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An investigation of the radionuclides of
arsenic produced by cyclotron bombardment
of germanium with 15 Mev neutrons

Watters, Harry J.; Fagen, John F.

Massachusetts Institute of Technology

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AN INVESTIGATION OF THE RADIONUCLIDES
OF ARSENIC PRODUCED BY CYCLOTRON
BOMBARDMENT OF GERMANIUM WITH
15 MEV DEUTERONS

—♦♦♦—
HARRY J. WATTERS
AND
JOHN F. FAGAN, JR.

Thesis
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AN INVESTIGATION OF THE RADIONUCLIDES OF ARSENIC
PRODUCED BY CYCLOTRON BOMBARDMENT OF GERMANIUM

with

15 Mev DEUTERONS

by

HARRY J. WATTERS
Lieutenant Commander, U. S. Navy
B. S. Purdue University
(1949)

and

JOHN F. PAGAN, JR.
Lieutenant, U. S. Navy
B. S., U. S. Naval Academy
(1945)

SUBMITTED IN PARTIAL FULFILLMENT OF THE
REQUIREMENTS FOR THE DEGREE OF
MASTER OF SCIENCE

ABSTRACT

Title: "An Investigation of the Radionuclides of Arsenic Produced by Cyclotron Bombardment of Germanium with 15 Mev Deuterons"

**Authors: Harry J. Watters, Lieutenant Commander, U.S. Navy
B. S., Purdue University (1949)**

and

**John F. Fagan, Jr., Lieutenant, U. S. Navy
B. S., U. S. Naval Academy (1945)**

Submitted to the Department of Physics on May 25, 1953 in partial fulfillment of the requirements for the degree of Master of Science.

RESUME

THIS AN INVESTIGATION OF THE EFFECTS OF STRESS
PERFORMED BY OFFICERS MEMBERS OF THE
NAVY IN THE PACIFIC

ANALYSIS MADE BY OFFICERS MEMBERS OF THE
U. S. NAVY (1942)

and

JOHN F. GARDNER, JR., COMMANDER, U. S. NAVY
U. S. NAVY (1942)

PRESENTED TO THE DEPARTMENT OF THE NAVY
IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE
DEGREE OF MASTER OF SCIENCE.

The arsenic produced by a deuteron bombardment of germanium has been studied to determine the nuclides present in the mixture. Identification of isotopes was made by comparing measured values of half life and maximum β energy with the accepted values. Yield values were determined for each isotope present by 4π solid angle counter measurements.

Counting rates were measured for a period of 53 days with 4π and coincidence counters, obtaining half lives which indicated that the nuclidic mixture was made up of As^{71} , As^{72} , As^{73} , As^{74} , and As^{77} . These indications were confirmed by maximum β energy values obtained by absorption measurements and from γ energies found using a γ -ray scintillation spectrometer. Measurements indicated that the 40 hour half life reported for As^{77} is in error by a significant amount, and that no As^{76} was obtained by this bombardment.

The 4π solid angle counter constructed was shown to have an efficiency of very nearly 100 percent for particles which escape the source. This counter has proven to be a very practical laboratory instrument and detailed instructions for its use are included as an appendix.

The current program by a business department of
Germany has been studied to determine the results
present in the future. Identification of factors was
made by comparing measured values of half life and ex-
cess with the expected values. This value was
determined for each sample measured by the same
method.

Results were measured for a period of 25 days
with the following procedure, starting with five
days before the end of the experiment was made up of
the following: $V_1, V_2, V_3, V_4, V_5, V_6, V_7, V_8, V_9, V_{10}$. These last five were
measured by means of a single value obtained by averaging
measurements and the 7 samples from the 2-5 day
relaxation experiment. Measurements indicated that
the 25 days half life reported for V_1 is in error by a
significant amount, and that the V_2 was measured by this
method.

The 25 days half life obtained was shown to
have an efficiency of very nearly 100 percent for particles
which escape the source. This constant has proved to be a
very important laboratory instrument and should be in-
cluded for its use and included as an appendix.

The following is a tabular summary of the results of the investigation:

Isotope	Method of decay	Energy (MeV)	$T_{1/2}$	Thick target yield* (uc/ μ amp-hr)
As ⁷¹	β^+	0.66	48.2 \pm 1.2 hrs.	7.6
As ⁷²	β^+	3.25	25.8 \pm 0.2 hrs.	64.9
	γ	0.85		
As ⁷³	β^-	0.11 $> E_{max} >$ 0.02	88.9 \pm 9.2 days	1.1
As ⁷⁴	β^+	0.99, 1.49	17.8 \pm 0.13 days	5.2
	β^-			
As ⁷⁶	Not present in the mixture			
As ⁷⁷	β^-	< 0.7	> 70 hours	5 < yield < 15**

* The thick target yield values specified apply if the deuteron beam current was exactly 36 μ amps and if the arsenic separation efficiency was 100 percent. Yield values quoted are based on β counting only and do not include orbital electron capture.

** Based on ratios of total β to β^+ counting rates.

Thesis Supervisor: Robley D. Evans

Title: Professor of Physics

The following is a partial summary of the results

of the investigation

Initial Value of Δ	Δ	Final Value of Δ	Number of Steps	Approximate Value of Δ
1.0	1.0 ± 0.0	1.0	1	1.0
0.5	0.5 ± 0.0	0.5	1	0.5
0.1	0.1 ± 0.0	0.1	1	0.1
0.01	0.01 ± 0.0	0.01	1	0.01
0.001	0.001 ± 0.0	0.001	1	0.001
0.0001	0.0001 ± 0.0	0.0001	1	0.0001
0.00001	0.00001 ± 0.0	0.00001	1	0.00001

See Appendix for details

The above table shows that the value of Δ remains constant for all values of the initial value of Δ . This is expected since the value of Δ is determined by the initial value of Δ and is independent of the number of steps. The results show that the value of Δ is constant for all values of the initial value of Δ .

See Appendix for details of the calculation of Δ .

TABLE I
RESULTS OF INVESTIGATION

MASSACHUSETTS
STATE OF MASSACHUSETTS

ACKNOWLEDGMENTS

MASSACHUSETTS INSTITUTE OF TECHNOLOGY
DEPARTMENT OF CHEMISTRY
77 MASSACHUSETTS AVENUE
CAMBRIDGE, MASSACHUSETTS 02139

The authors wish to express sincere thanks to their thesis advisor, Professor Robley D. Evans, for his interest and advice during the course of this work. Grateful acknowledgment is also made to Doctor Gordon L. Brownell, who suggested the problem, for his constant guidance and for the opportunity to utilize the facilities of the Massachusetts General Hospital Research Laboratory.

Thanks are due to all members of the group in the Radioactivity Center for their interest and suggestions. The opportunity for graduate study provided by the Radioactivity Center and its sponsors is greatly appreciated.

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MEMORANDUM

The subject also in review should relate to
 that basic subject, however being in force for
 his interest and other things the nature of this
 year. Special arrangements is also to be made
 under a special, but subject to review, the
 his interest in the subject and the committee is
 within the jurisdiction of the Department of State
 United States, Secretary.

There are no in all members of the group in the
 administrative order for their interest and organization.
 The committee for review also provided by the
 review center and the subject in review.

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I. INTRODUCTION

A. Importance of the Investigation

The fact that tumors of the brain take up a large amount of trace metal compared with that taken up by normal brain tissue makes it possible to detect and to actually determine the location of tumors in the human brain. Under the supervision of Dr. Gordon L. Brownell, a group at the Research Laboratory of the Massachusetts General Hospital has developed a suitable tracer technique for the diagnosis and preoperative location of brain tumors, using positron emitting isotopes. After intravenous injection of the tracer material, the patient's head is mechanically scanned in two dimensions by two scintillation counters connected in coincidence. Third dimensional location is obtained from the unbalanced single channel counting rates of the separate counters.

Since January 1953 a large number of patients have been examined using this technique. The results are

REPORT OF THE INVESTIGATION

The first part of the work was a
 large amount of time was spent in the
 to be made clear that it is not
 but in which the location of the
 the same data. Under the direction of Dr. G. L. Fisher,
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 has been organized to study the
 of the system and to determine
 the nature of the interaction between
 the system and the environment.
 The first part of the work was a
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 has been organized to study the
 of the system and to determine
 the nature of the interaction between
 the system and the environment.

outstanding. From many cases clinically diagnosed as borderline, the presence or absence of neoplastic brain tissue has been determined by this method. In all cases where surgery was performed, tumor location obtained by this technique has been confirmed. As yet no known incorrect diagnoses have been made. In addition to providing more quantitative information than is available from clinical diagnosis, this method provides the left-to-right localization which is difficult and often impossible to obtain clinically.

Radioactive arsenic was selected as the tracer metal because of several considerations. Arsenic is readily available from a deuteron bombardment of germanium. The half lives of arsenic isotopes fall within an acceptable range for tracer utilization. Most of the γ -rays emitted from arsenic isotopes are soft, thus decreasing harmful biological effects due to radiation. A very important advantage is that a large percentage of arsenic activity consists of positron emission. Precision measurements with very high resolution may be made on the resulting annihilation radiation.

The tracer arsenic is not injected until several days after bombardment. During this period any short-

outstanding. From many cases clinically diagnosed
as bacterial, the presence of bacteria in
brain tissue has been demonstrated in this method. In
all cases where surgery was performed, the bacteria
obtained by this method has been confirmed. In 1940

no known bacterial diseases have been seen. In
addition to providing more quantitative information
this is available from clinical records, this method
reveals the left-to-right localization which is often
not and often impossible to obtain clinically.

Inductive studies are selected as the most
total because of several considerations. It is in
positive results from a complete assessment of the
system. The full range of clinical features will
often be available only for these studies.
Most of the years which have been examined are
very few bacterial central nervous system cases
to include. A very important advance is that a
large percentage of central nervous system cases of
brain infection. Positive results with very little
evidence may be seen in the resulting localization
method.

The most common is not infrequently still several
days after diagnosis. During this period any acute

lived activity present decays to a negligible value compared with that of the 17.5 day isotope^(11,13) and does not affect the scanning measurements which require a period of approximately two hours. If the half lives and the relative activity percentages of the short-lived isotopes were accurately known, this waiting period could be decreased or even eliminated with a resultant increase in useful activity obtained from a given bombardment.

The purpose of the present investigation is to determine insofar as possible the methods of decay and associated decay energies, half lives, absolute activities, and isotopic yields of the arsenic obtained by the deuteron bombardment of germanium. In addition to decreasing the delay between bombardment and injection, this information may permit the use of short-lived isotopes as tracers. In effect this also decreases the bombardment time required to obtain a given amount of tracer material. It may be desirable to examine a single patient several times over a period of a few weeks. Accurate knowledge of the short-lived activity present may permit frequent injections of a lesser amount of tracer solution while avoiding harmful effects from the chemical toxicity of carrier arsenic present.

(11411) [11411] I have not yet received a copy of the report from the Bureau of the Census. It is the only report of the kind that has been published. The Bureau of the Census is the only one that has the data for the entire country. The Bureau of the Census is the only one that has the data for the entire country. The Bureau of the Census is the only one that has the data for the entire country.

The purpose of the present investigation is to determine the effect of the Bureau of the Census on the activities of the Bureau of the Census. The Bureau of the Census is the only one that has the data for the entire country. The Bureau of the Census is the only one that has the data for the entire country. The Bureau of the Census is the only one that has the data for the entire country.

B. Results of Previous Investigations

Prior to Sagan's investigations in 1938⁽¹⁾ very little was known about the radionuclides of arsenic. The principal results of his work on the arsenic produced by a deuteron bombardment of germanium, as modified by others, are tabulated below and include all data reported through 1941.

<u>Isotope</u>	<u>Type radiation</u>	<u>Energy (MeV)</u>	<u>Half life</u>	<u>Reference</u>
As ⁷¹	β^+		50 hour	2, 4
As ⁷³	β^+	0.6	68 min	2, 4
As ⁷⁴	β^+	0.9	18 day	1, 4
As ⁷⁶	β^-	1.1	26.8 hour	3, 5, 6
		1.7		
		2.7		
	γ	1.5		
		2.2		
		3.2	17.5 day	11, 28
As ⁷⁷	β^-		55-90 day	1, 2, 4
	γ			
As ⁷⁸	γ	0.57	68 min	7

Very little new information was published for several years but commencing in 1948 results were published which

5. Results of Previous Investigations

It is to be noted that the present investigation is a continuation of the work of the author in 1938 (1) and 1940 (2). The present results of his work on the effects of a constant temperature of 20°C. on the physiological reactions of the rat are given in Table I. It is to be noted that the data reported in Table I are all data reported in 1938 (1).

Experiment	Temp. (°C.)	Temp. (°C.)	Temp. (°C.)	Temp. (°C.)
1	20	20	20	20
2	20	20	20	20
3	20	20	20	20
4	20	20	20	20
5	20	20	20	20
6	20	20	20	20
7	20	20	20	20
8	20	20	20	20
9	20	20	20	20
10	20	20	20	20
11	20	20	20	20
12	20	20	20	20
13	20	20	20	20
14	20	20	20	20
15	20	20	20	20
16	20	20	20	20
17	20	20	20	20
18	20	20	20	20
19	20	20	20	20
20	20	20	20	20

Very little new information was obtained for these 20 experiments. In fact, the only new results were obtained when

conflicted with much of the previous data. The following is a tabulation of the most reliable data now available on the radioisotopes of arsenic without regard to their method of activation:

<u>Isotope</u>	<u>Type radiation</u>	<u>Energy (MeV)</u>	<u>Half life</u>	<u>Reference</u>
As^{70}	β^+		52 min	9
As^{71}	β^+ (33%)	0.6	50-60 hour	10, 11, 12
	K^+ (67%)	0.162		
As^{72}	β^+	0.27	26 hour	9, 11, 13
		0.67		
		1.84		
		2.5		
		3.34		
As^{72}	γ	0.702		
		0.835		
As^{73}	K^+	0.052	76-100 day	11
As^{74}	no β^+		17.5 day	11, 13
	β^-	0.69, 1.36		
	β^+	0.92, 1.53		
	$\beta^-/\beta^+ \sim 1.0\%$			
As^{76}	γ	0.593	27.6 hour	14, 18
		$\beta^+/\beta^- \leq 0.07\%$		
As^{77}	β^-	0.55, 1.21	40 hour	15, 16, 20
		0.679, 0.7		
As^{77}	no γ			

The following table shows the results of the analysis of variance for the different treatments. The values in parentheses are the values of the mean squares for the different treatments. The values in brackets are the values of the mean squares for the different treatments. The values in the last column are the values of the mean squares for the different treatments.

Treatment	Mean Square	Sum of Squares	D.F.	F-value
Control	1.0	1.0	1	
Treatment 1	1.0	1.0	1	
Treatment 2	1.0	1.0	1	
Treatment 3	1.0	1.0	1	
Treatment 4	1.0	1.0	1	
Treatment 5	1.0	1.0	1	
Treatment 6	1.0	1.0	1	
Treatment 7	1.0	1.0	1	
Treatment 8	1.0	1.0	1	
Treatment 9	1.0	1.0	1	
Treatment 10	1.0	1.0	1	
Treatment 11	1.0	1.0	1	
Treatment 12	1.0	1.0	1	
Treatment 13	1.0	1.0	1	
Treatment 14	1.0	1.0	1	
Treatment 15	1.0	1.0	1	
Treatment 16	1.0	1.0	1	
Treatment 17	1.0	1.0	1	
Treatment 18	1.0	1.0	1	
Treatment 19	1.0	1.0	1	
Treatment 20	1.0	1.0	1	

II. NUCLEAR PROPERTIES TO BE MEASURED

Time and equipment limitations prohibited conducting an investigation which could determine actual decay schemes of the active material. With a desire to extract as much information as possible in the time available, attempts were made to determine the following for each isotope of arsenic obtained from the bombardment:

1. Absolute β activity.
2. Half life.
3. Maximum β energies.

In addition it was desirable to obtain information regarding the γ -energies of the mixture of isotopes and the variation of the spectrum with time.

The two systems investigated exhibited similar behavior in the presence of the active material. It is noted that the extent of such information as possible in the literature available, attempts were made to determine the difference between the two systems obtained from the data-

and

1. Absolute activity.

2. Half life.

3. Half-life constant.

In addition it was desirable to obtain information regarding the percentage of the amount of isotope and the variation of the spectrum with time.

III. EXPERIMENTAL PROCEDURE

A. Preparation of Radioactive Arsenic

A chip of pure germanium metal $1/32$ inch thick with dimensions $3/8$ inch by 1 inch was used as a target in the M.I.T. cyclotron. This chip was bombarded with 15 Mev deuterons for a period of 20 minutes with an average beam current of 36 pamps.

After bombardment the germanium metal was oxidized to GeCl_4 in an evacuated system using gaseous Cl_2 .

To this was added HCl , H_2O_2 , and arsenic carrier after which the bulk of the GeCl_4 was distilled out. The arsenic remaining in the solution as As^{+5} was precipitated as a metal by the addition of ammonium hypophosphite. A detailed description of this separation procedure is contained in reference 21.

B. Schedule of Observations

Continuous observations were made of the disintegration rate of the active material by use of the 4 π and coincidence counters. In an attempt to ascertain

A. Preparation of Inactivated Vaccine

A quantity of pure glycerol was added to the vaccine with constant stirring for 1 hour and was then allowed to settle in the 500 ml. bottle. This was repeated with the vaccine for a total of 20 hours with an average time interval of 16 hours. After preparation the vaccine was allowed to settle for 24 hours in an inverted position using constant stirring. To this was added 10 ml. of 1% formalin solution after which the bulk of the vaccine was distilled over. The vaccine remaining in the solution as a residue was prepared as a solid by the addition of calcium hydroxide. A detailed description of this procedure is contained in reference 11.

B. Results of Observations

Continuous observations were made of the distribution of the active material by use of the 500 ml. and 100 ml. flasks. In an effort to determine

whether or not the arsenic contained any positron-emitting isotopes having half lives of the order of 1 hour or less (^{23}As , ^{24}As), coincidence counter measurements were made as follows: each minute during the third hour after bombardment, every 5 minutes during the fourth hour, every 10 minutes during the fifth hour, and every 15 minutes during the sixth hour. Thereafter the maximum interval between measurements was adjusted to approximately 1/10th the value of the half life indicated by a continuous plot of counting rate observations.

Due to the time required for preparation of ^{47}As counter sources and the time involved in making absorption measurements with the end window β counter, observations with these instruments were made hourly from the 6th through the 17th hour after bombardment, and thereafter in accordance with the schedule outlined above.

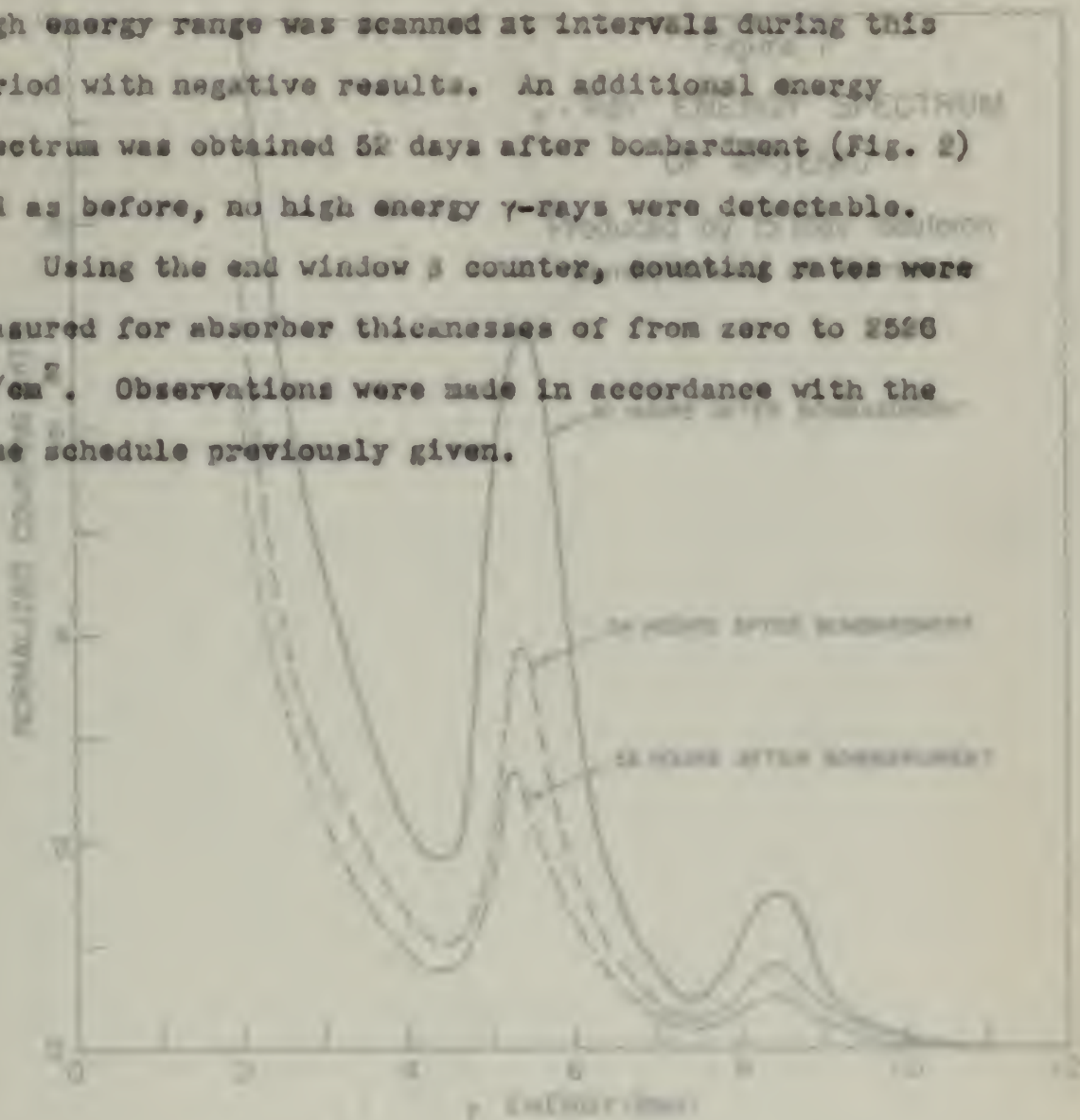
Using the sodium iodide scintillation spectrometer described in Appendix II an initial scan of the energy spectrum up to 3 Mev was made within three hours after bombardment in order to determine the maximum energy γ -rays emitted from the arsenic. With no detectable γ -energies present greater than 1 Mev, an operating range

whether or not the results obtained are similar
with the same having full view of the order of
how or less (1, 2, 3, 4, 5, 6, 7, 8, 9, 10)
with the same as follows: - some results during the
first two after each hour, every 2 hours during
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and every 15 minutes during the first hour. However,
the amount of time between measurements was adjusted
to approximately 15 min the value of the half life
indicated by a continuous list of results of the same
time.

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was chosen which included all γ -energies up to approximately 1.3 Mev. This energy range was scanned continually for the first 72 hours after bombardment (Fig. 1). The high energy range was scanned at intervals during this period with negative results. An additional energy spectrum was obtained 52 days after bombardment (Fig. 2) and as before, no high energy γ -rays were detectable.

Using the end window β counter, counting rates were measured for absorber thicknesses of from zero to 2526 mg/cm². Observations were made in accordance with the time schedule previously given.



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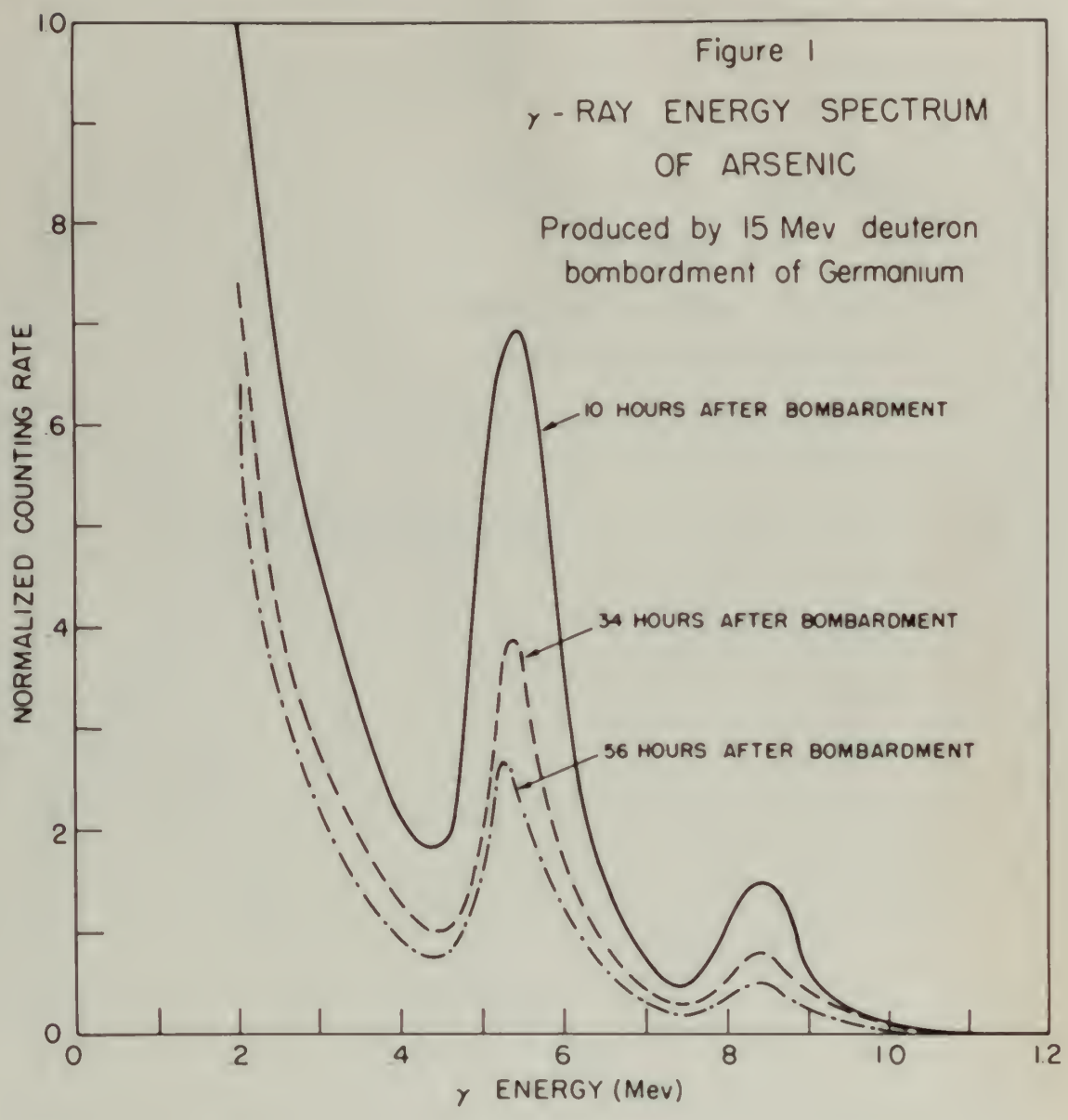
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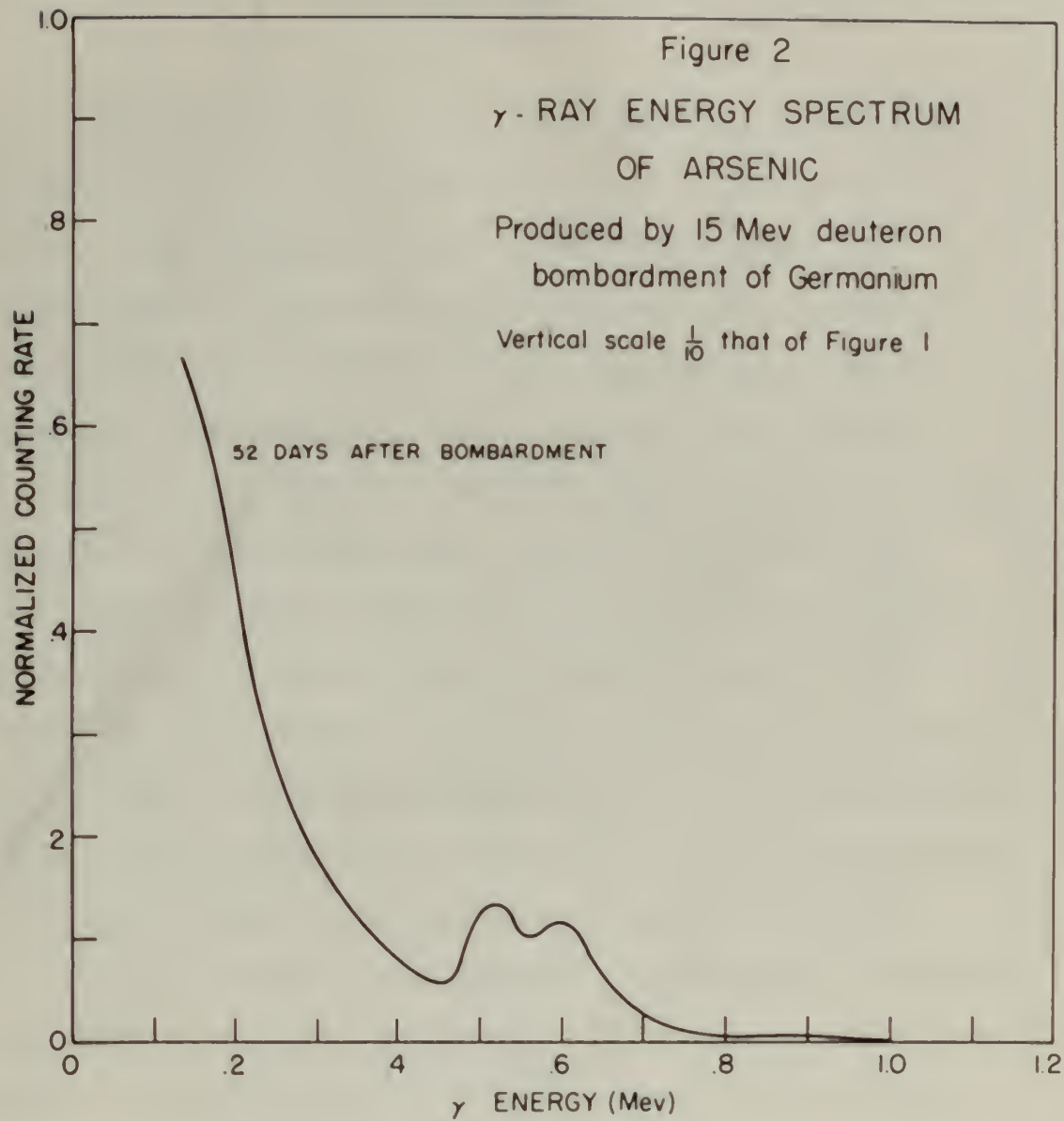
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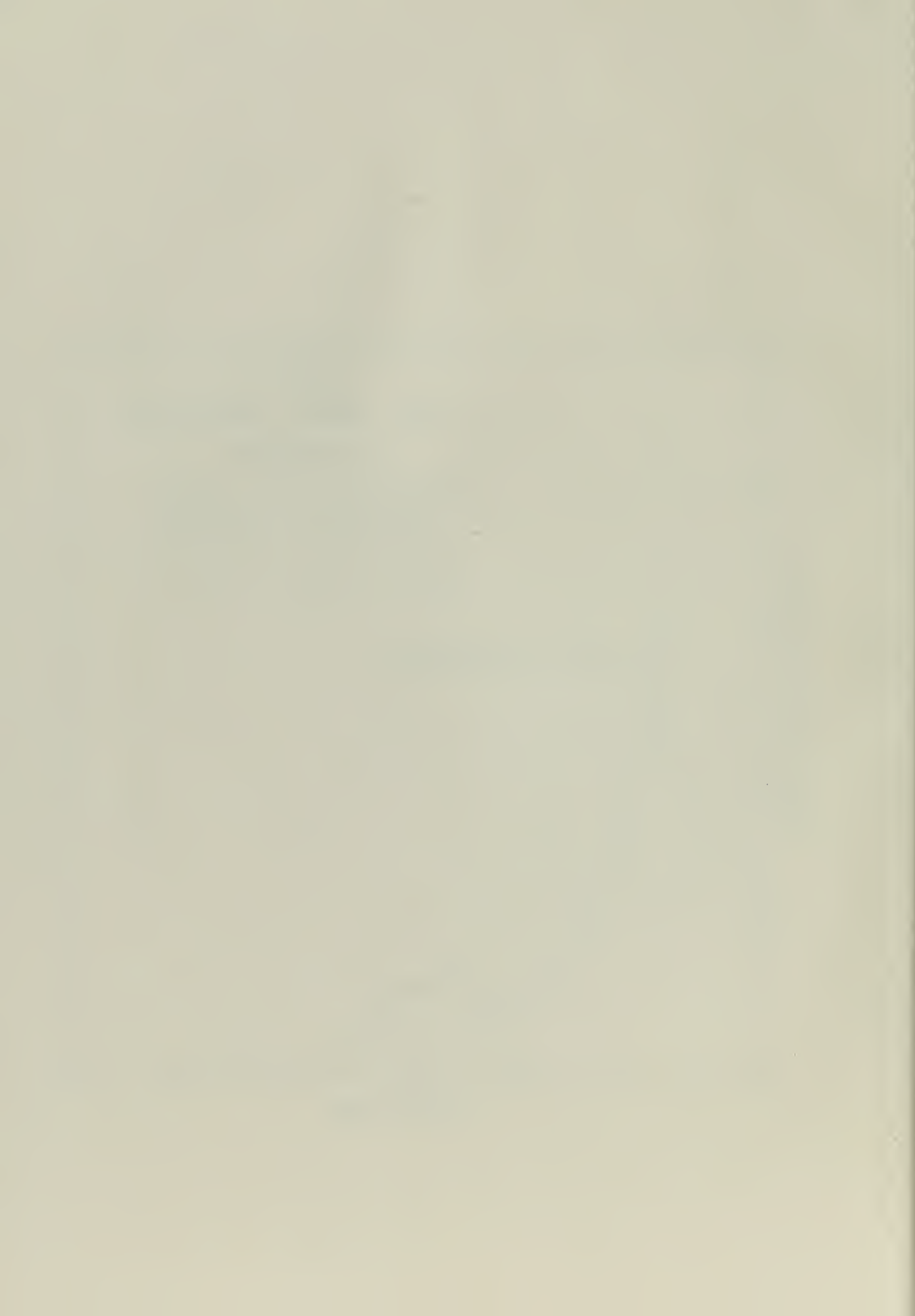
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IV. METHODS USED IN INTERPRETATION OF DATA

A. Half Life

Observed counting rate, corrected for instrumental error, was plotted on semilog paper as a function of time. Approximately 20 days after bombardment the curve obtained from coincidence measurements assumed a constant slope indicating the presence of a single isotope. Application of the method of least squares to data in the region of constant slope yielded a determination of half life, zero time activity, and their respective standard deviations. Subtraction of values thus obtained from the curve of total counting rate resulted in a residual curve also possessing a constant final slope. Successive application of this method permitted the resolution of 3 straight line components from the data obtained by coincidence counting (Fig. 3).

The 4 π counter data included a relatively long-lived component which was not apparent in coincidence measurements. Assuming this to be As⁷³ reported as a 0.05 Mev

A. INTRODUCTION

On a previous occasion, the interaction of light with a medium was studied. It was found that the interaction of light with a medium is a function of the frequency of the light. The present study is a continuation of the study of the interaction of light with a medium. The purpose of this study is to determine the interaction of light with a medium as a function of the frequency of the light. The results of this study are presented in the following sections.

The interaction of light with a medium is a function of the frequency of the light. It is found that the interaction of light with a medium is a function of the frequency of the light. The results of this study are presented in the following sections.

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The interaction of light with a medium is a function of the frequency of the light. It is found that the interaction of light with a medium is a function of the frequency of the light. The results of this study are presented in the following sections.

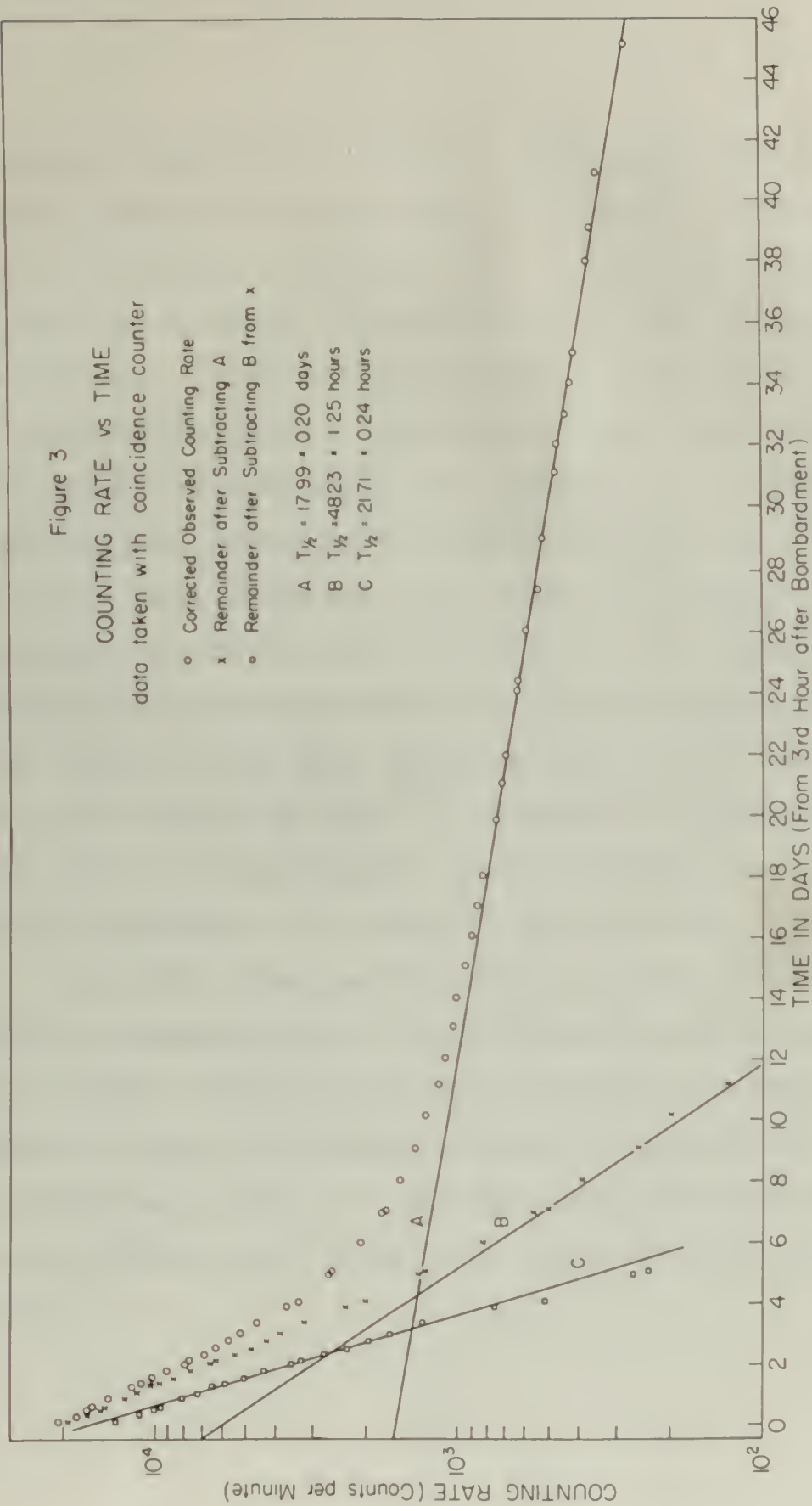
Figure 3

COUNTING RATE vs TIME

data taken with coincidence counter

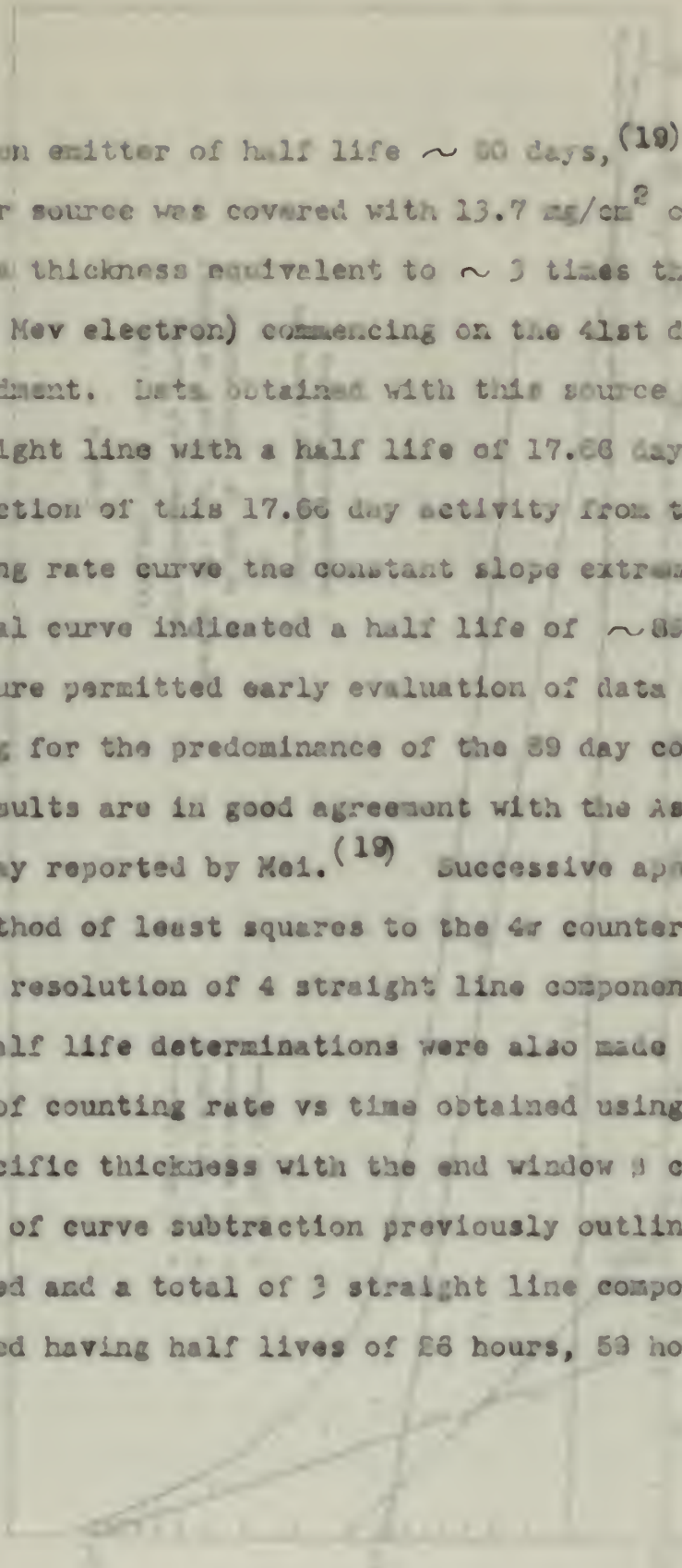
- o Corrected Observed Counting Rate
- x Remainder after Subtracting A
- o Remainder after Subtracting B from x

A $T_{1/2} = 17.99 \pm 0.20$ days
 B $T_{1/2} = 4823 \pm 125$ hours
 C $T_{1/2} = 2171 \pm 0.24$ hours



negatron emitter of half life ~ 80 days, ⁽¹⁹⁾ one 4 π counter source was covered with 13.7 mg/cm² of aluminum foil (a thickness equivalent to ~ 3 times the range of a 0.05 Mev electron) commencing on the 41st day after bombardment. Data obtained with this source plotted as a straight line with a half life of 17.66 days. After subtraction of this 17.66 day activity from the total counting rate curve the constant slope extremity of the residual curve indicated a half life of ~ 85 days. This procedure permitted early evaluation of data without waiting for the predominance of the 59 day component and the results are in good agreement with the As⁷³ method of decay reported by Mel. ⁽¹⁹⁾ Successive application of the method of least squares to the 4 π counter data resulted in the resolution of 4 straight line components. (Fig. 4)

Half life determinations were also made from semilog plots of counting rate vs time obtained using absorbers of specific thickness with the end window β counter. The method of curve subtraction previously outlined was employed and a total of 3 straight line components were resolved having half lives of 28 hours, 59 hours, and 16.7 days.



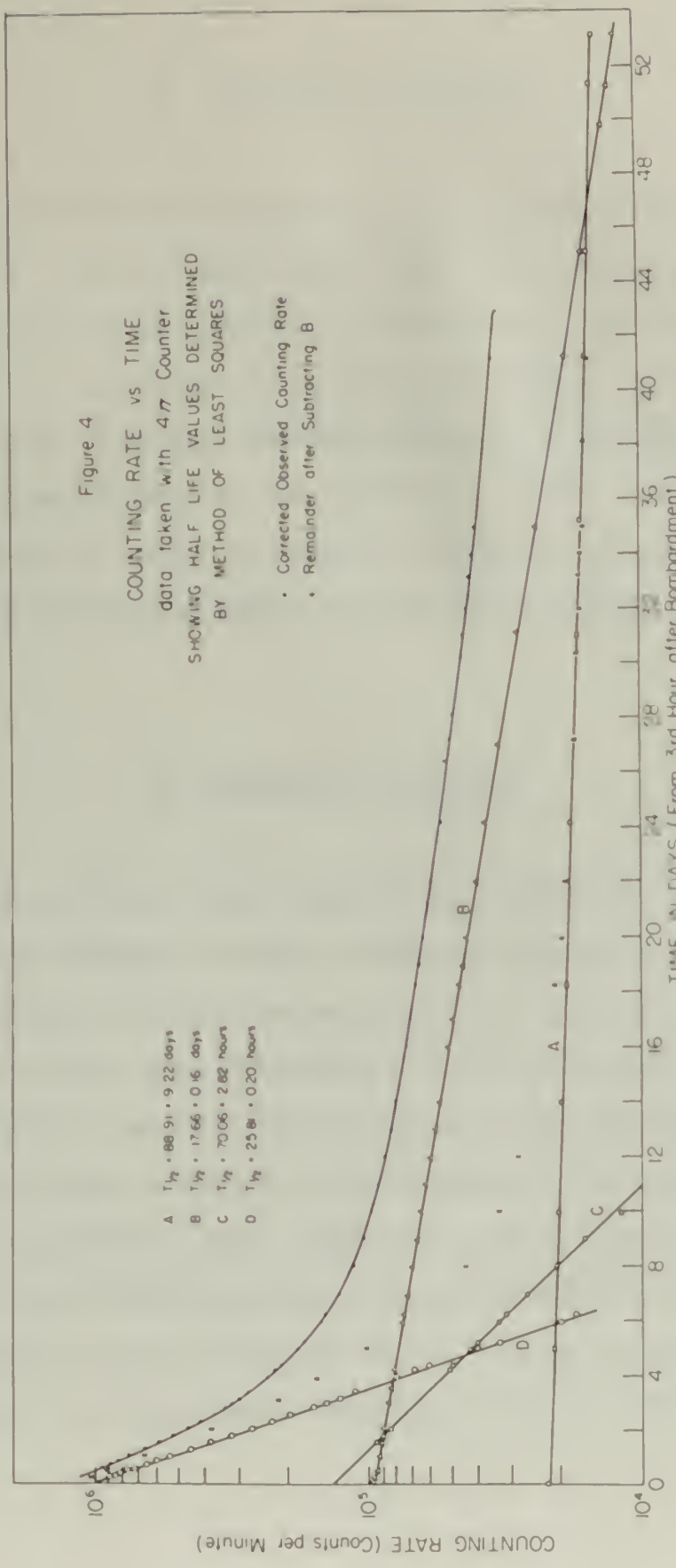
negative effect of half life of ^{131}I on the
 counter source was observed after 15.7 days of
 fall (a similar result was obtained in 3 days after
 a 0.05 day electron) according to the data after
 bombardment. This showed that the source activity
 a straight line with a half life of 15.7 days after
 extraction of this ^{131}I activity from the total
 counting rate curve was observed along activity of the
 residual curve indicated a half life of 15.7 days. This
 procedure permitted early evaluation of data without
 waiting for the termination of the ^{131}I component and
 the results are in good agreement with the ^{131}I half
 life of 15.7 days reported by ^{131}I . Successive evaluation of
 the amount of least squares to the counter data resulted
 in the resolution of a residual line component. (Fig. 4)
 Half life determinations were also made from scaling
 plots of counting rate vs time obtained using standards
 of specific thickness with the same window & counter. The
 method of curve resolution previously outlined was
 employed and a total of 3 residual line components were
 resolved having half lives of 15.7 days, 15.7 days, and 15.7
 days.

Figure 4

COUNTING RATE vs TIME
 data taken with 4π Counter
 SHOWING HALF LIFE VALUES DETERMINED
 BY METHOD OF LEAST SQUARES

• Corrected Observed Counting Rate
 * Remainder after Subtracting B

A $T_{1/2} = 88.91 \pm 9.22$ days
 B $T_{1/2} = 17.66 \pm 0.16$ days
 C $T_{1/2} = 70.06 \pm 2.82$ hours
 D $T_{1/2} = 25.81 \pm 0.20$ hours



B. Absolute β Activity

Since the efficiency of the 4π counter for β counting is quite high (Appendix I), these data were used in the determination of absolute β activities. The zero time activities obtained in applying the method of least squares to half-life determination were corrected to the time of completion of bombardment. These results can be specified in terms of yield if specific values of deuteron beam current and arsenic separation efficiency are assumed.

Counting rate (Counts per minute)

C. Maximum β Energies

$$E_{\text{max}} = 0.70 \text{ MeV} = 3.27 \text{ MeV}$$

These values were found from absorption curves obtained by use of the end window β counter (Appendix III). From measurements of maximum range made at various times the energy of the most energetic β was determined for both the 26 hour and the 17.5 day isotopes. The method is illustrated in Fig. 5 which is applicable to the 26 hour isotope.

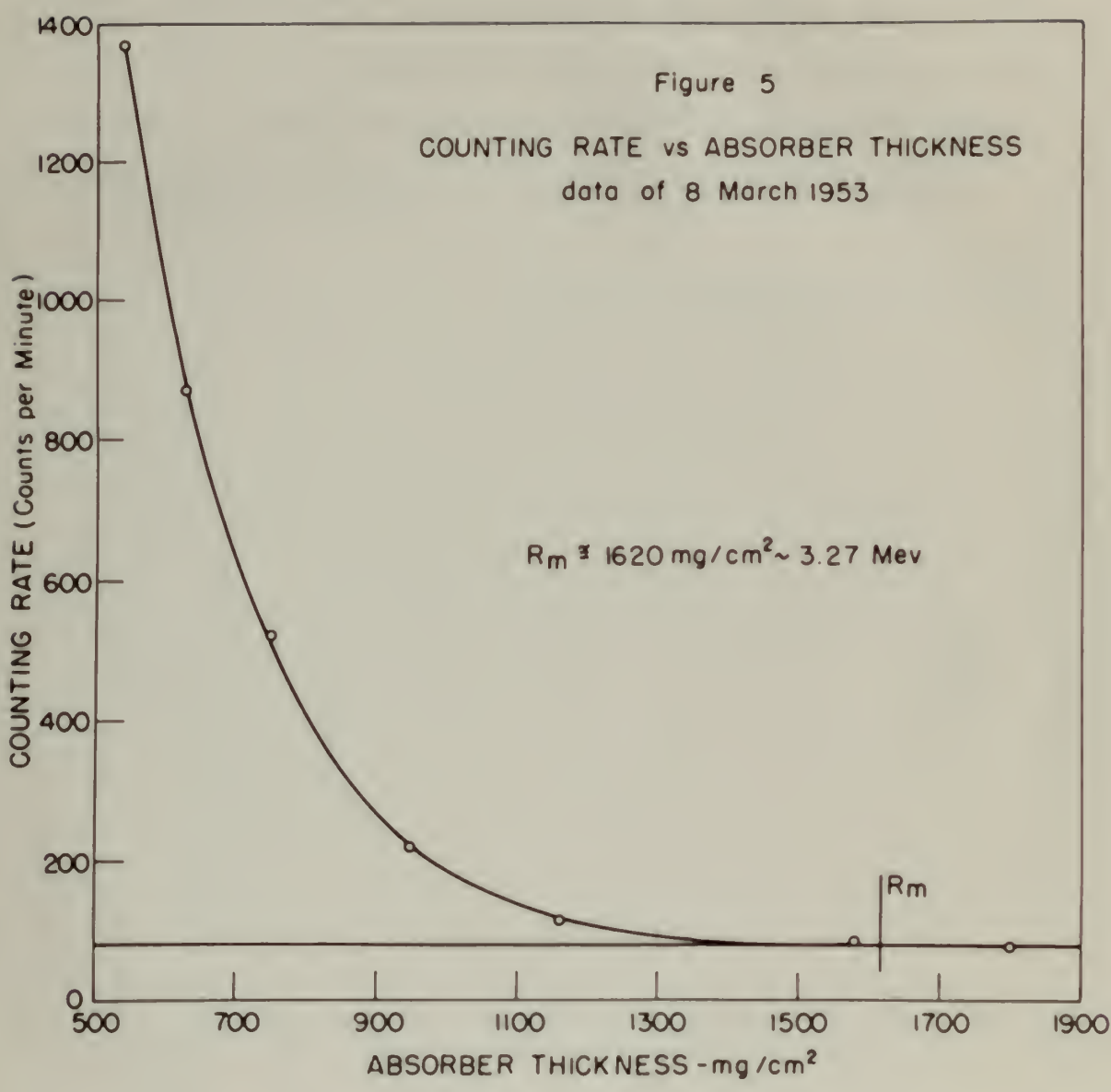
In addition, mass absorption coefficients were determined from semilog plots of counting rate vs absorber thickness taken at various times. Using these values maximum β energies were determined for the 17.5 day and

B. Absolute Activities

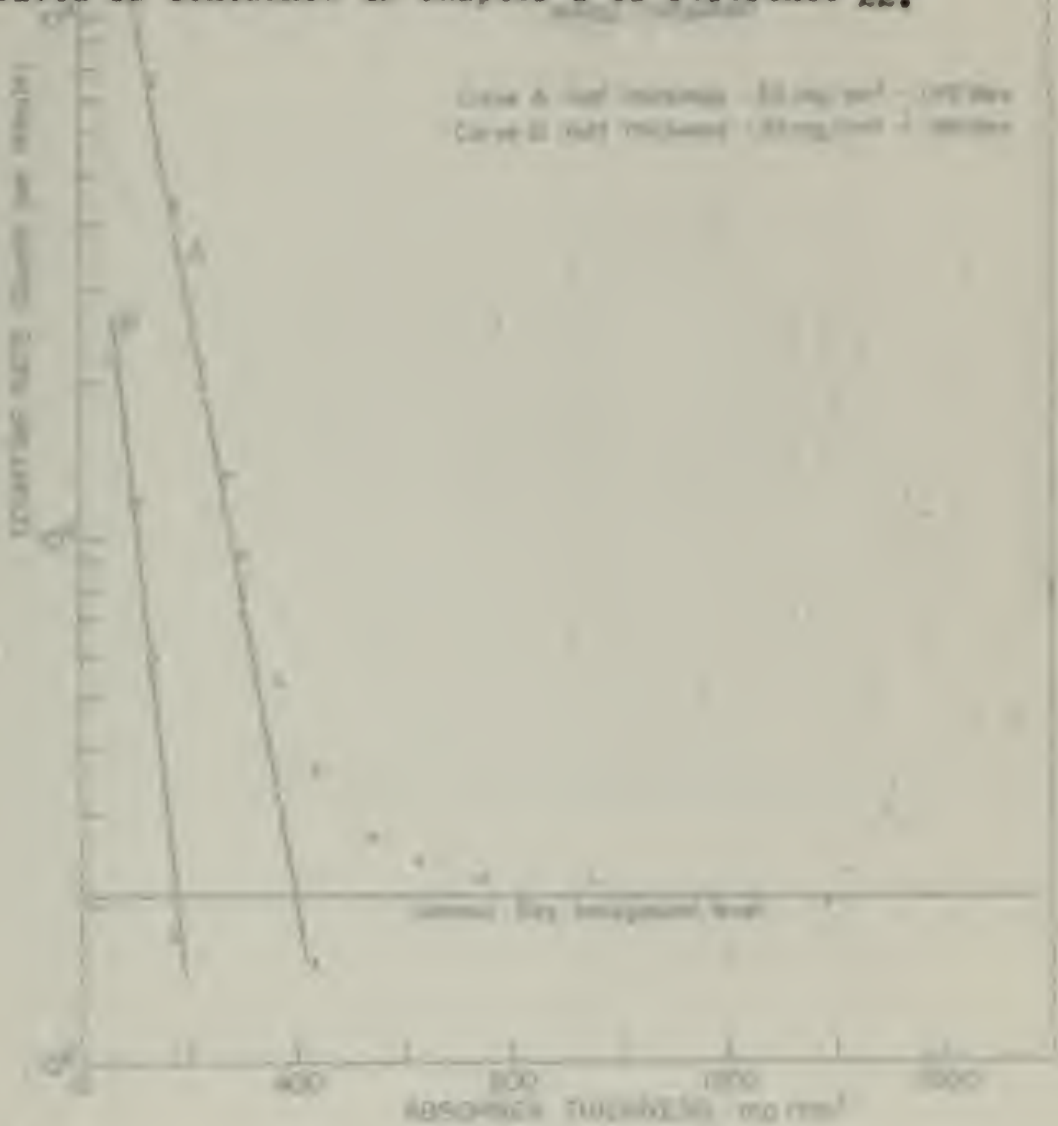
Since the efficiency of the β counter was constant in the range 100-1500 cps, the absolute activities of the samples were determined by comparing the count rate of the sample with that of a standard of known activity. The absolute activities obtained in applying the method of least squares to the life-time determination were corrected to the time of counting at composition. These values are specified in terms of yield of specific values of conversion. The present and previous absolute activities are compared.

C. Maximum Energies

These values were found from the β spectrum curves obtained by use of the end window β counter (Appendix II). From measurements of maximum range made at various times the energy of the most energetic β was determined for both the 20 hour and the 17.5 day samples. The method is illustrated in Fig. 2 which is applicable to the 17.5 day sample. In addition, mass absorption coefficients were used to find from peak heights of counting rate vs absorber thickness curves of various times. Data from these curves and maximum energies were determined for the 17.5 day and



the 59 hour isotopes (Figs. 6, 7). The value obtained by this method for the 17.5 day isotope agrees with that found by the maximum range measurement stated above. The curve obtained for the 26 hour isotope (Fig. 8) was concave toward the origin and could not be treated by this method. A detailed discussion of the method and theory involved is contained in Chapter I of reference 22.



The 10 hour isotherm (Fig. 4, V). The curve obtained by this method for the 10 hour isotherm agrees with that found by the maximum torque measurement method shown. The curve obtained for the 10 hour isotherm (Fig. 5) was compared favorably with the curve obtained by this method. A detailed discussion of the method and theory involved is contained in Chapter I of reference 2.

Figure 6

COUNTING RATE vs ABSORBER THICKNESS
data of 23 March 1953

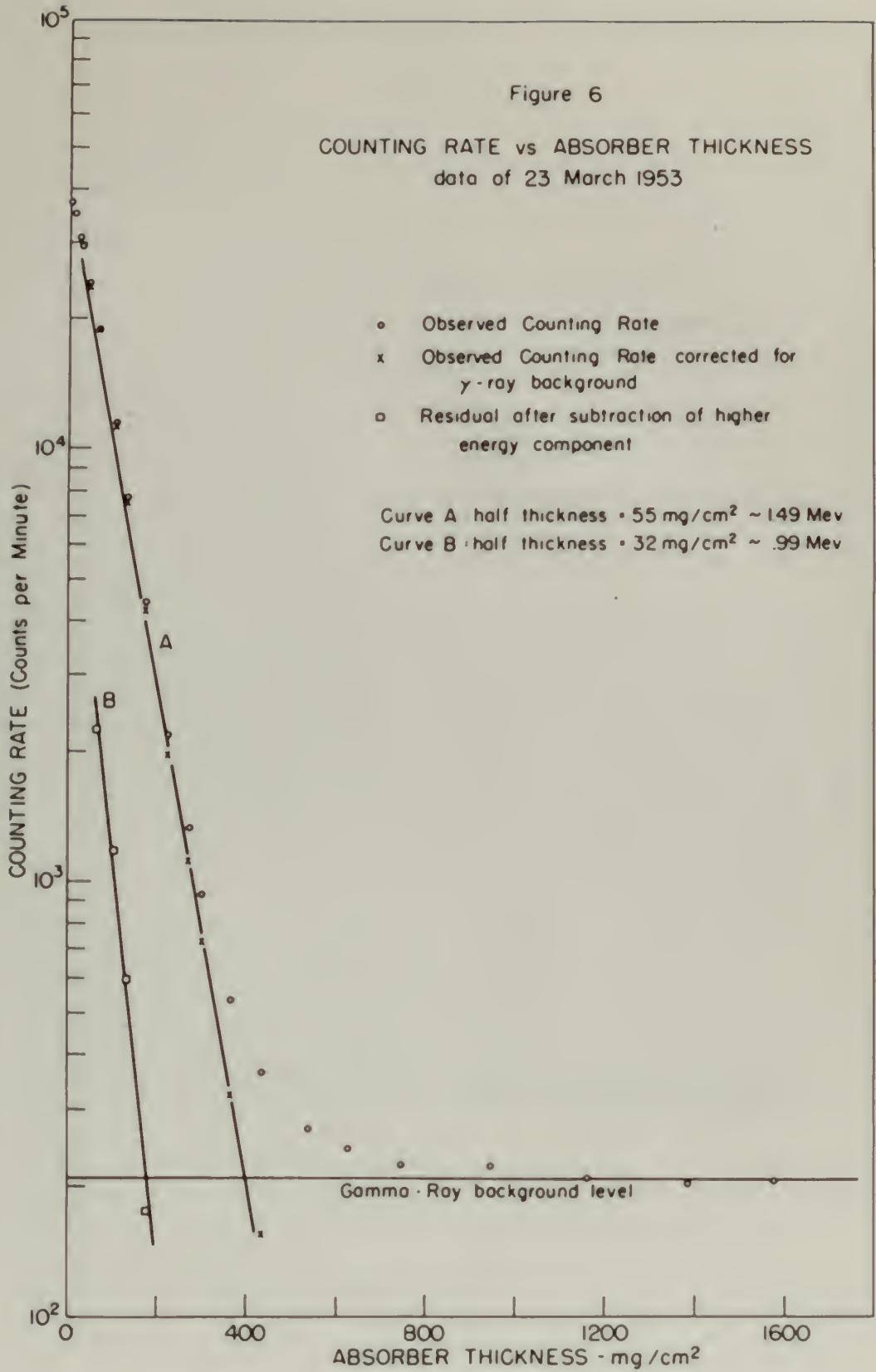
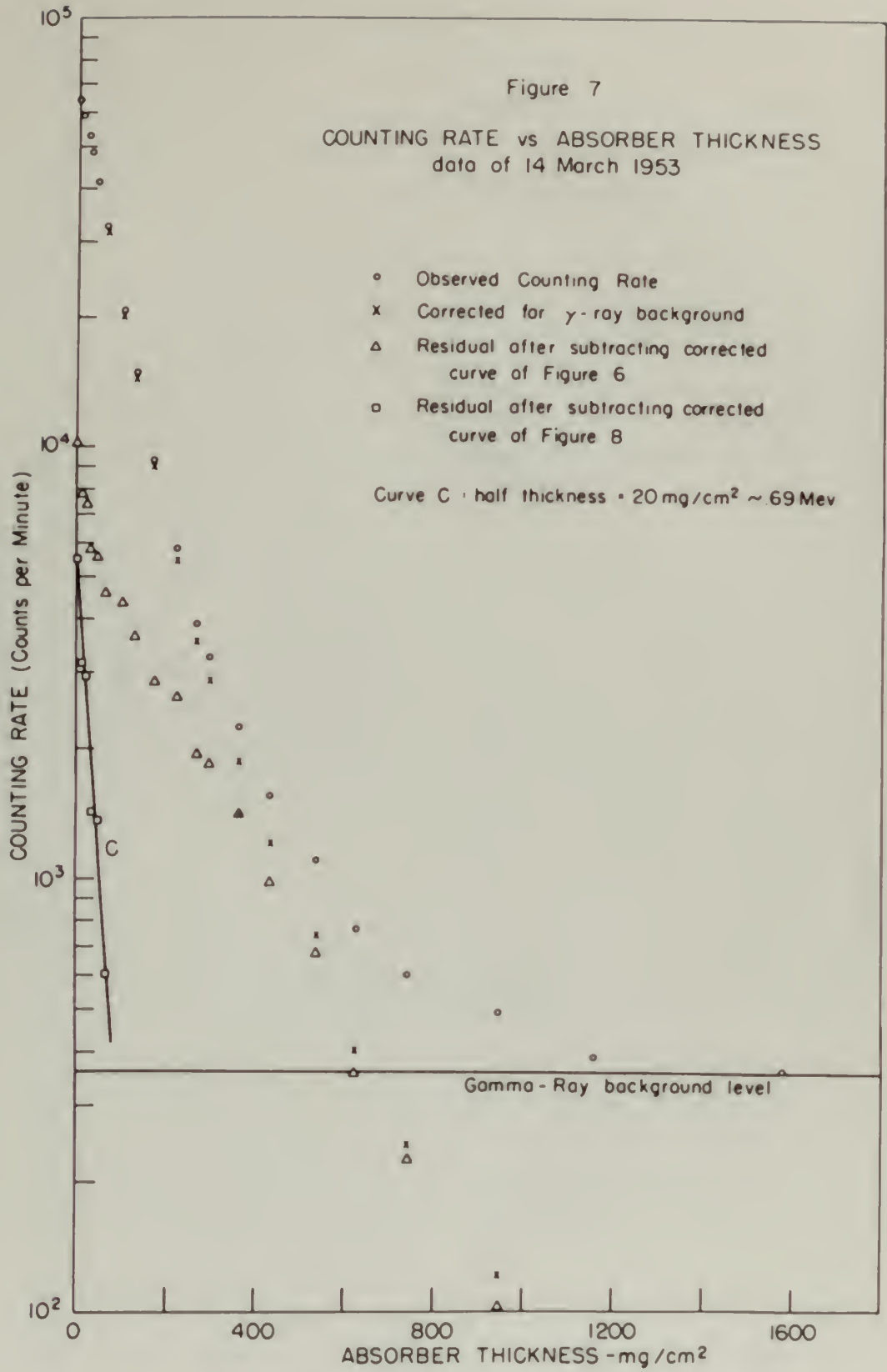
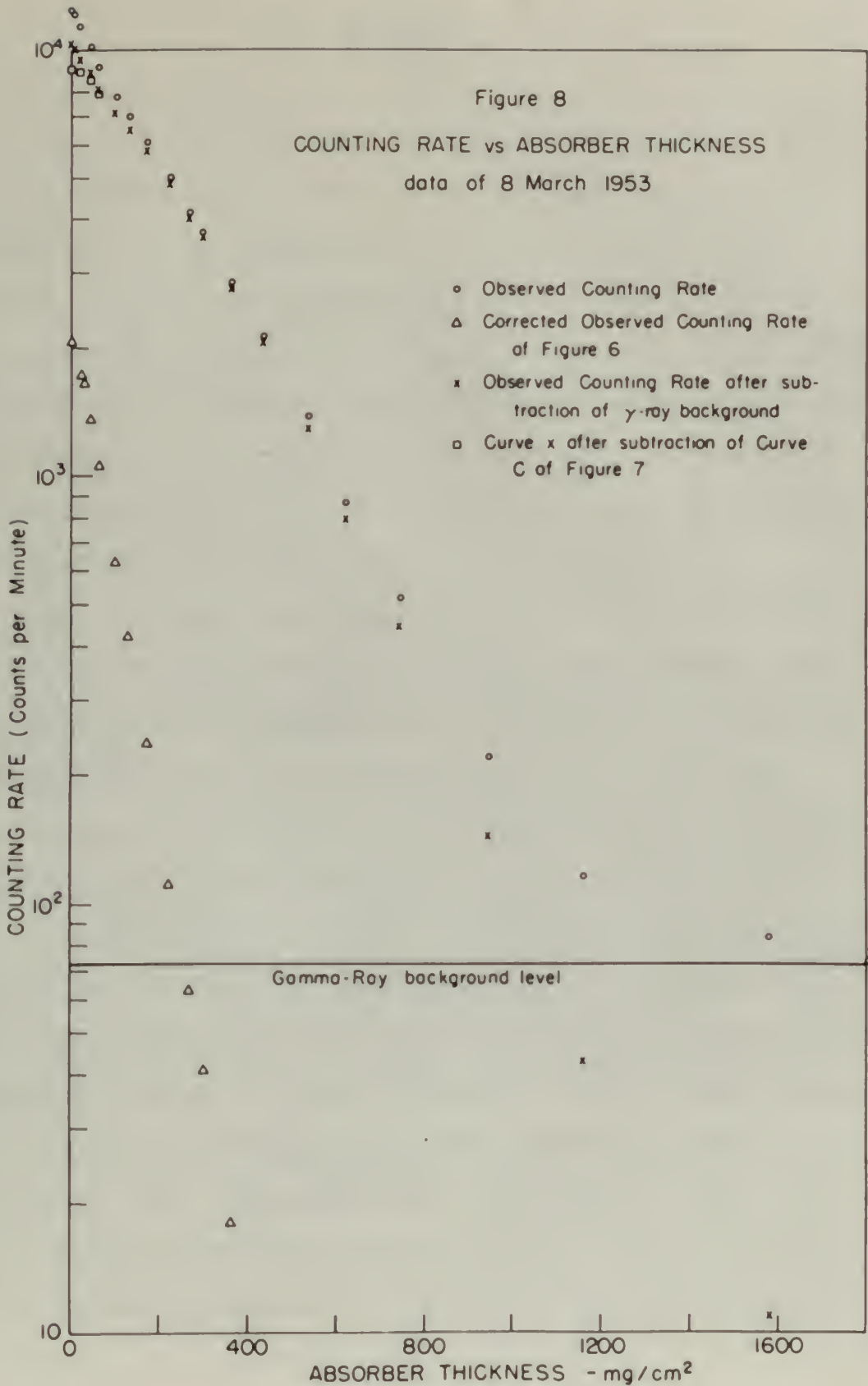


Figure 7

COUNTING RATE vs ABSORBER THICKNESS
data of 14 March 1953





V. RESULTS

1. The longest-lived isotope present in the mixture was detected by only the 4 π counter. The half life was determined to be 82.9 ± 0.2 days and by the filtering method described in Section IV A, the maximum β energy was found to be $0.11 \text{ Mev} > E_{\text{max}} > 0.02 \text{ Mev}$. Since this isotope was not detected with the coincidence counter it is assumed to be a pure negatron emitter. This nuclide is believed to be As⁷³ due to the close agreement with the reported characteristics of that isotope.⁽¹¹⁾

2. With 4 π , coincidence, and end window counters a half life of approximately 17.5 days was resolved. Two β energies of this nuclide were determined by absorption measurements to be 0.99 Mev and 1.49 Mev. The close agreement of these results with those previously reported^(9,11,19) seem to justify the assumption that this isotope is As⁷⁴.

3. End window β counter measurements using an absorber thickness of 224 mg/cm^2 indicated that no isotopes were present having β energies $> 0.7 \text{ Mev}$ and half lives in the range $16.7 \text{ days} > T_{1/2} > 26 \text{ hours}$. Using lesser amounts of absorber a half life of approximately 59 hours was resolved. The only other half lives found to have this approximate value were 70.1 ± 2.8 hours from 4 π counter data and 48.2 ± 1.2 hours from coincidence measurements.

1. The longest-lived species known in the mixture was detected by only the α counter. The half life was determined to be 0.11 ± 0.01 day and by the following method described in Section IV, the maximum energy was found to be 0.11 Mev $< E_{\alpha} < 0.02$ Mev. Since this isotope was not detected with the coincidence counter it is assumed to be a pure alpha emitter. This implies it is believed to be ^{212}Po due to the above agreement with

(ii)

2. With its coincidence, the α window contains a half life of approximately 17.5 days was recorded. The energies of this sample were determined by absorption measurements to be 0.99 Mev and 1.44 Mev. The direct

(er,rr,c)

agreement of these results with those previously reported was to justify the assumption that this isotope is ^{212}Po .

3. The window α counter measurements with an absorber thickness of 0.04 cm² indicated that no isotopes were present having a half life > 0.7 day and half lives in the range 10^{-7} days $< T_{1/2} < 10^{-6}$ days. Using these results of absorber a half life of approximately 30 days was resolved. The only other half lives found to date are approximately 70.1 \pm 1.5 hours from α counter data and 81.1 \pm 1.5 hours from coincidence measurements.

The latter value combined with the energy limitation previously specified justifies identification of this isotope as As^{71} . (10,11,12) The longer half life values obtained from 4π and end window counter measurements indicate that there is also present a neutron emitter having a $T_{1/2}$ longer than 70 hours with energy < 0.7 Mev. The 70 hour half life determined from 4π counter measurements is believed due to a mixture of As^{71} and As^{77} assuming that the single reported value for the half life of the latter (20) is in error. This apparent discrepancy is worthy of future study.

4. From data of the 4π and end window β counters a component of half life 25.8 ± 0.2 hours with a maximum β energy of 3.25 Mev was determined. A γ -ray energy of 0.85 Mev with half life approximately 29 hours was found from measurements made with the γ -ray scintillation spectrometer. These results confirm previously reported values (9,11,19) and identify this isotope as As^{72} . This half life determined from coincidence measurements was 21.7 ± 0.2 hours.

5. There was no indication of the presence of the 52 minute As^{70} isotope (9) in the mixture. In addition, since no γ -ray energies > 0.85 Mev were resolved it was

The latter value compares with the theory prediction
 previously reported for the characteristic of this
 isotope as ^{137}Ba . The largest half life values
 obtained from α and γ window counter measurements
 indicate that there is also present a component which
 yields a $T_{1/2}$ longer than 70 hours. The ratio > 0.7
 Now, the 70 hour half life determined from α window
 measurements is believed to be a mixture of ^{137}Ba and
 ^{137}La assuming that the alpha reported value for the half
 life of the latter ^{137}La is in error. This reported dis-
 crepancy is worthy of future study.

4. From data of the α and γ window counter
 a component of half life 10.5 ± 0.2 hours and a constant
 energy of 3.15 MeV was determined. A γ -ray energy of
 0.115 MeV with half life approximately 70 hours was found
 from measurements made with the γ -ray spectrometer
 spectrometer. These results confirm previously reported
 values ^{137}Ba and ^{137}La and ^{137}Ba . This
 half life determined from coincident measurements was
 21.7 \pm 0.7 hours.

5. There was no indication of the presence of the
 ^{137}La isotope ^{137}La in the mixture. In addition,
 since no γ -ray energy > 0.05 MeV was reported it was

apparent that the 27.6 hour As^{76} isotope having two reported γ energies > 1 Mev (14, 18) was not present.

6. Tabular summary of characteristics of the mixture of radionuclides determined by this investigation.

Isotopes	Method of decay	Energy (Mev)	$T_{1/2}$	Thick target yield* (uc/ μ amp-hr)
As^{71}	β^+	0.66	48.2 ± 1.2 hrs.	7.6
As^{72}	β^+	3.25	25.8 ± 0.2 hrs.	64.9
	γ	0.85		
As^{73}	β^-	$0.11 > E_{max} > 0.02$	88.9 ± 9.2 days	1.1
As^{74}	β^+	0.99, 1.49	17.82 ± 0.13 days	3.2
	β^-			
As^{76}	Not present in the mixture			
As^{77}	β^-	< 0.7	> 70 hours	$5 < \text{yield} < 15^{**}$

* The thick target yield values specified apply if the deuteron beam current was exactly 36 μ amps and if the arsenic separation efficiency was 100 percent. Yield values quoted are based on β counting only and do not include orbital electron capture.

** Based on ratios of total β to β^+ counting rates.

... that the ... χ^2 ... > 1 ...

... of ... of ...

Region	Number of events	Mean	Standard deviation	Mean \pm 1 s.d.	Mean \pm 2 s.d.
IV ₂₁	3*	6.00	1.00	5.00 ± 1.00	4.00 ± 2.00
IV ₂₂	3*	3.00	1.00	2.00 ± 1.00	1.00 ± 2.00
V ₂₁	1*	11.0	0.00	11.0 ± 0.00	11.0 ± 0.00
V ₂₂	3*	1.00	1.00	0.00 ± 1.00	-1.00 ± 2.00
VI ₂₁	3*	0.00	1.00	-1.00 ± 1.00	-2.00 ± 2.00
VI ₂₂	3*	0.00	1.00	-1.00 ± 1.00	-2.00 ± 2.00

* The ... χ^2 ... > 1 ...

... χ^2 ... > 1 ...

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DISCUSSION

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THE CHURCH

APPENDIX I

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11 Co^{60} counting rate vs discriminator voltage
for various counter voltages (electronic
gain constant) 61

18 Calibration curve of discriminator voltage
vs discriminator dial setting 68

A. Description of the Counter

The Co^{60} solid angle counter was designed as a
specialized laboratory instrument to be used for the
precise determination of Co^{60} activity, a photo-
graphic view of the cylindrical counter, Fig. 1,
(Detailed description of operation of counter and
pulse counting. Detailed specifications are
given in Figs. 2 and 3).

The sensitive volume of the counter is geometri-
cally similar to that of a well⁽¹⁾ and of a cylindrical⁽²⁾.
The counter was designed and operated as a flow
counter using a volume gas rather than as a fill
counter and in the fact that the former is more similar
with the latter geometry⁽³⁾. Since the counter
used an active gas that a source is changed, and as a
flow counter which eliminates the necessity for a
vacuum seal greatly simplifies the operating procedure
as compared with that of a fill-type counter. The de-

11	Co ²⁺ working with various organic ligands
12	For various number of ligands (bidentate)
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APPENDIX I
THE 4 π COUNTER

A. Description of the Counter

The 4 π solid angle counter was designed as a convenient laboratory instrument to be used for the absolute standardization of β emitters. A photographic view of the disassembled counter, Fig. 1, illustrates the important features of construction and source mounting. Detailed specifications are given in Figs. 2 and 3.

The sensitive volume of the counter is geometrically similar to that of Caswell⁽¹⁾ and of Borkowski⁽²⁾. The counter was designed and operated as a flow counter using n butane gas rather than as a fill counter due to the fact that the former is more stable with far better reproducibility⁽²⁾. Since the counter must be opened each time a source is changed, use as a flow counter which eliminates the necessity for a vacuum seal greatly simplifies the operating procedure as compared with that of a fill-type counter. The 0-

APPENDIX I

THE ...

A. Description of the Counter

The ... counter was designed as a
 omniscient laboratory instrument to be used for the
 absolute determination of ...
 graphic view of the assembled counter, Fig. 1,
 illustrates the important features of construction
 and source mounting. Detailed modifications are
 given in Figs. 2 and 3.

The sensitive volume of the counter is essentially
 fully similar to that of Caswell⁽¹⁾ and of ...⁽²⁾
 The counter was designed and operated as a thin
 counter using a ...
 counter due to the fact that the former is more suitable
 with the better ...⁽³⁾. Since the counter
 must be opened when the source is changed, and as a
 thin counter which eliminates the necessity for a
 vacuum bell source, standardizes the existing procedure
 as suggested with that of a Mill-type counter. The ...



Fig. 1. Diagram of apparatus for the experiment.

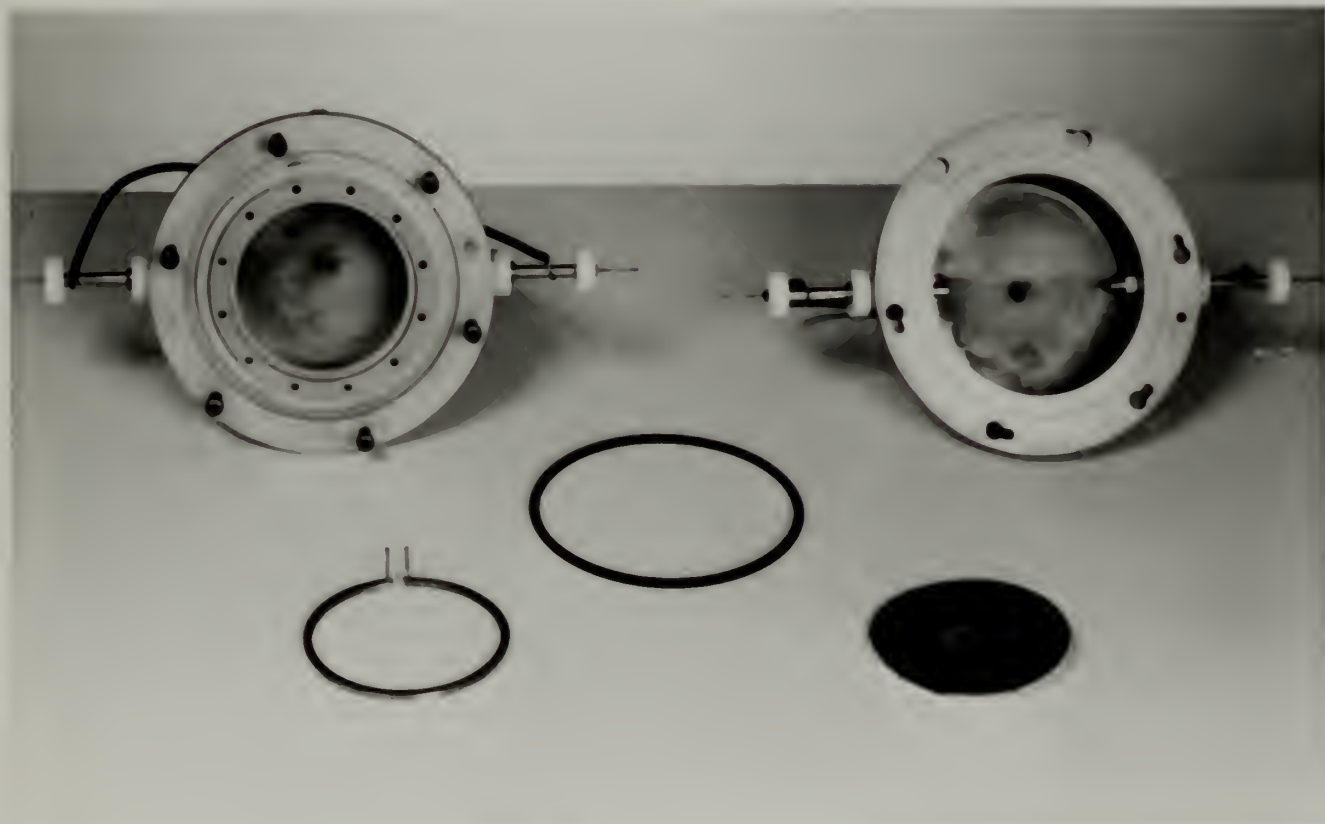
The upper glass shows the source ring in glass
 with the top half of the source covered. The
 water source is the low electrical-lead in the
 center of the thin film electrode as a
 lead ring is the source of the source ring.
 The lower glass shows the source ring in
 electrical contact, the electrical-lead the
 source ring source ring in glass in glass.
 The electrical-lead ring in glass.

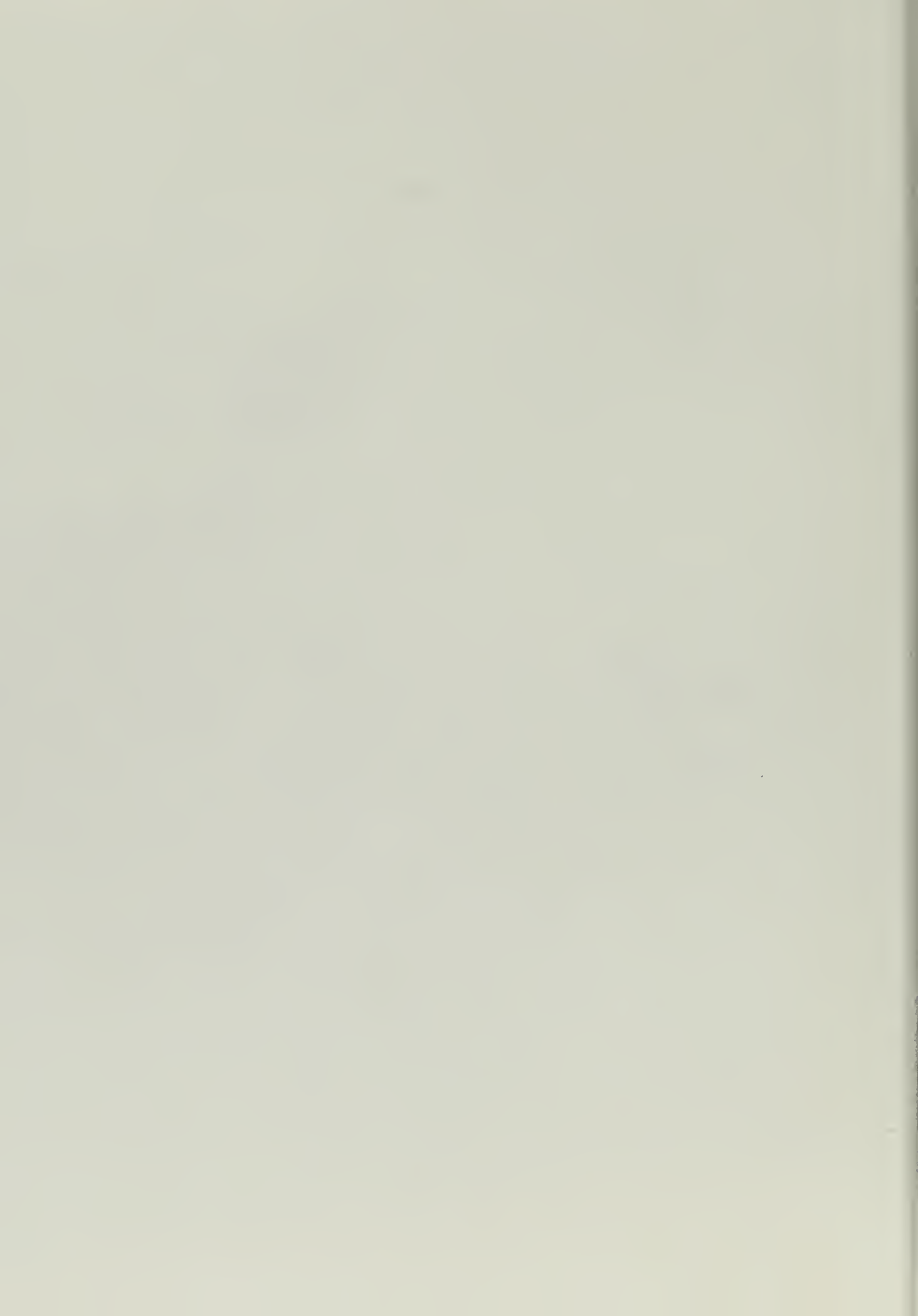


Fig. 1. Photograph of disassembled counter.

The upper view shows the source ring in place with the top half of the counter removed. The active source is the dark circular area in the center of the thin film which appears as a light area in the center of the source ring.

The bottom view shows the completely disassembled counter. The retaining ring for holding the source ring in place is shown with the removable handling pins in place.





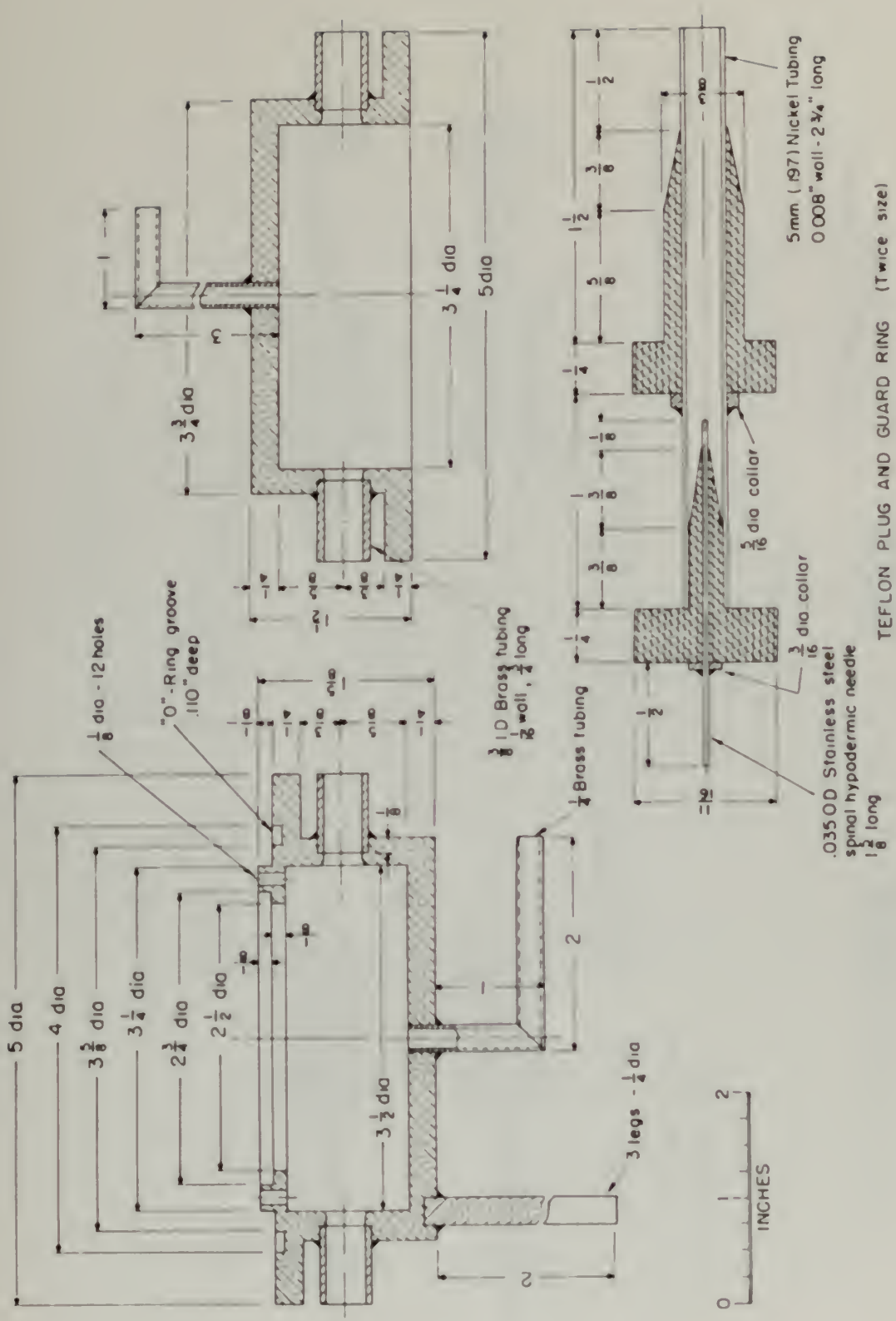
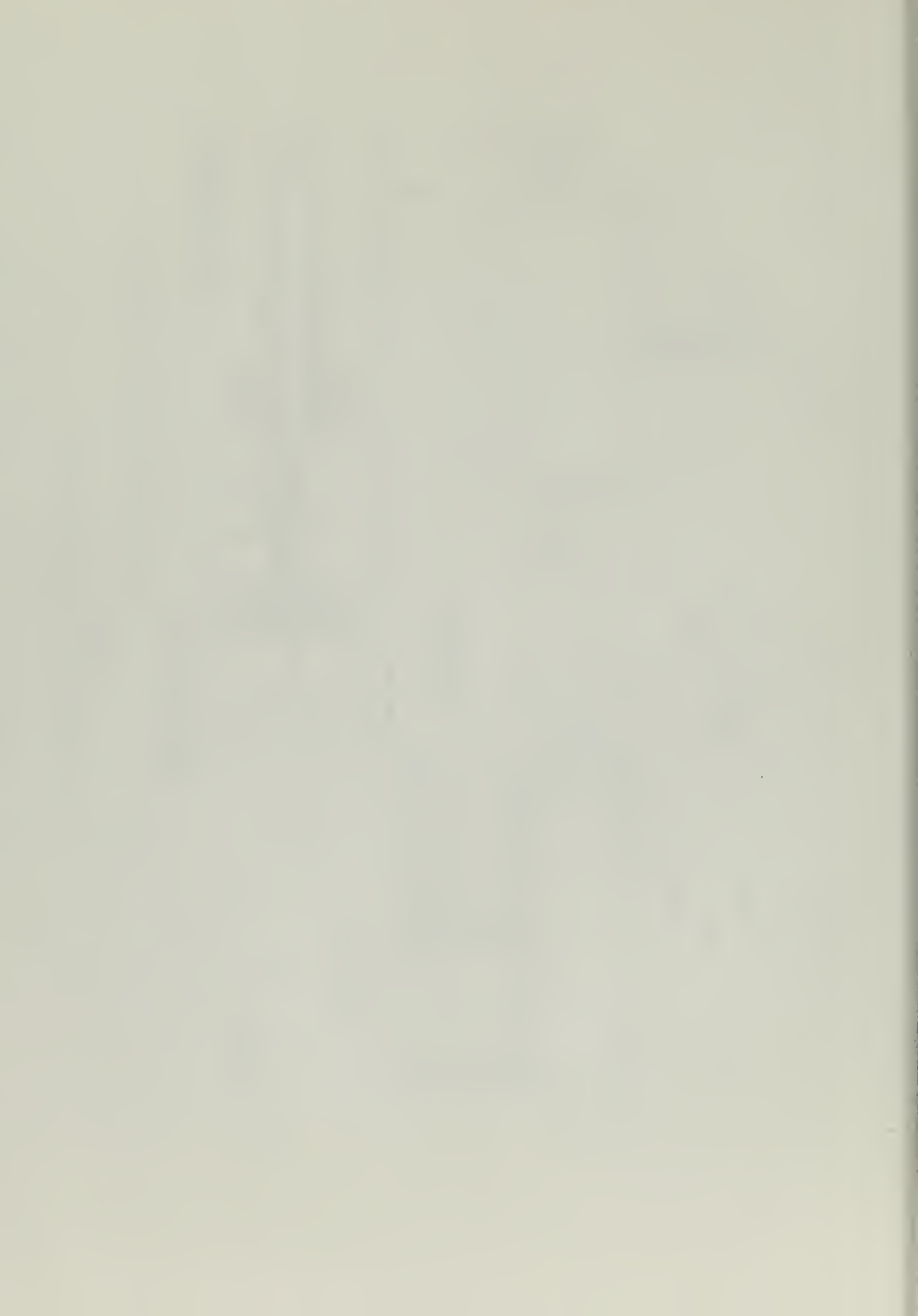


Figure 2
4 77 PROPORTIONAL FLOW COUNTER



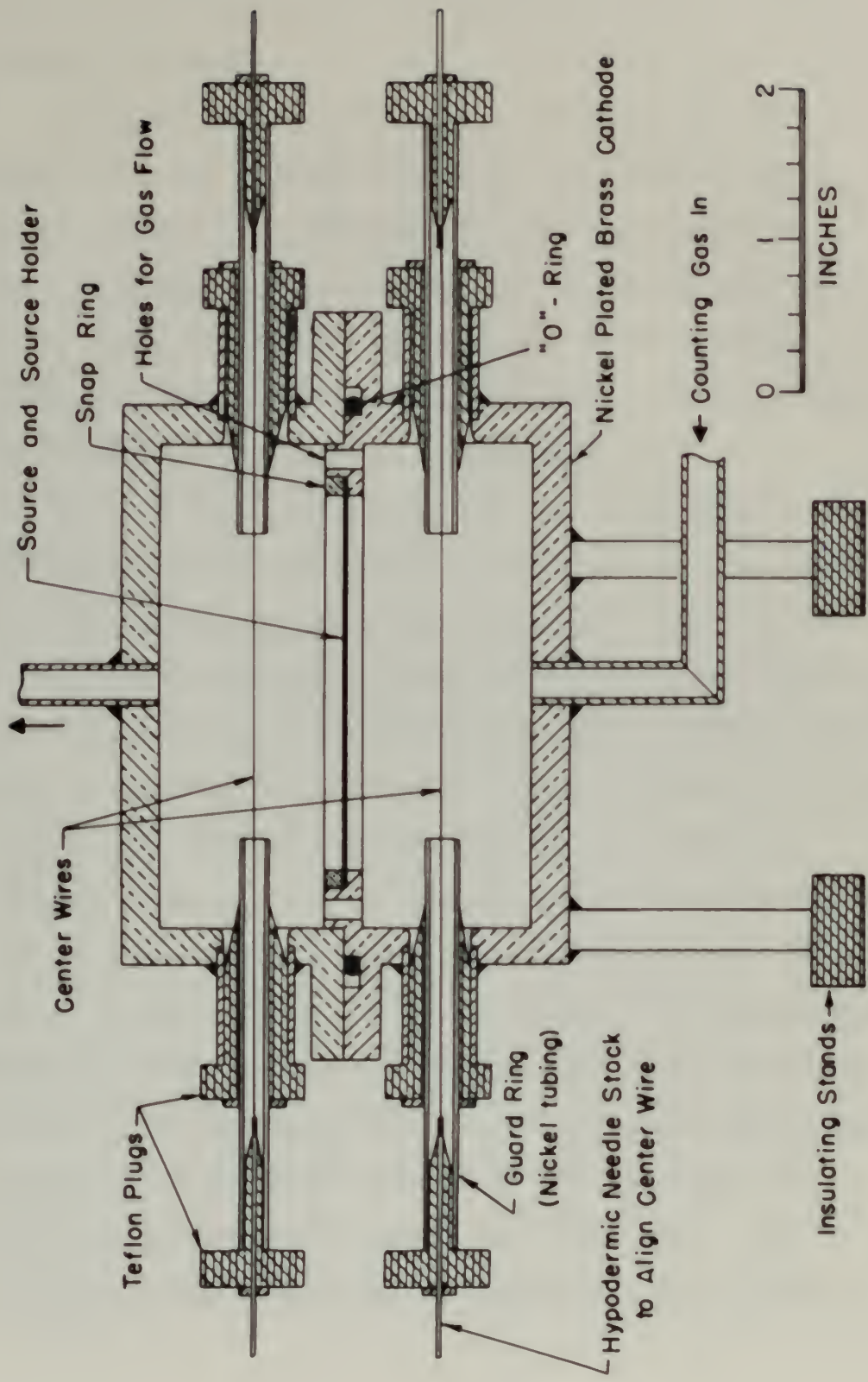


Figure 3
 CROSS SECTION OF 4π PROPORTIONAL FLOW COUNTER

ring seal is used to reduce leakage of flow gas to an absolute minimum.

Squeeze-fitted Teflon plugs are used instead of Kovar glass for counter case-to-guard ring and guard ring-to-center wire insulation, with all insulator surface leakage paths designed for approximately 5000 volts. This feature is for convenience in assembly, cleaning of the sensitive volume, and to minimize the possibility of breakage in handling.

The 1/4 inch thick counter case is machined from 5 inch brass stock, all sleeve entries to the case being silver-soldered and the entire assembly nickel-plated to facilitate cleaning. Center wires of 1 mil tungsten are aligned by 20 gauge hypodermic needle stock to which they are soldered at the extremities. Center wires are guard-ringed with the guard rings at the same high positive potential as the center wires. The tripod legs supporting the counter fit into insulating stands made of drilled polystyrene rod stock. A grounded brass shielding box contains the entire counter assembly and minimizes externally-caused electromagnetic interference. Electrical connections within the box are made with rubber-covered wire insulated for 5000 volts. All electronic connections to the shielding box are made

ring seal is used to reduce leakage of flow gas to an
absolute minimum.

Square-tipped Teflon pins are used instead of
brass pins for counter case-to-guard ring and guard
ring-to-center wire insulation, with all insulating
surface leakage paths designed for approximately 5000
volts. This feature is for convenience in assembly,
easing of the sensitive volume, and to minimize the
possibility of overage in handling.

The 1/8 inch thick counter case is machined from 3
inch brass stock. All sleeve entries to the case being
silver-soldered and the entire assembly nickel-plated
to facilitate cleaning. Center wire of 1 mil diameter
are aligned by 50 gauge hypodermic needle stock so when
they are soldered at the extraction. Center wire are
guard-tinged with the guard ring at the same high
positive potential as the center wire. The tripod
legs supporting the counter fit into insulating stands
made of drilled polystyrene rod stock. A grounded brass
shielding box contains the entire counter assembly and
minimizes externally-caused electrostatic interference.
Electrical connections within the box are made with
rubber-covered wire insulated for 5000 volts. All
electrical connections to the shielding box are made

by coaxial cable and associated fittings. A U-tube oil-filled bubbler external to the shielding box maintains gas pressure within the counter slightly above atmospheric and avoids changes in gas concentration.

Three mil shim steel stock* provides a sturdy source mounting ring. The steel is first cut into 3 inch squares and a 5/8 inch hole is punched in the center of these squares. The 2 3/4 inch outer diameter is then obtained by use of a jig and ordinary paper shears. The prepared source ring is held in place in the counter by use of a split brass ring (similar to a piston ring).

To retain some of the pulse limiting properties of the conventional Geiger counter while eliminating many of its objectionable features, the 4r counter is operated in the region of limited proportionality. (4, 5) The counter is operated at a well regulated 4300 volts with the cathode 2500 volts below ground and center wires and guard rings 1800 volts above ground to minimize corona and "spurty" noise effects. This operating point is approximately 500 volts above the beginning of a counting rate plateau which is better than 0.6 percent per 100 volts. The center wire output is fed

* Obtainable from Ward Steel Co., Arlington, Mass.

by special cables and associated lighting. A 10-amp
self-lit indicator assembly is also available for mounting
the pressure gauge. The counter assembly is shown in
figure 1 and a view thereof is shown in figure 2.
Three air line steel studs, spaced a study
apart, are mounted in the steel plate. The steel plate is
1/2 inch square and a 1/2 inch hole is located in the
center of each square. The 1/2 inch holes are spaced
is the distance of one of a 1/2 inch and center paper
apart. The pressure gauge is held in place in
the counter by use of a split screw cap (similar to a
plate cap).

To obtain some of the more limited results
of the experimental design counter while maintaining
some of its objectionable features, the counter is
operated in the range of 100 to 1000 cps.
The counter is operated at a well regulated 100 cps
with the output 1000 cps with known and counter
vibrations and known 1000 cps with known and counter
vibrations and "spurious" noise effects. This operating
range is approximately 100 cps with some low frequency
of a pulsating rate which is better than 0.5
percent per 100 cps. The counter rate output is 100
cps.

to a Model 100 amplifier through a cathode follower preamplifier (Atomic Instrument Co. Model 204-B) with the amplifier output driving an M.I.T. Model 400-R decade scaler. The counter with associated electronic equipment connected for normal operation is illustrated in Fig. 4.

B. Applicability to Absolute β Counting

If a counter is built which collects particles emitted from a source in all directions, it has many advantages for measurement of absolute activity. First, since all particles emitted from the source are counted, a direct measurement of the β disintegration rate is made without the need for precise knowledge of the solid angle with its accompanying scattering problems. Also, since the "efficiency for β particles" is now 100 percent, much smaller and thinner sources may be prepared thus reducing the self-absorption considerations. (8)

In the 4π counter, any β particle which produces an ion pair outside the source and source mounting will be counted unless this ion pair is formed in a region of low enough intensity that recombination occurs prior to

to a total 100 amplifier stages. A typical follower
amplifier (Radio Instrument Co. Model 100-0) with
the amplifier output driver as M.L.T. Model 100-3
is shown. The counter with associated electronics
equipment associated for total operation is illustrated
in Fig. 4.

B. Applicability to Problems of Counting

If a counter is built with various particles
collected from a source in all directions, it has many
advantages for measurement of radioactive activity. First,
since all particles emitted from the source are counted,
a direct measurement of the λ disintegration rate is
made without the need for previous knowledge of the solid
angle with the corresponding geometrical problems. Also,
since the "efficiency" for a counter is not 100 percent,
each particle not counted makes no contribution to the
reading for self-absorption considerations.⁽¹⁾

In the counter, any λ particle which produces
an ion pair outside the source and source housing will
be counted unless this ion pair is formed in a region of
low enough ionization that recombination occurs prior to

The first thing I noticed when I stepped
out morning in the early
The first thing I noticed when I stepped
out morning in the early
The first thing I noticed when I stepped
out morning in the early

Fig. 4. Photograph of counter with electronic equipment connected for normal operation.

The insulating polystyrene mounts are visible within the brass shielding box and the oil-filled bubbler is shown on the outside lower left corner.



initiation of the Townsend avalanche. Consideration of the geometry of and the fields existing in the sensitive volume indicates a very small probability for counting losses due to this effect. (1)

Any ionization produced by internal conversion electrons, branched spectra, γ -ray spectra, and electrons produced in the counter walls or in the gas will merely add to the total ionization per disintegration and will therefore be counted as a single pulse. This is also true of annihilation radiations and this fact makes the 4π solid angle method valid for the assay of positron emitters.

Deviations from 100 percent absolute efficiency will be due only to (a) absorption in the source and source mounting film, (b) areas of low field intensity mentioned above, and (c) resolving time losses.

C. Important Aspects of Source Preparation

The preparation of a thin source is the most difficult problem involved in the practical use of the 4π counter. It is essential that the source be quite thin and uniform for any isotope emitting soft β particles. The

indication of the downward adjustment, consideration of
the geometry of and the fields existing in the section
values indicate a very small probability for counting
losses due to side effects. (1)

The ionization produced by incident electrons
electrons, produced spectra, very spectra, and electrons
produced in the counter walls as in the gas will merely
add to the total ionization per disintegration and will
therefore be counted as a single pulse. This is also
true of secondary electrons and this fact causes the
to which walls extend walls for the case of position
counters.

Derivations from the present standard efficiency
will be due only to (a) absorption in the source and
source geometry (b) areas of low field intensity
mentioned above, and (c) receiving the losses.

C. Important Aspects of Counter Operation

The propagation of a thin source is the most difficult
problem involved in the practical use of the β counter.
It is essential that the source be kept thin and well-
level for any escape existing with β particles. The

chemistry involved in preparing uniform thin sources varies with the element involved. When a sample is simply allowed to dry, the active material has a tendency to crystallize out as one or more large particles or to dry in a thick ring of small crystals around the edge of the drop. Use of an infrared lamp speeds evaporation and reduces the tendency to "cluster" in every case attempted. It has been empirically determined that counting losses due to self-absorption can be neglected if the maximum solid content of the source is $\leq 5 \mu\text{gm}$ for β energies $\geq 0.6 \text{ Mev}$, but for β energies $\leq 0.4 \text{ Mev}$ solid content of the source should not exceed $0.1 \mu\text{gm}$. These approximate values are based on a total pipetted source volume of 0.085 ml . Within the specified limits, self-absorption losses are negligible compared to losses in the conductive layer on the source mounting film. Self-absorption can never be entirely eliminated by continued reduction of total solids since there is a finite particle size which the material must assume upon precipitation. It has been shown that below a certain very small concentration, a decrease in solids does not increase the observed counting rate. Also, a slight increase in solids above this value does not decrease the observed counting rate of a source.⁽³⁾

The tendency to crystallize out as one or more large particles or to stay in a stable state of small crystals depends on the size of the drops. Use of an electron microscope has shown that the tendency to crystallize is "strong" in every case investigated. It has been experimentally determined that crystallization does not occur in the case of small particles if the volume ratio of the particles is ≤ 0.5 but that it does occur if the volume ratio is ≥ 0.5 . These observations were made on a total particle volume of 0.001 ml. Within the specified limits, self-organized layers are possible compared to layers in the conventional type of surface coating film. Self-organized layers are directly eliminated by continuous reduction of total volume since there is a limit particle size which the material must exceed upon precipitation. It has been shown that below a certain very small concentration, decrease in volume does not increase the observed coating rate. Also, a slight increase in volume above this value does not decrease the observed coating rate of a surface.

(2)

D. Preparation of Source Mounting Film

A solution made by dissolving 5 grams of stick parlodion* in 85 ml of amyl acetate was found to produce the most durable very thin uniform films. A period of about two weeks, with frequent agitation, is required for the formation of the solution.

Thin films are made by dropping an appropriate amount of the above solution on a clean surface of distilled water. The water used should first be boiled to eliminate dissolved gases and an indicator such as phenolphthalein should be added in order to check pH. Water which is even slightly acidic seems to decrease the physical strength and life of the film produced. A room should be chosen which is as dust- and draft-free as possible and a strong light is essential for inspecting the films and the water surface.

The simplest and most expeditious method is as follows:

1. Fill an 8-10 inch diameter culture dish to overflowing with the water prepared as indicated above.
2. Express two drops of parlodion solution on the clean water surface and observe the color display under a strong white light as the film spreads.

* Obtainable from Central Scientific Co., N. Y.

B. Preparation of Glycerol Acetate Film

A solution made by dissolving 5 grams of acetic anhydride in 25 ml of dry acetone was found to produce the most durable film, with uniform thickness. A period of about two weeks, with frequent agitation, is required for the formation of the solution. This time can be made by dropping an appropriate amount of the above solution on a clean surface of distilled water. The water used should first be boiled to eliminate dissolved gases and an indicator such as phenolphthalein should be added in order to check pH. Water which is even slightly acidic seems to decrease the physical strength and life of the film produced. A room should be chosen which is as dark and draft-free as possible and a strong light is essential for inspecting the films and the water surface.

The simplest and most expedient method is as follows:

1. Fill a 5-10 inch diameter container with water overflowing with the water prepared as indicated above.
2. Express two drops of peroxide solution on the clean water surface and observe the color display under a strong white light as the film spreads.

3. When maximum color display is evident near the edges of the film, drop the prepared source ring horizontally from a height of about 1/2 inch onto the center of the floating paraffin film.

4. Holding one edge of the floating source ring and attached film, trim away the excess film with a very sharp knife. The ring is then slid from the surface of the water at a small angle to avoid surface tension film breakage and may be placed vertically in a drying rack.

5. Film thickness may be determined by observation of reflected color under white light and comparison with available curves which read directly in $\mu\text{g}/\text{cm}^2$ (Fig. 5). For more accurate determination, the α thickness gauge may be used. This consists of a collimated source of polonium fastened to a movable micrometer jaw which is mounted vertically above a thin window Geiger counter. A zero reading of the end of the α particle range is made, after which the film is placed over the counter window and the measurements repeated. The distance between the two curves so obtained gives the absorption of the film in air-cm, which can be translated directly into $\mu\text{g}/\text{cm}^2$. The gauge is capable of measuring thicknesses as small as 1 $\mu\text{g}/\text{cm}^2$ with less than 10 percent error.

3. When constant color change is evident near
 the edge of the film, say the purple color ring
 approximately from a width of about 1/4 inch from the
 center of the floating petal-like film.
 4. Holding one edge of the floating source film
 and attached film, turn over the source film with a
 very sharp knife. The ring is torn off from the
 outside of the source at a small angle to avoid contact
 between the two rings and may be placed vertically in
 a drying rack.
 5. This thickness may be determined by observation
 of reflected color under white light and comparison
 with available curves when this thickness is known.
 (See 7). For more accurate determination, the source
 must be very thin. This consists of a collapsed
 source of polymer material in a small diameter jar
 which is mounted vertically above a thin glass plate
 coated with a thin coating of the oil of the particles.
 The plate is held above the film in place over the
 source above and the measurement recorded. The
 distance between the two curves is obtained from the
 absorption of the film in air, when the film
 is held directly over the source. The curve is usually of
 constant thickness as well as 1/4 inch also less
 than in previous cases.

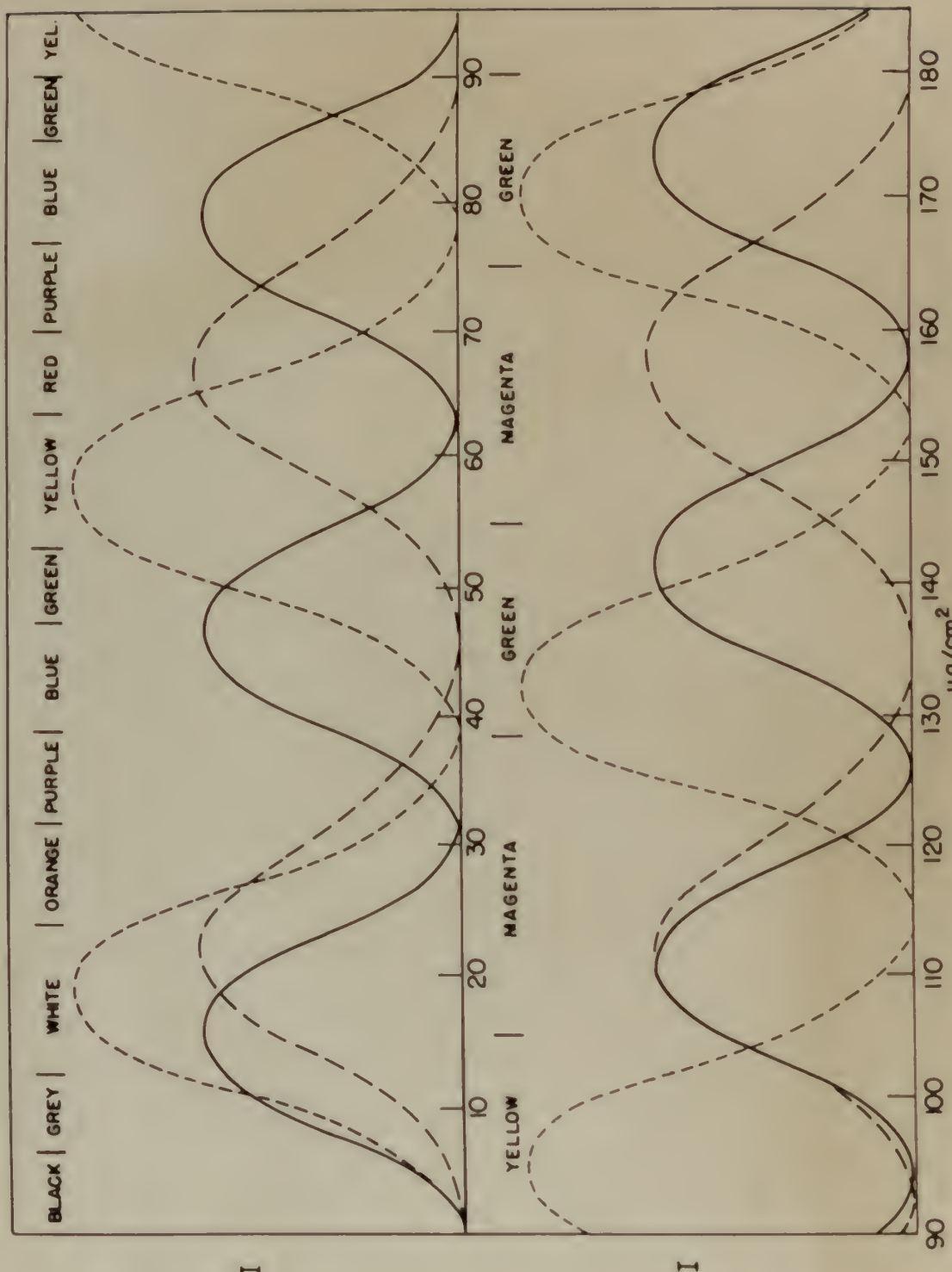


Figure 5 INTENSITY OF REFLECTED LIGHT vs FILM THICKNESS

E. Conducting Layer for Source Mounting Film

For absolute measurements, it is essential that the collecting field within the counter not be distorted by the dielectric-covered hole in the source ring. A thin conducting metallic layer which covers the entire source support may be evaporated from a heated tungsten filament, the evaporation being performed in a vacuum of approximately 1 micron. To insure electrical contact with the counter case, the layer should be deposited on the side of the source ring opposite that to which the parlodion film adheres. The apparatus used for metallic evaporation is illustrated in Fig. 6 where the method of supporting the source ring described below is clearly visible. Since the greatest danger of film breakage occurs in the metallic evaporation process, this operation should be performed prior to pipetting the active source material.

For source solutions which do not contain hydrochloric acid, aluminum produces a suitable conducting layer and the following procedure is recommended.

1. Place the prepared source ring horizontally atop a length of 50 mm diameter glass or pyrex tubing which encloses the prepared tungsten filament. The

For certain circumstances, it is essential that the venting pipe shall not be affected by the elastic-covered pipe in the same way. A thin conducting metallic layer which covers the entire surface should not be separated from a metal support. However, the expansion joint movement in a system of venting pipes is small. The layer should not separate with the venting pipe, but it should be separated on the side of the venting pipe possibly due to the expansion of the pipe. The expansion of the metallic expansion is illustrated in Fig. 2 where the method of expanding the venting pipe is shown. It is clearly visible that the greatest expansion of the venting pipe is the metallic expansion. This expansion should be performed prior to setting the entire sewer system.

The same technique should be used for the expansion of the venting pipe. The expansion of the venting pipe is shown in Fig. 3 where the expansion of the venting pipe is shown. The expansion of the venting pipe is shown in Fig. 3 where the expansion of the venting pipe is shown.

1. The expansion of the venting pipe is shown in Fig. 3 where the expansion of the venting pipe is shown. The expansion of the venting pipe is shown in Fig. 3 where the expansion of the venting pipe is shown.

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Fig. 6. Photograph of apparatus used in metallic evaporation.

The source ring lying on top of the cylindrical glass tubing under the bell jar is in proper position for aluminum evaporation. Tubing which carries cooling water for the filament electrodes is visible to the left of the bell jar.



source ring should be approximately 7 cm above the filament for a vacuum of 1 μ .

2. When a vacuum of 1 μ is reached, slowly increase filament current until the aluminum begins to evaporate from the filament. Observe the climb of deposited aluminum on the glass tube and when it reaches the top of the tube, shut off filament current. This procedure results in uniform conducting layers of from approximately 10 to 15 $\mu\text{g}/\text{cm}^2$ in thickness.

The hydrochloric acid in many source solutions will interact with the aluminum surface of the source mounting and frequently causes a decrease in counting efficiency. In such cases a thin layer (15-20 $\mu\text{g}/\text{cm}^2$) of gold produces a suitable conducting surface. A standard microscope slide placed at the same vertical distance above the evaporating filament as the source mounting film and coated simultaneously with the source film provides a measurement of the thickness of gold. A resistance across the length of the slide measuring between 50 and 100 megohms indicates a thickness of gold between 15 and 20 $\mu\text{g}/\text{cm}^2$. (3)

For the measurement of isotopes having β energies greater than 1 Mev, aluminum foil of 0.1 mil thickness*

* Obtainable from Frank H. Caffin and Son, 52 Elm St., Hyde Park, Mass.

source that should be approximately 7 cm above the
 filament for a vacuum of 1.4.
 3. When a vacuum of 1.4 is reached, slowly
 increase filament current until the aluminum begins
 to evaporate from the filament. Observe the film
 of deposited aluminum on the glass tube and when it
 reaches the top of the tube, shut off filament current.
 This procedure results in uniform evaporating layers of
 from approximately 10 to 15 μm^2 in thickness.
 The photochemical cell is now ready for use.
 will interact with the aluminum surface of the source
 mounted and frequently causes a decrease in counting
 efficiency. In such cases a thin layer (10-15 μm^2)
 of gold produces a suitable conducting surface. A
 standard microscope slide placed at the same vertical
 distance above the evaporating filament as the source
 mounting film and coated simultaneously with the source
 film provides a measurement of the thickness of gold.
 A test tube across the length of the slide measuring
 between 50 and 100 mm indicates a thickness of
 gold between 10 and 20 μm^2 . (3)
 For the measurement of isotopes having a specific
 greater than 100, aluminum foil of 0.1 μm^2 thickness*

* Obtained from Frank J. Gallo and Son, St. Louis, Mo.
 Type 700, Heraeus.

may be used instead of the evaporated metallic layer with no detectable counting loss. A fine mist of distilled water is deposited on the prepared source ring by use of an ordinary bulb type atomizer. The 0.1 mil foil is then laid over the moistened source ring, carefully brushed flat with a fine camel's hair brush, and the excess trimmed off with scissors. If the above is carefully performed the foil is then inseparable from the source ring and parlodion film.

No conducting layer need be applied to the source mounting if a high degree of accuracy is not required. Elimination of the metallic layer results in counting losses of from approximately 1 percent to 3 percent depending on the maximum β energy of the isotope used. For example, the observed counting rate from several non-conducting P^{32} sources increased by ± 0.5 percent with the addition of either foil or evaporated aluminum conducting coatings.

F. Preparation and Precipitation of Source Material

1. Isotopes emitting β particles of ≥ 0.6 Mev.

The best method found so far consists of adding

may be used instead of the unhardened metallic layer
with an adequate porous layer. A fine wire of
dissolved metal is deposited on the porous surface
and in part of an alloy with the element. The
0.1 mil foil is then laid over the metallic surface
and carefully bonded with a thin adhesive
film layer, and the upper surface cut with rollers.
If the above is carefully followed the foil is then
separated from the porous wire and position film.
The embossed layer may be applied in the same
manner as a thin layer of aluminum is not required.
Characteristics of the metallic layer results in coating
less of the porous layer a weight of 3 percent
depending on the surface energy of the lattice grid.
For example, the porous coating rate from several
microns to 20 microns measured by 0.5 percent
with the addition of silver foil or evaporated aluminum
conducting particles.

1. Preparation and investigation of porous material

2. Preparation and investigation of porous material

3. Preparation and investigation of porous material

4. Preparation and investigation of porous material

The above method is for the purpose of making

a small amount of Bentonite, a colloidal mud, to the pipetted drops of source solution. The source ions are adsorbed on the Bentonite which dries in a fairly uniform layer of fine particles. Microscopic observation of sources prepared in this manner yields a typical size of 1 micron for the largest particles, i.e., 0.1 mg/cm^2 for material of density 1.⁽¹⁾ The layer is much more uniform if instead of simply allowing the source to dry, an infrared lamp is used to decrease evaporation time.

Using the highest specific activity source material available to minimize source solid content, a solution of from 1 to $1.5 \text{ } \mu\text{e/ml}$ is prepared. This yields approximate counting rates from 55×10^3 to 89.5×10^3 dpm per 25 λ of active material. Since the resolving time of the counter is approximately 40 μsec , this range of activities limits resolving time losses to ≤ 2 percent.

In solutions of materials of high specific activity, considerable losses may be caused by adsorption of the active constituents on the walls of containing vessels and pipettes used in measurement.⁽²⁾ This effect results in a decrease of activity in solution, especially

a small amount of sediment, a colloidal one, in the
pipetted drops of source solution. The source ions
are adsorbed on the particles which arise in a fairly
uniform layer of fine particles. Microscopic exam-
ination of sources prepared in this manner yields a
typical size of 1 micron for the largest particles,
i.e., U_1 approx 10^2 for material of density 1. The
layer is some 1000 units in thickness of about
aligned the source to 5%, as determined by the
to determine approximate size.

Using the kinetic model for active source material
available for active source with constant, a solution
of from 1 to 1.5 units in prepared. This yields ap-
proximate counting rates from 10^2 to 10^3 and
does not give an active material. Since the resulting
size of the source is approximately 1000, this
range of activities limits resulting time factors to
 $\geq 10^2$.

In solution of material of high specific activity,
considerable losses may be caused by absorption of the
active constituents on the walls of contained vessels
and might also be accompanied. This effect
results in a decrease of activity in solution, especially

for carrier-free materials. This loss of activity may be reduced by the addition of inactive isotopes of the same chemical form as carriers prior to preparation of the source solution. For example, a small quantity of KH_2PO_4 is used with solutions of carrier-free P^{32} and KI is used with carrier-free I^{131} . The mass of carrier which may be added is determined by the permissible solid content of the solution but a desirable ratio to make adsorption negligible is approximately 10^6 inactive atoms per active atom.

The pH of the active solution is maintained so as to keep the active atoms in solution. For some isotopes the solution should be acidic while others require a basic solution. A general rule which has few exceptions is to prepare an acidic solution if the active atom is in the cation and a basic solution if it is in the anion. (8) In all solutions, any substance added to adjust the pH must be soluble when combined with the active material in order to prevent precipitation.

The following procedure is recommended for the actual source preparation:

- a. Express 25 μ (0.025 ml) of active solution on the center of the 5/8 inch diameter metallic coated parlodion film. The micropipette should be rinsed twice

for certain types of materials. This type of analysis may be required by the analysis of various features of the same material from an analytical point of view. The analysis of the same material, for example, a small quantity of K_2CO_3 is used with solutions of various types K_2CO_3 and K_2O is used with various types K_2CO_3 . The use of various which may be added is determined by the availability of this content of the solution and a suitable ratio of the various components. It is recommended that the analysis should be carried out.

The use of the active solution is essential to the study of the active state in solution. The use of various the solution should be made with the active state and the active solution. A general rule which may be applied is to prepare an active solution of the active state in the active and a weak solution. It is in the active. In all solutions, any reactions which are subject to the must be carried out with the active material in order to prevent precipitation.

The following procedure is recommended for the active state analysis. The active solution should be prepared in the active state (0.01 M) at active solution on the surface of the 5% lead chloride solution. The active state should be prepared in the active state. The active state should be prepared in the active state.

onto the source ring in order to remove all active material from the pipette. It has been experimentally determined that the following percentages of active material are contained in rinses of the pipette: -PIPETTE

1st rinse -- 3 percent of initial contents

2nd rinse -- 1 percent of initial contents

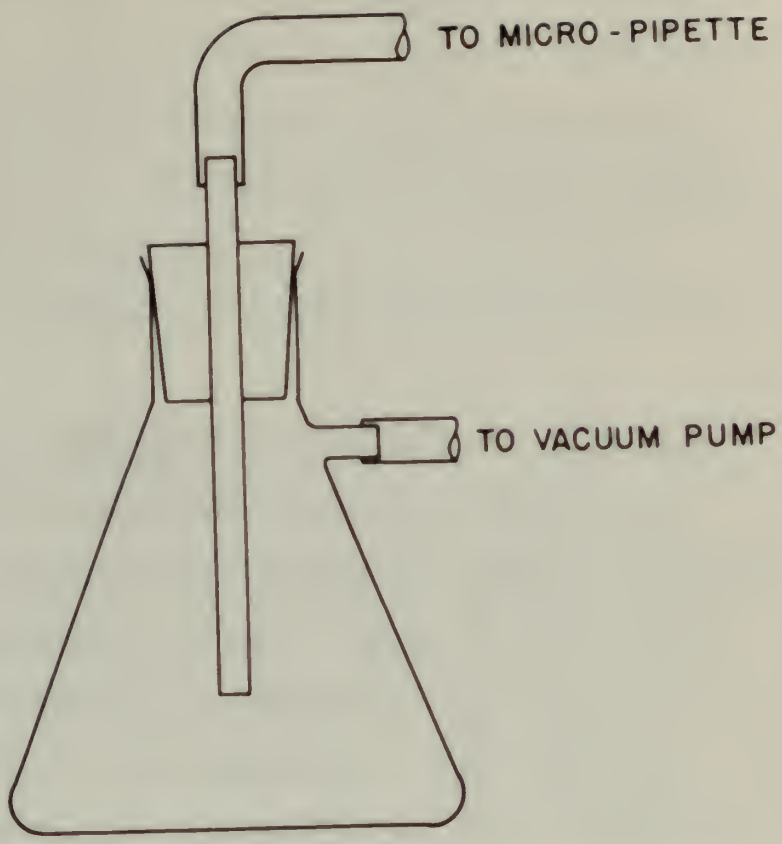
3rd rinse -- 0.5 percent of initial contents

Micropipettes used in source preparation must be calibrated with mercury since deviations of 1 percent from labeled volume are not uncommon. TO VACUUM PUMP

b. If several sources are to be prepared at a time, the vacuum trap arrangement illustrated in Fig. 7 is indispensable. After each source (consisting of one pipette volume plus two rinses) is expressed, the pipette must be thoroughly cleaned and dried prior to preparing the next source. Using the vacuum trap arrangement the pipette can be cleaned with an inactive carrier solution followed by flushing with pure distilled water and then dried by the air stream pulled through the pipette. VACUUM TRAP ARRANGEMENT FOR CLEANING AND DRYING
Figure 7 The total operation of cleaning and drying requires less than 3 minutes with this apparatus.

c. Express 10 μ l of Bentonite solution (approximately 25 mg Bentonite/ml H₂O) into the drop of source

* Obtainable from Radiation Counter Lab., 1814 W. 21st St., Chicago, Ill.



VACUUM TRAP ARRANGEMENT FOR
CLEANING AND DRYING PIPETTES

Figure 7

solution previously formed. Thoroughly disperse the Bentonite in the source solution using an air jet produced from an eye-dropper which has been flame-drawn to a fine capillary point. A strong light facilitates visual observation of the mixing which is complete when the entire drop takes on a cloudy appearance. The prepared source is then dried under a heat lamp.

2. Isotopes emitting β particles of ≤ 0.4 Mev.

In order to minimize self-absorption losses in the measurement of soft β particles extreme care must be taken in preparation of the thin sources, especially if the isotope solution is a chloride which tends to form large crystals on precipitation. The following procedure, applicable to preparation of Co^{60} sources, is cited as an example. (8)

a. Using Co^{60} of high activity (approximately 1 curie/gram), dilute to proper operating range using redistilled HCl. The solid content of ordinary HCl often exceeds the solid content of the source material. The carrier concentration should be of the order of 3 mg of CoCl_2 /liter giving a total solid content in a 25 μ aliquot of 0.075 μ g.

solution previously formed.

6. The weight of the substance in the
same solution using an air jet system from an eye-
dropper which has been fixed-down to a fine capillary
point. A strong light facilitates visual observation
of the rising liquid in capillary when the entire drop
falls on a shiny surface.

7. The weight of the substance in a
small test tube.

8. The weight of the substance in a small test tube.

In order to maintain self-consistency

in the measurement of self-consistency errors are made
to take in preparation of the test tubes, especially
if the test tubes solution is a mixture which tends to
form large crystals on cooling. The following
procedure, applicable to preparation of Co^{2+} solution,
is cited as an example. (8)

9. Weigh Co^{2+} of high purity (approximately
1.0000 gm), which is given equivalent weight being
redissolved in water. The solid content of ordinary HCl
often exceeds the solid content of the more refined.
The purity assessment should be of the order of 1
of $CoCl_2 \cdot 6H_2O$ giving a total solid content in a 250
ml of 0.075 gm.

b. After pipetting the required amount of active solution on the source film, evaporate the Co^{60} to dryness as CoCl_2 in order to get rid of the HCl . Then add a drop of water to the evaporated material to redissolve the CoCl_2 .

c. NH_3 , introduced as NH_4OH in a beaker, should be used to precipitate the cobalt which should cover the entire area of the original water drop quite uniformly.

Steps (b) and (c) above should be done in a desiccator with sodium hydroxide used as a desiccant. A Co^{60} source carefully prepared as outlined above will reduce self-absorption to the minimum value known to be obtainable at this time.

G. Technique Used in Absolute Counting

1. Sources.

Normally three sources are prepared as outlined in Section F from each solution to be counted. Comparison of counting rates of the three sources gives a measure of the precision in source preparation. With a little

practice the difference between sources due to all errors involved in preparation may be maintained at < 1 percent.

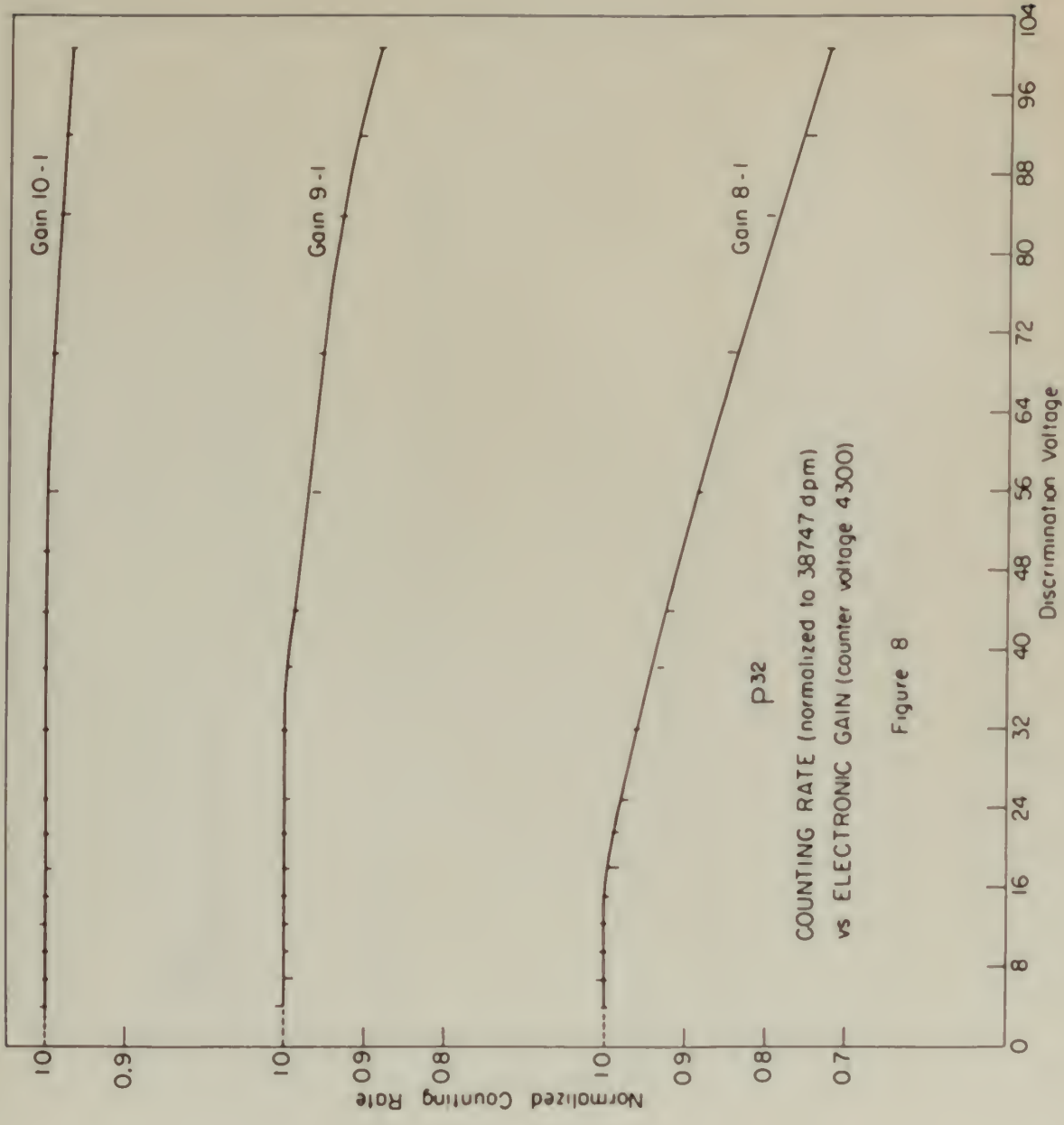
2. Counting procedure.

Background counts are taken before (and after if necessary) each run by inserting a plain shim steel disc in place of the source. Two comparison tests are made on all measurements made with the 4 π counter. First, with counter voltage fixed at 4300 volts, integral discriminator curves are plotted for gain settings of 10:1, 9:1, and 8:1. The latter two settings each decrease electronic gain by a factor of approximately 2 (Figs. 8 and 9). Secondly, with electronic gain held constant at 10:1, integral discriminator curves are plotted for counter voltages of 3900, 4100, and 4300 volts (Figs. 10 and 11). If in both cases the discriminator curves are flat over a discriminator range of \geq 10 volts (Fig. 12), we can assume that all β particles emitted into the sensitive volume are being counted. Figures 9 and 11 which are typical of Co^{60} clearly indicate the high percentage of collection and may be compared with Figs. 8 and 10 which are typical of P^{32} . The method of extrapolation to determine the true

provides the difference between counts due to all
events involved in processing and is maintained at
> 1 percent.

2. Counting procedure.

Background counts are taken before and after
if necessary) each run by inserting a thin lead shield
disc in place of the source. Two comparisons (each are
made on all measurements made with the counter.
First, the counter voltage is set at 500 volts, and
each discriminator counter is plotted for gain settings
of 10:1, 5:1, and 1:1. The latter two settings are
discussed elsewhere in a report of approximately 1950
(Fig. 8 and 9). Usually, with electronic gain held
constant at 10:1, integral discriminator curves are
plotted for counter voltages of 1000, 800, and 600
volts (Fig. 10 and 11). It is felt that the discrimi-
nator curves are flat over a discriminator range of \approx
10 volts (Fig. 11), we can assume that all particles
emitted into the sensitive volume are being counted.
Figures 6 and 11 which are typical of the ^{60}Co clearly indi-
cate the high percentage of collection and may be com-
pared with Fig. 8 and 10 which are typical of ^{226}Ra .
The method of extrapolation to determine the true



p_{32}
 COUNTING RATE (normalized to 38747 dpm)
 vs ELECTRONIC GAIN (counter voltage 4300)

Figure 8

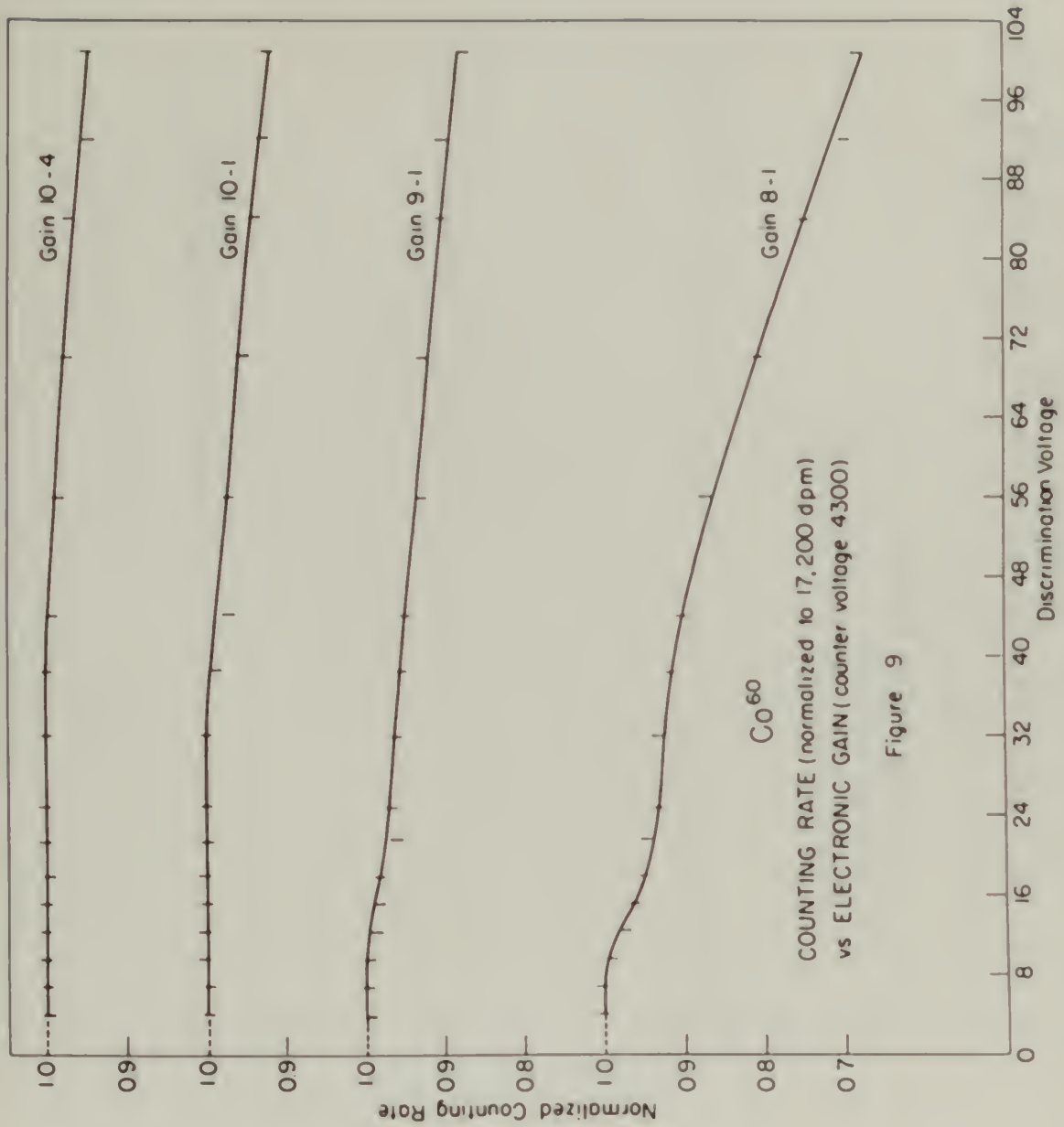
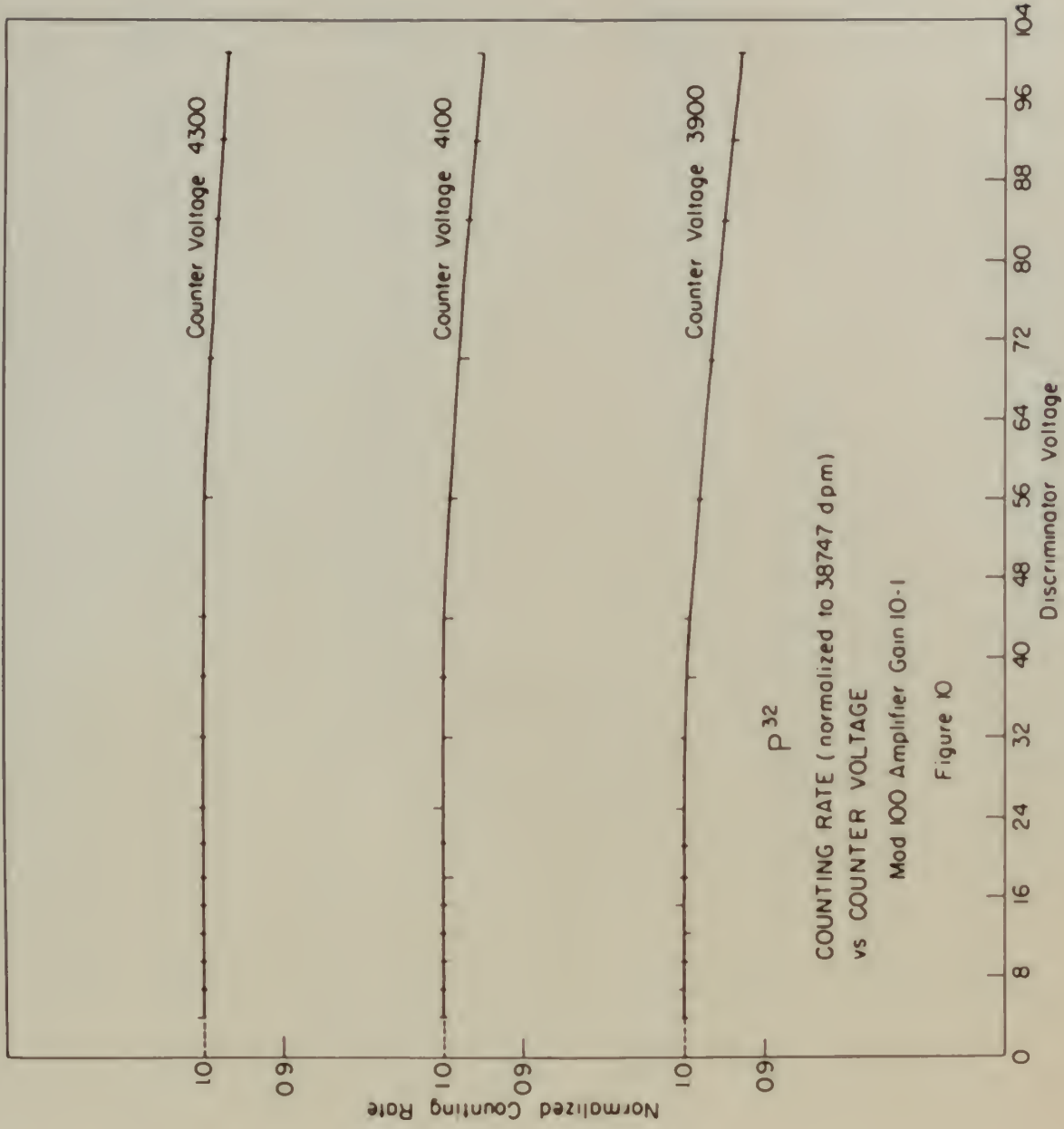


Figure 9



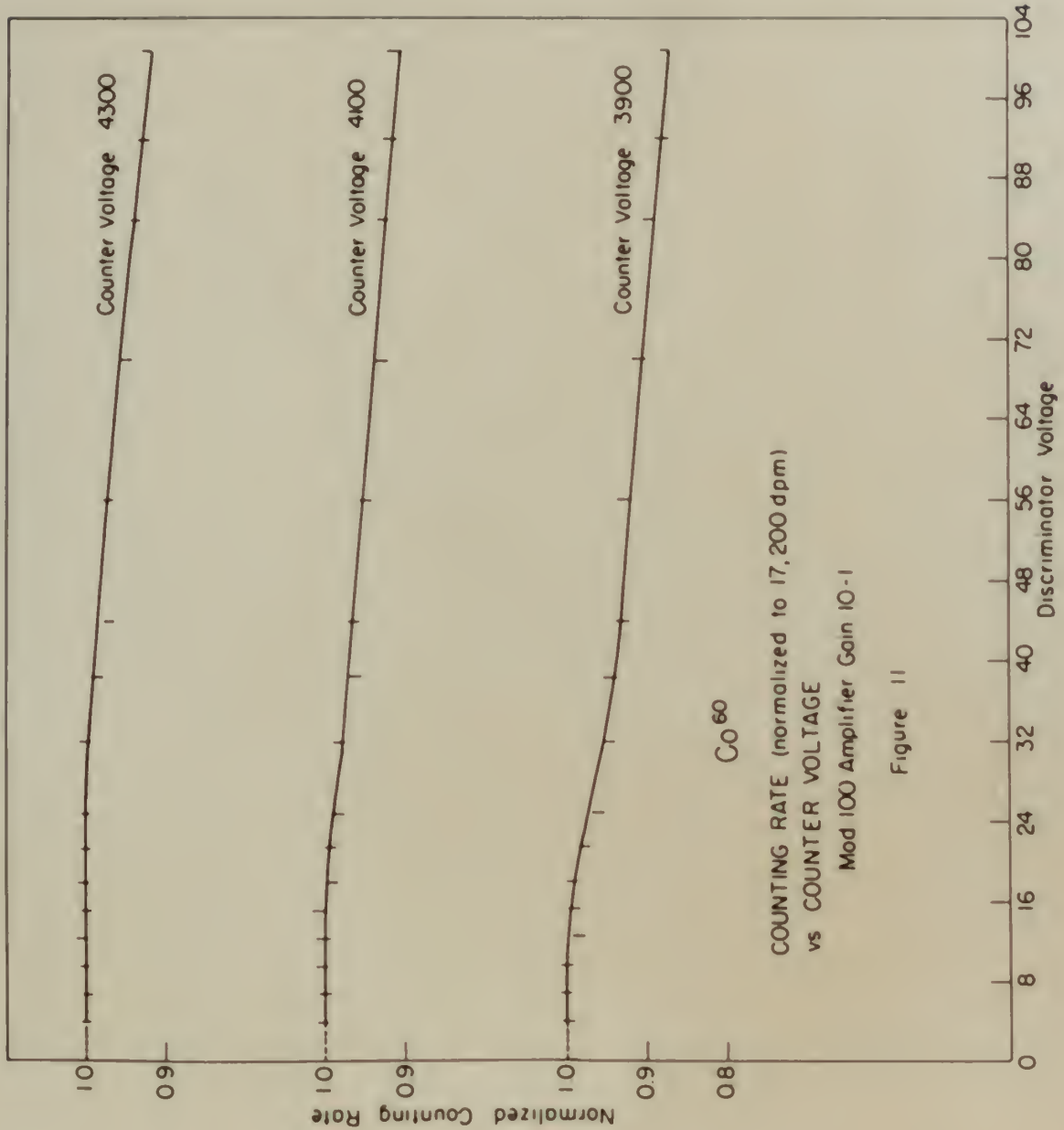
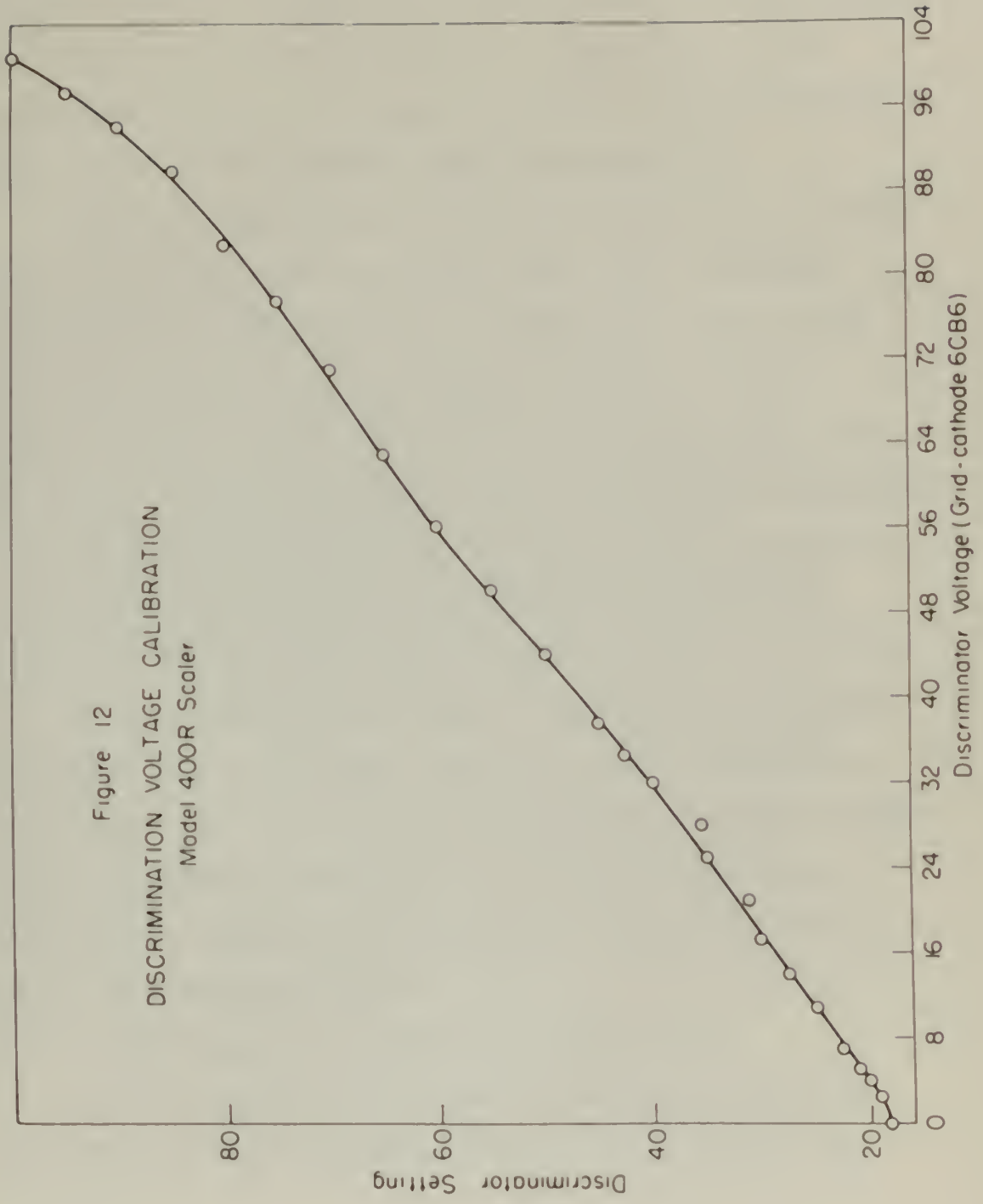


Figure 11

Figure 12

DISCRIMINATION VOLTAGE CALIBRATION
Model 400R Scaler



disintegration rate is indicated on all curves. A further check to verify proper operation is a statistical analysis of the counting rates for points on the discriminator plateau. The mean value thus obtained should agree with the extrapolated true counting rate.

Once the voltage range of the discriminator plateau has been determined for a given isotope the counting procedure is simplified. The discriminator may then be set at the midpoint of the voltage plateau and with counter voltage set at 4900 volts and an electronic gain of 10:1, a series of runs is made. A statistical analysis of these runs is then made to insure that the counter is operating properly and the observed counting rate is determined by the mean value thus obtained.

An electronic gain of 10:1 is chosen as the normal operating point. At this value the largest pulses in the counter just overdrive the Model 100 amplifier without causing counting losses, and the smallest pulses are sufficiently larger than noise to be detectable over a useful discriminator range.

3. Correction to observed counting rate.

The average background is subtracted from the mean value of the measured counting rate, to yield R_0 ,

Investigation into the effect of the
of the counting rate for points on the dis-
the mean value of the counting rate
with the measured counting rate.

One of the main points of the investigation
has been determined for a given length of the counting
procedure is simplified. The distribution may then be
of the midpoint of the voltage pulses and with
counter voltage set at 1000 volts and an electronic gain
of 100, a series of runs is made. A statistical analysis
of these runs is then made to insure that the counter is
operating properly and the observed counting rate is
determined by the mean value from several.

An electronic gain of 100 is chosen as the normal
operating point. At this value the largest pulses in
the counter loss envelope are about 100 millivolts
enough counting losses, and the smallest pulses are
sufficiently large that there is no significant count
rate discrimination.

3. Correction for counting rate loss.
The average background is subtracted from the
mean value of the measured counting rate, to yield R_0

the observed counting rate. For counting rates $\leq 60,000$ dpm, this value may be used as R_t , the true disintegration rate, if an error of 3 percent is acceptable.

Any or all of the following corrections may be made depending on the degree of precision desired:

a. Correction for resolving time losses. If N_o and N_A are defined respectively as observed and actual disintegration rates, then

$$N_A = \frac{N_o}{1 - N_o t}$$

when t is the counter resolving time which has been determined to be approximately 90 μ sec.

b. Correction for absorption due to source counting film. If it is desired to make this correction, then at the same time the source countings are prepared, cover layers of parlodien film of the same thickness should be prepared on source rings having a central hole of 1 3/8 inch diameter. The conducting film evaporated on these cover layers should be of the same thickness as that deposited on the ring containing the active source.

With the active source in place obtain K_{A1} , the actual counting rate. Then remove the source ring,

The general solution is $y = C_1 e^{2x} + C_2 e^{-2x}$. The boundary conditions are $y(0) = 1$ and $y(\pi) = 0$. This gives the system of equations $C_1 + C_2 = 1$ and $C_1 e^{2\pi} + C_2 e^{-2\pi} = 0$. Solving for C_1 and C_2 yields $C_1 = \frac{e^{-2\pi}}{e^{-2\pi} - 1}$ and $C_2 = \frac{1 - e^{-2\pi}}{e^{-2\pi} - 1}$.

$$y = \frac{e^{-2x}}{e^{-2\pi} - 1} + \frac{1 - e^{-2x}}{e^{-2\pi} - 1}$$

where x is the angle in radians. The function is periodic with period π .

2. Solution for $y'' + y = 0$. The general solution is $y = C_1 \cos(x) + C_2 \sin(x)$. The boundary conditions are $y(0) = 1$ and $y(\pi) = 0$. This gives the system of equations $C_1 = 1$ and $C_1 \cos(\pi) + C_2 \sin(\pi) = 0$. Solving for C_1 and C_2 yields $C_1 = 1$ and $C_2 = 0$. Thus, the solution is $y = \cos(x)$.

With the above results in hand, the general solution for the boundary value problem is $y = \cos(x)$.

carefully place the ring containing the cover layer directly over the source to form a sandwich. With this sandwiched source in place, again measure N_{A_2} . The percentage difference in N_{A_1} and N_{A_2} should be quite close to the true correction for absorption due to the mounting film.

A slightly more accurate determination of absorption in the mounting film⁽⁷⁾ is quoted below for completeness.

Experiments have been conducted to determine the amount of absorption, if any, due to the film between the source and the lower half of the 4π counter. The number of particles counted by the top half of the counter connected separately will be

$$N_{\text{top}} = \frac{N_0}{4} [1 + B_F + (1 - \tau) B_V(b)] \quad (1)$$

where N_0 is the true disintegration rate of the source, B_F is the percentage backscattering from the film, τ is the fractional absorption in the film, and $B_V(b)$ is the percentage backscattering due to the walls in the bottom half. The number of particles counted by the bottom half connected separately will be

$$N_{\text{bottom}} = \frac{N_0}{4} [(1 - \tau) + (1 + B_F) B_V(t)] \quad (2)$$

unregularly placed the ring containing the cover layer
 directly over the source to form a secondary shield.
 This secondary shield is placed at a distance $\frac{1}{2}r_0$
 The percentage difference in τ_1 and τ_2 should be
 quite close to the true correction for absorption due
 to the mounting film.

A slightly more accurate determination of absorption
 in the mounting film⁽⁷⁾ is given below for comparison.
 Experiments have been conducted to determine the
 amount of absorption, if any, due to the film between
 the source and the front half of the detector. The
 number of particles counted by the top half of the
 counter exposed separately will be

$$(1) \quad \left[\frac{1}{2} N_0 (1 - \tau) + \frac{1}{2} N_0 \tau \right] e^{-\mu_0 \frac{1}{2} r_0} = \frac{1}{2} N_0 e^{-\mu_0 \frac{1}{2} r_0}$$

where $\frac{1}{2} N_0$ is the true disintegration rate of the source,
 τ is the percentage scattering from the film,
 the treatment described in the film, and μ_0 is the
 percentage beam-absorbing due to the film in the region
 half. The number of particles counted by the bottom
 half exposed separately will be

$$(2) \quad \left[\frac{1}{2} N_0 (1 - \tau) + \frac{1}{2} N_0 \tau \right] e^{-\mu_0 \frac{1}{2} r_0} = \frac{1}{2} N_0 e^{-\mu_0 \frac{1}{2} r_0}$$

From symmetry considerations $B_V(t) = B_V(b) = B_W$.

The factor B_T can be neglected when the film is thin and of low atomic number, so that (1) becomes

$$N_{top} = \frac{N_0}{2}(1 + B_W - \tau B_W) \quad (3)$$

In the bottom half, again assuming $B_T = 0$, one obtains

$$N_{bottom} = \frac{N_0}{2}[(1 - \tau) + B_W] \quad (4)$$

Putting this in the form $y = ax + b$ gives

$$N_{bottom} = \frac{-N_0}{2} \tau + \frac{N_0}{2}(1 + B_W). \quad (5)$$

With the thin films under consideration it can be assumed that the absorption is directly proportional to the film thickness. Equation (5) can be used to determine the absorption correction graphically. A more direct method of determining τ can be deduced from eq. (3) and (4).

$$N_{top} - N_{bottom} = \frac{N_0}{2} \tau (1 - B_W). \quad (6)$$

The actual counting rate observed with top and bottom halves connected together is

$$N_{tb} = N_0(1 - \tau/2) \quad (7)$$

so that (6) becomes, if one lets $N_{top} - N_{bottom} = \Delta$

The velocity constant $k_p(t) = k_p(0) = k_p$
 The factor k_p can be neglected since the film is
 thin and of low atomic number, so that (1) becomes

$$(2) \quad \frac{dN}{dt} = -\lambda N + \frac{N_0}{t} \left(1 - \frac{t}{T} \right)$$

In the bottom half, again assuming $k_p = 0$, one obtains

$$(3) \quad \frac{dN}{dt} = -\lambda N + \frac{N_0}{t} \left(1 + \frac{t}{T} \right)$$

Putting this in the form $\dot{y} = ay + b$ gives

$$(4) \quad \dot{y} + \lambda y = \frac{N_0}{t} \left(1 \pm \frac{t}{T} \right)$$

With the thin film under consideration it can
 be assumed that the absorption is directly proportional
 to the film thickness. Equation (5) can be used to
 determine the absorption correction factor. A
 more direct method of determining λ can be derived
 from eqs. (1) and (4).

$$(5) \quad \frac{dN}{dt} = -\lambda N + \frac{N_0}{t} \left(1 - \frac{t}{T} \right)$$

The actual counting rate observed with thin film
 gives corrected pattern is

$$(6) \quad \frac{dN}{dt} = -\lambda N + \frac{N_0}{t} \left(1 - \frac{t}{T} \right)$$

so that (6) becomes, if one takes $N_{obs} = N$



$$\tau = \frac{N_{top} - N_{bottom}}{N_{tb} - N_b} = \frac{\Delta}{N_{tb} - N_b} \quad (8)$$

and similarly

$$B_V = \frac{N_{top} + N_{bottom}}{N_{tb}} - 1. \quad (9)$$

The absolute counting rate is then obtained by substituting (8) into

$$N_o = \frac{N_{tb}}{1 - \tau/\epsilon}. \quad (10)$$

"Thus by taking three different readings of the same source on a single film it is possible to determine the absorption by the film. This proves extremely useful for low energy β particles."

c. Corrections for self-absorption and back-scattering due to finite source thickness. In general the sources prepared are very thin compared with the half thickness for β absorption in source material. Since the resultant self-absorption and backscattering corrections are small (usually < 1 percent), approximate methods may be used to compute these corrections. The average source thickness is computed from the source area and the known mass of material contained therein.

Self-absorption may be estimated as follows:

Several measurements were made on the film and were corrected

$$(b) \quad \frac{\Delta}{\rho} = \frac{K_{eff} - K_{eff}^0}{K_{eff}^0} = \tau$$

and similarly

$$(c) \quad \lambda = \frac{K_{eff}^0 + K_{eff}}{2} = \lambda_0$$

The absolute counting rate is then obtained by substituting (b) into

equation (c) into

$$(d) \quad \frac{dN}{dt} = N \lambda$$

Thus by using these corrected readings of the count source on a scale that is possible to determine the absorption by the film, this method is very useful for low energy particles.

Correction for self-absorption and scattering in thin source detectors. In general the sources prepared are very thin compared with the self-absorption for a specimen in source material. Since the scattered self-absorption and penetrating corrections are small (usually < 1 percent), special care methods may be used to correct these corrections. The average source thickness is computed from the source area and the gross mass of material contained therein. Self-absorption may be estimated as follows:

Let t = half thickness for β in source material
 (in $\mu\text{g}/\text{cm}^2$)
 \bar{x} = average source thickness (in $\mu\text{g}/\text{cm}^2$)

then the true activity N_t is related to the observed activity N_A by

$$N_A \approx \frac{N_t}{\bar{x}} \int_0^{\bar{x}} \left(\frac{1}{2}\right)^{(\bar{x}/t)} d\bar{x}$$

$$\frac{N_A}{N_t} = \frac{t}{\bar{x}} \left[\frac{1 - \left(\frac{1}{2}\right)^{\bar{x}/t}}{\ln 2} \right] = \frac{1 - \left(\frac{1}{2}\right)^{\bar{x}/t}}{0.693 \frac{\bar{x}}{t}}$$

and $\frac{N_A}{N_t} \approx 1 - (0.346) \bar{x}/t$ for $\bar{x} \ll t$

If desired, backscattering corrections may be approximated from the results published by Zumwalt.⁽⁹⁾ In these corrections it is assumed that when the source mounting material is very thin, the percentage of saturation backscattering obtained is a function only of its thickness in terms of absorption half thickness. With this assumption, the Zumwalt data obtained for polystyrene can be applied to parlodion by comparing relative half thicknesses involved.

There is also a small loss for particles which travel transversely through the film and are absorbed

Let \bar{v} = mean velocity in cm/sec

(in cm³/sec)

\bar{v} = average velocity in cm/sec

Then the true activity a_{\pm} is related to the observed

activity a_{\pm}^o by

$$a_{\pm}^o = \gamma_{\pm} a_{\pm}$$

$$\frac{a_{\pm}^o}{\gamma_{\pm}} = a_{\pm} = \left[\frac{a_{\pm}^o}{\gamma_{\pm}} - 1 \right] \frac{1}{\gamma_{\pm}} = \frac{1}{\gamma_{\pm}^2} \left[a_{\pm}^o - \gamma_{\pm} \right]$$

$$1 \gg \gamma_{\pm} \approx 1 - \frac{1}{2} \left(\frac{1}{\gamma_{\pm}} \right)^2 \approx \frac{1}{2} \left(\frac{1}{\gamma_{\pm}} \right)^2$$

If desired, tabular activity coefficients may be
 calculated from the relative activities of ions.
 In these conditions it is assumed that when the limiting
 activity is very small, the percentage of ion-
 pair formation is negligible and is a function only of
 the distance in terms of ionic radii between ions.
 With this assumption, the limiting data obtained for
 relative half conductance values can be applied to relative half conductance
 values for other electrolytes.
 There is also a well known relative activity
 ratio relationship through the limiting activity

before emerging. An order of magnitude approximation of this loss can be made by consideration of the solid angle within which particles will traverse one half-thickness of the film before emerging. For P³² (assuming a half thickness of about 100 mg/cm²) this gives, for a uniform film 0.05 mg/cm² thick:

$$\delta \Omega \sim \frac{2r \times \frac{0.05}{100}}{4r} \approx \frac{1}{4000}$$

where $\delta \Omega$ = the fraction of emergent particles which traverse a path \geq one half-thickness of the film, which is negligible. For softer β particles, this correction may be large enough to require consideration.

H. Important Characteristics of 4r Counter

Counter voltage plateau: begins at 3700 volts, slope < 0.8 percent per 100 volts.

Settings for normal operation:

Counter voltage: 4300 volts

Cathode: 2500 volts below ground, 600 volts supplied by batteries.

Center wires and guard rings: 1800 volts above ground.

before reaching the order of magnitude approximation
 of this case can be made by consideration of the solid
 angle which these particles will traverse the solid
 thickness of the film before reaching the film surface
 a path thickness of about 100 mμ (for 100 mμ film, for
 a particle film 0.05 mμ thick)

$$\frac{1}{\Omega} \approx \frac{1}{\Omega_0} \left(1 + \frac{1}{\Omega_0} \right) \approx \Omega_0$$

where Ω is the fraction of particles which
 traverse a path \approx one half-thickness of the film, which
 is negligible for a particle film of thickness
 can be large enough to require consideration.

8. Experimental Considerations of the System
 The experimental system consists of a source of
 alpha particles of 5.0 MeV energy, a film of
 thickness 100 mμ, and a detector of 100 mμ thickness.
 The detector is a Geiger counter with a window
 of 100 mμ thickness. The detector is placed
 at a distance of 100 mμ from the film. The
 detector is connected to a counter which is
 connected to a scaler. The scaler is connected
 to a timer. The timer is connected to a
 power supply. The power supply is connected
 to the detector. The detector is connected
 to the counter. The counter is connected
 to the scaler. The scaler is connected
 to the timer. The timer is connected
 to the power supply.

Electronic gain (Model 100 amplifier, Atomic Instrument Co. preamplifier Model 2043):

Coarse gain: 10

Fine gain: 1

Resolving time: $20 \pm 5 \mu\text{sec}$ (determined by a series of measurements by the two-source method)

n-butane flow gas rate: approximately 2 bubbles/sec

Flushing time required for stable operation: 20 min

Length of discriminator plateau (at normal operating settings):

Co⁶⁰ 25 volts

I¹³¹ 30 volts

p³² 50 volts

Counter efficiency:

Co⁶⁰ \approx 87 percent (due to high solid content of available Co activity)

I¹³¹ \geq 98 percent

p³² \geq 99 percent

The following table shows the results of the experiment. The first column shows the number of trials, the second column shows the number of correct responses, and the third column shows the percentage of correct responses. The data is as follows:

Number of trials	Number of correct responses	Percentage of correct responses
10	7	70%
20	14	70%
30	21	70%
40	28	70%
50	35	70%

The results of the experiment show that the percentage of correct responses is constant at 70% regardless of the number of trials. This suggests that the subjects are able to learn the task quickly and maintain a high level of performance.

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The Commission on the Status of Women
and the Commission on Human Rights
are invited to meet in Geneva in 1975
for the purpose of discussing the
draft Declaration on the Elimination
of Discrimination against Women.

II. SUMMARY

The Commission on the Status of Women
and the Commission on Human Rights
are invited to meet in Geneva in 1975
for the purpose of discussing the
draft Declaration on the Elimination
of Discrimination against Women.

DECLARATION ON THE ELIMINATION OF DISCRIMINATION AGAINST WOMEN

Article 1
For the purpose of this Declaration, discrimination against women means any distinction, exclusion or restriction made on the basis of sex which has the effect or purpose of impairing or nullifying the equal rights of women in political, economic, social, cultural, civil or any other field.

Article 2
States should eliminate discrimination against women on all grounds, such as race, language, religion or political opinion.

Article 3
States should eliminate all forms of discrimination against women and should pursue a policy of equality of the sexes.

Article 4
States should take specific measures to protect women against all forms of discrimination and to ensure their equality with men in all spheres of life.

Article 5
States should take measures to modify the attitudes of their population and to eradicate all forms of prejudice against women and against equality of the sexes.

Article 6
States should take measures to ensure that women enjoy equal opportunities with men in the field of employment, including equal pay for equal work.

Article 7
States should take measures to ensure that women enjoy equal opportunities with men in the field of education and training.

Article 8
States should take measures to ensure that women enjoy equal opportunities with men in the field of political participation and representation.

Article 9
States should take measures to ensure that women enjoy equal opportunities with men in the field of economic and social participation.

Article 10
States should take measures to ensure that women enjoy equal opportunities with men in the field of cultural participation.

Article 11
States should take measures to ensure that women enjoy equal opportunities with men in the field of sports and recreation.

Article 12
States should take measures to ensure that women enjoy equal opportunities with men in the field of health and safety.

Article 13
States should take measures to ensure that women enjoy equal opportunities with men in the field of housing and social services.

Article 14
States should take measures to ensure that women enjoy equal opportunities with men in the field of rural development and agriculture.

Article 15
States should take measures to ensure that women enjoy equal opportunities with men in the field of urban development and housing.

Article 16
States should take measures to ensure that women enjoy equal opportunities with men in the field of family and social life.

Article 17
States should take measures to ensure that women enjoy equal opportunities with men in the field of international relations and cooperation.

Article 18
States should take measures to ensure that women enjoy equal opportunities with men in the field of science and technology.

Article 19
States should take measures to ensure that women enjoy equal opportunities with men in the field of culture and arts.

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States should take measures to ensure that women enjoy equal opportunities with men in the field of sports and recreation.

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States should take measures to ensure that women enjoy equal opportunities with men in the field of housing and social services.

Article 41
States should take measures to ensure that women enjoy equal opportunities with men in the field of rural development and agriculture.

Article 42
States should take measures to ensure that women enjoy equal opportunities with men in the field of urban development and housing.

Article 43
States should take measures to ensure that women enjoy equal opportunities with men in the field of family and social life.

Article 44
States should take measures to ensure that women enjoy equal opportunities with men in the field of international relations and cooperation.

Article 45
States should take measures to ensure that women enjoy equal opportunities with men in the field of science and technology.

Article 46
States should take measures to ensure that women enjoy equal opportunities with men in the field of culture and arts.

Article 47
States should take measures to ensure that women enjoy equal opportunities with men in the field of sports and recreation.

Article 48
States should take measures to ensure that women enjoy equal opportunities with men in the field of health and safety.

Article 49
States should take measures to ensure that women enjoy equal opportunities with men in the field of housing and social services.

Article 50
States should take measures to ensure that women enjoy equal opportunities with men in the field of rural development and agriculture.

APPENDIX II

THE SCINTILLATION γ -RAY SPECTROMETER

The counter consists of a sodium iodide thallium activated crystal 4.3 cm in diameter and 5 cm long mounted on an RCA type 6AW photomultiplier tube. The gain of the linear amplifier used is adjusted so that the spectrum is always represented by voltage pulses of from 0 to 100 volts. A single channel differential pulse height analyzer, fed by the linear amplifier, drives a precision counting rate meter of variable time constant and full-scale sensitivity of from 200 to 20,000 counts per minute. The discriminator base line is varied continuously from approximately 0 to 100 volts and a window of 2 volts is used for all observations. Calibration runs were made before and after obtaining each set of data by use of γ -emitters of known energy.

THE EXPERIMENTAL APPARATUS

The apparatus consists of a motor which is driven by a 220-volt AC supply. The motor is connected to a transformer which is connected to a 110-volt AC supply. The transformer is connected to a motor which is connected to a load. The load is connected to a voltmeter which is connected to a galvanometer. The galvanometer is connected to a scale which is graduated in millivolts. The scale is zeroed before the experiment is started. The motor is run at a constant speed and the voltage across the load is measured. The voltage across the load is found to be proportional to the current through the load. The current through the load is found to be proportional to the voltage across the load. The relationship between the voltage across the load and the current through the load is found to be linear. The slope of the line is found to be constant. The slope of the line is found to be independent of the voltage across the load. The slope of the line is found to be independent of the current through the load. The slope of the line is found to be independent of the resistance of the load. The slope of the line is found to be independent of the resistance of the motor. The slope of the line is found to be independent of the resistance of the transformer. The slope of the line is found to be independent of the resistance of the voltmeter. The slope of the line is found to be independent of the resistance of the galvanometer. The slope of the line is found to be independent of the resistance of the scale. The slope of the line is found to be independent of the resistance of the motor. The slope of the line is found to be independent of the resistance of the transformer. The slope of the line is found to be independent of the resistance of the voltmeter. The slope of the line is found to be independent of the resistance of the galvanometer. The slope of the line is found to be independent of the resistance of the scale.

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APPENDIX III

THE END WINDOW β COUNTER

A. Description of Equipment

The tube was operated at the potential of 1000 V. This equipment, used in determining β energies, consists of a conventional end window Geiger-Muller tube* and a set of calibrated aluminum absorbers.** The tube is mounted in a shielded sample changer*** which contains sliding trays for accurate positioning of source and absorber.

B. Experimental Technique

With a source inserted on the lower tray under the Geiger tube counting rates were recorded for various thicknesses of absorber contained on the upper tray.

* Tracerlab, Inc. Model EC-C1 Geiger Tube, window thickness 3 mg/cm².

** Tracerlab, Inc. Type N-3A Calibrated Absorbers.

*** Tracerlab, Inc. Model EC-90 Shielded Manual Sample Changer.

APPENDIX III
THE NEW VISION A COURSE

A. Description of Equipment

This equipment, used in determining a number of constants of a semiconductor and silicon diode-rectifier, consists of a set of calibrated silicon diodes, a voltmeter and a set of calibrated silicon diodes. The tube is mounted in a shielded metal container which contains a liquid crystal for accurate positioning of source and detector.

B. Experimental Techniques

With a source mounted on the inner face of the detector tube, the detector tube was rotated for various distances of detector constant on the outer face.

* Yonkers, Inc. Model 70-10 Silicon Tube, Vision
Vision 2 1/2" x 1 1/2"

** Yonkers, Inc. Type 2-12 Calibrated Diodes.

*** Yonkers, Inc. Model 70-10 Silicon Manual Tube

Chapter

Counting procedure varied depending upon the thickness of absorber. At high counting rates, the number of counts per 1 minute interval was recorded for at least three intervals. At intermediate counting rates, the preset count feature of the scaler was utilized and the time required for 10,000 counts was recorded. At very low counting rates, a preset count of 4000 was used.

The tube was operated at the mid-point of its voltage plateau to insure maximum stability. In addition a set of five standardized β emitters of known energy was periodically counted thus enabling the correction of observed counting rates for any changes in instrument sensitivity.

C. Corrections of Observed Data

Other than correcting for counter sensitivity fluctuations mentioned above, the only correction required was for resolving time losses. These corrections were made by adding to the observed counting rates the number of lost counts per minute (Fig. 2).

Y. S. Kim, Ph.D., Thesis, University of California, Berkeley, 1952, Chapter III.

Counting procedure varied depending upon the thickness of absorber. At high counting rates, the number of counts per 1 minute interval was recorded for at least three intervals. At intermediate counting rates, the present count feature of the scaler was utilized and the time required for 10,000 counts was recorded. At very low counting rates, a preset count of 4000 was used. The tube was operated at the mid-point of its voltage range to insure maximum stability. In addition a set of five standardized sources of known activity was periodically counted to enable the correction of observed counting rates for any changes in instrument sensitivity.

C. Corrections of Observed Rate

Other than correcting for counter sensitivity fluctuations mentioned above, the only correction required was for resolving time losses. These corrections were made by adding to the observed counting rates the number of lost counts per minute (Fig. 2).

D. Resolving Time Losses

It is generally assumed that the resolving time of an instrument is constant for all counting rates and corrections are usually made for resolving time losses by means of equations derived on the basis of two general counter types; the paralyzable and the non-paralyzable. A detailed treatment of these two cases* results in the following equations:

$$n = Ne^{-Np} \quad (\text{paralyzable type}) \quad (1)$$

$$n = N(1 - np) \quad (\text{non-paralyzable type}) \quad (2)$$

At low counting rates both equation (1) and equation (2) reduce to

$$N = n(1 + np) \quad (3)$$

where N and n are respectively the true and observed counting rates and p is the resolving time.

In this experiment it was considered necessary at times to count at very high rates (~ 50,000 cpm) because of the possible presence of short-lived isotopes and the desire to obtain complete sets of absorption data as quickly as possible. It was obvious that the approximate equation (3) could not be used and it was also found that

* Evans, R. D.: Class Notes for Course 8.512, Chapter 30.

D. Resolving Time Losses

It is generally assumed that the resolving time of an instrument is constant for all counting rates and corrections are usually made for resolving time losses by means of equations derived on the basis of two general counter types; the paralyzable and the non-paralyzable. A detailed treatment of these two cases results in the following equations:

$$(1) \quad n = n_0 e^{-2n\tau} \quad (\text{paralyzable type})$$

$$(2) \quad n = n_0 (1 - n\tau) \quad (\text{non-paralyzable type})$$

As for counting rates both equations (1) and equation (2) reduce to

$$(3) \quad n = n_0 (1 + n\tau)$$

where n and n_0 are respectively the true and observed counting rates and τ is the resolving time.

In this experiment it was considered necessary at first to count at very high rates ($\sim 30,000$ cps) because of the possible presence of short-lived isotopes and the desire to obtain complete sets of spectral data as quickly as possible. It was obvious that the approximate equation (3) would not be used and it was also found that

* Evans, G. I.: *Basic Data for Counting*, Chapter 10.

neither equations (1) nor (2) properly corrected the observed counting rates if a constant resolving time was assumed.

To obtain a useful relation between counting rate and resolving time loss, the response of the instrument to a series of standard sources of known activity was measured and a plot made of observed vs expected counting rate (Fig. 1).^{*} Two response curves are shown, one for a discriminator setting of 4, the other for a setting of 6.^{**} From the curve for a discriminator setting of 4, the setting used throughout the experiment, a plot of lost counts per minute vs observed counting rate (Fig. 2) was prepared to facilitate correction of the observed data.

To verify the accuracy of this procedure, several sources were counted with discriminator settings of 4 and 6. The following tabulation of the counting rates observed and the true counting rates computed from the applicable curve of Fig. 1, shows that the computed values agree within experimental error thus indicating that consistent corrections may be made by this method.

* Evans, R. D.: Class Notes for Course 8.512, Chapter 30, page 15.

** Data of E. Samuels, Physics Research Laboratory, Massachusetts General Hospital, Boston, Mass.

Figure 1

RESPONSE OF END WINDOW BETA COUNTER
COUNTING RATE vs STRENGTH OF SOURCE

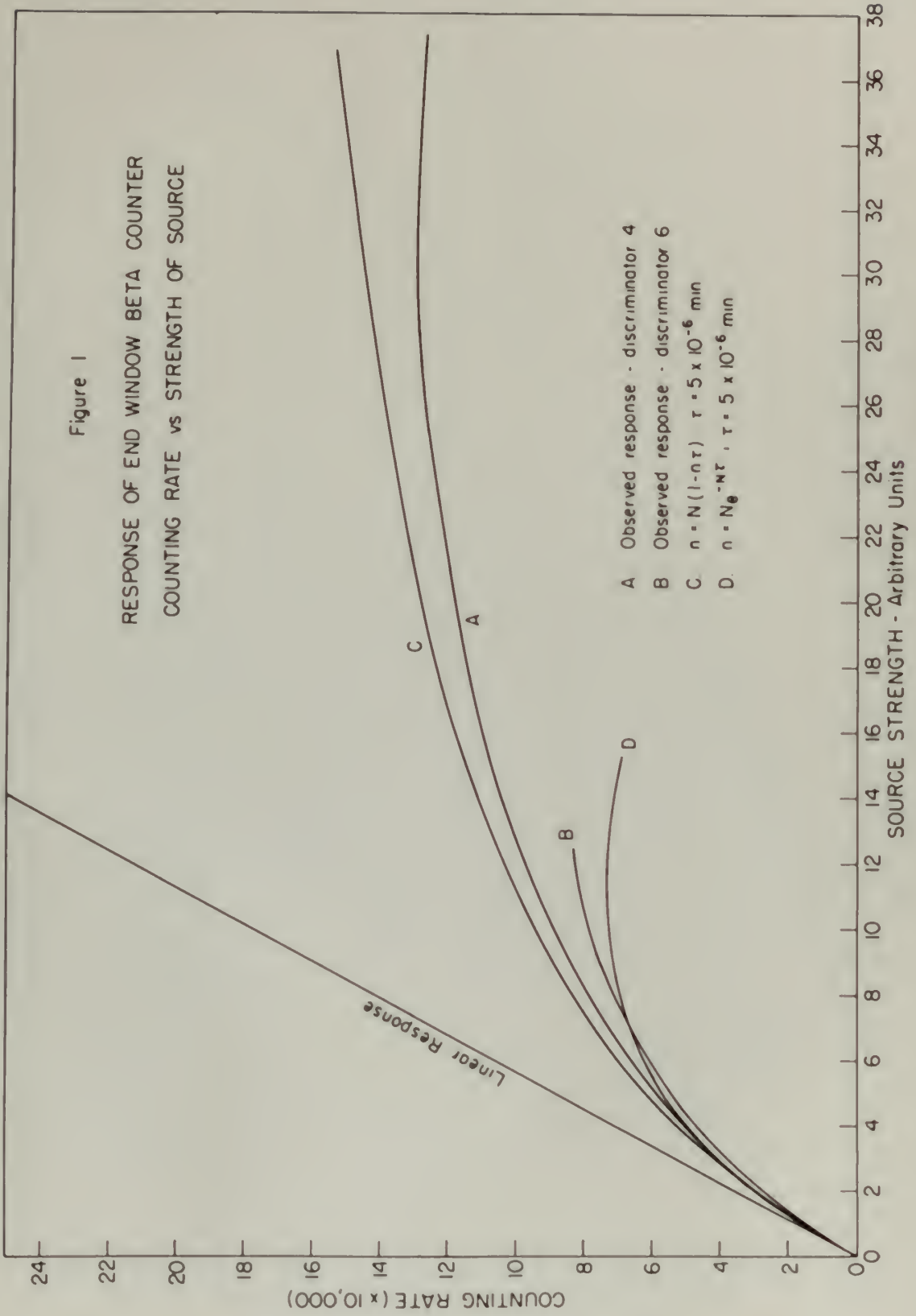
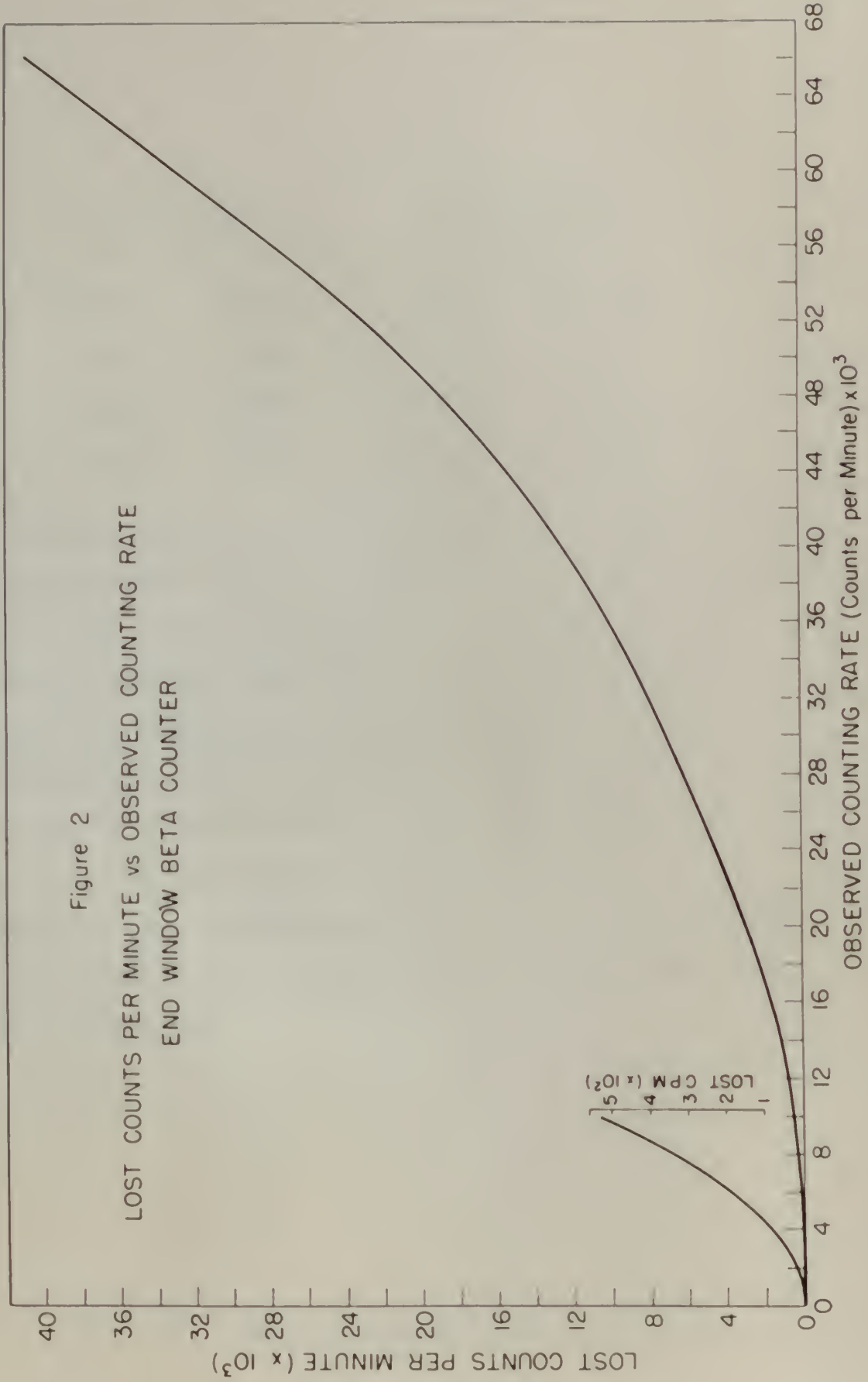


Figure 2

LOST COUNTS PER MINUTE vs OBSERVED COUNTING RATE
END WINDOW BETA COUNTER



<u>Observed counting rate cpm</u>		<u>Computed true counting rate cpm</u>	
<u>Disc. 4</u>	<u>Disc. 6</u>	<u>Disc. 4</u>	<u>Disc. 6</u>
10580	9850	10990	10700
41350	38910	54350	53900
67600	61100	110,000	109,000

Referring to Fig. 1, it is seen that neither response curve coincides with the theoretical curves of equations (1) and (2). It is apparent that the number of lost counts is strongly dependent upon the discrimination level. The results obtained may be explained by a consideration of the pulse height distribution as a function of counting rate.* This shows that as the counting rate is increased, many small pulses are formed and some fraction of these pulses is lost because of the discrimination level and not because of the dead time of the tube.

* McCall, R. C.: "Geiger-Muller Counters", M.I.T. Progress Report, 1953.

Observed counting rate cps		Computed true counting rate cps	
Class. A	Class. B	Class. A	Class. B
10880	9880	10880	10700
41300	38100	54300	52600
87000	81000	110,000	108,000

Referring to Fig. 1, it is seen that neither response curve coincides with the theoretical curves of equations (1) and (2). It is apparent that the number of lost counts is strongly dependent upon the discrimination level. The results obtained may be explained by a consideration of the pulse height distribution as a function of counting rate.* This shows that as the counting rate is increased, many small pulses are formed and some fraction of these pulses is lost because of the discrimination level and not because of the dead time of the tube.

* McCall, E. C. "Geiger-Müller Counters", W.I.T. Progress Report, 1951.

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1941-1942		1942-1943	
Actual	Budget	Actual	Budget
100	100	100	100
100	100	100	100
100	100	100	100

VI. SUMMARY

The following table shows the results of the operations of the company for the years 1941-1942 and 1942-1943. The table is presented in two columns, one for each year, and four rows, one for each item. The items are: Actual, Budget, Actual, and Budget. The results are as follows:

Actual 1941-1942: 100, Budget 1941-1942: 100, Actual 1942-1943: 100, Budget 1942-1943: 100.

The following table shows the results of the operations of the company for the years 1941-1942 and 1942-1943. The table is presented in two columns, one for each year, and four rows, one for each item. The items are: Actual, Budget, Actual, and Budget. The results are as follows:

Actual 1941-1942: 100, Budget 1941-1942: 100, Actual 1942-1943: 100, Budget 1942-1943: 100.

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APPENDIX AND IS INTENDED TO ASSIST THE READER IN LOCATING THE

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00

APPENDIX IV
THE COINCIDENCE COUNTER

A. Description

This equipment consists of two thallium activated sodium iodide scintillation counters connected in coincidence with single channel and coincidence scaling circuits driving mechanical registers. The crystals are 1.5 inches in diameter, 1 inch deep, and are mounted on RCA type 5819 photomultiplier tubes.

The counters are contained in lead shielded heads along with their cathode follower type preamplifiers shown schematically in Fig. 1. The two heads are mounted on a mechanical scanning device such that the two opposing crystals are coaxial and are separated by approximately 27 cm. A mounting bracket permits positioning of a source equidistant from the crystal faces and colinear with their common axis.

The coincidence circuit is of conventional design* providing both single channel and coincidence outputs which

* Dwg. No. B-1547-A, file 6425, Laboratory for Nuclear Science, M.I.T., 28 April 1950.

APPENDIX IV
THE COLLECTOR'S GUIDE

4. Description

This equipment consists of two main sections
which include a collector and a recorder in
addition to the alpha channel and coincidence counting
circuitry. The alpha channel is 1 inch long and is mounted
on a type 512 photomultiplier tube.
The recorder is mounted in lead shielded cans
along with their own type 512 photomultiplier
tubes. The two tubes are mounted
in a mechanical support which is the opposite
of the usual one and are rotated by approximately
90 degrees. A mounting bracket provides positioning of a source
independent from the crystal faces and collimator with their
common axis.
The coincidence circuit is of conventional design
providing both alpha channel and coincidence outputs which

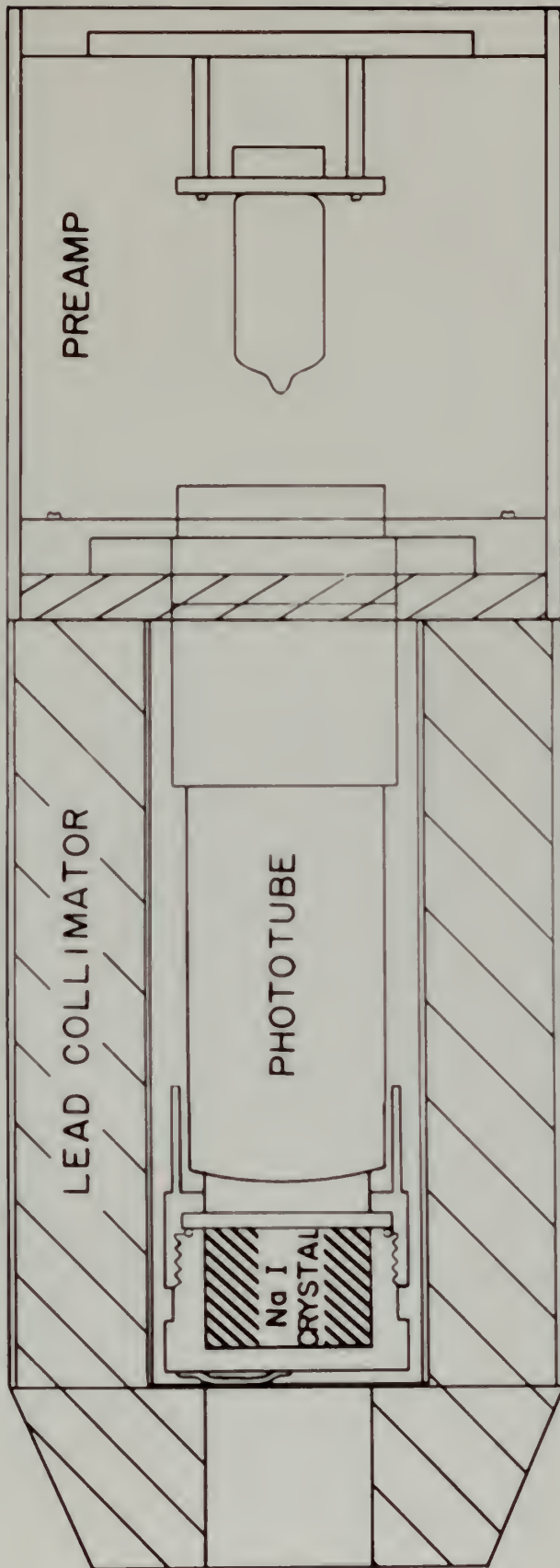
* Dept. No. 5-154-1, the 6050 Laboratory for Nuclear
Science, N.Y., in April 1960.



FIG. 1. Schematic diagram of detector assembly. The location of components within the lead collimator is as indicated.

Fig. 1. Schematic diagram of scintillation counter.

The location of components within the lead shielding head is as indicate.



0 10
SCALE = CM

are fed through linear amplifiers* to separate scaling circuits**. The equipment assembled for normal use is illustrated in Fig. 2.

B. Experimental Technique

The equipment is operated so that the individual channels register approximately equal counting rates when a source is at the mid-point on the axis between the counter heads.

Figure 3 illustrates that counting rates are only slightly affected by small displacements of the source from its central position. To minimize errors caused by variation in counter sensitivity due to other causes, a standard Na²² source was counted prior to each measurement and the correction thus determined was applied to the observed counting rate.

In all measurements the number of counts per 1 minute interval was recorded, each observation including at least three intervals for single channel counts and six intervals for coincidence counts. At least 10,000 events were included in each observation to insure a maximum fractional standard deviation of ≤ 1 percent.

* Atomic Instrument Co. Model 204B Linear Amplifier.

** Atomic Instrument Co. Model 1030 "Scale of 1000" Scaler.

are fed through linear amplifiers* to separate scaling
circuits**. The equipment assembled for tunnel use is
illustrated in Fig. 2.

B. Experimental Technique

The equipment is operated as first the individual
channels tested approximately equal counting rates
when a source is at the midpoint on the axis between
the counter heads.

Figure 3 illustrates that counting rates are only
slightly affected by small displacements of the source
from the central position. To minimize errors caused by
variation in counter sensitivity due to other causes, a
standard ^{22}Na source was counted prior to each measure-
ment and the correction thus determined was applied to
the observed counting rate.

In all measurements the number of counts per 1
minute interval was recorded, each observation including
at least three intervals for single channel counts and
six intervals for coincidence counts. At least 10,000
events were included in each observation to insure a
maximum fractional standard deviation of ≤ 1 percent.

* Model Instrument Co. Model 1003 Linear Amplifier.

** Model Instrument Co. Model 1000 Scale of 1000* Counter.



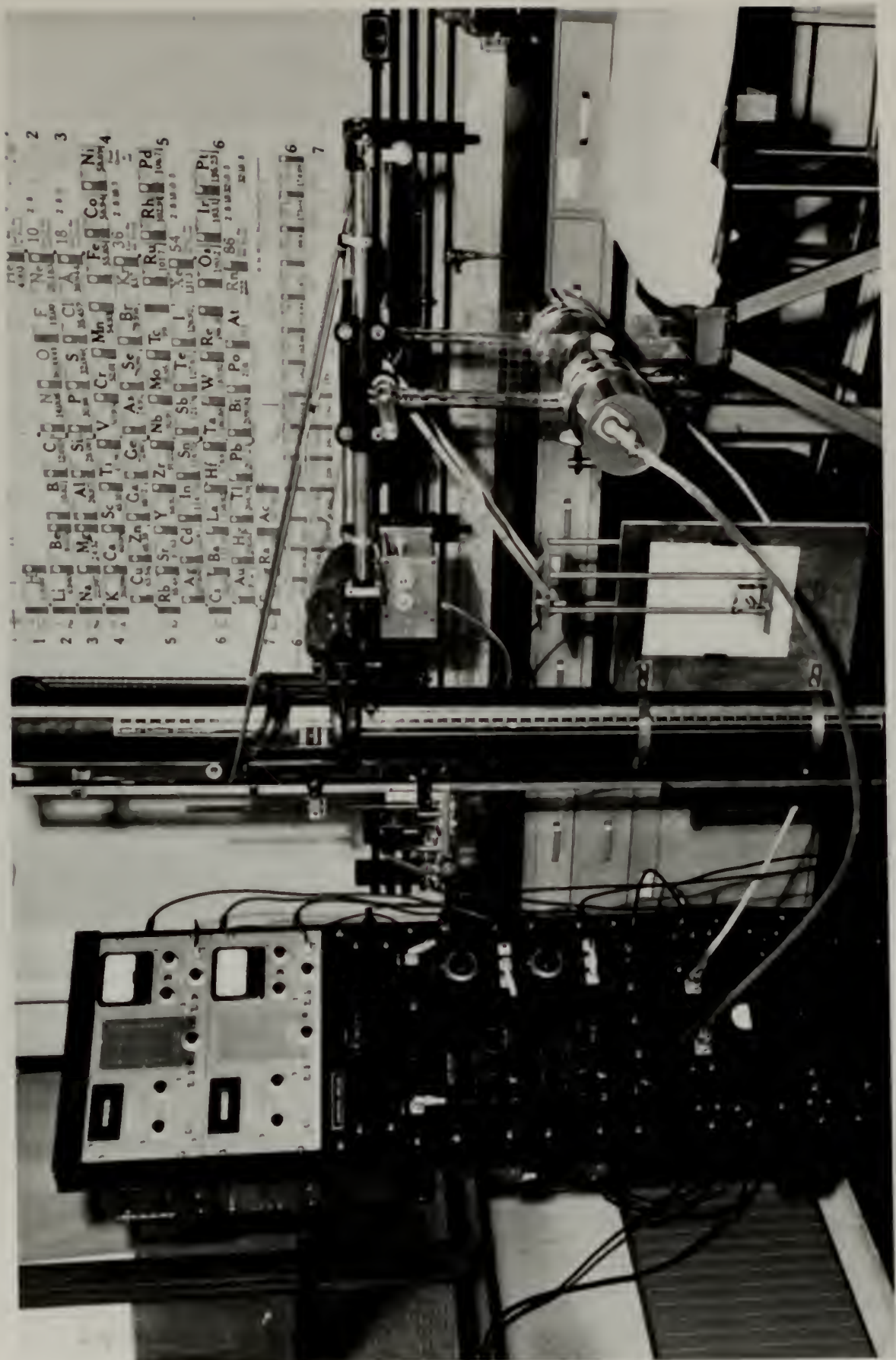
The following is a list of the
 names of the persons who
 were present at the meeting
 held on the 15th of the month
 of January, 1900, at the
 residence of Mr. J. C. [unclear]
 in the city of [unclear].

J. C. [unclear]

The following is a list of the
 names of the persons who
 were present at the meeting
 held on the 15th of the month
 of January, 1900, at the
 residence of Mr. J. C. [unclear]
 in the city of [unclear].

Fig. 2. Coincidence counter assembly.

The equipment is shown as assembled for use in scanning measurements at Massachusetts General Hospital. The scanning and plotting mechanisms are contained in the central section of the photograph. The opposing lead shielded counter heads are visible to the right of the plotting board.



1	H	Li	Be	B	C	N	O	F	Ne	10	20	2
2	Na	Mg	Al	Si	P	S	Cl	Ar	18	20	3	
3	K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	36	4
4	Rb	Sr	Y	Zr	Nb	Mo	Tc	Ru	Rh	Pd	54	5
5	Cs	Ba	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	86	6
6	Fr	Ra	Ac	Th	Pa	U	Np	Pu	Am	Cm	118	7

SENSITIVITY DISTRIBUTION-COINCIDENCE COUNTS

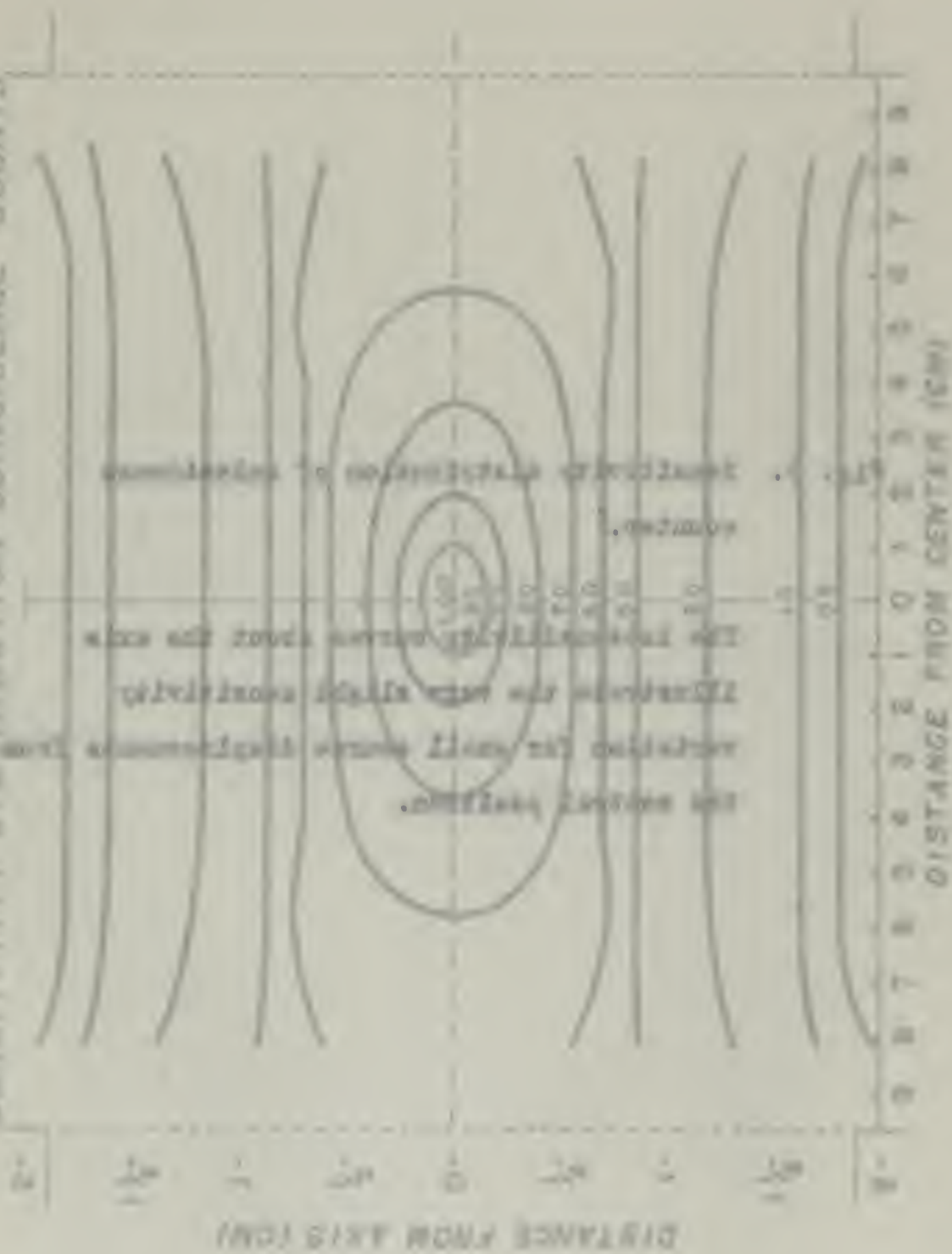
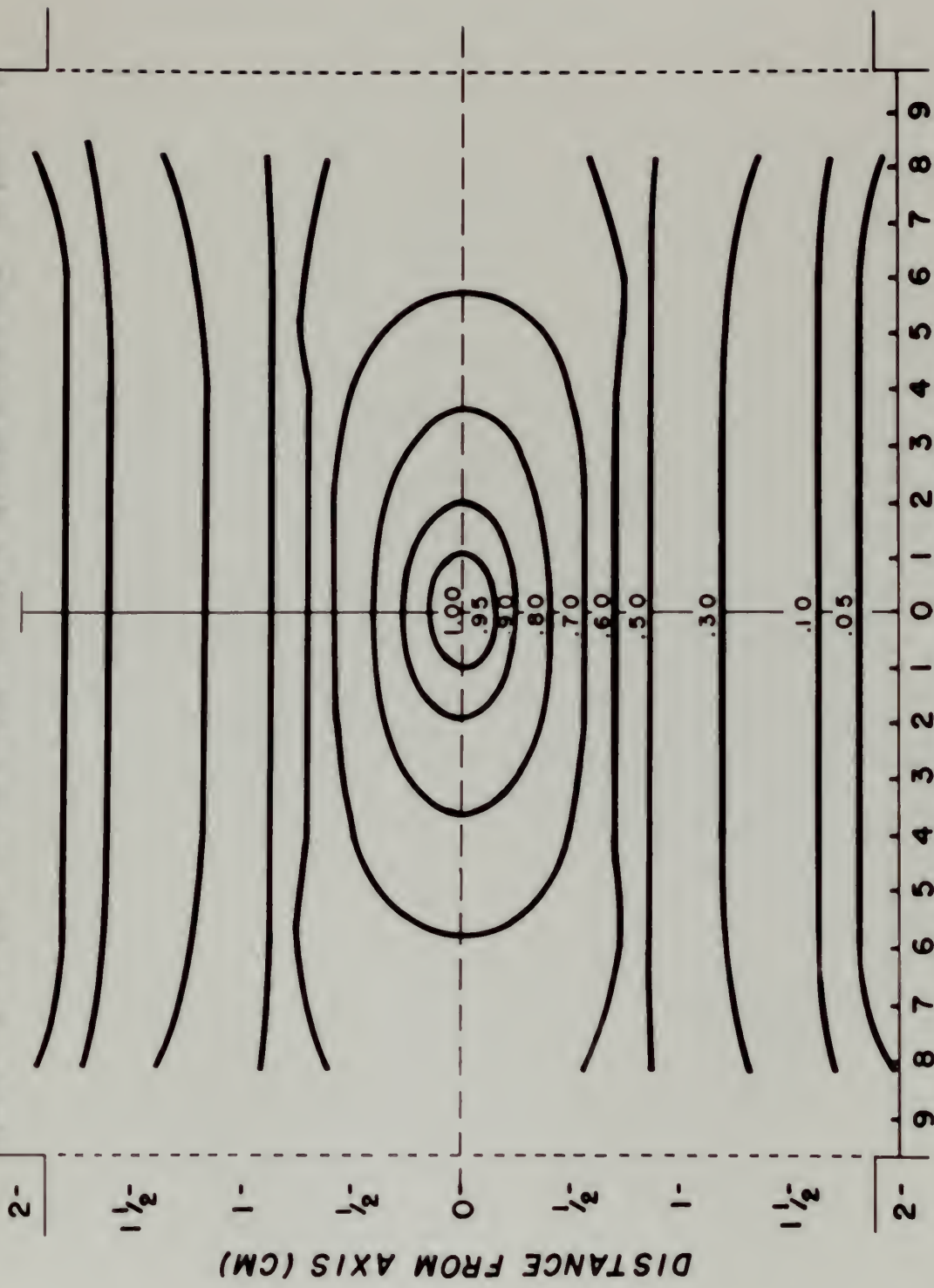


Fig. 3. Sensitivity distribution of coincidence counter.

The isosensitivity curves about the axis illustrate the very slight sensitivity variation for small source displacements from the central position.

SENSITIVITY DISTRIBUTION-COINCIDENCE COUNTS



DISTANCE FROM AXIS (CM)

DISTANCE FROM CENTER (CM)

C. Correction of Observed Counting Rates

Coincidence counting rates were corrected by subtracting from the observed values the chance coincidence rate. Chance rate was computed by means of the equation

$$C_{ch} = \tau N_a N_b$$

where N_a , N_b are the individual channel rates and τ is the resolving time of the coincidence circuit. By counting an essentially monoenergetic γ -ray emitter positioned off the axis of the crystals, τ was computed to be approximately 0.36 μ sec by use of the above equation.

Counter response appears to be linear for counting rates up to 140,000 cps on single channels and 14,000 cps for coincidences (Fig. 4). Consequently no corrections were applied to the data for resolving time losses.

C. Correction of Observed Counting Rates

Coincidence counting rates were corrected by subtracting from the observed values the chance coincidences rate. Chance rate was computed by means of the equation

$$C_{ch} = R_A R_B \tau$$

where R_A , R_B are the individual channel rates and τ is

the resolving time of the coincidence circuit. By

counting an essentially nonoverlapping γ -ray emitter

positioned off the axis of the crystals, τ was computed

to be approximately 0.34 nsec by use of the above equation.

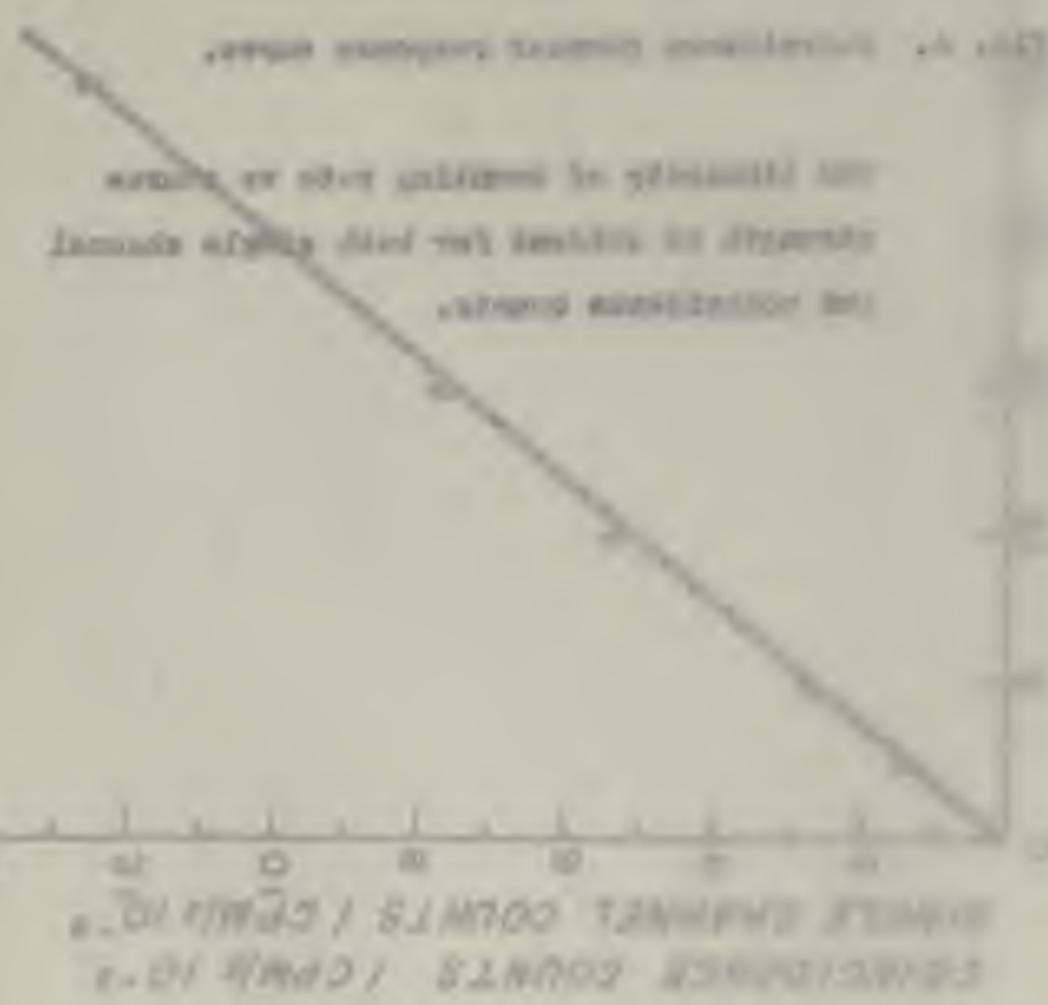
Counter response appears to be linear for counting

rates up to 140,000 cps on single channels and 14,000

cps for coincidences (Fig. 4). Consequently no corrections

were applied to the data for resolving time losses.

CORRECTION
 CURVE



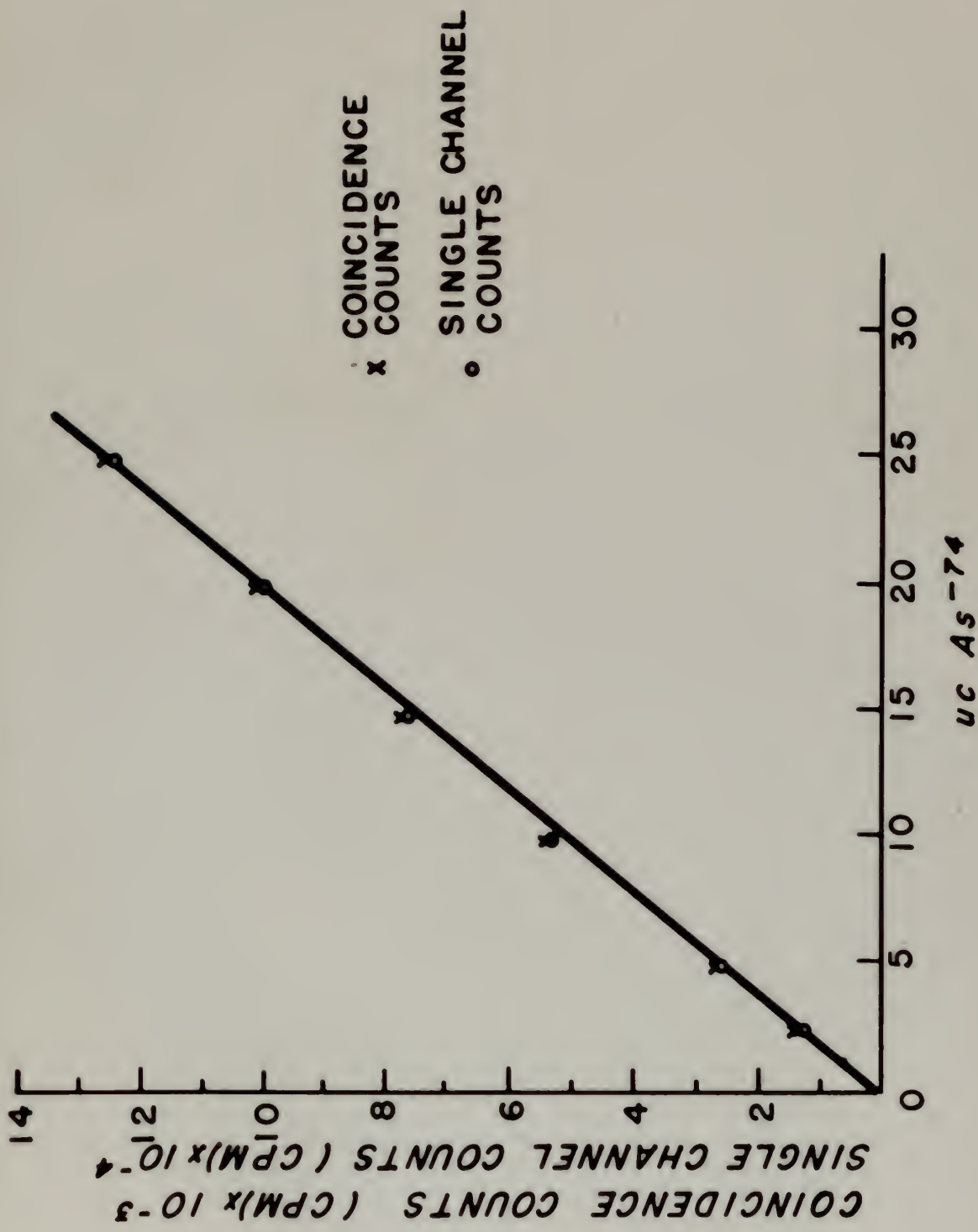
The correction curve is shown in the figure. The relationship of the correction curve to the measured counts is shown in the figure. The correction curve is shown in the figure.

The linearity of counting rate vs source strength is evident for both single channel and coincidence counts.

Fig. 4.

Fig. 4. Coincidence counter response curve.

The linearity of counting rate vs source strength is evident for both single channel and coincidence counts.



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An investigation of the radionuclides of arsenic produced by cyclotron bombardment of germanium with 15 Mev deuterons.

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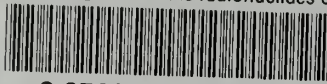
An investigation of the radionuclides of arsenic produced by cyclotron bombardment of germanium with 15 Mev deuterons.



Stanford Postgraduate School
Monterey, California

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