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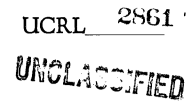
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Decay Properties of U^{232}

Frank Asaro and I. Perlman February 7, 1955

Berkeley, California

Decay Properties of U^{232⁺}

-2-

Frank Asaro and I. Perlman Radiation Laboratory and Department of Chemistry University of California, Berkeley, California

February 7, 1955

ABSTRACT

The alpha and gamma spectra of U^{232} have been studied with an alpha particle spectrograph and gamma-ray scintillation, proportional, and coincidence counters. Alpha groups of 5.318- (68 percent), 5.261-(32 percent) and 5.134-Mev (0.32 percent) and gamma rays of 57.9-(0.21 percent), 131- (0.075 percent), and 268- (0.004 percent) and 326-kev (0.004 percent) were observed. The half-life of the 58-kev first excited state of Th²²⁸ was found to be less than 50 microseconds. Spins and parities are assigned to the energy levels, and the results are evaluated with respect to the developing theory and systematics of complex alpha spectra and excited states of even-even nuclei.

^TThis work was performed under the auspices of the U.S. Atomic Energy Commission.

Decay Properties of U232[†]

Frank Asaro and I. Perlman Radiation Laboratory and Department of Chemistry University of California, Berkeley, California

February 7, 1955

INTRODUCTION

The uranium isotope, U^{232} , is a beta-stable alpha emitter with a half-life of 73.6 years¹ and was first identified following its growth from the shorter-lived β^- -emitter Pa^{232} which had been prepared by the d,2n reaction on Th^{232} .² The alpha particle energy has been determined by range measurement as 5.31 Mev³ and 5.27 Mev.⁴ It could be inferred that the U^{232} alpha spectrum was complex because prominent conversion electrons of an ~60-kev gamma ray were observed in coincidence with alpha tracks in a photographic emulsion.⁵ An early part of the present study, in which there was found an alpha group of 58-kev lower energy than the main group, has already been reported.⁶

There has been added recent interest in the alpha spectra of eveneven nuclei because they display prominently series of energy levels which have been interpreted as rotational states.^{7,8,9} The theory of Bohr and Mottelson which explains these rotational bands as a consequence of collective motions in highly deformed nuclei predicts a simple relationship of the energy spacing between members of the band: $E_{rot} \propto I(I + 1)$, where I refers to the spin which in this case is restricted to even integral numbers. As will be mentioned further in the

[†]This work was performed under the auspices of the U.S. Atomic Energy Commission. DISCUSSION, this simple expression applies to a limiting condition in which the rotational frequency is sufficiently slow to allow the nuclear structure to adjust adiabatically to the changing electric field. Where this situation does not apply, the rotational levels are perturbed and correction terms must be applied. It is noted that the heaviest nuclei show "pure" rotational bands and as the closed shell region around lead is approached the perturbation becomes more and more severe. In this context, it will be seen that Th^{228} (alpha decay of U^{232}) shows a just discernible departure from the ideal case and therefore represents the entry into the "light element" or closed shell region.

Another aspect of note in the decay of U^{232} is the appearance of a low-lying level which does not belong to the rotational band. This particular state with spin 1, odd parity, occurs in a limited region (around 136 neutrons). It was the subject of another publication¹⁰ and will be discussed further below.

METHODS

<u>Preparation of source</u>.--The U^{232} sources for the present study were prepared in two ways. In one of these Th²³² was bombarded with protons resulting in U^{232} by the following reactions:*

> $Th^{232}(p,n)Pa^{232}$ $Pa^{232} \xrightarrow{\beta} U^{232}$

*We wish to thank Dr. Louis M. Slater for making this sample available to us.

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The other method involved intensive neutron irradiation of ionium (Th^{230}) :

Th²³⁰(n,
$$\gamma$$
)Th²³¹
Th²³¹ $\xrightarrow{\beta}$ Pa²³¹
Pa²³¹(n, γ)Pa²³²
Pa²³² $\xrightarrow{\beta}$ U²³²

Inasmuch as Th^{232} was present in the irradiated ionium, U^{233} was also made by a first order neutron capture reaction. However, its contribution to the radioactivity was quite insignificant; the alpha spectrum showed a possible peak of the U^{233} position of maximum intensity 0.06 percent relative to the U^{232} groups. The factors which produced this favorable ratio were: (1) the ionium had been considerably enriched by electromagnetic separation¹¹ from the initial source materials, (2) the neutron capture cross sections for Th^{230} and Pa^{231} are much larger than that for Th^{232} , ¹² (3) the half-life for U^{233} is long compared with U^{232} , and (4) the uranium was separated before all of the intermediate Pa^{233} had decayed to U^{233} .

Alpha particle spectra.--All samples for the alpha-particle spectrograph were prepared by vacuum sublimation of the chloride onto a platinum plate which was masked to present a band 1 inch x 1/8 inch. This sample mounting technique, as well as the equipment and methods for taking alpha spectra, have been described in earlier reports.¹³⁻¹⁵ As before the alpha particles were caught on a photographic plate and the track count was plotted according to position on the plate. <u>Gamma ray measurement</u>.--Gamma-ray spectra were measured for the most part with a sodium iodide scintillation counter coupled to a 50channel pulse-height analyzer. In some experiments a xenon-filled proportional counter was used to produce the pulse for the analyzer.

Coincidence counting methods used in this and similar studies have been described elsewhere.^{10,16} Briefly, a gamma-ray pulse is fed into a 50-channel pulse-height analyzer which registers only when triggered through a gate circuit. For alpha-gamma coincidences the pulse produced by a zinc sulfide screen in optical contact with a photomultiplier tube is used to open the gate; for gamma-gamma coincidences, another sodium iodide crystal counter was used.

RESULTS

Alpha Spectrum

Four exposures, involving two U^{232} preparations, were made and the histories are given in Table I. The first measurement (Exp. 134) was made with a relatively weak sample before more active preparations had been made. Numbers 245 and 277 were moderately long exposures of an intense source aimed principally at finding low-intensity alpha groups. The more intense alpha groups register too many tracks for convenient counting. A shorter exposure (Exp. 278) was then madé to get more accurate data on the abundances and energies of the most prominent groups.

As a result of these several experiments, three alpha groups were found for U^{232} as summarized in Table II. Two of these are the characteristic high intensity groups of an even-even alpha emitter leading to the

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ground state and first excited state of the product. The energy of the main group was found to be 5.318 (±0.002) Mev and the second group was lower in energy by 57.2 ± 1.2 kev. (The 5.421-Mev¹⁷ alpha group of Th²²⁸, which grew into the U²³² sample, was used as the energy standard for these measurements.) When the differences in recoil energies from the two alpha groups are included, the first excited state is found to lie at 58.2 kev. The intensity of group α_{58}^* was found to be 32 percent.

Besides the two principal groups a weak alpha group of 0.32 percent abundance was identified as belonging to U^{232} . This group leads to a state 188 kev above the ground state. Gamma rays originating from this state and others will be described below. No other alpha groups belonging to U^{232} were noted but the limits of detection varied with the proximity to the observed groups. Over the energy range 5.07 Mev \rightarrow 4.83 Mev an upper limit of 0.01 percent could be set. At 4.822 Mev there was a small peak (0.06 percent of the U^{232} intensity) which is probably due to U^{233} as the energy agreement is good and it is expected that there would be some present.

The alpha spectrum of U^{232} with some of its daughter activities is shown in Fig. 1. The solid line curve was obtained from Exp. 277 in which the principal peaks were too strong to be counted. The positions and intensities of the strong groups taken from Exp. 278 are shown as the broken line curve of Fig. 1.

Refers to the alpha group leading to the 58-kev state.

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Gamma Rays and Decay Scheme

The partial decay scheme for U^{232} based on the present studies is shown in Fig. 2. It will be noted that a third excited level in Th²²⁸ is indicated by gamma radiation although the alpha group populating this state could not be seen. The experiments will be discussed in terms of the successive energy levels. Some of the gamma rays measured were extremely weak in intensity and a number of measurements of the gamma spectrum were made after repeated and varied chemical purification steps to make certain that these radiations did not arise from Th²²⁸ decay products. The only non-separable substance which could have been present was U^{233} and an upper limit to its intensity could be set on the basis of the alpha spectrum.

The measurements on the photons included: (1) identification and intensity measurements with a scintillation spectrometer, (2) alphagamma, and gamma-gamma coincidence determinations, (3) xenon-filled proportional counter measurements of energy. In all, 13 sets of measurements were made employing six different repurified samples of U^{232} from a common original stock.

<u>The 58-kev state</u>.--As indicated by the alpha spectrum, the first excited state lies at 58.2 ± 1.2 kev and is populated directly to the extent of 32 percent. Further possible population of this state by transitions from higher states cannot be much in excess of 0.32 percent.

Several measurements with the scintillation spectrometer in which the 59.6-kev gamma ray of Am^{241} was used as an energy standard showed a peak at 57 to 58 kev. Two measurements with a xenon-filled proportional

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counter gave the best energy determination as 57.9 ± 0.3 kev. The prominent Am²⁴¹ photon also served as an intensity standard. The best determination of the intensity was 2.1 x 10^{-3} relative to total alpha disintegrations. The total conversion coefficient is therefore 152 which agrees well with the value 120 for the L-shell conversion coefficient for an E2 transition which was estimated from available theoretical data.^{18,19} This assignment confirms the expected 2+ designation for the first excited state of Th²²⁸.

This same state has been observed by others²⁰⁻²² in the beta spectrum of MsTh₂ (Ac²²⁸). Brodie²¹ gave the energy of the transition as 57.0 kev and Kyles, Campbell and Henderson²² reported 56.75 kev. These two reports concluded that the transition is E2 although the conversion coefficients differed somewhat from those deduced here.

A puzzling feature of the 58-kev transition is an apparent discrepancy concerning the lifetime of the 2+ state. Two groups^{22,23} working on the beta spectrum of MsTh₂ have reported the lifetime of this state to be greater than 10 milliseconds. A third group has reported the lifetime to be larger than 0.5 second.²⁴ From the present results it is deduced that the lifetime of this state is less than 10 microseconds. In two measurements the coincidence rate between alpha particles and 58-kev gamma rays was 0.16 percent and 0.18 percent per alpha particle. These values correspond within experimental error with the above mentioned intensity of 0.21 percent for the 58-kev photon showing that it is in coincidence with the alpha particles within the resolving time of the coincidence circuit which was about 10 microseconds. In addition coincidences were observed between the 130 kev gamma ray and L x-rays. These L x-rays arise mainly from the conversion of the 58-kev gamma ray.

It is probable that the methods $^{22-24}$ by which delays between the events were noted could not distinguish between a long delay and a very rapid coincidence. It is to be expected 7,25 that in a region well away from closed shells such as this that the E2 transitions from the first excited state would be abnormally rapid rather than slow.

<u>The 188-kev state</u>.--A state at this energy is defined uniquely by the existence of a U^{232} alpha group with an energy lower than the ground state transition by the appropriate amount. As already mentioned this state is populated to the extent of about 0.32 percent.

The only gamma ray transition observed in the present study which depopulates this state is one of 131 kev. It leads from the 188-kev state to the 58-kev state as determined by gamma-gamma coincidence measurements between 131- and 58-kev photons and between the 131-kev gamma ray and L x-rays. This situation is similar to that of other eveneven nuclei among the heavy elements in which a series of states exist which are believed to belong to a rotational band and follow the sequence 0+, 2+, 4+,---. In none of these cases has a crossover transition been seen from the second even spin state to the ground state. It remains now to be seen if some of the other criteria for this sequence exist.

One of the properties of a pure rotational spectrum is that the energy spacings are proportional to I(I + 1), where I is the spin. The ratio of energies between the spin 4 and spin 2 states should be accordingly 20/6 or 3.33. In the present situation the ratio is 186/58 or 3.24.

If the 131-kev transition takes place between 4+ and 2+ states it must be of pure E2 character. Two measurements were made of the alphagamma coincidence rate for the 131-kev gamma ray. These gave intensities for the gamma ray of 0.05 and 0.06 percent. Two more precise measurements of the gamma spectrum gave a value of 0.075 percent for the best intensity of the 131-kev gamma ray. Inasmuch as the 188-kev state is populated to the extent of 0.32 percent the conversion coefficient becomes ~3.2. This is in good agreement with the expectations for an E2 transition, it is definitely too high for an El transition, and is about a factor of 3 too low for an Ml transition. There is some radiation observed at about 85 kev (see Fig. 3.). Although it may include scattered radiation from the 131-kev gamma ray, its abundance represents the maximum intensity of K x-rays in the decay of U^{232} . This abundance is 0.4 of the abundance of the 131-kev gamma ray, giving a maximum K conversion coefficient of 0.4. This is in good agreement with the expectations for an E2 transition but is a factor of 20 too low for an Ml transition.

A gamma ray transition of about this energy (128 kev) was studied by $\operatorname{Brodie}^{21}$ and by Kyles, Campbell, and Henderson²² in the conversion electron spectrum from MsTh_2 . As pointed out^{22} the L subshell conversion ratios and the low limit set on the K conversion indicates an E2 transition with which our conclusions agree. The same gamma ray (129 kev) was seen by Box and Klaiber²⁶ using a scintillation spectrometer.

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These workers^{20,22} also studied a transition of about 184 kev which they interpreted as an Ml transition and in their decay scheme was placed as the crossover transition from a level of this energy to the ground state for which the cascading transitions are those of 128 kev and 58 kev. In order to fit spins and parities to these conditions the dequence for the ground state and next two higher states was postulated²² to be 0+, 2+, 1+. There are two reasons why a state at 184 kev which meets this condition cannot be the same as our 188-kev state defined by the alpha decay of U^{232} . In the first place an even-even alpha emitter cannot populate directly a state of odd spin and even parity. The other argument has to do with intensities of gamma rays. According to the data and deductions of Brodie²¹ and of Kyles and coworkers²² the photon intensities of the 128-kev and 184-kev should be about the same. From our measurements the intensity of the 131-kev gamma ray is at least 30 times greater than a peak at 184 kev (see Fig. 3). In order to reconcile these results it is necessary to say that either there is a 1+ state at 184-kev state (populated from the beta decay of MsTh_o) which is different from our 188-kev state (from alpha decay of U^{232}) or the observed 184-kev transition involves higher lying states of Th²²⁸ which also would not be expected to be seen in alpha decay processes. An added datum which must be considered in the overall interpretation is the failure of Box and Klaiber²⁶ to observe a 184-kev photon in their scintillation counter study of MsTh. This could imply a high conversion coefficient for the 184-kev transition although it is rather difficult to set limits on the basis of their reported data.

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<u>The 326-kev state</u>.--From the present work a level at this energy (see Fig. 2) could only be inferred from gamma-ray measurements as the alpha particle intensity is apparently too weak for direct detection. The evidence for the state is based on the observation of two gamma rays of 326 and 268 kev (see Fig. 4). The energy differences alone suggest transitions from a 326-kev level to the ground state and the 58-kev state, respectively.

The intensities of the two peaks as determined with the scintillation spectrometer were both about the same and amounted to about 4×10^{-3} percent of the total alpha disintegrations for each. An alphagamma coincidence measurement for the 268-kev gamma ray gave an intensity of 5×10^{-3} percent. Similarly, if L x-rays were used to gate the scintillation spectrometer it was found that the 268-kev gamma rays were in coincidence and the intensity came out to be 3×10^{-3} percent per alpha particle. Much less abundant coincidences with L x-rays were found for the 326-kev gamma ray. Because the L x-rays arise predominantly from the 58-kev transition it is probable that the 268-kev gamma ray goes to the 58-kev state and the 326-kev gamma ray does not. The L x-ray coincidences with the 326-kev gamma ray could be due to impurities or to the decay of still higher excited states populated by Th²²⁸ alpha emission.

These intensity, coincidence and energy considerations imply that a single state at 326 kev is responsible for both of these photons as shown in Fig. 2. Box and Klaiber²⁶ also found a pair of gamma rays differing by 58 kev and having approximately the energies seen here.

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Their values were 336 and 278 kev. Furthermore, they found coincidences between both of these and a gamma ray of 790 kev which is also consistent with the supposition that both originate from a common level. However, the spectrum shown by these authors would indicate that the 336-kev gamma ray is at least twice as intense as the 278-kev gamma ray while in our measurements the 278-kev photon seemed to be somewhat more intense. This may suggest that there are two gamma rays of about 336 kev, one of which occupies the position as shown by us (and by Box and Klaiber) and the other is a transition between higher levels. Significantly, Kyles and co-workers²² report conversion Tines from a gamma ray of 336 kev but not from one at 278 kev.

In the absence of conversion coefficient data for our 326-and 268-kev transitions it is not possible to characterize them with certainty. We have been led to assign the 326-kev level 1- and consequently the transitions El by analogy with the well characterized 1-levels found from other even-even alpha emitters in this region.¹⁰

DISCUSSION AND SUMMARY

The decay scheme for U^{232} shown in Fig. 2 bears a strong resemblance to those of other even-even alpha emitters in this region. In each case a rotational band consisting of 0+, 2+, and 4+ states is seen. In some cases the 1- state lies between the 2+ and 4+ states but for U^{232} decay it lies above the 4+ state. Just as in other cases, two E2 transitions are seen which cascade from the 4+ to the 2+ and 0+ (ground state). The 1- state is de-excited by competing El transitions to the 2+ and 0+ states. To be sure, further information must be

-14-

obtained to get independent evidence that the transitions in question for $\text{U}^{2\,32}$ are indeed EL.

As has already been discussed there is a partial correlation of the energy levels found in the present study from the alpha decay of U^{232} and those from the beta decay of $MsTh_2$ found by others. There are also some points of apparent disagreement. Because of the extreme complexity of the $MsTh_2$ spectrum it is not possible to resolve these difficulties readily from existing data. Some of the information worth seeking in future studies on $MsTh_2$ has been mentioned in earlier discussion.

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REFERENCES

1.	Sellers, Stevens and Studier, Phys. Rev. <u>94</u> , 952 (1954).
2.	Gofman and Seaborg, National Nuclear Energy Series, Plutonium
	Project Record: Transuranium Elements Vol. 14B (McGraw-Hill
	Book Co. Inc., New York, 1949), p. 1427.
3.	A. H. Jaffe, unpublished data (1948).
4.	Kahn and Linenberger, reported in Los Alamos Scientific
	Laboratory Report LAMS-151 (October 1944).
5∘	Dunlavey and Seaborg, Phys. Rev. <u>87</u> , 165 (1952).
6.	Hollander, Perlman, and Seaborg, Revs. Modern Phys. 25, 469 (1953).
7.	Bohr and Mottelson, Phys. Rev. <u>89</u> , 316 (1953); <u>ibid</u> . <u>90</u> , 717 (1953)
	Kgl. Danske Videnskab Selskab, Matfys. Medd. <u>27</u> , No. 16 (1953).
8.	A. Bohr, Rotational States of Atomic Nuclei (Ejnar Munksgaard,
	Copenhagen, 1954).
9.	Asaro and Perlman, Phys. Rev. <u>91</u> , 763 (1953).
10.	Stephens, Asaro, and Perlman, Phys. Rev. <u>96</u> , 1568 (1954).
11.	For typical available compositions of enriched ionium see:
	Harmatz, McCurdy and Case, Oak Ridge National Laboratory Report
	ORNL-1724 (1954). We wish to thank members of the Electronuclear
	Research Division of Oak Ridge National Laboratory for making some
	enriched ionium available to us.
12.	See: <u>Neutron Cross-Sections</u> , U.S. Atomic Energy Commission Un-
	classified Report AECU-2040 (May 1952).
13.	F. L. Reynolds, Rev. Sci. Instr. <u>22</u> 749 (1951).

14. Asaro, Reynolds, and Perlman, Phys. Rev. 87, 277 (1952).

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- 15. Asaro, Thompson, and Perlman, Phys. Rev. <u>92</u>, 694 (1953).
- 16. Asaro, Stephens, Thompson, and Perlman, University of California Radiation Laboratory Unclassified Report UCRL-2766 (October 1954), Phys. Rev., to be published, April 1, 1955.
- 17. Asaro, Stephens, and Perlman, Phys. Rev. <u>92</u>, 1495 (1953).
- 18. Rose, Goertzel and Swift, <u>Tables</u> of <u>Conversion</u> <u>Coefficients</u>, privately circulated.
- 19. Gellmann, Griffith, and Stanley, Phys. Rev. 85, 944 (1952).
- 20. D. H. Black, Proc. Roy. Soc. (London) 106A, 632 (1944).
- 21. W. D. Brodie, Proc. Phys. Soc. <u>67</u>, 265 (1954).
- 22. Kyles, Campbell, and Henderson, Proc. Phys. Soc. <u>66</u>, 519 (1953).
- 23. Lewin, Perey, and Teillac, J. Phys. Radium 10, 33 (1949).
- 24. Suzor and Charpak, J. Phys. Radium 15, 682 (1954).
- 25. Goldhaber and Sunyar, Phys. Rev. <u>83</u>, 906 (1951).
- 26. Box and Klaiber, Phys. Rev. <u>95</u>, 1247 (1954).

Exposure number	Source of U ²³²	Slit width of spectrograph source (inch)	Duration of exposure	Activity of sample (dis/minute)	Time after Th ²²⁸ removal
134	$Th^{232}(p,n)Pa^{232}$ $Pa^{232} = U^{232}$	1/8 x 1	46 hr	10 ⁵	~l week
245	$Io^{230}(n,\gamma)Th^{231} \xrightarrow{\beta^{-}} Pa^{231}$ $Pa^{231}(n,\gamma)Pa^{232} \xrightarrow{\beta^{-}} U^{232}$	1/8 x 1	14 hr	6 x 10 ⁸	~l month
277	same sample as used in exposure 245	0.018 x 3/4	46 hr	•	7 months
278	same sample as us ed in exposure 245	0.018 x 3/4	47 min	• • • • •	7 months

Alpha-Particle Spectrograph Exposures

Table I

-18-

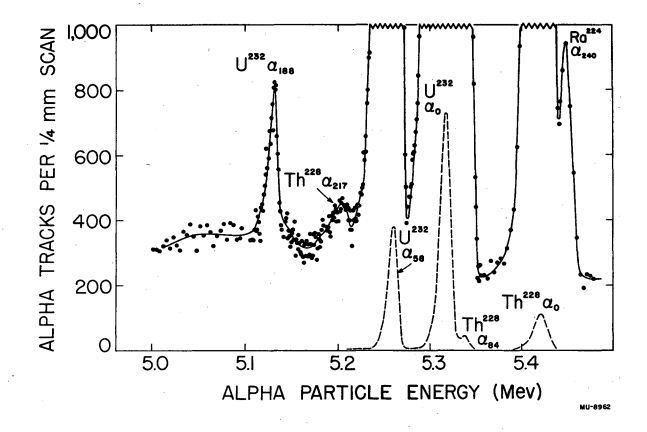
Best value	57.2	58.2	32		188	0,32	5.318	
							· · · · · · · · · · · · · · · · · · ·	<u> </u>
278	57.2 ± 1.2	58.2	32 ± 1	-	. 🛥	نە د	5.318 ± 2	5.261 ± 2
277	-	*	3	129. ₅	39 -	0.32 ± 0.0	3 -	5.260 ± 2
245	55 ± 4	56	35	۰. ۲۰۰۰ میں	189	0.3	5.317	-
134	58 ± 2	59 . 5	31		-		- .	· 🛶 –
			· · · · · · · · · · · · · · · · · · ·					
	(kev)	recoil (kev)	(%)	(kev)	(kev)	(%)	(Mev)	(Mev)
	between α_{0} and α_{58}	including difference in nuclear	to total $U^{232} \alpha$ particles	between α_{58} and α_{188}	separation between α_0 and α_{188}	to total U232 $lpha$	relative to Th ²²⁸ α_0 (5.421 Mev)	relative to Th ²²⁸ α_0
Exposure number	α particle energy separation	Separation between α_0 and α_58	Intensity of α58 relative	α decay energy separation	à decay energy	Intensity of α <u>188</u> relative	α particle energy of U ² 32 α ₀	α particle energy of $U^{2}3^2 \alpha_{5}8$

Results of Spectrograph Exposures

UCRL-2861

-19-

Table II



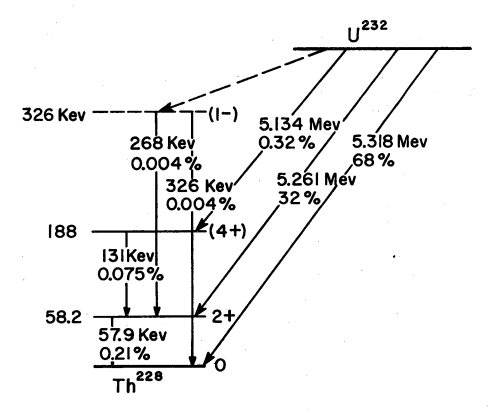
-20-

Fig. 1. U^{232} alpha spectrum with daughters

partially grown in.

- exposuré 277

----- exposure 278 Ordinate scale should be multiplied by 2.



MU-8963

Fig. 2. U^{232} decay scheme.

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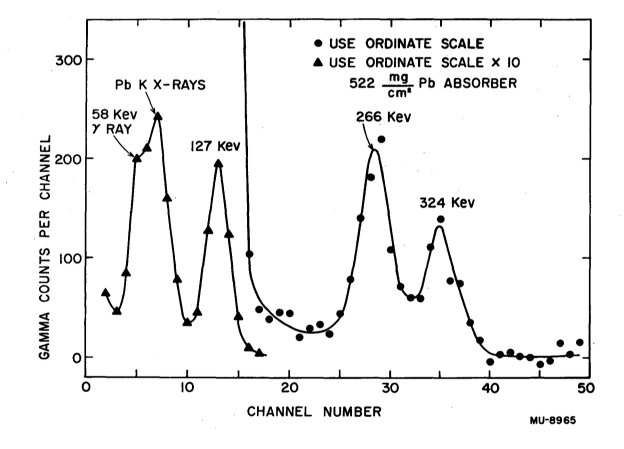


Fig. 3. U²³² low-energy gamma spectrum.

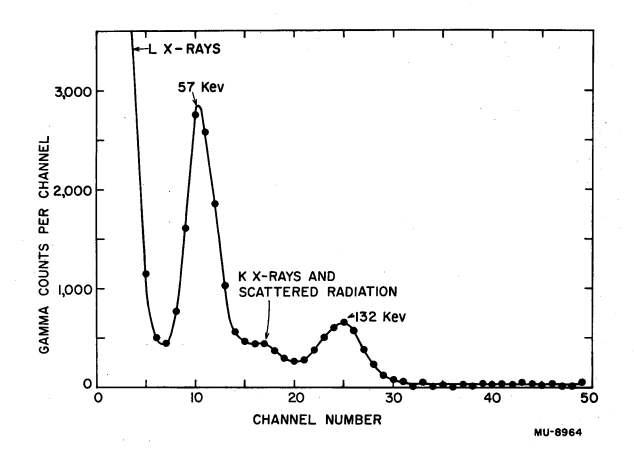


Fig. 4. U²³² high-energy gamma spectrum.