High-grade metamorphism in the Chapleau-Foleyet Area, Ontario

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Abstract

High-grade Archean rocks are exposed in the central Superior Province of the Canadian Shield in the Kapuskasing Structural Zone, a partly fault-bounded region up to 50 x 50 km. In migmatitic mafic gneiss, paragneiss, dioritic and tonalitic rocks, a lower-grade garnet-clinopyroxene-plagioclase (Gt-Cpx-Pl) zone and patchy, higher-grade orthopyroxene zones are distinguished. Grade decreases abruptly to greenschist in the Abitibi subprovince to the east across the Ivanhoe Lake cataclastic zone. Grade decreases gradually through amphibolite to greenschist facies in the Wawa subprovince to the west. Based on mineral-melt equilibria, minimum conditions for the Gt-Cpx-Pl zone are 750°C, 6 kbar, asr : 0.5-0.7 and for the orthopyroxene zone, 770°C, 6 kbar, asr : < 0.5.

Various garnet-biotite and garnet-pyroxene geothermometers and geobarometers yield apparent temperatures ranging from <600°C in the west to locally >800°C. Apparent pressure values derived from the pyrope-grossular-anorthite-diopside-quartz equilibrium are 5.4-8.4 kbar (average 6.3) and define a NNE-trending area of relatively high P in the eastern and central Kapuskasing Zone, supporting the interpretation of a tilted crustal section.

Introduction

Regional metamorphism variably affected rocks of the Superior Province over an area of ~3 x 10^6 km^2. The erosion level throughout the southern part of the province exposes metamorphic rocks of the greenschist and amphibolite facies except in the Kapuskasing Structural Zone, where pyroxene-bearing gneisses are associated with a linear gravity high (Innes, 1960; Bennett et al., 1967; MacLaren et al., 1968; Thurston et al., 1977). The Kapuskasing Zone in the relatively well-exposed Chapleau-Foleyet area consists of a variety of migmatitic pyroxene-garnet-hornblende-biotite-bearing rocks. The assemblages preserved at the time of metamorphic quenching provide insight into reactions leading to the decomposition of hornblende and biotite in an amphibolite-granulite facies transition zone. The metamorphic history and equilibration conditions of rocks in the Kapuskasing Zone and adjacent Wawa subprovince are examined in this study.

Metamorphism in the Kapuskasing Zone is Archean as inferred from U-Pb dates on zircon from leucosome in paragneiss (2627±5 Ma) and on metamorphic zircon in mafic gneiss (2650±2 Ma) (Percival and Krogh, 1983).

Geological setting

The Superior Province can be divided into metavolcanic-plutonic and metasedimentary subprovinces (Stockwell, 1970) (Fig. 1). In the south-central Superior Province, the easterly trends of the Abitibi-Wawa and Quetico-Opatica subprovinces are transected over a distance of 500 km by the north-northeast-trending Kapuskasing Structural Zone (Thurston et al., 1977) (Fig. 1). It is made up of gneisses in the upper amphibolite to granulite facies (Bennett et al., 1967; MacLaren et al., 1968) and is characterized by positive gravity and aero-magnetic anomalies over most of its length (Innes, 1960).

The Chapleau-Foleyet area straddles the Kapuskasing Zone and includes parts of the adjacent Abitibi and Wawa subprovinces. Supracrustal rocks of the Abitibi subprovince in the study area comprise mostly mafic metavolcanic flows and tuffs up to 6100 m thick (Goodwin, 1965). Easterly-trending belts of felsic metavolcanic and metasedimentary rock make up about 5 and 3% respectively of the supracrustal succession. Massive to foliated plutons of tonalite to granite composition (Streckeisen, 1976) range in diameter from ~1 to ~25 km. The intensity of metamorphism increases from the cores of supracrustal belts, where greenschist-facies assemblages are common (Thurston et al., 1977), to contact areas with large intrusive bodies, where hornblende-plagioclase±garnet assemblages prevail. Both primary structures, including bedding and volcanic features, and sub-vertical tectonic foliation, schistosity and lineation, are present in the supracrustal succession.

The contact between rocks of the Abitibi subprovince and the Kapuskasing Structural Zone to the west is the Ivanhoe Lake cataclastic zone (Fig. 2), comprising blast-
tomylonite, cataclasite and pseudotachylite veinlets (Percival and Coe, 1981). These fault rocks are sporadically developed in rocks of the eastern 2 km of the Kapuskasing Zone and have random orientation within individual outcrops. The overall NNE trend is deduced from the distribution of fault rocks in the field.

The Kapuskasing Zone consists of ENE-striking belts of paragneiss, mafic gneiss, tonalitic and dioritic rocks and units of the Shawmere anorthosite complex (Thurston et al., 1977; Percival, 1981a,b; Riccio, 1981). Paragneiss is layered, migmatitic, fine- to medium-grained, biotite-plagioclase-quartz rock, with some garnet and/or hornblende and/or orthopyroxene (Table 1). Concentrations of quartz, biotite, garnet, or graphite in some layers and the overall quartz-rich composition suggest that these rocks had a sedimentary origin. Mafic gneiss is a layered to homogeneous medium- to coarse-grained rock of high calcium (10.0–15.4 wt.% CaO), high-alumina (13.4–17.2 wt.% Al₂O₃) basaltic composition. It consists of Gt–Cpx–Hb–Pl–Qz–Il±Opx±Sp±Sc assemblages and contains tonalitic leucosome veinlets on the 1–20 mm scale. Flaser diorite to mafic tonalite occurs as homogeneous to layered medium- to coarse-grained Hb–Bt–Pl±Qz±Cpx±Opx assemblages with up to 10% quartz and locally up to 15% concordant quartz monzonite veinlets. Rare gabbro and hornblende layers may represent igneous differentiates. Tonalitic rocks are foliated to gneissic, locally xenolithic, consist of Bt–Pl–Qz±Hb±Gt assemblages, and contain paragneiss, mafic gneiss, and ultramafic (hornblende or Gt–Opx–Hb rock) inclusions. Late hornblende–biotite tonalite dikes up to 50 cm thick transect gneissic layering in mafic and tonalitic gneiss. In 20-cm wide zones adjacent to these dikes, garnet, pyroxene-bearing assemblages are replaced by hornblende ±biotite assemblages. The Shawmere anorthosite complex (Fig. 2) comprises a main northern body, ~50 × 15 km, and a southern lens, ~15 × 4 km. The southern intrusion is homogeneous gabbroic anorthosite whereas the larger body contains anorthosite through gabbro units as well as some ultramafic rocks. The northern body is crudely zoned, consisting of a thin marginal unit of migmatitic amphibolite, a layered anorthosite–gabbro unit and a core of plagioclase-megacrystic gabbroic anorthosite (Riccio, 1981). A thin unit of foliated tonalite within the complex yielded a minimum U/Pb zircon date of 2765 Ma (Percival and Krogh, 1983) which provides a minimum age for the paragneiss-mafic gneiss country rock.

The western boundary of the Kapuskasing Zone is defined by the change in structural style and orientation from domal in the Wawa subprovince to northeasterly belts to the east (Fig. 2). Rare metaconglomerate in this boundary area has tonalitic cobbles with a U/Pb zircon date of 2664±6 Ma and a K–Ar hornblende date of 2594±151 Ma (Percival et al., 1981).

The Wawa subprovince in the western part of the area (Fig. 2) is a tonalite to granodiorite gneiss terrane intruded by massive to foliated granite, quartz monzonite and tonalite plutons. The gneisses contain 0–25% amphibolitic enclaves as well as rare easterly-trending paragneiss units. Gneissosity is steep to subhorizontal, forming several domal structures, some with plutonic cores (Percival, 1981a). The Robson Lake dome, adjacent to the Kapuskasing Structural Zone (Fig. 2), has a core of mafic gneiss. Tonalite and granodiorite gneiss yield U/Pb zircon dates of >2707 and 2677±14 Ma respectively (Percival and Krogh, 1983).

The age of metamorphism in the Abitibi and Wawa subprovinces is constrained by zircon dates on deformed, metamorphosed volcanics of 2749–2696 Ma (Turek et al., 1982; Nunes and Pyke, 1980) and on post-metamorphic plutons of 2685–2668 Ma (Krogh and Turek, 1982; Krogh et al., 1982). Concordant U–Pb zircon dates of 2650 and 2627 Ma from high-grade rocks of the Kapuskasing Zone indicate either a discrete younger metamorphic event or sustained metamorphic effects of the pre-2685 Ma event (Percival and Krogh, 1983).

Metamorphic zones

Mineral assemblages in each lithologic unit are listed in Table 1 and plotted in Figure 3. Assemblage data in the Abitibi subprovince in this area are not sufficient for delineation of isograds. Chlorite–white mica–albite–quartz assemblages characterize metasediments and epidote–chlorite–albite–quartz–carbonate±white mica±hornblende assemblages are common in mafic rocks (Thurston et al., 1977). Hornblende–plagioclase±garnet assemblages occur in proximity to granitic plutons and locally near the Kapuskasing Zone.

Two high-grade metamorphic zones are recognized in the Kapuskasing Structural Zone and Wawa subprovince.
Fig. 2. Generalized geological map of the Chapleau–Foleyet area (modified after Percival (1981b)). Geological boundary between the Kapuskasing Zone and Wawa subprovince is gradational in this area. Approximate boundary is Chapleau River south to Robson Lake dome, then Nemegosenda River south to Lackner Lake complex.

The Gt-Cpx-Pl zone, with diagnostic assemblages in mafic gneiss, extends from the Ivanhoe Lake cataclastic zone in the east, westward into the Wawa subprovince. In addition to the diagnostic minerals, mafic rocks contain brown hornblende, quartz, ilmenite and/or sphene, rare scapolite and ubiquitous tonalitic leucosome veinlets comprising variable proportions of quartz and plagioclase with minor hornblende, garnet or clinopyroxene. Para-
Table 1. Mineral assemblages and densities of rocks in Kapuskasing Structural Zone and eastern Wawa subprovince

<table>
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<tr>
<th>Rock Type</th>
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<td>Anorthosite-suit rocks</td>
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Mineral abbreviations: Ol: olivine; Sp: sphene; Gt: garnet; Sl: sillimanite; St: staurolite; Sa: sapphire; Opx: orthopyroxene; Cpx: clinopyroxene; Hb: hornblende; Oa: orthoamphibole; M: muscovite; Bt: biotite; Pl: plagioclase; K: K-feldspar; Q: quartz; Sc: scapolite; I: ilmenite; Sn: spinel; Cc: calcite; Cu: cummingtonite; S.G.: specific gravity (bracketed numbers refer to average values); ± designation indicates that small quantities of the mineral are present as an additional phase in some assemblages.

The Kapuskasing structural zone is characterized by a transition from lower grade mafic rocks with Hb-Pl-Qz assemblages at the margins of the Kapuskasing Zone to higher grade felsic rocks with Gt-Cpx-Qz overgrowths. Typical assemblages at each grade are shown in Table 1. The Kapuskasing Zone is characterized by the presence of orthopyroxene and garnet at lower grade, which transform to higher grade assemblages of staurolite and muscovite. The zone is prograded with felsic intrusions and retrogression.

**Mineral chemistry**

All mineral analyses were carried out on an ARL model AMX microprobe equipped with an energy dispersive detector set up and supervised by P. L. Roeder and M. I. Corlett at Queen's University. Operating conditions were...
15 kV, 0.5 μA. Counting times of 120 seconds were used and data processed by an on-line minicomputer set up to apply Bence–Albee (1968) corrections and calculate weight per cent of oxides.

Between three and six spot analyses were made of each mineral in a probe section. Early detailed work showed that the average of three analyses is not significantly different from the average of six and therefore most of the results are averages of three analyses per mineral. Most minerals are homogeneous, with some chemical zonation within ~5 μm of the rim. Where rim composition is distinctly different from the interior analyses, it was excluded from the average.

Ten element analyses were recast into structural formulae according to the method of Deer et al. (1966). An estimate of Fe³⁺ content of garnet and pyroxene was made by assuming electroneutrality and adjusting Fe²⁺/Fe³⁺ accordingly. Maximum and minimum Fe³⁺ contents of hornblende were calculated by a method analogous to that of Stout (1972) and then averaged.
Mineral compositions from 25 paragneiss specimens are given in Appendix 1 and some compositional parameters are listed in Table 2. Garnet in paragneiss varies in Mg/(Mg+Fe2+) from 0.15 to 0.35. Grossular component constitutes 3–18 mol% and spessartine 1.1–11.7 mol%. Combined grossular-spessartine is generally 8–15 mol% with a few values as high as 25% (Table 2). Garnets are commonly zoned near the rim and the general pattern is toward increasing almandine (0–8 mol%) and spessartine (0–4 mol%) and decreasing pyrope (0–7 mol%) and grossular (0–3 mol%) components. The cores of grains are commonly homogeneous and the near-rim zonation may be a surface retrograde effect (cf. Lasaga et al., 1977). Orthopyroxene in paragneiss has a range of Mg/(Mg+Fe2+) of 0.44–0.62 and Al2O3 constitutes 0.9–3.5 wt%. Biotite is characterized by Mg/(Mg+Fe) = 0.35–0.68 and contains up to 4 wt.% TiO2. Plagioclase is in the range An24–35 with one exception of An66. Analyses of coexisting garnet, orthopyroxene, biotite, K-feldspar and plagioclase from specimen PG-21 are presented in Table 3.

Mineral analyses from 34 mafic gneiss specimens are given in Appendix 1, and compositional parameters are listed in Table 4. Two of the specimens in the MG-series, MG-1 and MG-2, are skarns, but are included because of their grossular-rich garnet-clinopyroxene assemblages. Garnet in mafic gneiss is in the Mg/(Mg+Fe2+) = 0.13–0.43 range and grossular component constitutes 18–34 mol%. Spessartine is less than 5 mol% and andradite component is less than 5 mol%, an exception being MG-2, with 20 mol% andradite. Garnets generally have homogeneous interiors but are weakly zoned immediately adjacent to the rim. Spessartine and grossular contents vary only slightly whereas almandine increases (0–5 mol%) and pyrope decreases (0–4 mol%) within 5–10 µm of the grain boundary. Clinopyroxene is salite, with Mg/(Mg+Fe2+) values of 0.45–0.80. Jadeite and Ca-tschermakite components are in the 0.22 and 0.78 mol% ranges respectively. Amphiboles are common hornblende except for some secondary actinolite. Amphibole which appears to be part of the equilibrium assemblage is brown and richer in Al2O3 (10–12 wt.%) and TiO2 (0.4–2.2 wt.%) than green amphibole of apparent retrograde origin (8–10 wt.% Al2O3; 0.03–0.56 wt.% TiO2). An exception is secondary sodic amphibole from MG-30, with 4.15 wt.% TiO2. Plagioclase is andesine to labradorite (An3FAn65) and contains up to 4.7 wt.% Al2O3. Scapolite has been noted in a few specimens, in one to the exclusion of plagioclase. Orthopyroxene is present in only two mafic gneiss specimens (MG-20, 22), and contains up to 4.7 wt.% Al2O3. Analyses of coexisting garnet, clinopyroxene, orthopyroxene and hornblende in specimen MG-20 are presented in Table 5.

Mineral analyses of dioritic, ultramafic and anorthosite-suite rocks are included in Appendix 1 and parameters listed in Table 6. Analyses of coexisting orthopyroxene, clinopyroxene, hornblende, biotite and plagioclase from a dioritic rock (OG-6) are presented in Table 7.

Graphical representation of assemblages

Minerals in mafic gneiss can be approximately represented in the system SiO2-Al2O3-FeO-MgO-CaO-Na2O-H2O, by ignoring the minor amounts of Fe2O3 calculated for the mafic phases. At fixed pressure, temperature and aH2O and in the presence of quartz and plagioclase of constant composition, phase relations can be shown in terms of three components:

\[ A = \text{Al}_2\text{O}_3 - (\text{CaO} + \text{Na}_2\text{O}) \]
\[ F = \text{FeO} \]
\[ M = \text{MgO} \]

Mineral compositions can thus be plotted on a diagram using A/F+M and M/F+M coordinates (Stout, 1972).

Projections of coexisting hornblende, clinopyroxene, orthopyroxene and garnet define a crossing tie-line (Fig. 4). Consequently, the four minerals are stable along a line in a P-T diagram if aH2O = f(P,T); at lower temperature, hornblende and garnet are stable and at higher temperature, orthopyroxene and clinopyroxene are stable (Fig. 4). If water activity varies independently of P,T, a decrease in aH2O at constant temperature would favor the orthopyroxene-clinopyroxene assemblage.

The assemblage garnet-clinopyroxene-hornblende is stable over a P-T range, represented by a shifting triangle on Figure 4. The garnet-clinopyroxene zone is reached when the bulk composition of the rock falls into the garnet-clinopyroxene-hornblende field. The orthopyroxene zone is reached when the path of metamorphism intersects the reaction

\[
garnet + \text{hornblende} \rightleftharpoons \text{orthopyroxene} + \text{clinopyroxene} \\
(1)
\]

The path of metamorphism extends only slightly beyond this reaction because the incompatibility of garnet–hornblende is never fully established.

Pressures and temperatures of migmatite formation

Tonalitic veins, consisting of plagioclase (An27–33), quartz and minor biotite, orthopyroxene, garnet, clinopyroxene or hornblende, occur in mafic rocks and in paragneiss as concordant layers and discrete pods. Several modes of origin are possible: 1) crystallization from...
injected melt; 2) metamorphic differentiation; or 3) crystallization from in situ anatectic melt. The injected melt hypothesis is improbable because the pods and veinlets are generally isolated. In addition, the mafic mineral content of the veinlets is the same as that in the host. Both characteristics suggest local derivation. Metamorphic segregation was invoked by Amit and Eyal (1976) to explain quartz-plagioclase leucosome in the Wadi Ma'grish migmatites. This process is probable at sub-solidus temperatures (~630°C; Amit and Eyal, 1976), but is a more tenuous hypothesis for the present suite of rocks where indicated temperatures are much higher (see Geothermometry). Above 690°C at 5 kbar, in the presence of an aqueous fluid, which would be required to
transport the quartz and plagioclase components into the metamorphic segregation, quartz-plagioclase–H₂O should form a melt phase (Yoder, 1967). The modal composition of tonalite veinlets in the Kapuskasing mafic gneiss (Anₓₙ₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋₅₋òng amount of water activity decreases. The overall negative slope of the vapor-absent melting curve is a function of the shapes of the haplogranitic solidi (concave toward high P, T) and amphibole dehydration curves (concave toward low P). These curves intersect at two P, T points where ae is 1 by definition. The vapor-absent melting curve joining these two points has a negative slope, which is in contrast to the experimental vapor-absent melting curves for hornblende in acid and intermediate compositions of Brown and Fyfe (1970), which show strong positive dP/dT values. The
Table 4. Mafic gneiss mineral composition and equilibria data

<table>
<thead>
<tr>
<th>No.</th>
<th>Assemblage</th>
<th>Metamorphic Zone</th>
<th>Garnet</th>
<th>Clinopyroxene</th>
<th>Ti*°C</th>
<th>P (bar)</th>
<th>H2O</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG-1</td>
<td>Sp Gt Cpx Pl Q Cc</td>
<td>3</td>
<td>0.11</td>
<td>0.209</td>
<td>0.677</td>
<td>0.103</td>
<td>0.546</td>
<td>0.028</td>
</tr>
<tr>
<td>MG-2</td>
<td>Gt Cpx Q</td>
<td>3</td>
<td>0.19</td>
<td>0.230</td>
<td>0.681</td>
<td>0.070</td>
<td>0.449</td>
<td>0.103</td>
</tr>
<tr>
<td>MG-3</td>
<td>Sp Gt Cpx Hb Q Sc I</td>
<td>3</td>
<td>0.078</td>
<td>0.310</td>
<td>0.327</td>
<td>0.083</td>
<td>0.676</td>
<td>0.052</td>
</tr>
<tr>
<td>MG-4</td>
<td>Sp Gt Cpx Hb Bt</td>
<td>4</td>
<td>0.10</td>
<td>0.626</td>
<td>0.237</td>
<td>0.04</td>
<td>0.599</td>
<td>0.024</td>
</tr>
<tr>
<td>MG-5</td>
<td>Sp Gt Cpx Cu Hb</td>
<td>4</td>
<td>0.092</td>
<td>0.572</td>
<td>0.304</td>
<td>0.031</td>
<td>0.571</td>
<td>0.061</td>
</tr>
<tr>
<td>MG-6</td>
<td>Sp Gt Cpx Hb Pl Q I</td>
<td>3</td>
<td>0.088</td>
<td>0.524</td>
<td>0.343</td>
<td>0.046</td>
<td>0.567</td>
<td>0.061</td>
</tr>
<tr>
<td>MG-7</td>
<td>Gt Cpx Hb Bt Pl Q I</td>
<td>1</td>
<td>0.105</td>
<td>0.607</td>
<td>0.235</td>
<td>0.033</td>
<td>0.568</td>
<td>0.050</td>
</tr>
<tr>
<td>MG-8</td>
<td>E Cpx Hb Pl Q I</td>
<td>1</td>
<td>0.123</td>
<td>0.644</td>
<td>0.211</td>
<td>0.022</td>
<td>0.700</td>
<td>0.050</td>
</tr>
<tr>
<td>MG-9</td>
<td>Gt Cpx Hb Pl Q I</td>
<td>4</td>
<td>0.11</td>
<td>0.587</td>
<td>0.260</td>
<td>0.039</td>
<td>0.576</td>
<td>0.032</td>
</tr>
<tr>
<td>MG-10</td>
<td>Sp Gt Cpx Hb Pl Q I</td>
<td>3</td>
<td>0.112</td>
<td>0.535</td>
<td>0.288</td>
<td>0.043</td>
<td>0.604</td>
<td>0.055</td>
</tr>
<tr>
<td>MG-11</td>
<td>Sp Gt E Cpx Hb Pl Q I</td>
<td>3</td>
<td>0.114</td>
<td>0.549</td>
<td>0.304</td>
<td>0.033</td>
<td>0.535</td>
<td>0.031</td>
</tr>
<tr>
<td>MG-12</td>
<td>E Gt Cpx Hb Pl Q I</td>
<td>4</td>
<td>0.118</td>
<td>0.561</td>
<td>0.276</td>
<td>0.046</td>
<td>0.622</td>
<td>0.077</td>
</tr>
<tr>
<td>MG-13</td>
<td>Gt Cpx Hb Pl Q I</td>
<td>3</td>
<td>0.124</td>
<td>0.582</td>
<td>0.254</td>
<td>0.061</td>
<td>0.603</td>
<td>0.037</td>
</tr>
<tr>
<td>MG-14</td>
<td>Sp Gt Cpx Hb Pl Q I</td>
<td>1</td>
<td>0.122</td>
<td>0.537</td>
<td>0.280</td>
<td>0.039</td>
<td>0.643</td>
<td>0.073</td>
</tr>
<tr>
<td>MG-15</td>
<td>Sp Gt Cpx Hb Pl Q I</td>
<td>3</td>
<td>0.126</td>
<td>0.570</td>
<td>0.272</td>
<td>0.031</td>
<td>0.632</td>
<td>0.044</td>
</tr>
<tr>
<td>MG-16</td>
<td>Gt Cpx Hb Bt Q I</td>
<td>3</td>
<td>0.131</td>
<td>0.581</td>
<td>0.260</td>
<td>0.029</td>
<td>0.680</td>
<td>0.099</td>
</tr>
<tr>
<td>MG-17</td>
<td>Gt Cpx Hb Bt Q I</td>
<td>3</td>
<td>0.137</td>
<td>0.606</td>
<td>0.227</td>
<td>0.030</td>
<td>0.607</td>
<td>0.073</td>
</tr>
<tr>
<td>MG-18</td>
<td>Sp Gt Cpx Hb Bt Q I</td>
<td>3</td>
<td>0.136</td>
<td>0.592</td>
<td>0.248</td>
<td>0.023</td>
<td>0.655</td>
<td>0.063</td>
</tr>
<tr>
<td>MG-19</td>
<td>Gt Cpx Hb Bt Q I</td>
<td>3</td>
<td>0.149</td>
<td>0.602</td>
<td>0.225</td>
<td>0.024</td>
<td>0.636</td>
<td>0.052</td>
</tr>
<tr>
<td>MG-20</td>
<td>Opx Cpx Hb Bt Q I</td>
<td>4</td>
<td>0.157</td>
<td>0.601</td>
<td>0.228</td>
<td>0.013</td>
<td>0.692</td>
<td>0.078</td>
</tr>
<tr>
<td>MG-21</td>
<td>Gt Cpx Hb Pl Q I</td>
<td>4</td>
<td>0.151</td>
<td>0.566</td>
<td>0.256</td>
<td>0.027</td>
<td>0.638</td>
<td>0.046</td>
</tr>
<tr>
<td>MG-22</td>
<td>Opx Cpx Hb Pl Q I</td>
<td>4</td>
<td>0.164</td>
<td>0.598</td>
<td>0.348</td>
<td>0.036</td>
<td>0.568</td>
<td>0.033</td>
</tr>
<tr>
<td>MG-23</td>
<td>Sp Gt Cpx Hb Bt Pl Q I</td>
<td>3</td>
<td>0.194</td>
<td>0.604</td>
<td>0.203</td>
<td>0.023</td>
<td>0.692</td>
<td>0.030</td>
</tr>
</tbody>
</table>
orthopyroxene + anorthite = garnet

$$\text{Fe}_2\text{SiO}_4 + \text{CaSiO}_3 + \text{NaAlSi}_3\text{O}_8$$

The $P-T$ position of $X_{\text{Mg}} = 0.09-0.33$ stability contours for Reaction 5 were estimated by first calculating the Fe:Mg ratio of orthopyroxene which would be in equilibrium with garnet using an average Grt-Opx Fe:Mg distribution coefficient of 3.46 then calculating the $P-T$ conditions defined by the assemblage according to Hansen (1981). This provides a minimum pressure at specified temperature in the absence of orthopyroxene. The intersection of stability contours for Reactions 4 and 5 defines the position of Reaction 1.

Paragneiss in the Grt-Cpx-Pl zone generally contains Grt-Bt-Pl-Qz assemblages and tonalitic leucosome. The first appearance of orthopyroxene can be related to a biotite-consuming reaction:

$$\text{biotite} + \text{quartz} \rightarrow \text{orthopyroxene}$$

$$\text{K-feldspar} + \text{H}_2\text{O}$$

The downward shift in the position of the dehydration curve as a result of coexistence with water-undersaturated melt has been calculated for (6) with natural mineral compositions in the same manner as for (2). The locus of intersection of equal $\Delta H_\text{O}$ for (6) and the tonalite solidus yields a vapor-absent melting curve along which the generalized reaction is:

$$\text{biotite} + \text{quartz} + \text{plagioclase}$$

$$\rightarrow \text{orthopyroxene} + \text{granodiorite}$$

The initial melt composition in the leucosomes is tonalitic but becomes granodioritic after K-feldspar is produced by decomposition of biotite, in accord with the observation of up to 10% antiperthite in leucosome of orthopyroxene-bearing paragneiss.

The curve (Fig. 6) has a steep positive $dP/dT$ slope and assuming $P$ of 6 kbar, orthopyroxene is stable above 770°C, in agreement with the experiments of Wendlandt (1981) and Clemens and Wall (1981).

The preceding discussion has assumed a closed system and equilibrium between melt and solids, however it is difficult to explain the presence of fresh orthopyroxene in tonalite veinlets if the system remained closed until crystallization. Reactions between water dissolved in the melt and orthopyroxene should yield hydrous phases. Hence water is considered to have been removed or diluted during or prior to crystallization. Removal could have been accomplished by collection and upward migration of tonalitic liquids leaving a water-depleted residue. The origin of many Archean tonalites has been ascribed to partial melting of mafic rocks containing garnet-clinopyroxene-quartz (Arth and Hansen, 1972) or garnet-clinopyroxene-orthopyroxene-plagioclase (Gower et al., 1982). Alternatively, dilution of the ambient fluid by externally-derived CO$_2$-rich vapors (Weaver, 1980; Friend, 1981; Janardhan et al., 1982) could induce crystallization and prevent retrogression, in the manner outlined by Wendlandt (1981).

Superimposed on Figure 6 is the vapor-phase absent melting curve for mafic rocks. The two curves intersect near 6 kbar, 775°C for natural compositions. This implies that in regional metamorphic terranes characterized by pressure above this intersection, hornblende will start to react to form garnet-clinopyroxene-melt at temperatures where paragneiss may be migmatitic but contains no orthopyroxene. In lower-pressure environments, biotite will become unstable at lower grade than hornblende, or hornblende may be involved in reactions producing orthopyroxene in mafic rocks (cf. Wells, 1979).

Minimum pressure-temperature conditions were de-
fined in the previous section by solid–melt equilibria for the Gt-Cpx-Pl zone and for the orthopyroxene zone. Regional variation in apparent P-T conditions is now examined, based on calibrated geothermometers and geobarometers applied to individual samples. (Tables 2, 4 and 6)

**Temperature estimation methods**

Geothermometers based on Fe–Mg exchange between mineral pairs are sensitive to temperature variation and only slightly pressure-dependent. Therefore, to evaluate temperature-variation on a first approximation basis, a value of 7 kbar was initially estimated for all specimens. This value cannot be so precisely estimated but is shown below to be a reasonable average for the area.

The mineral pairs most suitable for geothermometry by virtue of widespread distribution and unaltered character are garnet–biotite and garnet–clinopyroxene. The Ferry and Spear (1978) experimental calibration of the garnet-biotite thermometer has gained general acceptance (e.g., Ghent et al., 1979; Ganguly, 1979), the experimental calibration of Ellis and Green (1979), which takes account of the effect of calcium on Fe-Mg distribution, is well-suited for the study of the Kapuskasing Zone mafic rocks whose garnets contain up to 34 mol% grossularite. Mineral compositions meet the compositional restrictions imposed in both garnet–biotite and garnet–clinopyroxene calibrations (Appendix 1).

Figure 7 shows temperature estimates of the Ferry and Spear calibration plotted against estimates by the three most recent garnet–clinopyroxene calibrations for four rocks from the Kapuskasing Zone that contain all three minerals. Internal consistency is best for these samples for the Ellis and Green and Ferry and Spear calibrations. Both are experimentally-derived equations and take account of pressure effects. The agreement between the two techniques is in contrast to that reported for Adirondack granulites studied by Bohlen and Essene (1980).

Stoichiometric estimation of Fe$^{3+}$ in biotite from probe data is not possible because site vacancies in biotite leave the structural formula cation-deficient. However, wet chemical analyses of biotite consistently indicate the presence of Fe$_2$O$_3$. To assess the partitioning of Fe$^{3+}$
and temperatures up to 100°C lower result if the calculation assumes 8.4% more Fe\textsuperscript{3+}/Fe\textsuperscript{\textit{total}} in biotite than in garnet. The effect on geothermometric measurement of assuming 8.4% more Fe\textsuperscript{3+}/Fe\textsuperscript{\textit{total}} in biotite than in garnet is negligible (apparent change of -1°C using Ferry–Spear calibration). Assuming a 12% average, the difference is only -2°C.

The agreement between the garnet–clinopyroxene and garnet–biotite thermometers is best when all iron is assumed to be Fe\textsuperscript{2+} in garnet, biotite and clinopyroxene for both thermometers (Fig. 7) and the calculations in Tables 2, 4 and 6 are based on this procedure. Treating all iron as Fe\textsuperscript{2+} is actually an empirical correction used to achieve consistency between geothermometers (Fig. 7) and contrasts with the procedure of Raheim and Green (1975), who used calculated Fe\textsuperscript{2+} contents. The pyroxene commonly has larger Fe\textsuperscript{3+}/Fe\textsuperscript{\textit{total}} than coexisting garnet and temperatures up to 100°C lower result if the calculated Fe\textsuperscript{2+} value is used to calculate $K_D$. Further justification for using total Fe rather than Fe\textsuperscript{2+} comes from samples MG-17 and MG-25, whose garnet and clinopyroxene contain no Fe\textsuperscript{3+} as calculated stoichiometrically. Temperatures of 725 and 755°C are in the same range as values calculated by ignoring Fe\textsuperscript{3+} in other specimens (Table 4).

A calibration of the two-pyroxene thermometer, proposed by Powell (1978), yields results which are consistent with garnet–biotite and garnet–clinopyroxene temperatures for the Kapuskasing gneisses. The equations formulated by Kretz (1982) give averaged values in the same range but some pyroxene pairs yield temperatures that are up to 100°C discordant. The Wood and Banno (1973) and Wells (1977) calibrations consistently give temperatures some 200–300°C higher.

Additional results were obtained for rocks containing mineral assemblages without widespread distribution (Tables 2, 4 and 6). In general, oxide, feldspar and ilmenite–pyroxene thermometers yield temperatures some 200–300°C lower than those estimated by the Fe–Mg exchange thermometers. No generalizations regarding other thermometers are possible because of the limited number of occurrences.

### Regional temperature variation

Near-rim garnet analyses generally give results lower by 10–20°C than analyses of the relatively homogeneous interiors. The general analytical strategy was to check all

---

**Table 7. Microprobe analyses of minerals in OG-6**

| Mineral | Composition
<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>SiO\textsubscript{2}</td>
<td>53.28</td>
</tr>
<tr>
<td>TiO\textsubscript{2}</td>
<td>0.15</td>
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<tr>
<td>Al\textsubscript{2}O\textsubscript{3}</td>
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</tr>
<tr>
<td>Cr\textsubscript{2}O\textsubscript{3}</td>
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</tr>
<tr>
<td>Fe\textsuperscript{3+}</td>
<td>29.53</td>
</tr>
<tr>
<td>MnO</td>
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<tr>
<td>MgO</td>
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<td>CaO</td>
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<tr>
<td>Na\textsubscript{2}O</td>
<td>0.38</td>
</tr>
<tr>
<td>K\textsubscript{2}O</td>
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</tr>
<tr>
<td>Total</td>
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</tr>
<tr>
<td>Si</td>
<td>1.956</td>
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<tr>
<td>Al</td>
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<tr>
<td>Ti</td>
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<tr>
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<tr>
<td>Fe\textsuperscript{3+}</td>
<td>0.038</td>
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<td>Mn</td>
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<tr>
<td>Ca</td>
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<td>Na</td>
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<tr>
<td>K</td>
<td>0.00</td>
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<tr>
<td>(O)</td>
<td>(6)</td>
</tr>
</tbody>
</table>

1: orthopyroxene; 2: clinopyroxene; 3: hornblende; 4: biotite; 5: plagioclase
* Total iron as FeO
Specimen also contains quartz

between coexisting garnet and biotite, wet chemical analyses from several sources were used. Reinhardt’s (1968) analyses indicate that on the average the Fe\textsuperscript{3+}/(Fe\textsuperscript{2+} + Fe\textsuperscript{3+}) ratio in biotite is 2.4% larger than that in coexisting garnet. Lal and Moorhouse (1969) present data which indicate that the Fe\textsuperscript{3+}/(Fe\textsuperscript{2+} + Fe\textsuperscript{3+}) ratio in biotite is 8.4% higher than that of coexisting garnet, and Chinner’s (1960) data indicate a wide range, from 4 to 22%, with an average of 12.4% larger (Fe\textsuperscript{3+}/(Fe\textsuperscript{2+} + Fe\textsuperscript{3+}) in biotite than garnet. The effect on geothermometric measurement of assuming 8.4% more Fe\textsuperscript{3+}/Fe\textsuperscript{\textit{total}} in biotite than in garnet as calculated stoichiometrically, is negligible (apparent change of -1°C using Ferry–Spear calibration). Assuming a 12% average, the difference is only -2°C.

The agreement between the garnet–clinopyroxene and garnet–biotite thermometers is best when all iron is assumed to be Fe\textsuperscript{2+} in garnet, biotite and clinopyroxene for both thermometers (Fig. 7) and the calculations in Tables 2, 4 and 6 are based on this procedure. Treating all iron as Fe\textsuperscript{2+} is actually an empirical correction used to achieve consistency between geothermometers (Fig. 7) and contrasts with the procedure of Raheim and Green (1975), who used calculated Fe\textsuperscript{2+} contents. The pyroxene commonly has larger Fe\textsuperscript{3+}/Fe\textsuperscript{\textit{total}} than coexisting garnet and temperatures up to 100°C lower result if the calculated Fe\textsuperscript{2+} value is used to calculate $K_D$. Further justification for using total Fe rather than Fe\textsuperscript{2+} comes from samples MG-17 and MG-25, whose garnet and clinopyroxene contain no Fe\textsuperscript{3+} as calculated stoichiometrically. Temperatures of 725 and 755°C are in the same range as values calculated by ignoring Fe\textsuperscript{3+} in other specimens (Table 4).

A calibration of the two-pyroxene thermometer, proposed by Powell (1978), yields results which are consistent with garnet–biotite and garnet–clinopyroxene temperatures for the Kapuskasing gneisses. The equations formulated by Kretz (1982) give averaged values in the same range but some pyroxene pairs yield temperatures that are up to 100°C discordant. The Wood and Banno (1973) and Wells (1977) calibrations consistently give temperatures some 200–300°C higher.

Additional results were obtained for rocks containing mineral assemblages without widespread distribution (Tables 2, 4 and 6). In general, oxide, feldspar and ilmenite–pyroxene thermometers yield temperatures some 200–300°C lower than those estimated by the Fe–Mg exchange thermometers. No generalizations regarding other thermometers are possible because of the limited number of occurrences.

### Regional temperature variation

Near-rim garnet analyses generally give results lower by 10–20°C than analyses of the relatively homogeneous interiors. The general analytical strategy was to check all

---

**Fig. 4. Phase relations of mineral assemblages coexisting with tonalitic melt in mafic rocks.** Mineral compositions are generalized from Appendix I for a and c; plotted from Table 5 for b. (a) garnet–clinopyroxene–hornblende compositional triangles shift to the right along M/M+F axis in response to rising temperature; (b) garnet + hornblende react to produce orthopyroxene + clinopyroxene with increasing temperature (or decreasing $a_{H_2O}$ if $P$, $a_{H_2O}$ are independent of $T$); (c) stable mineral assemblages in the orthopyroxene zone.
the area as well as isolated occurrences in the 800°C range near the southeastern margin of the Kapuskasing Zone. Most of the Gt–Cpx–Pl zone is characterized by values above 700°C. Rocks which give apparent lower temperatures, in the 600°C range, are in the southwest corner of the area, where garnet-, biotite-bearing paragneiss schlieren occur as inclusions in tonalitic gneiss.

A high paleotemperature zone is present near the northern end of the Shenango complex (Fig. 2) and a possibly similar relationship exists east of the Nemegosenda Lake complex. The origin of the apparent thermal highs may relate to the late (1100 Ma) intrusive bodies. Samples from the vicinity of the Nemegosenda complex may have been affected by contact metamorphism, metasomatism, or crustal buckling or displacement adjacent to the body. Any of these processes could alter the apparent temperature of adjacent rocks, the first by readjusting Fe–Mg ratios of garnet and clinopyroxene, the second by increasing Ca content of garnet, and the third by displac-

Fig. 6. Vapor-phase-absent-melting curve for paragneiss. Construction as for Fig. 5. Stippled area is VPAM curve for mafic rocks. Note that the two VPAM curves intersect at 6–8 kbar, 775°C for natural compositions.

grains for zoning but exclude near-rim analyses from the average.

The results of the combined garnet–biotite, garnet–clinopyroxene and two-pyroxene thermometers are plotted in Figure 8, which shows an overall temperature range of 580–825°C. The extremes in this range are garnet–biotite temperatures; garnet–clinopyroxene results have a narrower spread (655–815°C). Much of the area without data points is underlain by diorite or tonalite orthogneiss without relevant assemblages.

The apparent temperature distribution of Figure 8 indicates two zones above 800°C in the northern part of
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Fig. 7. Comparison of temperature estimates derived from garnet-biotite and various garnet-clinopyroxene thermometers. The two thermometers are consistent if the Ferry and Spear (1978) and Ellis and Green (1979) calibrations are used and $K_p$ is calculated using the total iron content rather than stoichiometrically-determined Fe$^{2+}$.

Fig. 8. Paleotemperature map of part of the Chapleau-Foleyet area. Temperatures estimated by garnet-biotite (Ferry and Spear, 1978), garnet-clinopyroxene (Ellis and Green, 1979) and two-pyroxene (Powell, 1978) thermometry. Numbers to the left of dash are sample identifiers; numbers to the right are in °C. circles: PG-series; squares: MG-series; triangles: OG-series.

are at variance with temperature data, it is likely that the Fe–Mg ratios in minerals used in the thermometers have equilibrated under different conditions than those at which the assemblages were stable. Alternatively, low water fugacity may have stabilized orthopyroxene at relatively low temperatures. However, at temperatures as low as 650°C and with low water fugacity, the production of partial melts would not be possible (Robertson and Wyllie, 1971). Because migmatite veinlets are ubiquitous in paragneiss in this region, a low-temperature origin is unlikely and retrograde Fe–Mg exchange seems to be a better explanation. Alternatively, biotite may have been in solution in the liquid phase while rocks were at temperatures above the solidus and crystalized upon later cooling to ~650°C. However, there is no textural evidence that this process occurred.

Metagabbros of the Shawmere complex locally consist of the assemblage Opx–Cpx–Hb–Pl (e.g., OG-8, Table 6). Orthopyroxene occurs as homogeneous grains whereas clinopyroxene has two habits; large (up to 5 mm) grains have fine exsolution lamellae, and smaller matrix grains are optically and chemically homogeneous. Lamellae in the large grains are bent and offset, giving the complex grains undulose extinction. Chemically, the host clinopyroxene crystal ($Ca_{0.47}Fe_{0.10}Mg_{0.43}$) is identical in composition to the matrix clinopyroxene. The exsolved lamellae have the composition $Ca_{0.18}Fe_{0.21}Mg_{0.61}SiO_3$. By using a defocussed microprobe beam technique, analyses of the bulk composition of the clinopyroxene megacrysts were obtained. The bulk composition ($Ca_{0.40}Fe_{0.11}Mg_{0.49}$) is slightly poorer in calcium than matrix clinopyroxene. Bohlen and Essene (1978) interpreted pyroxenes with
exsolution lamellae in metamorphosed Adirondack anor-thosite massifs as being of igneous derivation. They derived temperatures both of igneous crystallization (−1100 ± 100°C) and metamorphism (−750°C) by reintegration of pyroxene compositions. The same treatment cannot be applied to the Shawmere metagabbro because orthopyroxene is homogeneous and plots in the metamorphic temperature range on Ross and Huebner's (1975) pyroxene solvus isotherm diagram. Assuming, however, that the exsolved clinopyroxene megacrysts at one time coexisted with a calcium-richer orthopyroxene, a temperature on the order of 1050°C can be estimated for igneous crystallization. Based on the composition of matrix clinopyroxene and homogeneous orthopyroxene, an estimated temperature of metamorphism of 750°C may be obtained from the Ross and Huebner isotherm plot.

**Pressure estimation methods**

Pressure estimates are dependent on the temperatures derived in the previous section. Several pressure-sensitive equilibria are relevant to assemblages with widespread distribution in the Kapuskasing Zone. The compositions of minerals in garnet-pyroxene-plagioclase-quartz parageneses are determined by equilibria, including:

\[
\begin{align*}
\text{CaAl}_{2}Si_{2}O_{8} & \rightleftharpoons \text{CaAl}_{2}Si_{2}O_{6} + SiO_{2} \\
2\text{CaAl}_{2}Si_{2}O_{12} + Mg_{3}Al_{2}Si_{3}O_{12} & \rightarrow 3\text{CaMgSi}_{2}O_{6} + 3\text{CaAl}_{2}Si_{2}O_{6}
\end{align*}
\]

(8)

\[
\text{CaAl}_{2}Si_{2}O_{8} + CaMgSi_{2}O_{6}
\]

(9)

\[
\begin{align*}
2\text{CaAl}_{2}Si_{2}O_{12} & + Mg_{3}Al_{2}Si_{3}O_{12} \\
\rightarrow 3\text{CaMgSi}_{2}O_{6} + 3\text{CaAl}_{2}Si_{2}O_{6}
\end{align*}
\]

(10)

\[
\text{CaAl}_{2}Si_{2}O_{8} + Mg_{3}Si_{2}O_{6}
\]

(11)

\[
\begin{align*}
\text{CaAl}_{2}Si_{2}O_{8} + Mg_{3}Si_{2}O_{6} & \rightleftharpoons 1/3\text{CaAl}_{2}Si_{2}O_{12} + 2/3\text{Mg_{3}Al_{2}Si_{3}O_{12} + SiO_{2}} \\
\text{in garnet} & \text{in garnet} & \text{in garnet}
\end{align*}
\]

(12)

Anorthite activity in plagioclase was estimated by the Al-avoidance model of Perkins and Newton (1981). For MG-20, a visual estimate of the proportions of An_{59} and An_{41} plagioclase was made to estimate a weighted average of An_{48}. Diopside and enstatite activities are derived from data of Wood and Banno (1973). Interaction parameters were used to estimate pyrope and grossular activity in garnet according to Perkins and Newton (1981).

Geobarometers based on (10) and (11) have recently been calibrated by Perkins and Newton (1981); results from mafic gneiss of the Kapuskasing Zone are presented in Table 4. Wood (1977) applied Equilibrium (8) to granulites as a geobarometer. Using a calibration of (8) based on thermochemical data of Helgeson et al. (1978), pressure estimates are seen to be controlled mainly by the formulation of the CaAl_{2}Si_{2}O_{6} activity model. Wood's (1978) formulation leads to a cluster of values in the 4 kbar range whereas Wood's (1979) model produces results averaging −15 kbar. Equilibria (9) and (12) are subject to the same uncertainty.

Wood's (1974) calibration of (12) gives additional pressure estimates for garnet, orthopyroxene-bearing assemblages. Pressure was estimated for a Gt-Si-Pl-Bt-Qz assemblage according to Ghent's (1976) calibration.

The accuracy of the various geobarometers is limited by the uncertainty in probe analyses, greatest where accurate analyses of small quantities of alumina in pyroxene are required ((8), (9), (12)). The low dP/dT slope of (10) and (11) and their use of major components recommend these equilibria as reliable geobarometers.

**Pressure estimates**

Estimates of peak metamorphic pressure depend on the assumption that present mineral compositions are not significantly different than those at the peak metamorphic conditions. In samples with retrograde amphibole, this assumption may not be valid because other minerals, including plagioclase, could have changed their compositions during retrograde reactions, presumably at lower P,T. Such reactions could also have occurred in response to changes in a_17r_6 at peak P,T.

The Perkins and Newton (1981) calibration of (10) yields pressure values generally in the range 5.4−8.4 kbar, averaging 6.3 kbar. The results are plotted on Figure 9 along with values derived from (11) and (12). The average pressure for the area from (11) is 7.7 kbar and individual values are commonly −2 kbar higher than those estimated from (10) for the same or proximal samples. Values from (10) are generally lower than those from (11), either because of different blocking temperatures in different mineral systems or uncertainty in the thermodynamic calibration (Newton and Perkins, 1982). A 6.3 kbar reference line separates areas with above-average apparent values from those with lower values. The line includes a roughly north-south-trending area of relatively high apparent pressure in the central and eastern Kapuskasing Zone (Fig. 9). This pattern supports the preliminary interpretation (Percival, 1981a) that the Kapuskasing Zone is at the base of a tilted, west-dipping crustal section. However, diffusion considerations may preclude this simple interpretation (see below).

Assuming that pressure is a function of depth, approximately 20−25 km of overburden has been eroded from the Kapuskasing zone. Previous estimates of relative uplift of
the Kapuskasing Zone with respect to adjacent regions where 8 km, based on heat-flow data (Cermak and Jessop, 1971) and at least several thousand feet (Thurston et al., 1977). Pressure-sensitive assemblages are lacking in the terranes to the east and west. However, slate some 5 km east of the Ivanhoe Lake cataclastic zone is at relatively low grade (chlorite-muscovite). Although dP/DT is generally not constant with depth (P. H. Thompson, 1977), assuming an average crustal geothermal gradient of -30°C/km, biotite should form at the expense of muscovite-chlorite at about 450°C, corresponding to a depth of about 15 km. Therefore, relative uplift along the Ivanhoe Lake cataclastic zone is estimated to be at least 5 and possibly greater than 10 km, based on metamorphic considerations.

Water activity calculations

Water activity was calculated for rocks containing assemblages corresponding to Mg end-member dehydration equilibria for which P-T estimates were available. Three equilibria have been used for this purpose, including (2):

\[
3\text{MgSiO}_3 + 2\text{CaMgSi}_2\text{O}_6 + \text{SiO}_2
\]

in calcic amphibole in orthopyroxene

\[
= \text{Ca}_2\text{Mg}_3\text{Si}_2\text{O}_8 + 2\text{MgSiO}_3 + \text{SiO}_2
\]

in biotite + orthopyroxene + quartz

\[
\Delta G^0 = -RT \ln K - \Delta V(P - 1)
\]

Activity of solids were estimated by ideal ionic solution models and thermochemical data are from Helgeson et al. (1978) and Haselton and Westrum (1980). Water activity is calculated as \( f_{H_2O} \) as defined by (15), divided by \( f_{H_2O} \) of pure water at \( P, T \) as defined by Helgeson and Kirkham (1974). Derived values are found in Tables 2, 4 and 6.

Equilibrium (13) yields values <0.1; (14) gives results <0.4 and (2) yields some values above 1.0. Values in excess of 1.0 indicate that the minerals were not in equilibrium at the specified P-T conditions. For example, brown hornblende in MG-11 yields \( a_{H_2O} = 0.853 \) whereas actinolite, presumed to be of retrograde origin, yields a mechanically-unsustainable value of 5.5. Green, secondary amphiboles indicate higher \( a_{H_2O} \) in every example (e.g., MG-4, 11, 20), but would yield lower \( a_{H_2O} \) values if a lower temperature were assumed.

No pattern of regional variation in water activity is obvious from values plotted on a map. Values are generally on the order of 0.1 near the western edge of the Kapuskasing Zone but do not show a regional trend for the rest of the area. There is no correlation between calculated water activity and the presence or absence of orthopyroxene. The estimates are in the range of values reported for granulite-facies terranes (e.g., Bohlen and Essene (1976b) for the Adirondacks; Horrocks (1980) for the Limpopo Belt) and for the Broken Hill area of Australia (Phillips, 1980).

Significance of pressure-temperature results

A comparison between P-T estimates for individual samples and probable P-T conditions in the Gt-Cpx-Pl and orthopyroxene zones shows that temperature estimates for rocks containing orthopyroxene are below the Gt-Cpx zone conditions deduced by mineral-melt equilibria. Eight of twenty-five estimates from the Gt-Cpx-Pl zone fall within the Gt-Cpx zone on the P-T diagram; the rest fall below. This pattern suggests that Fe-Mg exchange between mineral pairs continued during cooling and that derived temperature estimates are somewhat retrograde.

For the Conemarra region, Yardley (1977) suggested that temperature above 640°C had homogenized prograde zonation in garnets by intragranular diffusion. Lasaga et al. (1977) estimated that retrograde Fe-Mg exchange between the outer 10-15 μm of adjacent garnet and cordierite grains occurred down to ~450°C. In view of the quenching problem it may be realistic to regard the highest apparent temperatures, in the 800°C range, as closest to peak temperature. The lower values probably record quenching at later times. By this reasoning, the pressure and temperature estimates, based on mineral
systems with probable different blocking temperatures (O'Hara, 1977), may not even record a P-T condition through which the rock passed. Thus it is hazardous to attempt to deduce either a prograde or cooling metamorphic path for the area, based on geothermobarometry.

“High-pressure granulites”

The assemblage almandine garnet–clinopyroxene–plagioclase–quartz is diagnostic of the regional hypersthene zone according to Winkler (1979, p. 260, 267–268). de Waard (1965) and Green and Ringwood (1967) suggested that this assemblage forms as an alternative to orthopyroxene–plagioclase during high-pressure granulate-facies metamorphism. Turner (1981) attaches a different significance to the assemblage, regarding it as transitional from amphibolite to granulite facies based on Binns’ (1964) study. In the present study area, the location of the Gr-Cpx-Pl zone between Hb-Pl±Cpx rocks and orthopyroxene-bearing rocks suggests that it characterizes the amphibolite–granulite facies transition. Although the assemblage is the same as that in the Adirondacks (de Waard, 1965) and temperature conditions were similar (Bohlen and Essene, 1977), the path of metamorphism was different. In the Grenville Province, the development of garnet–clinopyroxene assemblages has been attributed to isobaric cooling of orthopyroxene–plagioclase granulites (Martingole and Schrijver, 1971; Whitney, 1978) whereas in the Kapuskasing Zone, garnet and clinopyroxene formed during prograde reactions.

Conclusions

In the eastern Wawa subprovince and Kapuskasing Structural Zone, three metamorphic zones are defined on the basis of assemblages in mafic rocks. The sequence Hb-Pl, Cpx-Hb-Pl, Gr-Cpx-Pl suggests easterly-increasing grade. Both hornblende- and biotite-bearing rocks in four areas surrounded by the Gr-Cpx-Pl zone contain metamorphic orthopyroxene, representing the highest grade attained. Tonalitic melt is thought to have been responsible for reducing $a_{H_2O}$ in the metamorphic fluid in mafic gneiss to values of 0.5–0.7 at temperatures of $\sim750^\circ$C, at which point Hb-Pl began to produce Gr-Cpx-melt assemblages by vapour-absent melting reactions. Biotite in adjacent migmatitic paragneiss units was stable at these conditions but began to break down to orthopyroxene–K-feldspar-melt by vapour-absent-melting reactions at $\sim770^\circ$C, $a_{H_2O}$ $\sim$0.5. Orthopyroxene first appears in mafic rocks at similar conditions.

Estimates of metamorphic pressure and temperature by various calibrations are 3.1–9.9 kbar, 600–825°C but most values are between 6.0 and 7.3 kbar and 650 and 775°C. The orthopyroxene-zone rocks yield average to below-average values suggesting that the peak-metamorphic distribution of elements has been altered, probably by ionic diffusion during cooling.

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