# Volume behavior of hydrous minerals at high pressure and temperature: II. Compressibilities of lawsonite, zoisite, clinozoisite, and epidote

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#### Abstract

The pressure dependence of the lattice parameters of natural zoisite  $[Ca_2Al_3Si_3O_{12}(OH)]$ , clinozoisite  $[Ca_2Al_3Si_3O_{12}(OH)]$ , and epidote  $[Ca_2Al_2FeSi_3O_{12}(OH)]$  as well as synthetic lawsonite  $[CaAl_2Si_2O_7(OH)_2 \cdot H_2O]$  have been measured at ambient temperatures by energy-dispersive X-ray diffraction in a diamond-anvil cell. The experimental results for each phase may be summarized concisely in terms of the ambient-temperature isothermal bulk modulus  $K_{298}$  (using the Murnaghan equation of state and assuming K' = 4): Lawsonite:  $K_{298} = 191 \pm 5$  GPa; zoisite:  $K_{298} = 279 \pm 9$  GPa; clinozoisite:  $K_{298} = 154 \pm 6$ GPa; epidote:  $K_{298} = 162 \pm 4$  GPa. These new measurements, together with the new thermal expansion data in the companion paper (Pawley et al. 1996), were used to calculate some phase equilibria for lawsonite dehydration to high pressures for comparison with experimental brackets. Important discrepancies between calculated and experimentally determined reactions become evident above 3 GPa.

## INTRODUCTION

Two factors have recently led to a renewed interest in the behavior of minerals at very high pressures. First, the ready availability of experimental apparatus capable of achieving pressures in the range of 20 GPa has led to an increase in the number and extent of experimental determinations of the stabilities of mineral phases, including hydrous silicates to these very high pressures (e.g., Pawley 1994; Schmidt and Poli 1994). Second, the increased attention of petrologists and geophysicists on the nature and behavior of the lithospheric slab as it descends into the mantle has raised several important questions concerning the amount and distribution of fluids at deep levels in subduction zones. Current thermodynamic models for hydrous and carbonate phases are not sufficiently complete to predict the temperatures and pressures of mineral decarbonation and dehydration reactions up to the pressures and temperatures prevailing in the deeper parts of subduction zones. Very basic data, such as the thermal expansivity and, more importantly, compressibility of common hydrous minerals likely to be important components in metamorphosed rocks in the descending slab, have not been measured, and simple linear extrapolations of molar volume, which are adequate for modeling crustal processes, are less accurate at the very high pressures of interest. Thus, we have selected four common mineral phases, which are believed to be likely hosts for transporting H<sub>2</sub>O into the mantle in subduction zones, to measure their bulk moduli and extend our capability of calculating mineral assemblage stabilities at these high pressures and temperatures. The significance of three of these phases as potential hydrated silicates in subducted slabs [lawsonite (*Ccmm*), CaAl<sub>2</sub>Si<sub>2</sub>O<sub>7</sub>(OH)<sub>2</sub>. H<sub>2</sub>O; zoisite (*Pnma*), Ca<sub>2</sub>Al<sub>3</sub>Si<sub>3</sub>O<sub>12</sub>(OH); and clinozoisite (*Pnma*), Ca<sub>2</sub>Al<sub>3</sub>Si<sub>3</sub>O<sub>12</sub>(OH)] was discussed in the companion paper (Pawley et al. 1996).

In addition to these three phases, we also investigated the compressibility of epidote. Fe<sup>3+</sup>-containing epidote  $(P2_1/m)$ , Ca<sub>2</sub>Al<sub>2</sub>FeSi<sub>3</sub>O<sub>12</sub>(OH), is common in rocks regionally metamorphosed under greenschist- and amphibolite-facies conditions. It also occurs in blueschists and eclogites crystallized in subduction-zone environments, and in some mafic igneous rocks. In experiments under possible subduction-zone *P*-*T* conditions, epidote of composition Ca<sub>2</sub>Al<sub>2,3</sub>Fe<sub>0.7</sub>Si<sub>3</sub>O<sub>12</sub>(OH) was found to be stable at pressures up to 3 GPa (Pawley and Holloway 1993). Its identification as monoclinic epidote, and not orthorhombic zoisite, was confirmed using Fourier-transform infrared spectroscopy.

## SAMPLES AND EXPERIMENTAL TECHNIQUE

The samples of natural zoisite, clinozoisite, and synthetic lawsonite used in our measurements are the same as those described in the companion paper (Pawley et al. 1996). The sample of epidote is from a monomineralic vein in high-pressure albite-amphibolites from the Gross Venediger area in the central Höhe Tauern, Eastern Alps (Holland 1979; Holland and Ray 1985). It is very close to the end-member composition; chemical analyses, by electron microprobe, of the epidote and clinozoisite sam-

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TABLE 1. Compositions of natural clinozoisite and epidote

	Clino	zoisite	Epidote		
	wt%	Cations	wt%	Cations	
SiO,	39.58	2.989	38.01	3.013	
TIO,	0.0	0.0	0.04	0.002	
Al <sub>2</sub> O <sub>3</sub>	33.65	2.996	21.41	2.001	
Cr <sub>2</sub> O <sub>3</sub>	0.0	0.0	0.09	0.006	
Fe <sub>2</sub> O <sub>3</sub>	0.34	0.019	16.12	0.961	
FeO	0.0	0.0	0.26	0.017	
MnO	0.0	0.0	0.06	0.004	
MgO	0.0	0.0	0.0	0.000	
CaO	24.70	2.000	23.48	1.996	
Na <sub>2</sub> O	0.0	0.0	0.0	0.0	
K,Õ	0.0	0.0	0.0	0.0	

ples used in this study are given in Table 1. All the small amount of Fe in clinozoisite was assumed to be  $Fe^{3+}$  in the recalculation of the microprobe data; for the epidote sample, however, only a small proportion of Fe was assumed to be  $Fe^{2+}$ , and this quantity was calculated on the basis of eight cations per formula unit with 12.5 O atoms. Clean and clear individual crystals of zoisite, clinozoisite, and epidote were hand-picked from the natural samples, and they and a batch of the synthetic lawsonite were crushed and ground to a fine powder for X-ray diffraction experiments. Individual powder samples were mixed with powdered NaCl, which acted as an internal standard and pressure calibrant for the high-pressure measurements.

High-pressure powder diffraction was performed, in a manner identical to that used in the study of Redfern et al. (1993), on the wiggler station 9.7 of the SERC synchrotron radiation source at Daresbury, U.K. The powdered sample was loaded into a preindented, heat-treated Inconel gasket (200 mm hole) in a lever-arm diamondanvil cell. A 4:1 nondried methanol-ethanol mixture was used as the pressure-transmitting medium. Measurements were made in energy-dispersive mode at room temperature between ambient pressure and 16 GPa. White radiation, collimated by a 150 µm pinhole, was diffracted at a fixed  $2\theta$  angle (accurately determined for each experiment from the diffraction pattern of NBS silicon powder) and further collimated using 500 mm long molybdenum receiving flats shimmed 100 µm apart. Spectra were collected between 5 and 120 keV using a lithium-drifted germanium detector with a resolution varying from 149 eV at 5.9 keV to 468 eV at 122 keV.

Pressure was calibrated with the use of the measured cell volume of the internal NaCl standard (determined from least-squares refinement of a minimum of three Bragg peaks) and the equation of state for NaCl given by Decker (1971). The estimated uncertainty in the pressure readings is  $\pm 2\%$ . Sample and NaCl diffraction peak positions were measured by Gaussian peak fitting of diffracted intensity. Values for unit-cell parameters were refined by a nonlinear least-squares method using the program UnitCell (Holland and Redfern in preparation), and from 7 to 15 peaks depending on sample and pres-

sure. Typical diffraction patterns for the four samples under pressure loading are shown in Figure 1.

### RESULTS

Values of cell parameters as a function of pressure are listed in Table 2 and plotted in Figures 2–5 as relative compressions. Uncertainties are shown only for relative volume compression for the sake of clarity in the figures. Values of the isothermal bulk modulus, K, and its pressure derivative, K', are commonly derived by fitting the pressure-volume data to the Birch-Murnaghan equation

$$P = \frac{3}{2} K \left[ \left( \frac{V_0}{V} \right)^{\frac{7}{3}} - \left( \frac{V_0}{V} \right)^{\frac{5}{3}} \right] \left\{ 1 - \frac{3}{4} (4 - K') \left[ \left( \frac{V_0}{V} \right)^{\frac{2}{3}} - 1 \right] \right\}$$

or to the simpler Murnaghan equation

$$P = \frac{K}{K'} \left[ \left( \frac{V_0}{V} \right)^{K'} - 1 \right]$$

which may be conveniently rearranged into volume-explicit form,

$$\frac{V}{V_0} = \left[1 + \frac{K'}{K}P\right]^{-\frac{1}{K'}}.$$

We fitted our measured volume data with both the Birch-Murnaghan and Murnaghan equations and found that, over the pressure ranges of this study, both equations yield identical results within error. Rather than fit the data to the Birch-Murnaghan equation in the form above by minimizing residuals in pressure, we adapted a nonlinear least-squares routine to refine initial guesses for  $V_0$ ,  $K_{298}$ , and K' by minimizing the residuals in volume (the measured parameter). The values for  $K_{298}$  and K' are often different when residuals in V rather than P are minimized, and the former is much the more useful because we wish to predict the volume from the pressure and not vice versa. Regression for three parameters,  $V_0$ ,  $K_{298}$ , and K', is not warranted by the quality of the data especially because the parameters are very highly correlated (e.g., increasing K' reduces  $K_{298}$  and  $V_0$ ). The scatter in our measured volumes is also too large to enable very precise values of K' to be determined, and attempts to obtain precise values for epidote and zoisite led to physically unrealistic (negative) values, the uncertainties of which were larger than the absolute values. Thus, we elected to set the pressure derivative K' to 4, a commonly accepted assumption that is based on the results of many measurements on silicate and oxide phases of geological interest and equivalent to reducing the Birch-Murnaghan equation to a simpler form by eliminating the final term (Anderson 1989; Liu and Bassett 1986). Nonlinear regression of the volume data can be found in Table 3.

### DISCUSSION

The two mineral end-members of the monoclinic epidote solid-solution series, clinozoisite and epidote, show virtually identical compression behavior, with c being

1

1



FIGURE 1. Energy-dispersive diffraction patterns of clinozoisite, zoisite, epidote, and lawsonite in the diamond-anvil cell at 1.5, 1.6, 0.4, and 0.7 GPa, respectively, displayed on a common energy scale. The step in the background (due mainly to Compton scattering in the diamonds) at ~20 keV is due to the absorption edge of diamond. The NaCl reflections are indicated, other peaks being indexed appropriately. The diffraction patterns of clinozoisite, zoisite, and epidote were collected with the detector at approximately 10.14°  $2\theta$ , and the diffraction pattern of lawsonite was collected with the detector fixed at 7.11°  $2\theta$ .

most compressible and b and a being less affected. The orthorhombic polymorph, zoisite, is rather different in its compression from the two monoclinic members in that its b axis is most compressible, with a and c much less affected. Lawsonite compression mainly results in shortening of the a and b axes, with less pronounced shortening of the c axis. Although zoisite has a smaller volume at room pressure, the larger compressibility of clinozoisite means that the monoclinic phase becomes the denser

ABLE 2.	Hig	h-pressure	cell	const	tants
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P (GPa)	a (Å)	b (Å)	c (Å)	β (°)	V (ų)
		Cline	zoisite		
0.1	8.859(14)	5.588(6)	10.151(12)	115.46(10)	453.7(7)
1.31	8.845(16)	5.570(8)	10.103(15)	115.40(11)	449.6(8)
1.50	8.848(7)	5.579(7)	10.092(11)	115.39(7)	450.1(5)
2.13	8.826(12)	5.559(5)	10.098(13)	115.41(9)	447.5(6)
2.33	8.814(11)	5.500(0)	10.060(9)	115.41(7)	447.0(5)
3 70	8.825(7)	5.535(4)	10.030(12)	115.30(7)	442.9(5)
4.58	8.790(16)	5.533(8)	10.027(27)	115.36(17)	439.9(10)
5.60	8.777(16)	5.531(7)	10.006(19)	115.24(12)	439.3(8)
5.69	8.779(13)	5.524(5)	9.966(22)	115.14(15)	437.5(8)
7.75	8.755(14)	5.504(5)	9.923(24)	115.18(16)	432.7(9)
3.20	8.757(31)	5.512(12)	9.918(75)	115.37(45)	432.6(28)
		Ep	idote		
0.0001	8.890(5)	5.641(3)	10.164(6)	115.55(7)	459.9(3)
J.U5	8.894(7)	5.030(5)	10.162(9)	115.53(6)	459.0(5)
0.10	8,896(7)	5.640(6)	10.165(1)	115.55(0)	459.7(4)
1.66	8.874(6)	5.623(5)	10.121(9)	115.62(6)	455.3(4)
2.95	8.865(4)	5.612(3)	10.109(5)	115.80(5)	452.8(2)
3.41	8.851(4)	5.608(3)	10.095(5)	115.80(6)	451.2(3)
4.24	8.836(5)	5.599(3)	10.060(6)	115.72(6)	448.4(3)
1.80	8.836(5)	5.596(4)	10.049(6)	115.83(6)	447.2(3)
5.21	8.827(8)	5.595(6)	10.030(8)	115.81(8)	445.9(6)
		Zo	oisite		
0.60	16.177(13)	5.554(7)	10.027(10)		901.0(12)
0.78	16.166(14)	5.552(7)	10.031(10)		900.3(12)
1.21	16.198(14)	5.549(6)	10.023(10)		900.8(12)
5.20	16 150(11)	5.494(3)	10.012(14)		888 4(6)
6.39	16 136(28)	5.503(9)	9.982(15)		886.3(19)
6.92	16.103(54)	5.487(19)	10.005(40)		884.0(37)
7.66	16.099(21)	5.504(7)	9.973(13)		883.7(15)
8.91	16.078(29)	5.476(12)	9.958(16)		876.8(21)
9.78	16.087(28)	5.473(9)	9.925(19)		873.8(20)
9.85	16.097(12)	5.459(5)	9.953(7)		874.5(9)
0.69	16.088(17)	5.462(10)	9.954(15)		8/4.0(10)
1.20	16 103(10)	5.431(10)	9.951(10)		868 7(13)
2 22	16.051(17)	5 444(13)	9.937(16)		868.4(12)
3.47	16.061(47)	5.390(16)	9.950(28)		861.2(30)
3.79	16.061(30)	5.418(30)	9.918(25)		863.1(42)
		Law	vsonite		
0.07	8.782(5)	5.833(2)	13.097(9)		670.9(4)
0.81	8.772(10)	5.825(4)	13.083(15)		668.5(8)
1.51	8.752(7)	5.815(4)	13.064(17)		664.8(8)
2.31	8.742(7)	5.800(4)	13.059(16)		662 5(4)
2.40	8.747(0)	5.807(3)	13.003(9)		658 4(8)
3.00 / 38	8 713(7)	5 787(3)	13.010(9)		655 9(5)
4.87	8.697(11)	5.776(5)	13.015(16)		653.8(9)
5.72	8.705(17)	5.772(8)	13.016(25)		654.0(14)
6.66	8.683(21)	5.763(8)	12.993(31)		650.3(17)
7.64	8.667(19)	5.754(9)	12.983(27)		647.4(16)
8.75	8.649(25)	5.746(12)	12.948(37)		643.4(21)
9.97	8.626(23)	5.729(11)	12.967(34)		640.8(19)
0.30	8.614(44)	5.740(14)	12.954(46)		630 5(11)
1 21	8 615/201	5 719(10)	12.930(79)		636.3(17)
1.40	8.625(31)	5.730(15)	12.958(54)		640.3(29)
2.05	8.613(44)	5.704(21)	12.930(75)		635.2(42)

of the two above  $\sim 2$  GPa; from the thermal expansion measurements [companion paper (Pawley et al. 1996)] it was concluded that clinozoisite also becomes denser than zoisite above ~300 °C at room pressure. The large difference in bulk modulus between zoisite and clinozoisite is noteworthy because it is somewhat unexpected. Our Relative Compression

1.010

1.00

0.990

0.980

0.960

0.950

0.940

0.930<sup>L</sup> 0  $\Box a/a_o$ 

c/co

V/V

2

◊ b/b₀

0

Lawsonite



6

8

10

12

14

K = 191(5)with K' = 4

data show that zoisite is far less compressible than clinozoisite. It would be convenient if we could offer a simple structural explanation for this difference, yet no other high-pressure study of these phases exists and the detailed structural responses to pressure must yet be identified. Comparing the Ca sites of zoisite and clinozoisite at ambient pressure and temperature, we observed that the Ca1 and Ca2 sites are significantly larger in the orthorhombic phase than in the monoclinic phase, and we would expect the Ca-O bond strength in zoisite to be weaker. Thus, the CaO<sub>n</sub> polyhedra in zoisite might at first be expected to



**FIGURE 3.** Relative compression of zoisite. Variation of cell parameters, normalized to values at 1 bar, as a function of pressure. The curve for volume is a least-squares fit, giving  $K_{298} = 279 \pm 9$  GPa (Murnaghan EOS); those for individual cell parameters are just guides for the eye. Error bars for  $V/V_0$  are two standard deviations; error bars on individual axial ratios are larger than symbols.



**FIGURE 4.** Relative compression of clinozoisite. Variation of cell parameters, normalized to values at 1 bar, as a function of pressure. The curve for volume is a least-squares fit, giving  $K_{298} = 154 \pm 6$  GPa (Murnaghan EOS); those for individual cell parameters are just guides for the eye. Error bars for  $V/V_0$  are two standard deviations; error bars on individual axial ratios are larger than symbols.

be more compressible, not less. Clearly the answer lies in some other aspect of the structure, possibly the influence of the AlO<sub>6</sub> octahedral chains and the silicate cross-linking. Furthermore, because the thermal expansion of zoisite is marginally greater than that of clinozoisite, increasing pressure in this phase cannot be thought of as volumetrically equivalent to decreasing temperature.

Zoisite is, therefore, the denser polymorph only close to ambient conditions. Under most metamorphic and all igneous conditions the denser phase is clinozoisite. These



**FIGURE 5.** Relative compression of epidote. Variation of cell parameters, normalized to values at 1 bar, as a function of pressure. The curve for volume is a least-squares fit, giving  $K_{298} = 162 \pm 4$  GPa (Murnaghan EOS); those for individual cell parameters are just guides for the eye. Error bars for  $V/V_0$  are two standard deviations; error bars on individual axial ratios are larger than symbols.

TABLE 3. Nonlinear regression of the volume data

	Murnagh	nan EOS	Birch-Murnaghan EOS		
	K298 (GPa)	V <sub>0</sub> (Å <sup>3</sup> )	К <sub>298</sub> (GPa)	V <sub>0</sub> (Å <sup>3</sup> )	
Lawsonite	191(5)	671.1(3)	191(6)	671.1(4)	
Zoisite	279(9)	904.4(6)	267(9)	905.6(6)	
Clinozoisite	154(6)	453.8(4)	154(7)	453.8(4)	
Epidote	162(4)	460.3(2)	161(5)	460.3(3)	

density relations should not be confused with the relative stability of the two polymorphs, which remains an unresolved problem and may require precise thermodynamic data to clarify given the sluggish kinetics of such transitions at low temperatures. However, although the arguments are beyond the scope of this paper and involve the relative heat capacities and entropies of the two polymorphs as well as natural Fe-Al partitioning, we believe that the transition has a steep positive dP/dT slope and occurs at temperatures near 350 °C, with zoisite as the stable phase on the high-temperature side (Holland in preparation). Thus, in the applications that follow, we calculate phase relations only at the elevated temperatures at which the orthorhombic form, zoisite, is stable.

To use both the compressibilities measured in this study and the thermal expansivities from the companion paper (Pawley et al. 1996) for thermodynamic calculations at high pressures and temperatures, we also need to estimate the temperature dependence of K,  $\partial K/\partial T$ . The bulk modulus K is approximately linear in T, and Anderson and Isaak (1993) showed that  $\delta_T = -(1/\alpha K)(\partial K/\partial T)_p$ . The quantity  $\partial K/\partial T$  is approximately constant for each mineral, and the Anderson-Gruneisen parameter  $\delta_T$  is a small positive number somewhat larger than K'. Taking  $\partial K/\partial T$ to be proportional to  $-\alpha K$  at 298 K, we found that the relationship  $\partial K/\partial T = -7.0(\pm 1.0)\alpha_{298}K_{298}$  reproduces most of the known experimental data. In addition, if the thermal expansivities are represented by  $\alpha = a_0 - 10a_0T^{1/2}$  as proposed in the companion paper (Pawley et al. 1996), then  $\partial K/\partial T = -3a_0K_{298}$ . In the thermodynamic calculations to follow we assume that  $\partial K/\partial T$  is constant, and so  $K_T = K_{298} - 3a_0K_{298}(T - 298)$ .

The Murnaghan equation is quite adequate for calculating volumes to 20 GPa, and thus the pressure integral for the Gibbs energy may be written as

$$\int_{0}^{P} V \, \mathrm{d}P = \frac{K_T V_T}{3} \left[ \left( 1 + \frac{4P}{K_T} \right)^{\frac{3}{4}} - 1 \right]$$

where  $V_T$  is the ambient-pressure molar volume found from the expressions given in the companion paper (Pawlev et al. 1996). The new formulation [above, and Pawley et al. (1996)] for thermal expansions and the Murnaghan equation for compressibilities are now used in a completely revised version of the internally consistent thermodynamic dataset and computer program THERMO-CALC described by Holland and Powell (1990). This updated version also incorporates a large body of new experimental data as well as the CORK equations for gases at high pressure and temperature (Holland and Powell 1991). The basic thermodynamic data used are presented in Table 4, and the phase relations at high pressures are shown in Figures 6 and 7 for comparison with the recent experimental data of Pawley (1994), Skrok et al. (1994), and Schmidt and Poli (1994) as well as earlier experiments of Newton and Kennedy (1963) and Chatterjee et al. (1984) on lawsonite breakdown. Figure 6a shows the lawsonite breakdown reaction to zoisite + kyanite + quartz or coesite +  $H_2O$  and shows that agreement between calculations and experiment is excellent to 3 GPa but steadily worsens as pressure increases. This behavior was noted in the companion paper (Pawley et al. 1996) for diaspore breakdown to corundum + H<sub>2</sub>O and therefore is probably a feature not specifically related to the thermodynamic properties of any of the solid phases in these reactions (none is involved in either reaction). The problem may be friction in the very-high-pressure

TABLE 4. Thermodynamic properties of selected phases in CASH

	Н	sd(H)	S	V	а	b	c	d	<i>a</i> 0	K <sub>298</sub>
Gr	-6640.81	3.39	0.256	12.535	0.6260	0	-5779.2	-4.0029	3.93	1680
Law	-4868.85	1.68	0.230	10.132	0.6878	0.1566	375.9	-7.1792	5.82	1910
Zo	-6897.65	2.82	0.296	13.575	0.5957	6.2297	-5921.3	-3.3947	6.77	2790
Kv	-2595.38	1.08	0.082	4.414	0.3039	-1.3390	-895.2	-2.9040	4.04	1590
Stv	-863.73	1.14	0.030	1.401	0.0681	0.6010	-1978.2	-0.0821	2.50	3100
Q	-910.75	0.45	0.0415	2.269	0.0979	-0.3350	-636.2	-0.7740	4.32	400
Ba	-908.41	0.45	0.04425	2.367	0.0979	-0.3350	-636.2	-0.7740	0.65	650
Coe	-907.64	0.45	0.0397	2.064	0.1087	-0.4387	0	-1.0725	1.94	1150
Tpz	-2905.44	1.42	0.1000	5.339	0.3877	-0.7120	-857.2	-3.7442	4.04	1315
Dia	-999.30	0.52	0.0353	1.776	0.1451	0.8709	584.4	-1.7411	7.97	2300
H <sub>2</sub> O	-241.81	0.03	0.1888	0	0.0401	0.8656	487.5	-0.2512	0	0

*Note:* Mineral phases: Gr = grossular, Law = lawsonite, Zo = zoisite, Ky = kyanite, Stv = stishovite, Q =  $\alpha$  quartz, Bq =  $\beta$  quartz, Coe = coesite, Tpz = hydroxy-topaz, Dia = diaspore, and H<sub>2</sub>O = water. Symbols as in Holland and Powell (1990) except  $a_0$  and  $K_{298}$ . Values for  $a_0$  should be multiplied by 10<sup>-5</sup>. The thermal expansions,  $\alpha_1$ , and compressibilities,  $K_7$ , at high temperature are expressed as  $\alpha_7 = a_0 - 10a_0 T^{-\alpha}$  and  $K_7 = K_{298} - 3a_0 K_{298}$  (T - 298), respectively (see text). The compressibility of diaspore is from Xu et al. (1994). The data for topaz were derived from the experiments of Wunder et al. (1993) using estimates of heat capacities and entropy [see Holland and Powell (1990) and Holland (1989) for methods]. Stishovite data were taken from Robie et al. (1979) and fitted to the experiments of Yagi and Akimoto (1976). High-pressure data for water were taken from the CORK equations (Holland and Powell 1991). Remaining data from Holland and Powell (1990), updated. Units are kilojoules, kelvins, and kilobars.



FIGURE 6. (a) Experimental brackets and calculated curves, using thermodynamic data from Table 4, for lawsonite breakdown. All pressures were corrected by -5% from the original experimental values to allow for a small friction effect in 0.5 in. piston-cylinder pressure cells. The coe = q curve was calculated from experiments of Bose and Ganguly (1995). (b) Experimental brackets and calculated curves for lawsonite + diaspore and lawsonite + corundum breakdown. H<sub>2</sub>O data were calculated using the CORK (Holland and Powell 1991) equations and the volume data of Rice and Walsh (1957). Abbreviations: law = lawsonite, zo = zoisite, ky = kyanite, dia = diaspore, cor = corundum, coe = coesite, and q = quartz.

piston-cylinder experiments [changing the correction from -5 to -10% would bring good agreement as in the diaspore-corundum case mentioned in Pawley et al. (1996)] or it may be the thermodynamic properties of H<sub>2</sub>O above 3 GPa. The calculated curves for the breakdown of lawsonite + diaspore are in quite good agreement (Fig. 6b) with the experiments of Schmidt and Poli (1994) at the comparatively low pressures of these experiments (<3.5



**FIGURE 7.** Experimental brackets and calculated curves for lawsonite breakdown at extreme pressures. Data used are described in the caption to Figure 6. Small squares = brackets of Schmidt and Poli (1994) on law =  $zo + ky + coe + H_2O$ ; circles = brackets of Schmidt and Poli (1994) on law =  $gr + ky + coe + H_2O$ ; large rectangles = experiments of Pawley (1994) (open = lawsonite growth, solid = gr + ky or tpz + stv or coe growth, hatch = law + gr + ky + coe). See text for discussion. Curve 1 is calculated using the high-pressure  $H_2O$  data of Brodholt and Wood (1993, 1994) and curve 2 using the data of Rice and Walsh (1957).

GPa). With these slight discrepancies appearing at higher pressures, we now consider the very-high-pressure experiments of Pawley (1994) and Schmidt and Poli (1994).

The calculated temperatures of dehydration reactions at very high pressures are rather sensitive to the volumes and fugacities assumed for water, and we show results (Fig. 7) for calculations based on extrapolations to 20 GPa using the Hugoniot-curve shock data of Rice and Walsh (1957) and the equation proposed by Brodholt and Wood (1993) from their molecular dynamics simulations. The Rice and Walsh measurements suggest smaller molar volumes at high pressures than the simulations of Brodholt and Wood, but the latter authors pointed out that (1) the shock temperatures are not well known, (2) their simulations are in good agreement with the PVT measurements of Burnham et al. (1969) at lower pressures, and (3) the simulations are in good agreement with the experimental fluid-inclusion volumes measured by Brodholt and Wood (1994). From Figure 7, the more reasonable agreement up to 8 GPa with the highpressure phase-equilibrium experiments of Pawley (1994) and Schmidt and Poli (1994) also suggests that the Brodholt and Wood simulations may lead to the more reliable estimates of the very-high-pressure properties of water. We refined the CORK equations of Holland and Powell (1991) so that a single expression can be used from 0 to 20 GPa, encompassing the low-pressure *PVT* data to 0.8 GPa and the high-pressure simulation results of Brodholt and Wood in the range 1–20 GPa. Only the virial-like terms in the CORK equation needed to be adjusted to fit all the data to within 0.6% average absolute deviation in volume, giving  $V = V_{\text{MRK}} + c_1(P - P^0) + c_2(P - P^0)^{1/2} + c_3(P - P^0)^{1/4}$ , with  $c_1 = 2.1111 \times 10^{-3} - 5.20907 \times 10^{-7}T$ ,  $c_2 = -9.17623 \times 10^{-2} + 1.12719 \times 10^{-5}T$ , and  $c_3 = 7.40395 \times 10^{-2}$ , where  $P^0 = 1.0$  kbar (0.1 GPa) and  $V_{\text{MRK}}$  is the molar volume from the Modified Redlich-Kwong equation calculated using Equations 3 and 6 of Holland and Powell (1991). All units for this EOS are in kilobars, kelvins, and kilojoules per mole.

Although the thermodynamic data for solids and water are in excellent agreement with the lawsonite breakdown reactions below 3 GPa shown in Figures 6a and 6b (as well as with several others, not shown, involving anorthite, pyrophyllite, and margarite), there are two serious discrepancies, which we cannot presently resolve, between calculated and experimental dehydration reactions at higher pressures. First, the calculated position of the reaction zoisite = grossular + kyanite + coesite +  $H_2O$  lies just above 4 GPa (Fig. 7), with small uncertainties ( $\pm 0.1$  GPa), in contrast with the experiments of Schmidt and Poli (1994), which locate it at  $\sim 6.7$  GPa. There may have been some overstepping of pressure in the latter experiments, as the critical experiments defining the upper pressure limit were (1) unreversed, containing lawsonite and not grossular in the starting material [the lma experiments in Table 1 of Schmidt and Poli (1994)], and (2) were of very short duration (between 5 and 57 min). Additional experiments of longer duration, using reversal mixes, are needed to confirm the position of the zoisite breakdown reaction. However, it is unlikely that they would result in a depression of the reaction of more than a few hundred megapascals. The discrepancy between calculated and experimental reactions is thus very disconcerting because extremely large changes in the free energy of any one phase involved in the reaction would be required to bring the calculated and experimental pressures into agreement. By way of example, the Gibbs energy of zoisite would have to be reduced by over 20 kJ/mol to stabilize it from 4.2 to 6.7 GPa at 1000 °C. To obtain this from one of the first-order properties (S, V) would require changing the standard-state entropy by  $+15.7 \text{ J/(mol \cdot K)}$  or the molar volume by nearly  $-3 \text{ cm}^3/\text{mol}$ . Alternatively, the same effect could be produced by lowering the activity of pure zoisite by an order of magnitude from a value of 1.0 to  $\sim 0.1$ . Appealing to one of the second-order parameters, such as heat capacity, thermal expansion, or compressibility, would require totally unbelievable and unrealistic changes in these properties. Because of the stoichiometry of the reaction, even more drastic changes would be required of the Gibbs energies of  $H_2O$  (+60 kJ) or grossular (+30 kJ) to force the change in calculated pressure from 4.2 to 6.7 GPa. We do not believe that a case can seriously be made for any such extreme alterations in the thermodynamic properties of zoisite, grossular, kyanite, coesite, or  $H_2O$  because the many equilibria in which they are involved in the range 300–1400 °C and 0–3 GPa are consistent with, and impose extremely tight constraints upon, their thermodynamic properties.

The second major discrepancy between calculated and experimental reactions concerns the breakdown of lawsonite at extreme pressures (Fig. 7). At pressures up to 8 GPa there is reasonable agreement between the calculated position of the reaction lawsonite = grossular + kyanite + coesite +H<sub>2</sub>O and the experimental results of Pawley (1994) and Schmidt and Poli (1994). However, although the experimental results suggest that the slope of the lawsonite breakdown reaction decreases gradually above 9 GPa, the calculations suggest a much sharper backbend. At present we can offer no solution to this dilemma, except to note the strongest remaining possibilities: (1) the properties of  $H_2O$ might change in hitherto unexpected ways at very high pressures, (2) the  $H_2O$  content of hydrous phases such as zoisite or lawsonite might change dramatically at high pressures, or (3) possible experimental problems, in the use of multianvil devices that require further investigation. Nevertheless, apart from the case of the very-high-pressure experiments, the thermodynamic calculations based on our new thermal expansions and compressibilities are in good mutual agreement with experimental observations.

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