Lawrence Berkeley National Laboratory

Lawrence Berkeley National Laboratory

Title

The High Energy Spallation Products of Copper

Permalink

<https://escholarship.org/uc/item/5ns4v2xw>

Authors

Batzel, Roger E. Miller, Daniel, R. Seaborg, Glenn T.

Publication Date 1951-01-09

BERKELEY \vert CALIFORNIA $\frac{L}{O}$ JNIVERSITY

UCRL-

Mill

 $\frac{1}{2}$

 $\sum_{i=1}^{n}$

TWO-WEEK LOAN COPY

This **is a Library Circulating Copy which may be borrowed for two weeks. For a personal retention copy. call Tech. Info. Division, Ext. 5545**

RADIATION LABORATORY

VCR&-1077 Unolassified - **Chemistry** Distribution

UNIVERSITY OF CALIFORNIA

Radiation Laboratory

Contract No. **W-7405-eng-48**

THE **HIGE ENERGY** SPALJLATION **PRODUCTS** OF COPPER

Roger **E,** Batzel, Daniel R, Miller, and Glenn **T,** Seaborg

January 9, 1951

THE HIGH ENERGY SPALLATION PRODUCTS OF COPPER

Roger **E,** Batzel, Daniel R, Miller, and Glenn T, Seaborg Department of Chemistry and Radiation Laboratory University of California, Berkeley, California

January 9, **1951 ABSTRACT**

Nuclear reactions induced in elemental copper by irradiation with charged particles accelerated to energies in the hundred-Mev range have been studied, and the reaction products cover a range from the region of the target nucleus to a region which is removed from the target nucleus by as many as fifty or more nucleons, The yields of the various profiucts were measured, and the results show that a large majority of the products result from reactions in which only a fraction of the total energy of the incident particle is left with the nucleus, **The** distributfon of the amounts of the reaction products is in agreement **with** a pictxwe of high energy nuclear reactions involving **nuclear** transparency and the idea that the nuclear reactions involve excitation following collisions and energy transfers between the individual nucleons in the impinging particle and the individual nucleons in the target nucleus. **The** results include recent work on the nuclear reactions induced by 340-Mev protons and some earlier qualitative work on the nuclear reactions induced by 190-Mev deuterons and 190- and 380-Mev helium ions.

THE HIGH ENERGY SPALLATION PRODUCTS OF COPPER

Roger E. Batzel, Daniel R. Miller, and Glenn T. Seaborg Department of Chemistry and Radiation Laboratory University of California, Berkeley, California

Jmyary **9, 1959** INTRODUCTION

The acceleration of charged particles to energies in the hundreds of Mev range by the Berkeley **1%-inch** cyclotron has made it possible to study nuclear reactions in which the reaction products cover a range fromthe region of the target nucleus to a region which is removed from the tar- \mathcal{C} . get nucleus by as many as thirty or more nucleons.¹ The term "spallation" has been suggested to describe this type of transformation in which the .< excitation of the nucleus is degraded by emission of small nuclear fragments such **as** neutrons, protons, deuterons and alpha particles.

Some of the results of spallation reactions induced with high energy deuterons and alpha particles have been reported previously. **"5 The** present study consists of the determination of the radioactive products formed by the irradiation of elemental copper with 340-Mev protons. The data also include the results of a cursory investigation of the products formed by the irradiation of copper **with 190-Mev deuterons am3 190- and 388-Mev** helium ions.

The observed spallation products include some 35 puclides from sodium through zinc and **the** dfstributfon of the amounts of the reaction products formed is in general agreement with the picture of high energy nuclear. re actions described by $Serber.^6$ The observed yields show the effects of nu elear transparency and are consistent with Serber's idea that the nuclear reactions **involve** excitation following collfsfons and energg. transfers, . between the individual nucleons in the impinging particle and the individual nucleons in the target nucleus. As would be expected on this picture, the products,which are.formed by reactions requiring **me amomts** of excitation

occur most frequently,

The spallation data from the deuteron and helium ion irradiations of copper represent the results of some early qualitative work done in **1947** and 1948, At the present time this laboratory is not planning any immediate further investigations of these sgallation distributions, and the data are published here for qualitative consideration only,

PROCEDURE

Irradiations with the high energy particles **were** carried out in the circulating beam of the 184-inch frequency modulated cyclotron. The targets consisted of 6 and 10 mil copper foil about **15** nrm long and 10 mm vide. The copper strips were clanped in target holders which could be attached to the movable probe head of the cyclotron, and the bombarding energies **were** adjusted by fixing the radial distance of the leading edge of the target strip from the origin of the beam. Since the degradation of the energy of the high energy particles in passing through the 10 mil copper foil is *small,* the targets could be considered as thin targets.

Because of the fluctuations in intensity of the circulating beam, **%he** exact amount of **the irradiation** of the copper target **could** not be determined directly. The amount of the nuclide Cu^{61} formed during any given irradiation of the **copper was** determined and by relating the amounts of the other nuclides formed to the amount of Cu^{61} produced, relative yields of nuclides from **any** series of proton, deuteron, ox helim ion irradiation8 could be compared,

The length of the irradiations varied from ten minutes to one hour depending on the half-lives of tht. nuclides which were to be studied, **One** approximately 30 hour proton bombardment, and one approximately 15 hour deuteron bombardment were done to determine the yields of the nuclides with

 $-l_{\iota}$.

very long half-lives,

An absolute cross section for the formation of the Cu^{61} from copper irradiated with 340-Mev protons was determined by irradiating a piece of **10 mil. copper foil in the external beam of the 184=inch cyclotron. The** beam intensity was measured directly by means of a Faraday Cup arrangement. In the case of the deuteron irradiations, the absolute cross section for the formation of Cu^{61} was calculated using an estimated value of one microampere for the deuteron beam intensity.

 $-5-$

and the sense of the sense of the form of

After the target was irradiated, it was dissolved and measured amounts of carriers were added to the target solution, The elemental fractions were separated by the chemical separation procedures to be discussed in the section on "Chemical Separation Procedures." For the proton irradiations, the **amount** of carries remaining after the chemical separation of **an** elemental fraction was determined by chemical analysis and the percent of the carrier recovered was taken as a measure of'the percent **of** the radioactive atoms ofthe element recovered. Ho correction **was mde** for lasses during the chemical separations in the cases of the deuteron and **he-'** lium ion irradiations, but usually duplicate experiments were performed in the case of the deuteron study, and the higher yields were given greater statistical weight in the final average value of the yields.

Counting of the activitfes **was** done on **an** end-window, alcohol-quenched, ,t argon-filled Geiger counter tube with a mica window of \sim 3 mg/cm^2 thickness used in conjunction with a **scale** of **6.4** counting cfreuit, **The** nuclides were characterized by half-life determinations, absorption measurements, and sign of particulate radiation emitted.

A crude beta-ray spectrometer **was** used in the determination of the sign of the beta particles and **was** especially useful where it **was** necessary **to**

resolve nuclides with similar half-life but differing in sign of particulate radiation emitted.

 $-6-$

The counting data were corrected for window and air absorption, and in the cases where the yields are based on the counting of K x-rays, corrections were made for the fluorescence yields of the x-rays. Backscattering corrections were made only in the case of the determination of the absolute cross section for the formation of the cu^{61} from the proton irradiation of the copper, The samples from the proton irradiations were mounted on thin cover glasses **or** 5 mil aluminm discs and the backscattering corrections for such backings are essentially equal and constant for the energy range of the beta particles observed, The samples from the deuteron and lielium ion irradiations were mounted on stainless steel and platinum plates, and although the backscattering corrections are not equal, no correction was applied since the correction is not too serious and the deuteron and helium ion data are presented for qualitative consideration only, **No** correction **was** made for self-absorption since the weights of the samples counted were almost negligible in **all** instances,

RESULTS

A, Radioactive Nuclides Identified

The characteristic activities of the nuclides discussed in this section were observed **in** the elemental fractions separated from the irradiated copper target. References for most of the nuclides characterized may be found in the 1948 edition of "Table of Isotopes" by Seaborg and Perlman.⁷ Only in the cases where information on the anclides was published in the literature subsequent to the data given in "Table of Isotopes" will references be cited. All activities identified in the elemental fractions are discussed below and include the results from all bombardments (190-Mev

deuterons, 190- and 380-Mev helium ions, and 340-Mev protons). A check of Figs. 3, 4, and 5 and the results for 190-Mev helium ions listed in the "Results" section will show which individual activities were identified **in** each of the elemental fractions from the different bombardments.

 $\frac{\text{Zino Fraction}}{\text{Area}}$ The characteristic activities of Zn^{65} , Zn^{63} , and $2n^{62}$ were observed in zinc. The $2n^{65}$ was counted through sufficient aluminum absorber to cut out completely the particulate radiation and x-rays, and the counting efficiency of the 1,1-Mev gamma-ray was taken as one percent. The zn63 **was** characterized by its half-life **and** the **nuclide** was assumed to decay 93 percent by positron emission and 7 percent by orbital electron capture. . The Zn^{62} was assumed to decay 10 percent by positron. emission and 90 percent by K-capture⁸ and was counted with the 10-minute $~cm^{62}$ daughter in equilibrium.

Copper Fraction.-- The activities resolved in the copper were those of Cu^{64} , Cu^{62} , Cu^{61} , and Cu^{60} . The 12.8-hour Cu^{64} was the longest lived $\mathcal{L}(\mathcal{L})$ activity identified and **was** assumed to have a counting efficiency of 50 percent. The 10.5-minute $\ddot{c}u^{62}$ was the shortest lived activity identified and the nuclide was considered as decaying completely by positron emission, **Cn61** decays **66** percent by positron emission9 **and** 66 percent **was** taken as the counting efficiency of the nuclide. The 25-minute Cu^{60} was resolved with precision only in the copper fractions from the 190-Mev deuteron and **380-Mev** helium ion irradiations,

Nickel Fraction.-- The ody observable activities in **nickel** were the 2.6-hour M_1^{65} and the 36-hour M_1^{57} . The formation of the small amount of $N1^{65}$ from copper irradiated with protons is probably due to the reaction ~u65(n,~)~i65 caused **by** secondary neutrons **and** will be neglected as **s**

 $-7-$

spallation product from 340-Mev protons. A counting efficiency of 50 percent was used for the M_1^{65} since the nuclide decays 50 percent by positron emission and **50** percent by orbital electron capture, 10

 $-8-$

Cobalt Fraction.-- The characteristic activities of Co^{61} and Co^{55} were identified in all elemental fractions of cobalt separated. The Co^{58} and 0.056 were identified definitely in the proton irradiations and the characteristic 72-day half-life of these activities was observed in the cobalt fraction separated from the copper irradiated with 190-Mev deuterons, The Co⁶¹ decayed with the characteristic 1.75-hour half-life and since it decays by negatron (negative beta partiole) emission, no correction **was** necessary for the counting efficiency. The Co^{55} was easily resolvable and since no ...electron capture branching has bee reported, the counting efficiency was taken as 100 percent,

The fact that the Co^{58} and Co^{56} decay with equal half-lives and are both positron emitters makes their resolution difficult. The resolution, of the two activities **was** done only for the cobalt fraction separated from the copper irradiated with 340-Mev protons. The shape of the composite positron spectrum from the cobalt fraction was **determined** by **means** of a crude beta-ray spectrometer after the 18-hour $Co⁵⁵$ had been allowed to decay. The shape of the positron spectrum for a sample of pure $0°^{56}$ which had been determined under similar conditions was available, and by subtracting the contribution of the $Co⁵⁶$ positron spectrum from the composite positron spectmm, **the** contribution of each isotope to the total activity was obtained. The counting efficiency of the Co⁵⁸ was taken as 15 percent and that of the $Co⁵⁶$ as 100 percent.

Iron Fraction.- **me** characteristic activities of the radionuclides Fe⁵⁹, Fe⁵⁵, Fe⁵³, and Fe⁵² were probably present in all iron fractions

separated from the copper irradiated with the protons, deuterons, and a separated from the copper irradiated with helium ions. All of these activities were identified definitely only in . . . the iron fractions separated from the copper irradiated with 340 -Mev protons. The Fe^{59} , Fe^{53} , and Fe^{52} were identified in the iron fractions from the copper irradiated with deuterons, and the yield for the iron from **380-Mev** helium ion irradiation is based only on the Fe^{59} . The 2.9-year $\mathbf{F}e^{55}$ was assumed to decay completely by orbital electron capture¹¹ and a counting efficiency of 10 percent was taken for the **x-rays.** The **Fe**⁵³ was assumed to decay 100 percent by positron emission. The Fe^{52} was counted in equilibrium with its 21-minute Mn^{52} daughter. By determining the amount of 21-minute Mn^{52} in equilibrium with the Fe⁵² and milking the 5.8-day Mn^{52} which grows into the iron fraction, an estimation was made of the amount of, positron emission and orbital electron capture branching for the $Fe⁵²$. It was found that the nuclide 3ecays approximately **65** percent by positron emission and 35 percent **by orbital** electron capture.

-9-

Manganese Fraction.-- The radionuclides Mn^{56} , Mn^{54} , Mn^{52} , and Mn^{51} were identified in manganese. The Mn^{56} was assumed to decay completely . . . by negatron $emission.$ The $M⁵⁴$ was counted through sufficient aluminum \cdots absorber to cut out all the particulate radiation **an&** x-rays, **and** a **value** of **0.8 percent was assumed as the counting efficiency of the 0.8-Mev gamma**ray. Thirty-five percent was taken as the positron branching of the 5.8-day Mn^{52} , and the yield for manganese mass number 52 is reported on the basis of this activity. **The** Mn5' was assumed to decay completely **by** positron emission.

Chromium Fraction.-- The two activities identified in chromium were those of Cr^{51} and Cr^{49} . Since the Cr^{51} decays by orbital electron capture and gamma-my emission the counting efficiency of this nuclide **would** have had to be estimated roughly, but it was possible to determine the counting efficiency directly by milking cr^{51} from a known amount of Mn⁵¹ which had decayed. With the assumption that Mn^{51} decays 100 percent by positron emission, the Cr⁵¹ was found to have a counting efficiency of 2.6 percent when counted in a precipitate of barium chromate. The cr^{49} was assumed to decay with no orbital electron capture branching.

Vanadium Fraction.-- The yield for vanadium is based on **v4'** which was **assumed** to decay **58** percent by positron emfssion.

Titanium Fraction,-- The only activity identified in titanium was that of the 3.08-hour Ti^{45} and the yield was calculated on the basis of the nuclide decaying entirely by positron emission.

Scandium Fraction.-- Several activities were observed in the scandium fraction separated from the copper irradiated with 340-Mev protons, and these activities were identified as belonging to sc^{48} , sc^{47} , sc^{46} , sc^{44m} , and a mixture of the 3.9 -hour Sc^{44} and Sc^{43} . The 3.9 -hour and 85 -day activities were **easily** resolvable from the decay curves, but the resolution of the 2.44-day $\text{Sc}^{44,\text{m}}$, the 3.4-day Sc^{47} , and 44 -hour Sc^{48} was more difficult. The contribution of the 2.44-day Sc^{44} , with its 3.92-hour positron daughter in equilibrium, to the total activity was resolved approximately with the crude beta-ray spectrometer. With the positron activity resolved, the remaining activity was almost entirely that of the 3.4 -day $3c⁴⁷$ and the small **amount** of se4* activity resolved **was** so **uncertain** that the **yield** of this nuclide is not reported. The yield for Se^{43} includes the 3.9-hour Sc⁴⁴ and these nuclides were assumed to decay completely by positron emission. The yield for Sc^{44} was calculated on the basis of the 2.44-day

 $-10-$

^fsomer ,

Calcium Fraction.-- Only two activities were observed in the calcium fraction separated from the copper irradiated with 34.0-Mev protons, One was the 150-day $Ca⁴⁵$ and the other was a 4.8 \pm 0.2 day beta emitter with an energy of about 1.2 Mev as determined by an aluminum absorption measurement. This activity is probably the 5.8-day scandium activity reported as Ca^{47} by Matthews and Pool.¹² The growth of a 3.4-day scandium was observed in the decay of the calcium fraction and the scandium daughter was milked from the fraction. The **aluminum** absorption measurement of this 3.4-day scandium daughter showed it to be the activity assigned to $5c^{47}$. The decay curve of the calcium fraction is shown in Fig. 1, and Fig, 2 shows the decay of the calcium fraction counted through sufficient aluminum absorber to cut out the beta particles of the $Ca⁴⁵$ and the $Sc⁴⁷$ daughter.

 $-11-$

Chlorine Fraction.-- The characteristic activities of $C1^{39}$ and $C1^{38}$ were identified in all the chlorine fractions separated from the irradiated copper, but **el3!"** was identified **only** in the chlorine fraction separated from the copper irradiated with 340 -Mev protons. The resolution of the $C1³⁸$ and $C1³⁴$ was accomplished by determining the relative amounts of positronsand negatronspresent by means of the crude beta-ray spectrometer. The $~0.1^{39}$ was resolved directly from the gross decay curve. The $~0.1^{39}$, $~0.1^{38}$, and $c1^{34}$ were assumed to have counting efficiencies of 100 percent.

Phosphorous Fraction.-- The only activity observed in phosphorous was that of the 14.3 -day P^{32} .

Sodium Fraction.-- Two activities were observed in the sodium fraction. The 14.8 -hour Na^{24} was definitely identified, but the 2.6-year Na^{22} could

 $-12-$

UCRL-1077

be identified only on the basis of the decay during a 120-day period.

1. The state of B. Wields of Spallation Products of Camberland State of a Plots of the observed yields of the spallation products are shown in Fig. 3, Fig. 4, and Fig. 5. The yields are given relative to Cu^{61} which was arbitrarily assigned a yield of 1.0. The numbers listed for the various nuclides represent the ratios of the numbers of atoms of the particular nuclides to the number of atoms of the cu^{61} formed in the irradiated copper, and hence the numbers represent the ratios of the cross sections for formation from elemental copper, The cross section for the formation of Cu61 from elemental copper irradiated with 340-Mev protons is 1.7×10^{-26} cm² as determined by a bombardment of copper in the external proton beam where the beam intensity **was** accurately measured. The cross section for formation of Cu^{61} from copper irradiated with 190-Mev deuterons is about 6×10^{-26} cm². This value is an average value of about six different deuteron irradiations, and the value was obtained by using an estimated value (one microampere) for the beam intensity. The cross sections for the formation of $Cu⁶¹$ from copper irradiated with 190and **380-Mev** helium ions were not estimated since only one bombardment was done at each energy,

The data for the spallation products **from** copper **irradiated with** 190-Mev helium ions **are** not plotted since **ody** a few values were determined. The relative yields were Cu^{61} , 1.0; Cu^{64} , 0.84; $C1^{38}$, 0.0006; and no 0.1^{34} was found.

The sum of the yields for the nuclides $60⁵⁶$ and $60⁵⁸$ from copper bombarded with 190-Mev deuterons was estimated to be about 1.0 on the basis of a counting efficiency of about 15 percent for the combination of the activities. This is consistent with assuming that the activity **is**

essentially all C_0^{58} .

It should be emphasized that the yields for 340-Mev protons are in no **way** relative to the yields for 190-Mev. deuterons, **190-Mev** alpha particles or 380-Mev alpha particles, since the cross sections for formation of Cu^{61} from elementel copper is known only in the case of the 34.0-Mev proton irradiations. Many of the values for the reported relative yields are ϵ very dependent upon the counting efficiencies assumed (e.g., orbital and analyzed) electron capture), and when these efficiencies are better known the yields can be recdculated,

DISCUSSION

A. General Observations

The wide distribution and large number of radionuclides formed as spallation products of copper are immediately apparent from a study of Fig. 3, Fig. 4, and Fig. 5. The identification of Cu^{64} , Co^{58} , Mn^{54} , and Se^{46} , shielded from formation by decay, is good evidence for believing that the observed nuclides are mainly primary products from the splitting up of the excited nucleus, The radionuclides with the largest neutron deficiencies or neutron excess are formed in lowest yield, and the yield for a given **Z** rises for the nuclides nearest the region of stability, Extrapblation of this effect to the region of stability indicates that the stable nuclides are formed in high yield. Thus the emission of almost exclusively neutrons (or protons) from a moderately excited nucleus, followed by a series of rapid positron (or negatron) decays, is a relatively rare event,

The majority of the observed yield is found in the region of the target nucleus indicating that reactions **requiring** mch less than the, **maximum** amount of excitation available to the nucleus are more probable;

 $-13-$

UCRL-1077

 $-14-$

UCRL-1077

this is discussed in some detail in the next section,

It should be possible, knowing the general distribution of spallation products for a given **2,** to extrapolate and fnterpolate yields for the nuclides not directly observed as spallation products of copper, The data from the spallation of copper irradiated with 340-Mev protons have been treated in this manner as shown in Fig. 6. It was assumed that the most probable yield for a given **Z** is a region 2 mass units wide and that the yields lmass unit on either side of this region are formed in yields abount 1.5 tines **lower** than the values in the region of the maximum yield, that spallation yields 2 mass units removed are about ten times lower than the maximm, and that yields **3** mass units removed are about a factor of 20 lower than the maximum.

^Asummation of these extrapolated and interpolated yields for *340-Mev* protons gives a total yield relative to Cu^{61} of about 30, and using the measured value of 1.7×10^{-26} cm² for the cross section for formation of the Cu^{61} , the total spallation cross section for copper is about 0.5×10^{-24} cm². The geometrical cross section for copper is about 1.1 \times 10⁻²⁴ cm^2 . The discrepancy corresponds in part to the nuclear transparency at these high energies, but probably to a larger extent it is explained by the approximate nature of this method of estimation of the reaction cross section, The data in Fig. 6 **show** that about 80 percent of the spallation yields are concentrated in the elements copper, nickel, and cobalt. Outside the immediate region of the nucleus, the yield values in the region of most probable yield for a given Z are a decreasing function of Z indicating that nuclear reactions requiring very high exitations of the nucleus are much less probable than reactions requiring perhaps some **25** to **50 Mev** of excitation.

أناعت المعدوي فالتركية المتناب المتناوب والتوارد

B. Mechanisms of High Energy Spallation and the state of \mathbb{R}

 $-15-$

The accepted and experimentally supported theory of nuclear reactions at low energies (< μ 0 Mev) involves the formation of a compound, excited nucleusi the incident particle is captured by the target nucleus to form a compound nucleus with an excitation energy equal to the kinetic energy plus the binding energy of the incident particle, The excitation is then dissipated as a separate step by the evaporation of nucleons to form the product nucleus. Using only this picture of the compound nucleus, one would expect that bombardment with particles of several hundred Mev of energy **would lead** to very low yields of nuclides which are within a few mass units of the target nucleus, since emission of a large number of particles from the highly excited compound nucleus would be much more probable, **On** the contrary, however, the data presented in the preceding sections show that a large majority of the reactions induced in copper irradiated with 340-Mev protons, 190-Mev deuterons, and 190- and 380-Mev helium ions lead to products which differ from the target nucleus by a loss of only several nueleons . 明新 计微处理图象 克

Serber has suggested a mechanism which very satisfactorily explains the observed facts *.6* Irradiation with high energy protons, deuterons, and helium ions can be considered on essentially the same basis since bombardment with 190-Mev deuterons or with 380-Mev helium ions can be regarded **as.** a simultaneous bombardment by the several individual nucleons (neutrons and protons) making up the incident particle. The binding of the nucleons in the incident particle is important mainly in giving a spatial correlation between them. Serber points out that the collision time between a high energy incident nucleon and a nucleon in the nucleus is short compared to \ldots the time of collision of the nucleons in the nucleus, suggesting that

coUisions between incident nucleons and the individual nucleons **in** the **nucleus** are of primary importance.

 $-16 - 16$

Since the individual nucleon-nucleon-collisions are important, the high energy nuclear reactions can be interpreted in terms of the high energy scattering between free nucleons, Consideration of high energy scattering leads to two conclusions. First, at sufficiently high energies. the nucleus becomes partially transparent to the bombarding particles, and second. the incident particle loses only a fraction of its energy in the collisions. Serber estuated that the mean kinetic energy transfer to the struck particle by a 100-Mev nucleon is about 25 Mev and that the mean free path for this 100-Mev nucleon traversing nuclear matter is about 4×10^{-13} cm. Since the struck particles have a shorter mean free path than the incident one, they will **usually** distribute their **energy** to other nucleons through collisions, but it is possible that these struck nucleons **can** escape from the nucleus with little or no energy loss, The subsequent behavior of the excited nucleus cen be described in terms of an evaporation model, with the excitation energy dissipated by boiling off particles with several Mev of kinetic energy each.

The high spallation yields in the immediate region of the target nucleus are probably formed by reactions in which only single nucleonnucleon collisions take place. In the case of the deuteron and helium ion yields, this means that only one of the incident nucleons undergoes a collision in these reactions. The minimum energy transfer from a single nucleon-nucleon collision would take place when such a collision occurs near the edge of the nucleus, and the struck nucleon escapes from the nucleus with little or no energy transfer to the nucleus,

UCRL-1077

UCRL-1077

A larger portion of the energy of the incident particle may be left with the nucleus if multiple nucleon-nucleon collisions take place. Both. of the nucleons in the deuterons may undergo nucleon-nucleon collisions (or, in the case of the helium ion, two or three or four of its **con**stituents may undergo such collisions) and each of these impinging nucleons after such collisions may either escape or collide with additional nucleons in the target nucleus. This would lead in the extreme to products resulting from nuclei excited to almost the full energy of the impinging proton, deuteron, or helium ion, amounting essentially to the formation of \sim a compound nucleus in the ordinary sense. Thus the target may receive sense. excitation energy from about zero up to the full energy of the projectile.

It is difficult to estimate the energetic requirements for formation of spallation products such as $c1^{38}$ and Na^{24} . It is apparent from energetic considerations that $c1^{38}$ can not be produced in copper irradiated with 190-Mev deuterons by a reaction in which only single nucleons (neutrons and protons) are emitted. The threshold for formation of $c1^{38}$ from copper by the reaction $Cu^{63}(p, pn6a)Cl^{38}$ in which the maximum number of alpha particles emitted is roughly 110 Mev, but recent experimental results indicate that fragments larger than alpha particles are emitted among the competitive products of nuclear reactions, ¹³⁻¹⁵ and the energetic requirements for these reactions are even lower than fop alpha'particle emission. The observed yield for any given spallation product of copper probably actually represents the sums of tho yields of several types of nuclear reactions which form the given nuclide. The fact that the observed yields for $C1^{38}$ from 190-Mev deuterons and 190-Mev helium ions are about a factor of six lower than those from 340-Mev protons and 380-Mev helium ions (the relative yields for chlorine from the copper irradiated with deuterons and \sim

 $-17-$

 $-18-$

UCRL-1077

helium ions should be accurate to at least a factor of two) indicates that nuclides in the region of $c1^{38}$ are probably-formed mainly by reactions which require an appreciable part of the entire energy of the whole incident particle.

It is not difficult to visualige, in terms of an average energy loss of about 25 Mev per nuclear collision and a mean free path $(4 \times 10^{-13} \text{ cm}^2)$ of about 2/3 the nuclear radius for copper, mechanisms by which the 190-Nev deuterons, and 190- and 380-Mev helium ions might impart large portions of their total energy to the copper nucleus. It is difficult, however, to understand how a **340-Mev** proton could impart large amounts of excitation to the nucleus with a fair probability if it transfers in a slngle nucleonnucleon collision an average kinetic energy only slightly higher than the **value** of 25 Mev given by Serber for 100-Hev nucleons and has a mean free path about **squal** to the nuclear diameter for copper as estimated by Y amaguchi , **26**

In the light of some recent results of high energy scattering experiments.^{17,18} it is possible to actually estimate the mean free path of 3.40-Mev protons in nuclear matter and to' estimate the average kinetic **en**ergy loss of a 340-Mev proton in one individual collision with a nucleon in a copper nucleus and to see if the values estimated for the average kinetic energy loss and mean free path are more **in** line with high energy transfers having a fair probability,

The experimental data on proton-proton scattering at $340 ~\mathrm{Mev}^{17}$ show that the scattering cross section does not follow an expected **1/E** dependence, and that the cross section is considerably higher than would be expected if it varied as $1/E$. Also the results indicate that the cross section is practically isotropic between 0° and 90° in the laboratory system. This

UCRL-1077

means that high energy transfers in single proton-proton collisions have a relatively high probability. By extrabolating the data from neutron- \sim proton scattering with 260-Mev neutrons¹⁸ to 340 Mev and averaging the $$ cross sections for **11-P** and P-P scattering at the different scattering angles, it is possible to estimzte a mean energy loss for the 340-Mev protons of about 90 Mev per collision in a copper nucleus, On the basis of these scattering data the mean free path of a 340-Mev proton in nu-经附属标准 clear matter has been estimated to be about the same as that for a 95 -Mev nucleon, ¹⁹ bout 4×10^{-13} cm. On the basis of the mean free path of the 340-Mev protons being about 2/3 the length of the nuclear radius for copper, and on the basis of the mean kinetic energy transfer per single nucleon-nucleon collision being about 70 Mev, it is not difficult to understand how large amounts of excitation are imparted to the nucleus.

Recent calculations based on an evaporation model for the emission of the particles from the excited nucleus indicate that the emission of alpha particles from the excited nucleus should be a fairly probable occurrence and that the evaporation **3f** particles should lead to products along the **region of stability.** ^{20,21} The observed spallation yields are in agreement with these calculations since the yields apparently do have a definite maximum of most probable yield for a given Z about **2 2ss unite** wide and a line **drawn** along **this** region of most probable yield **would** be drawn through the **region of stability.** The very high yield of Mn^{52} compared to Fe^{-52} for the isobaric pair $\rm Fe^{52-}Mn^{52}$, apparent in both the 190-Mev deuteron and the 340-Mev proton yields, indicates that there is a much larger difference in the yields than **would** be eXqected from the *regular* trends of the spallation yields, and it seems logical to explain the difference observed in terms

 $-20-$

of alpha particles being boiled off from the excited target nucleus as suggested by Helmholz $_{et}$ al.⁵ If the $_{Fe}^{52}$ and Mn⁵² are formed by reactions initiated by non-capture processes (processes in which the proton or the deuteron loses energy to the nucleus but a proton is not retained by the nucleus), the reactions for the formation of Fe^{52} and Mn⁵² should perhaps be written $\text{Cu}^{63}(\text{p}, \text{pp6na})\text{Fe}^{52}$ or $\text{Cu}^{63}(\text{d}, \text{dp6na})\text{Fe}^{52}$ and $\text{Cu}^{63}(\text{p}, \text{p3n2a})\text{Mn}^{52}$ or $~\text{Cu}^{53}$ (d.d3n2a)Mn⁵². On this basis the yields indicate that the probability of emission of two alpha particles plus three neutrons is much higher than that for the emission of one alpha particle, six neutrons, and a proton. Although this effect shows up only in special cases like this where such direct comparisons can be made, it is probably a general phenomenon.

It is also possible on the basis of the spallation yields to say something about the probability of exchange reactions, for example, reactions in which the incident proton or the proton of the incident deuteron undergoes an electron exchange with a neutron in the copper nucleus and emerges as a neutron. The high yields of copper relative to the corresponding zinc isotopes .indicate that' these exchange reactions do not predominate.

The decreasing yields of individual nuclei far removed from the target nuclei is of course due in part to statistical considerations and the greater number of possibilities. **A** larger number of different combinations of emitted particles is possible from a highly excited nucleus than from a nucleus excited to a smaller extent. Thus, even if the probabilities of exciting a nucleus to say 200 and 100 **Mev** were **the** same, the yield of an individual product resulting from the higher excitation would be lower than one resulting from the lower excitation,

It is interesting to note that no observable yield of \mathbb{C}^{67} was obtained from copper irradiated with 380-Mev hel2-m ions. **This** nuclide can

24. 忧心

be formed only by an $(a, 2p)$ reaction on $Cu⁶⁵$.

CHENICAL SEPARATION PROCEDURES

 $-21-$

The irradiated copper foil was dissolved in hot concentrated nitric acid or hydrochloric acid and hydrogen peroxide, and milligram amounts of the elements calcium through zinc were then added to act as carrier for the elemental fractions. Known amounts of the elemental fractions to be separated were added so that a quantitative estimation of the amounts of \cdot the original carrier lost durlng the chemicd separation procedures **could** be made in the case of the proton bombardment and the macro amounts of the other elements. were added to act as holdback carriers. The chemical separation procedures included distillation, extraction, and precipitation operations.

In general the degree of radiochemical purification required depends on the relative spailation yield and the counting efficiencies of the radionuclides in the elemental fraction. If the nuclides are formed in **high** yield and have a high counting efficiency, the degree of radiochemical. purification need not be too high, but if the spallation yields **are** low or the counting efficiencies are low, the degree of purification must be a high,

Usually the particular elements to be investigated were separated successively from the entire dissolved target solution. Since different combinations of elements were removed in each investigation, the chemical separation procedures as a whole varied from bombardment to bombardment. The essential steps for the separation and purification of the elemental fractions were the **same** and the over-all. procedures differed only in the **order** in which the separations were used, For this reason only the important steps necessary for the chemical separation and purification of

UCRL-1077

the various elemental fractions are discussed, The final precipitates, if the **final** step in the chemical. separation procedure is a precipitation, were **usually** dissolved, and aliquota were taken from the solution for counting,

 $-22-$

 $Zinc$.-- The dissolved target solution with carriers added was adjusted to **2g** in hydrochloric acid and the copper precipitated. as the sulfide. The hydrogen sulfide was expelled and the solution neutralized with amnonium hydroxide, The solution was then adjusted to pH **3** with dilute sulfuric acid, and the **zinc** was then precipitated as the sulfide. The precipitate **was** dissolved ih dilute hydrochloric acid and the hydrogen sulfide expelled by boiling, Three milligrams of iron carrier was added and the solution adjusted to 1 M in sodium hydroxide. The ferric hydroxide was removed by centrifugation, and the ferric hydroxide precipitation **was** repeated. The solution was then adjusted **to UJ** in hydrochloric acid and the zinc precipitated by adding 4 ml of ammonium mercuric \circ thiocyanate, The zinc mercuric thiocyanate precipitate was washed with a solution containing ammonium mercuric thiocyanate and then transferred to the an aluminum plate to be dried and weighed in this form.

Copper.-- The dissolved target solution with carriers added was adjusted to 2M in hydrochloric acid and the copper precipitated as the sulfide. **The** sulfide precipitate was dissolved in. hot concentrated nitric acid, the sulfide expelled or oxidized, and.the nitrate destroyed by boiling **with** concentrated hydrochloric acid. The solution containing the copper was adjusted to $0.5N$ in hydrochloric acid, sulfur dioxide bubbled through the solution to reduce the copper, and then sodium thiocyanate added to precipitate copper as the euprous thiocyanate, The cuprous

UCRL-1077

thiocyanate was washed with water and then dissolved in nitric acid, The nitrate was again destroyed by boiling with concentrated hydrochloric acid, and the solution again adjusted to 0.5 ^N in hydrochloric acid. The copper

 $-23-$

was then precipitated in the final form as the cuprous thiocyanate.

<u>Nickel</u>. - The solution remaining, after the copper had been rem Nickel.-- The solution remaining, after the copper had been removed as the sulfide, **was** boiled to expel. the hydrogen sulfide and the iron oxidized with a few drops of nitric acid, The solution **was** then made alkaline with **ammonium** hydroxide and the precipitated hydroxides removed. . The precipitate was washed with a hot solution of ammonium chloride and the wash combined with the supernate from the precipitation. Three **mg** of iron carrier was again added and the ferric hydroxide removed by centrifugation. The solution was made slightly acid with acetic **acid** and **²**ml of a 1 percent alcoholic solution of dimethylglyox3me **was** added **tb** precipitate the nickel as nickel dimethylglyoxime. The precipitate was washed with water and then dissolved in concentrated hydrochloric acid, One to 2 **mg** portions of copper, cobalt, and-manganese were added to act **as** holdback carriers, the solution neutralized with ammonium hydroxide, and then made slightly acid with acetic acid, Dimethylglyoxime **was** again added to make sure that the precipitation of the nickel dimethylglyoxime **was** complete, and the precipitate was washed again **with** water. The nickel was weighed **as** the nickel dimethylglyoxime,

Cobalt.-- After the copper had been removed as the sulfide, the same as the set solution **was** boiled to remove the hydrogen sulfide and then neutralized with potassium hydroxide. The solution **was** adjusted to **33** in acetic acid, and 3 to 4 ml of a saturated solution of potassium nitrite acidified with acetic acid was added to the hot solution containing the cobalt, The

precipitate of potassium cobaltinitrite was allowed to settle in a steam bath, removed **by** centrifugation, and washed with a 5 percent potassium nitrite solution acidified with acetic acid. The precipitate was dissolved in hydrochloric acid, and holdback carriers for zinc, copper, nickel, and manganese were added, The solution was then neutralized with potassium hydroxide and the cobalt again precipitated as potassium cobaltinitrite. !he cobalt **was** weighed as the potassium cobaltinitrite,

 $-24-$

注意学

UCRL-1077

Iron.-- The solutfon of the copper target, with 5 mg of iron carrier added, **was** adjusted to 7,758 in hydrochloric acid and the iron extracted : with isopropyl ether. The ether layer was washed four times with 6 ml ... portions of 7.75M hydrochloric acid, and the iron was extracted from the ether layer with water. The solution was made alkaline with ammonium hydroxide and the ferric hydroxide precipitate separated by centrifugation. The precipitate was dissoLved **En** hydrochloric acid and the iron again extracted from 7.75 M hydrochloric acid with isopropyl ether. The ether layer was again washed with 7,758 hydrochloric acid and the iron extracted from the ether layer with water. The iron was analyzed by a colorimetric method.

Manganese.-- The solution of the target, to which 5 mg of manganese carrier **and** holdback carriersfor the other elements had been added, **was** adjusted to 2M in hydrochloric acid and the copper removed as the sulfide. The supernate **was** made alkaline with amnonfum hydroxide and the'alkalfne sulfides precipitated. The **sulfide** precipitate was dissolved in concentrated nitric acid and **fwning** nitric acid added to make the volume up to about 5 ml. Two or three crystals of potassium chlorate were added, and the solution was boiled gently for a few minutes to precipitate manganese

dioxide. The precipitate was removed by centrifugation, washed with water, and' then dissolved in one drop nitric acid and one drop of hydrogen peroxide. Holdback carriers for zinc, copper, nickel, cobalt, iron, and chromium were added and the manganese was again precipitated as **manganese** dioxide from concentrated nitric acid, The mangaaese was analyzed by weighing as manganese dioxide.

Chromium.-- The steps for the* chemical separation of chromium were the same as those given for the sepaxation of the manganese down to the point where the manganese was first removed as manganese dioxide. At this point the chromium was in the form of the chromate and remained in the supernate after the manganese dioxide was removed by centrifugation, The nitric acid solution containing the chromate **was** adjusted to **0.2g** in nitric acid and the solution cooled in an ice bath, **Two** tothree drops of 30 percent hydrogen peroxide were added to form the blue peroyychromic acid, and the peroxychromic acid **was** extracted with diethy1 ether. The ether layer was washed twice with 5 ml portions of water containing a drop of nitric acid. The peroqchromic acid was **removed** from the ether layer by adding a solution of **O.5g** sodium hydroxide which breaks down the peroxychromic acid and forms the chromate, The ether was removed by evaporation, **and the** excess hydrogen peroxide was destroyed by boiling the solution. **The** solution was made slightly acid with acetic acid, and the barium was precipitated as the chromate by adding barium chloride dropwise to the hot solution of **the** sodium chromate. The chromium was weighed **as** barium chromate,

Vanadium,-- The vanadium was usually separated just subsequent to **the** extraction of the chromium as the blue peroxychromic acid as described in

 $-25-$

-26- UCRL-1077

the preceding section on chromium. The solution containing the vanadium was almost neutrdized with sodim hydroxide and then was poured into 10 ml of a hot solution of lM sodium hydroxide. The hydroxide precipitate was removed by centrifugation and then washed 'with 4 ml of hot **1i** sodium hydroxide, Two mg each of iron and titanium carrier were then added to the solution and the hydroxides precipitated, The supernatant solution containing the vanadium was boiled to reduce the volume and then made slightly acid with acetic acid. Lead acetate **was** added to precipitate the vmadiun as **lead** vanadate, and the precipitate was washed with a dilute solution of lead acetate made slightly acid with acetic acid. The lead vanadate was dissolved in nitric acid and the solution adjusted to 2N in nitric acid. The lead was precipitated as the sulfide, and the sulfide was expelled by boiling the solution. Five **mg** of chromium carrier was added and sulfur dioxide was passed through the solution to reduce the vanadium and chromium. The solution was almost neutralized with sodium hydroxide **and** then poured into a boiling solution of **1N** sodium hydroxide to precipitate the chromium hydroxide. The chromium hydroxide was removed by centrifugation, and the vanadyl ion was oxidized to the vanadate state with hydrogen peroxide. Vanadium was then precipitated as lead vanadate from a solution slightly acid with acetic acid. The vanadium was analyzed colorimetrically.

Titanium.-- After the copper had been removed as the sulfide, the solution with the carriers added was boiled to remove the hydrogen sulfide, and the solution was made alkaline with ammonium hydroxide to precipitate the hydroxides of iron, scandium, and **titanium.** The precipitate was washed with hot, dilute ammonim hydroxide and then dissolved in hydrochloric acid. The solution was then adjusted to **0,53** in hydrochloric acid and scandium precipitated as the fluoride by adding 0.3 ml of 27N hydrofluoric acid.

after **allowing** the precipitate to settle for ten minutes on a steam bath, the scandium fluoride was removed by centrifugation, and the supernatant **was** made alkaline with ammonium hydroxide. The combined titanium and ferric hydroxide precipitate **was** removed, dissolved in nitric acid and adjusted to **1.68** in nitric acid, Five **ml** of a solution of potassium iodate **was** added and the titanium separated as potassium titanium iodate. The precipitate was washed with a dilute solution of the potabsium iodate. The iodate was destroyed by bubbling sulfur dioxide into the precipitate mixed with a few ml of dilute hydrochloric acid. Ammonium hydroxide was added to precipitate the titanium hydroxide, the precipitate was removed by centrifugation and then dissolved in hydrochloric acid. Five **mg** of scandium carrier **was** added, and the scandium **was** precipitated as scandium fluoride from a 0.5M solution of hydrochloric acid. The titanium was again precipitated as the hydroxide and the precipitate dissolved in nitric acid. The solution was again adjusted to 1.6^M in nitric acid and the potassium titanium iodate again precipitated, The titanium **was** analyzed colorimetrically,

Scandium.- The steps in the separation procedure for scandium are the same as those for titanium down to the point where the scandium fluoride is removed. The fluoride precipitate was washed with 0.5 M hydrochloric acid containing one drop of 27N hydrofluoric acid. Sulfuric acid was used to: dissolve the scandium fluoride and the fluoride ion was removed by fuming the scandium solution with *a* **small amount** of concentrated sulfuric acid. The sulfuric acid solution was made alkaline with ammonium hydroxide and **the** scandium hydroxide precipitate removed by centrifugation. The hydroxide was dissolved in hydrochloric acid **anci** the solutian adjusted to **0,5J** in hydrochloric acid. Scandium fluoride was then precipitated by adding

-28- UCRL-1077

0.3 ml of 27N hydrofluoric acid, and the precipitate was allowed to settle on a stem **bath,** The precipitate was again dissolved in sulfuric acid, **tke** hydrogen fluoride distilled, and scandium hydroxide precipitated from an alkaline ammoniwn hydroxide solution, The scandium was weighed as scandium oxide .

Calcium.-- After the copper had been removed, the solution containing 5 **mg** of calcium carrier and holdback carriers for the other elements was **made** alkaline with mnomzim hydroxide, and the alkaline sulfides and hydroxides were precipitated. Holdback carriers for zinc through scandim were again added, and the alkaline sulfides and hydroxides were again precipitated. This last step was again repeated, and the solution was then boiled to remove the hydrogen sulfide. The solution was made slightly acid with oxalic acid and 5 ml of 4 percent ammonium oxalate was added to precipitate calcium oxalate. The precipitate was allowed to settle on a steam bath for ten minutes, was removed by centrifugation, and washed with water containing oxalic acid and ammonium oxalate. The calcium oxalate was dissolved in concentrated nitric acid, and a few crystals of **potassium** chlorate were added to oxidize the oxalate ions. Three mg of iron carrier was added, and the solution made alkaline with ammonium hydroxide. The ferric hydroxide was removed by centrifugation and the solution made acid with oxdic acid. Four *rnl* of a solution **L+.** percent in **ammonium** oxalate was added to precipitate calcium oxalate as the final purification step, and the precipitate was allowed to settle on a steam bath. The calcium was weighed as calcium oxalate.

Chlorine.-- The copper was dissolved in nitric acid and 5 mg of chlorine in the form of sodium chloride was added. The solution vas boiled to distill the hydrogen chloride and the gas was caught in a solution containing

two to three drops of nitric acid and sufficient silver nitrate to precipitate the chlorine as silver chloride. The solution was heated almost to boiling to coagulate the precipitate, and it **was** removed by centrifugation, The precipitate was washed with dilute nitric acid and then dissolved in dilute ammonium hydroxide. The solution was then acidified with nitric aeid and additional silver nitrate added to make sure that the precipitation of the silver chloride was complete. **The** chlorine was weighed as silver chloride,

 $-29-$

Phosphorous,-- The copper target **was** dissolved in nitric acid **md** 5 **mg** of phosphorous as phosphate ion was added to the solution. Holdback carriers were also added for the other elements zinc through calcium, The solution was adjusted to **la** in nitric acid and 10 ml of monim molybdate **was** added to the warmed solution to precipitate ammonium phosphonolybdate. The precipitate was allowed to settle for **15** minutes, removed by centrifugation, and washed with a solution of **1** percent nitric acid. The precipitate was dissolved in ammonium hydroxide containing citrate ions to complex titaniuq that interferes, Hydrochloric acid **was added** until the precipitate that formed dissolved **with** difficulty and *5* **mg** each of vanadiwn and titanium carrier were added, Sulfur dioxide was bubbled through the solution to reduce all the vanadium to the vanadyl state. Two ml of cold magnesium chloride was added, and the solution was allowed to stand for five minutes. Concentrated ammonium hydroxide was then added equal to one quarter of the original volume and the precipitate **was** allowed to settle for 15 minutes, The precipitate was removed by centrifugation and then washed with **3g** ammonium hydroxide, The precipitate was dissolved in **1i** nitric acid, and the phosphorous was again precipitated as the ammonium phosphonolybdate. The phosphorous was

weighed as the ammonium phosphomolybdate.

Sodium.-- The copper **wsls** dissolved in concentrated hydrochloric acid and hydrogen peroxide, and 5 mg of sodium chloride was added as carrier. Carriers for the elements zinc through potassium were added and the solution adjusted to 2M in hydrochloric acid and the copper precipitated as the sulfide. The solution was boiled to drymess to remove the excess acid, the residue dissolved, and then the solution made alkaline with ammonium hydroxide. Hydrogen sulfide was added and the sulfides and hydroxides were precipitated. Additional 3 mg portions of carriers zinc through scandium were added and precipitated. This scavenging process **was** repeated twice. Excess hydrochloric acid was added to the alkaline sulfide solution, the acidified solution boiled to dryness, and the ammonium chloride driven off, The residue **was** dissolved in water, the solution checked to make sure it was neutral, and the sodium precipitated as the sodium zinc uranyl acetate. Two 5 ml portions of zinc uranyl acetate were used to wash the precipitate, and the precipitate was then dissolved in absolute alcohol saturated with hydrogen chloride gas. The resulting mixture was cooled in an ice bath and the sodium chloride **separated** by centrifugation. The sodium chloride precipitate was washed with a *5* ml portion of the alcohol-hydrogen chloride solution. The sodium chloride was dissolved in water and the solution neutralized with potassium hydroxide. Zinc uranyl acetate **was** added to precipitate tho sodium, the precipitate removed by centrifugation, and then washed with additional zinc uranylesses are moved by centrifugation, and then washed with additional zinc uranylesses acetate, The precipitate **was** dissolved in absolute dcohol saturated with hydrogen chloride, the mixture cooled in an ice bath, and the sodium chloride removed by centrifugation, The sodium chloride was washed with an

-30-

UCRL-1077

Dealer State Company of Carry Care

ทั่วที่ไป และเปล่า ได้เป็นสมุทธศึกษณะรับทาง เป็นไปแบบที่ หรือกับไปร

 $\label{eq:2} \mathcal{F}^{(1)}_{\text{max}} = \mathcal{F}^{(1)}_{\text{max}} + \mathcal{F}^{(2)}_{\text{max}} + \mathcal{F}^{(3)}_{\text{max}} + \mathcal{F}^{(4)}_{\text{max}} + \mathcal{F}^{(5)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6)}_{\text{max}} + \mathcal{F}^{(6$

them are a communication whereas with path

 $\mathcal{L}^{\mathcal{A}}=\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A}}\mathcal{L}^{\mathcal{A}}+\mathcal{L}^{\mathcal{A$

start of the seal of the first response to a good control of the seal of the seal of the seal

Service State

additional portion of the ethyl'alcohol-hydrogen chloride solution, The sodium was weighed as sodium chloride,

ACKNOWLEDGMENTS

We wish to thank Mr. James **T,** Vale **and** the group operating the l&-inch cyclotron for the irradiations performed in the course of these studies. This work **was** performed under the auspices of the U. **S,** Atomic Energy Commission.

at any conservation of the company of the conservation and the

April 1998 - Francisco Maria Maria (1989), Antonio Maria (1999), and a finite state of the Maria (19

and the control of the control of the control of the second control of the control of the control of the control of

 $\label{eq:2.1} \frac{1}{2} \left(\left(\frac{1}{2} \right)^2 - \frac{1}{2} \left($

1. 经工作人员 医单位的 医单位

particularly application of the con-

 $\label{eq:2.1} \mathcal{L}(\mathcal{A}) = \mathcal{L}(\mathcal{A}) = \mathcal{L}(\mathcal{A}) = \mathcal{L}(\mathcal{A}) = \mathcal{L}(\mathcal{A})$

state of the community of the first state of the community of the community

a di Santa Maria (1918).
A di Santa Maria (1918) na mata mata sa mata m

good and the computation of the

a sa kacamatan ing kabupatèn Kabupatèn Kabupatèn Kabupatèn Kabupatèn Kabupatèn Kabupatèn Kabupatèn Kabupatèn K

at 1998 e destacados de la producción de la final de la contexta de la final de la producción de la caracterís
Constituídas de la contexta de la c

网络加速器 医二次原子体 **BIBLIOGRAPHY**

1. B. B. Cunningham, H. H. Hopkins, M. Lindner, D. R. Miller, P. R. O'Connor, I. Perlman, G. T. Seaborg, and R. C. Thompson, Phys. Rev. **72, 739, 740 (1947) (A).**

 $-32-$

- 2. D. R. Miller, R. C. Thompson, and B. B. Cunningham, Phys. Rev. $\frac{7}{4}$, 347 (1948)
- 3. H. H. Hopkins, Jr., Phys. Rev. 77, 717 (1950).
- *⁴***34.** Lindex and I. Perhian, Phys, Rev. **78,** 499 (1950)~
- 5, F. **D.** Bartell, **A. C,** Helmholz, **So** D. Softky, and **D.** D. Stewart, **U. So** Atomic Energy Commission Unclassified Document UCRL-757 (July, 1950).
- *6.* R. Serber, Phys. Rev. 72, 1114 (1947).
- 7. G. T. Seaborg and I. Perlman, Rev. Mod. Phys. 20, 585 (1948).
- 8, R, **W.** Hayward, Phys. Ret.. **s, S?** (1.950) **(A).**
- 9. G. E. Owen, C. S. Cook, and P. H. Owen, Phys. Rev. 78, 686 (1950).
- 10, **G.** Friedlander and **N,** L, Perlman, Brookhaven National Laboratory Report **51** (1950) ,
- 11, **G. L, Brownell, M.** I, **T,** Prog. Report 37 (1949).
- **12,** D, E, Matthews and M. L. Pool, Phys, Rev. *2,* **163** (1947).
- 1, S. Wright, Phys. Rev. 77, 742 (1950) **(A).**
- . L. **Marquez** and I. Perlman, *2;o* be published in Phys. Rev.
- 15. R. E. Batzel and G. T. Seaborg, Phys. Rev. 79, 528 (1950).
- 16. Y. Fujimoto and Y. Yamaguchi, Prog. Theor. Phys. 5, 141 (1950).
- 37, **0, Chamberlain** and C, Wiegand, Phys. Rev, *B,* 81 (1950).
- 18. E. Kelly, C. Leith, E. Segre, and C. Wiegand, Phys. Rev. 79, 96 (1950).
- 19. S. Fernbach, private communication.
- 20. K, J. Le Couteur, Proc, Phys. Soc, **A a,** ²⁵⁹**(1950).**
- 21. Y. Yanagxchi, Prog. Theor. Php. 5, **I42 (1950).**

LIST OF ILLUSTRATIONS

FIGURE

- **1, Decay** of c3lcium fraction separated from copper irradiated **with**
- **340-Mev** protons . 2. Decay of calciun fraction separated from copper irradiated with 340-Mev protons counted through sufficient absorber to cut **out** the beta particles of ca^{45} and the $3c^{47}$ daughter.
- *3,* Experimental yields from copper irradiated with **340-Mev** protons. Yields are relative to Cu^{61} .
- 4. Experimental yields from copper irradiated with 190-Mev deuterons. Yields are relative to Cu^{61} .
- 5. Experimental yields from copper irradiated with 380-Mev helium ions. Yields are relative to Cu^{61} .

 $-33-$

6, Gxperimental, interpolated, md extrapolated yields fron copper irradiated with 340-Mev protons. Yields are relative to cu^{61} .

Fig. 2

Fig. 3 *

 $Fig. 4$

Fig. 5

MU1223

Fig. 6