Specific Radioactivity of Neutron Induced Radioisotopes: Assessment Methods and Application for Medically Useful $^{177}$Lu Production as a Case

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Abstract: The conventional reaction yield evaluation for radioisotope production is not sufficient to set up the optimal conditions for producing radionuclide products of the desired radiochemical quality. Alternatively, the specific radioactivity (SA) assessment, dealing with the relationship between the affecting factors and the inherent properties of the target and impurities, offers a way to optimally perform the irradiation for production of the best quality radioisotopes for various applications, especially for targeting radiopharmaceutical preparation. Neutron-capture characteristics, target impurity, side nuclear reactions, target burn-up and post-irradiation processing/cooling time are the main parameters affecting the SA of the radioisotope product. These parameters have been incorporated into the format of mathematical equations for the reaction yield and SA assessment. As a method demonstration, the SA assessment of $^{177}$Lu produced based on two different reactions, $^{176}$Lu $(n,\gamma)^{177}$Lu and $^{176}$Yb $(n,\gamma)^{177}$Yb ($\beta^{-}$ decay) $^{177}$Lu, were performed. The irradiation time required for achieving a maximum yield and maximum SA value was evaluated for production based on the $^{176}$Lu $(n,\gamma)^{177}$Lu reaction. The effect of several factors (such as elemental Lu and isotopic impurities) on the $^{177}$Lu SA degradation was evaluated for production based on the $^{176}$Yb $(n,\gamma)^{177}$Yb ($\beta^{-}$ decay) $^{177}$Lu reaction. The method of SA assessment of a mixture of several radioactive sources was developed for the radioisotope produced in a reactor from different targets.
Keywords: specific radioactivity; target burn-up; isotope dilution; neutron capture yield; nuclear reaction; nuclear reactor; radioisotope production; targeting radiopharmaceutical; $^{177}\text{Lu}$, $^{175}\text{Lu}$, $^{176}\text{Lu}$, $^{177}\text{Yb}$, $^{176}\text{Yb}$, $^{175}\text{Yb}$, $^{174}\text{Yb}$

1. Introduction

State-of-the-art radiopharmaceutical development requires radioisotopes of specific radioactivity (SA) as high as possible to overcome the limitation of in vivo uptake of the entity of living cells for the peptide and/or monoclonal antibody based radiopharmaceuticals which are currently used in the molecular PET/CT imaging and endo-radiotherapy. The medical radioisotopes of reasonable short half-life are usually preferred because they have, as a rule of thumb, higher SA. These radioisotopes can be produced from cyclotrons, radionuclide generators and nuclear reactors. The advantage of the last one lies in its large production capacity, comfortable targetry and robustness in operation. This ensures the sustainable supply and production of key, medically useful radioisotopes such as $^{99}\text{Mo}/^{99m}\text{Tc}$ for diagnostic imaging and $^{131}\text{I}$, $^{32}\text{P}$, $^{192}\text{Ir}$ and $^{60}\text{Co}$ for radiotherapy. The high SA requirement for these radioisotopes was not critically considered with respect to their effective utilization in nuclear medicine, except for $^{99}\text{Mo}$. The current wide expansion of targeting endo-radiotherapy depends very much on the availability of high SA radionuclides which can be produced from nuclear research reactor such as $^{153}\text{Sm}$, $^{188}\text{W}/^{188}\text{Re}$, $^{90}\text{Y}$ and $^{177}\text{Lu}$. As an example, as high as 20 Ci per mg SA $^{177}\text{Lu}$ is a prerequisite to formulate radiopharmaceuticals targeting tumors in different cancer treatments [1,2].

So far in radioisotope production, reaction yield has been the main parameter to be concerned with rather than SA assessment and unfortunately, the literature of detailed SA assessment is scarcely to be found [3,4]. The SA assessment of radioisotopes produced in a reactor neutron–activated target is a complex issue. This is due to the influence of the affecting factors such as target burn-up, reaction yield of expected radionuclide and unavoidable side-reactions. All these depend again on the available neutron fluxes and neutron spectrum, which are not always adequately recorded. Besides, the reactor power-on time and target self-shielding effect is usually poorly followed up. Certainly, the SA of target radionuclides has been a major concern for a long time, especially for the production of radioisotopes, such as $^{60}\text{Co}$ and $^{192}\text{Ir}$, used in industry and radiotherapy. In spite of the target burn-up parameter present in the formula of reaction yield calculation to describe the impact of target depression, the SA assessment using the reaction yield was so significantly simplified that the target mass was assumed to be an invariable value during the reactor activation. Critically, this simplification was only favored by virtue of an inherent advantageous combination of the low neutron capture cross section (37 barns) of the target nuclide $^{59}\text{Co}$ and the long half-life of $^{60}\text{Co}$ (which keeps the amount of elemental Co unchanged during neutron bombardment) [4].

The targets used in the production of short-lived medical radioisotopes, however, have high neutron capture cross sections to obtain as high as possible SA values. This fact causes a high “real” burn-up of the target elemental content. Especially, the short half-life of the beta emitting radioisotope produced in the target hastens the chemical element transformation of the target nuclide and strongly affects the
SA of the produced radioisotope. The triple factors influencing the production mentioned above (target, neutron flux and short half-life of produced radionuclide) are also critical with respect to the influence of the nuclear side-reactions and impurities present in the target. Moreover, the SA of a radionuclide produced in nuclear reactor varies with the irradiation and post-irradiation processing time as well. All these issues should be considered for a convincing SA assessment of the producible radioisotope for any state-of-the-art nuclear medicine application. As an example, a theoretical approach to the SA assessment reported together with an up-to-date application for $^{177}$Lu radioisotope production is presented in this paper. This assessment can also play a complementary or even substantial role in the quality management regarding certifying the SA of the product, when it may be experimentally unfeasible due to radiation protection and instrumentation difficulties in the practical measurement of very low elemental content in a small volume solution of high radioactivity content.

High SA nuclides can be produced by $(n, \gamma)$ reaction using high cross section targets such as the $^{176}$Lu $(n, \gamma)^{177}$Lu reaction ($\delta = 2,300$ barns). $^{177}$Lu is a radioisotope of choice for endo-radiotherapy because of its favorable decay characteristics, such as a low energy beta decay of 497 keV (78.6%) and half-life of 6.71 day. It also emits gamma rays of 113 keV (6.4%) and 208 keV (11%) which make it useful for imaging in-vivo localization with a gamma camera.

$^{177}$Lu can be produced by two different routes, a direct route with the $^{176}$Lu $(n, \gamma)^{177}$Lu reaction and an indirect route via the $^{176}$Yb $(n, \gamma)^{177}$Yb ($\beta^-$ decay) $^{177}$Lu nuclear reaction-transformation. The direct route could be successfully performed in high neutron flux nuclear reactors but these are available in only a handful of countries in the world. Additionally, large burn-up of the target nuclide during high neutron flux irradiation may cause a degradation of the SA value of the produced nuclide if the target contains isotopic impurities. No-carrier-added (n.c.a) radioisotopes of higher SA can be produced via an indirect route with a nuclear reaction- followed- by- radioactive transformation process, such as in the process of neutron capture-followed-by- $\beta^-$ decay , $^{176}$Yb $(n, \gamma)^{177}$Yb ($\beta^-$ decay) $^{177}$Lu. In this case, the same reduction in SA is also be experienced if the target contains isotopic and/or elemental Lu impurities.

$^{177}$Lu production has been reported in many publications [5-9], but until now the product quality, especially the evaluation of $^{177}$Lu specific radioactivity in the product, has not been sufficiently analyzed. Based on the theoretical SA assessment results obtained in this report, the optimal conditions for the $^{177}$Lu production were set up to produce $^{177}$Lu product suitable for radiopharmaceutical preparations for targeting endo-radiotherapy.

### 1.1. Units of specific radioactivity, their conversion and SA of carrier-free radionuclide

The specific radioactivity is defined by different ways. In our present paper we apply the percentage of the hot atom numbers of a specified radioactive isotope to the total atom numbers of its chemical element present in the product as the specific radioactivity. This is denoted as atom %.

The following denotation will be used for further discussion. $N_{Ri(A)}$ is the hot atom numbers of radioisotope $R_i$ of the chemical element A and $\lambda_{Ri(A)}$, its decay constant. $N_A$ is the atom numbers of the chemical element A and $T_{1/2}$ (sec) the half-life of radioisotope $R_i$.

The SA unit of atom % is defined as follows:
SA (in unit atom %) = \frac{100 \times \text{Hot atom numbers of a specified radionuclide}}{	ext{Atom numbers of the chemical element of specified radionuclide}}

which can be formulated as follows:

\[
SA \text{ (atom %)} = 100 \cdot \frac{N_{R(A)}}{N_A}
\]

(1)

SA in units Bq/Mol and Bq/g are more currently used in practice. The conversion between the SA units is the following:

\[
SA \text{ (Bq/Mol)} = \frac{SA \text{ (Bq/g)} \cdot M_{\text{at}}}{100 \cdot (6.022 \cdot 10^{23})^{-1} \cdot M_{\text{ai}}}
\]

where \(M_{\text{ai}}\) is the atomic weight of the target or radioactive material of given isotopic composition of the chemical element A.

For a radioactive material containing \(n\) isotopes of the element A:

\[
M_{\text{at}} = \sum P_{n,A} / \sum (P_{n,A} / M_{N,A}),
\]

where \(P_{N,A}\) and \(M_{N,A}\) are the weight percentage and atomic weight of the isotope \(N,A\), respectively.

The specific radioactivity of the carrier-free radioisotope \(R_i\) is calculated as below:

\[
SA_{\text{Carrier-free}} \text{ (Bq/Mol)} = \frac{N_{R(A)} \cdot \lambda_{R(A)}}{N_{R(A)} / 6.022 \cdot 10^{23}} = 6.022 \cdot 10^{23} \cdot \lambda_{R(A)} \cdot \frac{4.1732 \cdot 10^{23}}{T_{1/2}}
\]

(3)

Identifying eq.2 with eq.3 (individualizing \(M_{\text{ai}}\) as the atomic weight of the concerned radioisotope), it is clear that the SA of a carrier-free radionuclide in unit atom % is 100%.

### 2. Theoretical Approach and Assessment Methods

Reactor-based radioisotope preparation usually involves two main nuclear reactions. The first one is the thermal neutron capture \((n, \gamma)\) reaction. This reaction doesn’t lead to a radioisotope of another chemical element, but the following radioactive \(\beta^-\) decay of this isotope during target activation results in a decrease in both the reaction yield and atom numbers of the target chemical element. The second reaction is the thermal neutron capture followed by radioactive transformation \(S (n, \gamma) R_x (\beta^- \text{ decay}) R_i\). This reaction leads to a carrier-free radioisotope of another chemical element than the target chemical element.

The SA assessment in the radioisotope production using the first reaction (with a simple target system) is simple. Careful targetry could avoid the side reaction \(S (n, \gamma) R_x (\beta^- \text{ decay}) R_i\) which could result in the isotopic impurities for the radioisotope intended to be produced using the first reaction. In this case the SA assessment in \((n, \gamma)\) reaction based production process can be simplified by investigation of the SA degrading effect of target nuclide burn-up, chemical element depression due to radioactive decay and isotopic impurities present in the target.

On the other hand the SA assessment in the radioisotope production using the second reaction (with complex target system) is more complicated. The complexity of the targetry used in \(S (n, \gamma) R_x (\beta^- \text{ decay}) R_i\) reaction based isotope production requires an analysis of the combined reaction system. This
system is influenced by both \((n, \gamma)\) reaction and neutron-capture-followed-by-radioactive transformation \(S (n, \gamma) R_x (\beta^- \text{decay}) R_i\). So the effect of side nuclear reactions in this target system will be assessed in addition to the three above mentioned factors that are involved in the simple target system. In this case the SA assessment is best resolved by a method of SA calculation used for the mixture of several radioactive sources of variable SA, which is referred to as a radioisotope dilution process.

For the calculation of SA and reaction yield of the radioisotope \(R_i\) in the two above mentioned reactions, the following reaction schemes are used for further discussion.

Reaction scheme 1:

\[
\begin{align*}
S_{1,A} & \xrightarrow{(n, \gamma) \text{reaction} \neq 1} S_{1,A} \xrightarrow{(n, \gamma) \text{reaction} = 1} R_i \xrightarrow{(n, \text{particle})} \lambda_{1,R_i} \\
\end{align*}
\]

Reaction scheme 2:

\[
\begin{align*}
S_{1,B} & \xrightarrow{(n, \gamma) \text{reaction} \neq 1} S_{1,B} \xrightarrow{(n, \gamma) \text{reaction} = 1} R_x \xrightarrow{(n, \text{particle})} \lambda_{1,R_x} \\
R_y & \xrightarrow{(n, \text{particle})} \lambda_{1,R_y} \\
\end{align*}
\]

Reaction scheme 3:

\[
\begin{align*}
S_{2,B} & \xrightarrow{(n, \gamma) \text{reaction} \neq 1} S_{2,B} \xrightarrow{(n, \gamma) \text{reaction} = 1} R_y \xrightarrow{(n, \text{particle})} \lambda_{1,R_y} \\
\end{align*}
\]

\(S_{1,A}\) is the target stable isotope of element A in the target; \(S_{g,A}\) (with \(g \geq 2\)) is the impure stable isotope of element A originally presented or produced in the target.

\(S_{1,B}\) is the target stable isotope of element B in the target; \(S_{2,B}\) is the stable isotope of element B in the target.

\(R_i, A\) or \(R_i\) is the wanted radioisotope of element A produced in the target from stable isotope \(S_{1,A}\).

\(R_x\) and \(R_y\) are the radioisotopes of element B produced in the target.

The particle emitted from reaction \((n, \text{particle})\) may be proton or alpha.

\(\sigma_{\text{th}}\), \(\sigma_{\text{epi}}\) and \(\sigma_{\text{fast}}\) are reaction cross sections for thermal, epi-thermal and fast neutrons, respectively.

\(\sigma_{1,\text{th}}\), \(\sigma_{2,\text{th}}\), \(\sigma_{2,\text{e}}\), \(\sigma_{2,\text{f}}\) are cross sections of thermal neutrons for the formation of isotopes i, x, y, from stable isotope 1, 2, 2, respectively.

\(\lambda\) is the decay constant.
The (n, γ) reaction yield and the specific radioactivity calculated from it depends on the neutron flux and reaction cross-section which is variable with neutron energy ($E_n$) or velocity ($v_n$). In the thermal neutron region, the cross-section usually varies linearly as $1/v_n$ (so called $1/v_n$ reaction), where $v_n$ is velocity of neutrons. The cross section-versus-velocity function of many nuclides is, however, not linear as $1/v_n$ in the thermal region (so called non−$1/v_n$ reaction.). As the energy of neutrons increases to the epithermal region, the cross section shows a sharp variation with energy, with discrete sharp peaks called resonance.

On other hand, the cross section values of the (n, γ) reactions tabulated in the literature present as $\sigma_0$ given for thermal neutrons of $E_n = 0.0253 \text{ eV}$ and $v_n = 2200 \text{ m/s}$ and as $I_0$ (infinite dilution resonance integral in the neutron energy region from $E_{Cd} = 0.55 \text{ eV}$ to 1.0 MeV) given for epithermal neutrons.

The symbols $\sigma_{th}$ and $\sigma_{epi}$ used in this paper are identified with the thermal neutron activation cross-section $\sigma_0$ and the infinite dilution resonance integral $I_0$, respectively, for the case of $1/v_n$ (n, γ) -reaction carried out with a neutron source of pure $1/E_n$ epithermal neutron spectrum (Epithermal flux distribution parameter $\alpha = 0$). Unfortunately, this condition is not useful any more for practical reaction yield and SA calculations.

In practice the target is irradiated by reactor neutrons of $1/E_n^{1+\alpha}$ epithermal neutron spectrum, so the value of $\Omega_i$ presenting as a sum ($\sigma_{1,(i(k))} + R_{epi,1,(i(epi))}$) in all the equations below has to be replaced by $\sigma_{eff(1/v_n)}$ for the "$1/v_n$"- named (n,γ) reaction and by $\sigma_{eff(non−1/v_n)}$ for the "non−$1/v_n$" - named (n,γ)-reaction. The detailed description of these $\sigma_{eff}$ values can be found in the ‘Notes on Formalism’ at the end of this section.

For the isotope production based on (n,γ) reactions the neutron bombardment is normally carried out in a well-moderated nuclear reactor where the thermal and epithermal neutrons are dominant. The fast neutron flux is insignificant compared to thermal and epithermal flux (e.g. $<10^7 \text{ n.cm}^{-2}.\text{s}^{-1}$ fast neutron flux compared to $>10^{14} \text{ n.cm}^{-2}.\text{s}^{-1}$ thermal one in the Rigs LE7-01 and HF-01 of OPAL reactor-Australia). Besides, the milli-barn cross-section of (n,γ) ,(n,p) and (n,α) reactions induced by fast neutrons is negligible compared to that of (n,γ) reaction with thermal neutron [11]. So the reaction rate of the fast neutron reactions is negligible. Nevertheless, for the generalization purposes the contribution of the fast neutron reaction is also included in the calculation methods below described. It can be ignored in the practical application of SA assessment without significant error.

2.1. The specific radioactivity of radionuclide $R_i$ in the simple target system for the (n, γ) reaction based radioisotope production

2.1.1. Main characteristics of the simple target system

The simple target system contains several isotopes of the same chemical element. Among them only one radioisotope $R_i$ is intended to be produced from stable isotope $S_{1,A}$ via a (n, γ) reaction $i = 1$ as described above in reaction scheme 1. Other stable $S_{g,A}$ isotopes ( with $g \geq 2$) of the target are considered as impure isotopes.
2.1.1.1. The target burn-up for each isotope in simple target system

The burn-up of the isotope $S_{1,A}$ is the sum of the burn-up caused by different $(n, \gamma)$ and $(n, \text{particle})$ reactions from reaction $i = 1$ to $i = k$, the cross sections of which are different $\sigma_{1,i}$ values. This total burn up rate could be formulated as follows:

$$-\frac{dN_{S_{1,A}}}{dt_{irr}} = \phi_{th} \cdot N_{S_{1,A}} \cdot \sum_{i=1}^{k} \sigma_{1,(th)}(i) + \phi_{epi} \cdot N_{S_{1,A}} \cdot \sum_{i=1}^{k} \sigma_{1,(epi)}(i) + \phi_{fast} \cdot N_{S_{1,A}} \cdot \sum_{i=1}^{k} \sigma_{1,(fast)}(i)$$

(4)

$\sigma_{1,(th)}$, $\sigma_{1,(epi)}$ and $\sigma_{1,(fast)}$ are the thermal, epithermal and fast neutron cross section of the $S_{1,i}$ nuclide for the reaction $i$, respectively. $\phi_{th}$, $\phi_{epi}$ and $\phi_{fast}$ are the thermal, epithermal and fast neutron flux, respectively. $t_{irr}$ is the irradiation time. $N_{S_{1,A}}$ is the atom numbers of the isotope $S_{1,A}$. By putting $R_{epi} = \phi_{epi} / \phi_{th}$ and $R_{fast} = \phi_{fast} / \phi_{th}$ ratios into eq.4, the following is deduced.

$$-\frac{dN_{S_{1,A}}}{dt_{irr}} = -\phi_{th} \cdot N_{S_{1,A}} \cdot \sum_{i=1}^{k} (\sigma_{1,(th)}(i) + R_{epi} \cdot \sigma_{1,(epi)}(i) + R_{fast} \cdot \sigma_{1,(fast)}(i))$$

By substituting:

$$\Omega_{1,i} = \sigma_{1,(th)}(i) + R_{epi} \cdot \sigma_{1,(epi)}(i) + R_{fast} \cdot \sigma_{1,(fast)}(i)$$

(5)

and:

$$\Delta_{S_{1,A}} = \phi_{th} \cdot \sum_{i=1}^{k} (\sigma_{1,(th)}(i) + R_{epi} \cdot \sigma_{1,(epi)}(i) + R_{fast} \cdot \sigma_{1,(fast)}(i)) = \phi_{th} \cdot \sum_{i=1}^{k} \Omega_{1,i}$$

(6)

the above differential equation is simplified as follows:

$$\frac{dN_{S_{1,A}}}{dt_{irr}} = -N_{S_{1,A}} \cdot \Delta_{S_{1,A}} \Rightarrow \frac{dN_{S_{1,A}}}{N_{S_{1,A}}} = -\Delta_{S_{1,A}} \cdot dt_{irr}$$

(7)

The un-burned atom numbers of the isotope $S_{1,A}$ at any $t_{irr}$ values ($N_{S_{1,A}}$) is achieved by the integration of eq.7 with the condition of $N_{S_{1,A}} = N_{0,S_{1,A}}$ at $t_{irr} = 0$. The result is:

$$N_{S_{1,A}} = N_{0,S_{1,A}} \cdot e^{-\Delta_{S_{1,A}} \cdot t_{irr}}$$

(8)

From this equation, the burned-up atom numbers of the isotope $S_{1,A}$ ($N_{b,S_{1,A}}$) is:

$$N_{b,S_{1,A}} = N_{0,S_{1,A}} - N_{S_{1,A}} = N_{0,S_{1,A}} \cdot (1 - e^{-\Delta_{S_{1,A}} \cdot t_{irr}})$$

(9)

The same calculation process is performed for any isotope $S_{g,A}$.

**Half-burn-up time of the target nuclide.** At half-burn-up time $T_{1/2-B}$ a half of the original atom numbers of the isotope $S_{1,A}$ are burned. Putting $N_{S_{1,A}} = N_{0,S_{1,A}} / 2$ into eq. 8, the $T_{1/2-B}$ value is achieved as follows:

$$T_{1/2-B} = 0.693 / \Delta_{S_{1,A}}$$

(10)
2.1.1.2. Reaction yield of radioisotope R\textsubscript{i} in the simple target system

By taking into consideration the un-burned atom numbers of the isotope S\textsubscript{1,\textit{A}} (eq. 8), the reaction rate of any isotope in reaction scheme 1 will be evaluated as follows. In this reaction process the depression of the atom numbers of radioisotope R\textsubscript{i} is caused by beta radioactive decays and (n, \gamma)/(n, particle) reaction-related destruction. The depression factor \( \Lambda_{R_i} \) of the radioisotope R\textsubscript{i} in reaction scheme 1 is formulated as follows:

\[
\Lambda_{R_i} = \sum_{m=1}^{m-j} \lambda_{m,R_i} + \Delta_i
\]

where \( \Delta_i = \phi_{th} \cdot \sum \Omega_i \) and \( \Omega_i = \sigma_{(th)} + R_{ep} \cdot \sigma_{(ep)} + R_{fast} \cdot \sigma_{(fast)} \).

Taking into account eq.5, R\textsubscript{i} radioisotope formation rate is the following:

\[
\frac{dN_{R_i}}{dt_{irr}} = \left( \phi_{th} \cdot \sigma_{(th)} + \phi_{th} \cdot R_{ep} \cdot \sigma_{(ep)} + \phi_{th} \cdot R_{fast} \cdot \sigma_{(fast)} \right) \cdot N_{0,S_{1,A}} \cdot e^{-\Delta_{S_{1,A}}t_{irr}} - N_{R_i} \cdot \Lambda_{R_i} \rightarrow
\]

\[
\frac{dN_{R_i}}{dt_{irr}} + N_{R_i} \cdot \Lambda_{R_i} = \phi_{th} \cdot \Omega_{1,i} \cdot N_{0,S_{1,A}} \cdot e^{-\Delta_{S_{1,A}}t_{irr}}
\]

(12)

By multiplying both sides of this equation with \( e^{\Lambda_{R_i}t_{irr}} \) and manipulating with the mathematical tool \( \frac{d(YX)}{dt} = X \cdot \frac{dY}{dt} + Y \cdot \frac{dX}{dt} \), this equation is converted into the following form:

\[
d \left( N_{R_i} \cdot e^{\Lambda_{R_i}t_{irr}} \right) = N_{0,S_{1,A}} \cdot \phi_{th} \cdot \Omega_{1,i} \cdot e^{(\Lambda_{R_i} - \Delta_{S_{1,A}})t_{irr}} \cdot d t_{irr}
\]

By integrating this equation and assuming \( N_{R_i} = 0 \) at \( t_{irr} = 0 \), the yield of radioisotope \( R_i \) at the irradiation time \( t_{irr} \) is the following:

The \( R_i \) atom numbers (N\textsubscript{Ri}):

\[
N_{R_i} = \frac{N_{0,S_{1,A}} \cdot \phi_{th} \cdot \Omega_{1,i}}{\Lambda_{R_i} - \Delta_{S_{1,A}}} \cdot (e^{-\Delta_{S_{1,A}}t_{irr}} - e^{-\Delta_{R_i}t_{irr}})
\]

(13)

The \( R_i \) isotope radioactivity (A\textsubscript{Ri}):

\[
A_{R_i} = \frac{N_{0,S_{1,A}} \cdot \phi_{th} \cdot \Omega_{1,i} \cdot \sum_{m=1}^{m-j} \lambda_{m,R_i}}{\Lambda_{R_i} - \Delta_{S_{1,A}}} \cdot (e^{-\Delta_{S_{1,A}}t_{irr}} - e^{-\Delta_{R_i}t_{irr}})
\]

(14)

These equations can be deduced from the well known Bateman equation [3,12]. The \( R_i \) atom numbers and radioactivity at the post-irradiation time \( t_c \) (N\textsubscript{R,i,tc} and A\textsubscript{R,i,tc}, respectively) are calculated by multiplying eqs.13 and 14 with the factor \( e^{-\Delta_{S_{1,A}}t_c} \).

Maximum yield of radioisotope \( R_i \). At the irradiation time (denoted as \( t_{irr-max} \)) where \( \frac{dA_{R_i}}{dt_{irr}} = 0 \), \( R_i \) radioactivity reaches maximum (\( A_{R_i-max} \)). By differentiating eq.14 and making it equal to zero:

\[
\frac{dA_{R_i}}{dt_{irr}} = \frac{N_{0,S_{1,A}} \cdot \phi_{th} \cdot \Omega_{1,i} \cdot \sum_{m=1}^{m-j} \lambda_{m,R_i} \cdot (e^{-\Delta_{S_{1,A}}t_{irr-max}} - e^{-\Delta_{R_i}t_{irr-max}} + \Lambda_{R_i} \cdot e^{-\Delta_{R_i}t_{irr-max}})}{\Lambda_{R_i} - \Delta_{S_{1,A}}} = 0
\]
the \( t_{\text{irr-max}} \) is deduced as follows:

\[
    t_{\text{irr-max}} = (\ln \frac{\Lambda_{R_i}}{\Delta_{S_{i,a}}}) / (\Lambda_{R_i} - \Delta_{S_{i,a}})
\]  

Equation (15) is useful for irradiation optimization to produce \( R_i \) radioisotope of highest yield. By introducing the value \( t_{\text{irr-max}} \) into eqs.13 and 14, we achieve the maximum yield of radioisotope \( R_i \) (\( N_{R_i-\text{max}} \) and \( A_{R_i-\text{max}} \)) as follows:

The maximum atom numbers \( N_{R_i-\text{max}} \) is:

\[
    N_{R_i-\text{max}} = \frac{N_{o,S_{i,a}} \cdot \phi_i \cdot \Omega_{1,j} \cdot (e^{-p} - e^{-h})}{A_{R_i} - \Delta_{S_{i,a}}} \]  

(16)

The maximum radioactivity \( A_{R_i-\text{max}} \) is:

\[
    A_{R_i-\text{max}} = \frac{N_{o,S_{i,a}} \cdot \phi_i \cdot \Omega_{1,j} \cdot f \cdot (e^{-p} - e^{-h})}{q} \]  

(17)

where \( D = \frac{\Lambda_{R_i}}{\Delta_{S_{i,a}}}; f = \left( \sum_{m=1}^{n} \frac{\lambda_{m,R_i}}{\Lambda_{R_i}} \right) / \Lambda_{R_i}; p = \ln \left( \frac{D}{(D - 1)} \right); h = D \cdot \ln \left( \frac{D}{(D - 1)} \right); q = (1 - D^{-1}) \)

As shown the maximum yield of radioisotope \( R_i \) is a function of the variable \( D \).

2.1.2. The SA assessment of radionuclide \( R_i \) in the simple target for \((n, \gamma)\) reaction based radioisotope production

2.1.2.1. General formula of SA calculation for the simple multi-isotope target

The simplification in the calculation is based on the fact that the target isotope \( S_{i,A} \) captures neutrons to form the wanted radioisotope \( R_i \) and the isotopic impurities in the target don’t get involved in any nuclear reactions whatsoever. The isotopic impurities may participate in some nuclear reactions to generate either stable isotopes of the target element or an insignificant amount of the isotopes of other chemical element than the target one. This simplified calculation process is supported by a careful targetry study regarding minimizing the radioactive isotopic impurities in the radioisotope product. The following is the SA of radioisotope \( R_i \) formed in a target composed of different stable isotopes:

\[
    \text{SA}_{R_i} = 100 \cdot N_{R_i} / (N_{R_i} + \sum_{i=1}^{g} N_{s_{i,a}}) \]  

(18)

\( \sum_{i=1}^{g} N_{s_{i,a}} \) is the sum of the remaining (unburned) atom numbers of \( g \) different stable isotopes of the same chemical element in the target. By placing the values \( N_{s_{i,a}} \) of different stable isotopes of the target from eq.8 into this equation, the following general formula is obtained for the SA of radioisotope \( R_i \):

\[
    \text{SA}_{R_i} = 100 \cdot N_{R_i} / \left( N_{R_i} + N_{o,s_{i,a}} \cdot e^{-\lambda_{s_{i,a}} t_{irr}} + N_{0,s_{i,a}} \cdot e^{-\lambda_{s_{i,a}} t_{irr}} + \ldots + N_{o,s_{i,a}} \cdot e^{-\lambda_{s_{i,a}} t_{irr}} \right) \]  

(19)

where \( N_{0,s_{i,a}} = 6.02 \cdot 10^{23} \cdot m \cdot P_{g} / (100 \cdot M_{g}) \); \( N_{o,s_{i,a}} = 6.02 \cdot 10^{23} \cdot m \cdot P_{i} / (100 \cdot M_{i}) \)

If the target contains impure isotopes of another chemical element, more stable isotopes of chemical element \( A \) generated via reaction scheme 3 above could be present in the denominator of this
formula. This amount may cause additional depression of \( S_{AS, irradiation} \). This small impurity will, however, bring about an insignificant amount of stable isotope \( S_{AS, stable} \) and its depression effect will be ignored. The eq. 19 is set up with an ignorance of insignificant amount of not-really-burned impure stable isotope which captures neutron, but not yet transformed into the isotope of other chemical element via a radioactive decay).

If the impure isotope \( S_{AS, stable} \) doesn’t participate in any nuclear reaction or its neutron capture generates a stable isotope of the target element, then zero value will be given to the parameter \( \Delta_{AS, irradiation} \) of eq.(19).

2.1.2.2. SA of radioisotope \( R_i \) in the simple two-isotope target

From the practical point of view, the target composed of two stable isotopes is among the widely used ones for radioisotope production. For this case the SA calculation is performed as follows:

\[
SA_{R_i, t_{irr}} = 100 \cdot \frac{N_{R_i} + N_{0, S_{1,i}} \cdot e^{-\Delta_{S_{1,i}, irradiation}} + N_{0, S_{2,i}} \cdot e^{-\Delta_{S_{2,i}, irradiation}}}{(N_{R_i} + N_{0, S_{1,i}}) \cdot e^{-\Delta_{S_{1,i}, irradiation}} + (N_{R_i} + N_{0, S_{2,i}}) \cdot e^{-\Delta_{S_{2,i}, irradiation}}}
\]

where \( N_{0, S_{1,i}} = 6.02 \cdot 10^{23} \cdot m \cdot P_1 / (100 \cdot M_1) \); \( N_{0, S_{2,i}} = 6.02 \cdot 10^{23} \cdot m \cdot P_2 / (100 \cdot M_2) \), \( R_i \) is the radioisotope expected to be produced from the stable isotope \( S_{1,i} \). \( P_1 \) and \( M_1 \) are for the isotope \( S_{1,i} \), \( m \) is the weight of the target.

By replacing \( \Delta_{AS, irradiation} \), \( \Delta_{AS, stable} \), \( \Delta_{AS, target} \) and the \( N_{R_i} \) value from eq.(13) into eq.(20), SA of radioisotope \( R_i \) in a two isotope target at the end of neutron bombardment, \( SA_{R_i, t_{irr}} \), is:

\[
SA_{R_i, t_{irr}} = \frac{100 \cdot M_2 \cdot P_1 \cdot \phi_{\beta} \cdot \Omega_{\beta} \cdot (e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}})}{M_2 \cdot \phi_{\beta} \cdot \Omega_{\beta} \cdot (e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}) + (\Lambda_{AS, irradiation} - \Delta_{AS, stable}) \cdot (M_2 \cdot P_1 \cdot e^{-\Delta_{S_{1,i}, irradiation}} + M_1 \cdot P_2 \cdot e^{-\Delta_{S_{2,i}, irradiation}})}
\]

\[
SA_{R_i, t_{irr}} = \frac{100 \cdot e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}}{a \cdot e^{-\Delta_{S_{1,i}, irradiation}} + b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}}
\]

where \( a = \frac{(\Lambda_{AS, irradiation} - \Delta_{AS, stable})}{\phi_{\beta} \cdot \Omega_{\beta}} \), \( b = \frac{M_1 \cdot (\Lambda_{AS, irradiation} - \Delta_{AS, stable})}{M_2 \cdot \phi_{\beta} \cdot \Omega_{\beta}} \)

SA at the post-bombardment time \( t_c \), \( SA_{R_i, t_{circ}} \), is:

\[
SA_{R_i, t_{circ}} = \frac{100 \cdot e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}}{(e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}) \cdot e^{-\sum_k^\infty \Delta_{S_{k,i}, irradiation}} + (\Lambda_{AS, irradiation} - \Delta_{AS, stable}) / \phi_{\beta} \cdot \Omega_{\beta} \cdot e^{-\Delta_{S_{1,i}, irradiation}} + b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}}}
\]

Maximum SA of radioisotope \( R_i \) in the simple two-isotope target. Rendering the differential of eq. 21 equal to zero offers the way to calculate the irradiation time at which the SA of nuclide \( R_i \) reaches maximum value ( \( SA_{R_i, max} \)):

\[
\frac{d(SA_{R_i, t_{irr}})}{dt_{irr}} = \frac{d(100 \cdot e^{-\Delta_{S_{1,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}})}{a \cdot e^{-\Delta_{S_{1,i}, irradiation}} + b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}} - e^{-\Delta_{S_{2,i}, irradiation}}) / dt_{irr} = 0
\]

The irradiation time where the SA reaches maximum is denoted as \( t_{irr, max} \). The equation for the calculation of the \( t_{irr, max} \) value, which is derived from the above differential equation, is the following:

\[
(\Delta_{S_{1,i}, t_{irr, max}} - \Lambda_{AS, irradiation} - a \cdot e^{-\Delta_{S_{1,i}, irradiation}} - b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}})
\]

\[
(\Delta_{S_{2,i}, t_{irr, max}} - \Lambda_{AS, irradiation} - a \cdot e^{-\Delta_{S_{1,i}, irradiation}} - b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}})
\]

\[
(\Delta_{S_{1,i}, t_{irr, max}} - \Lambda_{AS, irradiation} - a \cdot e^{-\Delta_{S_{1,i}, irradiation}} - b \cdot (P_2 / P_1) \cdot e^{-\Delta_{S_{2,i}, irradiation}} = 0)
\]
The solution of this equation performed by the computer software MAPLE-10 gives the value \( t_{irr,S_{\text{max}}} \). The analysis of the equation \( 24 \) and MAPLE-10 calculation result confirmed the SA of nuclide \( R_i \) reaches maximum at a defined characteristic irradiation time \( t_{irr,S_{\text{max}}} \) except for the case of \( P_2=0 \) or very large \( \Delta S_{2,t} \) value, which will be investigated in the following sections.

**SA of radioisotope \( R_i \) in the simple two-isotope target at the maximum reaction yield.** Replacing \( t_{irr} \) of eqs.\((21)\) and \( (22) \) with the \( t_{irr-max} \) from eq.\( 15 \) is to calculate SA at the maximum reaction yield \( S_{\text{A}_{\text{irr}}}(\text{achieved at the irradiation time } t_{irr-max}) \):

\[
S_{\text{A}_{\text{irr}}}(\text{irr}) = 100 \cdot \frac{(e^{-p} - e^{-h})}{(a \cdot e^{-p} + b \cdot (P_2 / P_1) - e^{-h})} \quad (25)
\]

where \( a = \frac{(\phi_0 \cdot \Omega_{j} + \Lambda_{R} - \Delta_{S_{\text{2,t}}} \cdot \Lambda_{R})}{\phi_0 \cdot \Omega_{j}} \); \( b = \frac{M_2 \cdot (\Lambda_{R} - \Delta_{S_{\text{2,t}}} \cdot \Lambda_{R})}{M_2 \cdot \phi_0 \cdot \Omega_{j}} \); \( p = \frac{\ln D}{(D-1)} \); \( h = D \cdot \frac{\ln D}{(D-1)} \).

These parameters are identical to that of the eq.(17) and (21).

**SA of radioisotope \( R_i \) in the target which is considered as a simple two-isotope target.** It is also a matter of fact that another very commonly used target system contains more than two stable isotopes (simple multi-isotope target system, \( g \geq 2 \)). Except \( S_{1,A} \) as shown in reaction scheme 1, all the impure isotopes of the same chemical element in the target don’t get involved in any nuclear reactions or they may participate in with very low rate giving insignificant burn-up \( (\Delta S_{g,A} = 0) \). This system is considered as a special two-isotope target system for which the non-depression of impure isotopes \( (\Delta S_{g,A} = 0) \) and the combined impure isotope percentage \( (P_{\text{imp},A}) \) and molecular weight \( (M_{\text{imp},A}) \) are applied. We will have the relevant equations for the calculation of the specific SA value of this target by putting \( \Delta S_{2,t} = \Delta S_{g,A} = 0 \), \( M_2 = M_{\text{imp},A} = \left(\sum_{g} P_{S_{g,A}} / \sum_{g} P_{S_{g,A}} \right) / M_{S_{g,A}} \) and \( p_2 = P_{\text{imp},A} = \sum_{g} P_{S_{g,A}} \) into eqs.\((21)-(25)\) \( (P_{S_{g,A}} \text{ and } M_{S_{g,A}} \text{ are the weight percentage and atomic weight of impure stable isotopes } S_{g,A}, \text{ respectively}). \)

2.1.2.3. SA of radioisotope \( R_i \) in the simple one-isotope target system

By introducing \( P_2 = 0 \) into eq.\((21)\), the SA of radioisotope \( R_i \) in the simple one-isotope target is the following:

\[
S_{\text{A}_{\text{irr}}}(\text{irr}) = 100 \cdot \frac{(1 - e^{-(\Lambda_{R} - \Delta_{S_{\text{2,t}}}) \cdot t_{irr}})}{(a \cdot e^{-p} - e^{-(\Lambda_{R} - \Delta_{S_{\text{2,t}}}) \cdot t_{irr}})} \quad (26)
\]

This equation doesn’t give the maximum value of \( S_{\text{A}_{\text{irr}}}(\text{irr}) \). The result is confirmed by a calculation with MAPLE 10 software. A double check is made by putting the differential of eq.\((26)\) equal to zero to investigate whether a maximum SA could be found:

\[
\frac{dS_{\text{A}_{\text{irr}}}(\text{irr})}{dt_{irr}} = \frac{d\left(100 \cdot (1 - e^{-(\Lambda_{R} - \Delta_{S_{\text{2,t}}}) \cdot t_{irr}})\right)}{dt_{irr}} = 0 \Rightarrow (\Delta S_{S_{\text{2,t}}} - \Lambda_{R}) \cdot (1 - a) \cdot e^{-(\Lambda_{R} - \Delta_{S_{\text{2,t}}}) \cdot t_{irr}} = 0 \quad (27)
\]

It is shown that the achieved differential eq.\((27)\) has no solution with the variable \( t_{irr} \) and gives a correct solution when \( t_{irr} \) value approaches to infinity. When \( (\Delta S_{S_{\text{2,t}}} - \Lambda_{R}) = 0 \) then a = 1 (as shown in eq.\((25)\)), hence the differential value is not defined, so the specific radioactivity has no maximum value.
at any time. This means that the SA of nuclide R_i in the stable isotope target of 100% isotopic purity never reaches maximum at any irradiation time.

It is also worth mentioning that when the value of $\Delta_{S_{g,A}}$ or $\Delta_{S_{g,A}}$ is very large, eq. (21) is converted to eq. (26). It means that the high burn-up of impure stable isotope S_{g,A} makes a multi-isotope target system change to a one-isotope target one. So, no maximum SA will be expected with this type of multi-isotope target system too.

As shown in eq. (26) the SA of these target systems increases with t_{irr}. This fact teaches us that a compromise between maximum yield achievable at t_{irr,max} and favorable higher SA at the time t_{irr}>t_{irr,max} is subject to the priority of the producer.

2.2. The specific radioactivity of radionuclide R_i in a complex target system for the S(n, $\gamma$) R_x ($\beta^-$ decay) R_i reaction based radioisotope production

2.2.1. Main characteristics of the complex target system

The complex target system contains several isotopes of different chemical elements. Among them only one radioisotope R_i is intended to be produced from stable isotope S_{1,B} of chemical element B via a S_{1,B} (n, $\gamma$) R_x ($\beta^-$ decay) R_i reaction i = 1 as described above in reaction scheme 2. Other stable S_{g,B} isotopes (with g $\geq$ 2) of the element B are considered as impure isotopes and they could be transformed into other isotopes (except R_i) of the chemical element A as described above in reaction scheme 3. Besides, the target could contain different isotopes of the element A as impure isotopes which could be involved in different nuclear reactions during target irradiation.

2.2.1.1. The yield of S_{1B}(n, $\gamma$) R_x ($\beta^-$ decay) R_i reaction

This reaction generates a carrier-free radioisotope R_i. The SAR_i value is 100 atom %. As shown in reaction scheme 2, the atom numbers (N_{R_i}) and the radioactivity (A_{R_i}) of R_i radioisotope of chemical element A are calculated based on the general Bateman equation[3,12]. This is detailed in the following equation:

$$N_{R_i,t_c} = N_{R_i,t_c}^{N_{R_i}^{th}} f_2 f_3 \left[ e^{-d_1 t_c} \left( \frac{e^{-d_1 t_c}}{d_1} \right) + e^{-d_2 t_c} \left( \frac{e^{-d_2 t_c}}{d_2} \right) + e^{-d_3 t_c} \right]$$

$$\Omega_{x} = \sigma_{x(th)} + R_{x,\text{epi}} \cdot \sigma_{x,\text{epi}} + R_{x,\text{fast}} \cdot \sigma_{x,\text{fast}}$$

$$\Omega_{i} = \sigma_{i(th)} + R_{i,\text{epi}} \cdot \sigma_{i,\text{epi}} + R_{i,\text{fast}} \cdot \sigma_{i,\text{fast}}$$

$$d_1 = \phi_{th} \cdot \sum_{x=1}^{g} \Omega_{x}$$

$$d_2 = \sum_{x=1}^{g} \lambda_{x,R_i} + \phi_{th} \cdot \sum_{x=1}^{g} \Omega_{x}$$

$$d_3 = \sum_{x=1}^{g} \lambda_{x,R_i} + \phi_{th} \cdot \sum_{x=1}^{g} \Omega_{x}$$

$$f_2 = \phi_{th} \cdot \Omega_{1,th}$$

$$f_3 = \lambda_{1,R_i}$$

The R_i atom numbers N_{R_i,t_c} present at post-irradiation time t_c is achieved by multiplying the

$$N_{R_i,t_c}$$

value with the decay factor $e^{-\sum_{x=1}^{g} \lambda_{x,R_i} t_c}$.

The AR_i value is simply derived by multiplying the N_{R_i} values with $\sum_{x=1}^{g} \lambda_{x,R_i}$.
2.2.1.2. SA-degradation effect of impure stable isotope generated from S_{2,B}(n, \gamma) R_y(\beta^- decay) S_{g,A} reaction

Referred to reaction scheme 3 involving the impure stable isotope S_{2,B} in the S_{1,B} target, reaction S_{2,B}(n, \gamma) R_y(\beta^- decay) S_{g,A} generates an amount of stable isotope S_{g,A} of the same chemical element to the wanted radionuclide R_{i,A}. This fact makes the SA of radionuclide R_{i,A} produced from stable isotope S_{1,B} lower, so the atom numbers of the stable isotope S_{g,A} should be evaluated for the purpose of SA assessment. The atom numbers of S_{g,A} is determined based on the activity of radioisotope R_y. Identifying eqs. (13) and (14) described for reaction scheme 1 with the process of reaction scheme 3, we get the following equations.

The atom numbers \( N_{R_y,t_{irr}} \) and the radioactivity \( A_{R_y,t_{irr}} \) of radionuclide R_y at irradiation time t_{irr} are calculated in the same manner as in Section 2.1.1.2 above (using eqs. (13) and (14)):

\[
N_{R_y,t_{irr}} = \frac{N_{o,S_{2,B}} \cdot \phi_m \cdot \Omega_{2,y,t_{irr}}}{\Lambda_{R_y} - \Delta_{S_{2,B}}} \cdot (e^{-\Delta_{S_{2,B}} \cdot t_{irr}} - e^{-\Lambda_{R_y} \cdot t_{irr}})
\]  

\[
A_{R_y,t_{irr}} = \frac{N_{o,S_{2,B}} \cdot \phi_m \cdot \Omega_{2,y,t_{irr}}}{\Lambda_{R_y} - \Delta_{S_{2,B}}} \cdot \sum_{m=1}^{m=n} \lambda_{m,R_y} \cdot (e^{-\Delta_{S_{2,B}} \cdot t_{irr}} - e^{-\Lambda_{R_y} \cdot t_{irr}})
\]  

\( \Delta_{S_{2,B}} \) is for stable isotope S_{2,B} and \( \Delta_{S_{1,A}} = \phi_{th} \cdot \sum_{y=1}^{y=8} \Omega_{2,y} \)

where \( \Omega_{2,y} = \sigma_{2,y(th) + R_{eq} \cdot \sigma_{2,y(eq)} + R_{fast} \cdot \sigma_{2,y(fast)} \)

\( \Lambda_{R_y} \) for radionuclide R_y and \( \Lambda_{R_y} = \sum_{m=1}^{m=n} \lambda_{m,R_y} + \Delta_{R_y} \), where \( \Delta_{R_y} = \phi_{th} \cdot \sum \Omega_{R_y} \)

The partial radioactivity of radionuclide R_y for the formation of S_{g,A} isotope is denoted as \( A_{R_y \rightarrow S_{g,A},t_{irr}} \). This quantity is calculated by either multiplying eq. (30) with a branch decay ratio \( f_{S_{g,A}} \) or using an individual decay constant \( \lambda_{R_y \rightarrow S_{g,A}} \) as follows:

\[
A_{R_y \rightarrow S_{g,A},t_{irr}} = \frac{N_{o,S_{2,B}} \cdot \phi_m \cdot \Omega_{2,y} \cdot \lambda_{R_y \rightarrow S_{g,A}}}{\Lambda_{R_y} - \Delta_{S_{2,B}}} \cdot (e^{-\Delta_{S_{2,B}} \cdot t_{irr}} - e^{-\Lambda_{R_y} \cdot t_{irr}})
\]

The S_{g,A} content \( N_{S_{g,A},t_{irr}} \) formed during neutron activation of the impure stable isotope S_{2,B} is calculated by integrating R_y nuclide radioactivity for the neutron irradiation time t_{irr} as below.

\[
N_{S_{g,A},t_{irr}} = \int_0^{t_{irr}} A_{R_y \rightarrow S_{g,A},t_{irr}} \cdot dt_{irr} = \int_0^{t_{irr}} \frac{N_{o,S_{2,B}} \cdot \phi_m \cdot \Omega_{2,y} \cdot \lambda_{R_y \rightarrow S_{g,A}}}{\Lambda_{R_y} - \Delta_{S_{2,B}}} \cdot (e^{-\Delta_{S_{2,B}} \cdot t_{irr}} - e^{-\Lambda_{R_y} \cdot t_{irr}}) \cdot dt_{irr} =
\]

At t_{irr} = 0, \( N_{S_{g,A},0} = 0 \), then \( C = \frac{\Lambda_{R_y} - \Delta_{S_{2,B}}}{\Delta_{S_{2,B}} \cdot \Lambda_{R_y}} \). Putting C value into the above equation we get:

\[
N_{S_{g,A},t_{irr}} = \frac{N_{o,S_{2,B}} \cdot \phi_m \cdot \Omega_{2,y} \cdot \lambda_{R_y \rightarrow S_{g,A}}}{(\Lambda_{R_y} - \Delta_{S_{2,B}}) \cdot \Lambda_{R_y} \cdot \Delta_{S_{2,B}}} \cdot \left( \frac{e^{-\Lambda_{R_y} \cdot t_{irr}} - 1}{e^{-\Lambda_{R_y} \cdot t_{irr}}} + C \right)
\]

At post-irradiation time (t_i) R_y radioactivity decreases as below:
\[ A_{R_y \rightarrow S_{g,A},t_c} = A_{R_y \rightarrow S_{g,A},t_{irr}} \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}}. \] (33a)

The radioactivity \( A_{R_y \rightarrow S_{g,A},t_c} \) is equal to the formation rate of \( S_{g,A} \) during decay time. By integrating this formation rate we get the \( S_{g,A} \) atom numbers \( N_{S_{g,A},t_c} \) collected at the time \( t_c \). Because the \( A_{R_y \rightarrow S_{g,A},t_{irr}} \) of nuclide \( R_y \) at the end-of-neutron-bombardment (E.O.B) time \( t_{irr} \) is independent on the variable \( t_c \), we get the following integral:

\[
N_{S_{g,A},t_c} = \int_0^{t_c} A_{R_y \rightarrow S_{g,A},t_{irr}} \cdot dt_c = \int_0^{t_c} A_{R_y \rightarrow S_{g,A},t_{irr}} \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} \cdot dt_c = \frac{A_{R_y \rightarrow S_{g,A},t_{irr}} \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}}}{-\lambda_{R_y \rightarrow S_{g,A}}} + C
\]

At \( t_c = 0 \), \( N_{S_{g,A},t_c} = N_{S_{g,A},t_{irr}} \), then \( C = N_{S_{g,A},t_{irr}} + \frac{A_{R_y \rightarrow S_{g,A},t_{irr}}}{\lambda_{R_y \rightarrow S_{g,A}}}. \)

Putting the \( C \) value into the above equation we have:

\[
N_{S_{g,A},t_c} = N_{S_{g,A},t_{irr}} + \frac{A_{R_y \rightarrow S_{g,A},t_{irr}}}{\lambda_{R_y \rightarrow S_{g,A}}} \cdot (1 - e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}})
\]

Further putting the \( A_{R_y \rightarrow S_{g,A},t_{irr}} \) value of eq. (31) and \( N_{S_{g,A},t_{irr}} \) of eq. (32) into this equation, we get the result below:

\[
N_{S_{g,A},t_c} = \frac{N_{o,s_{2,B}} \cdot \phi_{iB} \cdot \Omega_{2,F}}{(\lambda_{R_y \rightarrow S_{g,A}} - \Delta_{S_{2,B}}) \cdot \lambda_{R_y} \cdot \Delta_{S_{2,B}}} \cdot \left\{ (\lambda_{R_y \rightarrow S_{g,A}} \cdot \lambda_{R_y} - \lambda_{R_y \rightarrow S_{g,A}} \cdot \Delta_{S_{2,B}}) + (\lambda_{R_y \rightarrow S_{g,A}} \cdot \Delta_{S_{1,B}} - \lambda_{R_y} \cdot \Delta_{S_{1,B}}) \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} - \left( \lambda_{R_y \rightarrow S_{g,A}} \cdot \lambda_{R_y} - \lambda_{R_y} \cdot \Delta_{S_{2,B}} \right) \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} - \left( \lambda_{R_y \rightarrow S_{g,A}} \cdot \Delta_{S_{1,B}} - \lambda_{R_y} \cdot \Delta_{S_{1,B}} \right) \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} - \lambda_{R_y} \cdot \Delta_{S_{2,B}} \cdot (e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} - e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}}) \cdot e^{-\lambda_{R_y \rightarrow S_{g,A},t_c}} \right\} \] (33b)

As shown in this equation, the \( S_{g,A} \) content \( N_{S_{g,A},t_c} \) formed in the target from the \( S_{2,B} \) impure stable isotope is composed of a partial amount formed during neutron activation \( N_{S_{g,A},t_{irr}} \) of isotope \( S_{g,A} \) and its another part formed during post-irradiation decay of \( R_y \) induced in the target.

The SA of \( R_i \) in the above mentioned target containing both \( S_{1,B} \) and \( S_{2,B} \) (as described in reaction schemes 2 and 3) is calculated using eqs. (28) and (33b) as follows:

\[
SA_{R_i,t_c} = 100 \cdot N_{R_i,t_{irr}} \cdot e^{-\sum_{m=1}^{n_{m}} \lambda_{m,R_i} \cdot t_c} / (N_{R_i,t_{irr}} \cdot e^{-\sum_{m=1}^{n_{m}} \lambda_{m,R_i} \cdot t_c} + N_{S_{g,A},t_{irr}})
\] (34)

As a result of the analysis of the above equations, minimizing post-irradiation cooling/processing time is recommended to reduce the SA-degradation effect of the impure isotopes.

2.2.1.3. SA-degradation effect of impure isotopes of the chemical element A

The assessment of SA in system containing these impure isotopes can be found in the Section 2.1 for the simple target system.

2.2.2. The SA assessment of radionuclide \( R_i \) in a complex target system

The radioisotope dilution is involved in SA depression in a complex target system in which both the wanted radioisotope \( R_i \) and its unfavorable stable isotope are generated from different nuclear
reactions of both the target isotope and impurities. The complex target system is considered as a mixture of several radioactive sources of variable SA. The method of SA assessment for this mixture is formulated as below.

$SA_{j,Ri}$ is the SA of $R_i$ in the radioactive source $S_j$ the $R_i$ radioactivity of which is $A_{j,Ri}$. The radioactive source $S_j$ is produced in the target from a given nuclear reaction such as $S(n, \gamma)R_x (\beta^-$ decay) $R_i$ reaction (reaction scheme 2) or $(n, \gamma)$ reaction (reaction scheme 1). There are $n$ different radioactive sources $S_j (j=1\ldots n)$ in the target. So the target is a mixture of radioactive sources. The SA of this radioactive source mixture ($SA_{Mix,Ri}$) is calculated as follows:

$$SA_{Mix,Ri} = \frac{(A_{1,Ri} + A_{2,Ri} + \ldots + A_{j,Ri} + \ldots + A_{n,Ri})}{SA_{1,Ri} + SA_{2,Ri} + \ldots + SA_{j,Ri} + \ldots + SA_{n,Ri}}$$

where $SA_{j,Ri}$ is in unit of atom %. $A_{j,Ri}$ is either the hot $R_i$ atom numbers or $R_i$ radioactivity of the relevant radioactive source $j$.

This equation is valuable for all values of SA, except $SA_{j,Ri} = 0$. This situation excludes the unfavorable effect of some $(n, \gamma)$ reaction which generates a stable brother isotope $S_{g,A}$ of radioisotope $R_i$ in the target system (Reaction scheme 3). To solve this problem we have to combine the atom numbers of this stable brother isotope with the atom numbers of one specified radioactive source of the mixture to generate a new radioactive source of $SA \neq 0$, e.g. the combination of radioactive sources produced from the reactions in the scheme 2 and 3. This treatment will be detailed in a practical application for the $^{176}Yb$ target system in the following section.

2.3. Notes on formalism

During neutron bombardment of the target in the nuclear reactor of $1/E_n^{1.+a}$ epithermal neutron spectrum, the rate of $(n,\gamma)$ reactions is calculated based on either Westcott or Hogdahl formalism [13] depending on the excitation function of the target nuclide (the dependence of the reaction cross section on the neutron energy).

For the "non $-1/v_n"$ - named $(n,\gamma)$ reactions the modified Westcott formalism can be used to improve the accuracy of reaction yield calculation and the reaction rate in both thermal and epithermal neutron region for a diluted sample (both the thermal and epithermal neutron self-shielding factors are set equal to 1 or very close to unity) is:

$$r = \phi_{Westcott} \sigma_{Westcott} = n_n v_0 \sigma_0 \{g(T_n) + r s_0(\alpha)\} = \phi_{Westcott} k \sigma_0$$

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Westcott did not define a thermal neutron flux, but only a conventional (total) neutron flux was mentioned instead. It is the ascertainment from Westcott and Hogdahl formalism that:

\[ \phi_{th, Hogdahl} = \phi_{West, cot} \quad \text{and} \quad \phi_{epi, Hogdahl} \geq \phi_{epi, West, cot} \]

Practically, the values \( \phi_{th, Hogdahl} \) and \( \phi_{epi, Hogdahl} \) (usually known as thermal neutron flux \( \phi_{th} \) and epithermal neutron flux \( \phi_{epi} \), respectively) are determined by a specified nuclide (so called monitor) reaction based measurement of neutron flux in the spectral area below and above the cadmium cut-off energy \( (E_{Cd} = 0.55 \text{ eV}) \), respectively, while the Westcott neutron flux values \( \phi_{West, cot, th} \) and \( \phi_{West, epi, cot} \) are not usually available.

For the evaluation of the practical value of \( \phi_{th, Hogdahl} \) versus \( \phi_{West, cot} \), the ratio \( \left( \frac{\phi_{West, cot}}{\phi_{th, Hogdahl}} \right) \) was calculated using the extreme values of epithermal flux distribution parameter \( \alpha \), the neutron energy \( E_0 = 0.0253 \text{ eV} \) and \( E_{Cd} = 0.55 \text{ eV} \) and the Hogdahl ratio \( f_{th} = 0.02 \) (Rig LE7-01 of Australian OPAL reactor).

The values of \( \left( \frac{\phi_{West, cot}}{\phi_{th, Hogdahl}} \right) \) ratio of 1.011 for \( \alpha = -0.15 \) and 1.006 for \( \alpha = 0.3 \) were achieved. These results suggest that the "non-1/\( v_n \)" - named \( (n,\gamma) \) reaction yield will be around 1% less than the real value if \( \phi_{West, cot} = \phi_{th, Hogdahl} \) is used in the Westcott formalism based calculation. So the value \( \phi_{th} \) can be safely used in placement of \( \phi_{West, cot} \). This is agreed with calculation performed by other authors [5].

The k-factor in eq. N.1 is, however, generated from any values of Westcott’s factor \( g(T_n) \), so eq. (N.2) can be re-written as follows:

\[ r = \phi_{West, cot} \sigma_{West, cot} = \phi_{th} \{1 + f_H \xi(\alpha)\} \sigma_0 \{g(T_n) + r \cdot s_0(\alpha)\} \quad \text{(N.3)} \]

Because \( \left( \frac{\phi_{West, cot}}{\phi_{th, Hogdahl}} \right) = \{1 + f_H \xi(\alpha)\} \approx 1 \) as mentioned above, the reaction rate for the "non-1/\( v_n \)" - named \( (n,\gamma) \) reactions can be calculated as:

\[ r = \phi_{th} \sigma_{eff(non-1/v)} \quad \text{(N.4)} \]

Where \( \sigma_{eff(non-1/v)} = k\sigma_0 \) \quad \text{(N.5)}

For the "1/\( v_n \)" - named \( (n,\gamma) \) reactions, the reaction rate in both thermal and epithermal neutron region calculated based on the Hogdahl convention ion is:

\[ r = \phi_{th} \{\sigma_0 + f_H I_0(\alpha)\} \quad \text{(N.6)} \]

with \( \sigma_{eff(1/v)} = \sigma_0 + f_H I_0(\alpha) \) it is written as:

\[ r = \phi_{th} \sigma_{eff(1/v)} \quad \text{(N.7)} \]

For the above equations, \( \phi_{th, Hogdahl} \) or \( \phi_{th} \) is Hogdahl convention thermal neutron flux, \( f_H = \phi_{epi} / \phi_{th} \) is so-called Hogdahl conventional neutron flux ratio or epithermal to thermal (subcadmium) neutron flux ratio (in this paper \( R_{epi} \) is used instead i.e. \( f_H = R_{epi} \)). \( Q_0 = I_0 / \sigma_0 \) is infinite dilution resonance integral \( I_0 \) per thermal neutron activation cross-section \( \sigma_0 \) at the corresponding energy \( E_0 = 0.0253 \text{eV} \); \( I_0 = \int_{E_0}^{1.64} \frac{\sigma(E)}{E} dE \); \( I_0(\alpha) = Q_0(\alpha)\sigma_0 \).
\[ Q_0(\alpha) = \frac{Q_0 - u}{E_{c_d}^{\alpha}} + \xi(\alpha); \quad u = 2 \sqrt{\frac{E_0}{E_{c_d}}} = 0.429 \]

\[ \xi(\alpha) = 0.429 /(1 + 2\alpha)E_{c_d}^{\alpha} \] is for the cadmium cut-off energy \( E_{c_d} = 0.55 \text{ eV} \) correction, \( \alpha \) is epithermal flux distribution parameter, its extreme values \(-0.15 < \alpha < +0.3\), \( \sigma_{\text{eff}}(1/\nu) \) is Hogdahl convention effective cross-section. \( n_a \) is total neutron density.

\( v_n \) is neutron velocity and \( v_{0} \) is the most probable neutron velocity at 20 °C (2200 m/s).

\( g(T_n) \) is Westcott’s g-factor for neutron temperature \( T_n \), \( g(T_n) \neq 1 \) for "non \(-1/\nu_n\)" reactions.

\( r^* \) is a measure for the epithermal to total neutron density ratio in the Westcott formalism, \( r^* = r(\alpha) \sqrt{T_s/T_0} s_0(\alpha) \)

\( s_0 \) is ratio of the modified reduced resonance integral \((1/v_n - \text{tail subtracted})\) to the thermal cross-section \( \sigma_0 \),

\( s_0 = \frac{2}{\sqrt{\pi}} \{Q_0 - u\} \) and \( s_0(\alpha) = \frac{2}{\sqrt{\pi}} \{Q_0(\alpha) - \xi(\alpha)\} \)

\( \phi_{\text{Westcott}} \) is Westcott conventional (total) neutron flux,
\( \sigma_{\text{Westcott}} \) is Westcott convention effective cross-section,
\( \sigma_{\text{eff}}(\text{non}-1/\nu) \) is "non \(-1/\nu_n\)" effective cross-section.

### 3. Experimental

#### 3.1. Reagents and materials

The isotopically enriched \(^{176}\text{Yb}_2\text{O}_3\) and \(^{176}\text{Lu}_2\text{O}_3\) targets for neutron activation were purchased from Trace-Sciences International Inc. USA [10]. The \(^{176}\text{Yb}_2\text{O}_3\) target isotopic compositions were \(^{176}\text{Yb} (97.6\%), \(^{174}\text{Yb} (1.93\%), \(^{173}\text{Yb} (0.18\%), \(^{172}\text{Yb} (0.22\%), \(^{171}\text{Yb} (0.07\%), \(^{170}\text{Yb} (<0.01\%) \) and \(^{168}\text{Yb} (<0.01\%). \) The main Lanthanide impurities of this target were Er (50 p.p.m), Tm (50 p.p.m) and Lu (50 p.p.m). The \(^{176}\text{Lu}_2\text{O}_3\) target isotopic compositions were \(^{176}\text{Lu} (74.1 \%), \(^{175}\text{Lu} (25.9\%). \) The main Lanthanide impurities of this target were La (66 p.p.m), Yb (13 p.p.m), Tm (<1 p.p.m), Er (17 p.p.m), Dy (4 p.p.m), Gd (6 p.p.m), Eu (20 p.p.m), Sm (2 p.p.m) and Nd (1 p.p.m).

#### 3.2. Targets, reactor irradiations, chemical separation, elemental analysis and radioactivity calibration

The radioactive \(^{177}\text{Lu} + ^{175}\text{Yb}\) solutions were obtained by the reactor thermal neutron irradiation of \(^{176}\text{Yb}_2\text{O}_3\) and/or \(^{176}\text{Lu}_2\text{O}_3\) targets. A quartz ampoule containing an adequate amount of \(^{176}\text{Yb}_2\text{O}_3\) or \(^{176}\text{Lu}_2\text{O}_3\) target was irradiated with a thermal neutron flux in HIFAR reactor (Australia). A 24-hour cooling period was needed to let all \(^{177}\text{Yb} (112 = 1.911 \text{ hours}) \) radionuclides (which formed via \(^{176}\text{Yb} (n, \gamma)^{177}\text{Yb}\)) to be transformed to \(^{177}\text{Lu}\) via beta particle decay. The irradiated target was then dissolved in HCl solution and the radiochemical separation of \(^{177}\text{Lu}\) from the target solution was performed as reported in our previous papers [8,9]. The radioactivity of the different radioisotopes was calibrated using a CAPINTEC Dose calibrator and gamma-ray spectrometer coupled with ORTEC HP Ge detector. The gamma ray energy and counting efficiency calibration of this analyzer system were
performed using a radioactive standard source of $^{152}$Eu solution. Lutetium element and other metal content in the completely decayed $^{177}$Lu solutions (at least $> 10$ half-lives) was analyzed using ICP-MS instrument.

4. Results and Discussion

The methods developed in the above sections were evaluated and used for the assessment of SA values of two typical isotope target systems, enriched $^{176}$Lu and $^{176}$Yb targets. $^{177}$Lu produced from these targets is a representative for the state-of-the-art radioisotopes of high specific radioactivity used in targeted endo-radiotherapy.

4.1. The specific radioactivity of $^{177}$Lu radioisotope produced via $^{176}$Lu $(n, \gamma)$ $^{177}$Lu reaction

The $^{176}$Lu enriched target is used for $^{177}$Lu production. The main nuclear characteristics and nuclear reactions/ radioactive transformations of the $^{176}$Lu and $^{175}$Lu isotope are listed in Table 1. The production of $^{177}$Lu radioisotope is based on the reaction Lu-1. As shown in this data list, the target composes of two stable isotopes, $^{176}$Lu and $^{175}$Lu. In the reactions Lu-6 and Lu-3 the neutron captures yield the isotopes of another chemical element, so these reactions may cause a depression in elemental Lu atom numbers of the target during neutron bombardment.

Table 1. Nuclear characteristics of the radionuclides produced in the $^{176}$Lu enriched target [11].

| Target Stable Isotope (Denoted) | Conc. in target | Cross Sections, Barn | Nuclear reaction and product $(T_{1/2})$ | Reaction No. (Reaction branch) |
|--------------------------------|----------------|----------------------|------------------------------------------|-----------------------------|
| $^{176}$Lu $(S_{1,Lu})$        | $P_{1} = 74.1\%$ | $\sigma_{th}$ $\sigma_{epi}$ | $^{176}$Lu $(n, \gamma)$ $^{177}$Lu $(6.7d)$ $\beta^{-}$ $^{177}$Hf (stable) | Lu-1 $(i = 1)$ |
|                               |                | $2300$ $1200$         |                                          |                             |
| $^{175}$Lu $(S_{2,Lu})$        | $P_{emp,Lu} = 25.9\%$ | $<2.1 \times 10^{-3}$ | $^{176}$Lu $(n, \alpha)$ $^{172}$Tm $(8.2h)$ $\beta^{-}$ $^{172}$Yb (Stable) | Lu-3 $(i = 3)$ |
|                               |                |                      |                                          |                             |

* $^{177}$Lu depression caused by possible $(n, \gamma)$ and/or $(n, p)$ reactions is ignored compared to radioactive decay rate of $^{177}$Lu isotope

However, the effect of these reactions is ignored due to their low cross sections. The reactions Lu-2, Lu-4 and Lu-5 yield long-lived and stable isotopes of the Lutetium element, so the elemental Lu atom numbers of these isotopes are likely to be unchanged during neutron irradiation. This condition shows a similarity between the $^{176}$Lu enriched target and the multi-isotope target system of depression factor $\Delta_{S_{2,Lu}} = 0$ as described in Section 2.1.2.2 (Third bullet). So, eqs. 20–25 are adopted for the $^{177}$Lu specific radioactivity assessment of the Lu target. For this SA assessment process the relevant parameters should be identified to individualize the selected equations for the above mentioned $^{177}$Lu production reaction. These parameters are the following:

$S_{1,A}$ is $^{176}$Lu and re-denoted as $S_{1,Lu}$. $S_{2,A}$ is $^{175}$Lu and re-denoted as $S_{2,Lu}$. $R_i$ is $^{177}$Lu.
P1 is the weight percentage of $^{176}$Lu. $P_{imp,A}$ is weight percentage of $^{175}$Lu and re-denoted as $P_{imp,Lu}$. M1 is atomic weight of $^{176}$Lu. $M_{imp,A}$ is atomic weight of $^{175}$Lu and re-denoted as $M_{imp,Lu}$. $^\Delta_{S_{1},d}$ is $\Delta_{S_{1},d}$ for isotope $^{176}$Lu.

Because $^{176}$Lu has three neutron capture reactions (i=1, 2, 3) as listed in Tab. 1, the $\Delta_{S_{1},d}$ value is

$$\Delta_{S_{1},d} = \phi_{th} \sum_{i=1}^{3} \Omega_{1,i} \quad \text{or} \quad \Delta_{Lu-176} = \phi_{th} \sum_{i=1}^{3} (\sigma_{1,i,(th)} + R \sigma_{1,i,(res)})$$

where $\Omega_{1,i} = \sigma_{1,i,(th)} + R_{epl} \cdot \sigma_{1,i,(epl)} + R_{fast} \cdot \sigma_{1,i,(fast)}$ or $\Omega_{1,1} = (\sigma_{1,1,(th)} + R \sigma_{1,1,(res)})$ or $\Omega_{1,Lu-177} = (\sigma_{1,Lu-177,(th)} + R \sigma_{1,Lu-177,(res)})$

Because the $^{176}$Lu is a typical non-1/$\nu$ nuclide, the $\sigma_{1,i,(th)}$ cross section (of reaction Lu-1 in Table 1), which is tabulated for neutron of $E= 0.0253$ eV and $\nu_{n} = 2,200$ m/s, should be multiplied with a so called k-factor which is based on the Westcott convention equation as discussed in Section 2.3 [5,6]. The $k$ values ranging from 1.67 at 10 °C to 1.9 at 40 °C were calculated for the Munich reactor [5]. As generally accepted, we use the value $k = 1.74$ tabulated in reference [11] for our further calculation.

$$\Omega_{1,i} = \sigma_{1,i,(th)} + R_{epl} \cdot \sigma_{1,i,(epl)} + R_{fast} \cdot \sigma_{1,i,(fast)} = \sigma_{eff} (non-1/\nu) + R_{fast} \cdot \sigma_{1,i,(fast)} = k \cdot \sigma_{0} + R_{fast} \cdot \sigma_{1,i,(fast)}$$

or

$$\Omega_{1,1} = 1.74 \cdot \sigma_{0} + R_{fast} \cdot \sigma_{1,i,(fast)}$$

(the item $R_{fast} \sigma_{1,i,(fast)}$ can be ignored due to the insignificant value of fast neutron flux)

$\Delta_{S_{2},d}$ is $\Delta_{S_{2},d}$ for isotope $^{175}$Lu and $\Delta_{S_{2},d} = \phi_{th} \cdot \sum_{y=1}^{3} \Omega_{2,y} = 0$

where $\Omega_{2,y} = \sigma_{2,y,(th)} + R_{epl} \cdot \sigma_{2,y,(epl)} + R_{fast} \cdot \sigma_{2,y,(fast)} = 0$

$\lambda_{R_{i}}$ is $\Lambda_{Lu-177}$ for radionuclide $^{177}$Lu and $\lambda_{Lu-177} = \lambda_{Lu-177} + \Delta_{Lu-177}$$

where $\lambda_{Lu-177} = \phi_{th} \cdot \sum_{i=1}^{\nu-1} \Omega_{Lu-177}$ and $\lambda_{Lu-177} = \sum_{m=1}^{\nu-1} \lambda_{m,Lu-177}$

$^{177}$Lu specific radioactivities as a function of the target isotopic composition, neutron flux and irradiation time were formulated and calculated (as shown in Figures 1 and 2). The maximum values of $^{177}$Lu specific radioactivity and irradiation time were evaluated. These were used as optimal conditions for carrier-containing $^{177}$Lu production.

As shown in Figure 1a, the irradiation time for maximum yield ($t_{\text{irr,Yield-max}}$) and that for maximum specific radioactivity ($t_{\text{irr,SA-max}}$) are different. The results presented in Figure 3a state that the higher the $^{176}$Lu enrichment of the target, the bigger the difference between values $t_{\text{irr,Yield-max}}$ and $t_{\text{irr,SA-max}}$. The ratio of these times varies with thermal neutron flux applied and reaches a maximum value at the flux value of around $3 \cdot 10^{14}$ $n \cdot cm^{-2} \cdot s^{-1}$ for all the $^{176}$Lu enrichment values of the target.

This result is quite interesting in respect to the optimization of neutron flux utilization, irradiation time and target enrichment. Based on the result of further analysis of eqs. (15) and (24) the conclusion can be drawn that the coincident interaction based on the target burn-up / product depression ratio ($\Delta_{S_{1},d} / \lambda_{R_{i}}$) plays an important role in the creation of this maximum value of $t_{\text{irr,Yield-max}}$ and $t_{\text{irr,SA-max}}$. Although the higher neutron flux irradiation gives the higher SA as shown in Figure 3b, the bigger difference between values $t_{\text{irr,Yield-max}}$ and $t_{\text{irr,SA-max}}$. makes the outcomes of maximum yield and maximum SA incompatible.
Figure 1. a- $^{176}$Lu target nuclide depression, $^{177}$Lu build-up and $^{177}$Lu specific radioactivity in the $^{176}$Lu enriched target vs. irradiation time (Thermal neutron flux: $2.5 \cdot 10^4$ $n \cdot cm^{-2} \cdot s^{-1}$; Target composition: 74.1% $^{176}$Lu + 25.9% $^{175}$Lu; Target weight: 1.0 mg), $\bullet - \bullet - \bullet$: Specific radioactivity of $^{177}$Lu; $\Delta - \Delta - \Delta$: Atom numbers of $^{177}$Lu; $\Diamond - \Diamond - \Diamond$: Atom numbers $^{176}$Lu target nuclide. b- $^{177}$Lu specific radioactivity in the $^{176}$Lu enriched target vs. irradiation time and $^{176}$Lu enrichment of the target (Thermal neutron flux: $2.5 \cdot 10^4$ $n \cdot cm^{-2} \cdot s^{-1}$), $\bullet \bullet \bullet$: 100% purity $^{176}$Lu target; $\Diamond \Diamond \Diamond$: Target composition: 74.1% $^{176}$Lu + 25.9% $^{175}$Lu.
Figure 2. $^{177}$Lu specific radioactivity as a function of irradiation time and $^{176}$Lu isotopic purity in the target (Thermal neutron flux of $1.7 \times 10^4 \text{ n}\cdot \text{cm}^{-2}\cdot \text{s}^{-1}$ was applied. Nuclear data was extracted from literatures [11]).

Figure 3. a- Irradiation time ratio ($t_{\text{irr,SA-max}} / t_{\text{irr,Yield-max}}$) vs. thermal neutron flux and target composition. b- Maximum specific radioactivity of $^{177}$Lu vs. thermal neutron flux and target composition.
This is to say that the production of $^{177}\text{Lu}$ via $^{176}\text{Lu} (n,\gamma)^{177}\text{Lu}$ reaction with neutron flux of around $3\cdot10^{14} \text{ n cm}^{-2} \cdot \text{s}^{-1}$ could be awkward. Hence the production yield and desired SA should be compromised to achieve a cost effective production of clinically useful $^{177}\text{Lu}$ product. The $t_{\text{irr}, \text{SA-max}}$ values increase with the $^{176}\text{Lu}$ enrichment on the target (Figure 2) and the 100% purity $^{176}\text{Lu}$ target showed no-maximum SA value during neutron activation as seen in Figure 1b. This is confirmed by the analysis of differential equation 27, which was described in Section 2.1.2.3 above.

4.2. The specific radioactivity of $^{177}\text{Lu}$ radioisotope produced via neutron-capture-followed-by-radioactive transformation, $^{176}\text{Yb} (n, \gamma)^{177}\text{Yb} (\beta^- \text{decay})^{177}\text{Lu}$

The $^{176}\text{Yb}$ enriched target is used for $^{177}\text{Lu}$ radioisotope production. Based on the isotopic compositions of the elemental Lu -contaminated $^{176}\text{Yb}_2\text{O}_3$ target and the possible nuclear reactions listed in Tables 1 and 2, the total $^{177}\text{Lu}$ radioactivity in this activated $^{176}\text{Yb}$ target composes of one part (denoted as $A_{1-Lu-177}$) induced from the $^{176}\text{Yb}$ target nuclide via reaction $^{176}\text{Yb} (n, \gamma)^{177}\text{Yb} (\beta^- \text{decay})^{177}\text{Lu}$ and another part (denoted as $A_{2-Lu-177}$) from the $^{176}\text{Lu}$ impurity via $^{176}\text{Lu} (n, \gamma)^{177}\text{Lu}$ reaction. As shown in Table 2, the Lu-free $^{176}\text{Yb}$ target contains $^{174}\text{Yb}$ impure isotope, so the $^{175}\text{Lu}$ induced via reaction $^{174}\text{Yb} (n, \gamma)^{175}\text{Yb} (\beta^- \text{decay})^{175}\text{Lu}$ increases the atom numbers of elemental Lu in the target during neutron activation. Especially, due to long-lived $^{175}\text{Yb}$, the $^{175}\text{Lu}$ generation from beta decay of $^{175}\text{Yb}$ will be continued during post-irradiation processing/cooling of the target. The SA degradation effect of $^{174}\text{Yb}$ and Lu impurities on the carrier-free $^{177}\text{Lu}$ producible from an isotopically pure $^{176}\text{Yb}$ target is demonstrably predictable. The assessment of this effect measure is a showcase example for a
complex target system in which a stable brother isotope of radioisotope \( R_i \) may be generated from the elemental impurities.

**Table 2.** Nuclear characteristics of radionuclides produced in \(^{176}\)Yb target matrix [11].

| Stable Isotope (Denoted) | Conc. in target (%) | Cross Section, Barns | Nuclear reactions and products (T1/2) | Reaction No. (Reaction branch) |
|--------------------------|---------------------|----------------------|--------------------------------------|-------------------------------|
| \(^{176}\)Yb (S\(_1\)B) | 97.6                | 3.0                  | \(^{176}\)Yb(n,\(\gamma\)) \(^{177}\)Yb (1.9h) \(\beta^-\)\(^{177}\)Hf (stable) \(\beta^-\)\(^{177}\)Lu (6.7d) \(\beta^-\)\(^{177}\)Yb | Yb-1 (x = 1) |
| \(^{174}\)Yb (S\(_2\)B) | 0.93                | 63.0                 | \(^{174}\)Yb (n,\(\gamma\)) \(^{175}\)Yb (4.2d) \(\beta^-\)\(^{175}\)Lu (Stable) | Yb-2 (y = 1) |
| \(^{173}\)Yb (S\(_3\)B) | 0.18                | 17.4                 | \(^{173}\)Yb (n,\(\gamma\)) \(^{174}\)Yb (Stable) | Yb-3 (z = 1) |
| \(^{172}\)Yb (S\(_4\)B) | 0.22                | 1.3                  | \(^{172}\)Yb (n,\(\gamma\)) \(^{173}\)Yb (Stable) | Yb-4 (p = 1) |
| \(^{171}\)Yb (S\(_5\)B) | 0.07                | 50.0                 | \(^{171}\)Yb (n,\(\gamma\)) \(^{172}\)Yb (Stable) | Yb-5 (q = 1) |
| \(^{170}\)Yb (S\(_6\)B) | <0.01               | 10.0                 | \(^{170}\)Yb (n,\(\gamma\)) \(^{171}\)Yb (Stable) | Yb-6 (v = 1) |
| \(^{168}\)Yb (S\(_7\)B) | <0.01               | 2300                 | \(^{168}\)Yb(n,\(\gamma\)) \(^{169}\)Yb (32d) \(^{169}\)Tm (Stable) | Yb-7 (w = 1) |

* Elemental Lu content being of natural isotopic abundance in \(^{176}\)Yb target is assumed, i.e. 97.41 % \(^{175}\)Lu and 2.59 % \(^{176}\)Lu

Specific radioactivity \(SA_{\text{Lu-177}}\) achievable in the Lu- and \(^{174}\)Yb-contaminated \(^{176}\)Yb target will be calculated based on “isotopic dilution” equation (35) which is applied for a multi radioactive source system. The \(^{176}\)Yb enriched target discussed in this report is referred to as a system composed of two \(^{177}\)Lu radioactive sources, \(S_1\) and \(S_2\). The source \(S_1\) refers to the radioisotopes induced in the elementally pure (Lu impurity-free) Yb target, while \(S_2\) is the radioactive part produced from the Lu impurity in the target.

4.2.1. \(^{177}\)Lu radioactive source 1 (\(S_1\)): Radioactivity (\(A_{1-Lu-177}\)) and specific radioactivity (\(SA_{1-Lu-177}\)) of \(^{177}\)Lu isotope in the elementally pure (Lu impurity-free) Yb target

As shown in Table 2, the target is composed of seven stable Yb isotopes. Among them only \(^{176}\)Yb and \(^{174}\)Yb are involved in the neutron capture reactions (reactions Yb-1 and Yb-2, respectively) to produce the Lu isotopes (\(^{177}\)Lu and \(^{175}\)Lu). It is obvious that in the elementally pure (Lu impurity-free) Yb target, the \(^{177}\)Lu radioactivity is generated from reaction Yb-1. The elemental Lu atom numbers, however, result from both reactions Yb-1 and Yb-2. The following parameters should be evaluated for the SA assessment of \(^{177}\)Lu isotope in the given Yb target.

Radioactivity of carrier-free \(^{177}\)Lu (\(A_{1-Lu-177}\)) from the reaction Yb-1 in the Lu-free Yb target.

Radioactivity at post-irradiation time \(t_c\) (\(A_{1-Lu-177,t_c}\)) can be calculated using eq.28. The carrier-free \(^{177}\)Lu activity is the following:
\[
A_{1-Lu-177,\nu} = N_{0,Yb-176} \cdot f_2 \cdot f_3 \cdot \lambda_{Lu-177} \cdot \left( \frac{e^{-d_{1,Yb}}}{(d_2 - d_1)(d_3 - d_1)} + \frac{e^{-d_{3,Yb}}}{(d_1 - d_3)(d_3 - d_2)} + \frac{e^{-d_{5,Yb}}}{(d_1 - d_5)(d_2 - d_3)} \right) \cdot e^{-\sum_{m=1}^{\nu} \gamma_{m,Yb-177}} \tag{36}
\]

The relevant parameters should be identified to individualize this equation for the reaction Yb-1 mentioned in Table 2. Individualizing eq.28 with \( \Omega_{1,1} \) for \( \Omega_{Yb-176,1} \) (or \( \Omega_{Yb-176,Yb-177} \)), \( \Omega_y \) for \( \Omega_{Yb-177} \) and \( \nu_y \) for \( \Omega_{Lu-177} \), these parameters are the following:

\[
\begin{align*}
\Omega_{Yb-176,Yb-177} & = \sigma_{Yb-176,Yb-177} \cdot \sigma_{Yb-176,Yb-177} + R_{fast} \cdot \sigma_{Yb-176,Yb-177} \cdot \lambda_{Yb-177} \\
\Omega_{Yb-177} & = \sigma_{Yb-177} \cdot \sigma_{Yb-177} + R_{fast} \cdot \sigma_{Yb-177} \cdot \lambda_{Yb-177} \\
\Omega_{Lu-177} & = \sigma_{Lu-177} \cdot \sigma_{Lu-177} + R_{fast} \cdot \sigma_{Lu-177} \cdot \lambda_{Lu-177}
\end{align*}
\]

\((R_{fast} \cdot \sigma_{Yb-176,Yb-177}, \sigma_{Yb-177}, \sigma_{Lu-177}) \) and \( R_{fast} \cdot \sigma_{Yb-177,1,fast} \) can be ignored due to the insignificant value of fast neutron flux.

\[
\begin{align*}
d_1 &= \phi_{db} \cdot \sum_{s=1}^{s-4} \Omega_{Yb-176,Yb-177,s} + \phi_l \cdot \sum_{m=1}^{m-1} \lambda_{m,Yb-177} + \phi_l \cdot \sum_{m=1}^{m-1} \Omega_{Yb-177,m} \;
\end{align*}
\]

\[
\begin{align*}
d_2 &= \phi_{db} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-177,m} \;
\end{align*}
\]

\[
\begin{align*}
f_2 &= \phi_{db} \cdot \Omega_{Lu-177,1} \;
\end{align*}
\]

\[
\begin{align*}
f_3 &= \lambda_{1,Yb-177} \cdot \sum_{m=1}^{m-1} \lambda_{m,Yb-177} = \lambda_{Lu-177} \cdot \sum_{m=1}^{m-1} \Omega_{Lu-177,m}
\end{align*}
\]

\[
N_{0,Yb-176} \text{ denotes the atom numbers of } ^{176}Yb \text{ in the m grams weight target.}
\]

\[
N_{0,Yb-176} = 6.02 \cdot 10^{23} \cdot m \cdot P_{Yb-176} / (100 \cdot 176), \text{ where } P_{Yb-176} \text{ is the weight percentage of } ^{176}Yb \text{ in the } ^{176}Yb \text{ enriched target.}
\]

**Total elemental Lu atom numbers** \( N_{1-Lu} \) in the Lu impurity-free \(^{176}Yb \) target. This amount is the sum of the Lu atom numbers of carrier-free \(^{177}Lu \) radioisotope \( (N_{1-Lu-177}) \) from reaction Yb-1 and Lu atom numbers generated from the \(^{174}Yb \) impurity of the target \( (N_{1-Lu-175}) \) from reaction Yb-2.

\[
N_{1-Lu} = N_{1-Lu-177} + N_{1-Lu-175}
\]

\( N_{1-Lu-177} \) is calculated by a quotient of eq.36 and \( \lambda_{Lu-177} \):

\[
N_{1-Lu-177} = N_{0,Yb-176} \cdot f_2 \cdot f_3 \cdot \left( \frac{e^{-d_{1,Yb}}}{(d_2 - d_1)(d_3 - d_1)} + \frac{e^{-d_{3,Yb}}}{(d_1 - d_3)(d_3 - d_2)} + \frac{e^{-d_{5,Yb}}}{(d_1 - d_5)(d_2 - d_3)} \right) \cdot e^{-\sum_{m=1}^{\nu} \gamma_{m,Yb-177}} \tag{38}
\]

\( N_{1-Lu-175} \) calculation is adopted from eq.33b described in Section 2.2.1.2.

\[
N_{1-Lu-175} = \frac{N_{0,Yb-174}}{\left( \lambda_{Yb-175} - \phi_{db} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-174,m} \right)} \cdot \left( \lambda_{Yb-175} - \phi_{db} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-174,m} \right) \cdot \left( 1 - e^{-\phi_{db} \cdot \Omega_{Yb-174,1}} \right) \cdot \left( e^{-\phi_{db} \cdot \Omega_{Yb-174,1}} \cdot e^{-\phi_l \cdot \Omega_{Yb-174,Yb-175}} \right) \cdot \left( e^{-\phi_l \cdot \Omega_{Yb-174,Yb-175}} \cdot e^{-\phi_l \cdot \Omega_{Yb-174,Yb-175}} \right)
\]

Individualizing eq. (33b) for the \(^{175}Lu \) isotope generated from \(^{174}Yb \), the following is identified. 

\( S_{2,B} \) is stable isotope \(^{174}Yb \). \( \Delta_{Yb,Yb} \) is re-denoted as \( \Delta_{Yb-174} \) for isotope \(^{174}Yb \). Because \(^{174}Yb \) has only one neutron capture reaction \( (y = 1) \) as shown in Tab. 2, the \( \Delta_{Yb-174} \) value is

\[
\Delta_{Yb-174} = \phi_{db} \cdot \sum_{y=1}^{y} \Omega_{Yb-174,y} = \phi_{db} \cdot \Omega_{Yb-174,1}, \text{ where } \Omega_{Yb-174,y} = \sigma_{Yb-174,y} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-174,m} + R_{fast} \cdot \sigma_{Yb-174,y} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-174,m} \tag{39}
\]

(The items \( R_{fast} \cdot \sigma_{Yb-174,y} \) can be ignored due to the insignificant value of fast neutron flux).

\( \Delta_{Yb-175} \) is re-denoted as \( \Delta_{Yb-175} \) for radioisotope \(^{175}Yb \) and \( \Delta_{Yb-175} = \phi_{db} \cdot \sum_{m=1}^{m-1} \Omega_{Yb-175} \).
\( \Lambda_{R_y} \) is re-denoted as \( \Lambda_{Yb\rightarrow 175} \) as \( \sum_{m=1}^{\infty} \lambda_{m,R_y} \) and \( \Lambda_{Yb\rightarrow 175} \) as \( \sum_{m=1}^{\infty} \lambda_{m,Yb\rightarrow 175} \) for radionuclide Yb\(_{175}\) and \( \Lambda_{Yb\rightarrow 175} = \sum_{m=1}^{\infty} \lambda_{m,Yb\rightarrow 175} \). For simplification, \( \sum_{m=1}^{\infty} \lambda_{m,Yb\rightarrow 175} = \lambda_{Yb\rightarrow 175\rightarrow Lu\rightarrow 175} \) is applied and value \( \Delta_{Yb\rightarrow 175} \) is ignored due to unavailability of \( \sum_{m=1}^{\infty} \lambda_{m}\) value. So we get \( \Lambda_{Yb\rightarrow 175} = \lambda_{Yb\rightarrow 175\rightarrow Lu\rightarrow 175} = \lambda_{Yb\rightarrow 175} \cdot N_{0,5_{1,8}} \) is the initial \( ^{174}\text{Yb} \) atom numbers \( N_{0,5_{1,8}} \). This quantity is \( N_{0,5_{1,8}} = 6.02 \cdot 10^{23} \cdot m \cdot P_{Yb\rightarrow 174}/(100 \cdot 174) \). P \( \text{Yb}\rightarrow 174 \) is the weight percentage of \( ^{174}\text{Yb} \) impurity in the \( ^{176}\text{Yb} \) target.

\( ^{177}\text{Lu} \) specific radioactivity of the Lu impurity-free \( ^{176}\text{Yb} \) target. The specific radioactivity of \( ^{177}\text{Lu} \) source (the \( ^{177}\text{Lu} \) radioactive source 1) generated from the elemental Lu impurity-free \( ^{176}\text{Yb} \) target is the following:

\[
S_{L_{1}-Lu\rightarrow 177} = 100 \cdot N_{1-Lu\rightarrow 177,t_{1}}/(N_{1-Lu\rightarrow 177,t_{1}} + N_{1-Lu\rightarrow 175,t_{1}}) = 100 \cdot N_{1-Lu\rightarrow 177,t_{1}}/N_{1-Lu}
\]

4.2.2. \( ^{177}\text{Lu} \) radioactive source 2 \( (S_{2}) \): Radioactivity \( A_{2-Lu\rightarrow 177} \) and specific radioactivity \( S_{A_{2}-Lu\rightarrow 177} \) of \( ^{177}\text{Lu} \) generated from \( ^{176}\text{Lu} \) \( (n,\gamma) \) \( ^{177}\text{Lu} \) reaction of the elemental Lu impurity in the \( ^{176}\text{Yb} \) target.

Specific radioactivity. \( S_{A_{2}-Lu\rightarrow 177} \) of \( ^{177}\text{Lu} \) radioisotope produced in this target is calculated using the same process as described in the previous section 4.1 ‘The specific radioactivity of \( ^{177}\text{Lu} \) radioisotope produced via \( ^{176}\text{Lu} \) \( (n,\gamma) \) \( ^{177}\text{Lu} \) reaction’ of Results and Discussion. In this case the impure Lu content in the \( ^{176}\text{Yb} \) target is assumed to have natural abundance, so the values \( P_{1} = 2.59\% \) for \( ^{176}\text{Lu} \) and \( P_{imp,Lu\rightarrow 175} = 97.41\% \) for \( ^{175}\text{Lu} \) are put into calculation.

Radioactivity. Radioactivity of \( ^{177}\text{Lu} \) from the elemental Lu impurity in the \( ^{176}\text{Yb} \) target at post-irradiation time \( t_{c} \) is calculated using eq. (14):

\[
A_{2-Lu\rightarrow 177} = \frac{N_{0,5_{1,8}-176} \cdot \phi_{th} \cdot \Omega_{L_{1},Lu\rightarrow 176} \cdot \sum_{m=1}^{\infty} \lambda_{m,Lu\rightarrow 177} \cdot (e^{-\Delta_{Lu\rightarrow 177,t_{c}}} - e^{-\Delta_{Lu\rightarrow 177,t_{c}}} \cdot e^{-\sum_{m=1}^{\infty} \lambda_{m,Lu\rightarrow 177}})}{\Lambda_{Lu\rightarrow 177} - \Delta_{Lu\rightarrow 176}}
\]

\( N_{0,5_{1,8}-176} = 6.02 \cdot 10^{23} \cdot m \cdot P_{1}/(100 \cdot 176) \); \( P_{1}=2.59\% \) is natural abundance of \( ^{176}\text{Lu} \). \( m \) is the weight of elemental Lu impurity in the \( ^{176}\text{Yb} \) target.

4.2.3. \( ^{177}\text{Lu} \) specific radioactivity \( S_{A_{L_{1}}-177} \) and \( ^{177}\text{Lu} \) radioactivity \( A_{L_{1}-177} \) in the Lu-contaminated \( ^{176}\text{Yb} \) target as a combination of \( ^{177}\text{Lu} \) radioactive source \( S_{1} \) and \( S_{2} \).

\( ^{177}\text{Lu} \) specific radioactivity \( S_{A_{Lu}-177} \). The method of SA assessment of the mixture of several radioactive sources (referred to Section 2.2.2) is used for the SA calculation. The \( ^{177}\text{Lu} \) specific radioactivity \( S_{A_{Lu}-177} \) in the Lu-contaminated \( ^{176}\text{Yb} \) target is calculated using eq. (35) with relevant parameters of the mixture of two \( ^{177}\text{Lu} \) radioactive sources \( S_{1} \) and \( S_{2} \) as described above,

\[
S_{A_{Lu}-177} = \frac{S_{A_{1}-Lu\rightarrow 177} \cdot S_{A_{2}-Lu\rightarrow 177} \cdot (A_{1,Lu\rightarrow 177} + A_{2-Lu\rightarrow 177})}{A_{1-Lu\rightarrow 177} \cdot S_{A_{2}-Lu\rightarrow 177} + A_{2-Lu\rightarrow 177} \cdot S_{A_{1}-Lu\rightarrow 177}}
\]

\( ^{177}\text{Lu} \) radioactivity. Total \( ^{177}\text{Lu} \) radioactivity in the target is \( A_{Lu\rightarrow 177} = (A_{1-Lu\rightarrow 177} + A_{2-Lu\rightarrow 177}) \). \( ^{175}\text{Lu} \) isotope generated from \( ^{174}\text{Yb} \) \( (n,\gamma) \) \( ^{175}\text{Yb} \) \( (\beta^{-}\) decay) \( ^{175}\text{Lu} \) process and elemental Lu impurities
remaining in the $^{177}$Lu product (which is chemically separated from $^{176}$Yb target) makes the SA value of $^{177}$Lu strongly decreased.

Rendering the differential of eq. (42) equal to zero offers the way to calculate the irradiation time at which the SA of $^{177}$Lu reaches maximum value ($SA_{\text{Lu-177, max}}$). The maximum SA is predicted based on the opposite tendency of SA variation of elementally pure and Lu-contaminated Yb targets. This argument is supported by the results obtained below.

The SA of $^{177}$Lu radioisotope produced by the $^{176}$Yb (n, γ) $^{177}$Yb (β− decay) $^{177}$Lu process as a function of the elemental and isotopic impurities of the $^{176}$Yb enriched target and the times $t_{\text{irr}}$ and $t_c$ is shown in Figures 4 and 5. The experimental results reported in our previous publications agree well with the theoretical calculation results shown in Figure 5. The maximum SA value present on the curve D of Figure 4 shows a combined effect of $^{174}$Yb- and elemental Lu- impurities on the SA degradation. This tells us that the irradiation time should be optimized to obtain the highest SA for $^{177}$Lu produced via $^{176}$Yb (n, γ) $^{177}$Yb (β− decay) $^{177}$Lu reaction. While being the best theoretical way to produce carrier-free $^{177}$Lu, with this reaction we always obtain a $^{177}$Lu product of much lower SA due to the use of an isotopically/elementally impure target. The maximum SA value mentioned above is characterized for a specific target composition and its neutron irradiation conditions, so the theoretical assessment of SA developed in this paper is important before starting the neutron activation process. This avoids over-bombardment destroying the SA of $^{177}$Lu product and wasting expensive reactor operation time. Moreover, the post-irradiation processing time should be minimized to keep the SA as high as possible, the effect of which is shown in Figure 5.

**Figure 4.** Specific radioactivity of $^{177}$Lu radioisotope in the $^{176}$Yb target vs. irradiation time and content of $^{174}$Yb- and elemental Lu- impurities. (Thermal neutron flux: $5 \cdot 10^{13}$ n·cm$^{-2}$·s$^{-1}$, Nuclear data extracted from literatures [11]).

![Figure 4](image)
**Figure 5.** Specific radioactivity of $^{177}$Lu in the $^{176}$Yb target vs. post-irradiation time and content of $^{174}$Yb and elemental Lu impurities. (Thermal neutron flux: $5 \times 10^{13} \text{n cm}^{-2} \text{s}^{-1}$; Irradiation time: 240 hours; Nuclear data extracted from literatures [11]).

![Graph showing specific radioactivity of $^{177}$Lu](image)

a: SA of $^{177}$Lu isotope in the impurity-free $^{176}$Yb target; b: SA of $^{177}$Lu isotope in the $^{176}$Yb target containing 1.93% $^{174}$Yb; c: SA of $^{177}$Lu isotope in the $^{176}$Yb target containing 50 p.p.m Lu element impurities; d: SA of $^{177}$Lu isotope in the target containing 1.93% $^{174}$Yb and 50 p.p.m Lu element impurity. (*) is the experimental measurement result for this type of target.

### 5. Conclusions

Several factors affect SA of the radionuclide product which can be produced either by neutron capture reaction or by neutron-capture-followed-by-radioactive transformation processes. Among them the target composition (elemental and isotopic impurities), target nuclide and produced radioisotope depression causes (including target nuclide burn-up, reaction rate of target nuclide and the decay property of produced radionuclide) and the activation or post-irradiation time are most accounted for. With the method of SA assessment of multi radioactive source system the SA of a radioisotope produced in a reactor from different targets can be evaluated. The theoretical SA assessment of a radioactive nuclide has definitely given us a firm basis to set up an optimized process for the production of clinically useful radioisotopes and to evaluate the quality of the radionuclide product. A useful computer code based on the above developed SA assessment methods can be set up for a convenient daily use in the reactor-based production of medical radioisotopes such as $^{177}$Lu, $^{153}$Sm, $^{169}$Yb, $^{165}$Dy, $^{153}$Gd... This evaluation plays a complementary or even substantial role in the quality management system regarding certifying the SA of the product which may be experimentally inaccessible due to radiation protection and instrumentation difficulties in the practical measurement of very low elemental content (<0.01 μg/mCi·μL) of a radioactive solution of very high specific volume and specific radioactivity.
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