The phyllosilicates of Lafayette reveal a SAED pattern of distinct rings; sometimes appearing broken (see Fig. 6b), from which it is possible to take measurements. However, for SAED observations across the silicate gel of each sample, including the samples MIL 03346, NWA 817, and Nakhla, as well as the central gel regions of Lafayette olivine fractures, the pattern becomes a very diffuse ring with no distinct features from which to take a measurement, consistent with the absence of lattice spacings. The diffuse ring pattern confirms the predominantly amorphous nature of the silicate gel, whereas the ring patterns observed across the Lafayette phyllosilicates, prepared and analysed in the same way, implies polycrystallinity (Williams and Barry Carter, 1996).

The phyllosilicates of Lafayette olivine fracture deposits were measured to be an average of 4.3 Å, 3.2 Å, and 2.1–2.5 Å, from at least four measured observations, with a sum total of 9.6–10.0 Å across the three layers of the 2:1 T–O–T structure. In the mesostasis fracture deposits measured an average of 4.3 Å and 2.4 Å, from five measurements, with a sum of 6.7 Å across the two layers of the 1:1 T–O structure, and at least two measurements averaging 3.4–3.5 Å.

3.4. Fe³⁺/ΣFe determination

3.4.1. The calibration of Fe-K XANES in Fe-silicates

The 1s → 3d pre absorption edge centroid positions ferric-ferrous (Fe³⁺/ΣFe) ratios, obtained on the reference
Table 2
Compositions of saponite and serpentine phyllosilicates in Lafayette, and amorphous vein deposits across the nakhlites.

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<th>Lafayette</th>
<th>Lafayette</th>
<th>Lafayette</th>
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<td>26.0</td>
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<td>30.5</td>
<td>26.1</td>
<td>28.0</td>
<td>13.4</td>
</tr>
<tr>
<td>MgO (mole%)</td>
<td>11.1</td>
<td>10.1</td>
<td>9.6</td>
<td>8.0</td>
<td>11.3</td>
<td>7.5</td>
<td>6.9</td>
<td>2.5</td>
<td>4.8</td>
<td>3.1</td>
</tr>
<tr>
<td>MnO (mole%)</td>
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<td>1.1</td>
<td>0.5</td>
<td>0.5</td>
<td>0.6</td>
<td>0.6</td>
<td>0.9</td>
<td>1.0</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>CaO (mole%)</td>
<td>1.1</td>
<td>1.3</td>
<td>3.2</td>
<td>3.2</td>
<td>0.9</td>
<td>0.5</td>
<td>0.7</td>
<td>0.3</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Na₂O (mole%)</td>
<td>0.1</td>
<td>0.2</td>
<td>0.2</td>
<td>0.3</td>
<td>0.2</td>
<td>0.6</td>
<td>0.9</td>
<td>0.2</td>
<td>0.2</td>
<td>tr.</td>
</tr>
<tr>
<td>K₂O (mole%)</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>0.3</td>
<td>0.4</td>
<td>0.4</td>
<td>0.3</td>
<td>1.1</td>
<td>tr.</td>
</tr>
<tr>
<td>P₂O₅ (mole%)</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
</tr>
<tr>
<td>SO₃ (mole%)</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.4</td>
<td>1.9</td>
<td>3.4</td>
<td>0.5</td>
</tr>
<tr>
<td>Cl (mole%)</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>tr.</td>
<td>0.1</td>
<td>0.3</td>
<td>0.7</td>
<td>tr.</td>
<td>0.1</td>
<td>0.2</td>
</tr>
<tr>
<td>Total (mole%)</td>
<td>88.7</td>
<td>87.5</td>
<td>86.9</td>
<td>88.2</td>
<td>88.9</td>
<td>81.1</td>
<td>83.0</td>
<td>85.6</td>
<td>86.8</td>
<td>85.2</td>
</tr>
<tr>
<td>H₂O (mole%)</td>
<td>11.3</td>
<td>12.5</td>
<td>13.1</td>
<td>11.7</td>
<td>11.1</td>
<td>18.9</td>
<td>17.0</td>
<td>14.4</td>
<td>13.2</td>
<td>14.8</td>
</tr>
</tbody>
</table>

3.4.2. Fe-K XANES in the nakhlites

Based on this linear calibration (Fig. 7b), the Fe³⁺/ΣFe values for a different Y 000749 nakhlite olivine grain and a nontronite standard sample, Urgeirica Beira Alta, Portugal-MB 1972, 142 (Nontronite #142), were estimated to be Fe³⁺/ΣFe = 0.03 and Fe³⁺/ΣFe = 0.94 respectively. Thus using this method show, as expected, the Y 000749 olivine to be ferrous and the nontronite #142 to be ferric-rich.

All nakhlite samples with pre-terrestrial fracture filled veins were measured for Fe-K XANES. Based on the correlation between Fe-K XANES 1s→3d centroid position with Fe³⁺/ΣFe values, the oxidation state of the hydrothermal deposits across the nakhlite samples were determined. Fig. 8 shows at least one spectrum, over the pre-edge energy ranges, from the vein deposits of each nakhlite, including spectra for the amorphous and crystalline features of Lafayette, and also spectra from olivine and mesostasis fractures in Lafayette and NWA 817. This figure also shows a representative complete Fe K XANES spectrum. These deposits were found to be mostly ferric, with variable ratios of Fe³⁺/ΣFe values, and are stacked in Fig. 8 from top to bottom in order of estimated Fe³⁺/ΣFe, ranging from Fe³⁺/ΣFe = 0.3 in MIL 03346 to the highly ferric Fe³⁺/ΣFe = 0.9 in the olivine-filling gels of Lafayette (see Table 4 for the values). The serpentine mesostasis fracture-fills in Lafayette have an even higher ferric content.
A greater ferric content in the mesostasis veins compared to silicate veining in the olivine fractures is also seen in NWA 817.

The Lafayette veins become more ferric into their centre, from Fe$^{3+}$/ΣFe values of ~0.6 in the phyllosilicates increasing to a Fe$^{3+}$/ΣFe value of ~0.9 in the silicate gel. This increase in ferric content from the vein edges into the centre can also be seen in Fig. 9a, in which a map of normalised Fe-K XANES edge energy show a transition from dark to light representing the Fe$^{2+}$-rich to Fe$^{3+}$-rich compositions. Changela and Bridges (2010) had previously suggested, based on the Fe-rich content of the Lafayette

### Table 3

<table>
<thead>
<tr>
<th>Standard Sample</th>
<th>Fe Sample Type</th>
<th>1s → 3d Position (±0.05 eV)</th>
<th>Fe$^{3+}$/ΣFe (At%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y 000749 Olivine$^T$</td>
<td>Silicate</td>
<td>7111.3</td>
<td>0.0</td>
</tr>
<tr>
<td>Barwell L6 Olivine$^F$</td>
<td>Silicate</td>
<td>7111.9</td>
<td>0.0</td>
</tr>
<tr>
<td>San Carlos Olivine$^F$</td>
<td>Silicate</td>
<td>7111.9</td>
<td>0.0</td>
</tr>
<tr>
<td>San Carlos Olivine$^F$</td>
<td>Silicate</td>
<td>7111.7</td>
<td>0.0</td>
</tr>
<tr>
<td>Siderite$^F$</td>
<td>Carbonate</td>
<td>7112.5</td>
<td>0.0</td>
</tr>
<tr>
<td>NWA 817 Pyroxene$^F$</td>
<td>Silicate</td>
<td>7112.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Chromite$^F$</td>
<td>Oxide</td>
<td>7112.7</td>
<td>0.2</td>
</tr>
<tr>
<td>Peridotite JP-1$^T$</td>
<td>Igneous</td>
<td>7112.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Magnetite$^T$</td>
<td>Oxide</td>
<td>7113.3</td>
<td>0.7</td>
</tr>
<tr>
<td>Illite IMt-1$^T$</td>
<td>Silicate</td>
<td>7113.5</td>
<td>0.9</td>
</tr>
<tr>
<td>Nontronite NG-1$^T$</td>
<td>Silicate</td>
<td>7113.3</td>
<td>1.0</td>
</tr>
<tr>
<td>Nontronite NAu-2$^T$</td>
<td>Silicate</td>
<td>7113.4</td>
<td>1.0</td>
</tr>
<tr>
<td>Goethite$^T$</td>
<td>Oxy-hydroxide</td>
<td>7114.3</td>
<td>1.0</td>
</tr>
<tr>
<td>Hematite$^F$</td>
<td>Oxide</td>
<td>7114.2</td>
<td>1.0</td>
</tr>
<tr>
<td>Hematite$^F$</td>
<td>Oxide</td>
<td>7114.2</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Fe-K XANES data was measured in transmission ($^T$) or fluorescence ($^F$). Fe$^{3+}$/ΣFe values for the nontronites (NG-1 and NAu-2), illite (IMt-1), and peridotite (JP-1) were calculated from online reference material chemical compositions (Geological Survey of Japan/AIST, 1995; The Clay Minerals Society, 2013).

### Table 4

<table>
<thead>
<tr>
<th>Sample</th>
<th>Xanes 1s → 3d position (±0.05 eV)</th>
<th>Nature</th>
<th>Fe$^{3+}$/ΣFe (±0.03)</th>
<th>Fe-O bond length (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nontronite #142</td>
<td>7113.3</td>
<td>Fe$^{2+}$ silicate</td>
<td>0.9</td>
<td>2.00</td>
</tr>
<tr>
<td>Y 000749 Olivine</td>
<td>7111.8</td>
<td>Fe$^{2+}$ silicate</td>
<td>0.0</td>
<td>2.13</td>
</tr>
<tr>
<td>NWA 998</td>
<td>7112.7</td>
<td>Ol. Gel</td>
<td>0.6</td>
<td>2.03</td>
</tr>
<tr>
<td>Lafayette</td>
<td>7112.8</td>
<td>Saponite</td>
<td>0.6</td>
<td>2.05</td>
</tr>
<tr>
<td>Lafayette</td>
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<td>Saponite</td>
<td>0.6</td>
<td>2.04</td>
</tr>
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<td>Ol. Gel</td>
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<td>1.95</td>
</tr>
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<td>Ol. Gel</td>
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<td>2.05</td>
</tr>
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<td>Ol. Gel</td>
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<td>2.05</td>
</tr>
<tr>
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<td>7113.2</td>
<td>Ol. Gel</td>
<td>0.8</td>
<td>2.04</td>
</tr>
<tr>
<td>Lafayette</td>
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<td>Ol. Gel</td>
<td>0.7</td>
<td>2.03</td>
</tr>
<tr>
<td>Lafayette</td>
<td>7113.4</td>
<td>Serpentine</td>
<td>1.0</td>
<td>1.99</td>
</tr>
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<td>Serpentine</td>
<td>1.0</td>
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<td>Lafayette</td>
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<td>Serpentine</td>
<td>1.0</td>
<td>1.99</td>
</tr>
<tr>
<td>Lafayette</td>
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<td>Serpentine</td>
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<tr>
<td>Nakhla</td>
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<td>Ol. Gel</td>
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<td>–</td>
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<td>Ol. Gel</td>
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<td>2.10</td>
</tr>
<tr>
<td>NWA 817</td>
<td>7113.0</td>
<td>Meso. Gel</td>
<td>0.7</td>
<td>2.03</td>
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<td>NWA 817</td>
<td>7113.1</td>
<td>Meso. Gel</td>
<td>0.8</td>
<td>2.08</td>
</tr>
</tbody>
</table>

Nakhite measurements are of fracture filling deposits within olivine (Ol.) and mesostasis (Meso.). Fe–O bond lengths measured with EXAFS.

with Fe$^{3+}$/ΣFe = 1.0. A greater ferric content in the mesostasis veins compared to silicate veining in the olivine fractures is also seen in NWA 817.

Table 4

Fe-K XANES measurements (eV), Fe$^{3+}$/ΣFe ratios and Fe–O bond lengths.

Fe$^{3+}$/ΣFe values are of fracture filling deposits within olivine (Ol.) and mesostasis (Meso.). Fe–O bond lengths measured with EXAFS.
phylllosilicate composition, that the closest smectite was the saponite, griffithite, which has an unusually high abundance of $\text{Fe}^{3+}$ and would fit our $\text{Fe}^{3+}/\text{R}^{3+}$ results. The saponite in the Lafayette veins has similar $\text{Fe}^{3+}/\text{R}^{3+}$ values to terrestrial iron-rich saponite such as iron-rich sand beds and rhyolitic glassy Oya tuffs from Japan, which have $\Sigma \text{Fe}$ values ranging 0.73–0.95 (Sudo, 1954; Kohyama et al., 1972; Mackenzie and Berezowski, 1980); and saponites from

Fig. 3. EPMA-WDS analysis of the gel and phyllosilicates in the nakhlites. (a) Fe versus Si wt%. (b) Mg# = Mg/(Mg + $\Sigma \text{Fe}$ at.) versus Fe/Si wt%. (c) (Na + K) versus Si wt%. (d) Ca versus Si wt%. (e) Al versus Si wt%. (f) S versus Si wt%. (g) $\text{Al}_2\text{O}_3$ and $\text{MgO}$ compositions of phyllosilicate and gel within olivine and mesostasis. All nakhlite compositions are normalized to 100% anhydrous totals, and measured using EPMA and/or SEM-EDX. A previous Lafayette phyllosilicate and other nakhlite vein measurements are from: (1)Treiman et al. (1993); and (2)Changela and Bridges (2010).
American Canyon, California, with Fe\(^{3+}/\Sigma\)Fe = 1.00 (Abdel-Kader et al., 1978; Post, 1984).

Previous studies of Fe-K XANES have found similar trends, with pre-edge 1s → 3d centroid positions correlating with the Fe\(^{3+}\) content of Fe minerals (Waychunas et al., 1983; Wilke et al., 2001; Berry et al., 2003; Cottrell et al., 2009). Due to a lack of silicate standards of known intermediate Fe\(^{3+}/\Sigma\)Fe values, there was potential difficulty in characterising any nakhlite fracture deposits measured with 1s → 3d centroid positions within the energy region of 7111.9 eV to 7113.5 eV. However, the study by Berry et al. (2003) of silicate glasses with varying ferric content (determined by Mössbauer spectroscopy) does cover this Fe\(^{3+}/\Sigma\)Fe gap and so we were able to make comparisons between measurements of this earlier study and our own. By comparing the Fe-oxides of our study to those measured by Berry et al. (2003), including magnetite and hematite at 7113.6 eV and 7114.6 eV respectively, there is a minor negative shift of 0.3–0.4 eV required to bring the Berry et al. (2003) silicate glass measurements in-line with our silicate standards. A line of best-fit between the highly ferrous and the highly ferric silicate standards of our study, which was used for the linear calibration, closely resembles that of the linear trend between the 1s → 3d centroids and the
Fe$_{3+}$/R$_{Fe}$ values found by Berry et al. (2003). This comparison can be seen in Fig. 7b. We use the calibration to calculate the relative abundances of ferrous and ferric iron in the nakhlites samples.

### 3.4.3. EXAFS

EXAFS spectra have been obtained for reference materials and the nakhlites as a complementary way to Fe-K XANES of checking the ferric nature of the nakhlites.
The reference materials included the highly ferric nontronite #142 with $\text{Fe}^{3+}/\text{Fe} = 0.94$, similar in that respect to the nakhlite phyllosilicates and gel. Our analyses are able to constrain the Fe–O bond lengths, for example the ferric nontronite #142 reference sample was determined to have six Fe–O bond lengths of 2.00 Å. Due to iron atoms not being situated directly at the centre of an atomic octahedron structure, the Fe–O bond lengths are variable between $\text{Fe}^{3+}–\text{O}$ and $\text{Fe}^{2+}–\text{O}$ (Al-Hasni and Mountjoy, 2011) and can often be observed in EXCURV analysis as split peaks. In contrast to the ferric nontronite #142, the ferrous Y 000749 olivine was also determined to have six Fe–O bonds, but at lengths of 2.13 Å instead of the much shorter 2.00 Å Fe–O bond lengths for nontronite #142. These results are consistent with previous EXAFS studies for nontronite with Fe–O bond lengths of 1.97–2.04 Å (Manceau et al., 1998) and for olivine with Fe–O bond lengths of 2.08–2.15 Å (Redfern et al., 2000).

Fig. 10 shows how the data is used with EXCURV, with the $k^2$-weighted EXAFS spectrum (a and c), typically between ~7140 eV and ~7420 eV, and the Fourier-filtered EXAFS spectrum (b and d) of that energy region. The dashed curve (Theory) attempts to fit the solid curve of the “real” data (Experiment) based on approximate values of bond-type, bond-length, and bond-number, of near-neighbour atoms. In Fig. 10 the Fourier-filtering of the EXAFS spectrum reveals at least two prominent peaks representing the Fe–O bond (left peak) and the Fe–Si bond (right peak).

Fig. 9 (a) Fe-K XANES map of a Lafayette fracture fill vein in olivine. The map shows normalized intensity measured at 7120.0 eV. This Fe-K XANES map of normalized spectra intensity highlights the Fe$^{3+}$ (white) against the Fe$^{2+}$ (black) content of the hydrothermal vein. (b) The same Lafayette fracture fill hydrothermal vein with BSE imaging, with olivine (Ol), siderite (Sid), saponite (Sap) and the central silicate gel. A bright discontinuous patch of Fe oxides is present between the gel and phyllosilicate.
an average between the two bond lengths must be calculated, based on the number of atoms for each bond length. Therefore, the total number of Fe–O bonds averages a distance of 2.03 Å. There are also Fe–Si bonds measuring 3 at 3.23 Å and 3 at 3.42 Å, averaging 6 Fe–Si bonds at a distance of 3.33 Å.

The majority of the results from EXCURV analyses, like those of the Lafayette and NWA 817 veins seen in Fig. 10, have Fe–O bond lengths ranging between those of the ferric nontronite #142 and the ferrous Y 000749 olivine, and averaging at ~2.03 Å. A full list of the Fe–O bond length results determined by the EXAFS can be seen in Table 4. Fig. 11 compares the Fe–O bond length results to the Fe$^{3+}$/ΣFe values, determined via the ferric-ferrous calibration using Fe-K XANES measurements. This plot shows a trend in the nakhlite veining material between the Y 000749 olivine and the nontronite #142 samples, consistent with a negative correlation between Fe–O bond lengths and the Fe$^{3+}$/ΣFe ratio of that material, with ferric material having Fe–O bond lengths of ~2.00 Å.

4. DISCUSSION

4.1. The nakhlite hydrothermal fluid and depth of formation

Assuming that all of the nakhlites likely formed within a single cumulus pile, the variations between the different samples reveal likely burial depths. Mikouchi et al. (2003, 2012) compared nine nakhlite samples, observing variations in features such as the abundance of the mesostasis, olivine and pyroxene minerals, the Fe content of pyroxene rim zones, olivine Ca content and Mg-Fe content, from which the cooling rate for each nakhlite was deduced, and thus a burial depth estimated. The nakhlite cumulate pile had MIL 090030/032/136 and NWA 5790 near the top, just 1–2 m from the surface, ranging down in order through NWA 817, MIL 03346, Y 000749, Y 000593, GV, and Nakhla, to the bottom with Lafayette and NWA 998 at a burial depth of >30 m. The plot of Mg# against the Fe/Si ratio for the phyllosilicate and gel, (Fig. 3b), reveals a similar order of stacking from MIL 03346 with low Mg# and Fe/Si ratio. The only significant exception to this stacking order, as seen in Fig. 3b, is the plotting of NWA 817 between Y 000749 and Nakhla. The higher abundance of Ca in Lafayette gel and phyllosilicate relative to higher nakhlites suggests that the Ca precipitated early from the fluid, and this is consistent with the siderite composition which is most Ca-rich in Lafayette (Bridges and Grady, 2000). This correlation between the composition of the secondary silicates and the original depth of the surrounding nakhlite, was also found by (Changela and Bridges, 2010), but here with the addition of MIL 03346, NWA 817. NWA 998 is assumed to be laterally displaced from the fluid source, to explain the relatively deep origin in the nakhlite pile but low abundance of alteration veining.

Changela and Bridges (2010) proposed a model in which the secondary assemblages formed in an impact-induced hydrothermal system terminated by the precipitation of the silicate gel and evaporation of soluble salts. Lafayette was closest to the impact heat and fluid (buried H$_2$O–CO$_2$.
ice) source. Bridges and Schwenzer (2012) showed that the alteration assemblages described by both Changela and Bridges (2010) and here, were the result of selective dissolution of 20% Lafayette bulk, 10% mesostasis, 70% olivine with an additional albite component, together with 0.1 mol CO₂ derived from the ice. After brittle fracturing in the olivines and mesostasis of Lafayette, the fluid precipitated Ca,Fe-rich siderite at 150–200 °C followed in Lafayette by the crystalline saponite at 50 °C, then the poorly crystalline gel. The fluid varied from pH 6–8 with a water-rock ratio (W/R) ≤ 300, becoming more alkaline, pH 9 and W/R of 6 as the saponite crystallised. In addition, the increasingly ferric composition we identify from the Lafay-
ette saponite to the gel in the centre of the veins shows an oxidation trend.

Following this work, Lee et al. (2013) also noted that the likely origin for the fractures in augite and olivine is impact shock, suggesting that the secondary fluids preferentially exploited crystal faces parallel to [001] in these fractures. However, the textures we have described here and in previous work within the olivine and mesostasis are fracture fills, rather than the dominantly in situ alteration of olivine grains around the fractures as suggested by Lee et al. (2013). Such a scenario could not account for the layering within the fractures from siderite, replaced by saponite, followed by minor oxide and the gel which reflects the evolution of a single fluid. The composition of the alteration minerals and high-Ca saponite within Lafayette (Bridges et al., 2001) are also impossible to explain through isochemical alteration of adjacent low-Ca olivine. Although there is firm evidence for selective dissolution of olivine both texturally (Changela and Bridges, 2010; Lee et al., 2013), and from fluid modelling (Treiman and Lindstrom, 1997; Bridges and Schwenzer, 2012) within the nakhliites as a whole, the majority of fractures which contain secondary minerals (e.g. Figs. 2 and 8) do not show signs of extensive corrosion along the olivine grain boundaries. None the less an association between siderite and olivine is clear, though it is not exclusive as siderite is also sometimes present within the mesostasis of the nakhliites (Bridges and Grady, 2000). The olivine fractures may simply have provided the surfaces for the first hydrothermal minerals (C siderite) to crystallise upon.

The Fe-rich serpentine in the mesostasis of Lafayette identified here and in Changela and Bridges (2010) is notably more Al-rich than the saponite. Although the nakhlite hydrothermal fluid was derived from a mixture of nakhlite minerals (Bridges and Schwenzer, 2012) the association between feldspathic mesostasis and the Al-rich serpentine suggests that the local microenvironment did influence this phyllosilicate composition.

Lee et al. (2013) suggested on the basis of identification of some lattice spacings and SAED patterns that all of the gel in the centre of veins was 'nanocrystalline'. We suggest on the basis of the TEM studies here and previously (Changela and Bridges, 2010) that most of the material in the olivine fractures is poorly crystalline rather than having a homogenous crystalline structure, because crystallinity is only evident in small patches of nanometre-scale lattice spacings. Gooding et al. (1991) also suggested for Nakhl, the majority of the veining material in the nakhliites is poorly crystalline. This predominantly amorphous nature is a function of the rapid cooling of the fluid in its latter stages. The juxtaposition, at the micron scale, of amorphous material and crystalline smectite of similar composition has frequently been observed in terrestrial clay-bearing rocks e.g. Gaudin et al. (2005).

As ever with NWA and MIL desert finds the effects of terrestrial alteration need to be considered. The most obvious demonstration of this in the samples we have analysed here is the presence of high S enrichments (and see Hallis and Taylor, 2011) and calcite which have in places exploited the pre-existing martian veins.

### 4.2. Phyllosilicates on the martian surface

The CRISM Near InfraRed mapping spectrometer on Mars Reconnaissance Orbiter has been used to identify phyllosilicates similar to those we have identified in the nakhliites – serpentine, smectites (including montmorillonite) – in complex impact craters (Bibring et al., 2006; Mustard et al., 2008). This includes CRISM observations of deep Noachian craters with central peaks exposing Fe/Mg phyllosilicate-rich rocks at depths of 4–5 km. Schwenzer and Kring (2009) proposed that such secondary mineral assemblages were a result of the impacts creating hydrothermal systems. Although in most identified cases martian phyllosilicate deposits occur in terrain of Noachian age >3.5 Gyr, in contrast to the ≤670 Ma age of the nakhlite phyllosilicates (Swindle et al., 2000), there is growing evidence for later hydrothermal alteration associated with impact cratering (e.g. Carter et al., 2013). In contrast, rather than resulting from impact-induced hydrothermal activity, Carter et al. (2013) proposed that the secondary assemblages in Amazonian terrains were a result of the excavation by the impacts of older, pre-existing, phyllosilicate-bearing surfaces. The nakhliites' secondary mineral assemblage is strong evidence for impact-induced hydrothermal alteration initiated by fracturing (Bridges et al., 2001; Changela and...
Bridges, 2010; Bridges and Schwenzer, 2012). However, the rapidly cooled, metastable nature (e.g. as shown by the presence of poorly crystalline veins and the carbonate compositions) of the nakhlites’ secondary assemblage suggest that it was not part of a large, convecting hydrothermal system. For instance, the models in Abramov and Kring (2005) showed that impact craters of diameter 30–180 km produce convecting hydrothermal systems lasting for 70,000–380,000 years. In ≤7 km simple craters the lack of a central uplift and melt means that shock-emplaced heat dominates, with a low possibility of a convecting hydrothermal system developing (Abramov and Kring, 2005). The nature of the impact event associated with the nakhlite alteration is not yet clear but might be at the margins of a large impact crater or through a short-lived hydrous alteration event associated with a smaller, simple crater in an icy terrain. There is some evidence that smaller, simple craters can experience limited hydrous alteration. For instance, Osinski et al. (2013) described montmorillonite alteration in the 1.8 km diameter Lunar Crater of the Deccan traps. Schwenzer et al. (2012) suggested that craters as small as 5 km on Mars delivered sufficient heat to create liquid water.

5. CONCLUSIONS

(1). In Lafayette, the phyllosilicate filling the fractures of olivine grains is saponite, shown with HRTEM to be a 2:1 T–O–T lattice structure in a collapsed dehydrated state with d001-spacings of 9.6 Å.

(2). The fractures within the Lafayette mesostasis consist of crystalline phyllosilicates with a 1:1 lattice structure, and a d-spacing of 7.0 Å, confirming the presence of serpentine, together with lesser amounts of amorphous gel.

(3). There is a definite presence of crystalline phyllosilicates in the olivine fractures (typically with amorphous silicate gel in the centre of the veins) and mesostasis fractures of Lafayette, and minor, patchy crystallinity is found within the silicate gel, of saponite-like composition, within NWA 817 and Nakhla. The hydrothermal deposits, present throughout all of the nakhlite martian meteorites, except NWA 5790, are predominantly amorphous silicate of ferric saponite composition.

(4). An increasing shift in the measured Fe-K XANES 1s → 3d centroid positions of Fe-silicate standards, from 7111.3 eV to 7113.5 eV, coincides with an increased oxidation state from pure ferrous to pure ferric. This confirms the correlation between 1s → 3d centroid positions and ferric-ferrous (Fe³⁺/ΣFe) ratios, as found in previous studies (Waychunas et al., 1983; Wilke et al., 2001; Berry et al., 2003; Cottrell et al., 2009), allowing us to estimate the ferric content of the nakhlite Fe-silicates minerals.

(5). The crystalline saponite phyllosilicate of Lafayette has Fe³⁺/ΣFe values of ~0.6, compared to the highly ferric silicate gel at the centre of the veins which has Fe³⁺/ΣFe values of up to ~0.9. The Al-rich serpentine in Lafayette and NWA 817 was found to be highly ferric, more so than that of the saponite in those nakhlite samples, reaching Fe³⁺/ΣFe ~1.0. Fe–O bond lengths, determined by EXAFS, are 2.03 Å, consistent with ferric-rich compositions.

(6). The average chemical formula of the saponite minerals within the Lafayette olivine fractures is a stoichiometric, trioctahedral ferric saponite (Ca₀.₃Mg₁.₇FeIVO₄(OH)₆H₂O) (OH)₄·nH₂O. The Fe-rich serpentine mineral within the mesostasis fractures of Lafayette is (Ca₀.₁Mg₀.₉FeIVO₄Al₂O₅(Fe₄Al₈O₂₀(OH)₆·nH₂O. The overall correlation between nakhlite burial depth and the Mg# composition of the gel suggested previously is confirmed with more nakhlite samples and analyses. However, NWA 817 which has a high vol% of veining (10%, similar to Lafayette) and is expected to have been buried to only a few metres depth has a relatively high Mg# showing that there was more than one fluid pathway from the fluid source close to Lafayette.

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