EXPLOSIVE-DRIVEN HEMISPHERICAL IMPLOSIONS
FOR GENERATING FUSION PLASMAS

by

D. Sagie and I. I. Glass

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Summary

The IITIAS explosive-driven-implosion facility was used to produce stable, centered and focussed hemispherical implosions to generate neutrons from D-D reactions. A high resolution scintillator-detection system measured the neutrons and γ-rays resulting from the fusion of deuterium. Several approaches were used to initiate fusion in deuterium. The simplest and most direct proved to be in a predetonated stoichiometric mixture of deuterium-oxygen. The other successful method was a miniature Voitenko-type compressor where a plane diaphragm was driven by the implosion wave into a secondary small spherical cavity that contained pure deuterium gas at one atmosphere. A great deal of work still remains in order to measure accurately the neutron flux and its velocity distribution as well as the precise interactions of the neutrons with the steel chamber which produced the γ-rays. Nevertheless, this is the only known work where fusion neutrons were produced by chemical energy only in a direct and indirect manner.
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1. INTRODUCTION

The physics of thermonuclear reactions was studied intensively during and after World War II. During the last 40 years numerous papers were published giving the main parameters of several known thermonuclear reactions. Glasstone & Lovberg (1) gave a review of known results, which includes the collision cross-section of the fuel, by assuming a particle Maxwellian-energy distribution as a function of its temperature in a thermonuclear reaction. Collision cross-sections for D-D, D-T and D-He are shown in Fig. 1.1. It is clear that up to a temperature of a few keV the reaction intensity would be a very strong function of temperature. Therefore, the first goal of the explosive-driven implosion project was to reach this range of temperatures, rather than any other thermodynamic parameter. Since the nuclear fuel cannot be confined for an unlimited time, the actual period of confinement has a great influence on the efficiency of fuel burn up. Two main approaches to fusion are under intensive study in the scientific community. In "magnetic confinement" (2) the plasma is held by magnetic fields in the desired configuration for reaction times large (up to 1 s) compared to its disassembly time at the speed of sound, or the particle thermal speed. In "inertial fusion" (3) the reaction confinement is essentially at the sound speed or thermal disassembly time ($3 \times 10^{-11}$ sec at $T = 1$ keV for $p_1 = 10^6$). The numerical expression of the confinement term is given by the "Lawson criterion" (4), which is the product of the particle density and confinement time required to achieve an energy balance at a specific temperature. For example $n_0 > 10^{15}$ sec/cm$^3$ at 100 keV ($10^9$K) for a D-D reaction.

The present report deals with an investigation of the application of explosive-driven implosions to produce thermonuclear reactions. This approach basically belongs to the inertial-confinement family. Most of the current scientific activity in this area is dedicated to laser fusion. Fortunately, this work also consists of spherical implosions, so that the principles of this method and explosive-driven implosions are quite similar. Important research on laser fusion is being conducted at the University of California, Lawrence Livermore Laboratory (5), the University of Rochester, Lasers Energetics Laboratory (6), and K.M.S. Fusion, Ann Arbor (7). Their reports were of great interest and assistance during the present study.

Spherical sectors rather than full spheres were frequently used by Russian researchers to produce fusion reactions. Bogolyubskii et al (8) accelerated a polyethylene diaphragm (10\(\mu\) thick) up to 60 km/s by using a relativistic electron beam. They produced $3 \times 10^6$ neutrons with D$_2$ fuel contained in a conical lead capsule. Ziolkowski et al (9) used a solid explosive to implode a conical liner onto the surface of a copper cone. The generated shock wave was strong enough to accelerate a polyethylene layer up to a velocity of 50 km/s. The collapse of this layer in a gold cone filled with deuterium generated up to $3 \times 10^7$ neutrons. Recently, Anisomov et al. (10) reported that temperatures of 0.3 - 0.5 keV and a yield of $10^6$ neutrons had been achieved by using an explosively accelerated metal liner (up to 5.4 km/s) striking deuterium fuel contained in a conical lead capsule. For the present work the UTIAS hemispherical implosion chamber (11 - 17) was used in several modes to produce implosions appropriate for fusion.
The spherical implosion chamber was conceived by Glass (12) in the 1950's. Since then it has proved to be a unique facility to generate stable implosions in a safe and reusable facility in order to provide a small region of extremely high pressures and temperatures (13-25). The implosion chamber had originally been used as a driver for a projectile launcher (15), for generating strong shock waves in a constant area shock tube (16), and for the production of diamonds from graphite (17). Recently it was applied to producing fusion reactions in deuterium, as reported herein.

The principle of operation of the UTIAS implosion chamber is described in detail in the above-mentioned reports. In brief, PETN solid explosive is used to form a hemispherical shell (3 mm to 6 mm thick) in a 20-cm diameter hemispherical cavity milled in a massive steel chamber (Fig. 1.2). The remaining volume is filled with a stoichiometric mixture of H₂ or D₂ and O₂. This mixture is detonated by a very short thin exploding wire located at the geometric centre (Fig. 1.3). The arrival of the detonation wave at the spherical surface instantly and simultaneously fires the explosive liner. The detonation wave in the explosive liner hits the metal cavity, reflects, and implodes on the preheated burnt gases, focusses at the centre of the hemisphere (∼50 μsec after initiation of the exploding wire) and reflects, leaving behind a very small pocket (1 mm³) of extremely high-temperature, high-pressure and high-density plasma. In the present study H₂ is replaced by D₂. For some other fusion experiments this focus takes place on top of a base of a small capsule containing D₂ at ∼1 atm (Fig. 1.4). The cavity in the capsule serves to produce a second compression and heating stage in order to produce extremely high temperatures, pressures and densities in the deuterium plasma.

In the second section the experimental set-up is explained. Special consideration is given to the neutron-detection system. The third section describes the experiments that were conducted and their results. The fourth section discusses the results and lists the main conclusions. In Appendix A1 consideration is given to several approaches for the application of explosive-driven implosions to fusion with advantages and disadvantages of each approach. In Appendix A2 a review is given of the neutron detection system. In Appendix A3 some consideration is given to the possibilities of scaling for better performance and neutron yield.

2. EXPERIMENTAL EQUIPMENT AND INSTRUMENTATION

2.1 Driver Description

Basically, the experimental facility consists of the UTIAS implosion chamber (Fig. 1.2). The principle of operation of the implosion chamber is as follows. The explosive material forms a hemispherical shell (3-6 mm thick) in the 20-cm diameter hemispherical cavity milled in a massive steel back plate. The explosive shell of PETN weighs up to 200g and has a total energy of up to 1.2 megajoules. The remaining volume is filled with a
stoichiometric mixture of H\(_2\) and O\(_2\) with a pressure of up to 70 atm and an energy of up to 1 megajoule. This mixture is detonated by a very short (~1 mm), thin (0.127 mm dia) exploding nickel wire located at the geometric centre. The arrival of the detonation wave at the spherical surface instantly and simultaneously fires the explosive liner. It reflects as an implosion wave. The detonation wave in the explosive liner hits the metal cavity, reflects, overtakes the first implosion wave and combines to form a final imploding shock wave in the preheated gases. It then focusses at the centre of the hemisphere. When the implosion reflects at the centre it leaves behind a zone of extremely high pressure and temperature. If the H\(_2\) is replaced by D\(_2\) then the stage is set for D-D fusion at this focus. Alternatively, a small hemispherical capsule containing D\(_2\) at a few atm can be placed at the centre, with its base closed by a diaphragm and facing the focus (Fig. 1.4). This is a miniature Voitenko-type (36) compressor which brings the deuterium to extremely high pressures, temperatures and densities required to initiate a fusion reaction. Other set-ups using a small conical capsule or in combination with projectile compression have also been used but without successful results so far.

2.1.1 Implosion Chamber

The UTIAS implosion chamber was conceived by Glass (12) and has been used frequently for several applications in the last few years. It was used to drive projectiles to hypervelocities (15), to produce very strong shocks (16) and to convert graphite to diamond (17). The chamber itself was essentially the same for all these applications. The main modifications relate to a particular experiment. Briefly, the implosion driver consists of two halves, a rear plate and a front plate (Figs. 2.1 and 2.2). A 20-cm diameter, hemispherical cavity is machined in the rear plate and contains the explosive-liner package. The front plate consists of a cone which supports the barrel-capsule assembly and the liner plate. A 20-degree conical liner plate is placed on the top of the liner plate to give extra protection for the front plate section against any possible damage from an off-centred implosion (20). In the front plate, O-ring grooves are machined for pressure and vacuum sealing. Connections for the barrel, high-voltage feed-throughs for bringing in the ignition wires, and a pressure tube for venting and filling the chamber are installed in the front plate as well. The previous design was slightly modified to reduce possible damage during a run, and to ease disassembly afterwards.

2.1.2 Explosive Package

The explosive package that was used for the present experiments is described in detail by Saito (21). The preparation procedure is the final result of successive developments of the explosive-packaging technique which was studied by several researchers (15, 20, 22). Uniformity in thickness and especially in density is of prime importance to achieve centred, symmetrical implosions. The present scheme of initiating the explosive (superfine PETN) by a gaseous detonation wave in stoichiometric mixtures of hydrogen-oxygen or deuterium-oxygen at initial pressure of 14 to 70 atm, proved to be the most useful and was employed in all the runs in the present study. It was possible to
obtain nearly perfectly centred implosions from the reflected detona-
tion of the gas mixture alone. This indicates that the original
detonation wave was stable, symmetric, and well focussed with respect
to the origin and therefore could provide a symmetric source for
initiating the explosive liner.

To provide sufficient mechanical binding strength for the explo-
sive to be self-sustained, the superfine PETN powder was mixed with
1.5% by weight of cotton fibers (linters) and water to form a slurry.
A plastic honeycomb was glued to the accurately machined hemispherical
copper liner (Fig. 2.3). Then the slurry was worked and pushed into
the honeycomb to fill all air pockets. The cotton fibers serve as
a binder providing the powdery PETN with very positive support, while
the plastic honeycomb provides the support that allows the wet PETN
slurry to dry while retaining the proper geometry. In this manner,
the explosive package had sufficient strength to withstand handling
and was also sufficiently homogeneous to ensure a stable, focussed
implosion. The 20-cm diameter copper liner, with a 20-degree conical
liner, and a 6-mm thick honeycomb, contained about 200g of explosive.
The use of additional explosive was not advisable owing to possible
chamber damage.

2.1.3 Ignition System

In order to induce a spherical detonation wave in a combustible
mixture, sufficiently large amounts of energy must be released at a
very high rate. The initiation point must be located exactly at the
origin of the hemispherical explosive package to produce a symmetric
implosion. For a given amount of stored energy of the ignition capac-
itors, this criterion could be satisfied by keeping the resistance of
the exploding wire as high as possible, and the impedance of the rest
of the ignition system as low as possible. The system as used by
Roig (24) and Vasudevan (25) was adopted with minor modifications. A
0.127-mm (5-mil) diameter, 1-mm long nickel wire was used to ignite
the explosive-gas mixture. The transmission line of the high-voltage
circuit was a low inductance (16Ω) impedance) cable. The discharge
system consisted of a spark gap and a 1.0 μfd, 30 kV capacitor, as
shown in Fig. 2.4. The spark gap essentially consists of two brass
electrodes, a Plexiglas tube and an automobile spark plug. The oper-
ating cycle of the system included dehumidifying the spark gap with
compressed nitrogen and then charging the capacitor to 22 kV. A 5 kV
trigger pulse to the spark plug was used to initiate the discharge.

While the exploding wire should be placed exactly at the focus,
it's own physical dimensions, with the terminals and insulation,
practically eliminated that ideal position. Two designs were used
as shown in Fig. 2.5. The exploding wire was located about 1 mm and
0.2 mm above the origin for the first and second design, respectively.
Some other deviations from symmetry were introduced by the terminals
and insulation but these undesired materials are all evaporated
before the implosion arrival, causing some impurities to exist near
the focus. A photograph showing the assembly over a deuterium capsule
appears in Fig. 2.6.

2.1.4 Vacuum and Gas-Mixture-Inlet Systems

A schematic diagram of the pressure and vacuum manifolds to
the implosion chamber is given in Fig. 2.7, and the control panel is shown in Fig. 2.8. The control room has 12-inch concrete block walls, sand-filled and reinforced with steel rods. After assembly of the implosion chamber, the operation of the entire system, including the monitoring of pressures, vacuum pumps and the gas filling can be remotely controlled at the control panel in the control room. The control panel is designed for ease of operation and to eliminate accidents or damage to the instruments caused by following a wrong sequence of operation. The gases used were commercial bottles of compressed gas. They are shielded by heavy metal from direct damage through a failure of any connections. The \( \text{H}_2, \text{O}_2 \) or \( \text{D}_2 \) inlet is easily controlled by the operator who has just to open the appropriate valve.

2.2 Deuterium Capsule

As mentioned above some indirect approaches were tested as well. For those experiments pure deuterium was used, contained in a small capsule at the driver focus.

The main characteristics of the capsule were as follows: to contain the deuterium at the desired pressure, to maintain a good seal without a gas leak, to prevent impurities during assembly or the filling process, and to have the compression surface of the diaphragm located initially at the implosion focus. It must also provide a convenient way of filling the deuterium, maintain the desired geometry as long as possible during the gas-compression process, and reflect the "escaping" radiation back into the target. The design of the capsule was made in accordance with these requirements.

2.2.1 Capsule Design

A typical capsule design is shown in Fig. 2.9. Generally, the capsule was made of 440c stainless steel hardened up to 48R. For the design of a cone it was premachined with an apex radius of about 50 \( \mu \text{m} \), and the diaphragm was of 0.005" cold-rolled stainless-steel sheet, plastically formed to get the spherical cap at its centre. The diaphragm circumferential flat area was indium coated to provide a soft medium which served to seal the diaphragm against the main capsule body when firmly pressed with the fastening nut. A standard capsule having a hemispherical or cone basic diameter of 3.2 mm dia was capable of maintaining an interior gas pressure of 70 atm. The diaphragm was strong enough to withstand the \( 2\text{H}_2+\text{O}_2 \) detonation pressure without damage, as shown by primary experiments (3.1.2). Several experiments (3.1.3) also showed that this design was capable of containing the deuterium for at least 24 hours without a detectable leak. The capsule described above is easily modified to check the dominant parameters noted previously.

2.2.2 Preparation and Filling Procedure

The filling system for the capsules was developed, built and tested by Dr. A. K. Kudian. The system (Figs. 2.10 and 2.11) consists of a small vacuum chamber in which the open capsule is located. The fastening nut with the diaphragm inside are located in front of the capsule, fixed by two pins to a rotating shaft, and can be threaded in externally through a seal assembly without breaking the vacuum in
the chamber. The chamber itself is metal sealed and can be heated up to 400°C for degassing of all gaseous impurities that have penetrated into the capsule material. The organic seal of the rotating shaft is protected by a water cooling system. The chamber is evacuated by a primary rotating pump (Duo-Seal, Ref. 26) and secondary 2" diffusion pump (SpeediVac, Ref. 27). Two liquid nitrogen cold traps, between the rotary and diffusion pumps and between the diffusion pump and the chamber, provide further evacuation to less than 1 μ. A deuterium cylinder (Matheson, 28) with a maximum impurities level of 0.5% is connected to the system by high-vacuum standard copper tubing. With the capsule components all located inside, the chamber is pumped down to about 1 μ, then it is heated up to 200°C and maintained at that temperature for a few hours. After cooling down, the chamber is flushed by deuterium at 1 atmosphere and after a second evacuation is filled finally to the required pressure. The capsule is then closed firmly, the chamber is vented and then the capsule is taken out ready for use. McLeod and Pirani gages were used for precise vacuum measurements while gages installed on the deuterium-cylinder regulator were used to fill to the required pressure.

2.3 Neutron-Detection System

Among the few approaches that are available to detect a thermonuclear reaction, the detection of neutrons seems to be the most convenient. The neutrons being uncharged particles are the only particles that can penetrate the heavy-walled chamber that is required to confine the explosion inside. Their detection provides a positive unquestionable proof of a fusion reaction. Two independent neutron-detection systems were designed to be used simultaneously. The first is a scintillator-counting technique which is very sensitive and reliable. It makes it possible to detect both neutron intensity and their time of flight which is an indication of the neutron energy. As a second independent measurement, an indium activation technique was chosen to confirm the intensity of the neutron flux.

2.3.1 Scintillator Detection System

This system should cope with the following main constraints:

1. Neutrons are to be detected in a single pulse with a duration in the range of 10 nsec. This imposes on the electric circuit problems of transmitting and recording with low distortion, as well as very high reliability.

2. In order to record a signal shape with the above duration range, the oscilloscope is limited to a total detection time of about 1 μsec.

3. All the events in the combustion process previous to the neutron generation are not precisely fixed in time to provide a signal in advance that can trigger the detection system. Therefore, the system must be self-triggered by the neutron pulse itself.

4. The scintillator and the photomultiplier (PMT) constantly emit signals into the system. The scintillator is activated by high-energy particles moving around in the atmosphere (especially mesons) with a typical energy of 140 Mev. (Some cosmic particles carry energies of more than 1000 Mev.) The noise has a statistical voltage and flux distribution. Recalling that the neutrons
generated by D-D fusion have an energy of 2.45 MeV, the energies of the cosmic particles are much higher and, on average, would correspond to about 50 neutrons activating the scintillator at once. Shielding against these particles would be very heavy and expensive and therefore it was not practical to construct. The particles intersect with the scintillator at a rate of about 1 per second. In addition, electrical noise is an integral part of the operation of a photomultiplier (Fig. 2.12). The high potential that is maintained between the photocathode and the first anode attracts the electrons which are always "escaping" from the photocathode material. (This material is chosen a priori to have very loose electron bonds.) These electrons initiate the PMT amplification chain and in this way send a statistically distributed signal into the recording system. These dark current signals become less frequent for higher voltage levels (Fig. 2.12c). For our specific model of PMT, they are almost stable at a frequency of about 80 Hz, and a voltage level which is equivalent to a photon energy of about 1 MeV interacting with the scintillator material. The frequency of this noise as a function of its voltage is given in Table 2.1.

5. The ignition system that initiated the detonation-implosion cycle, radiates very high intensity electromagnetic waves. False triggering of the system by this noise must be eliminated.

The system is schematically shown in Fig. 2.13, while pictures of the main components are shown in Figs. 2.14 and 2.15. Technical specifications of the components are represented in Appendix A2. Two scintillator-photomultiplier assemblies are located in front of the implosion chamber. The first one is 30 cm from the neutron source. It provides for a low-speed sweep and a means of triggering the system. The second assembly is 80 cm from the source right behind the first one. The implosion chamber provides 10 cm of steel protection around the implosion space and has a steel window 2.5-cm thick thereby making a reasonable path for the neutrons (see Fig. 2.13). Both detectors are located within the solid angle emerging from this window. The short distance of the scintillators from the chamber increases the detection sensitivity. However, the close proximity may distort the signal. This short distance makes it possible for scattered neutrons and neutron-activated products from the steel chamber to reach the detectors. If a large enough signal could be obtained the detectors could be shifted further away from the chamber.

The recording system is based on two Hewlett-Packard Model 1744A single beam storage scopes, with a maximum frequency of 100 MHz and a sweep rate up to 5 nsec/div. Due to a special feature in the scope, it displays the entire triggering pulse with a 50 nsec backview. With a nominal writing speed of 1800 cm/μs, the oscilloscopes are capable of storing a 10 nsec single-shot sine wave with an amplitude of eight major divisions (full scale). The concept of the circuit operation is illustrated in Fig. 2.13. The recording system is armed and then it is ready to record and store a single signal. The detector assemblies are in an operating state. The ignition pushbutton, boosted by an amplifier, triggers scope I, the adjustable delay unit, and the counter simultaneously with the ignition of the chamber detonating-implosing cycle. The sweep rate of scope I is 100 μsec for full scale, enough to cover the fusion event, which is expected at about 50 μsec. The adjustable delay unit opens the gate after 40 μsec and enables signals from detector I to trigger scope II, externally. (The signal passage is blocked by the gate up to this moment.) If a signal from detector I now shows up, it is recorded by scope I and also triggers scope II.
Scope II is connected to detector II and so displays signals from this detector during 500 nsec after it has been triggered (its total sweep time). If a sufficient flux of 2.45 MeV neutrons is generated, both detectors would be activated. The second detector would respond 15 nsec after the first one, owing to the longer distance from the neutron source. Another delay unit extends this time difference even farther, and together with the "back in time" view of the oscilloscope, we are able to see the output of detector II back to 85 nsec before the trigger signal. This is essential, as in this manner it is possible to display the complete signal and any X-rays if they exist. (The X-rays propagate at the speed of light and would arrive before the neutrons. However, the signal would be too small to trigger the system.) The second scope system is blocked by the gate from receiving any signal up to about 100 nsec before the arrival of the expected neutron signal. However, this is a drawback as several noise signals could trigger that system from this moment on. Therefore, the trigger level must be set high enough to reduce the chance of false triggering. For example, if we chose a point such that the noise frequency is about 100 Hz, the probability of false triggering would be 1/1000. If the sensitivity were increased by lowering the trigger level, then the chance of false triggering would increase significantly; and vice versa, decreasing the sensitivity will reduce the probability of false triggering. Triggering by noise would eliminate some experimental data but it could not be mistaken for a neutron signal, since there is almost no chance for any two independent noise signals to stimulate both detectors simultaneously. Only neutrons with a short period and high flux could do that, and therefore the coincidence of the signals on both oscilloscopes is positive proof of a neutron flux. The counter starts to count with the ignition signal and stops with the triggering of the second scope, and strengthens the evidence of getting the signal at the right moment.

The scintillator-photomultiplier assemblies are identical in their mechanical design, but different in their internal electronic circuit (Appendix A2). The first detector serves both to trigger the system, and to record events through the whole process. Its signal is distributed into two main lines. Therefore, its circuit is built to provide a relatively high-voltage amplitude, but it is quickly saturated. The signal, being very nonlinear, can hardly provide an indication of the number of neutrons generated. The second detector has a more linear circuit. The signal is directly connected to the storage oscilloscope with no other components on the line, except the delay unit which is purely resistive and therefore transmits the high-frequency (100 MHz) signal without any distortion. The area confined by this signal rather than its amplitude provides an accurate indication of the amount of energy transmitted to the scintillator, which in turn is proportional to the number of neutrons that passed through the scintillator material. However, for a low-neutron yield, while only a few neutrons intersect with the scintillators, both detectors provide a good estimate of the neutron yield (referring to the signal amplitude). If in future more neutrons can be generated in the chamber, the detectors circuit should be changed or preferably, the detectors can be moved away from the source. Then the scattering effect of neutrons and neutron products from the chamber hitting the detectors will be minimized, and the overlap of the two detectors will be eliminated.

In the present set-up the effective diameter of the first
scintillator, taking only the solid angle emerging from the "window" into consideration, is 3cm, and the distance from the neutron source is 30 cm. For the second scintillator the effective diameter and distance are 7.5 cm and 80 cm, respectively. The scintillator material is 5cm thick and 10 cm in diameter, so that almost every intersecting neutron delivers some of its energy. A single neutron is equivalent to about 1650 neutron yield for the first detector and about 2960 neutron for the second one. However, even if a single neutron indicates that a fusion reaction has occurred, it cannot give a reasonable indication of the total yield, since the energy is delivered to the scintillator in a wide energy distribution. Considering 10 neutrons as the minimum-required flux for statistical estimation, then the minimum neutron yield that can be estimated is about 16,000 neutrons for the first detector and 30,000 for the second one. The lower limit of the system sensitivity is set by the electron-emission noise as was mentioned before since for too low a trigger level, this noise might trigger the system instead of the neutron signal. A very serious problem is presented by the giant electromagnetic noise generated by the discharge of the capacitor in order to explode the ignition wire. These electromagnetic waves propagate in the air and excite the detector to an amplitude which is equivalent to a few hundred neutrons intersecting the detector simultaneously. These electromagnetic waves oscillate and damp down (Fig. 2.16) and should diminish before the gate is open. Their damping rate is controlled by the ignition circuit conductors, and the oscillating period can be reduced to 35 usec. Unfortunately, the ignition noise is so high that several gates were not able to block it from penetrating and triggering the system. The solution was found by introducing a discriminator unit, which has an output logic signal (constant voltage and shape), for every input signal in a wide range. Although the discriminator is not able to maintain the logic signal for the ignition noise, it reduces it significantly, enough to eliminate it from penetrating through the gate. At the same time low signals are transmitted with only small attenuation.

2.3.2 Neutron-Activation Technique

To get an independent indication of neutron production, and an estimation of the total number as well, a neutron activation technique is used simultaneously. Several approaches for our specific chamber set-up are shown in Fig. 2.17. They are all designed not to block the neutron path to the scintillator. The expected performance is listed in Table 2.2. Most of the suggested approaches are based on activation by the fast neutrons, while the activation cross-section for fast neutrons is considerably lower than that of thermal neutrons. We have found it more convenient and reliable to attenuate the fusion neutrons down to the thermal region and then to activate indium (Fig. 2.18). The attenuation is done by paraffin with a scattering cross-section of 45.5 barns per (CH2) unit for fast neutrons (29). Through the 7cm path about 90% of the neutrons are attenuated, while the rest are attenuated by the paraffin at the back of the indium and are partially reflected back into the indium. The indium itself captures thermal neutrons by the reaction \( \text{In}^{115}(n,\gamma)\text{In}^{116m} \) with a cross-section of 145 barns (30). For the thickness of 2mm, 65% of the crossing thermal neutrons are captured. Having an effective area ratio of \( 3 \times 10^{-3} \) to the total space, 500 neutrons are required for every single reaction in the indium. The final detection sensitivity depends on the radioactive background in the laboratory. Assuming that a rate of 100 disintegration per hour is detectable, and for the activated \( \text{In}^{116m} \)
with a half-life of 54 min, a minimum neutron yield of $10^5$ can be detected. The activated indium can be drawn outside the chamber assembly within a minute, while the other approaches require an hour at least for the activated material to be taken out of the chamber, which makes materials with a half-life of minutes unreasonable to use. The indium is quickly cooled down and can be reused again for detection.

3. RESULTS AND DISCUSSIONS

3.1 Primary Functional Tests and Calibrations

3.1.1 Neutron Detection System Calibration

The neutron detection system was fully described in 2.3.1. The logics, main components and constraints were introduced. Here we will become familiar with the actual performance of the components and the integrated system. A typical signal of electron emission from the photomultiplier is shown in Fig. 3.1. A typical signal from a Co$^{60}$ radioactive source which has γ-rays (photons) of 1.1 MeV is shown as well. The signals are similar and cannot be distinguished even by sophisticated electronic equipment. A Co$^{60}$ radioactive source of 1 µCurie has been used to calibrate the detector. This source, when located at the front of the scintillator, stimulates it to about $10^4$ signals per second. The display of the exposed detector on the storage screen shows the envelope of the whole spectrum, which corresponds to the photons that transmitted all their energy to the scintillator material (Fig. 3.1c). Hence the area surrounded by this envelope represents energy of 1.1 MeV, and is used as the basis to compare with the energy that any other process delivers to the scintillator material. For short duration signals (less than 20 nsec, like the inertial-confinement fusion), we refer to the amplitude, rather than the area under the curve. A Co$^{60}$ photon, while completely delivering its energy of 1.1 MeV into the scintillator, produces a signal of amplitude A. A neutron of 2.45 MeV, when interacting with a scintillator made of "Pilot B" material (31) with diameter of 4" and thickness of 2", delivers on the average 0.295 MeV per interacting neutron, or 0.23 MeV per incident neutron (32). Hence $B = 0.23/1.1 \times A$ is the average amplitude for a single 2.45 MeV neutron and the total neutron yield is given by the ratio of the actual signal to B.

An estimate of the neutron yield as described above assumes a linear amplitude with energy delivered to the scintillator. This assumption might hold only for a low-neutron flux, while for a higher flux the PMT becomes saturated. Therefore, the detectors were calibrated directly with a neutron source (with assistance from the Laboratory for Laser Energetics, University of Rochester, N.Y.). Some calibration displays of detectors I and II are given in Figs. 3.2 and 3.3 respectively, and the amplitude as a function of distance is given in Table 3.1. The decay of the amplitude which is supposed to decrease like $r^{-2}$ decreases actually like $r^{-0.2}$ for detector I and $r^{-1}$ for detector II, owing to the high-saturation effect of the photomultiplier at that high level of radiation. The neutron source for the calibration (33) is a particle accelerator that accelerates deuterium ions into a static deuterium target. The total number of $5 \times 10^5$ neutrons are distributed in about 2 usec as shown in Fig. 3.4. (The total yield was measured by Dr. A. Entenberg using a silver activation technique.) For our specific PMT (RCAR575) we found the
largest electron-emission signals to be of about 80% of the \( \text{Co}^{60} \) highest signals. As to the signals from high-energy particles that are moving in the atmosphere (like a meson), they are displayed in Fig. 3.5 together with the electron emission and \( \text{Co}^{60} \) signals. The signals are distinguished by their energy. Typical frequencies are 100Hz for the electron emission in 10% of the upper-voltage range, 10^4Hz for the 1u Curie \( \text{Co}^{60} \) source, and a few Hz for the atmospheric particles. As was mentioned before neither a commercial nor a laboratory-made gate was able to block the high-energy signal generated by the atmospheric particles and the ignition noise. The solution was found by introducing a discriminator unit which was supposed to produce a constant amplitude output signal for a wide range of input signals. The actual performance is shown in Fig. 3.6. While the discriminator does not succeed to keep high input voltages at the nominal output of 0.6v, it does cut it off considerably. The very rare input signal of 18v, which is the maximum expected from the detector, is transmitted as 1.6v and this is now low enough to be blocked by the closed gate. On the other hand, low-voltage signals are transmitted through the discriminator with low attenuation (0.8v input and 0.6v output) such that, when the gate is open, they can pass through and trigger the system.

The electron-emission frequency of the first detector becomes almost stable at a level of 80Hz, while the signal amplitude is about 1.2v (Table 2.1). After distribution and some decay this signal emerges from the discriminator at 0.5v. The 1.2v signal compared with the \( \text{Co}^{60} \) maximum signal of 1.5v corresponds to \( \frac{1.2}{1.5} \times 1.1 \text{ MeV} = 0.88 \text{ MeV} \) of delivered energy. Since the 2.45 MeV neutrons deliver an average of 0.23 Mev per incident neutron (32), an output of 1.2v from the first detector is equivalent to \( \frac{0.88}{0.23} \times 4 = 10 \) neutrons. For the 80Hz frequency of electron emission and for an average gate opening period of 10usec before the imploding wave collapses, the chance for a false triggering will be: \( \frac{80 \text{Sec}^{-1} \times 10^{-5} \text{Sec}}{1/1250} \). Actually we chose to set up the level to trigger the second oscilloscope at 0.2v (0.5v at the first detector), which corresponds to \( \frac{0.5}{1.5} \times 1.1 \text{ MeV} = 0.367 \text{ MeV} \) or 1.6 neutrons. In the present geometric set-up this is equivalent to a total yield of about 2560 neutrons. The electron emission for this level is 125Hz putting the chance of false triggering at 1:570. The frequency of the dark current is unstable and sometimes might be greater by a factor of up to three. Some effort was made to reduce the "noise-to-signal ratio". The P.M.T. was cooled down to -50° C but an improvement of about 30% was not enough to justify the inconvenience. A potentiometer was introduced to control the potential between the photocathode and the first anode (Fig. 2.16), and the noise-to-signal ratio was taken for several voltage difference in a wide range of total high-voltage inputs. An input voltage of 2000v was the optimum, and the first anode to photocathode potential made no significant effect on the noise-to-signal ratio. To summarize: the detection system was set up to trigger at a total yield of 2600 neutrons with a reliability of 99.8% for the electron-emission effect. As was mentioned before, a signal in that case will be definite proof of a fusion reaction, but for an accurate estimation of the neutron yield a statistical number of neutrons are required to intersect the detector. Ten neutrons at the first detector are equivalent to a total yield of 16,000 neutrons.
The neutron source of the Laboratory for Laser Energetics, mentioned above, was used to display the signals of single neutrons. Figure 3.7a shows the display of detector II when exposed to the neutron source at a distance of 30 ft. This same signal when taken at a faster sweep rate is shown in Fig. 3.7b. At this distance the number of neutrons is low enough that the contribution of single neutrons can be recognized. With careful adjustment, the tail of the last signal can be displayed (Fig. 3.7c). Here single neutron signals are clearly recognized. As for the higher flux of neutrons, the actual calibration with the neutron source (Fig. 3.3, 3.4 and Table 3.1) showed very quick deviation from linearity due to saturation. A testing of the components which compose the system showed that the "bottle neck" was in the PMT unit. To study the PMT range of linear operation in more detail, the scintillators were replaced by light-emitting diodes (LED) type MV2 of Litronix (34). The specifications of both the LED and the PMT are given in Appendix A2. The relevant character is that the LED emits a light with a wave-length of 572 nm at the peak (green light). At that wave-length the PMT relative response is 23%; or the photocathode emits 23% of the electrons which would be emitted for the same light intensity but at 380 nm, which is the PMT peak wave-length. The maximum emission of the scintillator is generated at a wave-length of 425 nm, where the PMT relative response is 94%. The chosen LED is the closest to the scintillator wave-length that is available, having $\eta = 0.244$ of the scintillator coupling efficiency to the PMT. Knowing the power output of the LED, the power-response character of the PMT can be derived experimentally. (Neutron-yield calibration would follow, since the energy delivered into the scintillator is known and the coupling to the PMT as well. However, for the duration of about 10 nsec expected in our case, this calibration cannot be carried out practically. The rise-times of the LED and its power source (pulse generator) are of 2 - 3 nsec duration each.)

Figure 3.8 shows typical PMT-outputs for several power inputs into the LED, while Fig. 3.9 shows the output signal for constant-power input but for different durations. Different results were received for the two PMTs. Although the PMT are identical, their electronic circuits are different. The results are listed in Tables 3.2 and 3.3. The saturation effect, especially of PMT of detector 1, is well seen. The PMT output is still lower with the voltage for a constant-emission duration. The parameters of the PMT electronic circuit can be modified to maintain linearity for higher light emission. However, at the present stage of the project, we found it more useful to keep the PMT linearity at the lowest intensity and, as was explained before, above some minimum level it is always preferable to move the detector a certain distance away than to reduce its sensitivity.

3.1.2 Diaphragm Strength*

The diaphragm used for several indirect designs must contain the deuterium gas through the ignition and detonation stages, up to the moment when the implosion shock wave reaches it. The exploding wire adds enough energy to the $^2\text{D}_2+\text{O}_2$ to initiate a detonation wave with a pressure ratio of 22-fold the initial pressure (Chapman-Jouguet condition). The diaphragm, located close to the exploding wire,

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* This set of experiments was carried out with the kind assistance of Mr. T. Saito, which is appreciated very much.
is exposed to unsteady pressures and its strength under dynamic conditions is not well known. Therefore, it is difficult to predict analytically its survival up to the imploding-wave arrival. The spectrograph with an eight-photomultiplier system (polychromator) (Ref. 21) was used to determine experimentally when the diaphragm collapses. After being detonated (Fig. 3.10a) the gas reaches a temperature of about 5000 K, and radiates over the wavelength spectrum. Eight specific wavelength intensities as monitored by the photomultipliers and are recorded as a function of time. While the diaphragm blocks the light path, no signal is recorded by the system, but when the diaphragm breaks the light can be transmitted and signals appear. Typical results are shown in Fig. 5.3.10b and c. After the oscillations caused by the ignition noise light penetrates the broken 2-mil thick stainless-steel diaphragm, and is detected at 6 wavelengths out of the 8 over the range of 4128A to 6328A. Then the imploding wave arrives, generating a much higher temperature which is recognized by a sharp negative peak. For the 5-mil thick stainless-steel diaphragm it can sustain the gaseous detonation and no light penetrates before the imploding wave arrives. With some safety margin taken into account, this is the minimum thickness for a stainless-steel diaphragm. The diaphragm experiments were carried out with a vacuum inside the capsule, but even for a 400 psi deuterium pressure, the deviation from the nominal thickness was small.

3.1.3 Capsule Functioning

Owing to the very high-diffusive character of deuterium gas, and the filling procedure of the capsule, which necessitates external fastening of the diaphragm, it was important to confirm that the capsule really can contain the deuterium for a few hours at least. In order to check this, the filling system described in Sec. 2.2 was used. The volume of the vacuum chamber was reduced artificially to a minimum, to increase the sensitivity of the measurement. The capsule was installed and the regular filling procedure followed. When the capsule was full and closed, the chamber was evacuated again and then the capsule was opened. The capsule cavity volume of 0.1cm$^3$ is only 1:38000 of the chamber and tubing volume. The initial pressure at the capsule (100, 400 and 1000 psi) was enough for the equilibrium pressure to be clearly detected by the vacuum gages. The capsule was first filled up to 100 psi and immediately released for calibration. Then the procedure was repeated with 400 psi and 1000 psi pressure with immediate release and 24 hours of waiting. The indium plating that provides the metal seal was a subject of some concern, since its melting point is 156°C, while the procedure includes heating to 400°C for a few hours under high vacuum. At first we found indium traces inside the capsule. Therefore, the degassing temperature was reduced to 200°C and the indium was restricted to a narrow zone at the edge of diaphragm, with no indium plating on the capsule body. In all experiments, the capsule maintained its full pressure for 24 hours with no detectable change.

3.2 Fusion Experiments

A total number of 2 experiments were done during this study, as given in Fig. 3.4. The table includes a variety of designs, direct and indirect approaches as they were developed. The detailed design for each experiment is indicated by an appropriate figure in Figs. 3.11 to 3.17. Unfortunately, a completely reliable neutron detection
system was not available until December, 1980, owing to the necessity
to develop electrical components capable of coping with large noise
signals, and a lack of appropriate display equipment. Therefore, for
some of the earlier experiments, the present results are not decisive
and should be repeated.

Typical craters as produced by the imploding shock wave are
shown in Figs. 3.18 to 3.25. The final shape and dimensions of the
craters are a combination of several effects such as the pressure of
the first imploding wave, melting of the steel due to primary heating
by the shock wave and then heating by the very hot gases. A lot of
material is also ejected. A detailed study of crater generation can
be found in Ref. 35. The initial shape at the focus which is different
for each design also plays an important role in the final shape of the
crater. Therefore, the crater shape and dimensions cannot be used for
quantifying the imploding shock strength and focus. Only a rough estimate
can be made by referring to a deep, small diameter crater as resulting
from a stronger and better focussed implosion. As can be seen in Table
3.4, only in two very different experiments did neutrons appear as
randomly-distributed negative signals. One of these runs is a direct
detonation of the deuterium-oxygen stoichiometric mixture with a 97g
PETN shell, while the other made use of an indirect method using a
capsule of a miniature Voitenko design (36). The detection system for
the neutrons, and γ rays which were produced by fusion neutrons interact-
ing with the steel implosion chamber, consisted of the two scintillator-
photomultiplier assemblies previously described. The first detector
was located at the outer front-plate surface, 30 cm from the implosion
focus, while the second detector was 80 cm from the focus (Fig. 3.26).
The first oscilloscope displays the entire ignition-detonation-implosion
process lasting about 50 μsec as well as the subsequent events. The
second oscilloscope was designed to display the full undisturbed shape
of the first signal coming from the second detector with a sweep of 50
μsec/division. From its shape it would have been possible to obtain the
neutron-velocity distribution and its flux. However, the second
oscilloscope can only trigger if a large enough signal (≥ 0.5v) is
produced by the first detector. This requires more than one neutron to
cross the scintillator within 10nsec. The threshold level is essential
to prevent false triggering arising from the photomultiplier dark
current, cosmic rays and ignition noise.

Records of voltage vs time from the first detector for two runs
without fusion a) and with fusion b) are shown in Fig. 3.27. Initially,
there are large oscillations arising from the capacitor discharge
to the exploding wire, which are damped out in about 35 μsec. In
the case of no fusion when a stoichiometric mixture of 2H2+O2 is
detonated, no other signals appear. However, with a stoichiometric
mixture of 2D2+O2 when fusion occurs at about 50 μsec, about 20
negative signals appear in a random-time and amplitude distribution
over a period of about 50 μsec. The maximum amplitude of about 0.2v
corresponds to a single impact by a neutron or a γ photon in the MeV
range, obtained from a calibration of the scintillators at the University
of Rochester. In this case the second oscilloscope did not trigger
as none of the events generated a large enough signal. We estimate
that the total yield was a few thousand neutrons.

The distribution of events can be explained by the scattering of
neutrons as they encountered the steel chamber walls (37,38). Iron
has a total cross-section of about 3.5 barns at 2.45 MeV \((39,40)\), see Fig. 3.28 and Table 3.5 for details. Other components of the steel alloy like Mn and Co also contribute to the neutron scattering. Inelastic scattering by iron atoms produces mainly 0.85 MeV and 1.24 MeV \(\gamma\) rays. However, neutron capture by Fe\(^{56}\) produces metastable Fe\(^{57}\) with \(\gamma\)-ray energies up to 10 MeV. The total number of neutrons and photons generated per reacting neutron is about three. The coupling angle between the chamber and the detector is about tenfold greater for the scattered neutrons and \(\gamma\) rays than for the neutrons generated at the implosion focus. Consequently, the scintillator is far more efficient in detecting the indirect scattered neutrons and \(\gamma\) rays than the direct neutrons from the implosion focus. Owing to the large attenuation of the inelastically scattered neutrons, the phenomenon is spread out and delayed as recorded in Fig. 3.27. Figure 3.29 shows the display of the first detector for the case when a small hemispherical cavity, capsule containing \(D_2\) was placed at the implosion focus (miniature Voitenko compressor, Fig. 3.16a and 3.16b). The display is nearly identical with the \(2D_2\cdot O_2\) stoichiometric mixture case.

4. DISCUSSIONS AND CONCLUSIONS

The compression of hydrogen isotopes by accelerated shells or shock waves in order to achieve a fusion reaction was studied in some detail in the last decade, especially in conjunction with laser applications to fusion \((2-10, 41, 42)\). For planar shock heating, the final temperature can be readily calculated by assuming that all the kinetic energy that the shock has transmitted to the gas is turned into heat when the gas is suddenly stopped by the reflected shock. However, in the case of a spherical implosion this can also be done using numerical methods (Ref. 21) as long as one does not try to reach the discontinuity at the focus, where the continuum equations break down. If one uses these equations unlimited temperatures and pressures are reached at the origin. However, this would always be limited by the transport of energy through radiation and conduction. However, within a radius of a few microns from the origin very high pressures and temperatures are reached sufficient to obtain fusion reactions in deuterium. The Voitenko-type compressor is also capable of producing similar results.

For the direct approach a stoichiometric mixture of deuterium and oxygen must be used in the present experiments which introduces oxygen as an impurity and additional heat sink. This approach also uses a relatively large amount of deuterium to fill the hemispherical implosion chamber to pressures of tens of atmospheres. Yet, only a very small fraction at the origin can actually be used for fusion. Looking ahead for a possible commercial use of this process for energy production stimulated our study of the indirect approach where small amounts of pure deuterium could be confined right at the implosion origin. The challenge is to produce a secondary implosion to be extremely symmetrical. The stability characteristics of spherical implosions becomes very important at this stage. Several stability studies \((43,44,45,46)\) disagree about the stability criterion for spherical implosions. Fong and Ahlborn \((46)\) predicted stability for initial implosion conditions similar to our case. This was substantiated here experimentally by the generation of neutrons when the implosion takes place in a \(D_2\cdot O_2\) stoichiometric mixture. All the stability studies show significant reduction in implosion
stability as the rates of the radius of the perturbations to the radius of the shock wave increases. As a result a symmetrical collapse is all-important and is far more difficult to achieve for the indirect approach. From this point of view the projectile designs (Fig. 3.14 and 3.17) are preferred. In this case, the second implosion is not directly depended on the first implosion, but on its own initial geometry. Previous studies also showed (20) that by providing a proper entrance into the acceleration tube the projectile velocity is hardly reduced by small off-centering and defocussing effects. In some of the projectile experiments a multiple staging acceleration effect (47,48) was used to increase the final diaphragm velocity by a factor of two. In the final phase the shell velocity must be used to generate very strong shock waves to compress and heat the gas. This might be done by either a spherical implosion or a Voitenko compressor.

Although very encouraging results were obtained in generating neutrons and $\gamma$ rays from D-D fusion reactions using our implosion chamber, its full potential for fusion has not been realized so far. The present facility was built for various types of research applications. The geometrical accuracy of the assembly after years of service could be improved. The uniformity of the present hand-made explosive shells certainly could stand a re-evaluation. Asymmetry also results from gravitational effects due to the difference in density of the detonating gas components. A new vertical oriented implosion chamber built recently for making diamonds from graphite produces much better centered implosions (49).

The type of PETN used is very important. New batches from different suppliers were found to have a very fine structure and therefore had a higher density in the final shell. Following the usual explosive-package-preparation procedure the shells ended up with a density above 1g/cm$^3$, while for the older experiments, the density was usually about 0.65g/cm$^3$. Steele (50) pointed out in a figure (Fig. 41) taken from a Lawrence Livermore report (51) which shows that PETN at higher density is very difficult to initiate by a $2\text{H}_2$ and $\text{O}_2$ detonation wave. However, the present results definitely show that we are approaching the thermodynamic conditions for fusion. Therefore any improvement in the implosion strength and symmetry might be significant. For commercial use the explosive package can be cast and even machined to obtain accurate sphericity, uniformity in thickness and density. This would contribute significantly to the final performance. Consequently, some experiments that failed to show any neutron yield might do so with the above improved initial conditions.

The major result is that we have obtained fusion neutrons and $\gamma$ rays using chemical energy only. As far as we could determine it is the first time that this has been done. It would be gratifying for others to build an implosion chamber and improve on our initial results.
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<table>
<thead>
<tr>
<th>Voltage Level of Discrimination</th>
<th>Signals Frequency</th>
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<tbody>
<tr>
<td>0.22</td>
<td>1200</td>
</tr>
<tr>
<td>0.24</td>
<td>850</td>
</tr>
<tr>
<td>0.30</td>
<td>575</td>
</tr>
<tr>
<td>0.40</td>
<td>375</td>
</tr>
<tr>
<td>0.50</td>
<td>175</td>
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<tr>
<td>0.60</td>
<td>125</td>
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<td>0.70</td>
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<td>95</td>
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<tr>
<td>0.90</td>
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<td>1.25</td>
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<td>1.50</td>
<td>75</td>
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Table 2.2

Neutron activation detection

<table>
<thead>
<tr>
<th>Description</th>
<th>Solid Angle</th>
<th>Relative Flux Density</th>
<th>Counts for Detection</th>
<th>Minimum Detectable Yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Hemispherical envelope</td>
<td>$2\pi$</td>
<td>1/45</td>
<td>500/30 min.</td>
<td>$5 \times 10^6$</td>
</tr>
<tr>
<td>2. Capsule envelope</td>
<td>$2/3\pi$</td>
<td>1</td>
<td>300/30 min.</td>
<td>$10^6$</td>
</tr>
<tr>
<td>3. Barrel envelope</td>
<td>$1/2\pi$</td>
<td>1/4</td>
<td>300/30 min.</td>
<td>$10^7$</td>
</tr>
<tr>
<td>4. Foil in barrel; fast neutrons</td>
<td>$0.08\pi$</td>
<td>1/7</td>
<td>250/30 min.</td>
<td>$10^7$</td>
</tr>
<tr>
<td>5. Makrofol G/U$^{238}$ fission track foil</td>
<td>$0.08\pi$</td>
<td>1/7</td>
<td>10 sparks</td>
<td>$10^6$</td>
</tr>
<tr>
<td>6. Gold plated detector</td>
<td>$0.08\pi$</td>
<td>1/7</td>
<td>4/1st sec.</td>
<td>$10^6$</td>
</tr>
<tr>
<td>7. Foil in barrel; paraffin attenuated neutrons</td>
<td>$3 \times 10^{-3}\pi$</td>
<td>1/45</td>
<td>850/30 min.</td>
<td>$10^6$</td>
</tr>
<tr>
<td>8. Boron plated detector; paraffin attenuated neutrons</td>
<td>$3 \times 10^{-3}\pi$</td>
<td>1/45</td>
<td>34/1st sec.</td>
<td>$10^4$</td>
</tr>
</tbody>
</table>

Note: The activated foil is of In$^{115}$, if not specified.

$$\text{In}^{115}(n,\gamma) \xrightarrow{1.46 \text{ eV}} \text{In}^{116m} \xrightarrow{\gamma} \text{In}^{116}, T_{1/2} = 54 \text{ min.}$$
### Table 3.1

**Signal amplitude as a function of distance**

from D-D source of $3 \times 10^5$ neutrons

<table>
<thead>
<tr>
<th>Distance (ft)</th>
<th>Output (v)</th>
<th>Distance (ft)</th>
<th>Output (v)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.65</td>
<td>2.7</td>
<td>0.6</td>
<td>3 (Double)</td>
</tr>
<tr>
<td>5</td>
<td>2.5</td>
<td>10</td>
<td>1.9</td>
</tr>
<tr>
<td>10</td>
<td>2.2</td>
<td>20</td>
<td>1</td>
</tr>
<tr>
<td>20</td>
<td>1.9</td>
<td>27</td>
<td>0.5</td>
</tr>
</tbody>
</table>

### Table 3.2

**Output of PMT for several voltage inputs**

with 20 nsec duration to the LED

<table>
<thead>
<tr>
<th>Input to L.E.D. (v)</th>
<th>P.M.T. I Output (v)</th>
<th>P.M.T. II Output (v)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Output</td>
<td>Input</td>
</tr>
<tr>
<td>3</td>
<td>2.8</td>
<td>0.93</td>
</tr>
<tr>
<td>4</td>
<td>3.8</td>
<td>0.95</td>
</tr>
<tr>
<td>6</td>
<td>4.7</td>
<td>0.78</td>
</tr>
<tr>
<td>8</td>
<td>5.8</td>
<td>0.72</td>
</tr>
</tbody>
</table>
Table 3.3

Output of PMT for several durations at constant voltage input of 4V to the LED

<table>
<thead>
<tr>
<th>Input Duration (nsec)</th>
<th>P.M.T. I Output (v)</th>
<th>P.M.T. II Output (v)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>3.8</td>
<td>1.1</td>
</tr>
<tr>
<td>30</td>
<td>4.2</td>
<td>1.5</td>
</tr>
<tr>
<td>40</td>
<td>4.9</td>
<td>2.2</td>
</tr>
<tr>
<td>50</td>
<td>5.1</td>
<td>2.8</td>
</tr>
<tr>
<td>Run No.</td>
<td>Explosive State</td>
<td>Lag Pressure (ps)</td>
</tr>
<tr>
<td>----------------</td>
<td>----------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/7/79</td>
<td>100</td>
<td>Cone, flat 50</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2/7/80</td>
<td>50</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/7/80</td>
<td>40</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4/7/80</td>
<td>30</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5/7/80</td>
<td>20</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6/7/80</td>
<td>10</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7/7/80</td>
<td>5</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8/7/80</td>
<td>2</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9/7/80</td>
<td>1</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10/7/80</td>
<td>0.5</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11/7/80</td>
<td>0.25</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12/7/80</td>
<td>0.1</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13/7/80</td>
<td>0.05</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14/7/80</td>
<td>0.025</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15/7/80</td>
<td>0.0125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16/7/80</td>
<td>0.00625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17/7/80</td>
<td>0.003125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18/7/80</td>
<td>0.0015625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19/7/80</td>
<td>0.00078125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20/7/80</td>
<td>0.000390625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>21/7/80</td>
<td>0.0001953125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>22/7/80</td>
<td>0.00009765625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>23/7/80</td>
<td>0.000048828125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>24/7/80</td>
<td>0.0000244140625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25/7/80</td>
<td>0.00001220703125</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>26/7/80</td>
<td>0.000006103515625</td>
<td>Cone, hemispherical 70 dia, 150 ps</td>
</tr>
</tbody>
</table>

**Table 3.1** Exposure Data

Notes:

- Explosive batch: 1 = low resolution
- Explosive batch: 2 = medium resolution
- Explosive batch: 3 = high resolution
- Notes: XX = no signal

**Study of the main stratified**

The detection system failed to trigger at 15."
Table 3.5

Data on iron activation

Iron Activation

1. Possible Reactions (examples)

1. Elastic scattering

2. Inelastic scattering: 
\[ \text{Fe}^{56}(n,n')\text{Fe}^{56m} \xrightarrow{\gamma} \text{Fe}^{56} \]
\[ \text{Fe}^{56}(n,n')\text{Fe}^{56m} \xrightarrow{1.24} \text{Fe}^{56m} \xrightarrow{0.85} \text{Fe}^{56} \]

3. Absorption (capture): 
\[ \text{Fe}^{56}(n)\text{Fe}^{57} \xrightarrow{\gamma} \text{Fe}^{57} \]

4. 
\[ ^{26}\text{Fe} (n,p)^{25}\text{Mn}^{56} \xrightarrow{\gamma} ^{25}\text{Mn}^{56} , T_1 = 2.58h \]

2. Cross Sections

<table>
<thead>
<tr>
<th>Energy of n</th>
<th>$\sigma_{ab}$</th>
<th>$\sigma_{sc}$</th>
<th>$\sigma_{t}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.45 Mev</td>
<td>~1</td>
<td>~1</td>
<td>3.5</td>
</tr>
<tr>
<td>thermal</td>
<td>2.62</td>
<td>11</td>
<td>13.6</td>
</tr>
</tbody>
</table>

3. $\gamma$ Ray Spectra from thermal-neutron capture of Fe

<table>
<thead>
<tr>
<th>Photon Energy Mev</th>
<th>0-1</th>
<th>1-4</th>
<th>2-3</th>
<th>3-5</th>
<th>5-7</th>
<th>7-9</th>
<th>&gt;9</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of Photons per 100 captures</td>
<td>&gt;75</td>
<td>60</td>
<td>27</td>
<td>23</td>
<td>25</td>
<td>38</td>
<td>2.1</td>
</tr>
</tbody>
</table>

Highest energy $\gamma$-rays: 10.16 Mev

Average No. of photons per 100 captures >250
FIG. 1.1 PHYSICAL CONDITIONS FOR THERMONUCLEAR REACTIONS. VALUES OF $\bar{\sigma}v$ BASED ON MAXWELLIAN DISTRIBUTION FOR D-T, D-D (TOTAL), AND D-He$^3$ REACTIONS (REF. 1).
FIG. 1.2 UTIAS IMPLOSION CHAMBER FOR DIRECT 2D₂+O₂ DETONATION PETN IMPLOSIONS.
FIG. 1.3 SCHEMATIC DIAGRAM ILLUSTRATING THE PRINCIPLE OF OPERATION OF THE UTIAS IMPLOSION-DRIVEN CHAMBER.

(a) Ignition and detonation, (b) implosion phase, (c) reflection and generation of high pressure and temperature zone, (d) process shown in radius-time (r,t) plane.

D₁-Gaseous mixture detonation wave, S₁-Explosive liner is ignited and generates a strong imploding wave, D₂-Explosive detonation wave, S₂-Shock reflected from copper liner, S₂-S₃-Reflected refracted shock at explosive-gas interface, S₁-Final imploding shock, R-Reflected implosion wave at origin.
FIG. 1.4 UTIAS IMPLOSION-CHAMBER FACILITY INCLUDING A DEUTERIUM CAPSULE (VOITENKO-COMPRESSOR MODE).
MATERIAL: STEEL 4140 OR 4340

FIG. 2.1(a) IMPLOSION CHAMBER REAR PLATE.
FIG. 2.1(b) IMPLOSION CHAMBER FRONT PLATE.

MATERIAL: STEEL 4140 OR 4340
FIG. 2.2 IMPLOSION-DRIVEN CHAMBER – EXPLODED VIEW.

FIG. 2.3 VIEW OF A MOCK-UP SHOWING THE STEPS IN THE MANUFACTURE OF EXPLOSIVE PACKAGE.

1. Bare copper liner. 2. Thin coat of cement. 3. Plastic honeycomb matrix. 4. PETN and cotton fiber slurry forced into the matrix.
FIG. 2.4 IGNITION ASSEMBLY.

The spark gap, consisting of 2 brass electrodes in a Plexiglas tube with a spark-plug trigger, is dehumidified by flowing compressed nitrogen. The capacitor is then charged to 20 kV and a 5 kV trigger pulse to the spark-plug initiates the discharge (Ref. 24).
FIG. 2.5(a) EXPLODING-WIRE ASSEMBLY, MODEL I.
4. Exploding wire. 5. Ground terminal.

FIG. 2.5(b) EXPLODING-WIRE ASSEMBLY, MODEL II.
1. Capsule body. 2. Fastening nut. 3. Teflon tube.
FIG. 2.6  VIEW OF EXPLODING-WIRE ASSEMBLY OVER DIAPHRAGM OF DEUTERIUM CAPSULE.
FIG. 2.7 SCHEMATIC DIAGRAM OF PRESSURE AND VACUUM SYSTEM.
FIG. 2.8  CONTROL PANEL.
FIG. 2.9 TYPICAL CAPSULES.
FIG. 2.10  VIEW OF DEUTERIUM FILLING SYSTEM.
FIG. 2.11 DEUTERIUM FILLING SYSTEM — SCHEMATIC VIEW.
Fast Pulse Response Applications, to 3000 V
(Typical circuit values)

- C1: 0.005 μF, 20%, Ceramic Disc, 500 V dc
- C2: 0.01 μF, 20%, Ceramic Disc, 500 V dc
- C3: 0.02 μF, 20%, Ceramic Disc, 500 V dc
- C4: 0.05 μF, 20%, Ceramic Disc, 500 V dc

- R1: 400 kΩ (4-100 kΩ, 5%, 1/2 W in series)
- R2: 100 kΩ, 5%, 1/2 W
- R3: 130 kΩ, 5%, 1 W
- R4 through R13: 100 kΩ, 5%, 1/2 W

FIG. 2.12 SCHEMATIC DIAGRAMS OF RCA MODEL 8575 PMT.
(a) Electrical, (b) mechanical, current spectrum.
MEASURED UNDER THE FOLLOWING CONDITIONS: LIGHT ON CATHODE IS TRANSMITTED THROUGH A BLUE FILTER (CORNING CS No.5-38, POLISHED TO 1/2 STOCK THICKNESS). LIGHT ON FILTER IS 0.1 MICRONUMER VOLTAGE DISTRIBUTIONALPHAS USED AND SUPPLY VOLTAGE ADJUSTED TO OBTAIN AN ANODE CURRENT OF 2.6 MICROAMPERES. LIGHT IS EXCLUDED DURING MEASUREMENT. FOCUSING ELECTRODE IS CONNECTED TO DYNODE No.1 POTENTIAL ELECTRON MULTIPLIER SHIELD IS CONNECTED TO DYNODE - No. 8 POTENTIAL TUBE TEMPERATURE = 22°C. ONE PHOTOELECTRON PULSE HEIGHT > 8 COUNTING CHANNELS. INTEGRATING TIME CONSTANT = 10 µs (R=100 kΩ, C=1000 pF).

15 PHOTOELECTRONS

\[ \sum = 40,000 \text{ COUNTS PER MINUTE (667 cps)} \]

1/10 PHOTOELECTRON

PULSE HEIGHT - PHOTOELECTRON EQUIVALENTS

(c) typical dark
FIG. 2.13 SCHEMATIC VIEW OF NEUTRON-DETECTION SYSTEM.
FIG. 2.14 VIEW OF EQUIPMENT INSIDE EXPLOSION ROOM.
FIG. 2.15 CALIBRATION OF THE "BACK VIEW" SYSTEM. ORIGINAL COMPONENTS AND CABLES ARE USED.

Channel 1: Directly from the pulse generator
Channel 2: Signal from the wiring of detector I
Channel 3: Signal from the wiring of detector II
FIG. 2.16  TRACE WITHOUT FUSION SCOPE I. THE OSCILLATIONS ARE DUE TO THE CAPACITOR DISCHARGE AND VANISH WITHIN 35 μSEC.
FIG. 2.17 SOME NEUTRON ACTIVATION TECHNIQUES.
FIG. 2.18 ACTUAL DESIGN FOR INDIUM ACTIVATION.
FIG. 3.1 TYPICAL PHOTOMULTIPLIER-SCINTILLATOR ASSEMBLY SIGNALS (PMT II).
(a) Electron emission signal, (b) Co$^{60}$ signal, (c) spectra of Co$^{60}$ signals.
FIG. 3.2 DETECTOR I SIGNALS AS A FUNCTION OF DISTANCE FROM D-D SOURCE OF 3x10^5 NEUTRONS.
FIG. 3.3 DETECTOR II SIGNALS AS A FUNCTION OF DISTANCE FROM D-D SOURCE OF $3 \times 10^5$ NEUTRONS.
ONE DIVISION EQUALS 1μ SEC

NEUTRON PULSE AS A FUNCTION OF TIME

FIG. 3.4 KAMAN NEUTRON GENERATOR MODEL A-800. TOTAL YIELD: 3×10^5 NEUTRONS FROM D-D REACTION (REF. 33).
FIG. 3.5 "COSMIC RAYS" AS DETECTED BY DETECTOR II.
(a) Highest amplitude recorded in 10 min, (b) "cosmic rays" of several strengths, (c) "cosmic rays" and Co$^{60}$ photons imposed.
Note: Output is always smaller than input.

Fig. 3.6 Input vs. Output signals of Discriminator. Sweep rate: 50 nsec/div.
FIG. 3.7 DETECTOR II SIGNALS AT 30 FT-8 IN FROM NEUTRON SOURCE.  
(b) is equal to (a) but with higher sensitivity and faster sweep rate, while (c) is a further enlargement of signal tail of (b).
Source to L.E.D. 20nsec, 3v

Source to L.E.D. 20nsec, 6v

Source to L.E.D. 20nsec, 4v

Source to L.E.D. 20nsec, 8v

FIG. 3.8 OUTPUT OF PMT I FOR SEVERAL VOLTAGE INPUTS, WITH CONSTANT DURATION TO LED.
FIG. 3.9(a) OUTPUT OF PMT I FOR SEVERAL DURATIONS, AT CONSTANT VOLTAGE INPUTS TO LED.
FIG. 3.9(b) OUTPUT OF PMT II FOR SEVERAL DURATIONS AT CONSTANT VOLTAGE INPUTS TO LED.
Input to L.E.D. 20 n sec, 3v

20 n sec, 4v

0.5 v/div

20 n sec, 6v

20 n sec, 8v

0.5 v/div

FIG. 9 - CONTINUED.
FIG. 3.9(c) TYPICAL INPUTS TO LED.
FIG. 3.10(a) SCHEMATIC DIAGRAM OF EXPERIMENTAL FACILITY USED TO STUDY DIAPHRAGM SURVIVAL DURING DETONATION-IMPLSION PROCESS.

Note: The system was originally used to study temperatures near spherical implosion foci (Ref. 21).
FIG. 3.10(b) SPECTROSCOPE-PMT DISPLAYS SHOWING COLLAPSE OF A .002"
STAINLESS STEEL DIAPHRAGM. INITIAL PRESSURE IN THE CHAMBER:
400 PSI.
FIG. 3.10(c) SPECTROSCOPE-PMT DISPLAYS SHOWING COLLAPSE OF A .005" STAINLESS STEEL DIAPHRAGM. INITIAL PRESSURE IN THE CHAMBER: 400 PSI.
FIG. 3.11 FLAT DIAPHRAGM CAPSULE. MATERIAL: 440C STAINLESS STEEL - HARDENED.

FIG. 3.12(a) SPHERICAL SECTOR CAPSULE. MATERIAL: 440C STAINLESS STEEL - HARDENED.
FIG. 3.12(b) STANDARD BARREL AND EXTENSION TUBE.

FIG. 3.13 EXPLODING WIRE FOR DIRECT IMPLOSION OF $2D_2+O_2$ MIXTURE.
Note: For general view see Fig. 1.4.
FIG. 3.14 CAPSULE AND PROJECTILE.
(a) Projectile capsule. (b) Projectile. (c) Projectile capsule assembly details.
1. Capsule body, hardened stainless steel. 2. Projectile, Al6061. 3. Squeezing nut, steel. 4. Polyethylene seal, 0.008". 5. Diaphragm, 0.006" stainless steel. 6. Washer, 0.002" Mylar.

Material: S.P.S. Hardened to 48 Rc
FIG. 3.15 SHALLOW CONE CAPSULE.

Material: S.P.S.

Materials:
1-Hardened S.S.
2-Copper
3-Lead
4-Tungsten Carbide

FIG. 3.16(a) VOITENKO CAPSULE.

Steel

Dural
FIG. 3.16(b) VOITENKO CAPSULE – ASSEMBLY DETAILS.

FIG. 3.16(c) VOITENKO CAPSULE WITH FLYER PLATE ASSEMBLY DETAILS. SCALE 30:1.
1. Capsule body, hardened stainless steel. 2. Polyethylene seal, 0.006". 3. Mylar diaphragm, 0.028" (not to scale). 4. Mylar washer, 0.002". 5. Squeezing nut, steel. 6. Stem, aluminum, 0.04". 7. Conical liner. 8. Aluminum flyer, 0.04" (not to scale).
FIG. 3.17(a) 9" ACCELERATION TUBE. SCALE 1:1.

FIG. 3.17(b) COMMERCIAL DESIGN — DETAILS OF CAPSULE ASSEMBLY.
6. PRESSURE TUBE

10. BARREL

12. SHIELD-STEEL

II. BARREL EXTENSION

FIG. 3.17(c) COMMERCIAL DESIGN DETAILS.
FIG. 3.17(d) COMMERCIAL DESIGN DETAILS.
FIG. 3.17e BARREL OF COMMERCIAL DESIGN AFTER RUN. THE DAMAGE AT THE ACCELERATING TUBE ENTRANCE IS RELATIVELY SMALL, AND THE HOLE THROUGH IS ALMOST CLEAR.
FIG. 3.18a TYPICAL CRATERS: UPPER ROW — DEFOCUSED RUNS, CENTRE ROW — TYPICAL 100 + 120g RUNS, LOWER ROW — TYPICAL 160g RUNS.

FIG. 3.18b TYPICAL DEFOCUSED RUNS.
FIG. 3.19 TYPICAL WELL FOCUSED AND CENTERED CRATER.

EXPERIMENT: E1
CAPSULE DESIGN: VOITENKO (FIG. 3.16)
EXPLOSIVE WEIGHT: 113g
CRATER DIMENSIONS: 19mm DIA x 15mm DEEP
FIG. 3.20 CRATER BEFORE (a), AND AFTER (b) REMOVAL OF THE CONICAL LINER.

EXPERIMENT: A1
CAPSULE DESIGN: CONE (FIG. 3.11)
EXPLOSIVE WEIGHT: 162g
CRATER DIMENSIONS: 25mm DIA x 20mm DEEP
FIG. 3.21 CRATER BEFORE (a), AND AFTER (b) REMOVAL OF THE CONICAL LINER.

EXPERIMENT: A2
CAPSULE DESIGN: STANDARD CONE (FIG. 3.12)
EXPLOSIVE WEIGHT: 162g
CRATER DIMENSIONS: 20mm DIA x 15mm DEEP, OFF CENTRE BY 8mm
FIG. 3.22 CRATER BEFORE (a), AND AFTER (b) REMOVAL OF THE CONICAL LINER.

EXPERIMENT: A3
CAPSULE DESIGN: STANDARD CONE (FIG. 3.12)
EXPLOSIVE WEIGHT: 171g
CRATER DIMENSIONS: 21mm DIA x 18mm DEEP
FIG. 3.23  WELL-FOCUSED CRATER, (a) AS TAKEN FROM THE CHAMBER, (b) AFTER REMOVAL OF THE CONICAL LINER.

EXPERIMENT: C2
CAPSULE DESIGN: SHALLOW CONE (FIG. 3.15)
EXPLOSIVE WEIGHT: 111g
CRATER DIMENSIONS: 15mm DIA x 20mm DEEP
FIG. 3.24 WELL-FOCUSED CRATER, (a) AS TAKEN FROM THE CHAMBER,  
(b) AFTER REMOVAL OF THE CONICAL LINER.

EXPERIMENT: C3  
CAPSULE DESIGN: STANDARD CONE (FIG. 3.12)  
EXPLOSIVE WEIGHT: 132g  
CRATER DIMENSIONS: 13mm DIA x 13mm DEEP
FIG. 3.25 CRATER BEFORE (a), AND AFTER (b) REMOVAL OF THE CONICAL LINER.

EXPERIMENT: D2
CAPSULE DESIGN: SHALLOW CONE (FIG. 3.15)
EXPLOSIVE WEIGHT: 108g
CRATER DIMENSIONS: 13mm DIA x 14mm DEEP
FIG. 3.26 UTIAS IMPLOSION CHAMBER FACILITY AND SCINTILLATOR DETECTORS.
FIG. 3.27 OSCILLOSCOPE RECORD FROM DETECTOR I.

(a) Without fusion, $2\text{H}_2 + \text{O}_2$ mixture at 400 psi + 127g PETN explosive.
(b) With fusion, $2\text{D}_2 + \text{O}_2$ mixture at 800 psi + 97g PETN explosive.
(i) Arrival of implosion and beginning of events at 47 $\mu$sec.
(ii) Ignition noise.
FIG. 3.28 CROSS-SECTIONS FOR NEUTRONS AND $^{26}$Fe INTERACTIONS (REF. 39).
FIG. 3.29 DISPLAY OF DETECTOR 1 WITHOUT FUSION (a) AND WITH FUSION (b). EXPERIMENT E1 (VOITENKO COMPRESSOR), 113g PETN, 2H₂+O₂ AT 400 PSI, D₂ IN CAPSULE AT 17.65 PSI. DISTRIBUTED NEGATIVE SIGNALS DUE TO NEUTRONS OR γ-RAYS START AT 43 μSEC.
1.1 Direct Approach

The simplest means of obtaining fusion in the UTIAS Implosion Facility is to substitute deuterium for the hydrogen in the stoichiometric mixture \((2D_2 + O_2)\). [Further improvements could be obtained by using a mixture of deuterium and tritium.] Using some of the data from Saito (Ref. 21) estimates of temperatures can be obtained of about 440,000 K (38 eV) at a 10\(\mu\) radius and 2,000,000 K (170 eV) at 1\(\mu\) for this mixture having a low value of \(\gamma = 1.14\). The temperature after reflection is only about half the value behind the initial imploding shock wave. However, for gases at very high pressures (megabar) and high temperatures (100 eV), \(\gamma = 1.67\). In this case the density ratio drops from about 10 for \(\gamma = 1.14\) to 2.7 for \(\gamma = 1.67\). Consequently, the temperatures are much greater. At a radius of 10\(\mu\), the pressure is 10-fold greater and the temperature is about 18\(\times\)10\(^6\)K or 1580 eV. The reflected shock now supplies additional heating by a factor of 1.6 and yields a temperature of about 29\(\times\)10\(^6\)K or 2528 eV. The actual ideal attained temperature will lie between the two limits provided by \(\gamma = 1.14\) (where it starts) and 1.67 (where it ends).

Liquid densities can be reached as a result of shock and isentropic compression behind the implosion. Consequently, the simple gas law no longer applies and convolume and pressure effects must be considered. All in all temperatures in the range of a few hundred eV can be expected at a 10-\(\mu\) radius and in the KeV range at smaller radii.

Some limitations to the calculations should be mentioned:

(a) The self-similar solutions computed by Saito (Ref. 21) are valid as long as the gasdynamic equations do not break down when the implosion wave radius becomes small \(\sim 1\mu\). (At a particle number density of \(n = 10^{22}/\text{cm}^3\) and \(T = 10^6\text{K}\) the mean-free-path \(\lambda \sim 0.03\mu\); at \(n = 10^{22}/\text{cm}^3\) and \(10^7\text{K}\), \(\lambda \sim 2.3\mu\).)

(b) Any departure from spherical symmetry (caused by the boundary layer on the wall of the major diameter or by irregularities near the origin) could have a serious effect on accurate focussing.

(c) In the present hemispherical geometry, energy dissipation by viscous and radiative effects is enhanced.

(d) The compression process (compared to lasers) is relatively slow and the power density is low. Consequently, energy losses are high thereby limiting the final attainable temperatures.

(e) The use of oxygen introduces a large dissociation and ionization heat sink providing the means of radiating energy away from the fusion plasma.
1.2 **Indirect Approach**

A number of solid fuels could be placed at the focus of the implosion in order to initiate fusion. Some of these materials are listed in Table Al.1. Small hemispherical targets might be used or flat targets for the implosion to focus on (Fig. Al.a). Some initial calculations were done for such schemes but none was tried experimentally.

Other possibilities exist in using small hemispherical shells filled with deuterium and tritium placed at the implosion focus, either with the hemispherical surface above the major diameter of the facility (Figs. Al.b, c) or below it (like a Voitenko compressor, Fig. Al.d). In Fig. Al.c the fuel itself forms the hemispherical shell. Another variant is a small cone filled with the fusion fuel with its base parallel with the major diameter (Fig. Al.e) or with a spherical sector (Fig. Al.f). A projectile capsule was also tried (Fig. Al.g). However, only the Voitenko compressor type (Fig. Al.3) yielded fusion neutrons, as noted in the main text.
Table A1.1. Characteristics of Fusion Target Materials

<table>
<thead>
<tr>
<th>Target</th>
<th>Molecular Weight</th>
<th>Deuterium Density (cm^{-3})</th>
<th>(\bar{z})</th>
<th>(\bar{z}^2/\bar{z})</th>
<th>Deuteron Density Total Particle Density (g/cm^3)</th>
<th>Solid Density (g/cm^3)</th>
<th>Melting Point</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid deuterium (D_2)</td>
<td>4</td>
<td>4.2 \times 10^{22}</td>
<td>1</td>
<td>1</td>
<td>0.5</td>
<td>0.14</td>
<td>18°K</td>
</tr>
<tr>
<td>Lithium deuteride (LiD)</td>
<td>8.95</td>
<td>6.2 \times 10^{22}</td>
<td>2</td>
<td>2.5</td>
<td>0.166</td>
<td>0.92</td>
<td>680°C</td>
</tr>
<tr>
<td>Deuterium oxide (D_2O)</td>
<td>20</td>
<td>6.64 \times 10^{22}</td>
<td>3.33</td>
<td>6.6</td>
<td>0.154</td>
<td>1.10</td>
<td>0°C</td>
</tr>
<tr>
<td>Lithium borodeuteride (LiBD_2)</td>
<td>25.8</td>
<td>7.3 \times 10^{22}</td>
<td>2.0</td>
<td>3.17</td>
<td>0.22</td>
<td>0.79</td>
<td>275°C</td>
</tr>
<tr>
<td>Paraffin (C_{36}D_{7h})</td>
<td>580</td>
<td>7.2 \times 10^{22}</td>
<td>2.64</td>
<td>4.71</td>
<td>0.18</td>
<td>0.94</td>
<td>72°C</td>
</tr>
<tr>
<td>Polyethylene (CO_2-CD_2)_n</td>
<td>2000 \div 10^5</td>
<td>7.2 \times 10^{22}</td>
<td>2.66</td>
<td>4.75</td>
<td>0.18</td>
<td>0.94 \div 0.96</td>
<td>120 \div 130°C</td>
</tr>
</tbody>
</table>

\(\bar{z}\) - mean atomic number.
FIG. A.1.1 POSSIBLE INDIRECT CONFIGURATIONS FOR FUSION USING THE UTIAS IMPLOSION CHAMBER. THE BASIC CONFIGURATION IS SHOWN IN (a), AND (b) TO (f) INDICATED SECTION VIEWS OF THE CENTRAL AREA. CONFIGURATIONS (d), (e), (f) AND (g) HAD SOME INITIAL TRIALS AND ONLY (d) (VOITENKO-TYPE COMPRESSOR) WAS SUCCESSFUL.
APPENDIX A2

NEUTRON DETECTION SYSTEM

A2.1 Neutron Detectors

The detector, as designed by Dr. A. K. Kudian, is shown in Fig. A2.1. The main components are as follows:

(a) NaI(Tl) (thallium activated, sodium iodide) scintillator made by Nuclear Enterprises, Model NE102, 4 in diam, 2 in thick. Additional technical data are given in Table A2.1.

(b) Leading cone of highly polished aluminum to collimate the light leading to the PMT cathode.

(c) RCA photomultiplier (PMT) model 8575 with main specifications as follows:

- Quantum efficiency typical 31% at 3850Å, 27% at 4250Å
- Current amplification, \(1.4 \times 10^7\) typical at 2000 V
- Low dark noise
- Rise time, 2.8 nsec; transit time 37 nsec.

The electronic circuit of the scintillator for the first detector is shown in Fig. A2.2. The circuit is designed to operate with a positive power supply which eliminates the use of a capacitor. This prevents serious problems of high-frequency filtering. The circuit is also characterized by a very high output voltage so that it can be distributed to several components. The electronic circuit of the scintillator of the second detector is shown in Fig. A2.3. It also operates with a positive high-voltage power supply. It is more flexible for adjustment, but gives a lower output.

A2.2 Ancillary Equipment

Light-Emitting Diode (LED)


- Peak spectral emission 5650Å
- Typical forward volt 2.5V
- Forward current 20mA
- Luminance or output 1.0mcd
- Max. power dissipation 80mW (at 25°C)
- Max. rise time 1 nsec

For typical characteristics see Monsanto LED model MV5222.

Oscilloscopes

Hewlett-Packard model 1744A single beam storage scope.
Counter

Hewlett-Packard model 5325A universal counter. Accuracy up to 0.1 μsec.

Cameras

Hewlett-Packard model 197A camera.

Discriminator

The circuit is shown in Fig. A2.4.
### Table A2.1. Technical Data for NE 102

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value/Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>General description</td>
<td>Scintillation chemicals in polyvinyl-toluene</td>
</tr>
<tr>
<td>Light output</td>
<td>65% that of an anthracene crystal of the same geometry*</td>
</tr>
<tr>
<td>Light output vs temperature</td>
<td>Light output independent of temperature between -60°C and +20°C; light output at +60°C is 95% that at +20°C (Ref. 31)</td>
</tr>
<tr>
<td>Decay constant</td>
<td>2.2x10⁻⁹ seconds</td>
</tr>
<tr>
<td>Maximum emission</td>
<td>4250 AU</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>1.032</td>
</tr>
<tr>
<td>Softening temperature</td>
<td>75°C</td>
</tr>
<tr>
<td>Refractive index</td>
<td>1.581</td>
</tr>
<tr>
<td>Effects of liquids</td>
<td>Soluble in aromatic solvents, acetone, etc. Unaffected by water, dilute acids, dilute alcalis or lower alcohols</td>
</tr>
<tr>
<td>No. of atoms per cm barn</td>
<td>H: 0.0525, C: 0.0475, N: 1.8x10⁻⁴, O: 1.8x10⁻⁴</td>
</tr>
<tr>
<td>No. of electrons per cc</td>
<td>3.4x10²³</td>
</tr>
<tr>
<td>α to β ratio</td>
<td>0.072†</td>
</tr>
</tbody>
</table>

*Conditions of measurement: Source Cs 137 gamma. Basis of comparison: midpoint of falling Compton edge on differential curve, or 1% of maximum counting rate on integral curve. Standard: anthracene crystal of same weight and geometry mounted in M₂O reflector.
†Determined by J. B. Czirr.
The question always arises whether we could improve matters by going to a tenfold scale or a 100-cm diam hemispherical cavity. It can be shown that the required driver energy (derived from the Lawson criterion) is given by (for a D-T reaction at 10 KeV, but will be used here for convenience)

\[ E_s = 1.6\mu^3 \frac{1}{\varepsilon_g} \left( \frac{n_s}{n_d} \right)^2 \]  

(A3.1)

where \( E_s \) is the energy generated by the (driver) source, \( \mu \) is the energy multiplier for the entire process (or \( \mu = E_f/E_s \), where \( E_f \) is the fusion energy), \( \varepsilon_g \) is the coupling efficiency between the plasma and driver energies (or \( \varepsilon_g = E_p/E_s \)), \( n_s/n_d \) is the ratio of particles in solid deuterium to that of the final deuterium-particle density. The break-even point (no net energy production) is given by \( \mu = 1 \). For the present driver of \( \text{D}_2\text{O} + \text{PETN} \) explosive, \( E_s = 1220 \text{ kJ} \), \( n_s = 4.5 \times 10^{22}/\text{cm}^3 \), \( n_d = 5.2 \times 10^{22}/\text{cm}^3 \), \( T = 10^7 \text{K} \) (in view of the low neutron flux). Assume a fusion reaction diameter of 10 \( \mu \text{m} \), then the particle velocity is \( \sim 100 \text{ km/sec} \) [see Saito, (21)] and \( E_s = 0.1 \text{kJ} \), consequently \( \varepsilon_g = 0.1 \text{kJ}/1220 \text{ kJ} \sim 10^{-7} \) (a very low coupling efficiency). The difficulty arises from the fact that high shock velocities or temperatures are attained only close to the focus of the implosion in a very small volume. Therefore, by scaling up the facility by a factor of 10, the driver energy is increased 1000-fold, but the velocity will only go up by \( 10^{0.35} \) or 2.24-fold [Saito (21)]. The density will remain near the asymptotic condition and the temperature at 10 \( \mu \text{m} \) will increase as the pressure or as \( 100^{0.65} \) or only by a factor of 4.45. Consequently, the coupling efficiency, which occurs to the inverse fourth power in the above equation can be improved paradoxically enough by scaling down rather than by scaling up.

Of course the two successful runs using \( \text{D}_2\text{O} + \text{O}_2 + \text{PETN} \) directly and the Voitenko-type compressor with pure deuterium indirectly [Glass and Sagie (52)] out of twenty tries using configurations shown in Fig. A1.1 are only a beginning in order to investigate the details of the neutron production and the efficiency of the process. However, the foregoing considerations do point to the conclusions that scaling up the facility is not worthwhile and that perhaps some new ideas for an indirect method may be worth pursuing.
5. Fusion plasmas
Sagie, D. and Glass, I. I.

EXPLOSIVE-DRIVEN HEMISPHERICAL IMPLOSIONS FOR GENERATING FUSION PLASMAS

The UTIAS explosive-driven-implosion facility was used to produce stable, centered and focussed hemispherical implosions to generate neutrons from D-D reactions. A high resolution scintillator-detection system measured the neutrons and \( \gamma \)-rays resulting from the fusion of deuterium. Several approaches were used to initiate fusion in deuterium. The simplest and most direct proved to be in a predetermined stoichiometric mixture of deuterium-oxygen. The other successful method was a miniature Voitenko-type compressor where a plane diaphragm was driven by the implosion wave into a secondary small spherical cavity that contained pure deuterium gas at one atmosphere. A great deal of work still remains in order to measure accurately the neutron flux and its velocity distribution as well as the precise interactions of the neutrons with the steel chamber which produced the \( \gamma \)-rays. Nevertheless, this is the only known work where fusion neutrons were produced by chemical energy only in a direct and indirect manner.