Abstract. Hugoniot equation of state and release adiabat results are presented for c cut crystals of aragonite, the high-pressure polymorph of calcite, shocked to pressures of up to 40 GPa. A Hugoniot elastic limit is observed at 2.5±0.8 GPa and is similar to that of calcite, which, depending on orientation, ranges from 1.5 to 2.5 GPa. A phase transition, possibly displacive, occurs between 5.5 and 7.6 GPa. Above shock pressures of ~10 GPa, the aragonite and calcite Hugoniots are nearly coincident, suggesting transformation of both polymorphs to the same phase. Model calculations, attempting to characterize the high pressure CaCO_3 phase are presented. Aragonite release adiabats centered at pressures between 9 and 14 GPa indicate that states with apparent zero-pressure densities from 2.9 to 3.2 g/cm^3 are achieved upon decompression from progressively greater shock pressures. Observed unloading paths from shock pressures above 17 GPa are significantly and consistently shallower (in a density-pressure plane) than those from lower pressures, and zero-pressure densities up to 20% below that of the initial aragonite density are achieved upon unloading; these features suggest that vaporization is occurring upon unloading. According to theoretical shock temperature and entropy calculations, however, the minimum shock pressure for vaporization upon release for aragonite is 55 GPa (and 33 GPa for calcite), significantly higher than the observed value.

Introduction

Compressive behavior in the CaCO_3 system has generated interest since Bridgman's [1939] discovery of two low-pressure phase transitions in calcite. While subsequent static and dynamic compression experiments have extended the calcite pressure-density field to 94 GPa [Huang and Wyllie, 1976; Ahrens et al., 1966; Adadurov et al., 1961; Kalashnikov et al., 1973] only limited compressibility data exist on its high-pressure polymorph, aragonite. The static experimental results on aragonite by Bayuk et al. [1974] and Jamieson [1957] are limited to pressure of 1.5 and 2.4 GPa, respectively; until the present investigation, no shock compression aragonite data have been available.

The purpose of this investigation is two-fold: first, to determine the shock compression curve of aragonite, identify any possible phase transitions, and speculate as to their nature and to compare it to the calcite Hugoniot, and second, to measure release isentropes for aragonite and thus gain insight into the behavior upon release from shock pressure of carbonates in general.

The investigation of shock compression and release behavior of carbonates has direct application to descriptions of cratering mechanics and impact metamorphism of rocks containing carbonate minerals. Approximately 30% of the known or probable terrestrial meteorite impact craters occur at least partially in carbonate rocks [Grieve and Robertson, 1979]. Also, carbonate rocks may compromise a significant portion of the craters of the Martian surface [Booth and Kieffer, 1978], and have been suggested as being the primary phases which determine the partial pressure of CO_2 on the surface of Venus [Lewis, 1970]. In addition, a large class of craters for which energies are approximately known, the Pacific Test Site nuclear craters, were formed in calcite- and aragonite-containing coralline limestone. The shock compression and release data presented in this paper will be useful in understanding impact and explosion processes in these various carbonate terrains. Also, the release results should provide a firmer basis for estimating the contribution of shock-produced CO_2 to an evolving atmosphere.

Experimental Details and Data Reduction

The aragonite samples used in this study were pseudo-hexagonal twinned crystals collected by A. Jones near Minglanilla, Spain. The doubly terminated prisms, formed by an intergrowth of three individuals twinned on the (110) plane, display an approximately hexagonal cross section with small reentrant angles often appearing as prism faces at the twin boundaries. Guinier camera X-ray analysis verified the identity of the crystals and microprobe analysis indicated a relatively pure CaCO_3 composition (see Table 1).

Crystals with a minimum of cracking and discoloration (presumably due to iron and/or manganese), and measuring 15 to 20 mm in width were cut into 3- to 4-mm-thick sections and lapped to within uniform thickness tolerances of 0.1-0.3% (i.e., ±3-10 μm). Archimedean densities were determined using reagent grade toluene as a solvent and the temperature corrections of Berman [1939]. Individual density measurements varied by ±0.001 g/cm^3 while the average density for all 12 samples was 2.928 g/cm^3 with a standard deviation of 0.003 g/cm^3. The close agreement between this measured average density and the X-ray density of 2.930 g/cm^3 is probably due to a fortuitous cancellation of the effects of impurities and cracks. Owing to the irregular shape of the crystal sections used in the experiment, bulk density determinations were not performed each time. To obtain an approximate measure of sample porosity, a rectangle was cut from an aragonite crystal and lapped to within dimensional tolerances of 0.1%; the porosity determined from this measurement was 1.7%.

The samples were then mounted on either 2024 Al or Ta driver plates with the c axes, as determined...
Experimental results for aragonite Hugoniot and release states are listed in Table 2 and presented in particle velocity shock velocity ($U_p$, $U_s$) and density-pressure ($p$, $P$) diagrams in Figures 2 and 3, respectively.

In six of the experimental shot records, a two-wave structure, such as that displayed in Figure 1b, was observed. Three of these double-shock experiments, shots 383, 402, and 498, are interpreted as representing an elastic to plastic shock transition. (It is also possible that shot 498 represents a phase transition occurring at ~3.3 GPa. However, due to the lack of any corroborative evidence, it will be grouped, for discussion purposes, with the two lowest pressure experiments.) Comparing the shock velocity of the first wave to the longitudinal wave velocity along the c axis of aragonite identifies this first wave as elastic. The only single crystal data for aragonite are those of Voigt [1910]. Bayuk et al. [1974] report crystalline aragonite results. From the elastic stiffness ($c_{33}$) listed in Voigt's work, a compressional wave velocity along the c axis is computed from the appropriate equation for an orthorhombic crystal:

$$\frac{V_{caxis}}{P} = \frac{V_{caxis}}{P} = 5.39 \text{ km/s}$$ (1)

This agrees well with our recent ultrasonic measurement: $V_{caxis} = 5.39 \pm 0.035 \text{ km/s}$ [Kondo, personal communication, 1981]. The observed shock velocities of the assumed elastic wave, ranging from 5.67 to 6.00 km/s, are, as expected [Davison and Graham, 1979], slightly greater than the ultrasonically measured value. An additional argument for the interpretation of these velocities as elastic comes from the agreement of their corresponding shock pressures, 1.8 to 3.3 GPa, with Hugoniot elastic limit pressures of 1.5 to 2.5 GPa determined for single crystal calcite by Ahrens and Gregson [1964].

As seen in Figure 3, the deformational shock states following the elastic wave, indicated by an asterisk, are considerably displaced from the aragonite Hugoniot, as defined below 8 GPa by six first arrival states both in the elastic and plastic regime (specifically, first arrivals in shots 382, 383, 402, 423, 498, and 499). However, in double-shock experiments where the first wave is in the plastic regime (shots 382, 423, and 499), second arrival shock states (double asterisks in Figure 3) are consistent with both the low pressure Hugoniot, as defined in the last sentence, and the high pressure (above 17 GPa) Hugoniot. Thus, the deformational shock states indicated by asterisks in Figure 3 are anomalous with respect to all other experimental points. The reason for these deviations may lie in some unidentified phenomena associated with the elastic to plastic shock transition.

On a shock velocity-particle velocity plot, data for the first deformational waves, from those double-wave experiments in which both shock states are in the plastic regime (shots 382, 423, and 499), form a cluster at $U_s$ values greater than
those attributed to the elastic wave, and at low $u_p$ values (labeled "phase transitions" in Figure 2). As seen on the pressure-density plot (Figure 3), two of these values form a distinct cusp at ~7.5 GPa, which appears to divide the aragonite Hugoniot into a decidedly steeper (albeit poorly defined) low pressure part and a well-defined initially shallower high pressure part. On the basis of these observations, a phase change in aragonite is tentatively inferred to occur between shock pressures of 5.5 to 7.6 GPa.

As already mentioned, several aspects of the low pressure data remain enigmatic. It would appear from Figure 2 that $U_s-u_p$ values from the second wave arrivals in the six double-wave experiments together outline a poorly defined general trend. Such coincidence, however, would be unexpected in light of the interpretation that three of these points (with the lowest $u_p$ values) correspond to an intermediate shock state (following the elastic state), while the other three represent some higher pressure phase. The six second wave arrival points were fit to provide the linear least squares relationship:

$$U_s = 3.77 + 2.52 u_p, \quad r^2 = 0.81 \tag{2}$$

where 3.77 km/s and 2.52 represent the constants $c_0$ and $s$, respectively, of the $U_s-u_p$ Hugoniot representation, and $r^2$ is the square of the correlation coefficient. The intercept of 3.77 km/s is comparable to the bulk sound velocity of 4 km/s derived from the bulk modulus given by Voigt [1910]. Correcting the data for sample compression or wave interaction effects would move the points in Figure 2 to slightly lower $U_s$ and/or higher $u_p$ values. Due to the minimal effect of such adjustments [see, e.g., Ahrens et al., 1968], the large amount of scatter in our data, and the ambiguity in the constitutive relation of aragonite, the data are reported here without such correction. Further justification for the fit of (2) will be provided by the comparison of aragonite and calcite low pressure data (next section).

For particle velocities above 1 km/s, the $U_s-u_p$ data from a consistent set are described by the linear regression

$$U_s = 5.02 + 1.30 u_p, \quad r^2 = 0.99 \tag{3}$$

(Because of an unclear photographic record, shot 393 results are not included in this fit.) As seen in Figure 2, this latter trend is distinct from that observed at particle velocities below 1 km/s. The shift to a shallower slope, i.e., a smaller $s$ value, above a particle velocity of 1 km/s is similar to the pattern exhibited by many silicate minerals in mixed phase regimes [McQueen et al., 1967]. Further speculation on the higher pressure half of the aragonite Hugoniot and its relation to that of calcite will be presented in the next section.

Partially released states achieved by
TABLE 2. Summary of Hugoniot

<table>
<thead>
<tr>
<th>Shot</th>
<th>Flyer/Driver Material</th>
<th>Projectile Initial Velocity, km/s</th>
<th>Projectile Initial Density, g/cm³</th>
<th>Shock Particle Pressure, GPa</th>
<th>Particle Velocity, km/s</th>
<th>Pressure, GPa</th>
<th>Density, g/cm³</th>
</tr>
</thead>
<tbody>
<tr>
<td>382</td>
<td>2024 Al</td>
<td>1.194 ±0.005</td>
<td>2.925 ±0.001</td>
<td>6.26 ±0.04</td>
<td>0.41 ±0.02</td>
<td>7.45 ±0.15</td>
<td>3.13 ±0.01</td>
</tr>
<tr>
<td>383</td>
<td>2024 Al</td>
<td>0.782 ±0.005</td>
<td>2.929 ±0.001</td>
<td>6.00 ±0.08</td>
<td>0.11 ±0.01</td>
<td>7.45 ±0.15</td>
<td>2.98 ±0.01</td>
</tr>
<tr>
<td>393</td>
<td>Ta</td>
<td>1.376 ±0.005</td>
<td>2.926 ±0.001</td>
<td>6.85 ±0.09</td>
<td>0.11 ±0.01</td>
<td>20.97 ±0.4</td>
<td>3.45 ±0.01</td>
</tr>
<tr>
<td>402</td>
<td>2024 Al</td>
<td>0.934 ±0.005</td>
<td>2.923 ±0.001</td>
<td>5.2 ±0.04</td>
<td>0.03 ±0.14</td>
<td>2.33 ±0.04</td>
<td>2.99 ±0.1</td>
</tr>
<tr>
<td>423</td>
<td>2024 Al</td>
<td>1.059 ±0.005</td>
<td>2.925 ±0.001</td>
<td>6.29 ±0.06</td>
<td>0.30 ±0.11</td>
<td>5.55 ±0.07</td>
<td>3.07 ±0.01</td>
</tr>
<tr>
<td>491</td>
<td>2024 Al</td>
<td>1.998 ±0.001</td>
<td>2.928 ±0.001</td>
<td>6.34 ±0.04</td>
<td>1.02 ±0.01</td>
<td>18.16 ±0.4</td>
<td>3.51 ±0.01</td>
</tr>
<tr>
<td>494</td>
<td>Ta</td>
<td>1.237 ±0.005</td>
<td>2.933 ±0.001</td>
<td>6.25 ±0.04</td>
<td>0.96 ±0.16</td>
<td>17.54 ±0.4</td>
<td>3.46 ±0.01</td>
</tr>
<tr>
<td>496</td>
<td>Ta</td>
<td>2.414 ±0.005</td>
<td>2.928 ±0.001</td>
<td>7.40 ±0.05</td>
<td>1.83 ±0.3</td>
<td>39.74 ±0.4</td>
<td>3.89 ±0.01</td>
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<tr>
<td>497</td>
<td>Ta</td>
<td>1.736 ±0.001</td>
<td>2.932 ±0.001</td>
<td>6.75 ±0.05</td>
<td>1.33 ±0.17</td>
<td>26.32 ±0.4</td>
<td>3.65 ±0.01</td>
</tr>
<tr>
<td>498</td>
<td>2024 Al</td>
<td>0.712 ±0.005</td>
<td>2.930 ±0.001</td>
<td>5.66 ±0.09</td>
<td>0.20 ±0.1</td>
<td>3.26 ±0.04</td>
<td>3.04 ±0.01</td>
</tr>
<tr>
<td>499</td>
<td>2024 Al</td>
<td>1.550 ±0.001</td>
<td>2.930 ±0.001</td>
<td>6.42 ±0.09</td>
<td>0.41 ±0.1</td>
<td>7.64 ±0.11</td>
<td>3.13 ±0.06</td>
</tr>
<tr>
<td>502</td>
<td>Ta</td>
<td>2.063 ±0.001</td>
<td>2.932 ±0.001</td>
<td>7.05 ±0.03</td>
<td>1.38 ±0.18</td>
<td>32.53 ±0.5</td>
<td>3.78 ±0.01</td>
</tr>
</tbody>
</table>

*Final shock state unless otherwise noted.
†Intermediate deformational shock state.
‡Hugoniot elastic limit shock state.

Rarefaction wave propagation into the aragonite samples from the buffer-sample interface together with calculated maximum postshock densities achieved on complete release are plotted in pressure-density space in Figure 4. Two distinct trends are evident. Release paths originating at Hugoniot states at or below ~14 GPa are consistently steeper than the Hugoniot at these pressures, and the corresponding maximum postshock densities suggest density increases upon complete release of up to 10%. (For clarity, release paths for the three lowest pressure Hugoniot states are not shown.) Steep release adiabats, from shock pressures above the suspected onset of a phase transformation, have been observed for a number of silicates, including quartz [Grady et al., 1974], plagioclase feldspar [Ahrens et al., 1969a,b], and olivine [Jackson and Ahrens, 1979], and have been interpreted to indicate retention of the denser high pressure phase during the initial stages of unloading. The same interpretation may be invoked for aragonite released from pressures between 5 and 14 GPa. With one exception, shallow release paths are observed upon unloading from Hugoniot shock pressures above 17 GPa. The one exception, at ~26 GPa, displays a slope approximately parallel to that of the Hugoniot. Partial vaporization upon release might be expected to produce the observed release paths. Whether or not these phenomena can occur under the pressure-temperature states achieved upon unloading will be discussed in the next section. The calculated maximum postshock densities for those high pressure experiments do not form an internally consistent set; i.e., there is no continuous relationship
### Discussion of Aragonite Results and Comparison with Calcite Hugoniot

In this section, aragonite and calcite Hugoniot data are compared and used to draw some general conclusions on the shock compression and release behavior of carbonates.

The aragonite and calcite Hugoniots are plotted together in Figure 5. The single crystal calcite ($\rho = 2.71 \text{ g/cm}^3$) Hugoniot [Ahrens and Gregson, 1964; Ahrens et al., 1966] displays an orientationally dependent HEL between 1.8 and 2.4 GPa and a series of three, or possibly four, cusps, indicative of phase transitions, below ~10 GPa. It is the c cut calcite Hugoniot that is drawn in Figure 5; above approximately 10 GPa, this curve is representative of a and b cut calcite Hugoniot data as well. The authors relate two of these cusps to the calcite I to II and II to III transitions observed in static high pressure experiments [Bridgman, 1939; Singh and Kennedy, 1974]. In a detailed shock compression study of the low pressure phase changes in calcite, Grady et al. [1978] obtained stress-particle velocity profiles for two limestones and a marble which indicated the occurrence of one phase transition between 0.6 and 1.2 GPa and another beginning at 2.4 GPa; their interpretation of the first transition (calcite I...
Phase transitions in aragonite and calcite respectively, undergo a shift to shallower values within 2% of each other. In fact, the slopes of all the carbonate data above \( u_s \approx 1 \) km/s are subparallel. This pattern is typically displayed by data sets for materials of varying porosities [see, e.g., Carter and Marsh, 1980] whereby increasingly porous sample \( u_s - u_p \) fits lie at lower \( u_s \) values. Also, the apparent shift of the slope break toward higher \( u_p \) values with increasing CaCO\(_3\) starting density \( p \) is consistent with the observations made by Jackson and Ahrens [1979] on porous forsterite data. They speculate that increased internal energies at greater material porosities facilitate the onset of phase transitions at lower shock pressures.

Although the data presented in Figure 6 point strongly to attainment of an identical structure at high pressures by calcite and aragonite, they do not shed light on the nature of this transition. Three possibilities are (1) transformation to a high pressure phase at \( \sim 15 \) and \( \sim 18 \) GPa on the calcite and aragonite Hugoniots, respectively, (2) onset of a mixed phase regime at these same pressures, and (3) incipient melting. With regard to the third interpretation, there are, as discussed by McQueen et al. [1967], no features intrinsic to either a \( u_s - u_p \) or \( P - p \) representation of melting along a Hugoniot that would allow unequivocal identification of this phenomenon. The coincidence of a break in the \( u_s - u_p \) slope with the intersection of the melting curve and the Hugoniot to II) as displacive agrees with the static compression results of Merrill and Bassett [1975]. As previously noted, the aragonite Hugoniot has a HEL at 2.5 \( \pm 0.8 \) GPa and a cusp of 6.6 \( \pm 1 \) GPa. Unfortunately, static compression data on aragonite are too limited to aid in confirmation or characterization of the 6.6-GPa transition. Neither the volume compression studies of Bayuk et al. [1974] to 1.5 GPa nor the diamond anvil experiments of Jamieson [1957] to 2.4 GPa revealed any phase changes in aragonite at or below these pressures. Based on analogies with the calcite phase diagram and with the known phase relations of KNO\(_3\), an iso-structural analogue of CaCO\(_3\) [Rapport, 1966], we might speculate that the 6.6 \( \pm 1 \) GPa aragonite transition is displacive in nature, possibly involving rotation of carbonate groups and/or translation of calcium atom planes. Resolution of this issue, however, requires static high pressure experiments.

Above approximately 10 GPa, the Hugoniots of calcite and aragonite are, within the uncertainties in the data, indistinguishable in the pressure-density plane. However, a significant change in the \( u_s - u_p \) plots of both polymorphs is observed at corresponding pressures greater than 10 GPa. Our aragonite \( u_s - u_p \) data, together with that of nonporous and porous polycrystalline calcite, are presented in Figure 6; fits for the various data sets are summarized in Table 3. Note first of all the similarity in the slope of the fits to our aragonite and to the Adadurov et al. [1961] marble data at low (below \( \sim 1 \) km/s) particle velocities, which further reinforces our fit to the very scattered low pressure aragonite data. At particle velocities of 0.9 and 1.0 km/s, the \( u_s - u_p \) slopes of calcite (Adadurov et al. data) and aragonite, respectively, undergo a shift to shallower values within 2% of each other. In fact, the slopes of all the carbonate data above \( u_s \approx 1 \) km/s are subparallel. This pattern is typically displayed by data sets for materials of varying porosities [see, e.g., Carter and Marsh, 1980] whereby increasingly porous sample \( u_s - u_p \) fits lie at lower \( u_s \) values. Also, the apparent shift of the slope break toward higher \( u_p \) values with increasing CaCO\(_3\) starting density \( p \) is consistent with the observations made by Jackson and Ahrens [1979] on porous forsterite data. They speculate that increased internal energies at greater material porosities facilitate the onset of phase transitions at lower shock pressures.

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would, however, provide some indication of melting during shock compression. In an extensive investigation of phase transitions during shock processes, Carter [1973] showed that minor slope changes in the $U_{s}-u_{s}$ slopes of Eu, Er, and Pb do, in fact, coincide with the calculated melting and experimental Hugoniot curve intersections and concluded that shock melting occurs in these metals. However, the slope change in the aragonite $U_{s}-u_{s}$ Hugoniot is a major discontinuity, and as will be shown by Hugoniot temperature calculations presented in the following paragraphs, melting does not appear to be a feasible mechanism for the observed behavior. The second possibility is considered because, very typically, in the case of silicate mineral and rock Hugoniots, a decrease in $U_{s}-u_{s}$ slope indicates the onset of a mixed phase regime. The trends displayed by the carbonate data are not inconsistent with this interpretation; however, such an assumed mixed phase region would have an uncharacteristically long duration. As seen in Figure 6, even the most porous data of Kalashnikov et al. [1973] do not deviate from linearity at high $u_{s}$ values, and his nonporous results would indicate the existence of a mixed phase regime between 15 and 94 GPa; the improbability of such a phenomenon leads to rejection of the second possibility. The first possibility, i.e., transition to a high pressure phase, remains a viable explanation.

To evaluate more thoroughly the feasibility of these speculations, we will attempt to construct an isentrope for the high pressure CaCO$_3$ phase and use this reference curve to calculate temperatures along the calcite and aragonite Hugoniots. A third-order Eularian (Birch-Murnaghan) formulation is assumed for the high pressure phase isentrope:

$$P_S(p) = \frac{3K}{2} \left[ (\rho/\rho_0)^{7/3} - (\rho/\rho_0)^{5/3} \right] \left[ 1 + \frac{3}{4} \left( (\rho/\rho_0)^{2/3} - 1 \right) \right]$$

where $\rho_0$ is the zero-pressure density, and $K$ and $K'$ are the zero-pressure values of the bulk modulus and first pressure derivative, respectively, of the high pressure phase. This calculated isentrope pressure $P_S$ is related to the Hugoniot pressure $P_H$ at some given $p$ through the Mie-Grüneisen thermal equation of state:

$$P_H = \left[ P_S - \gamma \frac{E_S + E_{TR}}{\rho_0} \right] / \left[ 1 - \gamma (\rho/\rho_0 - 1) / 2 \right]$$

where $\rho_0$, $\gamma$, and $E_{TR}$ refer to the initial density of the shocked material, the Grüneisen parameter, and the energy of transition between the low and high pressure phases at STP, respectively. $E_S$ represents the internal energy change with isentrope compression from the zero-pressure density of the high pressure phase, $\rho_0$, to some given density, $\rho$:

$$E_S = \int_{\rho_0}^{\rho} \frac{P_S}{\rho^{\gamma/2}} d\rho$$

Using the appropriate input parameters of $\rho_0$, $K$, $K'$, $E_{TR}$, and $\gamma$, it should, therefore, be possible...
Fig. 6. Comparison of carbonate Hugoniot data, including porous polycrystalline calcite, in the $(u_p, u_s)$ plane. Parameters of fits to the data presented in Table 3.

to bring the $P_H$ values calculated from (5) into coincidence with the experimentally determined calcite and aragonite Hugoniot.

As already discussed, static compression data that would allow us to characterize the high pressure carbonate phase are not available; $\rho_o$, $K$, and $K'$, therefore, remain unconstrained parameters. The energy of transition to the high pressure phase is also unknown. However, a lower bound to $E_{TR}$ may be approximated by the sum of transition energies for the calcite I to II and II to III transitions. Using the data of Singh and Kennedy [1974], an $E_{TR}$ for calcite I to III of 24 $J/g$ is calculated. As listed by Ahrens et al. [1969a,b], typical estimated $E_{TR}$ for a variety of minerals range from 300 to $\sim$1000 $J/g$. However, since they refer to transitions occurring at pressures in excess of 30 GPa, where the carbonate transitions in question occur at pressures 50% lower, an upper $E_{TR}$ limit of 200 $J/g$ (for the carbonate transitions) appears reasonable. In our isentrope model calculations, therefore, $E_{TR}$ was varied between 20 and 200 $J/g$.

The Grüneisen parameter, assumed to be of the form

$$\gamma = \gamma_o \left( \frac{\rho_o}{\rho} \right)^n$$

was determined from the two sets of porous calcite data reported by Kalashnikov et al. [1973]. The $\gamma$ calculated for both data sets are plotted as a function of $\rho$ in Figure 7. An attempt was made to use Solenhofen limestone (4.6% porous) data, but the calculated $\gamma$ values showed extreme scatter. If inferences may be drawn from two data sets, the results of Figure 7 point to a direct dependence of $\gamma$ on porosity. Therefore, we used the $\gamma$ formulation determined from the less porous data set, i.e., $\gamma = 1.53 \left( \frac{\rho_o}{\rho} \right)^{0.6}$. The assumption

<table>
<thead>
<tr>
<th>Sample</th>
<th>Source</th>
<th>$\rho_o$ g/cm$^3$</th>
<th>$c_o$ km/s</th>
<th>$s$ GPa</th>
<th>Pressure Range, GPa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single crystal aragonite</td>
<td>present work</td>
<td>2.930</td>
<td>3.77</td>
<td>2.52</td>
<td>6-18</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5.02</td>
<td>1.30</td>
<td>18-40</td>
</tr>
<tr>
<td>Polycrystalline calcite</td>
<td>Adadurov et al. [1961]</td>
<td>2.703</td>
<td>3.40</td>
<td>2.00</td>
<td>5-13</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3.99</td>
<td>1.32</td>
<td>13-51</td>
</tr>
<tr>
<td>Polycrystalline calcite</td>
<td>Kalashnikov et al. [1973]</td>
<td>2.665</td>
<td>3.70</td>
<td>1.44</td>
<td>10-94</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2.020</td>
<td>1.74</td>
<td>10-71</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.705</td>
<td>1.15</td>
<td>10-59</td>
</tr>
<tr>
<td>Solenhofen limestone</td>
<td>van Thiel et al. [1977]</td>
<td>2.585</td>
<td>3.62</td>
<td>1.39</td>
<td>8-90</td>
</tr>
</tbody>
</table>
that $\gamma$ for the high pressure CaCO$_3$ phase is small is also consistent with the close coincidence of the calcite and aragonite Hugoniot at high pressure. Small variations in $\gamma$, however, do not significantly affect the model isentrope calculations.

The self-consistent sets of model parameters that best fit experimental calcite and aragonite Hugoniots are summarized in Table 4. It is evident from this table that varying the high pressure phase density by 0.1 g/cm$^3$ profoundly affects the bulk modulus and first pressure derivative values in the model calculations. Performing the calculation with $\rho_0$ greater than 3.1 g/cm$^3$ would constrain $K$ to very high values, above 100 GPa. On the other hand, the calculations are relatively insensitive to changes in $E_{TR}$. Therefore, even by allowing an $E_{TR}$ of 70 J/g for the calcite to aragonite transition (a maximum value calculated according to McQueen et al. [1963], assuming a calcite to aragonite transition at 9 GPa), it is not possible to force both calcite and aragonite to the same high pressure phase. Nevertheless, the $K/K'$ values for both polymorphs (at the same $E_{TR}, \rho_0$ conditions) are very similar, and transformation of both polymorphs to the same high pressure phase cannot be discounted. In the Hugoniot temperature calculations described in the following paragraph, the entire range of model isentrope parameters presented in Table 4 is considered.

Temperatures along the Hugoniot are determined from

$$T_H = 298 \exp \left[ \int \frac{\gamma}{\rho_0} d\rho \right] + \frac{E_E - E_S}{C_V}$$

(8)

The first term in this equation is the temperature rise along the high pressure phase isentrope, and the second is the temperature difference at some density between this calculated isentrope and the (experimental) Hugoniot. $E_S$ is given by (6), and $E_E$ by the Rankine-Hugoniot conservation equation

$$E_E - E_0 = 1/2(P_0 + P_H)(1/\rho_0 - 1/\rho)$$

(9)

where the standard $E_0$ and $P_0$ conditions are taken to be equal to zero and $\rho_0$ is the initial density of the shocked material. We assume that the heat capacity at constant volume $C_V$ is equal to the Dulong-Petit value of 3R, where R is the gas constant. According to experimental heat capacity measurements [Robie et al., 1978], 90% of this

![Fig. 8](image)

Fig. 8. Pressure versus temperature relation along aragonite and calcite Hugoniots. Numbers in parentheses represent ($E_{TR}$: $\rho_0$; $K$; $K'$) values used in model isentrope calculations along which temperature estimates are made. $\gamma$ was assumed to be of the form $\gamma = 1.53 (\rho_0/\rho)^{0.6}$. Calcite melting curve [Irving and Wyllie, 1973] appears in the right of the graph.

![Fig. 9](image)

Fig. 9. Plot showing systematic relationship between standard entropy, $S_{298.15K}$, and molar $S_298.15K$ for speculated high pressure CaCO$_3$ phase.
Hugoniot state. In turn, the entropy along the reached when calcite is shocked to a relatively pressure of 1 bar is the same as that of the melting temperature ($T_{local}/T_{melt} > 0.5$) is $\sim 18$ GPa on the calcite and aragonite Hugoniots, considering shock deformational behavior, on the vaporization at 1 bar. Since release is an several minerals including calcite. According to value along the release path with the entropy of intensity of local temperature excursions in vaporization upon unloading. As discussed above, passage of the shock wave. Thus, even though refer to the transition energy, initial density, waste heat approximation. The agreement adds the release of CaCO$_3$ to ambient pressures, the local shear bands. In fact, such zones of intense release, whereas release from pressures above of intense deformation and extreme temperature below $\sim 14$ GPa occurs along steep paths and to these authors, instabilities intrinsic to shock determinations. The calcite melting curve suggested interpretation that vaporization upon the bulk modulus, and first pressure derivatives of the high pressure phase values, respectively, used in the high pressure phase isentrope experimental evidence is consistent with the high pressure phase $O_o$ of $3.1 + 3.0$ g/cm$^3$ respectively, is based on the systematic relationship, shown in Figure 9, between $S_{298.15K}$ and the molar volume of carbonates [Robie et al., 1978]. The incipient vaporization criterion involves coincidence of the release state entropy value with the entropy increase achieved in raising a material to its vaporization temperature at 1 bar; according to the experimentally determined vaporization curve [Baker, 1962], vaporization of calcite at 1 bar occurs at 1171 K. Complete vaporization entropy is the sum of the entropy gained in raising the material to its vaporization temperature and the dissociation entropy at that temperature. Calculated postshock entropies for calcite and aragonite are presented in Figure 10; entropies for complete vaporization, $\Delta S \approx 3$ J/g K, do not appear to be reached below shock pressures of $\sim 100$ GPa for either calcite or aragonite. According to this model, then, minimum shock pressures for incipient vaporization are 33 and 55 GPa for calcite and aragonite, respectively. These values are similar to shock pressures required for vaporization as calculated by Kieffer and Simonds [1980] using the waste heat approximation. The agreement adds support to the contention that $\gamma$ is small. Thus, although the shock and release state experimental evidence is consistent with the suggested interpretation that vaporization upon high temperatures and pressures of carbonates at shock pressures in excess of $\sim 17$ GPa, theoretical considerations are in pronounced conflict with this interpretation. Possible causes for this discrepancy might be found by considering shock deformational behavior, on the microstructural scale, of brittle, thermally insulating solids. Grady [1980] and Horie [1980] extensively discuss the role of heterogeneous deformation during shock compression. According to these authors, instabilities intrinsic to shock compression processes result in localized regions of intense deformation and extreme temperature gradients which may persist for some time after passage of the shock wave. Thus, even though equilibrium temperature estimates are well below those required for melting, local 'hot spot' temperatures may be sufficient to melt material in local shear bands. In fact, such zones of intense local deformation have been observed in shock-recovered materials [see, e.g., Jeanloz et al., 1977]. Grady [1980] estimates the intensity of local temperature excursions in several minerals including calcite. According to his calculations, a significant fraction of the melting temperature ($T_{local}/T_{melt} > 0.5$) is reached when calcite is shocked to a relatively
modest pressure of 4.4 GPa. Although it is difficult to assess the extent to which processes of adiabatic shear and heterogeneous melting influence the Hugoniot results, their effects, at least in a qualitative sense, should be given serious consideration in evaluation of shock compression data.

Summary and Conclusions

The aragonite Hugoniot to 40 GPa can be summarized as follows. The lower portion of the curve is inferred to display a Hugoniot elastic limit between 1.8 and 3.3 GPa and a phase transition, possibly displacive, between 5.5 and 7.6 GPa. Above shock pressures of ~10 GPa, the density-pressure curve becomes initially shallower and remains smooth to 40 GPa; the calcite and aragonite Hugoniots in the 10 to 40 GPa range are, within the limits of experimental error, coincident, suggesting the transformation of both polymorphs to a common high pressure phase.

Measured release adiabats for aragonite form shock states below ~14 GPa show consistently steep initial unloading paths (in the density-pressure plane) and final zero-pressure densities up to 10% greater than the initial density. Unloading paths from pressures greater than ~17 GPa are unusually shallow, a pattern that is consistent with the interpretation that vaporization occurs upon unloading. Although the experimental results may suggest vaporization upon release from these pressures, theoretical shock temperature and entropy calculations decidedly conflict with this interpretation.

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