Cerenkov Luminescence Imaging for Radiation Dose Calculation of a ⁹⁰Y-Labeled Gastrin-Releasing Peptide Receptor Antagonist

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90Y has been used to label various new therapeutic radiopharmaceuticals. However, measuring the radiation dose delivered by 90Y is challenging because of the absence of suitable y emissions and its low abundance of positron emissions. For the treatment of prostate cancer, radiolabeled gastrin-releasing peptide receptor (GRPr) antagonists have yielded promising results in mouse models. In this study, we evaluated whether Cerenkov luminescence imaging (CLI) could be used to determine radiation doses of a 90Y-labeled GRPr antagonist in nude mice. Methods: Mice bearing subcutaneous prostate cancer xenografts were injected with 0.74-18.5 MBg of the 90Y-labeled GRPr antagonist DOTA-AR and underwent in vivo and ex vivo CLI at 1-48 h after injection. After imaging, animals were sacrificed, their tumors and organs were harvested, and the activity concentration was measured by liquid scintillation counting. In a second set of experiments, Cerenkov photon counts for tumor and kidney on in vivo CLI were converted to activity concentrations using conversion factors determined from the first set of experiments. Results: 90Y-DOTA-AR concentration in the 3 tumor models ranged from 0.5% to 4.8% of the injected activity per gram at 1 h after injection and decreased to 0.05%-0.15 injected activity per gram by 48 h after injection. A positive correlation was found between tumor activity concentrations and in vivo CLI signal (r^2 = 0.94). A similar correlation was found for the renal activity concentration and in vivo Cerenkov luminescence ($r^2 = 0.98$). Other organs were not distinctly visualized on the in vivo images, but ex vivo CLI was also correlated with the radioactivity concentration ($r^2 = 0.35$ – 0.94). Using the time-activity curves from the second experiment, we calculated radiation doses to tumor and kidney of 0.33 ± 0.12 (range, 0.21-0.66) and 0.06 \pm 0.01 (range, 0.05-0.08) Gy/MBq, respectively. Conclusion: CLI is a promising, low-cost modality to measure individual radiation doses of 90Y-labeled compounds noninvasively. The use of Cerenkov imaging is expected to facilitate the development and comparison of 90Y-labeled compounds for targeted radiotherapy.

Key Words: gastrin-releasing peptide receptor; Cerenkov luminescence imaging; targeted radiotherapy; $^{90}\mathrm{Y}$

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The β-emitter 90 Y is used for an increasing number of therapeutic radiopharmaceuticals, including the CD20 antibody ibritumomab tiuxetan, the somatostatin receptor ligands DOTATOC and DOTATATE, and microspheres for treatment of liver metastases (I–3). Because of its high β energy (Emax, 2.28 MeV), 90 Y can deliver therapeutically effective radiation doses at relatively low activity concentrations. Furthermore, the absence of γ emissions limits radiation exposure of personnel and individuals caring for the patient after radionuclide therapy. However, the lack of γ emission also represents a limitation of 90 Y-labeled pharmaceuticals, because quantitation of the activity delivered to the tumor and to dose-limiting normal organs is technically challenging (4). This is true not only for patient studies but also for preclinical studies in animals.

Rationale development of new ⁹⁰Y-labeled pharmaceuticals requires a comparison of radiation doses to tumor and normal tissues in animal models. To achieve this, the conventional approach is to perform serial biodistribution studies in large cohorts of animals to measure tissue activity concentrations at various time points and to determine the cumulated activity for each tissue of interest. Measuring activity concentrations at a single time point may be misleading because radiopharmaceuticals with different clearance rates and residence times may deliver different radiation doses despite a similar initial uptake. Noninvasive techniques to measure the concentration of ⁹⁰Y in animals are also needed to correlate the delivered radiation dose with growth inhibition or regression in individual animals.

 $^{90}\mathrm{Y}$ can be imaged by SPECT of the bremsstrahlung x-rays emitted after the β decay or by PET using positrons generated by internal electron-positron pair production (5,6). However, internal pair production is a rare event, limiting count statistics and requiring long acquisition times. Bremsstrahlung shows a broad, continuous spectrum with maximum x-ray energies of 2.28 MeV and is therefore difficult to image quantitatively using SPECT (7).

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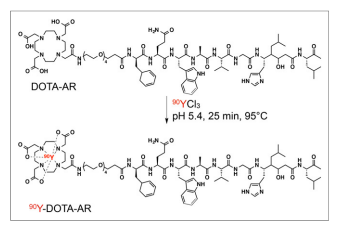


FIGURE 1. Structure of 90Y-DOTA-AR.

An alternative approach to measure β -emitting isotopes is Cerenkov luminescence (CL), a continuous spectrum of light emitted by B particles as they travel through a dielectric medium such as tissue (8,9). This light signal can be detected by sensitive luminescence imaging systems that are used in many laboratories for bioluminescence imaging (10–14). Because the intensity of CL produced is dependent on the energy of the β radiation, ⁹⁰Y is better suited for Cerenkov luminescence imaging (CLI) than lowenergy positron emitters such as ¹⁸F. Phantom studies by Liu et al. (15) have shown that 90Y emits about 10 times more detectable Cerenkov light per radioactive decay than ¹⁸F. These investigators have also demonstrated the feasibility of CLI of tumor xenografts with ⁹⁰Y-labeled peptides (15). An additional advantage of 90 Y is the absence of γ emissions, which generate noise in the optical detector system (16). In a recent study, the higher β energy and the lack of γ emissions resulted in an overall 270fold-higher sensitivity of Cerenkov imaging with 90Y as compared with ¹⁸F (16).

The purpose of the present study was therefore to evaluate CLI as a quantitative imaging modality to estimate the radiation dose of ⁹⁰Y-labeled radiopharmaceuticals in mice. Our hypothesis was that CLI will allow rapid, noninvasive, serial measurements of activity concentrations in situ radiation that can be used to estimate radiation doses to tumor and normal organs. We tested this hypothesis for a ⁹⁰Y-labeled gastrin-releasing peptide receptor (GRPr or bombesin receptor) antagonist in mice bearing prostate cancer xenografts. Prostate cancer remains the second-leading cause of cancer death in men in the United States (*17*), hence there remains an enormous need for new therapeutic strategies, especially for locally advanced and metastatic disease.

MATERIALS AND METHODS

The GRPr antagonist DOTA-AR (Fig. 1) was selected on the basis of its excellent GRPr-targeting efficacy and synthesized as previously described by Abiraj et al. (18). ⁹⁰Y-DOTA-AR was prepared by dissolving 20 µg of peptide (11.4 nmol) in ammonium acetate buffer (200 mL, 0.5 M, pH 5.4) (Fig. 1). The radiochemical yield and purity were typically 80%–90% (decay corrected) and more than 95%, respectively. After the addition of ⁹⁰YCl₃ (370 MBq), the solution was incubated for 25 min at 95°C (Fig. 1). A hydrophilic–lipophilic balance cartridge preconditioned with methanol (5 mL) and water (5 mL) was used to remove the free ⁹⁰Y from the ⁹⁰Y-DOTA-AR. After hydrophilic-lipophilic balance purification, the radioligand, with a specific activity

of 9.62-18.5 MBq/ μ g, was diluted in 0.9% NaCl with 0.1% bovine serum albumin for in vivo studies.

The GRPr-expressing human prostate cancer cell lines PC-3 (high expression of GRPr), VCaP (high expression of GRPr), and LNCaP (low expression of GRPr) were obtained from American Tissue Culture Collection and cultured in RPMI-1640 medium (MSKCC Media Preparation Facility), supplemented with 10% fetal bovine serum (HyClone) and penicillin (10,000 IU/mL) and streptomycin (10,000 µg/mL).

Experiments were performed according to a protocol approved by the Institutional Animal Care and Use Committee. Xenografts of PC-3, VCaP, and LNCaP cells were established by the subcutaneous injections of 5×10^6 tumor cells suspended in 150 μ L of 1:1 respective cell growth medium/Matrigel (BD Biosciences) in the area of the left shoulder of male athymic nude (outbred)/CrTac:NCr-Foxn1nu mice (7–9 wk; weight, ~20 g each; Taconic). When tumors reached a volume of approximately 150 mm³ (at 14–28 d after implantation), experiments were performed.

CLI

Fifteen mice bearing PC-3 prostate cancer xenografts were injected into the lateral tail vein with an activity of 11.1 MBq. In vivo and ex vivo CLI was performed at 1, 4, 24, and 48 h after injection. For determination of nonspecific uptake, a group of 3 PC-3 tumor–bearing mice was preinjected (5 min) with 154 nmol of unlabeled GRPr antagonist HZ201 ([D-Phe⁶]-BN(6-13)-ethylester) into the lateral tail vein, followed by 2.775 MBq of ⁹⁰Y-DOTA-AR. To test the feasibility of CLI for other prostate cancer models, we injected groups of 3 mice bearing VCap and LnCAp tumors with 2.775 MBq of ⁹⁰Y-DOTA-AR and performed in vivo and ex vivo imaging at 4 h after injection.

For in vivo imaging, CLI mice were anesthetized with isoflurane (1%–4%). CL images of 5-min duration were acquired with an IVIS 200 scanner (Caliper Life Sciences). For quantification of CLI signals, regions of interest were placed over the tumor and kidneys, which were both well visualized on the CL images. After the in vivo scan, the mice were sacrificed and dissected. Tumors and organs were placed in a tissue culture dish for ex vivo CLI, which was performed using the same scanner and acquisition parameters as for in vivo CLI. Regions of interest were manually drawn around tumors and normal organs on the in vivo and ex vivo images. CL intensity was expressed

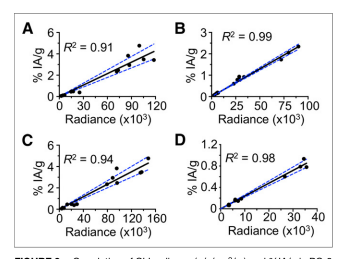


FIGURE 2. Correlation of CLI radiance (p/s/cm²/sr) and %IA/g in PC-3 tumor–bearing mice injected with 11.1 MBq of ⁹⁰Y-DOTA-AR. (A) Ex vivo data tumor. (B) Ex vivo data kidney. (C) In vivo data tumor. (D) In vivo data kidney. Graphs include data from tissues sampled from 1 to 48 h after injection.

TABLE 1Biodistribution Data of ⁹⁰Y-DOTA-AR in Nude Mice Bearing Tumors

				48 h
Tumor				
PC-3	3.62 ± 0.95	2.92 ± 0.57	0.45 ± 0.04	0.10 ± 0.05
LNCaP		0.29 ± 0.16		
VCaP		3.49 ± 0.50		
Liver	0.21 ± 0.05	0.12 ± 0.06	0.04 ± 0.01	0.02 ± 0.00
Kidney	1.76 ± 0.61	0.78 ± 0.14	0.17 ± 0.02	0.07 ± 0.01

as average radiance in number of photons (p) per second (s) per surface area (cm²) per steradian (sr), that is, p/s/cm²/sr.

After in vivo and ex vivo CLI, tumor and normal tissues were weighed, solubilized overnight in 1 mL of a SOLVABLE tissue solubilizer (PerkinElmer), and diluted to 5 mL with a commercial scintillation fluid cocktail (Insta-Fluor Plus; PerkinElmer) for liquid scintillation counting (LCS) (Tri-Carb 2910TR; PerkinElmer). The total counts injected per animal were determined by extrapolation from counts of an aliquot taken from the injected solution. The biodistribution data were expressed as percentage injected activity per gram of tissue (%IA/g).

Because we observed a linear relationship of the LCS-derived activity concentrations and the CL light signal with no obvious intercept (Fig. 2), we used linear regression to calculate a calibration factor, K, for conversion of radiance to activity concentration:

Activity concentration [MB q/g] = $K \times radiance [(p/s/cm^2/sr)]$.

Absorbed Dose Calculations

To evaluate the ability to measure individual tumor time–activity curves by CLI, 14 mice bearing PC-3 tumors were divided into 5 groups, and 90 Y-DOTA-AR was injected into the lateral tail vein with an activity of 0.74, 1.85, 3.7, 7.4, and 18.5 MBq (injected amount 0.02–0.57 nmol). In vivo CLI was performed in each group at 4, 24, 48, and 72 h after injection of 90 Y-DOTA-AR. The measured photon counts for Cerenkov radiation (CR) were converted to activity concentrations using the conversion factors determined in the experiments described above.

From these data, the ⁹⁰Y tumor- and kidney-absorbed doses (Gy/MBq) were derived as follows. The tumor and kidney time-activity data (%IA/g) were fit to biexponential functions, and the resulting

functions were analytically integrated, incorporating the effect of the physical decay of the ^{90}Y to derive the respective residence times (MBq h/g/MBq). The resulting residence time was then multiplied by the ^{90}Y equilibrium dose constant for nonpenetrating radiations (i.e., β particles), 0.538 g-Gy/MBq-h. To correct for partial absorption of β particles, these doses were multiplied by a volume-based absorbed fraction to calculate absorbed doses (Gy/MBq) to the tumor and kidney. Tumor and kidney volumes were derived from the 2-dimensional area on the white light images assuming ellipsoid geometry. Absorbed fractions for ^{90}Y were taken from data published by Bardiès and Chatal (19).

RESULTS

The biodistribution of $^{90}\text{Y-DOTA-AR}$ in tumor-bearing nude mice is briefly summarized in Table 1 and shown in detail in Supplemental Table 1 (supplemental materials are available at http://jnm.snmjournals.org). Activity concentrations in PC-3 tumors (mean \pm SD) were 3.62 \pm 0.95 %IA/g at 1 h and 2.92 \pm 0.57 %IA/g at 4 h, with washout by later time points (0.45 \pm 0.04 %IA/g at 24 h; 0.10 \pm 0.05 %IA/g at 48 h).

The pancreas, kidneys, and lung were the organs with the highest activity concentrations at the 1-h time point (Supplemental Table 1). Except for the lung, all normal organs showed washout of activity at the 4-h time point, resulting in increased tumor–to–normal organ ratios at the 4-h time point. $^{90}\text{Y-DOTA-AR}$ demonstrated a fast blood clearance, with only 0.03 ± 0.03 %IA/g remaining in the blood at 4 h after injection.

The ex vivo CLI radiance data for 90 Y-DOTA-AR (Table 2; Figs. 2A and 2B) show the same pattern as the activity concentration

TABLE 2Ex Vivo CLI Radiance of ⁹⁰Y-DOTA-AR in Nude Mice Bearing Tumors

Organ	1 h	4 h	24 h	48 h
Tumor				
PC-3	92,779 ± 1,886	$84,050 \pm 1,604$	14,087 ± 3,776	2,658 ± 1,267
LNCaP		$2,808 \pm 500$		
VCaP		52,941 ± 19,524		
Liver	$7,324 \pm 1,948$	3,885 ± 1,751	1,368 ± 217	681 ± 118
Kidney	67,834 ± 2,429	25,650 ± 2,661	4,513 ± 629	1,663 ± 201

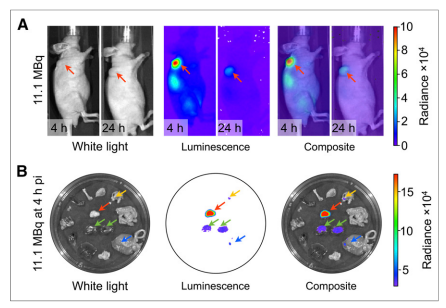


FIGURE 3. CLI of 90Y-DOTA-AR in PC-3 tumor-bearing nude mice. (A) Tumors in area of left shoulder (in vivo white light images) demonstrate high radiance (p/s/cm²/sr) at 4 after injection and moderate radiance at 24 h after injection. Kidney is also well visualized on 4-h images. (B) Ex vivo images of 4 h after injection clearly delineate radiance for tumor, kidneys, and in some organs not seen on in vivo images due to absorption of CL signal. Green arrows = kidneys; yellow arrow = stomach; blue arrow = large intestine; red arrow = tumor. pi = after injection.

measurements. Furthermore, linear regression analysis generally demonstrated a close correlation between 90Y-DOTA-AR activity concentration (%IA/g) and ex vivo CR (p/s/cm²/sr). For example, the correlation coefficient (R^2) between ex vivo CLI radiance and activity concentration was 0.91 for PC-3 tumors (Fig. 2A) and 0.99 for kidneys (Fig. 2B). Figure 3 shows an example of ex vivo CLI at 4 h. Correlations between ex vivo CLI and

VCap LnCap Α В Tumor Kidney 5 10 _ % IA/g MBq at 4 3 8 Radiance ×104 ω Composite Composite C ■ Tumor Ē Radiance x10³ Kidney .8 MBq at 4 h 2 Lucapucap Luminescence Luminescence

FIGURE 4. CLI of ⁹⁰Y-DOTA-AR in representative VCaP and LNCaP tumor-bearing nude mice. (A) Tumors in area of left shoulder demonstrate high radiance in VCaP and low radiance in LNCaP tumors on in vivo luminescence and composite images. (B) Graph demonstrates %IA/g for VCaP (n = 3) and LNCaP (n = 2) tumors. (C) Graph demonstrates radiance for VCaP (n = 3) and LNCaP (n = 2) tumors. Radiance is expressed in p/s/cm2/sr. pi = after injection.

activity concentrations for other organs are shown in Supplemental Figure 2.

The tumor and kidneys were also visualized by in vivo CLI (Fig. 3). For both tissues, there was a significant correlation between in vivo CLI radiance and ex vivo activity concentrations (PC-3 tumors, $R^2 =$ 0.94; kidneys, $R^2 = 0.98$) (Figs. 2C and 2D). The slope of the regression line was 2.9×10^{-5} (%IA/g)/(p/s/cm²/sr) or $3.145 \times$ 10^{-6} (MBq/g)/(p/s/cm²/sr) for the tumors and 2.4×10^{-5} (%IA/g)/(p/s/cm²/sr) or $2.627 \times 10^{-6} \text{ (MBq/g)/(p/s/cm}^2/\text{sr)}$ for the kidneys.

As expected, LNCaP tumors demonstrated a low CL, whereas uptake by VCaP tumors was comparable to PC-3, reflecting the relative GRPr expression levels of the 3 tumor cell lines; the CL correlated to a marked difference in radioactivity concentrations between LNCaP and VCaP tumors (Tables 1 and 2; Fig. 4). Cerenkov imaging was also able to document blocking of the binding of radiolabeled 90Y-DOTA-AR with an unlabeled GRPr ligand (Fig. 5). Both the Cerenkov signal and radioactivity concentrations were reduced by

about 80% by preinjection of an approximately 300-fold excess of the unlabeled GRPr antagonist HZ201(Fig. 5).

Figure 6A shows an example of a mouse that was imaged by CLI at various time points after injection of 0.74 MBq of 90Y-DOTA-AR. Both tumors and kidneys were visualized for injected activities ranging from 0.74 MBq to 18.5 MBq at 4 and 24 h after injection. At later time points, the kidney was only visible for mice

> that had been injected with at least 3.7 MBq. Using these serial images and the calibration factors from the previous set of experiments. we estimated the radioactivity concentration in the tumor and the kidneys over time (Figs. 6B, 6C, and 7). A dose-dependent decrease of the %IA/g in the tumor with increasing administered activities was apparent; presumably, the receptors are partially saturated at higher injected activities/ peptide masses. No such effect was observed for the kidneys, which do not express GRPrs. Overall, tumor doses showed a more than 3-fold variability and ranged from 0.21 to 0.66 Gy/MBq. Renal doses were also variable and ranged from 0.05 to 0.08 Gy/MBq. The calculated absorbed fraction doses to tumor and kidneys of individual animals are shown in Table 3.

DISCUSSION

The success of peptide receptor-targeted radiotherapy in neuroendocrine tumors (2) has renewed the interest in treating cancer with β-emitting radioisotopes. Radioloabeled GRPr antagonists have shown promise for

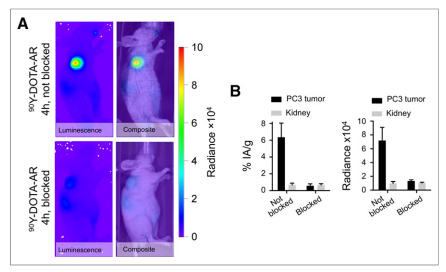


FIGURE 5. (A) CLI in blocked and nonblocked PC-3 tumor–bearing representative nude mice after injection of 90 Y-DOTA-AR. Tumors in area of left shoulder demonstrate high radiance in nonblocked tumors and low radiance in blocked tumors on in vivo luminescence and composite images. (B) Graphs demonstrate %IA/g and radiance for blocked (n=3) and nonblocked (n=3) tumors. Radiance is expressed in p/s/cm2/sr.

the treatment of prostate cancer in preclinical studies because of their favorable biodistribution and high tumor uptake (20–26). These findings were confirmed in the present study, which demonstrated a similar biodistribution profile of the GRPr antagonist ⁹⁰Y-DOTA-AR, as pre-

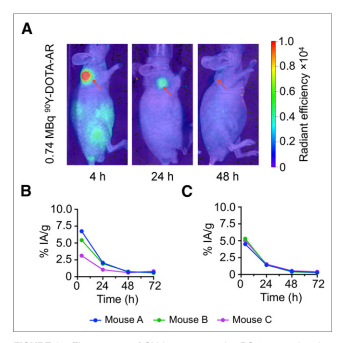


FIGURE 6. Time course of CLI in representative PC-3 tumor–bearing mouse after injection of 0.74 MBq of ⁹⁰Y-DOTA-AR. (A) In vivo composite CLI clearly demonstrates high radiance (p/s/cm2/sr) in PC-3 tumor at 4 h and still moderate radiance at 24 h after injection (red arrow). No increased CR signal was detected at 48 h after injection time point (red arrow). Corresponding CLI-based calculated time–activity curves for individual mice with injected doses of 0.74 MBq (B) and 7.4 MBq (C) of radioactivity. Percentage of activity accumulated by tumor decreases for higher injected activity due to partial blocking of receptor by higher injected peptide mass.

viously reported for other radiolabeled GRPr antagonists (18,22,25).

The biodistribution of radiopharmaceuticals is influenced by multiple factors, including target density, regional perfusion, vascular permeability, and renal and hepatic function. These parameters may well differ among individual animals and can change over time within an individual animal. Furthermore, labeling of the same molecule with different isotopes can markedly affect target binding affinity. For example, the affinity of 68Ga-DOTATOC for the somatostatin type 2 receptor is more than 4-fold higher than that of 90Y-DOTATOC (27). Therefore, it can be difficult to predict the uptake of a 90Y-labeled peptide based on the uptake of the same peptide labeled with a different radiometal. Thus, it is desirable to directly measure concentrations of ⁹⁰Y during therapy both noninvasively and repetitively.

One approach to achieve this goal is to visualize and quantify the CL of ⁹⁰Y. The

proof-of-principle work by Robertson et al. (28) used CLI in vitro and in vivo to image human prostate cancer xenografts after injection of different β-emitting radioisotopes. Several reports followed and demonstrated successfully the application of optical imaging with β^+ and β^- emitters (10–13,16,17,29,30) in preclinical models. CLI has also been used clinically during therapy of hyperthyroidism with ¹³¹I (31). Recently, it has been shown that CLI with ¹⁸F-FDG allows detection of superficial lymph node metastases in patients (32).

However, CLI is especially promising for imaging during therapy with 90 Y, because the amount of Cerenkov photons is dependent on the β energy. Although 18 F (β^+ with a positron endpoint energy of 633 keV) will create 1.4 photons per decay (in water), 90 Y (β^- with an endpoint energy of 2.28 MeV) can produce up to 57 photons per decay (29,33). The wavelength of CL lies mostly within the visible and ultraviolet part of the spectrum, which leads to strong attenuation in tissue. Furthermore, the scanning must be performed while carefully excluding external light sources because of the low overall amount of available photons, requiring low background photon counts. However, this procedure does not present a major obstacle for imaging subcutaneous tumors in nude mice, with widely available systems for bioluminescence imaging—a technique that also requires acquisition without ambient light.

In the present study, we show a close correlation between ex vivo CLI and radioactivity concentrations measured by LCS. Ex vivo CLI is much less time-consuming than LCS, which requires tissue homogenization. Therefore, CLI could facilitate comparison of the biodistribution of different ⁹⁰Y-labeled radiotracers in mice. More important, we also found a close correlation between in vivo CLI and concentration of ⁹⁰Y in subcutaneous tumors, allowing us to confirm receptor-specific uptake of the ⁹⁰Y-labeled peptide by blocking studies and in vivo imaging. We were also able to non-invasively study the effect of increasing peptide mass on tumor activity concentrations. Finally, serial CLI studies allowed us to noninvasively estimate the radiation dose for tumors and kidneys in individual mice. This noninvasive estimation represents a significant advantage over conventional biodistribution studies that pool

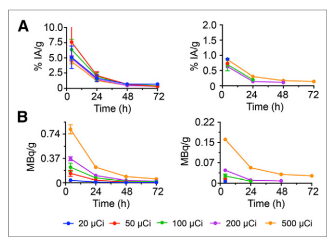


FIGURE 7. PC-3 tumors and kidneys in mice (n=14) injected with 0.74–18.5 MBq of ⁹⁰Y-DOTA-AR. (A) Calculated time–activity (%IA/g) curves for tumors (left) and kidneys (right). (B) Calculated time–activity (MBq/g) curves for tumors (left) and kidneys (right).

data from multiple animals. To our knowledge, CLI is the only modality capable of this kind of in vivo characterization of ⁹⁰Y-labeled peptides in such a facile and cost-effective way, as data of up to 5 animals can be acquired simultaneously within 5 min.

An alternative to CLI of ⁹⁰Y-labeled compounds is PET with ⁸⁶Y (*34*). However, average PET examination times in a preclinical setting account for approximately 20 min per mouse. Hence, many more mice can be evaluated using CLI than PET. Furthermore, ⁸⁶Y is an expensive radioisotope and is not commonly available, significantly limiting the ability to perform larger studies using ⁸⁶Y PET.

Although the findings of our study are encouraging for the use of CLI during preclinical development of 90 Y-labeled pharmaceuticals, the following limitations should be noted. Although superficial tumors and kidneys were well visualized by CLI, more deepseated organs, such as the pancreas, could not be delineated by CLI. Thus, the ability of CLI to noninvasively estimate doses to deep-seated normal organs or orthotopic tumors appears limited. For radiopeptides, however, the kidney is frequently the dose-limiting organ. Therefore, we expect that CLI can provide important information for the development of many 90 Y-labeled peptides.

It is also important to note that the rapid clearance of ⁹⁰Y-DOTA-AR resulted in low background activity concentrations, which facilitated visualization of tumors and quantification of intratumoral activity concentrations. Future studies are required to assess the accuracy of quantitative CLI for radiopharmaceuticals with less-selective tumor targeting. However, therapeutic radiopharmaceuticals will generally require high tumor–to–normal organ ratios and it is therefore likely that CLI will allow the quantitative assessment of intratumoral activity concentrations for other therapeutic radiopharmaceuticals as well.

Because of the short range of CL, the intensity of the CLI signal is mainly determined by the activity concentration at the surface of a tumor. Consequently, CLI will overestimate the total activity concentration in tumors with necrotic centers. For the same reasons, the tumor CL signal is dependent on the positioning of the animals, thus necessitating standardization for imaging studies.

CONCLUSION

CLI is a promising, low-cost modality to measure individual radiation doses by ⁹⁰Y-labeled peptides for tumor xenografts and kidneys in mice. The use of Cerenkov imaging is expected to

TABLE 3Absorbed Fraction Doses (Gy/MBq) of ⁹⁰Y-DOTA-AR in Tumors/Kidneys

Mouse no.	Injected activity (MBq)	Peptide mass (ng)	Tumor	Kidney
1	0.74	40	0.45	
2	0.74	40	0.35	
3	0.74	40	0.21	
			0.34 (average)	
4	1.85	100	0.66	
5	1.85	100	0.36	
6	1.85	100	0.26	
			0.43 (average)	
7	3.7	200	0.26	0.05
8	3.7	200	0.36	0.05
9	3.7	200	0.44	0.07
			0.35 (average)	0.06 (average)
10	7.4	400	0.24	0.05
11	7.4	400	0.22	0.05
12	7.4	400	0.25	0.04
			0.24 (average)	0.05 (average)
13	18.5	1,000	0.23	0.08
14	18.5	1,000	0.28	0.06
			0.26 (average)	0.07 (average)

facilitate the development and comparison of β emitters for targeted radiotherapy.

DISCLOSURE

The costs of publication of this article were defrayed in part by the payment of page charges. Therefore, and solely to indicate this fact, this article is hereby marked "advertisement" in accordance with 18 USC section 1734. This study was supported in part by the R25T Molecular Imaging Fellowship "Molecular Imaging Training in Oncology" (5R25CA096945-07) and by the NIH (1R01EB01494). J.G. was additionally supported by a NIBIB award 5R01EB014944-03 and NCI award 1R01CA183953. Technical services provided by the MSK Small Animal Imaging Core Facility were supported in part by NIH Cancer Center Support grant 2 P30 CA008748-48. No other potential conflict of interest relevant to this article was reported.

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