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130

THE RESPONSE OF NITROGUANIDINE TO A STRONG SHOCK

By

Donna Price and A. R. Clairmont, Jr.

ABSTRACT: In its range of detonability, nitroguanidine (NQ) behaves as a Group 1 explosive. In addition, it exhibits failure at the high TMD (> 94% at 5.08 cm diam) as well as the more common failure at a lower critical density. The high bulk density (HBD) form of NQ exhibits a critical diameter about three times that of the low bulk density form. This fact aided in studying the subcritical region of NQ (HBD) where a strong shock produces a subdetonation, supersonic, constant velocity front. This pseudo-detonation or LVD had failure characteristics similar to the detonability limits of Group 2 materials. That trend, the power of the LVD reactions, and the dimensions of the various gap tests can be combined to explain (1) a hump in the curve 50% pressure vs % TMD obtained for NQ(HBD) in the gap test and (2) a reversal in <u>apparent</u> shock sensitivity rating of NQ(HBD) and NQ(LBD) when tested on the largeand small-scale gap tests.

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2 February 1968

THE RESPONSE OF NITROGUANIDINE TO A STRONG SHOCK

This work was carried out under ORDTASK 033 102 F009 06 01 Prob 001 and MAT 03L 000 R011 01 01. Its results are particularly useful in the interpretation of gap test results as shock sensitivity of the test explosive, and in providing information about pseudodetonation (LVD) in granular charges.

> E. F. SCHREITER Captain, USN Commander

ALBERT LIGHTBODY

By direction

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TABLE OF CONTENTS

	Page
INTRODUCTION	1
EXPERIMENTAL	2
Nitroguanidine	2
Charge Preparation	2
Setup and Instrumentation	3
Records	3
RESULTS AND DISCUSSION	5
Ideal Detonation Velocity.	5
NQ (LBD)	10
NQ (HBD)	13
a. Detonation	13 20 29
Shock Sensitivity.	30
SUMMARY	37
REFERENCES	39
APPENDIX A, Additional Information on the Lots of NQ Used	41
APPENDIX B, Experimental Conditions for the Shots	46
ILLUSTRATIONS	
Figure Title	Page
1Smear Camera Record of Detonation.2Smear Camera Record of LVD	4 4
5 Comparison of Recent Experimental Values with Ideal Curve for NQ	8
4 Comparison of Detonation Behavior of NQ (LBD) with Ideal Curve	12

ILLUSTRATIONS (Cont'd)

Figure	Title	Page
5 6 7	Diameter Effect at $\rho_0 = 1.514$ g/cc for NQ (LBD) Section of Limit Curve for NQ (HED) X547.	14 14
8 9 10	NQ (LBD) Detonation Behavior of NQ (HBD), X530 Limit Curve for Detonation of NQ (HBD), X530 Pseudo Detonation (LVD) Behavior of NQ (HBD),	16 19 22
11	X530 Comparison of Diameter Dependence of LVD in Tetryl and NQ (X530)	23
12	Limit Curve for Pseudo Detonation of NQ (HBD), X530	20 28
13 14	Shock Sensitivity Curves for High and Low Bulk Density NQ	35
74	Sensitivity	35
A1-A3	Photomicrographs of NQ, X588	43-45
	TABLES	
Table	Title	Page
1 2 3	Comparison of Recent Data with Ideal Curve D_1 vs ρ_0 For NQ. Computed Values for NQ at $\rho_0 = 0.01$ g/cc Detonation Velocity Data for NQ (LBD) X547	6 9 10a
4 5 7 8 9 10 11 12	Diameter Effect at $\rho_0 = 1.514$ g/cc for NQ (LBD) X547 Detonability Limit Data For NQ (LBD) X547. Constant Velocity Data For NQ (HBD) X530 Lack of Diameter Effect on D of NQ (HBD) Critical Data for Detonation of NQ (HBD) X530. Limit Date for Pseudo Detonation of NQ (HBD) Data from NQ'S X530, X588, and X589 Previous Shock Sensitivity Tests on NQ Most Recent Shock Sensitivity Tests on NQ.	11 13 17-18 15 20 26 31 33 36

THE RESPONSE OF NITROGUANIDINE TO A STRONG SHOCK

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INTRODUCTION

In the course of comparing the shock sensitivity to detonation of a number of explosives, it was found that the high bulk density (HBD) form of nitroguanidine (NQ) failed to detonate in the small scale gap test.¹ Moreover, its apparent power in that test decreased with increasing loading density. This suggested the possibility that NQ might belong to Group 2 explosives typified by ammonium perchlorate.² Members of this group differ from TNT-like explosives (Group 1) in exhibiting a limit curve of increasing critical diameter with increasing critical density. As a result of this limit behavior, the typical detonation velocity (D) vs loading density(ρ_0) curve also differs from that of TNT-like materials in being non-linear and in exhibiting a maximum in D.

The purpose of the present study was to determine whether NQ is a Group 2 explosive. The study showed clearly that it is not, and also produced interesting new information about differences in behavior (detonation, detonability, and sensitivity) between the low bulk density (LED) NQ and NQ (HED). In particular, it provided a much more detailed picture of pseudo-detonation (or "low velocity detonation") in granular explosives than is available in the literature.

EXPERIMENTAL

Nitroguanidine

All samples of NQ, $H_2 N \cdot C(NNO_2) \cdot NH_2$ were supplied by NOS, Indian Head, Md. The NQ (LED) was manufactured by North American Cyanamid, Niagara Falls, Ontario and satisfies the military specification MIL-N-494A; its nitroguanidine content is 99.5% or greater. This material is in the form of needle-like, and frequently hollow, crystals with diameter of 5 to 10µ and length of 60-100µ. Only one lot of NQ (LED) was used and that was designated X547.

The NQ (HBD) was prepared at NCS from NQ (LBD). The preparation involves precipitation under controlled conditions and in the presence of a small amount of colloidally active material.³ The particles are chunky cylinders of ℓ/d of about 4. The average length is reported as about 20µ although particles of up to 200µ have been observed in the present work. The pour-density is about 0.7 g/cc as contrasted to less than 0.2 g/cc for NQ (LED). Two lots of NQ (HBD) were used; they were designated X530 and X589 and their sieve analyses are given in Appendix A.

The presence of the colloiding agent makes it difficult to grind the NQ (HED). Nevertheless, NOS was successful in grinding X589 to obtain a material in which 50-60% (by number) of the particles are under 1Qµ in dimensions. This lot was designated X588 and its sieve analysis and photomicrographs are also given in Appendix A.

Charge Preparation

For charge preparation, the NQ was dried at 50° C for 4 or more hours and then pressed to the required density, as shown in the tables. The lowest density charges were hand packed in cellulose acetate envelopes. The highest charge density for NQ (LBD) was 1.627 g/cc obtained in the isostatic press. NQ (HBD) can be pressed to at least 1.70 g/cc. The crystal density is 1.78 g/cc.

Cylindrical granular charges, 20.32 cm long, were prepared over the range of 1.27 to 7.62 cm in diameter. Preparation and procedures were comparable to those used in the ammonium perchlorate study.⁴

Set-up and Instrumentation

The experimental set-up of the previous work⁴ was also used here. Two inch long boosters (pentolite or tetryl) of the same diameter as the test charge were used for initiation, and a one-inch pellet of the same explosive was frequently placed as a witness at the end of the charge.

A 70 mm smear camera was used at a writing speed of 1 to 3 mm/ μ sec to follow the luminosity (or flasher enhanced luminosity) of the shock induced reaction. Film, flasher, length of charge observed, and writing speed for the photographs of each shot are tabulated in Appendix B.

Records

The records were reduced and interpreted by the methods described previously.⁴ The maximum error estimated for reading records of the lowest resolution is 1.5% in the detonation velocity. In the 14 sets of data which include replications, the average precision is 0.8%, but this value includes seven sets of low density charges (hand packed or hydrostatically pressed) for which the larger deviations (1.0 to 1.7%) most probable reflect non-uniformity of the charge, i.e., error in the effective value of ρ_0 .

The photographic records were of excellent quality and are illustrated by Figs. 1 and 2 typical of high and low velocities, respectively.

It should be mentioned that all velocity measurements will be somewhat high because a non-planar shock wave was used in the initiation. For D values comparable to that of the booster



FIG. I SMEAR CAMERA RECORD OF DETONATION (SHOT 279 OF NQ X530) AT $\rho_{o} = 1.30 \text{ g/cc}$ AND d=5.08 cm. LENGTH OF CHARGE OBSERVED 15.2 cm; WRITING SPEED 2 mm/ μ sec; NO FLASHER USED, MEASURED D 6.51 mm/ μ sec.)



FIG. 2 SMEAR CAMERA RECORD OF LVD (SHOT 26' F NQ X530 AT $\rho_0 = 1.00 \text{ g/cc}$ AND d = 2.54 cm. LENGTH OF CHARGE OBSERVED 6.35 cm; WRITING SPEED 2 mm/ μ sec; CELLULOSE ACETATE FLASHER. MEASURED LVD OF 2.5 mm/ μ sec.)

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NOLTR 67-169

(7.2 mm/ μ sec), this error varies from less than 0.2% at d = 2.5 cm to 1.5% at d = 7.6 cm.⁵ For lower D values, the error is less. No correction has been made for this effect, but it can be estimated whenever the size of the correction seems significant.

RESULTS AND DISCUSSION

Ideal Detonation Velocity

It is not necessary for the objectives of this work to investigate the validity of the currently accepted infinite diameter curve, D_i vs ρ_o , for NQ. Nevertheless, we must use the curve for reference and it seems worthwhile to see how well it fits recent data at relatively high and relatively low ρ_o .

NQ (LED) cannot be easily compressed to very low porosities. However, NQ (HED), which has been available in commercial quantities for more than ten years³, can be compressed to at least 95.5% TMD and probably higher. Materials like RDX and PETN can be detonated as single crystals, i.e., at 100% TMD, but NQ at 1.70 g/cc (95.5% TMD) cannot be detonated as a 5.08 cm diameter core in a 1.27 cm thick steel tube.⁶ There are consequently no data in the older literature for D measurements at high ρ_0 .

It is not porosity alone which accounts for this detonability phenomenon because 90/10 mixtures of NQ/RDX are detonable at 95% TMD in 5.08 c.1 diameter, unconfined charges.⁶* From the known ideal curve for RDX⁷ and the simple additivity principle, the D value for pure NQ at this porosity can be calculated from that measured for the mixture. The two highest density data points of Table 1 and Fig. 3 were obtained in this manner.

* It is not necessary that the material added be a more shock sensitive material. Ref. (6) also reports detonation of unconfined, 5.08 cm diameter NQ/AP, 69/31, at 96.8% TMD, a porosity at which neither component can be detonated alone. It is quite possible that an inert solid diluent would also increase detonability at low porosities.

TABLE 1 COMPARISON OF RECENT DATA WITH IDEAL CURVE* D₁ VS _{D6} FOR NQ

Comment	ad from Ref (6) values for NQ/RDX, at ca. 95% TMD. Simple additivit; sed. From present work, diameter negligible here.		work, Table 7, average of six g = 0.032); no detectable di- iffect.	Present work indicates very m. effect at this density.	rork, Table 4; very small di- fect.	Same comment as for 1.55 g/cc			Possibly low because of diameter	Possibly low because of diameter effect Required 15 cm run to stabilize			
	puted	LO, a Le use Le tra	ssent lues (eter e	(6).	sent ster e	. (9). mge.	(ot)	(01)	(01)	(11)	(21)	(11)	
ł	0 0 0 0		Pre Bre Bre	Rei Smê	Pre	Rei che	Ref	Rei	Ref	Ref	Ref	Ref	
Detonation Velocity, D mm/µsec	8.31	8.19	7.98	7.65	7.54	7.45	3.23	3.15	2.80	2.56	2.55	2.12	, Ref (8)
Charge diam d cm	5.08	5.08	3.8-5.1	Unknown	Extrapolated	Unkriown	7 to 10	7 to 10	7 to 10	5.08	7 to 10	~5	= 1.44 + 4.015 pg
Loading Density po g/cc	1.698	1.678	1.620	1.55	1.514	1.50	0.50	0.40	0*30	0.276	0.18	0.12	* D ₁ (mm/isec)

NOLTR 67-169

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Two sets of data in the present work were appropriate for extrapolation to D_1 (or for averaging at the higher density of 1.62 g/cc where the diameter effect cannot be detected). One point was obtained with NQ (HED); the second, with NQ (LED). Both fall on the ideal curve of Fig. 3. The other high density data are from the Army Manual as indicated in Table 1.

Table 1 and Fig. 3 also contain data for the very low density range (0.12-0.50 g/cc) which can be attained because of the needle like crystals of NQ (LED). There has been a recent interest in taking advantage of the low bulk density of this material to prepare charges exhibiting a correspondingly low detonation pressure. New low density data are available as a result.

Although the data from the very porous charges do not fall as close to the ideal curve as those at the higher densities, they lie as close to it as might be expected in view of the difficulty of controlling the uniformity of the very porous charges. However, the actual curve must diverge somewhat from the linear curve at low densities if the experimental data are to join smoothly with the value computed for $\rho_0 = 0.01$ g/cc. [The detonation parameters for $\rho_0 \cong 0$ were computed by H. Hurwitz on the Ruby code with the ideal gas law used as the equation of state for the detonation products. Results are shown in Table 2. At $\rho_0 = 0.01$ g/cc, $P_j = 187$ bars and $T_j = 2602^{\circ}$ K. The low pressure and high temperature indicate that the ideal gas law is applicable.] Divergence of the curve is shown as the dashed line in Fig. 3. According to this, the currently accepted D_i vs ρ_0 curve for NQ⁸

$$D_{1} (mm/\mu sec) = 1.44 + 4.015\rho_{0}$$
 (1)

holds down to $\rho_o \sim 0.3$ g/cc. At lower densities, it diverges to a value of $D_i (\rho_o = 0.01$ g/cc) = 2.05 mm/µsec.

Stesik and Shvedova¹³ first made use of such a computation $(\rho_0 \sim 0.01 \text{ g/cc})$ in studying nitrocelluolose (NC), another explosive in a physical form suitable for preparing charges of very low density.





TABLE 2

COMPUTED* VALUES FOR NQ AT $_{00} = 0.01 \text{ g/cc}$

	<u>C-J Values</u>
Velocity	2.048 mm/µsec
Pressure	186.7 bars
Temperature	2602 ⁰ K
Density	0.01795 g/cc
Gamma	1.258
Total gas	48.08 moles/1000g H.E.

Detonation Products (moles/1000g H.E.)

CO	8.604
^{co} 2	1.005
H	0.006
H ₂	10.585
н ₂ 0	8.591
N ₂	19.215
NH3	0.004
ОН	0.012
C (s)	0

Negligible (\leq 0.003 moles/1000g H.E.) concentration found for C(g), CH₄, N, N₂O, NO, O, and O₂. *Ruby code with ideal gas law as equation of state for products.

Their most porous charge of NC was 0.13 g/cc and their D vs ρ_0 curve for NC is very similar to that obtained here for NQ. Moreover, they computed low density values for TNT, tetryl, picric acid, RDX, and PETN, all materials difficult to prepare at densities less than 0.5 g/cc. The computed detonation velocity values were either close to the intercept ($\rho_0 = 0$) values or well above them, as is the case for NQ. E.g., computed values for TNT, RDX, and PETN were 1.996, 2.371, and 2.194 mm/µsec, respectively, whereas intercept values are⁷ 1.873, 2.532, and 1.550 mm/µsec.

NQ (LBD)

Table 3 contains the detonation velocity measurements made on the low bulk density NQ as a function of diameter and loading density. These values (or their averages, when given in Table 3) are plotted in Fig. 4. It is evident that the diameter effect is small for this material, but that it is larger at low ρ_0 than at high. In other words the higher density data are practically on the Eq(1) curve; the lower density data, below it. Since this is the case, it is reasonable to use the slope of the ideal curve to make small corrections in the high ρ_0 data for small variations in ρ_0 . Table 4 contains the D vs d data, at $\rho_0 = 1.514$ to 1.524 g/cc of Table 3, so corrected to $\rho_0 = 1.514$ g/cc; the corrected data are plotted in Fig. 5.

(page 10a follows)

TABLE 3

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DETONATION VELOCITY DATA FOR NQ (LED) X547

ρ <mark>ο</mark> (g/cc)	Detonation (Shock) Velocity mm/µsec	Shot No.	ρ _ο (g/cc)	Detonation (Shock) Velocity mm/µsec	Fade-out distance, diameters	Shot No.
<u>a</u> = 3.	310 cm; (1/d)) = 5•3	<u>d = 1.905</u>	cm ; (t/d)	= 10.7	
0.477h	3.122	77	1.5141	044.7	8	138
0.437h	- 20 - 0 - 9 - 66 		d = 1.588	cm; (1/d)	= 12. 8	
0.902H	4.03(4.848	103	0.604h	7.417	ı	541
1.3891 1.3891	7.070 7.070	1056	1.2111 1.5171	5.722 7.452	11	184 159
1.3891	7.143	107	d = 1.429	cm; (t/d)	= 14.2	
d = 3.	053 cm; (4/d	<u>= 5.0</u>	1.1991	F(4.46)	5.7	213
1.6271 1.6271	7.932 8.106	804 814	1.2161 1.5241	F(4.51) 7.403	4.5 -	215 162
d = 2.	540 cm; (1/d)) = 8.0	d = 1.270	cm ; (t/d)	= 16.0	
0.558h 0.999H 1.1851 1.5151	3.233 5.012 6.103 7.489	240 422A** 242 133	1.1921 1.1921 1.3201 1.5191	F(3.14) F(5.59) F(5.64) F	พรรณ พ อ ุพร _ิ จ์	11755 11755 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 11767 117777 117777 117777 117777 117777 1177777 1177777 1177777 1177777 11777777
Averaged a. 0.444.	values of p. 3.00(2); b. (and D, respective 0.90, 4.77(2); c.	1,521 1y: 1,39, 7,10(3)	(ردل.۲) ¹ غ ط. 1.63	ч. ст. 8.02(2).	COT

** Initiated by small booster (tetryl wafer of 1.27 cm diam x 0.635cm thick). Detonator alone failed to initiate a duplicate charge.

Symbols: F failed; h handpacked; H hydraulic press; 1 isostatic press.

* Stronger booster

NOLTR 67-169

TABLE 4

DIAMETER EFFECT AT $\rho_0 = 1.514$ g/cc for NQ (LBD) X547

đ	D#	10d ⁻¹
Cm	mm/µsec	
ω	7.54**	0
2.540	7.485	3.937
1.905	7.440	5.249
1.588	7.440	6.297
1.429	7.363	6.998
1.270	F	7.874

*Data from previous table corrected to present density by Eq (1) **By extrapolation

Fig. 5 shows that the diameter effect at this density (85% TMD) is very small. The velocity at d = 2.54 cm is only 0.06 mm/ μ sec lower than that at d = ∞ . The extrapolated value, D₁ = 7.54 mm/ μ sec, compares well with the Eq(1) value of 7.52 mm/ μ sec.

A few series of Table 3 were carried to the failure limit; those that were are summarized in Table 5, and the failure curve is illustrated in Fig. 6. This is typical Group 1 behavior for which the critical diameter increases with decreasing critical density. Fig. 7 compares the limit curve of NQ (LBD) with that of a coarse TNT.¹⁴ Although the trends are the same, the critical diameter of the NQ is 2 to 3 times that of the coarse TNT at the same \$TMD.



TABLE 5 DETONABILITY LIMIT DATA FOR NQ (LBD) X547

	de	(cm)
ρ <mark>ο (g/cc)</mark>	+	
1.00	2.54	-
1.21	1.59	1.43
1.52	1.43	1.27
	٩ _c (و	g/cc)
<u>d(cm)</u>	+	
ם את	1 50	1 01

NQ (HBD)

Table 6 contains the constant velocity measurements for high bulk density NQ, X530, as a function of diameter and loading density. The behaviors of this coarse particle sized explosive under shock are detonation, pseudo-detonation, and failure.

<u>a. Detonation.</u> The detonation pattern of NQ (HBD) is illustrated in Fig. 8. Like Group 1 explosives, the originating point for the D vs ρ_0 curves of different diameters is on the ideal curve at the high density end. (Points above the ideal curve exceed D₁ only by the order of magnitude of experimental error and also of that of the correction, which has not been made, for a non-planar initiating shock front.) From the originating point the curves fan out with slopes that increase as the diameter decreases.

The failure at highest densities (dead pressing) is not generally observed with the commoner Group 1 materials. It has been described and discussed in a previous section. The failure at the low density end of the 3.81 cm diameter curve was the usual one to be expected with approach to critical density. Failure at the low density end of the 5.08 cm curve is not as clear-cut. Of the four charges at $\rho_0 \sim 1.3$ g/cc, two were prepared in the isostatic press; their data





FIG. 6 SECTION OF LIMIT CURVE FOR NQ (LBD) X547

appear at the lower end of the linear portion of the 5.08 cm diameter D vs ρ_0 curve. The other two, prepared in the hydraulic press, had lower velocities. However, the apparent difference might arise from differences caused by different effective densities resulting from the two different methods of compaction. The points at 1.20 and 1.25 g/cc are also measurements made on charges prepared in the hydraulic press. Although the record for the lower density charge shows the familiar curvature obtained in failures, the record for the charge of 1.25 g/cc showed constant velocity. In other words, the D vs ρ_0 curve for d = 5.08 cm instead of terminating abruptly at the end of its linear portion, appears to fall rapidly as it approaches ρ_c . This may occur at other diameters and remain undetected because the experimental charge densities chosen were spaced too far apart.

The diameter effect on D, although somewhat more obvious for the NQ (HBD) data than for the NQ (LBD) data of Fig. 4, is still very small. When the D vs d data of Table 6 at $\rho_0 = 1.615 - 1.626$ g/cc are corrected to $\rho_0 = 1.620$ g/cc, they show no detectable trend (see Table 7). This is, in part, because the diameter effect at high ρ_0 is no larger than experimental error; in part, because NQ (HBD) is a coarse material with which it is hard to duplicate charges. The mean value is 7.98 mm/µsec ($\sigma = 0.030$) and compares well to the two point average for NQ (LBD), at d = 3.65 cm and corrected to $\rho_0 = 1.620$ g/cc, of 7.99 mm/µsec. Both are slightly higher than the ideal value of 7.94 mm/µsec from Eq(1).

TABLE 7

LACK OF DIAMETER EFFECT ON D OF NQ (HBD)

d(cm)	D	(mm/µsec)*	
5.08		7.963	
4.44		7.994	
4.13		7.969	
3.97		7.933	
3.81		8.026	
3.81		<u>7.997</u>	
	Average	$7.980 (\sigma = 0.030)$))

*po = 1.629/cc



FIG. 7 COMPARISON OF LIMIT CURVES FOR COARSE TNT AND NQ(LBD)

CONSTANT VELOCITY DATA FOR NQ (HBD) X530 TABLE 6

						Fade-out distance,	diameters	n. N. N.	00 44		(50 • •		
Shot No.	= 4.9	303 = 5.1	309	= 5.3	268 270 288	N609 N609	500 500 500	2675	260 290 290	331 371	212	271 315	N I N	
Front Velocity* mm/µsec	28 cm; (1/a)	7.969 69 cm; (1/d)	T+0.7	1 cm; (1/d)	2.942 2.874 3.188	3.210 3.208	3.109 F(2.15)	Fへ3。44 Fへ3。34	F(3.28) F(4.11)	6.723 6.841	8,034 7,997	下 (2・20) 日 (2・20)		
ρ <mark>ο</mark> (g/cc)	d = 4.12	1.6201 <u>d = 3.9</u> (1,6221	d = 3.8	0.735h 0.735h 0.851h	1.001h 1.000h	1.101h 1.202H	1.295H 1.3031	1.3071	1.4011	1.6221	1.0001 1.6651		
Shot No.	= 2.7	326 327 328	= 4.0	276	292 8 292 8 306	216°	305 310	330 318	279 317	282 304	281 273	280) = 4.6	294
Front Velocity* mm/µsec	cm; (t/d)	4.833 5.681 7.075	cm; (1/d)	3.211	v v v v v v v v v v v v v v v v v v v	3.754 3.784	F(4.88) 5.579	6.279 6.125	6.518 6.502	7.226	7.493	7,987	5 cm; (1/d	7.974
p <mark>o</mark> (g/cc)	d = 7.62	1.001h 1.198H 1.4221	d = 5.08	0.718h	0.05 1.001h 1.051	1.102h	1.200H 1.251H	1.292H 1.299H	1.2991 1.3041	1.4081	1.5121	1.6261	d = 4.44	1.6151

NOLTR 67-169

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TABL

						Averaged values of p _o and velocity, respectively:	a. 1.001, 3.752(2); b. 1.102, 3.769(2); c. 0.735, 2.908(2); d. 1.004, 3.209(2);	e. 0.72, 2.348(2); f. 1.001, 2.540(2); g. 0.709, 1.829(3); k. 0.850, 1.956(4).	* Number in parentheses after F is velocity of failing shock.	**Stronger booster. Other symbols as in Table 3.	
Fade-out Aistance	diameters	2°2		00000 00000 0000	2.2				99 • 4 7	10 4 0	0,0,0 0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,0,0,0 0,
Shot No.) = 5.6	313	= 8 . 0	ю ғ. н. <i>Ч. м</i> б 1 ғ. м б 1 в 1 в 1 в 1 в 1 в 1 в 1 в 1 в 1 в 1 в	250	d) = 12.8	2555 252 248***	2007 2007 2017 2017 2017 2017 2017 2017	262 257 257	544	245 245 246
Velocity* mm/µsec	l cm; (1/d	۴ı	cm; (t/d)	2.345 2.352 2.568 2.580 7.25) 7.<25)	Щ	75 cm; (1/	1.781 1.851 1.854	1.897 1.963	1.983 F(1.96)		* * £4 £4
ρ _o (g/cc)	d = 3.65	1.6171	d = 2.54	0.72h 0.72h 0.851h 1.001h 1.101h 1.2941 1.2941	1.6751	d = 1.58	0.704h 0.709h 0.715h	0.850h 0.850h 0.850h	0.850h 1.002h	1.009h 1.2831	1.2831 1.6781 6781







The limit data for detonation of NQ X530 are collected in Table 8 and plotted in Fig. 9 which also shows the failure curve for NQ (LBD) X547. It is evident that NQ (HBD), like NQ (LBD), shows the Group 1 trend of increasing d_c with decreasing ρ_c . In view of the difference in particle size and shape, it is not surprising that the NQ (LBD) curve lies well below that for NQ (HBD) nor that the "dead-pressing" phenomenon, if it exists in NQ (LBD), occurs at a higher density than in NQ (HBD). The highest density we achieved for the LBD material was 1.627 g/cc at d = 3.65 cm; it detonated at these conditions whereas the NQ (HBD) failed at 1.617 g/cc.

TABLE 8

CRITICAL DATA FOR DETONATION OF NQ (HBD) X530

ρ 。 (g	;/cc)	d	(cm)
+	-	+	
1.63	1.67	3.8	31
1.	62	3.81	3.65
1.40	1.36	3.8	31
1.25	1.20	5.0	08
l.	00	7.62	-

<u>b. Pseudo Detonation.</u> In the lower density region, shocked NQ (HBD) exhibits a diameter and density dependent constant front velocity well below that of the ideal curve. The behavior observed is illustrated in Fig. 10 which covers the region of what we shall call pseudo detonation or LVD (for the term "low velocity detonation" which frequently appears in the literature).

The phenomenon of LVD is well established for liquid explosives, particularly for nitroglycerin which has been extensively studied.

For solid explosives, many reported instances of LVD are obviously transient states encountered in the course of buildup to detonation. There are, however, a few instances of sub-detonation but supersonic constant velocities persisting for great distances.

Patterson¹⁵ first noted that a 1.5 in. diameter cartridge of blasting gelatin (NG and NC) exhibited a constant velocity of about 2.4 mm/ μ sec without any sign of fading over a length of at least 17 ft. He also found that at cartridge diameters of 7/8 to 2 in., either detonation (D ~ 8 mm/ μ sec) or LVD could be initiated according to the power of the booster used; that above d = 2 in. only detonation was stable, i.e., a reaction front initially moving at lower velocities would accelerate to D if the cartridge were long enough. Other workers noted that the blasting gelatin must have some aeration to exhibit LVD and that frequently some unburned explosive can be recovered after LVD has traversed the cartridge.

An earlier reference and one more relevant to the present work is that of Jones and Mitchell.¹⁶ They too used detonators of different power on coarse granular charges of TNT. They report, "by using a sufficiently narrow cartridge, or, alternatively, a sufficiently coarse grist of explosive, the low-order detonation can be made to travel throughout the length of the cartridge (at least a meter) at a rate which is both uniform and repeatable.----- On the other hand, if a sufficiently powerful initiator is used, the same cartridge propagates detonation at a greatly enhanced speed, the rate of detonation being again uniform and repeatable." They too comment on the presence of undecomposed explosive residue after LVD.

Recently, Parfenov and Apin¹⁷, have studied coarse granular low density charges of tetryl, TNT, and RDX. For each material, ρ_0 was fixed (~1.0 g/cc) and the diameter varied in the range of 5 to 40 mm. The ratio ℓ/d was kept ≥ 10 . The particle sizes used were 1000-1600µ, 630-1000µ, and 400-630µ for tetryl; 400-630µ for TNT, and 1000-1600µ for RDX. As in the previous two references, initiators of high and low power were used. In all three H.E., regions of detonation and of LVD were reported. For TNT and RDX, these regions showed





FIG. 10 PSEUDO DETONATION (LVD) BEHAVIOR OF NQ (HBD), X530

overlap*, but for tetryl they were sharply divided at $d = d_c$ for detonation. Consequently the division was also detectable when only the high powered initiator was used. Gurton^{1*}, on the other hand, reports an apparent overlap for another coarse tetryl (pour density of 0.9 g/cc, particle size not reported). At d = 1.11 cm (7/16 in.), Gurton reported 1.5 and 4.6 mm/µsec obtained respectively with weak and strong initiation. The latter figure is quite close to the ideal velocity for this density, and the former is in the range of LVD measurements of Ref. 17. It is also possible that 11.1 mm was about the critical diameter (it is near the d_c values of Ref. 17); if so, the two sets of data^{17,18} are completely consistent.

It is our opinion that the pattern of Fig. 10 is caused by a similar LVD phenomenon which is detected with our powerful initiator only at diameters below the critical value for detonation. It is possible, of course, that a weaker initiator might extend the LVD curves to higher ρ_0 , but the shock velocities in the failures (indicated as F in Fig. 10) are generally lower than the constant value of the LVD at d = 1.59-3.81 cm. For that reason a weaker initiator would not be expected to extend the LVD curves.

In detonation of a pure H.E., the diameter effect on D is found experimentally to be

 $D = \alpha + \beta/d \qquad \alpha, \beta, \rho, \text{ constant} \qquad (2)$ and d >> d_c If we let U_{LV} = the constant but pseudo detonation velocities of

Fig. 10 $U_{T,Y} - 1.42 = (d^n - 1) (3.29 - 1.08 \rho_0)$ (

 $U_{LV} - 1.42 = (d^{n} - 1) (3.29 - 1.08 \rho_{o})$ (3) where U, d, and ρ_{o} are in mm/µsec, cm, and g/cc, respectively, and

n \cong 0.129 + 0.316 $\rho_0 \cong$ 0.445 $\rho_0^{0.7}$ The ranges for Eq(3) are 0.75 to 1.0 g/cc and 1.59 to 5.08 cm in ρ_0 and d respectively.

*The overlap was in the diameter range of the velocity vs d curve, i.e., for some d values either detonation or LVD occurred according to the kind of initiation. It is interesting that the coarse TNT exhibited $d_c \cong 24$ mm at $\rho_o = 0.95$ g/cc (about twice the d_c for the finer material of Fig. 7) and that LVD was observed in the range 23-30 mm.

It is quite evident that the diameter dependence of $U_{LV} Eq(3)$ is very different from that of D Eq(2). Similarly the dependence on ρ_0 differs because, experimentally, Group 1 explosives show

 $D = A' + B'\rho_0, \qquad A', B', \text{ and } d \text{ constant}$ or $d > d_c$

 $D(d,\rho_0) = A + B\rho_0 - \beta(\rho_0)/d \qquad \text{where } A, \text{ B are constants}$ for the infinite diameter curve and $\beta(\rho_0)$ seems to be a linear function of ρ_0 .

For these reasons as well as the termination of the detonation curves shown in Figs. 8 and 10, the phenomena described by Eq(3) are considered pseudo-detonation or LVD. We do not mean to suggest by this term that the reaction involved in LVD is necessarily very weak. Even at its lowest velocities (~ 2 mm/µsec at d ~ 1.6 cm and $\rho_0 \sim 0.8$ g/cc) it was able to initiate an explosive witness placed at the end of the charge. At higher velocities it appears well able to damage steel plates.

Fig. 10 shows a previously unreported area of failure (1.20 to 1.36 g/cc in extent) separating the region of LVD from the region of detonation for the curves obtained with charges of d = 3.81 cm. But because previous investigators^{16,17,18} restricted themselves to a constant charge density (and a low one) for granular charges, they could not observe such a density dependent failure phenomenon. In view of our own difficulty in sharply dividing pseudo-detonation from detonation behavior in the moderately sized charges (d = 5.08 cm), it is fortuitous that we happened to choose the charge size of d = 3.81 cm at which such a failure area seems clearly evident. The smaller charges (d = 1.59 to 2.54 cm) exhibited only LVD and were evidently subcritical for detonation over the entire density range tested.

The trends of Fig. 10 indicate that as the charge diameter is increased, the LVD region approaches the detonation region. At sufficiently large diameter, therefore, only detonation should be observed. This trend is perhaps more evident when the data at fixed charged density are plotted as front velocity vs charge diameter.

This has been done in Fig. 11 for NQ, X530 at $\rho_0 = 0.85$ g/cc where it is compared with the Russian curve¹⁷ for coarse tetryl at $\rho_0 = 0.90$ g/cc. Parfenov and Apin¹⁷ found that the tetryl exhibited LVD at d \leq d_c, detonation at d \geq d_c. Because the critical diameter for detonation of NQ X530 at $\rho_0 = 0.85$ g/cc is much greater than that of the tetryl (See Fig. 9), the entire range of diameters in Fig. 11 is subcritical for NQ. At some large diameter, NQ, like the tetryl, would probably exhibit a discontinuous charge from LVD to detonation.

As Fig. 10 also shows, there exists a failure or limit curve for the LVD behavior. The critical data, collected in Table 9 and plotted in Fig. 12, emphasize the interesting fact that the limit curve for LVD (Fig. 12) is exactly opposite in trend to the limit curve for detonation (Fig. 9). Moreover the limit behavior for LVD (increasing diameter with increasing density) parallels the Group 2 limit behavior for detonation. It seems most likely, therefore, that it was an LVD reaction which was responsible for an increasing output with decreasing %TMD when NQ (HED) was tested in the small scale gap test.¹ It also seems highly probable from the dents produced that reactions in at least a part of the LVD region will be quite powerful enough to exhibit a positive result on the regular large scale gap test.

TABLE 9

LIMIT DATA FOR PSEUDO DETONATION OF NQ (HBD)

Diameter d(cm)	Density +	ρ _o (g/cc)
1.59	0.85	1.00
2.54	1.00	1.10
3,81	1.10	1.20
5.08	1.10	1.20

The usual qualitative explanation for an LVD reaction is that it is supported by the explosive decomposition of only a fraction of the H.E. present. If as Ref. 16 suggests, LVD can be ovserved in solids

only when they are coarse granular materials, it is possible that its propagation depends on a hot gas flow through a granular bed exhibiting a particular porosity. But it is difficult to consider such a mechanism applicable to blasting gelatin even when it contains some aerated areas. The concepts of partial reaction and surface reaction do seem applicable in both cases.

The mechanism of pseudo-detonation which seems most acceptable, however, is that suggested in 1962 by Bowden.¹⁹ It consists of an ignition wave coupled to a shock wave, e.g., a shock wave travelling through crystal of explosive, which contains a number of imperfections, ignites the explosive at the site of each imperfection.¹⁹ The ignition wave travels at the shockwave velocity although the deflagration, that continues at each ignition site after the shockwave has passed, progresses at a much slower, subsonic rate.

Imperfections in a single crystal serve as hetergeneous areas where energy concentration and hence ignition can occur. In granular explosives, there is probably a range in grain sizes and shapes which favor hot spot formation on grain surfaces when the grain bed is shocked. (Local concentration of energy could occur by reflection and reinforcement of the initial shock and by spalling or jetting of the shocked grains.) If the physical characteristics of the granular bed are such as to produce numerous, well distributed sites of ignition, such shock-produced hot spots can result in the continuous smear camera record of a pseudo-detonation. The grain size favoring pseudo-detonation will depend on the material. Thus, for the relatively sensitive tetryl ($\rho_0 = 0.90 \text{ g/cc}$), it is reported at a grain size of 400-1600 μ and d < d_c = 12 mm.¹⁷ But for the less sensitive NQ, X530 $(\rho_0 = 0.85 \text{ g/cc})$ it is found at a grain size of up to 200µ (average about 100μ) and d = 16 - 50 mm. It has not been observed at all in the low bulk density NQ* which has thin needle crystals of 50 - 100µ length.

*Attempts made to initiate LVD in NQ (LBD) with weak initiators were unsuccessful. Gurton¹⁸ refers to LVD of a NQ (LBD) at $\rho_0 = 0.5$ g/cc and d = 1.1 to 1.9 cm, but he measured the same velocities for both weak and strong initiation. Since the velocities were also high (80 - 90% ideal for $\rho_0 = 0.5$ g/cc), it is probable that the phenomenon he observed was detonation.











The instantaneous reaction and energy release at the hot spots, although amounting to only a small fraction of the total chemical energy of the explosive, is sufficient to contribute to the shock wave energy to the extent of producing an essentially constant velocity Moreover, in every case the energy release resulted in a front. pressure at the surface of the pentolite witness sufficiently high to initiate almost immediate detonation of the pentolite (See Fig. 2). Initiation of pentolite very near the shocked surface, i.e., a runlength of a few mm, requires shock pressures of about 40 kbar although the minimum initiating pressure (run length of up to two diameters from the shocked surface) is perhaps 10 kbar. The pressure of about 40 kbar, indicated by the behavior of the pentolite witness, is of the order of magnitude required to punch the steel witness plate in the standard gap test of shock sensitivity. It is therefore to be expected that some of the pseudo-detonation reactions will exhibit positive results in the gap test.

<u>c. Effect of Particle Size.</u> We have assumed that the major differences in behavior between NQ (LED) and NQ (HED) in the lower ρ_0 range arises from the difference in particle size. [In the higher ρ_0 range where both materials are detonable, they differ markedly in detonability (Fig. 9) but not in detonation velocity after initiation.] But because of the very different particle shapes in the two forms of NQ, it seemed desirable to compare a sample of NQ (HED) with the same material after it had been ground to a smaller particle size.

For this purpose, NQ (HBD) X589 was obtained; its screen analysis was similar to that of X530 (See Appendix A). A sample of X589 was then ground until it contained a large number (at least 50%) of fines; the ground material was designated X588. Table 10 contains the detonation velocity data obtained for comparision of these two materials.

X589 appears coarser than X530; it exhibits failure at $\rho_0 = 1.0$ g/cc whereas X530 shows LVD here. Grinding X589 makes it effectively

much finer than X530. For example, 5.08 cm charges of X588 exhibit velocities equal those of the 7.62 cm charges of X530. Moreover, at 5.08 cm diameter, NQ X588 is detonating at D of 0.88, 0.93, and 0.99 D_i , respectively, for ρ_0 values of 1.0, 1.2, and 1.35 g/cc. Hence, decreasing the particle size by grinding has eliminated the LVD region of Fig. 10 at this diameter. Increasing the diameter can also eliminate the LVD region. For any given material, the ratio of its particle size to its charge diameter and its shock sensitivity probably determine whether LVD appears.

Shock Sensitivity

The regular gap test²⁰ has been used to investigate the shockto-detonation sensitivity of various lots of NQ in the past. These earlier test results are collected in Table 11 and plotted in Fig. 13. Three production lots of NQ (HBD) from NOS fell on the same P_g vs ρ_o curve despite average particle size variation of 38-100 μ . Moreover, all three lots confirmed the existence of the unusual hump in the curve between 75 and 85% TMD. Heretofor we have had no suggested explanations for this peculiarity. In view of the present work we can surmise that it might be caused when the induced reaction is an LVD rather than detonation.

Aside from the small hump on the lower curve, the trends of Fig. 13 seem normal. NQ (HBD) approaches its "dead-pressed" density in the gap test configuration at about 92% TMD (1.64 g/cc); this compares well to slightly above 1.626 g/cc, the value found in the work on unconfined charges of d = 5.08 cm. At this high density, the diameter effect is negligibly small; hence the unconfined and confined charges should exhibit the same critical density for detonation. The sensitivity curve for NQ (LED) lies slightly above that for the high density material and approaches the lower curve at the greatest compaction. It, however, shows no hump and is very like P_g vs TMD curves of other H.E. (See, for example, Ref 1) The difference, NQ (LED) less shock sensitive than NQ (HED), is that expected when the chief difference between lots is a difference in particle size. Generally the finer material, if it differs at all from the coarser, shows lower shock sensitivity in a granular charge¹.

TABLE 10

DATA FROM NQ'S X530, X588, AND X589 Front Velocity Shot ρο g/cc NQ mm/µsec No. $d = 5.08 \text{ cm}; (\ell/d) = 4.0$ $F(3.54)^{a}$ x589 1.001h 397 F(4.14) 1.201H 396 1.4271 7.067 395 1.4331 7.169 399 **x**588^b 1.001h 4.803 411 1.202H 5.805 410 6.797 409 1.3511 Effect of Particle Size Front Velocity Relative Particle

NQ	<u>a</u>	ρ ₀ : 1.00	1.20	1.43	Size
x 589	5.08	F	F	7.12	Coarse
X 530	5.08	3.76	F	7.10 [°]	Less coarse
x 588	5.08	4.80	5.80		Least coarse

		Front	Veloci	ty	
NQ	<u>a</u>	ρ ₀ : 1.00	1.20	1.35	
x 588	5.08	4.80	5.80	6.80	Fine
X 530	7.62	4.83	5.68	6.74°	Coarse

^a F means failed. Number in parentheses is velocity of failing shock.

^b X589 ground. See Appendix A

^c Read from Fig. 8

The hump in the NQ (HBD) curve of Fig. 13 is at about 79% TMD or 1.41 g/cc. This is about the critical density for 3.81 cm diameter unconfined charges of NQ, X530 (See Figs. 8 and 9). The explosive core diameter in the gap test is 3.65 cm, but since the diameter effect is still small* at 1.41 g/cc, the effective diameter for detonation in the gap test should be close to 3.81 cm. Hence it is very near $\rho_0 \sim 1.41$ g/cc that detonation of this NQ would fail in the gap The confinement, by providing local shock enhancement by test. reflection from the steel walls, might well extend the density range of LVD up to the critical density for detonation. In other words, the containing steel tube which would have little effect on detonability at this density might so enhance the possibility of LVD that the pseudo-detonation can occur at $\rho_o < \rho_c$. If at charge densities lower than $\rho_0 \sim 1.41$ g/cc LVD does occur, its reaction would be sufficiently powerful to punch the steel witness plate, i.e., to produce a positive result in the gap test.

In accord with the above interpretation, the hump in the P_g vs % TMD curve of Fig. 13 results from a combination of detonability and shock sensitivity behavior near the critical density ρ_c for detonation. At $\rho_o > \rho_c$, NQ (HBD) detonates and P_g measures its shock <u>sensitivity</u> to detonation; at $\rho_o < \rho_c$ this NQ exhibits LVD and P_g measures its shock sensitivity to LVD.

If this explanation is correct, other samples of NQ (HBD) which exhibit $\rho_c \sim 1.41$ g/cc should also exhibit a hump in the P_g vs #TMDcurve. If such a NQ is ground until it is detonable at lower charge densities, then its shock sensitivity curve should resemble that for other H.E. and be concave upward over the #TMD test range with no hump present.

Table 12 contains the gap test data for X589, a recent production lot of NQ (HBD) and for X588, a sample from the same lot after grinding. Detonation velocity measurements (Table 10) showed that

*For 3.81 cm diameter charges, the D values at 1.5. and 1.41 g/cc are, respectively, 0.98 and $0.96D_4$.

TABLE 11

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PREVIOUS SHOCK SENSITIVITY TESTS ON NQ

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		Comment	Average particle size 51µ X311 screened to 38µ	Average particle size 64µ	Average particle size 100µ	
71170	പ്പ	kbar	90 55.5	88 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	24 27 24 29 29 29 29 20 20 20 20 20 20 20 20 20 20 20 20 20	00004111 000180 000000
1 200	•on	Cards	36 85	32 647 1282 196 196	94 86 128 128	성 194 215 _월 1 215 _월 1
	CIMIL	R	92 . 1 80 . 9	922 850 65 1 7 85 1 7 85 1 85 1 85 1 85 1 85 1 85	79.7 80.7 77.9 74.9	91 20 20 20 20 20 20 20 20 20 20 20 20 20
	ρ _o ‡	g/cc	1.641 1.441	1.641 1.511 1.5711 1.73H	1.421 1.441 1.59H 1.331	1.63 1.64 1.411 0.00 0.56h
		NQ(HBD)	X311*	X446*	X510	NQ(LEL) NQ(LEL) N

NOLTR 67-169

* 44 4-

*;

Reported in NOLTR 65-177 Reported in NOLTR 66-87 H hydraulic press, h handpacked, i isostatic press. Pentolite booster.

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X589 is apparently a coarser material than X530. It was therefore expected to exhibit a failure to detonate in the gap test and produce a hump in the P vs %TMD curve near the location of the hump in Fig. 13. The data taken in that location (range of 78 - 85% TMD) are plotted in Fig. 14 and show the expected hump. The ground material X588 was shown by detonation velocity measurements to be detonable in this range and should exhibit no hump in its sensitivity curve. As Fig. 14 shows, this is the case and the P_g vs %TMD curve for X588 in this region is concave upward and approximately parallel to that for NQ (LBD) which is also shown for comparison. The difference between NQ (LBD) and X588 (the latter is more sensitive) is that expected in comparing relatively fine with relatively coarse material. The difference between X588 and X589 is in the opposite direction. An apparent reversal in the usual particle size effect on shock sensitivity could occur if the witness damage is cause by detonation in the one case and LVD in the second.

Two additional aspects of gap testing are worth emphasizing. First, to judge by the velocity pattern of Fig. 10, there is a large range of conditions in which a single shot could not be adequately classified as detonation or LVD. This would be so even for an instrumented gap test (in which detonation velocity or pressure might be measured) because classification requires knowledge of the complete pattern of Fig. 10.

The second point is that the peculiarity in the measured shock sensitivity curve, which is, of course, caused by the test conditions and is not a peculiarity of the true sensitivity, could conceivably occur in any H.E. It is certainly not restricted to NQ although it may be restricted to H.E. which can exhibit LVD. It is not certain that all solid explosives can do this, but it does seem clear that a large grain size and a small charge diameter favor the appearance of LVD.

There is an example in the recent literature¹,²¹ in which the gap test results on a coarse (400 μ tetryl) closely paralleled those on NQ (HBD). In a small scale test (d = 12.7 mm, charge unconfined),







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TABLE 12

MOST RECENT SHOCK SENSITIVITY TESTS ON NQ

			<u>_50% I</u>	oint
NQ	ρ _o g/cc	TMD	No. <u>Cards</u>	Pg (kbar)
x 589	1.4371	80.7	80	58
	1.4391	80.8	79	58.5
	1.5161	85.1	68	65
	1.3881	78.0	97•5	50
x 588	1.3871	78.0	109	46
	1.4401	81.0	98	50
	1.5041	84.5	81	57.5

there was a decided peculiarity in the shock sensitivity curve (same qualitative direction as that of Fig. 13), and there was an <u>apparent</u> reversal in sensitivity ratings of the fine and coarse tetryl when the small rather than the large scale test results were used. The close parallelism with the behavior of NQ¹ coupled with the Ref. 17 results: (a) that tetryl (400 - 630 μ) exhibits LVD and (b) that at $\rho_{\circ} = 0.9$ g/cc, its d_c value is about 12 mm, i.e., approximately the gap test diameter used in Ref. 21, make it possible to explain these tetryl results in the same way as those for NQ.

SUMMARY

Recent detonation velocity data for NQ at high (1.7 g/cc) and low ρ_0 lie on the currently used ideal curve down to $\rho_0 \sim 0.3$ g/cc. Below this density the velocities lie above the straight line and appear to vary slowly with density to the terminal value of 2.05 mm/µsec computed for $\rho_0 = 0.01$ g/cc.

Work with NQ (LBD) showed that the diameter effect on D is small for this material, particularly at high densities. Its failure curve and detonation velocity pattern were typical of Group 1 explosives.

The high bulk density NQ also showed Group 1 behavior in the density range of 1.3 to 1.6 g/cc ($d \ge 5.08$ cm), coupled with failures to initiate at $\rho_0 > 1.62$ g/cc (dead pressing). However, NQ (HED) exhibited diameter- and density-dependent pseudo-detonations (LVD) in the density range of 0.7 to 1.1 g/cc. The functional relation between LVD, ρ_0 , and d differs markedly from that of D, ρ_0 , and d. In addition, the limit curve for d and ρ_0 in LVD shows the opposite trend to that of the failure curve in detonation, i.e., it shows that the critical density for LVD increases with increasing diameter.

It is to the existence of this LVD region that a previously unexplained hump in the shock sensitivity curve of NQ (HBD) is now attributed. [Although the LVD is of lower velocity than detonation, its driving reaction is frequently sufficiently powerful to damage

the steel witness used in the gap test.] Grinding NQ (HBD) to a finer material (and hence decreasing its critical diameter for detonation) resulted in a shock sensitivity curve showing no hump (Fig. 14).

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APPENDIX A

ADDITIONAL INFORMATION ON THE LOTS OF NQ USED

Sieve analysis is inappropriate for fine materials of high ℓ/d such as the NQ (LED). As remarked in the text, X547 was in the form of hollow needle-like crystals. A typical photomicrograph appears in Ref 10.

The Ro-Tap Sieve Analyser was used for the NQ (HBD) although, as the results show, this is not an ideal method of gaging grain size. Table Al contains the data which show 50% weight median values of 106, 95, and 117 μ for lots X530, 589, and 588, respectively. It seems highly unlikely that X538 (ground X589) is actually coarser than X589. Figs Al-3 are photomicrographs of X588 showing the large amount of fine material (< 10 μ); the coarse material is comparable to that of the original, unground NQ. Moreover, X588 detonates as a finer material than either X589 or X530, and X530 as finer than X589, but in both cases screen analysis indicated the reverse in relative fineness. Factors such as an unfavorable t/dand electrostatic retention of fines are evidently distorting the mean size as determined by the Ro-Tap. Under these circumstances, there seemed little point in running additional sieve analyses.

TABLE A1 RO-TAP SIEVE ANALYSES ON NQ (HBD)

Median	Bize, µ	100	106	95	ζττ
:Mesh(µ)	Total	96 . 9	99.5 97.2 98.3	9.99.0 99.0	99.8 99.4 99.6
₩	Pan	12.2	12.2	10.2 9.4 9.8	11.0 6.0 8.5
44	325	4.0 15.3	4.0 4.1 13.9	5.0 5.4 10.4	5.8 4.0 8.9 8.9
53	270	7.0 19.3	7.1 7.0 7.0 17.9	8.7 8.1 8.4 15.8	7.4 5.3 12.9
62	230	7.6 26.3	7.6 7.5 24.9	15.2 9.2 12.2 24.2	7.9 5.5 6.7 19.3
74	200	17.0 33.9	17.1 17.0 17.0 32.5	18.8 18.9 18.0 76.4	19.1 14.0 16.6 26.0
105	140	25.5 50.9	25.2 25.8 25.5 49.5	22.5 18.1 20.3 55.2	22.9 17.9 20.4 42.6
6' 1 '	No.: 100	23.6 76.4	26.3 23.7 25.0 75.0	19.5 29.5 24.5 75.5	25.7 48.4 37.0 63.0
	% on sieve	Я А С	Av < d:	Å ∧ d:	A A d
Sample	(300I)	н	н Q	ы и	н 0
	Material	X510	X530	X 589	x588 (x589 ground)

NOLTR 67-169







APPENDIX B

SUPPLEMENTARY DATA

Table Bl contains the optical details for each of the 70 mm smear camera records obtained. Table B2 contains the velocities obtained by fitting the position-time (x-t) data to the various functions:

- (a) linear
- b) hyperbolic (contains cross product xt)

(c) quadratic

(d) Spline fit (sequential cubic segments, continuous first and second derivatives.)

Whenever such a fit is made, the overall velocity for the interval can be found. The linear fit (a) has been used for all velocities reported in the tables of the text of this report. As Table B2 shows, from the sampling so far made, the overall velocity from the spline fits agrees very well with the velocity from (a) although there is no question that the spline treatment seems to give the best fit of the four to the actual x-t data.

In deciding whether the measured velocity is constant over the 63.5mm interval chosen near the end of the charge, we need some way of estimating the departure of the smear trace from a straight line. We have found by working with straight edges that variables in the record reading and reducing procedures can in troduce as much as 2% change in the velocity. We can compute that our use of a curved initiating shock will introduce up to 1% velocity change at the largest diameter (7.62 cm). Various other factors such as roughness and heterogeneity of the charge surface, over- or underexposure of the film, and stability of the front itself have effects that we have not yet been able to assess. It should be noted that for most of the shots in Table B2, the camera viewed the last six inches of the charge because the response (constant or attenuating velocity) was unknown. Consequently the time resolution is only about half

that attainable when a length only 2.5 in. of charge is viewed; 2.5 in. is used whenever a constant velocity front is expected. The poorer time resolution (6 in. view) very probably contributes something to the larger percentage changes observed. The absolute value of the change as well as its percentage value must also be kept in mind; random errors would be expected to show larger percentage effects at the lower velocities.

Again from the sampling, methods (b) and (c) generally measure about the same percentage change. The decision as to whether this indicates constancy or failure is still subjective . Up to 3% variation can be accounted for. Thus shot 305 (14%) is undoubtedly failing, shots 310 and 318 (2.7-3.2%) probably detonating, and shot 279 (6%) is also considered a detonation because of its location in the D vs ρ_0 family of curves in Fig. 8. It is possible that the factor of 2 difference in the velocity change for shot 279 as compared to shots 310 and 318 could arise because the latter two charges were prepared hydrostatically and consisted of four twoinch long pellets whose camera smear records are consequently made up of four arcs instead of the single one from the isostatically pressed charge No. 279. Moreover, the longitudinal density variation and hence detonation velocity would differ within the charges prepared in the different presses. Finally, any charge near its failure limit would be expected to show more instability of its front than a definitely super-critical charge.

In the low velocity regime, we are faced with both the lower absolute values of U_{LV} and the metastability of the disturbance. The latter fact means that propagation of the low velocity front will be much more sensitive to charge heterogeneity than would a detonation. Again our decision of constancy or failure has to be made, in part, from the U_{LV} pattern of Fig. 10. Thus because of the respective locations, Shot 288 (7.2%) is more apt to be a constant velocity than Shot 289 (4.4%). It is evident from the velocity changes shown in Table B2 that low ρ_0 and low d both

enhance the possibility of U_{LV} . The exact point at which ρ_0 and d have increased to the extent that this particular NQ(HBD) no longer exhibits a metastable reaction, but an unquestionably <u>unstable</u> one, cannot be selected without doing much more work on longer charges. The large values of some of the velocity **changes** (Table B2) suggest that some of the U_{LV} shown at higher ρ_0 and d in Fig. 10 are in fact attenuating shocks; if so, the limit curve of Fig. 12 should be shifted to the left. On the other hand, the **change** in velocity found for the more dubious cases is comparable to that found in detonation near the citical density (Shot 279); this suggests that the figures as now shown are correct.

The important result of the present work is the demonstration of the existence of an extensive low velocity region, not the exact boundaries of this region in the $d-\rho_0$ plane. For reasons discussed above, these boundaries would be very difficult to establish, would be of no theoretical interest, and would be almost impossible to verify (reproduction of a failure limit of a metastable phenomenon). Consequently, no further work on this topic is planned. TABLE B1

EXPERIMENTAL CONDITIONS FOR THE SHOTS

Writing Speed mm/µsec		N (N (N	م	เดดด	เดดด	20	N 0	ରା ରା ରା ର	101 0 1	໙	2	ц су M
Length Observed in.	ഗരഗ	oo (9	୰୰୰	୰୰୰୰	००	୰୰	୰୰୰୰	o o	9	9	ວ ບົງ
Flasher a	NN	i			NN	ZN	NN	NNEE	W W	M	M	
Shot No.	326 327 328	276 291*	277	292 * 306 316	203 305 310	330 318	279	282 281 281 281	280 294	303	309	268 270 288
50 C												
Writin Speed mm/µse	サエエ	ц м м м м	ŝ	44	0 0 0	V (V	Ч	ิงงง	0 0 H		0 0	ิดดด
Length Observed in.	~~~ ~~~	พ พ พ พ พ พ พ พ พ พ พ พ พ พ พ พ พ พ พ	ູ້	IJ	୰୰୰	7.5	7.5	7.5 7.5	~~~ ~~~~	7.5	5-2-	~~~~ ~~~~~
Flasher a	:	N N Lacquer	N	თ თ		M	М	NN	M	N	N M	zvz
Shot No.	*22 *82 *62	103* 105* 105*	107*	80 * 81*	545 455* 540	133	138	241 184 159	213 215 162	175	176 214	216 157 163

NOLTR 67-169

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Panatomic-X film used here; Tri-X film was used in remaining shots Writing mm/usec Speed Cellulose acetate used unless otherwise specified. Symbols 0000000000 t M N N キキキ are: M Magic Tape, N none, S Scotch tape. Observed Length ับ เภอออ ທີ່ທີ່ທີ່ ທີ່ມີທີ່ທີ່ in. Flasher ർ $\Sigma \Sigma \Sigma$ 222 ZZ Shot No. 395 395 395 411 409 409 с С * Writing mm/µsec Speed N HUHHUHHUHH **MHH** Observed Length S S in. ມູນ າບຸ່ມ 5000 005 0 6 0400000000000 6 Flasher ZZZZ ർ ZZZZZZZZZZ Σ Shot No. 313

TABLE B1 (Con't)

TABLE B2

RESULTS OF VARIOUS TREATMENT OF x.t DATA

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			ULN (mm/	/usec)		% Char	lge**	
Shot No.*	ရာ ရာ	0 0	Linear Fit*	Hyper. Fit	Spline	Quad.	Hyper.	
264 263	2.54 2.54	0.72 1.001	2.349 2.500	2.349	2.357	+3.1 - 0.1	-0.4	
268 270 288 289		0.735 0.735 0.851 1.101	2.942 2.874 3.188 3.109	2.959 2.959 2.068 2.058	2.949 2.841 3.174 3.092	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	+ - - - - - - - - - - - - - - - - - - -	
276 291 293 293	0000 0000 00000	0.718 0.851 1.001 1.102	3.211 3.554 3.735 3.735	3.212 3.492	3.213 3.552	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-0•0 -6•4	
	Deto	nating cr	Failing	Near Cri	tical Dens	itty		
305 310 279 279	بریزی 0.000 0.000 0.000 0.000 0.000 0.000	1.200H 1.251H 1.291H 1.2991	F(4.88) 5.579 6.518 6.518			-14.0 		
* Length o 270; for	f charge all oth	observed er shots,	by camer this len	a was 2. ngth was (5 in. for 6 in.	shots 26	3, 264,	ane

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** Change in $U_{\rm LV}$ over the portion of smear record which was read; this portion of the record corresponds to the last 63.5 mm of the charge.

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In its range of detonability Group 1 explosive. In addition TMD $(> 94\%$ at 5.08 cm diam) as lower critical density. The his exhibits a critical diameter all density form. This fact aided NQ (HBD) where a strong shock p constant velocity front. This characteristics similar to the materials. That trend, the pow dimensions of the various gap to a hump in the curve 50% pressun gap test and (2) a reversal in NQ(HBD) and NQ(LBD) when tested tests. (ty, nitrogua h, it exhibi well as the ligh bulk den bout three t in studying produces a s pseudo-deto detonabilit ver of the L tests can be re vs % TMD <u>apparent</u> sh i on the lar	nidine ts failu more co sity (H) imes that the sul ubdetona nation o y limits VD react combine obtained ock sens ge- and	(NQ) behaves as a ure at the high ommon failure at a BD) form of NQ at of the low bulk beritical region of ation, supersonic, or LVD had failure s of Group 2 tions, and the ed to explain (1) d for NQ(HBD) in th sitivity rating of small-scale gap
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	Low velocity detonation								
	Detonability								
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