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# Induced activation in accelerator components 

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#### Abstract

The residual activity induced in particle accelerators is a serious issue from the point of view of radiation safety as the long-lived radionuclides produced by fast or moderated neutrons and impact protons cause problems of radiation exposure for staff involved in the maintenance work and when decommissioning the facility. This paper presents activation studies of the magnets and collimators in the High Energy Beam Transport line of the European Spallation Source due to the backscattered neutrons from the target and also due to the direct proton interactions and their secondaries. An estimate of the radionuclide inventory and induced activation are predicted using the GEANT4 code.


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## I. INTRODUCTION

Activation induced by particle nuclear interactions in beam line components represents one of the main radiation hazards of high-energy accelerators. Although elements such as targets, collimators, magnets, and beam dumps are built to withstand radiation, they become themselves highly radioactive as a result of induced activation. Induced radioactivity is due either to direct interactions of the incoming beam or indirect interactions of secondary particles in the accelerator components leading to radionuclides production. This activation causes remnant ambient dose rates inside accelerator tunnels and target areas but also means that when components are being replaced at the end of their operational lifetime, they must be treated as radioactive waste. Large quantities of activated material arise when the whole accelerator is decommissioned as dismantling it will pose a major challenge for radiation protection. Exposure to radiation from induced activation can occur in connection with handling, transport, machining, welding, chemical treatment, and storage of irradiated items. These procedures can be extremely difficult because the accumulated doses can exceed the permitted limits and remote handling becomes necessary. In the field of radioactive materials and waste there are no internationally agreed recommendations while in radiation protection the

[^0]dose limits are dependent on the national regulations. Both the International and European Basic Safety Standards [1] only contain tables with radionuclide specific exemption limits [2] and do not make recommendations with respect to the clearance of radioactive material.

Because the accelerator components reveal high induced activation during normal operations and after accelerator shutdown, it is of primary importance to predict correctly their radionuclide inventory and residual activity before any handling and maintenance procedures. The technical challenge is first to ensure that the beam losses are small as residual activity depends greatly not only on material properties but also on the amount of beam loss. The main causes of beam loss in high-current accelerators are (i) space charge effects that arise due to Coulomb repulsion between particles; the Coulomb repulsion becomes more important as the beam current is increased and causes an increased emittance leading to beam losses; (ii) beam halo surrounding the beam core, caused by space charge induced emittance growth; (iii) emittance increase which could be due to several reasons like space charge effects, nonlinear resonances, chromatic aberrations in lenses; (iv) backstreaming neutrons coming from the target on the beam pipe; (v) mismatch of the beam across accelerator elements transitions; and (vi) low aperture to rms beam size ratio; this should be kept reasonably high to prevent the beam from hitting elements and getting lost.

The high-energy protons lost along the beam line generate secondary neutrons by spallation and these in turn activate the accelerator components through which they pass. Different nuclear fragments can be emitted as well converting the nucleus to an isotope most probably of
a different element with a high chance of being radioactive. The radionuclides that are produced are unstable and undergo radioactive decay. Although the induced radioactivity will depend on the beam loss history including cool down history, the probability of producing a particular isotope will also depend on the composition of the material struck, the spectrum of secondaries produced, and the production cross section of the isotope concerned [3].

The spallation reactions take place in two steps: a fast nuclear cascade followed by a slow evaporation process. In the first step the high-energy particle entering the nucleus is capable of knocking out one or more nucleons or even a fragment of the target nucleus. Another part of the projectile energy is transferred to the remaining nucleons which subsequently collide with each other, distributing the energy almost equally and in this way a cascade develops. Some of the nucleons reach the nuclear boundary and get lost; others are caught and distribute their kinetic energy among the remaining nucleons in the nucleus. The intranuclear cascade leads finally to a highly excited nucleus. The intranuclear cascade stage is not distinctively separated from the deexcitation stage and the preequilibrium emission follows. In this intermediate stage fast particles are emitted. In the deexcitation stage the remnant nucleus loses its residual excitation energy by evaporation processes emitting neutrons and low energy light particles (protons, deuterons, alpha particles). Some of the nucleons may leave the nucleus if their energy is higher than their binding energy. Nucleons at the surface in a less strongly bound state have binding energies of a few MeV , whereas nucleons in the deeper-lying shells need about 50 MeV to remove them from the nucleus. Charged particles such as protons and pions (produced mainly in the decay of the Delta resonance) have to overcome the Coulomb barrier in addition. If they overcome the strong force field of the nucleus, they even get additional energy. When the nucleus does not have enough energy to emit neutrons it deexcites by emitting gamma rays. At the end of this process the resulting nucleus is usually $\beta$ radioactive (although a bias to the production of $\alpha$ emitters exists for high A nuclei) and decays until it becomes stable. The disintegration is a random process being defined by the half-life of the produced radionuclides. Radioisotopes with a short lifetime of a couple of minutes can be considered as bringing no contribution to the residual radiation studies. The total number of the disintegrations of all sorts occurring per unit time represents the activity of the activated material.

## II. ACTIVATION FORMULA

The nuclear reactions are characterized quantitatively by their cross section. Assuming that the cross sections $\sigma_{T, \nu}$ of the possible nuclear reactions of converting a target nucleus $T$ into an isotope $\nu$ are known, the number of radioactive nuclei of isotope $\nu$ per gram of target material produced per unit time is

$$
\begin{equation*}
n_{\nu}=\Phi \frac{N_{0}}{A_{T}} \sigma_{T, \nu} \tag{1}
\end{equation*}
$$

where $\Phi$ is the fluence rate of projectile particles defined as the number of particles incident on a sample per unit surface, per second, $N_{0}$ is Avogadro's number, $A_{T}$ is the mass number of the target material, and $\sigma_{T, \nu}$ is the integrated cross section over the whole fluence spectrum. Once the radioactive nuclei have been produced, their number decays exponentially with time. Taking into account both the production rate $n_{\nu}$ and the decay rate, we can write

$$
\begin{equation*}
\frac{d N_{\nu}}{d t}(t)=n_{\nu}-\lambda_{\nu} N_{\nu}(t), \tag{2}
\end{equation*}
$$

where $\lambda_{\nu}$ is the decay constant of isotope $\nu$ and $N_{\nu}(t)$ is the number of nuclei $\nu$ as a function of time. If the target is irradiated during the irradiation time $t_{i}$, the number of radioactive nuclei $N_{\nu}\left(t_{i}\right)$ of an isotope $\nu$ is

$$
\begin{equation*}
N_{\nu}\left(t_{i}\right)=\Phi \frac{N_{0}}{\lambda_{\nu} A_{T}} \sigma_{T, \nu}\left(1-\exp \left[-\lambda_{\nu} t_{i}\right]\right) \tag{3}
\end{equation*}
$$

From Eq. (3) the activity after a cooling time $t_{c}$ can be obtained as

$$
\begin{equation*}
-\frac{d N_{\nu}}{d t_{c}}=\Phi \frac{N_{0}}{A_{T}} \sigma_{T, \nu}\left(1-\exp \left[-\lambda_{\nu} t_{i}\right]\right) \exp \left[-\lambda_{\nu} t_{c}\right] . \tag{4}
\end{equation*}
$$

This is the activation formula for one isotope that gives the disintegration rate of that particular isotope $\nu$ in one gram of target material $T$ which has been irradiated during a time $t_{i}$ and left to decay during a time $t_{c}$. The total specific activity of the target material will be the sum of the specific activities of all the particular isotopes $\nu$ producible. Although this is a simplified approach not considering daughter production, in the later chapter where we simulate the isotope production, when calculating the production rate we will take into account not only the production due to the direct interaction of various particles with the elements [Eq. (1)] but also the contribution to the production rate due to the decay of other isotopes produced in these interactions.

Quantification of the residual activity provides useful information regarding tolerable beam losses in the machine, choice of construction materials for accelerator components, and estimates of the necessary cooling time after switching off the accelerator. The current paper aims to give an estimate of the amount of radionuclides produced and consequently the radioactivity induced in the ESSHEBT components due to backstreaming neutrons coming from the target and direct proton interactions and their secondaries.

## III. THE GEANT4 MODEL

Owing to the complexity of the spallation reactions that take place inside the material of the accelerator components, a proper analytical treatment is hardly possible. Spallation reactions are complicated processes with many reaction channels and gamma spectroscopy for radionuclide identification proved in many cases to be not sufficient; therefore, Monte Carlo techniques are used to quantify the radioisotopes produced. One can apply the Monte Carlo method to the theoretical reaction model and thereby predict the outcome of the spallation reactions and then sum over all production channels to give an estimate of the total activity. The GEANT4 [4] code has been used in the current simulations to predict the spallation reaction products and to calculate the activity induced. It tracks the incident particle and simulates its interactions with nuclei of the material as well as the production of the secondary particles and their interactions. The geometry of the system and the characteristics of the particle beam must be also modeled in the code.

GEANT4 provides an extensive set of hadronic physics models for energies up to $10-15 \mathrm{GeV}$, both for the intranuclear cascade region and for modeling the evaporation processes. There are many different (data-based, parametrized, and theory-driven) models using different approximations and each has its own applicable energy range. Monte Carlo codes usually come with their own physics models and the user is offered default selections. Because of the vast range of applications, GEANT4 will not give the users any default physics models; the users themselves have to work out what models to use for what processes. In order to model the proton and neutron inelastic interactions in the energy range relevant for this study, the best physics models available are the three theoretical intranuclear cascade models provided by GEANT4: INCL/ABLA (Liégé) model, Binary cascade, and the Bertini model. The Liégé intranuclear cascade model together with the independent evaporation/fission code ABLA have been validated against experimental data for spallation processes in many different heavy elements [5]. However, the INCL/ABLA validation results presented at the IAEA benchmark for spallation reactions show that, for energies lower than 150 MeV , the results of the Liégé model are not so good as above this energy [6]. This is because the model does not have preequilibrium: INCL cascade is directly "coupled" to equilibrium deexcitation handled by ABLA and therefore it does not describe well enough low energy reactions (where nuclear structure effects start to play their role). Above 150 MeV , INCL/ ABLA works very well, being one of the best models available. On the other hand, the other two models available in GEANT4, Bertini and Binary cascade, do incorporate the preequilibrium model. The preequilibrium model in GEANT4 has been recently improved following a validation study against the transmutation by adiabatic resonance
crossing (TARC) experiment data, in order to improve several shortcomings in applying this model to neutron spallation processes [7]. All these recent developments have been considered and implemented in our code.

For the simulations presented in this paper, the Bertini model was selected. For neutron energies below 20 MeV , the high-precision models were selected. These models use the ENDF/B-VII [8], JENDL [9], MENDL-2 [10] and other data libraries [11]. The $S(\alpha, \beta)$ [12,13] coefficient, which takes into account the corrected treatment for neutron scattering on chemically bound elements in the thermal region, has also been implemented in the GEANT4 physics list used for this study.

## IV. RESULTS

The change in the lattice structure of the materials that are irradiated lead to the failure of the components and recently there has been a renewed interest in the topic of radiation damage as high-power accelerators are required for physics experiments. One such accelerator is the European Spallation Source (ESS) which will be built in Lund, Sweden. This is a high-power ( 5 MW ) proton accelerator that will produce high-intensity neutron beams through spallation. It is therefore essential to know what the residual dose rates will be in the most exposed elements like magnets and collimators placed in the close proximity of the target; therefore simulations had to be performed.

The calculations presented here refer to the linac layout such as it is presented in the conceptual design report [14]. This layout includes (i) an ion source, (ii) a low energy beam transport system, (iii) a radio frequency quadrupole accelerating from 75 keV to 3 MeV , (iv) a medium energy beam transport system, (v) a warm linac section used for acceleration to 50 MeV , (vi) a superconducting linac section for acceleration to 2.5 GeV , and (vii) the highenergy beam transport system (HEBT) which will transport the proton beam from the underground linac to the spallation tungsten target at the surface level [15]. The various elements of the transport system have been designed and optimized to obtain the desired beam sizes, phase advances, and Twiss parameters at specific locations. The HEBT line has four sections and Fig. 1 shows the layout of the HEBT sections. The HEBT starts with a straight 100 m long underground section (HEBT-S1) that accommodates a collimation system and space for additional cryomodules for a power and energy upgrade. The collimation system is placed at the beginning of the HEBT to remove possible halos generated in the upstream linac and subsequently to reduce losses in the downstream elements. The placement of the collimators at this location allows the reduction of the aperture of all magnetic elements throughout the HEBT. Only transverse collimation is proposed. In addition to capturing the halo, the collimators will additionally accept single short stray pulses and will be used as a diagnostic tool.


FIG. 1. The layout of a short section of the HEBT-S1, the HEBT-S2 and HEBT-S3. The layout includes the magnets-quadrupoles (green), dipoles (red), and octupoles (blue), the fixed collimator (cyan), and the target monolith (including the proton beam window and the target wheel) to shield the neutrons from the target.


FIG. 2. The horizontal (blue) and vertical (red) 1 rms beam envelopes throughout the HEBT. The colors denote different elements: red for dipoles, green for quadrupoles, blue for octupoles, orange for the fixed collimator, and grey for the proton beam window.

The HEBT-S1 is followed by a semivertical bending section denoted HEBT-S2 bringing the beam from the underground linac tunnel to the target, 1.6 m above ground level. The third horizontal section (HEBT-S3) includes the expansion system to provide the requested beam footprint at the target while keeping the peak current density minimal. The last section is a short horizontal section for a beam dump to be used for accelerator tuning and commissioning. The HEBT-S3 components were modeled in the GEANT4 simulation code since the elements of this particular beam line are the ones at risk because of the backstreaming neutrons coming from the target. HEBT-S3 is designed in order to fulfil the requirements for the beam footprint and peak current density at the target. In order to obtain a peak current density of less than $64 \mu \mathrm{~A} / \mathrm{cm}^{2}$ inside the requested footprint $(160 \times 60 \mathrm{~mm})$ while keeping the power deposited outside the footprint low, octupoles are utilized. Downstream of the last octupole, three quadrupoles are placed in order to spread the now folded beam over the desired footprint area. The input particle distribution is a superposition of two 6D Gaussian distributions, a core and a halo distribution, with widths determined by the linac output ( 1.90 mm horizontally and 2.74 mm vertically) [16]. The baseline rms envelopes are shown in Fig. 2 and the horizontal and vertical particle densities along the HEBT in Figs. 3(a) and 3(b).

## A. Radionuclides inventory in the HEBT-S3 magnets and collimators

Beam losses will inevitably occur along the HEBT line. These losses originate from weak tails of beam lost over
long operation periods in both transverse and longitudinal planes or short pulses (until the beam is shut down) of the main beam impacting major components. Experience has shown that activation caused by uncontrolled beam losses uniformly distributed along the beam line of $1 \mathrm{~W} / \mathrm{m}$ can be accepted for high-energy proton accelerators as a threshold for the hands-on maintenance [17]. Therefore the proton beam losses at the European Spallation Source were established to be on the average lower than $1 \mathrm{~W} / \mathrm{m}$. This value was used to calculate the direct beam impact on the magnets. We considered the direct impact of all the secondaries produced in the interactions between the incident protons and the beam pipe, as well as inside each element. At the collimator the beam losses are around 20 kW [18]. This value was used to simulate the direct beam impact effect on the particle collimator.

The HEBT line includes a large number of magnets with associated power supplies, and both normal conducting magnets as well as high-temperature superconducting magnets are considered for this beam line. The parameters for the elements in the HEBT line are given in the Table I.

The effect of the backscattered neutrons from the target on various magnets along the beam line (Fig. 4) has been considered in our activation calculations. The aperture of the octupole is reduced compared to the rest of HEBT magnets to obtain large field derivatives for a limited pole tip field [16]. This will unavoidably lead to higher losses at this element as compared to the upstream HEBT elements; however, these losses are expected to be small compared to the amount of neutrons from the target. Regarding the


FIG. 3. The normalized horizontal (a) and vertical (b) beam particle density along the HEBT. The projected beam profiles are shown on a linear scale. The vertical lines at 153 m (a),(b) represent the fixed collimator.
radioactive inventory originating from accelerator components experience has shown that the gamma emitting isotopes are dominant along with the relatively short-lived beta isotopes. Therefore, in order to study the beam line components activation, the code has been designed to register the production of any radioactive isotope that decays emitting gamma rays and also that has a lifetime longer than 24 hr . Shorter lived isotopes, like ${ }^{24} \mathrm{Na}$ for example, can pose a safety concern if the tunnel has to be accessed in emergency situations, but we limited our study to include only the isotopes that live longer than 24 hr . Therefore the isotopes that are recorded by our simulation code are ${ }^{7} \mathrm{Be},{ }^{46} \mathrm{Sc},{ }^{44} \mathrm{Ti},{ }^{51} \mathrm{Cr},{ }^{54} \mathrm{Mn},{ }^{59} \mathrm{Fe},{ }^{56} \mathrm{Co},{ }^{57} \mathrm{Co}$, ${ }^{58} \mathrm{Co},{ }^{60} \mathrm{Co},{ }^{65} \mathrm{Zn},{ }^{75} \mathrm{Se},{ }^{84} \mathrm{Rb},{ }^{85} \mathrm{Sr},{ }^{88} \mathrm{Y},{ }^{95} \mathrm{Zr},{ }^{94} \mathrm{Nb},{ }^{95} \mathrm{Nb}$, ${ }^{106} \mathrm{Ru},{ }^{109} \mathrm{Cd},{ }^{111} \mathrm{In},{ }^{113} \mathrm{Sn},{ }^{125} \mathrm{Sn},{ }^{124} \mathrm{Sb},{ }^{125} \mathrm{Sb},{ }^{125} \mathrm{I},{ }^{132} \mathrm{Cs}$, ${ }^{134} \mathrm{Cs},{ }^{137} \mathrm{Cs},{ }^{133} \mathrm{Ba},{ }^{139} \mathrm{Ce},{ }^{141} \mathrm{Ce},{ }^{144} \mathrm{Ce},{ }^{152} \mathrm{Eu},{ }^{154} \mathrm{Eu}$, ${ }^{153} \mathrm{Gd},{ }^{160} \mathrm{~Tb},{ }^{161} \mathrm{~Tb},{ }^{170} \mathrm{Tm},{ }^{169} \mathrm{Yb},{ }^{172} \mathrm{Hf},{ }^{182} \mathrm{Ta},{ }^{185} \mathrm{Os}$, ${ }^{192} \mathrm{Ir},{ }^{198} \mathrm{Au},{ }^{199} \mathrm{Au},{ }^{203} \mathrm{Hg},{ }^{210} \mathrm{~Pb},{ }^{207} \mathrm{Bi},{ }^{228} \mathrm{Th},{ }^{239} \mathrm{~Np}$, ${ }^{241} \mathrm{Am}$, and ${ }^{243} \mathrm{Am}$. One of the factors that contributes to the production of these isotopes inside the magnets is the

TABLE I. Parameters of the HEBT elements.

| Element | Length (mm) | Aperture (mm) | Strength (T) |
| :--- | :---: | :---: | :---: |
| Dipoles | 1570 | $40 \times 80$ (total gap) | 1.47 |
| Quadrupoles | 400 or 800 | 40 (radius) | 0.48 |
| Octupoles | 800 | 25 (radius) | 0.35 |

direct impact of the proton beam. When proton beam losses occur, the protons interact with the beam pipe producing hadronic and electromagnetic showers. Also when the protons interact with the magnets they will produce additional secondaries. We looked at all the secondaries that produce one of the isotopes of interest and Fig. 5 shows the number of each secondary particle that creates these isotopes per lost proton. It can be seen that even when considering only the direct proton beam interactions, the main contribution to the induced activation comes from the secondary neutrons produced either in the beam pipe or in the magnets. In fact the contribution of the secondary neutrons is about 3 times higher than the contribution of the lost protons.

The radioactive decay processes have been added to the standard physics list provided by GEANT4, such that the production rate takes into account not only the isotopes produced by the direct impact of incoming protons and by the backscattered neutrons on various accelerator components, but also the production of these isotopes following the decay of other isotopes produced in these interactions.

Apart from the geometry of the magnets shown in Fig. 1, the material composition also has to be known. For simplicity, the magnets were assumed in the simulations to be made of a homogeneous mixture of Fe and Cu in equal proportions. It is still an open question whether the magnets are suspended or supported and surely there will be some mechanical stainless steel structure holding each magnet in place. The radionuclide inventory present in the HEBT-S3 octupoles (OCT1 and OCT2) is presented in Fig. 6. The radioactive nuclide inventory of each of the HEBT-S3 quadrupoles is presented in Fig. 7. The isotope yield shown in these figures represents the total amount of a particular isotope given by direct proton interactions, backscattered neutrons, and also by secondary neutron interactions and other secondary interactions. There is a higher amount of isotopes produced by direct proton impact and its secondaries in the quadrupoles placed at the beginning of the HEBT-S3 beam line while for the quadrupoles placed in the close proximity of the target the isotopes produced by backscattered neutrons are predominant. In all octupoles and quadrupoles, the isotopes ${ }^{51} \mathrm{Cr}$ and ${ }^{54} \mathrm{Mn}$ are predominant although there are also significant amounts of isotopes ${ }^{57} \mathrm{Co},{ }^{58} \mathrm{Co}$, and ${ }^{60} \mathrm{Co}$.

Eventual beam tails formed in the HEBT will be caught by a set of fixed collimators just before the proton beam window in order to protect the edges of both the proton beam window and the main target. These collimators are copper cylinders of 1 m length with a diameter of 2 m and they have a rectangular opening of $(16 \times 6) \mathrm{cm}$. The copper collimators are coated with tungsten. Being placed in front of the target monolith, the collimators are the most radioactive elements in the beam line and therefore they will be exposed to a high flux of backstream neutrons coming from the target and will be at high risk due to the


FIG. 4. The magnets and collimator design for the HEBT-S3 beam line. For magnets, the Cu parts are shown in red while the Fe parts are shown in green. For the collimator, the Cu parts are shown in yellow while the W parts are shown in grey.
their position in the beam line. The radionuclide yields produced inside the Cu core and the W coating are shown separately in different colors in Fig. 8. The isotope yield is expressed per incoming proton and isotopes produced in all four mechanisms (by direct proton impact, by backscattered neutrons, by secondary neutron interactions, and other secondary interactions) are shown separately. Examination of the isotope chart shows a higher yield of all isotopes found in the magnets $\left({ }^{7} \mathrm{Be},{ }^{46} \mathrm{Sc},{ }^{44} \mathrm{Ti},{ }^{51} \mathrm{Cr},{ }^{54} \mathrm{Mn}\right.$, ${ }^{59} \mathrm{Fe},{ }^{56} \mathrm{Co},{ }^{57} \mathrm{Co},{ }^{58} \mathrm{Co},{ }^{60} \mathrm{Co},{ }^{65} \mathrm{Zn}$ ). There is a large range of other isotopes produced in a significant amount (i.e., ${ }^{109} \mathrm{Cd},{ }^{111} \mathrm{In},{ }^{113} \mathrm{Sn},{ }^{172} \mathrm{Hf},{ }^{182} \mathrm{Ta}$ ).

The isotopes found in the magnets and in the collimator are listed in Table II together with their half-lives, decay mode, and the level of gamma radiation they emit expressed as the gamma dose rate at 1 m when the decay rate is 1 Bq [19]. The radioactive isotopes may decay by beta ( $\beta^{-}$) emission or when proton rich isotopes are produced by positron $\left(\beta^{+}\right)$emission. In some cases the nucleus decays by capturing an orbiting electron. The


FIG. 5. Contribution of each particle type to the magnets activation due to proton beam losses in the beam pipe.
daughter nucleus resulting from decay may deexcite by emitting gamma radiation. It can be seen from the table that a number of isotopes like ${ }^{241} \mathrm{Am},{ }^{243} \mathrm{Am},{ }^{133} \mathrm{Ba},{ }^{207} \mathrm{Bi}$, ${ }^{109} \mathrm{Cd},{ }^{134} \mathrm{Cs},{ }^{137} \mathrm{Cs},{ }^{152} \mathrm{Eu},{ }^{154} \mathrm{Eu},{ }^{44} \mathrm{Ti},{ }^{60} \mathrm{Co},{ }^{94} \mathrm{Nb},{ }^{210} \mathrm{~Pb}$,


FIG. 6. The radionuclide inventory for the HEBT-S3 octupoles: (a) octupole OCT1, (b) octupole OCT2. The colors represent isotopes produced by backscattered neutrons from the target (blue), isotopes produced in direct proton interactions (dark blue), isotopes produced in secondary neutrons interactions (red), isotopes produced in other secondaries interactions (green).


FIG. 7. The radionuclide inventory for the HEBT-S3 quadrupoles: (a) quadrupole QP20, (b) quadrupole QP21, (c) quadrupole QP22, (d) quadrupole QP23, (e) quadrupole QP24, (f) quadrupole QP25, (g) quadrupole QP26, and (h) quadrupole QP27. The isotope yield represents the total amount of particular isotopes found in the magnets. The colors represent isotopes produced by backscattered neutrons from the target (blue), isotopes produced in direct proton interactions (dark blue), isotopes produced in secondary neutrons interactions (red), and isotopes produced in other secondaries interactions (green).


FIG. 8. The radionuclide inventory for the HEBT-S3 collimator: (a) isotopes produced due to direct proton interactions, (b) isotopes produced due to secondary neutron interactions, (c) isotopes produced due to other secondary interactions, and (d) isotopes produced due to backscattered neutrons from the target. The isotopes produced in the copper yield of the collimator and the tungsten coating are shown in different colors: red (copper), green (tungsten).
${ }^{106} \mathrm{Ru},{ }^{125} \mathrm{Sb}$, and ${ }^{172} \mathrm{Hf}$ pose significant risks for radiation protection as their half-life is more than one year.

## B. Activation of the magnets and collimators

The number of isotopes produced $N_{\text {iso }}$ inside the HEBTS3 accelerator components gives the production rate for each isotope. This production rate takes into account the direct impact of the proton beam on each accelerator component due to the proton beam spatial distribution, the interactions of the backscattered neutrons, and the isotopes production following the decay of other isotopes produced in these interactions. All these processes have been considered when counting $N_{\text {iso }}$ in the simulations output. The activation formula given in Eq. (4) offers the general theoretical approach in estimating the induced activity and relies on the knowledge and use of the cross section $\sigma_{T, \nu}$ of the inelastic process in which the target nucleus $T$ turns into the isotope $\nu$. However, in the GEANT4 code these cross sections are included in the database and the code can predict directly the number $N_{\text {iso }}$ of isotopes produced in these processes such that the user does not have to use each inelastic process cross section in
order to calculate the induced activity. The production rate for each isotope $N_{\text {iso }} \Delta t$ is simulated taking into account direct proton impact, backscatted neutrons, secondary interactions, and decay of other isotopes produced in these interactions.

GEANT4 calculates the isotope production using beside the particle induced production also the full Bateman solution considering the breeding of radioactive decay products. The majority of the radionuclide production is coming from the particle impact induced activation and only a smaller fraction from decay production. Therefore considering the production rate for radioactive isotope $\nu$ to be constant in time is a good estimate for the real situation.

The production rate for a radioactive isotope is given by

$$
\begin{equation*}
\frac{d N_{\nu}^{\mathrm{prod}}}{d t}=\frac{N_{\mathrm{iso}}}{\Delta t}=\frac{N_{\mathrm{iso}} I}{N_{p} e} \tag{5}
\end{equation*}
$$

where $N_{\text {iso }}$ is the number of isotopes produced, $I$ is the proton beam current from the accelerator, $N_{p}$ is the number of incident protons, and $e$ is the proton electric charge. Once the radioactive nuclei have been produced, their

TABLE II. List of radioactive isotopes produced inside the magnets and collimators by spallation reactions that contribute to the induced activity. The dose rates at 1 m are per Bq of the active isotope considering a point source.

| Isotope | (fSv $h^{-1} \mathrm{~Bq}^{-1}$ ) | Half-life | Isotope | (fSv $h^{-1} \mathrm{~Bq}^{-1}$ ) | Half-life | Isotope | (fSv $h^{-1} \mathrm{~Bq}^{-1}$ ) | Half-life |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ${ }^{241}$ Am | 84.79 | 432 y | ${ }^{137} \mathbf{C s}$ | 103.2 | 30.2 y | ${ }^{124} \mathbf{S b}$ | 288.3 | 60.2 d |
| ${ }^{243} \mathbf{A m}$ | 84.56 | 7388 y | ${ }^{152} \mathbf{E u}$ | 201.2 | 13.5 y | ${ }^{125} \mathbf{S b}$ | 102.8 | 2.7 y |
| ${ }^{198} \mathbf{A u}$ | 78.82 | 2.7 d | ${ }^{154} \mathbf{E u}$ | 204.2 | 8.6 y | ${ }^{46} \mathbf{S c}$ | 315.5 | 83.7 d |
| ${ }^{199} \mathbf{A u}$ | 18.66 | 3.1 d | ${ }^{59} \mathbf{F e}$ | 178.9 | 44.5 d | ${ }^{75} \mathbf{S e}$ | 232.3 | 119.7 d |
| ${ }^{133} \mathbf{B a}$ | 123.1 | 10.5 y | ${ }^{153} \mathbf{G d}$ | 46.59 | 240.4 d | ${ }^{113} \mathbf{S n}$ | 48.44 | 115 d |
| ${ }^{7} \mathrm{Be}$ | 9.292 | 532 d | ${ }^{172} \mathbf{H f}$ | 33.57 | 1.87 y | ${ }^{125} \mathbf{S n}$ | 46.74 | 9.64 d |
| ${ }^{207} \mathbf{B i}$ | 360.3 | 32.9 y | ${ }^{203} \mathbf{H g}$ | 68.41 | 46.6 d | ${ }^{85} \mathbf{S r}$ | 205.2 | 64.85 d |
| ${ }^{109} \mathrm{Cd}$ | 49.83 | 1.2 y | ${ }^{125}$ I | 74.32 | 59.4 | ${ }^{182} \mathbf{T a}$ | 208.6 | 114.4 d |
| ${ }^{139} \mathrm{Ce}$ | 55.54 | 137.6 d | ${ }^{111}$ In | 135.6 | 2.8 d | ${ }^{160} \mathbf{T b}$ | 178.8 | 72.3 d |
| ${ }^{141} \mathrm{Ce}$ | 19.79 | 32.5 d | ${ }^{192} \mathbf{I r}$ | 159.9 | 73.8 d | ${ }^{161} \mathbf{T b}$ | 18 | 6.9 d |
| ${ }^{144} \mathrm{Ce}$ | 6.30 | 285 d | ${ }^{54} \mathbf{M n}$ | 138.2 | 312 d | ${ }^{44} \mathbf{T i}$ | 39.09 | 60.2 y |
| ${ }^{56} \mathrm{Co}$ | 520.5 | 77.2 d | ${ }^{94} \mathrm{Nb}$ | 264.8 | 2.03 E 4 y | ${ }^{170} \mathbf{T m}$ | 1.673 | 128.6 d |
| ${ }^{57} \mathrm{Co}$ | 40.87 | 271.7 d | ${ }^{95} \mathrm{Nb}$ | 129.8 | 35 d | ${ }^{88} \mathbf{Y}$ | 481.9 | 106.65 d |
| ${ }^{58} \mathrm{Co}$ | 165.9 | 70.8 d | ${ }^{239} \mathbf{N p}$ | 138.6 | 2.35 d | ${ }^{169} \mathbf{Y b}$ | 88.37 | 32 d |
| ${ }^{60} \mathrm{Co}$ | 370.3 | 5.3 y | ${ }^{185} \mathrm{Os}$ | 131 | 93.6 d | ${ }^{65} \mathbf{Z n}$ | 89.24 | 244 d |
| ${ }^{51} \mathrm{Cr}$ | 6.32 | 27.7 d | ${ }^{210} \mathbf{P b}$ | 68.01 | 22.3 y | ${ }^{95} \mathbf{Z r}$ | 125.8 | 64 d |
| ${ }^{132} \mathbf{C s}$ | 155.6 | 6.5 d | ${ }^{84} \mathbf{R b}$ | 232.6 | 32.7 d |  |  |  |
| ${ }^{134} \mathrm{Cs}$ | 270.1 | 2 y | ${ }^{106} \mathbf{R u}$ | 365 | 1.02 y |  |  |  |

number decays exponentially with time. During the beamon period, the time evolution can be obtained by combining the production and decay rates:

$$
\begin{equation*}
\frac{d N_{\nu}}{d t}(t)=\frac{N_{\mathrm{iso}} I}{N_{p} e}-\lambda_{\nu} N_{\nu}(t) \tag{6}
\end{equation*}
$$

We consider first the boundary condition of having at $t=0$ a nonradioactive material. The solution of Eq. (6) gives the number of isotopes at any time $t$ during the beam exposure:

$$
\begin{equation*}
N_{\nu}(t)=\frac{N_{\mathrm{iso}} I}{N_{p} e \lambda_{\nu}}\left[1-\exp \left(-\lambda_{\nu} t\right)\right] \tag{7}
\end{equation*}
$$

However, after the beam is switched off, following a continuous exposure for a given time $t_{1}$, the number of isotopes after a time $t$, which includes both the beam-on period $t_{1}$ as well as the beam-off period, is given by

$$
\begin{equation*}
N_{\nu}(t)=\frac{N_{\mathrm{iso}} I}{N_{p} e \lambda_{\nu}}\left[1-\exp \left(-\lambda_{\nu} t_{1}\right)\right] \exp \left[-\lambda_{\nu}\left(t-t_{1}\right)\right] \tag{8}
\end{equation*}
$$

The induced activity for an accelerator element given by one particular isotope is given by Eq. (9):

$$
\begin{equation*}
A_{\nu}(t)=\lambda_{\nu} N_{\nu}(t) \tag{9}
\end{equation*}
$$

Using Eqs. (8) and (9), the activity of each isotope produced is given by Eq. (10) at any given time $t$ which includes the beam-on time $t_{1}$ and the beam-off period (during the beam-on period, $t=t_{1}$ ),

$$
\begin{equation*}
A_{\nu}(t)=\frac{N_{\text {iso }} I}{N_{p} e}\left[1-\exp \left(-\lambda_{\nu} t_{1}\right)\right] \exp \left[-\lambda_{\nu}\left(t-t_{1}\right)\right] \tag{10}
\end{equation*}
$$

The total induced activity inside each accelerator component is given by Eq. (11),

$$
\begin{equation*}
A(t)=\sum_{\nu} A_{\nu}(t) \tag{11}
\end{equation*}
$$

Short-lived isotopes decay rapidly and their presence is negligible after long cooling times. Considering a beam exposure of 365 days, and using Eqs. (10) and (11), the induced activity inside the magnets was calculated after a cooling-off period of 30 days considering a beam loss of $1 \mathrm{~W} / \mathrm{m}$ in the magnets and 20 kW at the collimator for a maximum beam current of 62.5 mA and a minimum beam current of 6.25 mA . The induced activity inside the quadrupoles after this period is shown in Fig. 9, inside the octupoles in Fig. 10, and inside the collimator is shown in Fig. 11. As expected, the worst affected beam line component is the collimator in front of the target monolith. There is a factor of approximately 100 difference in the total induced activity inside the quadrupoles, octupoles, and collimator.

The results so far have shown the activity after 365 days of exposure and 30 days of cooling off. However, Eq. (10) can be used to calculate the activity at any time $t$, both during the beam exposure time $t_{1}$ for which $\mathrm{t}=t_{1}$, as well as during the cooling-off period. The activity inside the accelerator components was calculated at various intervals during the 365 days of beam exposure and after the accelerator shut down. For each beam line element the total amount of Bq for each isotope was converted using


FIG. 9. Activity for the HEBT-S3 quadrupoles: (a) quadrupole QP20, (b) quadrupole QP21, (c) quadrupoles QP22, (d) quadrupole QP23, (e) quadrupoles QP24, (f) quadrupole QP25, (g) quadrupoles QP26, and (h) quadrupole QP27.


FIG. 10. Induced activity for the HEBT-S3 octupoles: (a) octupole OCT1 and (b) octupole OCT2.


FIG. 11. Total induced activity inside the HEBT-S3 collimator. Isotopes produced in the copper yield of the collimator are shown in red, isotopes produced in the tungsten coating of the collimator are shown in green.


FIG. 12. Dose rates calculated at 1 m from the element surface for the HEBT-S3 magnets: (a) quadrupoles and (b) octupoles.


FIG. 13. Time evolution of dose rates calculated at 1 m from the element surface for 365 days of exposure for various elements.
the Bq-to-dose conversion factors into dose (Table II). Since these conversion factors are for point sources, they do not account for self-absorption of the materials. Then the individual doses contributions from various isotopes were added up into a final result. The results are shown in Fig. 12 for various magnets and in Fig. 13 for the elements that are closest to the spallation target. The curves for different elements have been scaled for a direct and better comparison. The generic curves can obviously be divided into two separate parts corresponding to activation (data points before the end of irradiation) and decay (data points after the end of irradiation).

## V. CONCLUSION

The primary concern in building high-power accelerators is to minimize the beam losses because they produce first damage to the whole accelerator system due to the high beam energy deposition, and second they produce induced radioactivity. Predictions of the residual activity in the accelerator components is important for allowing access of maintenance personal after the beam is turned off. In this paper we presented a generic study of induced activation in accelerator components, and the same methodology can be applied to any future study of element activation from other accelerator facilities around the world. In this study we discussed the methodology of estimating the induced radioactivity in magnets and collimators of the ESS HEBTS3 beam line method, which relies on the computation of the production rate of each isotope of interest. All possible factors have been considered: the direct interaction of the proton beam with the elements, the interaction of the backscattered neutrons from the target, and the isotope production as a result of the decay of other isotopes produced in these interactions. The main factor proved to be the backscattered neutrons, the particle collimator placed in front of the target being the worst affected. The induced activity was calculated both during and after a

365-day beam exposure, reaching a maximum of $100 \mathrm{~Sv} / \mathrm{h}$ for this particular beam element.
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