Reprinted from the Journal of The American Ceramic Society, Vol. 58, No. 5-6 May-June, 1975 Copyright 1975 by The American Ceramic Society

Differential Thermal Analysis Investigation of the High-Low Cristobalite Inversion Under Hydrostatic Pressure to 7 Kbar

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The high-low inversion in polycrystalline cristobalite, synthesized from quartz at 1500°C, was investigated by differential thermal analysis to 7 kbar in hydrostatic apparatus. A region of anomalous curvature $(d^2T/dp^2 > 0)$ exists to ≈ 1 kbar; at higher pressure, the low-high and high-low transition temperatures vary linearly with pressure, with slopes of \approx 51.1 and 53.6 deg kbar⁻¹, respectively. Extrapolated 1-bar intercepts are $\approx 232^\circ$ and $\approx 209^\circ C,$ respectively. It is concluded that high cristobalite is less compressible than low cristobalite near the inversion. The hysteresis between the high→low and low→high transition temperatures decreases with increasing pressure.

I. Introduction

HE high-low inversion in cristobalite has been the subject of several quite careful investigations. Unfortunately, the lack of sufficiently large single crystals of cristobalite means that most current knowledge is based on experiments with polycrystalline material (also twinned in unknown degree), and the extent to which the various thermophysical properties are thus altered is essentially unknown. For this first-order transition, various data are available for the entropy of the transition and the volume change, so that the variation in transition temperature with pressure is presumably

predictable. A few high pressure data are already recorded^{1,2} for the transition temperature below 1 kbar. The present investigation was undertaken to obtain more extensive, accurate data on the variation with hydrostatic pressure of transition temperatures-both on heating and cooling, since appreciable hysteresis has been universally observed. Also, determination of the variation in hysteresis along the phase boundary should constrain the microscopic theory for this transition.

II. Experimental Procedure and Results

The cristobalite used had been synthesized from quartz heated 4 h at 1500°C in a Pt crucible in air.

In the first run, ≈ 12 mg of cristobalite was loaded into a Pt tube (OD 1.59 mm, wall thickness 0.13 mm) sealed at one end. For the second run, the cristobalite was mixed with an equal weight of fired BN, and ≈ 5 mg of this mixture was loaded into a similar capsule. Transitions were detected using DTA. In all runs, the measuring thermocouple, of Chromel and Alumel wires 0.13 mm in diameter, was lashed against the sample capsule with Chromel wire 0.13 mm in diameter; thermal contact was enhanced by use of silver conductive paint. The reference junction was always within $\leq 2 \text{ mm}$ of the

Received August 14, 1974; revised copy received December 27, 1974. *Now on leave at National Physical Research Laboratory, Pretoria, Republic of South Africa.

measuring junction. The arrangement of sample capsule, thermocouples, and hydrostatic pressure apparatus using Ar has been described previously.^{3,4}

Temperature was varied at ≈ 0.3 to 1.2 deg s^{-1} and the maximum temperature difference between measuring and reference thermocouples was usually $< 1^{\circ}$. Sensitivities in recording temperature and differential temperature were 0.2 mV cm^{-1} and as much as 8 $\mu \text{ V cm}^{-1}$, respectively. The temperature corresponding to the peak of the differential signal was taken as the transition temperature. Walker *et al.*⁵ discussed the preferred identification of peak temperature with inversion temperatures. On heating and on cooling, respectively, the observed transition temperatures usually agreed to within 1°. Temperatures for the onset of the transition were obtained with less precision than peak temperatures.

In the first run, the onset temperatures were $\approx 2^{\circ}$ to 4° lower on heating and higher on cooling than the peak temperatures at the lower pressures and $\approx 1^{\circ}$ to 2° different at the higher pressures. In the second run, there was $\approx 4^{\circ}$ to 6° difference throughout. Near the end of the second run, the transitions were observed near 3.0 kbar with heating/cooling rates varying from ≈ 0.2 to 2.7 deg s⁻¹; no dependence of transition temperature on rate was established within the experimental precision. From previous in situ calibrations^{4,6} of the same reels of thermocouple wire, accuracy in temperature is believed to be within 1° to 2°. Effects of pressure on thermocouple emf were considered small and no corrections were made.

Two or more cycles were made at each pressure and data were obtained on both increasing and decreasing pressures. Pressures were read to ± 5 bar on a 7-kbar pressure gage* and to $\pm 1/_2$ bar on a similar gage of 1-kbar range. In the first run, the 7-kbar gage was used exclusively; in the second run, the 1-kbar gage was also used, simultaneously, in its range. The 1-kbar gage has recently been calibrated to within 1 bar.

Data for the transition temperatures obtained in both runs, on heating and on cooling and on increasing and decreasing pressure, are shown in Fig. 1. Beyond the initial 0.8 to 1.0 kbar, the low \rightarrow high transition data fall on a line of slope $\approx 51._1$ deg kbar⁻¹, with 1-bar intercept $\approx 232^{\circ}$ C. Beyond the initial ≈ 1.5 kbar, the high \rightarrow low transition data fall on a line of slope $\approx 53._6$ deg kbar⁻¹, with 1-bar intercept $\approx 209^{\circ}$ C. Below these indicated limits to the initial pressure ranges, both trajectories show anomalous curvature $(d^2T/dp^2 > 0)$ with 1-bar intercepts $\approx 249^{\circ}$ C on heating and $\approx 225^{\circ}$ C on cooling. Agreement of the results of the two runs (Fig. 1) is within $\pm 1^{\circ}$.

Some of the original material was heated ≈ 24 h at ≈ 1500 °C in a covered Pt crucible in air. This annealed material, on examination in a differential scanning calorimeter, showed the same transition temperatures on heating and on cooling as did the original material and hence was not run at high pressures.

III. Discussion

Two previous studies of the high-low cristobalite inversion at high pressures have been found. Majumdar et al.¹ cite slopes of 5.0×10^{-2} deg bar⁻¹ for two types of cristobalite, one with a l-bar transition temperature of 268°C (text of Ref. 1) or 267°C (Table I of Ref. 1) and the other with a l-bar transition temperature of 242°C (text of Ref. 1) or 225°C (Table I in Ref. 1). Majumdar, in his thesis (Fig. 7 and Table II in Ref. 7), gives data for the inversion in "cristobalite (ordered)" on heating as: $p(\pm 2 \text{ atm}), T(\pm 1^{\circ}\text{C})=1$, 267°; 340, 285°; 580, 302°; 1000, 317°. These data correspond to the "maximum rate of heat absorption" (pp. 21, 22 of Ref. 7), a quantity whose temperature lies between the onset and the peak of the differential signal. In Fig. 7 of Ref. 7, a linear trajectory for "cristobalite (disordered)" is drawn from ≈ 234 °C at zero pressure to ≈ 283 °C at 1 katm; this trajectory is attributed to Roy and Hulbe, a reference which the present writers were unable to find. A linear regression fit of Majumdar's tabulated data7 yields a slope of 51.6 deg kbar⁻¹

The DTA investigation of Berger *et al.*² to ≈ 0.45 kbar was more extensive. They found a region of anomalous curvature below ≈ 0.1 kbar, whereas, at higher pressures, both the high \rightarrow low and



Fig. 1. Temperatures for low→high cristobalite inversion (upper set of points, obtained on heating) and high→low cristobalite inversion (lower set of points, obtained on cooling); data from first run (*circles*) and second run (*triangles*) were obtained on both increasing and decreasing pressures.

low \rightarrow high inversion trajectories had approximately constant slope of \approx 50 deg kbar⁻¹; although the data scatter, approximately constant hysteresis is indicated.

Various values have been obtained for the discontinuities in volume and in entropy for the transition. Because sufficiently large single crystals are not available for dilatometric measurements, X-ray diffraction data seem preferable for estimating the variation of volume with temperature. Extensive X-ray diffraction investigations have been made by Johnson and Andrews,8 using a powder camera and samples showing the inversion near $\approx 218^{\circ}$ C, and by Berger et al.⁹ using a diffractometer and samples inverting at $\approx 265^{\circ}$ to 270°C. Within their respective estimated accuracies, the volume of low cristobalite varies linearly with temperature up to the inversion temperature; for high cristobalite, similar results are obtained for at least some hundred degrees above the inversion. It is agreed^{8,9} that the thermal expansion coefficient is much greater below the inversion than above, but results from Ref. 9 are consistently lower than those from Ref. 8 for thermal expansion coefficients and for volumes at the same temperature. Majumdar's data7 tabulated for the "ordered" sample (there is a misprint for the lattice parameters given for 230°C) are consistent with data from Ref. 8 for low cristobalite (as extrapolated to higher temperatures, since there are differences between the inversion temperatures) but are higher both for volume and thermal expansion coefficient than the data from Refs. 8 and 9 for high cristobalite. Majumdar et al.¹ quote a volume discontinuity of 1.03 cm3 mol-1 both for the

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