The β-Ray Spectrum of ₉₁Ekatantalum²³³ *

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THE investigations of A. V. Grosse, E. T. Booth and J. R. Dunning¹ definitely established the existence of a β -emitting isotope of ekatantalum produced by irradiating thorium with slow neutrons. They found the halflife of this product to be 27.4 ± 0.3 days. It was assumed that the $_{91}$ Et²³³ decayed with the emission of a β -particle into $_{91}U^{233}$ and to test this hypothesis the β -ray spectrum of this substance was examined. It was found to consist mainly of conversion lines, some of the most intense of which were very close to those of 91Pa²³¹. To dispel any doubts as to the purity of the Et²³³ two separate curves were taken 27 days apart. Similar runs were made on a source of protactinium prepared by A. V. Grosse. The intensity of all the lines in the spectrum of Et²³³ decay at the same rate of approximately 27 days while the protactinium remained the same. Furthermore, closer examination revealed definite differences in the energies and structure of the lines of the spectra of ekatantalum and protactinium.

Because of the complex structure of the β -ray spectrum of ${}_{91}\text{Et}^{233}$, it is difficult to distinguish the nuclear disintegration electrons from the conversion electrons due to gamma-radiation. A very large fraction of the β -activity is due to conversion lines of which at least four can be distinguished at 63, 77, 192 and 293 kev. From Fig. 1 it appears that these lines are superposed on a continuous spectrum of relatively small intensity whose maximum energy end-point is masked by the line at 192 kev. By subtracting this line from the continuous background the end-point appears to lie in the neighborhood of 230 kev. Above 230 kev the spectrum consists most probably of a single conversion line at 293 kev.



FIG. 1. Beta-ray spectrum of spectrum of statistical spectrum of s

Because of the complexity of the spectrum it is planned to re-examine Et²³³ with greater resolving power (the present resolving power, i.e., half-width of line divided by $H\rho$ at maximum intensity, is 5 percent). The energies involved are in the region where corrections for absorption in the counter window (0.5-mil Cellophane) must be applied. A more detailed study with greater resolving power and thinner window is in progress which should yield more significant values of the end-point of the continuous spectrum and the energies of the gamma-rays.

¹A. V. Grosse, E. T. Booth and J. R. Dunning, Phys. Rev. **59**, 322 (1941). * Publication assisted by the Ernest Kempton Adams Fund for Physical Research of Columbia University.

The Fourth (4n+1) Radioactive Series^{*}

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 S_{90}^{110CE} the discovery by Fermi and his co-workers¹ of $_{90}^{203}$, a type 4n+1 radioactive product, numerous attempts^{2,3} have been made to determine its successive disintegration products. The discovery of fission by Hahn and Strassmann has generally cast doubt upon the interpretation of previous work in this field, and quite recently Hahn and Strassmann⁴ themselves questioned their original interpretation in regard to this series. We began work more than a year ago to renew these investigations.

Measurements were first made on the half-life of the thorium capture product, Th²³³, and the thickness of aluminum required to reduce the ionization-chamber readings to one-half. These were found, respectively, to be 23.0 minutes and 0.020 cm. Similar experiments with U²³⁹ yielded 23.5 minutes and 0.018 cm for the equivalent values. To dispel any doubts that these two decaying products might be the same, uranium surrounded by cadmium was irradiated with neutrons in a paraffin geometry near the cyclotron. To this irradiated sample a small amount of thorium was added as a carrier and precipitated as a fluoride. This thorium precipitate contained less than one percent of the uranium activity, thus proving definitely that the two products are not identical.

It has been found that the thorium captures neutrons by a strong resonance process similar to uranium. Three identical samples of thorium (50 mg ThO₂) painted on nickel sheets were exposed to neutrons from the cyclotron in a paraffin geometry. One sample was surrounded by no absorbers, the second by 0.40 g/cm² of cadmium, and the third by 1.4 g/cm² of boron. The initial activities of the 23-minute periods were in the ratio of 100 : 55 : 11. This is interpreted as evidence that thorium has a strong resonance capture above the cadmium cut off at an energy less than 25 ev.

A search was made for " $_{91}A^{233}$ " reported by I. Curie² to be the first disintegration product and to have a period of from 2 to 3 minutes. Thorium was irradiated with neutrons in a paraffin geometry, and element 91 separated with zirconium phosphate, the precipitate being placed upon an ion chamber in less than five minutes after the end of the irradiations. No short period was found in any of a number



FIG. 1. Decay curves of 91ekatantalum²³³.

of experiments. Under the same chemical conditions UX₂ (brevium) was separated from uranium practically quantitatively. A further search revealed a β -emitting isotope of ekatantalum (Et) from thorium irradiated with slow neutrons, with a period of 27.4 days. Since this isotope is excited by slow neutrons and the number of active atoms is quantitatively in approximate agreement with the number of active atoms of Th²³³ formed under the same conditions, it is presumed to be the daughter of Th²³³.

A strong preparation of 91Et²³³ was made by irradiating 7 kg of thorium nitrate in solution (kindly supplied by the Maywood Chemical Company) near the cyclotron for about 50 hours. It was concentrated by a factor of 350,000 by a procedure originally used for the isolation of protactinium.⁵ A practically quantitative separation was effected using 20 grams ZrO2 as a carrier. A final concentrate of 20 mg ZrO₂ was obtained by means of the spiral process⁵ and contained initially about 10¹⁴ atoms of 91Et²³³. This separation from Zr disproves the recent suggestion by Hahn and Strassmann⁴ that their 25-day activity from thorium precipitated by zirconium phosphate may be an isotope of zirconium, and definitely proves that their "pre-fission" interpretation was correct.6

The decay of the sample has been followed for nearly a year, some of the curves being shown in Fig. 1. Curves 1, 3, and 4 were made through a thickness of 2 mils of aluminum between the source and ion chamber, while curve 2 was made through 10 mils. The mean value of the half-life from these and other measurements is 27.4 ± 0.4 days; decay curves through 1.5 and 4.5 g/cm² of lead showed the same period. The radiation as shown by cloudchamber experiments consists of negative electrons7 and rather strong γ -rays. As has been shown by absorption measurements and more accurately by E. Haggstrom,⁸ the maximum energy of the electrons is somewhat less than 300 kev.

Further studies are being made of the decay products.

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¹ E. Fermi, E. Amaldi, O. D'Agostino, B. Pontecorvo, F. Rasetti and E. Segrè, Proc. Roy. Soc. A149, 522 (1935). ² I. Curie, H. v. Halban and P. Preiswerk, Comptes rendus 200, 1841 (1935).

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F. Strassmann and O. Hahn, Zeits. f. Physik 109, 538 (1938).
O. Hahn and F. Strassmann, Naturwiss. 28, 543 (1940).
A. V. Grosse, Ber. d. D. Chem. Ges. 61, 238 (1928).
A confirmation of this result has been obtained by G. T. Seaborg and J. W. Gofman at the University of California (private communi-

cation). We wish to thank Mr. G. Weil for making the cloud-chamber

observations

⁸ E. Haggstrom, Phys. Rev. **59**, 322 (1941). * Publication assisted by the Ernest Kempton Adams Fund for Physical Research of Columbia University.

Nomenclature of Nuclear Particles

C. Møller Institute of Theoretical Physics, University of Copenhagen, Copenhagen, Denmark December 10, 1940

FOLLOWING the original proposal of Belinfante,¹ the writer has in a recent note on the meson theory of nuclear forces² used the word "nuclon" as a common notation for the heavy nuclear constituents, neutrons and protons. In the meantime, however, it has been pointed out to me that, since the root of the word nucleus is "nucle," the notation "nucleon" would from a philological point of view be more appropriate for this purpose, and I am therefore glad to have the opportunity to call the attention of interested physicists to this point.

¹ F. J. Belinfante, Theory of Heavy Quanta, Proefschrift, s'Gravenhage, 1939. ² C. Møller, Phys. Rev. 58, 1118 (1940).

Fission Products of Uranium by Fast Neutrons

V. NISHINA AND T. YASAKI Nuclear Research Laboratory, Institute of Physical and Chemical Research, Komagome, Hongo, Tokyo, Japan AND K. KIMURA AND M. IKAWA Chemical Institute, Faculty of Science, Imperial University of Tokyo, Tokyo, Japan December 23, 1940 N continuing our work1 on the fission of uranium pro-

duced by fast neutrons, we found in the following way that rhodium and ruthenium isotopes are also produced by fission.

The uranium oxide, U₃O₈, carefully purified and freed from its disintegration products, was exposed for about 12 hours to fast neutrons produced by bombarding lithium with 3 Mev deuterons of some ten microamperes from our cyclotron. From the irradiated sample, rhodium was separated as metal and ruthenium as sulphide. Each fraction which was carefully freed from the known fission products such as molybdenum, palladium, silver, cadmium, antimony, tellurium, iodine, caesium, barium, lanthanum, etc., was examined for its activity.

The rhodium fraction shows a decay curve of a single period 34 hours, while the ruthenium fraction gives two