Radioactive disintegration spectra of some short-lived nuclides

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I. INTRODUCTION AND LITERATURE SURVEY

The purpose of this investigation was to determine the beta energy endpoints of some of the mirror nuclei which can be produced by the Iowa State College 70 Mev synchrotron. An improved scintillation spectrometer (1) permitted the attainment of much better statistical and calibration accuracies, and thereby better endpoint determinations than those obtained by previous investigators (2).

The mirror nuclei considered here are those pairs of nuclei with \( |N - Z| = 1 \) where \( N \) and \( Z \) are the number of neutrons and proton, respectively in these nuclei. The beta endpoints of these are of special theoretical interest as will be indicated by the following paragraphs.

In formulating the general Fermi theory of beta decay (3), there are five permissible Lorentz invariant interaction forms. Much experimental and theoretical work has been done to determine which of these forms are necessary to describe the observed beta spectra, since this information should be of assistance in determining the nature of nuclear forces.

Fermi (4), in his original theory, selected one of these forms and obtained a set of selection rules called the Fermi rules. Gamow and Teller (5) showed that if another one of these forms was chosen it would better fit
some of the existing data. The selection rules from this theory are called the Gamow-Teller or the G-T rules.

For comparison of the allowed spectra with experiment, the five forms can be combined into two terms, and the probability for decay, \( P(E) \), of an electron of energy \( E \) in an energy interval \( dE \) can be written as

\[
P(E) dE = (m_0^2 c^2 / 2\pi^2 \hbar^2) P(E, Z) (E^2 - 1)^{1/2} E(E_0 - E)^2 dE \times \left( G_F^2 |f1|^2 + G_{G-T}^2 |\sigma|^2 \right)
\]

where \( G_F \) and \( G_{G-T} \) are the Fermi and G-T coupling constants, respectively, \( E_0 \) and \( E \) are the maximum beta energy and the energy of the emitted beta particle, respectively, in \( m_0 c^2 \) units, \( Z \) is the nuclear charge of the daughter product, \( F \) is the Fermi function which corrects the observed electron energy for the effects of the Coulomb forces, \( m_0 \) is the rest mass of the electron, \( c \) is the velocity of light, \( \hbar \) is Planck's constant divided by \( 2\pi \), and the absolute value squared integral terms are the Fermi and G-T type interaction terms. The terms in this equation can be integrated with respect to \( E \) and evaluated, and the result can be written as

\[
\int t \left( G_F^2 |f1|^2 + G_{G-T}^2 |\sigma|^2 \right) = \text{constant}
\]

where \( t \) is the half-life of the transition and \( f \) is a dimensionless integral which corrects the observed half-life for the effects of the nuclear Coulomb field and the energy distribution of the transition. The unknowns in this equation are the terms in the parenthesis.
Wigner (6), on the basis of his supermultiplet model, first calculated the integral squared terms for the mirror nuclei and those nuclei with two odd nuclei over an even core of $N = Z$. Moszkowski (7) was the first to use the second equation to evaluate the ratios of the $G$'s for these transitions. Trigg (8) made a detailed analysis using all the available data on the superallowed transitions and obtained a value of $(G_{F}/G_{T})^{2}$ of 0.5. Winther and Kofoed-Hansen (9) plotted the equation

$$B = f(t \left[ (1 - x) \left| J_{1} \right|^{2} + x \left| J_{2} \left| \sigma \right|^{2} \right]$$

for the mirror nuclei and those nuclei which have closed shells in both neutrons and protons plus or minus one particle, where $x$ equals

$$\frac{G_{T}^{2}}{G_{T}/(G_{F}^{2} + G_{T}^{2})}.$$ 

They obtained a value for $B$ of $2600 \pm 85$ and for $x$ of $0.50 \pm 0.05$.

The values of the integrals in the equation for $B$ have been calculated for the superallowed transitions using various nuclear models and are listed for several nuclides in the papers of Wigner, Trigg, and Winther. If the data were consistent with a single value of $x$ and $B$ for a particular nuclear model, very strong support for this model would be obtained and the interaction strength constants, the $G$'s, could be calculated.
A large part of the uncertainties in the $x$ and $B$ are due to the uncertainties in the $f$ values used in the calculations. The purpose of this investigation was to obtain a better determination of the beta endpoint energies for the mirror transitions and thereby the $f$ values. The $f$ values found are combined with the best available $t_{1/2}$ values in Table 3.
II. EQUIPMENT, TESTS, AND ERRORS

Description of Equipment

A description of the equipment used in this investigation has been published as ISC-359 (1). Further description will be given of only the components not covered in that report. These are the cycle control circuit, the target transport system, the targets, and the scintillation pickup.

In order to produce enough activity in the element being investigated, it was necessary to have the target sample close to the "donut" of the synchrotron, since a thin target was used in order to minimize the energy loss of the electrons as they were emitted during radioactive decay. It also was necessary to have the photomultiplier tube used in the spectrometer at such a distance from the synchrotron that the synchrotron's magnetic field would not interfere with the operation of the tube. This was achieved by the fast pneumatic target transport system which was controlled by the cycle programming equipment.

The transport system consisted of a rectangular aluminum tube about ten feet long with one closed and one open end. A block of balsa, to which the target was attached by means of a nickel frame, was formed so that it would slide freely in the tube. A pipe was attached to the closed end of the tube so that by applying either more or
less than atmospheric pressure to the pipe, the target could be made to rapidly move to either end of the tube, i.e.; to the bombard or the record position. When the target was in the record position, the radioactive sample protruded from the end of the tube and was positioned accurately parallel to the scintillation crystal and about \( \frac{1}{8} \)" away.

All of the targets for the beta spectra except magnesium, silicon, and beryllium were prepared by pressing powder into a thin wafer with pressures of 5000 lb/in\(^2\). The wafers were 1.165 inches in diameter and less than 50 mg/cm\(^2\) in thickness. The thickness was checked by weighing the samples after pressing. Those that were too heavy were discarded. The acceptable samples were then mounted with diluted Duco cement between two sheets of rubber hydrochloride which had a thickness of 0.5 mg/cm\(^2\). The rubber hydrochloride was cemented to a rigid, light weight nickel frame which in turn was fastened to the balsa wood block mentioned previously. The silicon target was prepared by spreading over the same area enough powder to give it the desired thickness. It was bonded similarly. The magnesium target was prepared from strips of metal such that it presented a target one inch square by 0.005 inches in thickness. The beryllium target was similar to the magnesium one, except it was a solid sheet one inch square by 0.005 inches. The constituents of the targets will be listed as they are discussed under the name of the isotope.
In order to obtain enough counts per channel for good statistical accuracy, it was necessary to bombard and record many times. This was facilitated by the automatic control equipment (Figure 1) which cycled the synchrotron and the recording equipment. The following operations comprised one cycle:

1. The target was in bombard position with the synchrotron ON and the recording equipment OFF;
2. The target was in transit from the bombard position to the record position with the synchrotron OFF and the recording equipment either OFF or ON as desired;
3. The target was in record position with the synchrotron OFF and the recording equipment ON;
4. The target was in transit from the record position to the bombard position with the synchrotron OFF and the recording equipment OFF.

The time of each of these operations could be varied as desired. The usual procedure was to have the duration of operations 1 and 3 equal two half-lives. The transit time, step 2, could be varied from less than one-half to three seconds, with one second being the usual time. The time for step 4 could be varied from three to twelve seconds. The only long lived activity that interfered with the shorter half-life measurement was the 2.5 minute activity of P\textsuperscript{30} in the phosphorus sample.
Figure 1. Circuit Diagram of the Automatic Cycle Control.
The time required to run a calibration and the unknown spectrum varied from two to five hours. For these periods the equipment was as stable and drift-free as the equipment used for testing, which was reliable to one-half of one per cent.

The scintillation pickup consisted of an anthracene crystal and photomultiplier tube, a cathode follower, a pulse generator, and a step attenuator.

A typical scintillation crystal and phototube mount is shown in Figure 5, page 13 of Phipps' thesis (10) with the exception that no MgO coating was used on the top 3.1 mg/cm$^2$ aluminum reflector through which the beta-rays entered the crystal. A resolution of 15 per cent with a peak to valley ratio of three to one was obtained with the Cs$^{137}$ internal conversion line. This was consistent with the resolution reported by Phipps. These data were taken with a 1.5 inch crystal mounted on a Dumont 6177 photomultiplier tube, and also with Boley's (2) one inch crystal mounted on an RCA 5819 tube. Tests to determine the energy lost by electrons in transversing the 3.1 mg/cm$^2$ aluminum reflector indicated that the displacement of the apparent maximum of the 624 kev internal conversion electron peak of Cs$^{137}$ is less than 10 kev. These tests were carried out by scanning the cesium 137 conversion peak after placing zero through five 3.1 mg/cm$^2$ additional aluminum sheets between the source and the crystal.
The photomultiplier, step attenuator, and cathode follower circuit diagram is shown in Figure 2. The voltage pulse for a single event had a total duration of about two microseconds. This was determined by the capacity of the collector anode plus the input capacity of the cathode follower and the grid resistor. A pulse of nearly the same shape was produced by a simple pulse generator and was applied to the input of the cathode follower after attenuation by the decade resistor network.

The step attenuator was made with precision resistors mounted on an eleven position stepping relay. This type relay was selected in order that it could be mounted beside the cathode follower and be controlled remotely. As shown in Figure 2, the step attenuator could be disconnected from the cathode follower by pushing the RESET button. In order to eliminate electrical noise from the synchrotron, the pulse generator and the step attenuator control power cable were also disconnected while data were being taken.

In addition to the tests of the equipment reported in ISC-359, additional checks of the linearity of the scintillation pickup and the remainder of the equipment were made by determining the beta-endpoints of $^{32}P$, $^{80}Y$, $^{35}Cl$, and $^{18}Li$. The results of these runs, Figures 3, 4, 11, 12, 21, and 22, show that the equipment was linear to 13 Mev in agreement with the step attenuator calibration procedure.
Figure 2. Circuit Diagram of the Scintillation Pick-up.
Figure 3. Beta Spectra of $^{32}$P and $^{90}$Y.

- Counts $\times 10^{-3}$
- Step Attenuator Setting in Arbitrary Units
- Channel Number
- $^{32}$P
- $^{90}$Y
- Step Attenuator Calibration
Figure 4. Kurie plots of $^{32}P$ and $^{90}Y$. 

The diagram shows the relationship between channel number and the step attenuator setting, with annotations for $^{32}P$ at 1.70 MeV and $^{90}Y$ at 2.24 MeV. The equation $a = (W_n - W)^2 + A(W^2 - 1)$ is used for correction.
Experimental Procedure and Calibration

The camera used for recording the pulse information displayed on the oscilloscope was designed for only 100 feet of film at a loading. Since it was not practical to reload the camera with the assurance that the position of the camera with respect to the oscilloscope had not been changed, a run was concluded when 100 feet of film had been used. This was enough to obtain good statistical accuracy of the calibration and the unknown spectra.

The procedure used in all cases in which Cl^{34} served as a calibration spectrum was to bombard an unmounted NH_4Cl wafer for one hour and then to place it on the top aluminum reflector of the scintillation tube and crystal. In order to obtain the desired number of events from Cl^{34}, it was necessary to record for about 30 minutes and to use about 40 feet of film. Next, a linearity calibration was recorded from the pulse generator and step attenuator. As an added check, about 10 or 15 feet of Y^{90} was recorded with such a film speed as to yield a high density of dots near the endpoint of the spectrum. The remainder of the 100 feet of film was used to record the activity of interest.

The film was developed and processed through the multi-channel analyzer as described in ISC-359.
The data obtained from the automatic film scanner were number of counts and step attenuator positions versus the channel number. The channel number scale is later converted to an energy scale. These results are shown, with the exception of Figure 1 and 6, as the odd numbered Figures. When the positions of the step attenuator versus the channel numbers at which the calibration pulses appear are plotted, the nonlinearity of the equipment shows up as the deviation from the straight line. As can be seen, the equipment was linear within experimental error except at either end. This nonlinearity is due at the lower end to the nonlinear resistances of the crystal diodes in the clipping circuit, and at the upper end to the nonlinear characteristic of the oscilloscope for large deflections. By keeping the pulse sizes within the linear range of the equipment, the position of the "baseline" or zero pulse amplitude could be obtained by extrapolation of the calibration curve. That this zero was the true zero within experimental error was verified by the linearity experiments. The baseline can also be inferred from the statement by Birks (14) that if the voltage pulse height from an anthracene photomultiplier spectrometer is plotted versus energy for the electrons stopped by the crystal, the plot is linear above 125 kev and the linear portion, if extrapolated, crosses the energy axis at plus 25 kev. For most of the figures, 25 kev is of the order of one dot radius, hence the zero correction for any saturation effect is probably negligible.
With the zero amplitude "baseline" determined, it is necessary to determine the energy of at least one other point in the linear region of the equipment in order to have an energy calibration. For most of the data, the 4.45 Mev (12) beta-ray endpoint of Cl$^{34}$ and the 2.24 Mev (15) endpoint of Y$^{90}$ were used. The Y$^{90}$ served as an additional check when it was used with the Cl$^{34}$.

The endpoints were determined by making a Kurie plot from the spectrum. The Kurie plot consists of plotting the function $N\beta f$ versus the energy where $N$ is the number of counts at a given energy, $\beta$ is the ratio of the velocity of the electron to the velocity of light, and $f$ is the modified Fermi function as given in the National Bureau of Standard's Tables for the Analysis of Beta Spectra, Applied Mathematics Series, No. 13. In the following graphs, $N$ is the number of counts at a given channel number.

Because $f$ is a function of nuclear charge and energy, an energy scale must be assumed before a calibration Kurie plot can be made. This was done by selecting a suitable channel number from inspection of the observed spectrum and assigning to it the known endpoint energy. A Kurie plot was made with this assumption and a new channel number endpoint determined. This new endpoint was used for the next Kurie plot. This process converged rapidly to a unique endpoint.
An examination of the Kurie plots will show a fairly long straight portion near the end with the points bending up from the straight line at either end. The bending near the endpoint is caused by the resolution spread of the instrument. Calculations which make corrections possible have been made by Palmer and Laslett (16). With the exception of \( P^{30} \) (Figure 14) the uncorrected points show on the Kurie plots as open circles and the corrected points as solid dots below each of the open circles.

In making these corrections, it was necessary to know the resolution of the instrument at the energy at which the correction was being made. This was obtained for any energy by measuring the resolution for the Cs\(^{137}\) conversion line and using the fact that the resolution is known to go inversely as the square root of the energy.

Because of the bending of the Kurie plots at lower energies, several tests were run on \( Cl^{34}, Y^{90}, \) and \( P^{32} \) to determine over what portion of the spectrum the Kurie plots could be relied upon for endpoint determinations. Figure 4 shows the result of one such test. This may be compared with some of the \( Cl^{34} \) calibration Kurie plots. These Kurie plots and the Kurie plot of \( Li^{8} \) indicate that an allowed spectrum will give a straight Kurie plot for about the upper one-third of the spectrum from 2 to 13 Mev, when a scintillation counter is used as a detector.
In an attempt to determine the cause of this bending, the spectra shown in Figures 5 and 6 were run. The chlorine $^{34}\text{Cl}$ sources were extracted by the Szilard-Chalmers (17) process from a bombarded volume of CCl$_4$. The activity was deposited upon a 3.1 mg/cm$^2$ aluminum backing and placed three inches from the top of the crystal and on the axis of symmetry of the tube and crystal. There were four runs of 50 feet of film each. In the first run, a lead shield three eighths of an inch thick with a one-half inch hole was placed so that the shield was one inch from the top of the crystal and the hole was on the axis of symmetry of the system. In the second experiment the lead shield was removed and a lead cylinder three-eighths of an inch high and one-half an inch in diameter was placed in the position occupied by the hole. This was accomplished with the use of a sheet of 0.5 mg/cm$^2$ rubber hydrochloride which was also in place when the first run was made in order to have identical energy losses outside of the crystal. The third experiment was like the first except that the lead was replaced with aluminum. The fourth experiment was with no shield of any type.

In the spectra obtained with the lead cylinder versus the hole (Figure 5) it can be seen that the number of counts at a given energy for the cylinder is less than the number with the hole for high energies and is greater for low
Figure 5. Results of Tests to Determine the Effect Caused by Electrons Which Enter the Edge of the Crystal.
Figure 6. Results of Comparing the Crystal with No Shield to the Crystal with the Edges Shielded.
energies. This indicates that loss of electrons through the edges of the crystal caused some of the high energy electrons to be counted as low energy electrons which, in turn, caused the Kurie plot to bend upward at lower energies. The rapid turn up at about channel 34 is caused by a second beta group in Cl\textsuperscript{34}.

Because of the small solid angle permitted with the hole and the weak activities of the targets, it was not possible to use this method to straighten the Kurie plots.

Errors in Endpoint Determinations

The probable errors quoted with the beta endpoint energies are due to the uncertainties in the calibration energies, the zero energy point, and the endpoint channel numbers determined from the Kurie plots for both the calibration and unknown spectra.

As mentioned previously, a zero energy point could be determined by the use of a calibrated step attenuator. The probable error in each step attenuator calibration point is about the radius of the dot. The zero point was obtained by extrapolating the linear portion of the step attenuator setting versus channel number until the channel number axis was intersected. This zero had a probable error of less than one-half of a channel. One-half channel was used for error calculations.
Both \( Y^{90} \) and \( Cl^{34} \) have been measured by magnetic spectrometers, but the errors in the quoted endpoints are large enough that they must be considered in determining the probable errors in these measurements. The reported endpoints for \( Y^{90} \) are 2.27 ± 0.02 Mev (18), 2.25 Mev (19), 2.24 Mev (20), 2.23 Mev (21), and 2.18 ± 0.007 Mev (22). From these values, a value of 2.24 ± 0.04 Mev was assigned to \( Y^{90} \) for calibration purposes. The \( Cl^{34} \) endpoint has been measured as 4.45 ± 0.11 Mev (12) and 4.46 Mev (23). From these values, together with the values resulting from repeated comparisons of \( Cl^{34} \) and \( Y^{90} \) in this investigation, an endpoint value of 4.45 ± 0.10 Mev was assigned to \( Cl^{34} \).

In addition to these errors, additional errors were introduced in determining the endpoints from the Kurie plots. To determine the endpoint, only the high energy straight section of each Kurie plot was used. In order to determine what portion of the Kurie plot was the straight section, a straight edge was used to connect as many dots as possible, with an allowable deviation of one dot diameter. Then, a least squares best fit was made to the points along this region. The probable error from this method was not more than one-half of one channel. One-half channel was used as the probable error for calculations.
The quoted probable error, $e_n$, for the unknown endpoint energy, $E_n$, will be defined for the case of $n$ calibration points. Let $E_i$, $i = 0, 1, 2, \ldots, n-1$, be the $n$ energy calibration points; $V_i$ be the channel numbers corresponding to these points; $E_n$ and $V_n$ be the unknown energy and channel number, respectively; $e_i$ and $v_i$ be the probable errors in $E_i$ and $V_i$, $i = 0, 1, 2, \ldots, n$, respectively; and $A$ and $K$ be constants to be determined. Since the equipment was linear, 

$$E_i = K(V_i - A).$$

To make a least squares best fit, the function

$$\sum_{i=0}^{n-1} \left[ E_i - K(V_i - A) \right]^2$$

is minimized with respect to $K$ and $A$. $K$ and $A$ can be determined in terms of the $E_i$ and $V_i$. Then $E_n$ can be found as a function of the $E_i$, $i = 0, 1, 2, \ldots, n-1$, and the $V_i$, $i = 0, 1, 2, \ldots, n$ from the relation

$$E_n = K(V_n - A).$$

The probable error $e_n$ is defined as

$$e_n = \left[ \sum_{i=0}^{n} \left( \frac{\partial E_n}{\partial V_i} v_i \right)^2 + \sum_{i=0}^{n-1} \left( \frac{\partial E_n}{\partial E_i} e_i \right)^2 \right]^{\frac{1}{2}}.$$

The error determined in this manner agreed very well with that observed in the repeated runs of $\text{Cl}^{34}$ and $\text{Y}^{90}$. A further check was furnished by the several determinations of each unknown. From the close grouping of values in
Table 1 for each unknown determination, it can be seen that the uncertainties in the energies of the calibration points represent a large fraction of the probable error. The fraction of the probable error from this cause ranged from about one-third for Mg$^{23}$ to one-half for Ca$^{39}$. If Cl$^{34}$ and Y$^{90}$ could be determined more accurately, these quoted errors could be reduced.

Discrepancies in the values reported here and others will be discussed under the isotope in question.
III. APPLICATIONS TO SOME SUPERALLOWED TRANSITIONS

Magnesium 23

Mg$^{23}$ was formed by a ($\gamma$,n) reaction on stable Mg$^{24}$. The target was made from 0.005 inch foil. The longer half-life permitted the activity to be recorded with more ease than any of the other activities. For this reason, Mg$^{23}$ was used to test the equipment. The three runs shown in Table 1 were made in July, August, and October. The two earlier runs were made with the one inch crystal used by Boley. The statistics of the October run, the results of which are shown in Figures 7 and 8, were better than those of the other two. Y$^{90}$ was used as the calibration spectrum in all cases.

The three runs were averaged and the value of 2.95 ± 0.07 Mev was assigned to Mg$^{23}$. This agrees very well with the value by Boley and Zaffarano of 2.99 ± 0.10 Mev and the 2.8 Mev cloud chamber measurement (24). Except for a possible very weak high energy tail in the October run, there was no evidence for another beta group.

The agreement with Boley in this case and the disagreement in the other cases can be explained in part by the fact that no other beta transition is apparently excited by 70 Mev x-ray bombardment. Since no other noticeable beta group existed, Boley's assumed endpoint, which was probably based on an extrapolation of the straight portion
Table 1. Beta Endpoint Energies for Various Runs.

<table>
<thead>
<tr>
<th>Nuclei</th>
<th>Endpoint (Mev)</th>
<th>Bomb. Energy (Mev)</th>
<th>Crystal thickness (inches)</th>
<th>Calibration beta spectra</th>
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<td>y(^{90})</td>
</tr>
<tr>
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<td>40</td>
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<td>C(^{134}), y(^{90})</td>
</tr>
<tr>
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<td>13.4</td>
<td>65</td>
<td>1(\frac{1}{2})</td>
<td>C(^{134})</td>
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</table>
Figure 7. Beta Spectra of Mg$^{23}$ and Y$^{90}$. 
Figure 8. Kurie Plots of Mg$^{23}$ and Y$^{90}$. 
of the Kurie plot to the energy axis, was the correct end-point within experimental error. Discussions of the effects of a second beta group are made under the isotopes with which a second beta group was observed.

Silicon 27

$^{27}\text{Si}$ was formed by a ($\gamma$,n) reaction on $^{28}\text{Si}$. The target was of silicon powder better than 97 per cent pure with the greatest impurity being iron at 1.5 per cent maximum. Figures 9 and 10 show the results of one of the four runs. Two of these runs were made under identical conditions with the exception that one was made with the synchrotron at 65 Mev and the other at 30 Mev maximum beam energy. The other two runs were made with the synchrotron at 65 Mev. The first two runs were recorded with Boley's one inch crystal mounted on an RCA tube, and the other two runs were recorded with the one and one-half inch crystal mounted on the Dumont tube. The results of these runs are given in Table 1.

A second beta group was noticed in each of these runs. A subtraction was done in the two cases which had the better statistics, and the endpoints of 3.1 and 3.2 Mev were obtained. An estimate was made of the relative recorded intensities of the highest energy beta group and the second group obtained by the subtraction process from the Kurie
Figure 9. Beta Spectra of $\text{Si}^{27}$ and $\gamma^{90}$. 
Figure 10. Kurie Plots of $\text{Si}^{27}$ and $\gamma^{90}$. 
plot.* The ratios were about two to one for the 65 Mev bombardment and about four to one for the 30 Mev bombardment. The amount of the second group as compared to the first increased when the bombarding time was increased from eight to ten seconds, the delay before recording increased from about one-half a second to one second, and the recording time in both cases kept at ten seconds. It is known that, in addition to the (γ,n) reaction, all other reactions that are energetically possible are probably produced to some extent. After the (γ,n) and the (γ,p) reactions, the (γ,np) and/or the (γ,d) reactions are the next most probable. Edwards and MacMillian (25) found some reactions were reduced in intensity by a factor of about ten compared to the (γ,n) reactions.

According to Kofoed-Hansen (26), Si$^{26}$ should have an energy of 3.64 Mev and a half-life of 3.4 seconds. Since the higher energy beta group was the main group observed, especially at 30 Mev, and since the calculated threshold for the (γ,2n) reaction is 30 Mev, the higher energy group was assigned to Si$^{27}$. The preceding paragraph indicates that the 3.15 Mev beta group probably belongs to an activity with a longer half-life and a higher threshold than Si$^{27}$. All of the statements are consistent with

*For the method used, refer to the Appendix.
assigning the 3.15 Mev group to Al^{26} which has reported values ranging from 2.8 to 3.4 Mev and a half-life of about seven seconds (27).

The value of 3.76 ± 0.08 Mev for Si^{27} was obtained by averaging the endpoint values of the four runs. Three of these were with Y^{90} as the calibration spectrum and the other was with Y^{90} and Cl^{34}. This is in fair agreement with the 3.48 ± 0.1 Mev value by Boley and the two cloud chamber measurements of 3.5 Mev (28) and 3.74 Mev (29).

If a straight line is extended from the 3.15 Mev beta group until it crosses the energy or channel number axis, the intersection is about 3.5 Mev. Since Boley did not suspect the presence of the second beta group, he corrected all points above the second beta group. These corrections had the effect of making the data agree with the endpoint assumption.

Phosphorous 29 and 30

In order to produce P^{29}, it was necessary to obtain a (γ,2n) reaction on P^{31}. In addition to the P^{29}, P^{30} is produced. Usually, other activities can be ignored by energy or half-life considerations, but this was not so in this case. Since there were some discrepancies in the reported value of P^{30}, a spectrum of P^{30} was recorded and the Kurie plot made.
To obtain the $\text{P}^{30}$ spectrum, $\text{P}^{31}$ in the form of a pressed wafer of red phosphorous was bombarded for five minutes with the synchrotron at maximum energy and then allowed to decay for three minutes before the recording apparatus was started. Figures 11 and 12 show the spectrum and the 3.32 Mev value obtained from the Kurie plot.

$\text{P}^{30}$ was produced in large quantities and a strong background of it built up in the attempt to run $\text{P}^{29}$ giving another endpoint determination of 3.30 Mev for $\text{P}^{30}$ (Figures 13 and 14). While this was not the result sought, it was very useful as a check on the first run. Due to the strong background of $\text{P}^{30}$, the $\text{P}^{29}$ endpoint determination cannot be considered reliable. Since any value obtained by this method could not improve the accuracy of the $3.945 \pm 0.010$ Mev beta endpoint value (30), no further attempt was made to record the spectrum of $\text{P}^{29}$.

The assigned value of $3.31 \pm 0.07$ Mev for $\text{P}^{30}$ is in fairly good agreement with the values of previous work. The value of 3.5 Mev was obtained with a spectrometer in 1941 (31). No experimental error was reported for this value. The 3.4 Mev value was obtained by absorption (32) and the 3.0 value with a cloud chamber (28).

Figures 13 and 14 serve as good examples of the difficulty of observing the $(\gamma,2n)$ reaction over the $(\gamma,n)$ reaction and, also, the effects of endpoint corrections.
Figure 11 Beta Spectra of P^{30}, Cl^{34}, and Y^{90}. 
Figure 12. Kurie Plots of $^{p30}$, $^{Cl34}$, and $^{Y90}$. 
Figure 14. Kurie plots of $p_{29} + p_{30}$, $C^{134}$, and $y_{90}$. 

Step attenuator setting in arbitrary units.
In Figure 14, the open circles along the lower portion of the line labelled as $P^{30}$ are the result of applying the endpoint corrections on the basis of a 3.30 Mev endpoint. From the result of these corrections, it can be seen that a weaker second beta group may be overlooked if its endpoint lies near the endpoint of the stronger group.

**Sulfur 31**

$S^{31}$ was formed by a ($\gamma$,n) reaction on $S^{32}$. The target was pressed sulfur powder. All of the recordings were made with the beam at 65 Mev, the bombard and record time at five seconds, and the record delay at one second. Figures 15 and 16 are the spectrum and Kurie plot obtained for one of the runs. In addition to the endpoints recorded for $S^{31}$, there was another activity formed with an endpoint of 3.3 Mev. The recorded intensity of this group as compared to the higher energy group was estimated to be about one to two for the 65 Mev bombardments. This activity seemed to be long lived as compared to the cycle duration, and when measured by Kline (33), a half-life in agreement with the 2.5 minute half-life of $P^{30}$ was obtained. This agrees with the discussion in Silicon 27 regarding the amount of the ($\gamma$,pn) and ($\gamma$,d) reactions as compared to the ($\gamma$,n).

Kofoed-Hansen predicts a half-life of 1.8 seconds and a maximum beta energy of 4.19 Mev for $S^{30}$. With the cycle times used and the low initial production of $S^{30}$ expected, the $S^{30}$ should not be evident over the background of $S^{31}$. 
Figure 15. Beta Spectra of $S^{31}$ and $Cl^{34}$. 
Figure 16. Kurie Plots of $S^{31}$ and $Cl^{34}$. 
If any was observed, it could not be separated from the bending of the Kurie plot caused by the loss of high energy electrons through the edge of the crystal. Since Phipps found a $2.40 \pm 0.07$ second half-life in $S^{31}$, the principal beta group was assigned to this isotope.

The two runs with the Cl$^{34}$ calibrations were assigned most of the weight in determining the $4.50 \pm 0.10$ Mev endpoint value for $S^{31}$, because the $S^{31}$ endpoint value was very close to the Cl$^{34}$ endpoint value.

The value of $4.50 \pm 0.10$ Mev is in considerable disagreement with the value of $4.06 \pm 0.20$ Mev by Boley and the two cloud chamber measurements of $3.85 \pm 0.07$ (34) and $3.87 \pm 0.15$ Mev (35). The second beta group with an endpoint energy of $3.3$ Mev may have led Boley to assume too low an endpoint for endpoint corrections. An extrapolation of the middle portion of the Kurie plot until it crossed the energy axis gave a value of about $4.0$ Mev.

Chlorine 33 and 34

The targets for both the Cl$^{33}$ and the Cl$^{34}$ were made of pressed NH$_4$Cl. The Cl$^{34}$ calibration target was unmounted, but the target for the three attempts to measure Cl$^{33}$ was mounted in the usual manner. The synchrotron was run at 65 Mev maximum beam energy and the cycling equipment was set to bombard four seconds and to record four seconds after a delay of from one-half to one second. A short half-life was
observed and recorded. Only one beta group with an endpoint of $4.45 \pm 0.10$ Mev was observed. This is in agreement with the work of Arber and Stahelin (36) which showed that the 1.58 second triplet isotopic spin state of Cl$^{34}$ decays with a 4.45 Mev maximum beta energy.

The masking of the $(\gamma,2n)$ reactions by the $(\gamma,n)$ reactions is in agreement with the assignments in other parts of this work and is, also, in agreement with the findings of Edwards and MacMillian that the relative yields of the reactions are probably of the order of one to ten, respectively. This would be further decreased due to the delay before recording if the half-life of Cl$^{33}$ were shorter than that of Cl$^{34}$.

Potassium 38

The targets were made of KI for three of the runs. The target for the other run was made of KF. These were pressed powder wafers which were mounted as described earlier. The targets were bombarded for two seconds and recorded for two seconds after a delay of about one-half second with the exception of the 5.10 Mev $^{90}$Y determination which was bombarded for six seconds and counted for six seconds after a delay of about one second.

The intention in bombarding KI was to produce $^{37}$K which was reported by Phipps (10), Boley (2), and Langmuir (37) to have a half-life of about one second. The value by Phipps of $0.98 \pm 0.02$ seconds is considered the best of these
measurements. KI was bombarded and a half-life of about one second was observed. This was recorded and the endpoint was determined to be $5.06 \pm 0.11$ Mev. Figures 17 and 18 show the spectrum and Kurie plot of one of the determinations. The values of the five determinations agree within experimental error. In determining the assigned endpoint value of 5.06 Mev the endpoint values determined by the $\gamma^{90}$ was assigned half the weight of the values determined by the $\text{Cl}^{34}$ calibration.

There are several reasons for assigning the 5.06 Mev value to $\text{K}^{38}$ rather than to $\text{K}^{37}$. Stahelin's (38) recently reported half-life of $0.95 \pm 0.03$ seconds together with Kofoed-Hansen's expected half-life of 0.9 seconds and maximum beta energy of 4.84 Mev agree very well with this assignment of 5.06 Mev to $\text{K}^{37}$. Neither Phipps, Boley, nor Langmuir reported an attempt to determine if the half-life which they reported could be produced below the $(\gamma,2n)$ threshold. Because of the thin samples and the decrease in the beam intensity as the energy was lowered, it was not possible to run the synchrotron below the $(\gamma,2n)$ threshold and to produce enough activity to record.

Stahelin and Kofoed-Hansen state that the one second half-life of $\text{K}^{38}$ belongs to the triplet isotopic spin state and the 7.7 minute half-life to the singlet state. Since the triplet and singlet states are expected not to differ much
Figure 17. Beta Spectra of $K^{38}$, $Cl^{34}$, and $Y^{90}$. 
Figure 18. Kurie Plots of $K^{38}$, $Cl^{34}$, and $\gamma^{90}$.
in energy, as in the case of $^{34}\text{Cl}$, the 5.06 Mev assignment is in good agreement with the 2.8 Mev beta group plus the 2.16 Mev gamma-ray. The 7.7 minute $^{38}\text{K}$ probably decays by beta emission to the 2.16 Mev excited state of $^{38}\text{Ar}$ which decays by gamma emission.

Kofoed-Hansen predicts that if the isotopic spin is a "good quantum number", the ft values would be expected to be the same for all similar transitions. The ft value of $3300 \pm 400$ for $^{38}\text{K}$ is in good agreement with the values of $3300 \pm 900$ for $^{14}\text{O}$ (39) and $3000 \pm 400$ for $^{34}\text{Cl}$.

The work of Edwards and MacMillan indicates that the ratio of $^{37}\text{K}$ to $^{38}\text{K}$ produced should be about one to ten. There was some evidence of a second weaker beta group with an endpoint energy of $3.9 \pm 0.2$ Mev. In the three cases in which the bombard time and record time were about two seconds and the delay before recording about one-half of a second, estimates of the recorded intensities of the two groups gave ratios of about one to five. When the six second bombard, one second delay, and six second record cycle was run, the ratio of the recorded intensities was about one to two. This indicates that the 3.9 Mev group is longer lived than $^{38}\text{K}$ and is probably due to $^{37}\text{K}$. Besides being the strongest activity recorded another indication that the 5.06 Mev beta group belongs to $^{38}\text{K}$ was obtained from the calcium bombardment. A second beta group of 5.05 Mev was found in calcium. The ratios of the recorded intensities in calcium of the 5.05
Mev to the 6.10 Mev group was about one to two for the 65 Mev bombardments and about one to four for the 40 Mev bombardment. Also, since no gamma-ray lines of the intensities necessary for the 5.05 Mev group were observed in calcium, the 5.05 Mev group is assumed to be caused by a reaction other than the $(\gamma, n)$ reaction.

If Boley actually measured a mixture of $^{37}$K and $^{38}$K, he would have probably assumed an endpoint of about 4.5 Mev on which to base his resolution corrections and would get fairly good agreement with his data.

**Calcium 39**

The target was made of pressed CaF powder. $^{39}$Ca was produced by a $(\gamma, n)$ reaction on $^{40}$Ca. The $^{39}$Ca spectrum was recorded four times, of which three were at 65 Mev maximum beam energy and the fourth at 40 Mev maximum beam energy. Two of these runs and their Kurie plots are shown as Figures 19, 20, 21, and 22. Figures 21 and 22 show the linearity check which was obtained on the crystal and photomultiplier tube by running the $^{8}$Li spectrum. This permitted, in addition to the linearity check, a determination of the $^{39}$Ca and a check on the 13.4 Mev endpoint of $^{8}$Li. The $^{8}$Li was produced by a $(\gamma, p)$ reaction on $^{9}$Be which was in the form of 0.005 inch foil attached to one of the nickel and balsa holders.
Figure 19 Beta Spectra of Ca$^{39}$, Cl$^{34}$, and Y$^{90}$. 
Figure 20. Kurie Plots of Ca39, Cl34, and Y90.
Figure 21. Beta Spectra of Ca$^{39}$, Li$^8$, and Cl$^{34}$. 
Figure 22. Kurie Plots of Ca39, Li8, and Cl34.
Since the 6.10 Mev value was the best determination, 6.10 ± 0.11 Mev was chosen as the endpoint value of Ca³⁹. The two lower values are not considered to be as accurate as the two upper values. This is because for one of the determinations, the calibration was only by Y⁹⁰; and for the other, Figure 22, the scale was such that only the last four points of the Kurie plot determined the endpoint. The upper two values were both obtained by Cl³⁴ and Y⁹⁰ calibrations.

The value of 6.10 ± 0.15 Mev disagrees both with Boley's value of 5.13 ± 0.20 Mev and Bramms and Smith's value of 6.7 ± 0.5 Mev (40). The latter value was determined by absorption techniques. The difference between the value reported here for Ca³⁹ and Boley's value can be explained by the second beta group, discussed under Potassium 38 which Boley was unable to detect with his equipment. The number of channels for calcium in Figures 21 and 22 is about the same as used in Boley's determination. Since he had a reported 44 per cent resolution for the 624 kev Cs¹³⁷ line, the bends in the Kurie plot would have been obscured and attributed to resolution spread. This would have led him to assume too low an endpoint for resolution corrections and thereby obtain too low a value.

According to Kofoed-Hansen, Ca³⁸ should have a half-life of 0.7 seconds and a maximum beta energy of 5.21 Mev. This half-life and energy are not much different than those for K³⁸. The amount of Ca³⁸ produced as compared to the
amount of K^{38} is probably about one to two for the 65 Mev bombardments. Because of the delay of about 0.5 seconds between bombarding and recording, the amount of Ca^{38} recorded as compared to the amount of K^{38} recorded would be reduced by another factor of about two.

Since the second beta group's endpoint of 5.05 Mev agreed with the endpoint assigned to K^{38} and since the intensity of the 5.05 Mev beta group as compared to the 6.10 Mev beta group was reduced by a factor of two when the maximum bombarding energy was reduced, the 5.05 Mev energy was assigned to K^{38} and the 6.10 value to Ca^{39}.

**Gamma-rays**

In order to determine that the other beta groups were not caused by another branch in the decay process, the activities investigated were searched for gamma-rays. The results of this search indicate that if branching does occur, it occurs in such small amounts that it would not show in the beta spectra in this investigation.

The gamma-ray sources were prepared by filling a nickel box with the material to be bombarded. The box was made the size of the balsa block so that the same transport system could be used. This permitted the NaI(Th) crystal, which was used for recording the gammas, to be placed in a position where it would not be activated by the X-ray beam.
from the synchrotron. About 0.5 inches of aluminum absorber were placed between the target and the crystal. This was sufficient to stop the expected beta-rays.

Two methods were used in this part of the investigation. The first was to bombard the target and to take still pictures. The results, which were also visible on the oscillograph, gave the expected intense line at 0.51 Mev and the Compton edge resulting from it. Since only one beta group was observed with an endpoint within 0.5 Mev of the reported endpoints, and since there was no evidence of any strong gamma-ray lines with energy less than 0.5 Mev besides that due to the annihilation radiation, no further investigations were carried out in this region.

In order to determine if the second beta groups observed were accompanied by gamma rays, the following procedure was used. Instead of taking photographs of the spectra, the number of gamma-rays with energies above the annihilation peak was compared to the total number with energies above some fixed value in the neighborhood of 0.2 Mev. This was accomplished in the following manner. The equipment for recording was set up as described for the photographs except that, instead of taking pictures, the pulses from the amplifier were fed into a discriminator which, if fired, produced a pulse that was counted by a scaler. The discriminator used was a single channel differential
analyzer, of the Oak Ridge type, used in the integral position. With the gain control on the amplifier turned to the lowest of ten fixed positions which resulted in an attenuation of about three times, the discriminator was set just above the annihilation peak. This was checked by observing that it passed the Cs$^{137}$ 0.66 Mev gamma-ray. When the gain of the amplifier was turned to its maximum position, pulses of about 0.2 Mev were passed.

The cycling equipment was operated such that the sample was bombarded and then counted for a few cycles. If no background was being built up, the runs were averaged and the figures were rounded off to the values shown in Table 2. Runs were made with the amplifier gain set at minimum and maximum, with and without the target, resulting in four sets of data for each bombardment energy. Trial runs on magnesium were made at various bombardment energies. Due to the low intensity of the synchrotron X-ray beam at low energies, it was necessary to raise the beam energy to 30 Mev before a usable amount of activity could be produced. Runs were made at 30 and 35 Mev maximum beam energy. Table 2 gives the results of these runs.

The background for all the similar runs without a target was averaged to give the gamma-ray background in the room. If the backgrounds are subtracted from the appropriate columns, an estimate of the ratio of gamma-rays
above 0.6 Mev to those above 0.2 Mev can be obtained. For the 35 Mev bombardments the ratio is the order of two per cent in each case. Since these values represented integrated spectra, it can be concluded that if branching occurs with

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<td>30</td>
<td>No appreciable activity above phosphorous 30 background.</td>
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<tr>
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<td>230</td>
<td>200</td>
</tr>
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</table>

the emission of a gamma-ray of greater than 0.6 Mev energy, the intensity of these transitions is very small as compared to the annihilation radiation intensity. A further conclusion is that the second beta groups are not due to branching since the intensities would be so low that they would not record.
IV. SUMMARY AND CONCLUSIONS

A comparison of the results of this investigation and those of others is principally a comparison of this work and that of Boley and Zaffarano. Because of the large differences in some of the reported values and the values reported here, each spectrum in question was repeated several times, sometimes months apart, in order to assure that the results were not caused by some unknown change in the recording equipment between the time the calibration was placed on the film and the unknown spectrum was recorded.

Boley's dot spectra were counted by eye on a microfilm viewer. It was not practical to have many channels or to count large numbers of events. The number of channels along the straight portion of his Kurie plots varied between six and nine. Because of the 44 per cent resolution at 624 kev, three or four of these channels were corrected for resolution effect or endpoint corrections. As mentioned earlier, the endpoint corrections have the effect that if the endpoint is assumed too low, the corrections tend to make the corrected points agree with the assumption.

The endpoint energies obtained in this investigation and the best known values of the half-lives are combined in Table 3 to give the ft values.
With the exception of $K^{38}$ and $Cl^{34}$, the $ft$ values increase as the atomic number increases. This indicates that the $ft$ value assigned to $K^{38}$ does not seem to belong to the mirror

### Table 3. Beta Endpoint Energies and $ft$ Values Obtained in This Investigation.

<table>
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<tr>
<th>Nuclei</th>
<th>$E_{max}$ (MeV)</th>
<th>$t_\frac{1}{2}$ (sec)</th>
<th>$ft$ (sec)</th>
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<tr>
<td>$Mg^{23}$</td>
<td>$2.95 \pm 0.07$</td>
<td>$10.7 \pm 0.7$ (10)</td>
<td>$3600 \pm 500$</td>
</tr>
<tr>
<td>$Si^{27}$</td>
<td>$3.76 \pm 0.08$</td>
<td>$4.05 \pm 0.10$ (10)</td>
<td>$4000 \pm 450$</td>
</tr>
<tr>
<td>$S^{31}$</td>
<td>$4.50 \pm 0.10$</td>
<td>$2.40 \pm 0.07$ (10)</td>
<td>$5250 \pm 600$</td>
</tr>
<tr>
<td>$Cl^{34}$</td>
<td>$4.45 \pm 0.10$</td>
<td>$1.58 \pm 0.05$ (38)</td>
<td>$3000 \pm 400$</td>
</tr>
<tr>
<td>$K^{38}$</td>
<td>$5.06 \pm 0.11$</td>
<td>$0.95 \pm 0.03$ (38)</td>
<td>$3250 \pm 400$</td>
</tr>
<tr>
<td>$Ca^{39}$</td>
<td>$6.10 \pm 0.15$</td>
<td>$0.90 \pm 0.05$ (33)</td>
<td>$7150 \pm 1000$</td>
</tr>
<tr>
<td>$P^{30}$</td>
<td>$3.31 \pm 0.07$</td>
<td>$150$ (27)</td>
<td></td>
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</table>

nuclei, but to the triplet isotopic spin state where the Fermi interaction form, $|j_1|^2$, probably has the value of two (26) instead of one as in the case of the mirror nuclei. Doubling the $ft$ values for $K^{38}$ and $Cl^{34}$ brings them into agreement with the trend of the other $ft$ values.

The $ft$ values, combined with the values of $|\sigma|^2$ derived by Winther and Kofoed-Hansen do not lead to their unique value for the ratio of $G - T$ to Fermi interaction strength constants, which they obtained by using previous data.

Since improved equipment is being developed, these activities should be examined again for second beta groups
and for better endpoint determinations. The endpoints should be re-run if enough activity can be produced below the ($\gamma$,d) threshold to record.

Further investigations should be made for gamma-rays, especially below 0.51 Mev.
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It is also a pleasure to acknowledge the support given this work by the Ames Laboratory for the research assistantship and for the equipment provided.
VII. APPENDIX

An estimate of the relative intensities of two beta groups separated by the usual subtraction process with a Kurie plot can be made from the two slopes observed near the endpoint energies. From the Fermi theory of beta decay, it is evident that the ratio of the number of events in each group will be given by

$$\frac{N_2}{N_1} = \left[ \frac{T_2}{T_1} \right]^2 \frac{f_2}{f_1}$$

where the subscripts 1 and 2 refer to the first and second beta groups, respectively, N is the total number of events, T is the observed slope, and f is the dimensionless integral,

$$f = \int_{E_0}^{E_0} F(E,Z)(E^2 - 1)^{\frac{1}{2}} E(E_0 - E)^2 dE,$$

which has been evaluated and presented graphically by Feenberg and Trigg (41) and Moszkowski (42) for the allowed transitions.*

The use of the above relation for the relative intensities involves the assumption that the beta spectrum observed with a scintillation spectrometer is not appreciably affected near the endpoint by the degradation

*The terms in the latter equation have been defined in Chapter I.
of the energy spectrum caused by the electrons scattering out of the highest energy third of the spectrum, as demonstrated by Figures 4 and 6.