ENERGY SPECTRA OF FRAGMENTS FROM SILVER AND URANIUM BOMBARDED WITH 5.0 GeV PROTONS

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ABSTRACT

In this third paper in a series in which the characteristics of nuclear fragments produced in the interaction of 5 GeV protons with Ag and U targets were studied by means of dE/dx-E measurements with semiconductor detector telescopes new information was obtained on the energy spectra of light fragments. One set of measurements on fragments from a Ag target involved the use of a two-element telescope incorporating a ΔE detector as thin as 16 μm. A new algorithm for processing the ΔE and E data to extract particle identification was developed and the resulting particle spectra showed superior resolution for the elements from Li (Z = 3) to S (Z = 16). Segments of the energy spectra of each of these elements were measured at 20° and, for many of them, also at 45°, 90°, 135° and 160° to the beam direction. By use of 3-element telescopes and absorbers the high energy part of the energy spectrum for isotopes of He, Li, Be, B, and C ejected from Ag and U targets was measured at 20°. The measurements extended
beyond 300 MeV for $^6$Li and $^7$Li and to 400 MeV for $^7$Be. A distinct high-energy component was found in these cases.

The suitability of nuclear evaporation as a description of the emission of the low-energy fragments was tested with two simple theoretical models, one specifying isotropic fragment emission from a moving nucleus at a fixed nuclear temperature and one specifying isotropic fragment emission from a set of moving nuclei with a Maxwellian distribution of excitation energies and forward momenta. The second could describe rather well all the 90° data, provided a Coulomb barrier 0.4 that of the classical tangent spheres barrier was used. However, the measured intensity in the forward direction was much higher than predicted. Neither evaporation calculation was able to describe the highest energy part of the spectra, and the conclusion was drawn that these particles must be produced in the initial high-energy cascade.
I. INTRODUCTION

Fragments ejected from complex nuclei during the course of reactions induced by GeV protons can be studied with silicon semiconductor detectors incorporated in a particle-identifier system. Comprehensive studies of the characteristics of fragments ejected from uranium and silver targets bombarded by 5.5 GeV protons in the Bevatron have recently been published.\textsuperscript{1,2} The results described in the present paper are supplementary to these previous studies and extend them in two ways.

First, in the case of the silver target study\textsuperscript{2} the data have been extended and improved in the low energy region. For the reasons given in the discussion of the cited papers,\textsuperscript{1,2} it is important to determine the maximum in the energy spectrum of each of the light elements, Li, Be, B, etc. This maximum is determined by the effective Coulomb barrier felt by the fragment, and it is one of the characteristic features of the fragment ejection process that this effective Coulomb barrier is only 40-50 percent as high as the barrier obtained from a tangent spheres calculation. In the previous work the experimental low-energy cut off, which was fixed by the 20 micron thickness of the $\Delta E$ silicon detector, was so high that it exceeded the maximum in the energy spectrum for all fragments except He and Li. In the new study, a 16$\mu$ $\Delta E$ detector was used and, as a consequence, the turnover region of the energy spectrum was defined for Li, Be, and B. Measurements were made at 20°, 45°, 90°, 135°, and 160° to the beam. A special run was made at 20° to improve the identification of elements up through $Z = 18$ (argon), and a section of the energy spectra of these elements was determined. No attempt was made to remeasure the low energy spectra of fragments from uranium because the turnover regions of the spectra were well defined for Li, Be, and B fragments in the original study.
The second way in which the previous data were extended was in the remeasurement of the ultra-high energy portion of the fragment spectra. One of the interesting features of the previous work was the identification of Li and Be fragments with energies extending out to 160 MeV. In the new study these high-energy fragments were restudied in order to establish beyond question the emission of these ultra-high energy fragments and to measure the energy spectrum out to even higher energies. For this purpose measurements were made on fragments from uranium and silver targets with a three-element semiconductor telescope with a total Si thickness of 1100 to 2200 microns (257 to 513 mg/cm² Si). Aluminum absorbers of thickness ranging up to 1.3 g/cm² were placed in front of the first detector. In those experiments the energy spectra of isotopes He through C were measured at 20° to the beam. In the case of ⁶Li and ⁷Be the energy spectra were measured out to 340 MeV and 420 MeV, respectively.

Section II describes the experimental techniques for data taking and reduction. Section III presents the results of the restudy of the low energy portion of the energy spectra as well as a discussion of these results. The high energy data are presented and discussed in Section IV. Section V summarizes the main conclusions of the paper. Appendix A is devoted to a new method of determining particle identity from ΔE and E detector signals. Appendices B and C cover mathematical details on two evaporation formulas used in the analysis of the energy data.
II. EXPERIMENTAL

Inasmuch as the experimental techniques were nearly the same as those described in Ref. 1, they are presented here in shortened form except for details of the detectors or for changes from the earlier methods.

A. Bombardment Arrangement at Bevatron

Thin targets of metallic Ag or U were placed in the center of a 36 inch diameter target chamber located in one of the external beam lines of the Bevatron. This chamber could be isolated from the beam line by gate valves in the upstream and downstream ports. A mechanical pump system was provided for evacuation of the chamber to the 5-10 micron level, which was the normal pressure for the beam line. It was necessary to achieve a better vacuum for operation of the 2000eV detectors used in the second part of the study because bias voltages up to 1500 volts were needed. This better vacuum was obtained by installing 0.003 inch aluminum windows beyond the inlet and outlet valves and by pumping the chamber volume with a cryogenic pump. With this arrangement a much lower vacuum was maintained in the chamber and no problems with sparking or voltage breakdown occurred.

The silver target for the low energy experiments was prepared by volatilization of metallic silver to make a uniform foil of 1.1 mg/cm². This was glued across a hole in a 0.00025 inch mylar sheet which in turn was stretched across a 3.5 x 6 inch hole in an aluminum frame. The target was positioned at 55° to the beam for fragment measurements made at 20°, 45°, and 90°. It was placed at 135° for measurements at 135° and 160°. For measurements of fragments of higher energy silver foil targets with uniform thicknesses of 7 mg/cm² and 26 mg/cm² and a uranium foil target with a thickness of 27.5 mg/cm² were used in similar target frames.
The external beam of 5.0 GeV energy was delivered in pulses of 800 to 1000 msec duration repeated every 6 seconds and containing about $8 \times 10^{11}$ protons per pulse. The beam cross section at right angles to the beam integrated over a period of hours was determined by radioautographs to be roughly 1/2 inch vertically by 1-1/2 inch horizontally. The cross section of the target foils was somewhat larger.

The telescope of silicon semiconductor detectors was mounted on an aluminum block which in turn was clamped to a movable arm whose pivot point was at the chamber center.

B. Detector and Electronic Details for Measurement of Low Energy Fragments

The telescope had two elements: a 16 μm thick surface-barrier detector used as the ΔE detector and a 19 1/4 μm thick planar detector of phosphorus-diffused silicon used as the E detector. Behind these there was also a 381 μm thick planar detector of phosphorus diffused silicon used as the rejection detector to eliminate the recording of high-energy fragments which were not stopped in the E detector. Collimators with 5 x 7 mm openings cut into 0.125 inch copper were placed in front of the ΔE and the E detector. The distance between the two collimators was 2.8 cm. (In the 3-element telescope described in Sec. II.C. the collimators were positioned in front of the first and third detectors and were separated by 2.1 cm.)

The electronic circuitry was identical with that illustrated in Fig. 1 of Ref. 2. In brief, this circuitry accomplished the following purposes. Pulses from the ΔE and E detectors were not accepted unless they were received within a 50 nsec resolving time. If a pulse was simultaneously recorded in the
The reject detector event was discarded. A pile up rejector circuit inspected
the preamplifier output and excluded any events in which a second pulse was
received within an inspection period of 600 nsec. The output pulses from the
main amplifiers were passed through single channel analyzers which accepted all
pulses above 1.5 MeV in the ΔE detector and 2 MeV in the E detector. Pulses
surviving these tests were accepted for digitization by an analog-to-digital
converter interfaces to a PDP-8 Computer and were written on magnetic tape in
blocks of 128 events. These records were later processed at a larger computer
as discussed below. The surviving ΔE and E pulses were also transmitted to a
Goulding-Landis particle identifier system for analog development of a particle
identifier signal which was sent to a pulse height analyzer for recording and
display. The on-line display of a histogram of the analog particle identifier
signal was used for monitoring and trouble-shooting purposes during the data-
taking portion of the experiment, but all data presented in this paper resulted
from off-line computer calculations with the original digitized pairs of ΔE
and E pulse heights, as discussed in Appendix A and Sec. II.E. As discussed in
Ref. 1 precision pulsers were used for the energy calibration.

C. Detector and Electronic Details for Measurement of High Energy Fragments

The three different 3-element telescopes used in this part of the experiment
are listed in Table I. The ΔE and E detectors were all manufactured of
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phosphorus diffused silicon. The 1000μ and 2000μ E detectors were silicon surface-
barrier detectors. The 1500μ E detector was made of lithium drifted silicon.
It was our experience that lithium drifted detectors could be used only
for a short time (12 to 24 hours) in close proximity to the beam before radiation
damage effects, probably involving precipitation of lithium, caused deterioration
of the detector signals, particularly the timing characteristics of the signal. Data whose quality was seriously affected by these radiation damage effects were discarded. The surface barrier detectors also suffered radiation damage in this hostile environment, manifested mainly as a steady increase in leakage currents which rose steadily from a few μamps up to more than 50 μamps over several days of exposure. However, the energy and time characteristics of the pulses were not seriously affected and the quality of the data remained good.

The electronic circuitry associated with the 3-element telescope was identical with that displayed in Fig. 1 of Ref. 1 and fully discussed in that paper. Coincidence requirements, pile up rejection, and pulse height discrimination were in principle the same as those briefly mentioned in B above. One change in the use of the pile up rejector circuitry was the elimination of inspection of the E detector pulse because the long collection times for ionization pulses in these thick detectors produced preamplifier output pulses too slowly rising for the use in the pile up rejector. Also the role of the analog particle-identifier system was more important for these telescopes incorporating two ΔE detectors. By analog circuitry a particle identify pulse was generated from the ΔE₂ and the E = ΔE₁ + E pulses and compared with a second identity pulse generated from the ΔE₁ and E inputs. If the two identifications did not agree within a preset amount (in most cases 15%) the event was rejected and the ΔE and E signals were not sent to the computer for permanent recording. Not many events were rejected by this requirement (< 10%) but the quality of isotopic resolution was improved. The final particle identification was made off-line by use of the ΔE₂ and ΔE₁ sum as the ΔE value in a calculation performed according to the method described in Appendix A.
The total thickness of silicon in the detector stacks listed in Table I is not sufficient for exploring the range of fragment energies of interest in this study. We could have increased this thickness by using thicker $\Delta E$ detectors and thicker and more numerous $E$ detectors, but this would have introduced greater complexity in the electronic circuitry as well as special difficulties with the coincidence timing, and with the corrections for the dead layers on the extra detectors. We decided instead to extend the measured energy range by placing various thickness of absorber directly in front of the first detector in the telescope. Four absorbers were machined of aluminum stock with thickness 0.0052, 0.063, 0.100, and 0.189 inches (35.7, 432, 690 and 1295 mg/cm$^2$, respectively).

These thicknesses were chosen after detailed calculation of the expected energy losses for different particles in the absorber and in the detectors. A computer program called LAZY written by Dr. Creve Maples was modified for this purpose. In our version of the program the range relationships of P. G. Steward were incorporated. These calculations helped greatly in the choice of absorbers and detector thicknesses, energy thresholds for the detectors, and other parameters in the experimental setup.

Consideration was given to the possibility that the thick aluminum absorbers would cause multiple scattering of the fragments sufficiently great to affect the intensity of measured fragments. However, the fact that the absorber was mounted directly in front of the collimator attached to the front of the first $\Delta E$ detector greatly reduces the change of loss by scattering and even in the worst case, which is the scattering of Li by the 189 mil absorber, a calculation showed that the effect is negligibly small (< 1%).
The possibility of an appreciable reaction of the fragments with the aluminum absorber was also considered. For a reaction cross section assumed equal to the geometrical cross section for fragments above the Coulomb barrier, it was estimated that 5 percent or less of the fragments reacted during passage through the thickest absorber. This is within the experimental error for measurement of fragment intensity.

Table I summarizes the specific combinations of telescope and absorbers used to obtain the data discussed in Sec. IV.

The integrated beam through the target was measured with a monitor telescope identical to that described in Refs. 1 and 2. The factors needed to convert the ratio of counts in the fragment and monitor telescopes into an absolute reaction cross section were obtained from these earlier studies. The ultimate standard of intensity was the $^7$Be cross section determined by radiochemistry as discussed in the appendix of Ref. 1. In the runs with the Ag target and the 16 μm $\Delta E$ detector the energy windows in the monitor telescope were incorrectly set and the normalization was done in a more indirect manner; the $^7$Be spectrum at 90° was normalized to the corresponding spectrum of Ref. 2 and the resulting normalization factor was applied to all data from these runs.

D. Reduction of $\Delta E$ and E Data

**Generation of Particle Spectra.** The magnetic tapes with the event-by-event record of $\Delta E$ and E signal information were processed by the method described in Appendix A. A computer program called PICAL was used.

**Generation of Energy Spectra.** The output tape of the PICAL program, containing the summed data on PI vs $E_T$ in the form of a $1024 \times 1024$ matrix, was used as input to another program (SAVEX) which performed the following functions.
The data were sorted by PI windows, which defined specific particles, and the energy data for these selected particles were isolated. At this point the energy scale was an arbitrary one set by the gains in the amplifiers and by the analog-to-digital converter. The absolute energy scale for energy deposited in the detectors was computed by use of data taken with a calibrated pulser. The fragment energies were then corrected as in Ref. 1 for the losses incurred in escaping from the target and in passing through the thick aluminum absorbers. This correction was based on a power-law, range-energy relationship, \( R = aE^b \) with sets of parameters \((a \text{ and } b)\) fitted to different energy ranges as described by Bichsel and Tschalaer. 6 This correction was a major one in the case of the thick absorber experiments and it was complicated by the fact that in the process of working backward from the energy deposited in the detector to the energy incident on the absorber it was necessary to use several sets of power law parameters to cover the broad energy range. Cross checks were made between the results of this procedure and the entirely independent method employed in the above-mentioned LAZY program 4 which depends on the range-energy method of Stewart. 5 Agreement within a few MeV was found. In the case of element spectra the range-energy relationship of the most prominent isotope was used. The ambiguity in the correction caused by this procedure was negligible for all low-energy results for lighter elements discussed in Sec. III. The uncertainty in the correction was several MeV in the case of unseparated isotopes, such as unresolved \(^9\)Be + \(^{10}\)Be, in the thick absorber data discussed in Sec. IV.

The intensity scale was converted to an absolute cross section scale by multiplying the monitor telescope count by a calibration constant taken from the previous work. 1,2 The end result of the SAVEX program was a set of printed tables of double differential cross section data and plotted curves of the same data for selected fragments.
III. RESULTS AND DISCUSSION OF LOW ENERGY DATA

A. Particle Spectra

Figure 1 shows the particle spectrum derived from data taken at 20° to the beam with telescope 1 of Table I and analyzed according to the procedure discussed in Appendix A. This run represents a data-taking period of ~ 50 hours, the longest of any of the experiments. This length of time enabled us to obtain significant statistical accuracy for the elements up through sulfur (element 16). A comparison of this figure with Fig. 3 of Ref. 2 shows that better statistical accuracy and better elemental resolution was achieved in the present study. Beryllium-7 is well separated from $^9,^{10}$Be but otherwise no isotopic resolution is apparent; this is a consequence of the thinness of the $\Delta E$ detector.

Spectra similar to Fig. 1 were generated from data taken at 45°, 90°, 135°, and 160° with good resolution of the elements Li through Na. Figure 2 shows the 90° results which are typical of intensity levels for the low energy measurements at these 5 angles (about 12 hours per run).

B. Energy Spectra

Figure 3 shows the energy data for the element lithium at 5 angles. Data points from the present measurements are sums over 2 MeV energy intervals. A part of the data from Ref. 2 was included in Fig. 3 to indicate the trend of the energy spectra between 60 and 160 MeV at 5 angles. The dashed curves in the figure represent an attempt to fit the shape of the spectra up to 35 MeV to the predictions of a very simple evaporation expression taken from Refs. 1 and 2, which assumes a uniform nuclear temperature for the ensemble of evaporating nuclei. A brief statement of this evaporation expression is given in Appendix B.

As in Ref. 2 families of curves were computed for a variety of parameter choices and compared with the experimental data to arrive at the set of "best"
values listed in Table II. Values from Ref. 2 are also listed there. The differences in the two sets result from the fact that those from the present work were chosen for the best fit of the data up to 35 MeV whereas those from Ref. 2 were chosen for the best fit up to somewhat higher energy.

A similar presentation of energy spectra and theoretical curves for $^7\text{Be}$, $^9,^{10}\text{Be}$, boron, and carbon is given in Figs. 4 through 7. In all cases the parameters used for the calculated (dashed) curves are those given in Table II. Also, in all cases the data at the five angles are shown displaced for clarity by factors of 10 on the intensity scale. Figures 8 through 10 present data on N, Ne, and Na. Figure 11 is a composite of 20° data for the elements B through S.

The comparison of the data of Figs. 3-7 with the simple evaporation expression (dashed curves) confirms the main conclusions of the earlier studies: namely (1) an expression based on isotropic fragment emission from a moving nucleus and a constant nuclear temperature can fit the 90° data near the yield maximum, but (2) there is more forward peaking in the fragment emission than can be explained even after correction for center-of-mass motion of the emitting nucleus and (3) the effective Coulomb barrier is about 40-50 percent of that calculated by a classical tangent spheres estimate.

C. Evaporation Spectra Based on a Maxwellian Distribution in Excitation Energy

In Refs. 1 and 2 it was difficult to fit the entire range of energy in the measured energy spectra for Li, Be, and B with the evaporation expression based on constant nuclear temperature and it was noted that the changing slope of the high energy portions of the spectrum could be interpreted to require a steady increase in nuclear temperature to a value of 20 MeV or higher. It is
reasonable to expect a range of excitation energies for the collection of
struck nuclei from the conventional description of the initial high-energy
cascade. In principle, one might hope to determine the distributions in
excitation energy by examining the output from some detailed Monte-Carlo calcula-
tions of the high energy cascade of the type done by Chen et al.\textsuperscript{7} or Bertini et al.\textsuperscript{8}
This can be done for proton bombarding energies of several hundred MeV but is
inexact for GeV protons because of incomplete information on inelastic collisions
for the proton-nucleon interactions needed as input for the cascade calculations.
Even with reliable input data the calculation is a major undertaking and has
not been done up to now except for pioneering attempts by Metropolis et al.\textsuperscript{9}
and Bertini et al.\textsuperscript{8}

In the absence of a theoretical model, we decided it would be interesting
to explore the influence of an arbitrary distribution in excitation energy which
follows our intuition, as well as the preliminary indications from the cascade
calculations, as to its form: namely, that it rise rapidly to some most
probable value and then decrease slowly out to high values. A Maxwellian
distribution fits this description and was chosen for its many mathematical
advantages.

We programmed a new calculation, referred to as MAXVAP in the remainder
of this paper, in which the same formula as used in the above constant $\tau$ model
was used for the evaporation of particles. This was taken from LeCouteur's
treatment\textsuperscript{10} of the Weisskopf evaporation theory. The new feature introduced
in the MAXVAP calculation was the incorporation of a Maxwellian excitation energy
distribution. Another important feature introduced in the MAXVAP calculation
was the assumption that the velocity of the struck nucleus $v$ was directly
proportional to excitation energy, $E^*$. The details of this model are given
in Appendix C. Although it is known from published calculations of the cascade step that there is a distribution in atomic charge and mass number for excited nuclei present at the end of the initial cascade step, we did not complicate the MAXVAP program by introducing this additional distribution, but we selected an average emitting nucleus somewhat lower in Z and A than the target nucleus. To facilitate comparison with the earlier work we used the same choices: namely $^{96}$Tc as the emitting nucleus representative of the Ag target and $^{220}$Rn as representative of the U target. The selection of $^{96}$Tc as the emitting nucleus for a Ag target is reasonable on the basis of the calculation of Bertini et al. of 3 GeV protons on $^{100}$Ru. In Refs. 1 and 2 it was stated that sample calculations confirmed that the predicted properties of emitted fragments are not sensitive to changes of several units in Z or A of the emitting nucleus. The results of the MAXVAP calculation are illustrated by the solid curves in Figs. 3-10.

There are only three input parameters in the calculation; the most probable excitation energy, $E_m^*$, the fraction of the nominal Coulomb barrier, $k$, needed to fit the maximum of the energy distribution, and the level density parameter given by $a = M/b$, where $M$ is the mass number and $b$ is an adjustable constant. In the constant temperature evaporation model described in Appendix B it is necessary to specify two additional parameters; namely, the center of mass velocity, $v$, and a parameter $n$ which correlates $v$ with $V$, the velocity of the emitted fragment. In the MAXVAP model these parameters are unneeded because of the assumption that center of mass velocity is proportional to excitation energy.

Although there is no unique set of values for the three input parameters, a reasonable set was found to reproduce more or less well the characteristics of
the low-energy fragments from Ag. Generating families of curves to aid in
the selection of the "best" parameter was easier for MAXVAP than for the
5 parameter expression of Appendix B. A k value of 0.4 gave the best fit on
the Coulomb barrier section of the spectrum and this value was relatively
insensitive to variations in the $E^*_m$ and level density parameters. On the
other hand the calculated curves were quite dependent on the pairs of choices
of the last two parameters. If $E^*_m$ was decreased, no change in the calculated
curve was found if the level density was decreased at the same time. Best
fits to the data were obtained with $E^*_m$ and level density pairs which corresponded
to a nuclear temperature of about 11.5 MeV evaluated from the expression,$E^*_m = aT^2$. The best fit of the 90° Li data were obtained with $k = 0.4$,
$E^*_m = 400$ MeV and $a = 6$ ($b = 16$) and these choices were used for all the curves
drawn in Figs. 3-10. In each figure the curves were normalized to the 90° data.

In Figs. 3 and 4 there is clear improvement in fitting the 90° data
for 6Li and 7Be over that obtained with the constant temperature formula
(dashed curves). This improvement is even more marked on the higher energy
part of the spectrum discussed in Sec. IV. However, the MAXVAP expression
shows the same defect as the constant temperature formula in its inability to
account for a pronounced favoring of forward emission of the fragments in the
center-of-mass system. No theoretical model now in the literature is able to
account quantitatively for this feature of high energy reactions.

MAXVAP curves for B and C are not significantly better than the constant
temperature evaporation curves for the data shown in Figs. 6 and 7. We may
note also that the favoring of forward emission steadily decreases with $Z$ until
at oxygen the agreement of the data points with the curves is equally good at
all 5 angles. An error in beam monitoring during the 135° experiment probably
accounts for the fact that the 135° data for all fragments are high.
IV. RESULTS AND DISCUSSION OF HIGH ENERGY DATA

A. Particle Identification

In the experiments performed with the detector telescopes and absorbers described in Sec. II.C, only the fragments of lowest atomic number had sufficient range to penetrate the Al absorber and the two $\Delta E$ detectors. Figure 12 shows the particle spectrum obtained from the Ag target with telescope 2 (Table I) and the 690 mg/cm$^2$ Al absorber. Figure 13 shows the spectrum obtained from the U target with telescope 4 and the 690 mg/cm$^2$ Al absorber.

B. Energy Spectra for Li, Be, B and C

Energy data for $^6$Li and $^7$Li obtained in the runs with four different thicknesses of aluminum absorber are shown in Fig. 14. The solid lines show the trend of the data and have no theoretical significance. In the case of the Ag target data, these lines show the extension of the 20° data of Fig. 3, with the difference that Fig. 3 includes all lithium isotopes, whereas Fig. 14 shows $^6$Li and $^7$Li contributions, separately. The dashed lines represent curves computed by the MAXVAP code for the following parameter choices: $E_m^* = 400$ MeV, $k = 0.40$ and $a = 6.0$ ($b = 16$). The curve is the same as that shown in Fig. 3, except for a scale adjustment required by the fact that individual isotopes are displayed in Fig. 14. The MAXVAP curve lies below the 20° data at all energies but has the same slope out to about 90 MeV. On the other hand, the constant temperature evaporation expression of Appendix B generates a curve dropping steeply away from the data in the 60-90 MeV region. Above 100 MeV, however, there is a sharp break in the energy spectrum and the appearance of a distinct high-energy "tail" which cannot be represented by the evaporation calculation even with a maxwellian distribution of temperatures. If we examine the U target data we notice the same sharp break in the spectrum above 100 MeV.
Similar comments may be made about the $^7$Be data which are plotted in Fig. 15. In this figure we have included the low energy points for Ag near the Coulomb barrier and a dashed curve showing the MAXVAP computation. For the Ag data, the entire range of energy from 10 to 140 MeV is reasonably well reproduced by the evaporation calculation for an $E_m^*$ value of 400 MeV. However, if the MAXVAP calculation is normalized to the 90° data as shown in Fig. 4, the 20° data points lie well above the calculated curve for 20°. Beyond 140 MeV there is a distinct break in the spectrum revealing a high energy tail extending, at least to 400 MeV.

We have considered the possibility that the high energy component in the Li and Be spectra could be caused by some background or instrumental effect. We believe the observed effect is real. A strong argument to support this is that particle identification was made on those fragments after penetration of the thick absorbers. In all the experiments particle identification spectra similar to those shown in Figs. 12 and 13 were obtained and only the data appearing in the well defined Li or $^7$Be peaks were selected for plotting. We believe that the criteria set on the data are so stringent that there can be no doubt of the reality of the high energy component extending up to 300 MeV for Li and to 400 MeV for $^7$Be.

These high energy fragments cannot be described by any version of evaporation theory we have considered so far. In all likelihood they are not produced by evaporation but by some process occurring during the cascade. If so, they should be very strongly peaked in the beam direction. We regret we did not have sufficient beam time to make long runs at angles other than 20° in order to determine whether the angular distribution of 200 to 400 MeV fragments favors the forward direction much more strongly than that of fragments of lower energy, as known already from Refs. 1 and 2 and the present study.
For the $^7$Be fragments from U targets the spectrum shape is remarkably similar except for the shift in the location of the turnover near the Coulomb barrier. It is possible to generate a curve from the MAXVAP calculation which tracks the data points out to about 120 MeV before the transition into the high energy tail. However, the $E^*_m$ value necessary to achieve this agreement is greater than 2000 MeV when $^{220}$Rn is taken as a representative evaporating nucleus. This is an unphysically high value for $E^*_m$ because the average value of excitation would approach the bombarding energy and from all we know about GeV proton interactions with uranium the average excitation must be much lower than this. If we assume that $^7$Be is evaporated from a fission or fragmentation product of U with about half the charge and mass of U, the $E^*_m$ value can be reduced to 500 MeV. However, then the high energy fragment distribution would be isotropic in disagreement with the experimental fact.

We now consider the data for the heavier isotopes of Be shown in Fig. 16 and for B and C shown in Figs. 17 and 18. Because of the lower yield of these particles, a measurable number of them were observed only in the runs with the thinnest of the 4 aluminum absorbers, and it was not possible to determine whether the energy spectra of these fragments had high-energy components (tails) similar to those observed for $^6$Li, $^7$Li, and $^7$Be. A MAXVAP calculation with an $E^*_m$ value of 400 MeV agrees roughly with the $^9,^{10}$Be data from Ag out to 140 MeV and with the B data out to 180 MeV. However, in the case of the U target data, we encounter the same difficulty that the MAXVAP curve agrees with the data only if a much higher value of $E^*_m$ is chosen, namely $E^*_m = 2000$ MeV.

We compared our high-energy data on Be, B, and C fragments from uranium with those appearing in Ref. 1 in the energy region where they overlapped.
and found our cross section to be a factor of 2 lower. On the other hand, our cross section data on Li agree with the previously published data within the experimental error of about 25 percent.

C. High Energy Data for He Isotopes

In Refs. 1 and 2 it was observed that the high energy part of the $^3$He spectrum was flatter than that for $^4$He and that above a fragment energy of about 100 MeV the $^3$He yield was greater than that of $^4$He. Alard$^{12}$ has also noted a high $^3$He/$^4$He ratio in fragments from Al bombarded with 20 GeV protons and in C, Al, Au targets bombarded with 600 MeV protons. Piroué and Smith$^{13}$ measured high energy $^3$He produced in Pt bombarded with 2.9 GeV protons.

In our own research the measurement of He spectra was not a primary goal and the detector telescopes were not optimum for this purpose. Nonetheless, the He isotopes were clearly resolved in some of the thick absorber experiments (see Figs. 12 and 13) and a few points were determined in the high energy part of the energy spectra at 20° for $^3$He, $^4$He, and $^6$He. Figure 19 shows the results for the Ag target together with previous data from Ref. 2. Our $^4$He points do not join smoothly with the Ref. 2 data; our cross section is a factor of 2 lower in the overlap region near 100 MeV. A discrepancy exists also in the $^3$He data but it is somewhat less for this isotope. In the case of $^6$He the overlap is smooth within the experimental error. The dislocation of the points for $^3$He, $^4$He, and $^6$He in the Ag target run with the 690 mg/cm$^2$ aluminum absorber convince us there was a factor of 2 error in the monitoring or in the processing of the data for that run. We adjusted these points downward as shown by the arrows, and applied the same adjustment to the $^6$Li, $^7$Li, and $^7$Be data obtained in the same run and displayed in Figs. 14 and 15.
In the case of U we can report the following cross sections at 20°:
for \(^4\text{He}\) energies of 78, 100, and 135 MeV the cross sections are 600, 300, and 125 \(\mu\)b MeV\(^{-1}\) Sr\(^{-1}\), respectively. For \(^6\text{He}\) energies of 95, 117, and 160 MeV the cross sections are 20, 8, and 3.5 \(\mu\)b MeV\(^{-1}\) Sr\(^{-1}\), respectively. The \(^4\text{He}\) data of Ref. 1 extend to 120 MeV and at this energy the Ref. 1 cross section is double the one reported here. The \(^6\text{He}\) data in the two studies agree within 20 percent up to 120 MeV where the Ref. 1 data cut off.

The presence of a strong \(^3\text{He}\) contribution to the high energy He spectrum is a striking and interesting phenomenon which is discussed in detail by Alard\(^{12}\) in a thesis which reviews the literature on the observation of energetic deuterons, tritons, and \(^3\text{He}\) in high energy reactions and the explanations proposed for their production. Some theoretical papers propose the direct creation of deuterons and mass-3 particles in the interaction of the incident proton with a target nucleon,\(^{14}\) while others propose the formation of these particles out of the cloud of cascade neutrons and protons when 2 or more of these are produced with small relative momenta.\(^{15-17}\) Still others apply new statistical thermodynamic concepts to the calculation of particle production from a volume of hadrons strongly heated by the intranuclear cascade.\(^{12,18}\)
This paper describes new experimental data on energy spectra of identified fragments from Ag and U targets bombarded with 5.0 GeV protons. It may be considered an extension of the work described in 2 previous publications.\textsuperscript{1,2} For Ag targets, segments of the energy spectra of light elements up to sodium were measured at 5 angles to the beam with a detector telescope containing a 16 μm ΔE detector. For both Ag and U targets the high energy part of the spectrum was measured at 20° to the beam for He, Li, Be, B, and C fragments with telescopes including thicker detectors and absorbers.

Two evaporation models were tested to determine whether they could explain the main features of the light element spectra, with the most severe tests posed by the data on Li and Be. A simple evaporation expression based on a fixed nuclear temperature for the evaporating nucleus can reproduce the shape of the spectra only in the region of the maximum. This maximum defines an effective Coulomb barrier. At the higher energy, particularly above 100 MeV for Li and Be, the calculations in no way reproduce the slope of the data. A modified evaporation calculation called MAXVAP which involves an arbitrary Maxwellian distribution in excitation energy can reproduce the data rather well out to about 100 MeV for Li and Be fragments from Ag with acceptable choices of excitation energy and level density parameters. (E\textsuperscript{m*} = 400 MeV, a = 6.0). For fragments from U the MAXVAP calculations reproduces the 20° data reasonably well out to the same energy but only by use of a most probable excitation energy of 2 GeV which seems much too high, or by use of a level density parameter b \sim 80 which seems unreasonable.

Both evaporation models have difficulty in explaining the relative intensities of the particle emission at the 5 angles studied. Even after
correction for center-of-mass motion the data indicate a pronounced favoring of forward emission. This favoring decreases with atomic number and seems to disappear for oxygen and higher elements. Another curious feature, independent of the chosen evaporation expression, is the fact that the effective Coulomb barrier for fragment emission in these high energy reactions is only 40 to 50 percent that estimated from a simple tangent spheres model. An effect contributing to this reduction in the Coulomb barrier is discussed by Moretto.25

In the case of $^6$Li, $^7$Li, and $^7$Be the 20° data clearly reveal a high-energy component which cannot be described by the MAXVAP calculation and must represent fragments produced in some little understood way during the initial high energy cascade. This statement applies to $^6$Li fragments measured in this study between 120 and 320 MeV and $^7$Be fragments between 120 and 420 MeV. The He isotope yields as measured out to ~ 160 MeV show $^3$He/$^4$He ratio exceeding 1.0. This also is probably a consequence of the cascade step. Some suggestions for possible mechanisms for production of very high energy light fragments which have appeared in the literature, are cited but not discussed.

In the process of working up $\Delta E \times E$ data which included information on many particles over wide energy ranges the necessity for a better method of extraction of particle identity arose and this need was met by the method described in Appendix A.
ACKNOWLEDGMENTS

We acknowledge with thanks the many contributions of the operating crew and staff of the Bevatron to the success of these experiments. We are indebted to the semiconductor development group of LBL for fabrication of many of the detectors and to Gordon Steers for preparation of the 1 mg/cm$^2$ silver target. We owe thanks to our colleagues Arthur M. Poskanzer, Gilbert W. Butler, and J. David Bowman, and Richard E. Eppley for advice and assistance in the experimental and data analysis parts of the study. We are specially indebted to Arthur Poskanzer for a critical reading of the manuscript. The Canadian members of the team of authors wish to express their thanks to the Lawrence Berkeley Laboratory for its hospitality and support.
APPENDIX A: NEW TREATMENT OF $\Delta E/\Delta x$ AND $E$ DATA TO ACHIEVE IMPROVED PARTICLE IDENTIFICATION

A. Previous Treatments

Semiconductor detector telescopes consisting of a thin transmission detector to measure rate of energy loss ($\Delta E$) and a thicker detector to measure the residual energy $E$ have seen widespread use in recent years for the identification of charge and mass of the products of nuclear reactions. Several methods of treating the $\Delta E$ and $E$ information to achieve this purpose have been applied. One method uses the relationship $^1$ $(E + \Delta E) \times (\Delta E) = \alpha Z^2$ which is an approximation to the Bethe-Block formula for the rate of energy loss of a charged particle. Another $^3$ is based on the empirical fact that the ranges of light nuclei follow a power-law relationship of the form $R = aE^b$ where $R$ is the range of a particle of energy $E$, $a$ is a constant which is characteristic of the particle and the exponent $b$ is approximately equal to 1.6 but varies somewhat with $Z$. For particles with energy in the range well described by this power law it can be shown that the function $(E + \Delta E)^b - E^b = T/a$, where $T$ is the thickness of the $\Delta E$ detector and quantity $T/a$ is a constant characteristic of a specific particle. The calculation of this power law function can be done by the analog circuitry designed originally by Goulding and Landis $^3$ or it can be done in a computer by manipulation of the digital record of the $E$ and $\Delta E$ pulse heights. The latter offers the advantage that parameter $b$ can be varied until the best particle resolution is achieved.

The above treatments provide satisfactory resolution of the particle spectrum for experiments in which the number of species and the range of energies for each species are limited. However, in studies of high energy reactions where a great number of different species, each covering a wide energy range is present, the particle resolution is not completely satisfactory.
as is made evident in two-dimensional plots of the particle identification parameter versus total energy of the particle. Butler, Poskanzer, and Landis\textsuperscript{18} revised the treatment based on the power-law relationship and introduced an improved algorithm of the form

\[
\left[ \left( \frac{E + E}{k} \right)^m - \left( \frac{E}{k} \right)^m \right]^{1/2},
\]

where the exponent $m$ is not $b$ but $b - c\Delta E/T$. The second term is a correction on $b$ where $C$ has a typical value of 0.05 in units of mg/cm\(^2\)-MeV and $T$, as before, is the $\Delta E$ thickness in mg/cm\(^2\). The divisor $k$ is set equal to 300 but its exact value is not critical. This algorithm is applied by reading the $\Delta E$ and $E$ information from magnetic tape and performing the algebra in a computer. A range of choices of $b$ and $c$ can be tried until the best particle resolution is achieved.

This treatment of Butler, et al.\textsuperscript{20} is a considerable improvement on the other methods cited above for particle identification in a complex mixture. We found it to be quite satisfactory for the analysis of the high energy data in the present study. However, in the case of the data taken with the 16\(\mu\) $\Delta E$ detector, we found it was possible to get a more satisfactory resolution with the method now to be described.

\textbf{B. New Treatment}

The Bethe-Block equation\textsuperscript{21,22} can be written in the form

\[
MZ^2 \propto \frac{(E_T)(\Delta E)}{\ln E_T + \text{const.}}.
\]
It is customary to ignore the \( \ln E_T \) term, but we decided to retain it since we observed the \( E_T \) vs \( \Delta E \) data to follow an exponential form (see Fig. 20). We arbitrarily ignored the constant in the denominator and after several tests of Eq. (A-2) and minor variants with our data we chose the following expression as our favored algorithm,

\[
PI = k \left[ \left( \frac{E_T}{\ln E_T} \right) \left( \frac{\Delta E}{1024} \right)^b \right]^{1/2},
\]

(A-3)

where \( k = \left( \frac{\ln 1024}{1024} \right)^{1/2} (1024) \). This constant and the divisor 1024 in A-3 are normalization factors related to the fact that the input pulses were digitized on a scale of 1024. The square root was taken so that the PI scale would be linear in \( Z \) instead of \( Z^2 \) and the exponent \( b \) was introduced to correct for neglect of the constant in A-2. Generally, \( b \) has the value 1.0 for the lightest elements (He, Li, Be) and increases exponentially with \( Z \) to a value of about 10 for \( Z = 18 \). A fixed value of \( b \) results in good particle resolution over a range of several elements. Inclusion of the \( \ln E_T \) term holds PI constant over a wide range of \( E_T \) and use of a variation in \( b \) provides uniform particle resolution over a range of \( Z \).

C. Procedure in Use of New Algorithm

For each run the tape record of digitized \( \Delta E \) and \( E_T \) pulse heights on an event-by-event basis was read into the computer and the data were summed into a \( E_T \) vs \( \Delta E \) matrix of dimensions 1024 \( \times \) 1024. This matrix was written on a magnetic tape which was used as the source tape for subsequent data manipulation. This was the most time consuming step in the whole process. (For \( 10^6 \) events it required about 10 minutes of CDC-6600 central processor time.) In the case of data from 3-element telescopes the two \( \Delta E \) values were summed. A
sample result of the summation of the data in $E_T$ vs $\Delta E$ form is shown in topographic map form in Fig. 20 but with the data compressed into a $128 \times 128$ matrix.

A computer program, PICAL, using the $E_T$ vs $\Delta E$ matrix as input, calculated a PI value for each element in the matrix making use of the expression $(A-3)$ and a preselected value of the exponent $b$. This procedure converted the $E_T$ vs $\Delta E$ matrix into an $E_T$ vs PI matrix. The program generated a particle spectrum plot by projecting all data on to the PI axis. This process was repeated for other values of the exponent $b$ and the particle plots were inspected to determine which values of $b$ were best for each section of the PI scale. A final calculation was then made with different $b$ values applied to different regions of the data matrix and a final $E_T$ vs PI matrix and a final particle spectrum were generated. Figure 21 illustrates the influence of the $b$ parameter on the particle resolution, including the use of multiple values of $b$ in a final processing of the data. Figures 1 and 2 are also examples of final particle spectra obtained by use of multiple values of $b$. Figure 22 is a plot of the final $E_T$ vs PI matrix in topographical map form. Some details are lost in this representation because the $1024 \times 1024$ scale was reduced to $128 \times 128$ in the illustration. The success of the PI algorithm is measured by the straightness of the ridges (i.e. no variation in PI with increase in $E_T$).

By inspection of the final particle spectrum it was possible to select PI channel numbers to isolate the energy data belonging to a specific particle type. The use of such PI "windows" in the reduction of the energy data is described in Sec. III.E.
APPENDIX B: CONSTANT TEMPERATURE EVAPORATION EXPRESSION

The energy spectrum in the moving system of the evaporating nucleus is represented by the expression

\[ P(\epsilon) = \sum_{k=(k)_{-\Delta}}^{(k)+\Delta} (\epsilon-kB)e^{-(\epsilon-kB)/T}, \quad \epsilon > kB, \quad \text{(B-1)} \]

where \( \epsilon \) is the disintegration energy, \( B \) is the nominal Coulomb barrier and \( kB \) is the effective Coulomb barrier. \( \Delta \) is a small parameter which specifies the spread in \( k \) values caused by the fact that a variety of emitting nuclei are involved. The energies of fragments of mass \( A \) emitted by a nucleus of mass \( M \) were corrected for recoil of the residual nucleus and related to a velocity \( V \) in the moving system by the equation

\[ (1 - A/M)\epsilon = 1/2 mV^2 \quad \text{.} \quad \text{(B-2)} \]

At 90° the laboratory energy \( E \) was taken equal to \( (1 - A/M)\epsilon \). At the other angles the laboratory frame velocity \( V_L \) was corrected for the velocity of the moving system \( v \), which was typically about 0.006 times the speed of light. The double differential laboratory cross sections were calculated from \( P(\epsilon) \) by use of the relation

\[ \frac{d^2\sigma}{dE\,d\Omega} = P(\epsilon) \frac{E}{\epsilon} \frac{d\epsilon}{dE} \quad \text{,} \quad \text{(B-3)} \]

*See Ref. 1 for more complete description.*
which is proportional to $P(\varepsilon) \sqrt{E/\varepsilon} / (dV_L/dV)$. A positive correlation between $V$ and $v$ was introduced to fit the data. It allows for the fact that fragments which are emitted from a struck nucleus moving with greater than average velocity will be ejected with greater than average velocity in the moving system. The correlation takes the form

$$\frac{v - \langle v \rangle}{\langle v \rangle} = n \frac{V - \langle V \rangle}{\langle V \rangle},$$

where the average quantity $\langle V \rangle$ was taken to be the root mean square $V$ obtained from the average energy $\langle \varepsilon \rangle$, which is equal to $\langle k \rangle B + 2\tau$ for a Maxwellian distribution.
APPENDIX C: AN EVAPORATION CALCULATION BASED ON A MAXWELLIAN DISTRIBUTION IN EXCITATION ENERGY

A new expression for calculation of fragment evaporation spectra was devised in an effort to allow for the fact that fragment emission in nuclear reactions induced by GeV protons occurs from nuclei with a wide distribution in excitation energy. In the absence of clear theoretical guidance on the form of this distribution, a Maxwellian distribution of excitation energies was chosen, as mentioned in the main text Sec. III.C. In addition, the forward momentum of the emitting nucleus and the appropriate kinematics were included in a more straightforward way than in the calculation described in Appendix B. This new calculation, called MAXVAP, was developed as follows.

The forward momentum of the emitting nucleus was calculated for each value of excitation energy by the expression

\[
P_{\text{ll}}/P_{\text{inc}} = E^* / E_{\text{max}}
\]

obtained from Monte Carlo calculations and some experimental studies. The maximum incident momentum, \( P_{\text{inc}} \), was determined from the incident energy of the bombarding proton by relativistic expressions. The maximum excitation energy, \( E_{\text{max}}^* \), was set equal to the incident energy minus the small amount associated with the forward momentum of the hypothetical compound nucleus. The velocity of the emitting nucleus, commonly referred to as \( v \), can then be calculated by the expression:

\[
v = E^* \left( \frac{2E_0}{T} + 1 \right)^{1/2} \left( 1 + A_t / M A_t \right)^{-1/2}
\]

where \( E^* \) is the excitation energy, \( E_0 \) is the rest mass of the proton, \( T \) is the incident proton energy, \( A_t \) is the mass of the target nucleus, \( M \) is the mass of
the emitting nucleus, and \(c\) is the velocity of light. For a given set of conditions, \(v = \text{const} \times E\).

We assumed an isotropic distribution of the evaporated fragments in the center of mass system. To compare the predictions with our experimental results, we transformed the predicted energy distributions into the laboratory system by means of the non-relativistic relationship \(^{24}\)

\[
V = v \cos \theta + \left( \frac{V_c^2}{c} - v^2 \sin^2 \theta \right)^{1/2}, \tag{C-3}
\]

where \(V\) is the velocity of the emitted particle at an angle \(\theta\) in the laboratory system, and \(V_c\) is the velocity of the particle in the center of mass system. \(V\) is given by \(V^2 = 2E/A\) where \(E\) is the measured energy for the particle of mass \(A\). \(V_c\) is obtained from \(V_c^2 = 2E_c/A\) where \(E_c\) is the energy of the particle in the center of mass system. However, this center of mass energy must be corrected for the recoil of the emitting nucleus to obtain the true disintegration energy, \(\varepsilon\). The final expression for \(V_c\) is then \(V_c^2 = 2\varepsilon(M-A/M_A)\). It is then possible to calculate \(\varepsilon\) in terms of measured or calculable quantities by rearranging Eq. C-3 to give:

\[
\varepsilon = \frac{MA}{2(M-A)} \left[ (V - v \cos \theta)^2 + v^2 \sin^2 \theta \right]. \tag{C-4}
\]

The correction for the transformation of the angular distribution from the center of mass to the laboratory system was calculated by \(^{24}\)

\[
\frac{\sigma_L(\theta)}{\sigma_c(\theta)} = \frac{v^2}{V_c \left( \frac{V_c^2}{c} - v^2 \sin^2 \theta \right)^{1/2}}
= \frac{v^2}{(V - v \cos \theta)(V^2 + v^2 - 2Vv \cos \theta)}, \tag{C-5}
\]
which is again in terms of known quantities.

To recapitulate, the new calculation starts with the same LeCouteur formula\(^1\) for evaporation of particles as used in the constant \(\tau\) model,

\[
P(\epsilon) \, d\epsilon = \frac{e^{-kB}}{\tau} \exp \left( -\frac{\epsilon - kB}{\tau} \right) \, d\epsilon .
\]  

(C-6)

The Maxwellian excitation energy distribution (C-7), in which \(E_m^*\) is the most probable excitation energy, was folded into C-6 by summing over \(E^*\) for each value of \(E\)

\[
P(E^*) = E^* \exp \left( -\frac{E^*}{E_m^*} \right).
\]  

(C-7)

In addition the velocity of the struck nucleus, \(v\), was made proportional to \(E^*\).

The resulting expression is:

\[
P(\epsilon, E^*) = a(\epsilon - kB) \exp \left[ \left( -\frac{\epsilon - kB}{(E^*/a)^{1/2}} \right) + \frac{E^*}{E_m^*} \right].
\]  

(C-8)

where \(E^* = a\tau^2\) and \(a\) is the level density parameter given by \(M/b\). By calculating the value of \(\epsilon\) from Eq. C-4 at each value of \(E\), we obtained an expression for the emission probability in terms of \(E\) and \(E^*\). The laboratory cross section was then calculated by the expression:

\[
\frac{d\sigma}{dEd\Omega} \propto \sum_{E^*=1}^{E_{max}} P(E^*, E^*) \frac{\sigma_L(\theta)}{\sigma_c(\theta)} .
\]  

(C-9)

The upper limit on this summation represents a truncation of the high end of the Maxwellian energy distribution, which is not serious unless very high \(E_m^*\) values are used.
FOOTNOTES AND REFERENCES

* Work supported in part by the U. S. Atomic Energy Commission and in part by the National Research Council of Canada.


11. In the course of this comparison we found that the $^{11}$C and $^{15}$C data in Fig. 11 Ref. 1 are plotted a factor of 10 too high. Also in Fig. 16 of Ref. 2 there is a scale error in the plotting of the 20° and 90° data for $^{7}$Be.
Table I. Detector Telescope and Absorber Characteristics

<table>
<thead>
<tr>
<th>Telescope</th>
<th>Detector thicknesses (µm)</th>
<th>Al absorber thickness (mg/cm²)</th>
<th>Target element</th>
</tr>
</thead>
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<tr>
<td></td>
<td>ΔE₂</td>
<td>ΔE₁</td>
<td>E</td>
</tr>
<tr>
<td>1</td>
<td>-</td>
<td>16 SB</td>
<td>194</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>61</td>
<td>1000 SB</td>
</tr>
<tr>
<td>3</td>
<td>&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
</tr>
<tr>
<td>4</td>
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</tr>
<tr>
<td>5</td>
<td>135</td>
<td>96</td>
<td>2000 SB</td>
</tr>
</tbody>
</table>

SB = surface barrier.
<table>
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<tr>
<th>Fragment</th>
<th>Source of Data</th>
<th>$B$, Coulomb barrier (MeV)</th>
<th>$\langle k \rangle$</th>
<th>$\Delta$</th>
<th>Temperature $\tau$ (MeV)</th>
<th>n</th>
<th>Moving system velocity $\langle v \rangle 70$</th>
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<td>Li</td>
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<td>19</td>
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<td>0.1</td>
<td>11</td>
<td>2.0</td>
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<tr>
<td>$^7\text{Be}$</td>
<td>this work</td>
<td>24</td>
<td>0.40</td>
<td>0.05</td>
<td>12.0</td>
<td>1.5</td>
<td>0.0055</td>
</tr>
<tr>
<td>$^7\text{Be}$</td>
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<td>24</td>
<td>0.5</td>
<td>0.2</td>
<td>11</td>
<td>2.0</td>
<td>0.008</td>
</tr>
<tr>
<td>$^9, ^{10}\text{Be}$</td>
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<td>0.40</td>
<td>0.05</td>
<td>10.0</td>
<td>1.5</td>
<td>0.0055</td>
</tr>
<tr>
<td>$^9\text{Be}, ^{10}\text{Be}$</td>
<td>HBP</td>
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<tr>
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</tr>
</tbody>
</table>

$^+HBP$ denotes Ref. 2.
FIGURE CAPTIONS

Fig. 1. Spectrum of elements ejected at 20° to beam direction in bombardment of Ag target with 5.0 GeV protons. Fragments measured with 16 μm ΔE detector and 194 μm E detector. Particle spectrum generated from ΔE and E pulse heights by the technique described in Appendix A.

Fig. 2. Spectrum of elements ejected at 90° to beam direction in bombardment of silver target with 5.0 GeV protons.

Fig. 3. Energy spectra at 5 angles for lithium fragments ejected from Ag target. Vertical scale shift by factor 10 at each angle. Squares represent data from this study, dots represent data from Ref. 2. (Only a portion of the data from Ref. 2 are shown.) Dashed curve represents fitted curve calculated from simple evaporation theory (Appendix B) with fixed nuclear temperature normalized to 90° data. Solid curves represent fitted curves from evaporation code MAXVAP (Appendix C) incorporating Maxwellian distribution of excitation energies, also normalized to 90° data. E m* = 400 MeV, k = 0.4 and a = 6 in MAXVAP calculation.

Fig. 4. Energy spectra at 5 angles for 7Be ejected from Ag targets. See caption of Fig. 3 for meaning of curves.

Fig. 5. Energy spectra at 5 angles for 9,10Be fragments ejected from Ag targets. See caption of Fig. 3 for meaning of curves.

Fig. 6. Energy spectra at 5 angles for B fragments ejected from Ag targets. See caption of Fig. 3 for meaning of curves.

Fig. 7. Energy spectra at 5 angles for C fragments ejected from Ag targets. See Fig. 3 for meaning of curves.

Fig. 8. Energy spectra at 5 angles for N fragments ejected from Ag targets. Vertical scale shift by factor 10 at each angle. Solid lines represent evaporation expression MAXVAP.
Fig. 9. Energy spectra at 5 angles for Ne fragments ejected from Ag targets. Vertical scale shift by factor 10 at each angle. Solid lines represent evaporation expression MAXVAP.

Fig. 10. Energy spectra at 5 angles for Na fragments ejected from Ag targets. Vertical scale shift by factor 10 at each angle. Solid lines represent evaporation expression MAXVAP.

Fig. 11. Energy spectra for elements from B through S ejected from Ag target at 20° to the 5 GeV proton beam direction.

Fig. 12. Particle spectrum of fragments ejected at 20° to beam direction in bombardment of Ag target with 5.0 GeV protons. Fragments penetrating Al absorber (690 mg/cm²) were measured with 3-element telescope (see Table I). The particle spectrum was generated from the ΔE and E pulse information by the technique discussed in Appendix A.

Fig. 13. Particle spectrum of fragments ejected at 20° to beam in bombardment of U target with 5.0 GeV protons. Fragments penetrating Al absorber (690 mg/cm²) were measured with 3-element telescope (see Table I). The particle spectrum was generated from the ΔE and E pulse information by the technique discussed in Appendix A.

Fig. 14. High-energy part of energy spectrum of 6Li and 7Li fragments ejected at 20° to the proton beam. The 4 sets of points refer to measurements with different aluminum absorbers in front of first ΔE detector (35.7, 432, 690, and 1295 mg/cm², respectively). The U target data points for the 35.7 mg absorber are plotted 20% low. The solid curves were sketched in to represent the smoothed trend of the data. The dashed lines represent the same MAXVAP curves calculated for Ag as the one for 20° shown in Fig. 3, adjusted for 6Li and 7Li isotopic composition.
Fig. 15. High energy part of energy spectrum for $^7$Be fragments ejected at 20° to the proton beam. Lowest energy set of Ag points are reproduced from Fig. 4. The other sets of points came from measurements with 4 different aluminum absorbers in front of detector telescope. Solid curves represent smoothed trend of data. Dashed line is the MAXVAP curve calculated for Ag for 20° reproduced from Fig. 4. The 80-140 MeV points for U are plotted 20% low.

Fig. 16. Energy spectrum of $^{9,10}$Be fragments ejected at 20° to the proton beam. Lower energy set of points reproduced from Fig. 5 (Ag target). Higher energy sets of data taken with 3-element telescopes with 35.7 mg/cm$^2$ Al absorber in front. U data plotted 20% low. Dashed curve represents calculation based on evaporation code MAXVAP. This curve is the same as shown in Fig. 5 for 20°.

Fig. 17. Energy spectrum of B fragments ejected at 20° to the proton beam.

See caption of Fig. 16 for further details.

Fig. 18. Energy spectrum of C fragments ejected at 20° to the proton beam.

See caption of Fig. 16 for further details.

Fig. 19. Energy distributions at 20° to the beam for $^3$He, $^4$He, and $^6$He ejected from a Ag target. Solid curves represents data from Ref. 2. Squares and triangles show new data.

Fig. 20. Topographical display of particle total energy $E_T$ versus $\Delta E$ for particles emitted at 90° to beam from Ag target. $\Delta E$ data taken with 16 μm silicon detector. Display based on $128 \times 128$ grid, whereas $E_T$ versus $\Delta E$ data used in final analysis were stored in a $1024 \times 1024$ matrix. The data can be represented by an equation of the form $E_T = k \times \exp (-m\Delta E)$. 
Fig. 21. Particle spectra generated by the PICAL program using algorithm (A-5).
In the upper curves the exponent b was set at 1.0, 1.5, and 2.0, respectively.
In the bottom curve the above range of b values were used in the particle section in which they produced the best resolution.

Fig. 22. Topographical display of $E_T$ versus PI. Data are the same as those in Fig. 20. PI was calculated with algorithm (A-5) with the same b parameter range as given in Fig. 21.
Fig. 1
Fig. 2
Fig. 4
Fig. 5

$\frac{d^2\sigma}{dE d\Omega}$ (µb/MeV-sr)

Energy (MeV)

$9,10\text{Be}$

- MAXVAP
- Constant

$\tau = 100$

$20^\circ$
$45^\circ \times 10^{-1}$
$90^\circ \times 10^{-2}$
$135^\circ \times 10^{-3}$
$160^\circ \times 10^{-4}$
Fig. 6
Fig. 7

Energy (MeV)

$\frac{d^2\sigma}{dE d\Omega} \text{ (\mu b/MeV-sr)}$

- MAXVAP
- Constant
- $\tau = 11.5$

- $20^\circ$
- $45^\circ \times 10^{-1}$
- $90^\circ \times 10^{-2}$
- $135^\circ \times 10^{-3}$
- $160^\circ \times 10^{-4}$

Fig. 7
Fig. 8
Fig. 9

\[ \frac{d^2\sigma}{dE d\Omega} \quad (\mu\text{b/MeV-sr}) \]

Energy (MeV)

- 20°
- 45° x 10^{-1}
- 90° x 10^{-2}
- 135° x 10^{-3}
- 160° x 10^{-4}

Ne
Fig. 10

The graph shows the differential cross section $d^2\sigma / dE d\Omega$ (in $\mu$b/MeV·sr) for the Na isotope as a function of energy (MeV). The data is presented at different scattering angles: 20°, 45°, 90°, 135°, and 160°, with corresponding cross sections in $10^{-5}$, $10^{-4}$, $10^{-3}$, $10^{-2}$, and $10^{-1}$, respectively.
Fig. 11
Fig. 12
Fig. 13
Fig. 14
Fig. 16
Fig. 17
Fig. 18
Fig. 19
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