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Production of theranostic Tb isotopes: electromagnetic isotope separation before or after irradiation ?

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Terbium has a quadruplet of so-called theranostic isotopes useful for the preparation of radiopharmaceuticals: ^{152}Tb (PET imaging), ^{155}Tb (SPECT imaging), ^{161}Tb (beta- therapy) and ^{149}Tb (alpha therapy). All isotopes belong to the same element, thus assuring identical pharmacokinetics, an essential requirement for theranostics. $^{149},^{152},^{155}\text{Tb}$ with high radioisotopic purity is so far only available from spallation of Ta targets combined with on-line mass separation at CERN-ISOLDE or TRIUMF-ISAC.

Additional production at cyclotrons is urgently required to satisfy the great demand for medical applications. These isotopes could in principle also be produced by $^{155}\text{Gd}(p,n)^{155}\text{Tb}$, $^{152}\text{Gd}(p,n)^{152}\text{Tb}$ and $^{152}\text{Gd}(p,4n)^{149}\text{Tb}$ reactions respectively, provided targets of sufficient isotopic enrichment become available. Commercially available ^{152}Gd reaches only 30% enrichment, but $>90\%$ enrichment is required to minimize co-production of longer-lived Tb isotopes in (p,n) reactions.

We present a demo experiment performed at the tandem accelerator of the MLL Garching where ^{152}Tb was produced by irradiating a unique ion-implanted ^{152}Gd target ($>99\%$ enriched) with 8 MeV and 12 MeV protons respectively. At these energies only ^{152}Tb was observed while upper limits are derived for co-production of other Tb isotopes. This radioisotopic purity would enable direct use for human applications, only requiring a chemical Tb/Gd separation from the target material.

We will discuss prospects to efficiently separate more ^{152}Gd and ^{155}Gd with the SIDONIE mass separator at CSNSM Orsay and thus prepare cyclotron targets suited for high current irradiations.

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