RHODONITE-PYROXMANGITE PERITECTIC ALONG THE JOIN MnSiO₃–MgSiO₃ IN AIR

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ABSTRACT

In the MnSiO₃–MgSiO₃ join a pyroxmangite-rhodonite peritectic exists on the liquidus at a composition of approximately (Mn₀.₄₇Mg₀.₅₃)SiO₃ at 1350°C ± 5°C in air. Unit-cell dimensions are given for the continuous solid-solutions of pyroxmangite at compositional range (Mn₀.₇₄Mg₀.₂₆)SiO₃–(Mn₀.₁₀Mg₀.₹₀)SiO₃ and for rhodonite at compositional range (Mn₀.₆₀Mg₀.₴₀)SiO₃–MgSiO₃.

Single crystals of pyroxmangite, (Mn₀.₅₅Mg₀.₄₅)SiO₃ and rhodonite, (Mn₀.₈₉Mg₀.₁₁)SiO₃ were grown to a maximum size of 5mm × 3mm × 2mm by slow cooling of the melts. At 1200°C in air, the join MnSiO₃–MgSiO₃ consists of four phases, i.e., triclinic rhodonite, at MnSiO₃–(Mn₀.₅₅Mg₀.₄₅)SiO₃; triclinic pyroxmangite, at (Mn₀.₇₄Mg₀.₂₆)SiO₃–(Mn₀.₁₀Mg₀.₹₀)SiO₃; clinopyroxene, at (Mn₀.₃₂Mg₀.₆₈)SiO₃ and orthorhombic protoenstatite, at (Mn₀.₁₆Mg₀.₸₄)–MgSiO₃. The phase transformation at fixed P and T for (Mn, Mg)SiO₃ appears to be controlled by the mean cationic radii for the variable Mn and Mg contents. A narrow but distinct rhodonite-pyroxmangite coexistence region is found below the solidus around (Mn₀.₅₅Mg₀.₴₅)SiO₃–(Mn₀.₅₄Mg₀.₴₆)SiO₃.

INTRODUCTION

A discontinuous solid-solution (rhodonite-pyroxene²) in the join MnSiO₃–MgSiO₃ under atmospheric pressure, with a break in the liquidus profile at approximately 50 mol percent MgSiO₃ and 1330°C, was first noted by Lebedew (1911). The phase equilibrium study by Glasser and Osborn (1960) using a controlled atmosphere (H₂ 20 percent and CO₂ 80 percent) along this join defined an unbroken rhodonite solid-solution series. It extended from the MnSiO₃ end member (incongruent M.P. 1291°C) to compositions containing a maximum of 94.5 wt percent of MgSiO₃ at 1454°C. These authors, however, noted that the X-ray powder diffraction pattern of 10 MnSiO₃·90 MgSiO₃ wt percent appeared significantly different from that of rhodonite, but they presented no crystallographic data.

In the present study quenching and slow cooling carried out in the air at temperatures ranging from 1200°C to 1520°C reveals a pyroxmangite-rhodonite peritectic at 1350°C ± 5°C at approximately (Mn₀.₆₇Mg₀.₃₃)SiO₃ as found by Lebedew (1911) at (Mn₀.₅₀Mg₀.₵₀)SiO₃.

¹ Mineralogical Contribution No. 486.
² Since pyroxmangite was not established as a valid species until 1913 (Ford and Bradley), Lebedew described the MgSiO₃-rich phase as orthorhombic pyroxene, but this was no doubt pyroxmangite.
A tentative phase diagram along this join, the result of the crystal growth attempts for both rhodonite and pyroxmangite, and a crystal chemical discussion on the formation of pyroxenoids are given in this paper.

**Experimental Method**

Starting materials used for this study were mixtures of the analyzed C. P. grade reagents of silicic acid, manganese carbonate, and magnesium hydroxide. These were thoroughly ground with a small amount of ethyl alcohol in an agate mortar. The charges were placed in a platinum container and were heated in a pre-heated silicon carbide furnace. Temperatures were controlled within 5°C by a proportional controller using a Pt-Rh3% percent thermocouple. The products were quenched to room temperature, except in the case of single crystal growth experiments (Table 1) in which the crystals were slowly cooled to room temperature in order to avoid thermal shock. For the synthesis runs approaching two phase boundaries at temperatures considerably below (>50°C) solidus, at least two grindings and repeated heating were necessary to reach equilibrium. Mixtures of crystalline rhodonites and pyroxmangites with the different Mn/Mg ratio obtained by the slow cooling experiments (Table 1) were used as starting materials for the experiments in establishing subsolidus equilibrium between rhodonite and pyroxmangite at 1250°C. The reaction between the rhodonites and the pyroxmangites having the different chemical compositions took place within 100 hours to reach equilibrium. The samples were thoroughly ground every 24 hours. The detailed results of quenching experiments summarized in Table [I] have been deposited with the National Auxiliary Publication Service.

Unit-cell dimensions were measured using X-ray powder diffraction (CuKα with Si standard) and refined with a least square computer program written by Dr. Daniel Appleman of the U.S. Geological Survey. Indexing of the diffraction peaks was facilitated using both the existing data of Lindsley and Burnham (1970), Momoi (1969), and Chao et al. (1970) and the intensity data observed on the precession single crystal photographs. These were taken at three axes for the natural pyroxmangite from Taguchi, Japan (Momoi, 1964) and two synthetic pyroxmangite crystals grown from melts having the compositions of (Mn0.2Mg0.2)SiO3 and (Mn0.2Mg0.8)SiO3.

**Results and Discussion**

Unit-cell Data

The unit-cell dimensions for the eight pyroxmangites synthesized in air within the solid-solution range Mn0.15Mg0.85SiO3—Mn0.50Mg0.40SiO3, and the five pyroxmangites synthesized hydrothermally within the solid-solution range Mn0.85Mg0.15—MnSiO3 are listed in Table [II] together with the data for MnSiO3 II (pyroxmangite).

1 To obtain a copy of this material, order NAPS Document Number 01770 from National Auxiliary Publications Service of the A.S.I.S., c/o CCM Information Corporation, 866 Third Avenue, New York, New York 10022; remitting $2.00 for microfiche or $5.00 for photocopies, in advance, payable to CCMIC-NAPS.
Table 1. Results of slow cooling of melts.

<table>
<thead>
<tr>
<th>Melt Composition</th>
<th>Temperature range (°C)</th>
<th>Cooling rate °C per hour</th>
<th>Products</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>melt soaking end (hour)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Mg.80Mn.20)SiO₃</td>
<td>1500 3 1150</td>
<td>3</td>
<td>Pink pyroxmangite crystals 3x2x2mm.</td>
</tr>
<tr>
<td>(Mg.70Mn.30)SiO₃</td>
<td>1490 5 1200</td>
<td>3</td>
<td>Aggregates of small pyroxmangite crystals.</td>
</tr>
<tr>
<td>(Mg.60Mn.40)SiO₃</td>
<td>1470 5 1200</td>
<td>3</td>
<td>Light brown pyroxmangite crystals 1x1x0.2mm.</td>
</tr>
<tr>
<td>(Mg.50Mn.50)SiO₃</td>
<td>1450 5 1180</td>
<td>3</td>
<td>Large light brown pyroxmangite crystals 5x3x2mm with small rhodonite.</td>
</tr>
<tr>
<td>(Mg.45Mn.55)SiO₃</td>
<td>1500 3 1180</td>
<td>3</td>
<td>Massive pyroxmangite mixed with a small amount of rhodonite.</td>
</tr>
<tr>
<td>(Mg.43Mn.57)SiO₃</td>
<td>1500 2 1180</td>
<td>5</td>
<td>Massive mixture of pyroxmangite and rhodonite with pyroxmangite fringe and rhodonite dome.</td>
</tr>
<tr>
<td>(Mg.40Mn.60)SiO₃</td>
<td>1450 6 1180</td>
<td>6</td>
<td>Massive mixture of pyroxmangite and rhodonite with domes of pure rhodonite.</td>
</tr>
<tr>
<td>(Mg.35Mn.65)SiO₃</td>
<td>1400 3 1300</td>
<td>25</td>
<td>Massive rhodonite and pyroxmangite.</td>
</tr>
<tr>
<td>(Mg.34Mn.66)SiO₃</td>
<td>1400 3 1300</td>
<td>20</td>
<td>Massive rhodonite and pyroxmangite.</td>
</tr>
<tr>
<td>(Mg.32Mn.68)SiO₃</td>
<td>1400 3 1300</td>
<td>20</td>
<td>Massive rhodonite.</td>
</tr>
<tr>
<td>(Mg.30Mn.70)SiO₃</td>
<td>1420 6 1180</td>
<td>3</td>
<td>Massive rhodonite.</td>
</tr>
<tr>
<td>(Mg.25Mn.75)SiO₃</td>
<td>1420 5 1180</td>
<td>3</td>
<td>Rhodonite and cristobalite.</td>
</tr>
<tr>
<td>(Mg.20Mn.80)SiO₃</td>
<td>1440 5 1180</td>
<td>10</td>
<td>Crystals of rhodonite 5x4x2mm with opaque inclusion.</td>
</tr>
<tr>
<td>(Mg.10Mn.90)SiO₃</td>
<td>1400 1 1200</td>
<td>10</td>
<td>Crystals of rhodonite and small tridymite.</td>
</tr>
<tr>
<td>MnSiO₃</td>
<td>1350 1 1240</td>
<td>5</td>
<td>Massive rhodonite and tridymite.</td>
</tr>
<tr>
<td>0.55 MnO + 0.45 SiO₂</td>
<td>1300 6 1200</td>
<td>2</td>
<td>Rhodonite crystals and manganese oxides.</td>
</tr>
</tbody>
</table>

synthesized by Akimoto and Syono (1972) under high-pressure; the same data are presented graphically in Figures 2 and 3.

The unit-cell dimensions for the six synthetic rhodonites within the solid-solution range Mn₀.₈₅Mg₀.₁₅SiO₃-MnSiO₃ synthesized in air are given in Table [III] with the other data obtained for MnSiO₃ synthesized under H₂O total pressure (Momoi, 1969; present study), and
dry high-pressure (Akimoto and Syono, 1972); the data are also plotted graphically in Figure 4.

Tables [II] and [III] have been deposited with the National Auxiliary Publication Service.

The changes observed for the unit-cell dimensions of the pyroxmangite series (Figs. 2 and 3) are almost linear functions of Mn and Mg contents. Yet there is some evidence for lack of linearity. The observed small gap at the composition Mn$_{0.80}$Mg$_{0.40}$SiO$_3$ between the two series of pyroxmangites synthesized one in the air (left) and the other under water pressure (right) may not be significant enough to discuss unless more information on the vapor phase activities and the precise compositional data are obtained.

The unit-cell dimensions of the rhodonites series (Fig. 4) are more nearly linear with composition than those of the pyroxmangite series.

Phase Study

The tentative phase diagram for the join MnSiO$_3$-MgSiO$_3$ in air between 1200°C and 1520°C is constructed using the results of the present study (Table 1 and Table [1]). These distinctly show the rhodonite-pyroxmangite peritectic at 1350°C ± 5°C at approximately (Mn$_{0.47}$Mg$_{0.53}$)SiO$_3$. The region above 1520°C is excluded because of the complexity expected from the results of the detailed study by Glasser and Osborn (1960) at temperatures above 1520°C and compositions between MgSiO$_3$ and (Mg$_{0.90}$Mn$_{0.10}$)SiO$_3$. On the whole, the results of Glasser and Osborn (1960) and those of the present work agree well except for the peritectic, although the latter experiments were carried out in air; the diagram presented here does not represent a true binary system, since a small amount of higher valency ionic manganese (excess oxygen)—other than divalent manganese—is present, particularly in compositions over MnSiO$_3$ 60 mol percent. The amount of Mn$_3$O$_4$ in the bulk sample of (Mn$_{0.70}$Mg$_{0.30}$)SiO$_3$, mainly due to the admixed unidentified manganese oxides, was found to be 0.2 wt percent, and the amount increases progressively up to 1.5 wt percent at the MnSiO$_3$ composition. However, the compositions having Mg contents higher than 40 mol percent are virtually free from any appreciable amount of Mn$_3$O$_4$, especially in the silicate crystals. The amount of higher valency manganese in rhodonite, MnSiO$_3$, quenched at over 1200°C in air is in fact negligible (Muan, 1959). In any case, the phase diagram of this join may require complete restudy under controlled partial pressure of oxygen $P_o$.

1 See footnote on page 865.
A significant difference in the composition for the peritectic point, between Lebedew (1911) and the present study, is most likely due to the considerable difference of $P_0$, in the two experiments. The $P_0$, in a carbon crucible at high-temperature (used by Lebedew) is considerably lower than that for heating in a platinum container in air.

There is a narrow but distinct two-phase region for rhodonite and pyroxmangite at the solidus composition around $(\text{Mn}_{0.65}\text{Mg}_{0.45})\text{SiO}_3$. In the subsolidus, this two-phase region shifts toward Mn$_3$SiO$_3$ side as temperature lowers. At 1250°C, equilibrium was reached at the composition around $(\text{Mn}_{0.65}\text{Mg}_{0.45})\text{SiO}_3$. The fact is consistent with the results of the hydrothermal (Momoi, 1968 and present study) and the dry high-pressure (Akimoto and Syono, 1972) synthesis of rhodonite-pyroxmangite dimorph with composition Mn$_3$SiO$_3$.

A small subsolidus region for clinopyroxene was also found near the Mg side (Fig. 1) at the temperatures 1200°-1300°C and the composition Mn$_{0.20}$Mg$_{0.80}$SiO$_3$ of this diagram. This clinopyroxene belongs to the isomorphous series extending through $(\text{Mn}_{0.50}\text{Mg}_{0.50})\text{SiO}_3$ (hydrothermal) to the end member Mn$_3$SiO$_3$—III (high-pressure clinopyroxene) synthesized by Akimoto and Syono (1972).

No critical attempt was made to establish the subsolidus equilibrium boundaries between two pyroxenes and pyroxmangite. The two-phase areas given shaded in the left hand side of this diagram (Fig. 1) are synthesis boundaries.

**Single Crystal Growth**

Single crystals of $(\text{Mn}, \text{Mg})\text{SiO}_3$ (over Mg$_{50}$ mol percent) pyroxmangite have been obtained by slow cooling of clear melt for some of the compositions within the join Mn$_3$SiO$_3$—Mg$_3$SiO$_3$. The largest crystal measures $5 \times 3 \times 2$ mm at $(\text{Mn}_{0.50}\text{Mg}_{0.50})\text{SiO}_3$. Medium size $(3 \times 2 \times 2$ mm) crystals were obtained for $(\text{Mn}_{0.20}\text{Mg}_{0.80})\text{SiO}_3$. Only small crystals were grown of $(\text{Mn}_{0.30}\text{Mg}_{0.70})\text{SiO}_3$ and $(\text{Mn}_{0.40}\text{Mg}_{0.60})\text{SiO}_3$. All the crystals are light amber in color. The color intensifies progressively with increasing amount of manganese in the crystals. The color turns to pink by heating at 1250°C for several hours, due to reduction of the trace amount of higher valency manganese in the crystals.

Single crystals of rhodonite were grown to $5 \times 4 \times 2$ mm for the composition $(\text{Mn}_{0.80}\text{Mg}_{0.20})\text{SiO}_3$ by fairly rapid cooling (in air at 10°C/hour) of melt. Small crystals of $(\text{Mn}_{0.90}\text{Mg}_{0.10})\text{SiO}_3$ and $(\text{Mn}_{0.70}\text{Mg}_{0.30})\text{SiO}_3$ were also obtained. The crystals are dark red in color, and always contain some opaque dots which probably are hausmannite Mn$_2$O$_4$; the Mn$_2$O$_3$ content of the bulk crystalline aggregates for $(\text{Mn}_{0.80}\text{Mg}_{0.20})\text{SiO}_3$ was found to be 0.5 wt percent. These rhodonites
also turn to bright pink by heating in a reducing atmosphere, as mentioned above.

Single crystals of pure end member rhodonite were grown from a melt having the composition MnO 55 and SiO₂ 45 mol percent (congruent melting), in air, by slow cooling from 1300°C to 1200°C at a rate of 5°C per hour; this procedure is based on the phase diagram given by Muan (1959).

Crystal Chemistry

Presence of five and seven silicate chain repeat units in rhodonite and pyroxmangite respectively, has been first found by Liebau (Liebau, 1957, 1959; Liebau, Hilmer, and Lindemann, 1959). Chemical analyses of natural pyroxmangites show ratios of Ca²⁺ to (Fe²⁺ + Mg²⁺ + Mn²⁺) less than 1:6. No rhodonites have Ca contents more than that needed to completely fill one out of five cation sites (Deer, Howie, and Zussman, 1963). In view of its size and coordination, there is no doubt that Ca²⁺ has an important role in limiting compositional range of pyroxenoid structure type. Based on the chemical data, Liebau indicated Ca-ordered structures for both rhodonite and pyroxmangite, but no actual evidence for the ordered structure was given. He suggested the following formulae for rhodonite Caₓ(Mn, Mg)₉₋ₓ SiO₄ with 0 < x < 0.2 and for pyroxmangite (Ca, Mg)(Mn, Fe)₅ Si₂O₁₁.

In the recent crystal structure refinement of rhodonite, Peacor and Niizeki (1963) assigned Ca²⁺ in M₅ site with the largest metal oxygen distance, and Mg²⁺ and Fe²⁺ in M₄ site while the other three sites are predominantly occupied by Mn²⁺.

In a review paper on pyroxene and pyroxenoid crystal chemistry, Prewitt and Peacor (1964) indicated that pyroxmangite structure may also follow the same trend as rhodonite. In such a structure, Ca²⁺ may enter exclusively one out of seven cation sites.

Until recently, the above ordered structure hypothesis has been well accepted in most of the mineralogical literature. A complete summary made by Chao et al. (1970) for the existing chemical analysis of the terrestrial pyroxmangite did not, however, support the formula suggested by Liebau (1957). Furthermore, a rhodonite-pyroxmangite dimorph with the composition MnSiO₄ has been synthesized hydrothermally (Momoi, 1968) and under anhydrous high-pressure conditions (Akimoto and Syono, 1972), indicating that Ca or other divalent ions other than Mn²⁺ are not essential for chain repeat units of 5 or 7 silicon tetrahedra.

Single silicate chains that repeat after every nine silicon tetrahedra were found by Burnham (1966) in ferrosilite III, a new pyroxenoid,
FeSiO₃ (Lindsley, Davis and MacGregor, 1964). The fact led Lindsley and Burnham (1970) to believe that Ca²⁺ is not essential for the 7 repeats of structure formation of pyroxferroite. They predicted that Ca-free pyroxmangite may be stable on the FeSiO₃—MnSiO₃ join particularly at Mn-rich compositions.

Latest results of the crystal structure study of lunar pyroxferroite, \( \text{Ca}_{0.13}\text{Fe}_{0.88}\text{Mn}_{0.02}\text{Mg}_{0.02}\text{Si}_{3.00} \) (Burnham, 1971) showed that Ca is distributed over three different sites (\( M_5, M_8, \) and \( M_7 \)) of pyroxmangite type structure, not restricted in a single \( M_7 \) site.

The results of the present study now show solid solutions of pyroxmangite over the compositional range \( (\text{Mn}_{0.50}\text{Mg}_{0.50})\text{SiO}_3 \) to \( (\text{Mn}_{0.16}\text{Mg}_{0.84})\text{SiO}_3 \) and rhodonite over the compositional range \( (\text{Mn}_{0.55}\text{Mg}_{0.45})\text{SiO}_3 \) to \( \text{MnSiO}_3 \), both in air (Fig. 1), along the join \( \text{MnSiO}_3—\text{MgSiO}_3 \). The isomorphous pyroxmangite series extends all the way to \( \text{MnSiO}_3 \) under hydrothermal conditions (Figs. 2 and 3). The study made by Riboud

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**Fig. 1.** Phase diagram for the join MnSiO₃-MgSiO₃ at temperatures between 1200°C and 1520°C in air. Py = pyroxmangite, Rh = rhodonite, Tri = tri dysmite, Clpx = clinopyroxene, Pr = protoenstatite and L = liquid. Shaded area = two-phase coexisting region. Unidentified manganese oxides observed microscopically at the composition MnSiO₃ over 60 mol% are not included.
Fig. 2. Unit-cell edges and volumes of synthetic pyroxmangites, (Mn,Mg)SiO$_3$. Small open circles = pyroxmangites synthesized in air at high temperature. Open ellipses = pyroxmangites synthesized hydrothermally. Large open circles = pyroxmangites synthesized at high-pressure and high-temperature (Akimoto and Syono, 1972).

and Muan (1962) under CO$_2$/H$_2$ = 1:1, 1 atm. showed that rhodonite solid-solution extends to (Mn$_{0.52}$Fe$_{0.48}$)SiO$_3$ wt percent, but no pyroxmangite phase was observed by them in one atm experiments in the join MnSiO$_3$–FeSiO$_3$. It is known that the original specimen of pyroxmangite described by Ford and Bradley from Iva, South Carolina,
is in fact manganan pyroferroite (Chao et al., 1970; Lindsley and Burnham, 1970) with the formula \( \text{Fe}_{0.25}\text{Mn}_{0.41}\text{Ca}_{0.34}\text{SiO}_3 \), but neither rhodonite nor pyroxmangite containing more than 6 wt percent of MgO has been found in nature. Site occupancies of both Mg and Fe\(^{2+}\) in these synthetic pyroxenoids should be investigated.

In a study of the high pressure transformation of MnSiO\(_3\), Akimoto and Syono (1972) stated that

When MnSiO\(_3\) is subjected to high pressure, the Mn\(^{2+}\)/Si\(^{4+}\) ionic radius ratio is expected to become smaller since the larger Mn\(^{2+}\) ions tend to contract more than the small Si\(^{4+}\) ions. Hence, it may be reasonable to presume that high pressure phase of MnSiO\(_3\) is turned into the pyroxenoid and pyroxene (rhodonite-pyroxmangite-clinopyroxene) corresponding to the smaller average size of the octahedral cations.

The results of the present study have demonstrated analogous trans-
formations by progressively replacing octahedral Mn$^{2+}$ (0.83Å)$^1$ by smaller Mg$^{2+}$ (0.72Å)$^1$ at fixed temperatures under atmospheric pressure. These results thus confirm that the most important controlling factor for these phase transformations is average cation size at given

In addition, the presence of wide ranging solid-solution series for pyroxmangites and rhodonites in the join MnSiO$_3$-MgSiO$_3$ suggests that Mg$^{2+}$ ions must be distributed over several sites in the structure of rhodonite and pyroxmangite; they cannot be restricted to a single site.

The garnet-like tetragonal phase with 6-coordinated Si$^{4+}$ obtained by Akimoto and Syono (1972) under ultra-high pressure has not been synthesized under the conditions used for the present study. This suggests, as would be expected, that Si$^{4+}$ does not occupy 6-coordinated sites under low pressure conditions.

Primary crystallization of rhodonite in rhyolite was found at Kesebol, Sweden (Geijer, 1961), but most naturally occurring rhodonites and pyroxmangites have been found in contact metamorphic, metasomatic or pegmatitic rocks in which the crystallization temperatures of these minerals are much lower than those used in this study, and H$_2$O vapor is always present. Similar crystallization of manganese metasilicates under atmospheric conditions was found in various slags. No serious attempt was made to correlate the results of the experiments in air to any of the natural occurrences. Nevertheless, this study demonstrates that under the proper conditions of pressure, temperature and composition, rhodonite and pyroxmangite can coexist in equilibrium. Although these conditions are undoubtedly rare in nature, this assemblage when found could represent an approach to equilibrium. Such an example of intimate intergrowth of rhodonite and pyroxmangite has been described by Burrell (1942) in specimens from Broken Hill, New South Wales, Australia.

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**References**


LIEBAU, VON F. (1957) The crystal structures of rhodonite, CaMn₁₂₄(SiO₄), and pyroxmangite, (Ca, Mg)(Mn, Fe)₄₄₂(SiO₄). Acta Crystallogr. 10, 761.


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