Mutual exsolution in hornblende and cummingtonite: Compositions, lamellar orientations, and exsolution temperatures

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ABSTRACT

Exsolved pairs of hornblende and cummingtonite from two localities in southern New England, U.S.A., have been studied using transmission electron microscopy (TEM) to measure the crystallographic orientation of the exsolution lamellae. Both the cummingtonite and the hornblende show multiple generations or stages of exsolution. Electron microprobe analyses gave compositions of the pre-exsolved (averages) and the coarsest exsolved amphiboles.

In amphiboles, two orientations of lamellae are usually present and nearly parallel to {100} and {10T} of the host. The observed lamellar orientations are consistent with orientations predicted by the optimal phase-boundary (OPB) theory and are a function of small differences in a, c, and β of the host and exsolved amphiboles (Δa = a_host − a_amphibole, Δβ = β_host − β_amphibole, and Δc = c_host − c_amphibole). Because these lattice parameters vary with T and composition, the precise orientations of the lamellae are controlled by these variables. The widths of determined exsolution lamellae varied from microscopic (micrometer size) to submicroscopic (nanometer size). The TEM images show, in some cases, multiple generations of lamellae with lamellar orientations near {10T} differing up to 4°, whereas orientations near {100} differed up to 4°. For two pairs of coexisting amphiboles, the T dependence of the lattice parameters from 25 to 600 °C was measured using a Guinier camera. The OPB calculations indicate that for all the samples the different lamellar generations formed between about 780 and 300 °C (+50 °C, on the basis of maximum errors of the lattice-parameter determinations). Lamellae of the same generation all showed nearly the same width. Exsolution temperatures could not be derived from the “100” lamellae because relatively small variations in Δc, which controls exact orientation of the “100” lamellae, could not be measured accurately enough.

INTRODUCTION

A miscibility gap between the clinoamphiboles cummingtonite and hornblende was first proposed by Asklund (1923) and verified in 1962 (Asklund et al. 1962). Numerous workers have reported exsolution phenomena in coexisting cummingtonite and hornblende in amphibolite rocks from diverse geologic settings (e.g., Seitasaari 1952; Vernon 1962; Robinson 1963; Binns 1965; Boriani and Minutti 1965; Calleghari 1966; Jaffe et al. 1968; reviewed in Robinson et al. 1982a). In clinoamphiboles (space group C2/m), the host and the exsolved phases were commonly thought to share the {100} or {10T} planes. Using X-ray data, Ross et al. (1969) and Robinson and Jaffe (1969) found discrepancies between the actual and assumed lamellar orientations but could not interpret them. Later, Robinson et al. (1971) showed that these phase boundaries are not exactly parallel to the {100} or {10T} planes but are actually irrational planes close to {100} or {10T}. In the present study, these plane orientations are designated as “100” and “10T”. Bollman and Nissen (1968) termed such irrational planes the optimal phase boundaries. The “100” and “10T” orientations correspond to a minimum of lattice misfit and interfacial energy. Assuming nearly identical b lattice parameters for two intergrown phases, Robinson et al. (1971, 1977) showed that in hornblende-cummingtonite and augite-pigeonite pairs the orientations of exsolution lamellae are restricted to two exact phase boundaries in the a-c plane. These exact phase boundaries correspond to two vectors in the a-c planes of each lattice and that have the same lengths (see appendix of Robinson et al. 1977). The lengths of the vectors and, therefore, the orientations of these boundaries are functions of the differences in the lattice parameters (Δa, Δc, and Δβ) of the amphibole lamellae and host. Further discussion of optimal phase-boundary theory and application of the theory to other minerals can be found in Fleet (1982, 1985).
Recent work by Yang and Hirschmann (1995) suggests that Mg-rich clinoamphiboles, like the clinopyroxenes, show a phase transition from the $C$-centered to the primitive monoclinic unit cell in compositions at geologically relevant temperatures. However, in comparison with pyroxenes, there are much smaller changes in the amphibole lattice parameters both as a result of the phase transition and as a function of temperature (for pyroxenes, see Smith 1969, Cameron et al. 1973, Smyth 1974, Robinson et al. 1977; for amphiboles, see Cameron and Papike 1979, Yang and Hirschmann 1995). The exsolution processes in both mineral pairs are analogous, but because of the conversions used to assign group designations ($I2/m$ vs. $C2/m$) the approximate orientations of the exsolution planes are indexed differently: near $(100)$ and $(001)$ in augite and pigeonite, and near $(10\bar{1})$ and $(100)$ in hornblende and cummingtonite. The amphiboles that were studied here have space group $C2/m$. Discussion of the relationships between the $C2/m$ unit-cell setting and the $I2/m$ unit-cell setting can be found in Whittaker and Zussman (1961), Jaffe et al. (1968), Thompson (1978), and Robinson et al. (1982a, p. 60, Fig. 26).

Robinson et al. (1977) devised a geothermometer for exsolved monoclinic pyroxenes (augite and pigeonite) on the basis of the orientation of the exsolution lamellae and Fe-Mg content. The present study aims to determine if exsolved cummingtonite and hornblende can also be used as a geothermometer that would give temperatures of exsolution.

**Geologic setting of samples**

The exsolved cummingtonite and hornblende that were used in this study are from the amphibolites of Middle Ordovician rocks of the Partridge Formation and the Ammonoosuc Volcanics (Schumacher 1988; Hollocher 1993) from central Massachusetts and southwestern New Hampshire (Fig. 1). Metamorphism and structural development of this region occurred mainly during the Devonian Acadian orogeny, during which plutonism, nappetype tectonism, and syntectonic metamorphism that was followed by the formation of gneiss domes all occurred (e.g., Robinson 1986; Robinson et al. 1986, 1991).

Sample localities (Fig. 1) lay (1) within the Bronson Hill anticlinorium along the Massachusetts-New Hampshire border (samples 7A7C, 7A7D, 7A7R, and 7A7W, referred to here as the 7A7 series) and (2) on the eastern flank of the Bronson Hill anticlinorium in central Massachusetts (samples Q-603C, Q-798M, Q-J44R, WN27A, WD-H26C, and HFW-307W, designated as the CM series). For the rocks of the 7A7 series, the pelitic schists in this area yielded temperature estimates of 550-640 °C on the basis of garnet-biotite geothermometry (Tracy et al. 1976) and a minimum pressure estimate of 5-6.2 kbar on the basis of quartz + sillimanite + garnet assemblages (Robinson et al. 1982b). Peak metamorphic conditions for the CM series range between 635 and 675 °C and at least 5.5 and 7.2 kbar (Hollocher 1985).

**Experimental techniques**

The amphiboles were studied using transmission electron microscopy (TEM) and electron microprobe (EMP) analysis at Kiel University. The TEM samples were drilled out of standard petrographic thin sections with the use of an ultrasonic disc cutter and prepared by ion milling. Electron microscopy was performed with a Philips EM400T operating at 100 kV and equipped with an energy-dispersive X-ray analyzing system (Kevex, Unispec System 7000).

Heating experiments were performed with a Guinier film camera (Huber) at 50 kV and 20 mA using CuK$\alpha_1$ radiation. For Guinier camera measurements, amphibole powder was placed in quartz capillaries, which were closed after filling to minimize the amphibole oxidation and measured at various temperatures up to about 600 °C. Note that the prismatic habit of amphiboles led to their partial preferred orientation which reduced the accuracy of c-axis determinations. An internal silicon standard was used to correct for film shrinkage. T-dependent lattice parameters were determined by least-squares using the computer program FINAX (Hovestreydt 1983). For each sample, 11 reflections were used for lattice-parameter determination. Quantitative chemical analyses of amphibole were performed on a Camebax-MBX (Cameca) electron microprobe operated at 15 kV and 15 mA. Natural mineral standards were used.

**Mineral compositions**

The minerals of the assemblages were analyzed with the EMP. The EMP work concentrated on (1) the amphibole bulk chemistry and (2) the compositions of the coarsest exsolved end-members. In all ten samples the
bulk amphibole compositions were calculated from 60 to 90 point analyses collected in 1 μm steps across the grain perpendicular to one set of lamellae. These point analyses are mostly overlapping analyses. Because the diameter of the electron beam was about 2 μm, only coarser exsolution lamellae could be measured. About 10–20 point analyses were taken to get an average composition of the more coarsely exsolved amphibole.

Ferric estimates

Because of the fine scale of the exsolution, Mössbauer and wet-chemical analysis for amphibole Fe$^{3+}$ contents were not possible. Estimates of Fe$^{3+}$ contents for optically homogeneous cummingtonite samples were obtained by normalizing the sum of cations, excluding Na and K, to \(15 \text{[15eNK; see also Stout (1972), Robinson et al. (1982a, p. 3–12), and Schumacher (1991) for discussion]}\). For optically homogeneous hornblende, normalizing the cation sum, excluding Ca, Na, and K, to \(13 \text{(13eCNK)}\) yields the maximum ferric estimate but precludes the presence of cummingtonite component. Because these hornblendes coexist with cummingtonite, this procedure certainly overestimates the ferric content of the hornblende. Alternatively, the 15eNK correction gives a minimum Fe$^{3+}$ estimate, but, on the basis of Fe-Mg partitioning with cummingtonite, the 15eNK correction appears to be too conservative. By varying the amounts of Na that were assigned to the M4 position, it was possible to arrive at formulas that gave reasonable Fe-Mg values, Na content at M4, and cummingtonite contents.

Mineral assemblages and compositions

In addition to the exsolved hornblende and cummingtonite, all samples bore plagioclase, biotite, and quartz. Samples 7A7R and 7A7C (Fig. 1) also contained garnet. In sample HFW-307W, orthopyroxene was present.

Figure 2 shows \(^{57}\text{Al} \text{Al} \) plotted against A-site occupancy (Fig. 2A), Ca per 23 O atoms (Fig. 2B), and \(X_{\text{Ca}}\) for cummingtonite and hornblende in sample 7A7W (Fig. 2C) (see Table 1). Each plot suggests Fe$^{3+}$ correction is reasonable because in each case all six data points lie essentially on the tie line connecting the exsolved amphibole end-members.

The microprobe data for all the analyzed amphiboles are compared in AFM diagrams (Fig. 3). Plagioclase in
most samples is zoned between about An_{32} and An_{40}. The plagioclase in the orthopyroxene-bearing sample (HFW-307W) is An_{33}, and the plagioclase in sample Q-J44R is An_{32}. Garnets in 7A7R and 7A7C were nearly homogeneous and similar and had compositions of Py_{15.3}Alm_{4.4} Sp_{3.5}Gr_{0.8} and Py_{15.2}Alm_{5.8}Sp_{3.4}Gr_{0.7}, respectively. The coexisting amphiboles of the CM series have a much wider range of X_Mg compared with those of the 7A7 series. Samples 7A7W and all samples of the CM series apparently are too rich in Mg and too poor in Al to coexist with garnet. The hornblends of samples HFW-307W and Q-J44R plot differently than the other hornblends. The hornblende of HFW-307W has a low Al content (9.5 wt% Al_2O_3), whereas the hornblende of Q-J44R shows a higher Na content (2.09 wt% Na_2O). Both of these chemical traits would cause the hornblende to plot more negatively on the projection (Fig. 3) and may reflect differences in the bulk chemistry of these two samples.

**Characteristics of the exsolution lamellae**

**Orientation of the lamellae**

In the petrographic microscope, all the clinoamphiboles showed extensive exsolution (Fig. 4). The coarsest visible lamellae are 2.5–1 μm wide and generally show a “100” orientation. The range of thicknesses for various exsolution orientations is given in Tables 2 and 3.

Samples of 7A7R, Q-603C, and Q-798M were oriented in the TEM so that the a*-c* plane lay perpendicular to the electron beam, and thus lamellae were exactly parallel to the electron beam. In this orientation, the angles between the lamellae and the a or c axis as well as the...
TABLE 2. Angles (°) between “100” lamella and the c axis of the host phase and the average lamella thickness for samples Q-603C, Q-798M, and 7A7R

<table>
<thead>
<tr>
<th></th>
<th>Q-603C</th>
<th>Q-798M</th>
<th>7A7R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avg. “100” Lam ∩ c (°)</td>
<td>2.7</td>
<td>3.1</td>
<td>3.0</td>
</tr>
<tr>
<td>Avg. lamella thickness (µm)</td>
<td>2.0</td>
<td>0.1</td>
<td>0.2</td>
</tr>
<tr>
<td>Avg. “100” Lam ∩ a (°)</td>
<td>n.m.</td>
<td>n.m.</td>
<td>2.0</td>
</tr>
<tr>
<td>Avg. lamella thickness (µm)</td>
<td>0.0</td>
<td>0.1</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Note: n.m. = not measured.

lamellar width could be accurately measured. The orientations and widths of lamellae correlate and define natural groups of lamellae, which we interpret to be different generations or stages of exsolution. These data are shown in Table 2 for the “100” lamella and the host’s c axis (“100” ∩ c) and in Table 3 for the “101” lamella and the host’s a axis (“101” ∩ a) for samples 7A7R, Q-603C, and Q-798M. In each case, the lamellae exsolved at higher temperatures are coarser than the lamellae exsolved at lower temperatures because of the correlation between diffusion rate and temperature. An obvious difference between the lamellar orientations of these samples is that the angle of the lamellae to the a or c axes of the hosts could not be measured in all cases, and these lamellae are identified by “n.m.” in Tables 2 and 3. Most of the lamellar generations that are listed in Tables 2 and 3 are based on the characterization of the lamellar generations. Not every measurable amphibole could be oriented with the electron beam parallel to the b axis in the TEM. As a consequence, angles between the lamellae and the host’s a and c axes are not known. However, several problems were inherent to the characterization of the lamellar generations. Not every measurable amphibole could be oriented with the electron beam parallel to the b axis in the TEM. As a consequence, the angle of the lamellae to the a or c axes of the hosts could not be measured in all cases, and these lamellae are identified by “n.m.” in Tables 2 and 3. Most of the lamellar generations that are listed in Tables 2 and 3 are based on the correlation between lamellar width and lattice orientation. Further, because only a very small area of the ion-milled sample could be observed in the TEM, it is still possible that additional lamellar generations are present elsewhere in these samples.

Interlamellar spacings of the lamellae assigned to different generations were measured for samples 7A7D and Q-603C (Table 4), and these spacings also support the existence of different lamellar generations (Schumacher et al. 1993). On the basis of all these data, sample 7A7R (7A7 series) showed at least three lamellar orientations.

TABLE 3. Angles (°) between “101” lamella and the a axis of the host phase and the average lamella thickness for samples Q-603C, Q-798M, and 7A7R

<table>
<thead>
<tr>
<th></th>
<th>Q-603C</th>
<th>Q-798M</th>
<th>7A7R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avg. “101” Lam ∩ a (°)</td>
<td>3.0</td>
<td>3.1</td>
<td>3.0</td>
</tr>
<tr>
<td>Avg. lamella thickness (µm)</td>
<td>2.4</td>
<td>0.1</td>
<td>0.2</td>
</tr>
<tr>
<td>Avg. “101” Lam ∩ a (°)</td>
<td>n.m.</td>
<td>n.m.</td>
<td>3.0</td>
</tr>
<tr>
<td>Avg. lamella thickness (µm)</td>
<td>3.0</td>
<td>0.1</td>
<td>0.1</td>
</tr>
</tbody>
</table>

Note: n.m. = not measured.

“101” lamellar widths range from 1.5 to 0.1 µm in cummingtonite and from 1.5 to 0.1 µm in hornblende. For sample Q-798M, the observed hornblende “101” lamellar widths range from 1.0 to 0.07 µm.

TABLE 4. Distances between “100” lamellae with similar thicknesses for samples 7A7D and Q-603C

<table>
<thead>
<tr>
<th></th>
<th>7A7D</th>
<th>Q-603C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avg. “100” lamella width (µm)</td>
<td>1.50</td>
<td>12.50</td>
</tr>
<tr>
<td>Avg. lamella thickness (µm)</td>
<td>1.05</td>
<td>10.5</td>
</tr>
</tbody>
</table>

Note: n.m. = not measured.
and sample Q-603C (CM series) showed at least six generations of exsolution lamellae.

**HRTEM images**

High-resolution TEM (HRTEM) images (Fig. 5) show the coherent intergrowths of the host and guest phases. Figure 5A is viewed down [010] onto the a-c plane, showing an intergrowth between hornblende host and “100” and “101” cummingtonite lamellae of sample WD-H26C. The interface between the host and the lamellae is very sharp, so in this orientation the electron beam is parallel to the lamellar interfaces. The “100” lamella shares the a* axis with the host, and the c axis is common for the lamella and host. The “100” lamella shows a bending of the (001) fringes. The {101} lamella shares the c* axis with the host, and the a axis is common for the lamella and host. Where the lamellae approach each other, one always tapers down to a point at the end, possibly because of strain-field interactions (Livi and Veblen 1989; Smelik and Veblen 1991). The “100” lamella shows bowing at the area were the two lamellae meet. The bowing feature is very common where lamellae approach one another (Smelik and Veblen 1993). A possible explanation is that the Ca in these regions diffuses to or away from the site of the “100” lamella because Ca diffusion is faster parallel to the Si double chains (parallel to the c axis).

Figure 5B shows the (110) adjustment of cummingtonite host and disc-shaped, “101” hornblende lamellae in sample Q-603C. The adjustment of the two coherent intergrown lattices by a slight rotation of the (110) lattice fringes is very conspicuous. Compositions of these lenticular lamellae and the host were compared qualitatively in the TEM by energy-dispersive X-ray analysis. A significant difference was detected between the Ca contents of the host and the lamellae. Additionally, electron spectroscopic imaging (ESI) of these disc-shaped exsolution lamellae (Czank et al. 1993) showed significant relative differences between the Ca contents of the guest and host phases. Hornblende lamellae were also found inside an approximately 2 μm wide, early cummingtonite lamella within a hornblende host (sample 7A7D). This feature is further evidence that these amphiboles actually underwent multiple phases of exsolution.

**Optimal phase-boundary calculations**

**Influence of lattice parameters on lamellar orientation**

According to optimal phase-boundary (OPB) calculations, the differences between the pairs of lattice parameters of the host and exsolved phase control the lamellar orientation (i.e., the temperature dependence of Δa, Δc, and Δβ, where Δa = a_{host} - a_{lamella}, Δc = c_{host} - c_{lamella}, and Δβ = β_{host} - β_{lamella}). To determine the contribution of the
parameters $\Delta a$, $\Delta c$, and $\Delta \beta$ to lamellar orientation, two parameters were held constant and the third was varied in the calculation (Fig. 6).

The “100” lamellar orientation is most strongly affected by the differences in the $c$ lattice parameter ($\Delta c$) of the coexisting clinoamphiboles (Fig. 6A). A difference of only 0.03 Å between the two $c$ lattice parameters of host and guest phase would change the relative orientation of the lamellae by $7^\circ$. Differences in the monoclinic angle ($\Delta \beta$) and $a$ lattice parameter ($\Delta a$) have a lesser effect on the orientation of the “100” lamella.

Figure 6C shows unambiguously that the “101” orientation is most strongly affected by $\Delta \beta$. The $\Delta a$ parameter has a smaller and opposite effect on the “101” orientation of the lamella (Fig. 6B). Variation of the $c$ lattice parameter has practically no influence on the orientation of the “101” exsolution lamella.

Variations in lattice parameters with temperature

The temperature dependence of the amphibole lattice parameters was obtained by measuring $d$ values with a Guinier film camera over a $T$ range of 25–600 °C (Fig. 7). Figures 7A and 7B show different $T$-dependent changes in the lattice parameters of samples 7A7R and Q-603C, respectively, and indicate that the relative rates of change were not uniform. As a result, upon cooling, coherency strain must increase between lamella and host, but no features that can be attributed to this strain were observed.

The data on the $T$ dependence of lattice parameters combined with the OPB theory allow the prediction of lamellar orientations for both the “101” and “100” lamellae at a certain temperature for samples 7A7R and Q-603C. However, the “100” lamellar orientation is clearly the least reliable of the two lamellar orientations because it strongly depends on the temperature variations of the $c$ lattice parameters (Fig. 6), which are relatively small and the least certain of the measurements (Fig. 7). As a consequence, the rest of the discussion focuses on the temperature dependence of the “101” lamellar orientations.

Variations in lamellar orientation with temperature

For the amphiboles in Q-603C, the angle between the “101” lamellae and the $a$ axis of the host increases with higher temperature, whereas, surprisingly, the same angle decreases with higher temperature in 7A7R (Table 3). This result is somewhat unexpected because the lattice parameters of all the amphiboles showed the same general changes with temperature (Fig. 7). This feature, however, results from different rates of change with temperature between the two samples in the $\Delta a$ and $\Delta \beta$ of the
Temperature dependence of amphibole lattice parameters for sample 7A7R (A) and sample Q-603C (B). Lattice-parameter regression functions are as follows, in order of T: For 7A7R, \( \alpha_{\text{Cum}}(T) = 1.574 \times 10^{-5}T + 9.481(5) \), \( \beta_{\text{Cum}}(T) = 1.807 \times 10^{-5}T + 5.311(2) \), \( c_{\text{Cum}}(T) = 1.807 \times 10^{-5}T + 5.311(2) \), \( a_{\text{Cum}}(T) = 9.3573 \times 10^{-5}T + 5.318(2) \), \( c_{\text{Cum}}(T) = 2.501 \times 10^{-5}T + 5.318(2) \), \( A_{\text{Cum}}(T) = -4.224 \times 10^{-5}T + 105.19(2) \); and for Q-603C, \( \alpha_{\text{Cum}}(T) = 1.014 \times 10^{-5}T + 9.510(6) \), \( \beta_{\text{Cum}}(T) = 3.652 \times 10^{-5}T + 5.321(2) \), \( c_{\text{Cum}}(T) = 3.652 \times 10^{-5}T + 5.321(2) \), \( a_{\text{Cum}}(T) = -4.593 \times 10^{-5}T + 101.87(2) \), \( c_{\text{Cum}}(T) = 3.826 \times 10^{-5}T + 5.330(2) \), \( A_{\text{Cum}}(T) = -8.580 \times 10^{-5}T + 105.25(2) \). Errors given are indicated by parentheses and are based on averages of all measurements at different temperatures.

Exsolution and exsolution temperatures

Both cummingtonite and hornblende begin to break down to pyroxenes and other anhydrous phases at upper amphibolite- to granulite-facies conditions. As a result, the crest and the flatter parts of their heterogeneous solvus are metastable. This in turn renders supersolvus hornblende-cummingtonite compositions metastable relative to the pyroxene-bearing assemblages. As a consequence, mutual miscibility of hornblende and cummingtonite is much more restricted, and it is much more unlikely that, upon cooling, the spinodal regions of the cummingtonite-hornblende solvus will be encountered.

The style of exsolution that is normally ascribed to spinodal decomposition (fine, wavy intergrowths) was not observed in this study. The exsolution textures that have been described above are consistent with the overstepping of a coherent solvus and the precipitation and growth of new lamellae.

The scenario proposed is that, initially, homogeneous and coexisting cummingtonite and hornblende cooled below the coherent solvus. Subsequent intervals of undercooling resulted in various stages of coherent exsolution until at lower temperatures the kinetics became too sluggish for exsolution to continue. To reconstruct partially the exsolution history, the \( T \) dependence of the lattice parameters \( \langle a \rangle \) and \( \beta \) and OPB theory were used to calculate plots of \( \tan(10T) \) vs. temperature for samples 7A7R and Q-603C (calibration curves in Fig. 9). The actual measured orientations were then plotted, and the temperatures of exsolution were derived (Figs. 9A and 9B). The estimated exsolution temperatures are between 780 and 300 °C for both samples with an error of ±80 °C, which is based on the maximum errors of the lattice parameters that were used to construct the calibrations curves.

The highest exsolution temperature of 780 °C for the first generation of exsolution in Q-603C is the only estimate that lies above independent estimates of peak metamorphic temperatures. However, use of the maximum...
error of the lattice parameters for the determination of the OPB calibration curve indicates that the temperature of exsolution could have been as low as 700 °C (Fig. 9A). This is very close to and probably within the error of the estimated peak metamorphic temperatures of 675 °C for locality Q-603C, which reached temperatures near the breakdown of muscovite + quartz in associated pelitic rocks. It should also be noted that estimates of exsolution temperatures above 600 °C are based on linear extrapolations of the temperature dependence of the lattice-parameter data well beyond the temperature range of the measurements. If the extrapolations are not valid for higher temperatures, this could explain the unreasonably high temperature of exsolution for the first generation of exsolution. Consequently, the temperatures that are assigned to events that are above about 650 °C are more speculative, although the sequence of events that is outlined below must have occurred.

These results suggest the following sequence of events for both localities. At or around peak metamorphic temperatures the assemblages contained two homogeneous amphiboles, hornblende and cummingtonite. (In the optical microscope, epitaxial intergrowth of hornblende and cummingtonite parallel to {101} and {100} was common.) For sample Q-603C, the first hornblende exsolution lamellae (~2.4 μm wide) in cummingtonite must have formed shortly below peak metamorphic conditions. Further exsolution of the cummingtonite host formed hornblende lamellae that were about 0.5 μm thick (lamellar orientation and, therefore, T could not be de-
tended). A third phase of hornblende exsolution occurred at about 640 °C (0.26 μm). Lamellar size (0.13 μm) suggests an additional hornblende generation between 640 and 560 °C. These were followed by successive generations of hornblende lamellae formed at about 560, 480, 420, and 300 °C (Figs. 9A and 10, Table 3).

The lowest exsolution temperatures of about 300 °C were derived from the nanometer-sized, disc-shaped, hornblende exsolution in the cummingtonite host of sample Q-603C. Because these exsolutions do not have straight boundaries, their orientation cannot be measured accurately, so this temperature of exsolution is tentative at best. Additionally, the OPB theory may no longer be describing exsolution at these temperatures and with these dimensions.

Hornblende in sample Q-603C exsolved two generations of cummingtonite (thicknesses of about 1.5 and 0.45 μm) that could not be measured but must have formed above about 580 °C (Fig. 10). Two generations of cummingtonite lamellae formed at about 580 and 480 °C (Figs. 9A and 10, Table 3).

The cummingtonite of sample 7A7R exsolved two generations of hornblende (thicknesses of about 1.5 and 0.45 μm) that could not be measured but must have formed between about 580 °C and the peak metamorphic temperature of about 640 °C. Three additional generations of hornblende lamellae formed at about 580, 520, and 380 °C.

The earliest generation of cummingtonite lamellae in hornblende from sample 7A7R could not be measured (thickness of about 1.5 μm) but must have formed between 640 and 540 °C. Three generations of cummingtonite exsolution are found in hornblende of 7A7R, and these occurred during cooling at about 550, 480, and 340 °C (Figs. 9B and 10).

These two multigenerational exsolution histories are consistent with the evidence that the histories (P-T paths) for the two localities are different (e.g., Schumacher et al. 1989). The observations here suggest that amphibolite rocks of the CM series (Q-603C) cooled more slowly than amphibolite rocks of the 7A7R series (Schumacher et al. 1993).

Given the present state of our understanding of the thermal, baric, and compositional control of hornblende and cummingtonite lattice parameters, a general hornblende-cummingtonite geothermometer is not presently...
feasible. However, we have shown that detailed study of individual samples can produce results that can be predicted with OPB theory. As more amphibole data become available, routine determination of temperatures of exsolution in cummingtonite and hornblende may become possible.

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