

Equation of State of Forsterite¹

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Shock wave data for pure forsterite with initial bulk densities of 2.6 and 3.1 g/cm³ are obtained to 0.370 Mb by impacting series of specimens with tungsten alloy plates that are launched at speeds of up to 2.3 km/sec with a high-performance propellant gun. The onset of a shock-induced phase change, probably corresponding to the forsterite-'post spinel' phase change is observed at 0.280 ± 0.025 Mb. Because of the low shock temperatures, the transition is believed to be limited by the reaction rate and this pressure value should be taken only as an upper limit. Adiabats derived from the Hugoniot data for the forsterite phase are fit to the two-parameter finite strain Birch-Murnaghan equation and to two simple ionic equations of state. The Birch-Murnaghan form of the equation of state gives a zero-pressure bulk modulus (1.29 Mb) that agrees more closely with the ultrasonic data than the modulus obtained from the ionic equations of state. An unusual relaxation effect, in which the elastic shock precursor velocity varies from 5.8 to 9.5 km/sec, is also observed. The characteristic time of the relaxation process appears to be less than 1 μ sec.

The equation of state of the pure magnesium end member of the olivine series, forsterite, Mg₂SiO₄, plays a key role in the application of seismological and high-pressure ultrasonic and phase equilibrium data to investigation of the mineralogies and temperatures within the earth's upper mantle [Birch, 1952; Anderson, 1967; Ringwood, 1969]. In addition to the early pressure-density data of Bridgman [1948], complete elastic constant data to 400°C and 10 kb have recently been obtained ultrasonically on nearly pure crystals by Graham and Barsch [1969] and Kumazawa and Anderson [1969]. The previous shock-wave equation of state measurements of Hughes and McQueen [1958], McQueen et al. [1967], Trunin et al. [1966], and Ahrens and Petersen [1969] for various dunites have been useful in delineating the onset of a shock-induced phase change and, in some cases, the equation of state of a shock-induced high-pressure phase. For the dunites the shock-induced high-pressure phase has an apparent zero-pressure material density close to the density of an isochemical mix of the close-packed oxides SiO₂ (stishovite), MgO, and FeO. From the static high-pressure quenching experiments of Ringwood and Major [1970] it appears that at 1000°C pure forsterite

transforms to the distorted spinel or β olivine form (zero-pressure density, 3.48 g/cm³) at ~ 145 kb, the pure spinel phase of Mg₂SiO₄ (3.55 g/cm³) occurs at still higher pressures. Ahrens and Syono [1967] and Anderson [1967] have shown that above ~ 200 kb the pure spinel phase is unstable with respect to the denser oxides. On the basis of quenching experiments on analog compounds Ringwood and Reid [1968] have suggested that the 'post-spinel' phase of forsterite has the structure of either K₂NiF₄ or Sr₂PbO₄. The inferred zero-pressure density of the high-pressure shocked phases of the dunites [Ahrens et al., 1969], as well as the presumably ferromagnesian material of the mantle at depths below 900 km (~ 300 kb), is consistent with one of these structures [Anderson, 1970]. In addition to the data for dunites, a series of six Hugoniot points for the high-pressure phase of a ceramic forsterite has been obtained by McQueen and Marsh and is given in Clark [1966].

To obtain additional data on the equation of state and the onset of the shock-induced phase change in forsterite, we have conducted a series of shock-wave experiments on this material. Because of the stability of forsterite over a wide range of pressures under shock conditions, the apparent absence of significant strength effects, and the availability of precision ultrasonic data at low pressure, we have also fit the present data with several equations of state.

¹ Contribution 1775, Division of Geological Sciences, California Institute of Technology.

SPECIMEN MATERIALS

Polycrystalline forsterite obtained from Muscle Shoals Electrochemical Corporation was used to fabricate most of the specimens in the present study. This material, which is slightly porous, was fused from the oxides. It is light gray in appearance and has tabular grains 0.5–2 mm long. It came to us in grapefruit-sized pieces that were cored and sawed into 80 disks that were 22 mm in diameter and 4 or 6 mm thick. On the basis of visual and radiographic inspection 25 samples that were free from cracks, large pores, and obvious inclusions were finally used as specimens. Although the thin sections showed no phases other than forsterite to be present, a series of X-ray (shadowgraph) radiographs of some of the specimens showed them to contain a few ~ 0.1 -mm-thick slivers of a dense phase believed to be FeSi (density, 6.1 g/cm³). Using the apparent areas of the inclusion shown on the radiographs and assuming random orientation, we calculated that these slivers occupied about 0.02% of the sample volume; hence, they increased the bulk density by $\sim 0.04\%$. Microprobe analysis (Table 1) of the forsterite phase indicates that this material is stoichiometric and nearly iron free within the detectable limits of the instrument ($\sim \pm 0.5\%$). Using *Graham and Barsch's* [1969] value of 3.2217 g/cm³ for the density of forsterite and correcting for FeSi inclusions, we adopted a value of 3.223 g/cm³ for the zero-pressure crystal density of this material.

After machining the specimen to tolerances of ± 0.005 mm and removing cutting solvents in an ultrasonic cleaner, the samples were weighed and their dimensions recorded. Volume corrections of up to 0.01% were applied to take account of chips removed in the grinding process. The bulk density of all the specimens was deter-

mined to within ± 0.009 g/cm³ (Table 2). This uncertainty is wholly attributed to uncertainty in the volume determination.

In addition to the fusion cast material, three shots were fired at low pressure with a pressed aggregate of forsterite purchased from Atomergic Corp. This material had a bulk density of 2.6 g/cm³. Although binding agents were not used in preparing the samples, thin sections and electron microprobe scans indicated that a small amount of an isotropic aluminum-bearing phase, probably MgAl₂O₄, was present. Because of the uncertainties inherent in reducing the densities of the equation of state point by 0.02 ± 0.005 g/cm³ to take the impurities into account, further work on this material was discontinued.

EXPERIMENTAL DETAILS

The machined specimen disks were mounted on ground and polished 1.5-mm-thick driver plates of tungsten alloy (W, 90.4%; Ni, 4.65%; Cu, 4.90%; density, 16.9 g/cm³). Flat and inclined mirrors (with reflecting surfaces toward the direction of the incident shock) were mounted on the assembly (Figure 1) to record the shock arrivals at the specimen-driver interface and at the specimen's free surface. Various details of the technique, which also permits measuring the shock-induced free-surface velocity, were described by *Fowles* [1967] and *Ahrens et al.* [1968]. After using an optical lever setup to ensure that the flat mirrors lie within 4' of arc of planes parallel to the driver-specimen interface and after measuring the inclined mirror angle, contactor pins (E.G.&G. CA1038) were inserted in the driver plate and the sample was mounted in the semi-expendable assembly shown in Figure 1.

Shocks of different amplitude were driven into a series of specimens by impacting the specimens with flyer plates held in a Lexan plastic sabot [*Jones et al.*, 1966]. The flyer plate was made of the same alloy as the driver plate but was 3.8 mm thick, and the sabot was accelerated to speeds of 0.3 to 2.3 km/sec by a propellant gun 40 mm in diameter and 8 meters long. The assembly was suspended in a 5600-liter evacuated tank into which the gun tube protrudes. Before firing the projectile in the geometry depicted in Figure 2, the sample assembly was optically aligned with the axis of the gun tube. Details of this apparatus, the projectiles, and the alignment

TABLE 1. Composition of Muscle Shoals Synthetic Forsterite

| Element | Formula Proportions |
|---------|---------------------|
| Mg | 2.03 |
| Al | 0.001 |
| Si | 0.983 |
| Fe | 0.0004 |

Composition determined by electron microprobe method, A. Albee and A. Chodos.

and calibration procedures will be described in a forthcoming paper.

Because we employed the impedance-match method [Walsh and Christian, 1955] in the experiments to obtain the high-pressure shock state in the sample, the velocity of the flyer plate and hence the velocity of the projectile had to be measured. This was accomplished during the time that the projectile traveled the final 50 cm of the barrel. The projectile successively intersected three He-Ne laser beams that were oriented perpendicular to the bore axis and were detected by three photodiodes. The first two beams were brought through the barrel via holes 3 mm in diameter and the third beam was placed 1 cm from the gun-tube muzzle. After flying free in the vacuum chamber for a distance of 50 cm, the projectile struck the electrical trigger pins affixed to the target. The photodiode signals produced by interruption of

the laser beams and a signal produced by an external circuit upon the projectile's impact with the contactor pins were used to turn three time-interval counters on and off. The counters provided projectile travel times between the four stations to an accuracy of $\pm 0.1 \mu\text{sec}$. The distances required to calculate average projectile velocity within the three time intervals were separately measured to $\pm 0.1 \text{ mm}$, which provided an over-all precision of $\pm 0.2\%$ for the velocity between stations. A raster oscilloscope, triggered by the first photodiode, which later displayed the pin-contactor closure signal, provided redundant timing data.

To measure the velocity of the shocks induced in the sample upon projectile impact, the times between destruction of the mirror reflecting coatings were obtained by using an image converter camera (TRW model 1D). The optical path from the xenon flash tube (ignited when

TABLE 2. Hugoniot Data for Forsterite

| Shot | Thickness, mm | Initial Density, g/cm ³ | Projectile Velocity, km/sec | Elastic Precursor Velocity, km/sec | Shock Velocity, km/sec | Particle Velocity, km/sec | Pressure, Mb | Density, g/cm ³ |
|------|---------------|------------------------------------|-----------------------------|------------------------------------|------------------------|---------------------------|--------------|----------------------------|
| A82 | 4.80 | 2.634 | 1.201 ± 0.005 | 5.91 $\pm .03$ | 4.19 ± 0.05 | 1.01 | 0.113 | 3.463 |
| A70 | 4.83 | 2.627 | 1.17 $\pm .04$ | 5.82* ± 0.08 | 4.21 ± 0.04 | 1.03 | 0.116 | 3.489 |
| A75 | 4.83 | 2.633 | (1.42) | 5.83 ± 0.05 | 4.98 ± 0.08 | 1.21 | 0.159 | 3.469 |
| A68 | 6.27 | 3.093 | 1.11 ± 0.04 | 6.96 ± 0.12 | 6.83 ± 0.12 | 0.855 | 0.181 | 3.536 |
| A60 | 6.26 | 3.094 | 1.107 ± 0.005 | 7.03 $\pm .06$ | 6.48 ± 0.06 | 0.854 | 0.172 | 3.567 |
| A104 | 4.20 | 3.087 | 1.161 ± 0.006 | 8.22 ± 0.04 | (6.74) | 0.897 | 0.187 | 3.561 |
| A90 | 4.54 | 3.117 | 1.305 ± 0.01 | 8.48 ± 0.45 | 7.13 ± 0.14 | 0.995 | 0.221 | 3.623 |
| A105 | 4.20 | 3.119 | 1.285 ± 0.007 | 9.5 ± 0.5 | (7.28) | 0.975 | 0.222 | 3.602 |
| A65 | 6.25 | 3.104 | 1.53 ± 0.02 | 7.62 ± 0.34 | 7.47 ± 0.04 | 1.16 | 0.269 | 3.674 |
| A91 | 4.57 | 3.102 | 1.77 ± 0.02 | 8.29 ± 0.13 | 7.33 ± 0.09 | 1.35 | 0.307 | 3.801 |
| A93 | 4.54 | 3.115 | 2.132 ± 0.007 | 8.50 ± 0.14 | 7.34 ± 0.06 | 1.63 | 0.373 | 4.004 |
| A101 | 4.55 | 3.113 | (2.2) | (8.77) | † | † | † | |
| A89 | 4.43 | 3.118 | (0.315) | 5.2 ± 0.3 | † | † | † | |

Values in parentheses indicate a single measurement.

* Particle velocity obtained from free-surface motion of 0.031 km/sec gives elastic shock state; stress, 0.0048 Mb; density, 2.655 g/cm³.

† Second shock present but not recorded.

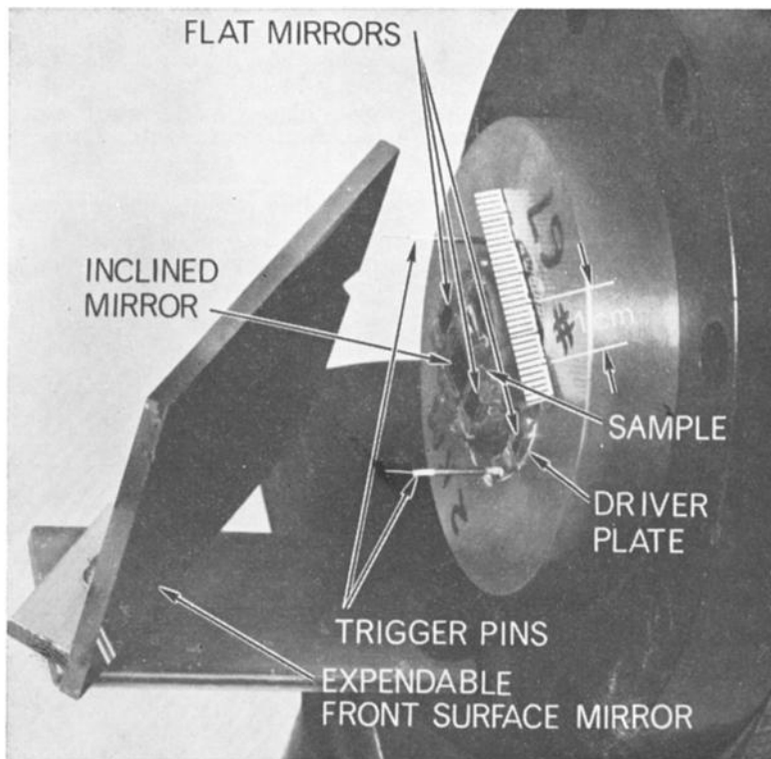


Fig. 1. Forsterite target assembly. Expendable front surface mirror transmits xenon flash tube illumination to flat and inclined mirrors and transmits light reflected from target mirrors to streak camera. Electrical contactor pins, activated upon projectile impact from left, actuate sweep of image converter streak camera.

the projectile passes the third laser beam) extended via an expendable mirror to the target, then to the expendable mirror, and then to an objective lens. The objective lens focused the image of the rear target surface on a slit that was external to the vacuum tank. The image converter camera, which was also focused on the slit, was sweep-triggered by the pin closure. A resulting streak record showing the typically rather ragged elastic precursor and the smooth following deformational, or main, shock in forsterite is shown in Figure 3. The reader is referred to papers by *Fowles* [1967] and *Ahrens et al.* [1968] for the various details of such records.

Application of image-converter streak cameras to precision shock transit-time measurements has previously been limited by the lack of suitable calibration methods for streak writing rates. Initially we used the oscilloscope calibration technique described by *Ahrens and Spetzler* [1969] for this purpose. A second method used

in the early experiments (prior to shot A70) was to time, by single sweep oscillography, the duration of the 'shutter' pulse that turns on the flow of electrons in the image converter tube for a time corresponding to the camera sweep interval. This procedure is easier to carry out than the *Ahrens and Spetzler's* [1968] method and yields an average writing rate over the streak length that is accurate to $\pm 0.5\%$. For later shots the intensity-modulated laser beam technique, depicted in Figure 2, was employed, which permitted the camera writing rate to be obtained to within $\pm 0.17\%$ at any point along the streak record (Figure 3). *Davies* [1970] has recently described a similar method.

RESULTS

The streak records obtained in the experiments performed on both the 2.6- and the 3.1-g/cm² forsterite showed that the initial shock-wave arrival (loss of mirror reflectivity) in the flat

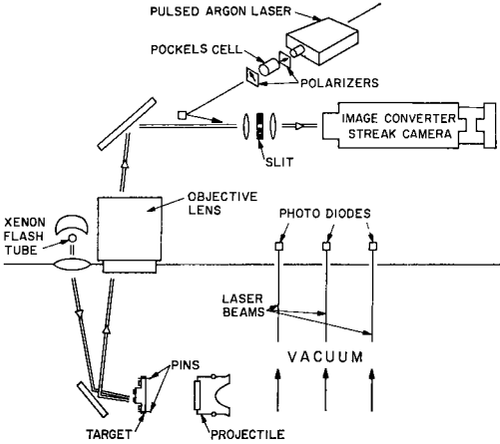


Fig. 2. Diagrammatic sketch of equation of state experiment. Projectile travels in vacuum. Velocity is determined by the projectile's cutting three laser beams and striking the contactor pins. Successive interaction with mirrors of resulting shock in sample, recorded as a function of time with streak camera, determines shock velocity. Pockels cell intensity modulates pulsed laser, providing timing marks for streak camera.

mirrors occurred well before the onset of appreciable free-surface motion was detectable in the inclined mirror image (Figure 3). Because it was also observed that the onset of free-surface motion coincided in time with a second usually complete loss of reflectivity in the flat mirrors, we inferred that an elastic shock or precursor was present in all the experiments. The fact that the free-surface velocities, which were measured with inclined mirrors, were approximately equal to twice the final shock particle velocity supported this conclusion. The elastic precursor is thought to represent a one-dimensional compression wave in which internal rearrangement of the material has not yet occurred. The second very much stronger shock corresponds to achievement of a state in which the material has undergone internal deformation and the compression is no longer one dimensional on a microscopic scale.

The motion of the free surface associated with the precursor arrivals in the 2.6-g/cm³ sample was just barely detectable. For shot A70, a precursor-induced free-surface velocity of 0.062 km/sec was measured. Assuming that the free-

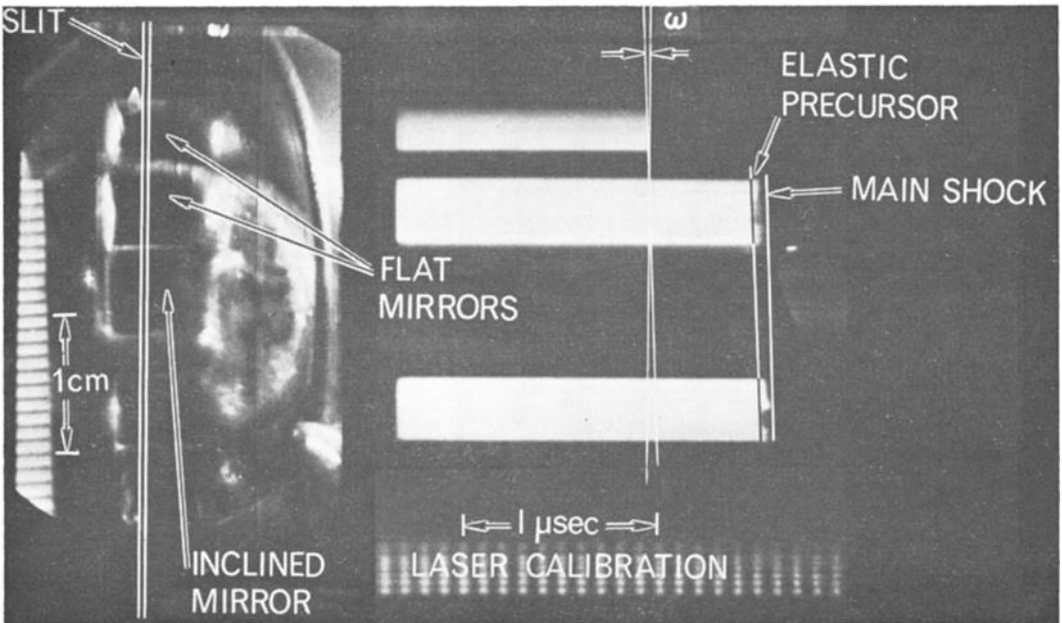


Fig. 3. Streak camera record obtained from rear surface of forsterite target assembly upon being impacted with flyerplate traveling at 2.1 km/sec. (Left) Image of target assembly as viewed statically through streak camera showing position of slit with respect to reflecting mirrors. (Right) Streak record showing irregular arrival of elastic precursor, followed by main shock.

surface velocity represents approximately twice the precursor particle velocity, we can calculate a precursor amplitude of ~ 0.0048 Mb. The surprising lack of a measurable amplitude from the precursor free-surface velocity for the denser 3.1-g/cm^3 forsterite forced us to conclude that the precursor amplitudes at the free surface were less than ~ 0.005 Mb. The inclined mirror angles used on both classes of samples were similar. A maximum stress offset of the deformational Hugoniot above the 'hydrostatic' curve of 0.0022 Mb was calculated from simple elastoplastic theory [Fowles, 1961] with a Poisson's ratio of 0.24 . We infer from the apparent rapid attenuation of the elastic precursor (discussed below) that even this 'maximum' value for strength effect is probably too high and that the deformational shock states achieved are effectively hydrostatic.

For some of the shots appreciable correction for shock tilts were applied in calculating the shock velocities by using the relations

$$\tan \epsilon = M \tan \omega V_a / W \quad (1)$$

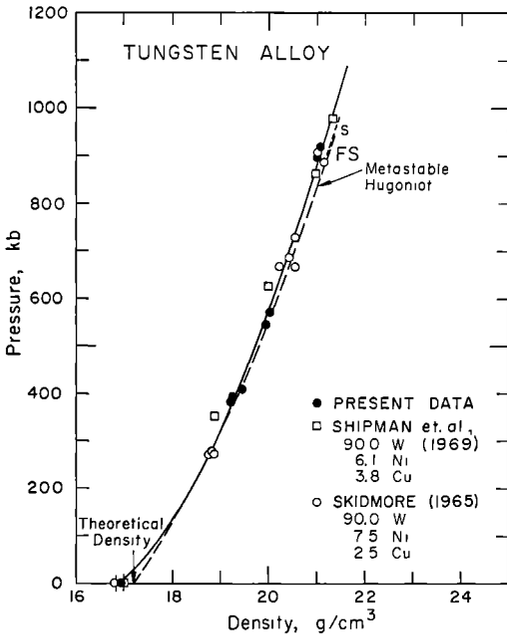


Fig. 4. Tungsten alloy Hugoniot data used for reduction of forsterite shots. Metastable Hugoniot indicates theoretical curves for samples having no porosity with compositions corresponding to Skidmore [1965] (S) and Shipman et al. [1969] (FS) data.

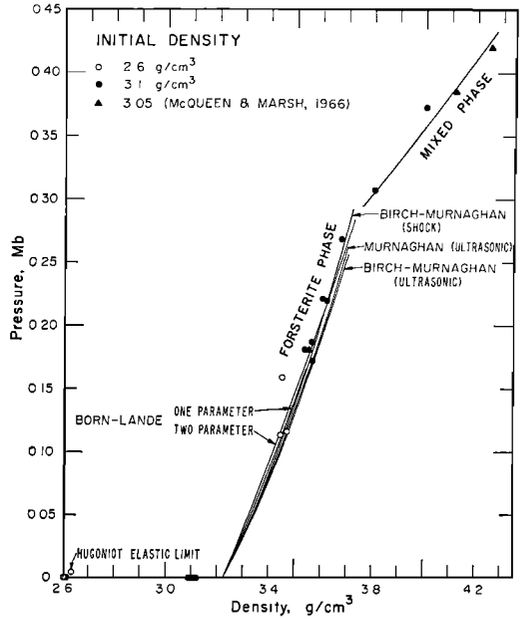


Fig. 5. Hugoniot data for pure forsterite, initial densities 2.6 and 3.1 g/cm^3 . Various principal adiabats for forsterite derived from shock and ultrasonic data are shown. Metastable transition, probably from forsterite to post-spinel phase, occurs at 0.280 ± 0.025 Mb.

$$V = V_a \cos \epsilon \quad (2)$$

where ϵ is the velocity component along the slit of the shock tilt angle with respect to the initial free surface, M is the magnification of the film, and W is the streak camera writing rate. The term ω is defined in Figure 3, and V_a and V are the apparent and actual shock velocities, respectively. In future experiments we expect also to correct for tilt in the direction perpendicular to the slit. The varying degrees of shock tilt experienced are reflected in the uncertainties assigned to the shock velocities in Table 2.

The final deformational shock states were calculated by using the impedance-match method and the Rankine-Hugoniot equations. The Hugoniot of the tungsten alloy flyer and the driver plate standard material used in the reduction of the data is shown in Figure 4. The measurements on this tungsten alloy will be reported separately.

In the analysis of the 2.6-g/cm^3 forsterite data a precursor particle velocity of 0.031 km/sec

and a precursor amplitude calculated from the observed shock velocity were assumed for the state in which the deformational or second shock wave impinged. For the denser 3.1-g/cm³ forsterite the second shock was assumed to be incident into material with zero initial velocity and pressure.

The major uncertainty in projectile velocity, and hence particle velocity in the shock state, arises not from the uncertainty in the absolute measurements of velocity between stations ($\pm 0.2\%$), but from the uncertainty in extrapolating the velocity to the time of impact over the last 50 cm of projectile flight. For shots fired before A90 both minor accelerations and decelerations of the projectile were measured in the last meter of travel. Later shots employing heavier projectiles (90 grams) all showed that the projectile was accelerating all the way to the target. This regularity reduced uncertainties where timing data from one of the stations were not obtained.

The resulting Hugoniot data are plotted in Figure 5 in relation to the pressure-density curves inferred from ultrasonic data. It is clear that the data for the 2.6-g/cm³ material (especially shot A75) are more scattered. Impurities may account for the scattering. Because the adiabats derived from the Hugoniot data are in acceptable agreement with the ultrasonic curves below 200 kb, finite strength effects, at least of a gross sort, are not measurably present. The reduced data for the equation of state in the forsterite regime (below ~ 0.280 Mb) are discussed below. Above 0.280 ± 0.025 Mb the present Hugoniot data, when taken with points obtained by McQueen and Marsh [given in *Anderson and Kanamori*, 1968], clearly indicate the onset of a region of anomalous compression that undoubtedly corresponds to a phase change taking place in the forsterite. The one low-pressure point obtained by McQueen and Marsh, which also closely fits the present data, supports this conclusion. The crystal structure of this high-pressure phase is not actually known, but the pressure at which the transition begins under shock suggests that forsterite is transforming to a phase in which the silicon ions are octahedrally coordinated by oxygen, although the onset of an overdriven olivine to spinel transition cannot be ruled out. At higher shock pressures the data of *McQueen et al.* [1967] indicate that a denser-

than-spinel phase forms under shock compression. *Ringwood and Reid* [1968], on the basis of high-pressure quenching experiments on analog compounds and on crystal-chemical considerations, suggested that the K_2NiF_4 or Sr_2PbO_4 structure will be assumed by this phase. Analysis of the Los Alamos high-pressure shock data for dunites and forsterite are consistent with zero-pressure densities of 4.1 and 4.0 g/cm³ for these structures [*Ahrens et al.*, 1969]. The present shock-induced transition pressure of 0.280 ± 0.025 Mb (shock temperature $\sim 160^\circ\text{C}$) could be taken as an upper bound to the forsterite-post spinel phase transition pressure. Because of the low shock temperatures the transition is probably reaction-rate limited. *Anderson* [1967] and *Ringwood* [1969] have suggested that this so-called post spinel phase is the predominant mineral of the lower mantle.

ELASTIC PRECURSORS

The behavior of the elastic precursor observed in the present experiments on the 3.1-g/cm³ forsterite is sufficiently unusual so as to deserve some comment. Examination of the apparent precursor velocities V_{1a} for the denser forsterite gives a surprisingly random series of values. In contrast, the precursor wave velocity for the 2.6-g/cm³ material (Table 2) is observed to be

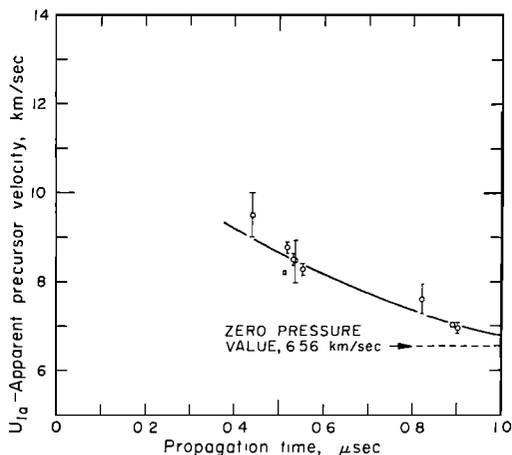


Fig. 6. Apparent precursor velocity versus shock propagation time for 3.1-g/cm³ forsterite. A relaxation process appears to be taking place in which precursor velocity approaches zero-pressure longitudinal velocity with an apparent characteristic time of less than 1 μsec.

nearly constant at 5.82–5.91 km/sec. (The range of pressures and specimen thicknesses for these samples is not large.) Since the sample thicknesses for the 3.1-g/cm³ material varies from 4.2 to 6.3 mm the existence of a stress relaxation effect could be indicated by plotting the parameter U_{ia} versus propagation time, as in Figure 6. This representation orders these data to a remarkable degree. At long propagation times the precursor velocity approaches the longitudinal elastic velocity 6.56 km/sec of the initially porous material (R. N. Schock, private communication). The time constant for the relaxation process is apparently less than 1 μ sec. This effect appears to be markedly different than the effect observed by *Wackerle* [1962] and *Ahrens and Duvall* [1966] in single and polycrystalline quartz, respectively. For the nonporous quartz, to first order, the shock velocity remains constant and the apparent particle velocity decays. Although not detectable in these experiments, it is likely that the particle velocity in the forsterite precursor may also decay.

EQUATION OF STATE PARAMETERS

We have reduced the shock data below 0.280 Mb (Table 2) to various adiabats, isotherms, and metastable Hugoniot forms of the equation of state by the methods outlined in *Ahrens et al.* [1969]. Because the ultrasonic data are related directly to the adiabats, they are discussed below. Because the two groups of samples differed significantly in their initial bulk density (2.6 and 3.1 g/cm³), the shock data were reduced specifically so as to take the different porosities into account. Rather than average initial densities and perform further reductions from a mean analytic Hugoniot curve, we calculated the pressure along the principal adiabat (or isentrope) for each density achieved along the Hugoniot. The principal adiabat is the adiabat centered at ambient pressure and at 25°C. For a Hugoniot state at a pressure and density p_H and ρ , respectively, the adiabatic pressure at that density is given by [*Ahrens et al.*, 1969]

$$p_a = p_H - \gamma \rho \cdot \left[p_H(1/\rho_{00} - 1/\rho)/2 - \int_{\rho_0}^{\rho} p_a d\rho/\rho^2 \right] \quad (3)$$

where ρ_{00} is the initial bulk density of the sample,

ρ_0 is the zero-pressure crystal density, and $\gamma = \alpha K_s/\rho C_p$ is the thermodynamic Grüneisen parameter. Here α is the thermal expansion, K_s is the adiabatic bulk modulus, and C_p is the specific heat at constant temperature. The assumption that γ depends only on volume is implicit in equation 3.

Because of the present lack of pertinent data or an adequate theory describing the variation of γ with pressure and temperature in silicates when they are below their Debye temperature, we assumed the volume dependence of γ to be given by

$$\gamma = \gamma_0(\rho_0/\rho)^n \quad (4)$$

The resultant series of adiabatic points were fit to the two-parameter Birch-Murnaghan, Born-Landé, and Born-Mayer forms of the equation of state given by

$$p = \frac{3}{2} K_0 \left[\left(\frac{r_0}{r} \right)^7 - \left(\frac{r_0}{r} \right)^5 \right] \cdot \{ 1 - \xi \left[\left(\frac{r_0}{r} \right)^2 - 1 \right] \} \quad (5)$$

$$p = \frac{3K_0}{n-1} \left[\left(\frac{r_0}{r} \right)^{n+3} - \left(\frac{r_0}{r} \right)^4 \right] \quad (6)$$

$$p = \frac{3K_0}{(r_0/\rho) - 2} \left[\left(\frac{r_0}{r} \right)^2 \cdot \left[\exp(r_0 - r)/\rho \right] - \left(\frac{r_0}{r} \right)^4 \right] \quad (7)$$

where K_0 is the appropriate zero-pressure bulk modulus and r_0 and r are characteristic lengths in the initial and compressed material, respectively. Hence $(\rho/\rho_0)^{1/3} = r_0/r$. The zero-pressure value of $\partial K/\partial P$ is related to the parameters ξ , n , and r_0/ρ by

$$\partial K/\partial P = 4 - 4\xi/3 \quad (8)$$

$$\partial K/\partial P = (n + 7)/3 \quad (9)$$

$$\partial K/\partial P = \frac{14 - (1 + r_0/\rho)(2 + r_0/\rho)}{6 - 3r_0/\rho} \quad (10)$$

Equations 5–7 are similar in form and all yield similar pressure-density curves for values of $r_0/r \lesssim 1.2$. Equation 5 is valid to third-order Eulerian strain [*Thomsen*, 1970] and can be used to represent an adiabatic curve if the strains are defined in an adiabatic sense. Equations 6 and 7 are derived from a simple static ionic centrosymmetric potential model of a crystal

TABLE 3. Equation of State Parameters Calculated from Forsterite Shock Data

| Equation of State | $\nu = 1.17$ $-(\rho_0/\rho)^n$ | ρ_0^* , g/cm ³ | K_H^* , Mbar | K_T^* | K_S^\dagger , Mb | K_S^\dagger | $(\partial K_S/\partial T)_P$ at 100°-200°C, 10 ⁻³ Mb/°K | $(\partial K_S/\partial T)_P$ at 200°-400°C, 10 ⁻³ Mb/°K | K_S^\ddagger , Mb | K_S^\ddagger |
|-----------------------|------------------------------------|-----------------------------------|-------------------|---------|-----------------------|---------------|---|---|------------------------|----------------|
| Birch-Murnaghan § (1) | 0 | 3.102 | 0.720 | 9.97 | 1.28 | 6.6 | -0.08 | -0.14 | 1.10 | 6.7 |
| Birch-Murnaghan § (2) | 1 | 3.102 | 0.720 | 9.97 | 1.30 | 6.4 | -0.10 | -0.17 | 1.08 | 6.6 |
| Born-Lande § (3) | 0 | 3.223 | 1.66 | 2.7 | 1.63 | 2.5 | | | | |
| Born-Lande § (4) | 1 | 3.223 | 1.66 | 2.7 | 1.44 | 2.7 | | | | |
| Born-Mayer § (5) | 0 | 3.223 | 1.65 | 2.8 | 1.62 | 2.5 | | | | |
| Born-Mayer § (6) | 1 | 3.223 | 1.65 | 2.8 | 1.62 | 2.6 | | | | |
| Born-Mayer § (7) | 2 | 3.223 | 1.65 | 2.8 | 1.62 | 2.7 | | | | |
| Born-Lande (8) | 0 | 3.223 | 1.45 | 5.17 | 1.41 | 5.17 | | | | |
| Born-Lande (9) | 1 | 3.223 | 1.45 | 5.17 | 1.41 | 5.17 | | | | |
| Born-Mayer (10) | 0 | 3.223 | 1.45 | 5.17 | 1.41 | 5.17 | | | | |

Solution number given in parenthesis after equation of state.

* Raw Hugoniot.

† Principal adiabat.

‡ Adiabat at 0.0 atm and 1200°C.

§ Two-parameter equation.

|| One-parameter equation.

and are strictly appropriate for the 0°K isotherm. These should be considered as representing only an empirical pressure-density relation for other thermodynamic curves, such as adiabats. The application of equations 6 and 7 to oxides has recently been discussed by *Anderson and Anderson* [1970]. For application to forsterite, more serious objections to the use of equations 6 and 7 are that the forces between ions are non-central and the Si⁴⁺-O⁻ bond in tetrahedrally coordinated silicates is known to be partially ionic [*Bloomfield et al.*, 1961]. Hence at low pressure the $(r_0/r)^4$ term, which reflects Coulombic type attractions, is likely to be in error, and at high pressures bond asymmetries will result in poor representations of the equation of state.

The suitability of different equations of state to the fitting of shock-wave data for a silicate, such as forsterite, is demonstrated by results summarized in Table 3. The zero-pressure values of K_S and $(\partial K_S/\partial P)_S$ are known from the ultrasonic measurements of *Kumazawa and Anderson* [1969] and *Graham and Barsch* [1969] to be 1.29 Mb and 5.0 to 5.4, respectively. Except for solutions 1 and 2, obtained with the Birch-Murnaghan equation, large variations from the ultrasonic values are indicated in Table 3. Although the Birch-Murnaghan solutions agree well with the zero-pressure bulk modulus, the values of $(\partial K_S/\partial P)_S$ of 6.6 and 6.4 for $n = 0$ and $n = 1$ (equation 4) obtained from the shock data are too high. The agreement with the ultrasonic data is better, however, than the agreement obtained from the Born-Lande or Born-Mayer two-parameter equation. The closer agreement of K_S and $(\partial K_S/\partial P)_S$ obtained by fitting the shock-wave data with the Eulerian Birch-Murnaghan equation is believed to be not accidental, but rather to result from the fact that the exponents in this equation are fixed, which has the effect of more seriously constraining the curvature of the resulting pressure-density curve. As shown by solutions 8-10, when the value of $(\partial K_S/\partial P)_S$ is fixed at the ultrasonic value, the Born-Lande and Born-Mayer equations of state give values of K_S that are in closer agreement with the ultrasonic values. It is clear from Table 3 that K_S and $(\partial K_S/\partial P)_S$ are affected only slightly by variation in the assumed value of n in equation 4.

The temperatures along the Hugoniot of Mg₂SiO₄ in the forsterite phase, which we calcu-

lated (e.g., 160°C at 280 kb) for the pressure range of the data are considerably lower than the temperatures of general geophysical interest. Shock temperatures were calculated by the method outlined in Ahrens *et al.* [1969]. The relatively low temperatures obtained result from the low value of the Grüneisen parameter and the incompressibility of forsterite. The pressure-density states of a series of adiabats centered at zero-pressure and high temperatures were calculated by using the present Hugoniot data and the zero-pressure specific heat and the thermal expansion data for forsterite [Clark, 1966]. Details of this type of calculation in which both the temperature variation of the specific heat and thermal expansion coefficient are taken into account are described by Davies and Ahrens (unpublished data, 1970). The values -0.8 and -0.10 kb/°C for $(\partial K_s/\partial T)_p$ that were calculated in the temperature interval 100°–200°C are somewhat lower than the values -0.14 and -0.17 kb/°C that were calculated for the interval 200°–400°C. The values for the interval 200°–400°C are in good agreement with the ultrasonic values of -0.150 and -0.176 kb/°C that were reported by Kumazawa and Anderson [1969] and Graham and Barsch [1969]. The values of K_s and $(\partial K_s/\partial P)_s$ for the adiabat centered at zero pressure and 1200°C are given in Table 3.

Acknowledgments. We appreciate the contributions of W. Miller, F. Lehner, and J. L. Blayney to the design of the electronics and optics, the help of D. Tanner, D. Johnson, H. P. Liu, and Rex Gibbons with the experiments, and the assistance of G. Davies with the reduction of data. We thank D. L. Anderson for reviewing the manuscript.

This research was supported by the National Science Foundation under grant GA 1630.

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(Received April 7, 1970.)