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Short-range order in synthetic aluminous tremolites: An infrared and triple-quantum MAS NMR study

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ABSTRACT

Fourier-transform infrared spectra (FTIR) were recorded on a series of synthetic amphiboles along the join Ca1.8Mg5.2Si8O22(OH)2–Ca1.8(Mg4.2Al)(Si7Al)O22(OH)2. The spectra were fitted by up to six component bands by optimization and non-linear least-squares techniques. 27Al MAS NMR and triple-quantum (3Q) MAS NMR spectra were recorded for the synthetic amphibole Ca1.8(Mg4.8Al0.4)(Si7.6Al0.4)O22(OH)2. The fitted FTIR and NMR spectra show that VIAl occurs at the M2 site and at the M1 or M3 sites; by analogy with previous crystal-structure refinement results on natural amphiboles, VIAl occupancy of M2 and M3 is presumed. The fine structure present in the FTIR spectra indicates that they are also affected by NNN (next-nearest-neighbor) interactions. There are two types of NNN arrangements: (1) SiSi or SiAl at adjacent T1T1 dimers; (2) permutation of Mg/Al over M2M2M3 sites. Discounting those arrangements unlikely on bond-valence grounds, there are two arrangements that give rise to five distinct bands in the infrared spectra. There are two principal conclusions: (1) infrared spectra of amphiboles in the principal OH-stretching region can be affected significantly by NNN effects; (2) the small number of bands due to NNN effects indicates that amphiboles show strong short-range order.

INTRODUCTION

The composition of calcic amphiboles, and the Al content in particular, has great potential in geobarometry of igneous and metamorphic processes. Both field evidence and experimental work suggest that Al content is correlated with the physical conditions (P and T) of amphibole formation (e.g., Spear 1981; Hammarstrom and Zen 1986; Hollister et al. 1987). This, together with the fact that amphiboles are ubiquitous in many geological environments (Hawthorne 1983), has prompted increased attention directed toward a more quantitative definition of Al solubility in both natural (Holland and Richardson 1979; Tribollet and Audren 1988; Kohn and Spear 1990; Mader and Berman 1992) and synthetic amphiboles (Jenkins 1988, 1994; Cho and Ernst 1991; Smelik et al. 1994). Two main problems have been encountered in these studies: (1) identifying the reaction exchanges responsible for Al incorporation in the amphibole structure; in calcic amphiboles, there is general consensus in considering the tschermak substitution as the main mechanism whereby one divalent cation and one Si are replaced by Al in both octahedral and tetrahedral coordination. (2) Thermodynamic modeling has been largely based on mixing models which did not take into account possible patterns of long-range order (LRO) and short-range order (SRO) involving substituting cations over non-equivalent sites. Most thermodynamic analyses have considered activity models based solely on Mg/Al ordering at M2 and Si/Al ordering at T1. Recent crystal-structure work has shown that these mixing models are inadequate. Long-range ordering patterns of Al in amphiboles are now quite well understood (Oberti et al. 1995a, 1995b), but we still know very little about SRO of Al. Infrared spectroscopy has potential with respect to SRO characterization in silicates, and there has been considerable effort recently to develop procedures to quantify SRO in synthetic amphiboles (Della Ventura 1993; Della Ventura et al. 1996a, 1996b, 1998; Hawthorne et al. 1996a, 1996b, 1997).

Here, we report Fourier-transform infrared (FTIR) and magic-angle spinning nuclear magnetic resonance (MAS NMR) spectroscopic studies of synthetic aluminous tremolites along the tremolite-tschermakite join, and show that IR spectroscopy in the OH-stretching region is a valuable tool in understanding particular features of Al incorporation in calcic amphibole.
**Materials and Methods**

The samples are described by Smelik et al. (1994) and Jenkins et al. (1997). Their nominal compositions, prepared by mixing appropriate amounts of reagent-grade chemicals, vary along the tremolite–magnesiohornblende join. In this series, all amphiboles have an empty A-site. Previous workers have reported a reproducible 5–10% enrichment of Mg at M4 in any attempt to synthesize end-member tremolite (Jenkins 1987; Graham et al. 1989; Pawley et al. 1993). Gottschalk et al. (1999) reported significant shift of amphibole compositions from the nominal stoichiometries. Electron-microprobe data also show that the synthesized amphiboles are on-composition, except for the more aluminous magnesiohornblende sample (AMPH 14–6), which is close to 90 mol% magnesiohornblende (Smelik et al. 1994; Jenkins et al. 1997).

**Infrared spectroscopy**

FTIR spectra in the OH-stretching region (Figs. 2 and 3) were collected using KBr disks (Robert et al. 1989). Spectra were recorded on a Nicolet 760 spectrophotometer equipped with a nitrogen-cooled InSb detector and a CaF2 beamsplitter, averaging 64 scans for a nominal resolution of 2 cm⁻¹. Spectra were fitted by interactive optimization followed by least-squares refinement according to the fitting procedure described in detail in Della Ventura et al. (1996a, 1998). A symmetrical Gaussian line-shape was used and all corresponding bands in the series were constrained to have the same widths as those derived from spectra in which the bands are prominent. The refined spectral parameters are given in Table 2. As is common with powder infrared spectra in the principal OH-stretching region, it is difficult-to-impossible to fit a baseline in an objective and yet persuasive manner. When dealing with narrow bands in a restricted energy range, this problem does not adversely affect the fitting procedure. However, when dealing with broad bands over a wide range of energy (as is the case here), a degree of uncertainty is introduced that is not necessarily reflected in the fit to the spectrum. For this reason, we do not attempt a quantitative interpretation of the fitted spectra.

**²⁷Al MAS NMR spectroscopy**

The ²⁷Al MAS NMR spectrum for AMPH 25–1 (A_{TOT} = 0.8) was obtained using a Chemagnetics CXP-400 multinuclear spectrometer operated at a frequency of 104.2 MHz. The MAS NMR experiment used very short (<10%) pulses and a 2 s pulse delay. The very short pulse length ensures that the spectrum is quantitatively reliable. Solid NaBr was used to set the magic angle accurately using the ⁷⁹Br NMR resonance. The sample was spun in nitrogen at 13.5 kHz and a spectral width of 40 kHz was used. A total of 1808 transitions were collected. ²⁷Al chemical shifts are reported relative to 1M Al(NO₃)₃ solution.

**Triple-quantum (3Q) ²⁷Al MAS NMR**

The triple-quantum (3Q MAS NMR) experiment uses three pulses: the first pulse excites the triple-quantum coherence (±3);

**Table 1. Nominal compositions (apfu) for the synthetic amphiboles**

<table>
<thead>
<tr>
<th>Sample</th>
<th>A</th>
<th>M4</th>
<th>M1, M2, M3</th>
<th>T1, T2</th>
<th>Total Al</th>
</tr>
</thead>
<tbody>
<tr>
<td>TREM 23-6</td>
<td>Ca₄Mg₅Si₈O₂₂(OH)₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMPH 25-1</td>
<td>Ca₄Mg₅Si₈O₂₂(OH)₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMPH 19-7</td>
<td>Ca₄Mg₅Si₈O₂₂(OH)₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMPH 28-1</td>
<td>Ca₄Mg₅Si₈O₂₂(OH)₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMPH 14-6</td>
<td>Ca₄Mg₅Si₈O₂₂(OH)₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
the second pulse converts the triple-quantum coherence back into observable magnetization ($\pm 1$); the third pulse equalizes the two coherence pathways (Z-filtering). Details are given in Baltisberger et al. (1996) and references therein. The three pulses had lengths of 3.3 $\mu$s, 1 $\mu$s, and 8 $\mu$s, respectively, and a 50 s pulse delay was used. A total of 96 scans per increment was required for $^{27}$Al 3Q NMR. To increase peak intensity and shorten the run time, rotor synchronization was used to eliminate spinning sidebands, and the interval between the two first pulses was incremented at 71.26 ms (corresponding to an MAS rate of 14.0 KHz). Under these conditions, 128 increments in $t_1$ were sufficient with 96 scans and a 50 s pulse delay. The 3Q MAS NMR experiment lasted about 3 days.

**Figure 2.** Infrared spectra in the principal OH-stretching region for synthetic amphiboles nominally along the join tremolite-magnesiohornblende.

**Figure 3.** Fitted infrared spectra in the principal OH-stretching region for synthetic amphiboles nominally along the join tremolite-magnesiohornblende. The component bands are vertically displaced for clarity; observed intensities are shown by hollow squares, and the line following the observed intensities is the envelope of the sum of the fitted component bands.

**Configuration notation**

In the amphibole structure, the OH group is directly bonded to two M1 and one M3 cations, and the H atom projects into the A-site cavity in which the A-site may be occupied or vacant (□). The NN (nearest-neighbor) configuration directly around the OH group may thus be represented by the general symbol M1M1M3–OH–A; for ordered end-member tremolite and pargasite, the symbols are MgMgMg–OH–A and MgMgMg–OH–Na, respectively. The M1M1M3 cluster is surrounded by two M2 and one M3 octahedra within the octahedral sheet (Fig. 4) and two T1 tetrahedra within the tetrahedral double-chain. The triplet of cations at M2M3M2 and the
doublet at T1T1 are thus NNN (next-nearest neighbor) cations to O3. The NNN configuration around the O3 group may be denoted by adding the cation doublet and triplet to the general symbol introduced above: M1M1M3–O3– AA:T1T1–M2M3M2. Note that the NN and NNN octahedra within the octahedral sheet have pseudo-trigonal point symmetry. For ideally ordered end-member tremolite and pargasite, the configuration symbols are MgMgMg–OH–ANa:SiAl–MgMgAl, respectively.

**POSSIBLE ARRANGEMENTS OF NN AND NNN CATIONS**

The NN and NNN arrangements of Figure 4a may be abstracted to the graph shown in Figure 4b. Possible NN and NNN cation arrangements may be derived by coloring the vertices of this graph, where the colors represent occupancy of the sites by different species of cations. The resulting graphs are shown in Figure 5. Some of the configurations shown in Figure 5 will not occur (or be very energetically unfavorable) due to significant deviation from the valence-sum rule (Brown 1981; Hawthorne 1992, 1994, 1997). For example, Hawthorne (1997) has shown that, in pargasite, the preferred configuration from a bond-valence viewpoint involves T1Si–T1Al linked to M2MgM3MgM2Al [or M2MgM3AlM2Mg], effectively preventing the occurrence of configurations involving more than one Al cation at adjacent M2M3M2 octahedra in the structure. The optimum arrangements from a bond-valence viewpoint are identified in Figure 5.

**MAS NMR**

The 27Al MAS NMR spectrum of AMPH 25–1 (Fig. 6) has three distinct maxima at 66.4, 3.8, and −3.1 ppm, respectively. The peak at 66.4 ppm is due to 3Al, whereas those at 3.8 and −3.1 ppm are due to 4Al (Welch et al. 1994; Jenkins et al. 1997). The 4Al component of the spectrum extends from about 10 to −50 ppm and clearly exhibits some quadrupolar character. Weak spinning sidebands occur at about δ±130 ppm. The spectrum is fully quantitative, as shown by the equal intensities (peak areas) of the 3Al and 4Al (total) components of the MAS NMR spectrum, as expected for Mg-tschermak exchange. This spectrum is essentially identical to that obtained by Jenkins et al. (1997) for AMPH 25–1, allowing for the different fields used (9.4 T vs. 11.7 T). These workers found that the 3.8 ppm peak has η = 0, whereas the −3.1 ppm peak has η = 1. These values are consistent with 4Al sites having high and low symmetries, respectively. M1 and M2 sites in C2/m amphiboles have symmetry 2, M3 has symmetry 2/m. Hence, the 3.8 ppm peak (low η) is reasonably assigned to M3 and the −3.1 ppm peak is due to M1 or M2. Below, we discuss the choice between M1 and M2.

The 3Q NMR spectrum has an isotropic component that allows Al peaks (Al sites) that overlap in the MAS spectrum to be resolved. This is usually not possible for the single-pulse
spectrum because of significant quadrupolar components of the spectrum that lead to considerable peak overlap, as we observe for $^{27}$Al. The 3Q spectrum (Fig. 7) clearly resolves two distinct $^{27}$Al peaks (FWHM = 2 ppm) with isotropic shifts of −44 and −59 ppm, respectively.

**INFRARED SPECTRA**

In this series, the spectrum of end-member tremolite (Fig. 2a) represents an obvious reference point as it is now well-characterized (Hawthorne et al. 1996a, 1997). It consists of a single very sharp absorption band (full-width at half-height maximum, FWHM, is in the range 3–4 cm$^{-1}$) at 3675 cm$^{-1}$. This band is assigned to the vibration of an O–H dipole bonded to three $^{27}$Mg cations and directed along $a^*$ toward an empty A-site: MgMgMg–OH–A$^\text{II}$:SiSi–MgMgMg, where $A^\text{II}$ represents a vacancy. All natural and synthetic tremolites show this band to be asymmetric toward the low-frequency side, suggesting the presence of a second component at around 3670 cm$^{-1}$. There is strong evidence (Hawthorne et al. 1996a, 1997) that this second band is due to Mg at M4 (a cummingtonite component) that seems ubiquitous in synthetic tremolite (Jenkins 1987; Ahn et al. 1991; Pawley et al. 1993; Maresch et al. 1994; Gottschalk et al. 1998, 1999). The intensity of this band in the spectrum of Fig. 2a (~10% of the total absorption) is more-or-less in accord with the stoichiometry of the tremolite sample examined here.

The introduction of Al into the tremolite structure correlates with the appearance of two more absorption features (labeled B and C in Fig. 2b to 2e), the intensities of which increase as a function of increasing Al. As is evident in the spectrum of Fig. 2b, the B and C absorptions are both doublets at 3655 + 3642 and 3624 + 3608 cm$^{-1}$. With increasing Al, the relative intensities of the B and C absorptions increase and the individual components of each doublet in Fig. 2b become less apparent. Furthermore, the very sharp band at 3675 cm$^{-1}$ in the spectrum of tremolite broadens considerably and shifts to slightly lower wavenumber.

**EFFECTS OF NNN CATIONS**

Hawthorne et al. (1996a) have shown that variation in the NNN (next-nearest-neighbor)-cation occupancy can also give rise to band splitting in the infrared spectra in the principal OH-stretching region. Pertinent in this regard for the amphib-
oles of this study are the adjacent T1 sites, which may be occupied by SiSi or SiAl, and the M2M3M2 triplet of sites (Fig. 4), which may be occupied by Mg and Al (Fig. 5).

Si and Al at the T sites

In the Al-bearing amphiboles of this work, half the constituent Al occurs at the T sites, but details of the relative ordering of Al and Si over the T1 and T2 sites are not known. By analogy with most natural amphiboles, Al should be ordered at the T1 site. However, natural and synthetic high-temperature amphiboles can show considerable disorder of Al over T1 and T2 (Oberti et al. 1995b; Welch et al. 1998; Welch and Knight 1999), but the intermediate synthesis temperatures and unusual composition of the synthetic amphiboles examined here make direct comparison ambiguous. Jenkins et al. (1997) proposed complete disorder of IVAl over the T1 and T2 sites, based on 29Si MAS NMR spectra; however, their calculations assume no SRO, a situation that the arguments of Hawthorne (1997) indicate to be most unlikely. Della Ventura et al. (1998) have proposed a mechanism for the interaction between OH at O3 and the O7 atom bridging adjacent T1 tetrahedra. The O7 atom acts as a hydrogen bond acceptor for the OH group at the adjacent O3 site (Fig. 8). When the O7 atom links to Si only [Si-O7-Si], the hydrogen bond is weak; when O7 links to Si and Al [Si-O7-Al], the hydrogen bond is much stronger and hence the principal stretching frequency of the OH group shifts to lower wavenumber. If Al were to occur at the T2 site, the H-atom would form a hydrogen bond with the O5 atom linking the T1 and T2 sites [Si-O5-Al], and hence the shift to lower wavenumber will be similar to that observed when Al occurs at the T1 site. Hence the spectra are unlikely to be affected significantly by any Al disorder over T1 and T2. For simplicity, we will formulate our model assuming all IVAl is ordered at the T1 site in these amphiboles, recognizing that neither the model nor the results should be significantly affected by any disorder of IVAl over T1 and T2.

Mg and Al at the M1,2,3 sites

Previous infrared and MAS NMR work on pargasite and M3+-substituted pargasites (Semet 1973; Raudsepp et al. 1987, 1991; Welch et al. 1994) has shown that the trivalent cations in these amphiboles are not completely ordered at M2 but also occur at M1 and/or M3. Oberti et al. (1995a) have shown by single-crystal structure refinement that Al occurs only at the M2 and M3 sites in pargasitic amphiboles. In a neutron powder-diffraction study of synthetic end-member pargasite, Welch and Knight (1999) also found that IVAl occurs at M2 and M3, and not at M1. Our interpretation of the 3Q NMR spectrum of AMPH 25–1 is made in the context of recent diffraction and spectroscopic studies of this and closely related amphiboles (Welch et al. 1994; Jenkins et al. 1997; Oberti et al. 1995a; Welch and Knight 1999). Although the M1 and M2 sites have the same point symmetry, their ligancy is different: M1 is O4(OH)2, M2 is O6. Therefore, we would expect these two sites to have distinctly different isotropic shifts. Furthermore, given the large separation (15 ppm) between the isotropic shifts of the IVAl peaks in the 3Q NMR spectrum, we are confident that the M1 and M2 sites would be resolved in the experiment. There is no suggestion in the 3Q MAS spectrum of any additional isotropic components to either of these two peaks. Thus, the 3Q MAS spectrum shows conclusively that there are just two different IVAl sites, not three. Moreover, the X-ray and neutron-diffraction studies of natural and synthetic Mg-rich hornblendes and pargasites indicate that Al is ordered at M2 and M3 only. We conclude that the two IVAl peaks in the 3Q MAS spectrum are most reasonably assigned to Al at M2 and M3, with none at M1. Single-pulse experiments cannot provide such conclusive proof. Thus, in considering the ordering of Mg and Al over the NN and NNN cation-sites in the amphibole structure, Mg can occupy the M1, M2, and M3 sites, whereas Al can occupy only the M2 and M3 sites. These observed long-range ordering patterns constrain possible local configurations about the O3 site.

Band assignment

If magnesiohornblende, $\square$Ca$_2$Mg$_4$Al$_3$Si$_3$AlO$_{25}$(OH)$_2$, were to show the maximum long-range order possible, then M1 = M2 = Mg, M3 = Al; this arrangement will have two local configurations around the O3 site: MgMgAl–OH–A–SiSi–MgAlMg
and MgMgAl–OH−→SiAl–MgAlMg, and would show two bands in its infrared spectrum in the principal OH-stretching region. If magnesiohornblende were to have the long-range ordering pattern M1 = M3 = Mg, M2 = 0.5 Al + 0.5 Mg, this arrangement will have the following two local configurations around the O3 site: MgMgMg–OH−→SiSi–MgMgAl and MgMgMg–OH−→SiAl–MgMgAl, and would show two bands in its infrared spectrum in the principal OH-stretching region. Previous interpretations of the OH-stretching spectra of aluminous amphiboles (Semet 1973; Raudsepp et al. 1987, Della Ventura et al. 1998) have assigned two broad bands to the spectrum (Fig. 2). It is instructive to consider the spectrum to slightly lower frequency, but still remains a major feature of their spectrum.

**MgMgAl–OH−→SiSi–MgMgMg** and **MgMgAl–OH−→SiAl–MgMgAl**, and would show two bands in its infrared spectrum in the principal OH-stretching region. Previous interpretations of the OH-stretching spectra of aluminous amphiboles (Semet 1973; Raudsepp et al. 1987, Della Ventura et al. 1998) have assigned two broad bands to this band is extremely sharp, whereas in amphiboles with significant Al, it splits into two components, one of which shifts to lower frequency, but still remains a major feature of the spectrum. Inspection of Fig. 2 indicates that there are additional bands in the spectra presented here, together with fine structure in the major bands, indicating that there is more disorder in these amphiboles than an ideal ordered model would indicate (Della Ventura et al. 1998).

**BANDS A AND A’**

The A band is the only feature (neglecting the minor cummingtonite component) in the spectrum of tremolite. It is notable that substitution 3 does not produce an additional band in the infrared region. If magnesiohornblende were to have the long-range ordering spectrum (Fig. 2). It is instructive to consider the spectrum to slightly lower frequency, but still remains a major feature of the spectrum. Previous interpretations of the OH-stretching spectra of aluminous amphiboles (Semet 1973; Raudsepp et al. 1987, Della Ventura et al. 1998) have assigned two broad bands to this band is extremely sharp, whereas in amphiboles with significant Al, it splits into two components, one of which shifts to lower frequency, but still remains a major feature of the spectrum. Inspection of Fig. 2 indicates that there are additional bands in the spectra presented here, together with fine structure in the major bands, indicating that there is more disorder in these amphiboles than an ideal ordered model would indicate (Della Ventura et al. 1998).

**BANDS B, B’, C, AND C’**

These bands must be assigned to configurations involving Al. There are four distinct substitutions involving Al in these amphiboles: (1) VAl → MgAl at the NN site; (2) IVAl → IVSi; (3) IVAl → IVMg; and (4) IVAl → VMMg at NNN sites. However, substitutions 3 and 4 are pseudosymmetrically equivalent, and hence will not give rise to different bands. Thus we cannot distinguish between M2M3M2 = MgMgAl and MgAlMg. Substitution 1 involves a change in bond-valence of 0.17 valence units (v.u.) at O1, a NN anion to O3, whereas substitution 2 involves a change of 0.25 v.u. at O1, hence substitution 2 is expected to cause a larger frequency shift than substitution 3. From the results of Raudsepp et al. (1987), we may make the following statement: replacement of Mg by Al at M3 (i.e., substitution 1) causes a downward frequency shift of ~30–35 cm⁻¹. The results of Hawthorne et al. (1996a) indicate that replacement of Si by Al at T1 (i.e., substitution 2) causes a downward frequency shift of ~20 cm⁻¹. These two results provide us with a basis for assigning bands in the spectra of Figure 2.

**Band A** at 3675 cm⁻¹ is MgMgMg–SiSi–MgMgMg; substitution 2 should produce a band corresponding to MgMgMg–SiAl–MgMgMg at 3655 cm⁻¹, which corresponds to band B (Table 2); substitution 1 should produce a band, corresponding to MgAlMg–SiSi–MgMgMg at 3645 cm⁻¹ which corresponds reasonably well with band B' (Table 2); combination of substitutions 1 and 2 should produce a band corresponding to MgMgMg–SiAl–MgMgMg at 3625 cm⁻¹, corresponding to band C (Table 2).

The remaining bands should then be due to substitution 3, which should produce smaller shifts than substitutions 1 and 2 (see above discussion). Table 3 shows the resulting band assignments. It is notable that substitution 3 does not produce a constant shift in frequency: from configurations 1 → 2, 3 → 4, 5 → 6, and 7 → 8, the frequency shifts are 4, 13, 17, and 17 cm⁻¹, respectively, for the VAl → VMMg substitution. Apparently the local relaxations associated with this substitution distribute themselves differently in the different local configurations.

**LONG-RANGE ORDER IN AL-BEARING AMPHIBOLES**

Recent results of X-ray crystal-structure refinement have established general patterns of long-range order of IVAl in calcic amphiboles (Oberti et al. 1995b): when IVAl ≤ 2 apfu, it is completely ordered at T1 in most metamorphic occurrences and partly disordered in some higher temperature igneous parageneses. The amphiboles of this study were synthesized in the temperature range 1096–1133 K (823–860 °C), and IVAl is likely to be ordered predominantly at T1.

Regarding VAl, Raudsepp et al. (1987) and Welch et al. (1994) have shown that VAl cannot be completely ordered at the M2 site in synthetic pargasite, and Oberti et al. (1995a) and Welch and Knight (1999) have shown that VAl is partly disordered over the M2 and M3 sites in Mg-rich pargasite. It is apparent from the infrared results given here and the NMR results of Jenkins et al. (1997) that VAl in magnesiohornblende is also disordered over the M sites. Indeed, comparison of the infrared spectra obtained here with the analogous infrared spectra of the synthetic potassium-richetterite-pargasite series of Della

**TABLE 3.** Possible configurations around the OH group in synthetic amphiboles of the tremolite-magnesiohornblende series, together with the observed band frequencies

<table>
<thead>
<tr>
<th>No.</th>
<th>Configuration</th>
<th>Observed</th>
<th>Band frequency (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[MgMgMg]–SiSi–MgMgMg</td>
<td>3675</td>
<td>A</td>
</tr>
<tr>
<td>2</td>
<td>[MgMgMg]–SiSi–MgMgAl</td>
<td>3671</td>
<td>A'</td>
</tr>
<tr>
<td>3</td>
<td>[MgMgMg]–SiAl–MgMgMg</td>
<td>3655</td>
<td>B</td>
</tr>
<tr>
<td>4</td>
<td>[MgMgMg]–SiAl–MgMgAl</td>
<td>3642</td>
<td>B'</td>
</tr>
<tr>
<td>5</td>
<td>[MgMgAl]–SiSi–MgMgMg</td>
<td>3642</td>
<td>B</td>
</tr>
<tr>
<td>6</td>
<td>[MgMgAl]–SiSi–MgMgAl</td>
<td>3625</td>
<td>C</td>
</tr>
<tr>
<td>7</td>
<td>[MgMgAl]–SiAl–MgMgMg</td>
<td>3625</td>
<td>C</td>
</tr>
<tr>
<td>8</td>
<td>[MgMgAl]–SiAl–MgMgAl</td>
<td>3608</td>
<td>C'</td>
</tr>
</tbody>
</table>

*M2M3M2 = MgMgAl and MgAlMg are indistinguishable.*
Ventura et al. (1998) suggests that VILAl behaves in a similar fashion in both series, i.e., partly disorders over the M2 and M3 sites.

**Quantitative considerations**

Hawthorne et al. (1996b) discussed some of the issues affecting band intensity in the principal OH-stretching region of the infrared. A key factor in this regard is the variation in transition moment as a function of transition energy; unless its relation is known, band intensities cannot be converted accurately to site populations. This relation is not well-known for amphiboles. It has been measured by Skogby and Rossman (1991) for single crystals of miscellaneous amphiboles, and by Hawthorne et al. (1997) for powder spectra of the series potassicrichterite-tremolite. In addition, Della Ventura et al. (1996b) have shown that it is not a factor in powder spectra of synthetic (Ni,Mg,Co)-potassicrichterite. Thus it is apparent that we do not yet understand this behavior sufficiently to derive accurate site-populations.

Another issue affecting derivation of site populations (Hawthorne et al. 1995) for the amphiboles examined here relates to accidental band degeneracy. As indicated in Table 3, five bands [(A + A'), B, B', C, C'] are caused by eight configurations. According to the configurations listed in Table 3,

\[ T1\text{Al} = 2(I^5 + I^6 + I^7 + I^8) / \sum_{n=1}^{8} I^n \]

\[ M3\text{Al} = 2(I^5 + I^6 + I^7 + I^8) / \sum_{n=1}^{8} I^n \]

According to the band assignment of Table 3, band B' corresponds to configurations 4 and 5, and hence we do not know \( I^4 \) or \( I^5 \), we only know \( I^4 + I^5 \). This means that we cannot calculate \( M3\text{Al} \) or \( T1\text{Al} \) as \( I^4 \) or \( I^5 \) only appears in the numerators of the above equations.

**SHORT-RANGE ORDER IN MAGNESIOHORNBLENDE**

The number of component bands observed in the infrared spectra corresponds closely to the number of bands predicted on the basis of local configurations that most closely satisfy the valence-sum rule (Hawthorne 1997), indicating that NNN arrangements in amphiboles can significantly affect the principal OH-stretching frequency. Furthermore, this correspondence provides experimental support for the prediction of Hawthorne (1997) that amphiboles are strongly short-range ordered, and that this ordering is induced by short-range (local) bond-valence requirements.

**THE OCCURRENCE OF VILAl AT THE M3 SITE**

As noted above, Oberti et al. (1995a) showed that VILAl is partly disordered over the M2 and M3 sites in Mg-rich pargasite; Welch and Knight (1999) found the same pattern of order in synthetic end-member pargasite. Hawthorne (1997) suggested that the occurrence of VILAl at M2 and VILAl at T1 are locally connected via short-range bond-valence requirements of the anions common to both sites; this short-range arrangement is shown diagrammatically in Figure 9. When the O1 (and O1') anion is linked to M2Mg and T1Si, the incident bond-strength at this anion is 1/3 × 3 + 1 = 2 v.u. When the same anion is linked to M2Al and T1Al (as for anions O1'' and O1''' in Fig. 9a), the incident bond-strength is 1/3 × 2 + 1/2 + 3/4 = 1.92 v.u., also close to the ideal value of 2 v.u. However, the local arrangement in Figure 9a involves the two Si-Al arrangements being related (in projection) by a horizontal mirror plane; there is no space-group constraint on this arrangement, and it is suscep-
tible to short-range order-disorder. The alternative arrangement is shown in Figure 9b, in which the two Si-Al arrangements are now related (in projection) by a twofold rotation axis orthogonal to the figure (Fig. 9b). If $^{VI}$Al occurs at M2, the bond-valence sum incident at O1 (2 v.u.) and O1" (1.92 v.u.) are in accord with the analogous values in Figure 9a, whereas the bond-valence sums incident at O1' and O1"" are 1/3 x 3 + 3/4 = 1.75 v.u. and 1/3 x 2 + 1/2 + 1 = 2.17 v.u., respectively. The large bond-valence deficiency at O1' can be alleviated by occurrence of $^{V}$Al at M3 rather than M2 (Fig. 9c), producing bond-valence sums of 2.17 v.u. at O1 and O1"" and 1.92 v.u. at O1' and O1". Thus the chain configurations above and below the octahedral strip seem to be implicated in the occurrence of Al-Mg disorder over the M2 and M3 sites.

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