Gem amphiboles from Mogok, Myanmar: crystal-structure refinement, infrared spectroscopy and short-range order-disorder in gem pargasite and fluoro-pargasite

Running title: Long- and short-range order in gem pargasite

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Abstract

The crystal structures of six gem-quality pargasites and fluoro-pargasites from Mogok, Myanmar, space group \( C2/m \), \( Z = 2 \), have been refined to \( R_1 \) indices of 2.20–2.78 % using MoK\( \alpha \) X-radiation. The unit formulae were calculated from the results of electron-microprobe analysis, and were used with the refined site-scattering values and the observed mean bond-lengths to assign site populations. \( ^7\text{Al} \) occurs at both the \( T(1) \) and \( T(2) \) sites but is strongly ordered at \( T(1) \). \( ^6\text{Al} \) is partly disordered over the \( M(2) \) and \( M(3) \) sites but does not occur at the \( M(1) \) site. \( ^8\text{Na} \) is split between the \( A(2) \) and \( A(m) \) sites and \( K \) occurs at the \( A(m) \) site. The infrared spectra in the principal (OH)-stretching region were measured and the fine structure was fit to component bands. The component bands were assigned to short-range ion arrangements over the configuration symbol \( M(1)M(1)M(3)–O(3)–A–O(3):T(1)T(1) \) using the refined site-populations and the expected frequencies from previously assigned spectra in more simple amphibole compositions, and correspond to the local arrangements: (1) MgMgMg–OH–Na–OH:SiAl; (2) MgMgMg–OH–Na–F:SiAl; (3) MgMgAl–OH–Na–OH:SiAl and (4) MgMgAl–OH–Na–F:SiAl.

Keywords: pargasite, amphibole, gem, crystal structure, infrared spectroscopy, Myanmar, short-range order.
Introduction

Short-Range Order-Disorder (SR-OD) is extremely common in amphiboles and describes the probability that local atomic arrangements occur more or less frequently than would be predicted by a random distribution of ions in the structure (Hawthorne and Della Ventura, 2007; Hawthorne et al., 2005). However, SR-OD does not obey translational symmetry and spectroscopic methods are generally required to derive such information. The intensity and position of bands in infrared spectra is dependent upon the (OH)-stretching frequency, which is sensitive to the local arrangement of all cations in the structure (Hawthorne and Della Ventura, 2007). Consequently, the interpretation of such spectra may reveal the preference of cations to form specific local arrangements within the C2/m amphibole structure (e.g. Hawthorne et al., 1996a; Della Ventura et al., 2014; Leissner et al., 2015). The local version of bond-valence theory (Hawthorne, 1997) may also be used to constrain local arrangements of ions, and a combination of crystal-structure refinement, infrared or MAS NMR spectroscopy, and local bond-valence calculations can derive information on SR-OD in amphibole provided that the composition of the amphibole is not overly complicated. The chemical variability of natural amphiboles gives rise to many overlapping absorption bands in the principal (OH)-stretching region of the infrared spectrum, and the spectra commonly do not contain enough information to allow the spectra to be fit correctly. This problem may be overcome by using [1] synthetic amphiboles (Raudsepp et al., 1987a,b, 1991; Boschmann et al., 1994; Della Ventura et al., 1996a,b, 1997, 1998a,b, 1999, 2001, 2003; Hawthorne et al., 1997, 2000; Robert et al., 1999, 2000; Najorka and Gottschalk, 2003), the compositions of which are usually sufficiently constrained that a combination of Rietveld refinement or single-crystal refinement, infrared and MAS NMR spectroscopy, and local bond-valence arguments are sufficient to derive SRO arrangements, or by using [2] gem amphiboles (Tait et al., 2001; Abdu and Hawthorne, 2009; Heavysege et al., 2015) for which the compositions are commonly simple. Thus the majority of work on SR-OD in amphiboles involves interpretation of the spectra of synthetic and gem-quality
samples where the composition either can be controlled or is simple enough to result in spectra that can be rigorously interpreted. In this study, we examine a series of gem-quality pargasites and fluoro-pargasites (Hawthorne et al., 2012) by single-crystal structure refinement, infrared spectroscopy and electron-microprobe analysis in order to characterize their local arrangements.

Experimental

Samples AM1, AM7 and AM8 are from the mineral collection of the U.S. Museum of Natural History and have the catalogue numbers USNM 109256, USNM 108917 and USNM 108413, respectively. Samples AM2, AM3 and AM6 were supplied by Dudley Blowett of Mountain Minerals, Colorado. Single-crystal Structure REFinement (SREF), Electron MicroProbe Analysis (EMPA) and Fourier-Transform InfraRed spectroscopy (FTIR) were done at the Department of Geological Sciences, University of Manitoba. Each sample was chosen for crystal clarity and uniform extinction under cross-polarized light. In preparation for SREF, crystals were cut to an approximate size-range 15–60 x 40–80 x 100–140 μm and attached to tapered glass fibres; similar samples were double-polished for FTIR analysis. In preparation for EMPA, the samples were mounted in epoxy in an acrylic ring, ground, polished and carbon coated.

X-ray diffraction (SREF)

Crystals were mounted on a Bruker D8 three-circle diffractometer equipped with a rotating-anode generator producing monochromatic MoKα X-radiation, multilayer optics and an APEX-II CCD detector. Totals of ~16,300 intensities were collected to 60° 2θ using 2–4 s per 0.2° frame with a crystal-to-detector distance of 5 cm. Empirical absorption corrections (SADABS, Sheldrick, 2008) were applied. Equivalent reflections were merged, resulting in ~5,320 reflections in the Ewald sphere. Intensities were averaged in the space group C2/m, resulting in ~1,370 unique reflections, and the intensities were reduced to structure factors. Unit-cell
dimensions were obtained by least-squares refinement of the positions of ~4,000 reflections with \( I > 10\sigma(I) \) and are given in Table 1. Each structure was refined in the space group \( C2/m \) with the SHELXTL version 6.14 program (Bruker AXS) to average \( R_1 \) indices of ~1.23% with anisotropic-displacement parameters at all sites. Atom positions and \( U_{eq} \) values are given in Table 2 and anisotropic-displacement parameters (Table S1) and crystallographic information files have been deposited with the Principal Editor of Mineralogical Magazine and are available as Supplementary material (see below). Selected interatomic distances are given in Table 3.

**Electron microprobe analysis (EMPA)**

Chemical analysis was done using a Cameca SX-100 electron microprobe operating in wavelength-dispersive mode using a voltage of 15 kV, a beam current of 20 nA, and a beam size of 1 \( \mu \)m. The following standards were used for \( K\alpha \) lines: Na, albite; Si, Ca, diopside; F, fluoro-riebeckite; Mg, forsterite; Al, andalusite; K, orthoclase; Ti, titanite; Fe, fayalite; Mn, spessartine; Cl, tugtupite; V, \( VP_2O_7 \); Cr, chromite; and Zn, gahnite. Ten analytical points were measured on each sample. Data were corrected using the PAP procedure of Pouchou and Pichoir (1985), and the mean compositions are given in Table 4, together with the unit formulae calculated on the basis of \((O,OH,F,Cl) = 24\) atoms per formula unit (apfu) with \((OH,F,Cl) = 2\) apfu.

**Infrared spectroscopy**

Fourier-Transform InfraRed (FTIR) spectra in the principal (OH)-stretching region were collected for all gem amphibole samples using a Bruker Tensor 27 FTIR spectrometer equipped with a KBr beam splitter and a DLATGS detector. It should be noted that the spectra of AM6 and AM7 contain relatively more noise due to lack of material.

Spectra over the range 4000–400 cm\(^{-1}\) were collected by averaging 100 scans at an operating resolution of 4 cm\(^{-1}\). Base-line corrections were done using the OPUS software and
spectra analysis in the 3800–3600 cm$^{-1}$ range was done using the OMNIC software. Spectra were fit using Gaussian curves of similar width (full-width-at-half-maximum) with the program FITSYK V0.9.8.

The $C2/m$ amphibole structure

The general formula for the amphiboles can be written as $A_{0.1}B_2C_5T_8O_{22}W_2$, where $A = Na, K, Ca, Li; B = Na, Li, Ca, Mn^{2+}, Fe^{2+}, Mg, Sr; C = Mg, Fe^{2+}, Mn^{2+}, Al, Fe^{3+}, Mn^{3+}, Ti^{4+}, Li; T = Si, Al, Ti^{4+}; W = (OH), F, Cl, O^{2-}$. The $C2/m$ amphibole structure is shown in Figure 1 and the correspondence between the general formula and the sites may be written as follows: $^A[A(2), A(m)]^B[M(4)]^C[M(1)_2 M(2)_2 M(3)]^T[T(1)_4 T(2)_4]O_{22}^\times O(3)_2$. A detailed summary of previous work on the structure and crystal chemistry of amphiboles was given by Hawthorne (1981, 1983a), Hawthorne and Oberti (2007) and Oberti et al. (2007).

Long-range site-populations

The $T$ sites

These amphiboles contain 1.79–2.41 apfu Al at the $T$ sites (Table 4). The scattering values of Si and Al are similar, which precludes accurate derivation of site populations by site-scattering refinement (Hawthorne, 1983c; Hawthorne et al., 1995). However, the difference in ionic radii of $^\text{\`T}$Si (0.26 Å) and $^\text{\`T}$Al (0.39 Å) (Shannon, 1976) allows derivation of site populations from $<T-O>$ distances (e.g. Papke et al., 1969, Hawthorne and Grundy, 1973a,b, 1977; Robinson et al., 1973; Bocchio et al., 1978; Hawthorne, 1981, 1983a,b; Oberti et al., 1995a). The preference of $^\text{\`T}$Al for $T(1)$ was confirmed by neutron diffraction (Welch and Knight, 1999) and by spectroscopic methods (Welch et al., 1994, 1998; Jenkins et al., 1997; Najorka and Gottschalk, 2003; Della Ventura et al., 2007). Hawthorne and Oberti (2007) showed that grand $<T-O>$ distances are also influenced by inductive effects where $^\text{\`T}$Al < 0.50 apfu, but vary linearly with $^\text{\`T}$Al content where $^\text{\`T}$Al > 0.50 apfu. As shown in Fig. 2a, these gem amphiboles accord very closely with this
relation with deviations of <0.001 Å. Hawthorne and Oberti (2007) also gave predictive relations for the Al site-populations of the $T(1)$ and $T(2)$ sites based on the observed $<T(1)–O>$ and $<T(2)–O>$ distances which were used here to derive site populations for the $T(1)$ and $T(2)$ sites. The resulting site-populations are in accord with the $^7$Al contents derived by electron-microprobe analysis (Fig. 2b).

**The $M(1,2,3)$ sites**

The total refined scattering at the $M(1,2,3)$ sites ($62.95–63.97$ electrons per formula unit [epfu]) (Table 5) shows an extremely small range which is consistent with the C-group cations indicated by the unit formulae of Table 4 (Fig. 3a). Similarly, the variation in grand $<M(1,2,3)–O>$ distances shows a linear correlation with the mean radius of the C-group cations from the unit formula in Table 4 (Fig. 3b). The compositions of the C-group are dominated by Mg ($3.59–4.14$ apfu) and Al ($0.56–1.28$ apfu), and the similarity of the scattering factors of Mg and Al ensure the very narrow range of refined site-scattering from the $M(1,2,3)$ sites. The net refined site-scattering values at $M(1,2,3)$ deviate from the effective scattering from the C-group cations of the unit formulae (Table 4) by an average of $0.31$ epfu. To facilitate assignment of site populations from the refined site-scattering values, the latter were adjusted (generally within one standard deviation, Table 5) to conform to the effective scattering from the C-group cations of the unit formulae indicated by the chemical formulae.

In most amphiboles, $[^6]$Al is ordered at $M(2)$ (e.g. Hawthorne, 1981, 1983a; Oberti et al., 2007). However, Semet (1973) showed by infrared spectroscopy that $[^6]$M$^{3+}$ cations disorder over the $M(2)$ and $M(1)$ and/or $M(3)$ sites in Mg-rich synthetic amphiboles, and Raudsepp et al. (1987a,b, 1991) showed by infrared spectroscopy and Rietveld refinement that this disorder occurs over the $M(2)$ and $M(3)$ sites. Oberti et al. (1995b) showed the same disorder of $[^6]$Al over the $M(2)$ and $M(3)$ sites in Mg-rich pargasites from peridotites, and this disorder of $[^6]$Al over the $M(2)$ and $M(3)$ has since been shown to be the case for Mg-rich synthetic (e.g. Welch and
Knight, 1999) and natural amphiboles (e.g. Tait et al., 2001; Heavysege et al., 2015). The absence of Al at the $M(1)$ site allows us to calculate the $M(1)$ site-populations directly from the refined site-scattering values (Table 5) using the scattering curves for Mg and the dominant transition-metal in the unit formulae. The resultant site-populations are also given in Table 5.

Preliminary site-populations for the $M(2)$ and $M(3)$ sites were calculated from the refined site-scattering values (Table 5) in terms of Mg* (= Mg + Al) and the dominant transition-metal. The results indicate that all V$^{3+}$, Cr$^{3+}$ and Ti$^{4+}$ occurs at the $M(2)$ site and the $<M(2)-O>$ bond lengths are in accord with this assignment. The X-ray scattering factors of Mg ($Z = 12$) and Al ($Z = 13$) are very close, and mean bond-lengths must be used to assign $[^6]$Al and $[^6]$Mg as the sizes of $[^6]$Mg ($r = 0.72$ Å) and $[^6]$Al ($r = 0.535$ Å) are significantly different (Shannon, 1976). There are equations relating $<M(2)-O>$ and $<M(3)-O>$ distances to constituent cation and anion radii and to other stereochemical aspects of the adjacent sites (Hawthorne and Oberti, 2007). Using these equations, $[^6]$Al and $[^6]$Mg were assigned to $M(2)$ and $M(3)$ such that there is an equally good fit between the observed and calculated $<M(2)-O>$ and $<M(3)-O>$ values; the deviations are within the error of estimate for each site given by Hawthorne and Oberti (2007). The resultant site-populations are given in Table 5.

The $M(4)$ site

The unit formulae (Table 4) indicate that the $M(4)$ site is occupied dominantly by Ca with minor Na, and the refined site-scattering values are in accord with this occupancy (Table 5). Thus the site populations were assigned directly from the formulae of Table 4.

The A site

In accord with previous work on local order-disorder in the A cavity of monoclinic amphiboles (e.g. Hawthorne and Grundy, 1972, 1973a,b; Hawthorne et al., 1996b), Na was assigned to the A(2) site and (Na + K) was assigned to the A(m) site in agreement with the refined site-
scattering values at these sites (Table 5) and the formulae derived from electron-microprobe analysis (Table 4).

**Short-range order in the C2/m amphibole structure**

The infrared spectra of amphiboles are strongly affected by Short-Range Order (SRO); the occurrence of local clusters of ions that may (or may not) give rise to discrete bands in the principal (OH)-stretching region of the infrared (Strens, 1966). Hawthorne et al. (1996a) showed that SRO is common in amphiboles, and that bands in the infrared can be assigned to specific local clusters of ions that centre on the O(3) site (Hawthorne and Grundy, 1976). Extensive work since then (Hawthorne *et al.*, 1997, 2000; Della Ventura *et al.*, 1999, 2001, 2003; Robert *et al.*, 1999, 2000; Hawthorne and Della Ventura, 2007; Leissner *et al.*, 2015) has shown that both nearest-neighbour (NN) and next-nearest-neighbour (NNN) cations affect the principal (OH)-stretching frequency of the locally associated (OH) group, and both infrared and Raman spectroscopy of the principal (OH)-stretching region may be used as a sensitive probe of NN and NNN short-range arrangements in amphiboles.

The arrangement of atoms around the O(3) site in the C2/m amphibole structure is shown in Figure 4a, b. The O(3) site is situated at the centre of the strip of octahedra and is bonded to two M(1) and one M(3) cations. Hawthorne and Grundy (1976) found that where O(3) = (OH), the H atom is ~1 Å (0.957(6) Å) away from the O(3) site and that the H atom forms a hydrogen bond with one of the O(7) oxygen atoms, shown by the dashed black line in Figure 4a, b. The strength of the bond O(3)-H bond controls the frequency of the principal (OH)-stretching vibration which is in turn related to the strength of the H…O(7) hydrogen bond (Hawthorne *et al.*, 2005). It follows that any variation in the occupancy of the sites locally associated with the O(3) and O(7) sites gives rise to fine structure in the principal (OH)-stretching region of the infrared. Such locally associated sites include the NN M(1)M(1)M(3)-, T(1)T(1)- and A(2/m)-sites. The M(1)M(1)M(3) sites are locally associated with the O(3)(OH) group which also interacts....
with the cation occupying the \( A \) site. Satisfaction of the bond-valence requirements at the \( O(7) \) site also influences the strength of the locally associated hydrogen bond and is controlled by the occupancy of the locally associated \( T(1) \)-sites.

*The configuration symbol and possible arrangements*

Hawthorne *et al.* (2005) introduced the idea of a *configuration symbol* which denotes the NN and NNN cation-sites associated with the \( O(3) \) site: \( M(1)M(1)M(3)–O(3)–A: T(1) T(1)–M(2)M(2)M(3)–M(2)M(2) \). Della Ventura *et al.* (1999) showed that the major bands in the spectrum of pargasite are slightly split by different occupancy of NNN sites. However, this splitting is small, and Robert *et al.* (2000) ignored this splitting in fitting of the spectra of pargasite–fluoro-pargasite solid-solutions. We use the same approach here, and consider only NN and NNN effects involving the configuration symbol \( M(1)M(1)M(3)–O(3)–A: T(1) T(1) \).

However, we need to incorporate the NNN effect of \( F \) across the occupied \( A \)-site into the configuration symbol (see below for a discussion of this effect). We shall do so in the following way: \( M(1)M(1)M(3)–O(3)–A–O(3): T(1) T(1) \) where the following arrangements about the \( A \) site are possible: \( M(1)M(1)M(3)–OH–A–OH: T(1) T(1) \); \( M(1)M(1)M(3)–OH–A–F: T(1) T(1) \); and \( M(1)M(1)M(3)–F–A–F: T(1) T(1) \); the last arrangement has no expression in the infrared.

*Constraints of composition*

[1] Inspection of Table 4 shows that the \( A \)-group cations completely occupy the \( A \) site, and hence local arrangements will involve only \( A = Na^+ \) (\( Na^+ = Na + K \)).

[2] For Mg-rich compositions, it has been shown both crystallographically (e.g. Raudsepp *et al.*, 1987a,b; Oberti *et al.*, 1995b; Welch and Knight, 1999) and spectroscopically (e.g. Semet, 1973; Raudsepp *et al.*, 1987b; Welch *et al.*, 1994, 1998; Della Ventura *et al.*, 1998b) that the trivalent C-group cations are partly disordered over the \( M(2) \) and \( M(3) \) sites, and Oberti *et al.* (1995b) showed how the \(<M(2)–O>\) and \(<M(3)–O>\) distances can be used to derive the site populations
associated with this disorder. Using the site populations of the \( M(3) \) site (Table 5), we may calculate the relative amounts of local arrangements at \( M(1)M(1)M(3) \) in the following manner:

\[
\text{MgMgMg:} \text{MgMgAl} = \text{M}^{\text{M(3)}} \text{M}^{\text{M(3)}} \text{Al (M}^{\text{M+}} = \text{Mg} + \text{Fe}^{2+}).
\]

[3] Arrangements involving \( T(1)–O(7)–T(1) \) are affected by the amount of \( ^{\text{T}}\text{Al} \) in the structure. Hawthorne and Oberti (2007) state that \( ^{(1)}\text{Al}–O(7)–^{(1)}\text{Al} \) arrangements do not occur unless \( O(7) \) is locally associated with \( \text{Ca} \) at the \( A \) site, as in fluoro-cannilloite (Hawthorne et al., 1996b). This is not the case for any of the amphiboles examined here (Table 4), \( ^{(1)}\text{Al}–O(7)–^{(1)}\text{Al} \) arrangements cannot occur, and the site populations of Table 5 may be used to calculate the frequencies of occurrence of arrangements involving the \( T(1)T(1) \) sites:

\[
^{(1)}\text{Al–O(7)–}^{(1)}\text{Si} = \frac{^{(1)}\text{Al}}{2} \text{ and } ^{(1)}\text{Si–O(7)–}^{(1)}\text{Si} = 1 – \frac{^{(1)}\text{Al}}{2}.
\]

[4] There is only minor occupancy of the \( C \)-group by divalent cations other than \( \text{Mg} \) (\( \text{Fe}^{2+} \)), and arrangements involving \( \text{Fe}^{2+} \) may be ignored (except perhaps for crystal AM1).

[5] Heavysege et al. (2015) proposed that arrangements involving \( M(1)M(1)M(3) = \text{MgMgAl} \) should be locally associated with the arrangement \( T(1)T(1) = \text{SiAl} \), as the bridging \( O(7) \) requires more bond-valence from the hydrogen bond when linked to \( \text{SiAl} \) than when linked to \( \text{SiSi} \). This local arrangement is possible as \( ^{(1)}\text{Al} > ^{\text{M(3)}}\text{Al} \) in all amphiboles of Table 5.

The nearest-neighbour effect of \( F \)

There are three possible local arrangements involving \( M(3) \) and \( \text{OH,F} \): [1] \( \text{OH–M(3)–OH} \), [2] \( \text{OH–M(3)–F} \), and [3] \( \text{F–M(3)–F} \); these arrangements gives rise to two, one and zero absorption events in the infrared, respectively. Thus the proportion of absorption events to null-absorption events is equal to the \( \text{(OH):F} \) ratio.

There is no disorder of trivalent cations over \( M(2) \) and \( M(3) \) in amphiboles where \( O(3) \) is occupied by \( F \) (Raudsepp et al., 1987b; Oberti et al., 1995c, 1998). This means that in amphiboles where both \( \text{OH} \) and \( F \) are present in major amounts, \( F \) will be strongly to completely associated with the local arrangement \( M(1)M(1)M(3) = \text{MgMgMg} \) and not with the
local arrangement MgMgAl. Above (point [2]), we showed that the $T(1)T(1)$ arrangements occur in the ratio $T(1)Al / 2 : 1 - T(1)Al / 2$. Inspection of Table 5 shows that $T(1)Al / 2 > M(3)Al$ for all crystals examined here, in accord with this short-range constraint. In addition, the number of arrangements, $T(1)T(1) = SiSi = 1 - T(1)Al / 2$, is always less than the amount of F in the amphibole, and hence all local arrangements MgMgMg–X–Na:SiSi will have X = F and have no expression in the infrared. The local arrangement MgMgMg–OH–Na:SiSi gives rise to a band at 3730–3740 cm$^{-1}$ (Della Ventura et al., 1999; Hawthorne, 2016). In accord with the above discussion, this band is not observed in the spectra of the amphiboles examined here (Fig. 5). A similar absence of this 3730–3740 cm$^{-1}$ band occurs in the spectra of fluoro-edenite and fluoro-pargasite (Della Ventura et al., 2014). In the Mogok amphiboles, the local arrangement MgMgMg–OH–Na:SiAl gives rise to a band at 3720–3710 cm$^{-1}$ (~3711 cm$^{-1}$) as $T(1)Si$ substituted for by $T(1)Al$ results in a relative band-displacement of $-20$ cm$^{-1}$ (Della Ventura et al., 1999; Hawthorne, 2016). The occurrence of such a band is in accord with the site populations of these amphiboles, and this is typically the highest-frequency band observed (Fig. 5, band A).

The next-nearest-neighbour effect of F

Infrared spectroscopy of amphiboles in the tremolite-fluorotremolite series show a single (OH)-stretching band at 3674 cm$^{-1}$ (Jenkins, 1987; Gottschalk et al., 1999; Robert et al., 1999), and the position of this band does not change with varying F content (apfu). This one-mode behaviour (Chang and Mitra, 1968) is due to a lack of coupling between O(3) sites adjacent across the A cavity, in accord with the fact that the A-site in tremolite is vacant and cannot couple O(3) arrangements across the A-site cavity (Hawthorne et al., 2005).

Amphibole with an occupied A site shows two-mode behavior (Chang and Mitra, 1968) with regard to the variation in (OH) and F in the structure (Robert et al., 1999, 2000). Work on the richterite-fluororichterite series shows three distinct bands in the infrared, one at 3730 cm$^{-1}$ due to the arrangement MgMgMg-OH-Na-OH:SiSi, one at 3674 cm$^{-1}$ due to the arrangement
MgMgMg-OH\(\bullet\)-OH:SiSi, and one at 3711 cm\(^{-1}\) due to the arrangement MgMgMg-OH-Na-F:SiSi (Della Ventura et al., 1998a, 1999; Hawthorne et al., 2005). This two-mode behaviour (Chang and Mitra, 1968) suggests that the local (OH)-group arrangement is affected by the adjacent arrangement involving F; the occurrence of an additional band in the richterite spectrum (\(A = \text{Na}\)) compared to the tremolite spectrum (\(A = \square\)) indicates that coupling of adjacent O(3) arrangements occurs only in the presence of an occupied A-site. It should also be noted that the intensity of the band at 3674 cm\(^{-1}\) decreases with increasing F, further evidence of coupling (OH)-F coupling across the occupied A-site. Hawthorne et al. (2005) suggested that this coupling is due to differential repulsion between H and the A-site cation, which forces the A-cation to occupy a more central position. Where the adjacent O(3) site is occupied by F, no such repulsion occurs and thus the A-group cation moves further away from the (OH)-group. As a result, the vibrational frequency between the H and A-group cation is reduced, and the OH-\(^4\)Na-F arrangement will absorb at a relatively lower wavenumber.

The frequency of local arrangements OH-\(^4\)Na-OH, OH-\(^4\)Na-F and F-\(^4\)Na-F may be calculated from the F content of the crystal (Heavysege et al., 2015) where \(x = F/(F + OH)\):

\[
\begin{align*}
\text{OH-}\(^4\)Na-OH: & \quad (1 - x)^2/(1 - x + x^2) \\
\text{OH-}\(^4\)Na-F: & \quad x(1 - x)^2/(1 - x + x^2) \\
\text{F-}\(^4\)Na-F: & \quad x^2/(1 - x + x^2)
\end{align*}
\]

The arrangement F-\(^4\)Na-F is not visible in the infrared, and the relative intensities for any arrangement involving \(A = \text{Na}\) (or K) and its F-shifted equivalent may be calculated from the relevant expressions above and the F content of the crystal. Moreover, the relative displacement of the F-shifted equivalent band is \(~19\) cm\(^{-1}\) (Robert et al., 1999, 2000).

Accordingly, the F-shifted MgMgMg-OH-Na:SiAl arrangement should give rise to a band at 3700–3690 cm\(^{-1}\), which is indeed the case (Fig. 5, band B).

Above in point [2], we saw that the NN configuration has two principal arrangements, MgMgMg and MgMgAl. Hence the arrangements derived above, MgMgMg-OH-Na-OH:SiAl
and (F-shifted) MgMg–OH–Na-F:SiAl, should have $M(1)M(1)M(3) = MgMgAl$ equivalents:

$$MgMgAl–OH–Na–OH:SiAl$$ and (F-shifted) $MgMgAl–OH–Na–F:SiAl$. Thus we expect to see four bands in the infrared spectra of these amphiboles. These $MgMgAl–OH–Na–OH:SiAl$ and $MgMgAl–OH–Na–F:SiAl$ arrangements should give rise to bands at 3687–3677 cm\(^{-1}\) and 3667–3657 cm\(^{-1}\), respectively, as $^{M(1,3)}Mg$ substituted by $^{M(1,3)}Al$ results in a relative band displacement of $\sim 33$ cm\(^{-1}\) (Della Ventura et al., 1999). Calculating the frequencies of occurrence of these last two arrangements via the relation $MgMgMg:MgMgAl = ^{M(3)}Mg^*:^{M(3)}Al$ ($Mg^* = Mg + Fe^{2+}$) indicates that they are significantly less common than their MgMgMg analogues.

However, their absorption is shifted to lower frequency, and as shown by Skogby and Rossman (1991), the transition probability of absorption increases strongly with decreasing absorption frequency, and hence the ratios of local arrangements do not one-to-one correspond with the absorption intensities of the corresponding bands in the infrared.

**Description and interpretation of spectra**

The observed band intensities and the amounts of the local arrangements assigned to these bands in the spectra of each amphibole (Fig. 5) are given in Table 6. In general, the observed band intensities and the calculated amounts of the corresponding local arrangements are in reasonable accord. However, the disagreement between observed and calculated values tends to increase among arrangements corresponding to bands at lower frequencies. This is probably due to increased transition probabilities resulting in the lack of one-to-one correspondence between the relative amounts of local arrangements and the intensities of their associated absorption bands (Skogby and Rossman, 1991; Della Ventura et al., 1996a,b; Hawthorne et al., 1997). The results of this effect are apparent for AM2 and AM6. For these crystals, $^{M(3)}Al$ is low (0.01–0.05 apfu) (Table 5), and one expects the band intensity of any local arrangement involving $^{M(3)}Al$ (bands C and D) to be relatively low. However, this is not the case as the observed absorption intensities of the bands are significantly higher than predicted (Table 6).
The NNN effects of band-splitting, which have been ignored here, could also cause slight discrepancies between band intensities and the calculated amounts of different local arrangements (Della Ventura et al., 1999; Robert et al., 2000).

AM1–3 and 6–8 were each fit using four peaks, corresponding to the local arrangements: (1) MgMgMg–OH–Na–OH:SiAl; (2) MgMgMg–OH–Na–F:SiAl; (3) MgMgAl–OH–Na–OH:SiAl and (4) MgMgAl–OH–Na–F:SiAl (Table 6). Figure 5 shows that most of the observed bands corresponding to the same local arrangement are centered reasonably close to the same wavenumber (cm\(^{-1}\)), as one would expect from the arguments previously proposed. However, not all bands corresponding to the same local arrangement are centered at exactly the same wavenumber (cm\(^{-1}\)) in all samples. For example, Figure 5 shows that band A is centered at ~3707 ±4 cm\(^{-1}\) in all samples; however, it is tightly centered at ~3710 cm\(^{-1}\) in AM1, 3 and 8, and at ~3707 cm\(^{-1}\) in AM2 and 7. A similar correlation is observed for band B, which is tightly centered at ~3692 cm\(^{-1}\) in AM1, 3, 8 and 6 and at ~3684 cm\(^{-1}\) and ~3688 cm\(^{-1}\) in AM2 and 7, respectively (Table 6).

Most of the curves used to fit the observed spectra have band centres that fall well within the theoretical frequency ranges previously calculated. Nonetheless, certain chemical features may explain minor differences in band position. \(^{M(1,3)}\)Fe\(^{2+}\) ranges from 0.02–0.12 apfu (Table 5) and results in a frequency shift of ~15 to ~18 cm\(^{-1}\) where substituting for \(^{M(1,3)}\)Mg, only half the frequency shift produced by \(^{M(1,3)}\)Mg \(\leftrightarrow\) \(^{M(1,3)}\)Al (~33 cm\(^{-1}\)) (Iezzi et al., 2005; Della Ventura et al., 1999). Even though the relatively low amount of Fe\(^{2+}\) may have discernible effects in the infrared, it is probably masked by the MgMgMg–OH–Na–F:SiAl arrangement (~3692 cm\(^{-1}\)) as the arrangement MgMgMg–OH–Na–OH:SiAl may be shifted ~15 to ~18 cm\(^{-1}\) via the substitution \(^{M(1,3)}\)Mg \(\leftrightarrow\) \(^{M(1,3)}\)Fe\(^{2+}\), from 3711 cm\(^{-1}\) to 3693–3696 cm\(^{-1}\). The amount of \(^{A}\)K in most samples is significant and ranges from 0.05–0.44 apfu (Table 5). Although all local arrangements are written with Na as the A-site constituents, K partly occupies the A-site in all samples and may account for some minor differences in band position between samples as \(^{A}\)Na \(\leftrightarrow\) \(^{A}\)K results in a
frequency shift of +5 cm\(^{-1}\) (Robert et al., 1989).

**Summary**

In all samples, \(^{7}\)Al occupies both \(T(1)\) and \(T(2)\) but is strongly ordered at \(T(1)\). \(M(1,2,3)\) are dominated by \([6]\)Mg, \([6]\)Al is partly disordered over \(M(2)\) and \(M(3)\) and is absent at \(M(1)\). \([6]\)Fe\(^{2+}\) is disordered over \(M(1,2,3)\) and minor tri- and tetravalent cations (Cr\(^{3+}\), V\(^{3+}\) and Ti\(^{4+}\)) are weakly ordered at \(M(2)\). \(M(4)\) is completely occupied by Ca, except for AM1 which contains minor \([M(4)]\)Na. \(A(2)\) is dominated by Na and \(A(m)\) is occupied by both Na and K (Table 5). The \(O(3)(OH):F\) ratio ranges from 1.42:0.58 (AM3) to 0.84:1.16 (AM6) (Table 4).

Band A at \(~3707\) cm\(^{-1}\) is the most intense (Fig. 5) and corresponds to the local arrangement MgMgMg–OH–Na–OH:SiAl. Band B at \(~3692\) cm\(^{-1}\) is typically the second most intense band and corresponds to the arrangement MgMgMg–OH–Na–F:SiAl, the F-shifted equivalent of band A. Bands C and D correspond to the arrangements MgMgAl–OH–Na–OH:SiAl and MgMgAl–OH–Na–F:SiAl, respectively. There is reasonable agreement between observed band intensities and calculated ratios of the corresponding arrangements (Table 6), supporting the assignment of these bands to these specific local arrangements.

**Acknowledgements**

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**Supplementary material**

To view supplementary material for this article, please visit https://doi.org/10.1180/mgm.2018.149
References


Jenkins, D.M. (1987) Synthesis and characterization of tremolite in the system H₂O-CaO-MgO-


Strens, R.G.J. (1966) Infrared study of cation ordering and clustering in some (Fe,Mg) amphibole solid solutions. *Chemical Communications (Chemical Society of London)*, 15, 519–520.


**Figure Captions**

**Figure 1.** The monoclinic $C2/m$ amphibole structure viewed along 001.
Fig. 2. (a) grand $\langle T-O \rangle$ (Å) plotted against $^{T}\text{Al}$ (apfu) with associated error; (b) calculated $^{T}\text{Al}$ (apfu) plotted against observed (empa) $^{T}\text{Al}$ (apfu) for each sample (shown as red dots). Dashed black line in (a) represents the equation given by Hawthorne & Oberti (2007), and in (b) the 1:1 relation between the variables.
Fig. 3. (a) effective scattering from C-group cations (epfu) plotted against refined $M(1,2,3)$ site-scattering (epfu); (b) mean radius of the $M(1,2,3)$ sites (Å) ($<r[M(1,2,3)]>$) plotted against the grand mean $M(1,2,3)$–O bond-lengths (Å) ($<M(1,2,3)$–O>) for each sample (shown as red dots). Dashed black lines represent a linear relation between both variables in (a) and (b).
Fig. 4. (a) the local structure surrounding the O(3) site (red circle); (b) connectivity around the O(3) site in C2/m amphiboles projected onto (010). The dashed black line represents the H…O(7) bond in (a) and (b). T(2) sites have been omitted in (b) for simplicity. Modified from Della Ventura et al. (1999).
Fig. 5. Comparison of the fitted spectra of each sample in the principal (OH)-stretching region. Vertical lines represent the peak positions of bands A, B, C and D. Black curves represent the observed spectra and red curves represent the sum of the fitted bands. Note the slight deviations in band centres corresponding to the same local arrangements in each sample.
TABLE 1. Structure-refinement information for gem amphiboles from Mogok, Myanmar.

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TABLE 2. Atom coordinates and equivalent–displacement parameters (Å²) for gem amphiboles from Mogok, Myanmar.

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M(2) & 0 & 0.17609(3) & 0 & 0.00477(19) \\
M(3) & 0 & 0 & 0 & 0.0076(3) \\
M(4) & 0 & 0.28064(2) & 1/2 & 0.00949(12) \\
A(2) & 0 & 0.4878(2) & 0 & 0.0059(16) \\
A(m) & 0.5363(6) & 0 & 0.0743(11) & 0.0204(12) \\
O(1) & 0.10448(10) & 0.08913(6) & 0.21492(19) & 0.0097(2) \\
O(2) & 0.11842(10) & 0.17378(5) & 0.73711(19) & 0.00862(19) \\
O(3) & 0.10634(14) & 0 & 0.7146(2) & 0.0096(4) \\
O(4) & 0.36824(11) & 0.25181(6) & 0.79307(19) & 0.0107(2) \\
O(5) & 0.35096(10) & 0.14074(6) & 0.11572(19) & 0.0111(2) \\
O(6) & 0.34365(10) & 0.11791(6) & 0.61223(19) & 0.0117(2) \\
O(7) & 0.33793(16) & 0 & 0.2848(3) & 0.0143(3) \\
\end{array}
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**AM7**
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T(1) & 0.27907(4) & 0.08597(2) & 0.30456(8) & 0.00842(13) \\
T(2) & 0.29038(4) & 0.17390(2) & 0.81518(7) & 0.00677(13) \\
M(1) & 0 & 0.08948(4) & 1/2 & 0.0086(2) \\
M(2) & 0 & 0.17604(4) & 0 & 0.0057(2) \\
M(3) & 0 & 0 & 0 & 0.0077(3) \\
M(4) & 0 & 0.28018(2) & 1/2 & 0.01024(15) \\
A(2) & 0 & 0.4875(2) & 0 & 0.0090(16) \\
A(m) & 0.5374(6) & 0 & 0.0767(12) & 0.0194(15) \\
O(1) & 0.10487(11) & 0.08861(6) & 0.2156(2) & 0.0102(2) \\
O(2) & 0.11900(11) & 0.17359(6) & 0.7351(2) & 0.0089(2) \\
O(3) & 0.10711(16) & 0 & 0.7151(3) & 0.0109(4) \\
O(4) & 0.36742(12) & 0.25134(6) & 0.7923(2) & 0.0111(2) \\
O(5) & 0.35041(11) & 0.14023(7) & 0.1135(2) & 0.0115(2) \\
O(6) & 0.34387(11) & 0.11779(7) & 0.6116(2) & 0.0121(2) \\
O(7) & 0.33855(18) & 0 & 0.2847(3) & 0.0150(3) \\
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**AM8**
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M(1) & 0 & 0.08991(4) & 1/2 & 0.0083(2) \\
M(2) & 0 & 0.17608(3) & 0 & 0.0047(2) \\
M(3) & 0 & 0 & 0 & 0.0074(3) \\
M(4) & 0 & 0.28086(2) & 1/2 & 0.00995(14) \\
A(2) & 0 & 0.4809(3) & 0 & 0.0294(17) \\
A(m) & 0.5379(9) & 0 & 0.0805(19) & 0.025(2) \\
O(1) & 0.10402(11) & 0.09017(6) & 0.2124(2) & 0.0100(2) \\
O(2) & 0.11801(11) & 0.17418(6) & 0.7411(2) & 0.0092(2) \\
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TABLE 4. Chemical composition (wt.%) and unit formula (apfu) for gem amphiboles from Mogok, Myanmar.

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apfu

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| M(3) | 1 | 0.79 Mg + 0.17 Al + 0.04 Fe$^{2+}$ | 1.00 | 12.80(6) | 12.73 | 2.057 |
| 2 | 0.96 Mg + 0.01 Al + 0.03 Fe$^{2+}$ | 1.00 | 12.32(7) | 12.43 | 2.074 |
| 3 | 0.77 Mg + 0.19 Al + 0.04 Fe$^{2+}$ | 1.00 | 12.36(7) | 12.75 | 2.055 |
| 6 | 0.93 Mg + 0.05 Al + 0.02 Fe$^{2+}$ | 1.00 | 12.34(7) | 12.33 | 2.070 |
| 7 | 0.84 Mg + 0.11 Al + 0.05 Fe$^{2+}$ | 1.00 | 12.64(8) | 12.81 | 2.068 |
| 8 | 0.86 Mg + 0.12 Al + 0.02 Fe$^{2+}$ | 1.00 | 12.27(7) | 12.40 | 2.074 |

| M(4) | 1 | 1.97 Ca + 0.03 Na | 2.00 | 39.69(11) | 39.73 | 2.485 |
| 2 | 2.00 Ca | 2.00 | 39.97(11) | 40.00 | 2.472 |
| 3 | 2.00 Ca | 2.00 | 40.14(11) | 40.00 | 2.485 |
| 6 | 2.00 Ca | 2.00 | 40.60(10) | 40.00 | 2.482 |
| 7 | 2.00 Ca | 2.00 | 40.38(11) | 40.00 | 2.485 |
| 8 | 2.00 Ca | 2.00 | 40.15(11) | 40.00 | 2.472 |

| A(m) | 1 | 0.48 Na + 0.06 K | 0.54 | 6.51(2) | 6.42 | 2.555 |
| 2 | 0.04 Na + 0.28 K | 0.32 | 5.33(2) | 5.76 | 2.611 |
| 3 | 0.40 Na + 0.05 K | 0.45 | 5.55(3) | 5.35 | 2.589 |
| 6 | 0.07 Na + 0.44 K | 0.51 | 7.98(3) | 9.13 | 2.631 |
| 7 | 0.05 Na + 0.42 K | 0.47 | 7.29(3) | 8.53 | 2.633 |
| 8 | 0.16 Na + 0.18 K | 0.34 | 5.16(3) | 5.18 | 2.589 |

<p>| A(2) | 1 | 0.46 Na | 0.46 | 5.56(2) | 5.06 | 2.646 |
| 2 | 0.68 Na | 0.68 | 7.09(3) | 7.48 | 2.720 |
| 3 | 0.55 Na | 0.55 | 5.95(2) | 6.05 | 2.620 |
| 6 | 0.49 Na | 0.49 | 5.36(2) | 5.39 | 2.767 |
| 7 | 0.53 Na | 0.53 | 5.82(2) | 5.83 | 2.759 |
| 8 | 0.66 Na | 0.66 | 7.09(3) | 7.26 | 2.694 |</p>
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