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Fred S. Goulding and Yvonne Stone

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SEMICONDUCTOR RADIATION DETECTORS

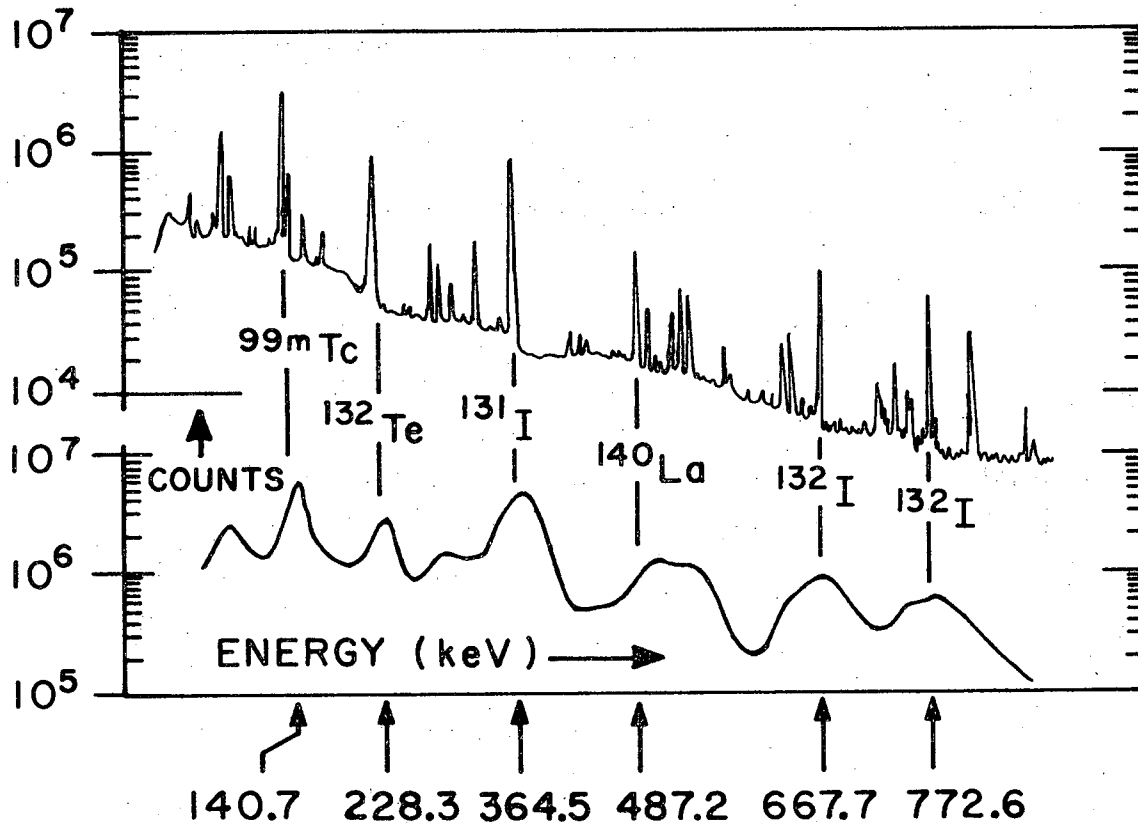
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July 1970

ABSTRACT

The basic physics of semiconductor radiation detectors is reviewed, including energy determination by ionization, charge production, charge collection, and detector materials. Use of these detectors in X-ray fluorescence analysis, high-energy nuclear spectroscopy, determination of very short decay lifetimes, particle identification, and measurement of mesonic X rays is described.



XBL 706-1196

Fig. 1

A complex gamma-ray spectrum, due to gross fission products, observed by a germanium detector (upper curve) and a scintillation detector (lower curve).

Just as the theory of atomic structure grew from spectroscopic observations of transitions in the atomic shell, so nuclear theory has developed from studies of the radiations emitted in nuclear transitions. But while the light produced or absorbed in atomic transitions is easy for us to see and measure, we have no simple way of observing nuclear radiations, which may be charged particles (electrons, positrons, protons, and heavier nuclear constituents), or electromagnetic radiations of very short wavelength (gamma rays). Necessarily then, the progress of nuclear physics has closely followed the development of new means of detecting and measuring nuclear radiations.

In the past decade, we have witnessed a major revolution in nuclear experiments as a new device, the semiconductor radiation detector, has appeared on the scene, providing relatively simple and accurate methods of measuring the energy of many of the radiations produced in nuclear processes. Knowledge of fission and of the structure of nuclei, detection of new transuranic elements and determination of their properties, and exploration of the nuclear surface all have advanced due to the development of these new detectors.

Even outside the field of nuclear physics, their influence is being strongly felt. In archeology, because of semiconductor radiation detectors, specimens can now be analyzed in fine detail by observing the gamma rays emitted from a sample after neutron bombardment in a reactor, or by observing the characteristic fluorescent X-rays produced when a sample is exposed to an X-ray or gamma-ray source. Biology, geology, mining, criminology, and many types of industrial processing have all taken advantage of this new analytical tool.

The power of semiconductor detectors in analysis is graphically illustrated by comparing the gamma-ray spectra shown in Fig. 1. The ordinate is a measure of the intensity of the gamma rays emitted from the sample; the abscissa shows their energy. The wealth of fine

detail seen in the spectrum produced by the semiconductor detector is totally absent in that produced by its predecessor, the scintillation detector.

Radiation Energy Determination by Ionization

All our methods of detecting and measuring the energy of nuclear radiation depend finally on the effects of charged particles, even if this radiation is electromagnetic in nature (gamma rays) or consists of uncharged particles (neutrons, neutrinos, neutral mesons). The conversion process by which charged particles are produced from the others might be a "knock-on" effect, such as the production of high-energy protons by neutrons passing through hydrogenous material; or a photoelectric or other interaction in which gamma rays release electrons; or possibly a nuclear reaction in which a charged particle is one of the end products. In all cases, the conversion process must be taken into account in interpreting the results.

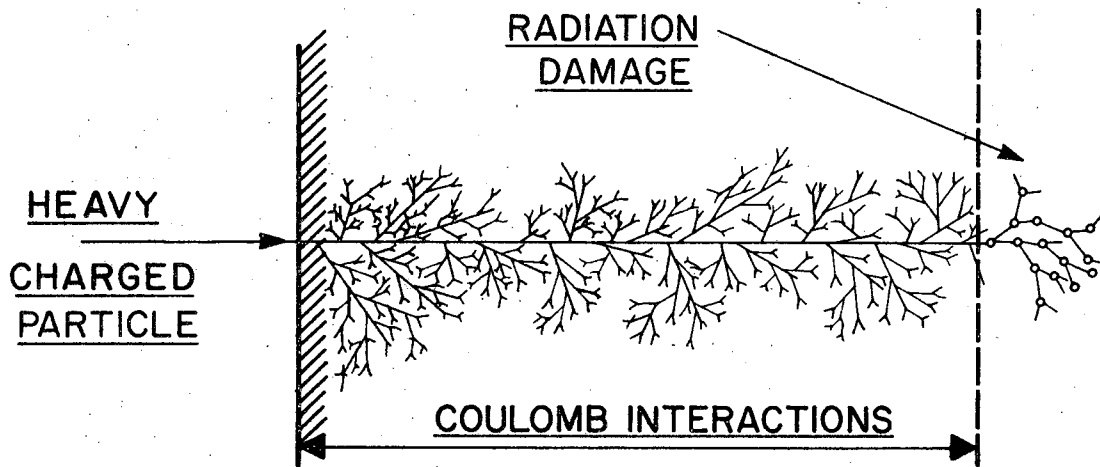
Only two fundamental methods of determining the energy of charged particles are available to us. If we bend the particles in a precisely known uniform magnetic field, as in a magnetic spectrometer, we obtain a measure of their momentum; then if their mass is known, their energy is determined. While this technique can provide very precise determinations of energy (accuracies $\sim 0.01\%$), magnetic spectrometers are large and expensive, and cover a limited range of energies. The second basic method of finding the energy of charged particles utilizes the ionization produced by their passage through matter. Although individual exchanges of energy between a particle and the electrons in matter vary greatly in size, a well defined average amount of energy is absorbed from the particle for each elementary charge it releases along the ionization track. Therefore, if the particle's energy is totally absorbed in a piece of material (i.e. if the particle stops in it), the ionization produced in the material is a direct measure of the particle's energy.

Ionization occurs in all materials. In making detectors, our problem is to select materials in which the products of ionization--positive and negative free charges--can be measured. Sometimes, a secondary effect of the free charge can be utilized; the operation of scintillation detectors, for example, depends on light emission from certain crystals, plastics, or liquids following excitation by the ionization process. In certain cases, direct collection of the charge to produce a measurable current in an external circuit can be achieved: only gaseous ionization detectors and semiconductor detectors have adequate charge collection properties to warrant their being considered for spectrometers. Semiconductor detectors have many advantages over gaseous ionization chambers and scintillation detectors; the fact that virtually all nuclear spectroscopy now uses semiconductor detectors is proof of their value. The basic reasons are:

(1) Solids are about 1000 times as dense as gases, and therefore much smaller thicknesses are needed to absorb radiation. Because of this, gaseous ionization chambers can be used only for measurements of slow-moving highly-charged particles which produce short dense ionization tracks, but semiconductor detectors can be used for a much wider range of types of radiation.

(2) The energy required to produce a single pair of elementary charges is much smaller in solids than in gases; in fact, the solids normally used in detectors produce about 10 times the amount of ionization, compared with gases, for the same amount of absorbed energy. Consequently, electrical signals are larger, statistical fluctuations less important, and the energy resolution significantly better.

(3) In principle, the above advantages also apply to scintillation detectors; indeed heavy materials of high atomic number used as scintillators give even better absorption efficiencies than can be achieved with our two common semiconductors, germanium and silicon.



XBL 706-1204

Fig. 2

Passage of a heavy charged particle through a solid. Most of the energy is absorbed in producing electron-hole showers; at the end of the track, the particle becomes neutral and loses its remaining energy in atomic collisions.

But, unfortunately, inefficient processes are involved both in the conversion of free electrons into light quanta by the scintillator, and in the conversion of light quanta into electrons by the electron multiplier tube. Statistical fluctuations are very large, and the energy resolution suffers accordingly. Semiconductor detectors usually exhibit one to two orders of magnitude improvement in energy resolution compared with scintillators.

The Charge-Production Process

Production of ionization in a detector material is vital to detector operation and determines the basic limitations on performance. As it passes through a detector, a heavy charged particle interacts with atomic electrons via Coulomb forces, producing a large number of small energy-exchanges (~ 100 eV) all along its track. Each electron set free initiates a shower of holes and electrons (Fig. 2). Apart from occasional small-angle scattering by nuclei, or involvement in a nuclear reaction (processes that become important only at high energies), a heavy particle travels along a straight line until its velocity becomes small enough for electrons to become attached to it. After becoming neutral, its remaining energy is dissipated in atomic collisions, displacing atoms, which collide with other atoms, producing intense damage to the detector material.

The action of fast neutrons is similar to that of charged particles near the end of their track, which makes them especially effective in producing radiation damage. In many experiments, neutron damage limits detector life. Light-weight charged particles such as electrons are almost unable to impart sufficient energy to the atoms of the detector material by collision to displace them, so that radiation damage is negligible.

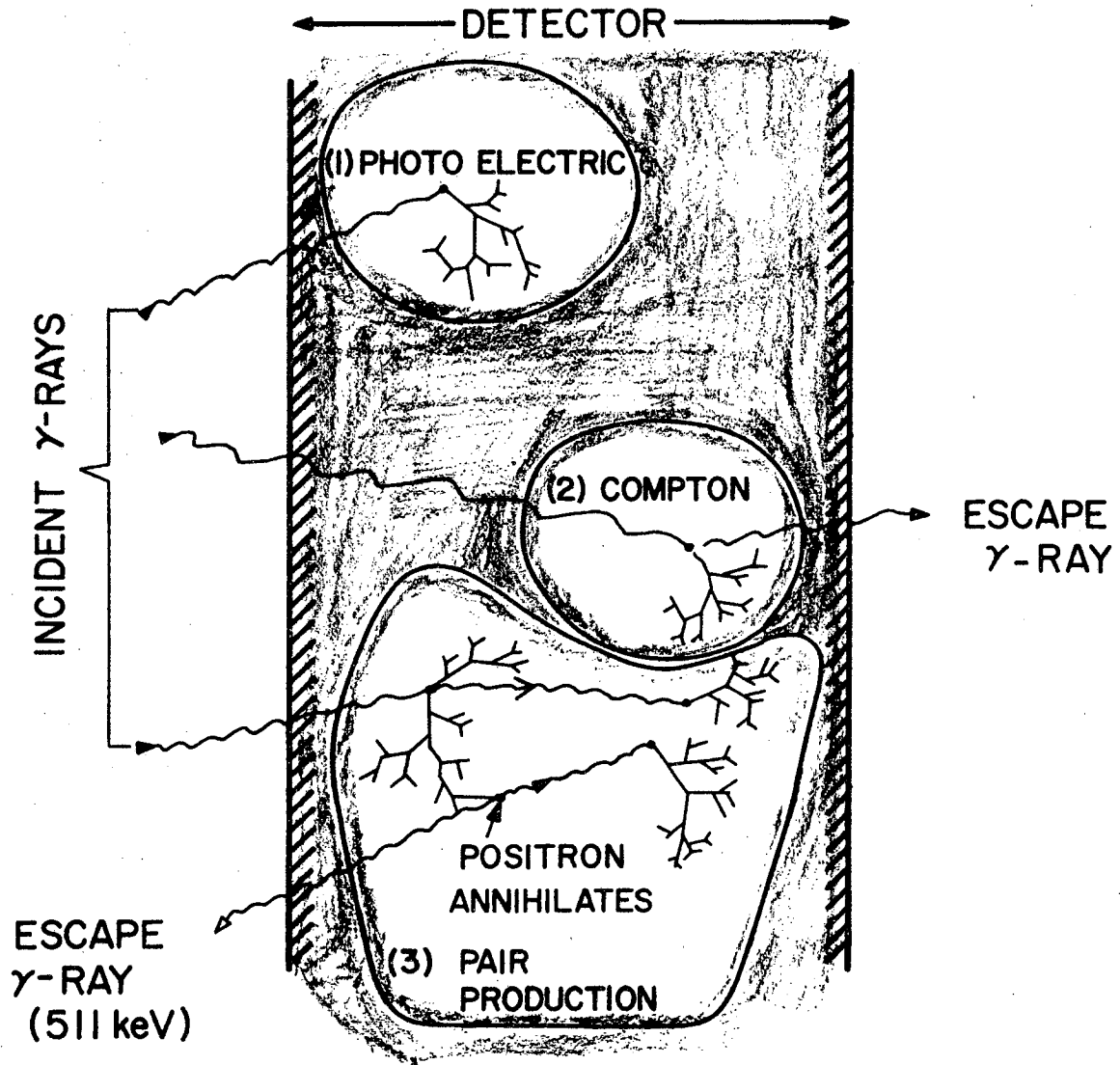


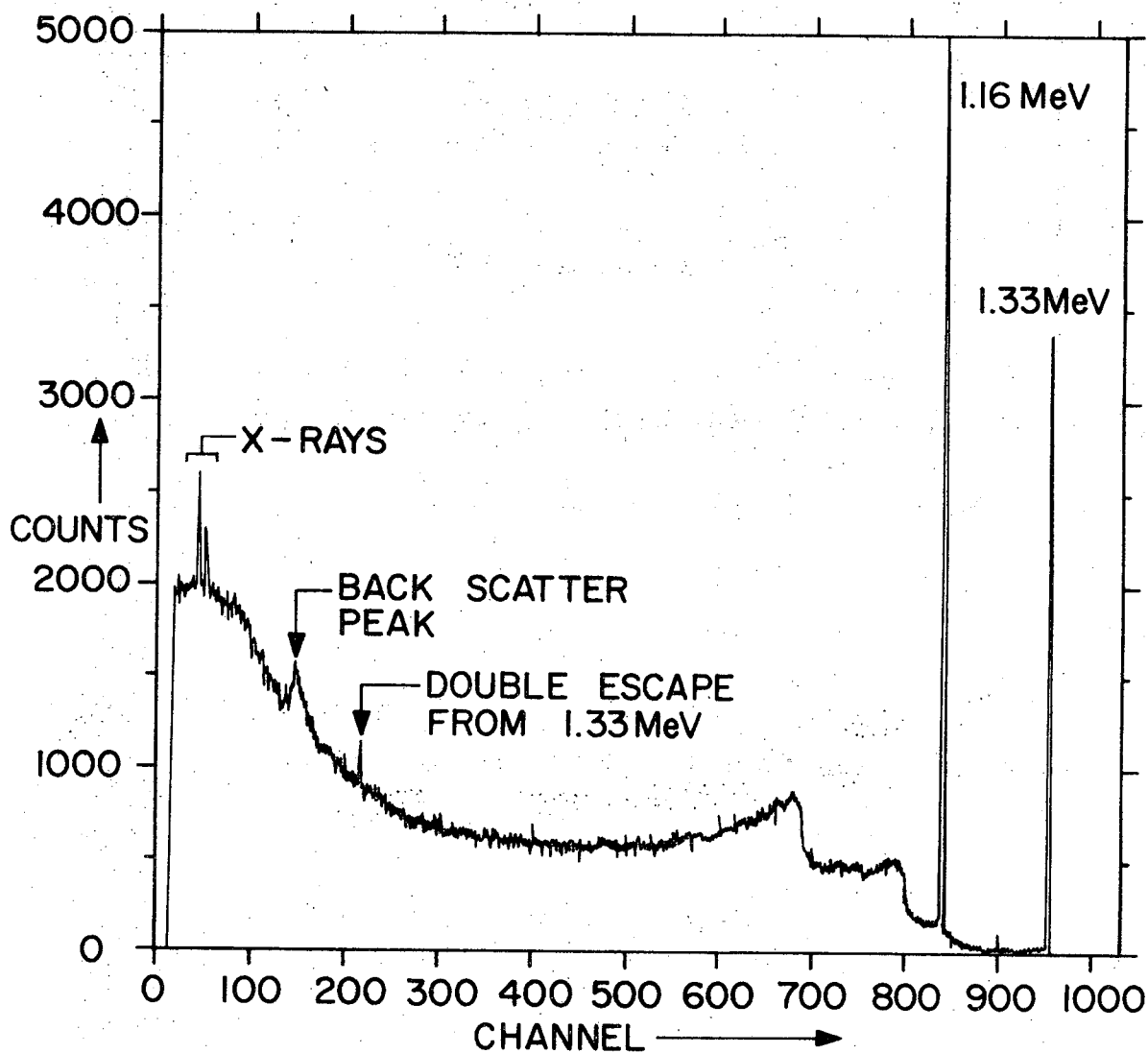
Fig. 3

XBL 706-1203

Three interaction mechanisms between gamma rays and the electrons in a material. In the photoelectric process nearly all the gamma-ray energy is transferred to the electron, which then produces a hole-electron shower. In the Compton process (billiard-ball collision), only a portion of the gamma-ray energy is given to the electron. Pair-production requires the expenditure of 1.02 MeV to produce an electron-positron pair, which can happen only with high-energy gamma-rays. The 1.02 MeV reappears as gamma rays when the positron annihilates.

When gamma rays enter a detector, any of three primary interactions may take place between the gamma rays and electrons: a photoelectric interaction, a Compton collision, or pair-production. These interactions are illustrated in Fig. 3. Whereas a photoelectric event produces an amount of ionization corresponding directly to the gamma-ray energy, Compton events produce a variable amount of ionization. Only if the degraded (less energetic) secondary gamma-ray is fully absorbed can we get useful information from Compton interactions about the distribution of gamma-ray energies. Therefore, unlike the case of heavy charged particles, where the amount of ionization is almost the same for each member of a monoenergetic group of particles, the ionization produced by monoenergetic gamma rays is not constant. Instead, a statistical distribution of pulses of ionization is produced, containing well defined peaks that can be used to determine gamma-ray energies. The rest of the ionization pulses constitute an undesirable background which can be reduced only by having a larger detecting volume to prevent the escape of degraded gamma rays. The difference between the ionization-pulse distribution for gamma rays and that for heavy charged particles is seen by comparing Figs. 4 and 5.

Another striking difference between gamma rays and charged particles is that gamma rays may pass through a detecting medium without interacting at all. In fact, the efficiency for detecting gamma rays and the probability of the energy of a gamma ray being totally utilized to produce ionization may be quite small, as shown in Fig. 6. On the other hand, all heavy charged particles are detected, even by very thin detectors. For their total energy to be absorbed, however, the thickness of the detector must exceed the range of the particle in the detector material. Range-energy curves for a range of energies and several particles in silicon and germanium are shown in Figs. 7 and 8.



XBL 706 1202

Fig. 4

The gamma-ray spectrum from ^{60}Co observed by a germanium detector. Note the two full-energy peaks due to the 1.16-MeV and 1.33-MeV gamma rays, and the Compton distribution at energies less than that of the peaks.

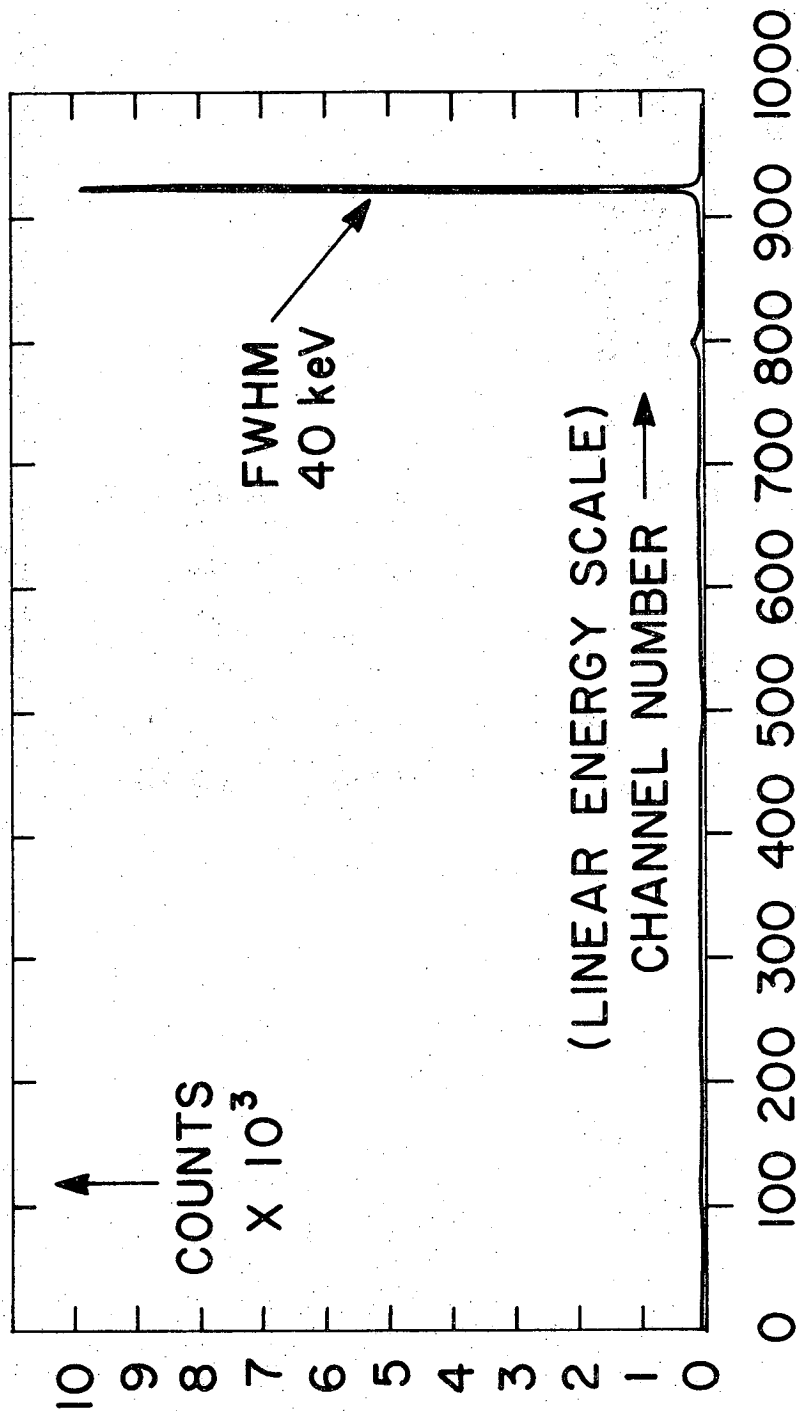


Fig. 5 XBL 706-1205

42-MeV proton spectrum in germanium. Note the virtual absence of background at energies lower than the peak.

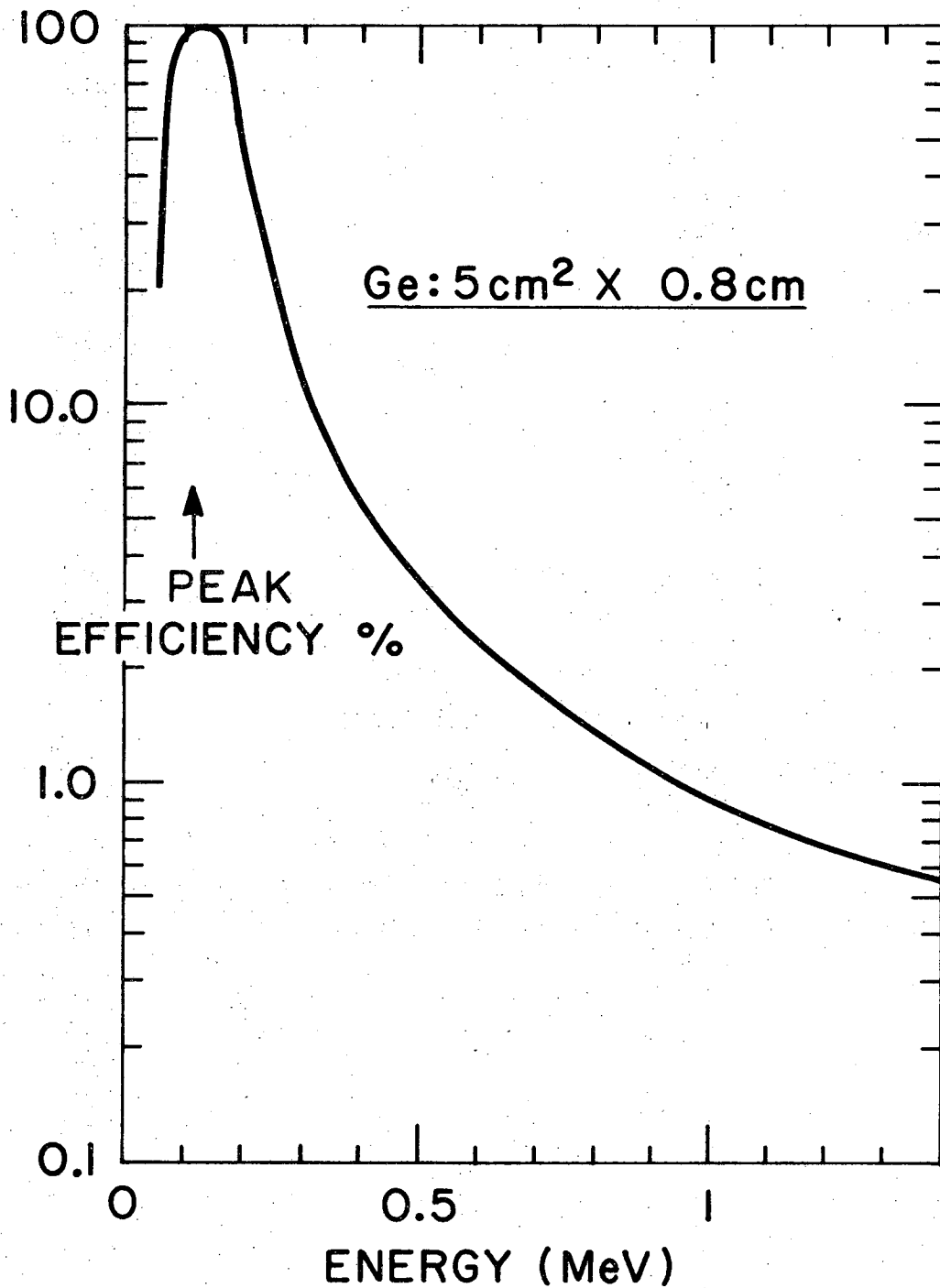
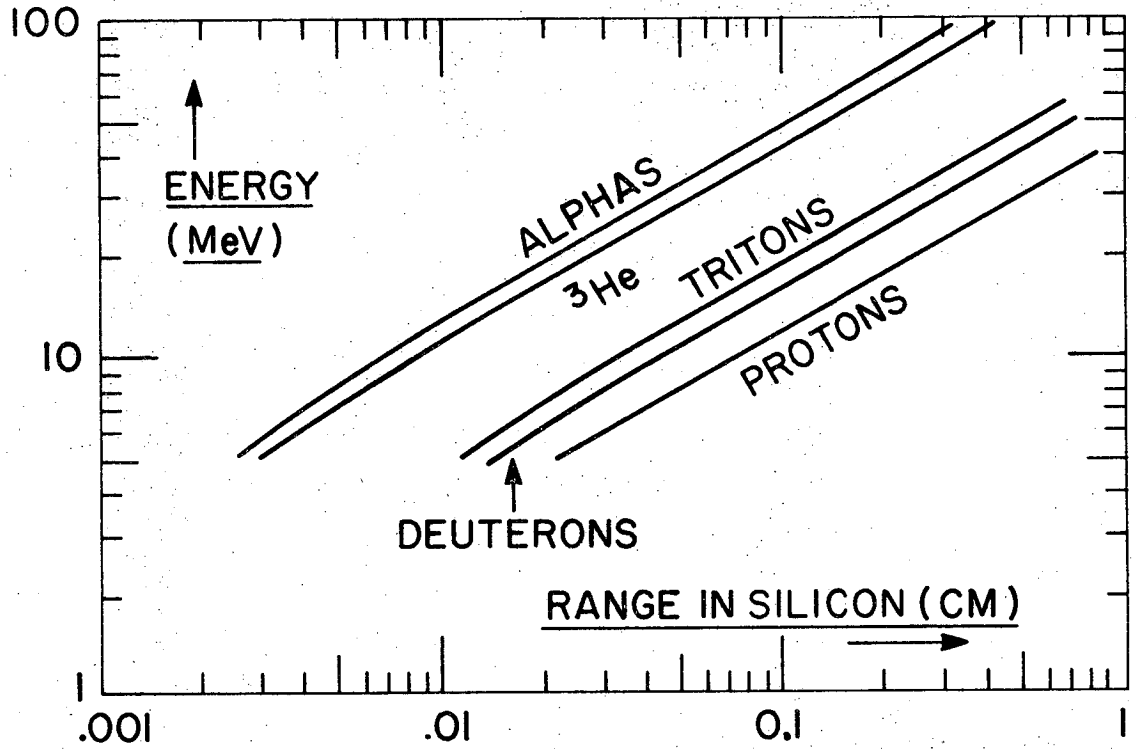


Fig. 6

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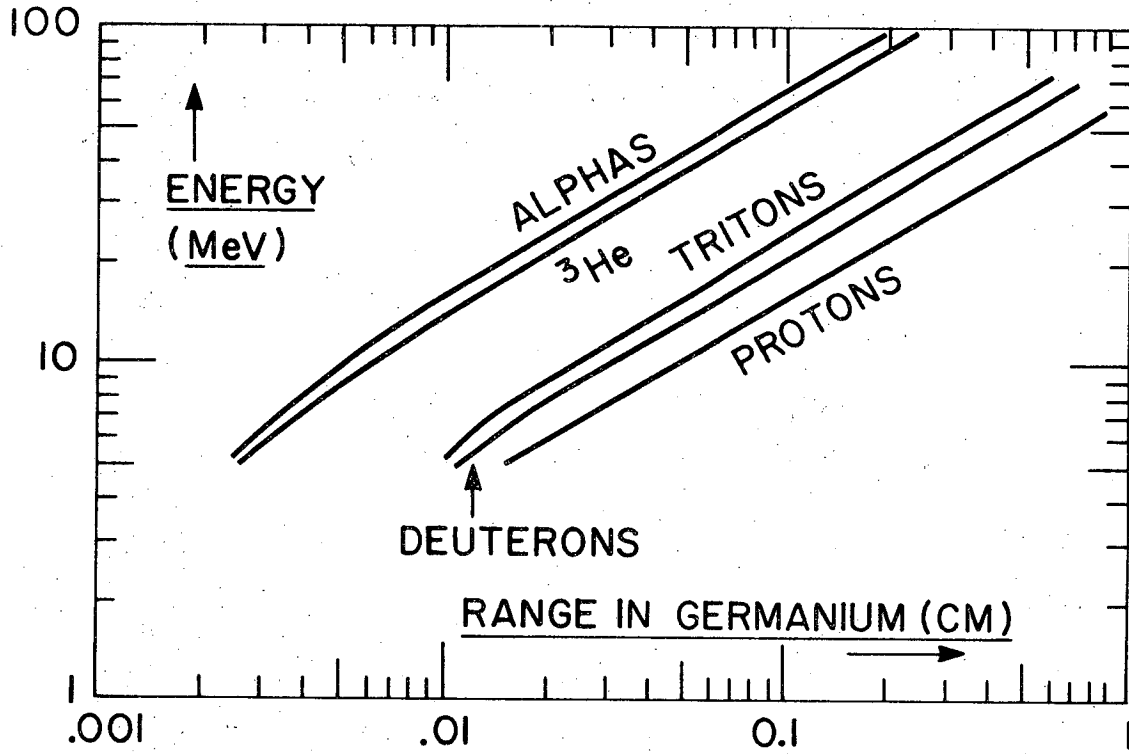
Gamma-ray efficiency as a function of energy for a germanium detector 5 cm² and 0.8 cm thick. Peak efficiency is defined as the ratio of counts appearing in the peak to the total number of photons entering the detector.



XBL 706 - 1195

Fig. 7

The range of protons, deuterons, tritons, ³He, and alpha particles in silicon.



XBL 706-1194

Fig. 8

The range of protons, deuterons, tritons, ³He, and alpha particles in germanium.

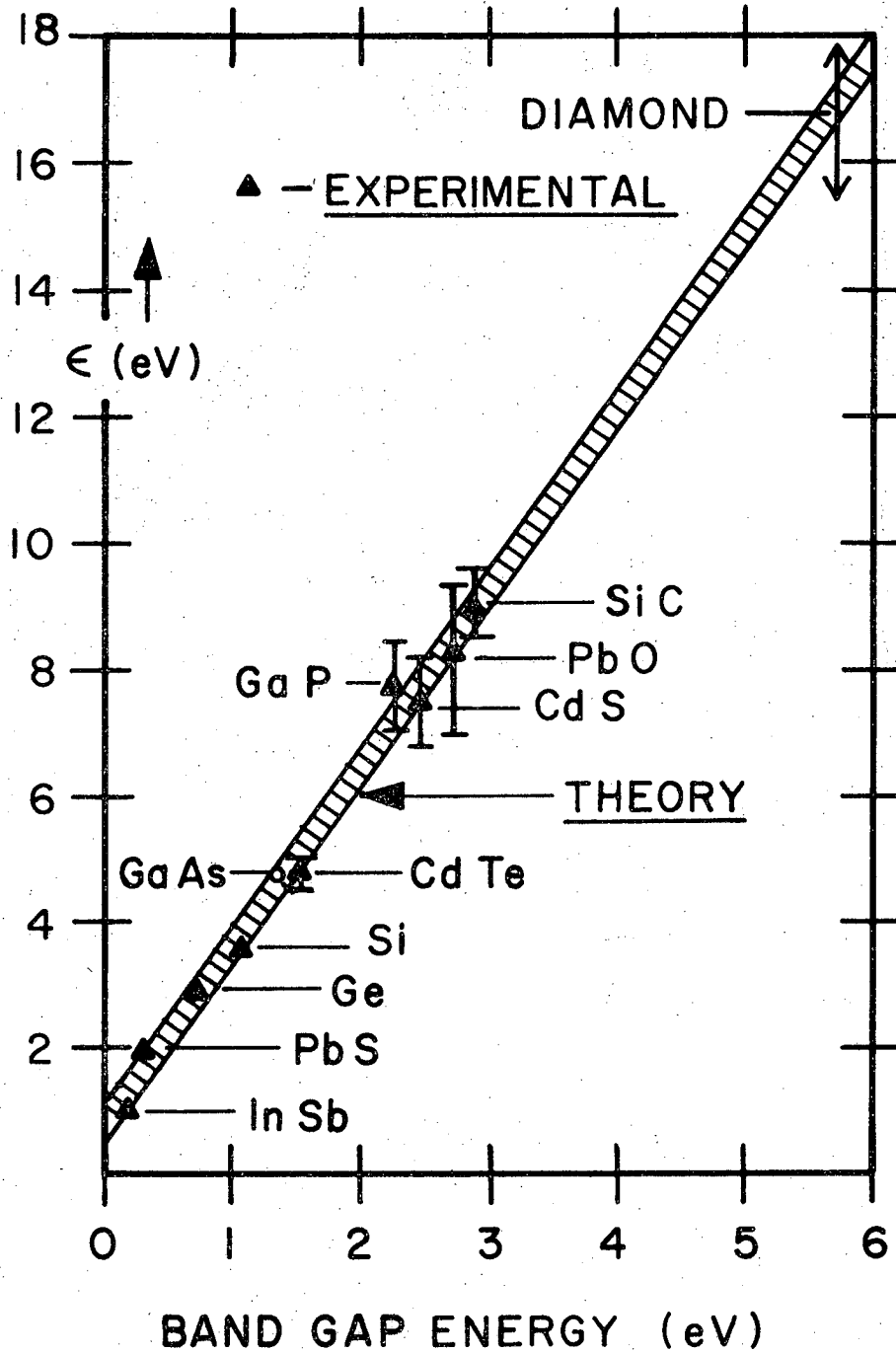


Fig. 9

XBL 706-1193

The variation in value of ϵ as a function of band-gap of a semiconductor. Experimental values are shown and compared with the theory of Klein (1).

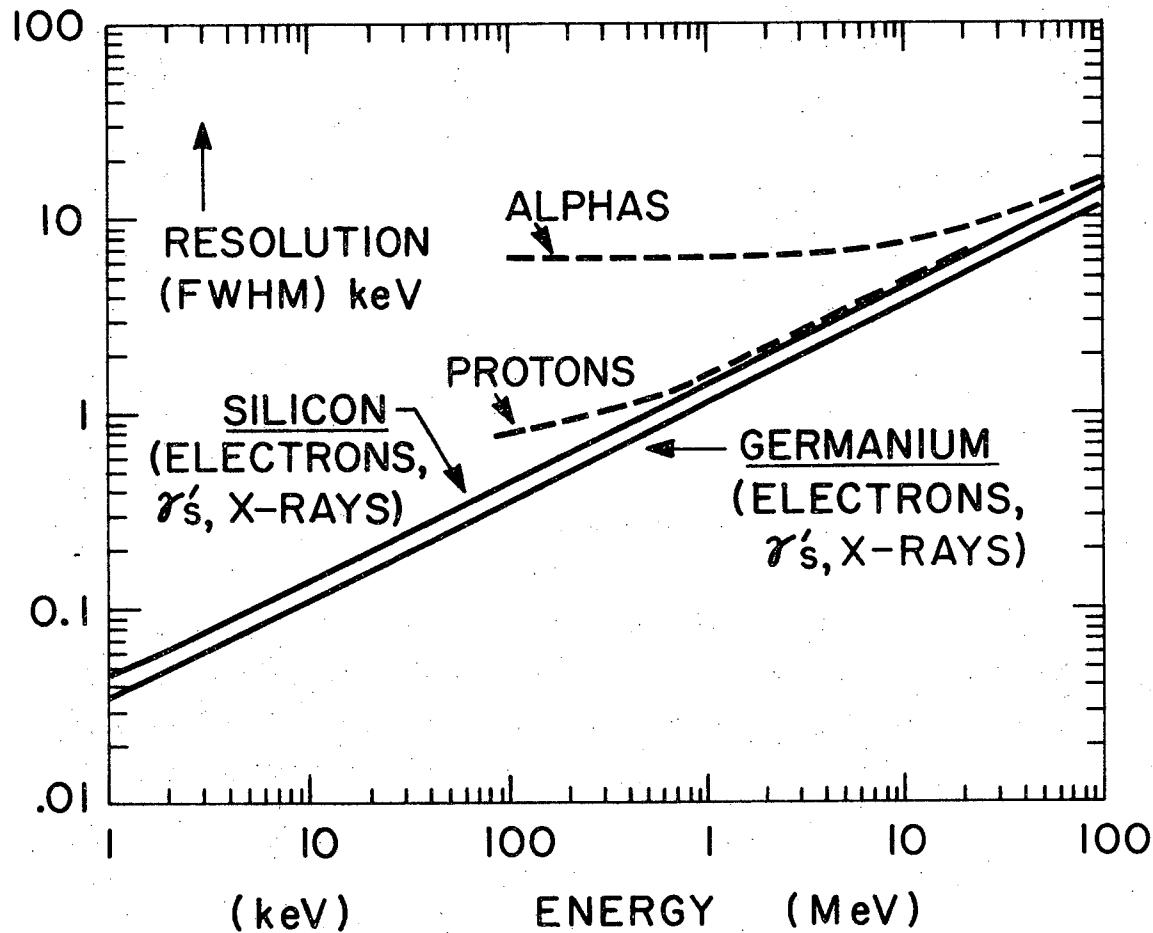


Fig. 10

XBL 706-1210

The energy resolution as a function of energy for silicon and germanium detectors. These curves take into account the basic statistical limitations involved in the charge-production process and in atomic collisions. The Fano factor F is assumed to be 0.08 for germanium, and 0.1 for silicon--the best experimental values yet observed. No account is taken of the amplifier-noise contributions to resolution.

The primary effect of radiation is to release electrons and holes which travel at high velocities initiating showers of free electrons and holes, the final members of which are travelling at thermal velocities in the lattice. The energy of the radiation is expended partly in exciting electrons from the valence-bond of the solid--this we may consider useful work from the point of view of detectors--and partly in wasteful excitation of lattice vibrations. The performance of a detector material depends on how energy is shared between useful and wasteful work, and on the statistical fluctuations in this process. We can characterize the performance in terms of two variables:

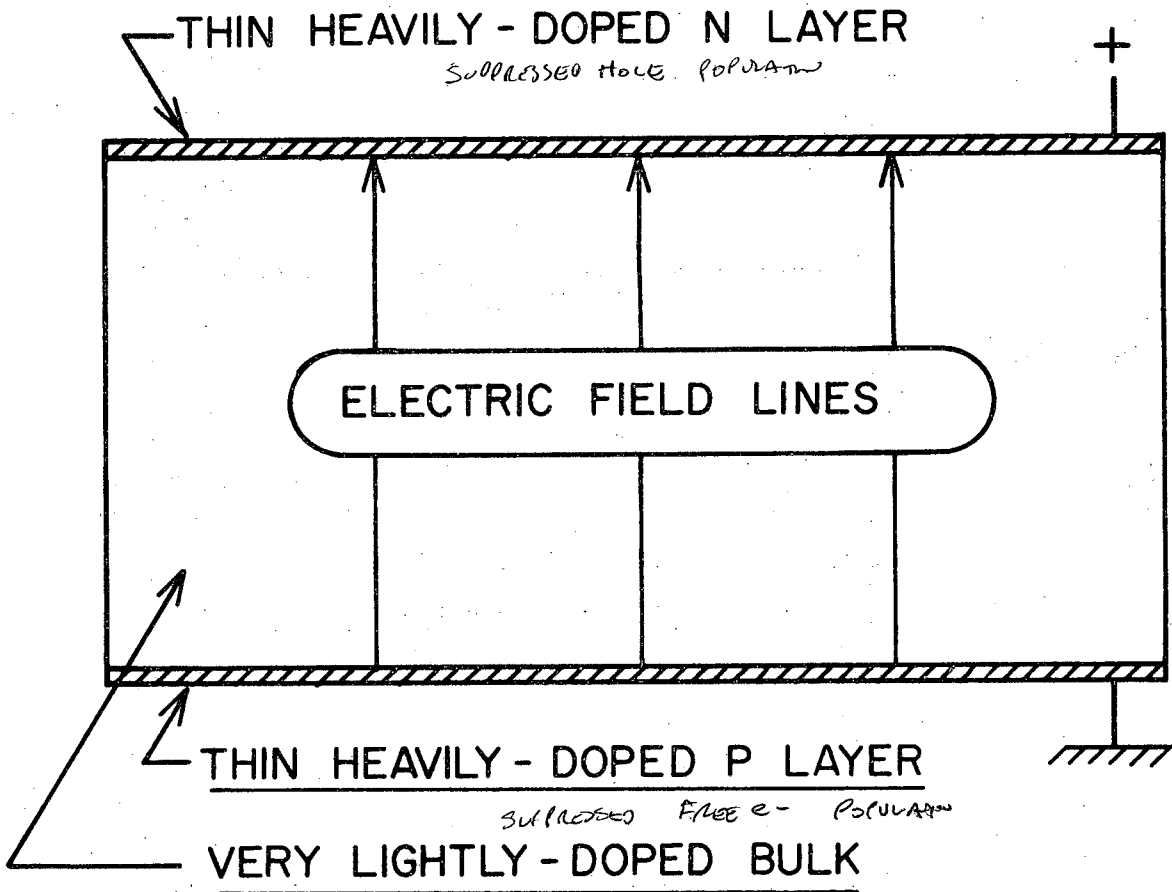
- (1) ϵ , the mean energy required to produce a hole-electron pair.
- (2) F , the Fano factor, a smoothing factor applied to Poisson statistics to obtain the value of the fluctuations in the ionization produced in a material. If the average number of carriers produced is N , the mean square deviation, $\overline{n^2}$, is given by

$$\overline{n^2} = FN.$$

Theoretical (1) and experimental (2) studies of these parameters permit prediction of the value of ϵ for different materials (Fig. 9), and accurate estimates of the energy resolution attainable in semiconductor detectors (Fig. 10). Allowance is made, in the case of heavy particles, for the statistics associated with the atomic-collision processes at the end of the track as analyzed by Linhard (3). No allowance has been made for the effects of noise in the electronic amplifier used with the detector.

The Charge-Collection Process

A detector can be thought of as a block of material with electrical contacts attached to opposite faces, a structure analogous to the parallel-plate gaseous ionization chamber. Free charges produced in the detector are collected by applying a high voltage across the block.



XBL 706-1209

Fig. 11

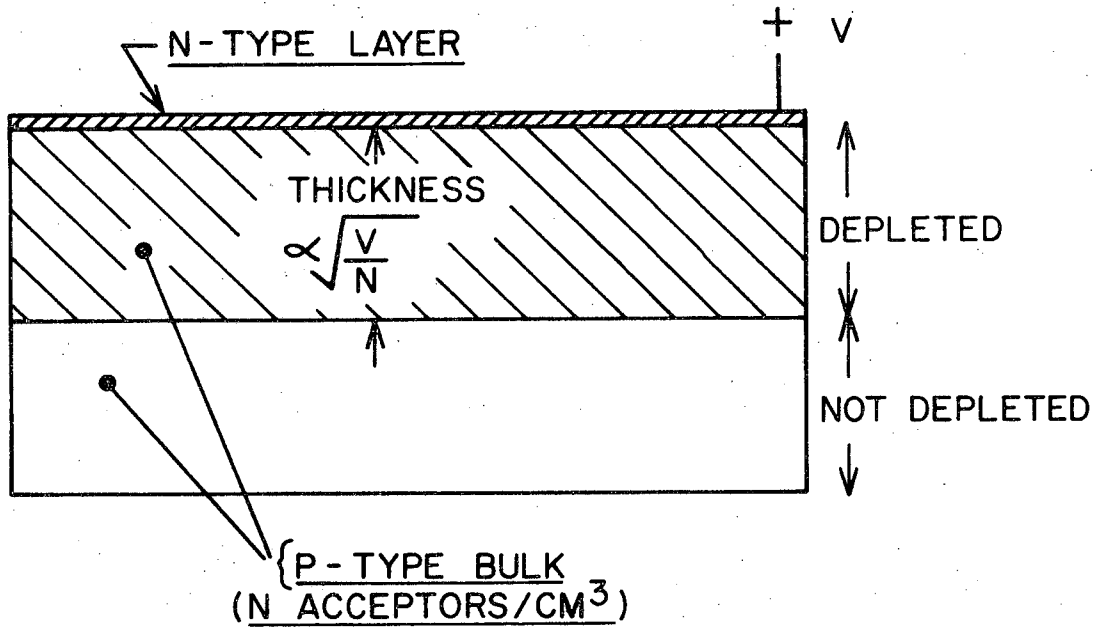
An ideal, fully depleted detector with heavily doped surface layers of opposite types.

The impulse of current flowing in the external circuit while the free charges are in transit measures the energy of the photon or particle producing the signal. This simple picture of the operation of a semiconductor detector focusses attention on two important requirements:

(1) The free charges--electrons and holes--must not recombine or be lost by becoming trapped before being collected; otherwise the total charge flowing in the external circuit will not be a true measure of the original ionization. In other words, the average lifetime of a carrier before recombining, or being trapped, must be long compared with its transit time across the sensitive region, which can range from 10^{-9} sec to almost 10^{-6} sec, depending on the design of the detector. The detector material must be such that the mobility of charge carriers is high, so that their velocity will be high in reasonable electric fields.

(2) Spurious currents in the detector must be very small because signals produced in the external circuit may be minute. As an illustration of the magnitude of the quantities involved, we can now measure 1-keV X-rays, each releasing an average of only 300 electrons, with a resolution (FWHM) of about 100 eV. This implies a RMS error of only about 15 electrons in the whole signal-measuring process, including contributions from the signal amplifier and those arising from spurious charges flowing in the detector. The need to reduce spurious current flow determines the structures chosen for detectors and influences the types and quality of the materials used.

The ideal semiconductor detector consists of a thick slice of very pure material having almost no free carriers of either type, with a positive electrode containing no free positive holes and a negative electrode containing no free electrons. A close approximation to such a structure is shown in Fig. 11. Heavy doping of the n layer almost totally suppresses its hole population; doping the p layer suppresses its free electrons.



XBL 706-1208

Fig. 12

The common junction detector in which the material is not fully depleted.

When voltage is applied to this structure in the direction shown in Fig. 11, an electric field is produced throughout the bulk of the material, with the field lines terminating on the heavily doped surface layers. The direction of the electric field is such as to oppose injection of majority carriers from either contact into the bulk, although aiding injection of the minority (almost nonexistent) carriers. Contacts of the above type reduce the currents injected at the contacts almost to zero. The main sources of leakage in a detector then become charge generation and injection at the side surfaces of the material where the lattice suddenly terminates, and generation of holes and electrons in the main bulk of the material by thermal lattice vibrations that excite carriers from the valence into the conduction band (usually through the intermediary of traps). Lowering the temperature of the device, commonly to liquid-nitrogen temperature (77°K), reduces the current due to thermal excitation although surface effects may still be present. Well-made small silicon detectors, for example, cooled to 77°K and shielded from infrared radiation from the warm walls of the enclosure, exhibit leakage currents as low as 10^{-14} amps, corresponding to less than ten electron-hole pairs flowing across the detector in the few microseconds needed for measuring the ionization induced by radiation.

A less ideal structure is shown in Fig. 12. Here the bulk p-type material is used for one noninjecting contact and the electric field only partially penetrates the material. In so doing, it drives free electrons and holes away from a region called the "depletion layer" whose thickness is proportional to the square root of the applied voltage. The depletion layer forms the sensitive region of the detector, from which the free carriers produced by ionization can be collected.

Detector Materials

Our choice of materials for semiconductor detectors is very limited, principally by charge-collection considerations. For charge-carrier losses to be small during their transit across the detector, crystals that are nearly perfect are required; lattice imperfections can trap charge in transit and cause excessive leakage currents by acting as intermediaries in the thermal excitation of carriers into the conduction band. So far, only silicon and germanium have performed adequately, although cadmium telluride shows promise (4). Materials with constituents of high atomic number are desirable as they are efficient absorbers of gamma rays by the photoelectric process. However, other considerations enter into the choice of material: for example, germanium cannot be used if room-temperature operation is required, because thermal excitation of carriers into the conduction band produces excessive leakage. If operation at still higher temperatures ($>100^{\circ}\text{C}$) is necessary, even silicon is unusable; then silicon carbide or cadmium telluride become possibilities.

Another factor has played a big part in the development of semiconductor detectors: namely, that the highest purity silicon and germanium produced by standard zone-levelling and crystal-growing techniques contain about 10^{12} impurities (either acceptors or donors) per cm^3 . Application of reasonable voltages to a detector of the type shown in Fig. 12, made from this material, produces a depletion layer only 0.5 to 2 mm thick. Such detectors can be used in many applications; but their gamma-ray efficiency is low, and they are not thick enough to stop many of the particles of interest in nuclear physics.

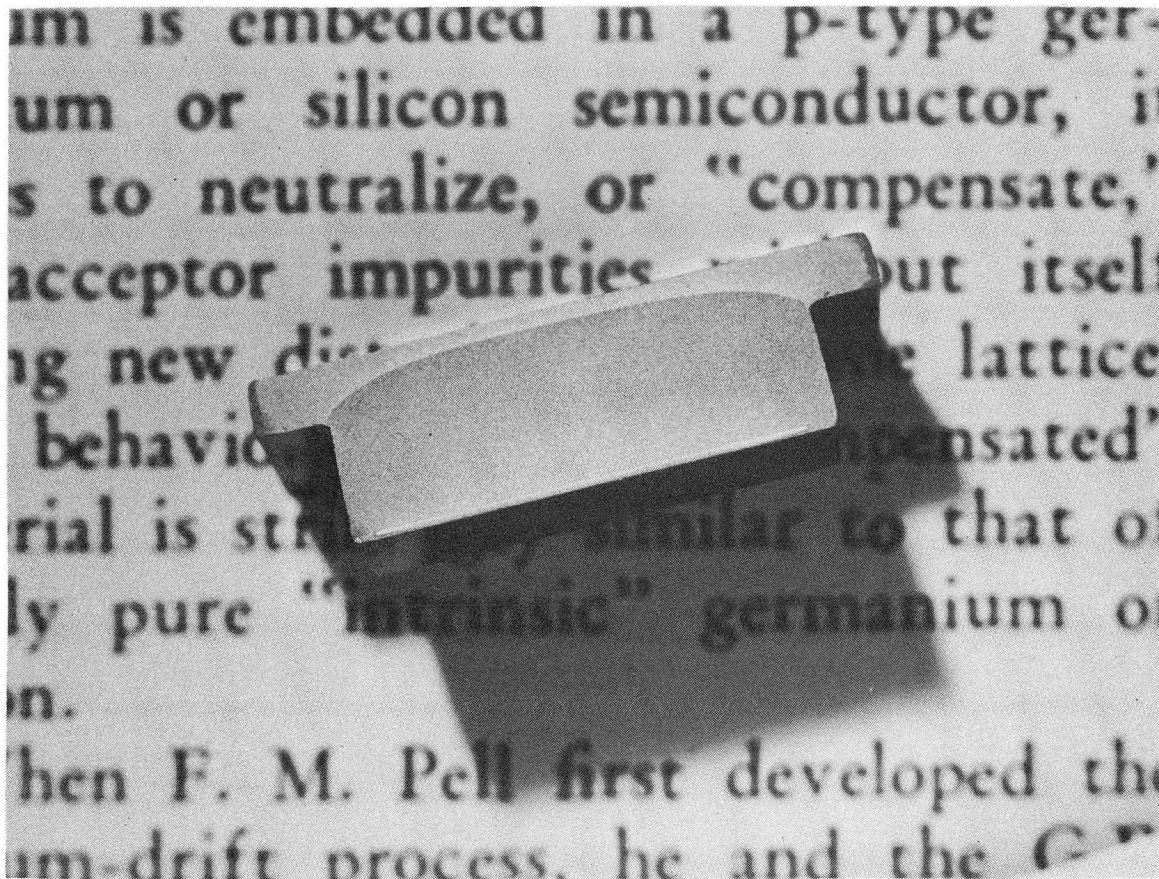
The inherent limitations of this material have been circumvented by "lithium drifting", a process invented by Pell (5) for compensating acceptor impurities with an interstitial donor (lithium). By this method both germanium and silicon can be produced with thick regions

that exhibit almost no net impurity concentration. To make lithium-drifted detectors, lithium is evaporated onto and diffused into the face of a slice of p-type material at about 400°C to produce an n-p junction; then with the temperature held high enough for the lithium ions to be reasonably mobile (20 to 50°C for Ge and 120 to 150°C for Si), a positive voltage (500V) is applied to the lithium-doped side of the junction. Lithium ions drift in the electric field in such a way as to almost exactly compensate the acceptor impurity level in the material; in a week or two, a compensated region about a cm thick is produced (Fig. 13).

Such detectors are now used in a variety of applications; lithium-drifted silicon detectors are particularly important in nuclear-reaction studies with particles produced by accelerators, and in low-energy X-ray work, while lithium-drifted germanium detectors are used mostly for gamma-ray spectroscopy. Germanium detectors having very large volumes can be made using the coaxial-drift technique (6), whereby lithium is drifted from the outer surface of a cylindrical block of germanium toward the middle; sensitive volumes of 50 cc are common, resulting in reasonably high gamma-ray detection efficiencies. In all very-high-resolution work, cooling the detector and the first signal-amplifying stage is necessary; liquid-nitrogen-cooled cryostats are usually employed for this purpose. In other cases thermoelectric coolers are sometimes adequate.

In spite of the value of lithium for compensating impurities, interest is increasing in programs aimed toward producing materials pure enough to make lithium drifting unnecessary, for several difficulties have arisen that are due specifically to the presence of lithium:

- (1) Radiation damage is exaggerated by the tendency of lithium ions to precipitate at damage sites and thereby to become electrically inactive.



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Fig. 13

Photograph of a lithium-drifted region in a detector. The region was revealed by copper plating from a CuSO_4 solution containing a trace of HF with a bright light shining on the solution.

(2) Charge trapping is now believed to be due largely to lithium-defect pairs (7).

(3) Since lithium ions are quite mobile in germanium even at room temperature, handling problems are serious, and storage of lithium-drifted germanium detectors at room temperature is out of the question.

Recent work has shown that germanium crystals can be produced with impurity concentrations substantially below 10^{11} impurity atoms per cm^3 (8). If 10^{10} atoms per cm^3 can be achieved, detectors with sensitive thicknesses of one cm can be made without lithium drifting. This advance will change many aspects of the use of germanium detectors, enabling them to be applied in areas where only silicon has been used so far.

Spectrometer Systems

Fully utilizing the properties of semiconductor detectors has demanded advances in signal amplifiers and the associated electronics (9). In particular, electrical noise in the amplifier, which results in broadening of peaks in pulse-height spectra, has been reduced over the past four years by a factor of ten. This improvement has principally come from the use of field-effect transistors at low temperatures (10). Significant advances have also been necessary in the data-acquisition aspects of nuclear electronics. The fine detail produced by semiconductor detectors has resulted in a large increase in the number of channels in pulse-height analyzers; the large data-flow has made the use of on-line computers almost a necessity.

Semiconductor detectors are now used extensively for spectrometer systems in all areas of nuclear physics. In the following brief account of some of these applications, we have avoided the standard ones and have chosen, instead, a few cases where use of the detectors has resulted in totally new capabilities in physics or other sciences,

and cases that illustrate the present and future potential of these detectors.

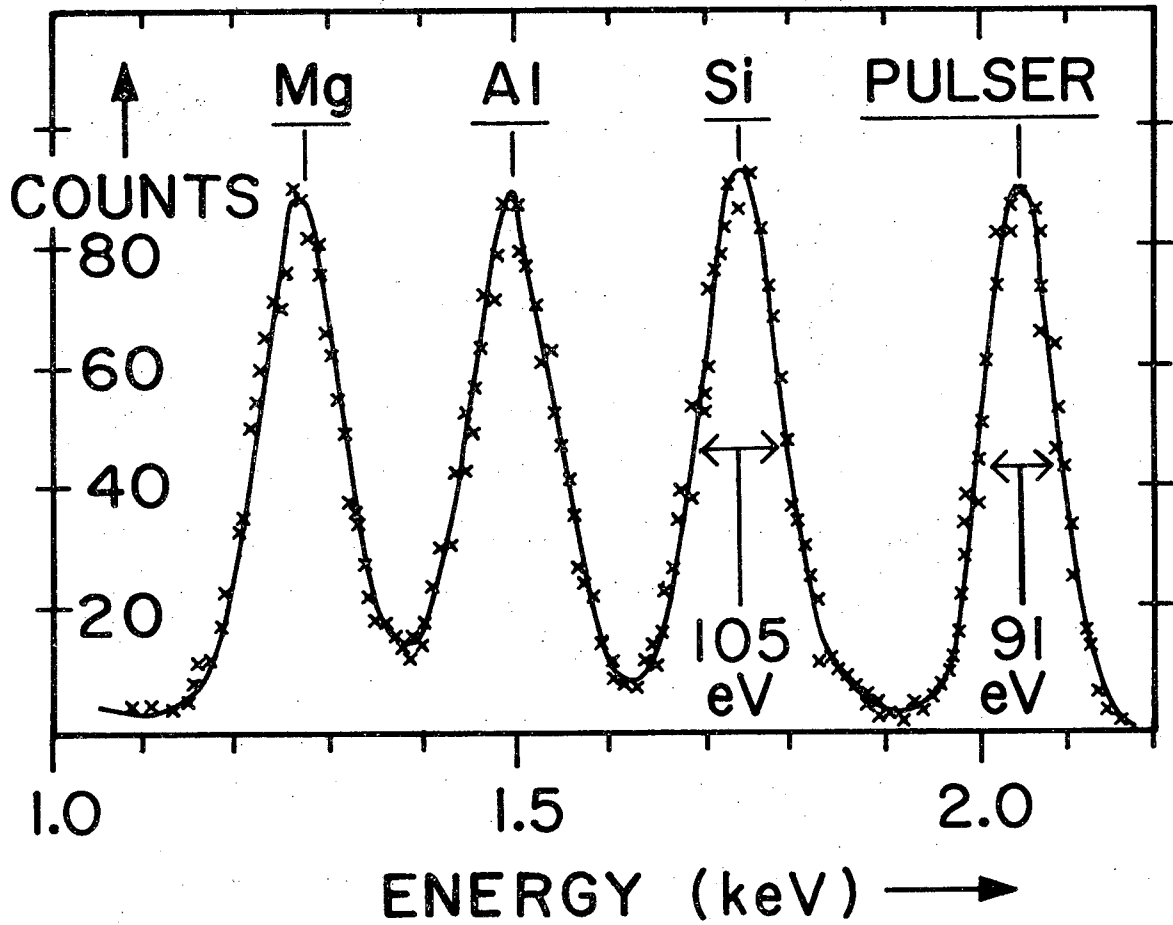
X-Ray Fluorescence Analysis

Determining the chemical elements in a sample by means of their characteristic X-rays has been an analytical tool for many years, but the advent of high-resolution X-ray spectrometers using semiconductor detectors, combined with the availability of low-energy X-ray sources such as ^{55}Fe , have converted an expensive, cumbersome tool into a convenient one. The method relies on radiation to remove electrons from the inner atomic shells of atoms in the sample; when the resulting vacancies fill, X-rays characteristic of the elements in the sample are emitted. The characteristic X-ray spectra are much simpler, and therefore easier to interpret, than optical spectra since only the few electrons in the inner shells are involved. For this type of analysis, semiconductor-detector X-ray spectrometers are now replacing crystal-diffraction spectrometers, giving much higher detection efficiency, and permitting simultaneous observation of a wide range of energies. The increase in detection efficiency makes possible the use of much smaller exciting sources than before--radioactive materials such as ^{55}Fe or ^{241}Am , small X-ray tubes, electron-beam tubes, and small particle-accelerators are all being used to advantage. Since 1965 when semiconductor detectors were first suggested for this application (11), a substantial industry has grown up to exploit the potential of X-ray fluorescence systems for a wide variety of analytical problems.

Using a scanning electron microscope or an electron microprobe for excitation, elemental distributions can be determined in regions as small as one micron. Typical applications include determining the distribution of chemical elements in biological tissue and studies of grain boundaries and microscopic nonuniformities in materials. Using

proton beams, or preferably heavy-ion beams, which are more efficient than X rays or electrons in exciting fluorescence, one can detect sub-nanogram quantities of many elements. A Swedish group has recently described the use of this technique for measuring the elemental composition of particulate matter precipitated from the atmosphere (12). For medical research or diagnosis, an "X-ray photograph" of the thyroid gland can be obtained by scanning the gland with a detector system while exciting the iodine it contains with a suitable gamma-ray source, such as ^{241}Am . Determining the composition of pottery sherds has helped archeologists to trace the trade routes of ancient peoples. X-ray fluorescence analysis has even been used to check the authenticity of "old" paintings, by nondestructive analysis of pigments, and also to detect fake coins.

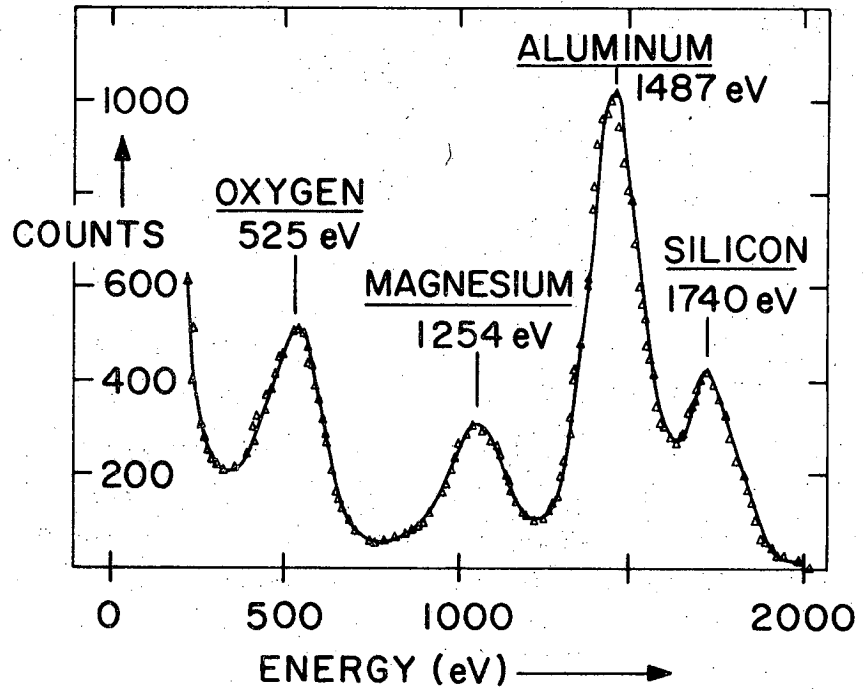
The latest performance capabilities of detectors for X-ray analysis are illustrated in Figs. 14 and 15. In the first of these spectra (13), separation of the X-rays of magnesium, aluminum, and silicon is shown, together with a peak produced by a constant-amplitude electronically generated pulse. The electronic-pulse peak shows the contributions of electronics to the resolution, while the difference between it and the X-ray peaks indicates the charge statistics and collection properties of the detector. The spectrum in Fig. 15 illustrates the abilities of a silicon X-ray detector at very low energies (14). Here the oxygen X-ray (525 eV) produced by X-ray excitation of a Al_2O_3 target is clearly separated from noise counts at the bottom end of the spectrum, despite the difficulty of producing characteristic X-rays in elements of low atomic number. Detection of carbon X-rays appears likely to be achieved in the very near future. For measurements of the characteristic K X-rays of the heavier elements, thin-window germanium detectors must be used since the efficiency of silicon detectors falls rapidly for X-rays with energies greater than 30 keV. In this energy range, and with good low-noise amplifiers, charge-production statistics determine the energy resolution of the system, as shown in Fig. 10.



XBL 706-1197

Fig. 14

Fluorescence spectrum of X-rays from a target containing Mg, Al, and Si as seen by a silicon detector. The exciting source was a small X-ray tube (13).



XBL 706-1207

Fig. 15

Fluorescence spectrum of X-rays from a Al_2O_3 target bombarded with Al X-rays from the Al anode of a small X-ray tube. The target was contained in the same vacuum chamber as the detector so that no absorbing material was present between the target and the detector (14).

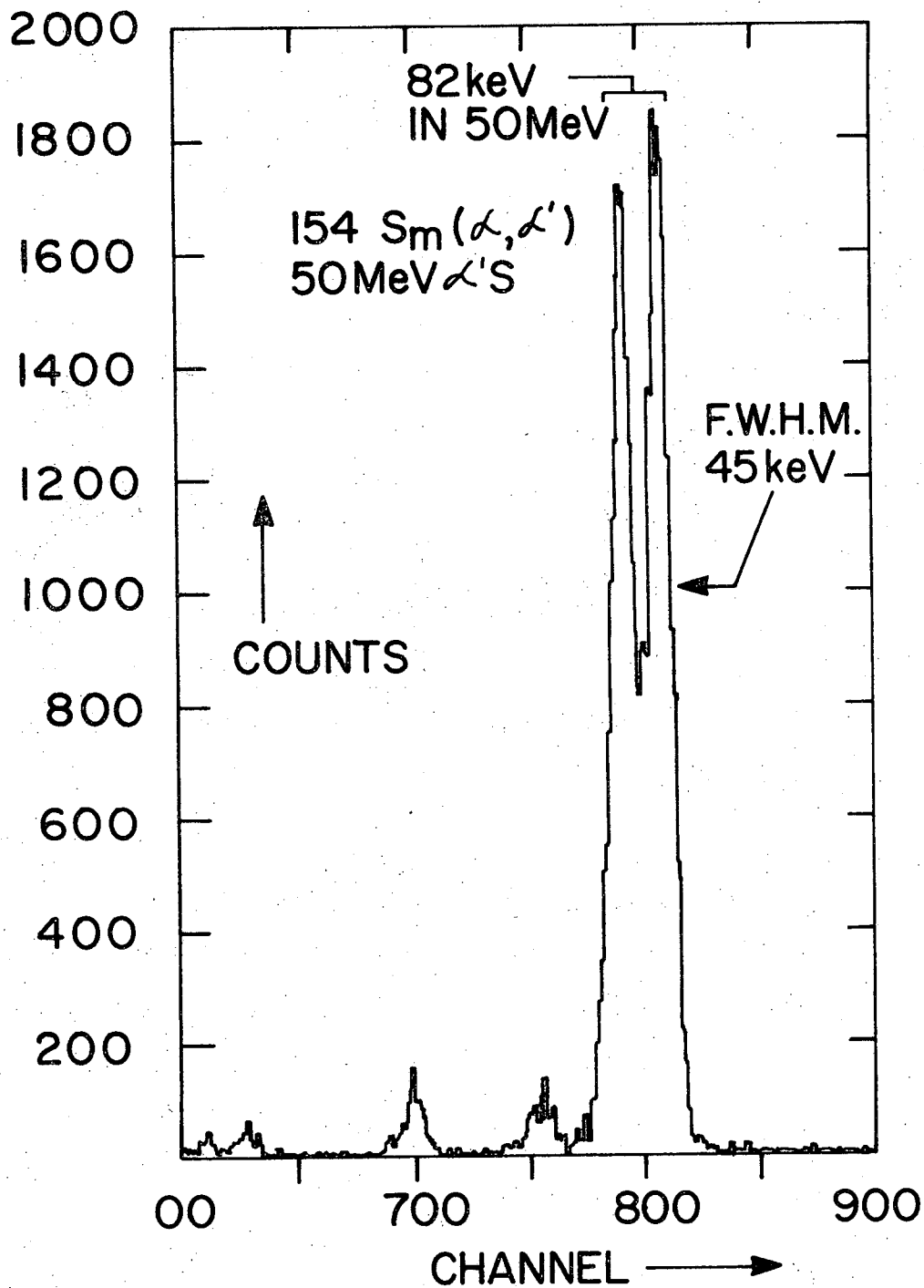


Fig. 16

XBL 706-1201

The spectrum of alpha particles scattered from a ^{154}Sm target. (Beam energy 50 MeV.)

High-Energy Nuclear Spectroscopy

Many of the experiments at cyclotrons and Van de Graaff accelerators involve bombarding a target material with a beam of particles of fairly high energy (~ 50 MeV) and measuring the energy and angular distribution of the particles that emerge. Studies of this nature serve to elucidate the structure of nuclei, and their energy levels. Fig. 16 shows the energy spectrum of emerging alpha particles, observed with a silicon detector at a fixed angle when a ^{154}Sm target was bombarded with 50-MeV alpha particles. The peak of higher energy is due to alpha particles elastically scattered from ^{154}Sm nuclei; the almost equally intense peak lying 82 keV below the main peak is caused by alpha particles whose energy has been reduced by giving up 82 keV to excite the samarium nuclei.

The energy resolution of the system used in this experiment was just adequate to separate the peak due to the 82-keV excited state from that due to the elastic group. Much better energy resolutions (20 keV FWHM at 50 MeV, i.e., 0.04%) have been obtained in very carefully controlled experiments where the beam quality, target thickness, and slit scattering were all ideal, where the detector was carefully selected and operated at constant low temperature, and where the associated electronics was used under optimum conditions.

It is very difficult to make silicon detectors of adequate thickness to detect particles having a longer range, such as 50-MeV protons. Entry of particles into the side of a detector has been used, but the variable energy-absorbing layer at the surface worsens the energy resolution. Experiments using germanium detectors for long-range particles have been fairly successful--resolutions of 18 keV have been obtained for 42-MeV protons--but radiation damage and the problems associated with handling and cooling germanium detectors have inhibited their use. However, the development--probably quite soon--of very

pure germanium requiring no lithium drifting may well change the situation drastically. The use of germanium detectors for even higher energies has been demonstrated (15) and promises exciting applications in the future.

Measurement of Very Short Decay Lifetimes

Although nuclear reactions produce excited states with lifetimes ranging from 10^{-20} sec to many years, standard electronic timing methods permit determination only of those longer than 10^{-9} sec. A novel method for determining some lifetimes in the range 10^{-13} to 10^{-9} sec utilizes the high-resolution capabilities of a germanium gamma-ray detector system to measure the Doppler shift in the energy of gamma rays emitted by nuclei that are in high-velocity flight. The method, proposed by Litherland and described by Alexander and Allen (16), has been used, for example, to study the lifetime of the 871-keV state of ^{17}O produced by bombarding a deuterium target with 20-MeV ^{16}O . In the $d(^{16}\text{O}, p\gamma)^{17}\text{O}$ reaction, the emerging ^{17}O nuclei travel predominantly in a narrow forward cone. A plunger stops them at an adjustable distance (ranging from 20 microns to about 2 cm) from the target. A germanium gamma-ray detector, situated in a backward direction with reference to the beam, observes the gamma rays emitted when the excited ^{17}O nuclei decay to their ground state. If decay occurs after the nucleus is stopped, the detected gamma-ray energy is 871 keV; the energy is less if decay occurs while the nucleus is in flight (Fig. 17). By measuring the ratio of the intensities of the two gamma rays, and knowing the fragment velocity and the distance from target to plunger, one can determine the lifetime of the excited level in ^{17}O .

This technique has been extended by the same experimenters to even shorter lifetimes by accurately observing the shape of the gamma-ray peak that results when a nuclear species stops inside a target.

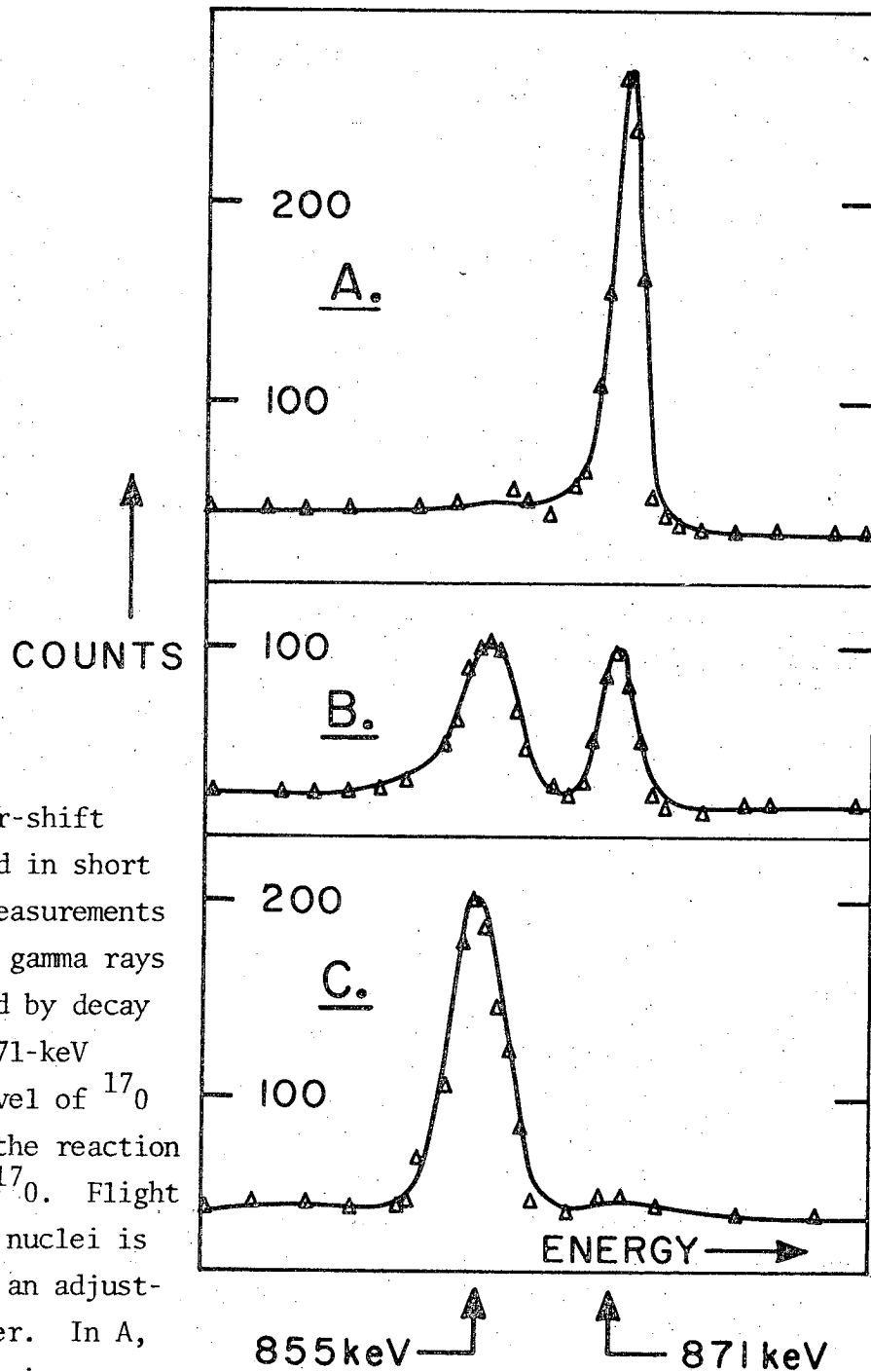


Fig. 17

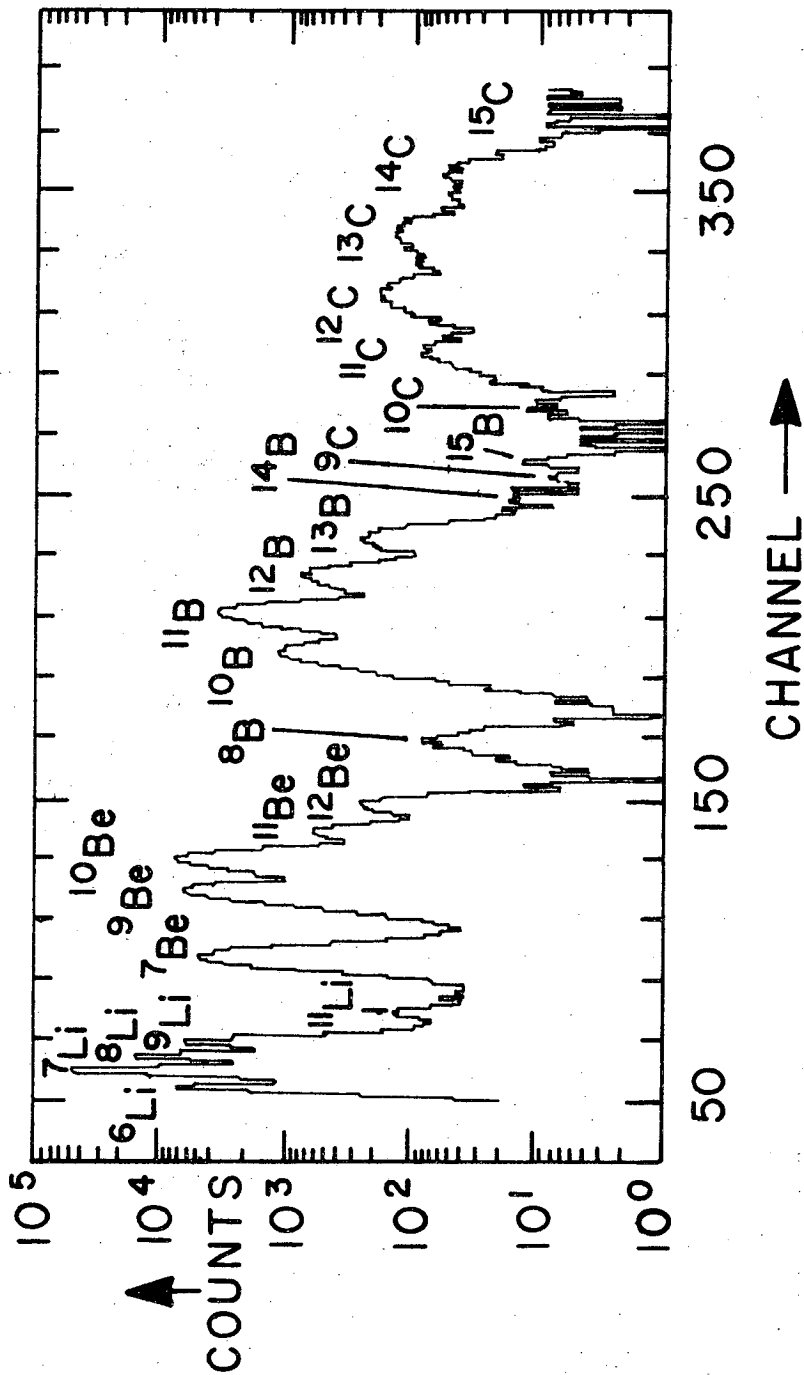
The Doppler-shift effect used in short lifetime measurements (16). The gamma rays are emitted by decay from the 871-keV excited level of ^{17}O formed in the reaction $d(^{16}\text{O}, p\gamma)^{17}\text{O}$. Flight of the ^{17}O nuclei is stopped by an adjustable plunger. In A, the plunger is very

close to the target and nearly all decays occur with the ^{17}O nuclei at rest. In B, the plunger is 2.54 mm away from the target. About 50% of the decays occur while the nuclei are flying away from the detector, producing a Doppler shift to lower energies in the gamma-ray spectrum. In C, the plunger is 12.5 mm away from the target; nearly all gamma rays are then Doppler shifted.

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Fig. 18

The spectrum of isotopes produced in fragmentation of uranium nuclei by 5-GeV protons as observed by a detector telescope and particle identifier. This spectrum represented the first observation of the short-lived isotopes ^{11}Li , ^{14}B , ^{15}B (18).



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Correlation of the observed and the calculated peak-shapes gives information on the lifetime. The method has also been used in studies of the types of fragments produced in nuclear fission.

Particle Identification

High-energy nuclear reactions generally produce many types of particles (Fig. 18), making it necessary to select the specific reaction products desired for study. Passing each product particle through a very thin silicon transmission detector (often less than 20 microns thick) generates a signal proportional to the rate at which the particle loses energy in silicon; absorbing the remainder of its energy in a second detector, and suitably processing the two signals, permits unique identification of each particle up to a mass number of about ten (17).

For mass numbers above ten, this identification is not unique: signals due to the heaviest isotopes of one element overlap those due to the lightest isotopes of the next. To eliminate this uncertainty, the first transmission detector in a telescope is moved some distance in front of the remaining detectors, and the time of flight of the particles between detectors is measured. The additional information permits unique identification of mass numbers through twenty (18).

Mesonic X-Rays

When a target material is placed in a flux of negatively charged mesons, hybrid atoms are formed in which a meson replaces an electron and goes into orbit very close to the nucleus, far inside the inner shell of atomic electrons. Because mesons are so much heavier than electrons, some of these orbits are so small that they overlap the nucleus itself, acting as a sensitive probe of the fields close to the nucleus and of the nuclear surface. Germanium and silicon

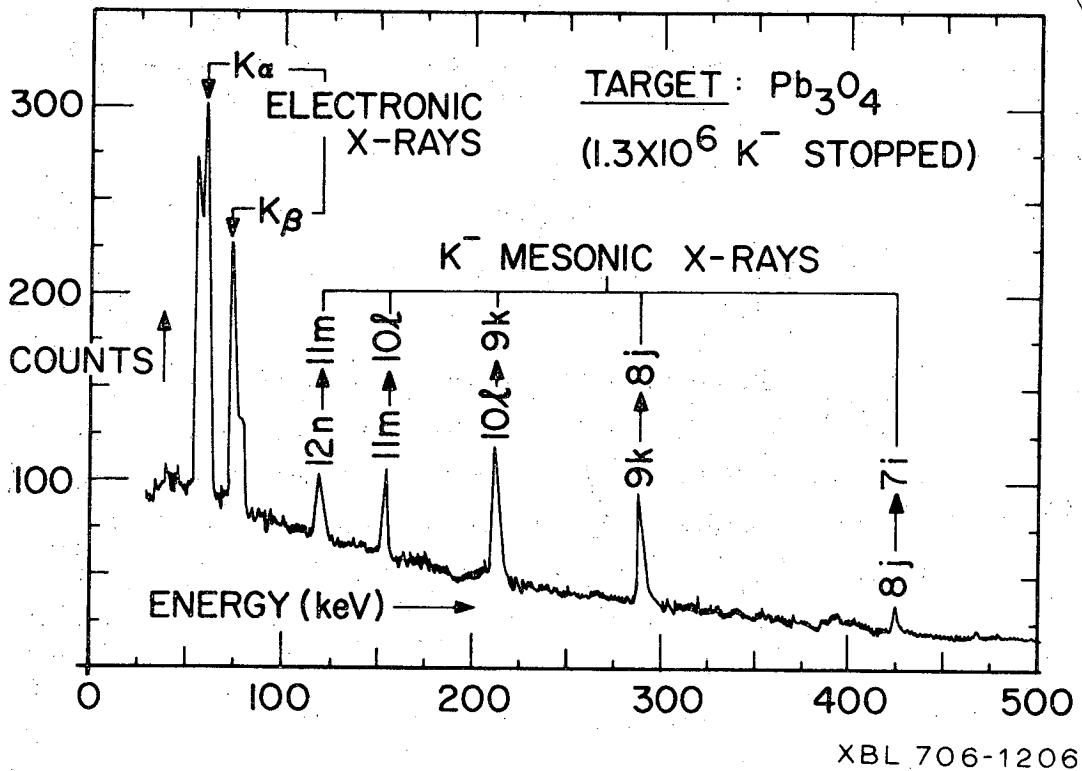


Fig. 19

K $^-$ -mesonic X-ray spectrum for lead produced by kaon bombardment of a lead-oxide target (19). The highest energy X-ray seen represents a transition to the innermost kaon orbit outside the nuclear surface. The transition involved in producing the X-rays are indicated on the figure.

detectors give adequate energy resolution to observe the X-ray spectrum generated as these mesons cascade inward, from orbit to orbit, toward the nucleus.

Recent work has studied π^- -mesonic and K^- -mesonic atoms, and Σ^- -hyperonic atoms, in isotopes covering a wide range of atomic number. X-rays from K^- -mesonic atoms have yielded interesting data on the distribution of nuclear matter near the surface of nuclei and have shown that a "halo" of nuclear matter extends well beyond the conventional nuclear radius (19). As the K^- -mesons fall inward, the X-rays produced are seen as a series of lines of increasing energy in the spectrum recorded by the semiconductor-detector system (Fig. 19). Finally, as the meson reaches the nucleus, it is absorbed by a surface particle, and the X-rays suddenly stop. Hence the X-ray line having the highest energy indicates the energy of the innermost transition; a relatively simple calculation yields the radius of the orbit closest to the nucleus. By carrying out this measurement for a range of nuclei of slowly increasing atomic number, the nuclei for which a given X-ray line is just "snuffed out" can be determined, which gives the precise radius of these nuclei.

Activation Analysis

For many years nuclear reactors have been used to "activate" materials. After capturing slow neutrons from the reactors, materials can be analyzed for trace elements by observing the gamma rays emitted upon decay of excited nuclear states. Other methods of activation are also possible, and have specific advantages for the detection of certain elements; for example, ^3He activation is a powerful tool for detecting traces of oxygen in materials.

Until the advent of germanium gamma-ray detectors, decay energies could be observed with reasonable efficiency only by using scintillation detectors. However, as can be seen by comparing the gamma-ray

spectra of Fig. 1, germanium detectors increase by a large factor our ability to analyze complex samples, and therefore have made a significant difference to all types of activation analysis. Recently, these methods have been applied to the analysis of pottery and other artifacts of interest to archeologists (20).

Conclusion

The past decade has seen the rapid development and exploitation of one of the most significant tools of nuclear physics, the semiconductor radiation detector. Applications of the device to the analysis of materials promises to be one of the major contributions of nuclear research to technology, and may even assist in some aspects of our environmental problems. In parallel with the development of these applications, further developments in detectors for nuclear research are taking place: the use of very thin detectors for heavy-ion identification, position-sensitive detectors for nuclear-reaction studies, and very pure germanium for making more satisfactory detectors for many applications suggest major future contributions to physics.

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