Accelerator Mass Spectroscopy (AMS) is a highly sensitive technique that is useful in isotopic analysis of specific elements in small samples (1mg or less of sample containing 10^6 atoms or less of the isotope of interest).\[1\]

**Accelerator Mass Spectroscopy**

AMS requires a particle accelerator, originally used in nuclear physics research, which limits its widespread use due to high costs and technical complexity. Fortunately, UC Davis researchers have access to the Lawrence Livermore National Laboratory Center for Accelerator Mass Spectrometry (CAMS LLNL), one of over 180 AMS research facilities in the world. AMS is distinct from conventional Mass Spectrometry (MS) because it accelerates ions to extremely high energies (millions of electron volts) compared to the thousands of electron volts in MS (1keV=1.6×10^{-16} J). This allows AMS to resolve ambiguities that arise in MS due to atomic and molecular ions of the same mass. AMS is most widely used for isotope studies of $^{14}$C, which has applications in a variety of fields such as radiocarbon dating, climate studies, and biomedical analysis.\[2\] Some of the most fascinating applications of AMS range from exposure dating of surface rocks, $^{14}$C labeled drug tracer studies, and even radiocarbon dating of artifacts such as the Shroud of Turin and the Dead Sea Scrolls.\[3\]

**Theory**

In conventional atomic mass spectrometry, samples are atomized and ionized, separated by their mass-to-charge ratio, then measured and/or counted by a detector. Rare isotopes such as $^{14}$C present a challenge to conventional MS due to their low natural abundance and high background levels. Researchers were challenged by isobaric interference (interference from equal mass isotopes of different elements exemplified by $^{14}$N in $^{14}$C analysis), isotopic interference (interference from equal mass to charge isotopes of different elements), and molecular interference (interference from equal mass to charge molecules, such as $^{12}$CH$_2^-$, $^{12}$CD, or $^{13}$CH$^-$ in $^{14}$C analysis). Most AMS systems employ an electrostatic tandem accelerator that has a direct improvement in background rejection, resulting in a 10^8 time increase in the sensitivity of isotope ratio measurements. As the natural abundance of $^{14}$C in modern carbon is 10^{-12} (isotopic ratio of $^{14}$C:$^{12}$C), a sensitivity of 10^{-15} is a prerequisite for $^{14}$C analysis.
Figure 1. A schematic of the AMS system at Lawrence Livermore National Laboratory Center for Accelerator Mass Spectrometry.

Figure 1, above, starts with a negative ion sputter source, which commonly consists of a stream of Cs\(^+\) with energies of 2-3 keV focused on the surface of a solid sample in order to transfer enough energy to the target material to produce free atoms and ions of the sample material. This process, called sputtering, separates neutral, as well as positive and negative ions from the sample surface. The sample is held at a negative potential, and negatively charged ions are accelerated away from the sample, resulting in a beam of negative ions (Figure 2, below). Cs\(^+\) is particularly useful in \(^{14}\)C studies because it does not form a negative ion from \(^{14}\)N, thereby eliminating isobar interference.\(^4\) It is important to have a beam of negative ions entering the accelerator because the negative ions are attracted to the high-voltage terminal which results in their net acceleration.

Figure 2. Cs sputter ion source.

The low energy (~5-10 keV) diverging beam that leaves the ion source is accelerated, focused and transported to the accelerator by the injector system.\(^2\) CAMS LLNL employs a low-energy mass spectrometer that selects for the desired atomic mass\(^5\) that separates ions by their mass to charge ratio (\(^{12}\)C, \(^{13}\)C, and \(^{14}\)C ions pass through separately). Most AMS systems use sequential injection, a process that switches between stable and rare isotopes via the application of
varying voltages to the electrically insulated vacuum chamber of the analyzer magnet. In sequential injection, typical injection repetition rates are 10 sec$^{-1}$ to minimize variations in the electrical load.[2] This process allows the development of more versatile systems, allowing for analysis of a wide range of isotopes.[1] The alternative to sequential injection is simultaneous injection, a process adopted in accelerators dedicated to $^{14}$C analysis. A recombinator is used following sequential injection, which is a sequence of magnetic analyzers and quadrupole lenses that focus the stable and rare isotopes so they recombine and enter the accelerator together.

The traditional accelerator was first developed in the early 1930s for nuclear physics research. In 1939, UC Berkeley scientists Luis Alvarez and Robert Cornog were the first to use AMS in the detection of $^3$He in nature using the 88-inch Berkeley cyclotron.[5] Now, over 70 years later, cyclotrons have been replaced by an accelerator type with greater energy stability: the tandem electrostatic accelerator. An electrostatic accelerator works by acceleratting particles though a magnetic field generated by high voltages using a mechanic transport system that continuously transports charges from ground to the insulated high-voltage terminal. All tandem accelerators with a maximum terminal voltage above 5 MV use such a mechanical system.[2] The negative ions that enter the accelerator are attracted to the high-voltage terminal, which is what accelerates the CAMS LLNL employs a tandem Van de Graaff accelerator, in which a second acceleration of millions of volts is applied. In all tandem accelerators, atoms are stripped at the high-voltage terminal using either a thin Carbon foil or Argon gas. Stripping is the process in which two or more electrons are removed. The Van de Graaff accelerator removes at least four electrons. It is preferrable to remove at least three electrons because by this process that molecular isobars of $^{14}$C (such as $^{12}$CH$_2^-$, $^{12}$CD, or $^{13}$CH$^+$) are destroyed due to the high instability of their positively charged forms, and atomic C$^+$ ions such as $^{12}$C$^+$, $^{13}$C$^+$, and $^{14}$C$^+$ are separated due to their different mass to charge ratios.[4] The negative ions are changed to positively charged ions and are thus accelerated back to the ground potential in the high-energy part of the accelerator. Transmission through a foil changes with time due to radiation damage and foil thickening, thus gas strippers are used in all modern analyzers due to their increased transmission stability.[2]

Magnetic lenses focus the high energy particles leaving the accelerator into a magnetic dipole, (the high energy analyzing magnet). Stable isotopes can be collected at off-axis beam stops where secondary focusing lenses and additional analyzing equipment remove unwanted ions and molecular fragments to eliminate background. At CAMS LLNL, a magnetic quadrupole lens focuses the desired isotope and charge state to a high-energy mass spectrometer which passes $^{12}$C$^+$ and $^{13}$C$^+$ into Faraday cups and further focuses and stabilizes $^{14}$C in a quadrupole/electrostatic cylindrical analyzer that leads to a gas ionization detector.[5] The magnetic quadrupole and electrostatic selectors coupled together ensure high selectivity and sensitivity, respectively. Other detectors commonly found in AMS systems include surface barrier, time-of-flight, gas filled magnets, and x-ray detectors.

**Interpretation**

Rare isotopes analyzed by AMS are always measured as a ratio of a stable, more abundant (but not too abundant) isotope. For example, the ratio in $^{14}$C studies is generally shown as $^{14}$C/$^{13}$C. Less abundant isotopes are preferable in AMS because the decreased flux of ions reduces background and wear on the instrument, which is of particular concern due to the quick deterioration of particle detectors (performance deteriorates at rates higher than a few thousand
particles per second\(^{(1)}\).

## Applications

Common radioisotope elements measured with AMS and their applications are shown in Table 1\(^{(4)}\), below. Because \(^{14}\)C analysis is by far the most popular application of AMS, the methods discussed below are all techniques used involving \(^{14}\)C.

<table>
<thead>
<tr>
<th>Element (Common Isotope)</th>
<th>Radioisotope with AMS</th>
<th>Natural abundance</th>
<th>Half-life (yr)</th>
<th>Study application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen ((^{1})H)</td>
<td>3H</td>
<td>trace</td>
<td>12.33</td>
<td>Biological/biomedical, Nutritional trace</td>
</tr>
<tr>
<td>Beryllium ((^{9})Be)</td>
<td>10Be</td>
<td>trace</td>
<td>1,510,000</td>
<td>Geochronology, Hydrogeological study, Exposure dating</td>
</tr>
<tr>
<td>Carbon ((^{12})C)</td>
<td>14C</td>
<td>1 x10-10%</td>
<td>5730</td>
<td>Biological/biomedical, Nutritional trace</td>
</tr>
<tr>
<td>Aluminum ((^{27})Al)</td>
<td>26Al</td>
<td>trace, synthetic</td>
<td>720,000</td>
<td>Biological/biomedical, Exposure dating</td>
</tr>
<tr>
<td>Chlorine ((^{35})Cl)</td>
<td>36Cl</td>
<td>7x10−11%</td>
<td>301,000</td>
<td>Earth Science, Hydrogeological study, Exposure dating, Migration of nuclear waste</td>
</tr>
<tr>
<td>Calcium ((^{40})Ca)</td>
<td>41Ca</td>
<td>trace, synthetic</td>
<td>116,000</td>
<td>Biological/biomedical, Nuclear weapon testing</td>
</tr>
<tr>
<td>Nickel (58Ni)</td>
<td>59Ni</td>
<td>trace, synthetic</td>
<td>112,000</td>
<td>Nutritional trace</td>
</tr>
<tr>
<td>Iodine (127I)</td>
<td>129I</td>
<td>trace, synthetic</td>
<td>15,700,000</td>
<td>Biological/biomedical, Migration of nuclear waste, Environmental study</td>
</tr>
</tbody>
</table>

Radiocarbon dating is an analytical method based on the rate of decay of \(^{14}\)C, a radioactive carbon isotope formed in the atmosphere by the reaction between neutrons from cosmic rays and \(^{14}\)N (neutron + \(^{14}\)N = \(^{14}\)C + proton).\(^{(2)}\) Resultant \(^{14}\)C atoms are taken up by plants in the form of \(^{14}\)CO\(_2\), then transferred to animals though the food chain. When animals and plants die, they cease to uptake \(^{14}\)C, and a steady decay of \(^{14}\)C continues in their tissues over time. \(^{14}\)C atoms decay via electron emission (β radiation) to form \(^{14}\)N, a process which has a half life of 5,730 years.\(^{(5)}\)
Radiocarbon levels in the atmosphere change according to complex patterns which are affected by a variety of fluctuations ranging from the sun’s solar activity and the earth’s magnetic field, to ocean ventilation rate and climate. $^{14}$C analysis of tree rings, corals, lake sediments, ice cores, and other sources has led to a detailed record of $^{14}$C variations through time, allowing researchers to establish an official radiocarbon calibration curve (also referred to as a radiocarbon clock) dating back 26,000 calendar years. In the 1960s, nuclear weapons testing released large amounts of neutrons into the atmosphere, nearly doubling $^{14}$C activity.[2] Samples taken after this time period can be radiocarbon dated using a $^{14}$C bomb curve like the peak shown below in Figure 3, can retrieve very precise dates (within 1 year at the steepest part of the curve).

Figure 3. The New Zealand curve (red) is representative of atmospheric $^{14}$C in the Southern Hemisphere, and the Austrian curve is representative of the Northern Hemisphere.

$^{14}$C analysis provides valuable information in the radiocarbon dating of the world’s most priceless artifacts. One such example of the monumental impact of $^{14}$C AMS is the radiocarbon dating of the Dead Sea Scrolls to dates from 300 BC to AD 61 by labs in Zurich and Arizona. AMS has also contributed greatly to environmental and atmospheric studies by providing information regarding particle composition and origin. In the biochemical field, synthesized $^{14}$C labeled compounds can be administered as a tracer dose for in-vivo human metabolic and drug studies which require AMS analysis of graphitized biological samples.

AMS is a highly sensitive method for isotopic analysis that has numerous key applications that are only growing with advances in technology. High costs and technical complexities that arise with the use of a particle accelerator are the only limits to the widespread use of AMS. Recent times have seen the emergence of commercially available compact accelerators that use as low as 200 kV for radiocarbon dating and biomedical applications, and as particle accelerators become more commonplace, modifications to the instrument have also broadened the number of isotopes the instrument can measure.

References

Contributors and Attributions

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