# Calculation and application of clinopyroxene-garnet-plagioclase-quartz geobarometers\*

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Abstract. Recently published thermodynamic and experimental data in a variety of chemical systems have been evaluated to derive Gibbs free energies for hedenbergite and pyrope. These were used to calculate the geobarometric equilibria

Hedenbergite + Anorthite = Grossular + Almandine + Quartz: "HD barometer",

Diopside + Anorthite = Grossular + Pyrope + Quartz: "DI barometer".

We have compared the pressures obtained from these equilibria for garnet-clinopyroxene-orthopyroxene-plagioclasequartz assemblages with the geobarometer

Ferrosilite + Anorthite = Almandine + Grossular + Quartz: "FS barometer".

Pressures calculated for 68 samples containing the above assemblage from a variety of high grade metamorphic terranes indicate that, in general, the HD and DI barometers yield values that are in good agreement with the FS barometer, and that the three barometers are generally consistent with constraints from aluminosilicate occurrences. However, in some samples the HD barometer yields pressures up to 2 kbar greater than constraints imposed by the presence of an aluminosilicate phase. Relative to the FS barometer, the HD barometer overestimates pressure by an average of  $0.2 \pm 1.0$  (1 $\sigma$ ) kbar and the DI barometer underestimates pressure by an average of  $0.6 \pm 1.6 (1 \sigma)$  kbar. The pressure discrepancies for the HD and DI barometers are likely to be a result of imprecision in thermodynamic data and activity models for silicates, and not a result of resetting of the clinopyroxene equilibria. The relative imprecision of the DI barometer relative to the FS barometer results from overestimates of pressure by the DI and FS barometers in Fe-rich and Mg-rich systems, respectively. Application of the HD and DI barometers to high grade Cpx-Gt-Pg-Qz assemblages yields pressures that are generally consistent with other petrologic constraints and geobarometers. It is concluded that the HD and DI barometers can place reasonable constraints on pressure ( $\pm 1$  kbar relative to the FS barometer) if not extrapolated to mineral assemblages whose compositions are extremely far removed from the end member system for which the barometers were calibrated.

## Introduction

Major advances in the accuracy and precision of geobarometers have been made in the last decade, in part a result of careful experimental reversal of pressure dependent equilibria (Bohlen et al. 1980; Bohlen and Boettcher 1982; Bohlen et al. 1983a, 1983b; Gasparik 1984a, 1984b; Gasparik and Newton 1984; Bohlen and Liotta 1986; Koziol and Newton 1988), more precise thermodynamic data (e.g. Haselton and Newton 1980; Metz et al. 1983; Bohlen et al. 1983; Robie and Hemingway 1984; Haselton et al. 1987; Robie et al. 1987), and more accurate modeling of activitycomposition relations for mineral phases involved in geobarometric equilibria (e.g., Newton et al. 1980; Newton and Haselton 1981; Ganguly and Saxena 1984; Davidson and Lindsley 1985; Anovitz and Essene 1987a). Reasonably accurate and precise geobarometers now exist for most granulite facies metabasites, charnockites and high grade metapelites, and for some upper amphibolite facies metabasites and metapelites. The above experiments also serve as important constraints on thermodynamic data for mineral phases involved in the particular reaction. In concert with precise thermodynamic data for other phases, the experimentally constrained thermodynamic data may be used to calculate geobarometers that are not easily reversed experimentally. Using this approach we have calculated the location of the pressure dependent reactions

 $Hedenbergite + Anorthite = Grossular + Almandine \\ + Quartz$ 

$$3 \text{ CaFeSi}_{2}O_{6} + 3 \text{ CaAl}_{2}\text{Si}_{2}O_{8} = 2 \text{ Ca}_{3}\text{Al}_{2}\text{Si}_{3}O_{12} + \text{Fe}_{3}\text{Al}_{2}\text{Si}_{3}O_{12} + 3 \text{ Si}O_{2}$$
(1)

(the "Hedenbergite (HD) barometer") and,

$$Diopside + Anorthite = Grossular + Pyrope + Quartz$$
  

$$3 CaMgSi_2O_6 + 3 CaAl_2Si_2O_8 = 2 Ca_3Al_2Si_3O_{12}$$
  

$$+ Mg_3Al_2Si_3O_{12} + 3 SiO_2$$
(2)

(the "Diopside (DI) barometer"), in order to expand the range of assemblages for which there exist reasonably pre-

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cise geobarometers. Because garnet, plagioclase and quartz are phases common to both barometers we have chosen to distinguish them on the basis of the particular pyroxene present. Reaction 2 was previously calculated by Newton and Perkins (1982) based on the best available thermochemical data. Their version of Reaction 2 typically underestimated pressure by an average of 2.2 kbar relative to their orthopyroxene geobarometer (Newton and Perkins 1982), which was ascribed to imprecise thermodynamic data.

This paper presents calculations of new calibrations for clinopyroxene-garnet-plagioclase-quartz geobarometers in the system  $CaO-Al_2O_3-FeO-MgO-SiO_2$ , based on thermodynamic data derived from a number of sources and constrained by a variety of experimental equilibria. The precision and accuracy of these calibrations are tested by comparing pressures obtained for Reactions 1 and 2 against the equilibrium

Ferrosilite + Anorthite = Almandine + Grossular + Quartz

$$6 \text{ FeSiO}_{3} + 3 \text{ CaAl}_{2}\text{Si}_{2}\text{O}_{8} = 2 \text{ Fe}_{3}\text{Al}_{2}\text{Si}_{3}\text{O}_{12} + \text{Ca}_{3}\text{Al}_{2}\text{Si}_{3}\text{O}_{12} + 3 \text{ SiO}_{2}$$
(3)

(the "Ferrosilite (FS) barometer"), and the location of the polymorphic transition

Sillimanite = Kyanite 
$$(4)$$

(Richardson et al. 1968; Holdaway 1971; Robie and Hemingway 1984). Reaction 3 is derived by addition of the experimentally reversed reactions

Fayalite + Quartz = 2 Ferrosilite $Fe_2SiO_4 + SiO_2 = FeSiO_3$ (5)

(Bohlen et al. 1980) and

Fayalite + Anorthite = Almandine + Grossular  $3 \operatorname{Fe}_2 \operatorname{SiO}_4 + 3 \operatorname{CaAl}_2 \operatorname{Si}_2 \operatorname{O}_8$ =  $(2 \operatorname{Fe}_3 \operatorname{Al}_2 \operatorname{Si}_3 \operatorname{O}_{12} + \operatorname{Ca}_3 \operatorname{Al}_2 \operatorname{Si}_3 \operatorname{O}_{12})_{SS}$ 

(where SS is a 2/3 almandine + 1/3 grossular solid solution produced in the experiments, Bohlen et al. 1983a), and has proven to be an extremely useful geobarometer for granulites. Calculations of Reaction 3 as a geobarometer have been presented by Bohlen et al. (1983a), Perkins and Chipera (1985), and Anovitz and Essene (1987a) that differ in the choice of thermodynamic data and garnet mixing model used to calculate the position of the end member reaction (Reaction 3). For these purposes we have used the garnet mixing model of Ganguly and Saxena (1984) with Ca – Fe mixing parameters derived by Anovitz and Essene (1987a). Sources of thermodynamic data are discussed in detail in the following sections.

#### Thermodynamic data

A variety of sources and experimental studies were used in the compilation of thermodynamic data for the present study. Carefully reversed experiments on Reactions 5, 6 and the reactions

Almandine + Rutile = Ilmenite + Sillimanite + Quartz  $Fe_3Al_2Si_3O_{12} + 3TiO_2 = 3FeTiO_3 + Al_2SiO_5 + 2SiO_2$  Almandine + Grossular + Rutile = Ilmenite + Anorthite + Quartz

$$2 \operatorname{Fe_{3}Al_{2}Si_{3}O_{12} + Ca_{3}Al_{2}Si_{3}O_{12} + 6 \operatorname{Ti}O_{2}}_{=6 \operatorname{FeTi}O_{3} + 3 \operatorname{CaAl_{2}Si_{2}O_{8}}_{+3} \operatorname{Si}O_{2}}$$
(8)

(Bohlen et al. 1983b, 1983c; Bohlen and Liotta 1986)

Anorthite = 
$$Grossular + Kyanite + Quartz$$

$$3 \text{ CaAl}_{2}\text{Si}_{2}\text{O}_{8} = \text{Ca}_{3}\text{Al}_{2}\text{Si}_{2}\text{O}_{12} + 2 \text{ Al}_{2}\text{Si}\text{O}_{5} + \text{Si}\text{O}_{2}$$
(9)

(Koziol and Newton 1988) and

$$Wollastonite + Anorthite = Grossular + Quartz$$
  
2 CaSiO<sub>3</sub> + CaAl<sub>2</sub>Si<sub>2</sub>O<sub>8</sub> = Ca<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub> + SiO<sub>2</sub> (10)

(Newton 1966; Hays 1967; Boettcher 1970; Huckenholz et al. 1975) combined with heat capacity functions and volume data serve to constrain the 1 bar Gibbs free energy  $(\Delta G_{298})$  of fayalite, ferrosilite, kyanite, grossular, almandine and anorthite. The latter data were compiled by Anovitz and Essene (1987a; Table 1) from a number of sources and comprise part of an internally consistent thermodynamic data set for selected phases in the system CaO–FeO  $-Al_2O_3-SiO_2(-TiO_2)$  (Table 1 and 2). Thermodynamic data for diopside (Tables 1 and 2) have been compiled and evaluated by Sharp et al. (1986) from various sources and experiments in the system CaO–MgO–SiO<sub>2</sub>–CO<sub>2</sub>. The reader is referred to these sources for a detailed discussion of the methods used in deriving the respective data. Thermodynamic data for enstatite are taken from Robinson et al. (1982), and derivation of thermodynamic data for pyrope and hedenbergite is described below.

There has been considerable discussion concerning the presence of Al-Si disorder in synthetic anorthite produced in phase equilibrium experiments such as Reaction 9, and the necessity of adding a configurational entropy term to  $S'_{298}$  of anorthite in order to fit the experimental reversals (e.g., Gasparik 1984a; Wood and Holloway 1984; Koziol and Newton 1988). Anovitz and Essene (1987a) also address this problem, concluding that use of different values for  $S_{298}^{\circ}$  of grossular, and lack of application of thermal expansion and compressibility in phase equilibrium calculations may require an  $S_0^{\circ}$  term of anorthite in order to fit the reversals. However, the thermodynamic data set derived by Anovitz and Essene (1987a: Tables 1 and 2 of this study) uses an alternative  $S_{298}^{\circ}$ for grossular and includes the effects of expansivity and compressibility of solids, providing an adequate fit to the reversals for Reaction 9 and other equilibria. Following their conclusion we do not believe a configurational entropy term is warranted for anorthite.

The loci of relevant equilibria were calculated with a computer program (EQUILI, Wall and Essene unpubl) that solves the relation

$$\Delta G_{T_{2}}^{P_{2}} - \Delta G_{T_{1}}^{P_{1}} = \int_{P_{1}}^{P_{2}} \Delta V dP - \int_{T_{1}}^{T_{2}} \Delta S dT$$

(6)

(7)

with an experimental reversal or a known Gibbs free energy of reaction as the starting point for the calculation. Using EQUILI, reactions are then calculated at 1 bar, 298 K in order to obtain the Gibbs free energy  $(\Delta G_{298}^{\circ})$  of each reaction. Algebraic combination of the reactions allows calculation of the  $\Delta G_{298}^{\circ}$  of each phase assuming that the  $\Delta G_{298}^{\circ}$  of quartz and other simple phases (e.g., hematite, magnetite, wollastonite) are known (Robie et al. 1979; Robinson et al. 1982). Equilibria for which there are no experimental reversals can then be calculated at high P and T using the 1 bar, 298 K data as a starting point.

Thermodynamic data for hedenbergite (Tables 1, 2) are constrained by the following experiments in the system Ca-Fe-Si-O:

Andradite = Wollastonite + Hematite

$$Ca_3Fe_2Si_3O_{12} = 3 CaSiO_3 + Fe_2O_3$$
(11)

(Huckenholz et al. 1974, Suwa et al. 1976),

Table 1. Molar volume, entropy,	entropy coefficients, and	Gibbs free energies (relative	e to elements) of phases	involved in thermodynamic
calculations				

Phase	V <sub>298</sub> cc/mol	Ref	S <sup>°</sup> 298 J/mol K	А	В	C	D	Ref	⊿G°298 kJ/mol	Ref
Ni	6.59	1	29.87	29.397	4.410	1.473	-187.3	1	0	_
NiO	10.97	1	37.99	41.660	11.707	5.954	-234.2	1	-211.6	17
α-Quartz	22.69	1	41.46	73.488	0.782	15.376	-436.1	11	-856.3	2
$\beta$ -Quartz	22.30	2	41.46	70.601	4.255	32.715	-421.1	11	-853.2	18
Corundum	25.57	1	50.92	116.336	11.983	19.514	-688.3	1	-1582.2	1
Hematite	30.28	2	87.49	84.429	92.839	1.987	-511.0	2	-745.3	2
Magnetite	44.52	2	160.33	376.660	-139.963	122.478	-2242.1	2	-1014.1	2
Kyanite	44.21	3	82.42	177.000	24.849	29.564	-1049.1	3	-2445.8	4
Sillimanite	50.02	3	95.77	169.439	28.903	25.623	-1002.8	3	-2440.7	4
Wollastonite	39.79	2	81.67	102.236	28.447	13.443	-605.9	2	-1549.2	2
Enstatite	31.35	2	66.32	118.248	11.159	20.280	-699.6	1	-1457.4	2
Ferrosilite	32.99	4	95.82	116.495	15.201	16.159	-686.3	12	-1116.2	18
Fayalite	46.15	5	152.13	156.620	35.066	15.619	-920.4	2	-1377.1	18
Diopside	66.11	6	142.72	230.948	22.916	34.840	-1361.3	13	-3025.0	19
Hedenbergite	67.85	7	173.59	214.116	48.359	24.698	-1262.0	14, 15	-2677.7	20
Anorthite	100.79	2	199.28	263.705	62.848	31.468	-1556.7	2	-4010.9	18
Grossular	125.30	2	255.98	494.298	7.895	83.893	-2912.6	4	-6282.3	18
Andradite	131.67	8	316.82	470.395	46.903	63.743	-2765.6	16	-5413.2	20
Almandine	115.11	9	342.63	429.345	90.458	53.133	-2532.8	9	-4940.8	18
Pyrope	113.27	10	266.27	452.332	44.807	65.538	-2664.2	10	- 5936.4	20

 $S_T^{\circ} = S_{298}^{\circ} + A Ln T + B 10^{-3} T + C 10^{5} T^{-2} + D$ 

1: Robie et al. 1979; 2: Robinson et al. 1982; 3: Robie and Hemingway 1984; 4: Anovitz and Essene 1987b; 5: Essene unpbl; 6: Levien and Prewitt 1981; 7: Cameron et al. 1973; 8: Huckenholz et al. 1974; 9: Metz et al. 1983; 10: Haselton and Newton 1980; 11: Hemingway 1987; 12: Bohlen et al. 1983; 13: Krupka et al. 1985a, 1985b; 14: Bennington et al. 1984; 15: Haselton et al. 1987; 16: Robie et al. 1987; 17: Holmes et al. 1986; 18: Anovitz and Essene 1987a; 19: Sharp et al. 1986; 20: this study

And radite = Wollastonite + Magnetite +  $O_2$ 

$$6 \operatorname{Ca}_2 \operatorname{Fe}_2 \operatorname{Si}_3 \operatorname{O}_{12} = 18 \operatorname{Ca}_3 \operatorname{SiO}_3 + 4 \operatorname{Fe}_3 \operatorname{O}_4 + \operatorname{O}_2$$
(12)

(Gustafson 1974),

And radite + Magnetite + Quartz = Hedenbergite +  $O_2$ 

$$3 \text{ Ca}_{3}\text{Fe}_{2}\text{Si}_{3}\text{O}_{12} + \text{Fe}_{3}\text{O}_{4} + 9 \text{ SiO}_{2} = 9 \text{ CaFeSi}_{2}\text{O}_{6} + 2 \text{ O}_{2}$$
(13)

(Burton et al. 1982), and,

And radite + Quartz = Hedenbergite + Wollastonite +  $O_2$ 

$$2 \operatorname{Ca_3Fe_2Si_3O_{12}} + 4 \operatorname{SiO_2} = 4 \operatorname{CaFeSi_2O_6} + 2 \operatorname{CaSiO_3} + O_2$$
 (14)  
(Liou 1974).

The calculated position of these reactions in  $fO_2 - T$  space at 2 kbar (relative to MH) are shown in Fig. 1, along with the calculated position of the NNO and FMQ buffers. The MH and FMQ buffers are calculated from data in Robinson et al. (1982, Table 1), and the NNO buffer is calculated from the solid state electrochemical measurements of Holmes et al. (1986, Table 1). All buffers were corrected for pressure and temperature using the expansivity and compressibility data in Table 2.

Thermodynamic data for hematite, magnetite, and wollastonite are taken from Robinson et al. (1982), and data for andradite and hedenbergite are compiled from sources listed in Tables 1 and 2. The  $\Delta G_{298}^{\circ}$  for andradite was calculated from Reaction 11 using the experimental reversal of Huckenholz et al. (1974) (1 bar,  $1137 \pm 5^{\circ}$  C) as a starting point, yielding  $\Delta G_{298}^{\circ}$  (And) = -5413.2 kJ/mol. If the reversal of Suwa et al. (1976: 1 bar, 1165° C) is used as a starting point, the free energy of andradite changes by -0.3 kJ. Use of the latter value for  $\Delta G_{298}^{\circ}$  (And) shifts Reaction 12 to higher temperature, increasing the discrepancy between the experimental reversal for Reaction 12 and its calculated position (Fig. 1). Therefore we have used the reversal of Huckenholz et al. (1975) as the reference reaction for  $\Delta G_{298}^{\circ}$  (And).

The experimental reversal at 800° C of Burton et al. (1982) for Reaction 13 was selected as the starting point for the calculation

of  $\Delta G_{298}^{\circ}$  (Hd), and the calculated position of Reaction 13 is in good agreement with the reversals at 600 and 700° C (Fig. 1). The position of Reaction 14 was calculated using the  $\Delta G_{298}^{\circ}$  (Hd) (-2677.7 kJ/mol) determined from these latter experimental constraints. Assuming that the reversals of Burton et al. (1982) are reliable, there is a discrepancy of 1.5 log units between the calculated position of Reaction 14 and the experimental reversals of Liou (1974) (Fig. 1) that is not ascribable to errors in the thermodynamic data. The direction of the shift suggests that the synthetic andradite used in the experiments of Liou (1974) may be non-stoichiometric with possible substitutions of the nature Fe<sup>+2</sup>Fe<sup>+3</sup>-CaFe<sup>+3</sup> or  $(OH)_4^{-4} - (SiO_4^{-4})$ . The reactants used in the experiments of Burton et al. (1982) were stoichiometric, but analyses of run products for Reaction 13 are not given. More thorough characterization of the run products for all experiments, or new experiments in the Ca--Fe-Si-O system, are required in order to resolve the discrepancy. Further calculation of reactions involving hedenbergite rest on the assumption of the choice of experimental constraints used here.

Using a slightly different experimental and thermodynamic data base, Robie et al. (1987) obtained values for  $\Delta G_{298}^{\circ}$  (And) =  $-5414.8 \pm 5.5$  and  $\Delta G_{298}^{\circ}$  (Hd) of  $-2674.3 \pm 5.8$  kJ/mole. The values differ from ours because we have included the effects of thermal expansion and compressibility in our calculation, and have started with a different thermodynamic data base. Helgeson et al. (1978) report values of  $\Delta G_{298}^{\circ}$  (And) = -5428.7 kJ/mol and  $\Delta G_{298}^{\circ}$  (Hd) = -2674.5 kJ/mol.

Volume and entropy data for pyrope are taken from Haselton and Westrum (1980) and Haselton and Newton (1980). The Gibbs energy of pyrope was calculated from experimental data on the reaction

$$3 MgSiO_3 + Al_2O_3 = Mg_3Al_2Si_3O_{12}$$
(15)

(Gasparik and Newton 1984), using thermodynamic data for enstatite from Robinson et al. (1982) and data for corundum from Robie

Table 2. Compressibility and thermal expansion data used in thermodynamic calculations

Phase	а	b	Ref	с	d	e	f	Ref
Ni	0.535	0.900	21	1.092 E - 01	4.207 E - 03	1.663 E - 06	- 5.856 E-10	30
NiO	0.600	1.000	22	-3.570  E-02	1.347 E-03	-2.295 E-08		31
α-Quartz	4.300	24.000	21	1.944 E-03	3.274 E-03	2.876 E-06	6.922 E-10	2
$\beta$ -Quartz	1.776	0	21	3.861 E-01	1.983 E-02	2.562 E-05	$1.044 \mathrm{E}{-}08$	2
Corundum	0.380	0	21	-7.526  E-02	2.034 E-03	0.876 E-06	-3.150  E-10	30
Hematite	0.509	0.454	2	9.682 E-02	3.870 E-03	1.056 E-09	-1.402  E-12	2
Magnetite	0.560	0	2	-6.156 E-02	2.320 E-03	3.481 E-06	-1.486 E-09	2
Kyanite	0.790	5.000	23	5.427 E-02	2.144 E-03	8.101 E-10	-3.296 E-10	32
Sillimanite	0.788	4.810	23	1.964 E-02	7.618 E-04	1.476 E-06	-7.665 E-10	32
Wollastonite	1.465	9.819	2	1.118 E-01	5.009 E-03	-4.166 E-06	1.554 E-09	2
Enstatite	1.010	0	2	4.569 E-03	-3.2922 E-04	5.928 E-06	-2.534 E-09	2
Ferrosilite	0.990	0	24	9.590 E-02	3.822 E-03	5.737 E-11	2.778 E-13	4
Fayalite	0.767	0.978	2	-2.810  E-01	1.017 E-03	3.881 E-06	-1.672 E-09	2
Diopside	1.108	11.020	6, 21, 25	9.535 E-02	4.005 E-03	$-1.420 \mathrm{E}{-}06$	7.625 E-10	7
Hedenbergite	2.117	31.313	26	-5.224  E-02	4.148 E-03	-4.245 E-06	2.999 E-09	7
Anorthite	1.087	0	27	1.566 E-01	1.311 E-03	-8.876 E-08	2.075 E-10	2
Grossular	0.721	3.200	2	4.468 E-02	1.761 E-03	9.401 E-07	2.674 E-10	2
Andradite	0.725	0	28	-4.731 E-02	2.107 E-03	5.732 E-07	7.252 E-01	30
Almandine	0.571	0	4	4.541 E-02	1.654 E-03	1.231 E-06	-3.023 E-10	4
Pyrope	0.137	2.650	29	4.876 E-02	2.084 E-03	8.206 E-07	$-2.678 \mathrm{E}{-10}$	30

$$\begin{split} V^{P}_{298} &= V^{\circ}_{298} \, (1 - aP \, 10^3 + bP^2 \, 10^6), \, P \, \text{kbar} \\ V^{-}_{T} &= V^{\circ}_{298} + V^{\circ}_{298} / 100 \, [c + dT + eT^2 + fT^3], \, T^{\circ} \, \text{C}, \end{split}$$

2, 4, 6, 7: Table 1; 21: Birch 1966; 22: Hazen and Prewitt 1977; 23: Brace et al. 1969; 24: Bass and Weidner 1984; 25: Hazen and Finger 1981; 26: Vaidya et al. 1973; 27: Liebermann and Ringwood 1976; 28: Babuska et al. 1978; 29: Hazen and Finger 1978; 30: Skinner 1966; 31: Nielson and Leipold 1965; 32: Winter and Ghose 1979

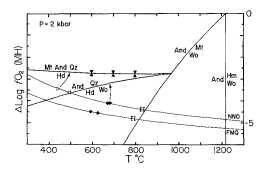


Fig. 1.  $fO_2 - T$  equilibria at 2 kbar in the system Fe-Si-Ca-O (relative to MH buffer) that constrain  $\Delta G_{298}^{\circ}$  of hedenbergite and andradite. Filled triangles: reversals of Burton et al. (1982) for Mt + And + Qz = Hd + O<sub>2</sub>; open triangles reversals of Gustafson (1974) for Mt + And +  $Qz = Hd + O_2$  at  $fO_2$  defined by NNO buffer; vertical bars: reversals of Gustafson (1974) for And =  $Wo + Mt + O_2$  at NNO and FMQ buffers; diamonds: reversals of Liou (1974) for  $Hd + Wo = And + Qz + O_2$  at NNO and FMQ buffers. Note disparity between calculated and experimental position of the latter reactions, denoted by arrows between reversals and reactions to which they correspond

et al. (1979). In the system MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> (MAS), enstatite contains significant solid solution of Mg-Tschermak's component (Mg<sub>0.5</sub>AlSi<sub>0.5</sub>O<sub>3</sub>) (Boyd and England 1964; Hensen and Essene 1971; Anastasiou and Seifert 1972; MacGregor 1974; Danckwerth and Newton 1978; Lane and Ganguly 1980; Perkins and Newton 1980; Perkins et al. 1981; Perkins 1983). In order to calculate the location of the end member Reaction (15) one must correct for the reduction in enstatite activity. Aranovich and Kosyakova (1987) have recently published activity-composition relations for orthopyroxene that are based on experimental equilibria in the MAS and FMAS systems. For the range of aluminum contents encountered

in the above experimental studies the model of Aranovich and Kosyakova (1987) yields activity coefficients for the MgSiO<sub>3</sub> component in enstatite-Mg-Tschermak's solid solutions ( $\gamma_{MeSiO_3}^{Opx}$ ) that are slightly less than one, suggesting only slight departures from ideality. However, the difference between the ideal model and the model of Aranovich and Kosyakova (1987) may yield significant differences in the Gibbs energy of pyrope because reactions involving pyrope and enstatite are extremely sensitive to changes in pressure. We have employed the latter model in the calculations to follow.

Using the midpoint of the reversal for Reaction 15 (Gasparik and Newton 1984, 850° C and  $16.25 \pm 0.25$  kbar,  $X_{MaTs}^{En} = 0.06$ ) as a starting point for the calculation, the shift in position of Reaction 15 due to orthopyroxene solid solution was calculated from the relation

$$RT\ln(K_2/K_1) = \int_{P_1}^{P_2} \Delta V dP$$

where  $K_2 = 1$  for the position of the corrected reversal,

$$K_1 = (1/a_{MgSiO_3}^{Opx})^3$$

and pyrope, sillimanite and quartz are assumed to be pure phases. For these calculations we have followed the convention of Hensen and Essene (1971) and Aranovich and Kosyakova (1987) by writing the formula of enstatite and Mg-Tschermak based on 2 cations (see Appendices I and II). The location of Reaction 15, adjusted to pure enstatite, is at 12.7 kbar, 850° C, and serves as the starting point for calculation of the Gibbs energy of pyrope. The calculated value of  $\Delta G_{298}^{\circ}$  (Reaction 15) is 18.1 kJ/mol, yielding  $\Delta G_{298}^{\circ}$  (Py) = - 5936.4 kJ/mol. As in deriving the Gibbs energy of hedenbergite, the Gibbs energy for pyrope is dependent on our choice of experiments and Gibbs free energies for enstatite and corundum.

Wood and Holloway (1984) present an analysis of equilibria in the CMAS system, deriving an  $\Delta H_{1000}^{\circ}$  for pyrope (relative to oxides) of -84.9 kJ/mol (S<sub>1000K</sub> = 777.8 J/mol, Haselton and Newton 1980). Using enthalpy and entropy data for the oxides (Robie et al. 1979), this yields  $\Delta G_{1000}^{\circ}$  (Py) for the study of Wood and

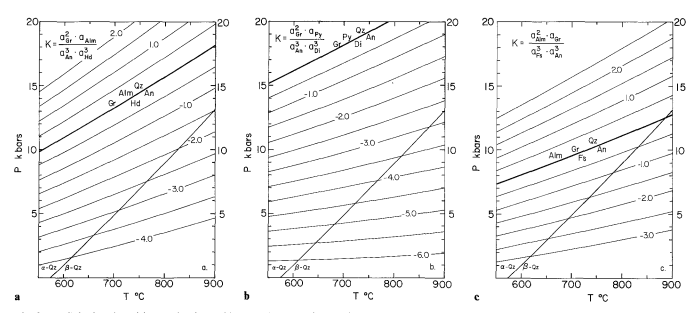


Fig. 2a–c. Calculated position and values of  $\log_{10} K$  for Reactions 1 through 3; a HD barometer; b DI barometer; c FS barometer

Holloway of -87.8 kJ/mol, in fair agreement with the value derived in this study ( $\Delta G_{1000}^{\circ} = -83.4$  kJ/mol,  $\Delta H_{1000}^{\circ} = -79.3$  kJ/mol).

Newton (1987) has re-evaluated experiments in the system  $MgO-Al_2O_3-SiO_2$  that constrain  $\Delta H_{298}^{\circ}$  and  $\Delta G_{298}^{\circ}$  of pyrope and enstatite. Based on analysis of the available experimental and thermochemical data base Newton (1987) obtains values for  $\Delta H_{298}^{\circ}$  (Py) and  $\Delta H_{298}^{\circ}$  (En) (relative to oxides) of -74.2 and -32.7 kJ/mol, respectively. Combined with entropy data for these phases the calculated values of  $\Delta G_{298}^{\circ}$  are in very good agreement with those of Newton (1987): -77.7 and -31.9 for pyrope and enstatite, respectively. The same experimental data base was used in both studies, but different values of  $S_{298}^{\circ}$  for enstatite were used and the mechanics of calculating the location of the equilibria using EQUILI differ slightly from the method used by Newton (1987).

Other thermodynamic compilations yield Gibbs free energies for enstatite that are in good agreement with the values used in this study. Berman et al. (1986) give a value for  $\Delta G_{298}^{\circ}(En) =$ -1458.5 kJ/mol, and Helgeson et al. (1978) report a value of -1459.9 kJ/mol.

The free energies of pyrope and enstatite have also been evaluated by fitting the experimentally reversed equilibrium

Enstatite + Sillimanite = Pyrope + Quartz

$$3 MgSiO_3 + Al_2SiO_5 = Mg_3Al_2Si_3O_{12} + SiO_2$$
(16)

(Hensen and Essene 1971; Perkins 1983), with various estimates of the high temperature heat capacity of pyrope. There are significant disparities between the experiments of Hensen and Essene (1971) and Perkins (1983) on Reaction 16 at 1150 and 1300° C, and among the possible high temperature extrapolations for the entropy of pyrope in the temperature range of the experiments (Haselton and Newton 1980; Robinson and Haas 1983; Berman and Brown 1985). However, both experimental data sets (adjusted for the Al content of the orthopyroxene using the methods discussed above) and the various heat capacity expressions converge at 1000° C (Moecher 1988). The thermodynamic data used in the present study for pyrope (with the heat capacity of pyrope from Haselton and Newton 1980) and enstatite are in very good agreement with both sets of experiments at 1000° C. More complete characterization of synthetic reactants and experimental products for Reaction 16, or high temperature (>1200 K) calorimetry on pyrope are needed in order to resolve the discrepancy at temperatures above the 1000° C reversals.

## Geobarometry

#### Clinopyroxene barometers

The positions of Reactions 1, 2 and 3 were calculated using 1 bar, 298 K as a starting point (Figs. 2a–2c). The clinopyroxene reactions have average slopes ( $\Delta P/\Delta T$ ) of 23 (HD) and 20 bar/° C (DI). These slopes are not as low as Reaction 3 (14.5 bar/° C), but are still useful for geobarometry, with the HD barometer having the greatest temperature dependency of the three barometers. The calculated slopes become less steep at lower pressure and more negative values of  $\log_{10} K$  (Fig. 2), particularly for the FS and DI barometers. An uncertainty of 5 kJ in the value of  $\Delta G_{298}^{\circ}(rxn)$  results in an uncertainty of approximately 0.7 kbar for both Reactions 1 and 2. Uncertainties of 5 kJ in the  $\Delta G_{298}^{\circ}$  for any phase involved in the reaction will yield an uncertainty of 0.7 kbar per mole of that phase in Reactions 1 and 2.

# Activity models

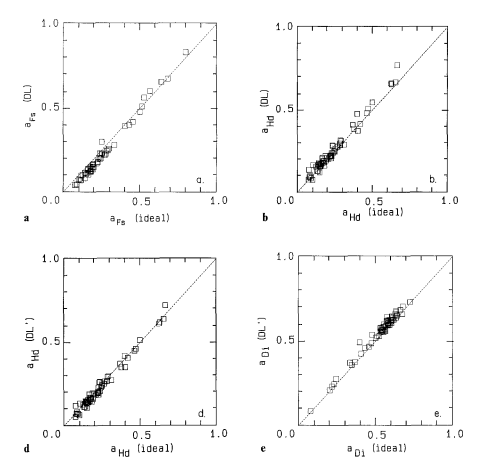
Application of these equilibria to natural systems requires consideration of the reduction in activity of end member components due to solid solutions in plagioclase, garnet, orthopyroxene, and clinopyroxene. Included in Figs. 2a, 2b, and 2c are the calculated positions for values of  $\log_{10}$  of the equilibrium constant K for each reaction. For example K is defined for Reaction 1 by the relation:

$$K = (a_{\text{Ca}_{A}\text{A}_{2}\text{Si}_{3}\text{O}_{12}}^{\text{Gi}})^{2} a_{\text{Fe}_{3}\text{A}_{2}\text{Si}_{3}\text{O}_{12}} / (a_{\text{Ca}_{A}\text{L}_{2}\text{Si}_{2}\text{O}_{8}}^{\text{Pg}})^{3} (a_{\text{Ca}_{F}\text{e}\text{Si}_{2}\text{O}_{6}}^{\text{Cpx}})^{3}.$$

Variations in  $\log_{10} K$  with pressure were calculated from the relation:

$$RT\ln(K_2/K_1) = \Delta G_{T_2}^{P_2} - \Delta G_{T_2}^{P_1} = \int_{P_1}^{P_2} \Delta V dP$$

Given appropriate activity models for plagioclase, garnet, and pyroxene, the equilibrium constant can be calculated from chemical analyses of the coexisting phases, and with



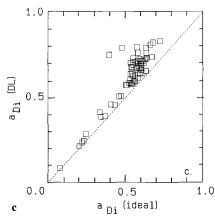


Fig. 3. a  $a_{Car}^{px} = s_{12O_6}^{cp}$  for an ideal two-site approximation versus activities calculated from model of Davidson and Lindsley (1985) (DL); b  $a_{CareSi_2O_6}^{cpx}$  with ideal model vs.  $a_{CareSi_2O_6}^{px}$  with DL model; c  $a_{Car}^{cpx} = s_{12O_6}^{cpx}$  with ideal model vs.  $a_{CareSi_2O_6}^{cpx}$  with DL model; d  $a_{CareSi_2O_6}^{cpx}$  with ideal model vs.  $a_{CareSi_2O_6}^{cpx}$  with adjusted DL activities (DL); e  $a_{CareSi_2O_6}^{cpx}$  with ideal model vs.  $a_{CareSi_2O_6}^{cpx}$ with adjusted DL activities (DL);

an estimate of temperature the pressure can be read from Fig. 3.

The activity models for plagioclase, garnet, or pyroxene will ultimately limit the accuracy and precision of a geobarometer. The models of Newton et al. (1980) or Orville (1972) for plagioclase, Perkins (1979), Newton and Haselton (1981), or Ganguly and Saxena (1984) for garnet, and an ideal two site approximation (e.g., Wood and Banno, 1973) for pyroxenes are typically selected by most workers. For this study the plagioclase activity model of Newton et al. (1980) (Appendix II) was used in evaluation of the geobarometers. At 700° C and constant garnet and pyroxene activities, this latter plagioclase activity model yields slightly lower pressures (on the order of 0.1 kbar) than that of Orville (1972).

The quaternary garnet mixing model of Ganguly and Saxena (1984) with modified values for Ca – Fe mixing parameters (Anovitz and Essene, 1987a) was used to calculate the activity of Ca<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub>, Fe<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub>, and Mg<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub> in garnet, and to calculate the position of the FS reaction (Reaction 3). The derivation of these parameters are discussed by Anovitz and Essene (1987a), and analytical expressions for calculating activities are outlined in Appendix II. The model used here typically yields Ca<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub> and Fe<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub> activities 1 to 5% greater than values obtained from the original formulation of Ganguly and Saxena (1984).

Activity coefficients for Mg<sub>3</sub>Al<sub>2</sub>Si<sub>3</sub>O<sub>12</sub> in garnet ( $\gamma_{Mg}^{Gt}$ ) obtained from the model of Ganguly and Saxena (1984) (GS) are significantly greater than those obtained from the model of Haselton and Newton (1980: HN). For a garnet of composition Alm<sub>50</sub>Py<sub>25</sub>Gr<sub>25</sub> at 700° C,  $\gamma_{Mg}^{Gt}$ (GS)=1.7

and  $\gamma_{Mg}^{Gt}(HN) = 1.3$ . This difference is due to the sign for the ternary constants as discussed by Ganguly and Saxena (1984, p 94, eq 16). If the sign of the ternary constant in the expression for  $\gamma_{Mg}^{Gt}$  is changed from that given in Ganguly and Saxena (1984), one obtains similar values for  $\gamma_{Mg}^{Gt}$  for the two models. This discrepancy needs further evaluation. In general, use of the GS model as it now stands yields significantly lower pressures than the HN model for the DI barometer, all other factors being equal.

Chatillon-Colinet et al. (1983) proposed that an ideal mixing approximation for Mg-Fe orthopyroxenes is consistent with solution calorimetric data on orthopyroxene solid solutions. Therefore the activity of FeSiO<sub>3</sub> in orthopyroxene was calculated using the ideal model of Wood and Banno (1973) (Appendix II). Davidson and Lindsley (1985) have modeled the phase equilibria of quadrilateral pyroxenes in order to derive a pyroxene activity model. Although Davidson and Lindsley do not present an explicit analytical formulation for activity coefficients similar to garnet and plagioclase, we obtained a computer program (PM Davidson pers comm 1987) that calculates activities of  $Fe_2Si_2O_6$ , Mg<sub>2</sub>Si<sub>2</sub>O<sub>6</sub>, CaFeSi<sub>2</sub>O<sub>6</sub>, and CaMgSi<sub>2</sub>O<sub>6</sub> for quadrilateral pyroxene. The pyroxene projection scheme of Lindsley (1983) was used to calculate mole fractions of Wo. En and Fs, the components upon which the activity model of Davidson and Lindsley (1985) is calculated. The pyroxene activity model of Davidson and Lindsley (1985) yields activities of  $a_{Fe_2Si_2O_6}^{Opx}$  that are similar to or only slightly less than ideal two site activities calculated with the Wood and Banno (1973) model (Fig. 3a).

Activities of CaFeSi2O6 and CaMgSi2O6 were also cal-

Terrane	Samples	Ref	ALS <sup>a</sup>	T° C/P kbar <sup>b</sup>
Grenville Province				
Parry Sound Ont Otter Lake Que	S3B 015B S10B M65 S32 S86E33 A12 DL2	1 2	Sil Sil	800/<9.8 700/<7.8
Adirondack Mts Highlands	BM2 BM13 ET15 ET24 LL6 MM4 MM18 PH4 SL5 SL26 SR31	3	Sil	800/<9.8
Lowlands	W 9 74 C 248 B	4	Sil	700/ < 7.8
Pikwitonei Belt	A41 C104 D50 D110 E18	5	Sil	800/<9.8
India Sargur Belt Karnataka	S33 S3 S4 113E GN4A	6 7 8	Ky/Sil	700/<7.8
Nilgiri Hills Mysore Bengal	DT MB GV2 332 352 680 K 21 SM4B SM44B	9 10 11 12	Sil Sil	750/<8.8 700/<7.8
Furua Complex	MF283.3 MF268.1 WE322.3 ZC.8 DMa40 C247.1 C311.1 C352.2	12	Ky/Sil	800/9.8
Westchester PA	74-67, -71, -207 A, -322 C	14	Ку	650/>6.8
Wind R. Range WY	A22	15	Sil	700/<7.8
SW Minnesota	DM4C2	16	Sil	700/<7.8
Lesotho	LT2	17		·
Oaxacan Complex	16076 25277 16976	18	Sil	730/<8.3
West Greenland				
Buksefjorden	174087 174102	19	Sil Ky/Sil	800/<9.8 600/6.3
Isortoq	88589 91141	20	Ky/5fi	000/0.5
Finnish Lapland	47111 6611	21	Sil	750/<8.8
Doubtful Sound	36461 36468	22		
Broken Hill	911 10770	23	Sil	800/<9.8

Table 3. Sources of analytical data on Gt-Cpx-Opx-Pg-Qz assemblages and quoted aluminosilicate occurrences used to evaluate geobarometers of this study

<sup>a</sup> ALS = aluminosilicate: Ky = kyanite, Sil = Sillimanite.

<sup>b</sup> Quoted T with P constraint.

1: Moecher (1988); 2: Perkins (1979); 3: Bohlen (1979); 4: Stoddard (1976); 5: Paktunc and Baer (1986); 6: Janardhan and Gopalkrishna (1983); 7: Srikantappa et al. (1985); 8: Janardhan et al. (1982); 9: Hansen et al. (1984); 10: Harris et al. (1982); 11: Devaraju and Coolen (1983); 12: Bhattacharya and Mukherjee (1987); 13: Coolen (1980); 14: Wagner and Srogi (1987); 15: Sharp (1988); 16: Moecher (1984); 17: Griffen et al. (1979); 18: Mora and Valley (1985); 19: Wells (1977); 20: Glassley and Sorensen (1980); 21: Hörmann et al. (1980); 22: Oliver (1977); 23: Phillips (1978)

culated using a modified ideal model, in comparison with the model of Davidson and Lindsley (1985). Ideal CaFeSi<sub>2</sub>O<sub>6</sub> activities were approximated by the relationship

$$a_{\text{CaFeSi}_{2}\text{O}_{6}}^{\text{Cpx}} = [X_{\text{Ca}}^{\text{M2}}] [X_{\text{Fe}^{2+}}^{\text{M1}}]$$
(17)

where  $X_{Fe^{2+}}^{M1} = Fe^2 - (1 - Ca - Na - Mn)$ ,  $X_{Ca}^{M2} = Ca$ , and Ca, Na, Mn, and Fe<sup>2+</sup> are the number of atoms of the respective cations. Pyroxene analyses taken from the literature are renormalized to four cations, and values of ferric and ferrous iron were calculated from charge balance and stoichiometry. Analyses taken from the literature were of varying quality, and not all of these pyroxenes were analyzed for Na, which usually occurs in significant quantities in high grade clinopyroxenes. Failure to analyze for Na will affect the calculation of Fe<sup>2+</sup> and Fe<sup>3+</sup>, and ultimately the value of  $X_{Fe^{2+}}^{M2}$ . Ideal CaMgSi<sub>2</sub>O<sub>6</sub> activities were calculated as

$$a_{\text{CaMgSi}_{2}O_{6}}^{\text{Cpx}} = [X_{\text{Ca}}^{\text{M2}}] [X_{\text{Mg}}^{\text{M1}}].$$
(18)

where  $X_{Ca}^{M2} = Ca$  and  $X_{Mg}^{M1} = Mg$ , based on a 4-cation pyroxene formula. The clinopyroxene model is based on crystal chemical observations on igneous pyroxenes that iron and magnesium do not mix ideally on the M1 and M2 sites, with iron showing a greater tendency relative to magnesium to partition into the M2 site (Cameron and Papike 1980; Dal Negro et al. 1982). Although there are no cation partitioning data on granulite facies augites, one might predict that the ordering would be even more pronounced in granulite facies augites compared to igneous augites, as tempera-

Locality	REF	BWB-FS	PC-FS	MAE-FS	NP-EN	PC-EN	MAE-HD	MAE-DI
Parry Sound	1	nd	nd	7.5–11.7	nd	nd	8.1–12.8	8.2–12.0
Otter Lake	2	8.1, <8.4	8.0, <7.5	8.4, <8.5	6.6, <8.1	9.7, <9.1	8.5, <7.9	7.8, <7.2
Adir. High.	3	7.1-8.2	7.1-8.6	6.5-8.8	8.5–9.9	10.4-11.5	5.8-9.5	7.4–13.1
Adir. Low.	4	6.6	nd	6.4	6.1	nd	5.6	6.1
Pikwitonei <sup>a</sup>	5	6.6–9.2	7.9–11.7	6.6-9.1	7.1–9.5	11.1-12.8	5.3–9.4	4.7-8.3
Sargur <sup>a</sup>	6 7	nd nd	nd nd	9.0 10.5, 10.5	9.0 9.1, 9.1	nd nd	10.1 9.7, 10.1	8.3 7.0, 7.1
Karnataka <sup>a</sup>	8 9	9.0, 9.2 nd	10.0, 9.5 nd	9.5, 10.4 8.4–9.1	8.3, 9.5 9.1, 9.1	9.4, 9.8 nd	8.7, 10.7 8.0–8.7	8.1, 8.8 7.0–7.7
Nilgiriª	10	nd	nd	9.0-10.2	6.9-8.8	nd	9.1–10.0	8.5-9.5
Mysore <sup>a</sup>	11	nd	nd	12.1	9.9	nd	11.8	9.5
Bengal <sup>a</sup>	12	nd	nd	7.5, 7.2	6.2, 6.3	nd	8.9, 7.9	7.9, 7.6
Furua Complex	13	9.6–11.4	9.9–12.1	10.1–12.6	9.4–12.7	9.8–11.6	9.9-12.8	7.4–10.2
Westchester <sup>a</sup>	14	8.9–9.6	9.8-10.8	9.6-10.0	5.6-9.4	9.1–12.3	9.9–11.5	7.3–8.8
Wind R Range <sup>a</sup>	15	5.0	nd	5.3	nd	nd	6.1	8.2
Minn R Valley <sup>a</sup>	16	nd	nd	6.0	nd	nd	5.8	7.0
Lesotho	17	nd	nd	11.8	10.1	nđ	12.6	10.0
Oaxaca Complex <sup>a</sup>	18	6.3–7.3	nd	7.3-8.0	7.5–7.8	nd	7.0-8.0	6.7-8.0
Buksefjorden	19	8.3, 7.4	9.3, 8.5	9.5, 8.7	7.6, 7.5	9.2, 8.6	11.6, 7.4	9.1, 4.8
Isortoq	20	nd	nd	7.7, 7.7	7.6, 8.3	nđ	8.1, 8.7	7.4, 7.6
Finnish Lapland	21	7.6, nd	7.9, 8.0	9.0, 9.0	7.3, nd	9.1, 9.1	8.8, 9.6	8.5, 9.2
Doubtful Sound	22	11.2, 9.6	11.2, 12.7	13.0, 11.8	12.1	12.5, 12.7	16.1, 15.4	14.0, 12.6
Broken Hill	23	nd	nd	4.5, 4.0	nd	nd	3.0, 2.5	3.8, 3.0

**Table 4.** Comparison of quoted pressures from other Opx-Pg-Gt-Qz barometers with orthopyroxene and clinopyroxene barometers of this study (using ideal pyroxene activities). References as in Table 3. nd = pressure not determined

<sup>a</sup> pressure quoted by reference for a particular locality otherwise pressure quoted by authors of calibration.

*NP*: Newton and Perkins (1982) Mg - Opx + Pg = Gt + Qz barometer

BWB: Bohlen, Wall and Boettcher (1983a) Fe - Opx + Pg = Gt + Qz barometer

PC: Perkins and Chipera (1985) Fe-Opx and Mg-Opx+Pg=Gt+Qz barometer

MAE: Moecher, Anovitz and Essene, this study, FS, HD, and DI barometers

tures are lower and cooling histories are likely to be slower for the former. Single crystal refinements and Mössbauer studies of homogeneous natural pyroxenes from the granulite facies are needed to evaluate this assumption further.

Activities of  $CaFeSi_2O_6$  in  $Cpx(a_{CaFeSi_2O_6}^{Cpx})$  calculated with the model of Davidson and Lindsley (1985) yield  $a_{CaFeSi_2O_6}^{Cpx}$  that are usually similar to or slightly greater than values obtained using the ideal approximation (Fig. 3b). The values for  $a_{CaMgSi_2O_6}^{Cpx}$  calculated from the model of Davidson and Lindsley (1985) are generally greater than ideal activities (Fig. 3c). The scatter in  $a_{CaMgSi_2O_6}^{Cpx}$  is an artifact of the projection scheme used to correct natural compositions to those appropriate for the activity model. The effect of subtracting non-quadrilateral components (mainly Al. Na, and Fe<sup>3+</sup> in granulite facies pyroxenes) in the projection scheme is to overestimate the amount of quadrilateral pyroxene components, and this projection scheme is not strictly valid for clinopyroxenes with a large fraction of nonquadrilateral components (Lindsley 1983). The greatest departure from quadrilateral space is for aluminous and sodic pyroxenes in granulites from Doubtful Sound, N.Z., in granulite xenoliths from Lesotho, and from the Westchester Prong, PA, granulites. When the Davidson and Lindsley (1985) activities are reduced by an amount equal to the mole fraction of non-quadrilateral components  $(1-Al^{vi}-2Ti-Fe^{3+}-Mn)$ , the degree of scatter is significantly reduced (Figs. 3d, 3e).

#### Evaluation of barometers

Pressures were calculated using the Gt-Cpx-Pg-Qz(HD, DI) barometers (Reactions 1 and 2) and the Gt-Opx-Pg-Qz (FS) geobarometer (Reaction 3) for 68 samples with Gt-Cpx-Opx-Pg-Qz assemblages (Tables 3 and 4). Widespread application of the FS barometer in the Central Gneiss Belt of the Grenville Province of Ontario has yielded pressures that are in good agreement with aluminosilicate occurrences and other geobarometers (Anovitz and Essene 1987b). The FS barometer also yields pressures that are consistent with reported aluminosilicate occurrences for the terranes studied here (Table 3) and previous estimates of pressure for other terranes (Table 4).

Shown in Fig.4 a are pressures obtained from the FS barometer as calculated for the different models for  $a_{\text{FerSi}_2O_6}^{\text{Opx}}$ .

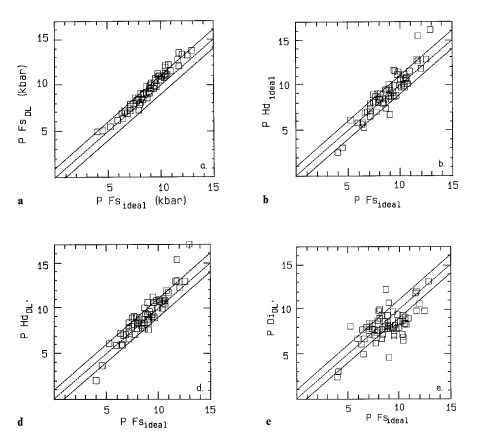


Fig. 4. a Pressures obtained for FS barometer with ideal activities for ferrosilite ( $P_{\rm Fs}$  ideal) vs. pressure obtained with model of Davidson and Lindsley ( $P_{\rm Fs}$  DL); **b** Pressures obtained for HD barometer vs. those for FS barometer, both with ideal approximation for pyroxene activities; **c** Pressures obtained for DI barometer vs. those for FS barometer, both with ideal approximation for pyroxene activities; **d** Pressures obtained for HD barometer vs. those for FS barometer, with  $a_{\rm CaFeSi_2O_6}^{\rm Cop}$  from DL adjusted for non-quadrilateral components (DL') and ideal model for ferrosilite; **e** Pressures obtained for DI barometer vs. those for FS barometer, with  $a_{\rm CaMgSi_2O_6}^{\rm Cpx}$  from DL adjusted for non-quadrilateral components (DL') and ideal model for ferrosilite

Using the Davidson and Lindsley (1985: DL) model one obtains slightly greater pressures than with the ideal model, with the greatest departures being at high pressures (low  $a_{Fe_2Si_2O_6}^{Opx}$ ). The pressure difference between the two models is well within a liberal estimate of precision for the FS barometer of  $\pm 1$  kbar. If the DL ferrosilite activities are adjusted for non-quadrilateral components in a manner similar to clinopyroxene activities, the pressure difference does not change or increases on the order of only 0.1 kbar, because of the lower amount of non-quadrilateral components in Opx relative to Cpx. For convenience we will compare the HD and DI barometers using ideal ferrosilite activities for the FS barometer.

Pressures obtained for the HD and DI barometers are plotted against pressures obtained for the FS barometer in Figs. 4b and 4c, using ideal activities for clinopyroxene and orthopyroxene. There appears to be a significant difference between the two clinopyroxene barometers relative to the orthopyroxene barometer in terms of apparent relative pressure differences and apparent precision. Some 53 of the 68 pressures obtained for the HD barometer fall within 1 kbar of equal pressure, with an average pressure difference  $(P_{\rm FS} - P_{\rm HD})$  of  $-0.2 \pm 1.0 (1\sigma)$  kbar. For the DI barometer 28 of the 68 samples fall within 1 kbar of the values obtained from the orthopyroxene barometer, and the average pressure difference  $(P_{\rm FS} - P_{\rm DI})$  is  $0.6 \pm 1.6 (1\sigma)$  kbar. The same general distribution of pressures with slightly less scatter is observed if clinopyroxene component activities are calculated using the model of Davidson and Lindsley (1985), corrected for non-quadrilateral components (Figs. 4d and 4e).

15

10

5

5

P Fs<sub>ideal</sub>

10

15

Di<sub>ideal</sub>

٦

с

We have also evaluated the HD and DI barometers by comparison with pressure constraints from reported aluminosilicate occurrences in high grade terranes (Table 4). For this purpose pyroxene activities were calculated using the ideal model. In general the HD barometer is consistent with aluminosilicate constraints when a reasonable temperature uncertainty is included, although pressures are overestimated for some samples from Parry Sound, Ontario, the Sargur Belt, India and the Furua Complex, Tanzania. The DI barometer yields pressures consistent with aluminosilicates, but pressures are lower than both the FS and HD barometers.

To a first approximation, the HD and DI barometers yield reasonable estimates of pressure for most garnet twopyroxene granulites, assuming the FS barometer is recording accurate pressures. Considering the possible sources of error in deriving the thermodynamic data, the agreement with FS pressures is satisfactory. The HD barometer shows a slight tendency to overestimate pressure relative to the FS barometer for some samples, and the DI barometer shows an opposite trend. In testing the barometers it was generally observed that the greatest pressure discrepancies using the HD barometer were for pyroxenes with a high

Mg number  $(100 (Mg/[Mg + Fe^{2+}]), e.g., Doubtful Sound,$ NZ, sample 36461. In general pyroxenes in the assemblage Gt-Cpx-Opx-Pg-Qz become more magnesian and aluminous with increasing pressure. Therefore higher pressures are accompanied by greater extrapolations from the end member Fe system and the pressures will be less reliable. Aside from the general tendency for the DI barometer to underestimate pressure, the DI barometer overestimates pressure in Fe-rich systems such as high grade banded iron formation from the Wind River Range and samples of Ferich metabasites from the Adirondack Highlands and Minnesota River Valley. This accounts for the samples that fall 3 to 4 kbar above the line of equal pressure in Figs. 4c and 4e. Conversely, Mg-rich samples with Fe-poor Opx yield high FS pressures causing the largest excursions below the line of equal pressure (Fig. 4c, 4e). The apparent disagreement between the DI and FS barometers is in part an artifact of the comparison scheme and is not seen when comparing the HD and FS barometers (Fig. 4b, 4d), because in Mg-rich rocks both  $X_{CaFeSi_2O_6}^{Cpx}$  and  $X_{FeSiO_3}^{Opx}$  will be low, yielding high pressures for both HD and FS. Therefore the highest pressures in Figs. 4b and 4d are likely to be overestimates of pressure. The data are consistent with a pressure dependence for extreme compositions, and in application of these barometers one should be aware of the possible errors involved in extrapolation of the HD barometer to Mg-rich systems, and extrapolation of the DI barometer to Fe-rich systems.

The observed pressure difference for the DI barometer relative to the FS barometer cannot be a result of the clinopyroxene barometers equilibrating at lower pressure relative to the FS barometer. If the agreement in pressure between the HD and FS barometers is real, it suggests that the DI barometer is vielding lower pressures as a result of errors in thermodynamic data and/or activity models. The difference  $(P_{\rm FS} - P_{\rm DI})$  is well within a reasonable error in  $\Delta G_{298}^{\circ}$ for any of the phases involved in the reaction, and we cannot identify any one phase as having an erroneous  $\Delta G_{298}^{\circ}$ . We believe the pressure discrepancy can be tied to ideal diopside activities. By assigning all the Mg to the M1 site, the ideal model for diopside (Eq. 18) may overestimate the activity of diopside, as some of the Mg is likely to partition into the M2 site (Cameron and Papike 1980; Dal Negro et al. 1983). This will tend to lower the  $X_{Mg}^{M1}$ , decreasing the  $a_{CaMgSi_2O_6}^{Cpx}$  and raising the pressure calculated from Reaction 2. For example, diopside activities calculated using the Wood and Banno (1973) approximation (in which Mg and  $Fe^{2+}$  are equipartitioned between the M1 and M2 sites) raises pressures by 0.1 to 0.6 kbars, with the greatest pressure increase being for the most Mg-rich clinopyroxenes.

The temperature dependence of the HD and DI barometers has been evaluated by carrying through a  $\pm 50^{\circ}$  C temperature uncertainty in the calculation of  $\log_{10} K$  and pressure. For this temperature range the value of  $\log_{10} K$ changes by only 1 to 2%, but pressure varies by  $\pm 1.0$  and  $\pm 0.3$  kbar for the HD and DI barometers respectively. The former is a relatively high temperature dependence for a barometer and illustrates the need for reasonably precise temperatures ( $\pm 30^{\circ}$  C) when applying the HD barometer. Compositional variations on the order of 1 mol% anorthite in plagioclase, hedenbergite and diopside in clinopyroxene, and almandine and pyrope in garnet correspond to pressure variations on the order of 0.1 kbar. A compositional variation of 1 mol% grossular has approximately twice the effect 101

as other components, as grossular activities are raised to a higher exponent in the expression for  $\log_{10} K$ .

#### Application of barometers

Orthopyroxene barometry is the preferred technique for calculating pressure in Gt-Opx-Cpx-Pg-Qz granulites. The available FS geobarometers have a sound experimental and thermodynamic basis, and modeling of a-X relations for orthopyroxene is relatively straightforward. However, clinopyroxene barometry is required for rocks containing Gt-Cpx-Pg-Qz assemblages without orthopyroxene. To evaluate the utility of the clinopyroxene barometers we have applied them to Gt-Cpx-Pg-Qz assemblages in rocks from several high grade metamorphic terranes.

The Whitestone Anorthosite (WSA) is a 170 km<sup>2</sup> gabbroic anorthosite body metamorphosed to granulite facies in the western Grenville Province of Ontario (Thompson 1983). Gt-Cpx-Pg-Qz assemblages occur within the main body of the anorthosite and within segments of the WSA intersected by the Parry Sound Shear Zone (PSSZ), a high grade ductile shear zone separating a predominantly amphibolite facies tectonic domain from one at granulite facies (Davidson 1984, 1986). Calculation of pressure (using ideal activities) for samples within the main body of the WSA (Moecher 1988) average 9.8 (HD) and 9.5 (DI) kbar (at 750° C, Gt-Cpx thermometry), identical to regional metamorphic pressures deduced from a variety of geobarometers ( $10 \pm 1.0$  kbar, Anovitz and Essene 1987b). In comparison, pressures from assemblages within the PSSZ average 10.8 (HD) and 11.4 kbar (DI) at 700° C (Gt-Cpx thermometry), or essentially the same as the regional data considering a 1 kbar pressure uncertainty. The pressures are consistent with the occurrence of kyanite in pelitic gneisses within the PSSZ ( $P > 7.8 \pm 1.0$  kbar at  $700 \pm 50^{\circ}$  C), and indicate that ductile shearing occurred in a deep crustal setting.

 $Gt-Cpx-Pg-Qz-(\pm Wo\pm Cc\pm Scap)$  assemblages are also common in calc-silicate rocks from the western Grenville Province. The primary compositional difference between calc-silicate and mafic granulite assemblages is that garnet in the calc-silicates has higher  $X_{Ca}^{Gt}$ , and lower  $X_{Fe^{2+}}^{Gt}$ and  $X_{Mg}^{Gt}$  than in mafic rocks. Clinopyroxene is typically a diopside-hedenbergite solid solution with minor Al, Fe<sup>3+</sup> and Na, and plagioclase shows the same range of composition as in samples of mafic granulite. Calc-silicate assemblages from the Parry Sound-Muskoka area of Ontario (Moecher, unpbl data) with high  $X_{Ca}^{Gt}$  (>0.90) yield unrealistic pressures (13 to 15 kbar) using the HD barometer and low pressures using the DI barometer (5-6 kbar), with pressure varying widely with slight variations in  $X_{\text{Fe}^{2+}}^{\text{Gt}}$  and  $X_{\text{Mg}}^{\text{Gt}}$ . It is likely that the garnet activity model used here is not accurate at such extreme garnet compositions, and in garnet with large amounts of andradite component. However, some calc-silicate samples with intermediate garnet compositions (e.g.,  $X_{Ca}^{Gt} = 0.35$  to 0.40,  $X_{Fe^{2+}}^{Gt} = 0.45$  to 0.55) yield pressures (8.9 (DI) to 11.9 kbar (HD)) that are consistent with regional pressure estimates (Anovitz and Essene 1987b). To avoid the subjective choice of which samples yield accurate pressure, calculations based on Reaction 10 would be more appropriate for high grade calc-silicate rocks.

Percival (1983) describes  $Gt-Cpx-Pg-Qz-(\pm Hbl \pm Ilm \pm Ti)$  assemblages in mafic gneisses from the Chap-

leau-Folevet area of the Wawa Belt of western Ontario, which are regarded as a cross section of deep crust upthrust along the Kapuskasing Structural Zone (Percival and McGrath 1986). Sillimanite is the only aluminosilicate found and is reported in one sample from this terrane. Application of the DI barometer of Newton and Perkins (1982) yields pressures in the range 3.9 to 8.3 kbar, with an average of 6.3 kbar for peak temperatures of 800° C (Percival 1983). These values do not include the +1.6 kbar correction suggested by Newton and Perkins (1982) which would raise average pressure to approximately the 7.9 kbar. Gt-Opx-Pg-Qz barometry by Percival and McGrath (1986) yield pressures that average 9.4 kbar (at 800° C) using the FS barometers of Bohlen et al. (1983a) and Perkins and Chipera (1985). The FS barometry indicates that the DI barometer of Newton and Perkins (1982) underestimates pressure by 3.1 kbar for these samples. We have applied the present clinopyroxene barometers to 27 samples of Gt - Cpx - Pg - Qz metabasites from this terrane, obtaining pressures from the HD and DI barometers that average  $9.6 \pm 1.3$  kbars (HD,  $1\sigma$ ) and  $8.9 \pm 1.4$  kbar (DI). Some of the pressure variation may be real variations in regional pressure. The pressures are consistent with a maximum pressure of approximately 10 kbar imposed by the occurrence of sillimanite at 800° C, and are in excellent agreement with the orthopyroxene barometry.

Tella and Eade (1986) report Gt-Cpx-Pg-Qz assemblages in fragments of mafic gneisses entrained within the Tulemalu fault zone, a high grade ductile shear zone in the Northwest Territories, Canada. The samples are interpreted as deep crustal relicts brought to the surface in granitic melts along the shear zone during ductile displacements. At temperatures of  $750 \pm 50^{\circ}$  C, the DI barometer of Newton and Perkins (1982) yielded pressures that average 10.9 kbar (unadjusted for the 1.6 kbar pressure correction). The clinopyroxene barometers of this study yield 12.9 (HD) and 11.7 (DI) kbar. It is difficult to evaluate the accuracy of these values without other petrologic constraints on pressure.

Ghent et al. (1983) have mapped a prograde Gt-Cpxisograd in sillimanite zone metabasites (Hbl-Pg-IIm  $\pm Qz \pm Gt \pm Cpx \pm Cc \pm Ti$ ) from Mica Creek, British Columbia. The Gt-Cpx isograd nearly coincides with a kyanite-sillimanite isograd, placing excellent constraints on maximum pressure of  $7.8 \pm 1$  kbar at an upper temperature limit of 700° C (based on Gt-Cpx thermometry). The DI barometer of Newton and Perkins (with correction) yields pressures of  $7.2 \pm 0.3$  kbar at 650° C (Ghent et al. 1983, 8 samples). In lower grade rocks containing kyanite, Ghent et al. (1979) and Newton and Haselton (1981) quote pressures of 6.6 to 8.9 kbar at approximately  $575 \pm 50^{\circ}$  C using the reaction

Anorthite = Grossular + Kyanite + Quartz  

$$3 \operatorname{CaAl_2Si_2O_8} = \operatorname{Ca_3Al_2Si_3O_{12}} + 2 \operatorname{Al_2SiO_5} + \operatorname{SiO_2}$$
(18)

(Ghent 1976). The clinopyroxene barometers of the present study yield apparently high values of  $9.8 \pm 0.9$  (HD) and  $10.5 \pm 0.6$  (DI) kbar at 650° C. The reason for this discrepancy is not immediately apparent. Ghent et al. (1983) see no structural evidence for an increase in pressure for the higher grade rocks near the sillimanite isograd, relative to lower grade rocks below the sillimanite isograd and that yield pressures approaching those calculated for the mafic gneisses. It appears for these rocks that the clinopyroxene barometers overestimate pressure by about 2–3 kbar, although the upper limit of pressure obtained from Reaction 18 overlaps the lower limit of pressure for the clinopyroxene barometers assuming an error estimate of about 1 kbar.

Sanders et al. (1987) describe high pressure granulite facies gneisses from the Northeast Ox Inlier of northwestern Ireland. Gt-Cpx-Pg-Qz-Ilm-Ru metabasites interpreted as decompressed eclogites occur with metapelites, metapsammites, and ultramafites. Sieve-texture intergrowths of clinopyroxene and plagioclase are interpreted to be a result of exsolution of an original omphacitic clinopyroxene. As with the Doubtful Sound, NZ samples, the presently preserved clinopyroxenes are magnesian  $(X_{HD}^{Cpx})$ = 0.03 to 0.10) and relatively high in  $Al_2O_3$  (up to 8 wt % Al<sub>2</sub>O<sub>3</sub>). Peak temperatures are estimated at 850-900° C, based on Gt-Cpx thermometry and the presence of mesoperthite, and the occurrence of kyanite in pelites (Gt-Ky-Pg-Kfs-Qz-Ru) places a minimum constraint on pressure of 10.8 to 11.8 kbar at 850-900° C. Pressure estimates for the metabasites from Gt-Cpx core compositions range from 19 (HD) to 15 (DI) kbar, and 16 (HD) to 13 (DI) kbar for rim compositions. In view of the tendency of HD barometer to overestimate pressure for Mgrich clinopyroxenes, the HD values are likely to be upper limits on pressure. However pressures calculated from Reaction 18 for the Gt-Ky-Pg-Qz assemblage reported by Sanders et al. (1987), using the calibration of Newton and Haselton (1981) and Essene (unpl. data), are also in excess of 15 kbars (at 800° C), in agreement with the DI barometer. Pressures calculated from Gt-Pg-Ilm-Ru-Qz assemblages in the metabasite (Reaction 8: Anovitz and Essene 1987a) yield minimum pressure estimates of 12 to 13 kbar (assuming  $a_{\text{FeTiO}_3}^{\text{IIm}} = 1$ ). If equilibrium can be demonstrated for the sieve-texture clinopyroxenes then the pressures indicated by the geobarometry are consistent as a whole with the extremely deep crustal history for these samples suggested by Sanders et al. (1987).

Wood (1975) presents analyses for coexisting garnet, clinopyroxene inclusions within garnet, and plagioclase from metagabbros from the South Harris, Scotland, Igneous Complex. The presence of kyanite in metapelites associated with the meta-igneous lithologies places a lower limit on pressure of 9 kbar. Pressures were estimated to range from 10 to 13 kbar at 800 to 860° C (two-pyroxene and Gt-Cpx thermometry), based on constraints from Gt-Ol-Pg and Gt-Opx-Pg-Qz barometry. We have recalculated the Gt-Cpx temperatures to be 750° C using the Ellis and Green (1979) thermometer, and these are likely to be lower limits on temperature as garnet and clinopyroxene in contact with one another tend to re-equilibrate with falling temperature after the peak of metamorphism (Johnson et al. 1983; Moecher et al. 1986). Pressures for three samples average 12.3 (HD) and 10.6 (DI) kbar. One sample with  $X_{Hd}^{Cpx}$ =0.07 yields the highest pressure, and as with the samples from Doubtful Sound, NZ and the Northeast Ox Inlier, Ireland, this is likely to yield an overestimate of pressure. Excluding this sample the average HD pressure is 11.3 kbar. Metamorphic pressure for the South Harris area is likely to be  $11 \pm 1$  kbar, essentially that predicted by Wood's earlier analysis.

Evaluation of the HD and DI barometers in previous sections indicated that the DI barometer tended to yield somewhat lower and more scattered pressures than the HD and FS barometers. The results from the six areas above indicate that the two clinopyroxene barometers are in better agreement with one another. This is not inconsistent with the former observation, as some of the garnet two-pyroxene granulites yield similar pressures for all three barometers. From the foregoing applications it is apparent that the clinopyroxene barometers of this study can yield reasonable pressures for most Gt-Cpx-Pg-Qz assemblages. In the absence of assemblages for which experimentally determined geobarometers are available, the HD and DI barometers can be applied with reasonable precision in the granulite facies. However, the thermodynamic, temperature, and compositional dependencies of these barometers, and the inherently less precise nature of calibrations based largely on thermodynamic data compared to reversed experiments, must be kept in mind when applying the barometers in natural settings. More accurate calibrations of these barometers will require careful experiments on Reactions 1 and 2, or on the reactions that constrain the  $\Delta G_{298}^{\circ}$  of hedenbergite and pyrope.

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#### Appendix I

Mineral formulae, abbreviations, and symbols

Albite (Ab):	NaAlSi <sub>3</sub> O <sub>8</sub>
Almandine (Alm):	Fe <sub>3</sub> Al <sub>2</sub> Si <sub>3</sub> O <sub>12</sub>
Anorthite (An):	CaAl <sub>2</sub> Si <sub>2</sub> O <sub>8</sub>
Andradite (And):	$Ca_3Fe_2Si_3O_{12}$
Corundum (Cor):	$Al_2O_3$
Diopside (Di):	CaMgSi <sub>2</sub> O <sub>6</sub>
Enstatite (En):	MgSiO <sub>3</sub>
Fayalite (Fa):	Fe <sub>2</sub> SiO <sub>4</sub>
Ferrosilite (Fs):	FeSiO <sub>3</sub>
Grossular (Gr):	$Ca_3Al_2Si_3O_{12}$
Hematite (Hm):	Fe <sub>2</sub> O <sub>3</sub>
Kyanite $(Ky)$ :	$Al_2SiO_5$
Mg-Tschermakite (MgTs):	Mg <sub>0.5</sub> AlSi <sub>0.5</sub> O <sub>3</sub>
Magnetite ( <i>Mt</i> ):	Fe <sub>3</sub> O <sub>4</sub>
Pyrope (Py):	$Mg_3Al_2Si_3O_{12}$
Quartz $(Qz)$ :	SiO <sub>2</sub>
Sillimanite (Sill):	$Al_2SiO_5$
Wollastonite (Wo):	CaSiO <sub>3</sub>
	-

Cc: calcite Cpx: clinopyroxene Gt: garnet Hbl: hornblende Kfs: potassium feldspar Opx: orthopyroxene Pg: plagioclase Scap: scapolite Ti: titanite

 $\Delta G_{298}^{\circ}$ : Gibbs free energy of reaction at STP (kJ/mol)  $V_{298}^{\circ}(j)$ : molar volume of phase *j* at STP (cm<sup>3</sup>/mol)  $S_{298}^{\circ}(j)$ : molar entropy of phase *j* at STP (J/mol·K)  $\Delta V$ : volume change of reaction  $\Delta S$ : entropy change of reaction  $X_j^i$ : mole fraction of component *i* in phase *j*  $\gamma_i^i$ : activity coefficient of component *i* in phase *j*  $a_j^i$ : activity of component *i* in phase *j* P: pressure, kbar T: temperature, K or °C where stated

K: equilibrium constant

R: ideal gas constant, 8.3145 J/mol·K

# Appendix II

Activity models

*Pyroxene*. The ideal activity models for clinopyroxene used in this study are summarized in the text, equations 17 and 18 for hedenbergite and diopside, respectively. The model used for orthopyroxene is that of Wood and Banno (1973) in which the activity of ferrosilite is taken as:

# $a_{\text{Fe}_2\text{Si}_2\text{O}_6}^{\text{Opx}} = [X_{\text{Fe}^{2+}}^{\text{M1}}] [X_{\text{Fe}^{2+}}^{\text{M2}}].$

Pyroxene analyses are normalized to 4 cations and  $Fe^{3+}$  is calculated from charge balance and stoichiometry. After subtracting the atomic amounts of Ca, Na, and Mn from the M2 site, and subtracting Al<sup>vi</sup>, Ti, Cr, and Fe<sup>3+</sup> from the M1 site, Fe<sup>2+</sup> is partitioned into each remaining site in the same ratio as  $Fe^{2+}/(Fe^{2+} + Mg)$  in the mineral.

The equilibrium constant for the FS barometer includes a term for  $a_{\text{Fe}_2\text{Si}_2\text{O}_6}^{\text{Opx}}$  that is raised to the 3rd power. Because the two site model squares  $X_{\text{Fs}}$ , the activity of ferrosilite calculated above is taken to the 3rd power in calculating the equilibrium constant for Reaction 3, although the thermodynamic data for ferrosilite are based on a 2 cation formula (6 moles of Fs for Reaction 3). When calculating the activity of  $FeSiO_3$  simply as mole fraction of  $Fe^{2+}$ , the exponent for Fs in Reaction 3 is raised to the 6th power.

*Plagioclase.* Anorthite activities were calculated according to the model of Newton et al. (1980), also given in Newton and Haselton (1981), Newton and Perkins (1982), and Newton (1983). Plagioclase analyses were normalized to 5 cations, and  $X_{An}^{Pg} = Ca/(Ca + Na)$ . The activity of anorthite is calculated from the following relations:

 $RT \ln \gamma_{An}^{Pg} = (X_{Ab}^{Pg})^2 [2050 + 9392 X_{An}^{Pg}]$  $a_{CaAl_2Si_2O_8}^{Pg} = \gamma_{An}^{Pg} [X_{Mg}^{Gt} - (1 + X_{An}^{Pg})^2]/4$ 

Garnet. Grossular, pyrope, and almandine activities were calculated using the quaternary garnet mixing model of Ganguly and Saxena (1984), with the Ca – Fe mixing parameters for the grossular-almandine join calculated by Anovitz and Essene (1987a). The Ca – Mg, Ca – Mn, Fe – Mg, Fe – Mn, and Mg – Mn mixing parameters are the same as in Ganguly and Saxena (1984). Use of these parameters yields the following analytical expressions for the activity coefficient of grossular, pyrope, and almandine:

 $RT \ln \gamma_{C_a}^{Gt} = (X_{M_a}^{Gt})^2 (4047 - 1.5 T - 6094 X_{C_a}^{Gt})$  $+(X_{Fe^{2+}}^{Gt})^{2}(150-1.5T+7866X_{Ca}^{Gt})$  $+ X_{Mg}^{Gt} X_{Fe^{2+}}^{Gt} [3290 - 3.0 T + 886 X_{Ca}^{Gt} + 2300 (X_{Mg}^{Gt} - X_{Fe^{2+}}^{Gt})]$  $+4640(1-2X_{Ca}^{Gt})$ ]  $+X_{\text{Fe}^{2+}}^{\text{Gt}}X_{\text{Mn}}^{\text{Gt}}[2117-1.5T+3933X_{\text{Ca}}^{\text{Gt}}-1967(1-2X_{\text{Ca}}^{\text{Gt}})]$  $+ X_{Mg}^{Gt} X_{Mn}^{Gt} [2524 - 1.5 T - 3047 X_{Ca}^{Gt} + 1524(1 - 2 X_{Ca}^{Gt})]$  $+2300 \left[ X_{Mg}^{Gt} X_{Fe^{2+}}^{Gt} X_{Mn}^{Gt} \right]$  $RT \ln \gamma_{Mg}^{Gt} = (X_{Ca}^{Gt})^2 (1000 - 1.5 T + 6094 X_{Mg}^{Gt})$  $+(X_{\rm Fe^{2+}}^{\rm Gt})^2(2500-4600X_{\rm Mg}^{\rm Gt})+3000(X_{\rm Mg}^{\rm Gt})^2$  $+ X_{Ca}^{Gt} X_{Fe^{2+}}^{Gt} [1757 + 747 X_{Me}^{Gt} - 3933 (X_{Ca}^{Gt} - X_{Fe^{2+}}^{Gt})]$  $+4640(1-2X_{Mg}^{Gt})$ ]  $+ X_{Fe^{2+}}^{Gt} X_{Mn}^{Gt} [4350 - 2300 X_{Mg}^{Gt} + 1150(1 - 2X_{Mg}^{Gt})]$  $+ X_{Ca}^{Gt} X_{Mn}^{Gt} [5524 + 3047 X_{Mn}^{Gt} - 1524(1 - 2X_{Mn}^{Gt})]$  $+3933 [X_{Ca}^{Gt} X_{Fe^{2+}}^{Gt} X_{Mn}^{Gt}]$  $RT \ln \gamma_{\text{Fe}^{2+}}^{\text{Gt}} = (X_{\text{Mg}}^{\text{Gt}})^2 200 + 4600 X_{\text{Fe}^{2+}}^{\text{Gt}})$  $+(X_{C_a}^{Gt})^2(4083-1.5T-7866X_{Fe^{2+}}^{Gt})$  $+ X_{M_{e}}^{Gt} X_{C_{a}}^{Gt} [943 - 1633 X_{Fe^{2+}}^{Gt} - 3047 (X_{M_{e}}^{Gt} - X_{C_{a}}^{Gt})$  $-4640(1-2X_{\rm Fe^{2+}}^{\rm Gt})$ ]  $+ X_{Ca}^{Gt} X_{Mn}^{Gt} [2117 - 1.5 T - 3933 X_{Fe^{2+}}^{Gt} + 1917(1 + 1917)]$  $-2X_{\rm Fe^{2}+}^{\rm Gt}$ +  $X_{Mg}^{Gt} X_{Mn}^{Gt} [-1650 + 2300 X_{Fe^{2+}}^{Gt} - 1150(1 - 2X_{Fe^{2+}}^{Gt})]$  $+ 3048 [X_{Mg}^{Gt} X_{Ca}^{Gt} X_{Mn}^{Gt}].$ 

Because there are three equivalent 8-coordinated sites in garnet, the activity of  $Ca_3Al_2Si_3O_{12}$ ,  $Mg_3Al_2Si_3O_{12}$ , and  $Fe_3Al_2Si_3O_{12}$  are calculated as

 $\begin{aligned} & a_{\text{Cs}_{3}\text{Al}_{2}\text{Si}_{3}\text{O}_{12}}^{\text{Gt}} = (X_{\text{Ca}}^{\text{Gt}}\gamma_{\text{Ca}}^{\text{Gt}})^{3}, \\ & a_{\text{Mg}_{3}\text{Al}_{2}\text{Si}_{3}\text{O}_{12}}^{\text{Gt}} = (X_{\text{Mg}}^{\text{Gt}}\gamma_{\text{Mg}}^{\text{Gt}})^{3}, \\ & a_{\text{Fe}_{3}\text{Al}_{2}\text{Si}_{3}\text{O}_{12}}^{\text{Gt}} = (X_{\text{Fe}^{2}}^{\text{Ft}}\gamma_{\text{Fe}^{2}}^{\text{Gt}})^{3}. \end{aligned}$ 

Garnet analyses are normalized to 8 cations with  $Fe^{3+}$  calculated form charge balance and stoichiometry. For garnets with Si > 3.00,  $Fe^{3+}$  was calculated from the relation  $Fe^{3+} = 2 - Al - 2$  Ti - Cr.