

Applications of a compact ionization chamber in AMS at energies below 1 MeV/amu

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Available online 6 March 2008

Abstract

The increasing demand for measuring long-lived radionuclides with small AMS machines at energies below 1 MeV per nucleon raises the need for compact detectors which still have a decent energy resolution and allow for a clear identification of the incident particles. Based on a design by the AMS group at the ETH Zurich a compact gas ionization chamber was built and installed at the 3 MV tandem AMS facility VERA (Vienna Environmental Research Accelerator). The main challenge in AMS is the detection of rare isotope species in the presence of strong isotopic and isobaric interferences. The task of the ionization chamber is the suppression of the unwanted isobar by separating the ions via their different stopping powers. Measurements of ^{36}Cl at VERA showed an achieved suppression of the unwanted stable isobar ^{36}S of 3×10^{-4} and measurements of ^{10}Be showed an achieved suppression of ^{10}B of at least 3×10^{-6} . Additional suppression of the isobaric ions can be achieved by a degrader foil technique applied to ^{10}Be measurements by G.M. Raisbeck. In combination with the new ionization chamber the achieved suppression of ^{10}B is at least 10^{-10} . Measurements of blank samples at VERA show that the background for AMS with ^{10}Be is below 2×10^{-15} .

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PACS: 07.75.+h; 29.40.Cs

Keywords: Accelerator mass spectrometry; Ionization chamber

1. Introduction

Accelerator Mass Spectrometry (AMS) is a powerful technique for the detection of long-lived radioisotopes at typical abundances in the range of 10^{-12} – 10^{-15} . The precise determination of such low abundances has found various applications as tracers or chronometers in geology, archaeology and biomedicine. Tuniz et al. give a detailed overview of the technique as well as a discussion of its numerous applications [1].

One of the main tasks in AMS measurements is the separation of the rare radioisotope from stable isobaric background. There are various ways to achieve this separation. In some cases this task can be efficiently fulfilled inside the ion source using negative ions. Examples for these cases are

^{14}C , ^{26}Al and ^{129}I where the stable isobars ^{14}N , ^{26}Mg and ^{129}Xe do not form negative ions. In other cases negative molecular ions can be used. One example is the measurement of ^{41}Ca where the interference of ^{41}K is strongly suppressed by using $^{41}\text{CaH}_3^-$ ions through the instability of $^{41}\text{KH}_3^-$. Another possibility is the separation after acceleration using different methods like energy loss in matter, using the mean charge state in a gas-filled magnet or full stripping in a subsequent stripper stage.

The scope of this work is to show the application of a compact ionization chamber to achieve the separation of different isobars in AMS.

2. Setup of the detector

At the laboratory for ion beam physics at the ETH Zurich a compact ionization chamber was developed for use

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with their sub-MeV tandem AMS machines [2]. The aim of such detectors is to separate the stable isobar from the radioisotope via their different energy loss in a gas. Based on this design a similar detector was built at the VERA laboratory for use at the VERA 3 MV tandem AMS facility.

Owing to its compact design the detector fits into a standard DN100CF cross piece. It can be inserted and retracted without breaking the vacuum. The active detector volume has a length of about 6 cm. A scheme of the inner part is shown in Fig. 1. The anode is split into two consecutive regions with equal lengths, which allows a simultaneous measurement of the energy loss in two different regions. The preamplifier electronics is built with CR-110 charge sensitive preamplifier modules from Cremat Inc. (Watertown, MA, USA). These compact hybrid modules have a small footprint, which allows mounting the preamplifiers for the two signals inside the active detector volume close to the anodes. This reduces the electronic noise due to short cable lengths. A major breakthrough for the use of ionization chambers at energies below 1 MeV/amu was the development of thin silicon nitride foils [3]. These foils are pinhole free and commercially available from Silson Ltd. (Northampton, UK) with thicknesses down to 30 nm. They are remarkably homogeneous, which produces almost no low energy tails. In our setup windows of square shape with a side length of 5 mm are used. They come with a native silicon frame of 10 mm × 10 mm that can be glued onto the entrance of the ionization chamber. The 50 nm foil used in our setup withstands easily a gas pressure above 100 mbar inside the detector. The Frisch grid [4] consists of gold coated tungsten wires (20 μm thickness) glued or soldered onto a stainless steel frame. The applied voltage between anode and cathode is 200 V. The Frisch grid was set to 60% of the full voltage using a resistive voltage divider. The distances between the cathode, the Frisch grid and the anodes are 25 mm and 5 mm, respectively. As counting gas isobutane is used. A separate gas regulation system is used to maintain a constant pressure inside the detector. The pressure was chosen such that the best possible separation for the measured isobars was achieved.

The output of the preamplifiers is sent to a dual channel spectroscopic amplifier (Ortec model 855) for amplification and pulse shaping. The resulting signals are sent, via two ADCs gated by the signal from the first anode, to the VERA data acquisition.

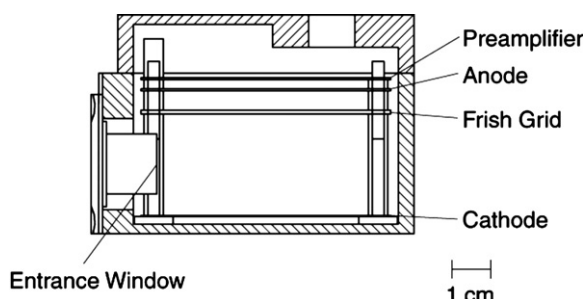


Fig. 1. Schematic of the active detector volume.

3. Results

For the first performance tests of the detector a ^{13}C particle beam with an energy of 12 MeV was produced. The detector was filled with 95 mbar of isobutane to fully stop the particles inside the active volume. The energy resolution (FWHM) of the measured signal was found to be 100 keV (0.8%). This remarkably high resolution outperforms a silicon detector by far. Also measurements with heavier ions were performed. For a 12 MeV Cu beam an energy resolution of 220 keV (1.8%) and for a 16.4 MeV Au beam a resolution of 400 keV (2.4%) were measured.

3.1. ^{41}Ca

In the case of ^{41}Ca the stable isobar is ^{41}K . As mentioned earlier this can be suppressed by using CaH_3^- ions. But as the hydride is highly hygroscopic special care has to be taken during preparation and handling of the sample material. Another approach is using CaF_3^- ions, which are easy to handle but show a much smaller suppression of KF_3^- . The CaF_3^- ions are extracted from CaF_2 sample material. After the acceleration stage ions of charge state 4^+ with an energy of 16 MeV were selected. The spectra of a typical sample material and a blank are shown in Fig. 2. The measurement time was 422 s. The measured isotope ratio of $^{41}\text{Ca}/\text{Ca}$ of the unknown sample was 5.1×10^{-12} . The blank sample was measured to have 2.4×10^{-13} .

3.2. ^{36}Cl

In ^{36}Cl measurements the stable isobar is ^{36}S . A suppression to a certain degree of sulfur can be achieved by chemical treatment of the sample material and putting it into a special sample holder containing virtually no sulfur, but a relatively high background of ^{36}S remains. To achieve a decent separation a high particle energy of 28 MeV using the 7^+ charge state of the ions was chosen. Instead of fully stopping the particles inside the active region of the detector a low pressure of 35 mbar of isobutane was used. In this way the counts of ^{36}Cl ions show up in a region of the spectra which is empty of events of scattered ^{36}S particles. The spectra of a typical sample material and a blank are shown in Fig. 3. The bin for the ^{36}Cl events was chosen in such a way that only the right half of the peak area was used. This reduces the detection efficiency by half but in this way a suppression of the unwanted isobar ^{36}S of 3×10^{-4} could be achieved. The measured isotope ratio of $^{36}\text{Cl}/\text{Cl}$ was 4.6×10^{-12} . The blank was measured to have 5.5×10^{-15} .

3.3. ^{10}Be

AMS measurements of ^{10}Be are influenced by a strong interference of the stable isobar ^{10}B . Previous measurements at VERA were performed using BeO as sample material and extracting BeO^- beams from the sputter ion source. The interfering isobar is stopped in front of a silicon

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