## Letters to the Editor

 $oldsymbol{D}$  UBLICATION of brief reports of important discoveries in physics may be secured by addressing them to this department. The closing date for this department is five weeks prior to the date of issue. No proof will be sent to the authors. The Board of Editors does not hold itself responsible for the opinions expressed by the correspondents. Communications should not exceed 600 words in length.

## Mass Spectrographic Identification of Radioactive Lanthanum Isotopes\*

R. A. NAUMANN,\*\* F. L. REYNOLDS, AND I. PERLMAN Radiation Laboratory and Department of Chemistry, University o California, Berkeley, California December 12, 1949

WO neutron deficient isotopes of lanthanum, one new and the other previously tentatively assigned, have been given mass assignments by the technique of dispersing the isotopes on a photographic plate with a mass spectrograph. These and other light isotopes of lanthanum were prepared and some of their radioactive characteristics determined by absorption methods and with a low resolution beta-ray spectrometer.

In some experiments, pure CsNO<sub>3</sub> was irradiated with 30-Mev alpha-particles and this was followed by the isolation of the lanthanum fraction. The mass spectrograph showed a single radioactivity at mass number 135; and by following the decay of this line on the photographic plate through a slit arrangement, a 19-hour half-life was obtained. This confirms the mass assignment previously made by other methods.1 Since this irradiation should have produced La<sup>136</sup> and La<sup>134</sup> as well as La<sup>135</sup>, it was inferred that La<sup>136</sup> and La<sup>134</sup> have half-lives less than about 20 minutes since it required several hours to make the exposure on the mass spectrograph. The isotope La<sup>136</sup> had previously been reported as a 2-hour positron emitter,<sup>1</sup> but the observed activity has since been shown to be F<sup>18</sup> which partially went through the particular chemical separation employed. A search was made for La<sup>136</sup>, and it proved to be the 10-minute period reported by Maurer<sup>2</sup> from the irradiation of barium with deuterons. The assignment was made to La<sup>136</sup> since the ratio of the 10-minute period to the 19-hour period increased with decreasing alpha-particle energy on cesium in the range below 30 Mev, as would be expected for the  $(\alpha, n)$ reaction in relation to the  $(\alpha, 2n)$  reaction. Furthermore, the preparation of this activity in good yield by Maurer<sup>2</sup> using 5.8-Mev deuterons on barium would now point to La136 in view of the present knowledge of La<sup>138</sup>, La<sup>137</sup>, and La<sup>135</sup>, unless one assumes the 10-minute activity to be a metastable state of one of these isotopes. Very recently Robertson, Carss, and Pool,3 using separated stable barium isotopes as targets, have definitely proved this period to be associated with La<sup>136</sup>. The characteristics of La<sup>136</sup> which we determined are a half-life of 9.5 minutes, positron of 2.1 Mev determined with the spectrometer, and x-rays in high abundance from which it is estimated that about  $\frac{1}{3}$  of the decay is by positron emission and  $\frac{2}{3}$  by electron-capture.

The irradiation of cesium with 50-Mev alpha-particles resulted in a new lanthanum activity of 4.0-hour half-life. This activity has positrons of about 1.2 Mev, a conversion electron line of 0.26 Mev determined with the spectrometer, a harder gamma-ray of 0.8 Mev measured by lead absorbers, and K x-rays in high yield. This activity apparently decays predominantly by electroncapture since the positrons were only a few percent as abundant as the x-rays.

The 4-hour period was assigned to La133 by the mass spectrograph method. Figure 1 shows the original photographic plate

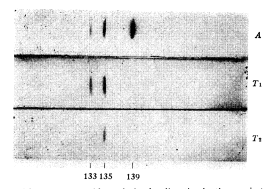


FIG. 1. Mass spectrographic analysis of radioactive lanthanum isotopes by transfer technique. (Exposures for transfer plates  $T_1$  and  $T_2$  given in the text.)

(A) and two transfer plates  $(T_1 \text{ and } T_2)$  made by exposing successively blank photographic plates with the receiver plate. The upper picture, A, shows the receiver plate with the line for stable lanthanum at mass number 139 (actually measured as the singly ionized oxide), and two other lines at 135 and 133. The middle plate,  $T_1$ , is the first transfer plate exposed for 24 hours after analysis showing that the lines at 133 and 135 are caused by radioactivities. The bottom plate,  $T_2$ , made by a 72-hour exposure following the first one, shows only the 19-hour La135 since the 4-hour La<sup>133</sup> had decayed.

Our thanks are due Mr. B. Rossi, Mr. J. Vale, and the members of the groups operating the 60-inch and 184-inch cyclotrons, and Mr. J. G. Conway for showing the CsNO3 used to be spectrographically pure.

\* The work described in this paper was performed under the auspices of the AEC.

a AEC.
\*\* Now at Princeton University, Princeton, New Jersey.
1 J. B. Chubbuck and I. Perlman, Phys. Rev. 74, 982 (1948).
2 W. Maurer, Zeits. f. Naturforschung 2a, 586 (1947).
3 Robertson, Carss, and Pool, Bull. Am. Phys. Soc. 24, No. 7, 14 (1949).

## Measurement of the Earth's Magnetic Field at High Altitudes at White Sands, New Mexico\*

S. F. SINGER Applied Physics Laboratory, Johns Hopkins University, Silver Spring, Maryland AND

E. MAPLE AND W. A. BOWEN, JR. Naval Ordnance Laboratory, White Oak, Maryland December 19, 1949

 $\mathbf{A}^{\mathrm{N}\,\mathrm{Aerobee^l}}$  rocket containing a total field magnetometer was fired at the White Sands Proving Ground, New Mexico (geomagnetic latitude 41° North) on April 13, 1948, at 1441 MST. The rocket reached a height of 372,000 feet (114 km) above sea level. Telemetered data gave the value of the earth's magnetic field during flight. The magnetometer measured the sum of the squares of three perpendicular components of the field so as to give an output independent of its orientation and thus obviate the need for stabilization of the instrument inside the rolling and precessing rocket. The magnetic effects of the rocket and instrumentation were quite small and were canceled by means of a system of compensating coils placed around the magnetometer. The residual error was mainly due to the inherent orientational error of the magnetometer itself and produced a roll-modulation of amplitude less than one milligauss. In analysis of the data, this error was averaged graphically over each roll period (about 1 second). A

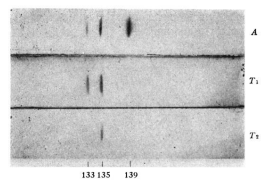


FIG. 1. Mass spectrographic analysis of radioactive lanthanum isotopes by transfer technique. (Exposures for transfer plates  $T_1$  and  $T_2$  given in the text.)