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LOS ALAMOS SCIENTIFIC LABORATORY OF THE UNIVERSITY OF CALIFORNIA O LOS ALAMOS **NEW MEXICO**

THE HYDRODYNAMIC HOT SPOT AND SHOCK INITIATION OF HOMOGENEOUS EXPLOSIVES

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LOS ALAMOS SCIENTIFIC LABORATORY OF THE UNIVERSITY OF CALIFORNIA LOS ALAMOS NEW MEXICO

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THE HYDRODYNAMIC HOT SPOT AND SHOCK INITIATION OF HOMOGENEOUS EXPLOSIVES

by

Charles L. Mader



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ABSTRACT

The shock initiation of nitromethane, liquid TNT, and single-crystal PETN has been computed using one-dimensional numerical reactive hydrodynamics and realistic equations of state. The computed pressures and velocities as functions of time agree with the experimental values to within the probable experimental error.

The initiation of shocked nitromethane by spherical hydrodynamic hot spots was investigated. When a hydrodynamic hot spot has decomposed, it sends a shock wave into the undetonated explosive and heats it. What occurs thereafter depends upon the initial strength of the shock wave and how well it is supported from the rear. Whether the explosion of the hot spot propagates to the rest of the fluid or not depends primarily on the initial size of the hot spot.

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I. INTRODUCTION

The shock initiation of homogeneous explosives has been extensively investigated by Campbell, Davis, and Travis¹. They observed that the detonation of the heated, compressed explosive begins at the interface, where the explosive has been hot the longest, after an induction time which is a decreasing function of the shock strength. The detonation proceeds through the compressed explosive at a velocity greater than the steady state velocity in uncompressed explosive, overtaking the initial shock and overdriving the detonation in the unshocked explosive.

The reactive hydrodynamics of shock initiation has been investigated by Hubbard and Johnson² and by Enig³. The results of their calculations qualitatively resembled the experimental observations of Campbell, Davis, and Travis¹. However, quantitative agreement between the experimental and computed pressures and velocities as a function of time was not obtained because of the unrealistic equations of state. We have attempted to use the best numerical reactive hydrodynamics and as realistic equations of state as are available to determine if we could obtain quantitative agreement with the experimental observations.

Since any realistic equation of state is somewhat involved, it was necessary to use electronic computers and considerable amounts of machine time to obtain the desired quantitative agreement.

Having found this comparison reasonably successful, we investigated the hydrodynamics of hot spots. Experimentally 1 it has been observed that

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0.05 to 0.02 cm radius bubbles in shocked nitromethane caused reacting hot spots in the liquid nitromethane that either exploded and propagated or failed to propagate in times of the order of 0.1 μ sec. Evans, Harlow, and Meixner⁴ have computed that the interaction of a shock with a bubble in nitromethane produces hot spots in the liquid nitromethane of the same order of magnitude in size as the original bubbles. Thus, reacting hot spots of the order of 0.05 to 0.01 cm radius should explode and propagate or fail to propagate in times of the order of 0.1 μ sec.

Zinn⁵ has shown that the thermal conductivity model hot spot does not result in propagation of an explosion. He has also shown that chemical reaction must be well under way by the end of the interval $0.04 (R^2)/k$, or cooling by heat conduction prevents the hot spot from exploding. The thermal diffusivity k, is $\lambda/\rho c$, where λ is heat conductivity, ρ is density, and c is heat capacity. A reasonable value for k for nitromethane is 0.001. To fail in 0.1 μ sec, a thermal conductivity hot spot must have a radius smaller than 5×10^{-5} cm. Quenching of a 0.01 cm radius hot spot takes 4000 μ sec. It is immediately apparent that the thermal conductivity hot spot is unsatisfactory as a model to explain the experimental observations¹ and computed shock interaction results⁴ for hot spots in shocked nitromethane. A new model is required, and we propose that the hydrodynamic hot spot model is at least qualitatively satisfactory.

The introduction of kinetics into a numerical hydrodynamic calculation raises the question of how satisfactorily the calculation reproduces the interaction of kinetics and hydrodynamics. Presently, it is necessary to perform numerical experiments and compare the results with known steady state results. For example, comparison of gamma-law Taylor waves and matched pressures across boundaries with numerical results indicate that the kinetics does not disturb the nonreactive portion of the hydrodynamics. The interaction

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of kinetics and hydrodynamics appears to be reasonably well, but certainly not exactly, reproduced by our numerical scheme.

II. THE COMPUTATIONAL METHOD

The hydrodynamic equations are presented and discussed in Appendix A. They were solved using the numerical difference technique described by Fromm⁶. The equation of state is described in Appendix B. The Fickett and Wood beta equation of state $\frac{7}{100}$ form was chosen for the detonation products off the Chapman-Jouguet (C-J) isentrope. The pressure, volume, temperature, and energy values along the C-J isentrope were computed using the BKW codes⁸. The BKW codes use the Kistiakowsky-Wilson equation of state as modified by Cowan and Fickett⁹. The Grüneisen equation of state¹⁰ was chosen for the undetonated explosive off the experimental Hugoniot. Pressures and volumes on the Hugoniot are available from experimental data. and Hugoniot temperatures were calculated using the technique of Walsh and Christian¹⁰. The equation of state for mixtures of condensed explosive and detonation products was computed assuming pressure and temperature equilibrium. The use of a first order reaction of the Arrhenius type described $Z_{inn}^{11,12}$ has shown that the experiin Appendix A is apparently realistic. mentally observed thermal initiation of explosives can be explained for the common explosives using Arrhenius first order reaction kinetics for adiabatic explosion times of hours down to 10 μ sec or less. Our computed explosion times are in the 1- μ sec range; thus the Arrhenius type kinetics is a reasonable choice.

The boundary conditions for the shock initiation calculations were chosen to approximate the experimental conditions that the plate ceases to push soon after initiation occurs at the plate-explosive interface; however, the essential features of the calculation are not changed if one permits the particle velocity

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of the rear boundary to remain at its initial value throughout the calculation. The boundary conditions for the hot spot calculations were the usual ones for continuous systems.

The calculations were performed on an IBM 7090 electronic computer using the "SIN" code. A thousand mesh points could be used in the SIN code. Most of the computations in this report were performed with 500 mesh points. The graphs were prepared directly from computer output using a Stromberg-Carlson SC-4020 Microfilm Recorder.

III. RESULTS OF THE SHOCK INITIATION CALCULATIONS

The pressure-distance profiles for the shock initiation of nitromethane are shown in Figure 1. The shock travels into the explosive, causing shock heating and resultant chemical decomposition. Explosion occurs at the rear boundary, since it has been hot the longest. A detonation develops which has a C-J pressure and velocity characteristic of the explosive at the shock pressure and density. The detonation wave overtakes the shock wave and then decays to the normal density C-J pressure and velocity.

The computed and experimental data for the shock initiation of nitromethane, liquid TNT, and single-crystal PETN is shown in Table I.

The computed induction time (time to explosion) is primarily dependent upon the shock temperature. We probably know the shock temperature only to $\pm 100^{\circ}$. In the case of nitromethane and PETN, we were lucky, and the computed shock temperatures were remarkably close to those required to reproduce the experimental induction times. However, we were not so lucky in the case of liquid TNT. The computed shock temperature was about 100° too high (1200 instead of 1100). We recomputed the shock temperatures using a heat capacity of 0.48 cal/g/°C instead of the literature value of 0.383 cal/g/°C, so as to obtain a shock temperature that would give the experimental induction time.



Fig. 1. Shock initiation of nitromethane by a 92 kbar shock.

Table I

SHOCK INITIATION OF NITROMETHANE, LIQUID TNT, AND PETN

Parameter	Input	Calculated	Experimental	BKW					
NITROMETHANE									
Particle Velocity (cm/µsec)	0.171		0.171 ± 0.01						
Initial Temperature (°K)	300		300						
Shock Pressure (mbar)		0.087	0.086 ± 0.005						
Shock Velocity (cm/ μ sec)		0.445	0.450						
Induction Time (µsec)		1.34	1.4 ± 0.7						
Det. Vel. + Part. Vel.* (cm/µsec)		1.044	1.022 ± 0.03						
Det. Vel. (Steady State)* (cm/ μ sec)		0.860	0.851 ± 0.02	0.847					
Pressure* (mbar)		0.350	0.300	0.355					
	LIQUID	TNT							
Particle Velocity (cm/µsec)	0.176		0,176						
Initial Temperature (°K)	358.1		358,1						
Shock Pressure (mbar)		0.126	0,125						
Shock Velocity (cm/µsec)		0.490	0.490						
Induction Time** (µsec)		0.68	0.70						
Det. Vel. + Part. Vel.* (cm/ μ sec)		1.034	1.10 ± 0.1						
Det. Vel. (Steady State)* (cm/µsec)		0.858	0.925 ± 0.1	0.850					
Pressure* (mbar)		0.460		0.466					
	SINGLE-CRYS	TAL PETN							
Particle Velocity (cm/ μ sec)	0.111		0.117						
Initial Temperature (°K)	298,17		298,17						
Shock Pressure (mbar)		0.115	0,112						
Shock Velocity (cm/ μ sec)		0.564	0,588						
Induction Time (µsec)		0.337	0.300						
Det. Vel. + Part. Vel.*** (cm/ μ sec)		0.985	1.090 ± 0.1						
Det. Vel. (Steady State)*** (cm/ μ sec)		0.870	0.973 ± 0.1	0.882					
Pressure*** (mbar)		0.481		0.473					

*In compressed explosive.

 $\ensuremath{^{**}C_V}$ adjusted to give agreement between computed and experimental induction time.

***In compressed explosive, computed using small cell size.

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The results of the computation of the shock initiation of PETN are of special interest. Our initial calculations were satisfactory until the shocked explosive detonated; then we obtained a flat-topped pressure profile with a very large detonation velocity. Because of the large reaction rate for PETN, the undetonated explosive next to the detonation front was shocked to a value well below the C-J volume when it completely decomposed. The pressure increased, and the velocity decreased, with decreasing mesh size until the C-J pressure and velocity was obtained. Thereafter the pressure and velocity did not change with further decrease in mesh size. It can be shown that decreasing Z (Arrhenius frequency factor) by a factor of 10 is approximately equivalent to decreasing ΔX and Δt by a factor of 10. Thus, we could keep the mesh size constant and decrease Z after the explosion started, or we could decrease the mesh size and keep Z constant. For the PETN calculation it is necessary to decrease the mesh size used in our calculations for nitromethane by a factor of 10^7 . Satisfactory results may also be obtained by decreasing Z by 10^7 after the detonation has started, using the same mesh size as that used for nitromethane.

We conclude that the computed shock initiation of nitromethane, liquid TNT, and single-crystal PETN agrees qualitatively and quantitatively with the experimental data within experimental error.

IV. HYDRODYNAMIC HOT SPOT CALCULATIONS

The SIN code was used to investigate the propagation of hydrodynamic hot spots. The concept was first suggested by Francis Harlow of Group T-3.

Campbell, Davis, and Travis¹ have reported that "bubbles" about 0.7 mm in diameter cause hot spots in shocked nitromethane that will propagate, while bubbles about 0.4 mm in diameter cause hot spots that will not propagate. Evans, Harlow and Meixner⁴ have investigated the interaction of a shock wave with a bubble in nitromethane. The collapse of the bubble was accompanied by the generation of high temperatures in the liquid just above where the bubble had been. While calculational fluctuations precluded determination of the exact temperature profiles, the size of the hot region was of the same order of magnitude as the bubble. Increasing the bubble size results in increased hot spot size. Thus, a hot spot in shocked nitromethane may or may not propagate, depending primarily upon its size. When a hot spot has decomposed it sends a shock wave into the undetonated explosive and heats it. What occurs thereafter depends upon the initial strength of the shock wave and how well it is supported from the rear.

A. Temperature Hot Spots

The model for a temperature hot spot is a volume of uniform-density explosive containing a centered sphere of explosive at a higher energy than that of the surrounding medium. Thus, we have a hot spot of considerably higher temperature and only slightly higher pressure than the rest of the explosive.

Figure 2 shows the pressure-distance plots for a 0.292 cm radius hot spot in shocked nitromethane. Initially, at the boundary between the hot spot and the shocked nitromethane, a small shock is sent into the shocked nitromethane, and a rarefaction is sent back into the hot spot. At 0.03 μ sec, part of the hot spot explodes and propagates through the remainder of the hot spot, which has been cooled by the rarefaction. A strong shock goes into the undetonated nitromethane, and a rarefaction goes back into the detonation products. The strongly shocked nitromethane at the hot spot boundary explodes at about 0.06 μ sec, and the detonation propagates through the rest of the nitromethane. The hot spot propagates at a velocity of 0.856 cm/ μ sec, which is the computed equilibrium detonation velocity of the shocked nitromethane. The experimental detonation velocities of the Campbell, Davis,

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Fig. 2. Development of a detonation in shocked nitromethane (94.7 kbar, 1200°K) from a 0.292 cm radius spherical temperature hot spot (1404°K).

and Travis¹ hot spots in shocked nitromethane are 0.8 ± 0.1 cm/ μ sec.

Figure 3 shows in considerable detail the mechanism of propagation of a 0.06 cm radius hot spot. Initially, at the hot spot boundary, a small shock is sent into the shocked nitromethane, and a rarefaction is sent back into the hot spot. The pressure of the hot spot increases as a result of chemical reaction. At 0.0288 μ sec, about 0.045 cm of the original hot spot explodes. At 0.035 μ sec, the entire hot spot has exploded. A shock is sent into the undetonated explosive, and a rarefaction is sent into the detonation products. The undetonated explosive at the hot spot boundary does not explode until 0.08 μ sec, or after an induction time of 0.045 μ sec. In 0.1 μ sec, the detonation is propagating at full velocity and pressure.

Figure 4 shows the pressure-distance profiles for a 0.0292 cm radius hot spot in shocked nitromethane. The behavior computed is essentially the same as for the larger hot spots except that the divergence of shock waves and the convergence of the rarefactions are sufficient to lower the strength of the shock at the hot spot interface so that propagation does not occur. The nitromethane at the boundary of the hot spot is cooling at 0.06 μ sec, having a temperature of 1393°K, which is less than the initial hot spot temperature of 1404°K. At 0.08 μ sec, the temperature of the boundary nitromethane has decreased to 1370°K.

Figure 5 shows the failure of a 0.27 cm radius hot spot in unshocked nitromethane. The pressure of a constant volume detonation in nitromethane initially at 1450°K is 81.5 kbar. A rarefaction travels into the detonation products, and a shock travels into the undetonated nitromethane. The shock temperature is sufficiently low that no reaction occurs, and the hot spot does not propagate. However, if an explosive has a constant volume detonation pressure that causes a shock in the undetonated explosive strong enough to give adequate shock heating, the hot spot propagates. An example is



Fig. 3. Development of a detonation in shocked nitromethane (94.7 kbar, 1200°K) from a 0.06 cm radius spherical temperature hot spot (1404°K).



Fig. 3 Continued



Fig. 4. Failure of a 0.0292 cm radius spherical temperature hot spot (1404°K) to develop into a detonation in shocked nitromethane (94.7 kbar, 1200°K).



Fig. 5. Failure of 0.27 cm radius spherical temperature hot spot (1450°K) to develop into a detonation in unshocked nitromethane (300°K).

single-crystal PETN. The computed constant volume detonation pressure is 163 kbar for a PETN hot spot initially at 1500°K. A shock strong enough to propagate the detonation may be sent into the undetonated explosive.

Since we do not consider heat conduction in our calculations, it is of interest to determine if it might have any effect on our results for hot spots. $Zinn^5$ has shown that chemical reaction must be well under way by the end of the interval $t = 0.04 (R^2)/k$, or heat conduction will prevent the hot spot from exploding. If we consider a hot spot with radius R of 0.06 cm, then t is 0.144 seconds. This is about 10⁶ larger than the maximum times we are considering in our hot spot hydrodynamics.

The characteristics of temperature hot spots may now be described. Temperature hot spots of 0.06 cm radius and larger in shocked (90 kbar) nitromethane will propagate if the initial hot spot temperature is high enough that the hot spot explodes before the rarefaction arrives at its center. The hot spot might explode even after the rarefaction arrived at its center, since the rarefaction is weak.

Temperature hot spots of 0.03 cm radius and smaller in shocked (90 kbar) nitromethane will not propagate whether they explode or fail to explode before the rarefaction reaches their centers. The shock is not adequately supported from the rear.

As the pressure and temperature of the shocked nitromethane decrease, the critical size of the hot spots increases. The shock temperature in the undetonated explosive is less and, hence, must be maintained for a longer time if the boundary explosive is to explode. A larger hot spot is required to maintain the shock temperature for a longer time.

As the pressure and temperature of the shocked nitromethane increase, the critical size of the hot spot slowly decreases. The shock temperature in the undetonated explosive is higher and does not need to be maintained so long for explosion to occur; however, the strong divergence of the shock wave and convergence of the rarefaction at the small diameters minimize the effect.

For nitromethane, a shock pressure and temperature exist below which any hot spot of reasonable size will not propagate by this mechanism. Large hot spots that explode in unshocked explosive may or may not propagate, depending upon the constant volume detonation pressure and the magnitude of the shock sent into the undetonated explosive.

B. Pressure Hot Spots

The model for a pressure hot spot is a centered sphere whose density and energy have been increased to a Hugoniot value larger than the density and energy of the surrounding explosive. This results in a hot spot of considerably higher pressure and temperature than the rest of the explosive.

Figure 6 shows the pressure-distance profiles for a 0.03 cm radius pressure (112 kbar, 1395°K) hot spot in shocked nitromethane (94.7 kbar, 1230°K). The rarefaction travels into the hot spot until, at 0.032 μ sec, about 0.013 cm of the hot spot explodes. The rarefaction has cooled the hot spot sufficiently that it does not explode almost instantaneously, as it does in the case of the temperature hot spot. A shock is sent into the rest of the undecomposed hot spot, and at 0.034 μ sec the shocked hot spot explosive decomposes and builds up to a C-J detonation pressure and velocity characteristic of the high density nitromethane. Upon arriving at the hot spot boundary, the shock pressure is so high that the detonation front continues unchanged.

Figure 7 shows the pressure-distance profiles for a 0.0175 cm radius pressure hot spot in shocked nitromethane. At 0.032 μ sec, the first 0.0005 cm explodes; however, the converging rarefaction is sufficiently strong that the hot spot fails to propagate.

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Fig. 6. Development of a detonation in shocked nitromethane (94.7 kbar, 1230°K) from a 0.03 cm radius spherical pressure hot spot (112 kbar, 1395°K).



Fig. 7. Failure of a 0.0175 cm radius spherical pressure hot spot (112 kbar, 1395°K) to develop into a detonation in shocked nitromethane (94.7 kbar, 1230°K).

The characteristics of pressure hot spots may now be described. Pressure hot spots of 0.03 cm radius and larger in shocked (90 kbar) nitromethane will propagate if the initial hot spot temperature is high enough that the hot spot explodes before the rarefaction arrives near its center.

Pressure hot spots of 0.015 cm radius in shocked (90 kbar) nitromethane at temperatures of 1400°K and lower fail because the rarefaction arrives at the center of the hot spot before it explodes.

Pressure hot spots of about 0.0175 cm radius in shocked (90 kbar) nitromethane at 1400°K explode but fail to propagate because of the strong convergence of the rarefaction.

C. Hydrodynamic Hot Spot Summary

Over a considerable range of hot spot temperatures, the critical radius of a temperature hot spot in shocked (90 kbar) nitromethane is 0.03 to 0.06 cm. Over a considerable range of hot spot temperatures and pressures, the critical radius of a pressure hot spot is 0.015 to 0.03 cm. The computed critical radii of the hot spots are of the same order of magnitude as one expects from bubbles of 0.02 cm radius that were observed to form hot spots that fail, and bubbles of 0.035 cm radius that were observed to form hot spots that propagate. The hydrodynamic hot spot appears to be a satisfactory model for explaining the experimental observations.

V. CONCLUSIONS

The experimentally observed shock initiation of homogeneous explosives may be quantitatively reproduced using one-dimensional, numerical, reactive hydrodynamics, and realistic equations of state. A shock travels into the explosive, causing shock heating. Explosion occurs at the rear boundary, since chemical decomposition has been occurring there longest. A detonation develops and overtakes the shock wave.

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The hydrodynamic hot spot is reasonably successful as a model for computing the critical sizes of hot spots in shocked nitromethane. Nitromethane hot spots of a size expected to result from interactions of a shock with the bubbles used experimentally may propagate or fail to propagate within the experimentally observed times of the order of 0.1 μ sec. A hydrodynamic hot spot may fail to explode if the rarefaction reaches its center before it can explode adiabatically. If the hot spot explodes, it sends a shock wave into the undetonated explosive, which heats the explosive. Whether or not it propagates depends upon the initial strength of the shock wave and how well it is supported from the rear.

APPENDIX A

THE HYDRODYNAMIC EQUATIONS

The Lagrangian conservation equations in one dimension for slabs, cylinders, and spheres are:

> $\frac{\mathrm{d}U}{\mathrm{d}t} = -\mathrm{R}^{\alpha-1} \frac{\mathrm{d}P}{\mathrm{d}M}$ Conservation of momentum $V = R^{\alpha - 1} \frac{dR}{dM}$ Conservation of mass $\frac{\mathrm{d}\mathbf{E}}{\mathrm{d}\mathbf{t}} = -\frac{\mathrm{d}\mathbf{P}\mathbf{U}\mathbf{R}^{\alpha-1}}{\mathrm{d}\mathbf{M}}$

where $E = I + 0.5U^2$

$$\frac{dR}{dt} = U$$
$$dM = \rho_0 r^{\alpha - 1} dr = \rho R^{\alpha - 1} dR$$

where dM = element of mass per unit angle The chemical reaction equation is

$$\frac{\mathrm{dW}}{\mathrm{dt}} = \mathrm{ZW} \, \mathrm{e}^{-\mathrm{E}*/\mathrm{R}_{\mathrm{g}}\mathrm{T}}$$

The nomenclature is identical with that given at the end of this

Appendix, except that R is Eulerian distance, r is Lagrangian distance, and E is total energy. The treatment of total energy is further discussed in Appendix B.

The differencing of the differential equations was investigated using the Lax^{13} , Harlow and Daly¹⁴, and Fromm⁶ techniques.

The Lax scheme has an effective viscosity which varies inversely with the time increment. This scheme must be run with the largest time increment possible (the Courant Δt) to prevent serious diffusion. In any calculation involving both undetonated shocked explosive and its detonation products, the smaller time increment associated with the higher sound speed of the detonation products must be used in order to satisfy the Courant condition. The time increment is too small to give a reasonable effective viscosity for the undetonated shocked explosive. In addition, the calculation suffers from small oscillations, and in this region the two effects can result in a second explosion ahead of the main detonation wave. A solution to the problem can be obtained by using a differencing scheme in which the artificial viscosity is not a function of the time increment. Numerous such schemes are available; however, most of them suffer from fluctuations at the front of the shock wave which render them useless for systems in which the reaction rates have an exponential dependence on temperature. This problem was studied by $Zovko^{15}$ and Enig, and they chose the Lax scheme to avoid the fluctuations. They, however, were not attempting the fine resolution described here and therefore did not observe the other difficulties associated with the Lax scheme.

We investigated two attractive difference schemes using the artificial viscosity method to determine if we could dampen the fluctuations and still retain the desirable characteristics.

The first method investigated was the cell-centered method of Harlow and Daly 14 . In this method the parameters are cell-centered, the energy is

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conserved locally, entropy is changed only by the viscosity, and the system is accurate and stable to infinitesimal fluctuations. However, it was not possible to dampen the fluctuations sufficiently using any of the usual forms for the artificial viscosity¹⁶.

The second difference scheme we investigated was that described by Fromm⁶. The Richtmyer-von Neumann type of viscosity did not sufficiently damp the small fluctuations associated with the Fromm difference equation. The Landshoff type of viscosity¹⁶ required careful adjustment of the arbitrary constant, but at best was still not successful. However, the particle in cell (PIC) type of viscosity¹⁶ was successful in damping the fluctuations.

In the Fromm type of difference equations, the pressure, temperature, energy, and specific volume of the cells are considered to be located at the centers of mass of the elements, and the particle velocity is considered to be located at the boundary between the cells.

The energy is conserved locally in the Fromm type of difference equations. This may be shown by summing the total energy in Equation (6) over all the mesh points, resulting in

$$\sum_{j=1}^{j=N} \left[E_j^{n+1} - E_j^n \right]$$

where $E = (I + 0.5U^2)M$. All the terms of the sum cancel except at the end points. Thus, the difference of the total energy of the space at two consecutive times is just the flux at the two boundaries.

In the Fromm type of difference equations, one finds internal consistency for the case of slabs. For example, from Equations (4) and (5), one may derive the difference form of the conservation equation dV/dt = dU/dM.

One can show, by approximating the parameters as Taylor expansions, that the Fromm difference equations formally approach the differential equations as the time increment and cell mass decrease. Using the Fromm difference equations, one can show, by substituting the difference equations into the thermodynamic equation TdS = dI + PdV, that entropy is not conserved in the absence of viscosity.

Fromm⁶ has shown that the difference equations are conditionally stable to infinitesimal fluctuations, using the von Neumann method, in which the perturbation is expanded in a Fourier series and a given Fourier component is examined.

Thus, the difference equations are sufficiently accurate and stable to infinitesimal fluctuations, and energy is conserved locally. The equations can be shown not to conserve entropy in the absence of viscosity. Numerically we have shown that the difference equations with the PIC viscosity are conditionally stable.

I. THE SIN DIFFERENCE EQUATIONS FOR SLABS, CYLINDERS, OR SPHERES

A. Initial Conditions

$$R_{j+\frac{1}{2}} = J\Delta R \tag{1}$$

where

$$j = 0, 1, 2, \dots$$

$$R_{-\frac{1}{2}} = U_{\text{piston}}(\Delta t)$$

$$M_{j} = \rho_{0} \left(R_{j+\frac{1}{2}} \right)^{\alpha-1} \Delta R$$
(2)

B. Conservation Equations

$$U_{j+\frac{1}{2}}^{n+\frac{1}{2}} = U_{j+\frac{1}{2}}^{n-\frac{1}{2}} + \frac{(\Delta t) \left(R_{j+\frac{1}{2}}^{n} \right)^{\alpha-1}}{0.5 \left(M_{j} + M_{j+1} \right)} \left[\left(P_{j}^{n} - P_{j+1}^{n} \right) + \left(q_{j}^{n} - q_{j+1}^{n} \right) \right]$$
(3)

$$\begin{aligned} \mathbf{R}_{j+\frac{1}{2}}^{n+1} &= \mathbf{R}_{j+\frac{1}{2}}^{n} + \mathbf{U}_{j+\frac{1}{2}}^{n+\frac{1}{2}} \left(\Delta t \right) \end{aligned} \tag{4} \\ \mathbf{V}_{j}^{n+1} &= \left(\mathbf{R}_{j+\frac{1}{2}}^{n+1} \right)^{\alpha-1} \frac{\left(\mathbf{R}_{j+\frac{1}{2}}^{n+1} - \mathbf{R}_{j-\frac{1}{2}}^{n+1} \right)}{\mathbf{M}_{j}} \end{aligned} \tag{5} \\ \mathbf{I}_{j}^{n+1} &= \mathbf{I}_{j}^{n} + \frac{\left(\Delta t \right)}{\mathbf{M}_{j}} \left\{ \left[\frac{\mathbf{M}_{j} \mathbf{P}_{j-1}^{n} + \mathbf{M}_{j-1} \mathbf{P}_{j}^{n}}{\mathbf{M}_{j} + \mathbf{M}_{j-1}} + \mathbf{0.5} \left(\mathbf{q}_{j}^{n} + \mathbf{q}_{j-1}^{n} \right) \right] \mathbf{U}_{j-\frac{1}{2}}^{n+\frac{1}{2}} \left(\mathbf{R}_{j-\frac{1}{2}}^{n+1} \right)^{\alpha-1} \\ &- \left[\frac{\mathbf{M}_{j+1} \mathbf{P}_{j}^{n} + \mathbf{M}_{j} \mathbf{P}_{j+1}^{n}}{\mathbf{M}_{j+1}} + \mathbf{0.5} \left(\mathbf{q}_{j}^{n} + \mathbf{q}_{j+1}^{n} \right) \right] \mathbf{U}_{j+\frac{1}{2}}^{n+\frac{1}{2}} \left(\mathbf{R}_{j+\frac{1}{2}}^{n+1} \right)^{\alpha-1} \\ &+ \frac{1}{8} \left[\left(\mathbf{U}_{j+\frac{1}{2}}^{n-\frac{1}{2}} + \mathbf{U}_{j-\frac{1}{2}}^{n-\frac{1}{2}} \right)^{2} - \left(\mathbf{U}_{j+\frac{1}{2}}^{n+\frac{1}{2}} + \mathbf{U}_{j-\frac{1}{2}}^{n+\frac{1}{2}} \right)^{2} \right] \end{aligned} \tag{6}$$

C. Chemical Reaction Equation

$$W_{j}^{n+1} = W_{j}^{n} - \Delta t Z W_{j}^{n} e^{-E^{*}/R_{g}T_{j}^{n}}$$
(7)

where $1 \ge W \ge 0$

D. Equation of State

 P_j^{n+1} , T_j^{n+1} , and C_j^{n+1} are computed using the HOM equation of state subroutine from V_j^{n+1} , I_j^{n+1} and W_j^{n+1} .

E. Artificial Viscosity

$$q_{j}^{n+1} = \frac{K}{V_{j}^{n+1}} (0.5) \left(U_{j-\frac{1}{2}}^{n+\frac{1}{2}} + U_{j+\frac{1}{2}}^{n+\frac{1}{2}} \right) \left(U_{j-\frac{1}{2}}^{n+\frac{1}{2}} - U_{j+\frac{1}{2}}^{n+\frac{1}{2}} \right)$$
(8)

$$\text{if } \left(U_{j-\frac{1}{2}}^{n+\frac{1}{2}} - \ U_{j+\frac{1}{2}}^{n+\frac{1}{2}} \right) \text{is positive, otherwise } q_{j}^{n+1} = 0.$$

F. Time Increment

II.

NOMENCLATURE

One may use a constant Δt or compute it for each cycle.

$$\Delta t = (CK)\Delta R \frac{V_j^{n+1}}{V_0 C_j^{n+1}}$$
(9)

G. Boundary Conditions for Shock Sensitivity Calculations

$$U_{-\frac{1}{2}}^{n} \text{ and } U_{-\frac{1}{2}}^{n+1} = U_{\text{piston}} \text{ where boundary is at } j = 0$$

if $W_{j+\frac{1}{2}}^{n} > 0.5$, otherwise = 0
$$q_{-1}^{n} = q_{1}^{n}$$
$$P_{-1}^{n} = P_{1}^{n}$$

R_{-\frac{1}{2}} is computed using Equation (4)

 $\alpha = 1$ for slabs, 2 for cylinders, and 3 for spheres $C = \text{sound speed in cm/}\mu\text{sec}$ CK = constant of about 0.1 $\Delta t = \text{time in }\mu\text{sec}$ $E^* = \text{activation energy}$ I = internal energy in mbar-cc/g j = net point of Lagrangian netK = constant of about 2

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- M = mass
- n = time cycle
- P = pressure in mbar
- q = artificial viscosity
- R = radius in cm

- ρ_0 = initial density in g/cc
- T = temperature in[°]K
- $U = particle velocity in cm/\mu sec$
- V = volume in cc/g
- $V_0 = 1/\rho_0$
- W = mass fraction of the undecomposed explosive
- Z = frequency factor

APPENDIX B

THE EQUATION OF STATE

Initially we investigated the equation of state used by Enig³. He used the Tait equation for the undetonated explosive and the gamma-law equation of state for the detonation products. We adjusted the parameters in the Tait equation of state to give pressures and volumes in reasonable agreement with the experimental Hugoniot values over the range of interest. The parameters were also adjusted to give temperatures that agreed with the values computed using the Walsh and Christian technique¹⁰.

The growth of detonation from an initiating shock was computed using the Tait and gamma-law equation of state for nitromethane, liquid TNT, and PETN. The computed detonation velocities in the compressed explosive were less than those experimentally observed. The major cause of this discrepancy is that the detonation velocity is not a function of density with a gammalaw equation of state, but is only a function of energy. Likewise, the C-J temperatures and pressures were in marked disagreement with the observed values.

Thus, a more realistic equation of state was necessary if we were to reproduce the experimental results. The one we used is called the HOM equation of state.

The Fickett and Wood⁷ beta equation of state was chosen as the form

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for the detonation products. The pressure, volume, temperature, and energy values along the C-J isentrope were computed using the BKW codes⁸ which use the Kistiakowsky-Wilson equation of state as modified by Cowan and Fickett⁹.

The Grüneisen equation of state¹⁰ was chosen for the undetonated explosive. Hugoniot pressures and volumes are available from experimental data. Temperatures on the Hugoniots were calculated using the technique of Walsh and Christian¹⁰.

The equation of state for mixtures of condensed explosive and detonation products is derived in Equations (1) through (19) of this Appendix. The equations require a single iteration for each problem, hence the equation of state for mixtures requires considerable computer time.

The equation of state and rate parameters are shown in Table BI. The fits to the detonation products isentrope are good from 1 to 5×10^{-4} mbar. The heat capacity of the detonation products was computed using the BKW C-J parameters (shown in Table BII) for the shocked explosive; Equations (7) through (11) to compute $E - E_i$ and T_i ; and Equation (12) to compute C_V^i . The Hugoniot data used for liquid TNT were reported by $Garn^{17}$. The Hugoniots used for nitromethane and single-crystal PETN were approximated by W. Davis from available experimental data. The fits to the Hugoniot pressures and temperatures are good from 1 to 350 kbar. We used the activation energy and frequency factors of Cook and Taylor¹⁸ for PETN; of Cottrell¹⁹ for nitromethane; and of Zinn¹² for liquid TNT.

I. THE CONDENSED EXPLOSIVE AND DETONATION PRODUCT EQUATION OF STATE

Knowing E, V, and W, we calculate P, T, C.

Table BI

EQUATION OF STATE AND RATE PARAMETERS

Parameter	Nitromethane		Liquid TNT		PETN	
Detonation						
Product		Exp		Exp		Exp
A	-3.0177389	00	-3.5356454	00	-3,4346496	00
в	-2.4028066	00	-2.4873789	00	-2.4498765	00
С	+1.9981153	-01	+2.7864444	-01	+2.1221113	-01
D	+9,9666715	-03	+5.3344652	-02	+6.0697867	-03
Е	-4.4261494	-03	-2.7491409	-02	-5.7596766	-03
K	-1.5762640	00	-1.7153682	00	-1.5954648	00
L	+5.3780073	-01	+5.7252810	-01	+4.8780484	-01
М	+1.2202885	-02	+4.6863081	-02	+4.6779526	-02
N	-2.6100200	-03	+1.0161346	-03	+9.1462157	-04
0	-2.9775278	-04	-7.8496783	-05	-7.5662542	-05
ର୍	+7.7365662	00	+7.6317317	00	+7,4428658	00
R	-5.2925794	-01	-4.7735596	-01	-5.2085389	-01
S	+9.2967764	-02	+1.1398787	-01	+1.0458042	-01
т	+1.2878571	-02	+2.6182746	-02	+1.4337016	-02
U	-5.9641444	-03	-1.5211680	-02	-1.1391835	-02
C_V^{\dagger}	+5.5600000	-01	+5.0000000	-01	+4.6700000	-01
z	+6.0000000	-02	+6.0000000	-02	+1.0000000	-01
Condensed						
Explosive						
Ag	-8.6335924	00	-5.6489393	01	-9.8284779	00
B	-1.6490344	01	-2.8785482	02	+3.4325262	01
Cs	-1.5367448	01	-5,9739302	02	+1.3815194	02
$\tilde{D_8}$	-1.1037889	01	-5.5607584	02	+1.3998427	02
E	-4.2114837	00	-1.9124016	02	+4.5106916	01
F	+6.0426166	00	-1.0712113	00	-1.3504381	02
Gs	+4.4825758	00	-4.8502353	01	-8.0905992	02
$\tilde{H_s}$	+1.4829189	01	-1.2282460	02	-1.7206747	03
Is	+7.6203073	00	-1.3376445	02	-1.5994725	03
Js	-2.4398587	-02	-5.0515048	01	-5,4551108	02
$\gamma_{\rm S}$	+6.8050000	-01	+1,6600000	00	+7.7000000	-01
Cv	+4,1400000	-01	+4.8000000	-01	+2_6000000	-01
v ₀	+8.8652000	-01	+6.9444000	01	+5.6497000	-01
Chemical						
Reaction						
E*	+5.3600000	04	+4.1100000	04	+5.2300000	04
\mathbf{Z}	+4.0000000	08	+1.7100000	07	+1.2600000	17

Table BII

BKW COMPUTED C-J PARAMETERS

Explosive	Density (g/cc)	E ₀ (kcal/mole)	P _{C-J} (mbar)	Det. Vel. (cm/µsec)	^Т С-Ј (°К)	V _{C-J} (cc/mole)	γ
Nitromethane (shocked to	1.128	-14.92	0.151	0.687	2960	15.33	2.53
81 kbar)	1.762	3.91	0.355	0.847	2810	10.83	2.56
Liquid TNT (shocked to	1.447	3.41	0.173	0.675	2960	15.54	2.81
125 kbar)	2.261	87.76	0.466	0.850	2860	10.68	2.50
PETN (shocked to	1.770	-95.5	0.330	0.832	2890	12.59	2.72
112 kbar)	2.210	-47.89	0.473	0.882	2610	10.95	2.63

<u>A. Condensed Explosive</u> $(W = 1, \text{ then } E = E_s, V = V_s)$

Hugoniot pressures and volumes are known experimentally, Hugoniot temperatures are computed using the Walsh and Christian technique¹⁰. The Hugoniot data is fitted by the method of least squares to Equations (1) and (2).

$$\ln P_{\rm H} = A_{\rm s} + B_{\rm s} \ln V_{\rm s} + C_{\rm s} \left(\ln V_{\rm s} \right)^2 + D_{\rm s} \left(\ln V_{\rm s} \right)^3 + E_{\rm s} \left(\ln V_{\rm s} \right)^4$$
(1)

$$\ln T_{\rm H} = F_{\rm s} + G_{\rm s} \ln V_{\rm s} + H_{\rm s} \left(\ln V_{\rm s} \right)^2 + I_{\rm s} \left(\ln V_{\rm s} \right)^3 + J_{\rm s} \left(\ln V_{\rm s} \right)^4$$
(2)

$$E_{\rm H} = \frac{1}{2} P_{\rm H} \left(V_0 - V_{\rm s} \right) \tag{3}$$

$$P_{s} = \frac{\gamma_{s}}{V_{s}} \left(E_{s} - E_{H} \right) + P_{H}$$
(4)

where $\gamma_{s} = V \left(\frac{\partial P}{\partial E}\right)_{V}$

$$\Gamma_{s} = T_{H} + \frac{\left(E_{s} - E_{H}\right)(23,899)}{C_{V}}$$
 (5)

$$C_{s} = \sqrt{-PV_{s} \left[B_{s} + 2C_{s} \ln V_{s} + 3D_{s} \left(\ln V_{s} \right)^{2} + 4E_{s} \left(\ln V_{s} \right)^{3} \right]}$$
(6)

(assumes that the slope of the Hugoniot curve is approximately the same as the slope of the adiabat at the same V_s)

B. Detonation Products
$$(W = 0, \text{ then } E = E_g, V = V_g)$$

The pressure, volume, temperature, and energy values of the detonation products along the C-J isentrope are computed using the Kistiakowsky-Wilson equation of state⁸ and fitted by the method of least squares to Equations (7) through (9).

$$\ln P_{i} = A + B \ln V_{g} + C \left(\ln V_{g} \right)^{2} + D \left(\ln V_{g} \right)^{3} + E \left(\ln V_{g} \right)^{4}$$
(7)

$$\ln E_{i}^{\prime} = K + L \ln P_{i} + M \left(\ln P_{i} \right)^{2} + N \left(\ln P_{i} \right)^{3} + O \left(\ln P_{i} \right)^{4}$$
(8)

$$E_i = E'_i - Z$$

where Z is a constant used to change the standard state to make it consistent with the condensed explosive standard state

$$\ln T_{i} = Q + R \ln V_{g} + S \left(\ln V_{g} \right)^{2} + T \left(\ln V_{g} \right)^{3} + U \left(\ln V_{g} \right)^{4}$$
(9)

$$-\frac{1}{\beta} = R + 2SlnV_{g} + 3T(lnV_{g})^{2} + 4U(lnV_{g})^{3}$$
(10)

$$\mathbf{P}_{\mathbf{g}} = \left(\frac{1}{\beta \mathbf{V}_{\mathbf{i}}}\right) \left(\mathbf{E}_{\mathbf{g}} - \mathbf{E}_{\mathbf{i}}\right) + \mathbf{P}_{\mathbf{i}}$$
(11)

where $\beta = \frac{1}{V} \left(\frac{\partial E}{\partial P} \right)_V$

$$T_{g} = T_{i} + \frac{\left(E_{g} - E_{i}\right)(23,899)}{C_{V}^{\dagger}}$$
(12)

$$C_{g} = \sqrt{-PV_{g} \left[B + 2ClnV_{g} + 3D(lnV_{g})^{2} + 4E(lnV_{g})^{3} \right]}$$
(13)

(assumes that the slope of the adiabat at P_i , V_g is the same as at P, V_g).

C. Mixture of Condensed Explosive and Detonation Products

$$V = WV_{s} + (1 - W)V_{g}$$
⁽¹⁴⁾

$$E = WE_{s} + (1 - W)E_{g}$$
 (15)

$$P = P_g = P_s$$
(16)

$$T = T_g = T_s$$
(17)

One may derive an expression for V or V as a function of E, V, and W by multiplying Equation (5) by (W/C_V) and Equation (12) by $[(1 - W)/C_V]$ and adding the resulting equations. Substituting T for T and T, per Equation (17), and E for $[WE_s + (1 - W)E_g]$, per Equation (15), one obtains

$$T = \frac{23,899}{C_V W + C_V'(1 - W)} \left\{ E - \left[WE_H - E_i(1 - W) \right] + \frac{1}{23,899} \left[T_H C_V W + T_i C_V'(1 - W) \right] \right\}$$
(18)

Equating Equations (4) and (11), one obtains an equation into which Equation (18) may be substituted to obtain

$$P_{H} - P_{i} + \left(\frac{\gamma_{s}^{C}V}{V_{s}} - \frac{C_{V}^{\dagger}}{\beta V_{g}}\right) \left(\frac{1}{C_{V}^{W} + C_{V}^{\dagger}(1 - W)} \left\{ E - \left[WE_{H} + E_{i}(1 - W)\right] + \frac{1}{23,899} \left[T_{H}C_{V}^{W} + T_{i}C_{V}^{\dagger}(1 - W)\right] - \frac{1}{23,899} \left(\frac{\gamma_{s}^{C}V_{H}^{T}}{V_{s}} - \frac{C_{V}^{\dagger}T_{i}}{\beta V_{g}}\right) = 0$$
(19)

With Equation (14), we now have an expression for V_g or V_s as a function of the known parameters E, V, and W. In the HOM code, if W was between 0.999 and 0.7, we iterated on V_s , and if W was less than 0.7, we iterated on V_g . Knowing V_g and V_s , one may compute the other unknowns. The largest sound speed was assumed to be that of the mixture.

II. NOMENCLATURE

C = sound speed in cm/µsec C = heat capacity of condensed explosive in cal/g/°C C' = heat capacity of detonation products in cal/g/°C E = total internal energy in mbar-cc/g

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- P = pressure in mbar
- T = temperature in^oK
- V = total volume in cc/g
- V_0 = initial volume of condensed explosive in cc/g
- W = mass fraction of undecomposed explosive

Subscripts

- g = detonation products
- H = Hugoniot
- i = isentrope
- s = condensed explosive

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