Cross sections of proton-induced reactions on ¹⁵²Gd, ¹⁵⁵Gd and ¹⁵⁹Tb with emphasis on the production of selected Tb radionuclides

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Abstract

Cross sections are presented for various Dy, Tb and Gd radionuclides produced in the proton bombardment of ¹⁵⁹Tb as well as for the reactions ¹⁵²Gd(p,4n)¹⁴⁹Tb and ¹⁵⁵Gd(p,4n)¹⁵²Tb up to 66 MeV. The experimental excitation functions are compared with theoretical predictions by means of the geometrydependent hybrid (GDH) model as implemented in the code ALICE/ASH, as well as with values from the TENDL-2012 library and previous literature experimental data, where available. Physical yields have been derived for the production of some of the medically important radioterbiums, namely ¹⁴⁹Tb (radionuclide therapy), ¹⁵²Tb (PET) and ¹⁵⁵Tb (SPECT). The indirect production of high-purity ¹⁵⁵Tb via the decay of its precursor ¹⁵⁵Dy is reported. The possibility of a large-scale production facility based on a commercial 70 MeV cyclotron is also discussed.

Keywords:

66 MeV protons, radionuclide production, ¹⁴⁹Tb, ¹⁵²Tb, ¹⁵⁵Tb, excitation functions, integral yields

1. Introduction

Several members of the radiolanthanides have been studied in recent years for their potential in nuclear medicine. Recently, Müller *et al.* [1] reported on a preclinical study of new tumour-targeting radiopharmaceuticals labeled with a unique quadruplet of Tb radionuclides, namely ¹⁴⁹Tb, ¹⁵²Tb, ¹⁵⁵Tb and ¹⁶¹Tb. These radionuclides are unique in that they collectively contain properties suitable for all thee major modalities of nuclear medicine, namely PET, SPECT and radionuclide therapy, from the same element. Müller *et al.* reported excellent results with a folate-based targeting agent containing a DOTA chelator for binding the Tb to the biomolecule. Both ¹⁵²Tb (PET) and ¹⁵⁵Tb (SPECT) imaging of folate receptor (FR)-positive human tumours xenografted into mice were shown to be of high quality. In addition, the same compound labeled with the therapeutic Tb radionuclides (¹⁴⁹Tb and ¹⁶¹Tb) demonstrated conclusive results on remission of the disease. While independent studies were performed with the α -particle emitter (¹⁴⁹Tb) and the β^- emitter (¹⁶¹Tb), the authors speculated that cocktails of these two isotopes to optimize the efficacy of

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the treatment might be an intriguing option, which required further systematic investigation. For further information as well as an account of prior work, the reader is referred to Ref. [1].

The majority of the radioterbiums is not yet generally available. Except for the reactor produced ¹⁶¹Tb, the large-scale production of the other radioterbiums, with a radionuclidic purity suitable for medical use, has not yet been demonstrated. In one of the earlier studies in the literature, Levin et al. [2] demonstrated the photonuclear production of ¹⁵⁵Tb via the reaction ¹⁵⁶Dy(γ ,n)¹⁵⁵Dy \rightarrow ¹⁵⁵Tb, using bremsstrahlung with a maximum energy of 25 MeV at an electron accelerator. A radionuclidic purity > 99.9% could be obtained, which is substantially higher than what can be obtained using reactions suitable for a cyclotron. While their electron beam current was rather low and the corresponding yield relatively small, these authors speculated about large-scale production possibilities with future electron accelerators having orders of magnitude higher beam intensities. A number of studies investigated the use of ¹²C beams. Allen *et al.* [3] and Zaitseva et al. [4] studied the ^{nat}Nd(${}^{12}C,xn$) ${}^{152}Dy \rightarrow {}^{152}Tb$ and ${}^{nat}Nd({}^{12}C,xn) {}^{149}Dy \rightarrow {}^{149}Tb$ reactions, respectively. While useful yields could be obtained, the end product inevitable contained a mixture of Tb radionuclides. This is clearly a general problem when charged-particle induced reactions are utilized, even on highly enriched targetry. Electromagnetic (EM) isotope separation is used at the ISOLDE facility of CERN for the collection of radioterbiums produced in proton-induced spallation reactions on Ta [1, 3, 5]. EM separation may yet prove to be the only feasible way to ensure a radionuclidic purity close to 100%. It is interesting to contemplate this form of separation, in conjunction with nuclear reactions at energies low enough to be suitable for commercial cyclotrons, as an option for the large-scale production of radioterbiums in a dedicated facility. Compared to the large scale of ISOLDE, a facility based on a commercial cyclotron would indeed be small and compact.

This present study reports on the work of a collaboration established to measure production cross sections for the direct formation of 149,152,155 Tb in 152,155 Gd + p as well as for the indirect formation of 155 Tb via the decay of its precursor 155 Dy in 159 Tb + p. Enriched 152 Gd and 155 Gd target materials were used as well as natural Tb, which is mono-isotopic 159 Tb (100%). The new measurements were prompted, in part, by a previous study on proton-induced reactions on nat Gd [6] up to 66 MeV. This is close to the energy region covered by the new generation of commercial 70 MeV cyclotrons [7] which can deliver beams of high intensity (several hundred μ A). A particular aspect of the present investigation is to establish whether a commercial 70 MeV cyclotron will be suitable for the large-scale production of 149 Tb, 152 Tb and 155 Tb.

The relevant part of the nuclide chart is presented in Fig. 1. This a simplified diagram in that many of the shorter-lived metastable states, which feed the respective ground states, have been omitted. Also, the data were taken from the compilation of Firestone and Eckström [8]. The relevant proton-induced nuclear reactions are listed in Table 1. As already discussed in Ref. [6], this part of the nuclide chart is particularly unfriendly (from a radionuclide production point of view) due to seven stable Gd nuclei, which lead to many open reaction channels. In addition, a relatively large number of Tb radionuclides have half-lives in the region of several hours to a few days. Consequently, the production of the medically important Tb radionuclides in a radionuclidically pure form, and not only in a no-carrier-added (NCA) form, is rather compromised. Another complicating factor is that the location of ¹⁴⁹Tb is quite far from the line of stability. As will be discussed later, it may yet transpire that EM isotope separation may have to be a compulsory component of a large-scale production facility for these Tb radionuclides.

Cross sections are presented for 153,155,157,159 Dy, 153,154m2,155,156 Tb and 151,153 Gd in 159 Tb + p as well as for 152 Gd(p,4n) 149 Tb and 155 Gd(p,4n) 152 Tb. For most of these reactions, no previously measured data could be found in the literature. Physical yields have also been derived for the reactions relevant to 149,152,155 Tb production.

2. Experimental

2.1. Irradiations

The well-known stacked-foil technique was employed to measure the excitation functions of radionuclides produced in ¹⁵⁹Tb + p from the (p,n) reaction threshold up to 66 MeV. In the case of ¹⁵²Gd + p and ¹⁵⁵Gd + p, only the energy region relevant to the (p,4n) reaction was investigated. Three foil stacks were irradiated in this study, henceforth referred to as Stacks 1, 2 and 3, using 66 MeV proton beams supplied by the separated-sector cyclotron (SSC) facility of iThemba LABS. The foil stacks were bombarded in an accurately calibrated Faraday chamber mounted at the end of an external beam line. This irradiation chamber, based on the design of the RERAME (REcoil RAnge MEasurements) facility of the Laboratory Nazionale del Sud (LNS) in Catania, Italy, has been described in detail elsewhere [9]. The beam was collimated to a spot of 4 mm in diameter. The average beam current was 50 nA and the irradiation time was 2 h for each stack. The beam current and accumulated charge were measured with a Brookhaven Instruments Corporation Model 1000C current integrator. The current and charge values were also logged in 1 second intervals to an analysis computer using the LabVIEW software. The beam energy was accurately measured using a calibrated 90° bending magnet.

Stacks 1 and 2 contained thin targets prepared from oxides of Gd and Tb using the sedimentation technique developed by Rösch *et al.* [10]. Gadolinium trioxide (Gd_2O_3) and tetraterbium heptaoxide (Tb_4O_7) powders were obtained from Isoflex USA and Koch Chemicals Ltd., respectively. Due to limited quantities of the enriched Gd powder, only relatively few targets could be made. In each case, fine powder of the target material was sedimented onto a 26 μ m thick Ti backing and covered with a 10 μ m thick Al foil. The diameter of a sediment layer was 13 mm, defined by the inner diameter of the sedimentation cells used. The thicknesses of the sediment layers varied between 1.4 and 4.4 mg/cm² but nominally 2.5 mg/cm² for ¹⁵⁹Tb₄O₇, 4 mg/cm² for ¹⁵²Gd₂O₃ and 1.5 mg/cm² for ¹⁵⁵Gd₂O₃. The thickness of each target sediment and foil was determined accurately by weighing. Stack 1 contained seven ¹⁵²Gd and seven ¹⁵⁹Tb targets while Stack 2 contained seven ¹⁵⁵Gd targets, covering the energy region 30-62 MeV. The target foils were interspersed with Cu monitor foils of nominally 108 mg/cm² thickness (99.99%, Goodfellow, UK) as well as Cu and Al degrader foils of various thicknesses, as required. Just prior to the bombardment, it was decided to degrade the 66 MeV proton beam by a few MeV to better match the expected energy region of the (p,4n) reaction for both 152 Gd + p and 155 Gd + p with the assembled energy region of each foil stack. This was done with a single Cu degrader placed at the front of each stack. Irradiations of similar stacks at a beam energy of 30 MeV, to cover the energy region relevant to reactions with lower thresholds, have been planned but have not yet been performed. Consequently, as the data sets for (p,xn) reactions where x < 4are still incomplete, only results for the (p,4n) reactions in 152,155 Gd + p are presented at this stage.

Stack 1 had another inherent limitation, namely that the ¹⁵²Gd targets had an enrichment level of only 30.60%. The content of the other stable Gd isotopes are as follows: ¹⁵⁴Gd (9.3%), ¹⁵⁵Gd (18.1%), ¹⁵⁶Gd (14.8%), ¹⁵⁷Gd (8.6%), ¹⁵⁸Gd (11%) and ¹⁶⁰Gd (7.6%). Although not ideal, these measurements are nevertheless expected to provide useful information for the important reaction ¹⁵²Gd(p,4n)¹⁴⁹Tb, while the interpretation of several of the other (p,xn) reactions may be rather complicated. Note that the natural abundance of ¹⁵²Gd is only 0.2%, the reason why ¹⁴⁹Tb was not observed in the ^{*nat*}Gd + p study [6]. An increase of the ¹⁵²Gd content by more than two orders of magnitude should rectify this. In the case of the ¹⁵⁵Gd targets, however, no such complications are expected as the level of enrichment was 99.82%.

The decision to irradiate a third stack containing metallic Tb foils was made later. This was done, in part, as a check on the results from the Tb_4O_7 sediment targets. Results from experiments using the sedimentation technique sometimes show unusually large scatter, which could potentially be a problem in stacked-foil experiments having so few targets as Stacks 1 and 2 in this work. The scatter is thought to be a result of

sediment targets being rather fragile, sometimes suffering damage either before or during bombardment. A second reason was that for 159 Tb + p, almost no excitation function data could be found in the literature, which made it an interesting study in its own right. Also, because of the delay in the investigation of the lower energy region, it was decided to design Stack 3 to cover the entire energy region down to the threshold of the (p,n) reaction. Normally, we prefer not to degrade the 66 MeV beam so far down due to the increasing energy uncertainty with increasing penetration depth in the stack. On the other hand, the lower energy region for 159 Tb + p could always be re-investigated later with a lower energy incident beam (e.g. 30 MeV), should compelling reasons exist to do so. Stack 3 consisted of elemental Tb foils of nominally 18.6 mg/cm² thickness (99%, Goodfellow Metals, UK), Cu monitor foils of nominally 45.8 mg/cm² (99.9%, Goodfellow, UK) throughout the stack, as well as Ti monitor/degrader foils of nominally 115 mg/cm² in the region between 20 and 35 MeV. A selection of Al and Cu foils of various thicknesses were used as degraders.

2.2. Radionuclide assays

After bombardment, the activated foils and sediments were repeatedly assayed by means of standard off-line γ -ray spectrometry. Two calibrated HPGe detectors were used. Above about 120 keV, a coaxial Ge detector with a relative efficiency of 18% and a resolution of 1.8 keV at 1.33 MeV was used. For lower photon energies, in particular the 58 keV γ -ray of ¹⁵⁹Dy and the 97 keV γ -ray of ¹⁵³Gd, a planar Ge detector containing a 10 mm thick crystal and a thin Be window was used. This was done because the planar detector has a higher efficiency and a significantly lower Compton background for these low-energy γ -rays. The photopeak areas were determined by means of the quantitative Canberra Genie 2000 analysis software in conjunction with a Canberra DSA 1000 multi-channel analyser system. The efficiency curves for these detectors were determined by means of standard calibrated γ -ray sources traceable to either the BIPM or NIST.

2.3. Data analysis

The cross sections of the observed activation products were calculated from their measured γ -ray emissions using decay data from the online compilation of Firestone and Eckström [8]. The γ -lines used to identify the nuclides of interest are listed in Table 2. The well-known activation formula (see e.g. [11]) was used and corrections were made for decay losses during and after bombardment as well as during counting.

The accumulated charge was directly measured by means of a calibrated current integrator. Consequently, the cross sections extracted from the Cu and Ti monitor foils served only as a consistency check. The results for the nat Cu(p,x) 62 Zn monitor reaction are shown in Fig. 2 for all three stacks, together with the standard excitation function recommended by the IAEA [12]. Only the logged beam currents from the electronic current integrator were used for deriving the experimental cross sections. These measurements reproduce the recommended curve satisfactorily, therefore, there was no need to make any corrections based on the monitor reaction results. A similar quality of agreement was found for the nat Ti(p,x) 48 V monitor reaction.

Corrections for beam current losses due to nonelastic nuclear interactions were made according to the prescription and tables of Janni [13]. The average proton energy in each foil or sediment was calculated using the code STACK, which contains the stopping power formulae of Anderson and Ziegler [14]. These stopping powers were checked against the newer compilation of Berger [15] and found to be in excellent agreement. A number of independent checks were also performed using the Monte Carlo code SRIM [16] and found to be consistent with the published stopping power tables.

The total uncertainties in the measured cross sections were obtained by summing all the contributing uncertainties in quadrature and are expressed with a 1σ (68%) confidence level. In addition to statistical

uncertainties, a systematic uncertainty was estimated to be about 7%, including the uncertainty in beam current integration (3%), detector efficiency (5%), counting geometry (1%), decay corrections (2%) and foil thickness (3%) [17]. The uncertainty in energy of each measured data point was estimated from the uncertainty in incident beam energy, foil thickness and energy straggling in the stack.

3. Theoretical calculations

In this work, theoretical excitation functions were calculated using the Geometry Dependent Hybrid (GDH) model as implemented in the code ALICE/ASH [18]. The calculations were performed using the recommended values for the input parameters according to the comments in the preamble to the code. Level densities were calculated according to the generalized superfluid model (GSM). The normal pairing shift was selected. Experimental nuclidic masses were used where available, else calculated using the built-in Myers and Swiatecki mass formula of the code. A level density parameter of a = A/9 and an initial energy bin size of 0.5 MeV were used. The calculations were performed within the frame of the GDH model for the pre-equilibrium emission of protons and neutrons. The subsequent evaporation during the equilibrium stage was calculated according to the Weiskopf-Ewing formalism as implemented in the code. The evaporated particles included protons, neutrons, deuterons and α -particles.

To avoid the known scatter in the high energy "tail" regions of some of the excitation functions, especially in the case of the (p,n) reactions, we adopted the practice of performing sets of calculations, varying the energy bin size (parameter *ED* in the code) from 0.4 to 0.6 MeV in steps of 0.01 MeV, followed by taking the arithmetic average over all sets. This has been reported on extensively in a previous paper [19]. In cases where the scatter was not problematic, an energy bin size of 0.5 MeV was adopted.

In addition to the ALICE/ASH predictions, the measured excitation functions are also compared with the relevant cross sections compiled in the TENDL-2012 library [20, 21].

4. Results and Discussion

The results from Stacks 1 and 2 (i.e. the ${}^{152}\text{Gd}_2\text{O}_3$, ${}^{155}\text{Gd}_2\text{O}_3$ and ${}^{159}\text{Tb}_4\text{O}_7$ sediment targets) are presented in Table 3. The results from Stack 3 (i.e. the Tb metal foils) are presented in Tables 4, 5 and 6. The excitation functions are shown in Figs. 3 though 14 and the derived physical yield curves and other related quantities (e.g. radionuclidic purity) in Figs. 15 through 18.

4.1. The ${}^{152}Gd(p,4n){}^{149}Tb$ reaction

Because the ¹⁵²Gd₂O₃ targets only had a 30.6% level of enrichment of ¹⁵²Gd, several other reactions also contributed to the direct formation of ¹⁴⁹Tb, such as ¹⁵⁴Gd(p,6n)¹⁴⁹Tb (Q = -43.36 MeV) and ¹⁵⁵Gd(p,7n)¹⁴⁹Tb (Q = -49.80 MeV). The ¹⁵²Gd(p,4n)¹⁴⁹Tb (Q = -28.21 MeV) reaction, however, is expected to dominate the excitation function. The measured data are shown in Fig. 3, scaled for targets of 100% enrichment. Comparisons are made with ALICE/ASH predictions for the (p,4n) reaction on its own, as well as the sum of the contributions from the (p,4n), (p,6n) and (p,7n) reactions mentioned above, weighed according to the relative ¹⁵²Gd, ¹⁵⁴Gd and ¹⁵⁵Gd abundances in the target material, respectively. The ALICE/ASH predictions overestimate the cross sections considerably and have been renormalized (with a scale factor of SF = 0.31) for purposes of comparison with the experimental data. One can clearly see in the figure that both ALICE/ASH curves reproduce the shape of the excitation function quite well, even though the data points are rather few. The excitation function peaks at about 250 mb at an energy of 42 MeV. Also, the contributions from the (p,6n) and (p,7n) reactions are indeed small and feature only at energies above about 55 MeV. Figure 3 also shows the TENDL-2012 values for the (p,4n) reaction which are substantially larger than the measured values and have been renormalized to the measured data for purposes of comparison (scale factor SF = 0.39). The TENDL-2012 data do not reproduce the shape of the experimental excitation function nearly as well as the ALICE/ASH predictions and appear to be shifted towards lower energies by about 4–5 MeV.

4.2. The ${}^{155}Gd(p,4n){}^{152}Tb$ reaction

Concerning the measurements, only the (p,4n) reaction on ¹⁵⁵Gd contributed significantly to ¹⁵²Tb formation as the ¹⁵⁵Gd enrichment level was close to 100%. The experimental cross sections are shown in Fig. 4, together with ALICE/ASH predictions and the relevant TENDL-2012 values. The ALICE/ASH and TENDL-2012 values both underestimate the excitation function and have therefore been renormalized to the measurements for purposes of comparison (scale factors SF = 1.7 and SF = 1.3, respectively). Both predictions are somewhat shifted relative to the measurements – TENDL-2012 towards lower energies and ALICE/ASH towards higher energies. The spline fit through the measured data was used for calculating the integral yield curve, which will be presented later. The excitation function reaches a maximum of about 900 mb at energy of about 39 MeV.

4.3. The ${}^{159}Tb(p,7n){}^{153}Dy$ reaction

The ¹⁵⁹Tb(p,7n)¹⁵³Dy (Q = -49.6 MeV) reaction could only be observed in three of the Tb metal foils of Stack 3, at the highest energies, as shown in Fig. 5. It is presented here for completeness only. Both ALICE/ASH and TENDL-2012 do not reproduce the onset of the excitation function in the threshold region very well, therefore, a spline fit through the measured data, extrapolated to 70 MeV, has been used to calculate the physical yield curve. This will be further discussed later.

4.4. The ${}^{159}Tb(p,5n){}^{155}Dy$ reaction

In the case of the 159 Tb(p,5n) 155 Dy (Q = -33.45 MeV) reaction, both the ALICE/ASH prediction, the TENDL-2012 data and the experimental data agree reasonably well in magnitude at the maximum of the excitation function. However, as shown in Fig. 6, the TENDL-2012 results seem to be systematically shifted towards lower energies, while the ALICE/ASH prediction is shifted towards higher energies. There is also good agreement in shape between the ALICE/ASH prediction and the measurements, the main difference being the approximately 2 MeV shift. In contrast, the TENDL-2012 excitation function shows a significantly narrower width in the peak region. While not perfect, the overall agreement between the measurements from Stacks 1 and 3 is acceptable – the majority of the experimental points from the two data sets agree within the experimental uncertainties. This is very reassuring. The excitation function reaches a maximum of about 520 mb at 48 MeV. As before, a spline fit through the measured data, shown in the figure, was used for calculating integral yields.

4.5. The ${}^{159}Tb(p,3n){}^{157}Dy$ reaction

The results for the ¹⁵⁹Tb(p,3n)¹⁵⁷Dy (Q = -17.04 MeV) reaction are shown in Fig. 7. The agreement with the ALICE/ASH prediction is very satisfactory, both in absolute magnitude and in shape (the theoretical maximum only slightly higher), while the TENDL-2012 values are somewhat lower with the peak position shifted towards the lower energy side by about 4 MeV. The overall agreement between the measurements from Stacks 1 and 3 is also acceptable. The excitation function reaches a maximum of 1040 mb at about 27 MeV.

4.6. The ${}^{159}Tb(p,n){}^{159}Dy$ reaction

Figure 8 shows the results for the ¹⁵⁹Tb(p,n)¹⁵⁹Dy (Q = -1.148 MeV) reaction. The excitation function peaks just over 100 mb at an energy of about 10.4 MeV. The agreement with the only set of previously published data found in the literature, by Hassan *et al.* [22], is reasonable. The ALICE/ASH calculations and TENDL-2012 values largely overpredict the excitation function, however, ALICE/ASH reproduces the peak region quite well when the values are scaled with a factor SF = 0.42. We also noted a strange anomaly in the (p,n) data of TENDL-2012 for a range of target nuclei. There is a notable but spurious second local maximum towards higher energies, which is not characteristic of (p,n) reactions. Previous versions of TENDL also show this anomaly (see Fig. 8). We checked the TENDL data of the (p,n) reaction for many lighter and heavier nuclei as well. The anomaly is notably present for the majority of nuclei between Eu and Bi but seems to be absent below Z = 61. In fact, for some nuclei between Eu and Bi the TENDL-2012 values show a prominent second peak.

4.7. The ${}^{159}Tb(p,X){}^{153}Tb$ process

Several reaction channels are open for the formation of ¹⁵³Tb ($T_{1/2} = 2.3$ d) e.g. ¹⁵⁹Tb(p,p6n)¹⁵³Tb (Q = -46.65 MeV), ¹⁵⁹Tb(p,d5n)¹⁵³Tb (Q = -44.42 MeV), ¹⁵⁹Tb(p,t4n)¹⁵³Tb (Q = -38.16 MeV) as well as indirectly via the the decay of the precursor ¹⁵³Dy ($T_{1/2} = 6.4$ h). The measurements were performed after the complete decay of the shorter-lived precursor and the resulting cross sections are therefore cumulative. The measurements are presented in Fig. 9, as well as the ALICE/ASH and TENDL-2012 predictions. It is impossible to make a judgment on the quality of the theoretical predictions as the measurements reflect only the steep slope near an effective threshold somewhat above 50 MeV. It seems, however, as if the TENDL-2012 values are shifted towards lower energies and the ALICE/ASH values towards higher energies. Neither curves can be successfully scaled to the data, therefore, the scale factors were kept at a value of SF = 1. This may be an indication that energy shifts are more likely than over- and underpredictions.

4.8. The ${}^{159}Tb(p,X){}^{154m2}Tb$ process

Similar to the results of the ^{nat}Gd + p [6], it was only possible to extract cross sections for the m2 state in ¹⁵⁴Tb. Both the ALICE/ASH and TENDL-2012 curves had to be scaled considerably for purposes of making a comparison with the measurements (SF = 0.075), as shown in Fig. 10. In this particular case, the cross sections are for the direct production of ^{154m2}Tb due to the very long half-life of the precursor, ¹⁵⁴Dy ($T_{1/2} = 3 \times 10^6$ y). Even though the relevant reaction Q-values are all below 40 MeV (¹⁵⁹Tb(p,p5n)^{154m2}Tb, Q = -39.93 MeV; ¹⁵⁹Tb(p,d4n)^{154m2}Tb, Q = -37.51 MeV; ¹⁵⁹Tb(p,t3n)^{154m2}Tb, Q = -31.25 MeV) no ^{154m2}Tb were detected below 53 MeV. It is difficult to make a conclusive judgement about the agreement between the experimental measurements and the theoretical predictions. It seems, however, that the TENDL-2012 values reproduce the shape of the excitation function better in this case, albeit with an order of magnitude overprediction.

4.9. The ${}^{159}Tb(p,X){}^{155}Tb$ process

The excitation function of ¹⁵⁵Tb ($T_{1/2} = 5.42$ d) contains contributions which are directly produced (¹⁵⁹Tb(p,p4n)¹⁵⁵Tb, Q = -30.57 MeV; ¹⁵⁹Tb(p,d3n)¹⁵⁵Tb, Q = -28.34 MeV; ¹⁵⁹Tb(p,t2n)¹⁵⁵Tb, Q = -22.09 MeV) as well as an indirectly produced contribution via the decay of a shorter-lived precursor, ¹⁵⁵Dy ($T_{1/2} = 9.9$ h). These measurements were performed after the decay of the precursor, thus, the resulting cross sections are cumulative. As shown in Fig. 4.11, the data span several orders of magnitude and are therefore plotted on a log scale. The agreement with the ALICE/ASH prediction is satisfactory, both in shape and absolute magnitude, although the theoretical values are shifted towards higher energies by about

2 MeV. In contrast, the TENDL-2012 values are shifted somewhat towards lower energies and underestimate the maximum cross section. The excitation function peaks at a value of about 670 mb at 51 MeV.

4.10. The ${}^{159}Tb(p,X){}^{156}Tb$ process

Two relatively long-lived metastable states, ^{156m1}Tb ($T_{1/2} = 1.02$ d) and ^{156m2}Tb ($T_{1/2} = 5$ h), both decay to the ground state of ¹⁵⁶Tb ($T_{1/2} = 5.4$ d). The measurements were performed after the near-complete decay of the metastable states. There is no Dy precursor (¹⁵⁶Dy is stable). The experimental results are shown in Fig. 12, together with the ALICE/ASH calculations and the TENDL-2012 values. The TENDL-2012 results are in excellent agreement with the measured data. The ALICE/ASH prediction seems to start at a too-high threshold and also slightly overpredicts the excitation function maximum.

4.11. The ${}^{159}Tb(p,X){}^{151}Gd$ process

Only a few values could be measured above 50 MeV, dominated by the ¹⁵⁹Tb($p,\alpha 5n$)¹⁵¹Gd (Q = -30.84 MeV) reaction as the ¹⁵⁹Tb(p,2p7n)¹⁵¹Gd (Q = -59.89 MeV) is not contributing significantly in this energy region. The results are shown in Fig. 13. The ALICE/ASH prediction is reasonable while the TENDL-2012 values are shifted towards lower energies.

4.12. The ${}^{159}Tb(p,X){}^{153}Gd$ process

The contributions from both the ¹⁵⁹Tb($p,\alpha 3n$)¹⁵³Gd (Q = -15.99 MeV) and ¹⁵⁹Tb(p,2p5n)¹⁵³Gd (Q = -44.30 MeV) reactions are clearly visible, as shown in Fig. 14. The sharp rise starting just below 60 MeV is from the latter reaction, which is not well reproduced by the ALICE/ASH predictions, nor by TENDL-2012. This kind of behaviour in an excitation function is not unusual and has been described before, e.g. for the ⁶⁸Zn(p,X)⁶⁴Cu reaction, where the contributions from the ⁶⁸Zn($p,\alpha n$)⁶⁴Cu and the ⁶⁸Zn(2p,3n)⁶⁴Cu reactions show a similar trend [9].

4.13. Integral yield calculations

Thick target yields for ¹⁴⁹Tb and ¹⁵²Tb, derived from the excitation functions for the ¹⁵²Gd(p,4n)¹⁴⁹Tb and ¹⁵⁵Gd(p,4n)¹⁵²Tb reactions, respectively, are shown in Fig. 15. Note that these are *physical yields*. In the case of ¹⁴⁹Tb, the scaled ALICE/ASH prediction shown in Fig. 3 (solid curve) was used for this purpose, while the spline fit in Fig. 4 was used for ¹⁵²Tb. With an energy window of 66 MeV down to threshold (about 30 MeV), a physical yield of 2556 MBq/ μ Ah (69.1 mCi/ μ Ah) can be expected for ¹⁴⁹Tb and 1924 MBq/ μ Ah (52.0 mCi/ μ Ah) for ¹⁵²Tb from the above (p,4n) reactions, respectively. These are very high yields but, unfortunately, do not lead to radionuclidically pure products, as already mentioned. The value for ¹⁵²Tb is higher than the rather conservative estimate of 1200 MBq/ μ Ah (32.4 mCi/ μ Ah) from the *nat*Gd + p study [6].

Thick target yields for ^{153,155,157,159}Dy in ¹⁵⁹Tb +p, derived from spline fits through the measured excitation function data (see Figs. 5–8) are shown in Fig. 16. It is evident that the ¹⁵⁵Tb produced via ¹⁵⁵Dy decay will contain the decay products of a considerable amount of ¹⁵⁷Dy as well, in any energy window. However, ¹⁵⁷Tb is long-lived with a half-life of 71 y [8], therefore only low levels of ¹⁵⁷Tb activity are to be expected in the final product. The yield of ¹⁵⁹Dy is quite low as it is also quite long-lived ($T_{1/2} = 144.4$ d) and furthermore it decays to stable ¹⁵⁹Tb, thus, it is no contributor to any radiocontaminant. Nevertheless, stable Tb will affect labeling efficiency, which should be taken into consideration. Lastly, ¹⁵³Dy will contribute ¹⁵³Tb as a radiocontaminant but only above 59 MeV.

To produce ¹⁵⁵Tb via ¹⁵⁵Dy decay would require two chemical separations. The Dy should be separated from the Tb target material as soon as possible after the end of bombardment (EOB). An optimum waiting

time would then be required to allow for maximum growth of ¹⁵⁵Tb, followed by a second chemical separation to isolate the Tb from the remaining Dy. In order to evaluate this production route, one needs to investigate the growth and decay characteristics of all the relevant Dy radionuclides on a common timeline. It is also necessary to make some assumptions on the duration of the chemical separations. Let t = 0 be the time when an isolated quantity of pure ¹⁵⁵Dy starts to decay. The growth and decay curve of ¹⁵⁵Tb formed via the decay of the ¹⁵⁵Dy is shown in Fig. 17, where $A_d(t)$ denotes the activity of the daughter at time t and $A_m(0)$ denotes the activity of the mother radionuclide at time t = 0. The curve reaches a maximum at t = 39.6 h, which should be close to the optimum waiting time after the first chemical separation before performing the second chemical separation, depending on the effective duration of the separation procedure. The dashed line in Fig. 17 shows the points in time where the ¹⁵⁵Tb activity reaches 90% of the maximum. These times are t = 21.6 h and t = 72.5 h during the periods of net growth and net decay, respectively. This is rather convenient, as it indicates that there is an interval of many hours during which the second chemical separation can be performed. A FORTRAN code developed previously [23] was adapted to follow the growth and decay of all the produced Dy radionuclides on a common timeline, taking the durations of chemical separations into account. In fact, the bombardment time, waiting times and processing times are all variables. Different production scenarios can therefore be investigated. The code calculates the yield and radionuclidic purity as well as the ¹⁵⁵Tb purity as a percentage of all Tb nuclei produced.

As an example of the kind of results one can obtain, the conditions of Table 7, which we consider to be realistic, will be used in the following calculations. A bombardment time of 10 h is just over one half-life of the mother radionuclide, ¹⁵⁵Dy. Waiting time 1 is the time interval between EOB and the start of the first chemical separation and includes tasks such as the removal of the target from the beamline, transport to a reception hot cell, decapsulation and transfer of the irradiated material to a processing hot cell. The actual ion-exchange procedure of the chemical separations is assumed to have a duration of 1 h each. Waiting time 2 is the period required for the accumulation of ¹⁵⁵Tb from ¹⁵⁵Dy decay and is taken as 39 h. Waiting time 3 is a short period between the end of the second chemical separation and the reference time, allowing for the removal of the product from the hot cell and performing a measurement of the activity. The exit energy of the energy window is taken as 35 MeV and the entrance energy is varied between 50 and 70 MeV. The results are shown in Fig. 18, plotted versus the entrance proton energy. The yield rises from 39.8 GBq (1074 mCi) at an incident energy of 50 MeV to 116.6 GBq (3151 mCi) at an incident energy of 70 MeV. It is evident that this production route is suitable for the production of Ci quantities of 155 Tb. The radionuclidic purity is better than 99.9% at all incident energies below 60 MeV but decreases monotonically above 60 MeV to a value of about 88% at 70 MeV. The number of ¹⁵⁵Tb nuclei, expressed as a percentage of all produced Tb nuclei, has a value of 57.5% at 40 MeV, increasing towards higher energies until reaching a maximum of 68.1% at 62 MeV, then decreases slightly to a value of 65.4% at 70 MeV.

It is evident that this indirect production route can provide high yields of ¹⁵⁵Tb with excellent radionuclidic purity, however, nuclei of other stable and long-lived Tb species are also produced. This reduces the purity in terms of the fraction of all Tb nuclei produced, which may affect the labeling efficiency.

5. Electromagnetic Isotope Separation

Electromagnetic (EM) isotope separation (also called EM mass separation) may be the only method to obtain highly radionuclidically pure ¹⁴⁹Tb. In principle, the ¹⁵²Gd(p,n)¹⁵²Tb and ¹⁵⁵Gd(p,n)¹⁵⁵Tb reactions can provide high purity ¹⁵²Tb and ¹⁵⁵Tb, respectively, provided that target material with a sufficiently high level of enrichment (close to 100%) are used. EM isotope separation will, however, be required if the respective high-yield (p,4n) reactions are utilized. In fact, it might be possible to collect sufficient quantities

of pure ¹⁵²Tb and ¹⁵⁵Tb simultaneously with the ¹⁴⁹Tb, depending on the chosen target material. It is, therefore, interesting to contemplate the possibility of using EM isotope separation as the technology of choice to remove isotopic impurities. Recently, Lapi *et al.* [24] and D'Auria *et al.* [25] demonstrated efficient EM isotope separation for ⁹⁹Mo. These authors also speculated about using EM isotope separation for many other medically important radionuclides, some of which can be produced in sufficient quantities via reactions such (n,γ) and (γ,n) but which result in products with a low specific activity. While the principles of EM isotope separation are well known for many decades, practical issues remain, e.g. ion source reliability and efficiency as well as the challenges associated with handling high levels of radioactivity after extensive operation at high beam intensities.

It is the belief of the majority of the present authors that should ¹⁴⁹Tb prove to be important enough to nuclear medicine at some time in the future, the present engineering problems, which preclude largescale productions with good radionuclidic purity, will be solved. This happened before, e.g. in the case of ¹⁸F. Furthermore, future dedicated facilities may be small and compact, in stark contrast to the large spallation + ISOL facilities such as ISOLDE at CERN. A possible scenario may include a commercial 70 MeV cyclotron, such as the IBA Cyclone 70 XP, Best BSCI 70p or CIAE CYCCIAE70 [7] in conjunction with a compact EM isotope separator, such as that of the Ion Source Test Facility (ISTF) at TRIUMF [25]. This last-mentioned facility is a fantastic step in the right direction and may be ideally suited for proof-of-principle studies. In fact, prior chemical processing of batch targets may provide feedstock of already relatively high specific activity for the EM separator system, which will then provide the final purifying step.

6. Conclusion

Measured cross sections for the 152 Gd(p,4n) 149 Tb, 155 Gd(p,4n) 152 Tb and 159 Tb(p,xn) 153,155,157,159 Dy reactions have been presented. In the case of 159 Tb + p, cross sections for various Tb and Gd radionuclides have also been extracted, some directly produced and others cumulatively. Integral yields for 149 Tb and 152 Tb in the proton bombardment of enriched Gd targets have been calculated, as well as the yield and purity to be expected from the indirect production route 159 Tb(p,5n) 155 Dy $\rightarrow {}^{155}$ Tb. It is shown that excellent yields can readily be obtained with a commercial 70 MeV cyclotron, however, EM isotope separation may be the only way to achieve 149,152,155 Tb products with a high level of radionuclidic purity and the desired specific activity. Furthermore, it is speculated that dedicated, compact facilities for their production may be feasible. However, this would require a great deal of further experimental investigation and development work. A facility such as the ISTF at TRIUMF may be ideally suited to pursue such studies.

Concerning nuclear data needs, a consistent experimental investigation of the 152 Gd(p,n) 152 Tb and 155 Gd(p,n) 155 Tb reactions is still lacking. This will require 152 Gd + p and 155 Gd + p stacked-foil experiments in the energy region from the respective (p,n) reaction thresholds up to about 30 MeV.

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Reaction	<i>Q</i> value (MeV)	Natural Isotopic Abundance ^a	Reaction	Q value (MeV)	
152 Gd(p,4n) 149 Tb	-28.21	0.20%	¹⁵⁹ Tb(p,n) ¹⁵⁹ Dy	-1.148	
154 Gd(p,6n) 149 Tb	-43.36	2.18%	159 Tb(p,3n) 157 Dy	-17.04	
155 Gd(p,7n) 149 Tb	-49.80	14.80%	159 Tb(p,5n) 155 Dy	-33.45	
156 Gd(p,8n) 149 Tb	-58.33	20.47%	159 Tb(p,7n) 153 Dy	-49.60	
152 Gd(p,n) 152 Tb	-4.77	0.20%	¹⁵⁹ Tb(p,p3n) ¹⁵⁶ Tb	-23.66	
154 Gd(p,3n) 152 Tb	-19.91	2.18%	¹⁵⁹ Tb(p,d2n) ¹⁵⁶ Tb	-21.43	
155 Gd(p,4n) 152 Tb	-26.35	14.80%	¹⁵⁹ Tb(p,tn) ¹⁵⁶ Tb	-15.17	
156 Gd(p,5n) 152 Tb	-34.89	20.47%	¹⁵⁹ Tb(p,p4n) ¹⁵⁵ Tb	-30.57	
157 Gd(p,6n) 152 Tb	-41.25	15.65%	¹⁵⁹ Tb(p,d3n) ¹⁵⁵ Tb	-28.34	
158 Gd(p,7n) 152 Tb	-49.18	24.84%	¹⁵⁹ Tb(p,t2n) ¹⁵⁵ Tb	-22.09	
160 Gd(p,9n) 152 Tb	-62.58	21.86%	¹⁵⁹ Tb(p,p5n) ^{154m2} Tb	-39.73	
154 Gd(p, γ) 155 Tb	4.83	2.18%	159 Tb(p,d4n) 154m2 Tb	-37.51	
155 Gd(p,n) 155 Tb	-1.61	14.80%	159 Tb(p,t3n) 154m2 Tb	-31.25	
156 Gd(p,2n) 155 Tb	-10.14	20.47%	¹⁵⁹ Tb(p,p6n) ¹⁵³ Tb	-46.65	
157 Gd(p,3n) 155 Tb	-16.50	15.65%	¹⁵⁹ Tb(p,d5n) ¹⁵³ Tb	-44.42	
158 Gd(p,4n) 155 Tb	-24.44	24.84%	¹⁵⁹ Tb(p,t4n) ¹⁵³ Tb	-38.16	
160 Gd(p,6n) 155 Tb	-37.83	21.86%	159 Tb(p, α 3n) 153 Gd	-15.99	
160 Gd(p, γ) 161 Tb	6.81	21.86%	¹⁵⁹ Tb(p,2p5n) ¹⁵³ Gd	-44.30	
			159 Tb(p, α 5n) 151 Gd	-30.84	
			159 Tb(p,2p7n) 151 Gd	-59.89	

Table 1: Proton-induced reactions for the relevant Tb, Dy and Gd radionuclides.

^aTaken from Firestone & Eckström [8].

Nuclide	Half-life	Decay mode	γ -rays (keV)	Intensity (%)	
¹⁴⁹ Tb	4.118 h	$\epsilon + \beta^+: 83.3\%$ $\alpha: 16.7\%$	352.24	29.43	
¹⁵² Tb	17.5 h	$\epsilon + \beta^+$: 100%	271.13	8.6	
		$\beta^+: 17\%$	344.27	65.0	
			586.26	9.4	
¹⁵³ Tb	2.3 d	$\epsilon + \beta^+$: 100%	212.00	31.0	
^{154m2} Tb	22.7 h	$\epsilon + \beta^+$: 98.2%	225.94	26.8	
		IT: 1.8%			
¹⁵⁵ Tb	5.32 d	<i>ϵ</i> : 100%	180.10	7.45	
			367.23	1.48	
¹⁵⁶ Tb	5.35 d	$\epsilon + \beta^+$: 100%	199.21	40.9	
			356.42	13.6	
			534.32	66.6	
¹⁵³ Dy	6.4 h	$\epsilon + \beta^+$: 99.99%	213.75	10.9	
¹⁵⁵ Dy	9.9 h	$\epsilon + \beta^+$: 100%	226.92	68.4	
¹⁵⁷ Dy	8.14 h	$\epsilon + \beta^+$: 100%	326.16	92.0	
¹⁵⁹ Dy	144.4 d	<i>ϵ</i> : 100%	58.00	2.22	
¹⁵¹ Gd	124.0 d	<i>ϵ</i> : 100%	153.60	6.20	
			243.28	5.60	
¹⁵³ Gd	240.4 d	<i>ϵ</i> : 100%	97.43	29.0	
			103.18	21.1	

Table 2: Investigated Tb, Dy and Gd radionuclides and their decay properties used for experimental cross-section determinations^a.

^aTaken from Firestone & Eckström [8].

Proton energy (MeV)	Cross section (mb)							
	$^{152}{ m Gd}(p,\!4n)^{149}{ m Tb}$	¹⁵⁵ Gd(p,4n) ¹⁵² Tb	¹⁵⁹ Tb(p,5n) ¹⁵⁵ Dy	¹⁵⁹ Tb(p,3n) ¹⁵⁷ Dy				
31.73 ± 1.11	_	$(2.24 \pm 0.21) \times 10^{+2}$	_	_				
35.69 ± 0.98	_	- ´	_	$(3.73 \pm 0.34) \times 10^{+2}$				
35.52 ± 0.99	$(1.71 \pm 0.19) \times 10^{+2}$	_	_	- ´				
36.71 ± 0.96	–	$(8.04 \pm 0.74) \times 10^{+2}$	_	_				
41.31 ± 0.84	$(2.48 \pm 0.27) \times 10^{+2}$	-	_	_				
41.46 ± 0.83	- ´	_	$(1.80 \pm 0.17) \times 10^{+2}$	$(2.03 \pm 0.19) \times 10^{+2}$				
42.22 ± 0.82	_	$(8.21 \pm 0.75) \times 10^{+2}$	_	_				
46.55 ± 0.71	$(1.63 \pm 0.19) \times 10^{+2}$	_	_	_				
46.69 ± 0.71	_	_	$(5.07 \pm 0.47) \times 10^{+2}$	$(1.33 \pm 0.12) \times 10^{+2}$				
47.25 ± 0.70	_	$(3.77 \pm 0.35) \times 10^{+2}$	_	_				
52.25 ± 0.58	$(9.43 \pm 1.28) \times 10^{+1}$	_	_	_				
52.38 ± 0.58	_	_	$(5.52 \pm 0.51) \times 10^{+2}$	$(1.12 \pm 0.10) \times 10^{+2}$				
52.76 ± 0.57	_	$(2.33 \pm 0.21) \times 10^{+2}$	_	_				
57.52 ± 0.48	$(5.92 \pm 0.88) \times 10^{+1}$	_	_	_				
57.64 ± 0.48	_	_	$(4.16 \pm 0.38) \times 10^{+2}$	$(1.05 \pm 0.10) \times 10^{+2}$				
57.87 ± 0.47	_	$(1.63 \pm 0.15) \times 10^{+2}$	_	_				
62.46 ± 0.39	$(4.41 \pm 0.69) \times 10^{+1}$	_	_	_				
62.58 ± 0.39	_	_	$(2.10 \pm 0.19) \times 10^{+2}$	$(8.79 \pm 0.82) \times 10^{+1}$				
62.68 ± 0.39	_	$(1.20 \pm 0.12) \times 10^{+2}$	_	_				

Table 3: Measured cross sections for the production of Tb and Dy radionuclides in the irradiation of ^{152,155}Gd and ^{nat}Tb with protons – results from Stacks 1 and 2.

Proton energy (MeV)	Cross section (mb)						
	¹⁵³ Dy	¹⁵⁵ Dy	¹⁵⁷ Dy	¹⁵⁹ Dy			
8.35 ± 2.53	_	_	_	$(6.45 \pm 0.46) \times 10^{+1}$			
8.76 ± 2.45	_	_	_	$(7.73 \pm 0.55) \times 10^{+1}$			
10.43 ± 2.22	_	_	-	$(1.01\pm 0.07)\times 10^{+2}$			
11.91 ± 2.06	_	_	-	$(9.10 \pm 0.64) \times 10^{+1}$			
13.28 ± 1.94	-	_	-	$(7.16 \pm 0.51) \times 10^{+1}$			
15.47 ± 1.78	_	_	$(1.17 \pm 0.11) \times 10^{+0}$	$(4.23 \pm 0.30) \times 10^{+1}$			
17.45 ± 1.66	-	_	$(2.81 \pm 0.20) \times 10^{+1}$	$(2.89 \pm 0.21) \times 10^{+1}$			
19.28 ± 1.56	-	_	$(1.63 \pm 0.11) \times 10^{+2}$	$(2.54 \pm 0.18) \times 10^{+1}$			
20.98 ± 1.48	-	—	$(4.41 \pm 0.31) \times 10^{+2}$	$(2.36 \pm 0.17) \times 10^{+1}$			
23.75 ± 1.37	-	—	$(8.47 \pm 0.59) \times 10^{+2}$	$(2.05 \pm 0.15) \times 10^{+1}$			
26.89 ± 1.25	-	-	$(1.04 \pm 0.07) \times 10^{+3}$	$(1.90 \pm 0.14) \times 10^{+1}$			
29.21 ± 1.17	-	—	$(9.70 \pm 0.68) \times 10^{+2}$	$(1.70 \pm 0.13) \times 10^{+1}$			
31.41 ± 1.11	-	-	$(7.53 \pm 0.53) \times 10^{+2}$	$(1.57 \pm 0.11) \times 10^{+1}$			
33.49 ± 1.04	-	-	$(5.23 \pm 0.37) \times 10^{+2}$	$(1.47 \pm 0.11) \times 10^{+1}$			
35.48 ± 0.99	-	-	$(3.38 \pm 0.24) \times 10^{+2}$	$(1.32 \pm 0.09) \times 10^{+1}$			
39.01 ± 0.89	-	$(3.90 \pm 0.28) \times 10^{+1}$	$(2.11 \pm 0.15) \times 10^{+2}$	$(1.31 \pm 0.09) \times 10^{+1}$			
42.31 ± 0.81	-	$(2.16 \pm 0.15) \times 10^{+2}$	$(1.61 \pm 0.11) \times 10^{+2}$	$(1.18 \pm 0.09) \times 10^{+1}$			
45.42 ± 0.74	-	$(4.29 \pm 0.30) \times 10^{+2}$	$(1.38 \pm 0.10) \times 10^{+2}$	$(1.13 \pm 0.08) \times 10^{+1}$			
48.39 ± 0.67	-	$(5.19 \pm 0.36) \times 10^{+2}$	$(1.17 \pm 0.08) \times 10^{+2}$	$(1.05 \pm 0.08) \times 10^{+1}$			
51.25 ± 0.61	-	$(5.06 \pm 0.35) \times 10^{+2}$	$(1.04 \pm 0.07) \times 10^{+2}$	$(1.00 \pm 0.07) \times 10^{+1}$			
54.00 ± 0.55	-	$(4.75 \pm 0.33) \times 10^{+2}$	$(9.97 \pm 0.70) \times 10^{+1}$	$(9.65 \pm 0.80) \times 10^{+0}$			
56.65 ± 0.50	-	$(3.75 \pm 0.26) \times 10^{+2}$	$(9.37 \pm 0.66) \times 10^{+1}$	$(8.86 \pm 0.72) \times 10^{+0}$			
59.87 ± 0.44	$(6.97 \pm 0.71) \times 10^{+0}$	$(2.89 \pm 0.20) \times 10^{+2}$	$(9.15 \pm 0.64) \times 10^{+1}$	$(8.46 \pm 0.59) \times 10^{+0}$			
62.97 ± 0.36	$(3.36 \pm 0.25) \times 10^{+1}$	$(2.14 \pm 0.15) \times 10^{+2}$	$(8.37 \pm 0.59) \times 10^{+1}$	$(8.14 \pm 0.66) \times 10^{+0}$			
65.95 ± 0.30	$(7.75 \pm 0.55) \times 10^{+1}$	$(1.65 \pm 0.12) \times 10^{+2}$	$(7.45 \pm 0.52) \times 10^{+1}$	$(7.16 \pm 0.72) \times 10^{+0}$			

Table 4: Measured cross sections for the production of Dy radionuclides in the irradiation of *nat*Tb with protons.

Proton energy (MeV)	Cross section (mb)							
	¹⁵³ Tb	154m Tb	¹⁵⁵ Tb	¹⁵⁶ Tb				
23.75 ± 1.37	_	_	_	$(2.19 \pm 3.33) \times 10^{-1}$				
26.89 ± 1.25	_	_	_	$(1.12 \pm 0.64) \times 10^{+0}$				
29.21 ± 1.17	_	_	_	$(2.39 \pm 0.59) \times 10^{+0}$				
31.41 ± 1.11	_	_	_	$(3.86 \pm 0.29) \times 10^{+0}$				
33.49 ± 1.04	_	_	$(4.18 \pm 1.72) \times 10^{-1}$	$(7.46 \pm 0.54) \times 10^{+0}$				
35.48 ± 0.99	_	_	$(1.27 \pm 0.21) \times 10^{+0}$	$(1.41 \pm 0.10) \times 10^{+1}$				
39.01 ± 0.89	_	_	$(4.99 \pm 0.36) \times 10^{+1}$	$(4.11 \pm 0.29) \times 10^{+1}$				
42.31 ± 0.81	_	_	$(2.61 \pm 0.18) \times 10^{+2}$	$(7.41 \pm 0.52) \times 10^{+1}$				
45.42 ± 0.74	_	_	$(5.34 \pm 0.37) \times 10^{+2}$	$(1.05 \pm 0.07) \times 10^{+2}$				
48.39 ± 0.67	_	_	$(6.63 \pm 0.46) \times 10^{+2}$	$(1.12 \pm 0.08) \times 10^{+2}$				
51.25 ± 0.61	_	_	$(6.73 \pm 0.47) \times 10^{+2}$	$(1.14 \pm 0.08) \times 10^{+2}$				
54.00 ± 0.55	$(3.08 \pm 1.59) \times 10^{-1}$	$(1.12 \pm 0.30) \times 10^{+0}$	$(6.65\pm 0.47)\times 10^{+2}$	$(1.24 \pm 0.09) \times 10^{+2}$				
56.65 ± 0.50	$(1.99 \pm 0.22) \times 10^{+0}$	$(1.41 \pm 0.18) \times 10^{+0}$	$(5.63 \pm 0.39) \times 10^{+2}$	$(1.23 \pm 0.09) \times 10^{+2}$				
59.87 ± 0.44	$(1.38 \pm 0.10) \times 10^{+1}$	$(2.92 \pm 0.26) \times 10^{+0}$	$(4.71 \pm 0.33) \times 10^{+2}$	$(1.30 \pm 0.09) \times 10^{+2}$				
62.97 ± 0.36	$(4.95 \pm 0.35) \times 10^{+1}$	$(4.99 \pm 0.51) \times 10^{+0}$	$(3.92 \pm 0.27) \times 10^{+2}$	$(1.23 \pm 0.09) \times 10^{+2}$				
65.95 ± 0.30	$(1.14 \pm 0.08) \times 10^{+2}$	$(5.76 \pm 0.55) \times 10^{+0}$	$(3.30 \pm 0.23) \times 10^{+2}$	$(1.16 \pm 0.08) \times 10^{+2}$				

Table 5: Measured cross sections for the production of Tb radionuclides in the irradiation of *nat*Tb with protons.

Proton energy (MeV)	Cross section (mb)					
	¹⁵¹ Gd	¹⁵³ Gd				
33.49 ± 1.17	_	$(2.36 \pm 0.22) \times 10^{-1}$				
35.48 ± 1.11	_	$(9.67 \pm 0.70) \times 10^{-1}$				
39.01 ± 1.04	_	$(4.57 \pm 0.32) \times 10^{+0}$				
42.31 ± 0.99	_	$(8.80 \pm 0.62) \times 10^{+0}$				
45.42 ± 0.89	_	$(1.17 \pm 0.08) \times 10^{+1}$				
48.39 ± 0.81	_	$(1.44 \pm 0.10) \times 10^{+1}$				
51.25 ± 0.74	_	$(8.92 \pm 0.63) \times 10^{+0}$				
54.00 ± 0.68	$(9.06 \pm 1.14) \times 10^{-1}$	$(8.83 \pm 0.62) \times 10^{+0}$				
56.65 ± 0.50	$(1.59 \pm 0.15) \times 10^{+0}$	$(8.34 \pm 0.56) \times 10^{+0}$				
59.87 ± 0.46	$(5.71 \pm 0.41) \times 10^{+0}$	$(2.07 \pm 0.15) \times 10^{+1}$				
62.97 ± 0.39	$(9.26 \pm 0.65) \times 10^{+0}$	$(3.98 \pm 0.28) \times 10^{+1}$				
65.95 ± 0.30	$(1.11 \pm 0.08) \times 10^{+1}$	$(7.80 \pm 0.55) \times 10^{+1}$				

Table 6: Measured cross sections for the production of Gd radionuclides in the irradiation of nat Tb with protons.

Production step	Duration (h)
Bombardment time	10.0
Waiting time 1	0.5
Chemical separation 1 ^{<i>a</i>}	1.0
Waiting time 2	39.0
Chemical separation 2^b	0.5
Waiting time 3	0.5

Table 7: Production conditions for calculating the ¹⁵⁵Tb yield via ¹⁵⁵Dy decay in the proton bombardment of a thick ¹⁵⁹Tb target.

Exit proton energy: 35 MeV

Incident proton energy: Variable in range 50 - 70 MeV

^{a 155}Dy from ¹⁵⁹Tb target matrix
 ^{b 155}Tb from recovered ¹⁵⁵Dy

Dy-150	Dy-151	Dy-152	Dy-153	Dy-154	Dy-155	Dy-156	Dy-157	Dy-158	Dy-159	Dy-160	Dy-161	Dy-162
7.17 m	17.9 m	2.38 h	6.4 h	3.0E6 y	9.9 h	0.06%	8.14 h	0.10%	144.4 d	2.34%	18.9%	25.5%
ε+β⁺, α	ε+β⁺, α	ε, α	ε+β⁺	α	ε+β⁺	stable	ε	stable	ε	stable	stable	stable
Tb-149	Tb-150	Tb-151	Tb-152	Tb-153	Tb-154	Tb-155	Tb-156	Tb-157	Tb-158	Tb-159	Tb-160	Tb-161
4.12 h	3.48 h	17.61 h	17.5 h	2.34 d	m2 :23 h	5.32 d	g :5.35 d	71 γ	180 y	100%	72.3 d	6.88 d
ε+β+, α	ε+β⁺, α	ε+β⁺, α	ε+β⁺, α	ε+β ⁺	ε +β ⁺ , IT	ε	ε+β ⁺	ε	ε+β⁺, β⁻	stable	β ⁻	β ⁻
Gd-148	Gd-149	Gd-150	Gd-151	Gd-152	Gd-153	Gd-154	Gd-155	Gd-156	Gd-157	Gd-158	Gd-159	Gd-160
74.6 y	9.28 d	1.8E6 y	124 d	0.20%	240.4 d	2.18%	14.80%	20.47%	15.65%	24.84%	18.48 h	21.86%
α	ε+β⁺, α	α	ε, α	α >1E14	ε	stable	stable	stable	stable	stable	β ⁻	stable

Figure 1: Simplified presentation of the relevant section of the nuclide chart containing data from Firestone & Eckström [8].



Figure 2: Monitor excitation function for the production of 62 Zn in the bombardment of nat Cu with protons. Open triangles: this work (Stack 1). Open squares: this work (Stack 2). Solid circles: this work (Stack 3 – see text for details).



Figure 3: Excitation function for the production of 149 Tb in the irradiation of 152 Gd with protons. Solid circles: this work (Stack 1). The solid curve shows the results from ALICE/ASH calculations [18] and the dotted curve the corresponding values from the TENDL-2012 library [20, 21] for the (p,4n) reaction on 152 Gd only. The dashed curve also shows the ALICE/ASH calculations for the (p,4n) reaction with inclusion of co-produced contributions from the (p,6n) reaction on 154 Gd and the (p,7n) reaction on 155 Gd (see text). Note that the dashed curve has been lowered slightly relative to the solid curve to make it visible in the region of energy where the two curves overlap.



Figure 4: Excitation function for the production of ¹⁵²Tb in the irradiation of ¹⁵⁵Gd with protons. Solid circles: this work (Stack 2). See also caption to Fig. 3.



Figure 5: Excitation functions for the production of ¹⁵³Dy in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 6: Excitation functions for the production of ¹⁵⁵Dy in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). Open circles: this work (Stack 1). See also caption to Fig. 3.



Figure 7: Excitation functions for the production of ¹⁵⁷Dy in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). Open circles: this work (Stack 1). See also caption to Fig. 3.



Figure 8: Excitation functions for the production of ¹⁵⁹Dy in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). Open diamonds: Hassan *et al.* [22]. See also caption to Fig. 3.



Figure 9: Excitation functions for the production of ¹⁵³Tb in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 10: Excitation functions for the production of 154m2 Tb in the irradiation of 159 Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 11: Excitation functions for the production of ¹⁵⁵Tb in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 12: Excitation functions for the production of ¹⁵⁶Tb in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 13: Excitation functions for the production of 151 Gd in the irradiation of 159 Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 14: Excitation functions for the production of ¹⁵³Gd in the irradiation of ¹⁵⁹Tb with protons. Solid circles: this work (Stack 3). See also caption to Fig. 3.



Figure 15: Integral physical yields for the production of ¹⁴⁹Tb and ¹⁵²Tb via (p,4n) reactions on targets of ¹⁵²Gd and ¹⁵⁵Gd, respectively. A 100% enrichment of the targets is assumed.



Figure 16: Integral physical yields for the production of ^{153,155,157,159}Dy in the proton bombardment of ¹⁵⁹Tb (natural Tb).



Figure 17: Growth and decay of ¹⁵⁵Tb formed in the decay of ¹⁵⁵Dy. A_d and A_m denote daughter and mother activities, respectively (see text).



Figure 18: Production yield, radionuclidic purity as a percentage of total Tb radioactivity, and purity as a percentage of total Tb nuclei produced of ¹⁵⁵Tb in the proton bombardment of a thick Tb target, plotted as a function of incident beam energy. The exit proton energy is 35 MeV. The production conditions are those of Table 7.