

A Unique Matched Quadruplet of Terbium Radioisotopes for PET and SPECT and for alpha- and beta-Radionuclide Therapy: An in vivo Proof of Concept Study with a New Receptor-Targeted Folate Derivative

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Background: Compared to other lanthanides, terbium (Tb) comprises clinically attractive radionuclides such as ^{152}Tb and ^{155}Tb with suitable decay properties for PET and SPECT imaging and ^{149}Tb and ^{161}Tb suitable for targeted alpha- and beta-radionuclide therapy. To evaluate these 4 radioisotopes we employed the folic acid/folate receptor (FR)-targeting strategy which has been thoroughly investigated over years by our group. Thus, the goal of this proof-of-concept study was to produce all 4 Tb-radioisotopes and assess their diagnostic and therapeutic features in vivo when labeled with a folate-based targeting agent.

Methods: ^{161}Tb was produced by irradiation of ^{160}Gd targets with neutrons at Paul Scherrer Institute or Institut Laue-Langevin. After neutron capture, the short-lived ^{161}Gd decays to ^{161}Tb . ^{149}Tb , ^{152}Tb , and ^{155}Tb were produced by proton-induced spallation of tantalum targets, followed by an online isotope separation process at ISOLDE/CERN. The isotopes were purified by means of cation exchange chromatography at PSI. A long-circulating DOTA-folate conjugate (cm09) was radiolabeled with all four Tb-radioisotopes under standard labeling conditions. Biodistribution studies were performed over seven days using ^{161}Tb -cm09. Diagnostic PET/CT (^{152}Tb -cm09) and SPECT/CT (^{155}Tb -cm09 and ^{161}Tb -cm09) and therapy experiments with ^{149}Tb -cm09 and ^{161}Tb -cm09 were performed in KB tumor bearing mice.

Results: Carrier-free Tb-radioisotopes were obtained after purification, with activities ranging from approximately 6 MBq (for ^{149}Tb) to approximately 15 GBq (for ^{161}Tb). The radiolabeling of cm09 was achieved in >96% radiochemical yield for all 4 Tb-radioisotopes. The tissue distribution of ^{161}Tb -cm09 resulted in a high tumor uptake (~20% ID/g, 24 h p.i.) which was retained over several days. PET/CT and SPECT/CT studies allowed excellent tumor visualization in mice even 24 h after injection of the ^{152}Tb -cm09 and ^{155}Tb -cm09. Targeted radionuclide therapy studies performed with ^{149}Tb -cm09 and ^{161}Tb -cm09 revealed a significant inhibition of tumor growth and a prolonged survival (>1.7-fold and >2-fold) of treated mice compared to untreated controls.

Conclusions: This is the first comprehensive study with all 4 Tb-radioisotopes using one and the same targeting agent (Müller et al. J Nucl Med 2012, in press). Excellent in vivo tumor visualization was enabled with ^{152}Tb -cm09 and ^{155}Tb -cm09 using PET/CT and SPECT/CT. Thus, ^{152}Tb and ^{155}Tb could become ideal diagnostic matches for their therapeutic counterparts ^{149}Tb and ^{161}Tb providing identical chemical properties. Regarding the therapeutic approach, promising results were obtained after application of ^{149}Tb -cm09 and ^{161}Tb -cm09 with respect to a prolonged overall survival of the mice and an effective tumor growth inhibition.

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