

NOV 13 1974

High-Pressure Transformations in CaSiO_3 *

Eric Essene

Department of Geology and Mineralogy, The University of Michigan,
Ann Arbor, Michigan 48104

Received March 25, 1974

Abstract. Wollastonite transforms to a triclinic high-pressure polymorph, wollastonite II, at pressures > 25 kb. The equilibrium boundary [$P(\text{bars}) = -4.7 \times T(^{\circ}\text{C}) + 32.810$] has a rather flat negative $P-T$ slope. The rapid reactivity of the transition over at least a 1000°C range, its insensitivity to T and its location around 30 kb are indicative of its potential for a pressure calibration curve at high T .

Wollastonite reacts to form pseudowollastonite at 1125°C and 1 atm, and pseudowollastonite congruently melts at 1544°C (Osborn and Schairer, 1941). Kushiro (1964) followed these reactions to 25 kb finding steep positive slopes on a $P-T$ diagram.

Wollastonite itself has several closely related monoclinic and triclinic polytypes (Peacock, 1935; Trojer, 1968; Wenk, 1969; Ito *et al.*, 1969; Jefferson and Brown, 1973) which have very small differences in volumes. These polytypes are sensitive to shear (Wenk, 1969; Coe, 1970) and are insensitive to $P-T$ of formation (Wenk, 1969), as is characteristic of non-reconstructive and polytypic reactions where ΔG_r is small. The identification of a specific polytype requires single-crystal X-ray data impossible to obtain on the fine-grained run products of this study, and it is fortunate indeed that the specific polytype is unlikely to petrologically and experimentally significant.

Ringwood and Major (1967) converted wollastonite at high pressures to a new polymorph which was about 5% denser than wollastonite at room $P-T$. They placed the equilibrium at about 30 kb for $T = 900^{\circ}\text{C}$, an excellent estimate in light of these experiments. Trojer (1969) refined the structure of this phase finding "...irregular layers of Ca atoms...interconnected by pairs of Ca atoms and $\text{Si}_3\text{O}_9^{-6}$ rings...located in between the layers". Trojer's data yield a cell volume of 37.59 cc, 6.2% denser than wollastonite. The dense polymorph is here designated wollastonite II. Wollastonite II was synthesized at 27 kb and 1600°C by the writer and W. Hibberson while preparing a probe standard of CaSiO_3 . Reversals were then sought on the reactions wollastonite I = wollastonite II and wollastonite I = pseudowollastonite.

The experiments were conducted on the 5/8" Australian National University piston-cylinder apparatus modified from the design by Boyd. Talc/BN cells were used except at 600°C where the pressure medium was entirely talc. Pressures were measured on an advancing ram and were corrected uniformly by -10% (Green *et al.*, 1966). Temperatures were measured with a Pt/Pt-10% Rh thermocouple and are nominal values. Because of the exceptionally

* Contribution No. 317 from the Mineralogical Laboratory, Department of Geology and Mineralogy, The University of Michigan.

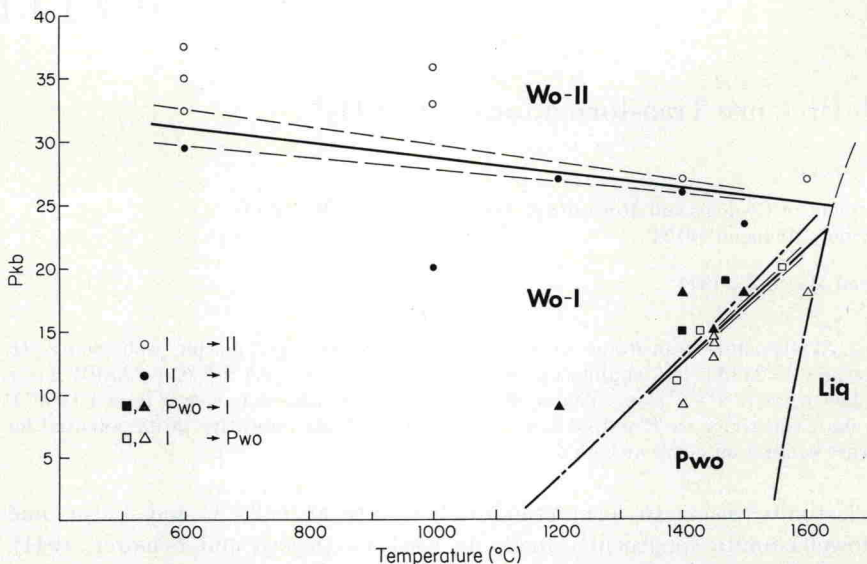


Fig. 1. P - T phase diagram for CaSiO_3 . Inferred equilibria are shown by solid lines and error limits are dashed. These experiments are represented by open and closed circles and triangles. Kushiro's inferred equilibria are shown by dash-dot lines and his reversals by open and closed squares. *Wo-I* wollastonite I; *Wo-II* wollastonite II; *Pwo* pseudowollastonite; *Liq* liquid

flat slope of the wollastonite I—wollastonite II reaction, systematic errors of even 50°C yield only 0.24 kb error in the location of this curve. Run times varied from two days at 600°C , to 8 hrs at 1200°C , to hrs at 1600°C . Starting materials for the runs plotted on the figure were 50/50 mixtures of wollastonite I—wollastonite II and wollastonite I—pseudowollastonite for each transformation. The wollastonite and pseudowollastonite used were purchased from Tem-Pres and had about a 10–30 micron grain size. The wollastonite II strating material was synthesized from Tem-Pres wollastonite and had about the same grain size. 5% excess water was added for the 600°C runs, the 900°C runs were slightly dampened and higher T runs were dry. The runs proceeded to at least 90% of completion as determined by both optical and X-ray examination of run products.

The wollastonite-pseudowollastonite equilibrium obtained in this study agrees well with reversals given by Kushiro (1964) (see Fig. 1). The experimental slope of this transition may be combined with S_T data (Robie and Waldbaum, 1968) to calculate ΔV_T from the relation $\Delta S/\Delta V = dP/dT$. Since expansion data are available for pseudowollastonite (Skinner, 1966), one may calculate the volume (40.73 ± 0.25 cc) of wollastonite at 1125°C , and its average thermal expansion between 25° and 1125°C ($+2.0 \pm 0.7\%$). This agrees well with extrapolations of thermal expansion data on wollastonite—ferrosilite solid solutions (Skinner, 1966), which give a 2.3% expansion at 1125°C for pure wollastonite.

Wollastonite I reacts to form wollastonite II above 25 kb (see Fig. 1). The equation of the boundary is $P = -4.7 \text{ bars}/^\circ\text{C} \times T^\circ\text{C} + 32,810 \text{ bars}$. The slope, $-4.7 \text{ bars}/^\circ\text{C}$, is rather flat and negative. Combination of the experimental data with ΔV_{298}° allows estimation of ΔS_r if the effect of P and T on ΔV is neglected. By this calculation, wollastonite II has an entropy of 0.3 e.u. greater than wol-