Dynamic compression of SiO₂: A new interpretation

Joseph A. Akins and Thomas J. Ahrens
Lindhurst Laboratory of Experimental Geophysics, Seismological Laboratory 252-21, California Institute of Technology, Pasadena, CA, USA

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[1] In light of recent discoveries of post-stishovite phases of SiO₂, with the CaCl₂ and α-PbO₂ structures, we have reassigned the regimes along the Hugonioti for initial quartz, coesite, cristobalite, porous coesite, and fused silica. Calculated Hugoniots for fused silica, cristobalite and porous coesite indicate transition to stishovite, and then melt. Hugoniots for crystal quartz and coesite indicate that transition occurs to stishovite, then the CaCl₂ structure and finally to melt.


1. Introduction

[2] The recent discoveries that pure silica transforms to the CaCl₂ (ρo = 4.291 g/cm³) and α-PbO₂ (ρo = 4.334 g/cm³) structures, slightly more dense than stishovite (ρo = 4.287 g/cm³), have led to a reexamination of the dynamic compression of SiO₂. Although the first observation of the CaCl₂ structure indicated it was stable above ~100 GPa [Tsuchida and Yagi, 1989] later theoretical [Cohen, 1991] and experimental [Andraut et al., 1998; Dubrovinsky et al., 1997; Kingma et al., 1995] studies have constrained the onset of the stishovite to CaCl₂ transition to ~50 GPa at 298 K. The α-PbO₂ structure is found to be stable above ~80 GPa, based on experiments and theoretical calculations [Dubrovinsky et al., 2001; Dubrovinsky et al., 2001; Dubrovinsky et al., 1997, 2001; Teter et al., 1998]. The α-PbO₂ structure has also been observed in the Shergotty meteorite, a Martian basalt thought to have experienced multiple shock events [El Goresy et al., 2000].

[3] In addition to these high-pressure silica phases, with oxygen in 6-fold coordination, transition to an 8-fold coordination structure is predicted near 200 GPa [Dubrovinsky et al., 1997; Teter et al., 1998]. Although it is unclear whether substantial quantities of a phase of SiO₂ exists in the Earth’s mantle, the equation of state of the high-pressure SiO₂ phases play a crucial role in mantle-core chemical reactions which sequester iron from mantle silicates into the core [Luo et al., 2002].

[4] Crystal and fused silica were the first silicates studied by dynamic compression methods [Adadurov et al., 1962; Wackerle, 1962]. Thereafter McQueen et al. [1963] demonstrated that the Hugoniots of crystal (ρo = 2.65 g/cm³) and fused silica (ρo = 2.2 g/cm³), above 40 GPa, correspond to the properties of the rutile structure of SiO₂ [Stishov and Popova, 1961], which was later found in shocked quartz-bearing rocks of Meteor Crater [Chao et al., 1962]. Subsequently, small amounts of stishovite were recovered from peak pressures in the 15–28 GPa range [DeCarli and Milton, 1965].

[5] Additional studies of quartz and fused silica, as well as cristobalite (ρo = 2.13 g/cm³), porous coesite (ρo = 2.4 g/cm³), coesite (ρo = 2.92 g/cm³), and stishovite (ρo = 4.31 g/cm³) have been conducted by Fowles [1967], Trunin et al. [1971], Podurets et al. [1990], Borshchevskii et al. [1998], Lyzenga et al. [1983], Zhugin et al. [1999] and Luo et al. [2002]. With the exception of the more scattered fused silica results, we take these data together and point out that strong evidence exists for the transitions to stishovite, the CaCl₂ structure and melt along the Hugoniots of quartz and coesite, whereas, cristobalite and porous coesite melt directly from the stishovite regime.

2. Dynamic Compression of Crystal Quartz

[6] The outline of our interpretation of the crystal quartz Hugoniot is shown in Figure 1. Theoretical calculations for Hugoniot pressure-density and pressure-temperature for crystal quartz (and other SiO₂ phases) of Figures 1–4 are based on the Mie-Grüneisen offset from the 3rd order Birch-Murnaghan isentropes (e.g. McQueen et al. [1963]). We infer from the phase diagram of Figure 4 and the independent pressure-density and pressure-temperature Hugoniot data for various silica phases, there are the following eight regimes along this Hugoniot: 1) The elastic shock Hugoniot extending to 5.5–15.0 GPa depending on orientation [Fowles, 1967; Wackerle, 1962]. 2) The quartz deformational Hugoniot extending from 15 to 23 GPa [Zhugin et al., 1999].

Figure 1. New interpretation of Hugoniot of crystal SiO₂ (quartz). Data (circles) from Marsh [1980], Fowles [1967], Trunin et al. [1971], Lyzenga et al. [1983], Podurets et al. [1976, 1990], Borshchevskii et al. [1998]. Static compression data for initially stishovite (x’s) from Andraut et al. [1998]. Curves in stishovite, CaCl₂ and melt regime calculated from parameters of Table 1.

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The compression of stishovite and the CaCl₂ structure, observed in the diamond anvil study of Andrault et al. (1998) (and references therein) are plotted below the shock wave data in Figure 1. Notably the diamond anvil data displays a transition at ~54 GPa from stishovite to the CaCl₂ structure in approximate agreement with the deviation of the calculated 298K isotherms of stishovite and the CaCl₂ structure.

3. Coesite, Porous Coesite, and Cristobalite Data

Shock data for coesite (ρ₀ = 2.92 g/cm³) and porous coesite (ρ₀ = 2.4 g/cm³) are plotted in Figure 2a. The theoretical Hugoniot curves are consistent with the stishovite, CaCl₂ and melt EOS parameters of Tables 1 and 2. Coesite and porous coesite are assessed to transform to stishovite above 30 GPa. The single datum for porous coesite at 85 GPa is assumed to correspond to a point in the stishovite-melt regime which achieves a higher density than stishovite. In contrast the single-crystal coesite achieves a state at 62 GPa, consistent with coesite transformed to stishovite, whereas the two-higher pressure states at 85 and 111 GPa appear to lie along the CaCl₂ regime of the coesite Hugoniot; melt is inferred above 137 GPa. The coesite data of Luo et al. [2002] is consistent with this interpretation.

4. Shock Temperatures

Complete shock temperature data for both fused silica and crystal quartz, as well as calculations in the solid and melt phases, are shown in Figure 3 and for the solid only in Figure 4 relative to our newly inferred high-pressure phase diagram of SiO₂. Shock temperatures were calculated using the method of Ahrens et al. [1982]. The calculated shock temperatures of Wacketle [1962] for both fused silica and crystal quartz are in remarkable agreement with Lyzenga et al. [1983] and McQueen and Fritz [1982] data. The fused silica shock temperatures below 55 GPa are in the stishovite stability field, the data between 55 and 65 GPa correspond to superheated stishovite, and above 75 GPa lie along the melt region of the Hugoniot. For crystal quartz the Hugoniot between 70 and 115 GPa is in the CaCl₂ structure rather than

[9] In the case of cristobalite data, Figure 2b, we interpret the point at 46 GPa as still being in the cristobalite-stishovite mixed-phase regime, the datum at 64 GPa as in the stishovite regime and the data at 75 and 86 GPa as in the dense SiO₂ melt regime.

Figure 2. (a) Hugoniot data, initially porous (ρ₀ = 2.4 g/cm³) and polycrystalline (ρ₀ = 2.92 g/cm³) coesite. (b) Hugoniot data, cristobalite (ρ₀ = 2.13 g/cm³).

Figure 3. Hugoniot temperatures, fused quartz (FO) and quartz (QTZ), Lyzenga et al. [1983] (with minor corrections for Al absorption, Boslough [1988]). Melting curve is shown for comparison.

Figure 4. Proposed phase diagram of SiO₂. Only data and calculations for solid phases are shown. Shock temperature calculations are terminated at pressures corresponding to observed, or expected, drops in temperature due to melting. The field of α-PbO₂ is indicated for reference only.
the superheated stishovite phase. Shock temperature states above 115 GPa are completely melted. The interpretation of super-heated solid followed by a drop in temperature due to melting is also supported by the post-shock temperature analysis of Boslough [1988]. Therefore the drop in shock temperature at 70–75 GPa for fused silica corresponds to catastrophic homogeneous melting [Lu and Li, 1998] of stishovite, whereas the drop between 115 and 120 GPa for shocked crystal quartz results from catastrophic homogeneous melting of crystal quartz transformed to the CaCl2 structure.

[11] The shock temperature calculations (Figure 4) for cristobalite indicate that it enters the regime of transition to stishovite above 20 GPa and the molten SiO2 stability field at 65 GPa. The pressure-density Hugoniots data clearly demonstrates that like fused silica transformed to stishovite, cristobalite also is superheated along the stishovite branch of its Hugoniot to shock pressures of 65 GPa (see Figure 2b) at which point it enters a stishovite-melt mixed-phase regime. The pressure-density data suggest that at ~70 GPa complete melting has occurred. Therefore we predict future shock temperature experiments will see a drastic decline in shock temperature from ~6000 to ~4000 K at ~70 GPa in cristobalite.

[12] Similarly, porous coesite appears to begin to transform to the stishovite structure above 30 GPa and is completely transformed into the stishovite regime at 55 GPa. Between 55 and 80 GPa it appears to remain in the stishovite structure although above 55 GPa the stishovite is superheated. The pressure-density data (Figure 2a) suggest that melting of the superheated stishovite occurs at ~85 GPa and above ~90 GPa the Hugoniot is inferred to be in the complete melt regime.

[13] We note that coesite also demonstrates the phase transition to stishovite starting at 30 GPa, and is driven into the coesite-stishovite mixed phase regime from 30 to 50 GPa and in the pure stishovite region from 50 to ~62 GPa. Transition from stishovite to the CaCl2 structure appears to occur at ~65 GPa and superheating occurs up to ~7000 K. We infer that at ~137 GPa transition to the melt occurs. Coesite shocked into the CaCl2 structure is superheated and extends metastably into the melt from ~80 to ~137 GPa. Given this new interpretation of the transformation from stishovite to the CaCl2 structure along the quartz and coesite Hugoniots, at ~70 and ~65 GPa, respectively, the Clapeyron slope of the transition is constrained to be ~180 K/GPa compared to 250 K/GPa predicted by Kingma et al. [1995].

[14] Stishovite is too cold to undergo any of the above mentioned solid-solid phase transitions along its Hugoniot [Lu et al., 2002]. We predict that stishovite will melt at ~425 GPa along its principal Hugoniot. To observe the solid-solid transitions, future shock experiments on porous stishovite, stishovite/coesite, or stishovite/paraffin mixtures are needed. For \( \rho_0 = 3.6 \text{ g/cm}^3 \) we predict CaCl2 to \( \alpha\text{-PbO}_2 \) transition will be observed at ~100 GPa and ~3000 K, and melting at ~200 GPa and ~3600 K. For \( \rho_0 = 3.95 \text{ g/cm}^3 \) we predict CaCl2 to \( \alpha\text{-PbO}_2 \) at ~100 GPa and ~1600 K, \( \alpha\text{-PbO}_2 \), 8-fold coordination solid at ~200 GPa and ~350 K and melting at ~315 GPa and ~6500 K.

5. Hugoniot Sound Velocities

[15] The measurements of sound velocity behind the shock front in dynamically compressed crystal quartz (\( \rho_0 = 2.65 \text{ g/cm}^3 \)), quartzite (\( \rho_0 = 2.65 \text{ g/cm}^3 \)) and novaculite (\( \rho_0 = 2.64 \text{ g/cm}^3 \)), shown in Figure 5, support the above interpretation of the combination of pressure-density and shock temperatures. Here \( V_p \) is the longitudinal (non-zero rigidity) elastic velocity of the solid, whereas \( V_p \) is the bulk plastic velocity of the deformed solid and above ~107 GPa, the velocity of shock-induced melt. The rarification velocities for elastic and deformational unloading demonstrate that the stishovite compressional wave velocity increases to about 12.9 km/s, at 35 GPa. At this pressure mode softening (decreasing shear modulus), begins to occur [Cohen, 1991]. With increasing pressure the sound velocity decreases in anticipation of

Table 2. Transition Energies

<table>
<thead>
<tr>
<th>Phase</th>
<th>Phase Initial Density, ( \rho_o ) (g/cm³)</th>
<th>Transition Energy (MJ/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystobalite</td>
<td>( \alpha\text{-quartz} )</td>
<td>0.056</td>
</tr>
<tr>
<td>Fused quartz</td>
<td>( \alpha\text{-quartz} )</td>
<td>0.70</td>
</tr>
<tr>
<td>( \alpha\text{-quartz} )</td>
<td>Stishovite</td>
<td>0.86</td>
</tr>
<tr>
<td>Coesite</td>
<td>Stishovite</td>
<td>0.82</td>
</tr>
<tr>
<td>Stishovite</td>
<td>CaCl2</td>
<td>0.5</td>
</tr>
<tr>
<td>( \alpha\text{-quartz} )</td>
<td>Melt</td>
<td>2.5</td>
</tr>
</tbody>
</table>

\( \Delta E \geq \Delta H \) at ambient conditions.

References:
[1982], [2000], [1993], [1986], [1998], [2000].

Figure 5. Sound speed measurements in SiO2 phases with \( \rho_o = 2.65 \text{ g/cm}^3 \). \( V_p \) versus pressure curve was calculated by Carpenter et al. [2000]. Quartz data of McQueen [1992] below 90 GPa not plotted.
the ~65 GPa transition to the CaCl₂ structure. The compressional wave velocity in the CaCl₂ structure (metastably shocked into the melt region above 80 GPa) then rises again until at 115 GPa catastrophic melting occurs. At this pressure the longitudinal velocity becomes equal to the bulk velocity, as is expected upon melting. In contrast, the bulk sound velocity steadily increases at the onset of the stishovite to CaCl₂ structure transition and continues to increase with pressure. Quite interestingly the bulk sound velocities data appear to change little upon catastrophic melting of CaCl₂ structure at ~115 GPa. This implies the bulk modulus of solid SiO₂ in the CaCl₂ structure is essentially equal to that of the melt. If other silicates behave similarly, then in contrast to the conclusion of Karato and Karki [2001] the presence of a small fraction of melt in the lowermost mantle may remain a viable explanation for the high values of (∂ ln Vp/∂ ln ρ) as discussed by Duffy and Ahrens [1992].

6. Conclusions

[17] The shock wave pressure-density data for cristobalite and porous coesite are consistent with these phases transforming from their initial state to the stishovite structure upon shock compression. Above 40–50 GPa, transformation to stishovite is complete and shock-induced transformation to the melt begins at 65 GPa in cristobalite, and ~85 GPa in porous coesite. Based on shock temperature measurements in fused and crystal silica, we infer that in cristobalite, fused silica and porous coesite the solid is superheated in the stishovite structure starting at 45, 55 and 60 GPa, respectively. In contrast, shock induced transformation to stishovite from initially crystal quartz and coesite begins at 22 and 30 GPa, respectively. Complete transformation to stishovite is completed at 40–50 GPa along their Hugonioti and transformation to the CaCl₂ structure begins at 65–75 GPa. The CaCl₂ phase for crystal quartz and coesite is metastably superheated above the 80 GPa, 4000 K melting line of the CaCl₂ structure. Catastrophic melting occurs at ~115 GPa along the quartz Hugoniot, and is expected to occur at ~137 GPa along the coesite Hugoniot shocked into the CaCl₂ structure. At higher pressures the Hugoniot of crystal quartz and coesite lie in the melt regime.

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References


J. A. Akins and T. J. Ahrens, Lindhurst Laboratory of Experimental Geophysics, Seismological Laboratory 252-21, California Institute of Technology, Pasadena, CA 91125, USA. (jaja@caltech.edu)