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A new approach to monitor ^{13}C -targets degradation in situ for $^{13}\text{C}(\alpha,n)^{16}\text{O}$ cross-section measurements at LUNA

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Abstract. Direct measurements of reaction cross-sections at astrophysical energies often require the use of solid targets able to withstand high ion beam currents for extended periods of time. Thus, monitoring target thickness, isotopic composition, and target stoichiometry during data taking is critical to account for possible target modifications and to reduce uncertainties in the final cross-section results. A common technique used for these purposes is the Nuclear Resonant Reaction Analysis (NRRA), which however requires that a narrow resonance be available inside the dynamic range of the accelerator used. In cases when this is not possible, as for example the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction recently studied at low energies at the Laboratory for Underground Nuclear Astrophysics (LUNA) in Italy, alternative approaches must be found. Here, we present a new application of the shape analysis of primary γ rays emitted by the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ radiative capture reaction. This approach was used to monitor ^{13}C target degradation *in situ* during the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ data taking campaign. The results obtained are in agreement with evaluations subsequently performed at Atomki (Hungary) using the NRRA method. A preliminary application for the extraction of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction cross-section at one beam energy is also reported.

Key words. ^{13}C enriched solid target, NRRA, ion beam, γ -shape analysis, nuclear astrophysics

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1 Introduction

Knowledge of the stoichiometric composition of solid state targets and their behaviour during ion beam irradiation is of great importance in various fields of ion beam physics, from material analysis to nuclear astrophysics [1–4]. The main goal of the latter is to measure nuclear reaction cross-sections at, or near, the energy region of astrophysical interest (the so-called Gamow window), typically of the order of hundreds of keV or less. Since cross-sections drop exponentially with decreasing energy in this energy region, counting rates can be of the order of one event per hour or lower. Therefore, high beam currents (hundreds of μA) and long irradiation times (weeks or months) are often necessary to achieve high enough signal-to-noise ratios for a successful cross-section measurement at low energies. Yet, target modification processes (such as diffusion, melting, sputtering or contamination of target surface [5, 6]) that occur under intense beam irradiation may result in significant changes of target composition and/or stoichiometry as a function of irradiation depth [7] and an *in-situ* monitoring of target properties is generally required.

Typically, this is achieved by using the well-established Nuclear Resonant Reaction Analysis (NRRA) (see, for example, [8,9] and refs. therein), which requires a narrow resonance¹ to exist in the reaction of interest and to be accessible within the dynamic range of the particle accelerator. If no resonance is present or accessible, for example because of beam energy restrictions, other methods must be employed.

This was the case of the astrophysically important $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction [10] recently studied in direct kinematics at the Laboratory for Underground Nuclear Astrophysics (LUNA) [11,12] of the Laboratori Nazionali del Gran Sasso (LNGS), INFN, Italy. Because of the small cross-sections involved at the energies investigated ($E_\alpha = 305 - 400$ keV), intense α -particle beams were needed, leading to severe target degradation and frequent target replacements. Unfortunately, no resonances exist in the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction at $E_\alpha < 400$ keV and the NRRA method could not be used to monitor the state of ^{13}C targets during irradiation. Alternatively, one could use a proton beam, also available at LUNA, on the same targets and exploit the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ reaction for NRRA analysis. However, also in this case no resonance exists that can be accessed with the 400 kV accelerator, hence a new approach to monitor the deterioration of ^{13}C targets during α -beam irradiation had to be used.

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¹ A narrow resonance is defined as one whose total width Γ is much smaller than the target thickness ΔE in energy units. The latter represents the energy lost by the ion beam in going through the target and depends on the initial beam energy as well as on the target composition and physical thickness.

In this paper, we report about an innovative application of the so-called γ -shape analysis [13]. The approach consists in a detailed study of the shape of the γ -ray lines emitted in the radiative proton-capture process $^{13}\text{C}(p,\gamma)^{14}\text{N}$ so as to periodically check both the thickness and stoichiometry of ^{13}C targets used during the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ campaign at LUNA. To validate the approach developed here, complementary NRRA measurements were also performed off-site (at Atomki in Debrecen, Hungary) on some targets, both before and after α -beam irradiation at LUNA.

The paper is organized as follows: first, we describe the NRRA technique used to characterize ^{13}C targets at Atomki (sect. 2); then, we present the γ -shape approach applied to a primary transition in the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ reaction to assess target deterioration during the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ campaign at LUNA (sect. 3); and finally, we report the results of the validation procedure (sect. 4), together with a preliminary application of the γ -shape analysis to the evaluation of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction cross-section (sect. 5).

2 Reaction yields and target properties: The NRRA approach

The NRRA method is frequently used in measurements of reaction cross-sections of astrophysical interest and has already been extensively exploited in previous studies at LUNA [13–17].

Briefly, the yield Y of a nuclear reaction can be determined from experimental quantities as [18]:

$$Y = \frac{N_R}{N_b} \quad (1)$$

where N_R is the number of reactions (producing either particles or γ rays) and N_b is the number of beam particles incident on the target. The latter quantity can be determined as Q/eq , where Q is the charge accumulated on target during beam irradiation, e is the elementary charge and q is the charge state of the projectile. On the other hand, Y is a function of the reaction cross-section σ and the number N_A of active nuclei² (per square centimetre) in the target.

For targets of thickness ΔE , corresponding to the energy lost by a beam of initial energy E_0 in traversing the target, and taking into account the energy dependence of the cross-section, the relationship between Y and the cross-section σ (at an energy E within the target) can be expressed as [18]:

$$Y(E_0) = \int_{E_0 - \Delta E}^{E_0} \frac{\sigma(E)}{\epsilon(E)} dE \quad (2)$$

² For targets consisting of chemical compounds, active nuclei are defined as those of a given species that take part in the nuclear reaction under study. All other nuclear species present in the target do not contribute to the reaction yield and are regarded as inactive.

Here, ϵ is the so-called *stopping power* which, for a given beam ion and energy, depends only on the chemical composition and stoichiometry of the target. For compound targets containing both active and inactive nuclei, the effective stopping power ϵ_{eff} is used instead, which can be parametrized using the Bragg's addition rule³ [19]:

$$\epsilon_{\text{eff}}(E) = \epsilon_{\text{A}}(E) + \sum_i \frac{N_i}{N_{\text{A}}} \epsilon_{\text{I}_i}(E) \quad (3)$$

Here N_i/N_{A} is the ratio between inactive and active nuclei, and ϵ_{A} and ϵ_{I} are the stopping powers of the corresponding (active and inactive) pure materials. Their values are available in the literature and can be calculated using SRIM [20].

If the nuclear reaction cross-section is well known, a measurement of the yield (eq. 2) can be used to experimentally determine the effective stopping power and thus to monitor the degree of deterioration of the target during beam irradiation. In particular, the NRRA method exploits the existence of a narrow and isolated resonance in a given reaction, whose cross-section is known and can be well described by the Breit-Wigner expression, σ_{BW} [19]. By measuring the yield as a function of beam energies in the proximity of the resonance and for targets of thickness ΔE much larger than the resonance width Γ , a characteristic resonance yield curve is obtained (see for example Fig.1), which contains information about the target thickness and composition. Specifically, the height of the yield plateau depends on the target stoichiometry, while the FWHM of the yield profile provides a measure of the target thickness. If either or both the target thickness and stoichiometry change as a result of intense ion beam bombardment, so will the shape of the (thick-target, resonant) yield profile and repeated resonance scans can be used to quantify the degree of target deterioration.

2.1 NRRA measurements at Atomki

Solid targets were produced by evaporating 99% enriched ^{13}C powder (by Sigma Aldrich) on 4 cm diameter tantalum backings. In order to remove traces of light elements from the Ta surface, a cleaning procedure [21] with citric acid solution was used before evaporating the targets. The evaporation was performed by the electron gun technique using a Leybold UNIVEX 350 vacuum evaporator at Atomki. The vacuum chamber of the evaporator consists of a copper melting pot, an adjustable arm used to hold the tantalum disk at 10 cm from the melting pot, and an electron gun (similar to the setup described in [22]). An oscillator quartz mounted inside the vacuum chamber at 15 cm from the melting pot was used to monitor the evaporation.

NRRA measurements were carried out at the 2 MV Medium-Current Plus Tandem Accelerator [23] at Atomki

³ For the present work a target composed of ^{13}C and Ta was assumed (see sect. 2.1) and further corrections to Bragg's rule, typically required for carbon compounds with O and H, can safely be neglected.

immediately after target production. For these measurements, a narrow resonance in the $^{13}\text{C}(\text{p},\gamma)^{14}\text{N}$ reaction ($Q = 7550.56$ keV) was used. The resonance is located at a proton beam energy $E_{\text{p}} = (1747.6 \pm 0.9)$ keV and has a width $\Gamma = (135 \pm 8)$ eV [24]. Thus, resonance scans were performed at beam energies in the range $E_{\text{p}} = 1742 - 1770$ keV.

Targets were irradiated with typical proton beam currents of $i = 500$ nA, covering a beam spot size of about 5 mm diameter. Given the low beam intensity on target, neither a cooling system nor a cold trap were needed for this setup. The target chamber was isolated from other beam-line components and acted as a Faraday cup for charge integration. An electrically insulated collimator biased to -300 V was placed at the entrance of the chamber to suppress secondary electrons. A 100% relative efficiency n-type coaxial HPGe detector was mounted in close geometry, at a distance of about 3 cm from the target, and at 0° with respect to the beam axis.

Spectra of the emitted γ rays were collected with an ORTEC MCA (model ASPEC 927) and the ORTEC MAESTRO software. The region of interest (ROI) in the γ -ray spectra was set to $E_{\gamma} = 8.0 - 9.4$ MeV ($E_{\gamma} \approx E_{\text{c.m.}} + Q$) so as to include both the full-energy peak and the single- and double-escape peaks of the direct capture transition to the ground state of the ^{14}N compound nucleus. Given the magnitude of the resonant cross-section ($\sigma_{\text{BW}} \simeq 10$ mb [25]), it was possible to reach a statistical uncertainty below 1% in less than 3 minutes of proton irradiation at the given currents, with negligible environmental background.

At a proton beam energy $E_{\text{p}} = 1747$ keV, the average target thickness was found to be 5 keV, corresponding to a physical thickness of about 170 nm and to an areal density $N_{^{13}\text{C}} \approx 10^{18}$ atoms/cm². The heights of the yield plateau of all fresh targets were consistent with each other within experimental uncertainties, indicating that all targets had the same initial stoichiometry and confirming the reproducibility of the evaporation procedure.

For some targets, the thickness uniformity was also verified by repeating the resonance scan on three different spots of the same target, 6 mm apart from each other. This requirement was especially important for the LUNA experiment because the α -particle beam has a typical diameter of about 15 mm on target, so uniformity of the evaporated layer had to be guaranteed over the whole beam-spot area. In the three spots examined, the shapes of the resonance profile were consistent within the uncertainties [26]. Based on the test measurements, no modification of stoichiometry was observed during irradiation at Atomki. In addition, NRRA was performed also on a few natural carbon targets, whose ^{13}C content is known to be 1.1%. The comparison of the plateau heights confirmed a ^{13}C abundance in the enriched targets compatible with the 99% value guaranteed by Sigma Aldrich [27].

2.2 The NRRA results

Figure 1 shows a typical resonance yield curve obtained on a fresh target (upper panel) and on a target exposed to

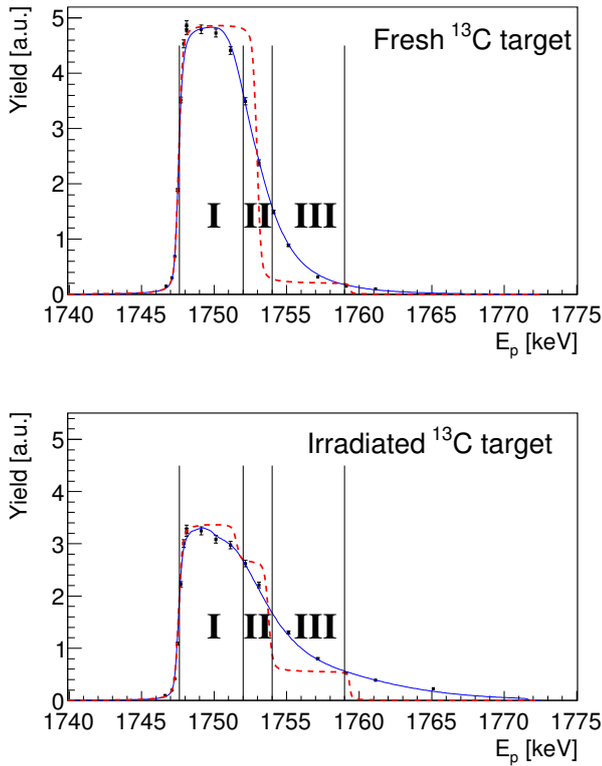


Fig. 1: $^{13}\text{C}(p,\gamma)^{14}\text{N}$ thick-target resonance yields obtained on a fresh ^{13}C target (upper panel) and on the same target after 2.1 C of accumulated α -beam charge (lower panel). Experimental data (black squares) were fit taking into account beam spread with (blue line) and without (red dashed line) beam straggling effects. Vertical lines indicate the boundaries of layers with different stoichiometries (see text for details).

about 2.1 C of α -beam irradiation at LUNA (lower panel). As can be seen, the shapes of the resonance profiles differ significantly as a result of beam exposure, both in height and FWHM of the yield plateau.

In order to quantify the degree of deterioration, experimental data (black points in fig. 1) were fit taking into account a number of experimental effects, such as beam energy resolution and beam straggling [28] within the target. These factors can be folded into the expression of the yield (eq. 2) as [19]:

$$Y(E_0) = k \int_{E_0-\Delta E}^{E_0} dE' \int_{E_i=0}^{\infty} dE_i \int_{E=0}^{E_i} \frac{\sigma(E)}{\epsilon_{\text{eff}}(E)} g(E_0, E_i) f(E_i, E, E') dE \quad (4)$$

Here, k is a normalization constant that includes the branching ratio of the transition and the γ -ray detection efficiency at the resonance energy; $g(E_0, E_i)dE_i$ describes the energy distribution of particles in the beam; and $f(E_i, E, E')dE$ describes the beam energy loss and straggling through the target (see [19] for more details). Provided all other quantities are known, a measurement

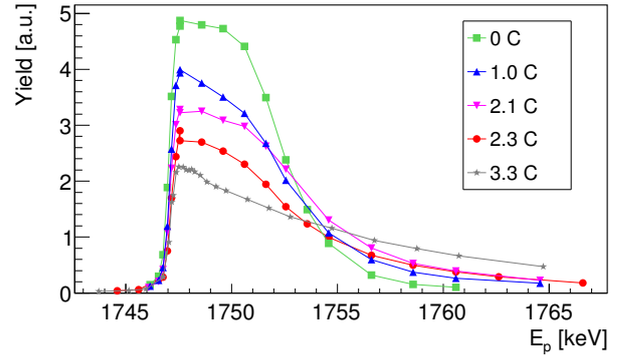


Fig. 2: (Colour online) Resonance yield profiles measured on targets with different accumulated (α -beam) charge. Lines are drawn to guide the eye.

of the resonance yield profile can be used to determine $\epsilon_{\text{eff}}(E)$ (*i.e.*, the stoichiometric ratio $N_{\text{I}}/N_{\text{A}}$) at the resonance energy.

For the present analysis, the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ reaction cross-section $\sigma(E)$ was taken from the TENDL-2017 nuclear data library [25] and evaluated as the sum of a non-resonant and a resonant component described by a second order polynomial and the Breit-Wigner formula, respectively. The stopping power $\epsilon_{\text{eff}}(E)$ was assumed to be constant over the total width ($\Gamma \simeq 135$ eV) of the resonance. The $g(E_0, E_i)dE_i$ function was assumed to follow a normal distribution with a FWHM of ~ 350 eV [23].

As for the calculation of the $f(E_i, E, E')dE$ function, assumptions on some target properties were needed. Here, it was assumed that the targets initially consisted of ^{13}C and Ta only, but with varying stoichiometric ratios as a function of depth. The TRIM software [29] was then used to calculate the energy loss and energy straggling of the beam for a given $N_{\text{Ta}}/N_{^{13}\text{C}}$ ratio. It was found that the resonance profiles could be well-reproduced by assuming three layers of different $N_{\text{Ta}}/N_{^{13}\text{C}}$ stoichiometric ratios (calculated using a χ^2 minimization), but with homogeneous composition within each layer. For fresh targets we assumed $N_{\text{Ta}}/N_{^{13}\text{C}} = 0$. The calculated yield curves obtained including the beam spread with and without beam straggling effects are shown in fig. 1 as solid (blue) and dashed (red) lines, respectively. Vertical lines indicate the boundaries of the layers with different stoichiometries. In fitting the yield profiles, the $N_{\text{Ta}}/N_{^{13}\text{C}}$ ratios in the various layers and their thickness were treated as free parameters.

Finally, NRRA measurements were also repeated on a sample of four targets after target irradiation at LUNA with different accumulated charges. The results from these measurements were used to validate the γ -shape analysis method (see sect. 4). Figure 2 shows the NRRA profiles obtained on targets with different amounts of accumulated α -beam charges. A significant modification of the resonance yield curve was observed with increasing accumulated charge during α -beam irradiation at LUNA.

Table 1: Stoichiometric ratios fitted with NRRA (third column) and corresponding effective stopping power values [$\text{keV}/10^{18}$ atoms cm^{-2}] on targets with different accumulated (α -beam) charge.

Target	Charge [C]	$N_{\text{Ta}}/N_{^{13}\text{C}}$	$\epsilon_{\text{eff}} \pm \Delta\epsilon_{\text{eff}}$
T29	0	0.000	3.12 ± 0.16
T26	1.00	0.047 ± 0.001	3.84 ± 0.19
T29	2.10	0.101 ± 0.002	4.62 ± 0.23
T28	2.34	0.149 ± 0.003	5.41 ± 0.27
MT10	3.30	0.202 ± 0.004	6.32 ± 0.32

In particular, the plateau becomes lower and the falling edge becomes longer. The observed depth profile indicates strong diffusion of ^{13}C into the Ta backings. However, the position of the leading edge of the yield curves does not change appreciably, indicating negligible carbon build-up on target surface during irradiation.

The extracted effective stopping power values given in table 1 (see sect. 4.3 for an evaluation of the uncertainties) are those corresponding to layer I⁴.

3 The γ -shape analysis method

For a non-resonant radiative capture reaction $A(x, \gamma)B$ at sub-Coulomb energies, the shape of a primary γ -ray transition is governed by the behaviour of the reaction cross-section $\sigma(E)$ over the energy range covered by the incident beam as it loses energy in traversing the target [13]. For a thick target, the shape is also influenced by the energy dependence of the stopping power, and by the concentration profile of active nuclei as a function of target depth (which may change during irradiation).

Additional experimental effects may further contribute to the exact shape of the γ -ray line and must be taken into account. Specifically, the high-energy rise of the peak may be Doppler-shifted by the recoil of the compound nucleus, while its low-energy tail may be affected by beam straggling effects. Thus dY_i , the number of counts per unit of charge in channel i of the acquired γ -ray spectrum, with central value E_{γ_i} (E_i is the corresponding projectile energy) and width ΔE_{γ} is given by the expression [13]:

$$dY_i = A \frac{\sigma(E_i)}{\epsilon_{\text{eff}}(E_i)} \Delta E_{\gamma} \zeta(E_{\gamma}) P(E_i) f(E_i, E, E') dE_i, \quad (5)$$

where A is a normalization constant that includes the branching ratio of the transition and the γ -ray detection efficiency, $\zeta(E_{\gamma})$ is a Gaussian function accounting for the experimental broadening effects: the HPGe energy resolution (roughly 10 keV at $E_{\gamma} = 7840$ keV) and the Doppler

broadening (about 6 keV) caused by the finite angular range covered by the detector in close geometry. The function $P(E_i)$ describes the concentration profile of active nuclei within the target (see below), and $f(E_i, E, E')dE_i$ describes the energy broadening due to beam straggling effects.

The target concentration profile $P(E)$ can be modelled as the product of two Fermi functions [4]:

$$P(E) = \left[\exp\left(\frac{E - E_0}{\delta_1}\right) + 1 \right]^{-1} \left[\exp\left(\frac{E_0 - E - \Delta E}{\delta_2}\right) + 1 \right]^{-1} \quad (6)$$

where E_0 is the incident beam energy, ΔE the target thickness, and δ_1 and δ_2 are two parameters accounting, respectively, for the slopes of the falling and leading edges of the thick-target profile.

The analysis of γ -ray line shapes has been extensively used in the past to extract information on unknown cross-sections of astrophysical reactions ([30–32]), provided that the target profile $P(E)$ could be measured independently (*e.g.*, through NRRA analysis).

In the present study, we exploited instead the γ -shape analysis approach to determine $P(E)$ and the effective stopping power ϵ_{eff} using the well-known cross-section of the $^{13}\text{C}(p, \gamma)^{14}\text{N}$ reaction, as explained in the following sections.

3.1 The γ -shape measurements at LUNA

In order to monitor the target degradation during the $^{13}\text{C}(\alpha, n)^{16}\text{O}$ measurements, data taking at LUNA consisted of long α -beam runs with accumulated charges of ≈ 1 C per run, interspersed by short proton-beam runs with typical accumulated charges of 0.2 C at most, so as to minimize possible changes in target stoichiometry caused by the proton irradiation itself.

Proton beam runs were all performed at the same reference energy, $E_p = 310$ keV. The choice for this energy was dictated by the need to maximize counting statistics while minimizing beam-induced background from a broad resonance at $E_p \simeq 340$ keV in the $^{19}\text{F}(p, \alpha\gamma)^{16}\text{O}$ reaction on always present ^{19}F contaminants in the experimental setup. Note that at such a low proton-beam energy the resulting target thickness is $\Delta E \simeq 15$ keV and neither the $^{13}\text{C}(p, \gamma)^{14}\text{N}$ reaction cross-section nor the effective stopping power ϵ_{eff} can be regarded as constant.

Primary γ rays ($E_{\gamma} = 7840$ keV) arising from the $^{13}\text{C}(p, \gamma)^{14}\text{N}$ direct capture transition into the ^{14}N ground state (hereafter, DC \rightarrow GS transition) were detected using a HPGe detector with a relative efficiency of 120% and FWHM of 2.8 keV at $E_{\gamma} = 1460$ keV. The detector was mounted at 55° to the beam axis and brought to a distance of 5 mm from the target holder [33]. The same type of electronics and DAQ used in the NRRA measurement was used to acquire the γ -spectrum at LUNA.

⁴ Indeed, as the cross-section of $^{13}\text{C}(\alpha, n)^{16}\text{O}$ reaction drops exponentially with energy, the outermost layers of the target gives the main contribution to the reaction yield. Thus, only the stoichiometric ratio of layer I are of interest here.

3.2 The γ -shape analysis and results

Figure 3 shows the DC \rightarrow GS peak ($E_\gamma = 7840$ keV) of two γ -ray spectra acquired on a fresh target (upper panel) and after an α -beam irradiation of 3.3 C of total accumulated charge (lower panel).

Experimental spectra (blue crosses) were fit using eq. (5), where the low-energy trend of the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ reaction cross-section was taken from King *et al.* [34] and Genard *et al.* [35], and the beam straggling distribution function $f(E_i, E, E')dE_i$ was evaluated by Monte Carlo simulations using TRIM.

For runs on fresh targets, parameters A , ΔE , δ_1 and δ_2 in eq. 5 and 6 were left free to vary, while the stoichiometric ratio $N_{\text{Ta}}/N_{^{13}\text{C}}$ was set to 0, as no degradation had yet occurred. For runs on irradiated targets, parameters A and δ_1 were fixed to the fit values of the “fresh” target, leaving ΔE , δ_2 and $N_{\text{Ta}}/N_{^{13}\text{C}}$ as free parameters.

The results of the fitting procedure are shown as red curves in Fig. 3, while dash-dotted green curves show target profiles $P(E)$ defined in eq. (6) in arbitrary units. Note the change in the shape of the target profile $P(E)$ following irradiation with the α -beam. A linear background (dashed line) was included in the ROI of the fit to account for multiple Compton-scatter events in the HPGe detector. The χ^2 was minimized in the region delimited by vertical lines, for a number of degrees of freedom $\nu = 40$. We obtained a reduced $\tilde{\chi}^2 \approx 1.6$ for both plots shown.

Table 2 reports the values of the fit parameters for both spectra shown in fig. 3. As expected, $N_{\text{Ta}}/N_{^{13}\text{C}}$ and δ_2 show a significant change, indicating a strong modification in the target stoichiometry and a likely diffusion of ^{13}C nuclei into the backing.

Table 2: Parameter values obtained from the γ -shape fits to the peaks in fig. 3. The normalization constant is $A = (3.16 \pm 0.01) \times 10^{-4}$ (in a.u.) for both profiles.

Q [C]	$N_{\text{Ta}}/N_{^{13}\text{C}}$	ΔE [keV]	δ_1 [keV]	δ_2 [keV]
0	0	21.7 ± 0.1	0.33 ± 0.03	4.13 ± 0.22
3.3	0.16 ± 0.011	22.2 ± 0.1	0.33 ± 0.03	10.19 ± 0.22

4 Validation and discussion

4.1 Role of inactive nuclides in the γ -shape analysis

In order to check the effect of possible light contaminants (*e.g.*, H, He, C, O) on the effective stopping power, we performed several SRIM calculations for proton energies $E_p = 280 - 310$ keV, and alpha energies $E_\alpha = 300 - 400$ keV (relevant to the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ data taking campaign). In the energy ranges considered, the energy dependence of stopping power for each element (H, He, C, O), assumed as the only contaminant in the target, changes by less than 3% for proton projectiles and less than 5% for alpha particles. Similar conclusions can be drawn in the case where

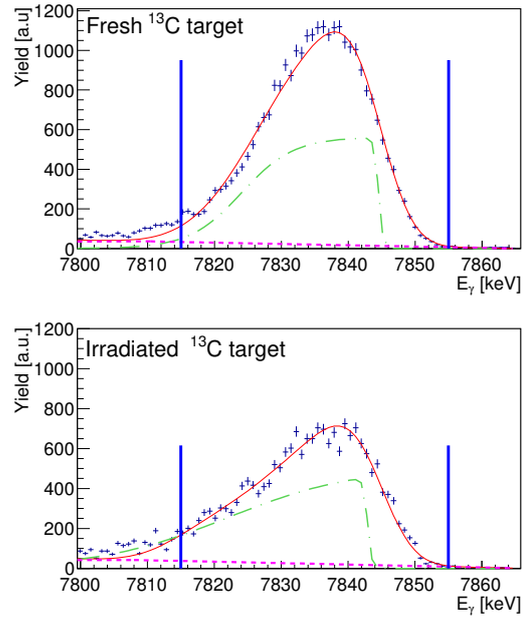


Fig. 3: (Colour online) Gamma-ray peak from the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ DC \rightarrow GS transition as obtained on a fresh ^{13}C target (upper panel) and on the same target irradiated with α -beam for 3.3 C of accumulated charge (lower panel). Experimental data were fit (red line) using eq. 5 and including a linear background (dashed line). The χ^2 was minimized in the region delimited by the vertical lines. In dash-dotted green, the target profile $P(E)$ (in arbitrary units), as defined in eq. 6, shows the concentration of active nuclei as a function of depth (*i.e.* beam energy within the target).

more than one contaminant is present at the same time. These conclusions were further supported by additional ERDA analysis performed on irradiated targets at the Ion Beam Center of Helmholtz-Zentrum Dresden-Rossendorf. The analysis confirmed that the concentration of elements such as H, He and O after the α -beam irradiation at LUNA was at most 10% [36]. We conclude that, for our γ -shape analysis, the effective stopping power is essentially insensitive to the actual species of inactive nuclei present in the target [27]. Stoichiometric values N_I/N_A obtained from the γ -shape fit are reported in table 3 for each one of the inactive species considered, together with the associated stopping powers for proton and α beams.

4.2 Comparison of NRRA and γ -shape analysis results

To validate the results of the γ -shape analysis approach, a comparison to the results obtained with the well-established NRRA method was made. To this end, the effective stopping powers arising from the stoichiometric ratios obtained with the NRRA at $E_p = 1747.6$ keV (table 1) were recalculated at $E_p = 310$ keV using eq. (3) assuming that the targets consist of a compound of only ^{13}C and Ta.

Table 3: Stoichiometric ratios for possible inactive nuclei (H, He, C, O and Ta), as obtained from a γ -shape fit of the primary γ ray in $^{13}\text{C}(p,\gamma)^{14}\text{N}$. The corresponding effective stopping powers are calculated for a proton beam at $E_p = 310$ keV and an α beam at $E_\alpha = 400$ keV.

Inactive species	N_I/N_A	$\epsilon_{\text{eff}}(p)$ [keV/ 10^{18} atoms/cm 2]	$\epsilon_{\text{eff}}(\alpha)$ [keV/ 10^{18} atoms/cm 2]
H	1.92 \pm 0.15	14.36 \pm 1.44	59.96 \pm 3.21
He	1.19 \pm 0.11	14.51 \pm 1.45	54.79 \pm 4.14
^{12}C	0.55 \pm 0.023	14.51 \pm 1.45	57.67 \pm 3.15
O	0.48 \pm 0.016	14.65 \pm 1.46	56.65 \pm 2.80
Ta	0.16 \pm 0.011	14.53 \pm 1.45	53.45 \pm 2.64

401 Table 4 reports the values of the effective stopping
402 powers obtained with the two methods for different accu-
403 mulated (α -beam) charges. The results obtained are in
404 agreement within uncertainties (see sect. 4.3 for the un-
certainties evaluation).

Table 4: Effective stopping powers [keV/ 10^{18} atoms/cm 2] calculated at $E_p = 310$ keV using the NRRA and the γ -shape analysis approach for targets of different accumulated (α -beam) charge.

Target	Charge [C]	γ -shape $\epsilon_{\text{eff}} \pm \Delta\epsilon_{\text{eff}}$	NRRA $\epsilon_{\text{eff}} \pm \Delta\epsilon_{\text{eff}}$
T29	0	9.38 \pm 0.48	9.37 \pm 0.47
T26	1.00	10.53 \pm 1.05	10.83 \pm 0.54
T29	2.10	12.15 \pm 1.21	12.51 \pm 0.63
T28	2.34	13.49 \pm 1.35	14.01 \pm 0.70
MT10	3.30	14.53 \pm 1.45	15.64 \pm 0.78

4.3 Uncertainties budget

407 The overall uncertainty on the effective stopping power
408 evaluation has three main contributions: a 4-5% system-
409 atic error on SRIM tabulated values of stopping powers
410 for pure materials (common to both methods); a 1% and
411 a 3% systematic error on the charge integration on tar-
412 get for measurements at Atomki (NRRA) and LUNA (γ -
413 shape), respectively; a 2% and an 8% fit uncertainty on the
414 extracted stoichiometric ratios from the NRRA and the γ -
415 shape approaches, respectively. In both approaches, fit un-
416 certainties were calculated [37] by varying the $N_{\text{Ta}}/N_{^{13}\text{C}}$
417 within a range $[N_{\text{Ta}}/N_{^{13}\text{C}} \pm \delta]$ until the χ^2 value increased
418 by a fixed amount $\Delta\chi^2$ (which depends on the number of
419 fit parameters, 3.2 in this specific case) around its mini-
420 mum value. The overall uncertainty on the effective stop-
421 ping power was then obtained by summing in quadrature
422 all sources of errors and resulted in an overall 5% error
423 for the NRRA measurements and an overall 10% error for
424 the γ -shape analysis, respectively. A summary of the main
425 uncertainties for the two techniques is presented in table
426 5.

Table 5: Summary of main uncertainties for the two tech-
niques.

Uncertainty source	NRRA	γ -shape
Charge accumulation	1%	3%
Stopping power from SRIM	5%	5%
Evaluation of stoichiometric ratio	2%	8%
Total uncertainty	5%	10%

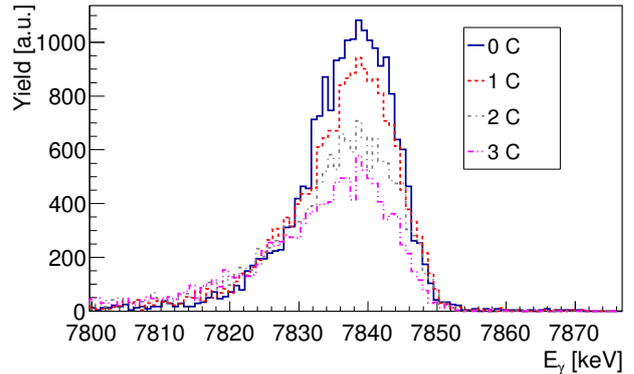


Fig. 4: (Colour online) Overlay of γ -ray spectra for the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ DC \rightarrow GS transition acquired on the same target at different accumulated (α -beam) charge. Both the height of the peak and its FWHM change with increased charge as expected, indicating severe target modification during α -beam irradiation.

5 Target degradation correction applied to the evaluation of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction cross-section

427 During the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ data taking campaign, over a
428 hundred ^{13}C targets were used for an overall accumulated
429 α -beam charge of about 300 C. For each target, γ -ray spec-
430 tra acquired during short proton runs, taken before and
431 after long α -beam irradiation runs, were analyzed follow-
432 ing the procedure described in sect. 3 to correct for target
433 degradation effects in the evaluation of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$
434 reaction cross-section.
435
436
437

438 As an example, fig. 4 shows a superposition of the
439 DC \rightarrow GS peak in four γ -ray spectra acquired on the
440 same target at increasing values of accumulated (α -beam)
441 charge. As expected, the higher the accumulated charge,
442 the lower the (p,γ) yield and the broader the shape of the
443 γ -ray line.

444 From fits to each peak, we extracted values of $N_{\text{Ta}}/N_{^{13}\text{C}}$
445 at the target surface and plotted them as a function of
446 the accumulated charge Q (see fig. 5, where for clarity,
447 results are displayed for three targets only). Open sym-
448 bols in the figure correspond to stoichiometric ratios de-
449 termined with the γ -shape analysis method on reference
450 proton runs, while filled symbols correspond to linearly
451 interpolated values. The latter were used to calculate av-
452 erage effective stopping powers (eq. 3) to be used in the

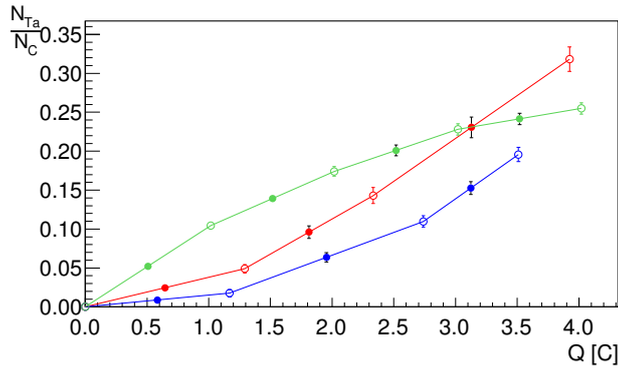


Fig. 5: (Colour online) Stoichiometric ratios $N_{\text{Ta}}/N_{^{13}\text{C}}$ as a function of accumulated charge Q on three different targets (represented by different colours). Open symbols correspond to values obtained with the γ -shape analysis; filled symbols represent linearly interpolated values.

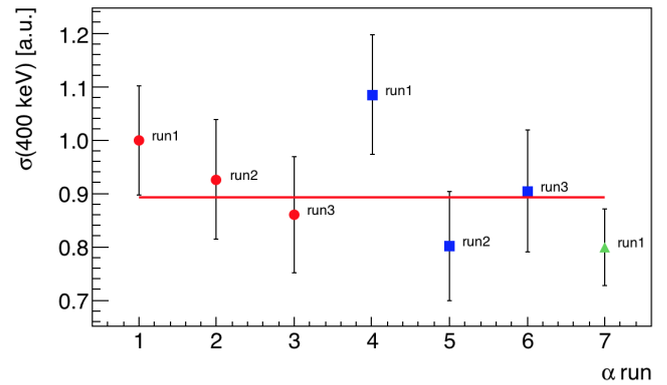


Fig. 6: (Colour online) $^{13}\text{C}(\alpha,n)^{16}\text{O}$ cross-section (in a.u.) extracted from different α -beam runs on three different targets (indicated in red, blue and green). The solid red line represents the weighted average of the data points shown. Error bars include statistical and systematic uncertainties in the ϵ_{eff} evaluation.

evaluation of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction cross-section, thus accounting for target degradation during long α -beam irradiation in-between successive proton runs.

Figure 6 shows the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ cross-sections (in arbitrary units) evaluated from measurements performed at the same beam energy ($E_{\alpha} = 400$ keV) on three different targets. The error bars shown arise from a combination of statistical and systematic uncertainties in the ϵ_{eff} evaluation. All data points are within 2σ from the weighed average (red line). Final results on the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ cross-section over the full energy range ($E_{\alpha} = 305 - 400$ keV) covered at LUNA will be presented in a forthcoming publication.

6 Conclusions

In this paper we reported on a new application of the γ -shape analysis used to monitor *in situ* the degradation of ^{13}C targets exposed to intense α -beam irradiation during the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction study at LUNA. Specifically, fits to the peak shape of the $\text{DC} \rightarrow \text{GS}$ γ -ray transition in the $^{13}\text{C}(p,\gamma)^{14}\text{N}$ reaction were used to obtain quantitative information on target degradation as a function of accumulated (α -beam) charge on target.

The γ -shape analysis was used as an alternative to the standard NRRA, whose application at LUNA was precluded by the lack of appropriate resonances in the energy range accessible with the 400 kV accelerator. NRRA measurements were, instead, performed at Atomki, both to characterize initial target thickness and stoichiometry and, for a subset of targets, as a way to validate the γ -shape analysis. A comparison of the stoichiometric values obtained with both methods shows agreement within experimental uncertainties.

We also verified that the effective stopping powers used in the evaluation of the $^{13}\text{C}(\alpha,n)^{16}\text{O}$ reaction cross-sections were independent from the assumption of inactive contaminant(s) present in the target.

The effective stopping power values obtained with the γ -shape analysis were extracted with an overall 10% uncertainty. While the use of the γ -shape analysis was validated specifically for ^{13}C targets in the present study, this approach may have wider applications especially where the use of traditional analytical methods, such as the NRRA is not possible.

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