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ABSTRACT

The polarization of the neutrons produced by photodisintegration of deuterium by 2.75-Mev gamma rays has been measured and the results compared to theoretical predictions. The photoneutrons were taken at five gamma-ray-neutron angles between 45° and 135° in the laboratory and scattered from magnesium which has a broad resonance at these neutron energies. The polarization was determined from the asymmetry in the left-right scattering from magnesium. The neutrons are found to be negatively polarized as predicted. The magnitude of the polarization and the dependence on the gamma-ray-neutron angle are found to be in agreement with the theory within experimental error.
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I. INTRODUCTION

In this paper, we present the results of an experiment to measure the polarization of the neutrons from the reaction $D(\gamma,n)p$ using 2.75-Mev gamma rays. The only other experiment\(^1\) on the polarization of photonucleons which has appeared in the literature was performed at 240 Mev where theories are not available at present.

That the nucleons produced by photodisintegration of deuterium should be polarized was first pointed out by Rosentsveig\(^2\) and independently by Czyz and Sawicki.\(^3\) At low gamma-ray energies, the polarization arises from an interference between the electric dipole transition and the spin-flipping magnetic dipole transition. The spin of the outgoing neutron (or proton) is normal to the reaction plane defined by the momenta of the incoming gamma ray and the outgoing neutron.

Calculations of the polarization to be expected at various gamma-ray energies have been published by several authors.\(^4\)-\(^9,20\) At medium gamma-ray energies, the polarization was found to be sensitive to the n-p wave functions as well as to the interaction with the electromagnetic field. For gamma-ray energies within a few Mev of threshold, the n-p interaction depends only on the depth and range of the potential and can be treated in the effective-range
approximation. The polarization at low energies still depends on the assumptions concerning the electromagnetic transitions. Thus, experimental measurements of the polarization would, for medium gamma-ray energies, supply information concerning the n-p system and the interaction of nucleons with the electromagnetic field. At low energies, polarization measurements would test the consistency of the current interpretation of the electromagnetic transitions derived mainly from the differential cross sections for the photodisintegration of the deuteron.

II. POLARIZATION AT VERY LOW GAMMA-RAY ENERGIES

The photodisintegration of the deuteron just above threshold is usually assumed to proceed predominantly by dipole transitions. The transitions are:

\[ ^3S \to ^3P \quad \text{electric dipole} \]
\[ ^3S \to ^1S \quad \text{magnetic dipole} . \]

The differential cross section is written

\[ \frac{d\sigma}{d\Omega} = a + b \sin^2 \theta , \]

where the isotropic term, \( a \), results from the magnetic dipole transition and \( b \) derives from the electric dipole transition.

Rosentsveig\(^4\) treated the polarization at low energies in the zero-range approximation. Kawaguchi\(^6\) arrived at polarization formulae containing the coefficients of the angular distribution for photodisintegration and the nucleon-nucleon scattering-phase shifts. The derivation is based on the unitarity of the S matrix which results in a connection between scattering and photodisintegration. Kawaguchi assumed that, at low energies, only dipole transitions took place and that no noncentral force was involved. Then the photoproton polarization for unpolarized gamma rays and unpolarized deuterons was given by
\[ P = \pm \frac{(2ab/3)^{1/2} \sin \theta \sin [\delta(1^1S) - \delta(3^3P)]}{a + b \sin^2 \theta}, \]

where \( a \) and \( b \) are the angular distribution coefficients mentioned above, and \( \delta(1^1S), \delta(3^3P) \), are the singlet-S and triplet-P n-p scattering-phase shifts. At 2.75 Mev \( \delta(3^3P) \) is negligible compared to \( \delta(1^1S) \). The ambiguity in the sign of the polarization is a consequence of the mode of derivation of the expression.

By dividing numerator and denominator of the fraction by \( b \), one finds that \( P \) depends only on the ratio \( a/b \) and on \( \delta(1^1S) \). The polarization is symmetric about 90° in the center of mass. \( P \) is zero at 0° and 180°, maximum at \( \sin \theta_{\text{max}} = \sqrt{a/b} \), and has a shallow minimum at 90°. Kawaguchi derived the above formula for \( P \) for the proton, however, at low energies, the neutron polarization is given by the same expression.

Kramer and Müller⁹ have treated the low-energy polarization extensively in the effective-range approximation. They write

\[ P \cdot \frac{d\sigma}{d\Omega} = (\gamma_0 + \gamma_1 \cos \theta + \gamma_2 \cos^2 \theta) \sin \theta. \]

At 2.75 Mev, \( \gamma_2 \) is negligible, while \( \gamma_1 \) is approximately 2% of \( \gamma_0 \). For the accuracy to which the present experiment can be made, the equation may be written:

\[ P \cdot \frac{d\sigma}{d\Omega} = \gamma_0 \sin \theta. \]

The expression above is the same as that derived by Kawaguchi where

\[ \gamma_0 = (2ab/3)^{1/2} \sin \delta(1^1S). \]

However, the ambiguity in sign is removed. Kramer and Müller calculated \( a, b, \) and \( \gamma_0 \) at several energies from 3 to 11 Mev and under various assumptions for the parameters of the effective-range theory. Kramer¹⁰ has also evaluated the polarization at 2.75 Mev for comparison to this experiment. A plot of the polarization from this calculation is given in Fig. 9.
III. POLARIZATION EXPERIMENT USING 2.75-Mev GAMMA RAYS

The difficulties in measuring the polarization at low energies are mainly due to the small photodisintegration cross section of deuterium (~1 mb) and the fairly low neutron polarization expected (~30%). That is, a strong source of gamma rays and an efficient polarization analyzer are required. In the present experiment, we have used the 2.75-Mev gamma rays from a Na$^{24}$ source to disintegrate the deuterium. The neutron polarization was determined from the left-right asymmetry in the scattering from magnesium. The entire feasibility of the experiment is due to the fortunate fact that the neutrons from a Na$^{24}$-D source have energies which fall on a broad scattering resonance in magnesium having a high polarization-analyzing ability.

Radioactive sources have been used frequently in the past to supply gamma rays for the measurement of the photodisintegration cross sections of deuterium at low energies. A radioactive source has the advantage of providing nearly monochromatic gamma rays in contrast to the bremsstrahlung spectrum from an electron accelerator. Na$^{24}$ is an excellent source for the present experiment. The 2.753-Mev gamma ray is the only gamma ray from Na$^{24}$ of appreciable abundance above the 2.225-Mev$^{11}$ threshold for the photodisintegration of the deuteron. The gamma-ray energy is known to within 1 kev.$^{12}$ The 15.0-hour half-life$^{13}$ is known to better than 0.1 hr and is convenient for the preparation of the source and for counting with the source. The neutron activation cross section of 0.54 b is sufficient to permit the preparation of large sources in available neutron fluxes. As mentioned above, the energies of the photoneutrons are crucial to the experiment. Because of the momentum of the center of mass in the laboratory system, the neutron energy depends on the angle between the incoming gamma ray and the outgoing neutron.
For a Na\textsuperscript{24}-D source, the neutron energy in the laboratory is given in terms of the laboratory angle by

\[ E_n(\text{kev}) = 262 + 33 \cos \theta. \]

Natural magnesium is composed of three isotopes: Mg\textsuperscript{24} (78.6\%), Mg\textsuperscript{25} (10.1\%), and Mg\textsuperscript{26} (11.3\%). The total cross section for the scattering of neutrons from magnesium has been measured by Fields and Walt,\textsuperscript{15} and recently by Newson et al.\textsuperscript{16} In the range from 15 to 370 kev, Newson et al. find a resonance at 260 kev with a width of 75 ± 15 kev which they assign to Mg\textsuperscript{24}. Comparison of the energy of this resonance to the energies of the Na\textsuperscript{24}-D photoneutrons given in the equation above shows a remarkable match. The 260-kev level in Mg\textsuperscript{24} was assigned \( J^\pi = 1/2^- \). Mg\textsuperscript{24} has another prominent resonance at 83 kev, assigned \( J^\pi = 3/2^- \), a p-wave level. Since Mg\textsuperscript{24} has zero spin, the resonances at 83 kev and 260 kev evidently correspond to p-wave scattering in the two neutron spin orientations. The total cross section at the peak of the 260-kev resonance is about 11 b. The minor isotopes have cross section curves similar to that of Mg\textsuperscript{24}. Mg\textsuperscript{25} has some undetermined structure in the region of 250 to 350 kev and Mg\textsuperscript{26} has a wide resonance near 300 kev, assigned \( J^\pi = 1/2^- \). The peak cross sections of the minor isotopes in the 300-kev region are about 11 b.

Recently, Elwyn, Lane, and Langsdorf\textsuperscript{17,18} have measured the left-right ratios of neutrons scattered from magnesium. They employed neutrons emerging at an angle of 51° from the Li\textsuperscript{7}(p,n)Be\textsuperscript{7} reaction. The polarization, \( P_1 \), of the neutron beam in the energy region in which we are interested was determined from the measured \( P_1 P_2 \) for scattering from Li\textsuperscript{7} targets. The beam polarizations so determined were in agreement with those calculated directly from the magnesium measurements; there was some uncertainty due to the minor isotopes of magnesium. Elwyn, Lane, and Langsdorf analyzed
their data to give the polarization, \( P_2 \), of the neutron scattering from natural magnesium from 206 kev to 312 kev. \( P_2 \) is quite large, varying from 89% at 240 kev to 47% at 312 kev. The findings of Elwyn, Lane, and Langsdorf motivated our use of magnesium as a polarization analyzer which in turn made the present experiment possible.

The scattering of polarized neutrons from \( \text{Mg}^{24} \) is analogous to the well-known case\(^{19}\) of scattering from \( \text{He}^4 \). Both are spin-zero nuclei (\( \text{Mg}^{24} \) is in fact an alpha-particle nucleus) and the \( P_{3/2} \) and \( P_{1/2} \) states are split by the spin-orbit nuclear force, the \( P_{1/2} \) being at higher energy. The present experiment is similar to the familiar double-scattering experiment except that the first scattering which produces the polarization is replaced by the photodisintegration reaction. Whereas, in the case of double scattering, the polarization is produced by the spin-orbit interaction which depends on the spin orientation, the photodisintegration involves the spin-flipping magnetic dipole and the non-spin-flipping electric dipole transitions. That is, the polarization is produced by an interference between the electromagnetic transitions and we analyze the resulting neutron polarization by a nuclear scattering.

IV. EXPERIMENTAL DETAILS

A. Layout of the Experiment

The experiment was performed by activating a \( \text{Na}^{24} \) gamma source in the core of the Livermore Pool-Type Reactor and then transferring it in a lead container to an experimental setup on the floor of the reactor building. Figures 1 and 2 show the layout of the experiment. The \( \text{Na}^{24} \) gamma source was pivoted about the deuterated polyethylene (\( \text{CD}_2 \)) to provide for a range of \( \gamma \)-n angles. The \( \text{Na}^{24} \) and \( \text{CD}_2 \) were located in a lead cave designed to
Fig. 2. Side view of the experimental arrangement.
shield against gamma rays. A water wall placed against one side of the cave shielded the magnesium analyzer and the counter from neutrons. The desired beam of photoneutrons was allowed to emerge from the cave via a collimator in the shielding and to strike the magnesium polarization analyzer. A shielded neutron counter was pivoted about the magnesium to allow counting on the left and right sides of the magnesium.

The major considerations in the design of the experiment were as follows:

1. The scale of the experiment was determined largely by the necessary shielding — about 30 cm of lead and 60 cm of water. In addition, the magnesium was located 91 cm from the water wall to prevent neutrons from travelling from the magnesium to the wall and then scattering into the counter. Thus the CD$_2$ – Mg distance was over 2 meters.

2. The Na$^{24}$, CD$_2$, and Mg had thickness limitations:
   (a) The Na$^{24}$ source thickness had to be limited to avoid excessive degradation of gamma rays by Compton scattering. Those degraded gamma rays which remained above the photoneutron threshold would produce neutrons of the wrong energy and of a different degree of polarization.
   (b) The CD$_2$ thickness was limited to control the amount of neutron scattering within the plastic. Such scattering can change the direction, energy, and polarization of the neutrons.
   (c) Finally, the magnesium thickness must be chosen to avoid excessive multiple scattering which reduces the measured left-right ratio.

3. The Na$^{24}$ – CD$_2$ and Mg – counter distances were chosen as practical compromises between angular resolution and counting rates. The vertical dimensions of these components were made approximately 50% greater than horizontal dimensions because the angular resolution is much more dependent on angular spreads in the reaction plane.
B. Detailed Descriptions of Experimental Components

1. Lead cave. The inside dimensions were 91-cm \times 76-cm \times 41-cm high. Side walls were 30.5-cm thick, made of three widths of lead bricks offset to prevent cracks. A 2.54-cm-thick steel plate supported a 30.5-cm lead roof. A 20.3-cm lead floor was supported by 70 cm of concrete to raise the mid-plane of the reaction to 113 cm above the floor of the reactor room. The roof contained two 12.7-cm diameter holes. One was used to load the CD$_2$ and the other to load the Na$^{24}$ source.

The lead served to protect personnel from radiation hazard and also to prevent the generation of photoneutrons from the deuterium contained in the hydrogenous materials of the bulk neutron shielding.

2. Arm for Na$^{24}$ gamma-source movement. In order to vary the \( \gamma-n \) angle and the Na$^{24}$ - CD$_2$ distance within the cave, a 40.6-cm aluminum radial arm was constructed with a pivot on the neutron beam axis directly below the CD$_2$ neutron source. It was turned by a hand crank which extended up out of the cave through the CD$_2$ loading hole. A small platform was installed on top of the radial arm. The platform was positioned by a long screw driven by a reversible motor. On the platform was a funnel which received and centered the Na$^{24}$ source from the loading hole at the 90° position from the CD$_2$. The funnel was constructed of aluminum with a 1.6-mm-thick wall. The Na$^{24}$ could be positioned from 13.2 to 40.3 cm from the CD$_2$. Most of the polarization measurements were made at 30.2 cm. The outer end of the rotating arm rolled on two ball bearings. Nine equispaced angles of the arm from 45° to 135° were determined by the dropping of the ball bearings into small drilled holes in an aluminum platform under the rotating arm. Figure 3 is a photograph of the inside of the cave; Fig. 4 is an end view.
Fig. 3. Photograph showing the inside of the experimental cave.
Fig. 4. Detail of the source-handling equipment.
3. Water wall and collimator. The neutron shielding consisted of 61 cm of water contained in aluminum tanks. The wall extended 2 meters beyond the sides of the lead cave and 40 cm above the roof of the cave. The collimator was 10.2-cm wide and 15.2-cm high. The first 45.7 cm was through lead. The lead collimator was extended 15.2 cm into the inner cavity of the cave to reduce leakage of gamma rays out of the collimator. The outer 61 cm of the collimator was constructed of polyethylene fitted between the water tanks.

C. Na\textsuperscript{24} Source

NaF was used as a convenient salt of sodium. Fluorine is a light element producing no long-lived activities. Disks of NaF were cold-pressed under 2700 atmospheres and stacked in cylindrical aluminum cans. A space 0.16-cm radially and 1.27-cm axially was allowed for possible expansion under activation. The aluminum containers were welded tight against leaks. The cans had a 6.03-cm diameter base to center the source in its position on the radial arm. Actual dimensions of the sources are given in Table I.

<table>
<thead>
<tr>
<th>No.</th>
<th>NaF height cm</th>
<th>NaF diam cm</th>
<th>Al Wall Thickness cm</th>
<th>NaF wt g</th>
<th>Al wt g</th>
<th>Activity (curies)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>15.2</td>
<td>3.18</td>
<td>0.32</td>
<td>296</td>
<td>236</td>
<td>530 to 900</td>
</tr>
<tr>
<td>2</td>
<td>15.2</td>
<td>5.72</td>
<td>0.32</td>
<td>679</td>
<td>319</td>
<td>1600</td>
</tr>
</tbody>
</table>

A typical sample of the materials used in the sources was activated to a few microcuries and a gamma-ray spectrum made with a NaI spectrometer to test for impurities having gamma rays above 2.225 Mev. No appreciable impurities were detected. Also, the decay of the sources was checked during
the actual runs by taking neutron counts periodically. Over about two half-lives, the sources decayed with a 15.0-hour half-life within about 2%.

D. CD₂ Cylinders

The deuterium neutron sources were molded out of deuterated polyethylene. This makes possible thin, nearly self-supporting sources. The thin CD₂ cylinder was stiffened by the insertion of three 1-cm-thick disks of styrofoam. In order to have a reasonable solid angle subtended by the CD₂ from the Na²⁴ and at the same time to keep the CD₂ thin, the CD₂ was made into a hollow cylinder. The cylinder also preserves the source geometry when the γ-n angle is varied. Although one side of the CD₂ cylinder is closer to the Na²⁴ than the other, the magnesium scatterer is so far away that this effect is not important.

The cylinders were molded from granular CD₂ by heating steel molds to 125°C, then applying 1000 atm pressure to the melted plastic. The inner mold had a slight taper to facilitate removal. Both inner and outer sections of the mold were coated with silicon grease and then lined with aluminum foil prior to loading. After removal of the cylinder from the mold, the aluminum foil was peeled off and the cylinder trued in a lathe. The two finished cylinders had the dimensions listed in Table II.

Table II. Dimensions of CD₂ cylinders.

<table>
<thead>
<tr>
<th>Cyl.</th>
<th>o.d. cm</th>
<th>Wall cm</th>
<th>Weight g</th>
<th>Height cm</th>
</tr>
</thead>
<tbody>
<tr>
<td>thin</td>
<td>9.68</td>
<td>0.152</td>
<td>79.4</td>
<td>15.40</td>
</tr>
<tr>
<td>thick</td>
<td>9.84</td>
<td>0.267</td>
<td>137.0</td>
<td>15.40</td>
</tr>
</tbody>
</table>
The hydrogen in the CD₂ was more than 99% deuterium. Since beryllium is the only other element with a photoneutron threshold below 2.75 Mev, there was no problem with impurities.

E. Magnesium and Carbon Scatterers

The scatterers were supported on the collimator axis above the counter pivot point. In order to facilitate rapid changes of scatterer, a light platform consisting of a cross of steel strips was held in place by four piano wires. The wires were attached to the pivot point, to the four corners of the cross, and then flared out slightly to an overhead support well out of the counter's view. A styrofoam disk was pressed into the cross from the bottom and mechanical stops attached to positively position the scatterers. Total weight of the scatterer platform was approximately 30 grams.

A variety of shapes and thicknesses for the scatterers was tried during the experiment. A description of the scatterers is given in Table III. Most of the data were taken with the magnesium thin "X" and the carbon "V". However, the other thicknesses served to give an indication of the multiple scattering effect. The "X" presents a uniform thickness to the beam and is completely symmetrical to the beam and counter when oriented as shown in Fig. 1. The carbon "V" was placed with the apex on the beam axis and the "V" bisected by the beam axis. The scatterers were designed to cover an area just equal to the cross section of the collimator.

The magnesium metal used for the scatterers had several impurities, the principal ones being Al (3%), Zn (1%), and Mn (0.2%). A small correction to the data was made for these impurities.
Table III. Description of magnesium and carbon scatterers.

<table>
<thead>
<tr>
<th>Material</th>
<th>Shape</th>
<th>Height cm</th>
<th>Dimensions cm</th>
<th>Wall Thickness cm</th>
<th>Weight g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnesium</td>
<td>Thick Cyl.</td>
<td>15.24</td>
<td>10.16 o.d.</td>
<td>0.630</td>
<td>499.5</td>
</tr>
<tr>
<td>&quot;</td>
<td>Thin Cyl.</td>
<td>15.24</td>
<td>10.16 o.d.</td>
<td>0.328</td>
<td>265.5</td>
</tr>
<tr>
<td>&quot;</td>
<td>Thick &quot;X&quot;</td>
<td>15.24</td>
<td>Two plates ea. 15.24 wide</td>
<td>0.635</td>
<td>469.3</td>
</tr>
<tr>
<td>&quot;</td>
<td>Thin &quot;X&quot;</td>
<td>15.24</td>
<td>Two plates ea. 15.24 wide</td>
<td>0.318</td>
<td>238.2</td>
</tr>
<tr>
<td>&quot;</td>
<td>Thin &quot;V&quot;</td>
<td>15.24</td>
<td>Two plates ea. 7.62 wide</td>
<td>0.318</td>
<td>119.0</td>
</tr>
<tr>
<td>Carbon</td>
<td>Cylinder</td>
<td>15.24</td>
<td>10.16 o.d.</td>
<td>0.508</td>
<td>376.1</td>
</tr>
<tr>
<td>&quot;</td>
<td>Solid Rod</td>
<td>15.24</td>
<td>4.45 diam</td>
<td>---</td>
<td>395.7</td>
</tr>
<tr>
<td>&quot;</td>
<td>&quot;V&quot;</td>
<td>15.24</td>
<td>Two plates ea. 7.62 wide</td>
<td>1.270</td>
<td>450.0</td>
</tr>
</tbody>
</table>

F. Neutron Counter

Because of the requirements of high efficiency and low gamma-ray sensitivity, a counter consisting of BF$_3$ tubes surrounded by a neutron moderator was constructed. The inner counter (portion within the thermal neutron shielding) consisted of nine BF$_3$ tubes connected in parallel and embedded in polyethylene 20.3-cm wide by 40.6-cm high by 15.3-cm thick. The BF$_3$ tubes were filled with enriched BF$_3$ (96% B$^{10}$) at 120-cm Hg pressure. Their arrangement (Fig. 5) was found empirically by tests with the BF$_3$ tubes submerged in a water tank. The tubes were placed near the front face to prevent excessive fall of efficiency at low neutron energies. Over the small range of neutron energies encountered in the present experiment the variation in counter efficiency should be negligible, judging by the performance of other
Fig. 5. Top view of the inner neutron counter.
similar moderated counters. A thermal neutron shield of 0.076 cm of cadmium and 0.635 cm of Boral was placed around the inner counter.

The inner counter was surrounded by a massive (6800-kg) laminated shield of polyethylene. A hole in the shield served as a collimator for the inner counter which was recessed 28 cm. The shield serves two purposes, to shield against background neutrons diffusing in the room from all directions, especially from overhead, and to limit the neutrons scattered from the magnesium to direct paths to the counter, not via surrounding objects.

The entire counter was mounted on a triangular aluminum cart with a bearing for the pivot point and two ball-bearing wheels on the corner legs. The pivot point and wheels were directly on the concrete floor. Positioning of the counter was accomplished by rotating the counter up to mechanical stops on either side of the beam.

The electronics used was standard except for a transistorized preamplifier powered by mercury cells designed by Mr. Howard Spracklen of this Laboratory (see Fig. 6). The BF$_3$ tubes were operated as proportional counters and the scaler discriminator set to cut out almost all gamma-ray counts. Stability of the counter was such that counts taken under the same conditions were found to be reproducible within the statistics.

V. EXPERIMENTAL PROCEDURES

A. Irradiation of Na$^{24}$ and Source Transfer

Since Na$^{24}$ sources in the kilocurie range were required for this experiment, special procedures had to be developed to facilitate safe handling. The Na$^{24}$ source was attached to a 0.159-cm diameter stainless steel aircraft control cable. This type of cable is flexible, strong, withstands radiation, and does not activate objectionably. For activation, the Na$^{24}$ was
suspended in the center of the core of the reactor in order to obtain maximum flux and uniformity of activation. Typically, the source was exposed to a flux of $5 \times 10^{13}$ neutrons cm$^{-2}$ sec$^{-1}$ for 22 hours. After irradiation, the source was loaded into a large, lead, fuel-element coffin. The coffin had 20.3-cm-thick walls, was 1.5-m long, and weighed 3180 kg. The top was equipped with a plug with a helical hole which passed the source cable. On the bottom, an eccentric door could be rotated to open the coffin. The coffin was initially submerged in the reactor pool, the bottom door opened, and the source drawn up into the coffin. Then the door was closed and the coffin hoisted by crane clear of the water where it was allowed to drip dry. Afterwards the coffin was lowered onto the experimental cave, the bottom door opened and the source lowered into the funnel on the radial arm. Slack cable allowed movement of the arm within the cave. It was possible to withdraw the source back into the coffin at any time to make changes in the cave. The setup is illustrated in Fig. 4.

The strongest source (1600 curies) gave a reading of 23 r/hr on the surface of the coffin. However, by maintaining a reasonable distance and minimizing the time, it was handled safely. No personnel ever received more than the A.E.C. tolerance dose during the experiments.

B. Counting Procedures

1. Left-right ratio determination. An experiment normally consisted of determining the left-right ratios at from five to nine $\gamma$-n angles. Since the differential cross section for photodisintegration is peaked at $90^\circ$, lower counting rates would be encountered at angles near $45^\circ$ or $135^\circ$. A fairly steady count rate was maintained by alternating angular positions from the outside angles towards $90^\circ$. Thus, the decay of the source was balanced by
the increase of the differential cross section as $90^\circ$ was approached. Typically, ten minute counts were taken in the following order at each angular position:

<table>
<thead>
<tr>
<th>Count</th>
<th>Scatterer</th>
<th>Counter Position</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>None</td>
<td>Left</td>
</tr>
<tr>
<td>2.</td>
<td>Mg</td>
<td>Left</td>
</tr>
<tr>
<td>3.</td>
<td>Mg</td>
<td>Right</td>
</tr>
<tr>
<td>4.</td>
<td>None</td>
<td>Right</td>
</tr>
<tr>
<td>5.</td>
<td>Carbon</td>
<td>Right</td>
</tr>
<tr>
<td>6.</td>
<td>Carbon</td>
<td>Left</td>
</tr>
<tr>
<td>7.</td>
<td>None</td>
<td>Left</td>
</tr>
<tr>
<td>8.</td>
<td>Mg</td>
<td>Left</td>
</tr>
<tr>
<td>9.</td>
<td>Mg</td>
<td>Right</td>
</tr>
<tr>
<td>10.</td>
<td>None</td>
<td>Right</td>
</tr>
</tbody>
</table>

After correction for decay, a Mg L/R ratio could be calculated. For example:

$$\frac{L}{R} = \frac{\text{count 2} - \text{count 1}}{\text{count 3} - \text{count 4}}.$$  

Typical counting rates with thin CD$_2$ and thin Mg "X" were 1000 cpm, with no-scatterer rates 50% to 60% as high. No-scatterer counts taken with the collimator plugged with polyethylene showed that approximately 70% of the counting rate was due to neutrons which did not come through the collimator. Most of these probably came over the top of the water wall by air scattering.

The experiment to measure the left-right ratio was performed four times, each run lasting about 30 hours or two half-lives of the source. The conditions for the runs are summarized in Table IV.
### Table IV. Summary of experimental runs.

<table>
<thead>
<tr>
<th>Run No.</th>
<th>Source No. (See Table I)</th>
<th>Na$^{24}$ Strength (curies)</th>
<th>CD$_2$ Cyl. (See Table II)</th>
<th>Scattering and Shape: (See Table III)</th>
<th>Measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Thin</td>
<td>Thick Cyl.</td>
<td>Thin Cyl.</td>
<td>Thick &quot;X&quot;</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>50</td>
<td>Thin</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>530</td>
<td>Thin</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>733</td>
<td>Thin</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>1600</td>
<td>Thin</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>900</td>
<td>Thin</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Thin</td>
<td>Thick</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>
2. "Off-center" counts. Neutrons from the CD$_2$ could scatter from the inner walls of the cave (especially the back wall) and eventually emerge from the collimator. In order to derive an empirical correction for this source of background, the CD$_2$ was displaced 20.2 cm on the radial arm so that neutrons could not leave through the collimator directly. The Na$^{24}$ source was also displaced, the counts being corrected to the usual Na – CD$_2$ separation. Counts were taken at several angles, the procedure being the same as outlined in paragraph 1 above for normal counts. The resulting "off-center" counts were subtracted from the corresponding normal counts as a background. The "off-center" background was 8% of the normal count at 90°. It was fairly constant in absolute value for all angles, as one would expect for neutrons randomly scattered.

3. Angular distribution measurement. The angular distribution of the photoneutrons from the CD$_2$ was measured by rotating the counter into the direct beam from the collimator. The Na$^{24}$ – CD$_2$ distance was 40.3 cm. Counts were taken for the nine angles normally available on the arm. Then the measurements were extended to forward angles by displacing the CD$_2$ along the collimator axis towards the magnesium. Several angles were set up by a combination of angular and radial displacements of the Na$^{24}$. Similarly, backward angles were reached by displacing the CD$_2$ along the collimator axis away from the magnesium. "Off-center" counts were taken with the CD$_2$ displaced out of the direct acceptance of the collimator and subtracted as a background. All counts were corrected to a common time. Statistics were 1% or better. The resulting angular distribution is plotted in Fig. 7.

C. Experimental Tests and Checks

1. Gamma-ray background. A count was taken with the Na$^{24}$ in place but no CD$_2$ in the cave. The result, 15 cpm, indicated that the counter was
Fig. 7. Angular distribution of Na$^{24}$ - D photoneutrons.
not sensitive to gamma rays and that neutrons were not generated outside of the \( \text{CD}_2 \). The gamma radiation level on the face of the counter was approximately 1 mr hr\(^{-1} \).

2. Carbon scatterer. The most conclusive test for built-in asymmetries in the experimental setup consisted of replacing the magnesium scatterer by carbon. Carbon has no resonances in this energy region and has been found by other observers\(^{18,21} \) to give no left-right asymmetry in the scattering of polarized neutrons. As indicated above, the carbon runs were alternated with the magnesium runs. No significant asymmetries were found with the carbon.

3. Beryllium neutron source. For one measurement, the \( \text{CD}_2 \) was replaced by a beryllium cylinder (15.24-cm high, 10.16-cm o.d., 0.40-cm thick, weighing 348.7 g) and the left-right ratio determined for the beryllium photoneutrons at 90°. The left-right ratio for the thick magnesium "X" scatterer was 0.91 ± 0.14. Na\(^{24} \)–Be photoneutrons at 90° have nearly 1-Mev energy. Since the polarization is not known from theory and the analyzing properties of magnesium at this energy are also unknown, the measured left-right ratio is not significant except that it is different from that found with the Na\(^{24} \)–\( \text{CD}_2 \) neutrons, as one would in general expect if the effect of the 260-kev magnesium resonance is specific to the deuterium neutrons.

4. Collimator shape. Some neutrons from the \( \text{CD}_2 \) are scattered by the walls of the collimator. This may change the polarization and the energy of those neutrons. In order to investigate this effect experimentally, we changed the shape of the collimator and looked for changes in the measured left-right ratio. In one trial, the collimator was reduced to 5.1-cm wide and 12.7-cm high by inserting more polyethylene in the polyethylene end of the collimator. In another trial, the collimator was reduced to 10.2-cm wide and
7.6-cm high. In neither trial was any statistically significant change in the left-right ratio noted. Thus, despite the relatively crude collimator used, scattering from the collimator was apparently not important. Some factors which tend to suppress this effect are: the fact that the magnesium scattering cross section for neutrons with degraded energy is smaller than for unscattered neutrons; that some of the scattered neutrons fan out and miss the magnesium scatterer which covers only the area of the collimator; and, finally, neutrons scattered through small angles may not lose their polarization. On the other hand, a more refined collimator design might be desirable in a future experiment.

5. Counter tests. To investigate possible effects associated with very low energy neutrons, a left-right ratio was measured with 0.96 cm of Lucite absorber on the face of the counter. The counting rate was reduced by 23% but the usual left-right asymmetry was observed.

The polarization-scattering of magnesium varies fairly slowly with the scattering angle judging by the measurements by the Argonne Group \(^1\) at 90° and 45°. In addition, the vertical extent of the counter should have only a small effect on the asymmetry. However, we made one asymmetry measurement with the counter aperture reduced to 20.3 cm by 19.1 cm. No significant change in the left-right ratio was noted.

VI. RESULTS

A. Left-Right Ratios and Polarization

The counts from Runs 3 and 5 were combined for the five angles common to both runs. The results were corrected for "off-center" background only. Those ratios are listed in Table V, and plotted in Fig. 8. The errors shown are statistical only.
### Table V. Measured left-right ratios and calculated photoneutron polarization.

<table>
<thead>
<tr>
<th>$\theta$ c.m. (deg.)</th>
<th>Neutron Energy Lab. (kev)</th>
<th>Measured Left-Right Ratio for Mg Scatterer</th>
<th>Corrected Left-Right Ratio = $\varepsilon$</th>
<th>$P_1P_2^*$</th>
<th>$P_2$ for Mg</th>
<th>Photoneutron Polarization $P_1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>49.6</td>
<td>284.5</td>
<td>1.324 ± 0.025</td>
<td>1.547</td>
<td>0.224</td>
<td>-0.658 ± 0.045</td>
<td>-0.34 ± 0.05</td>
</tr>
<tr>
<td>71.8</td>
<td>274.0</td>
<td>1.348 ± 0.027</td>
<td>1.390</td>
<td>0.170</td>
<td>-0.725 ± 0.051</td>
<td>-0.23 ± 0.06</td>
</tr>
<tr>
<td>93.6</td>
<td>262.0</td>
<td>1.323 ± 0.023</td>
<td>1.394</td>
<td>0.171</td>
<td>-0.794 ± 0.065</td>
<td>-0.22 ± 0.06</td>
</tr>
<tr>
<td>114.8</td>
<td>250.0</td>
<td>1.373 ± 0.031</td>
<td>1.635</td>
<td>0.251</td>
<td>-0.851 ± 0.067</td>
<td>-0.29 ± 0.05</td>
</tr>
<tr>
<td>135.7</td>
<td>239.5</td>
<td>1.370 ± 0.037</td>
<td>1.526</td>
<td>0.217</td>
<td>-0.888 ± 0.084</td>
<td>-0.24 ± 0.06</td>
</tr>
</tbody>
</table>

$P_1P_2^* = \frac{1}{0.96} \times \frac{\varepsilon - 1}{\varepsilon + 1}$ where 0.96 = $\left[ \frac{\vec{P}_1 \cdot \vec{P}_2}{|\vec{P}_1| |\vec{P}_2|} \right]$
Fig. 8. Measured left-right ratios. Errors shown are statistical only.
The left-right ratios were then corrected for several effects which reduce the observed ratio from the true ratio. These corrections are discussed in detail in Section VII. From the corrected left-right ratio, \( \varepsilon \), we calculated \( P_1 P_2 \), the product of the polarization, \( P_1 \), of the incident beam by the polarization, \( P_2 \), for the scattering from magnesium, using the following relation:

\[
P_1 P_2 = (1/0.96)[(\varepsilon - 1)/(\varepsilon + 1)]
\]

where 0.96 is the average cosine of the angle between the reaction and scattering planes arising from the finite extent of the sources and counter. \( P_2 \), the polarization for scattering by magnesium, was taken from the work of Elwyn, Lane, and Langsdorf\(^ {18} \) and listed in Table V. The errors shown were quoted by them. The sign convention used here is that the polarization is positive for a net spin orientation in the direction of the vector \( \vec{k}_i \times \vec{k}_f \) where the k's are the wave numbers of the incident and final produced particles.

Finally, \( P_1 \) was calculated from \( P_1 P_2 \) and \( P_2 \). The error in \( P_1 \) includes the major known uncertainties as explained in Section VII.

Figure 9 is a plot of \( P_1 \). The curve shown is a plot of the theoretical curve derived by Kramer.\(^ {10} \) Kramer's calculation, carried out for \( E_\gamma = 2.759 \) Mev, depends on two parameters, the singlet scattering length, and the singlet range. The singlet scattering length is experimentally known;\(^ {22} \) \( a_s = -23.69 \pm 0.055 \times 10^{-13} \) cm. On the other hand, the singlet range is not well known. Kramer's results for three values of the singlet range are given below, where \( \gamma_0 \) is the parameter in the expression for the polarization given in Section II, \( a \) and \( b \) are the coefficients of the angular distribution, and \( \sigma_T \) is the total cross section for photodisintegration.
Fig. 9. Polarization of the neutrons from the reaction D(νn)H. The errors shown include the statistical errors from this experiment and the uncertainty in $P_2$ for magnesium. The theoretical curve is from the calculation by G. Kramer.
We have chosen case 2 above since the \( a/b \) is fairly close to the experimental value. We note that \( \gamma_0 \) is not very sensitive to the assumed value of the singlet range. It should be pointed out that \( \sigma_T \) for all three cases is lower than experimental values by about 4\%, although the quoted experimental errors are nearly large enough to overlap the theoretical value.

B. Angular Distribution

The measured angular distribution for the Na\(^{24}\) – D neutrons from our source is plotted in Fig. 7, together with a curve which was fitted to the data. The angular distribution was useful mainly as a check on the scattering of the neutrons in the CD\(_2\). This is discussed in Section VII.

VII. CORRECTIONS AND UNCERTAINTIES

A. Compton Scattering in the Na\(^{24}\) Source

Compton scattering of the gamma rays degrades the gamma-ray energy. One finds from the kinematics of Compton scattering that after scattering through an angle of 17° or greater, a 2.753-Mev gamma ray has an energy below 2.225 Mev, the photodisintegration threshold. This fact shows that those gamma rays which have been scattered by the inner walls of the cave cannot produce photoneutrons. However, some gammas are scattered in the source itself and in the surrounding aluminum funnel.
For Source No. 1, using the total Compton-scattering cross section, we find that 14.4% of the gamma rays are Compton-scattered in the source and funnel. From the differential cross section for Compton scattering we estimate from a rough numerical integration that, of those gammas which are scattered, 16% are scattered through an angle of 17° or less. The scattered gammas which are above the photodisintegration threshold therefore constitute \((0.16)(14.4\%) = 2.3\%\) of the total. Since the source possesses cylindrical symmetry, the angular distribution of scattered as well as unscattered gammas is isotropic in azimuth. Photoneutrons produced by the degraded 2.3% of the gammas will have the wrong energy for the magnesium resonance scattering and will be scattered with little left-right asymmetry, except that those neutrons with energies near the 83-kev resonance will probably be scattered with the opposite asymmetry. For the purpose of correcting the experimental data, we assume that 2.3% of the gammas produce unpolarized neutrons. This means that the measured left-right ratio was low by about 1% from Compton scattering.

Experimentally, Run 4 with Source 2 (approximately twice as thick as Source 1) gave an average left-right ratio, over nine angles, of 1.268. The corresponding average for Run 5 with Source 1 was 1.304. On the other hand, Run 3 with Source 1 gave an average of 1.331 for 5 angles. Thus, any possible source thickness effect is obscured by statistical fluctuations and reproducibility difficulties.

A test with 5 cm of aluminum absorber placed between the Na\(^{24}\) and \(\text{CD}_2\) failed to show any appreciable change in the left-right ratio.

B. Neutron Scattering in the \(\text{CD}_2\)

A certain fraction of the photoneutrons may be scattered within the \(\text{CD}_2\), undergoing possible changes in direction, energy, and polarization. We
considered this effect too complicated to calculate, therefore, an empirical correction was derived from the measurements with two CD$_2$ cylinders having wall thicknesses of 0.152 cm and 0.267 cm. The correction was based on a linear extrapolation (of the left-right ratio) to zero CD$_2$ thickness. This procedure was applied to the data for each of the five angles. The resulting corrections correspond to a fraction of neutrons scattered (and depolarized) equal to 21%, when averaged over the 5 angles. This correction is the largest of the experiment and furthermore affects the angular dependence of the polarization. The correction depends on two asymmetry measurements which results in an increased uncertainty, due to statistics, in the corrected left-right ratio.

The angular distribution of Fig. 7 can be used to give an independent estimate of the CD$_2$ scattering. A previous accurate measurement of the angular distribution for a Na$^{24}$-D source based on counting photo protons in deuterium gas gave the result:

$$\frac{d\sigma}{d\Omega} = C(0.167 + \sin^2\theta).$$

Our angular distribution was fitted by the curve

$$\frac{d\sigma}{d\Omega} = K(0.38 + \sin^2\theta).$$

If one assumes that in our case a fraction $f$ of the total flux of neutrons was scattered isotropically, then, from the above angular distributions, it follows that $f = 21\%$. This isotropic scattered component (assumed unpolarized) results in a 16% unpolarized component at 90° and 33% at 45°, the factor of 2 arising from the change in differential cross section. Thus, the correction is expected to produce a curvature in the calculated polarization which is convex upward as plotted in Fig. 9. The correction derived here was found to be in surprising agreement with the empirical correction from the source-thickness extrapolation. However, we have not considered how the magnesium
scatterer modifies the effect. The modification depends on the energy spectrum of the scattered neutrons and the magnesium cross section variation with energy. Therefore we can only consider the angular distribution as qualitative evidence for the correction.

C. Magnesium Impurities

The abundances of the magnesium impurities (see Section IV, E) were weighted by their respective average neutron-scattering cross sections in the energy range of this experiment. These weighted abundances were divided by the weighted abundances for all constituents, including the magnesium. Assuming that the impurities have no scattering asymmetry, the result is equivalent to 1.6% of the neutrons being scattered without asymmetry. The effect on the left-right ratio is about 0.8%.

D. Finite Geometry

From the left-right ratio, one calculates $P_1 P_2 \cos \psi$, where $\psi$ is the angle between $\vec{P}_1$ and $\vec{P}_2$, or, in other words, the angle between the photo-production and Mg scattering planes. We have calculated an average $\cos \psi$ considering only the vertical dimensions of the Na$^{24}$, CD$_2$, Mg, and counter. Using this rough approximation, it is possible to carry out the resulting integrals analytically. The result is $\cos \psi = 0.96$.

E. Multiple Scattering in Magnesium

It is difficult to calculate the multiple scattering correction, including polarization effects and nonisotropic angular distributions. In fact, we do not at present have adequate data on the angular distribution for this purpose. The Argonne group$^{18}$ has investigated the multiple scattering experimentally by measuring the left-right ratios for scattering of polarized neutrons from
various thicknesses of Mg. They find a trend towards higher left-right ratios in going down to a thickness of 3.2 mm (a single slab at 45° to the beam), but for smaller thicknesses the trend was not clear, the changes of asymmetry with thickness being of the order of the statistical uncertainties. The Argonne data were taken with thicknesses of 3.2 mm or less, and no correction for multiple scattering was applied.

Our own measurements of asymmetry vs Mg thickness were qualitatively in agreement with the Argonne results. We also have not applied any multiple scattering to the data. Fortunately, the correction is canceled if it is applied to both the Argonne data and to our data for equal Mg thicknesses. Our data were taken with two 3.2-mm slabs forming a cross. We estimate that such a cross is about 60% thicker than a single 3.2-mm slab for multiple scattering purposes. Thus, a small multiple-scattering effect remains, but we consider that less uncertainty is involved if we do not attempt a correction.

F. Errors

The corrections described above were applied to the data. Mechanical asymmetries in the apparatus are believed to affect the left-right ratio by less than 1%. The carbon results can be taken as confirmation of this estimate.

The uncertainties in the measurements made in this experiment are believed to be largely statistical, namely the statistics of the measured left-right ratio and of the ratio for the thick CD₂ which entered through the thickness correction. Other sources of error were smaller by comparison. The statistical errors were combined with the quoted error on the Mg P₂ as independent errors to calculate the error in P₁ listed in Table V.
VIII. CONCLUSIONS

This experiment has shown that the neutrons from the photodisintegration of deuterium by 2.753-Mev gamma rays are negatively polarized. The sign, magnitude, and angular dependence of the polarization are in agreement with the theoretical prediction, verifying the assumptions of the theory within the experimental uncertainties. These assumptions were that only magnetic dipole and electric dipole transitions are important at this energy and that only central nuclear forces are involved. The parameters derived from effective range theory gave a satisfactory fit to the data. The present results provide confirmation for the theory of the low-energy photodisintegration of the deuteron derived from cross-section measurements.

In the future, it should be possible to improve on the present results, especially by using thinner deuterium sources. In addition, the same setup can be used to measure the polarization of the photoneutrons from beryllium by replacing the magnesium with a scatterer having a suitable resonance.

IX. ACKNOWLEDGEMENTS

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* This work was done under the auspices of the U.S. Atomic Energy Commission.


10 G. Kramer, private communication.


18. A. J. Elwyn, R. O. Lane, and A. S. Langsdorf, Jr., private communication.


