RADIOISOTOPE DETECTION WITH THE ARGONNE FN TANDEM ACCELERATOR*

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ABSTRACT. A standard heavy ion nuclear structure facility has been used to detect several long-lived radioisotopes by counting the accelerated ions. The problem of eliminating strong isobaric background beams has been solved by combining the energy loss dispersion through a uniform Al foil stack with the high momentum resolution of an Enge split-pole magnetic spectrograph. Radioisotope concentrations in the following ranges have been measured: ${}^{14}C/{}^{12}C = 10^{-12}$ to 10^{-13} , ${}^{26}Al/{}^{27}Al = 10^{-10}$ to 10^{-12} , ${}^{23}Si/Si = 10^{-8}$ to 10^{-14} , ${}^{36}Cl/Cl = 10^{-8}$ to 10^{-11} .

INTRODUCTION

The Argonne National Laboratory has operated an electrostatic FN tandem accelerator since 1967. In 1973 and 1977, some essential parts of the accelerator were replaced in order to meet the stringent requirements as an injector for the newly developed super-conducting linear accelerator. The conversion of the tandem to metal-bonded ceramic-titanium accelerator tubes with straight electrode geometry, a pelletron charging system and an enclosed corona-tube voltage distribution system, and the use of pure SF₆ insulating tank gas resulted in a machine of ion-optic simplicity, high voltage stability up to 9 MV terminal voltage, and an ultra high vacuum system ($\approx 10^{-8}$ Torr). At present, both the tandem alone and the combined system tandem-linac are almost entirely devoted to heavy ion nuclear physics research. Negative heavy ion beams are routinely produced from an "inverted" cesium-beam sputter source (Chapman, 1976), which can also take small sputter samples of a few milligrams. A major analyzing instrument for heavy ion spectroscopy at the tandem is an Enge split-pole magnetic spectrograph with an ionization chamber focal-plane detector (Erskine, Braid, and Stoltzfus, 1976). This instrument is well suited to identify tandem-accelerated ions by mass and atomic number up to mass 40 and with some improvements probably up to mass 60.

The conditions described above encouraged us to start experiments in the newly revived field of accelerator mass spectroscopy, which is exactly 40 years old (Alvarez and Cornog, 1939a,b). Reviews on the recent developments are given by Muller (1979), Purser, Litherland, and Gove (1979), and Bennet (1979). Our own activity in this field started about nine months ago. The intention was to use as much of the existing experimental facility as possible in order to fit the measurements smoothly into the nuclear physics research program. In addition, rather than concentrating only on ¹⁴C and trying to set up a system for radiocarbon dating, we aimed towards developing a technique applicable for a large variety of radioisotopes.

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Accelerator Techniques

One of the major obstacles in accelerator spectroscopy is the sometimes overwhelming background from stable isobaric beam components. Fortunately, this problem does not exist for ¹⁴C measurements with the tandem since ¹⁴N does not form stable negative ions. However, the detection of various other radioisotopes of interest suffers severely from 36 36the problem. Examples are the isobaric pairs Cl - S and, in partic-

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ular, Si - Si = S. We will describe briefly our technique to solve this 14 16

problem, below. A more detailed description will be given elsewhere (Henning and others, ms in preparation).

EXPERIMENTAL TECHNIQUE

Figure 1 shows a schematic view of the essential parts of the radioisotope detection system at Argonne. It illustrates the measurement of 36

Cl in the presence of a S background. The basic concept of our 17 16 following here is a pain through a uniform

method is to disperse the energy of the isobaric pair through a uniform foil stack (Clerc and others, 1975) and measure the energy difference



Fig 1. Schematic illustration of a ³⁶Cl measurement with the radioisotope detection system at Argonne. Additional negative-ion background components not shown in the figure are ${}^{18}O_2^{-}$ and ${}^{12}C_2^{-}$.

808

accurately with a magnetic spectrograph. The energy-loss difference due to different atomic numbers, results, then, in energy- and spatiallyseparated components at the focal plane of the spectrograph. If the background lines are high in intensity, they can be conveniently shielded from the focal plane detector. This turned out to be an important option in some cases since the resolution of the detector starts to deteriorate if the total counting rate exceeds 1000 counts/sec. The detector provides additional energy-loss, total energy, and position (B ρ) signals which allow a clear identification of mass and atomic number.

The separation through the foil stack is only useful if the energy straggling is small compared to the energy-loss difference. We investigated several foil materials, including C, Al, Si, and Au in different foil arrangements. Single thick foils (1 to $4mg/cm^2$) and stacks with up to 40 thin foils were studied. Non-uniformity and pinholes were the major obstacles. The best peak-to-valley ratio for adjacent components and an energy



Fig 2. A) Gated position $(B\rho)$ spectrum from an NaCl sample with ${}^{36}Cl/Cl \approx 10^{-8}$ as measured in the focal-plane detector of the spectrograph. Window set on the $\Delta E-E$ spectrum of figure 3. B) Gated position spectrum with an additional window set on Cl^{13+} ions as shown in figure 3.

straggling close to the one expected from theory was obtained for a stack of 15 Al foils, each $100\mu g/cm^2$ thick. All measurements, except for ¹⁴C for which a thin carbon foil is sufficient, have been performed with this Al stack. Figure 2 shows the total position spectrum in the spectrograph for an incident beam of 88MeV ³⁶Cl⁹⁺ + ³⁶S⁹⁺. The difference in energy loss for the two components is 2.5MeV and corresponds to a spatial separation of 4cm. Additional information is provided by the data in the ΔE vs E and E vs B ρ plane (fig 3). Appropriate windows in these spectra clean up the position spectrum as demonstrated in the lower part of figure 2.

Measurement of the radioisotope concentration from a sample in the ion source, requires a reliable normalization to the beam current of the stable isotope(s). We decided to make this measurement as close to the spectrograph as possible. For this purpose, a small Faraday cup with suppressor ring and a 3mm entrance aperture was interchanged in posi-



Fig 3. ΔE vs E and E vs $B\rho$ spectra for the same sample as in figure 2. Windows set in these spectra are indicated. The low-lying group in the ΔE vs E spectrum is due to lighter beam contaminants (probably carbon and oxygen).

810

tion with the energy-loss foil stack in front of the spectrograph (fig 2). The whole focusing system from the ion source to the spectrograph was set up first with the stable-isotope beam by requiring the current on the entrance aperture of the Faraday cup to be close to zero. Then the 40° inflection magnet (see fig 1) was set to the radioisotope mass and the terminal voltage was changed in such a way as to match the magnetic rigidity of the stable-isotope beam. None of the other beam-optical elements were touched. The final settings of the inflection magnet and the terminal voltage were verified by scanning in small steps around the precalculated values and optimizing the counting rate of the radioisotope in the spectrograph. The terminal voltage control was provided by a generating voltmeter (Brookshier, Wallace, and Langsdorf, 1971). The entrance and exit slits of the 90° analyzing magnet were wide open. An exception was the run with 32Si, when the 32S background beam was strong enough to allow tandem stabilization by slit control. In these measurements, the energy was kept constant and the complete accelerator system was adjusted separately for ³²Si + ³²S and ³⁰Si by optimizing the beam current in the spectrograph Faraday cup.

Until now, all changes were made manually as in the routine operation of the tandem. It takes approximately 5 to 10 minutes to switch between radioisotope and stable isotope running condition. Although a

Radio- isotope	Sample	Ratio	Measured (10 ¹²)	Expected (10 ⁻¹²)
чC	Charcoal "Fermi" graphite** Graphite	¹⁴ C/ ¹² C ¹⁴ C/ ¹² C ¹⁴ C/ ¹² C	$egin{array}{r} 1.06 \pm 0.15 \ 4.5 \pm 0.7 \ 0.20 \pm 0.05 \end{array}$	$\begin{array}{c} 1.144 \pm 0.011 * \\ 5.137 \pm 0.041 * \\ \mathrm{low} \end{array}$
²⁶ Al	$^{27}Al(\gamma, n)$ $^{27}Al(^{16}O, ^{17}O)$ ^{27}Al	²⁶ Al/ ²⁷ Al ²⁶ Al/ ²⁷ Al ²⁶ Al/ ²⁷ Al	$290 \pm 60 \\ 91 \pm 30 \\ \smile 4$	$410 \pm 50*** \ \simeq 100 \ \mathrm{low}$
³² Si	Si (t, p) Si (¹⁸ O, ¹⁶ O)	⁸² Si/Si ⁸² Si/Si	-10^4 0.095 ± 0.050	$10^{3}-10^{4}$ < 0.1 §
⁸⁶ C <i>l</i>	$\begin{array}{l} \operatorname{NaC}l\left(\mathrm{n},\gamma\right)\\ \operatorname{NaC}l\left(\mathrm{n},\gamma\right)\\ \operatorname{NaC}l\left(\mathrm{n},\gamma\right)\\ \operatorname{NaC}l\end{array}$	²⁶ Cl/Cl ³⁶ Cl/Cl ³⁶ Cl/Cl ³⁶ Cl/Cl	$\begin{array}{c} 14,\!900 \pm 2200 \\ 1360 \pm 200 \\ 100 \pm 20 \\ 60 \pm 20 \end{array}$	$\begin{array}{c} 12,\!900\pm700 \\ 1450\pm100 \\ 36.8\pm2.0 \\ 10 \\ 10 \end{array}$

 TABLE 1

 Summary of radioisotope measurements

* Measured from β activity with a sample from the same material by D D Coleman, Illinois State Geological Survey, Urbana.

** Graphite probably originating from the first reactor pile in Chicago built by E Fermi in 1942.

*** Al beam stop from the 20MeV electron linac in Argonne. ${}^{20}Al$ content determined independently from activity (1.8MeV γ -ray) and ${}^{27}Al(\gamma, n)$ cross-section.

+ Al sample irradiated with 1×10^{17} ¹⁰O ions (56MeV) from the tandem. ²⁰Al content estimated assuming $\sigma({}^{19}O, {}^{17}O) = 1$ mb. **Part** (1.3mg) of a K₂SiF₆ sample produced by D Alburger and G Harbottle,

[‡] Part (1.3mg) of a K₂SiF₆ sample produced by D Alburger and G Harbottle, Brookhaven National Laboratory, from the irradiation of ³⁰SiO₂ with 2.58 × 10¹⁹ tritons (3.4MeV). ³²Si content estimated from β activity.

§ Si single crystal irradiated with $4 \times 10^{15^{-18}}$ O ions (56MeV) from the tandem. ²²Si content estimated assuming σ (1⁸O, 1⁹O) = 1mb.

¶ NaCl samples irradiated in the Argonne CP5 reactor. ³⁸Cl content determined from ²⁴Na activity, ²³Na(n, γ) and ³⁵Cl(n, γ) cross-section.

computerized system might improve the precision in the switching procedure, other factors such as the current measurement of the stableisotope beam and the background subtraction are the dominant contributions to the errors given in table 1.

RESULTS

Table 1 summarizes the results of our measurements. As mentioned previously, our aim was to determine how well our method works for a variety of radioisotopes. We shall briefly discuss each radioisotope.

¹⁴C. The charcoal sample originated from a piece of wood burned in a fireplace. Agreement with the β activity measurement is satisfactory. The so-called "Fermi" graphite sample was obtained from stock graphite at Argonne. Originally intended to be used for background determination, it turned out to be fairly radioactive. If all the ¹⁴C was produced from ¹³C(n, γ), this graphite must have been exposed to a total thermal neutron



Fig 4. Gated position spectrum from the three Al samples (see table 1). Windows have been set on ${}^{26}Al^{11+}$ and ${}^{29}Al^{10+}$ ions, respectively. Some ${}^{27}Al$ probably originating from double charge exchange in the tandem can be seen.

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flux of 10^{17} neutrons/cm². Another graphite sample showed a relatively high ¹⁴C background. It seems unlikely that this is the inherent background from the ion source. Typical running conditions were: ¹²C⁻ = 1 μ A, TV (terminal voltage) = 6 MV, ¹³C⁴⁺ = 15 nA in the spectrograph cup (used for normalization).

²⁶Al. Figure 4 shows ²⁶Al spectra measured for the three Al samples. Typical running conditions were: ${}^{27}Al^{-} =$ few nA, TV = 8 MV, ${}^{27}Al^{8+} = 2$ nA. The spectra shown in figure 4 were obtained in about 30 minutes each. The low background observed in the blank sample suggests that measurements below 10^{-12} are feasible. However, for such measurements, it is desirable to increase the output of the ion source by at least an order of magnitude. No indication of the existence of ${}^{26}Mg^{-}$ has been observed.

³⁶Cl. Several samples with known ³⁶Cl concentrations were prepared. Results on the highly enriched samples agree well with expected values. The origin of the ³⁶Cl content in the blank sample is not known. The NaCl material was taken from a window for infrared spectroscopy. Typical running conditions were: ³⁷Cl⁻ = 40 nA, TV = 8 MV, ³⁷Cl¹⁰⁺ = 1.5 nA. For the highest enriched sample, the ³⁶Cl¹³⁺ counting rate in the focal-plane detector was 10 counts/sec while the ³⁶Sl³⁺ rate was about twice as high (see fig 2).



Fig 5. Gated position spectrum of ${}^{32}Si^{8+}$ ions from the K_2SiF_6 sample (see table 1). A 4cm wide Ta slit in front of the focal-plane detector shields the ${}^{32}S$ background. Windows are set on ${}^{32}Si^{8+}$ ions (see fig 6).

³²Si. All Si samples produced a ³²S⁹⁺ current between 0.03 to 0.3 nA at the spectrograph cup. Since this corresponds to 107 to 108 32S ions, careful shielding of the focal-plane detector is mandatory. Even so, the tails of the sulfur peaks forced us to select a low intensity charge state emerging from the $\overline{A}l$ foil stack. We chose the ³²Si⁸⁺ component which contains only 0.76 percent of the incident ³²Si⁹⁺ ions (80MeV). Figure 5 shows the ³²Si peak as obtained with the K₂SiF₆ sample [see footnote ‡ in table 1]. A total of 122 ³²Si⁸⁺ ions were counted in 56 minutes. A normalizing current of only 0.074 nA of ³⁰Si⁹⁺ was measured in the spectrograph cup. The isotopic abundance of ³⁰Si in this sample was approximately 5 percent. A background sample of enriched K₂³⁰SiF₆ material (95.6 percent ³⁰Si) produced ≈500 times higher ³⁰Si currents. In 15 minutes, zero counts were observed in the region for ³²Si⁸⁺ ions (see fig 6) which corresponds to an upper limit of ${}^{32}Si/Si < 5 \times 10^{-12}$. The highest source current, 10 μ A of ²⁸Si⁻, was obtained from the Si single crystal sample [see footnote § in table 1]. From this sample with natural Si abundance (ie, 3.1 percent ³⁰Si) a normalizing current of 35 nA ³⁰Si⁹⁺ was measured in the spectrograph cup. Since the 32S background was lowest with this sample, we measured ³²Si⁹⁺ ions (6.7 percent charge state probability) emerging from the Al foil stack. In 33 minutes, 10 counts of ³²Si⁹⁺ ions were detected. The ³²Si/Si ratio from these values (table 1) indicates that measurements in the 10⁻¹⁴ range are possible for ³²Si under favorable conditions.

CONCLUSION

It appears that accelerator mass spectroscopy can be performed at a tandem accelerator used normally for heavy ion nuclear physics research without major additional investments in equipment. The precision and sensitivity obtained with the described technique should allow the investigation of a number of interesting problems.



Fig 6. ΔE vs E spectra for the K₂SiF₆ samples. The irradiated sample (left) shows a clear ³²Si peak, whereas the background sample (right) has zero counts in the same region.

814

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