Neutron capture cross section measurements of ²⁴¹Am at the n_TOF facility

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Abstract. Neutron capture on ²⁴¹Am plays an important role in the nuclear energy production and also provides valuable information for the improvement of nuclear models and the statistical interpretation of the nuclear properties. A new experiment to measure the ²⁴¹Am(n, γ) cross section in the thermal region and the first few resonances below 10 eV has been carried out at EAR2 of the n_TOF facility at CERN. Three neutron-insensitive C₆D₆ detectors have been used to measure the neutron-capture gamma cascade as a function of the neutron time of flight, and then deduce the neutron capture yield. Preliminary results will be presented and compared with previously obtained results at the same facility in EAR1. In EAR1 the gamma-ray background at thermal energies was about 90% of the signal while in EAR2 is up to a 25 factor much more favorable signal to noise ratio. We also extended the low energy limit down to subthermal energies. This measurement will allow a comparison with neutron capture measurements conducted at reactors and using a different experimental technique.

1 Introduction and motivation

The design and modelling of advanced nuclear systems and nuclear fuel cycles need detailed knowledge of the nuclear data of the most relevant isotopes, including nuclear reaction cross sections and decay data. In particular, accurate knowledge of the neutron capture cross section of 241 Am(n, γ) is required since it plays an important role in the nuclear energy production and fuel cycle scenarios. For this reason, the isotope 241 Am is listed in the NEA High Priority Request List [1, 2]. Cross section data can also provide valuable nuclear structure data necessary for the improvement of nuclear models and the statistical interpretation of the nuclear properties.

The neutron capture cross section of ²⁴¹Am is an important quantity for nuclear technology applications. The nuclei ²⁴¹Am and ²³⁷Np are the minor actinides which contribute most to the long-term hazard of nuclear waste, as ²⁴¹Am (half life $T_{1/2} = 432$ y) decays via α decay into ²³⁷Np ($T_{1/2} = 2$ My). After Np, Am is the second most abundant minor actinide present in spent nuclear fuel, of which ²⁴¹Am is the most abundant isotope. The build-up of heavier actinides like Cm, goes through (n, γ) reactions on Am isotopes. The ²⁴¹Am(n, γ) reaction leads to ²⁴²Cm production via the decay of the nucleus ²⁴²Am ($T_{1/2} = 16$ h). ²⁴²Cm decays via α emission into ²³⁸Pu, which is fissile in a fast spectrum.

The discrepancies observed between experimental and evaluated data sets in the ²⁴¹Am neutron capture cross section has triggered an interest for new measurements. Several time-of-flight measurements of ²⁴¹Am(n, γ) reaction have been performed in the last years: Jandel *et al.* [3] in Los Alamos National Laboratory, using the 4π calorimeter of BaF₂ crystals DANCE. Lampoudis *et al.* [4] at the

In order to improve the accuracy on the neutron induced cross sections of ²⁴¹Am in the low energy region, a new neutron capture measurement has been performed at the neutron time-of-flight facility n_TOF at CERN. This time the measurement was done at the new flight path EAR2 [7], located 20 m vertically from the neutron spallation source. Due to the much higher neutron flux, EAR2 is more suited for highly radioactive isotopes. The focus of this measurement is on the resolved energy region and in particular the thermal region and the first resonances below 10 eV. A comparison with previous reported results is further presented.

2 Experimental setup

2.1 The n_TOF facility

The n_TOF facility is located at CERN (European Organization for Nuclear Research) and has been described else-

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GELINA facility in Geel, Belgium used two liquid scintillator detectors based on deuterated benzene (C₆D₆). At the n_TOF facility at CERN, two measurements have been performed in the past, both at the 185 m flight path of EAR1. One measurement was done with C_6D_6 detectors by Fraval et al. [5], and the other using the BaF₂ total absorption calorimeter (TAC) by Mendoza et al. [6]. Both experiments were using the same ²⁴¹Am sample. The results of the two EAR1 measurements present sizeable differences at low energy below 10 eV, but were still in agreement within the quoted uncertainties. Both measurements suffer from a large systematic uncertainty at low neutron energy caused by the large background component due to the radioactivity of ²⁴¹Am. Indeed for these measurements in EAR1 the gamma-ray background at thermal energies was about 90% of the signal.

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Figure 1. A schematic view of the ²⁴¹Am sample used in the experiment.

where [8]. The neutron beam is generated through spallation of 20 GeV/c protons extracted in pulses from the Proton Synchrotron (PS) and impinging on a 40 cm thick and 60 cm diameter cylindrical lead target. The proton pulses are delivered with a time spread of 7 ns (rms), having a nominal intensity of 7×10^{12} protons per pulse at a minimum repetition rate of 1.2 s, leading into a very high instantaneous neutron flux. After moderation of the initially fast neutrons, a wide-spectrum neutron beam is produced from thermal to a few GeV. Two experimental areas where experiments are mounted are available at the n_TOF facility: EAR1 [9], having a horizontal flight path of about 185 m from the spallation lead target, and EAR2 [7], with a vertical flight path of about 20 m.

Since the beam line of the vertical beam is about 10 times shorter, the available neutron flux per proton pulse in EAR2 is considerably higher than in EAR1 [10]. Therefore EAR2 is particularly suited for measuring highly radioactive samples. In addition, due to the approximately 10 times shorter flight distance, the time-of-flight interval needed to cover a range of neutron energies, is 10 times smaller. This results in an additional increase of a factor 10 of the signal to noise ration due to radioactivity. On the other hand, the excellent energy resolution at EAR1, due to the large flight distance, cannot be reached at EAR2.

2.2 Data acquisition

The Data Acquisition System (DAQ) has been upgraded to 175 MSample digitizers from Teledyne SP Devices with a sampling frequency of up to 2 GHz and an amplitude resolution of 12 or 14 bit. The acquired waveforms are, after zero-suppression, automatically transferred from the DAQ computers to the CERN Advanced STORage manager (CASTOR) for long-term storage and off-line analysis. The larger on-board memory has increased the exploitable time-of-flight range with respect to the previous DAQ system, and is now expanded down to thermal neutron energies for both EAR1 and EAR2. The DAQ is triggered by each proton pulse and records detector signals during a time window of 100 ms.



Figure 2. The γ -ray spectrum of the ²⁴¹Am sample measured with HPGe setup BANEX of the actinide facility in CEA Cadarache. Several characteristic γ rays from α -induced reactions are shown as well. The ²⁴¹Am sample activity in linear scale is shown in the inset illustrating the very strong 59.5 keV line dominating the activity spectrum.

and immobilized in an aluminum oxide (Al_2O_3) pressed pellet forming a rigid disk of 12.26 mm diameter. The disk was then encapsulated in an aluminum container and sealed with Stycast as illustrated in figure 1.

The first sample, labelled IRMM1, has a mass of 32.23 \pm 0.19-mg and was the same sample as used in the previous n_TOF capture experiments [5, 6]. A second sample, labelled IRMM4, has a 40.98 \pm 0.25 mg content of ²⁴¹Am. The masses were determined by calorimetry [11]. The radioactivity of the IRMM1 sample was about 4.1 GBq while for the IRMM4 was about 5.1 GBq. More details on the ²⁴¹Am sample composition are presented in table 1.

Several nearly identical dummy (or blank) samples, without americium oxide inside the Al_2O_3 disk, and also encapsulated inside an aluminium canning, were also produced. The one that was used in the previous measurements at n_TOF appeared to contain a broken Al_2O_3 pellet which seemed to be assembled with a samarium-containing glue during the production process. This became visible in the neutron capture spectra from that measurement [5] and was later confirmed by an X-ray analysis of the dummy samples. Therefore, for the present measurement we have used one of the other dummy samples, without a noticeable samarium content, for the background measurements.

The α -activity of the ²⁴¹Am sample generated several γ -rays which could be identified as shown in figure 2. In an independent measurement with HPGe setup BANEX of the actinide facility in CEA Cadarache, the full gamma-ray

Table 1. The material quantities of the two used ²⁴¹Am samples.

sample ID	²⁴¹ Am	Al_2O_3
	(mg)	(mg)
IRMM1	32.23±0.19	305
IRMM4	40.98 ± 0.25	388

2.3 The $^{\rm 241}{\rm Am}$ sample

Two 241 Am samples were used for this experiment. Each sample consists of americium oxide (241 AmO₂), infiltrated



Figure 3. The three C_6D_6 scintillation detectors used to measure the capture γ -rays in n_TOF-EAR2. The additional fourth monitoring detector is also visible on the picture.

spectrum from 40 keV up to 5 MeV was measured. The the γ -rays by α -induced reactions are clearly visible. On the linear scale of the inset of figure 2 it can be seen that the main component of the activity of the ²⁴¹Am sample is concentrated at the 59.5 keV γ -ray, coming from ²³⁷Np following the α -decay of ²⁴¹Am.

2.4 The C₆D₆ detectors

Three C_6D_6 liquid scintillator detectors were used for the γ -ray detection. The main characteristics of these detectors are a very good time resolution and a low sensitivity to scattered neutrons. The very high activity of the ²⁴¹Am samples draws a large current over the voltage divider of the photomultiplier. To avoid an undesirable detector response due to a strong flux of 59.5 keV γ -rays, a Pb shielding between sample and detector was used in the previous n_TOF measurements. For the present measurement we have chosen to increase the distance between sample and detector, avoiding the need for a Pb shielding and allowing the possibility to use a lower gamma-ray threshold.

The three detectors and the sample were mounted as shown on the picture in figure 3, together with a schematic view as in figure 4. The sample (shown in red) was placed on a horizontal sample holder, which was fixed with three vertical thin rods of 15 cm long (two are shown on the picture) to the end flange of the evacuated beam pipe. In this way the material directly around the sample was minimized. The detectors were hold in place using carbon fibre rods (not shown). The distance between the front face of the scintillator cells and the center of the sample was approximately 40 cm. This distance was chosen to optimize the detector performance in presence of the large radioactivity and the deliberate absence of shielding between the detector and sample. A fourth C_6D_6 detector was placed even further away from the sample and was only used for monitoring purposes.

Each C₆D₆ detector was energy calibrated using standard γ -ray sources (¹³⁷Cs, ⁸⁸Y, ⁶⁰Co and a mixed Am-Be source).



Figure 4. A schematic view of the detectors and sample setup used for the present experiment. While the distances in the figure are on scale, for clarity the detectors are not presented in perspective. The distance between detectors and sample is about 50 cm. The inset shows a top view of the set-up with the three-detector configuration.

Supplementary measurements with other samples (dummy sample: Al canning plus the Al₂O₃ matrix, only Al canning, natural C and Pb) were carried out to determine the background. Measurements without beam and without sample (no sample, beam off), with beam but with the sample in place (²⁴¹Am, beam off), and with neutron beam but without any sample were also performed. For the absolute normalisation the low-energy saturated resonances of five different samples (¹⁹⁷Au, ^{nat}Ag, ¹⁰³Rh, ^{nat}Ir and ^{nat}U) were measured.

3 Preliminary results

The main objective of the present work is a precise measurement of the 241 Am(n, γ) reaction cross section in the thermal energy region and for the first few resonances below 10 eV. In the region below 10 eV the uncertainties of the two previous n_TOF experiments were very large. This region is particular important for nuclear fission technology. Experimental data either cover only thermal energies, or are time-of-flight data, but do not include the full thermal region. With the present measurement we com-



Figure 5. Comparison of the ²⁴¹Am spectra together with the ²⁴¹Am coutn rate from radioactivity at EAR1 (2010) and EAR2 (2017). The fitted spectrum due to the constant radioactivity, is represented by the blue $1/\sqrt{E}$ line.

prise both the full thermal region and the resonance region. In addition, the background due to high radioactivity of ²⁴¹Am is drastically reduced in n_TOF-EAR2.

In figure 5 we show the counting spectrum of the 241 Am(n, γ) for the present EAR2 experiment. For comparison, also the previous EAR1 C₆D₆ results [5] are shown on the same neutron energy scale. In order to compare the present results to the previous n_TOF experiments, in addition to using the same sample (IRMM1), also the same energy threshold of 300 keV was applied to each C₆D₆ detector. Still an important difference is the use of a 2 mm Pb shielding for the EAR1 setup, and no shielding in the EAR2 setup. We clearly see the complete thermal peak from the neutron flux present in the EAR2 counting spectrum. In the EAR1 experiment, while the data acquisition system could not record down to this low energy, the thermal neutrons are strongly suppressed by the borated water moderator. The ²⁴¹Am spectra from EAR1 presents a visible better energy resolution as compared to EAR2.

In the figure we also see the fit of the radioactivity background. The number of counts per time unit is constant for this spectrum, and shows as a $1/\sqrt{E}$ in the units used here. Also, compared to the previous measurements performed in EAR1, there is a substantial increase in the

ratio between the capture signal and the radioactivity background of more than a factor 200 in EAR2 due to the increased neutron flux. This is one of the most important features of the n_TOF facility and essential for the measurement of neutron-induced cross section of highly radioactive targets. Nevertheless, even if the resolution is much less favourable in EAR2, the peak-to-valley ratio of the first strong resonance and the region between the first and second strong resonances, is significantly better for the EAR1 experiment, indicating that the time-dependent background is much better in EAR1. The strong thermal component of the neutron flux in EAR2, which is efficiently suppressed in EAR1, results in a high flux of inbeam 2.2 MeV gamma rays from neutron capture on hydrogen of the water moderator near the spallation target. Those gamma-rays contribute significantly to the timedependent background in EAR2.

4 Conclusion

The here presented counting spectrum shows the capability of n_TOF EAR2 to peform neutron capture measurements in the low energy range, covering both the full thermal region and the first resonances. The excellent ratio of signals due to neutron capture and due to the radioactivity background is particularly beneficial for a nucleus like ²⁴¹Am. At present all data are verified for integrity and consistency and a full analysis of the data is ongoing.

References

- [1] https://www.oecd-nea.org/dbdata/hprl/
- [2] E. Dupont et al. (2019), these proceedings
- [3] M. Jandel et al., Phys. Rev. C 78 (2008)
- [4] C. Lampoudis et al., Eur. Phys. J. Plus 128 (2013)
- [5] K. Fraval et al., Phys. Rev. C 89 (2014)
- [6] E. Mendoza et al., Phys. Rev. C 97 (2018)
- [7] C. Weiß et al., Nucl. Instr. Meth. A 799, 90 (2015)
- [8] E. Chiaveri et al. (2019), these proceedings
- [9] C. Guerrero et al., Eur. Phys. J. A 49 (2013)
- [10] M. Sabaté-Gilarte et al., Eur. Phys. J. A 53 (2017)
- [11] H. Tagziria et al., Nucl. Instr. Meth. A 691, 90 (2012)