Preliminary Therapy Evaluation of $^{225}$Ac-DOTA-c (RGDyK) Demonstrates that Cerenkov Radiation Derived from $^{225}$Ac Daughter Decay Can Be Detected by Optical Imaging for In Vivo Tumor Visualization

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Abstract

The theranostic potential of $^{225}$Ac-based radiopharmaceuticals continues to increase as researchers seek innovative ways to harness the nuclear decay of this radioisotope for therapeutic and imaging applications. This communication describes the evaluation of $^{225}$Ac-DOTA-c(RGDyK) in both biodistribution and Cerenkov luminescence imaging (CLI) studies. Initially, La-DOTA-c(RGDyK) was prepared as a non-radioactive surrogate to evaluate methodologies that would contribute to an optimized radiochemical synthetic strategy and estimate the radioactive conjugate’s affinity for $\alpha_v\beta_3$, using surface plasmon resonance spectroscopy. Surface plasmon resonance spectroscopy studies revealed the IC$_{50}$ and K$_d$ of La-DOTA-c(RGDyK) to be 33 ± 13 nM and 26 ± 11 nM, respectively, and suggest that the complexation of the La$^{3+}$ ion to the conjugate did not significantly alter integrin binding. Furthermore, use of this surrogate allowed optimization of radiochemical synthesis strategies to prepare $^{225}$Ac-DOTA-c(RGDyK) with high radiochemical purity and specific activity similar to other $^{225}$Ac-based radiopharmaceuticals. This radiopharmaceutical was highly stable in vitro. In vivo biodistribution studies confirmed the radiotracer’s ability to target $\alpha_v\beta_3$ integrin with specificity; specificity was detected in tumor-bearing animals using Cerenkov luminescence imaging. Furthermore, tumor growth control was achieved using non-toxic doses of the radiopharmaceutical in U87mg tumor-bearing nude mice. To our knowledge, this is the first report to describe the CLI of $\alpha_v\beta_3$+ tumors in live animals using the daughter products derived from $^{225}$Ac decay in situ. This concept holds promise to further enhance development of targeted alpha particle therapy.

Key words: Actinium-225, Targeted Alpha Particle Therapy, Cerenkov Luminescence Imaging, $\alpha_v\beta_3$ integrin.

Introduction

The recent approval of Radium-223 ($^{223}$Ra) by the United States Food and Drug Administration for palliative treatment of bone metastasis associated with hormone-refractory prostate cancer has renewed interest in alpha particle (α)-emitting radionuclides and targeted alpha particle therapy (TAT) [1-3]. This
type of therapy derives its efficacy from α-particles, which are essentially helium (He) atom nuclei. Once ejected from the decaying nucleus, they travel less than 100 microns in a linear manner depositing a very large amount of energy through their path by ionization. This energy deposition, also referred to as Linear Energy Transfer (LET), is responsible for their enhanced cytotoxicity at activities far below those needed to achieve the same cell-killing efficiency of low LET β-emitters. Furthermore, when high-LET occurs within the cell nucleus irreparable double-stranded (ds) DNA breaks occur and result in activation of autophagy, necrosis, and cell cycle arrest pathways [4, 5]. Importantly, α-particles do not rely on the generation of indirect reactive oxygen species; thus efficacy of TAT is not diminished by tumor hypoxia or chemoresistance that commonly develops after traditional chemotherapy. Finally, when properly targeted, the high LET of α-particles can therapeutically target small clusters of malignant cells among normal tissues [2, 6-11].

Except for Radium-223 (223Ra), few α-particle emitting radionuclides are used clinically due to incompatible half-lives, cost of production, and limited availability. 225Ac (t1/2 = 10 d; Eαmax = 6-8 MeV) has gained considerable attention in TAT research because 1) it is readily available from Oak Ridge National Laboratories, 2) it has a 10-day half-life, 3) it functions as a nanogenerator at the tumor site, emitting 4 α-particles that release approximately 30 MeV of total kinetic energy per nuclear decay, 4) it can be chelated to DOTA, which is contained in numerous radiopharmaceuticals approved for clinical use or currently undergoing clinical trials, and 5) it is currently being used successfully in preclinical studies and clinical trials [2, 6-12]. Despite these attributes, biodistribution and pharmacokinetic data of 225Ac-based radiopharmaceuticals cannot be acquired using traditional pre-clinical imaging techniques such as positron emission tomography (PET) without significantly modifying the TAT radiopharmaceutical to accommodate a PET isotope. Although preclinical SPECT imaging with 225Ac-radiopharmaceuticals has been attempted through the detection of the 440 keV γ emission that occurs after 213Bi decay, data are limited since animals are typically euthanized 1 h after receiving the radiopharmaceutical and imaged 24 h after euthanasia to ensure isotopic equilibrium [13]. Thus, these experiments do not address the potential of longitudinal SPECT studies to inform development of 225Ac-radiopharmaceuticals. However, the ability to image and visualize 225Ac-based radiopharmaceutical biodistribution, metabolism, and clearance in animal models through longitudinal imaging studies would be advantageous, since it is currently unavailable for preclinical stages of TAT development [14, 15].

The Cerenkov effect, observed nearly a half-century ago, describes the emission of ultraviolet light when certain charged particles exceed the phase velocity of light in a given medium [16, 17]. This effect can be observed using standard optical imaging systems originally designed to detect bioluminescence and fluorescence. Since first being described in molecular imaging research, the imaging and visualization of Cerenkov radiation, or Cerenkov luminescence imaging (CLI), has been demonstrated preclinically and clinically with various radioisotopes [16, 18]. Recently, several groups described the Cerenkov emissions derived from medically relevant isotopes, including 225Ac, which yields the largest optical signal among all isotopes examined [15, 19]. Since the α-particle emitted during nuclear decay travels with low velocity, it was postulated that the observed emission resulted from the beta decay of 213Bi, 209Tl, and 209Pb in the 225Ac decay chain. Subsequent reports using theoretical and experimental means described the association between Cerenkov radiation and 225Ac decay, but did not use an 225Ac-based radiopharmaceutical in an animal model to demonstrate its applicability to drug or radiopharmaceutical development [19, 20]. In this communication, we provide initial evidence that the secular equilibrium between 225Ac and its daughter products can be harnessed to visualize TAT delivery in live animals, using standard optical imaging techniques. We accomplished this by synthesizing 225Ac-DOTA-c(RGDyK) and evaluating its stability, biodistribution, and potential use as an imaging agent in CLI in a murine model expressing human glioblastoma U87mg tumors, which overexpress the α3β1 integrin. This receptor is a member of the integrin superfamily and is overexpressed in malignancies and on neovasculature necessary for tumor growth [21, 22]. Finally, preliminary toxicity and therapy studies were conducted to demonstrate the effect of 225Ac-DOTA-c(RGDyK) on tumor growth, to further assess the theranostic potential of this approach.

**Materials and Methods**

Unless otherwise noted, all chemicals were purchased from Sigma-Aldrich Chemical Co. (St. Louis, MO, USA), and solutions were prepared using ultrapure water (18 MΩ-cm resistivity). DOTA-c-c(RGDyK) was purchased from CPC Scientific, Inc. (Sunnyvale, CA), and its purity and mass were confirmed at Wake Forest using HPLC and electrospray ionization (ESI) mass spectrometry as described below. Recombinant human α3β1 integrin was purchased from R&D Systems, Inc. (Minneapolis,
was purchased from AnaSpec (Freemont, CA). 225Ac(NO3)3 (t1/2 = 10 d; Edmax = 6-8 MeV) was received from Oak Ridge National Laboratory (Oak Ridge, TN) and dissolved in 0.2 M Optima grade hydrochloric acid (HCl, Fisher Scientific) before use. Electrospray ionization (ESI) mass spectra were obtained on an Agilent 1100 LC/MSD in the Wake Forest University Department of Chemistry.

The radiochemistry reaction progress and purity were analyzed using a Waters analytical HPLC (Milford, MA), which runs Empower software and is configured with a 1525 binary pump, 2707 autosampler, 2998 photodiode array detector, 2475 multichannel fluorescence detector, 1500 column heater, fraction collector, Grace Vydac 218MS C18 column (5 µm × 250 mm, Grace Davidson, Deerfield, IL) and a Carrol Ramsey 105-s radioactivity detector (Berkeley, CA). DOTA-c(RGDyK) and the associated La-DOTA-c(RGDyK) complex were monitored at 220 nm using a mobile phase consisting of 0.1% TFA/H2O (solvent A) and 0.1% TFA/acetonitrile (solvent B), and a gradient consisting of 0% B to 70% B in 20 min at a flow rate of 1.2 mL/min. In addition, radio-TLC was conducted on a Bioscan AR 2000 radio-TLC scanner equipped with a 10% methane:argon gas supply and a PC interface running Winscan v.3 analysis software (Eckert & Ziegler, Berlin, DE). Varian ITLC-SG strips and Merck C18 TLC plates were employed using a 0.9% NaCl/10 mM NaOH and 30:70 10% NH4OAc/methanol solution as eluents, and 225Ac(NO3)3 as a standard control. Activity of radioactive samples was measured using previously described procedures with either a CRC-25R radioisotope calibrator (Capintec, Inc.) or a Perkin Elmer 2480 Wizard® gamma counter (Waltham, MA) [23].

### Synthesis of La-DOTA-c(RGDyK)

A 1.5 mL vial was charged with DOTA-c(RGDyK) (200 µg, 0.17 µmol), LaCl3 (76.1 µg, 0.20 µmol), 0.2 mL of water, and 6 µL 0.1 M Na2CO3. The pH of the resulting solution was 5.5-6, and it was stirred for 1 h at 70 °C. The product was obtained in quantitative yield and lyophilized to a white solid. Formation of La-DOTA-c(RGDyK) was confirmed by ESI-MS analysis. Calculated for C51H72N14O16SLa, 1307.4 [(M+2H)+] Observed: 1307.6 [(M+2H)+].

### Radiochemical synthesis of 225Ac-DOTA-c(RGDyK), Quality Control, and in vitro Serum Stability

The complexation of 225Ac with DOTA-c(RGDyK) was achieved by reacting DOTA-c(RGDyK) (5-10 µg (5-10 µL 1.0 mg/mL in water)) with 225Ac(NO3)3 (3.4 MBq) that was diluted in 100 µL of water containing 10 µL of 20% L-ascorbic acid. The pH of the resulting solution was adjusted to 5.5-6 using 1 M Tris buffer (10-12 µL), and then incubated at 60°C for 1 h. Reaction progress and radiochemical purity of 225Ac-DOTA-c(RGDyK) were measured without further purification using ITLC with gamma counting, radio-TLC, gamma counting of radio-HPLC fractions, or CLI. In vitro serum stability was carried out by adding 50 µL of 225Ac-DOTA-c(RGDyK) (2 MBq) to 900 µL of human serum. The solutions (n = 4) were incubated at 37 °C for 10 days and were analyzed daily using radio-TLC, ITLC with gamma counting or size exclusion chromatography using a Superdex 200 10/300 GL™ column (GE Healthcare Life Sciences, Piscataway, NJ) and phosphate buffered saline (PBS) as eluent with a flow rate of 0.5 mL/min.

### Surface Plasmon Resonance Spectroscopy (SPR) of La-DOTA-c(RGDyK)

Lyophilized samples stored at -20 °C were reconstituted to 100 µg/mL in 130 mM NaCl, 10 mM HEPES, 1 mM CaCl2, 1 mM MgCl2, pH 7.4 (HBS) for SPR measurements. Samples of the lyophilized peptide were dissolved in water (5 mg/mL) and then diluted in sodium acetate buffer (50 µg/mL; pH 5.0) for coupling to biosensor chips. CM-5 biosensor chips (Biacore, Inc., Piscataway, NJ) were activated for amine coupling by reaction with 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide and N-hydroxysuccinimide in a Biacore T100 instrument [24, 25]. c(RGDyK) (50 µg/mL) in 0.1 M sodium acetate buffer (pH 5.0) was delivered to the sample chamber, yielding immobilization densities of 222 response units (RU) for the αvβ3(La-DOTA-c(RGDyK)) binding series and 196 RU for the La-DOTA-c(RGDyK) competition series. Residual active groups were blocked with ethanolamine. Interactions between immobilized c(RGDyK) and recombinant human integrin and La-DOTA-c(RGDyK) were measured by collecting the signals from both reference and sample channels at 10 Hz. Reagents were maintained at 25.0 ± 0.1°C in the sample compartment; experiments were performed at 25.00 ± 0.01°C in the analysis chamber. Two start-up cycles were performed in which HEPES buffered saline (HBS) was delivered, followed by a 2-step regeneration cycle (20 mM EDTA in HBS for 180 sec then 5 M NaCl in water for 180 sec). During the binding steps, increasing concentrations (0 - 300 nM) of αvβ3 in HBS were delivered at 30 µL/min for 1000 sec. Dissociation of the αvβ3 integrin:ligand complex was monitored for 1000 sec, as HBS flowed over the biosensor surface at 30 µL/min, and residual bound integrin was displaced by the regeneration process.
cycle. The biosensor was then equilibrated for 300 sec with HBS before delivery of the next integrin aliquot. Sensorgram data (sample – reference signals) were further corrected by subtracting the time-dependent profiles obtained with a buffer blank. These doubly corrected signals were then fit globally by nonlinear regression (Biacore Evaluation software); first to a single-site binding model, and then to a ligand-induced conformational change model [25, 26]. In each case, the quality of the fit was judged by the residuals and the reduced χ² value. For competition experiments, the αvβ₃ concentration was fixed at 83 nM and samples containing increasing concentrations of La-DOTA-c(RGDyK) were prepared for delivery to immobilized c(RGDyK) (vide supra).

Animal Models

All animal experiments were conducted in compliance with Institutional Animal Care and Use Committee guidelines established by Wake Forest University Health Sciences. Normal BALB/c and nude mice (weight, 20–22 g; age, 6–8 wk) were purchased from Jackson Laboratories (Bar Harbor, ME). Thirty-six female athymic nu/nu mice were subcutaneously injected into the flank with U87mg human glioblastoma tumor cells (1x10⁶ cells; ATCC, Alameda, CA). The average radiance (p/s/cm²/sr) field of view B) with no light interference from the excitation lamp. Images were analyzed using the Living Image 2.6 software (Caliper Life Sciences, Alameda, CA). The average radiance (p/s/cm²/sr) was used for quantitative region of interest (ROI) analysis from each image. Background correction was performed through (a) use of dark images acquired at the equivalent instrument integration setting immediately before experimental image collection, or (b) ROI analysis of a region in the same experimental image but remote from the area of interest.

Therapy Studies

Maximum tolerated dose studies (MTD) were conducted using eight cohorts of normal BALB/c mice. After a single intravenous injection of ²²⁵Ac-DOTA-c(RGDyK) or saline, cohorts (n = 5 mice/cohort) were weighed 3 times/week, and monitored for 110 days for signs of distressed behavior. At the end of that period, animals were euthanized and tumors and organs of interest were removed and imaged ex vivo. Optical images were collected using a Xenogen IVIS 100 optical imager (f/stop: 2; binning 1, field of view B) with no light interference from the excitation lamp. Images were analyzed using the Living Image 2.6 software (Caliper Life Sciences, Alameda, CA). The average radiance (p/s/cm²/sr) was used for quantitative region of interest (ROI) analysis from each image. Background correction was performed through (a) use of dark images acquired at the equivalent instrument integration setting immediately before experimental image collection, or (b) ROI analysis of a region in the same experimental image but remote from the area of interest.
c(RGDyK) or \(^{225}\text{Ac-DOTA-c(RGDyK)}\). Doses were determined from the results of the MTD study. Animals were monitored daily for signs of distressed behavior, and tumor growth was evaluated thrice weekly using manual tumor volume (volume = 0.52 X [width]^2 X [length]) measurements. Mice were sacrificed 18 days after tumor cell inoculation as previously described [31].

**Statistical Methods**

All data are presented as mean±SD or mean (95% confidence intervals). For statistical classification, Student’s t test (two-tailed, unpaired) was performed either using Prism 6.0 (GraphPad Software, Inc., San Diego, CA) or SigmaPlot (Systat Software, Inc. San Jose, CA). For therapy data, a one-way ANOVA test (including Dunnett’s multiple comparison test) was performed. A p<0.05 was considered significant.

Results and Discussion

The reliable availability of carrier-free \(^{225}\text{Ac}\) from Oak Ridge National Laboratories or The Institute of Transuranium Elements has allowed the theranostic potential of \(^{225}\text{Ac}\) to be tested [2, 9]. However, unlike many radionuclides used in imaging and therapy, the high cost of actinium and the lack of stable isotopes to facilitate even the most basic studies for ligand development or conjugate performance at an early preclinical stage have retarded progress. To facilitate our studies in development of \(^{225}\text{Ac-DOTA-c(RGDyK)}\), we chose to prepare La-DOTA-c(RGDyK) as a non-radioactive surrogate to develop an initial synthetic route to the \(^{225}\text{Ac}\)-radiopharmaceutical and evaluate the conjugate’s affinity for \(\alpha_v\beta_3\) integrin once the trivalent ion was chelated to the conjugate (see Scheme S1 of the Supporting Information (SI)). We hypothesized that the use of La\(^{3+}\) was appropriate, since the reaction conditions needed to prepare many \(^{225}\text{Ac}\) compounds or complexes parallel those used to synthesize analogous La\(^{3+}\) compounds and complexes. Since their chemistries are highly similar in both preparative and analytical experiments, La\(^{3+}\) is considered a useful surrogate for \(^{225}\text{Ac}\)\(^{3+}\) [32, 33]. Accordingly, we prepared La-DOTA-c(RGDyK) in quantitative yield using a modified literature procedure [34], and confirmed the identity of this complex using analytical HPLC (Figure 1) and electrospray ionization mass spectrometry (ESI*-MS) (Figure S1).

We then determined the affinity of this conjugate for \(\alpha\beta_3\) using SPR, a sensitive optical technique that provides real-time kinetic data for binding a receptor (delivered in solution by microfluidics) to a ligand (immobilized on a biosensor) [35, 36]. This powerful analytical tool has many applications in environmental protection, biotechnology, medical diagnostics, and drug discovery [37]. Initially, we sought to determine the rate, affinity, and specificity of the interactions between \(\alpha\beta_3\) and c(RGDyK). Increasing concentrations of \(\alpha\beta_3\) were delivered to a biosensor derivatized with c(RGDyK). SPR data showed time- and concentration-dependent increases in RU when \(\alpha\beta_3\) interacted with c(RGDyK) (Figure S2). RU signals were reduced to near-background values when c(RGDyK) was included at 11- and 115-fold molar excess over \(\alpha\beta_3\) (30 nM). These competition results demonstrate that our SPR approach reports specific \(\alpha\beta_3:c(RGDyK)\) binding [24].

The \(\alpha\beta_3\) binding data (10 concentrations ranging from 3 - 300 nM) were fit to a reversible two-state bimolecular interaction model, to determine the forward (\(k_{d1}, \text{L/mol-sec}\)) and reverse (\(k_{d2}, \text{sec}^{-1}\)) rate constants for the initial binding step and forward (\(k_{d1}, \text{sec}^{-1}\)) and reverse (\(k_{d2}, \text{sec}^{-1}\)) rate constants for the subsequent stabilizing conformational change (Figure S2) [25]. The quality of the fit can be judged by the close correspondence between the solid lines and the data in Figure S3 and the tightly distributed residuals. This data set yielded \(k_{d1} = 1.58 \pm 0.01 \times 10^4\ \text{L/mol-sec}\)
and a $k_d$ of $9.91 \pm 0.03 \times 10^{-3}$ sec$^{-1}$ at 25 °C. These studies also revealed the kinetic parameters for the conformational change step ($k_{d2}$ of $1.22 \pm 0.01 \times 10^{-3}$ sec$^{-1}$ and $k_{f2}$ of $1.23 \pm 0.01 \times 10^{-3}$ sec$^{-1}$; Figure S2). The overall $K_d$ for the reaction was $3.16 \times 10^{-7}$ mol/L. Attempts to fit the $\alpha_3\beta_3\gamma_3$ binding data to a single-site kinetic model proved problematic, yielding large deviations between the biphasic association and dissociation profiles and the fitted lines [26, 35, 36, 38]. Recognizing that binding RGD ligands perturb $\alpha_3\beta_3$ resting conformation, we focused on a ligand-induced conformational change model that others have successfully applied to analyze multiphasic SPR kinetic data [39-44].

Our next goal was to determine how tightly $\alpha_3\beta_3$ binds La-DOTA-c(RGDyK). However, amine coupling of this conjugate to a biosensor chip was precluded because its lysine residue was blocked by derivatization. Hence, we followed a competition approach in which $\alpha_3\beta_3$ (83 nM) binding to immobilized c(RGDyK) was first measured, followed by two samples with increasing concentrations of La-DOTA-c(RGDyK) [26]. This three-step sequence was followed to examine competitor concentrations of 9, 26, 94, 300, 915, 2515, 10061, 37729 nM, covering a range of 0.1 – 450-fold molar excess. The resultant kinetic traces (Figure S4) show a concentration-dependent decrease in $\alpha_3\beta_3\gamma_3$ binding signals (RU), which approached baseline values as the concentration of La-DOTA-c(RGDyK) approached 300 nM. Plotting RU vs La-DOTA-c(RGDyK) molar excess over $\alpha_3\beta_3$ (log scale) yielded a sigmoid profile characterized by an IC$_{50}$ equivalent to $0.40 \pm 0.16$ fold-molar excess [26]. This corresponds to an IC$_{50}$ of $33 \pm 13$ nM and yields a K$_d$ of $26 \pm 11$ nM, which align with literature reports describing $^{68}$Ga-or $^{64}$Cu-RGD based radiopharmaceuticals [45, 46]. Furthermore, the similarity of our results with published values for DOTA-c(RGDyK) implies that complexation of the La$^{3+}$ ion by the conjugate does not interfere with integrin binding and suggests that efficient binding of $^{225}$Ac-DOTA-c(RGDyK) to $\alpha_3\beta_3$ is likely to occur in vivo.

Based on our synthetic strategy to prepare La-DOTA-c(RGDyK), we prepared $^{225}$Ac-DOTA-c(RGDyK) with a radiochemical purity greater than 99.8%, a radioyield greater than 95%, and specific activity of 680 kBq/µg (Scheme S2). These values are based upon results obtained from ITLC in conjunction with gamma counting that revealed a single radiolabeled product, which eluted with the solvent front. Although few $^{225}$Ac-labeled peptides have been reported, the specific activity for our tracer is well within the spectrum reported for similar radiopharmaceuticals [31, 47]. Additionally, we sought to determine if $^{225}$Ac-radiopharmaceuticals could be analyzed using radio-TLC, HPLC with gamma counting or CLI, since their use in this arena has not been reported previously (Figure 1 and S5). Although ITLC has been used traditionally for $^{225}$Ac radiopharmaceutical purity determination, all these techniques can be informative during radiopharmaceutical synthesis and may enhance development of TAT. We obtained quality control samples and then waited 24 hours so that the secular equilibrium between $^{225}$Ac and its daughter products was established. Based on radio-TLC analyses, we observed that unchelated $^{225}$Ac remained at the origin ($R_f = 0$), whereas $^{225}$Ac-DOTA-c(RGDyK) eluted near the solvent front ($R_f = 0.86$). Additionally, CLI of the same TLC plates indicated only one spot at the origin and one near the solvent front (Figure S5). Although CLI resolution is poor compared to that of the radio-TLC scanner, two samples with significantly different $R_f$ values could be distinguished; thereby providing qualitative confirmation of the radio-TLC results in excellent agreement with our ITLC data. Further corroboration of purity was provided by measuring radioactivity in HPLC fractions, which was determined by gamma counting using a 190-230 keV window. Plotting the results per unit time demonstrated we synthesized $^{225}$Ac-DOTA-c(RGDyK) in 99.8% purity, which agreed with the gamma counting results from ITLC plates. Using this technique, $^{225}$Ac-DOTA-c(RGDyK) had a retention time of 11.02 minutes, similar to the UV-HPLC chromatograms of DOTA-c(RGDyK) and the non-radioactive analogue, La-DOTA-c(RGDyK) (Figure 1).

For initial stability studies, we incubated $^{225}$Ac-DOTA-c(RGDyK) in an aliquot of human serum for 10 days at physiological temperature (see Table S1 for results). After 10 days, less than 5% of the radiopharmaceutical had degraded in the presence of serum proteins. We observed similar results when analyzing samples using radio-TLC or ITLC with traditional gamma counting, suggesting that these techniques can accurately describe the stability of $^{225}$Ac radiopharmaceuticals. However, since daughter nuclei could be bound to serum proteins that do not migrate in our ITLC system, we elected to perform size exclusion chromatography to further characterize the stability of our radiotracer and the results of these experiments are depicted in Figure S6. When samples of serum which contained unchelated $^{225}$Ac were analyzed using SE chromatography, we observed a large peak, whose elution started at 28 minutes and ended at 38 minutes. The peak patterns and elution profile matched that of serum protein components, which were observed in the UV chromatogram at 220
nm. In contrast, in serum samples containing the radiotracer, a minor peak (5%) was observed at 30 minutes, which correlated with serum proteins in the UV profile; the major peak (95%) eluted between 41-45 minutes and might reflect the intact radiopharmaceutical, which further corroborates the stability results obtained using ITLC and radio-TLC. While few $^{225}$Ac-peptide based radiopharmaceuticals are available for comparison, $^{225}$Ac-DOTA conjugated antibodies demonstrate similar stabilities in human serum, and further reinforce the observed stability of $^{225}$Ac-DOTA-c(RGDyK) in vitro \[48, 49\].

We evaluated $^{225}$Ac-DOTA-c(RGDyK), by conducting biodistribution studies in $\alpha_\beta^3$ U87mg tumor bearing mice, since U87mg tumor cells overexpress $\alpha_\beta^3$ integrin. This model was used successfully in previous studies to evaluate several PET and SPECT radiopharmaceuticals designed to image $\alpha_\beta^3$ expression \[46\]. $^{225}$Ac-DOTA-c(RGDyK) showed rapid blood clearance, with 99% of the activity present at 1 h removed from the blood by 24 h (Figures 2 and S7 and Table S2). In contrast, activity remained higher in the liver and kidney, with slower clearance from these tissues over time. From 1-24 h, activity in the liver fell by 52%, while 68% of the 1 h activity was excreted from the kidney by the end of the study. In addition, although appreciable accumulation (3.72±0.5%ID/g) of activity in the tumor was observed at 1 h, it decreased by 54% at 24 h post-injection (p.i.). This yielded average tumor:blood and tumor:muscle ratios of 12, 95 and 42, 2.4, 5.7 and 3.6 at 1, 4 and 24 h, respectively.

To examine the in vivo specificity of $^{225}$Ac-DOTA-c(RGDyK) for the receptor, blocking studies were performed at 4 h p.i. by co-injecting excess c(RGDyK) along with the analogous radiotracer (Figure 3). Injection of c(RGDyK) reduced accumulation of $^{225}$Ac-DOTA-c(RGDyK) in the tumor by 91% (non-block vs. block (%ID/g): 2.68±0.037 vs. 0.26±0.085; p < 0.0001), suggesting that the radiopharmaceutical is being retained at the tumor site through specific interactions between the targeting RGD ligand and the $\alpha_\beta^3$ integrin. These results are in excellent agreement with previously published data that describe the biodistribution of analogous PET and SPECT radiopharmaceuticals designed to target this integrin \[45, 50\]. Furthermore, it suggests that the conjugation of the radiometal chelate did not interfere with ligand:receptor interactions, as predicted by our SPR experiments using La-DOTA-c(RGDyK).

Cerenkov luminescence imaging continues to evolve as an optical imaging technique with potential applications in preclinical drug development, radiopharmaceutical development, and surgical sciences by enhancing intraoperative techniques \[16\]. Only a few reports have described CLI in the context of the $\alpha$-emitting isotope $^{225}$Ac, which does not demonstrate prompt Cerenkov emissions, but relies upon the beta decay of $^{213}$Bi, $^{209}$Tl and $^{209}$Pb to generate the optical signal that is detected using standard optical instrumentation \[15, 19, 20\]. Initial reports using in silico modeling and in vitro experimentation described a mandatory data acquisition delay of 10 hours for $^{225}$Ac to achieve equilibrium with its daughter radionuclides so that the measured light output in a unit volume would be proportional to the
activity of Ac-225 in that unit volume [19, 20]. With this criterion in mind, we injected U87mg tumor-bearing mice, with $^{225}\text{Ac}$-DOTA-c(RGDyK) or with $^{225}\text{Ac}$-DOTA-c(RGDyK) and excess c(RGDyK) as a blockade (Figure 4). We relied on the well-documented ability of RGD-containing ligands to be internalized on integrin binding to sequester the $^{225}\text{Ac}$ and the daughter products within the tumor. Although not used as part of an imaging strategy, this technique, often referred to as the nanogenerator approach, has been used successfully to sequester $^{225}\text{Ac}$-based radiotherapeutics and daughter decay products within a cell to increase their therapeutic effectiveness [51-53]. Surprisingly, all animals tolerated $^{225}\text{Ac}$-DOTA-c(RGDyK) extremely well, demonstrating no signs of distress during these experiments. Twenty-four hours after radiopharmaceutical injection, whole-animal CLI revealed Cerenkov radiation emissions from the subcutaneous, $\alpha_\text{v}\beta_3^+$ tumors and the liver and kidneys, which have been shown in related biodistribution studies to be involved in the clearance and excretion of this radiopharmaceutical [46, 54, 55]. In vivo imaging and ROI analysis (Figure 4A-C) revealed that tumors demonstrated an average radiance of $5.8 \times 10^3 \pm 0.5 \times 10^3$ p/s/cm$^2$/sr, while tumors of animals receiving $^{225}\text{Ac}$-DOTA-c(RGDyK) and blockade demonstrated a significant reduction in average radiance ($1.1 \times 10^3 \pm 0.08 \times 10^3$ p/s/cm$^2$/sr; $p < 0.0001$). These results suggest that activity delivered by the radiopharmaceutical and internalized through specific receptor-interactions is being retained in the tumor. Our data agree well with the biodistribution and blocking data reported for similar RGD systems radiolabeled with PET and SPECT isotopes [21, 50].

On ex vivo imaging (Figure S8 and Figure 4D), tumors had an average radiance of $8.8 \times 10^3 \pm 1.6 \times 10^3$ p/s/cm$^2$/sr; tumors of animals receiving the radiopharmaceutical and blockade demonstrated significantly less Cerenkov emission, with an average radiance of $1.1 \times 10^3 \pm 0.25 \times 10^3$ p/s/cm$^2$/sr ($p < 0.0001$). All other tissues examined did not demonstrate significant Cerenkov emission. These results contrast with the biodistribution results that show some radioactivity retention in all tissues. This difference may be attributed to the volume-dependent nature of Cerenkov emissions within tissues of different sizes, and the differences in detection sensitivity between the optical imaging and gamma counting techniques employed in these studies.

![Figure 4. Cerenkov luminescence imaging with $^{225}\text{Ac}$-DOTA-c(RGDyK). Images were acquired 24 h after intravenous injection to allow for equilibrium to occur between $^{225}\text{Ac}$ and its daughter products, which generate Cerenkov luminescence. (A) Efficient tumor localization was observed in U87mg tumor bearing animals at 24 h p.i. (B) Localization was reduced upon administration of blockade demonstrating in vivo specificity. (C, D) In vivo (C) and ex vivo (D) quantification based upon ROI analysis demonstrates that Cerenkov radiation in the tumor is significantly reduced upon the administration of blockade. (★) denotes tumor.](http://www.thno.org)
Based upon the results extrapolated from biodistribution data (Table S3), tumor clearance of radioactivity was approximately 3 fold slower than the clearance of radioactivity from the kidney. However, both tissues received a similar dose, which was not predicted by the CLI results. This finding corroborates an earlier publication that reported a similar observation in a murine model of leukemia [56].

We also completed preliminary therapy studies with 225Ac-DOTA-c(RGDyK) to further assess the theranostic potential of our approach (see Figure S9). Toxicity studies were completed in normal mice in order to determine the maximum tolerated dose of 225Ac-DOTA-c(RGDyK). By the end of the 16 week study, none of the animals achieved any of the predefined study endpoints, which included distressed behavior or average group weight loss of 15%, and only the cohort receiving the highest dose (0.16 MBq) demonstrated an average weight loss of 10±4% compared to control animals. Nephrotoxicity is a well-documented consequence of targeted alpha particle therapy; thus, we measured blood urea nitrogen (BUN) and creatinine content in serum, as indicators of renal health [57, 58]. BUN retention was elevated in all cohorts compared to the control animals suggesting impairment. However, inter-individual values were high among all groups, so that the differences were not statistically significant (Figure S9C). We also measured serum levels of creatinine, a byproduct of muscle metabolism eliminated from the blood by glomerular filtration and proximal tubular secretion. Compared to the control cohort, creatinine levels were elevated in all groups that received doses of 225Ac-DOTA-c(RGDyK) above 0.04 MBq. In contrast to the BUN results, inter-individual values were not as divergent; suggesting that elevated levels of creatinine in serum could reflect real kidney damage and a true reduction in kidney function (Figure S9A). Finally, histopathologic examination indicated that only the cohorts receiving the two highest doses developed multi-focal tubular nephrosis with lymphocytic infiltration, consistent with radiation-induced damage to the tubular epithelium (Figure S10). All other cohorts did not exhibit pathological changes consistent with acute kidney damage or impaired renal function. Furthermore, liver pathology was normal among all cohorts despite the enhanced retention of the radiotracer. All other tissues were similar in treatment versus control groups, consistent with findings of other groups investigating 225Ac-labeled peptides for TAT [31].

Overall, we established the MTD of 225Ac-DOTA-c(RGDyK) to be 0.04 MBq, and initiated therapy studies using female nude mice inoculated with human U87mg cells in the flank. Four days after inoculation, we randomly divided the mice into cohorts and treated them with saline, DOTA-c(RGDyK) or 225Ac-DOTA-c(RGDyK) at 0.04 MBq (1·MTD), 0.02 MBq (0.5·MTD) or 0.01 MBq (0.25 MTD). We then monitored tumor growth for 14 days after treatment to assess tumor burden. Based upon our results, a statistically significant reduction in tumor volume was achieved in animals receiving radiotherapy (Figure S9D). This suggests that tumor growth control can be achieved using doses of 225Ac-DOTA-c(RGDyK), which do not cause toxicity in non-target tissues. Furthermore, this growth control is observed with much lower levels of radioactivity than would be needed with β-emitting radiopharmaceuticals that target the same integrin [50], and illustrates the cytocidal potency of α-emitting radiotherapeutics.

This report demonstrates through proof-of-concept studies that the secular equilibrium that exists between 225Ac and its daughter products can be used for in vivo tumor detection. Additionally, we provide data illustrating the low toxicity of 225Ac-DOTA-c(RGDyK) in normal mice and its ability to retard tumor growth compared to controls. Our findings suggest that 225Ac-based radiopharmaceuticals have theranostic potential that should be explored pre-clinically for translation into the clinic. However, several limitations of these studies should be considered [19, 20].

Although CLI is demonstrated here using an 225Ac-based radiopharmaceutical, not all α-emitting radioisotopes will produce sufficient Cerenkov radiation to be useful for imaging. Secondly, given the 10 h time delay needed for equilibrium to be established between 225Ac and its daughter products, imaging the pharmacokinetics of small molecules and peptides at early time points is currently not possible. However, using this technique to monitor the biodistribution of 225Ac-labeled antibodies and nanoparticles, which require extended circulation time for effective tumor targeting and blood clearance, remains a possibility and worthy of exploration. Finally, the high recoil energy associated with 225Ac and daughter product release from the original conjugate must be considered. Emission of Cerenkov radiation may not reflect the actual biodistribution of the radiopharmaceutical in question, but of the daughter products producing the Cerenkov emissions [59]. We utilized a thoroughly established murine model and relied upon the nanogenerator approach to keep 225Ac-DOTA-c(RGDyK) and its daughter products within tumor cells. However, inability to retain every daughter

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product after injection highlights the need for new materials, techniques, and mechanisms to effectively sequester the daughter products with the original conjugate to minimize the effects of this recoil energy. Several of these materials (including liposomes and LaPO4-based nanoparticles) have been described, and appear moderately effective in this arena [13, 60-64].

Preliminary toxicity and therapy studies demonstrated that low doses of 225Ac-DOTA-c(RGDyK) retarded tumor growth in mice 14 days after treatment. The potential of treating αβ3+ tumors with 225Ac-RGD-based therapies remains unknown, since larger studies are needed. αβ3+ malignancies demonstrate a diverse biology; understanding the effectiveness of these therapies among different cancer types will need to be elucidated [65]. Secondly, numerous reports have indicated that RGD multimerization increases tumor retention of several radiopharmaceuticals [54, 66-71]; 225Ac-radiopharmaceuticals containing RGD monomers and multimers will need to be compared in the TAT setting. Finally, the use of these radiopharmaceuticals with other therapies or technologies such as pulsed high intensity focused ultrasound may enhance uptake and retention of this therapeutic in tumors [72-74]. This is a future direction to explore so that the maximum clinical benefit of this technology can be achieved.

While renal impairment was only observed in animals receiving the highest doses of 225Ac-DOTA-c(RGDyK), we did not conduct long-term toxicity studies. In earlier work that described 212/213Bi-based radiopharmaceuticals, late-onset nephrotoxicity was observed in mice [75-77]. Diminished renal function is one of the most critical factors hindering TAT therapy development and its clinical implementation. With respect to 225Ac-radiotherapeutics, the damage caused by radiotracer reabsorption in the proximal tubules or by the localization of 213Bi and 221Fr daughter products in the renal tubule cells needs to be addressed, since damage caused by these daughter products can lead to nephropathy and ultimately, renal failure. Several strategies have been proffered to minimize kidney toxicity [57, 58]. These include amino acid co-infusion to block peptide reabsorption; angiotensin 1 blockade; chelation therapy to sequester 225Ac daughter products; and diuresis using spironolactone, furosemide or chlorothiazide. However, the choice of remediation and its efficiency will depend on the therapeutic under investigation [48].

In summary, this communication describes the evaluation of 225Ac-DOTA-c(RGDyK) in both biodistribution and Cerenkov luminescence imaging studies. This radiopharmaceutical was prepared in excellent radiochemical purity and specific activity that is in agreement with other 225Ac-based radiopharmaceuticals. It was highly stable in vitro and in vivo biodistribution studies confirmed the ability of the radiotracer to target αβ3+ integrin with specificity. Our results confirm earlier reports that CLI with certain α-emitting radiopharmaceuticals is possible in live animals as predicted by their theoretical and in vitro experiments [19, 20]. Furthermore, this concept may have broader implications for promising technological advancements in radiochemistry, radiopharmaceutical development, molecular imaging and the material sciences, and is worthy of further study. To our knowledge, this report is the first to describe Cerenkov luminescence imaging of αβ3+ tumors in live animals using the in situ decay of 225Ac and its daughter products. This work could provide new ways in which to further enhance targeted alpha particle therapy.

Supplementary Material
Supplementary schemes, figures and tables. http://www.thno.org/v06p0698s1.pdf

Abbreviations
CLI: Cerenkov Luminescence Imaging; ESI-MS: Electrospray Ionization Mass Spectrometry; HBS: HEPES Buffered Saline; ITLC: Instant Thin Layer Chromatography; kBq: Kilobecquerel; Kic: Inhibitory Constant; IC50: Half Maximal Inhibitory Concentration; MBq: Megabecquerel; %ID/g: Percent Injected Dose Per Gram of Tissue; P.I.: Post-injection; ROI: Region of Interest; RU: Response Unit; SPR: Surface Plasmon Resonance; TAT: Targeted Alpha Particle Therapy; TLC: Thin Layer Chromatography

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Competing Interests
The authors have declared that no competing interest exists.

References


