Gold Coated Lanthanide Phosphate Nanoparticles for Targeted Alpha Generator Radiotherapy

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Abstract

Targeted radiotherapies maximize cytotoxicity to cancer cells. In vivo α-generator targeted radiotherapies can deliver multiple α particles to a receptor site dramatically amplifying the radiation dose delivered to the target. The major challenge with α-generator radiotherapies is that traditional chelating moieties are unable to sequester the radioactive daughters in the bioconjugate which is critical to minimize toxicity to healthy, non-target tissue. The recoil energy of the 225Ac daughters following α decay will sever any metal-ligand bond used to form the bioconjugate. This work demonstrates that an engineered multilayered nanoparticle-antibody conjugate can deliver multiple α radiations and contain the decay daughters of 225Ac while targeting biologically relevant receptors in a female BALB/c mouse model. These multi-shell nanoparticles combine the radiation resistance of lanthanide phosphate to contain 225Ac and its radioactive decay daughters, the magnetic properties of gadolinium phosphate for easy separation, and established gold chemistry for attachment of targeting moieties.

Introduction

Beta emitting radionuclides have found widespread use in cancer therapy. A major advance in nuclear medicine was the development of targeted endo-radiotherapies with two targeted radiotherapy agents approved for clinical use. BEXXAR® 131I, labeled with 131I, is used to treat follicular lymphoma while Zevalin® 90Y, containing 90Y, is used for treatment of B cell non-Hodgkins lymphoma [1–2]. Other targeted radiotherapy agents labeled with β− emitters 131I, 90Y, 177Lu, and 188Re are showing promise in ongoing clinical trials [3–4]. One of the challenges associated with β− emitting targeted radionuclide therapies is, however, the inherent toxicity from the death of normal, healthy cells resulting from the crossfire radiation damage from the relatively long ranges of the β− particles in tissue [5]. For example, β− particles from 177Lu (β_{max} \approx 0.5 MeV) have a range of 1.5 mm in tissue and β− particles from 90Y (β_{max} = 2.3 MeV) deposit their energy over a range of 12 mm. Targeted radiotherapies based on α particles are a promising alternative to β− particles because the α particles deposit all of their energy within a few cell diameters (50–100 μm). Because of their much shorter range, targeted α-radiotherapy agents have great potential for application to small, disseminated tumors and micro metastases and treatment of hematological malignancies consisting of individual, circulating neoplastic cells [6]. Compared with β− particles, α particles provide a very high relative biological effectiveness, killing more cells with less radioactivity. The high linear energy transfer of α particles induces significantly more DNA double strand breaks than β− particles [7]. Also, the biological effectiveness of α particles does not depend upon hypoxia or cell cycle considerations [8–9]. Most α emitters also have a relatively low γ-ray component in their decay allowing for outpatient treatments and lower radiation doses to nuclear medicine personnel [10].

A number of targeted alpha therapy (TAT) agents based on the single alpha emitting radionuclides 211At (t_{1/2} = 7.2 h), 212Bi (t_{1/2} = 60 min), 212Pb (t_{1/2} = 10.6 h), and 212Bi (t_{1/2} = 61 min) have been developed and are showing promise in pre-clinical and clinical trials [11]. The radiotherapeutic efficacy of TAT could, however, be further enhanced by use of in vivo α-generator radionuclides like 225Ac, which emits four α particles in its decay chain (Figure 1). The median lethal dose of 225Ac constructs is one to two orders of magnitude lower than the LD50 values for the corresponding single α emitting 212Bi labeled antibodies in vivo with a number of cancer cell types [12]. Moreover, the longer half-life of 225Ac (t_{1/2} = 10 d) reduces activity loss during radiopharmaceutical synthesis and allows greater time for localization of antibodies to receptor sites. Despite these advantages, there is a distinct challenge associated with targeted in vivo α-generator radiotherapy. If the α-emitting daughter products in the 225Ac decay chain are not sequestered at the target site, they can migrate and deliver a potentially toxic dose.
to non-target tissue [11]. The recoil energy of the $^{225}$Ac daughters following alpha decay (100 keV) will sever any metal-ligand bond used to form the bioconjugate, releasing the daughter radionuclides from the targeting agent. Renal toxicity is currently the dose-limiting factor in clinical use of $^{225}$Ac. In the recent work of Schwartz et al., almost 80% of the absorbed dose to the renal medulla was delivered by free $^{213}$Bi when using a metal-ligand bioconjugate to deliver $^{225}$Ac in a mouse model [13]. Metal-ligand bioconjugates fail to sequester the daughter products of $^{225}$Ac ($^{221}$Fr, $^{217}$At, and $^{213}$Bi) and the released $^{213}$Bi accumulates in the kidney [14–15]. An alternative strategy to this challenge, incorporating $^{225}$Ac in engineered liposomes, was found to retain less than 10% of the $^{213}$Bi activity from the decay of $^{225}$Ac in vitro [16].

The in vivo $\alpha$ generator $^{223}$Ra, which also emits four alpha particles in its decay chain, is an effective treatment for metastatic bone cancer [17]. Radium-223 chloride has been granted Fast Track designation by the U.S. Food and Drug Administration for the treatment of hormone-refractory prostate cancer in patients with bone metastases [18]. It is effective in this case because radium is a calcium mimic with a high affinity for bone tissue and the daughter products either have short half-lives or have a high affinity for bone ($^{211}$Pb, $t_{1/2}$ = 36 m). Translation of in vivo $\alpha$ generators to anything besides metastatic bone cancer with a calcium mimic will require a different mechanism for both delivering and retaining the radioactive daughters in the target tissue.

In this work, we demonstrate that a multilayered nanoparticle (NP) can contain the recoiling daughters of the in vivo $\alpha$ generator, and when coupled to a targeting antibody, can bind to biologically relevant receptors and deliver multiple $\alpha$ particles to each receptor site. While the range of the recoiling daughters is less than the diameter of the layered NPs, the $\alpha$-particles lose less than 0.2% of their energy as they exit from the center of the NPs. The layered NPs consist of $^{225}$Ac-doped $\{\text{La}_{0.5}\text{Gd}_{0.5}\}_{\text{PO}_4}\$@GdPO$_4$@Au. A schematic of the layered NP design is illustrated in Figure 2. These multi-shell particles combine the radiation resistance of crystalline lanthanide phosphate [19], [20–21] to encapsulate and contain atoms of the therapeutic radionuclide $^{225}$Ac and its radioactive daughters, the magnetic properties of gadolinium phosphate NPs, and established surface chemistry of gold NPs for attachment of targeting agents [22].

Nanoparticles offer a number of favorable properties with regard to drug delivery. The ability to hold multiple copies of a therapeutic or imaging moiety provides the ability to generate efficacious results even against targets with low receptor numbers in vivo [23]. Multi-functional, layered NPs allow for synergistic combinations of properties exhibited by the individual NPs including containment, purification and conjugation. Moreover, in the system described in this work, GdPO$_4$ could function as a magnetic resonance imaging (MRI) contrast agent [24] and Au can be doped with the single photon emission computed tomography (SPECT) radionuclide $^{199}$Au for $\gamma$-ray imaging [25].

Results and Discussion

$\{\text{La}_{0.5}\text{Gd}_{0.5}\}_{\text{PO}_4}$$^{225}$Ac core particles were synthesized by hydrolysis of sodium tripolyphosphate (Na-TPP) in the presence of equimolar mixtures of La and Gd salts with $^{225}$Ac present at the tracer level. Nanoparticles of diameter ca. 4 nm precipitated out of solution after heating for 3 hours at 90°C. In order to improve retention of the decay daughters, four additional shells of pure GdPO$_4$ were added, each by dispersing the precipitated NPs in a solution of Gd$^{3+}$ and Na-TPP and heating for an additional 3 hours. Gold was then added to the NPs by reduction of NaAuCl$_4$ with sodium citrate.

Rietveld refinement of x-ray diffraction (XRD) patterns indicated that LaPO$_4$ NPs exhibited the rhabdophane phase consistent with the description of Buisset et al. [26]. However, the $\{\text{La}_{0.5}\text{Gd}_{0.5}\}_{\text{PO}_4}$, $\{\text{La}_{0.25}\text{Gd}_{0.75}\}_{\text{PO}_4}$, and GdPO$_4$ systems crystallized in the anhydrous monazite phase [27]. The monazite phase for LaPo$_4$ NPs was previously observed for crystalline...
synthesis in organic solvents [28]. The XRD measurements yielded NP grain sizes of 4.04 nm for LaPO₄, 2.79 nm for \{La₀.₅Gd₀.₅\}PO₄, 2.91 nm for \{La₀.₂₅Gd₀.₇₅\}PO₄ and 3.11 nm for GdPO₄. Size estimates of the \{La₀.₅Gd₀.₅\}PO₄ NPs from transmission electron microscopy (TEM) images (Figure 3) match the grain sizes predicted by XRD, indicating that the core particles were a single crystal phase. Neutron activation analysis of magnetically separated \{La₀.₅Gd₀.₅\}PO₄ core NPs gives a La to Gd mole ratio of 1.11±0.03. Pure LaPO₄ and pure GdPO₄ exhibited larger grain sizes than their mixed counterparts.

Addition of GdPO₄ shells to the core \{La₀.₅Gd₀.₅\}PO₄ NP causes epitaxial growth of the particle. Mean diameters increase sequentially with each shell addition (Table 1). Addition of four GdPO₄ shells to the core \{La₀.₅Gd₀.₅\}PO₄ produces 22 nm diameter NPs and addition of an outer gold layer increases the particle diameter to 27 nm. Electron energy loss spectroscopy (EELS)-TEM images of the NPs are shown in Figure 4.

Gold coated NPs with four epitaxially added GdPO₄ shells were further characterized by dynamic light scattering. Hydrodynamic diameters and zeta potentials are shown in Table 2. An increase of the hydrodynamic diameter on addition of polyethylene glycol (PEG) and antibody is common for NPs. The highly negative zeta potentials should lead to stability in water which was confirmed by monitoring changes in the UV-Vis spectrum of the particles over a 1 month period in both 18 MΩ water and saline solution. No shift was observed in the plasmon resonance over this time period.

Nanoparticles with GdPO₄ shells followed by Au coating dramatically increased radioactive daughter retention in vitro compared with previously published results for core-only lanthanum phosphate NPs [28]. Adding 2 shells increased retention of the \(^{221}\)Fr daughter from 50% for the LaPO₄ core to 70%. With four shells of GdPO₄, the initial retention of the \(^{221}\)Fr daughter was 98%. Daughter retention decreased by roughly 2% per day over the course of a week, and stabilized at 88%. Further, the presence of the Au/4 GdPO₄ shells increased the retention of the \(^{225}\)Ac parent itself by roughly an order of magnitude. Over the course of 3 weeks, the multi-layered particles retained greater than 99.99% of the \(^{225}\)Ac parent radionuclide. Particles with more than 4 shells of GdPO₄ settled out of solution rapidly and were difficult to manipulate. Monitoring the plasmon resonance indicated that the multi-layered particles remained stable towards aggregation in PBS over the course of one month.

For in vivo biodistribution testing, the NPs were conjugated to the mAb 201b monoclonal antibody via a lipoic acid-PEG₁₂-COOH linker [29]. MAb 201b targets thrombomodulin receptors which are highly expressed in lung endothelium. The antibody quickly localizes to its vascular target and clears from circulation with a half-life of 40 hours [30]. 3-sulfo-N-hydroxysuccinimide (sulfo-NHS) and 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC) activated the carboxylate of the PEG for coupling to amine groups on the antibody, leading to the formation of an amide bond. The reaction was quenched with glycine and conjugates were purified by centrifugation. The conjugated NPs were redispersed in phosphate buffered saline (PBS) containing bovine serum albumin (BSA). The antibody conjugation process is summarized in Figure 5.

**Table 1. Growth of NP diameter as a function of shell addition as measured by TEM.**

| Particle System                  | Diameter (nm) |
|----------------------------------|---------------|
| \{La₀.₅Gd₀.₅\}PO₄ Core           | 5.0±1.5       |
| \{La₀.₅Gd₀.₅\}PO₄@1 shell GdPO₄  | 7.8±2.8       |
| \{La₀.₅Gd₀.₅\}PO₄@2 shells GdPO₄ | 9.9±2.6       |
| \{La₀.₅Gd₀.₅\}PO₄@3 shells GdPO₄ | 13.3±1.8      |
| \{La₀.₅Gd₀.₅\}PO₄@4 shells GdPO₄ | 22.4±7.7      |
| \{La₀.₅Gd₀.₅\}PO₄@4 shells GdPO₄@Au | 26.8±4.9     |

Figure 3. TEM image of \(\{\text{La}_{0.5}\text{Gd}_{0.5}\}\)PO₄ core NPs.
doi:10.1371/journal.pone.0054531.g003

Figure 4. TEM of a characteristic cluster of NPs. EELS analysis indicates the presence of La, Gd, and Au in all particles in the cluster. doi:10.1371/journal.pone.0054531.g004

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Renal toxicity concerns associated with 213Bi relocation to the tissue (1 h, 72% (After 24 hours, this number further decreased to 1.5% from the injected dose migrated to kidney tissues after 1 hour.

PLOS ONE | www.plosone.org 4 January 2013 | Volume 8 | Issue 1 | e54531

Retention of 211Bi, from the decay of 225Ac in the a-generator NPs, was 69% ± 3% in lung tissue after 1 hour and increased to 84% ± 3% after 24 hours. Similar 211Bi retention values were observed in liver (1 h, 81% ± 4%; 24 h, 92% ± 1%) and spleen tissue (1 h, 72% ± 3%; 24 h, 82% ± 16%). Despite the widespread renal toxicity concerns associated with 211Bi relocation to the kidney from 225Ac a-generator therapies, only 2.8% of the 211Bi from the injected dose migrated to kidney tissues after 1 hour. After 24 hours, this number further decreased to 1.5%.

A larger dose (ca. 80 μCi/animal) of 225Ac NPs was imaged using CT/SPECT of the 218Fr γ-ray (218 keV, 11.6%). Mice injected with this larger dose were sacrificed 1 hour post-injection and imaged 3 hours post-sacrifice to allow the daughter products of 225Ac to reach their equilibrium activities. The CT/SPECT images (Figure 7) clearly show large uptake in the lung for the {La0.5Gd0.5}(225Ac)PO4@GdPO4@Au-mAb-201b NPs which is in agreement with the biodistribution data. When competed with unconjugated mAb 201b antibody, the images showed high uptake in the lung. If the antibody conjugated NPs cannot bind their in vivo target, they are cleared from circulation via the reticuloendothelial system. Finally, PEG coated NPs without antibody also show high uptake in the reticuloendothelial system (Figure 7), further indicating that the lung uptake is not due to particulate trapping in the small capillary system.

To improve in vivo biodistribution experiments of the 225Ac containing NPs (ca. 2 μCi/animal) demonstrated that the antibody-targeted NPs localized in the lung consistent with the binding properties of mAb 201b. The NPs exhibit high lung uptake with the antibody conjugate after 1 hour (51%ID/g). This high lung uptake dropped to 16.8%ID/g when the antibody conjugated NPs were competed with unconjugated antibody (Figure 6). These results demonstrate that the antibody retained its binding affinity and specificity even after conjugation to the NPs and that the NPs localized in the lung through antibody binding. While the antibody-labeled NPs cleared rapidly from the lungs in these proof-of-principle experiments (after 24 hours, 225Ac activity was predominant present in the liver and spleen), previous strategies used to reduce reticuloendothelial functioning such as treatment with clodronate liposomes could be applied to mitigate the rapid clearance [31], [32–33].

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Table 2. Dynamic light scattering of NPs in 18 mΩ water.

| Particle | Hydrodynamic diameter (nm) | Zeta potential (mV) |
|----------|---------------------------|---------------------|
| {La0.5Gd0.5}(225Ac)PO4@GdPO4@Au | 101.4±1.5 | −63.2±1.6 |
| {La0.5Gd0.5}(225Ac)PO4@GdPO4@Au-PEG | 382.3±6.5 | −56.4±0.1 |
| {La0.5Gd0.5}(225Ac)PO4@GdPO4@Au-mAb-201b | 1498±77 | −27.9±2.4 |

doi:10.1371/journal.pone.0054531.t002

Figure 5. MAb 201b antibody conjugation to multi-layered NPs.

doi:10.1371/journal.pone.0054531.g005
which is difficult to access with larger constructs that promote endocytosis. Attempts to reduce 213Bi toxicity through targeted, metal-chelate based internalizing antibodies have shown only moderate success [35].

The NP construct described in this work improves 225Ac daughter retention relative to both chelate approaches and previous NP constructs. \(\text{La}_{0.5}\text{Gd}_{0.5}\{225\text{Ac}\}PO_4@\text{GdPO}_4@\text{Au}\) NPs contain 88% of the 221Fr daughter in vitro, compared with 50% retention observed with \(\text{La}(225\text{Ac})\text{PO}_4\) NPs [28]. Additionally, the in vivo \(\alpha\)-generator delivery agent has a negligible effect on the energies of the emitted \(\alpha\)-particles. A 6 MeV \(\alpha\)-particle loses less than 0.2% of its energy in the layered NP whereas the range of the 100 keV recoiling daughters is \(\sim\)20 nm in bulk LnPO\(_4\). Moreover, a portion of the kinetic energy of the daughter particle may be transferred to the entire particle. If a portion of the recoil energy is distributed throughout the highly structured crystalline lattice, the recoiling range of the daughter radionuclides will be significantly decreased [36].

In vivo, the increase of retention of 213Bi in the target tissue over time results from a combination of the ability of the layered NPs to retain the daughter products and endocytosis of the TAT NP. In this work, 213Bi daughter retention in vivo with the layered NP showed improvement over the LaPO\(_4\) core NP [28]. The 213Bi retention is lower than the 221Fr retention because prior decays of 225Ac, 221Fr, and 217At can move the remaining \(\alpha\)-emitting nuclides towards the surface of the NP. From this position nearer to the surface, subsequent \(\alpha\) decays are likely to release the daughter nuclide from the NP. The amount of 213Bi which relocates to the kidney from other tissues shows marked improvement with the \(\text{La}_{0.5}\text{Gd}_{0.5}\{225\text{Ac}\}PO_4@\text{GdPO}_4@\text{Au-mAb-201b}\) system compared with the \(\text{La}(225\text{Ac})\text{PO}_4@\text{mAb-201b}\) system. Only 2.8% of the injected dose migrated to the kidney after 1 hour and 1.5% after 24 hours in the layered NPs while 10% of the ID relocated to the kidney after 1 hour and 5% after 24 hours with the core only lanthanum phosphate NPs.

These experiments demonstrate that multi-functional, layered NPs can be used to deliver and retain 225Ac and its daughter radioisotopes at a target site thereby reducing the absorbed dose to non-target organs. TAT experiments in a model tumor system are in progress to directly assess the efficacy of the constructs.

Materials and Methods

All chemicals were used as received from Sigma-Aldrich and were at least ACS grade unless otherwise noted. Water originated from an in house 18 MΩ MilliQ system. Radioactivity measurements were performed with \(\gamma\)-ray spectroscopy employing a calibrated high purity germanium detector employing a PC-based multichannel analyzer (Canberra Industries) windowed on 221Fr (212 keV) and 213Bi (440 keV). 225AcCl\(_3\) was prepared as previously described from a 229Th cow [28]. A Spectra/Por 10 kDa molecular weight cutoff (MWCO) regenerated cellulose dialysis membrane was used to separate NPs from solutions. Dialysis membranes were washed of preservatives before use against 18 MΩ water. A large NdFeB magnet (3” O.D. ×0.5” thick, surface field = 0.4 T) was obtained from United Nuclear.

Preparation of \(\{\text{La}_{0.5}\text{Gd}_{0.5}(225\text{Ac})\}\text{PO}_4\) Core Particles

Core particles were made by modifying a methodology developed by Buisette et al. [26]. Briefly, 50 µL each of 0.1 M LaCl\(_3\) and GdCl\(_3\) were mixed in a 1 mL V-bottom vial with spin vane. For the synthesis of radioactive NPs, 5.2 mCi of 225AcCl\(_3\) in 0.1 M HCl was added to the lanthanide mixture. Next, 200 µL of 0.1 M sodium tripolyphosphate (Na-TPP) was added to give a
total Ln:Na-TPP ratio of 1:2 resulting in a clear, colorless solution. If the solution remained turbid after addition of Na-TPP, it was vortexed with small (10 μL) additions of Na-TPP until the solution appeared clear. The resulting solution was then capped and heated at 90°C for 3 hours giving a turbid, white suspension of particles. Particles were purified via dialysis overnight. This preparation produced monodisperse particles of ~4 nm diameter which were characterized by transmission electron microscopy (TEM, JEOL 1400), neutron activation analysis (NAA) and x-ray diffraction (XRD, Scintag X2).

Layering of Particles
Core particles described above were centrifuged at 3,000 g for 3 minutes and the supernatant was removed. The particles were redispersed in a solution consisting of 200 μL of 0.05 M GdCl₃ and 400 μL 0.05 M Na-TPP. The resulting mixture was vortexed briefly then sonicated using a bath sonicator for 10 minutes before heating at 90°C for three hours. This process was repeated for up to four shell additions, at which point the solution becomes a thick milky white. Particles were purified by dialysis as above before gold coating.

The dialyzed particles (12 mg) were collected and split evenly between three 5-mL V-bottom vials. 300 μL of 0.1 M tribasic sodium citrate was added to each vial along with 1.5 mL of 18
MΩ water. Next, 2.5 mL of 1 mM NaAuCl₄⁻ was added dropwise to the solution slowly at the rate of 1 mL every 10 minutes. After the final addition, the solution was kept at 90°C for 30–45 minutes. A large NdFeB magnet (surface field = 0.4 T) was placed next to the V-bottom vial for 16 hours to separate the particles from solution. The supernatant was decanted to isolate the magnetically-activated particles. In the radiotracer labeling experiments, the separation efficiency was determined by γ-ray spectrometry of the removed supernatant and magnetically collected particles [28]. Non-radioactive analogs of the particles were characterized by EELS-TEM (Zeiss Libra 120) and NAA.

In Vitro Testing of \( ^{221}\text{Fr} \) Retention

In order to test retention of the \( ^{225}\text{Ac} \) decay products in vitro, the \( ^{225}\text{Ac} \)-NPs were loaded into a dialysis membrane and dialyzed against 400 mL of 18 MΩ water. The dialysis tube was stirred for a sufficient time for daughter equilibrium to be established (>3 hours), then a 5 mL aliquot was taken for γ-ray spectrometry analysis. Each sample was re-analyzed at a later time to determine the level of \( ^{225}\text{Ac} \) in the removed dialysate fraction. The measured activities were corrected for decay and dialysate loss from prior aliquot removals. The \( ^{213}\text{Bi} \) activity in the dialysate was used as a measure of the \( ^{221}\text{Fr} \) that was released from the NP, as \( ^{213}\text{Bi} \) which escaped from the particles did not move across the dialysis membrane [28].

Surface Modification

Surfaces were modified using a lipooamide-dPEG₂₉-acid linker (Quanta Biodesign). Two mg of dPEG were added, followed by 6 mg of Tris(2-carboxyethyl)phosphine reducing agent to cleave the disulfide bond. The pH of the solution was adjusted to 7 using 0.1 M NaOH and the reaction mixture was stirred for 4 hours. Connection of the linker was determined by a shift in the plasmon resonance near 530 nm as monitored by UV-Vis spectroscopy before and after the addition of the linker.

Antibody Conjugation

To attach antibodies to the linker, the carboxylate group on the PEG linker was first activated using 8 µL of 10 mg/mL aqueous 3-sulfo-N-hydroxyxuccinimide (sulfo-NHS) and 80 µL of 10 mg/mL 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC). 4 mg of EDC and 0.4 mg of sulfo-NHS were used per mg of NPs. After 15 minutes, the solutions were centrifuged to remove excess EDC/NHS, the supernatant removed, and the particles redispersed in phosphate buffered saline (0.01 M NaPO₄ pH 7.6 in 0.15 M NaCl, PBS). MAb 201b was added (~1 mg of mAb/mg of NP), and the mixture was mixed by rotation overnight. The reaction was then quenched with 1 mg of glycine, which was allowed to react for 15 minutes. The particles were then centrifuged to remove excess antibody. The supernatant was removed and the particles were redispersed in PBS containing 5 mg/mL bovine serum albumin (BSA/PBS). The particles were sonicated with a Branson microprobe for 10 sec and vortexed prior to injection. The final product of the mAb conjugated NP was ~3 mg/mL NP with 400 µCi of \( ^{225}\text{Ac} \) and ~1 mg mAb 201b.

Biodistribution Studies

All experiments involving mice were performed according to the Institutional Animal Care and Use Committee of the University of Tennessee approved protocol 1502. Female BALB/c mice (body mass ~20 g) were used for all biodistribution and imaging experiments. Biodistribution and daughter retention assays were done on three groups, consisting of three mice per group, were each injected intravenously (tail vein). Groups 1 and 2 were injected with \( ^{225}\text{Ac} \)-Gd₄O₄(\( ^{225}\text{Ac} \))PO₄@mAb-201b outer shell/inner shell/core conjugates, while group 3 was treated with \( ^{225}\text{Ac} \)-Gd₄O₄(\( ^{225}\text{Ac} \))PO₄-pEG NPs as a control. Group 1 mice received 14.6 µg of NP with 1.95 µCi of Ac-225 and ~5 µg of attached mAb 201b (this value was estimated from data in a parallel experiment wherein about 30% of added radioiodinated mAb was incorporated in NP under similar conditions). Group 2 received the same amount of targeted NP but with the addition of 750 µg of free mAb 201b as competitor. Group 3 received the same amount of NP and Ac-225, but with no targeting agent conjugated. Mice were housed with food and water ad libitum in a light/dark cycle environment before sacrificing at 1 and 24 h post-injection for biodistribution and in vivo retention studies. Biodistribution studies were performed on lungs, liver, spleen, and kidneys to evaluate the amount of both \( ^{221}\text{Fr} \) and \( ^{213}\text{Bi} \) in target organs by measuring weighed tissue samples in a γ-ray scintillation counter at a specific time postsacrifice and again after the radioisotopes had achieved decay equilibrium (>3 h). Quantities of \( ^{221}\text{Fr} \) and \( ^{213}\text{Bi} \) present at the time of animal sacrifice were determined by appropriate crossover and decay corrections as previously described [28].

MicroSPECT/CT Imaging

Small animal imaging was performed using a microCAT II SPECT dual modality platform (Siemens Preclinical Imaging, Knoxville, TN). Mice were injected with approximately 40 times more NP than were the mice for biodistribution studies. Thus the mice for competition with cold mAb 201b did not have the same ratio of cold competitor to radiolabeled NP and competition was not as complete. Animals were sacrificed by overdose of isoflurane at 1 h postinjection and imaged via microSPECT/CT 3 h later when the \( ^{225}\text{Ac} \) and its daughters had reached equilibrium. SPECT data for the final images were acquired as previously described by Woodward et al. [28].

Author Contributions

Conceived and designed the experiments: MFM JW SJK SM JDR. Performed the experiments: MFM JW RAB SJK. Analyzed the data: MFM SJK. Contributed reagents/materials/analysis tools: AJR. Wrote the paper: MFM SJK SM JDR.

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