Aqueous alteration of the Martian meteorite Northwest Africa 817: Probing fluid–rock interaction at the nakhlite launch site

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Abstract–The nakhlite meteorites characteristically contain iddingsite, a hydrous iron–magnesium silicate that formed by aqueous alteration on Mars. Iddingsite is most abundant in Northwest Africa (NWA) 817, and alteration products in this meteorite also have the lowest deuterium/hydrogen ratio of any nakhlite. Taken together, these distinctive properties could be interpreted to show that NWA 817 was altered under different physico-chemical conditions than the other nakhlites and by liquid water from a separate reservoir. Here this interpretation is tested through a petrographic, mineralogical, chemical, and isotopic study of NWA 817. We find that its iddingsite occurs as olivine-hosted veins of nanocrystalline smectite and Fe-oxyhydroxide. Strong similarities in the mineralogy of iddingsite between NWA 817 and other nakhlites suggest that these meteorites were altered under comparable physico-chemical conditions, with the Fe-rich composition of NWA 817 olivine grains rendering them especially susceptible to aqueous alteration. Analyses of NWA 817 bulk samples by stepwise pyrolysis confirm that its iddingsite has unusually low deuterium/hydrogen ratios, but owing to terrestrial weathering of this meteorite, the hydrogen isotopic data cannot be used with confidence to infer the origin of Martian aqueous solutions. NWA 817 was most probably altered along with the other nakhlites over a short time period and in a common aqueous system. One interpretation of a correlation between the eruption ages of three of the nakhlites and the chemical composition of their iddingsite is that water originated from close to the surface of Mars and flowed through the nakhlite lava pile under the influence of gravity.

INTRODUCTION

Meteorites of the nakhlite group sample the mid-Amazonian igneous crust of Mars (e.g., Bogard and Johnson 1983; McSween 1994; Nyquist et al. 2001). Strong similarities between them in bulk chemical composition, mineralogy, and cosmic ray exposure age indicates that these meteorites were impact-ejected simultaneously, and from the same place on Mars’s surface (e.g., Treiman 2005). The launch site has been suggested to contain a single thick lava flow/intrusion (e.g., Friedman-Lentz et al. 1999; Mikouchi et al. 2006, 2012; Corrigan et al. 2015), or multiple lavas (Jambon et al. 2016). Recent 40Ar/39Ar dating of six meteorites indicates that they were sourced from at least four lava flows that formed over a time span of ~93 Ma (i.e., between 1416 ± 7 Ma and 1322 ± 10 Ma; Cohen et al. 2017).

The nakhlites are composed principally of augite and olivine phenocrysts, between which is a finely crystalline (rarely glassy) mesostasis (Bunch and Reid 1975). Most of them also contain a very fine-grained material called iddingsite (Ashworth and Hutchison 1975) (Fig. 1), whose constituents include hydrous
silicates (smectite, serpentine, opal-A, amorphous gel, stilpnomelane), Fe-oxides, and Fe-oxyhydroxides (Bunch and Reid 1975; Gooding et al. 1991; Treiman et al. 1993; Bridges and Grady 2000; Gillet et al. 2002; Noguchi et al. 2009; Changela and Bridges 2011; Chatzitheodoridis et al. 2014; Kuebler 2013; Lee et al. 2013, 2015a; Lee and Chatzitheodoridis 2016). The iddingsite is commonly intergrown with sidereite, gypsum, and halite (e.g., Gooding et al. 1991; Bridges and Grady 2000; Lee et al. 2013; Tomkinson et al. 2013). As iddingsite has formed from aqueous solutions, its presence in the nakhlites shows that liquid water must have been available in the crust of Mars relatively late in the planet’s history (Ashworth and Hutchison 1975).

The origin of the aqueous fluids that were responsible for alteration of the nakhlites can be constrained using the hydrogen isotopic composition of water/OH in iddingsite because various reservoirs of Martian hydrogen differ in their deuterium/hydrogen (D/H) ratio (hereafter expressed as δD [(D/H)_{sample}/(D/H)_{V-SMOW} - 1] × 1000). The planet’s atmosphere is D-rich (present-day δD is 4950 ± 1080‰; Webster et al. 2013), whereas the Martian mantle (δD ≤ 275‰; Hallis et al. 2012; Usui et al. 2012) is close in composition to terrestrial reservoirs (δD –300 to 0‰; Lécuyer et al. 1998). Usui et al. (2015) suggested that Mars has a third hydrogen reservoir within hydrated crust or ground ice with an intermediate D/H ratio (δD 1000–2000‰). The hydrogen isotopic composition of iddingsite in Governador Valadares, Lafayette, Miller Range (MIL) 00346, Nakhla, and Yamato (Y)-000593 is in excess of terrestrial values. Nakhla, the only nakhlite fall, has yielded the highest δD of 1165‰ (Leshin et al. 1996; Hallis et al. 2012). Thus, water/OH in the iddingsite is interpreted to have equilibrated with Mars’s atmosphere, but could equally have been derived from the hydrated crust/ice reservoir that was postulated by Usui et al. (2015). Iddingsite in Lafayette and Y-000593 has been dated to 633 ± 23 Ma (Borg and Drake [2005] and references therein), meaning that with regard to those two meteorites at least, the liquid water that was responsible for aqueous alteration cannot have been sourced from the same episode of igneous activity that formed their parent rocks.

Northwest Africa (NWA) 817 has relatively abundant iddingsite that formed by aqueous alteration on Mars (Gillet et al. 2002). The δD value of its iddingsite (−170 ± 14‰) has been interpreted to show that the water responsible for alteration was derived from the Martian mantle (Gillet et al. 2002; Meunier et al. 2012). Taken together, the hydrogen isotopic data therefore suggest that the nakhlite launch site experienced two episodes of aqueous alteration—one by crustal/atmospheric fluids (recorded by iddingsite in Governador Valadares, Lafayette, MIL 00346, Nakhla, and Y-000593) and the other by mantle-derived solutions (recorded by NWA 817 alteration products). Furthermore, given that NWA 817 iddingsite has not been radiometrically dated, it is possible the liquid

Fig. 1. An area of N8-1 containing augite (Aug), olivine (Ol), and mesostasis (M). a) X-ray map overlain on a BSE image. The X-ray map has been made by blending three element maps: Si Kα (red), Mg Kα (blue), and Fe Kα (green). These color combinations render augite grain cores purple, the narrow Fs-rich rims to augite grains orange, olivine grain cores blue, olivine grain rims light green, iddingsite brown, equant grains of Fe-Ti oxide lime green, and the mesostasis predominantly dark red. Both the augite and olivine grains are compositionally zoned, and the mesostasis contains many acicular Fe-rich crystals. b) X-ray map of the same field of view as (a) that was made by blending maps for Mg Kα (blue) and Ca Kα (yellow). This element combination renders olivine mid blue to dark blue, augite turquoise, calcite yellow. Open fractures are black, and the mesostasis contains a variety of fine crystals. Note that the calcite veins are almost entirely restricted to augite. (Color figure can be viewed at wileyonlinelibrary.com.)
water was sourced from the same episode of igneous activity that formed the nakhlite lavas at 1.4–1.3 Ga (Meunier et al. 2012). Here we test the idea that the nakhlites were altered by multiple fluids from different sources by comparing the petrographic, mineralogical, chemical, and isotopic properties of iddingsite in NWA 817 with other members of the group.

**MATERIALS AND METHODS**

NWA 817 was found in Morocco in December 2000 as a single 104 g stone (Grossman and Zipfel 2001), and was first described in detail by Gillet et al. (2002) and Sautter et al. (2002). This study has used three pieces of a commercially obtained sample, hereafter referred to as N8-1, N8-2, and N8-3. None of these pieces has a fusion crust. N8-1 was used for petrographic, mineralogical, and chemical analysis, whereas N8-2 and N8-3 were used for deuterium/hydrogen (D/H) measurements by stepwise pyrolysis. After the D/H work, a polished block was made from N8-3 in order to identify those components that had lost water/hydroxyls during pyrolysis.

The polished blocks made from N8-1 and N8-3 were coated with carbon and backscattered electron (BSE) images were acquired using two field-emission gun scanning electron microscopes (FEG-SEM), a Zeiss Sigma and a FEI Quanta 200F, both operated at 20 kV and under high vacuum. X-ray maps and quantitative chemical analyses were acquired using the Zeiss equipped with Oxford Instruments INCA and AZtec microanalysis systems operating through a X-Max silicon-drift energy-dispersive X-ray (EDX) detector. Mapping was undertaken at 20 kV/1–5 nA, whereas the quantitative analyses were acquired at 20 kV/1 nA with a 60 s count time. X-ray spectra were calibrated using the following mineral standards: jadeite (Na, Al), pericline (Mg) olivine (Si), apatite (P), pyrite (S), orthoclase (K), wollastonite (Ca), rutile (Ti), rhodonite (Mn), almandine garnet (Fe), and Ni-metal (Ni). The “feature mapping” function of AZtec was used to locate barite in N8-1 by automatically searching for grains falling within a predefined contrast range in BSE images. By setting this contrast band to detect only the brightest pixels (i.e., grains with a high mean atomic number, Z), grains of barite could readily be located, and their size, shape, petrographic context, and chemical composition could be determined. For electron backscatter diffraction (EBSD) work, N8-1 was polished for 4 h using 0.01 μm colloidal silica in a NaOH suspension, then coated with 5 nm of carbon. EBSD and EDX maps were acquired simultaneously from a 250 × 200 μm area of mesostasis at Oxford Instruments Nano-analysis, High Wycombe, using a Hitachi SU-70 field-emission SEM equipped with an Oxford Instruments CMOS-based Symmetry EBSD detector and X-Max 80 mm² EDX detector. The SEM was operated at 15 kV/12 nA, and mapping was undertaken at a step size of 200 nm. The data were acquired using AZtec EDX-EBSD software v. 3.4., and the electron backscatter patterns (EBSP) were collected at resolutions of 156 × 128 pixels/EBSP to allow the acquisition of 247 patterns/s. The data were processed and interpreted using Oxford Instruments HKL Channel 5 software. Noise reduction using a single “wildspike” correction was followed by an iterative 6 point nearest neighbor zero solution correction, which is routine for this type of data (Watt et al. 2006; Forman et al. 2016, 2017). Grains were identified using the Channel 5 grain detection algorithm where a single grain is defined as <1° misorientation across adjacent pixels.

Electron-transparent samples for transmission electron microscopy (TEM) were extracted from N8-1 using a FEI DuoMill dual-beam focused ion beam (FIB) instrument. Milling employed a 30 kV Ga⁺ ion beam at various currents following the protocol of Lee et al. (2003). The ~100 nm thick samples were studied initially by low-voltage scanning TEM (LV-STEM) imaging in the Quanta SEM (Lee and Smith 2006), then bright-field and high-resolution images, and selected area electron diffraction (SAED) patterns were acquired using a FEI T20 TEM operated at 200 kV. Subsequently, high-angle annular dark-field (HAADF) images and EDX analyses were obtained from the same foils using a JEOL ARM200F aberration-corrected TEM that was operated at 200 keV/~100 pA, giving a nominal spot size of <0.2 nm. The EDX analysis used a Bruker 60 mm² SDDEDX spectrometer, and spectra were quantified using proprietary software.

The D/H work on N8-2 and N8-3 used chips (77.5 and 108 mg, respectively) in preference to powders in order to minimize contamination by the terrestrial atmosphere. These samples were placed in an all-glass vacuum line and pumped down overnight. They were incrementally heated in a temperature-controlled induction heater to 200, 500, and 1050 °C (±5 °C). Samples were held at each designated temperature for a minimum of 15 min. The gases released were initially stored in a cold trap at −196 °C using a liquid nitrogen bath. This bath was then replaced by a mixture of dry ice and acetone at −78 °C, trapping water but releasing other gases. The yield of gas not trapped at −78 °C was measured using an Edwards PR10-C Pirani vacuum pressure gauge. The dry ice and acetone bath was then removed from the cold trap, which was gently heated, and the water was released. This water was reduced to hydrogen via exposure to chromium powder at 850 °C.
(Donnelly et al. 2001). The hydrogen gas yield for each step was measured using the Pirani vacuum pressure gauge, and the hydrogen collected using a mercury topopump. Hydrogen isotopes were measured using a VGI Optima mass spectrometer. In-run repeat analyses of water standards (international standards GISP, V-SMOW, and internal standard LT-STD) gave a reproducibility better than ± 2‰ for δ²H.

RESULTS
Petrography and Mineralogy of NWA 817

The modal mineralogy of NWA 817, in vol%, is: 69% pyroxene, 10–15% olivine, 15–20% mesostasis, and 1% Fe-Ti oxide (Gillet et al. 2002; Sautter et al. 2002). Olivine grains in the sample studied are euhedral or subhedral single crystals (Fig. 2). They are zoned, with a measured core-to-rim compositional range of Fa₅₇ to Fa₈₅, respectively, which is in good agreement with previous analyses (Sautter et al. 2002; Richter et al. 2016). Grain cores are compositionally uniform and with an oval outline (Fig. 2). Augite crystals are also zoned, with narrow Fs-rich rims (Fig. 1a). EBSD shows that the mesostasis is composed primarily of elongate laths of feldspar, many of which are twinned, and EBSP patterns acquired from these laths were indexed using an albite reference (Fig. 3). Other constituents include apatite, augite, cristobalite, olivine (including 5–10 µm size fayalite dendrites), and titanomagnetite grains containing ilmenite lamellae (Fig. 3). A few regions of the mesostasis produced EBSD with no Kikuchi bands, suggesting the presence of glass, a nanocrystalline phase, or an electron beam-sensitive material (Prior et al. 2009). Minerals within the mesostasis and the rims of the phenocrysts have been deformed (Fig. 3). The active slip systems that have accommodated deformation within the fayalite dendrites are [100] and [010], dominated by (010)[100] and (001)[100], which is consistent with A-Type and E-Type slip, respectively (Karato et al. 2008) (Fig. 3).

N8-1 is crosscut by 1–3 µm wide fractures, many of which contain calcite (Figs. 1b and 4a–c). These fractures are most common in augite, can also occur in olivine, but are rare in the mesostasis (Figs. 1b and 4b). Grains of barite, some Sr-rich, occur throughout N8-1, and the 7.1 × 10⁴ µm² area that was analyzed by feature mapping contains 47 barite grains >300 nm in size. The hosts of these grains and their total size (expressed as µm²) are as follows: olivine (14 grains/26 µm²), augite (14 grains/17 µm²), mesostasis (12 grains/38 µm²), and fractures (7 grains/219 µm²). Barite also occurs within veins of iddingsite outside of the mapped region (Fig. 5).

Olivine-Hosted Iddingsite Veins

Vein Microstructure and Microstratigraphy

Iddingsite occurs as patches in the mesostasis and within grains of pyroxene (Gillet et al. 2002), but is most abundant as veins in olivine (Figs. 1a, 2, and 4a).
SEM point counting of four grains shows that they contain 5.7–20.0 vol% iddingsite (mean 11.6 vol%; 2024 points counted). The abundance of olivine in NWA 817 (10–15 vol%) means that iddingsite veins comprise ~1.2–1.7 vol% of the whole rock. These veins are red-orange in color when viewed in thin section using plane polarized transmitted light (Sautter et al. 2002), and have a lower Z than their host olivine. They are most abundant in the Fe-rich rims of olivine grains, where they are typically a few tens of micrometers in length by submicrometer in width, and have finely serrated walls (Fig. 2). Veins that crosscut olivine grain cores are up to 40 μm in width, frequently curved, and may have coarsely serrated walls (Figs. 4a and 5a). Almost all veins terminate abruptly at olivine grain margins, and so do not continue into adjacent augite grains or the mesostasis (Fig. 2).

Three components of the iddingsite can be recognized by differences in atomic number (Z) and location within the veins (1) low-Z axial bands; (2) mid-Z body of the vein; (3) high-Z rims (Fig. 5). The axial bands are ~1 μm in width, and straight or gently curving, regardless of the shapes of the vein walls (i.e., the axial bands do not zig-zag or pinch and swell in step with teeth and notches of the serrations) (Figs. 5a and 5b). Mid-Z iddingsite is the volumetrically
dominant vein constituent, and is featureless in BSE images (Figs. 5a and 5b). High-Z iddingsite forms a 1–4 μm wide selvage along vein walls, and crosscuts the mid- and low-Z components as narrow veins (Fig. 5b). Selvages of high-Z iddingsite can occur where the mid-Z iddingsite is juxtaposed with veins of calcite (Fig. 4c), but high-Z iddingsite is also present in the absence of calcite. Fibers of high-Z iddingsite penetrate into olivine (Fig. 5c), and can also crosscut the axial bands to make a ladder-like structure (Fig. 5d). All three iddingsite components are crosscut by veins of calcite, and the high-Z iddingsite contains narrow veins of barite (these veins were too small and rare to have been identified by Feature mapping; Fig. 5c).

Iddingsite Chemical Composition and Mineralogy

The low- and mid-Z iddingsite is composed principally of Si, Mg, and Fe, and yields analytical totals of ~89 wt%; the remaining 11 wt% is likely comprised of water and other volatiles not measured by SEM-EDX (Table 1). The low-Z axial bands have a higher Mg/Fe ratio than mid-Z iddingsite (Table 1; Fig. 6). Both the low- and mid-Z iddingsite are featureless in bright-field TEM images, although the two components can be distinguished by subtle differences in their degree of electron scattering, which reflects contrasts in Z (Fig. 7a). High-resolution TEM shows that the axial bands contain crystals with a ~1 nm lattice spacing that are 3–4 layers thick (i.e., 3–4 nm) (Fig. 7b). Mid-Z iddingsite also contains crystals with a ~1 nm basal layer spacing, but which are smaller than those in the axial bands (i.e., 2–3 layers thick). SAED patterns acquired from the mid-Z iddingsite have continuous rings, which is consistent with the randomly oriented nanocrystals observed by high-resolution TEM. High-Z iddingsite has greater concentrations of Ca, Mn, and Fe, and lower Si and Mg than the low- and mid-Z iddingsite, and its analytical totals are also lower (~86 wt%) (Table 1; Fig. 6). TEM images of a vein of high-Z iddingsite that crosscuts the two other
components shows that it comprises a central compact band that is rimmed by curved fibers. The interface between the curved fibers and the low- and mid-Z iddingsite is crenulated (Fig. 7c). The fibers have a basal layer spacing of 1 nm, and contain packets up to ~20 layers in thickness (Fig. 7d). The structural formula, (Ca,K)$_{0.6}$(Al,Mg,Fe,Mn)$_{4.6}$(Si,Al)$_8$O$_{20}$(OH)$_4$, is consistent with the dioctahedral smectite nontronite (Table 1).

**Iddingsite Hydrogen Isotopic Composition**

N8-2 and N8-3 evolved almost identical amounts of water during stepped pyrolysis (0.45 and 0.47 wt%, respectively), and δD values are similar between the two samples (Table 2). SEM imaging of N8-3 shows that heating to 1050 °C has led to almost complete loss of calcite, and shrinkage and fracturing of the iddingsite veins. The mean atomic number of the low- and mid-Z iddingsite in the heated samples is similar to that of the olivine, and SEM-EDX analyses give totals that are close to 100 wt% (Table 3).

**DISCUSSION**

We have sought to answer the question of whether NWA 817 was altered under different physico-chemical conditions than the other nakhlites, by fluids of a different provenance (i.e., derived from the Martian mantle as opposed to the atmosphere/crust of Mars), and possibly also at a different time. We first explore
the processes and products of aqueous alteration of NWA 817, and compare results with work on the other nakhlites. We then evaluate the hydrogen isotopic data and conclude with a discussion of the implications of our findings for understanding the history of liquid water at the nakhlite launch site.

Environments of Aqueous Alteration of NWA 817

NWA 817 contains a suite of alteration products whose crosscutting relationships describe a relative chronology of water/rock interaction. Iddingsite formed first, and was followed by calcite (Fig. 4c). Barite occurs within high-Z iddingsite and so formed after all three iddingsite components (Fig. 5c), although the relative timing of barite and calcite precipitation could not be resolved. Calcite and barite are commonplace in hot desert ordinary chondrite finds where they are interpreted to be products of terrestrial weathering (Ash and Pillinger 1995; Barrat et al. 1998; Lee and Bland 2004; Al-Kathiri et al. 2005). As outer parts of Saharan meteorites (and, by extension, other hot desert meteorites) are often enriched in barium, it will have been sourced externally (Stelzner et al. 1999). The veins calcite and barite that occur in hot desert shergottite and nakhlite finds (Coulson et al. 2007; Tomkinson et al. 2015; Hallis et al. 2017), including NWA 817, are likewise interpreted to be terrestrial in origin. The association of calcite with augite in NWA 817 (Fig. 1b) implies that a proportion of the calcium could have been derived from the dissolution of clinopyroxene. In later sections, we discuss the implications of this hot desert weathering of NWA 817 for interpreting the hydrogen isotopic results.

Formation of NWA 817 Iddingsite

The microstratigraphy of the olivine-hosted iddingsite veins shows that there are three components formed in the order of low-Z, mid-Z, and then high-Z.
The low-Z axial band is interpreted to have filled hairline fractures (e.g., Figs. 5a and 5b). The walls of these narrow veins were then replaced to make the mid-Z iddingsite, during which time serrations developed in response to a crystallographic control on the trajectory of the replacement front (Lee et al. 2013). The higher abundance of iddingsite veins in the rims than cores of olivine grains probably reflects the greater susceptibility of Fa-rich olivine to aqueous alteration (e.g., Wogelius and Walther 1992). A control on iddingsite vein formation by olivine chemistry can also account for curved veins, whose intragranular distribution is controlled by the shape of the boundary between the core and rim of zoned phenocrysts (Fig. 4a).

The compositional evolution of iddingsite is characterized by depletion in Mg from low-Z to mid-Z iddingsite followed by decreases in Mg, Al, and Si in the high-Z iddingsite, with complementary enrichment with respect to Ca, Mn, and Fe (Fig. 6, Table 1). One of the hosts of these elements is smectite, and our identification of this clay mineral agrees with Gillet et al. (2002), who described ferromagnesian smectite in NWA 817 using X-ray diffraction and TEM. However, smectite cannot be the sole constituent of low- and mid-Z iddingsite because the veins have considerably lower concentrations of Al and/or Si, and higher Fe, than terrestrial Fe-Mg smectites such as nontronite, saponite, and montmorillonite (Deer et al. 1992). Structural formulae highlight the lack of octahedral Al (Table 1). The low- and mid-Z iddingsite is similar in chemical composition to iddingsite that has replaced olivine in terrestrial metagabbros (Kendrick and Jamieson 2016; Table 3). Raman spectroscopy showed that this terrestrial iddingsite contains amorphous hydrated silica and goethite; smectite was not recorded by Kendrick and Jamieson (2016), but this is perhaps unsurprising given the difficulty in identifying nanoscale crystals of such minerals by Raman spectroscopy. We therefore suggest that the low- and mid-Z iddingsite in NWA 817 contains an Al-poor and Mg-rich smectite, probably saponite, intergrown with Fe-oxyhydroxide (goethite/ferrihydrite) and some silica. The temperature range over which water is evolved from NWA 817 during pyrolysis (i.e., ~89% at <500 °C; Table 2) is also consistent with such a mineralogy (Archer et al. 2014; Garenne et al. 2014).

The high-Z component has replaced earlier iddingsite, and its occurrence as a selvage along vein walls shows that aqueous solutions gained access to olivine grain interiors via these intragranular boundaries. The veins of high-Z iddingsite comprise a
compact core lined by nontronite fibers. The core is rich in Fe and Si (~48 wt% Fe₂O₃, ~25 wt% SiO₂) but its chemical composition does not match that of a single stoichiometric mineral. We suggest therefore that the core is composed of ferrihydrite or goethite (which have 85 and 90 wt% Fe₂O₃, respectively) with Si absorbed on crystal surfaces. Fe-oxyhydroxides can have up to 19 wt% SiO₂ as an absorbed component (Childs 1992; Andersen et al. 2009). Some of the silica detected in the high-Z core could also be present as inclusions of low- and mid-Z iddingsite, or unaltered olivine. As discussed below, the properties of low- and mid-Z iddingsite are consistent with a Martian origin, which was originally proposed by Gillet et al. (2002). As it crosscuts the two earlier components, high-Z iddingsite clearly formed during a later phase of water/rock interaction which could have been hot desert weathering given that Fe-oxyhydroxides are abundant in meteorites recovered from northwest Africa (Lee and Bland 2004).

Table 2. Stepwise pyrolysis results from N8-2 and N8-3.

| Sample | Step (°C) | Yield (mmols/mg) | H₂O (wt%) | δD (%/V-SMOW) |
|--------|-----------|------------------|-----------|---------------|
| N8-2   | 200       | 0.04             | 0.07      | −31.0         |
|        | 500       | 0.18             | 0.32      | −29.6         |
|        | 1050      | 0.03             | 0.05      | −23.8         |
| Total  |           | 0.25             | 0.45      |               |
| N8-3   | 200       | 0.10             | 0.20      | 28.7          |
|        | 500       | 0.12             | 0.22      | −103.1        |
|        | 1050      | 0.03             | 0.05      | −90.0         |
| Total  |           | 0.25             | 0.47      |               |

*aSample mass 77.5 mg.
*bSample mass 108 mg.

Fig. 7. TEM images of iddingsite in N8-1. a) HAADF STEM image of all three iddingsite components. A vein of high-Z iddingsite (HZ) crosscuts mid-Z (MZ) iddingsite and the low-Z axial band (LZ). b) High-resolution image of the low-Z iddingsite. Many small crystals are present, whose lattice fringes have a ~1 nm spacing. c) Bright-field image of high-Z iddingsite crosscutting mid-Z iddingsite. Between the two iddingsite components are curved smectite crystals and pores (white). d) High-resolution image of a smectite crystal between mid- and high-Z iddingsite. The lattice fringes have a ~1 nm spacing.
Comparison of Alteration Products in NWA 817 with Other Nakhlites

From this understanding of the chemical composition and mineralogy of NWA 817 iddingsite, we now compare it with iddingsite from the other nakhlites in order to help answer the question of whether these meteorites were altered in the same aqueous system.

Iddingsite is most common within nakhlite olivine grains, which reflects the susceptibility of this mineral to low temperature aqueous alteration (Leschin and Vicenzi 2006). The microstratigraphy of the olivine-hosted veins is similar between NWA 817, Lafayette, and Nakhla. In all three meteorites, the veins possess an axial band that has a lower Z than the rest of the vein, and this internal structure is consistent with an origin by centripetal dissolution of mesostasis glass and olivine over a brief period of time, at low temperatures (<100 °C) and low water/rock ratios. The presence of smectite suggests a moderate to alkaline pH (Mustard et al. 2008).

In order to compare the chemical composition of iddingsite between different nakhlites, a compilation was made of published SEM-EDX/electron probe analyses (Fig. 8). Significant numbers of analyses (i.e., >12) are available for only four of the nakhlites: Lafayette, Nakhla, NWA 817, and MIL 00346 (by contrast between two and four analyses are available for each of NWA 998, Governador Valadares, Y-000593, and Y-000749). The data for Lafayette, Nakhla, NWA 817, and MIL 00346 show that iddingsite in each meteorite has a narrow range in Mg concentrations, but a considerable spread in Al+Si and Fe values (Table 5; Fig. 8). Magnesium concentrations increase in the order of MIL 03346–NWA 817–Nakhla–Lafayette (Table 5). This variation in Al+Si and Fe at a fairly constant Mg could be accounted for by differences between analysis points in the relative proportions of smectite/silica and Fe-oxyhydroxide/Fe-oxide.

Martian Versus Terrestrial Sources of Aqueous Solutions

The similarities between nakhlites in the chemical composition and mineralogy of their iddingsite imply

Table 3. Chemical composition of iddingsite.

|                  | NWA 817 low-and mid-Z iddingsite | Terrestrial iddingsite |
|------------------|----------------------------------|------------------------|
|                  | N8-3<sup>a</sup> | N8-1<sup>b</sup> |                 |
| SiO₂             | 45.89              | 42.49              | 41.31            |
| TiO₂             | 0.01               | bd                 | 0.01             |
| Al₂O₃            | 0.78               | 0.36               | 0.10             |
| Cr₂O₃            | –                  | bd                 | 0.01             |
| Fe₂O₃            | 43.92              | 37.92              | 41.69            |
| MnO              | 0.60               | 0.46               | 0.49             |
| MgO              | 6.01               | 6.07               | 5.38             |
| CaO              | 0.69               | 0.06               | 0.53             |
| K₂O              | 1.28               | 0.50               | 0.07             |
| Na₂O             | 1.16               | 0.35               | 0.03             |
| SO₃              | 0.09               | 0.05               | –                |
| Total            | 100.43             | 88.25              | 89.62            |
| n                | 20                 | 31                 | 33               |

Analyses of NWA 817 by SEM-EDX.

<sup>a</sup> Analyzed after stepwise pyrolysis to 1050 °C (so water and other volatile phases would have been removed). Expressing Fe as FeO reduces the total to 96.01 wt%.

<sup>b</sup> Analyses of unheated sample N8-1 (from Table 1, column 1).

<sup>c</sup> Analyses of C (so water and other volatile phases would have been removed). Expressing Fe as Fe₂O₃ reduces the total to 96.01 wt%.

In all analyses, Fe is expressed as Fe₂O₃ to aid comparison.

– not measured; bd below detection limits. 

Iddingsite is most common within nakhlite olivine grains, which reflects the susceptibility of this mineral to low temperature aqueous alteration (Leschin and Vicenzi 2006). The microstratigraphy of the olivine-hosted veins is similar between NWA 817, Lafayette, and Nakhla. In all three meteorites, the veins possess an axial band that has a lower Z than the rest of the vein, and this internal structure is consistent with an origin by centripetal dissolution of mesostasis glass and olivine over a brief period of time, at low temperatures (<100 °C) and low water/rock ratios. The presence of smectite suggests a moderate to alkaline pH (Mustard et al. 2008).

In order to compare the chemical composition of iddingsite between different nakhlites, a compilation was made of published SEM-EDX/electron probe analyses (Fig. 8). Significant numbers of analyses (i.e., >12) are available for only four of the nakhlites: Lafayette, Nakhla, NWA 817, and MIL 00346 (by contrast between two and four analyses are available for each of NWA 998, Governador Valadares, Y-000593, and Y-000749). The data for Lafayette, Nakhla, NWA 817, and MIL 00346 show that iddingsite in each meteorite has a narrow range in Mg concentrations, but a considerable spread in Al+Si and Fe values (Table 5; Fig. 8). Magnesium concentrations increase in the order of MIL 03346–NWA 817–Nakhla–Lafayette (Table 5). This variation in Al+Si and Fe at a fairly constant Mg could be accounted for by differences between analysis points in the relative proportions of smectite/silica and Fe-oxyhydroxide/Fe-oxide.

Martian Versus Terrestrial Sources of Aqueous Solutions

The similarities between nakhlites in the chemical composition and mineralogy of their iddingsite imply
that they were altered under comparable conditions (e.g., Eh, pH), but it does not necessarily follow that fluids came from the same reservoir. Here we explore whether the hydrogen isotopic data are consistent with a single source or multiple sources of the aqueous solutions, and whether the fluids were solely Martian or both Martian and terrestrial.

The hydrogen isotopic composition of NWA 817 iddingsite shows similarities but also some differences to the other nakhlites (Fig. 9). All of the analyzed
meteorites yield some low δD values, which reflect terrestrial contamination (Leshin et al. 1996; Hallis et al. 2012). However, NWA 817 differs from the other nakhlites in that its δD values do not exceed 30‰ (Table 2; Fig. 9). Gillet et al. (2002) interpreted the unusually low δD values of NWA 817 to indicate that its iddingsite had formed from fluids derived from the Martian mantle. They discounted the possibility that the hydrogen isotopic system had been disturbed by terrestrial alteration because they found little evidence for hot desert weathering. However, the samples analyzed for the present study have been moderately weathered, producing calcite, barite, and possibly also the high-Z iddingsite. The presence of calcite along olivine vein walls (Fig. 4c) is clear evidence that terrestrial water had been in contact with iddingsite. As hydrogen is a very light and mobile atom, the hydrogen isotopic system is vulnerable to alteration, and indeed previous work has shown that terrestrial weathering can significantly change the hydrogen isotopic composition of nakhlite iddingsite. For example, Hallis et al. (2012) obtained very different δD values from iddingsite in the paired Antarctic nakhlites MIL 03346 and MIL 090030 (Fig. 9), presumably reflecting contrasting intensities of weathering. Even in Nakhla, which fell in 1911, individual analysis points range in δD from −117 to 1165‰, due to spatially variable terrestrial contamination (Hallis et al. 2012). Exchange of hydrogen isotopes between iddingsite and terrestrial water would have been facilitated by the high temperatures that NWA 817 is likely to have experienced during hot desert weathering; temperatures beneath the surfaces of low albedo rocks such as basalt in hot deserts can reach ~70 °C (Warke and Smith 1998).

This evidence for terrestrial fluids having been in contact with NWA 817 iddingsite means that the hydrogen isotopic data cannot be used with confidence to constrain the source of water that was responsible for Martian aqueous alteration. While it remains possible that the low δD values reflect fluids derived from the Martian mantle, it is also conceivable that prior to its fall to Earth NWA 817 iddingsite had δD values of >1000‰, which were then lowered to <30‰ by hot desert weathering. Our EBSD results from the mesostasis give some weight to the argument that NWA 817 was not altered by aqueous fluids with a mantle composition derived from the igneous activity that formed the NWA 817 parent lava. Olivine slip systems are sensitive to temperature, pressure, strain rate, and water content (Karato et al. 2008). The dominant A-type slip system observed in the fayalite dendrites (Fig. 3) would have formed by rapid crystallization of the mesostasis during cooling under very dry conditions and at high temperatures >1000 °C (Carter and

| Table 5. The chemical composition of iddingsite and its host rock. |
|---------------------------------------------------------------|
| **Nakhlite** | **Mean Mg/(Al+Si+Fe)** | **Mean MgO (wt%)** | **40Ar/39Ar age ± 2σ (Ma)** |
| **Iddingsite** | **Bulk rock** | **Iddingsite** | **Bulk rock** |
|---------------------------------|-----------------|-----------------|-----------------|
| Lafayette | 0.241 (0.033) | 20.282 | 12.01 (2.22) | 12.90 |
| Nakhla | 0.153 (0.040) | 20.259 | 7.10 (1.75) | 11.91 |
| NWA 817 | 0.122 (0.025) | 20.216 | 5.83 (1.24) | 10.31 |
| MIL 03364 | 0.065 (0.014) | 20.198 | 3.26 (0.78) | 9.30 |
|---------------------------------|-----------------|-----------------|-----------------|
| **Ages from Cohen et al. (2017). NWA 817 was not analyzed by them.** | | | |
| **Iddingsite mean compositions are from the following number of analyses: Lafayette 20, Nakhla 12, NWA 817 41, MIL 03346 16. Values in parentheses are standard deviations.** | | | |
| **Bulk rock mean compositions are from the following number of analyses: Lafayette 1, Nakhla 3, NWA 817 1, MIL 03346 2. Standard deviations not quoted owing to the small number of analyses.** | | | |
| **The sources of all analyses are listed in the caption of Fig. 8.** | | | |
Ave’Lallemant 1970; Jung and Karato 2001; Karato et al. 2008). While this evidence argues against aqueous alteration being linked to the phase of volcanic activity that formed NWA 817, it does not preclude the possibility that fluids could have been sourced from later magmatism at the same location. However, McCubbin et al. (2013) calculated that the nakhlite and chassignite magmatic fluids contained a minimum of 1.36 wt% Cl (within the Si-rich evolved melt that crystallized to form the intercumulus glass of MIL 03346). Therefore, if NWA 817 iddingsite did form from magmatic fluids, this Cl enrichment should be evident within its chemical composition. The absence of detectable Cl within NWA 817 iddingsite (Table 1) thus argues against a magmatic origin for this alteration product.

**Implications for Understanding Aqueous Alteration at the Nakhlite Launch Site**

The simplest interpretation of the mineralogical, chemical, and isotopic results obtained from NWA 817 is that it was altered on Mars, at the same time as the other nakhlites, and by D-rich solutions that were sourced from the Martian crust or atmosphere. The water was probably stored in permafrost, which could have melted in response to heating by a nearby impact or volcanic activity (Borg and Drake 2005). Volcanic heating is favored here because the low shock stage of the nakhlites (Fritz et al. 2005) is inconsistent with them having undergone two impacts (i.e., one to drive aqueous alteration at 633 Ma, and the other to eject the nakhlites from Mars). The crustal/atmospheric hydrogen isotopic signature was then overprinted by hot desert weathering. The low- and mid-Z iddingsite are interpreted to have formed on Mars, whereas the high-Z component could equally be terrestrial in origin.

Differences in $^{40}$Ar/$^{39}$Ar ages between nakhlites can be used to reconstruct the stratigraphy of the nakhlite launch site by assuming that eruption age corresponds to stratigraphic position (i.e., the youngest nakhlite, Lafayette, samples the flow that was closest to the Martian surface; Cohen et al. 2017). The correspondence between the mean MgO value of iddingsite and the eruption age of its host meteorite (Table 5; Fig. 10) could therefore suggest that fluids evolved in chemical composition as they moved through the lava pile. The sense of compositional change could have been a decrease in Mg as the fluids flowed from shallower to deeper levels, or an increase in Mg as the water moved from deeper to shallower levels. Empirical support for the shallower-to-deeper scenario comes from the compositional evolution of NWA 817 iddingsite. The MgO values of the three generations (i.e., low-Z, mid-Z, high-Z) are 11.5 wt%, 5.9 wt%, and 1.3 wt%, respectively. Thus, Lafayette iddingsite is compositionally similar to NWA 817 low-Z iddingsite, suggesting that the Lafayette parent lava flow was altered by “primitive” near-surface fluids. MIL 03346 iddingsite is compositionally similar to NWA 817 mid-Z iddingsite, implying that the MIL 03346 parent lava flow interacted with more “evolved” and deeper fluids. A caveat to this model is that the $^{40}$Ar/$^{39}$Ar age of the nakhlite NWA 5790 implies that it was in the middle of the lava pile (Cohen et al. 2017), yet in contrast to the overlying and underlying meteorites has little evidence for aqueous alteration (Tomkinson et al. 2015). Although this discrepancy could argue against the model of gravity-driven fluid flow, inhomogeneities in porosity and permeability of the lavas could readily explain why some meteorites escaped Martian aqueous alteration.

Another interpretation of differences between meteorites in iddingsite MgO values is that they are controlled by the chemical composition of their host rock. Iddingsite and bulk rock compositions correlate well in Nakhla, NWA 817, Lafayette, and MIL 03346.
Aqueous alteration of NWA 817

This study has sought to answer the question of whether NWA 817 was altered in the same aqueous system as the other nakhlites. The principal findings are as follows.

1. NWA 817 contains more iddingsite than any other nakhlite, thus testifying to a relatively high intensity of aqueous alteration on Mars. Even if the high-Z iddingsite is terrestrial in origin, it is insufficiently abundant to change this conclusion. The relatively high degree of aqueous alteration of NWA 817 could be explained by the unusually Fe-rich composition of its olivine grains, making them highly reactive in the presence of Martian groundwater.

2. In common with the other nakhlites, NWA 817 iddingsite occurs mainly as veins within olivine grains, and is composed of nanocrystalline smectite and a Fe-rich phase that is most likely to be Fe-oxyhydroxide. These petrographic and mineralogical similarities between iddingsite in the different meteorites suggest that the nakhlites were aqueously altered under comparable physico-chemical conditions.

3. NWA 817 iddingsite has lower δD values than any other nakhlite. However, as this meteorite has been terrestrially weathered to form calcite, barite, and possibly also high-Z iddingsite, the hydrogen isotopic data cannot be used with confidence to constrain the provenance of Martian aqueous solutions.

4. The nature of deformation of olivine crystals in the mesostasis of NWA 817 suggests that the rock was emplaced dry so that aqueous alteration is unlikely to have been linked to igneous activity associated with eruption of its parent lava flow. The possibility of alteration by later generations of magmatic fluids cannot be discounted, although the absence of detectable Cl within NWA 817 iddingsite argues against such a scenario.

5. The correspondence between the chemical composition of iddingsite and the stratigraphic position of its host rock could be interpreted to indicate that the aqueous solutions were sourced from close to the surface of Mars and evolved chemically down-flow. Alternatively, this relationship may reflect a control on the chemistry of iddingsite by the composition of its host rock in a system with very low water/rock ratios.

6. NWA 817 was probably aqueously altered along with the other nakhlites at ~633 Ma and by water derived from crustal or atmospheric reservoirs.

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