Evaluation of a setup for pNRA at LIBAF for applications in geosciences

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A new setup for photon tagged nuclear reaction analysis pNRA is being developed at Lund’s ion beam analysis facility LIBAF. Particle induced gamma ray emission PIGE and nuclear reaction analysis NRA are two methods that have been extensively used for light isotope measurement in ion beam analysis IBA. There is an abundance of nuclear reactions between light elements and MeV protons, deuterons and alpha particles. This means that in principle all elements from lithium all the way up to chlorine can be analyzed using those techniques. Detection limits can be improved for some elements, if those two methods are fused together into pNRA.

The new setup for pNRA will benefit from advances in detector technology that occurred during the last 20 years. A LaBr$_3$ scintillator detector and an annular double sided silicon strip detector DSSSD are used in coincidence to detect a gamma and a charged particle respectively. Both detectors are connected to a VME based data acquisition system. Of primary interest in this work is the analysis of isotopic ratios of light elements in geological samples, which are usually thick with a complex matrix. This setup can be for instance used to measure isotopic fractionation of oxygen and boron. We will present the setup and discuss its capabilities.

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1. Introduction

There is a multitude of methods available within ion beam analysis IBA [1]. A new setup for photon tagged nuclear reaction analysis pNRA [2] is currently under development at Lund’s ion beam analysis facility LIBAF. pNRA combines Particle Induced Gamma ray Emission PIGE and Nuclear Reaction Analysis NRA into one technique. Those two aforementioned methods have been extensively used for light isotope spectroscopy in IBA [3]. The principle of pNRA is to analyze isotopic content of a sample by detecting both the gamma quanta and the charged particle produced in a nuclear reaction [4]. This technique was first proposed 20 years ago, it was suggested that it could be used for trace element analysis in thin samples [2] and depth profiling of thick samples [5].

The availability of many nuclear reactions for interaction of light ions with matter means that with the right choice of projectile and projectile energy all elements from lithium all the way up to chlorine can be observed and quantified [6]. Previous pNRA setup at LIBAF has been used for boron measurements in biological samples [7]. It was also suggested that this method could be successfully combined with a nuclear microprobe setup [6]. Important for this study is that pNRA can be used to measure isotopic fractionation of light elements.

The primary goal for the new setup is the analysis of geological material, which entails thick samples with a complex matrix. Isotopic fractionation of light elements is relevant in many contexts within geosciences. The application that inspired the current project is stable oxygen isotope quantification in extraterrestrial geological material. Oxygen isotopic fractionation can be used to both recognize and classify the extraterrestrial minerals [8]. In complex thick and semi-thick samples background and interferences can be a problem for PIGE and NRA. Here pNRA can prove to be superior [6].

2. Method

The minimum equipment required for pNRA is a source of MeV ions, a particle detector, a gamma detector and electronics for creating a coincidence [2]. The tagging of the charged particle with the corresponding gamma is a way of suppressing the background and separating the interesting reactions from interferences expected in complex targets. The interesting nuclear reactions in this case are inelastic scatterings and resonances, which leave the nucleus in an excited state, so that $\gamma$-emission is possible.

The setup discussed in the current work benefits greatly from advances in detector technology that occurred during the last few decades. Description of the old setup can be found in [6]. This
new setup was installed and tested at LIBAF, which is a facility based around a single ended 3 MV accelerator of Van der Graaff type with a nuclear microprobe and p, d and α beams [9].

The ability to capture as large as possible fraction of reaction products is always desirable in Nuclear Physics. This is especially true here since the limiting factor for the detection of each element is the cross sections of the nuclear reactions involved; many of those are on the order of some mb [1]. For the charged particle this means primarily large solid angle coverage is desired. Due to slow integration times for standard shaping electronics and large cross section for elastic scattering, the ability to suppress or prevent pile-up is also highly desirable. For the gamma particles the relevant parameter to maximize the yield is the gamma capture efficiency, which is related to both the detector size and the scintillator material. In simple terms an efficient pNRA experiment requires large detectors and a compact geometry. Other important parameters for this kind of setup are a good timing and a good energy resolution. Improving those greatly improves background suppression which means a gain in signal to noise ratio.

3. Experiment

Fig. 1 provides a schematic description of the experimental setup. The detectors used are a large 1.5 × 2 inches cylindrical LaBr$_3$ scintillator detector [10] and a double sided silicon strip detector DSSSD, previously described in [11]. They are used in coincidence to detect a gamma and a charged particle respectively. The analog electronics on the DSSSD side consists of Mesytec preamplifier MPR-16 and shaper STM-16 [11,12]. Shaping amplifiers provide both the energy and timing signals. Signals from the PMT can be used without amplification and split to provide both timing and energy. The rest of the analog electronics is a standard coincidence setup constructed out of NIM modules. The only complication is aligning the timing of gamma and particle signals to create a common gate for both of them in the data acquisition system DAQ. Passive delay cables are used for that purpose. The faster, gamma branch starts the coincidence. A 1 μs long coincidence gate is used, which is more than enough to observer all, prompt and nearly prompt reactions. The shorter gate allows us to increase current and get stronger signal for the same background. In principle a TOF of less than 5 ns is expected between the gamma start and particle stop, for prompt particles. In reality the leading edge discriminators built into the shaping electronics on the DSSSD introduce an amplitude walk into the spectrum [13]. The setup is very flexible since there is always the option of switching the coincidence off which reduces the method back to PIGE, RBS and NRA.

LaBr$_3$ is a new scintillator material intended for applications where both the timing and energy resolution are important. This inorganic crystal has density of 5.3 g/cm$^3$, which is more than NaI, CsI and BaF$_2$. That means it has a higher absorption efficiency then all three of them [14]. This material has a very good energy resolution (3% @ 660 keV) and timing properties (250 ps pulse rise time) [14]. The Hamamatsu PMT attached to this particular detector is fast, small and vacuum compatible, which means that the detector can be placed in the reaction chamber, less than 1 cm behind the target. The count rate in the gamma detector is nowhere near enough to saturate the detector, so as compact as possible geometry is desired. In this geometry the solid angle of the detector is 3.6 sr.

The DSSSD is placed 3 cm in front of the target. This DSSSD is annular with 64 radial electrodes on the front side and 32 ring shaped electrodes on the back side of the detectors as described in [11]. A real particle is defined as an event that generates equal amount of charge in two electrodes, one on the front and one on the backside of the detector, this means there are 2048 possible pixels. This large segmentation removes the pileup and adds angular information, which can be further used to separate interesting reactions. This version of DSSSD has an inner active diameter of 14 mm, active outer diameter of 85 mm and a dead layer of 2 μm. For the current geometry the size of the detector is 2.5 sr, each individual pixel is on the order of 1 msr.

Together those two detectors have large solid angle coverage for both the particle and the gamma quanta. The true coincidence rate for this system will be defined by the smallest detector or rather smallest solid angle times interaction efficiency which is almost 100% for a particle detector and which is an energy dependent quantity for the gamma detector.

The main drawbacks of the system are the internal activity of the LaBr$_3$ crystal, which leads to a characteristic spectrum seen in Fig. 2 (dotted line). This activity is due to decay of $^{138}$La, a naturally occurring La isotope, and due to contamination of the crystal with $^{227}$Ac, which is part of actinium series decay chain [10]. The

Fig. 1. Experimental setup consists of a large area segmented particle detector, DSSSD and a fast gamma detector, a LaBr$_3$ scintillator. DSSSD is placed in the backward geometry, LaBr$_3$ in the forward geometry, both detectors capture large portion of the solid angle. Coincidence setup is built out of standard NIM electronics. Scintillator energy output is split and used to generate start for the coincidence gate of 1 μs length. Triggers from the silicon shapers are used as the stop signals. It is necessary to introduce delay to get both energy signals into the gate.
4. Results

To evaluate the systems performance data has been taken for some representative samples. The event mode data acquisition provides us with three correlated parameters for each event that are used to find and isolate interesting reactions. The parameters are the energies of the particle and of the gamma quanta, and their relative arrival time at their corresponding detectors. This information can be combined into a number of figures commonly used in coincidence experiments. The timing information gives the coincidence peak, which is used as the main filtering condition in a coincidence experiment. The plot of energies of the entities involved in the coincidence gives a two-dimensional energy–energy plot. This plot provides an immediate identification and separation of the interesting reactions.

The focused proton beam at 2.5 MeV was scanned over the samples. Materials used as test samples were thin PIXE standards originally intended for calibration of a PIXE or an XRF setup [19]. They each consist of 2 μm Mylar backing with known amounts of the element of interest deposited onto them. Since the intended target for the system are thick geological samples, those standards were fixed onto a thick brass backing. The sample materials were CaF₂, NaCl and Al. This sample configuration is intended to approximate real physical situation, at least as far as background level is concerned.

In Fig. 3 reactions induced in CaF₂ sample are studied. A reaction marked as D in the figure is ¹⁹F(p,pγ)¹⁹F for Eγ = 197 keV [20] and is an interesting case to observe. This reaction is a nearly prompt decay for which the decay time is slow enough to measure. The previously measured half-life of this transition is 90 ns and this value can be easily recreated from the projection of the figure onto the time axis. A truly prompt ¹⁹F(p,γ)¹⁰O marked as P can be seen as well, this reaction is induced above a number of high lying levels in the ¹⁰O nucleus which means that several strong gamma rays can be observed when a cut is placed on this reaction [21].

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A hardware coincidence condition with 1 μs long gate is used as the trigger for the data acquisition system, but this gate can be easily shortened to a few hundred nanosecond. Unfortunately the actual prompt coincidence peak in the timing spectrum without any further context would have rather poor time resolution. As can be seen in Fig. 3 projection of the prompt peak, marked as P, onto the time axis is more than 50 ns thick and clearly asymmetric. Thus extending coincidence time spectra into a two dimensional histogram, where the energy of the particle is plotted as the function of arrival time in each detector, is an important tool. This way we see that the coincidence time is not a straight line. It changes with the energy of the particle. This is a direct effect of the leading edge discriminators built into the shaping amplifiers. It is well understood that the leading edge timing circuits are affected by the noise of the signal [13]. This problem can be solved in hardware, for instance constant fraction discriminators do not suffer from that problem, but that would require additional investment in electronics. Making a banana shaped cut in the particle energy-time spectrum, which follows the behavior of the leading edge discriminators, is sufficient to separate the real coincidences from accidental background. The same cut was placed on each one of the timing spectra. Actual FWHM of timing peak when leading edge behavior is taken into account is less than 16 ns.

Positions of the three peaks observed in the energy-time spectra during the measurements on Al and Na rich samples have been superimposed onto the current spectrum. Those 3 processes are inelastic scattering reactions of protons and mark a line parallel to the one created by the $^{19}$F(p,p$\gamma$)$^{20}$Ne reaction. This suggests that it should be possible to separate protons from alpha particles with this setup.

An example of a particle-gamma energy spectrum can be seen in Fig. 4. A sample of NaCl was irradiated with 14.4 μC of protons. Two reactions can be clearly seen at the energy of 2.5 MeV. They are identified as $^{23}$Na(p,p$\gamma$)$^{23}$Na with $E_c = 440$ keV and $^{23}$Na(p,γγ)$^{26}$Ne with $E_c = 1636$ keV [22]. Energy resolution of the LaBr$_3$ during this run was around 9% much worse than the expected 3%. This is most likely due to noise in analog electronics. Still both reactions can be clearly seen and separated. In Regular NRA experiment those two interactions would interfere with each other, which complicates the quantification. This is not a problem for pNRA. The stronger one of the two reactions is used to estimate minimum detectable limit for sodium.

Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Charge (μC)</th>
<th>Sensitivity (counts/(μC * μg/cm$^2$))</th>
<th>MDL ((μg/cm$^2$)/μC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>14.4</td>
<td>145.6</td>
<td>0.36</td>
</tr>
<tr>
<td>F</td>
<td>12.7</td>
<td>20.9</td>
<td>4.84</td>
</tr>
<tr>
<td>Al</td>
<td>16.5</td>
<td>1.32</td>
<td>20</td>
</tr>
</tbody>
</table>

Compilation of the relevant results such as minimum detectable limit for the samples described above and for the Al standard can be found in Table 1. Cross section data for relevant reactions are available in the IBANDL database [23]. Minimum detectable limit depends on the background counts under the peak ($N_b$) and systems sensitivity ($S$), MDL = $3 \times N_b / S$, where sensitivity $S$ is yield per μC per amount of substance. It can be seen in the table that both of those numbers vary two orders of magnitude between the samples. That is an expected behavior for the pNRA method. They can also vary rather strongly with beam energy and detector geometry used. This means that these numbers can be improved for individual isotopes, but never for an entire range of elements simultaneously.

5. Conclusions

As was shown in figures and explained in the previous section almost background free spectra can be produced with straightforward cuts on the data. The standard samples with known areal concentrations used in the experiment allow us to estimate minimum detectable limit, for a number of elements, for the current experimental settings. The internal activity of the LaBr$_3$ detector is the primary factor defining the detection limit for most samples of interest, which will be thick with dense matrix, often consisting of heavy atoms. The accidental background seen is small and can
be decreased even further with a choice of a shorter coincidence gate.

pNRA is a method useful in specific situations only. It will not replace existing techniques but it can easily complement them. NRA and PIGE are both hampered by lack of cross section information together with errors in energy loss. This makes analysis of complex spectra difficult and often ambiguous. pNRA spectra by the very nature of this method promise to be cleaner with less background and fewer interferences, desired resonances can be studied with greater ease.

A more precise experimental run is necessary to fully determine the limitations of the system and account for them. The particular pNRA setup under construction at LIBAF could be potentially used to provide additional information. Position sensitivity of the DSSSD could be used as an additional criterion in data evaluation since the cross section of many resonances has a strong angular dependence. Even mass separation should in principle be attainable although this might require an investment in faster electronics. The next step for this setup is to apply it to a set of oxygen standard samples and real geological samples and measure oxygen content and isotopic ratio.

References