Seeing through metamorphic overprints in Archean granulites: Combined high-resolution thermometry and phase equilibrium modeling of the Lewisian Complex, Scotland

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Abstract

The Lewisian Complex in northwest Scotland presents a record of the transition from the Neo-Archean to the Paleoproterozoic. However, this record is complicated by a long and varied history after peak metamorphism that has erased and/or partially reset much of the early history of the rocks. Such overprinting is a common feature of Archean granulites and poses a substantial problem when trying to understand the tectonic processes that were active prior to the onset of modern plate tectonics.

By combining careful petrography with phase diagram modeling and a range of exchange thermometers we obtain the peak and retrograde temperature history of the Lewisian Complex from a single, well-preserved, representative sample of garnet-bearing mafic granulite. We present the application of high-resolution electron probe microanalysis (HR-EPMA) to characterize sub-micrometer orthopyroxene exsolution lamellae in clinopyroxene. We discuss ways to mitigate issues associated with HR-EPMA including surface contamination, beam drift, standards, and the need to correct for secondary fluorescence effects. The resulting compositions from our HR-EPMA analyses provide an independent measure of the retrograde temperature conditions and can also be used to back-calculate the compositions of clinopyroxene in the peak assemblage.

We obtain peak metamorphic conditions for the Lewisian of >11 kbar and >1025 °C, and constrain subsequent metamorphic overprints to 850 °C (Grt-Cpx), 590 °C (Opx-Cpx), and 460 °C (Mag-Ilm). These peak and retrograde temperatures span the range of those found in the literature. Whereas recent phase equilibrium studies assume equilibrium among all preserved high-*T* minerals, this study considers microstructural and mineral-chemical evidence for corona formation that reflects post-peak decompression with partial equilibriation at ~850 °C, as recognized in some earlier studies.

Keywords: Lewisian, granulites, HR-EPMA, phase equilibria, Archean, Scotland

INTRODUCTION

The Lewisian Complex of northwest (NW) Scotland comprises a sequence of Archean igneous rocks, which have undergone a series of metamorphic events. At \geq 2.8 Ga, these are not only the oldest rocks in the British Isles, but also have a history spanning the crucial transition from neo-Archean to the Paleoproterozoic, and the initiation of modern-style tectonics.

The Lewisian Complex can be divided into three regions: northern, central, and southern (Peach et al. 1907), and it comprises mainly tonalite-trondhjemite-granodiorite (TTG) orthogneisses with subordinate blocks and lenses of intermediate, mafic, and ultramafic rocks (Sutton and Watson 1950; O'Hara 1961; Rollinson and Windley 1980). This study focuses on the central region, specifically the mafic rocks of Scouriemore (Fig. 1), that underwent (ultra)high-temperature metamorphism during the Archean (Badcallian) regional metamorphic episode (Park 1970; Sills and Rollinson 1987). Despite extensive study since the time of Peach and Horne (Peach et al. 1907), there still exists much

This study focuses on one particularly well-preserved mafic

debate as to the peak metamorphic conditions attained during the Badcallian metamorphic event. Estimates in the older literature are derived from several different geothermobarometers (Wood 1975, 1977; O'Hara and Yarwood 1978; Barnicoat and O'Hara 1979; Savage and Sills 1980; Rollinson and Windley 1980; Pride and Muecke 1980; Rollinson 1981) and span a wide range from 820 to 1150 °C and 6 to 15 kbar (Barnicoat 1983; Sills and Rollinson 1987). The scatter in *P*-*T* estimates is largely due to the one (or more) subsequent metamorphic events that have wholly erased or partially reset the peak mineral compositions (Barnicoat 1983; Sills and Rollinson 1987). Modern phase equilibrium modeling techniques (e.g., de Capitani and Brown 1987; de Capitani and Petrakakis 2010; Connolly 1990; Connolly and Petrini 2002; Powell et al. 1998; Holland and Powell 2011; Green et al. 2016) have been applied to the Badcallian metamorphism (e.g., Johnson and White 2011; White et al. 2017; Feisel et al. 2018; Forshaw et al. 2019), but this approach has been hampered by the large P-Tfield in which the peak mineral assemblage occurs, by domains of partial re-equilibration that exist within samples, and issues relating to partitioning of minor elements among the dominant minerals.

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FIGURE 1. Simplified geological map of the area surrounding Scourie Bay, showing the location of sample S98-14 on Scouriemore (after O'Hara 1961).

granulite sample (Fig. 2) from Scouriemore, NW Scotland. This sample is representative of the relatively Fe-rich metabasites found across the region, which are dominated by garnet and clinopyroxene (cf. Savage and Sills 1980). It exhibits similar features to other samples characterized by the authors (Online Materials¹ A) and to samples analyzed by other studies in the Lewisian (Wood 1975, 1977; O'Hara 1961; Barnicoat and O'Hara 1979; O'Hara and Yarwood 1978; Savage and Sills 1980; Rollinson and Windley 1980; Pride and Muecke 1980; Feisel et al. 2018; Johnson and White 2011). However, samples in the literature generally show higher degrees of amphibolite-facies retrogression and hydration compared with the rock sample on which we focus. Breakdown microstructures (i.e., post-peak net-transfer reactions) preserved within this rock can be linked to a succession of metamorphic processes and thus be used to determine the P-T conditions of the retrograde path. Importantly,

FIGURE 2. QEMSCAN (left-hand side) and plane-polarized optical thin section image (right-hand side) of S98-14. Black dashed line denoted border between left and right, and white dashed line shows the outline of a pyroxene grain that crosses between the two images. Locations of EPMA analyses and transects are noted in black on the ppl image. QEMSCAN image colors: red = garnet; green = clinopyroxene; blue = plagioclase; purple = orthopyroxene; orange = ilmenite; reddish-orange = magnetite; gold = pyrite. Optical image colors: reddishorange = garnet; green = clinopyroxene; white = plagioclase; brown = orthopyroxene; black = oxide or altered plagioclase.

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this allows us to back-calculate the compositions of the peak assemblage, here interpreted to be only garnet and clinopyroxene (±plagioclase/quartz). The microstructures of particular importance to this study, as also described in Barnicoat (1983) and Sills and Rollinson (1987); are the breakdown of peak garnet (Fig. 3) to form corona textures composed of plagioclase and orthopyroxene (±magnetite-ilmenite), as well as the exsolution of (peak) coarse granoblastic clinopyroxene to form lamellae of orthopyroxene, which do not reach the edge of the host grain.

We combine phase equilibrium modeling with the careful application of exchange thermometers applied to these microstructures to provide a more precise record of Badcallian peak metamorphism and a better understanding of Archean lower crustal *P-T* conditions. In the process we present an adapted analytical technique using HR-EPMA that allows accurate determination of sub-micrometer orthopyroxene (Opx) exsolution lamella in clinopyroxene (Cpx). Abbreviations here and throughout are after Whitney and Evans (2010).

To analyze phases below $\sim 5 \ \mu m$ by EPMA, low accelerating potential and a field emission gun source are required. It is a known issue that using accelerating potentials below $\sim 10 \ keV$ induce multiple analytical issues, including problems with using the *L* X-ray transition lines of transition metals, increased sensitivity to surface contamination, enhanced sample damage, electron beam drift over the course of the analysis, and issues with secondary fluorescence (Gopon et al. 2013, 2017; Llovet et al. 2016; Armstrong 2012). We present a way that uses careful standard matching, combined with a correction scheme for secondary fluorescence, to accurately determine the compositions of micrometer and sub-micrometer orthopyroxene exsolution lamellae.

The analysis of sub-micrometer lamellae allows determination of a two-pyroxene retrograde temperature, but more importantly allows us to recombine the exsolved orthpyroxene into the clinopyroxene host to determine the composition of the "proto-Cpx" (defined as the original composition of the Cpx at peak *P-T*). This proto-Cpx can then be used to determine the peak





Badcallian temperature using the graphical pyroxene thermometer of Lindsley (1983). These widely applicable techniques allow us to see through subsequent metamorphic overprints to determine the peak granulite P-T conditions.

Methods

Sample selection and petrography

Garnet-bearing mafic granulites were selected from a suite of >100 samples collected by the authors since 1991 from the central region of the Lewisian Complex. At Scouriemore, the mafic bodies range from garnet-clinopyroxene-bearing granulites with minor plagioclase through to two-pyroxene granulites. In addition to any record of re-equilibration during slow cooling from peak conditions, they have been subjected to varying degrees of retrograde overprinting during subsequent events. Thin sections from Scouriemore were noted to contain retrograde hydrous phases; principally amphibole and biotite but epidote and chlorite were observed in heavily retrogressed mafic-intermediate gneisses (Online Materials1 A). After re-examination of all samples to assess the state of equilibrium, effects of deformation, and evidence for reaction relationships among minerals, a sample (S98-14) from Scouriemore (locality 3 of O'Hara 1961), showing excellent preservation of petrographic relationships and minimal retrogression, was selected for detailed study. This representative, and near totally anhydrous, sample shows similar features to those present in all samples characterized (Online Materials1 A), as well as those found in the literature (O'Hara 1961; Rollinson and Windley 1980; Feisel et al. 2018; Johnson and White 2011).

QEMSCAN analysis was conducted on a FEI Wellsite SEM at Rocktype Ltd. (Oxford, U.K.), at a resolution of 10 μ m across an entire thin section of S98-14 to determine the phases present and their modal abundance. A further high-resolution scan (1 μ m pixel size) was conducted surrounding a well-preserved garnet and its corona. The modal abundance from the 10 μ m scan, combined with microanalysis, was used to construct the bulk composition of the rock.

A grain size analysis of sample S98-14, using the approach in Abrams (1971), was conducted to quantify the size and abundance of the major minerals present. This involved drawing random lines across five thin sections from the same hand sample of S98-14 and measuring the diameter of the long and short axes of each crystal that the lines intersected. The diameter of the long and short axis was then used to calculate the area of the corresponding ellipse, which is assumed to estimate the volume of the crystal. The diameter of a sphere of equivalent volume to this was then taken to represent the grain size.



FIGURE 3. High-resolution QEMSCAN image (1 µm pixel size) of a corner of a representative garnet corona (see inset for location). Note the association of magnetite and ilmenite within the corona and that the orthopyroxene present in the matrix in the top right of the image is in fact connected to the garnet corona by orthopyroxene present along the Cpx-Cpx grain boundary. Image colors: red = garnet; green = clinopyroxene; blue = plagioclase; purple = orthopyroxene; orange = ilmenite; reddish-orange = magnetite.

Electron probe microanalysis (EPMA)

EPMA analyses were conducted in the Department of Earth Sciences, University of Oxford, U.K., on a CAMECA SXFive-FE. The CAMECA Peaksight Version 6.2.0.2064 software was used for quantitative analyses.

15 keV electron microprobe analysis. Standard EPMA analysis used an acceleration voltage of 15 keV, a beam current of 20 nA, a spot size of 1 µm, as well as 30 s on-peak and 15 s background count times. For all silicate minerals silicon, titanium, aluminum, iron, manganese, magnesium, calcium, sodium, and potassium were analyzed using the following X-ray lines and diffracting crystals: SiKa (TAP), TiKa (LPET), AlKa (TAP), FeKa (LLIF), MnKa (LLIF), MgKa (TAP), CaKa (PET), NaKa (TAP), KKa (PET). Additionally, CrKa (LLIF) and NiK α (LLIF) were analyzed in orthopyroxene. For oxide minerals sodium, magnesium, aluminum, silicon, calcium, titanium, chromium, manganese, iron, and nickel were analyzed using the following X-ray lines and diffracting crystals: NaKa (TAP), MgKa (TAP), AlKa (TAP), SiKa (TAP), CaKa (PET), TiKa (PET), CrKa (LLIF), MnKa (LLIF), FeKa (LLIF), NiKa (LLIF). Oxygen was calculated by stoichiometry for silicate and oxide analyses. Standards for silicate analyses included: albite for Si, Al, and Na; orthoclase for K; andradite for Fe; 99.9% purity manganese metal for Mn; 99.99% purity chromium metal for Cr; 99.999% purity nickel metal for Ni; synthetic MgO for Mg; wollastonite for Ca; and synthetic TiO2 for Ti. Standards for oxide analyses included albite for Si, Al, and Na; synthetic MgO for Mg; magnetite for Fe; orthoclase for K; andradite for Ca; 99.99% purity chromium metal for Cr; 99.999% purity nickel metal for Ni; and 99.9% purity manganese metal for Mn. Point analyses provided an overview of the modal oxide weight percentages for each mineral, while line scans identified any heterogeneity within minerals expressed as elemental zoning between cores and rims. Analytical uncertainty for individual EPMA analyses is reported in the Online Materials1 B and is based on the counting statistics of the detectors. Because we report averages of multiple analyses in Table 1, the uncertainty from counting statistics is in the third or fourth decimal place. Because we only report to two decimal places in Table 1 the uncertainty is not shown there, but we report the standard deviation of the averaged analyses.

High-resolution EPMA (HR-EPMA). To analyze the small analytical volumes of the $<5 \mu$ m pyroxene exsolution lamella, lower accelerating potentials (<10 keV) were required. An accelerating voltage of 7 keV was selected to minimize the analytical volume of the analysis region (~400 nm diameter, ~500 nm penetration at 7 keV; Fig. 4), while still providing enough overvoltage to excite the CaKa line (Gopon et al. 2013). The estimates of analytical volume (Fig. 4) were determined using a Monte Carlo simulation of electron scattering using the CASINO (v2.1) software (https://www.gel.usherbrooke.ca/casino) and modeled using the average composition of clinopyroxene in our thin section (Table 1). Lower accelerating potentials were attempted to decrease the interaction volume further but were noted to give incorrect Ca concentrations of secondary standards due to low overvoltage of the CaKa X-ray transition below 7 keV. A current of 10 nA was used, with 30 s on-peak and 15 s background count times. During analyses a liquid nitrogen cooled cold plate was used to keep contamination levels to a minimum (Gopon et al. 2015).

Iron, sodium, calcium, titanium, aluminum, silicon, and magnesium were analyzed using the following X-ray lines and diffracting crystals: $FeL\alpha$ (PC1), NaK α (TAP), CaK α (LPET), TiK α (LPET), AlK α (TAP), SiK α (TAP), MgK α (TAP). Oxygen was calculated by stoichiometry. Standards used were: UW-7239 hedenbergite for Fe and Ca; University of Michigan synthetic ferrosilite for Si; University of Oxford synthetic MgO and TiO₂ for Mg and Ti, respectively; albite for Na and Al in all samples.

Measurements of lamella composition were performed in a series of points along the length of each lamella. Stage reproducibility (the ability of stage automation to return to the desired site) below 1 µm is not possible on electron microprobes, therefore each point was acquired without the aid of any stage automation. The median composition of these points was determined, and any data points that were significant outliers (outside of 5 relative percent) from this median composition of a lamella were considered erroneous due to beam drift and removed. Also, any analysis with an oxide total of more than 102% and <98% was considered erroneous and removed. Analytical uncertainty for individual and averaged 7 keV analyses are reported in the Online Materials¹ B. It should be noted that the number of analyses averaged to generate the host and lamellae pyroxene compositions are much less than in the 15 keV portion of this study, and the uncertainty from counting statistics therefore lies in the second and third decimal place (Online Materials¹ B).

Mitigating issues with HR-EPMA analysis. The combination of a small beam size from a field emission-sourced microprobe, together with the low accelerating

Mineral	Aineral Cpx 1									Cpx 2						
Location No. of analyses	Cpx Hos 6	t Op>	Opx Lam (3 µm) 9		Cpx Recomb N/A	Garnet Pair 20		Cpx Host 11	Opx Lam (5 µ 9		μm)	Cpx Recomb N/A	Garnet Pair 20			
Analysis type	7 keV	St.dev.	7 keV	St.dev.		15 keV	St.dev.	7 keV	St.dev.	7 keV	St.dev.		15 keV	St.dev.		
SiO ₂	50.25	0.39	51.26	1.86	50.33	39.07	0.13	50.84	0.20	51.17	1.23	50.87	39.22	0.10		
TiO ₂	0.70	0.06	0.00	0.12	0.64	0.11	0.01	0.55	0.02	0.08	0.06	0.51	0.08	0.01		
Al ₂ O ₃	5.84	0.28	3.67	0.19	5.67	22.14	0.06	4.72	0.17	3.70	0.22	4.64	22.21	0.07		
Cr ₂ O ₃	-	-	-	-	-	-	-	-	-	-	-	-	-	-		
Fe_2O_3	-	-	-	-	-	-	-	-	-	-	-	-	-	-		
FeO	7.76	0.27	24.26	0.26	9.07	22.59	0.16	8.46	0.46	25.54	0.72	9.80	22.51	0.15		
MnO						0.84	0.04						0.84	0.03		
MgO	11.95	0.26	20.16	0.69	12.61	9.14	0.08	11.81	0.11	19.05	0.99	12.38	8.96	0.07		
CaO	22.39	0.24	0.16	0.05	20.62	6.03	0.07	22.32	0.34	0.18	0.05	20.59	6.12	0.07		
CaO (uncorr.)	-	-	0.45	0.05	20.64	-	-	-	-	0.48	0.05	20.61	-	-		
NiO	-	-	-	-	-	-	-	-	-	-	-	-	-	-		
Na ₂ O	0.85	0.05	0.01	0.02	0.79	0.01	0.01	0.79	0.03	0.01	0.02	0.73	0.01	0.01		
K ₂ O	-	-	-	-	-	0.00	0.00	-	-	-	-	-	0.00	0.01		
Total	99.74		99.52		99.73	99.93		99.49		99.73		99.52	99.95			
Si	1.87		1.93		1.87	2.98		1.90		1.93		1.90	2.99			
Ti	0.02		0.00		0.02	0.01		0.02		0.00		0.01	0.00			
Al	0.26		0.16		0.25	1.99		0.21		0.16		0.20	2.00			
Cr	-		-		-	-		-		-		-	-			
Fe ³⁺	0.03		0.00		0.03	0.04		0.02		0.00		0.01	0.02			
Fe ²⁺	0.21		0.76		0.26	1.40		0.25		0.81		0.30	1.42			
Mn	-		-		-	0.05		0.00		0.00		0.00	0.05			
Mg	0.66		1.13		0.70	1.04		0.66		1.07		0.69	1.02			
Ca	0.89		0.01		0.82	0.49		0.89		0.01		0.83	0.50			
Ni	-		-		-	-		-		-		-	-			
Na	0.06		0.00		0.06	0.00		0.06		0.00		0.05	0.00			
К	-		-		-	0.00		0.00		-	-	-	0.00			
Sum	4.00		3.99		4.00	8.01		4.00		3.98		4.00	8.00			
X _{Mg}	0.76		0.60		0.73	0.42		0.73		0.57		0.70	0.41			

 TABLE 1. Low 7-keV EPMA analyses (wt% oxide and cations per formula unit) for host Cpx, lamella Opx (both the CaO values before and after the secondary fluorescence), and recombined "proto-Cpx" (using the corrected CaO values)

Notes: Normal 15-keV EPMA analyses (wt% oxide and cations per formula unit) for garnet pairs and mean Mag/Ilm/Opx/Grt/Pl analyses. Bulk composition determined by combining these mineral analyses with QEMSCAN modes. "--" denotes elements not analyzed in a particular mineral. All iron is assumed to be FeO in this table, with the exception of the bulk composition where Fe_2O_3 was calculated in minerals using the method of Droop (1987). (Continued on next page)



FIGURE 4. Backscatter electron image of pyroxene exsolution lamella (bright lines) in S98-14, overlain with the modeled electron interaction volume of a 7 keV (dark gray), 15 keV (orange), and 20 keV (blue) electron beam in orthopyroxene, using the CASINO (v 2.1) electron interaction modeling software. Image was acquired at 20 keV on a FEI Quanta650-FEG-SEM.

potentials necessary for analysis of sub-micrometer phases, introduces several issues that must be addressed (Gopon et al. 2013, 2015). These include but are not limited to problems with the generation of low-energy X-ray lines, difficulty in aiming the electron beam, increased sample damage and enhanced sensitivity to surface contamination, and issues with secondary fluorescence.

Issues at low accelerating voltage due to bonding effects changing the X-ray generation characteristics of a sample (especially for the *L* lines of transition metals), have been noted by several researchers (McSwiggen 2014; Heikinheimo et al. 2016; Buse and Kearns 2018), with numerous techniques being developed to mitigate these issues. In previous studies, the authors used the FeL/ lines for low-voltage work for measurement of Fe in Fe-Si alloys (Gopon et al. 2013, 2017). However, because of the low-count rates of this X-ray line, the relatively low abundance of Fe in these samples compared to the previous study, and the presence of appreciable amounts of Mg in pyroxenes (the II MgKa² peak interferes on the FeL/ peak), the FeLa peak was used instead in combination with careful standard matching for the analysis of iron.

Environmental and user-induced vibrations were noted to cause shifts in the beam position during analysis. Therefore, analyses were conducted in the evening, to reduce the effect of environmental vibrations and motion of the machine operator was limited during analysis. Spectrometer motion during the analysis was also noted to cause the beam to drift (up to ~50 nm for the parameter used here) from the intended analysis position. This drift is thought to be caused by electromagnetic interference from the spectrometer motors (personal communications with CAMECA Inst.) and has been noted in all FEG-EPMAs the authors tested (four CAMECA instruments and four JEOL). CAMECA instruments were found to generally have half of the drift compared to other manufacturers (Gopon et al. 2014). Therefore, an effort was made by the authors to limit spectrometer motion during the course of the analysis and the spectrometers move and count in tandem. Secondary standards were run at the beginning of each day of measurement to test the validity of our calibrations.

In HR-EPMA we use a highly focused electron beam at low accelerating potentials. This means that while the same number of electrons interact with the sample, as in more traditional EPMA, the flux is much higher. This increased flux means that the potential for sample damage is also increased (Gopon et al. 2013). For this reason, a relatively low beam current of 10 nA was used in our analyses. An additional complication of operating at low accelerating potentials is that the smaller interaction volume means that any surface contamination will have a larger effect on the resultant analysis, as well as have a greater proportional effect in de-

TABLE 1.—CONTINUED

Mineral	Mag		llm		Орх		Grt		PI		Bulk comp.
Location	Mean										
No. of analyses	5		32		246		299		381		
Analysis type	15 keV	St.dev.									
SiO ₂	0.06	0.01	0.08	0.09	50.74	0.63	38.88	0.33	53.19	0.51	46.66
TiO ₂	0.31	0.28	49.81	0.85	0.08	0.09	0.08	0.04	0.01	0.02	0.07
AI_2O_3	0.43	0.04	0.21	0.04	3.50	0.59	22.06	0.18	29.33	0.43	14.56
Cr ₂ O ₃	3.10	0.34	0.82	0.07	0.06	0.02	-	-	-	-	-
Fe ₂ O ₃	-	-	-	-	-	-	-	-	-	-	0.82
FeO	88.00	0.61	45.38	0.96	23.50	0.88	23.44	1.14	0.18	0.23	15.16
MnO	0.03	0.02	1.71	0.71	0.32	0.07	0.84	0.09	0.00	0.01	-
MgO	0.05	0.02	0.45	0.55	21.21	0.83	8.36	0.79	0.03	0.11	10.59
CaO	0.07	0.01	0.08	0.07	0.36	0.12	6.51	0.49	12.05	0.51	11.30
CaO (uncorr.)	-	-	-	-	-	-	-	-	-	-	-
NiO	0.13	0.02	0.02	0.01	0.04	0.02	-	-	-	-	-
Na ₂ O	0.00	0.02	0.01	0.02	0.01	0.01	0.01	0.01	5.04	0.22	0.82
K ₂ O	0.00	0.01	0.01	0.01	0.00	0.01	0.00	0.01	0.17	0.19	0.02
Total	92.17		98.58		99.81		100.18		99.99		100.00
Si	0.00		0.00		1.90		2.98		2.41		
Ti	0.01		0.95		0.00		0.00		0.00		
Al	0.02		0.01		0.15		1.99		1.57		
Cr	0.10		0.02		-		-		-		
Fe ³⁺	1.86		0.07		0.00		0.00		0.00		
Fe ²⁺	1.00		0.90		0.74		1.50		0.01		
Mn	0.00		0.04		0.01		0.05		0.00		
Mg	0.00		0.02		1.18		0.95		0.00		
Ca	0.00		0.00		0.01		0.53		0.59		
Ni	0.00		0.00		-		-		-		
Na	0.00		0.00		0.00		0.00		0.44		
K	0.00		0.00		0.00		0.00		0.01		
Sum	3.00		2.00		4.00		8.02		5.03		
X _{Mg}	_		_		0.62		0.39		_		

celerating the electron beam (Gopon et al. 2015). To minimize this contamination all samples were thoroughly cleaned in a series of alcohol baths, a liquid nitrogen cold trap was used to minimize carbon buildup during analysis, and oil-free scroll pumps were used in the instrument.

Secondary fluorescence can potentially have a negative effect on the accuracy of EPMA measurements due to the primary generated X-rays exciting atoms outside of the intended excitation volume and generating additional spurious X-rays (Llovet et al. 2012; Bastin et al. 1984). For the example of the analysis of a 2 μ m wide Ca-free Opx lamella in the center of a Cpx, PENELOPE modeling shows that 0.30 wt% CaO would be detected in the Opx, which would be entirely due to secondary fluorescence. Therefore, analyzing lamellae of varying witht would have the effect of changing the apparent Ca concentration (Fig. 5). Note error bars in Figure 5 are determined based on the counting statistics for the simulated X-rays that reach our simulated detector, which was modeled using the size and geometry of our SX5-EPMA.

In the pyroxenes studied, all elements measured were investigated for issues of secondary fluorescence using X-ray generation modeling software (Llovet and Salvat 2016). Of the elements analyzed, only Ca was deemed to have an appreciable secondary fluorescence influence. Because the Ca content has the largest influence on the temperature produced by the thermometry methods used (Fig. 5; Lindsley 1983), a set of corrections was developed to adjust the Ca measurements for the influence of secondary fluorescence.

These corrections involved using the pyPENELOPE software (Pinard et al. 2010), to model the influence of secondary fluorescence on the composition measured by EPMA. The software allows the user to define complex compositions and geometries, including that of the lamella in relation to its host, to simulate the movement and generation of electrons and X-rays in the analytical target, its neighboring phases, and the detector. Tailored simulations using the uncorrected pyroxene compositions, lamellar widths and orientations, as well as spectrometer geometries were run on each lamella-host pair to properly remove the excess Ca measured in the low-Ca pyroxene. For each host-lamella pair a simulation was run for the lamella composition modeled as a bulk material and the lamella composition modeled as an appropriately sized lamella (derived from BSE images; Fig. 6; Table 1) in the host composition. The difference in the modeled Ca X-ray intensity from the lamella simulation and the bulk simulation was then used to determine the excess Ca attributed to secondary fluorescence. An adjustment was then made to the uncorrected lamella analysis to remove this excess Ca (Table 1).

Thermobarometry

This study attempts to use a combination of thermobarometric techniques, alongside careful petrography, to better constrain the peak metamorphic and retrograde conditions experienced by the Lewisian Complex.

Conventional thermometry. First, conventional Grt-Cpx thermometry, using the calibrations of Powell (1985) and Ravna (2000), was completed using an average of 20 analyses from the middle of the garnet transcets from standard EPMA analysis and proto-Cpx compositions (see "Single proto-Cpx thermometry" section). Garnet core analyses were used because they are assumed to be least susceptible to equilibration through diffusion (Caddick et al. 2010). This was followed by Grt-Cpx thermometry from the rims of garnet and clinopyroxene, which should provide a minimum temperature for the end of corona formation. Two-pyroxene thermometry



FIGURE 5. Influence of lamellar width on apparent CaO concentration and subsequent two pyroxene (2-pyx) temperature using the QUILF program, for a (Ca free) Opx exsolution lamella in a Cpx grain. Modeled using PENELOPE and spectrometers at 45° to the lamellar orientation and 45° take-off angle, and the compositions listed in Figure 1 with the exception of Ca, which, to highlight the issue, was modeled as 0.00 wt%. (Andersen et al. 1993) was applied to single crystals using the lamella and host compositions obtained from HR-EPMA analysis. This single crystal two-pyroxene thermometer should record the temperature of pyroxene exsolution on the retrograde path. Finally, Mag-Ilm thermometry, using the calibrations of Carmichael (1966), Powell and Powell (1977), and Spencer and Lindsley (1981), was calculated using the ILMAT spreadsheet of Lepage (2003) and oxide compositions from standard (15 keV) EPMA analyses. The uncertainty in the temperature derives mostly from the equations in the thermometer itself, and we therefore report the uncertainty of the techniques as cited in the original papers (Table 2). The estimated uncertainty of the Ravna (2000) thermometer is higher than the rest, as it is the only work that did not estimate error from equations, but instead used the deviation from experimentally made mineral pairs. The largest uncertainty of the Mag-Ilm thermometer comes from the experimental calibrations of ulvospinel and ilmenite activities. We therefore report the uncertainty of Mag-Ilm as the range of temperatures obtained from using the activities calculated by Carmichael (1967), Anderson (1968), Spencer and Lindsley (1981), and Stormer (1983).

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Phase equilibrium modeling. Phase equilibrium modeling presents an alternative method for determining the peak P-T conditions of granulite facies rocks. The recent development of activity-composition (a-X) relations for minerals in upper amphibolite and granulite facies mafic rocks (Green et al. 2016), has allowed P-T paths to be constrained using this technique, for several high-grade terranes including the Lewisian. Observed mineral assemblages are matched to those on thermodynamically predicted phase diagrams to find the P-T conditions of formation. Phase diagrams were constructed to provide a pressure constraint to the temperatures obtained from



FIGURE 6. Backscattered electron image showing the exsolution of orthopyroxene (Opx), from a clinopyroxene (Cpx) host. Note the dashed line, which shows the extent of lamellar growth.

exchange thermometers. In our sample, these were produced in the 9-component Na₂O-CaO-K₂O-FeO-MgO-Al₂O₃-SiO₂-TiO₂-O₂ (NCKFMASTO) system using Theriak/Domino version 11/02/2015 (de Capitani and Petrakakis 2010) and the internally consistent data set of Holland and Powell (2011; update ds62, 06/02/2012). Note that H₂O was excluded from the system, because the selected sample contains no primary hydrous phases. The following activity-composition (*a-X*) relations were used: melt, clinopyroxene (Green et al. 2016); garnet, orthopyroxene, and biotite (White et al. 2014); epidote (Holland and Powell 2011); magnetite-spinel (White et al. 2002); ilmenite-hematite (White et al. 2000); plagioclase and K-feldspar (Holland and Powell 2003). Pure phases included quartz and rutile.

Effective bulk compositions were calculated by combining modal proportions of phases, from QEMSCAN, with representative mineral compositions, from EPMA. EPMA-derived mineral compositional analyses (in wt% oxide) were converted to molecular mineral formulas using a standard number of oxygen atoms per formula unit. The proportion of Fe^{3+} in pyroxene and garnet was determined using the method of Droop (1987). For consistency, when building effective bulk compositions, we disregarded any minor elements from EPMA analyses that were not incorporated in a mineral's *a-X* relation. (e.g., Ti in Cpx or Mn in Grt). This method allows a petrographically based assessment of which phases are participating in any specific equilibration process, and therefore of determining an effective bulk composition for different stages of the sample's evolution, on petrological criteria.

The general assumption in phase equilibrium modeling is that the appropriate composition to use is the total bulk composition of the sample. This will be the case if equilibrium is achieved for major elements among all minerals present, and this is most likely at peak granulite conditions. If retrograde processes have subsequently altered the mineral compositions, or changed the mineral assemblage, the total rock composition is still applicable if the whole-rock system is essentially closed. As described earlier, the sample in question is an anhydrous rock with a distinctive microstructure of finer-grained coronas composed of plagioclase and orthopyroxene surrounding garnet. Therefore, we first calculate a phase diagram for the total bulk composition to constrain the pressures for the peak conditions in the sample, before corona development. Second, after demonstrating that certain domains in the rock, i.e., the core regions of coarse-grained clinopyroxene, were isolated from the retrograde reaction that formed the coronas, we investigate the effect of a bulk composition modified by a varying contribution of clinopyroxene, to determine the limits on pressure for corona formation.

The importance of bulk composition information for determining the stability limits of mineral assemblages was emphasized by Powell and Holland (2008). However, the uncertainty on the boundaries of mineral assemblage fields is not often assessed. Here we test the sensitivity to bulk composition variation using a Monte-Carlo type randomization that perturbs the volume percent of each mineral with a $\pm 20\%$ relative uncertainty, following the approach of Palin et al. (2016a). For sample S98-14, 20 new bulk compositions were created, and 20 phase diagrams produced to assess the uncertainty on certain critical assemblage field boundaries. We take the range in field boundaries as the best estimate of pressure uncertainty.

Single proto-Cpx thermometry. Frost and Chacko (1989) suggested that "the best way to recover high temperatures in granulite terrains is through the reintegration of exsolution lamellae or the use of relict mineral assemblages." Following this approach, we attempt to calculate the original pyroxene composition, prior to exsolution, using a two-stage process. First, the percentages of host and lamella in the crystal were determined from backscattered electron images of the high-Ca pyroxene grains using the thresholding tool in the freely available ImageJ Software (https:// imagej.nih.gov/ij/). The recombined proto-Cpx composition was then calculated using these percentages, together with the average lamella and host compositions from HR-EPMA analysis. Single pyroxene thermometry was then conducted on these recombined analyses using the experimental calibrations of Lindsley (1983).

Method	Reference	Analyses used	Uncertainty	Cpx Crystal 1 T (°C)	Cpx Crystal 2 T (°C)	Other T (°C)	Average T (°C)
Single pyroxene	Lindsley (1983)	Cpx recomb	±50 °C	1025	980	_	1003
Phase equilibrium modeling	Green et al. (2016)	Grt; Cpx; Opx; PI; Mag; Ilm	±150 ℃	-	-	950	950
Grt-Cpx	Powell (1985)	Cpx recomb; Grt pair	±20 °C	885	892	-	884
·		Cpx rim avg; Grnt rim avg	±70 °C	-	-	755	
	Ravna (2000)	Cpx recomb; Grt pair	±100 °C	859	828	-	846
		Cpx rim avg; Grnt rim avg	±70 °C	-	-	670	
Орх-Срх	Andersen et al. (1993)	Opx lam (uncorr Ca); Cpx host	±16 °C	742	737	-	740
	Andersen et al. (1993)	Opx lam (corr Ca); Cpx host	±16 °C	590	594	-	592
Mag-Ilm	Powell and Powell (1977)	Mag; Ilm	±12.5 °C	-	-	382	382
-	Spencer and Lindsley (1981)	Mag; Ilm	±13 °C	-	-	508	503
	Anderson and Lindsley (1985)	Mag; Ilm	±14.5C	-	-	509	502

This thermometer is graphical and requires one to plot the clinopyroxene composition in ternary pyroxene space, using a specific set of calculations to determine wollastonite, enstatite, and ferrosilite contents. These calculations can be completed using the pyroxene component algorithm in the QUILF program (Andersen et al. 1993). The uncertainties of the temperatures obtained using this technique are taken from the original paper.

A simpler but flawed approach to determining the "bulk" clinopyroxene composition would be to use a large, defocused electron beam that would analyze the host and lamella simultaneously. This method was attempted and deemed to give erroneous numbers (see Online Materials¹ B) because it ignores the assumption in the matrix corrections of EPMA that only one phase is analyzed at a time, as well as analyzing different proportions of host and lamellae in each analysis spot.

RESULTS

Petrographic and modal analysis

QEMSCAN analysis of S98-14 determined that the thin section is composed of (in area%) 35.6% garnet, 35.6% clinopyroxene, 11.1% orthopyroxene, 14.6% plagioclase feldspar, 0.14% quartz, 0.19% magnetite, and 0.11% ilmenite. Hydrous minerals, including biotite and hornblende, as well as accessory pyrite, are found in small quantities (<1%). Grain size analysis of the four major minerals (Fig. 7) determined that garnet ranges in size from 0.5–3.5 mm (2.1 mm average), clinopyroxene ranges from 0.5–2.5 mm (0.9 mm average), and orthopyroxene and plagioclase feldspar do not exceed 0.5 mm in size (0.21 and 0.24 mm average). The spread and uneven distribution of garnet sizes is due to the comparatively small number of (large) garnets and the difficulty separating individual garnet grains from the large garnet aggregates in the thin sections considered.

Garnet and clinopyroxene occur within a coarse granoblastic texture, with relatively smaller clinopyroxene grains sometimes appearing as inclusions in garnet (Fig. 2). Plagioclase feldspar is generally found in contact with garnet as part of a corona texture (cf. O'Hara 1961; Barnicoat 1983; Johnson and White 2011; Fig. 3). A few examples of plagioclase not in contact with garnet can be found throughout the thin section, but these still occur as circular masses, which we interpret as surrounding a garnet occurring outside the plane of the thin section. Orthopyroxene occurs in two habits, as granular minerals (0.1-0.5 mm) generally found in contact with plagioclase feldspar on the outer edges of the corona textures, and as exsolution lamellae (<10 µm) within granoblastic clinopyroxene. The lamellae of orthopyroxene do not reach the edge of most of the host clinopyroxene crystals (Fig. 6), which we take as an indicator of exsolution of the original pyroxene in a closed system. Rare examples of orthopyroxene not adjacent to plagioclase are found, but are interpreted as being in contact with plagioclase in the third dimension or noted to be connected with the corona textures via narrow bands of orthopyroxene. Oxide minerals are relatively rare, but are exclusively contained within the plagioclase portion of the corona textures (Figs. 2 and 3). Oxides are mostly magnetite and ilmenite and are always found in contact with each other, with ilmenite generally being volumetrically smaller. Quartz is generally found on grain boundaries in coronas, and in rare cases as inclusions. The general mineral association within the corona textures is a core of garnet, surrounded by a corona of plagioclase feldspar, oxide minerals found within the plagioclase ring, and granular orthopyroxene located on the edge of the plagioclase ring away from the garnet (Fig. 3).



FIGURE 7. Grain size analysis of the 4 major minerals present in sample S98-14, taken as a representative sample for the metabasic rocks at Scouriemore.

EPMA

Table 1 shows a compilation of the mean weight percent oxide analyses and calculated cations per formula unit (cpfu) taken during both the 7 and 15 keV analytical sessions. The full analyses from which these averages are derived are included in the Online Materials1 B. Error estimates of individual analyses are reported in the Online Materials¹ B and are not shown in Table 1, due to the large number of analyses combined to generate each value, making the error from counting statistics smaller than the significant digits reported. All minerals show minimal withingrain compositional variation: plagioclase is labradoritic [X_{An} = Ca/(Ca+Na+K) = 0.59; matrix-orthopyroxene has $X_{Mg} = Mg/$ $(Mg+Fe^{2+}) = 0.63$ and Al = 0.17 cpfu; magnetite has Cr = 0.10 cpfu; ilmenite has Mn = 0.04 cpfu and Cr = 0.02 cpfu; garnet is dominantly almandine (49%), with lesser pyrope (31%), grossular (18%), and spessartine (2%) components. Line scans across two garnet crystals elucidate slight heterogeneity, with a <0.2 wt% increase in CaO and decrease in MgO from core to mantle, transitioning into more moderate (~0.5 wt%) increase and decrease at the rim in contact with plagioclase coronas (Fig. 8). It is noted that the compositions do not plateau in the core of the garnet. Proto-Cpx has $X_{Mg} = 0.70$, Na = 0.06 cpfu, Al = 0.20 cpfu, and Ca = 0.82 cpfu, classifying as diopside-augite (Morimoto 1988). When comparing Na, Ca, and Fe contents of the cores and rims of matrix clinopyroxene and clinopyroxene included in garnet, slight compositional heterogeneity exists (Fig. 9; Online Materials¹ B). Rims are typically Na-poor and Ca-rich compared to matrix cores and included clinopyroxene; while recombined proto-Cpx is relatively Ca-poor and Fe-rich. The heterogeneity in clinopyroxene is due to varying amounts of host clinopyroxene and orthopyroxene lamellae in the analytical volume.

Thermobarometry

Table 2 shows the results of the application of different thermometers to S98-14. A range of temperatures was produced depending on the mineral system chosen for analysis.

Grt-Cpx thermometry, using the calibrations of Powell (1985) and Ravna (2000), obtained temperatures of ~830–880 °C, using recombined (proto-Cpx) clinopyroxene compositions and



FIGURE 8. Garnet line profiles (locations shown in Fig. 5) exhibiting the change in CaO and MgO (wt%). Note the gap in the Garnet 2 data, which corresponds to a Cpx inclusion.

garnet cores, and ~670-760 °C using clinopyroxene rim and garnet rim compositions. Proto-Cpx compositions were calculated from analyses of exsolution lamellae of orthopyroxene and host clinopyroxene using the methods outlined in "Single proto-Cpx thermometry" section. Garnet core, garnet rim, and clinopyroxene rim compositions are noted in the Online Materials¹ B. Two-pyroxene thermometry from non-recombined matrix clinopyroxene and exsolved orthopyroxene (Andersen et al. 1993) derived temperatures of ~740 °C, when considering Opx lamellae that had not been corrected for excess Ca contents due to secondary fluorescence. In contrast, when fluorescence-corrected orthopyroxene analyses were used, this calibration returned lower temperatures of ~590 °C ("High resolution EPMA (HR-EPMA)" section). Mag-Ilm thermometry of the exsolved oxides within the garnet coronas, yielded temperatures ~460 °C.

The effective bulk composition for phase equilibrium modeling, as well as the mean mineral compositions used for its calculation, are presented in Table 1. Figure 10a shows the isochemical *P*-*T* phase diagram for this specific bulk composition. "Petrographic and modal analysis" section outlines mineral associations that suggest a garnet-clinopyroxene (±plagioclase feldspar) peak mineral assemblage free from orthopyroxene and the oxide minerals currently present in the sample. Based on these associations, the closest match to the peak assemblage is the field Cpx-Grt-Pl-Rt-Qtz, which spans a range of pressures above 10 kbar and temperatures up to the dry solidus at



FIGURE 9. Na and Fe (total) cations per formula unit (cpfu) plotted vs. Ca cpfu. Note the slight decrease in Na content for the non-lamellae containing rims, as well as the general trend in the data toward the recalculated proto-Cpx composition (blue). Error bars are shown but are smaller than the symbol, with the exception of Na for the Cpx; proto-Cpx composition are the average of 12 analyses so have a smaller error than the symbol even for Na.

~1050–1100 °C. There is no direct evidence in the rock for rutile at peak conditions, despite a small modal amount (<1%) being predicted by the phase diagram for our bulk composition (see "Obtaining peak *P*-*T* conditions" section for discussion of this).

The three major constraints on our rock are the garnet, quartz, and orthopyroxene mode-zero lines, which denote the boundaries in P-T space where these phases disappear from the predicted assemblage (Fig. 10). The Opx mode-zero line (Opx-out) provides a minimum pressure for our assumed peak assemblage, the Qz mode-zero line (Qz-out) provides a minimum pressure for the end of corona growth, while the Grt mode-zero line (Grt-in) represents a minimum pressure for the final corona stage. Additionally, the corona mineral assemblage (Opx-PI-IIm) is a constraint itself, as it falls within the phase field Opx-Cpx-Grt-Pl-Ilm. To show the sensitivity of the Grt, Qz, and Opx mode-zero lines to bulk composition, the range of P-T conditions over which they lie for the 20 randomly generated bulk compositions is shown by the blue (Opx), gray (Qz), and red (Grt) shaded regions (Fig. 10a). We use this as the best estimate on the uncertainty of our phase equilibrium modeling *P*-*T* work.

Because textural evidence demonstrates that the core regions of coarse-grained clinopyroxene may have been isolated from the retrograde reaction that formed the coronas, we produced a P-X diagram in which the volume percent clinopyroxene inputted into the calculated bulk composition is varied between 35.6 and 0% (Fig. 10b). This allowed assessment of how the predicted proportion of corona minerals changes for a range of different inputted clinopyroxene volumes (Fig. 10b). Figure 10b shows that for the full range of bulk compositions there is little change in the pressure at which the corona forming reaction initiates (i.e., Opx-out line). It also highlights that considerable change in the volume percent of the corona minerals occurs between the orthopyroxene and quartz mode-zero lines, with almost no change once quartz has left the assemblage (Fig. 10b).

Finally, single pyroxene thermometry (Lindsley 1983) using the corrected and proto-Cpx composition ("Single proto-Cpx



FIGURE 10. (a) Isochemical *P*-*T* phase diagram for S98-14. The large, bolded assemblage (Cpx-Grt-Pl-Rt-Qz) represents the closest match to our interpreted peak assemblage. The red, gray, and blue shaded regions depict the uncertainty on the garnet, quartz, and orthopyroxene mode-zero lines (Grt-in, Qz-in, and Opx-out) determined from 20 randomly generated bulk compositions. (b) *P*-*X* phase diagram showing the effect of varying the contribution (vol%) of clinopyroxene to the calculated bulk composition. The scientific color map "batlow" (Crameri 2021; Crameri et al. 2020) is used to depict the predicted ratio among the volumes of corona minerals [Grt/(Grt+Pl+Opx+Ox)].

thermometry" section) obtained temperatures ranging from 980 to 1025 °C (Fig. 11; Table 2). Pyroxene thermometry assumes coexisting clino- and orthopyroxene. In the absence of a coexisting Ca-poor pyroxene, the clinopyroxene is not saturated in the orthopyroxene component. Therefore, the temperature obtained will be a minimum estimate for the inferred orthopyroxene-free peak assemblage, and we take the highest temperature obtained as the best estimate of peak.

DISCUSSION

Microstructural significance

Understanding the mineral relationships that are preserved in the breakdown microstructures is crucial if the aforementioned thermometers and barometers are to be applied correctly, and accurate P-T estimates are to be obtained for the different stages of metamorphism. The coronas are perhaps the most striking microstructural feature of the mafic granulites found at Scouriemore (Fig. 2) and have previously been observed by several authors (O'Hara 1961; Savage and Sills 1980; Barnicoat 1983; Johnson and White 2011), who attribute their formation to the breakdown of a high-temperature garnet (possibly reacting with clinopyroxene) to form plagioclase feldspar, orthopyroxene, and magnetite. This reaction is thought to occur due to retrograde decompression, either as part of a separate metamorphic event or during slow cooling from peak conditions. Thus, the key questions that arise are, how extensive was this reaction, what was the peak metamorphic assemblage, and how do we best use this information to re-interpret the P-T history of these rocks?

Johnson and White (2011) indicated that while orthopyroxene

and plagioclase are present within both the coronas and matrix, magnetite generally only exists in the coronas and thus advocated a peak assemblage of Cpx-Grt-Pl-Opx-Ilm. In contrast, Savage and Sills (1980) found a complete lack of plagioclase within the matrix and therefore proposed an initial assemblage of garnetclinopyroxene. They bolstered this hypothesis by confirming that garnet was absent in samples with the highest X_{Mg} ratio, and best preserved in those with the lowest X_{Mg} ratio (Savage and Sills 1980). These two models represent end-members of a spectrum with a high-pressure, high-variance assemblage (Grt-Cpx) on one side and a moderate-pressure, lower-variance assemblage (Grt-Cpx-Pl-Opx) on the other.

In our assessment of the peak metamorphic assemblage, we consider the textural setting, relative size, and compositional heterogeneities of the minerals. Figures 2 and 7 illustrate the striking bimodal grain size distribution between coronal plagioclase-orthopyroxene and the relatively larger clinopyroxene and garnet. This size discrepancy was noted to varying degrees in all samples the authors characterized (Online Materials¹A), with clinopyroxene generally greater than twice as large as orthopyroxene, when they exist in the same sample. This size discrepancy can also be seen in the images from previous studies (Feisel et al. 2018; Johnson et al. 2012; Johnson and White 2011). The comparable size and spatial relationship of plagioclase and orthopyroxene (Fig. 7), as well as their textural relationship with garnet, would suggest that they grew over a similar temperature and/or time window. Although plagioclase and orthopyroxene do also sometimes appear to coexist in the matrix, seemingly distant from any garnet, this probably represents a cut effect due to slicing the rock in such a fashion to only see the corona. Some of this "matrix" orthopyroxene is observed connecting with corona textures via a narrow band of orthopyroxene (Fig. 3). This shows that mineral growth from the garnet breakdown can occur outside of the immediate area of the corona texture. In other cases, they may result from all garnet within a local region degrading as the corona forming reaction runs to completion.

Our findings generally agree with those of O'Hara (1977), Barnicoat (1983), and Savage and Sills (1980) in that orthopyroxene, plagioclase, and magnetite-ilmenite only grew during the retrograde path. A caveat to this interpretation is that the peak garnet composition cannot be retrieved from the corona products (Opx-Pl-Ilm-Mag) alone. The Na content of the labradorite in the corona requires an extra source for Na, either from pre-existing Na-rich plagioclase, and/or from clinopyroxene. The decrease in Na content in the rims of matrix clinopyroxene (Fig. 9) suggests that clinopyroxene was, at least in part, involved in this reaction. However, the Na content and vol% of the corona plagioclase is too high to have been solely sourced from the clinopyroxene. As such peak plagioclase must be a reactant in the full coronaforming reaction. Because we do not know the composition of the peak plagioclase or the amount of clinopyroxene and/or peak plagioclase consumed, it is difficult to write a balanced reaction that considers all of the phases that petrography suggests. We instead write a simplified volume balance 1 for the minimum phases required for a garnet breakdown reaction.

$$10Grt + 3Cpx + 2Qz \rightarrow 10Opx + 9Pl$$
(1)

We note that this volume balance has excess SiO_2 in the products, and therefore requires quartz as a reactant to balance. The crossing of the quartz mode-zero line (Fig. 10) is therefore potentially important in the corona-forming reaction. We note that our thin section does include rare examples of quartz as inclusions, strengthening our argument of peak quartz, and that a quartz consuming retrograde reaction 1 is possible. We know from petrographic and mineral chemical evidence that the actual breakdown reaction likely included precursor plagioclase as well as the consumption of small amounts of clinopyroxene, as evidenced by the high-Na content of the corona plagioclase. We therefore suggest the following generalized reaction 2 based on mineral relationships, abundances, compositions, and our volume balance 1.

$$Grt + Cpx + peak - Pl + Qz \rightarrow Opx + corona - Pl + Ti - Mag$$
 (2)

The presence of clinopyroxene as a reactant in 1 and 2 might imply that it has lost its peak composition. However, Figure 6 shows that the lamellae do not reach the edge of most of the clinopyroxene grains. This carries two implications: that the lamella boundaries did not provide pathways for the interior to communicate with the grain-boundary network, and that the exsolution occurred within a system isolated from the rest of the rock. This minimizes the possibility that the interiors of clinopyroxene grains have changed their bulk composition since peak, or near-peak, conditions. Consequently, the contribution of clinopyroxene to the corona-forming reaction is probably from the consumption of clinopyroxene grains around the immediate perimeter of the corona, together with some compositional exchange via the grain-boundary network, with lattice diffusion over a limited distance affecting clinopyroxene rims (Fig. 9). We can place constraints on the amount of clinopyroxene that may have reacted by assuming that the titanium in the corona assemblage comes entirely from the breakdown of clinopyroxene. There is reason to believe that that the peak mineral assemblage did not contain a titanium-dominant phase, as would be suggested from phase equilibrium modeling (see "Obtaining peak P-T conditions" section), and that all Ti-oxide phases came from the breakdown of clinopyroxene (±garnet) in reaction 1. This would mean that a mass balance of ilmenite and clinopyroxene places an upper limit on the amount of clinopyroxene that might have reacted. Using the measured TiO₂ content of clinopyroxene and the current vol% ilmenite, suggests that 20-30% of the clinopyroxene was consumed in reaction 2.

In summary, using microstructural evidence, we interpret the peak assemblage to be clinopyroxene and garnet (±plagioclase and quartz) with no orthopyroxene as either lamellae or individual grains. Subsequently, coronas consisting of plagioclase and orthopyroxene formed around garnets, at the expense of

FIGURE 11. Ternary plot (adapted from Lindsley 1983) showing host Cpx, lamella Opx, and recombined "proto-Cpx" compositions for two grains from S98-14 and the subsequent calculated temperatures. Pyroxene end-member compositions were calculated using the QUILF program and temperatures calculated using the Lindsley (1983) calibrations.



garnet, parts of the adjacent clinopyroxene, and the peak plagioclase and quartz predicted by phase equilibrium modeling. This Fe-Ti oxide later exsolved to form the magnetite-ilmenite intergrowth we see today.

Obtaining peak P-T conditions

Determining the peak *P*-*T* conditions of high-temperature rocks has presented a challenge to metamorphic petrologists for some time (O'Hara 1977; Frost and Chacko 1989; Harley 1989; Fitzsimons and Harley 1994; Pattison and Bégin 1994; Chacko et al. 1996; Pattison et al. 2003). This is a result of exchange thermometers having an apparent closure temperature below that of peak granulite facies conditions, meaning that the mineral compositions have probably been affected by down-temperature diffusional resetting during the initial stages of cooling (O'Hara 1977; Frost and Chacko 1989; Pattison and Bégin 1994). Aside from the obvious implications for temperature estimation, this also has a feedback effect (Harley 1989), because calculated pressures depend on temperature. Frost and Chacko (1989) coined this problem of unreliable *P*-*T* estimation in high-grade rocks the "granulite uncertainty principle."

In our case, the rock records a continuum of retrograde metamorphic processes in which minerals behaved differently. Therefore, linking mineral compositions to the same event can be problematic and determining P-T conditions using conventional thermobarometry, without petrographic evidence for local equilibrium relationships, is inappropriate.

Some have suggested that phase equilibrium modeling may be a superior technique for determining the peak metamorphic conditions in granulite facies rocks (e.g., Palin et al. 2016b; Feisel et al. 2018). Nevertheless, when simply matching observed mineral assemblages to those predicted on phase diagrams this technique suffers from high-variance assemblage fields providing little constraint on temperature. Some authors compare observed modal abundances or mineral compositions to those predicted within these fields to tighten their *P-T* condition (Powell and Holland 2008). However, recent studies have shown that the current *a-X* relations fail to reproduce natural mineral compositions for some phases, which in turn affects modal abundance predictions and phase diagram topology (Forshaw et al. 2019; Santos et al. 2019; Starr et al. 2020).

Another issue in phase equilibrium modeling relates to the majority of activity-composition relations for silicates only accounting for major element chemistry. Minor and trace components, such as Ti, are not accommodated in all mineral solution models (e.g., Cpx in Green et al. 2016; Grt in White et al. 2014). This is problematic in our sample because titanium cannot be allocated to the two major peak minerals (clinopyroxene and garnet) for which EPMA analyses reveal a minor titanium component (Table 1; Online Materials¹ B). As a result, a stoichiometric Ti-bearing mineral (in our case rutile) is predicted to occur, even though the amount of titanium in the rock could be accommodated as trace components in the existing silicate minerals. We attempted to circumvent this issue when calculating our effective bulk composition by removing titanium contents of silicate minerals and only including titanium from ilmenite. In comparison, an XRF-derived bulk composition would include titanium that is "locked" up in silicates. Even so, we predict

rutile in low modal abundance (<1%) at our peak *P-T* conditions. We hypothesize that rutile either: (1) reacted to form ilmenite/ magnetite during decompression, or (2) did not exist at peak and this small amount of TiO_2 was incorporated as a trace element in the large quantities of clinopyroxene and/or garnet. We note that all of the Ti-bearing minerals occur within garnet coronas and that analyses of clinopyroxene combined with modal proportion estimates indicate that it contains four times as much titanium as that contained in oxide minerals. Additionally, in 1 we invoke the breakdown of a small amount of clinopyroxene (20–30 vol%; "Microstructural significance" section) in the corona forming reaction, which would be a source for titanium in the corona mineral assemblage. It is therefore possible that all titanium was contained in garnet and clinopyroxene at peak, and that rutile need not have been part of the predicted peak assemblage.

Both this and previous studies suggest that minerals in the Scouriemore metabasites equilibrated, partly or wholly, by reaction or interdiffusion, at different stages in the metamorphic cycle. Even those that may well have reached equilibrium at some stage, such as garnet and clinopyroxene at peak conditions, have behaved differently during the retrograde evolution. The application of phase equilibrium modeling is complicated by these disequilibrium textures (cf. White et al. 2008). However, when combined with detailed optical microscopy, these petrographic relationships can be used to determine likely peak and retrograde assemblages.

As a result, we have examined these issues, and argue that it is possible to determine valid constraints on pressure and temperature by considering individual features in terms of local equilibrium, diffusional closure, and reconstructing the peak compositions of minerals in local domains. Disequilibrium textures can be used to determine retrograde conditions, but when using phase equilibrium modeling to do so the uncertainty regarding composition and volume percent of the disequilibrium textures must be considered. Fortunately, for our sample changing the effective bulk composition has a negligible effect on the predicted phase fields (Fig. 10b) and means that the corona assemblage can be used to obtain retrograde pressures. The observed garnet and clinopyroxene chemical zonation can be used to obtain upper and lower estimates on the temperature diffusional closure for garnet breakdown. Temperatures obtained from exsolution features in pyroxene and oxides place further temperature constraints on retrograde conditions. The petrographic observations and complex retrograde zoning patters, show that recombined single pyroxene thermometry is the best predictor of peak temperature conditions.

This single pyroxene thermometry of a recombined proto-Cpx analysis is not limited to a specific assemblage, requiring only the presence of clinopyroxene, a mineral that is stable for most upper-amphibolite and granulite facies mafic rocks. It also does not require chemical equilibrium between minerals, just that the clinopyroxene as a whole retains its peak composition during exsolution (Fig. 6). However, single pyroxene thermometry is only applicable to high-Ca pyroxenes, due to the close spacing of temperature contours for low-Ca pyroxenes (Fig. 11). The need to correct for the effects of secondary fluorescence should also be considered prior to analysis and can be achieved using PENELOPE modeling (Pinard et al. 2010). In these samples, the need for correction was apparent due to the specific geometries ($<5 \mu m$ Opx lamella in Cpx) and the large influence that Ca has on pyroxene thermometry (Lindsley and Frost 1992; Fig. 5). It is also important to consider when this technique should be used over other existing thermobarometric methods. Calculating a recombined proto-Cpx analysis for single pyroxene thermometry is most useful: (1) when the peak composition is no longer retained in one of the minerals and (2) when an applicable two-mineral exchange thermometer is absent.

Implications for the Lewisian Complex

In the case of the mafic rocks from Scouriemore investigated here, the restricted mineral assemblage means that a limited number of exchange thermometers can be used to study the metamorphic history, but also that determining local equilibrium assemblages is simpler. A range of temperatures have been obtained from a single, representative, well preserved, sample (S98-14; Table 2), which highlights the need for careful petrography and analytical parameters if geologically meaningful temperatures are to be obtained. Here, we attempt to relate the temperatures obtained from different thermometers to the stages in the metamorphic evolution of the rock discussed above.

Above, we propose that single pyroxene thermometry is the best predictor of the peak temperature conditions of ultrahigh-temperature metamorphism, hence we take >1025 °C to represent peak conditions. When assessing pressures, we have inferred that orthopyroxene is observed only as part of the corona texture and thus the peak assemblage was likely garnet-clinopyroxene (±plagioclase-quartz). Phase equilibrium modeling constrains this assemblage and thus peak pressures to >11.5 kbar for the original bulk composition, and >11 kbar for the 20 random bulk compositions (lower extent of blue region on Fig. 10a). Therefore, we interpret peak Badcallian metamorphic conditions to lie at >1025 °C and >11 kbar.

A subsequent metamorphic event and/or cooling and decompression from peak produced the corona textures of the garnets (O'Hara 1961; Barnicoat 1983; Johnson and White 2011). Line scans through garnet highlight the change in composition from core to rim (Fig. 8), which results in garnet cores returning higher Grt-Cpx temperatures than rims, as was noted by previous authors (Barnicoat 1983; Sills and Rollinson 1987). The notable lack of a flat interior plateau in these profiles shows that the peak composition has been lost, probably due to late diffusional exchange related to corona formation. Therefore, the core of the garnet cannot be used to determine peak conditions and the ~850 °C garnet(core)/proto-Cpx temperatures is interpreted to record some intermediate time during the exchange process. The garnet(rim)/ clinopyroxene(rim) temperature then represents the closure of this system at ~700 °C. Phase equilibrium modeling provides a constraint on pressure with the observed assemblage (Opx-Cpx-Grt-Pl-Ilm) found at 7-9 kbar, consistent with the results of Johnson and White (2011) and Feisel et al. (2018). The large pressure and temperature difference (>2 kbar/~200 °C) between peak and corona formation are suggestive of a subsequent metamorphic event.

In clinopyroxene crystals, where orthopyroxene lamellae do not reach the proto-grain boundary, we are able to assume exsolution has occurred in a closed system. Thus, we interpret the two-pyroxene temperature of \sim 590 °C as recording the end of clinopyroxene exsolution. Magnetite-ilmenite thermometry generates even lower temperatures (~460 °C), but due to the lower closure temperature of oxides relative to silicates, this temperature is probably recording the late stages of the metamorphism, that caused clinopyroxene exsolution.

Figure 12 exhibits a *P*-*T* diagram that summarizes the *P*-*T* estimates given above and shows a proposed *P*-*T*-*t* path for the rocks of Scouriemore. For those temperatures where independent pressure estimates could not be obtained (Mag-Ilm and Cpx-Opx) we use literature estimates for the correlating pressures (Cartwright and Barnicoat 1989). We infer that peak pressures were much higher than have been suggested by recent phase equilibrium modeling results, in line with those suggested by Cartwright and Barnicoat (1989).

Although we have described a sequence of events for our sample S98-14 that is applicable to garnet-bearing Lewisian metabasites, our data do not constrain the absolute timing of metamorphic events. The time of peak Badcallian granulite-facies metamorphism is generally quoted as ~2.7 Ga (e.g., Cartwright and Barnicoat 1989), but the evidence is still equivocal after decades of study, mostly based on zircon geochronology. Whereas some recent studies (e.g., Crowley et al. 2015) provide evidence for two distinct periods of high-temperature metamorphism at ~2.7 and ~2.49 Ga, others such as Taylor et al. (2020) point to an apparent continuum of zircon ages as implying that supra-solidus metamorphism and melt generation lasted for more than 200 Ma. Nevertheless, if conditions at 2.49 Ga reached or remained at granulite facies, it is reasonable that the coronas formed at this time. The hydrous amphibolite-facies retrogression and cooling, commonly referred to as the Inverian, that immediately followed (Cartwright and Barnicoat 1989; and references therein) has not directly affected S98-14, but this interval marks the first opportunity for exsolution phenomena to appear in clinopyroxene and Fe-Ti oxides over the T interval 600-450 °C. However, the Laxfordian



FIGURE 12. P-T diagram summarizing the P-T estimates of this study and showing the proposed P-T path, compared to those of previous studies.

event at ~1.76 Ga, most clearly recorded in shear zones, evidently reached similar conditions at Scourie (Beach 1973), and therefore exsolution could be related to post-Laxfordian cooling. We cannot resolve this question, but we note that the orthopyroxene lamellae in clinopyroxene are very narrow, with planar interfaces that show no indication of annealing or later modification.

IMPLICATIONS

When appropriate exchange thermometers are absent, minerals have been partially or wholly overprinted, and temperatures obtained from phase equilibrium modeling are poorly constrained, we show that single proto-Cpx thermometry can provide a viable alternative for determining peak temperature estimates. Clinopyroxene commonly exsolves in a closed system that can be interrogated to determine: (1) peak conditions by phase recombination and (2) exsolution temperatures, if the appropriate analytical precautions are taken. Exsolution is a prevalent feature in granulitic rocks, so the technique presented has wide ranging applications to other terranes (e.g., Bohemian Massif, Grenville Orogen, etc.).

Our study combines several old and new thermobarometric techniques with careful petrography to determine the peak and retrograde mineral assemblages of the mafic granulites of Scouriemore. We obtain a much higher estimate of the Bacallian peak *P-T* conditions (>11 kbar and >1025 °C) compared to recent studies, which relied on phase equilibrium modeling alone, but one that is consistent with earlier work in the Lewisian (Barnicoat and O'Hara 1979; Sills and Rollinson 1987; Cartwright and Barnicoat 1989; Wheeler et al. 2010). In the process of generating our proto-Cpx temperature we also obtain a range of temperatures (850 °C Grt-Cpx; 590 °C Opx-Cpx; 460 °C Mag-Ilm) that represent estimates of the metamorphic overprints on the Lewisian. We note that a more nuanced approach is necessary in the application of phase equilibrium modeling to retrogressed mafic granulites, highlighting the need for a careful consideration of effective bulk composition.

Using these data, we have re-interpreted the *P*-*T* history of the Lewisian, with our study differing from previous work in two main regards: (1) we have an independent estimate of temperature from single proto-Cpx thermometry and (2) we use the petrologic evidence to argue for an orthopyroxene-free peak mineral assemblage in samples with corona textures. Our work sheds new light on rocks from the crucial transition from Archean to modern style subduction, and at the same time developed a new technique (single pyroxene thermometry) that can be widely used to study similarly overprinted high-grade terranes.

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