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RESEARCH LABORATORIES

TECHNICAL REPORT

STUDY OF DETONATION BEHAVIOR OF SOLID PROPELLANTS

FIRST QUARTERLY REPORT

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ABSTRACT

The present status of the one-dimensional computer routine is discussed including recent modifications in the grain burning phase of the reaction process. Calculations of minimum pressure for initiation of TNT are presented and compared with experimental values. Use of such comparisons for determining the rate parameters of the ignition process are considered.

Calculations are also presented on the detonation behavior of a typical composite double-base propellant. Minimum pressures for initiation, reaction and pressure profiles through the wave, and wave trajectory, are shown. The material is found to be marginal for support of a steady state detonation.

The theoretical background for the planned experiments on the equation of state of solids is discussed.

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SECTION 1

INTRODUCTION

This report presents the most recent developments in the one-dimensional treatment of detonation. In particular, the procedures for handling the energy distribution between solid and gaseous phases now appear to be satisfactory. Provision has been made for two grain burning processes which may conform either to a surface ignition regressive grain burning model, or to a point ignition progressive grain burning model. It is next planned to incorporate these provisions into the two-dimensional computer routine, and when this is accomplished, it is believed that the procedures for pressure-initiated detonation will be essentially complete. During the remainder of the program it is planned to undertake calculations designed to investigate the conformity between computed and experimental behavior. An important part of this latter step is the work on experimental study of the equation of state of propellant and explosive materials which is presently in progress.

Work on thermal initiation of detonation has been held in abeyance pending the development of operational procedures capable of providing acceptable computer running times.

SECTION 2

MATHEMATICAL DEVELOPMENTS

Two modifications were made to the one-dimensional computer routine during this report period. The first of these is concerned with the computation of energy distribution between solid and gaseous phases in the detonation waves; the second is concerned with adaptation to a point ignition internal grain burning model which, as discussed in the last progress report, appears to be more pertinent to the detonation of homogeneous materials than the external surface burning model presently used.

The problem of energy distribution between phases was discussed to some extent in the last quarterly report. The specific methods used to solve this problem were found to be unsatisfactory because the solution became unstable a short time after ignition of the charge. This instability was subjected to considerable study during the reporting period, and was found to be caused by a coupling of small oscillations in the values of v_s , e_s and p . In order to avoid this behavior, it now appears necessary to compute v_s , v_g by the integration of equations defining the relation between dv_s , dv_g and dv . These are derived as follows.

Considering first the dv_g , the term is assumed to depend upon the sum of two effects, the first to be derivable from the compression or expansion of a mixture of solid and gas, with simultaneous transfer of material from the solid to the gas phase, the second to be derivable from the addition of heat to the gas phase, at constant average volume. (Some changes have been made in the nomenclature used in this program. New definitions are given at the end of the report.)



First, one has:

$$de_s = - p dv_s \quad (1)$$

From the solid equation of state:

$$(p + \beta)v_s = (\gamma_s - 1) e_s + C \quad (2)$$

$$(p + \beta)dv_s + v_s dp = (\gamma_s - 1) de_s = - (\gamma_s - 1) p dv_s$$

Then:

$$\left(\frac{\beta}{p} + \gamma_s \right) p dv_s = - v_s dp$$

$$\frac{d \ln p}{d \ln v_s} = - \left(\frac{\beta}{p} + \gamma_s \right) \quad (3)$$

And for the gas ($\beta = 0$)

$$\frac{d \ln p}{d \ln v_g} = - \gamma_g \quad (4)$$

Also:

$$(1 - f_s) v_g + f_s v_s = \bar{v}$$

$$(1 - f_s) dv_g - v_g df_s + f_s dv_s + v_s df_s = d\bar{v} \quad (5)$$

From (3) and (4)

$$\frac{dv_g}{dv_s} = \frac{v_g}{v_s} \frac{(\gamma_s + \beta/p)}{\gamma_g}$$

Combining with (5) and solving for dv_g ;

$$dv'_g = \frac{d\bar{v} - (v_s - v_g) df_s}{1 - f_s \left[1 - \frac{v_s \gamma_g}{v_g (\gamma_s + \beta/p)} \right]} \quad (6)$$

The effect of heat addition to the gas phase at constant average volume has been discussed in previous reports.¹ Considering the following diagram for one gram of material:

gas phase	solid phase
vol = $v_g f_g$	vol = $v_s f_s$

$$f_g de_g = dE - f_g pdv_g \quad (7)$$

$$f_s de_s = pf_s dv_s \quad (8)$$

From the equation of state:

$$de_s = \frac{1}{\gamma_s - 1} \left[(p + \beta) dv_s + v_s dp \right] \quad (9)$$

$$de_g = \frac{1}{\gamma_g - 1} \left[pdv_g + v_g dp \right] \quad (10)$$

Also:

$$dv_s = - \frac{1 - f_s}{f_s} dv_g \quad (11)$$

Elimination of dp , de_s , de_g and dv_s between equations 7-11 gives

$$dv_g'' = \frac{(\gamma_g - 1) dE}{(1 - f_s) p \gamma_g + (p \gamma_s + \beta) \frac{v_g}{v_s} \frac{(1 - f_s)^2}{f_s}} \quad (12)$$

One then has

$$dv_g = dv_g' + dv_g''$$

By similar methods, the expression for dv_s is found to be:

$$dv_s = \frac{d\bar{v} - (v_s - v_g) df_s}{f_s + (1 - f_s) \frac{v_g}{v_s} \frac{(\gamma_s + \beta/p)}{\gamma_g}} \quad (13)$$

and

$$de_s = -pdv_s$$

In order to treat materials with a self-reactive binder, an additional grain burning equation has been added to the scheme, identical in form with the one presently used but having different rate parameters and associated with a different heat of reaction. This requires an additional term in the calculation of the chemical heat release. The previous differential equations are then differenced to give the following computational procedure:

IGNITION PHASE

Treated as before

GRAIN BURNING PHASE

Set initial conditions:

$$\begin{aligned} e_s^0 &= \bar{e}^0 \\ f_2^0 &= 1 ; \quad f_2^1 = 1 - F_1 \\ f_3^0 &= 1 ; \quad f_3^1 = 1 - F_1 \\ f_4^0 &= 1 - F_1 \end{aligned} \quad (14)$$

$$(f_2^{n+1})^{1/3} = (f_2^n)^{1/3} - \frac{B_2 \Delta t^n p^m}{p^m/w + 1} \exp \left[\frac{-A_2}{e_s^n + Q_2' / \gamma_g^n} \right] \quad (15)$$

$$(f_3^{n+1})^{1/3} = (f_3^n)^{1/3} - \frac{B_3 \Delta t^n p^m}{p^m/w + 1} \exp \left[\frac{-A_3}{e_s^n + Q_3' / \gamma_g^n} \right] \quad (16)$$

$$f_4^{n+1} = 0.8 \exp(-D^{n+1}) + 0.2 \exp(-4D^{n+1}) \quad (17)$$

$$D^{n+1} = (D_s + \frac{D_o}{p^n}) \pi^2 (t^{n+1} - \tau) \quad (18)$$

$$f_s^{n+1} = F_2 f_2^{n+1} + F_3 f_3^{n+1} + F_4 \quad (19)$$

$$v_s^{n+1} = v_s^n + \frac{(\bar{v}^{n+1} - \bar{v}^n) - (v_g^n - v_s^n)(f_s^n - f_s^{n+1})}{f_s^{n+1} + (1 - f_s^{n+1}) \left(\frac{v_g^n}{v_s^n} \right) \left(\frac{\gamma_g^n + \beta/p^n}{\gamma_g^n} \right)} \quad (20)$$

$$v_g^{n+1} = v_g^n + \frac{(\bar{v}^{n+1} - \bar{v}^n) - (v_g^n - v_s^n)(f_s^n - f_s^{n+1})}{1 - f_s \left[1 - \frac{v_s^n}{v_g^n} \frac{\gamma_g^n}{\gamma_s^n + \beta/p^n} \right]} \quad (21)$$

$$+ (\gamma_g^n - 1) \left[(e_s^n - e_g^n)(f_s^n - f_s^{n+1}) + F_2 Q_2 (f_2^n - f_2^{n+1}) + F_3 Q_3 (f_3^n - f_3^{n+1}) + Q_4 (f_4^n - f_4^{n+1}) \right] \\ \frac{(1 - f_s^{n+1}) \left[p^n \gamma_g^n + (p^n \gamma_g^n + \beta) \left(\frac{v_g^n}{v_s^n} \right) \left(\frac{1 - f_s^{n+1}}{f_s^{n+1}} \right) \right]}{}$$

$$\gamma_s^{n+1} = \gamma_{go} + \left[\frac{b}{v_s^{n+1}} + \left(\frac{b}{v_s^{n+1}} \right)^2 \right] (\gamma_{go} - 1) \quad (22)$$

$$\gamma_g^{n+1} = \gamma_{go} + \left[\frac{b}{v_g^{n+1}} + \left(\frac{b}{v_g^{n+1}} \right)^2 \right] (\gamma_{go} - 1) \quad (23)$$

$$\begin{aligned}
 p^{n+1} = & (\gamma_g^{n+1} - 1) \left[e^{-n} - \frac{f_s^{n+1}}{\gamma_s^{n+1} - 1} (\beta v_s^{n+1} - C) + F_2 Q_2 (f_2^n - f_2^{n+1}) + F_3 Q_3 (f_3^n - f_3^{n+1}) \right. \\
 & \left. + Q_4 (f_4^n - f_4^{n+1}) - (p^n + 2q^{n+1/2}) \left(\frac{\bar{v}^{n+1} - \bar{v}^n}{2} \right) \right] / \left[\bar{v}^{n+1} \right. \\
 & \left. + \left(\frac{\gamma_s^{n+1} - \gamma_s^n}{\gamma_s^{n+1} - 1} \right) f_s^{n+1} v_s^{n+1} + (\gamma_g^{n+1} - 1) \left(\frac{\bar{v}^{n+1} - \bar{v}^n}{2} \right) \right] \quad (24)
 \end{aligned}$$

$$\begin{aligned}
 \bar{e}^{n+1} = & \bar{e}^n - (p^n + p^{n+1} + 2q^{n+1/2}) \left(\frac{\bar{v}^{n+1} - \bar{v}^n}{2} \right) + F_2 Q_2 (f_2^n - f_2^{n+1}) \\
 & + F_3 Q_3 (f_3^n - f_3^{n+1}) + Q_4 (f_4^n - f_4^{n+1}) \quad (25)
 \end{aligned}$$

$$e_s^{n+1} = e_s^n + (p^n + p^{n+1} + 2q^{n+1/2}) \left(\frac{v_s^{n+1} - v_s^n}{2} \right) \quad (26)$$

$$e_g = \frac{p^{n+1} v_g^{n+1}}{\gamma_g^{n+1} - 1} \quad (27)$$

Other details of the program remain the same.

The second modification concerning adaptation of the routine to an internal grain burning model has turned out to be relatively straightforward. It was shown in the previous progress report² that the grain burning rate laws for the two models are identical if f is defined in the external burning case as the fraction of material unburned, and in the internal burning case as the fraction of material burned. It follows very readily that conversion from the former to the latter problem can be accomplished by changing the sign of Q_2 , Q_3 , B_2 and B_3 in the input, and redefining the initial values of the f 's as follows:

$$\begin{aligned}
 f_2^0 &= 0 & f_2^1 &= F_1 \\
 f_3^0 &= 0 & f_3^1 &= F_1
 \end{aligned} \quad (28)$$

Integration is terminated when $f \geq 1$.

The proper procedure for f_4 is somewhat in doubt with this model. However, the diffusion rate law is significant here only in the case of composite double-base propellants with grain size of the order of 5 microns or less. If it can be assumed that the centers of ignition are located in the interface between oxidizer grain and binder, then the most reasonable procedure is to use the same geometry for the diffusion process in both models. Although this may falsify to some extent the form of the diffusion reaction profile through a detonation wave, it would not significantly affect the reaction zone length and so should not appreciably alter detonation behavior.

SECTION 3

CALCULATIONS

The one-dimensional program in the form described in Section 2 has been used to examine the detonation behavior of TNT and of a representative composite double-base propellant. Input data for the two materials is as shown in Table I except as noted in the illustrations. The external-regressive grain burning formulation of the grain burning reaction was used in these cases.

Of particular importance in calculating the detonation behavior of real materials is the selection of proper values for the rate parameters A_1 , A_2 , B_1 and B_2 . Data for TNT are available from two sources. The first of these consists of thermal decomposition studies; this source has been reviewed and used by Cook.^{3,4} The second consists of experimental work carried out at the Naval Ordnance Laboratory on the shock pressures necessary for initiation of detonation in TNT.⁵ Such pressures are strongly dependent upon the values used for B_1 and A_1 and the data can therefore be used to evaluate these parameters if the other needed information about the material is available. As has been discussed in previous reports, the assumption that the ignition zone is a high-temperature region in the vicinity of a bubble or void which has been passed over by a shock, leads to the postulate

$$A_1 = A_2/2$$

$$B_1 = B_2$$

TABLE I
INPUT DATA FOR CALCULATION

TNT	Common	Composite Double Base
$C = 795 \text{ cal/gm}$	$a = 4.0 \times 10^{-8}$	$C = 664 \text{ cal/gm}$
$\beta = 50 \text{ kbar}$	$\gamma_o = 1.3$	$\beta = 50 \text{ kbar}$
$Q_1 = 1130 \text{ cal/gm}$	$b = 1.03$	$Q_1 = 622 \text{ cal/gm}$
$Q'_2 = Q_2 = 1130 \text{ cal/gm}$	$D_s = 10^{-4} \text{ cm}^2/\text{sec}$	$Q'_2 = Q_2 = 380 \text{ cal/gm}$
$Q'_3 = Q_3 = 0$	$D_o = 10^{-1} \text{ cm}^2/\text{sec}$	$Q'_3 = Q_3 = 864 \text{ cal/gm}$
$Q_4 = 0$	$w = 20 \text{ kbar}$	$Q_4 = 0$
$A_1/R = 4715^\circ\text{K}$	$m = 1$	$A_1/R = 5500^\circ\text{K}$
$A_2/R = 9430^\circ\text{K}$		$A_2/R = 11,000^\circ\text{K}$
$B_1 = 1.08 \times 10^{13}$		$A_3/R = 19,400^\circ\text{K}$
$B_2 = 1.08 \times 10^{13}$		$B_1 = 1.6 \times 10^{12}$
$B_3 = 0$		$B_2 = 1.6 \times 10^{12}$
$F_1 = 0.01$		$B_3 = 4.6 \times 10^{15}$
$F_2 = 0.99$		$F_1 = 0.01$
$F_3 = 0$		$F_2 = 0.205$
$F_i = 0$		$F_3 = 0.470$
$Z = 0.14 \text{ cm}$		$F_i = 0.315$
$P_a = 1.5 \text{ gm/cm}^3$		$Z_2 = 2.5 \times 10^{-3} \text{ cm}$
$P = 93.8 \text{ kbar}$		$Z_3 = 12 \times 10^{-3} \text{ cm}$
$t_p = 1.5 \mu\text{sec}$		$\rho_a = 1.80 \text{ gm/cm}^2$
		$P = 100 \text{ kbar}$
		$t_p = 4 \mu\text{sec}$

The procedure for obtaining values of the rate parameters is to compare computed values of the minimum pressure for initiation of detonation obtained with different values of A_1 and B_1 with the experimental value. The computed values are obtained by setting B_2 and B_3 equal to zero so that the grain burning reactions are suppressed, and by using a low value of F_1 so that the ignition reaction alone is not capable of supporting a detonation wave. The calculation is initiated by an imposed shock of ~ 100 kbars having a duration of $1.5 \mu\text{sec}$, which is heavy enough to cause ignition. This disturbance then propagates into the material as a fading wave, and the maximum wave pressure at the last zone which ignites is the minimum pressure for initiation.

Figure 1 shows the trajectory of such a fading wave obtained with values of A_1 and B_1 derived from the data of Cook. This gives directly

$$A_2 = 43,400 \text{ cal/mol}$$

$$B_2 = 1.08 \times 10^{13}$$

whereupon

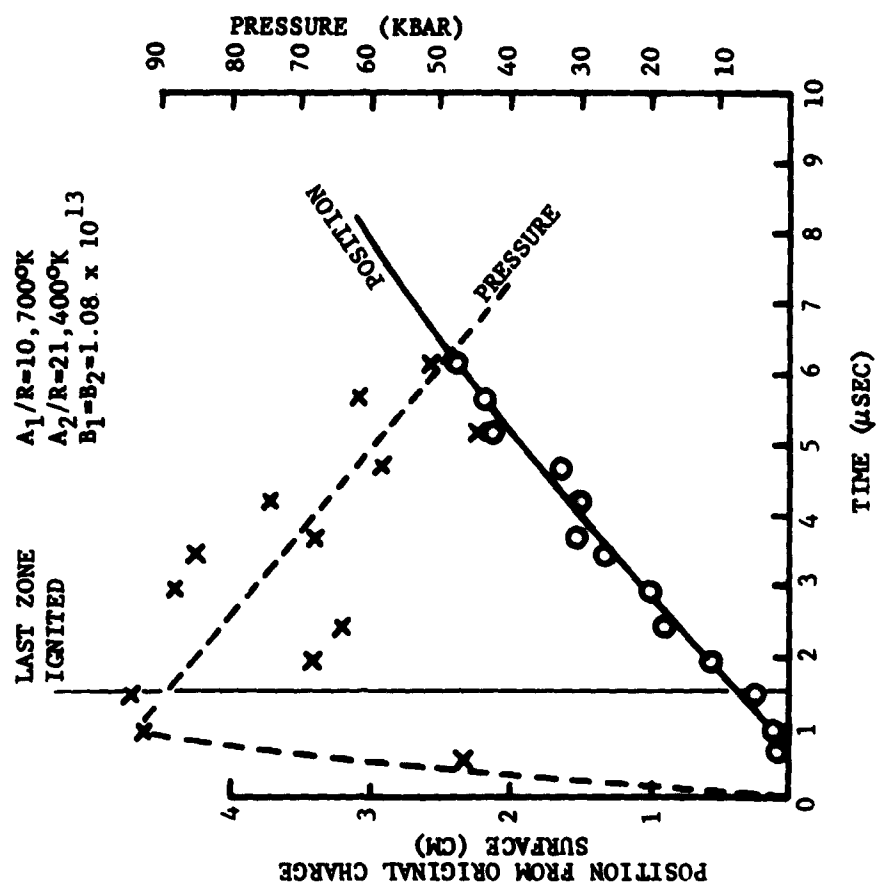
$$A_1 = 21,700 \text{ cal/mol}$$

$$B_1 = 1.08 \times 10^{13}$$

For purposes of the calculation B_2 and B_3 were set equal to zero. Data on the heat of explosion, density, and specific heat of TNT were obtained from Cook.⁴ Values of the other input parameters were approximations obtained as discussed in previous reports.

Comparison of the fade point with the curve of wave pressure shows that fading occurs at a pressure of ~ 90 kbars. This value is somewhat uncertain because of the rather wide fluctuations which occur in the computed maximum pressure. Accuracy could be increased to any desired value by increasing the stability conditions imposed on the integration and decreasing the mesh size. This would, however, increase the running time, and was not deemed to be of importance to the present calculations. The experimental value reported by NOL is about 30 kbar.

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FIGURE 1. POSITION AND MAGNITUDE OF PEAK PRESSURE IN FADING WAVE IN TNT ($A_1/R = 10,700^\circ K$ - $A_2/R = 21,400^\circ K$)

It is deduced that either the rate parameters obtained from thermal decomposition data predict reaction rates much lower than are actually obtained in a detonation process, or that the assumed equation of state parameters lead to a value of energy density behind the lead shock much lower than that actually obtained. At the present time it is not possible to choose between these two explanation.

If the value taken for A_1 is 9500, then the results are as shown in Figure 2. The minimum shock pressure for initiation here agrees very well with the experimental value. The A_1 corresponds to an A_2 for TNT of ~ 20 kcal which is not regarded as unreasonable.

Figures 3, 4 and 5 show additional results, using these values of A_1 and A_2 . Figure 3 shows the reaction profile through the wave at a time of $12 \mu\text{sec}$ after initiation when the wave has propagated about 5 cm into the charge. Complete reaction behind the wave is not being achieved at this point, and therefore it would not be expected that the steady state velocity would have been reached. This is shown by a plot of wave trajectory in Figure 4, which indicates a velocity of about $4 \text{ mm}/\mu\text{sec}$. It is expected that upon further propagation, the wave velocity would increase and the reaction zone length would decrease. However, since the overall length was set at 10 cm, it was regarded as unlikely that growth to steady state would occur before reaching the end of the charge. This wave will be examined further by rerunning the problem with a longer charge length.

Figure 5 shows plots of v_g , v_s and e_g through the wave. The reduced amount of scatter shown in the points is a consequence of the imposition of increased stability criteria on the integration. These results together with those discussed above indicate that the mathematical procedures now being used are quite well behaved.

Calculation has also been carried out on the propagation of a detonation wave through a representative composite double-base propellant using the input data indicated in Table I, obtained from information provided by the Hercules Powder Company. In this problem, two grain burning reactions were used, one for the ammonium perchlorate oxidizer, and another for the double-base binder. As in the case of TNT, the rate parameters for the binder were obtained from thermal decomposition and adiabatic self-heating data. Those for ammonium perchlorate were obtained from data published by the Aerojet General Corporation on the regression rates of APC surfaces in their hot plate experiments, which have been discussed in a previous progress report.¹ The aluminum was treated as inert. Grain size taken for the ammonium perchlorate was the average size of the

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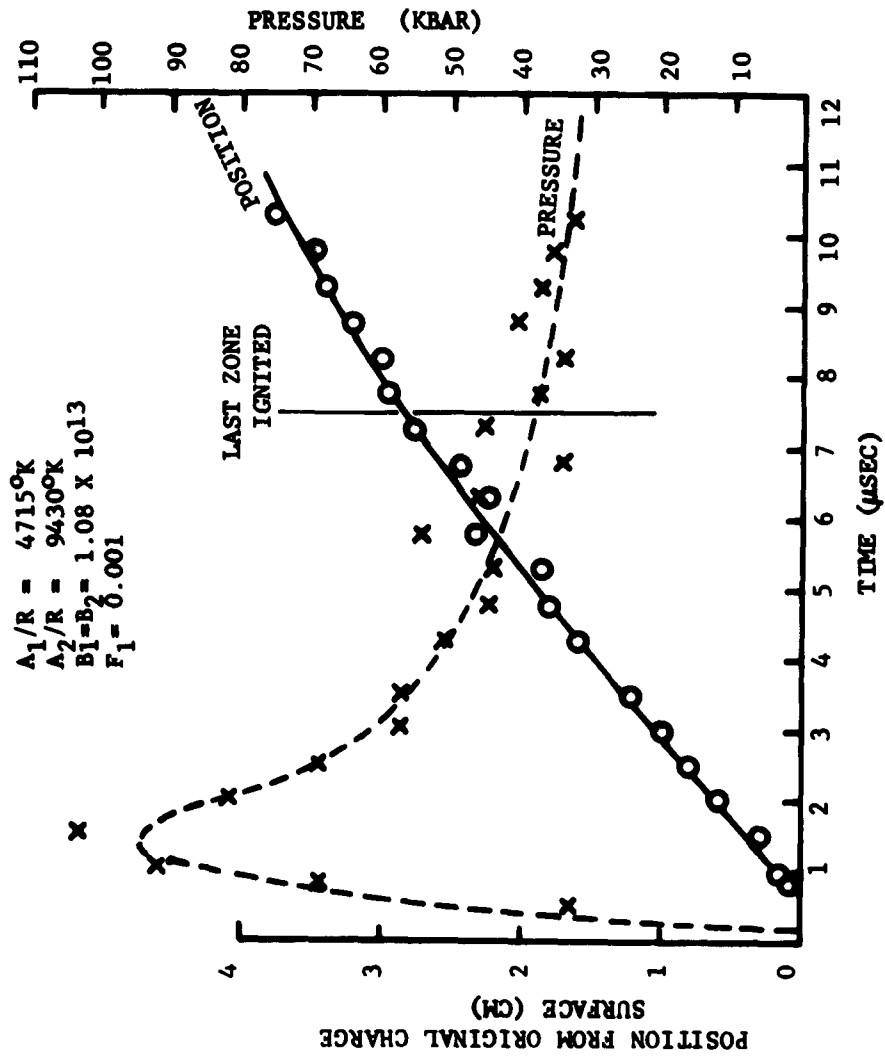


FIGURE 2. POSITION AND MAGNITUDE OF PEAK PRESSURE IN FADING WAVE IN TNT ($A_1/R = 4715^\circ K$ - $A_2/R = 9430^\circ K$)

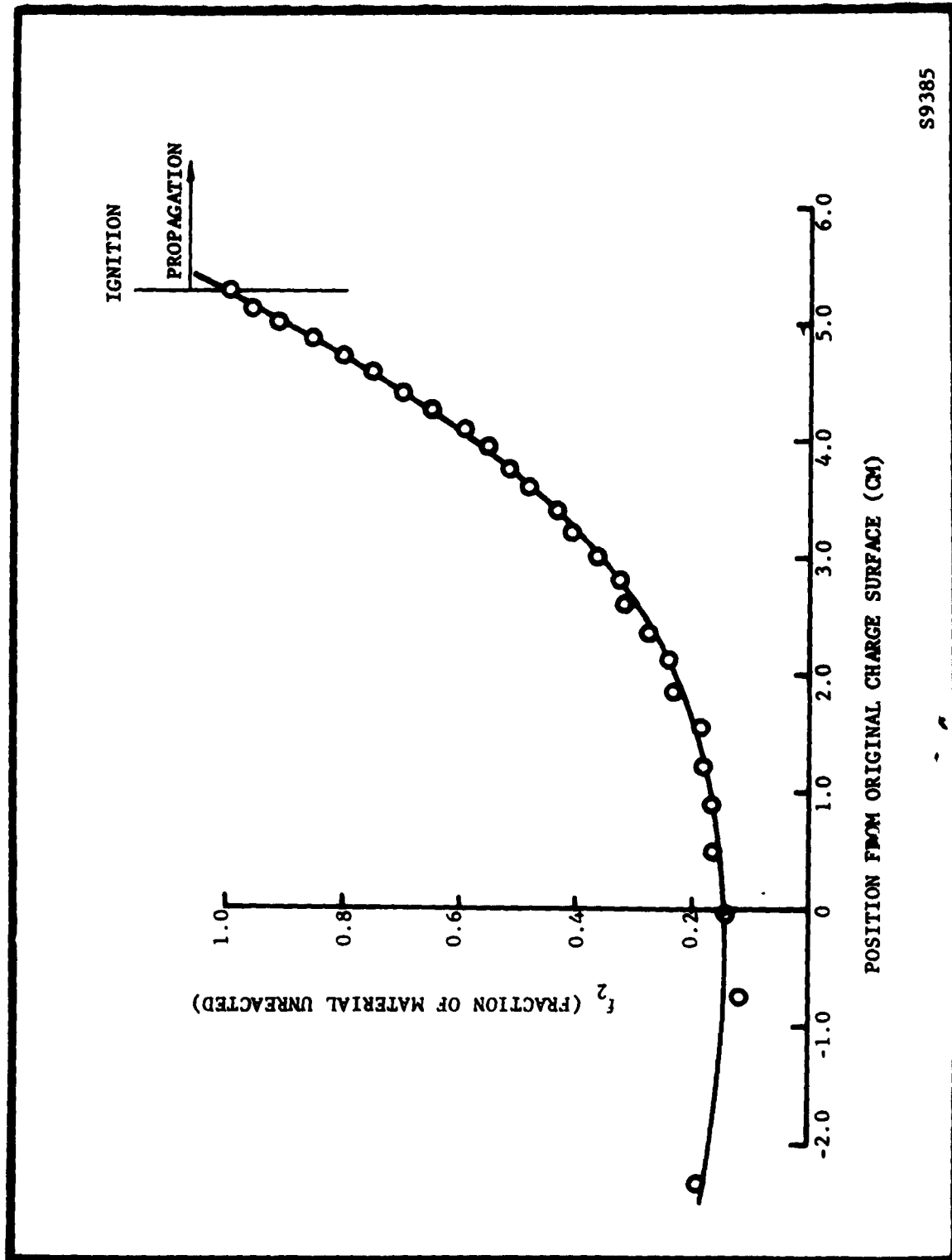


FIGURE 3. REACTION ZONE PROFILE THROUGH DETONATION WAVE IN TNT

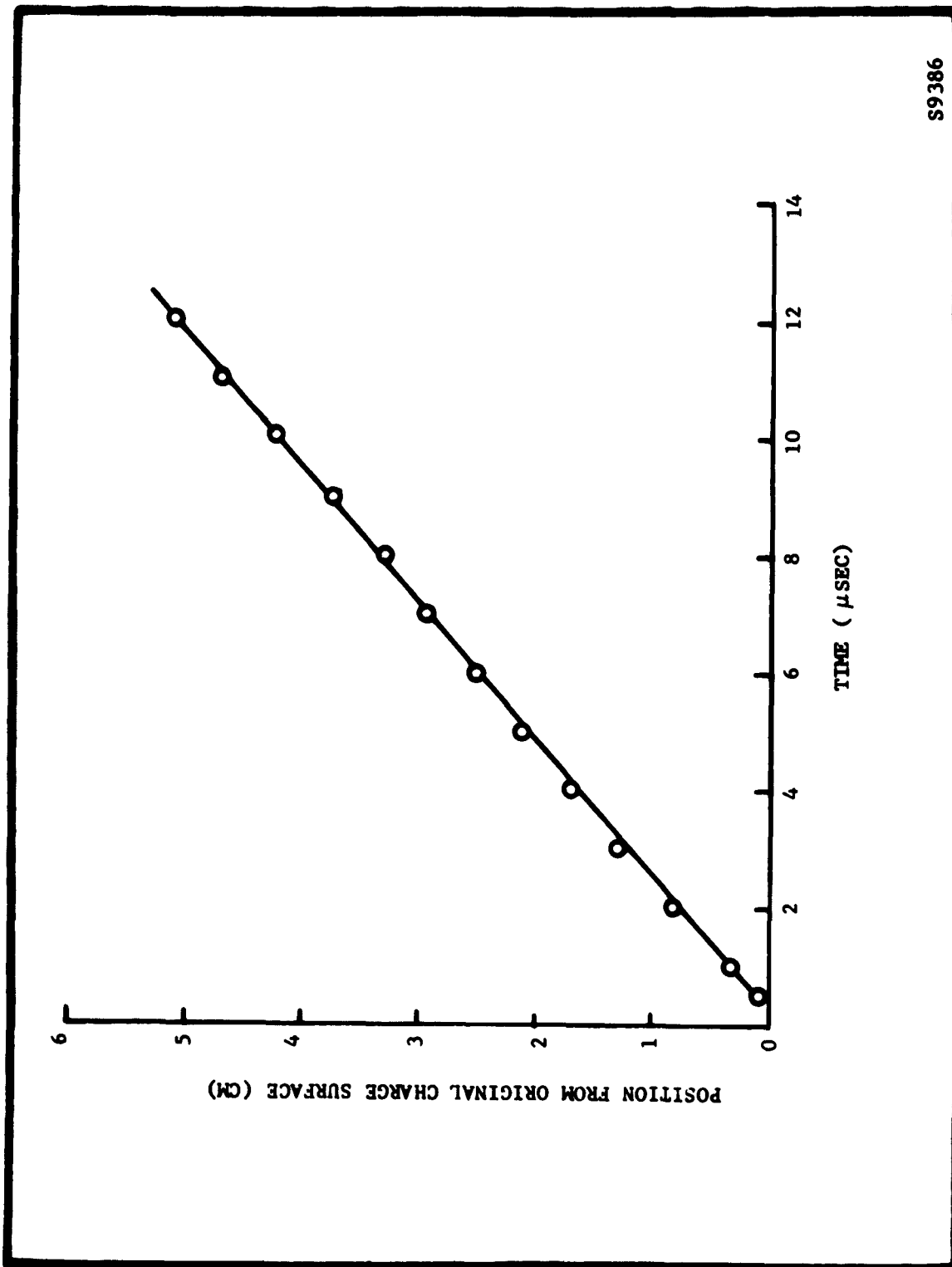


FIGURE 4. INITIAL TRAJECTORY OF DETONATION WAVE IN TNT

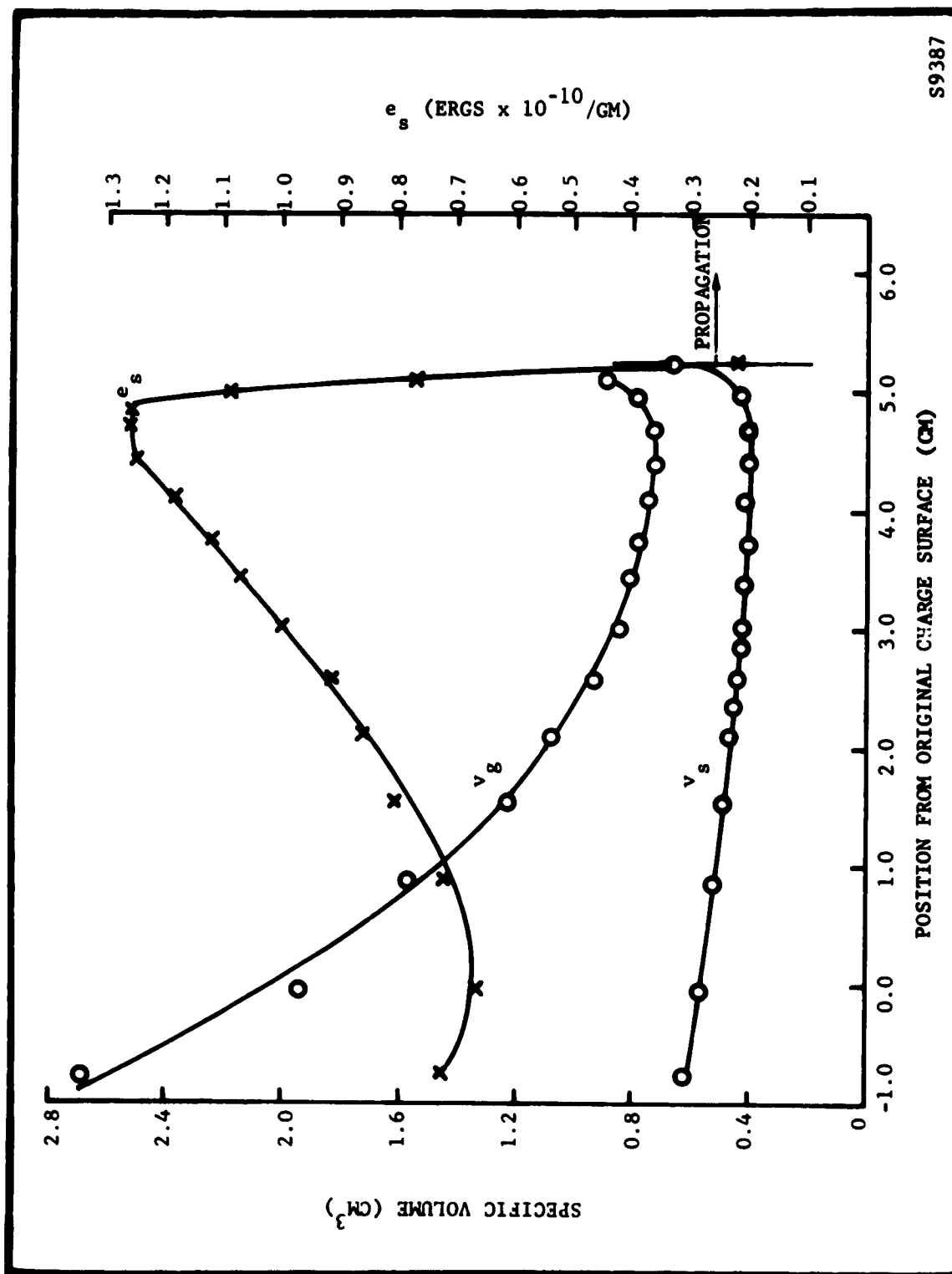


FIGURE 5. SOLID-PHASE ENERGY DENSITY AND SOLID AND GAS SPECIFIC VOLUMES THROUGH A DETONATION WAVE IN TNT

material used in the original mix; that for the binder was 1/2 the average distance between APC grains. The rate parameters for the ignition reaction were taken as those of the ammonium perchlorate for two reasons. First, under conditions of low shock intensity the APC rate law gives a faster reaction than that for the binder. This rate would therefore be most important in the ignition process. Second, NOL data from card gap experiments^{6,7,8} support this conclusion. The heat of the ignition reaction was taken as the average of the heats of reaction of the APC and binder since they are assumed to contribute equally to the ignition reaction.

Figure 6 shows the trajectory and peak pressure in a computed wave initiated with a 100 kbar shock. Initial wave velocity is found to be about 5 mm/ μ sec slowing to about 4 mm/ μ sec after 10 μ sec. Peak pressures decay rapidly from the initial imposed value of 100 kbar to about 50 kbar. It is again pointed out that the scatter shown in the computed points is an inherent consequence of numerical integration methods, which, for these problems, is particularly severe in the region of peak pressure. It could be reduced, as desired, by tightening the stability criteria for the integration; however, at an increase cost in computing time.

Figure 6 indicates that this wave approached a steady-state condition. However, at the point at which the calculations were terminated, the pressure and velocity still seemed to be decreasing slightly, and it is possible that fading would ultimately have occurred. The conclusion is that the material is marginal for detonability. It is of considerable interest here that experimental data on similar propellant materials obtained at the Naval Ordnance Laboratory are consistent with the conclusions to be drawn from these calculations that a pressure of 50 kbar is somewhat above the maximum pressure required for initiation.^{6,7,8} It would be premature to speculate as to the significance of this agreement at the present time. However, an important difference between these results and those discussed previously for TNT lies in the fact that the rate data used for the ignition reaction in the present case were obtained by the hot plate technique, while those for TNT were derived from thermal decomposition data. Further study of this point appears desirable because it may shed some light on the question of which experimental methods are most suited for obtaining kinetic information appropriate for predicting detonation behavior.

Figure 7 shows some data on reaction and pressure profiles through the detonation wave in the above composite double-base propellant. It is of interest here that in the surface layers, the binder has burned completely and the APC only to a slight extent. In the interior of the charge, neither binder or APC has reacted significantly.

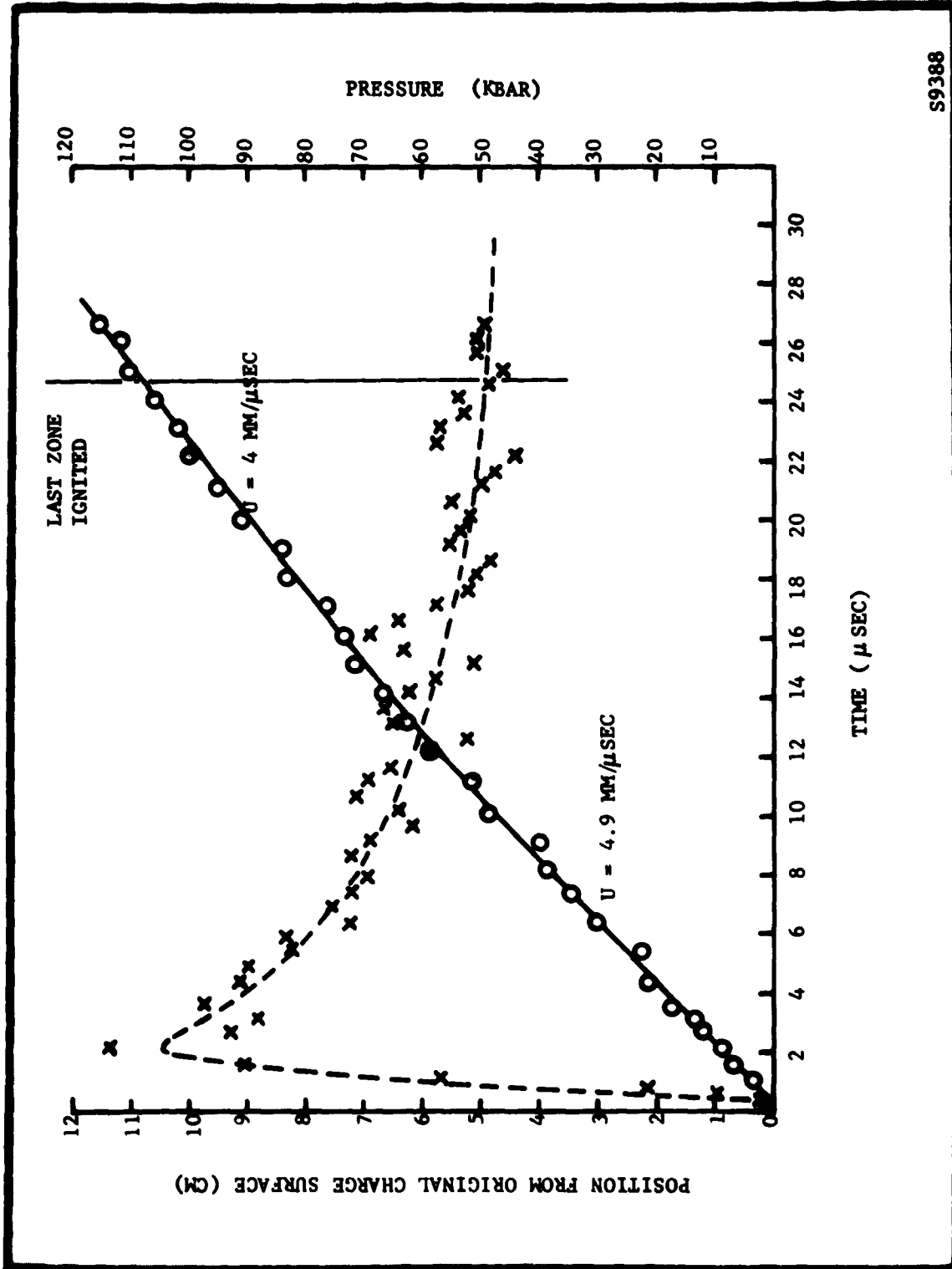


FIGURE 6. TRAJECTORY AND PEAK PRESSURES OF DETONATION WAVE IN COMPOSITE DOUBLE-BASE PROPELLANT

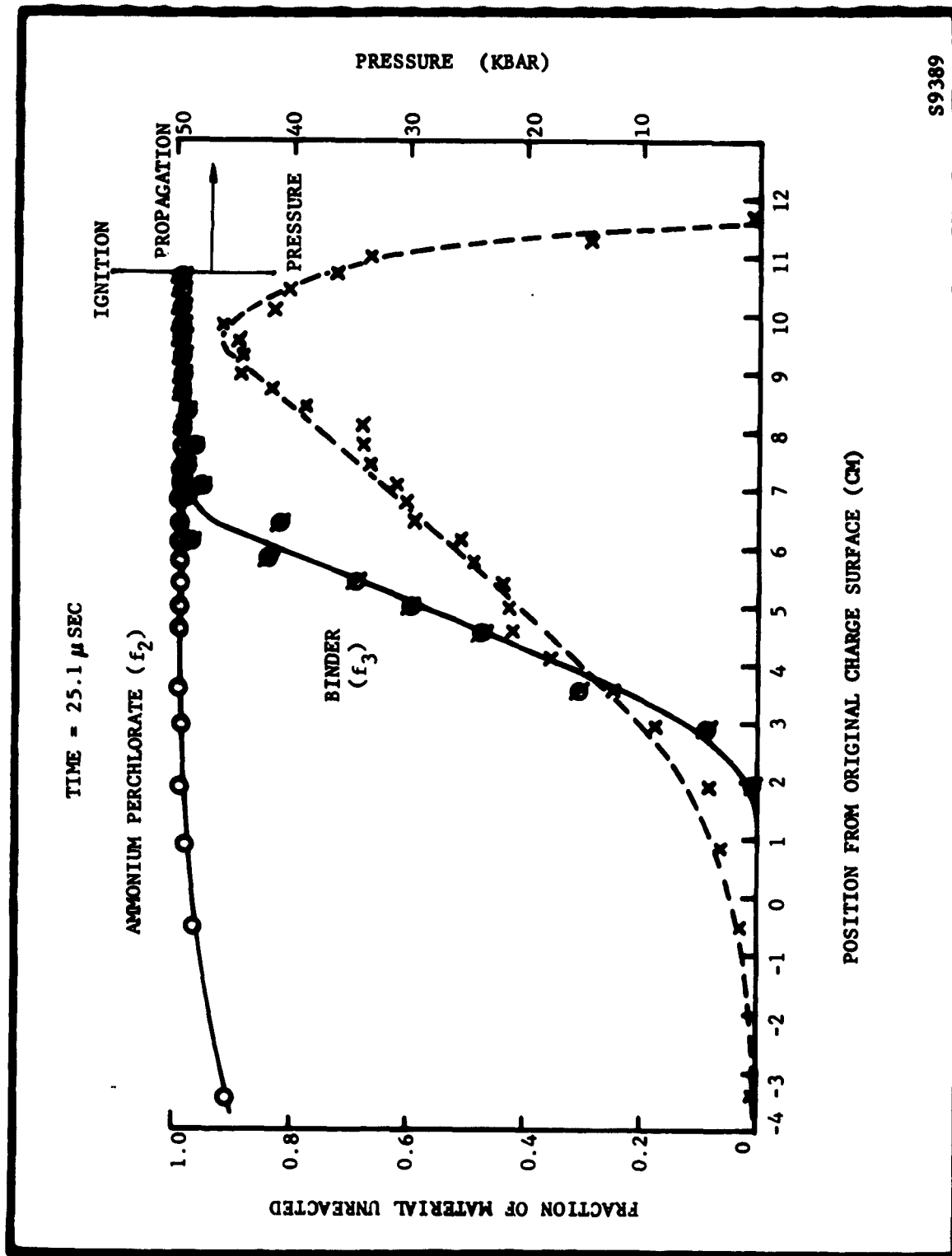


FIGURE 7. REACTION PROFILES FOR BINDER AND OXIDIZER AND PRESSURE PROFILE THROUGH A DETONATION WAVE IN COMPOSITE DOUBLE-BASE PROPELLANT

The explanation for this behavior appears to reside in the high activation energy, (A_2), and thus the high temperature coefficient for the binder reaction. Reaction in the surface is rapid because of the high energy deposited in the material by the initial shock. However, the energy released by binder reaction is not sufficient to drive a shock of this intensity so that following release of the surface pressure the wave decays. As a consequence of the decreasing pressure in the wave, the energy in the solid drops to a value at which the binder no longer reacts at a significant rate. Similar behavior is not observed on the part of the ammonium perchlorate reaction because its lower activation energy makes it less sensitive to changes in solid energy density. Following the point at which the grain burning reactions subside, the wave is supported only by the ignition reaction, and this constitutes the observed steady state process.



SECTION 4

EQUATION OF STATE STUDIES

The equation of state presently used for the solid phase is similar in form to that derivable by representation of the solid as a collection of harmonic oscillators. If the possible frequencies are ν_g , the equation obtained from such a model is the following:⁹

$$\left[p + \frac{\partial \phi}{\partial v} \right] v = \sum \gamma_g \frac{\frac{h \nu_g}{e^{RT}} - 1}{\frac{h \nu_g}{e^{RT}} - 1} \quad (29)$$

where

$$\gamma_g = - \left(\frac{\partial \ln \nu_g}{\partial \ln v} \right)_T \quad (30)$$

Here the summed terms represent the internal thermal energy of the material and ϕ is a potential function such that the internal potential energy is given by

$$e_\phi = \int_{\infty}^v \frac{\partial \phi}{\partial v} dv \quad (31)$$

For present purposes, internal thermal energy is meant to include the kinetic and potential energy associated with atomic and molecular vibration; and internal potential energy represents that energy due to atomic and molecular distortion.

At high temperature the summation in equation (29) reduces to

$$RT \sum_g \gamma_g - \xi \quad (32)$$

where ξ is the zero point energy. If all of the γ_g are equal, substitution of (32) in (29) gives:

$$\left[p + \frac{\partial \phi}{\partial v} \right] v = \gamma_G \left[3NkT - \xi \right] \quad (33)$$

The oscillator model upon which equation (33) is based is closely approximated only by a perfect single crystal. However, the form of the equation is a consequence of the condition that the potential and thermal forms of internal energy are respectively dependent upon the volume only and upon the temperature only, and are therefore separable into different terms. If it is assumed that however poorly the simple oscillator model may represent reality in other respects, this separation of thermal and potential energy is valid for real materials, then the equation is of general application, provided only that the functions $\frac{\partial \phi}{\partial v}$ and γ_G , which now must be regarded as somewhat empirical in nature, are known.

In treating detonation behavior, it is more convenient to express the equation of state in terms of total energy, since the conservation equation refers to total energy. Adding equation (31) to both sides of (33), one has ultimately

$$\left[p + \frac{\partial \phi}{\partial v} + \frac{\gamma_G}{v} \int_{v_0}^v \frac{\partial \phi}{\partial v} dv \right] v = \gamma_G e + \gamma_G \left[3 N k T_0 + \xi \right] \quad (34)$$

This is now the form of the solid equation of state presently in use where

$$\beta = \left[\frac{\partial \phi}{\partial v} + \frac{\gamma_G}{v} \int_{v_0}^v \frac{\partial \phi}{\partial v} dv \right] \quad (35)$$

$$\gamma_G = \gamma - 1$$

$$c = \gamma_G [3NkT_0 + \xi]$$



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$3NkT_0$ is the thermal energy content at ambient temperature.

e = total internal energy referred to zero at ambient conditions.

An objective of the present program is to determine the functions β and γ_G for the materials and the condition of significance to detonation phenomena. These can be obtained from p-v-T data. Partial differentiation of equation (33) gives:

$$\gamma_G = \frac{v}{3Nk} \left(\frac{\partial P}{\partial T} \right)_v \quad (36)$$

or

$$\gamma_G = \frac{v}{C_v} \left(\frac{\partial P}{\partial T} \right)_v \quad (37)$$

Thus γ_G at any v is given by the slope of the p-T curve. Substitution in (33) then provides $\frac{\partial \phi}{\partial v}$.

The necessary p-v-T data will be obtained by static pressure methods, making use of a piston-cylinder device designed by Kennedy¹⁰ capable of an ultimate pressure in the order of 80 kilobars. This apparatus has been delivered and is in the process of being set up and checked out. Some preliminary development of methods and techniques will be necessary. It is estimated that initial equation of state data will be available near the end of the next report period.



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NOMENCLATURE

t = time	ϕ = potential function
p = pressure	f = fraction of material unreacted
q = von Neumann "q"	F = fraction of material in zone
v = volume	A = activation energy
u = material velocity	B = pre-exponential constant
e = specific energy	m = reaction order
h = Planck's constant	Q = heat of reaction
k = Boltzman constant	E = chemical energy released
τ = ignition delay	ξ = zero point energy
w = empirical pressure constant	C_v = specific heat
β = equation of state constant	L = charge length
C = equation of state constant	Z = grain radius
γ = equation of state constant	ρ = density
T = temperature	a = molecular diameter
R = gas constant	P = applied pressure
ν = vibration frequency	U = wave velocity

Subscripts

o	ambient condition
1	ignition process
2	oxidizer process
3	binder process
4	diffusion process
g	gas phase
s	solid reaction phase
i	inert phase
G	Grüneisen constant
a	initial conditions
z	final conditions

Superscripts

n	time mesh number
j	space mesh number
o	ignition point
bar	average value



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