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1 **REVISED VERSION #2**

2 **A NEW CALIBRATION TO DETERMINE THE CLOSURE TEMPERATURES OF Fe-Mg**
3 **ORDERING IN AUGITE FROM NAKHLITES**

4
5 Alvaro M.^{1*}, Domeneghetti M.C.², Fioretti A.M.³, Cámara F.^{4,5}, Marinangeli L.⁶

6
7 ¹Dipartimento di Geoscienze, Università degli Studi di Padova, Italy

8 ²Dipartimento di Scienze della Terra e dell' Ambiente, Università degli Studi di Pavia, Italy

9 ³Istituto di Geoscienze e Georisorse CNR, UOS di Padova, Italy

10 ⁴Dipartimento di Scienze della Terra, Università di Torino, Italy

11 ⁵CrisDi, Interdepartmental Centre for the Research and Development of Crystallography, Torino, Italy

12 ⁶Int'l Research School of Planetary Sciences, Università G. d'Annunzio, Chieti, Italy.

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15 *Corresponding author e-mail: matteo.alvaro@unipd.it

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21 **ABSTRACT**

22 Recently it has been shown that the relatively low closure temperature (T_c) of 500(100)°C calculated for augite from
23 Miller Range nakhlite (MIL 03346,13) using the available geothermometers would correspond to a slow cooling rate
24 inconsistent with the petrologic evidence for an origin from a fast cooled lava flow. Moreover, previous annealing
25 experiments combined with HR-SC-XRD on an augite crystal from MIL 03346 clearly showed that at 600°C the Fe²⁺-
26 Mg degree of order remained unchanged, thus suggesting that the actual T_c is close to this temperature.

27 In order to clarify this discrepancy we undertook an *ex situ* annealing experimental study at 700, 800 and 900 °C,
28 until the equilibrium in the intracrystalline Fe²⁺-Mg exchange is reached, using an augite crystal from Miller Range
29 nakhlite (MIL 03346,13) with composition ca. $En_{36}Fs_{24}Wo_{40}$. These data allowed us to calculate the following new
30 geothermometer calibration for Martian nakhlites:

31
32
$$\ln k_D = -4421(\pm 561)/T(K) + 1.46(\pm 0.52) \quad (R^2=0.988), \text{ where } k_D = [(Fe^{2+}_{M1})(Mg_{M2}) / (Fe^{2+}_{M2})(Mg_{M1})].$$

33
34 The application of this new equation to other Martian nakhlites (NWA 988 and Nakhla) suggests that for augite
35 with composition close to that of MIL 03346, the T_c is up to 170°C higher with respect to the one calculated using the
36 previous available geothermometer equation, thus suggesting a significantly faster cooling in agreement with petrologic
37 evidence.

38

1 **Keywords:** augite, closure temperature, Martian nakhlite, single crystal X-ray diffraction, thermal history,
2 geothermometer.

4 INTRODUCTION

5 The broadest application of intracrystalline Fe²⁺-Mg partitioning between the *M1* and *M2*
6 crystallographic sites in the pyroxene structure is the determination of the closure temperature (T_c) of
7 the exchange reaction, that provides important constraints on the cooling rate of the pyroxene-bearing
8 host rocks (e.g. Ganguly and Domeneghetti 1996). The partition coefficient, k_D , of the order-disorder
9 Fe²⁺-Mg reaction depends on the closure temperature of the exchange equilibrium, which in turn is
10 affected by the sample cooling rate. Although this technique has been successfully developed (Virgo
11 and Hafner 1969; Saxena and Ghose 1971; Smyth 1973; Sueno et al. 1976; Ganguly 1982; Molin and
12 Zanazzi 1991; Sykes-Nord and Molin 1993; Ganguly and Domeneghetti 1996; Stimpfl et al. 1999;
13 Pasqual et al. 2000; Alvaro et al. 2011) and applied for orthopyroxene and pigeonite-bearing rocks,
14 relatively few data are available for clinopyroxenes (McCallister et al. 1976; Dal Negro et al. 1982;
15 Ghose and Ganguly 1982; Molin and Zanazzi 1991 and Brizi et al. 2000). For clinopyroxenes, the
16 extent of Fe-Mg exchange is limited, because the *M2* site is mainly occupied by Ca and Na [Ca + Na
17 \approx 0.7–1.0 atoms per formula unit (apfu) in magmatic clinopyroxenes]. The most recent calibration for
18 clinopyroxenes has been provided by Brizi et al. (2000). The geothermometer based on Fe²⁺-Mg
19 exchange in calcic clinopyroxenes has been tested in some Earth and planetary geological contexts
20 (Malgarotto et al. 1993a,b; Abdu et al. 2009), providing T_c consistent with the other geological
21 evidence. However, when applied to augites extracted from MIL 03346 and other nakhlites, this
22 calibration yielded T_c for augite (Domeneghetti et al. 2013) that appears inconsistent with the fast
23 cooling rates inferred from: (i) petrographic textures (Treiman, 2005); (ii) pyroxene morphologic
24 characters (Hammer, 2006); (iii) olivine Fe-Mg and Ca zoning profiles and ilmenite exsolution
25 (Mikouchi, et al. 2012) and (iv) experimental results on mineral equilibria (Herd and Walton, 2008).
26 In order to account for these evident discrepancies Domeneghetti et al. (2013) suggested that either
27 Brizi et al. (2000) calibration was, for some reason, unsuitable for the special composition of nakhlite
28 clinopyroxenes or the augite geothermometer was disclosing some complexity in the nakhlites final
29 cooling history. To further investigate this issue we undertook a new '*ex situ*' equilibrium annealing
30 study combined with high-resolution single-crystal X-ray diffraction (HR-SC-XRD) experiments on
31 augite crystals from Miller Range nakhlite (MIL 03346,13) with composition ca. $\text{En}_{36}\text{Fs}_{24}\text{Wo}_{40}$, in
32 order to obtain a new thermometric calibration for nakhlites.

33 The reliability of our newly proposed geothermometer over a wide range of temperatures, has
34 been evaluated through its application to the fast cooled terrestrial sample FON39 (Brizi et al. 2000),

1 a sample with composition close to that of MIL 03346. Furthermore these data have been compared
2 with those reported by Domeneghetti et al. (2013) for sample Theo's flow, regarded as a terrestrial
3 analogue for MIL 03346 (Lentz et al. 1999; 2011) in order to gain comparative information on the
4 possible stratigraphic and geological setting for nakhlites. Finally, a tentative calculation of the
5 cooling rates of these samples has been performed combining the newly collected data and the kinetic
6 data from literature (e.g. Brizi et al. 2001 and Domeneghetti et al. 2013).

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9 MATERIAL AND METHODS

10 *Samples*

11 Miller Range (MIL 03346,13), Northwest Africa (NWA 998) and Nakhla are augite-rich
12 igneous rocks formed in flows or shallow intrusions of basaltic magma on Mars (see Treiman 2005).
13 Samples MIL 03346, NWA 998 and Nakhla have been found in Miller Range, Antarctica, Northwest
14 Africa and El Nakhla al Baharia, Egypt, respectively (Treiman 2005 and references therein).

15 All the augite samples from these rocks show homogeneous cores and iron-enriched rims (Treiman
16 2005). The core compositions are ca. $Wo_{40}En_{36}Fs_{24}$, $Wo_{39}En_{38}Fs_{23}$, $Wo_{38}En_{38}Fs_{24}$, respectively for MIL
17 03346, NWA 998 and Nakhla. The terrestrial sample considered in this work, FON39, is a dacite lava
18 flow from Fonualei Island (Tongan archipelago, S.E. Pacific Ocean) containing augite crystals with
19 composition ca. $Wo_{37}En_{36}Fs_{27}$.

20 A small fragment (0.10 g) of MIL 03346,13 was obtained from the meteorite sample curator of
21 NASA Johnson Space Center, whereas crystals from the NWA 998, FON39 and Nakhla samples have
22 been kindly provided by A.J. Irving, G.M. Molin and the meteorite curator at the Natural History
23 Museum of London, C. Smith, respectively.

24 A careful selection of pyroxene single crystals under the polarizing microscope was
25 performed. Moreover, for MIL 03346, NWA 998 and Nakhla small core single crystals were obtained
26 by cutting off the zoned rims and have been labelled MIL N.14 (see Domeneghetti et al. 2013) and
27 N.19, NWA 998 N.11 and Nakhla N.1. One single crystal from FON39 (here labelled FON39 N.1)
28 has been selected from the abovementioned batch of crystals used by Brizi et al. (2000). All the
29 selected crystals showed sharp extinction and sharp diffraction profiles and were therefore considered
30 to be suitable for X-ray data collection. Crystal MIL N.19 was selected for the annealing experiments.

32 *Electron microprobe analysis*

33 Crystals NWA 998 N.11 and Nakhla N.1 were embedded in epoxy resin and polished for
34 electron microprobe analysis. A Cameca-SX50 electron microprobe with a fine-focused beam (1

1 μm diameter) operating in the wavelength-dispersive (WDS) mode was used. Operating conditions
2 were 15 kV accelerating voltage and 15 nA beam current; counting times were 20 s at the peak and
3 20 s at the background. The following synthetic end-member mineral standards were used: diopside
4 for Mg, ferrosilite for Fe, wollastonite for Si and Ca, chromite for Cr, corundum for Al, MnTiO_3 for
5 Mn and Ti, and a natural albite (Amelia albite) for Na. X-ray counts were converted into oxide
6 weight percentages using the PAP correction program. Analyses are precise to within 1% for major
7 elements and 3-5% for minor elements. The results of the chemical analysis are reported in Table 1.
8 The crystal chemical formula was calculated on the basis of six oxygen atoms,. Only those spot
9 analyses with total cation contents of 4.000 ± 0.005 atoms on the basis of six oxygen atoms and
10 charge balance $3^{[4]}\text{Al} + \text{Na} - 3^{[6]}\text{Al} - 4\text{Ti} - 3\text{Cr} - 3\text{Fe}^{3+} \leq |0.005|$ were selected and averaged. The
11 Fe^{3+} content was calculated by stoichiometry following Droop (1987).

12 The values of 37.09(14), 36.64(10), 36.68(14) e.p.f.u. calculated from the analysis of crystal
13 MIL N.1 (Domeneghetti et al. 2013), NWA 998 N.1, Nakhla N.1, respectively, are in very good
14 agreement with the sum of the observed m.a.n.s for the *M1* and *M2* sites obtained from the structure
15 refinement, before introducing chemical constraints, i.e. 37.07(6), 37.21(6), 36.66(6), 36.72(6)
16 e.p.f.u. for crystals MIL N.14 and 19, NWA 998 N.11, Nakhla N.1, respectively (see Table 1 and
17 Table 2).

18

19 *Annealing experiment and X-ray diffraction*

20 The annealing experiments have been carried out at 700, 800 and 900°C using MIL N.19
21 sample until the equilibrium in the Fe^{2+} -Mg exchange reaction was reached. The crystal was sealed
22 into a silica vial, after alternate flushing with nitrogen and evacuating, together with an iron-wüstite
23 buffer to control the oxygen fugacity $f\text{O}_2$. Inside the silica tube, the crystal and the buffer were put
24 into two small separate Pt crucibles to avoid contact between them. After equilibrium in the Fe-Mg
25 exchange reaction was reached quenching was performed by dropping the tubes into cold water.
26 Further details on the annealing protocol used are given in Alvaro et al. (2011) and Domeneghetti et
27 al. (2013).

28 HR-SC-XRD data (i.e. up to 0.434 \AA^{-1}) were collected on crystal MIL N.19 before and after
29 each annealing experiment using a three-circle Bruker AXS SMART APEX diffractometer,
30 equipped with a CCD detector and 0.3mm MonoCap collimator (graphite-monochromatized $\text{MoK}\alpha$
31 radiation, $\lambda = 0.71073 \text{ \AA}$ operating 55 kV, 30 mA) following the same procedure described in
32 Domeneghetti et al. (2013). The same data collection protocol has been adopted for crystals NWA
33 998 N.11, Nakhla N.1 and FON39 N.1. Data reduction has been performed for each sample
34 following the procedure described in detail by Domeneghetti et al. (2013) for MIL 03346 crystals.

1 The samples' site distribution were obtained through full-matrix least-squares refinements carried
2 out with SHELX program (Sheldrick 1997) as described in Domeneghetti et al. (2013). Chemical
3 constraints have been taken from the microprobe analysis as reported in Table 1¹ for NWA 998
4 N.11 and Nakhla N.1 and those reported in literature for MIL03346, FON39 (Domeneghetti et al.
5 2013; Brizi et al. 2000, respectively), assuming 1σ as the error.

6 For all crystals the constraints reported in Domeneghetti et al. (2013) were also introduced
7 into the refinement. For each crystal considered in this study the results obtained from the structural
8 refinement (i.e. unit-cell parameters, discrepancy indices R_{all} and R_w based on all the F_o^2 , the
9 goodness of fit) are reported in Table 2. The site populations obtained from the structural
10 refinements with chemical constraints are reported in Table 3.

11 12 RESULTS AND DISCUSSION

13 *Determination of the Fe²⁺-Mg ordering state*

14 The Fe²⁺-Mg ordering state was estimated from the site population (Table 3) by means of
15 the intracrystalline distribution coefficient k_D , using the same expression adopted by Brizi et al.
16 (2000): $k_D = [(Fe^{2+}_{M1})(Mg_{M2})/(Fe^{2+}_{M2})(Mg_{M1})]$. The k_D values and relative propagated errors
17 obtained are reported in Table 3.

18 An attempt **at** structural refinement was also performed by considering Mn fully ordered in
19 the M2 site in agreement with the stronger preference for the M2 site of Mn compared to Fe²⁺,
20 observed by Stimpfl (2005a, 2005b) in a donpeacorite sample. Because of the low Mn contents of
21 the selected samples (see Domeneghetti et al. 2013 and Brizi et al. 2000) this procedure did not
22 significantly affect the k_D values.

23 24 *New geothermometer and evaluation of the closure temperatures for augite samples*

25 For the two MIL 03346 crystals k_D ranges from 0.026(5) to 0.028(5), in very good
26 agreement with those found for the other two naxhlite samples of 0.027(5) and 0.025(4) for NWA
27 998 N.11 and Nakhla N.1, respectively, confirming the very similar rock history and evolution of
28 these samples. The k_D value obtained on crystal FON39 N.1 [$k_D = 0.080(5)$] is identical within
29 estimated standard deviations (e.s.d.'s) to that reported by Brizi et al. (2000).

30 In Fig. 1 $\ln k_D$ is plotted against $1/T$ for MIL 03346 crystal N.19 (this study), N.14 from
31 Domeneghetti et al. (2013), together with the literature data from Brizi et al. (2000). Weighted
32 linear regression of $\ln k_D$ versus $1/T$ for the four temperatures (600, 700, 800 and 900°C) at which
33 crystals MIL N.14 and N.19 were annealed yields the following equation:

¹ Table 2 has been deposited as supplementary material.

1
$$\ln(k_D) = -442(\pm 561)/T(K) + 1.46(\pm 0.52)(R^2 = 0.988)$$

2 where $k_D = [(Fe^{2+}_{M1})(Mg_{M2})/(Fe^{2+}_{M2})(Mg_{M1})]$.

3 The closure temperature obtained for MIL 03346 N.14 and N.19, NWA 998 N.11, Nakhla
4 N.1 and FON39 N.1 with this new geothermometer are reported in Table 3, together with those
5 obtained using the calibration by Brizi et al. (2000) on a crystal from sample FON39. As already
6 expected from their comparable k_D values, the T_c calculated for the three naxhlites are identical
7 within estimated standard deviation, ranging from 585(83)°C for Nakhla N.1 to 594(87)°C for MIL
8 N. 14. The precision of our method does not permit to reliably assess if there is a significant
9 difference between the closure temperatures of Fe-Mg ordering in clinopyroxenes in MIL 03346
10 and NWA 998 samples, unlike as indicated by petrological and textural evidence (see Treiman
11 2005; Hammer 2009, Mikouchi et al. 2012).

12 At the same time this study shows that the geothermometer calibrated by Brizi et al. (2000)
13 underestimates the naxhlites pyroxene T_c by ca. 170°C (see Table 3 and Fig. 1). In particular, the T_c
14 calculated using the equation obtained by Brizi et al. (2000) for crystal FON39 N.1 (which has a
15 composition close to that of the naxhlite pyroxenes investigated in this study) leads to the
16 discrepancies up to about 170°C (i.e. far beyond their e.s.d.'s). The $\ln k_D$ vs. $1/T$ relation for sample
17 FON39 N.1, as determined in this study, is illustrated in Fig. 1 and compared with that of Brizi et
18 al. (2000). A possible explanation for the large disagreement between the two calibrations is that
19 there was a large error in the determination of the furnace temperature in the experiments of Brizi et
20 al. (2000). However, the T_c calculated for FON39 N.1 using our geothermometer ($T_c = 836^\circ\text{C}$) is
21 still well below their first recalculated annealing T , being 922°C instead of the published 750°C.
22 Therefore, such a temperature difference would explain why the crystal was actually disordering.
23 Moreover, our T_c of 836°C seems to be reasonable considering the presence of volcanic glass in the
24 groundmass of FON39 dacite host rock .

25 Further evidence of the mismatch is provided by the calculation of the closure temperature
26 for other augite samples with different X_{Fe} and degree of order available in the literature regardless
27 of the clinopyroxene composition, i.e. Theo's flow clinopyroxene (TS7 by Domeneghetti et al.
28 2013), KC (andesitic dike) and PD30 (basaltic dike) by Brizi et al. (2000). In fact, the T_c calculated
29 with our new geothermometer for Theo's flow clinopyroxene (ca. 700°C, with k_D ca. 0.05), KC and
30 PD30 (ca. 900°C, k_D ca. 0.1) are about 100, 89 and 52 °C higher, respectively, than those obtained
31 using the equation by Brizi et al. (2000).

32 In order to evaluate the effect of such differences in closure temperature on the thermal
33 history of these samples a reliable Arrhenian relation for augite is needed. However, the only
34 available kinetic data for clinopyroxenes are those published by (i) Brizi et al. (2001) for an augite

1 sample with composition $Wo_{43}En_{46}Fs_{11}$ and (ii) Domeneghetti et al. (2005) for a $P_{21/c}$ pigeonite
2 sample with composition $Wo_{10}En_{47}Fs_{43}$. Therefore, bearing in mind that these Arrhenian relations
3 are not suitable for our Martian sample composition, a tentative calculation of the cooling rates for
4 all samples has been done. Because of the abovementioned discrepancies between our equilibrium
5 data and those by Brizi et al. (2000) we decided to calculate the cooling rate using both the kinetic
6 data of Brizi et al. (2001) and those by Domeneghetti et al. (2005) after correcting for the different
7 Ca content. The cooling rate have been modelled using an asymptotic cooling law: $1/T(K) = 1/T_0 +$
8 ηt (where η is the cooling time constant, Ganguly 1982) and assuming fO_2 conditions for Martian
9 samples equal to that of IW +2.65 (Domeneghetti et al., 2013). The calculation was carried out
10 using the program CRATE (Ganguly pers. comm.). The $C_{0(augite)}$ for each sample has been obtained
11 through a correction factor, that accounts for their Ca content, starting from $C_{0(pigeonite)}$ following the
12 procedure reported in Ganguly (1982) for diopside from Lesotho Kimberlite pipe. The resulting
13 cooling rates calculated using Brizi et al. (2001) Arrhenian relation at their respective T_c are 6.8,
14 4.2, 2.7, 4.8 and 51.3°C/h for MIL 03346, NWA 998, Nakhla, Theo's Flow and Fon39,
15 respectively. On the other hand, the cooling rates calculated at their respective T_c using the
16 Arrhenian relation from Domeneghetti et al. (2005) corrected for the different Ca contents resulted
17 in cooling rates one order of magnitude slower, being 0.16, 0.13, 0.13, 0.09 and 3°C/h for MIL
18 03346, NWA 998, Nakhla, Theo's Flow and Fon39, respectively. However, the slower cooling rate
19 obtained on TS7 compared to that of naxhlites, despite its higher T_c (720°C for TS7 vs. ca. 600°C
20 for the naxhlites), could be due to the differences in composition (i.e. Fe and Ca contents) that affect
21 both equilibrium and kinetic behavior.

22 Cooling rates alone cannot be used to calculate the precise depths at which MIL 03346,
23 NWA 998, Nakhla were cooling to around their respective T_c , because this calculation heavily
24 depends on the choice of boundary conditions (Sears et al., 1997; Nabelek et al., 2002; Vorsteen
25 and Schellschmidt, 2003) which are still fairly unconstrained for naxhlites. However, as a tentative
26 exercise, assuming a $T_0=1150$ °C for the naxhlite magma (Stockstill et al. 2005); $T_{air} = T_{bedrock} = 0$
27 °C at the time of magma extrusion (Treiman, 2003; Shuster and Weiss, 2005), a thermal diffusivity
28 of 31.5 m²/y, and using the simplified mathematical model for cooling of volcanic bodies proposed
29 by Jaeger (1968), we obtained a burial depth in the range 2.5 - 2.7 m for the three naxhlite samples,
30 at the T_c and cooling rates calculated with the new calibration. The pyroxene closure temperatures
31 of the three naxhlites and their cooling rates at the T_c are identical within error and therefore it
32 would be meaningless to try to distinguish the individual burial depths. As a further exercise we
33 calculated the burial depth of Theo's flow sample TS7 and compared the resulting burial depth
34 with the actual position of the sample within the lava sequence, as observed in the field.

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1 Assuming a single magma unit of 120 m erupted at $T_0=1140$ °C (Lentz et al., 2011), we could not
2 find any solution for a cooling rate of $= 0.091$ °C/h at a $T_c = 720$ °C . We observe, however, that
3 these cooling conditions are fulfilled in a lava flow of ca. 50 m (the same thickness as the
4 pyroxenite layer), at a burial depth of ca. 40 m, which is close to the position of this sample within
5 the pyroxenite layer. This appears to suggest that Theo's flow is not a single 120m magma unit, but
6 possibly represents multiple injections. It is worth noting that using the cooling rate values obtained
7 with Brizi et al. (2001) calibration, the same exercise cannot provide a realistic solution for the
8 burial depth of pyroxene in either the nakhlites or Theos' flow.

11 CONCLUSIONS

12 Our new calibration of the Fe-Mg exchange geothermometer, experimentally obtained on
13 augite from the nakhlite MIL 03346, provides a significant revision of the Brizi et al. (2000) augite
14 geothermometer. The new calibration yields closure temperatures (T_c) of the augite Fe-Mg
15 exchange significantly higher than those calculated with the previous calibration.

16 Tentative calculation of the cooling rates of the host lava, at the T_c of augite, allow
17 evaluation of burial depths and yield values of 2-3 m for the three nakhlite samples. Closure
18 temperatures and cooling rates for the three nakhlites, which are identical within errors, do not
19 permit any meaningful comparison between their burial depths.

20 However, these calculated cooling rates allow reconciliation of the relatively low augite T_c
21 obtained from MIL 03346 (and other nakhlites), with the petrographic and textural evidence for a
22 fast cooling. Moreover, it is clear that the nakhlites T_c (about 600°C) is lower than that calculated
23 for TS7 (720 °C) sample, which was supposed to be cooled within Theo's lava flow at a burial
24 depth of 85 m.

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Figures and Tables

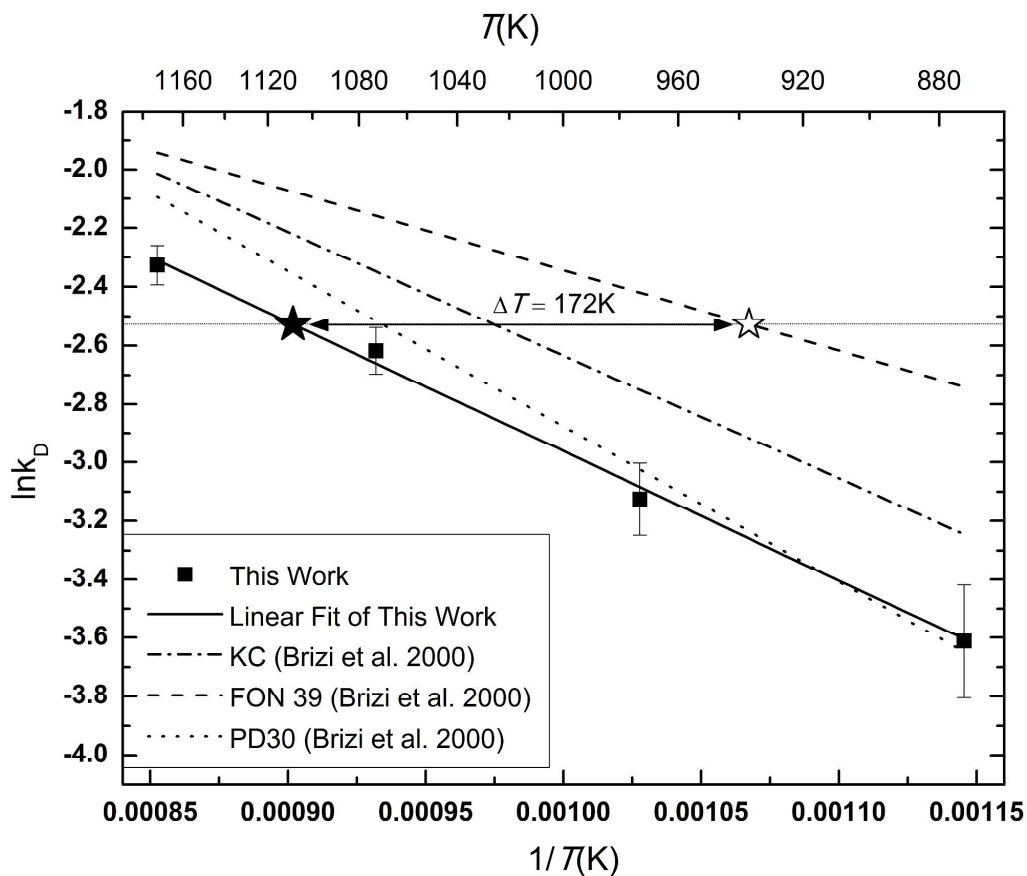


Fig. 1. $\ln k_D$ versus $1/T$ (K^{-1}) for the augite samples considered in this work together with those reported by Brizi et al. (2000) for FON39, PD30 and KC samples. Solid line represents geothermometer equation calibrated by linear fitting of MIL N.19 data. Filled star and open star represent the closure temperature calculated for FON39 N.1 with our geothermometer calibration and with that of Brizi et al. (2000), respectively.

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Table 1. Electron microprobe analyses and formulae in atoms per formula unit (apfu) based on six oxygen atoms.

| | Cpx Nakhla N.1 (averaged spots (44)) | Cpx NWA 998 N.11 (averaged spots 8) |
|--------------------------------|---|--|
| | % oxides | |
| SiO ₂ | 52.30 (22) | 51.14(18) |
| TiO ₂ | 0.11(3) | 0.34(2) |
| Al ₂ O ₃ | 0.45 (2) | 1.02(3) |
| Cr ₂ O ₃ | 0.43(5) | 0.44(3) |
| FeO | 14.33 (23) | 13.40(31) |
| Fe ₂ O ₃ | - | - |
| MnO | 0.45(4) | 0.42(4) |
| MgO | 13.39(15) | 12.99(15) |
| CaO | 18.35(13) | 18.84(14) |
| Na ₂ O | 0.14(4) | 0.27(4) |
| K ₂ O | 0.01(1) | 0.00 |
| Total | 97.97(41) | 99.06(13) |
| | a.p.f.u. | |
| Si | 1.983(6) | 1.953(4) |
| Ti | 0.003(1) | 0.010 (1) |
| Al | 0.020(1) | 0.046(2) |
| Cr | 0.013(1) | 0.013(1) |
| Fe ²⁺ | 0.448(6) | 0.401(9) |
| Fe ³⁺ | 0.006(6) | 0.033(5) |
| Mn | 0.014(1) | 0.014(1) |
| Mg | 0.757(8) | 0.740(8) |
| Ca | 0.745(5) | 0.771(5) |
| Na | 0.011(3) | 0.020(3) |
| K | 0.000(1) | 0.000(2) |
| Total | 4.000(2) | 4.001(2) |
| m.a.n.* | 36.68(14) | 36.68(17) |

m.a.n.*: calculated total mean atomic number for M1 and M2 sites, in electrons per formula unit (a.p.f.u).

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Table 2. Unit cell parameters and information on data collection and structure refinement for untreated NWA 998 N.11, Nakhla N.1 MIL N.19 and FON39 N.1. Data for MIL N.19 obtained after each annealing temperature (700, 800 and 900°C) are also reported.

| | NWA 998 | Nakhla N.1 | MIL N.19 | | | FON39 N.1 | |
|---------------------------|---------------------------------------|---------------------------------------|--|--|--|--|--|
| | N.11 | | | | | | |
| | Untreated | Untreated | Untreated | 700°C | 800°C | 900°C | Untreated |
| Crystal sizes (mm) | 0.170 x 0.128 x 0.050 | 0.185 x 0.185 x 0.090 | | 0.170 x 0.120 x 0.080 | | | 0.100 x 0.087 x 0.060 |
| <i>a, b, c</i> (Å) | 9.7468(4), 8.9397(4), 5.2509(2) | 9.7539(9), 8.9542(8), 5.2536(5) | 9.7589 (5), 8.9484 (4), 5.2537 (2) | 9.7559 (5), 8.9505 (4), 5.2538 (2) | 9.7575 (4), 8.9507 (4), 5.2551 (2) | 9.7603 (5), 8.9501 (4), 5.2553 (2) | 9.7474 (4), 8.9385 (4), 5.2536 (2) |
| β (°) | 106.2677(13) | 106.382(3) | 106.2246(17) | 106.1946(17) | 106.2057(15) | 106.2451(18) | 106.4000(18) |
| V (Å ³) | 439.21(3) | 440.21(7) | 440.52(3) | 440.56(3) | 440.73(3) | 440.75(3) | 439.11(3) |
| μ (mm ⁻¹) | 1.14 | 1.13 | 1.13 | 1.13 | 1.13 | 1.13 | 1.14 |
| I_{ind} | 2735 | 2724 | 2777 | 2785 | 2778 | 2785 | 2766 |
| R_{int} | 0.016 | 0.017 | 0.022 | 0.024 | 0.024 | 0.024 | 0.03 |
| R_{all}, R_w, S | 0.031, 0.080, 1.21 | 0.023, 0.058, 1.14 | 0.028, 0.073, 1.06 | 0.029, 0.076, 1.04 | 0.027, 0.073, 1.06 | 0.029, 0.071, 1.07 | 0.034, 0.100, 1.15 |
| m.a.n. ^(a) | 36.66(6) | 36.72(6) | 37.21(6) | 37.20(6) | 37.13(6) | 37.17(6) | 36.73(6) |

Standard deviations are given in parentheses. I_{ind} is the number of independent reflections used for structure refinement; $R_{\text{int}} = \sum |F_o^2 - F_c^2(\text{mean})| / \sum [F_o^2]$ where F_o and F_c are the observed and calculated structure factors; $R_{\text{all}} = \sum ||F_o^2| - |F_c^2|| / \sum [F_o^2]$; $R_w = \{ \sum [w(F_o^2 - F_c^2)^2] / \sum [w(F_o^2)^2] \}^{1/2}$; $S = [\sum [w(F_o^2 - F_c^2)^2] / (n-p)]^{0.5}$, where n is the number of reflections and p is the total number of parameters refined. ^(a) m.a.n. is the mean atomic number (in electrons per formula unit) before introducing the chemical constraints. Crystal system monoclinic $C2/c$; radiation type $\text{MoK}\alpha$.

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Table 3. Site populations and k_D for Martian nakhlites obtained in this work (NWA 998 N.11 , Nakhla N.1, MIL N.19 and FON39 N.1) together with those reported by Domeneghetti et al. (2013) for sample MIL N.14

| | | NWA 998 N.11 | NAKHLA N.1 | MIL N.14 | | MIL N.19 | | | FON39 N.1 ^(a) | |
|-----------|------------------------------|-----------------|---------------|-----------|--------------|--------------|--------------|--------------|-----------------------------|--------------|
| | | Untreated | Untreated | Untreated | 600° C | Untreated | 700° C | 800° C | 900° C | Untreated |
| T | Si | 1.954 | 1.983 | 1.964 | 1.965 | 1.964 | 1.965 | 1.965 | 1.964 | 1.976 |
| | Al | 0.047 | 0.017 | 0.036 | 0.035 | 0.036 | 0.036 | 0.035 | 0.036 | 0.024 |
| M1 | Mg | 0.721(2) | 0.732(2) | 0.688(3) | 0.689 (2) | 0.688(2) | 0.680 (2) | 0.670 (2) | 0.660 (2) | 0.703(2) |
| | Fe | 0.214(3) | 0.229(3) | 0.256(4) | 0.257 (3) | 0.257(3) | 0.265 (3) | 0.275 (3) | 0.285 (3) | 0.269(2) |
| | Fe³⁺ | 0.035 | 0.013 | 0.026 | 0.021 | 0.028 | 0.029 | 0.027 | 0.027 | 0 |
| | Al | 0 | 0.003 | 0.007 | 0.01 | 0.004 | 0.003 | 0.004 | 0.004 | 0.012 |
| | Cr | 0.013 | 0.013 | 0.007 | 0.007 | 0.007 | 0.007 | 0.007 | 0.007 | 0.002 |
| | Ti | 0.01 | 0.003 | 0.008 | 0.008 | 0.008 | 0.008 | 0.008 | 0.008 | 0.005 |
| | Mn | 0.007 | 0.007 | 0.008 | 0.008 | 0.008 | 0.008 | 0.009 | 0.009 | 0.009 |
| | M2 | Mg | 0.018(3) | 0.018(3) | 0.015(3) | 0.016 (3) | 0.014(3) | 0.021 (3) | 0.032 (3) | 0.039 (3) |
| | Fe | 0.196(2) | 0.224(2) | 0.191(4) | 0.190 (4) | 0.196(3) | 0.188 (3) | 0.178 (3) | 0.170 (3) | 0.223(3) |
| | Ca | 0.758 | 0.733 | 0.768 | 0.769 | 0.765 | 0.767 | 0.766 | 0.767 | 0.722 |
| | Mn | 0.006 | 0.008 | 0.006 | 0.006 | 0.006 | 0.006 | 0.005 | 0.005 | 0.008 |
| | Na | 0.022 | 0.017 | 0.02 | 0.02 | 0.02 | 0.019 | 0.02 | 0.019 | 0 |
| | k_D | 0.027 | 0.025 | 0.028 | 0.031 | 0.026 | 0.043 | 0.073 | 0.098 | 0.080 |
| | σk_D | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.006 | 0.007 | 0.005 |
| | T_c (°C) ^(b) | 411 | 397 | 405 | 416 | 434 | 497 | 636 | 734 | 664 |
| | T_c (°C) ^(c) | 601 | 585 | 594 | 605 | 623 | 684 | 811 | 895 | 836 |
| | σT_c | 87 | 83 | 87 | 84 | 83 | 67 | 43 | 31 | 33 |

Note: $k_D = [(Fe^{2+}_{M1})(Mg_{M2})/(Fe^{2+}_{M2})(Mg_{M1})]$, $R^{3+} = Fe^{3+} + Al + Cr + Ti$. The site occupancy values represent atoms per six oxygen atoms. (a) Chemical constraints introduced are based on the chemical analysis provided by Brizi et al. (2000). Closure temperature calculated using the geothermometer reported by Brizi et al. (2000) using a crystal from sample FON39 (b) and calculated using equation from this study (c). Standard deviations on closure temperatures have been calculated accounting for the linear regression errors.

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