Quality analysis of *in vivo* near-infrared fluorescence and conventional gamma images acquired using a dual-labeled tumor-targeting probe

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Abstract. The cyclic peptide, cyclopentapeptide cyclo(lys-Arg-Gly-Asp-phe) (c(KRGDF)), which is known to target αvβ3 integrin, is dual-labeled with a radiotracer, 111In, for gamma scintigraphy as well as with a near-infrared dye, IRDye800, for continuous-wave (cw) imaging of αvβ3 positive human M21 melanoma in xenografts. Twenty-four hours after administration of the dual-labeled peptide at a dose equivalent to 90 μCi of 111In and 5 nmol of near-infrared (NIR) dye, whole-body gamma scintigraphy and cw imaging was conducted. Image acquisition time was 15 min for the gamma scintigraphy images and 800 ms for the optical images acquired using an NIR sensitive intensified charge-coupled device. The results show that while the target-to-background ratio (TBR) of nuclear and optical imaging were similar for surface regions of interest and consistent with the origin of gamma and NIR radiation from a common targeted peptide, the signal-to-noise ratio (SNR) was significantly higher for optical than nuclear imaging. Furthermore, an analysis of SNR versus contrast showed greater sensitivity of optical over nuclear imaging for the subcutaneous tumor targets. While tomographic reconstructions are necessary to probe TBR, SNR, and contrast for interior tissues, this work demonstrates for the first time the direct comparison of molecular optical and planar nuclear imaging for surface and subsurface cancers. © 2005 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.2114748]

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1 Introduction

Nuclear imaging is currently the most effective available medical imaging modality for measuring tracer uptake associated with disease markers as well as with tissue metabolism. Nuclear medicine is prominent in cancer diagnosis, treatment, and prognosis and generally involves the injection of a radionuclide that is targeted for disease specificity. Typically, gamma ray emitting radionuclides are chelated to moieties that are biocompatible, interact with disease markers, or have affinity towards tumor sites.1–10 Gamma emission can be collected by a gamma camera, producing a scintigram, or a planar projection image that depicts the location of the radionuclide. In addition to planar projection images, tomographically reconstructed images are possible with single photon emission computed tomography (SPECT) and positron emission tomography (PET).

The nuclear tomographic technique of SPECT is based on detecting individual gamma rays emitted at random from the radionuclide, and the tomographic technique of PET is based upon positron emitters, which interact with the electron rich environment subsequently annihilating to produce two high-energy photons that travel in the opposite direction away from the position of their generation. Their colinear emission enables back-projection to form an image of the radionuclide location. These tomographic imaging approaches may provide millimeter resolution and are limited by the need for ionizing radiopharmaceuticals with finite half-lives and long camera integration times.

Near-infrared (NIR) fluorescence-enhanced optical imaging is similar to nuclear imaging in that a disease targeting molecule conjugated to a radiation-emitting (albeit non-ionizing radiation) agent is injected into the body for localization to the diseased site. Upon activation with excitation light, the agent emits fluorescent photons which are then detected externally by sensitive photon detectors. In contrast to nuclear agents, fluorescent agents do not become inactive signal generators after emission but instead are available for repeated activation and relaxation. Excitation and emission in the NIR range are required for imaging tissue, as these energies have low absorption in blood, melanin, and water en-
abling penetration in tissues. In contrast to nuclear imaging, high photon count rates enable increased temporal resolution for dynamic imaging.\textsuperscript{11,14} Optical imaging may become a significant molecular imaging tool since it provides digital photo-quality images with low noise and high photon signal emission that can be quantified for pharmacokinetic information\textsuperscript{11,12} as well as tomography.\textsuperscript{13,14}

Optical imaging in tissues using fluorescence can be employed via several strategies. The imaging strategies generally encompass: (i) detection of endogenous fluorescence; (ii) detection of exogenous fluorescence that is targeted to disease via antibody, polymer, peptide, recombinant protein, or other biocompatible conjugates, or otherwise nontargeted; and (iii) detection of light emitted from quantum dots or other inorganic particles. Detection of bioluminescence represents an optical imaging technique, although not an excitation-emission process. The measurement of fluorescent light emitted from the tissue surface by these strategies may be possible with sensitive image intensifiers coupled to charge-coupled device cameras to provide planar images over the whole body of small animals. Optical imaging by intensified charge-coupled device (ICCD) systems is comparable to conventional nuclear scintigraphy. As ICCDs collect fluorescent photons for image generation, gamma cameras detect gamma emissions and render images that are processed postacquisition to identify diseased tissue sites. Herein we directly compare optical and nuclear imaging using a dual labeled, targeting fluorescence and nuclear agent.

There are presently few examples reported in the literature involving the combination of nuclear and optical imaging. Most studies with small animals have involved reporter gene constructs that either express (i) a visible or red fluorescent protein for excitation and emission (fluorescence); (ii) Renilla or firefly luciferase, which produces visible light in the presence of its exogenous substrate, coelenterazine or D-luciferin (bioluminescence); and/or (iii) express the wild type or mutant herpes simplex virus type 1 thymidine kinase (genotype: HSV1-tk and HSV1-sr39tk, respectively), which phosphorylates PET labeled substrates such as FIAU (2′-[18F]fluoro-2′-deoxy-1-β-arabinofuranosyl-5-iodouracil), FHBG (9-[4-18F]fluoro-3-hydroxymethylbutyl)guanine), and FPCV (8-[18F]fluoropenciclovir).\textsuperscript{15–19} These nuclear-optical reporter genes may permit enhanced specificity of the disease with a selectively activated upstream promoter. Unfortunately, the near-term clinical application of gene reporter systems is limited by the need for stable gene transfection and transformation in patients. In this work, we seek to directly compare optical and nuclear imaging using a single targeting moiety containing both a fluorophore and a radionuclide that is exogenously introduced in trace diagnostic concentrations. In previous work, Frangioni and coworkers compared optical and nuclear imaging modalities by evaluating osteoblastic activity in a male nude mouse using a targeting nuclear probe and a second targeting optical probe in order to provide complementary optical and nuclear images for comparison.\textsuperscript{20} While other investigators\textsuperscript{4} have employed nuclear and optical probe combinations, to date there have been no studies that employ a single, dual-labeled exogenous agent for direct comparison of optical and nuclear imaging modalities. Herein, we use a single probe, dual labeled to provide direct comparison of optical and nuclear molecular imaging techniques using established figures of merit.

We present a dual-labeled \textsuperscript{111}indium (\textsuperscript{111}In) and NIR dye, IRDye800, on the cyclopentapeptide cyclo(lys-Arg-Gly-Asp-phe), (c(KRGDf)), which is known to target \(\alpha\beta\) integrin expressed in human melanoma.\textsuperscript{21} The use of this probe enables us to image the interaction between a tumor receptor and a targeting ligand in a mouse model by both conventional nuclear imaging and fluorescence-enhanced optical imaging techniques. Since the melanoma is located subcutaneously on the animal, diffusion-based tomography is not valid over the small volumes and short length scales.\textsuperscript{22} Furthermore, since direct comparison of fluorescent intensity to gamma ray signals is desired, we conduct the optical imaging using simple, continuous-wave (cw) measurements. The acquired nuclear scintigraphy and cw optical images enable an analysis on the differences between optical and nuclear image qualities.

## 2 Materials and Methods

Fluorescence enhanced optical imaging with an ICCD camera and planar gamma imaging with a small animal gamma camera were performed on a total of six nude mice bearing human melanoma tumors.

### 2.1 Dual-labeled Contrast Agent

For the dual-labeled contrast agent, a peptide containing the amino acid sequence arginine-glycine-aspartic acid, which is known to bind to \(\alpha\beta\) integrin,\textsuperscript{23,24} was used. Specifically, Lys-c(KRGDf) was synthesized on linker-PL-DMA resin using Fmoc solid phase chemistry as previously described.\textsuperscript{21,25} The radiometal chelator, P-succinimidobenzyl diethylenetriaminepenta-acetic acid, (DTPA) derivative, was first reacted with the \(\omega\)-amino group of the Lys unit in Lys-c(KRGDf). Subsequently IRdye800 was conjugated to the \(\omega\)-amino group of the Lys unit in DTPA-Lys-c(KRGDf) to give DTPA-Lys(IRDye800)-c(KRGDf). Lastly, the DTPA-Lys(IRDye800)-c(KRGDf) was mixed with \textsuperscript{111}InCl\textsubscript{3} to form the final product, \textsuperscript{111}In-DTPA-Lys(IRDye800)-c(KRGDf). The NIR dye (IRDye800) excitation/emission wavelengths were measured at 785/830 nm.

### 2.2 Animal Studies

For animal preparation, four- to six-week-old female athymic nude mice (nu/nu 13-21 g) were purchased from Harlan Sprague Dawley, Inc. (Indianapolis, IN). Well-characterized human melanoma tumor cells positively expressing the \(\alpha\beta\) integrin receptors (M21) and human melanoma tumor cells not expressing the \(\alpha\beta\) integrin receptors (M21-L) were maintained at 37°C in a humidified atmosphere containing 5% \(\text{CO}_2\) in Dulbecco’s modified Eagle’s medium with F12 nutrient and 10% fetal bovine serum. Cells from the positive and negative integrin lines were implanted subcutaneously into the right and left hind region of the six nude mice, respectively. The M21 and M21-L tumor cell lines were kindly provided by Dr. Cheresh (The Scripps Research Institute, La Jolla, CA). The \(\alpha\beta\) integrin positive and integrin negative melanoma cell lines have been used in several studies to in-
vestigate integrin expression and activity in cell signaling, angiogenesis, and proliferation. Cheresh and Spiro originally explored the M21 variants by looking at receptor affinity to the RGD amino acid sequence. This investigation led to the isolation of the M21-L, or “M21 Lows,” variant via affinity to the RGD amino acid sequence. This investigation led to the isolation of the M21-L, or “M21 Lows,” variant via affinity to the RGD amino acid sequence. This investigation led to the isolation of the M21-L, or “M21 Lows,” variant via.

Imaging commenced 24 h after administration of the dual-labeled conjugate. Before imaging, the mice were tagged for identification and anesthetized with Nembutal (50 mg/kg body weight). Animals were then imaged individually with the NIR optical imaging system. After optical imaging, the animals were transferred approximately 15 min later to the nuclear gamma camera system to acquire planar scintigraphy images. The nuclear imaging system and NIR optical imaging system are described below.

Animals were maintained in a pathogen-free mouse colony in the Department of Veterinary Medicine (The University of Texas M. D. Anderson Cancer Center, Houston, TX). The facility is accredited by the American Association for Laboratory Animal Care and all experiments were performed in accordance with the guidelines of the Institutional Animal Care and Use Committee.

2.3 M.C.AM Small Animal Gamma Imaging System

Conventional gamma imaging is accomplished in the following manner: (1) gamma rays upon $^{111}$In decay are emitted isotropically from within the animal, (2) the emitted gamma rays are collimated, (3) the collimated gamma rays strike a scintillation crystal where visible photons (scintillation light) are produced, and (4) the photons are then guided towards a photomultiplier tube (PMT), which collects a ray sum of scintillation light used to generate the projection image.

Conventional gamma imaging was performed in this study with an M.C.AM small animal gamma camera (Siemens, Hoffman Estates, IL). This gamma camera is controlled by two central processing units for nuclear acquisition and data processing. The M.C.AM houses a high-definition digital detector, which is mounted on a stand that can be manually adjusted and rotated over 90 deg. A collimator is fitted over the entire detector plane and can be removed for replacement. The collimator used for this study was a medium energy low penetration (MELP) with a resolution of 4 to 20 mm and sensitivity of 237 cpm/μCi. A placement template, covering the collimator, defined the area of the scintillation crystal for placement of the animals over the fixed field of view (FOV) of $54.4 \times 39.7$ cm$^2$. Images were acquired at a 237 keV energy window and by integrating the detector over a time interval of 15 min for which a static planar gamma image was acquired.

The M.C.AM system produces a total imaging matrix size of $1024 \times 1024$ pixels, and the image type was read as a 16-
bit multi pixel array providing intensity counts on the order of 2\(^{16}\) levels of gray.

2.4 NIR-Fluorescence-enhanced ICCD Imaging System

Continuous-wave optical imaging of fluorescent NIR photons was performed with a previously developed small animal optical imaging system.\textsuperscript{11} The system consists of two complementary detectors, a NIR sensitive image intensifier (model FS9910C ITT Night Vision, Roanoke, VA) and a 16-bit dynamic range full frame Photometrics (model CH350, Roper Scientific, Trenton, NJ) CCD camera. The image intensifier captured the fluorescent NIR signal upon emission from the melanoma tumor site via a 28- to 80-mm zoom Nikon lens (Nikon, Japan). The animal’s whole body, an equivalent FOV detection diameter of 11.5 cm (or 8.1 cm for three animals), was focused onto the entire 18-mm-diameter intensifier tube. The image intensifier was selected based upon its optimum photocathode photoresponse [\(\mu\text{A/lumen}\)] in the NIR. The full-frame CCD camera imaged the 550-nm amplified light signal output at the intensifier’s phosphor screen via a 40-mm Nikon lens (Nikon, Japan). Likewise, to optimize the signal obtained by the NIR system, the CCD camera was designed for a favorable 80% quantum efficiency at 550 nm.

Fluorescence imaging was performed first by expanding a 785-nm excitation source over the body. Here, we used a Thorlabs (80 mW 785 nm, model Sanyo DL7140-201, Newton, NJ) laser diode operating at a dc current of 85 mA and lasing at 785 nm, a suitable wavelength for IRDye800 light absorption. The beam was expanded with a plano-convex lens to a desired circumference over the whole body for area illumination. The diode current was kept constant by a driver (Thorlabs, Inc. Newton, NJ, model LDCC500, and the temperature was controlled (Thorlabs, Inc. Newton, NJ, model TEC2000) at 15°C. A holographic optical diffuser was placed before the plano-convex lens to provide a uniform excitation light field.

Optical filters were fixed to the 28- to 80-mm zoom lens to control fluorescence light input. A 785-nm holographic notch band rejection filter (Kaiser Optical Systems Inc., Ann Arbor, MI, model HNPF-785.0-2.0) was placed in front of the Nikon lens to selectively block the excitation wavelength. Additionally, an 830-nm bandpass filter (Image quality, Andover Corp., Salem, NH, model 830.0-2.0) was stacked with the band rejection filter to selectively pass the emission signal (±10 nm). White-light imaging was acquired by the CCD system, with no optical filtering and with a low power lamp.

2.5 Image Processing

Images acquired from the ICCD camera system and the nuclear M.CAM gamma camera were processed on a 2.6-GHz Windows PC using both the gratis imaging software, ImageJ, (National Institutes of Health, Washington, DC), and the Precision Digital Imaging System software, V++ (Digital Optics, Auckland, New Zealand). ImageJ is an image analysis and processing program in Java and supports both the optical and nuclear image file formats. The nuclear imaging file format, namely an Interfile (containing the file types .hdr and .img) was opened with support from an ImageJ Java script, or “plugin” (http://www.med.harvard.edu/JPNM/jif/plugins/NucMed.html). V++ is the Roper Scientific CCD camera software interface with programmable modules for ICCD operation. The processed nuclear images were opened and read as unsigned 16-bit intensity matrices. That is, each pixel was represented by a positive floating point integer up to a total of 2\(^{16}\) possible levels of gray. Likewise, the optical file formats (.tif files) were imported into ImageJ or V++ for the intensity calculations described below. Similarly, the optical images were of unsigned data type with integer values ranging from 0 to 65,536.

2.6 Image Analysis by Figures of Merit

In order to assess relative performance of optical imaging and conventional gamma imaging, the mean and standard deviation of intensity counts associated with five primary regions of interest (ROI) were computed (see Fig. 1). Of the five ROI evaluated, one region represented the image “target” and the remaining four regions represented the image “background” for various calculations presented herein.

First, the “hot” area selected for the “target” ROI was the positive tumor, M21. The M21 tumor was fixed as the “target” ROI for all calculations throughout this analysis. The second ROI selected encompassed the negative tumor, M21-L, and this region represented one of the “background” ROI in selected computations. The third ROI surrounded nontumor, or normal tissue (NT). The NT ROI encircled an area on the chest region of the mouse body, and similar to the M21-L ROI, the NT ROI represented a “background” in the computations presented herein. Fourthly, the mean and standard deviation of intensity counts were computed for a ROI that delimited the entire mouse body (WB). The last “background” ROI selected was of the true image background (BK), an area located adjacent to the animal in the surrounding background and not over the body.

The same selected regions of interest corresponding to the M21, M21-L, NT, WB, and BK were used for analysis of optical and nuclear images. That is, the true area of each region remained constant between the nuclear and optical images.

The mean intensity, or mean projected signal, \(\bar{P}\), is defined below where the integers \(m\) and \(n\) represent the pixel values that make up the \([i \times j]\) image intensity matrix for the respective regions of interest selected:

\[
\bar{P} = \frac{1}{mn} \sum_{i=1}^{m} \sum_{j=1}^{n} P_{(i,j)}
\]

(1)

The optical and nuclear images were evaluated by target-to-background (TBR) and signal-to-noise (SNR) ratios. TBR and SNR are image quality figures of merit used to evaluate signal detection and are often reported in medical imaging analyses.\textsuperscript{31–35} The TBR and SNR are computed by:

\[
\text{TBR} = \frac{\bar{P}^t}{\bar{P}^b}
\]

(2)

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where the superscript $t$ represents the mean projected target signal and the superscript $b$ represents the mean projected background signal.

In addition to correlating the variability in emission signals obtained between optical and nuclear modalities, an analysis of image contrast was assessed. The contrast was calculated by the following relationship:

$$ C = \frac{\bar{P}_t - \bar{P}_b}{\bar{P}_b} = \text{TBR} - 1. $$

Using these figures of merit described above, we compared: (i) the TBR for both nuclear and optical techniques; (ii) the SNR as a function of integrated area for optical and nuclear images; and (iii) the contrast achievable from both the optical and nuclear images.

## 3 Results

Figure 1 shows typical nuclear gamma and fluorescence-enhanced optical images of a nude mouse bearing the M21 and M21-L tumors 24 h postinjection of the dual-labeled contrast agent. Both images are presented on equivalent pseudo-color schemes and were acquired with the imaging specifications listed in Table 1. The optical image resolution was 380 $\mu$m/pixel (or 303 $\mu$m/pixel, for three animals imaged with a slightly decreased field of view) and the nuclear image resolution was 609 $\mu$m/pixel. Notably, the total integration time required for the acquisition of the optical image was 800 ms while the projected gamma scintigram took a total of 15 min to acquire.

Despite an equivalent dynamic range, the conventional scintigram and optical image (see Fig. 1) have obvious differences in appearance. The M21 tumor is clearly delineated in the optical image; the M21 ROI intensity is much higher in contrast to the background M21-L tumor. Note however the elevated intensity in the optical image most apparent in the animal’s neck and abdomen region. The signal at those regions can be attributed to elevated fluorescence signals owing to an irregularity in the expanded excitation beam as well as excitation light leakage from the reflected laser light. Alternatively, the nuclear scintigram of the same animal appears noisy and the tumor is hard to delineate. The scintigram displays low contrast between the M21 ROI and the rest of the body. This reduction in contrast, common in nuclear imaging, arises due to the scatter of gamma rays which are captured into the collimator wells and contribute to the high background signal.

In addition, in a few of the images (as can be noted by example of Fig. 1) the uptake of the dual-labeled peptide in the liver and kidneys is evident in the nuclear scan (i.e., the saturated intensity signal in the abdomen region) but not in the optical scan. Owing to the lower energetics and greater attenuation experienced by NIR photons in comparison to gamma emission, cw planar imaging is surface weighted as evidenced by Fig. 1. While tomographic diffusion approaches for 3-D interior imaging have been developed in large tissue volumes,

Table 2 summarizes six image quality factors computed from the nuclear and optical images: (1) the WB ROI area ($\text{cm}^2$), (2) the M21, M21-L, and NT ROI areas ($\text{cm}^2$), (3) the mean projected signal from the M21 tumor ROI, (4) the mean projected signal from the WB ROI, (5) the mean projected signal from the M21-L tumor ROI, and (6) the mean projected signal from the NT ROI. The mean projected signals were computed from Eq. (1), and the total tumor and whole body areas ranged from 0.8 to 1.2 $\text{cm}^2$ and 40 to 45 $\text{cm}^2$, respectively. Owing to the successful targeting of the dual-labeled probe, the mean intensity in the M21 ROI was higher than the M21-L ROI for both the gamma scintigram and optical image. The average intensity (a.u.) in the M21 ROI for the six optical images was 1138.4 ($\pm$529) and 1257.5 ($\pm$507), for the six scintigrams. In comparison, the average intensity in the M21-L ROI for the six optical images was 701.6 ($\pm$330) and 860.9 ($\pm$352), for the six scintigrams.

The TBR figures of merit resulting from each imaged animal are summarized in Table 3. The TBR values, computed by Eq. (2), are presented for the three background regions of interest: (1) M21 target to M21-L background, (2) M21 target to NT background, and (3) M21 target to WB background. The average M21 tumor to M21-L tumor ratios for optical

### Table 1 Parameters used to collect images using the intensified CCD camera system and the M.CAM small animal gamma imager. All images were obtained under the instrumentation parameters outlined, with the exception of one M.CAM image that was arbitrarily cropped to 648×900 pixels (resolution unchanged) and three animals that were imaged with a smaller field of view during the optical imaging study (values in parentheses).

<table>
<thead>
<tr>
<th>Modality</th>
<th>Integration time (s)</th>
<th>Total pixels</th>
<th>Field of view</th>
<th>True resolution size (µm/pixel)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCAM</td>
<td>900</td>
<td>652 896</td>
<td>39.7 54.4</td>
<td>609</td>
</tr>
<tr>
<td>CCD</td>
<td>0.800</td>
<td>300 300</td>
<td>11.4 (8.1)</td>
<td>380 (303)</td>
</tr>
</tbody>
</table>

$$ SNR = \frac{\bar{P}_t - \bar{P}_b}{\sqrt{\sum_{i=1}^{m} \sum_{j=1}^{n} (P_{i,j} - \bar{P}_t)^2}} $$

$$ TBR = \frac{C}{C_b} - 1 $$
(1.4 ± 0.3) and nuclear (1.5 ± 0.2) were similar based on a statistical significance analysis using a paired t-test (α = 0.05). The average M21:NT ratios from the acquired optical and nuclear images were 1.3 (±0.2) and 1.5 (±0.5), and the similar TBR ratios with the whole body as the “background” (M21:WB) for optical and nuclear resulted in 1.6 (±0.3) and 1.6 (±0.7), respectively.

Table 3 also lists the SNR computed by Eq. (3) from the optical and nuclear image data. The SNR using the M21 and M21-L tumor ROIs was greater from measurements acquired by the ICCD system than from measurements acquired by the M.CAM (statistically significant, P value = 1.3e-6). In fact, the maximum SNR value among all six optical images was 7.7 dB. The SNR from the scintigram corresponding to the same optical SNR was 0.3 dB. The average SNR (a.u.) from the optical images was 4.7 (±0.7) and 0.78 (±0.5) from the nuclear images. The SNR and TBR for the six animals are presented by a bar plot in Fig. 2.

The image contrasts computed by Eq. (4) for all six xenografts are listed in Table 3. The contrast was determined by the mean projected signal in the M21 tumor and the mean projected signal in the M21-L tumor. The tumor was detected over the background at an elevated contrast by the M.CAM system with an average contrast of 0.5 (±0.3) as opposed to 0.4 (±0.2), the average contrast obtained from the ICCD system.

Figure 3 is a plot of the SNR versus contrast for selected contrast values arranged in increasing magnitude and computed from all six mice. The contrast and SNR displayed by Fig. 3 were computed using the M21 tumor and BK ROIs. That is, the true background signal was compared to the highest signal present in the images, the tumor intensity. This analysis permits a comparison between nuclear and optical image noise floor. Because SNR and contrast are proportional, their linearity provides an outlook on the tumor “detectability” for various levels of image contrast. The SNR required to achieve a specified contrast was lower for the nuclear system than for the optical system. The ICCD camera system was estimated to have a 200+ fold greater sensitivity at the lower contrast values than the nuclear M.CAM imager.

4 Discussion

To date, optical techniques have yet to demonstrate the robust detection of cancer in humans that nuclear systems can clinically provide. In this work, we identify the features of fluorescence optical imaging that give rise not only to improvements in optical image quality but also to sensitive acquisition of whole body fluorescent signals using a simple ICCD system. Herein, we have shown that surface melanoma tumors can be easily delineated in all six optical images, while the tumor boundary could not be identified (on a qualitative level) by the gamma images using a single, dual-labeled targeting agent. Several factors that contribute to these differences in measured signal are discussed below.

First, although the functional imaging aims of optical and nuclear techniques are similar, it should be prefaced that the purpose of nuclear imaging modalities used in clinical practice is not strictly to define superficial tumor margins. Rather, nuclear systems play dominant roles for bone scans, lymphoscintigraphy, immunoscintigraphy, and the tracing of diseases (e.g., cancers, Alzheimer’s, Parkinson’s, AIDS dementia, etc.). PET and SPECT are employed over conventional gamma imaging and provide improved image quality and sensitivity. In this study we compare conventional gamma imaging to NIR optical imaging for superficially localized tumors in small animals; both are nontomographic methods with instrumentation designed for small animal imaging.

Second, in order to provide an impartial comparison of nuclear and optical signals, a well-documented and equally detectable contrast agent dose must be established. The dual-labeled peptide, 111In-DTPA-Lys(IRDye800)-c(KRGDF), used in this study was synthesized for a 1:1 molar ratio of DTPA to conjugated IRDye800. After addition of the radiolabel, 111In, a final product equivalent to 20 μCi of radioisotope per nmol of fluorophore was obtained. Indeed the specific activity can be altered by varying the amount of 111In to be chelated to DTPA-Lys(IRDye800)-c(KRGDF). The total specific activity however, should be balanced for adequate detection of both radio and fluorescent signals. When applied to the clinical setting, radionuclide dosimetry is calculated based on realistic integration times (for patient comfort), radiation exposure (for patient safety), half-life and energy of the isotope, and the minimum detectable signal. Typical clinical intravenous injection doses of Octreoscan (111In-D-Phe-DTPA-octreotide) change depending on imaging circumstances (3 mCi to 6.0 mCi), and as clinical doses vary so do experimental doses in small animals. Despite a > 105 lower bodyweight, mice may be administered 100 times lower doses than humans in order to obtain an adequate signal at realistic integration times. It is noteworthy that the tracer dose does not linearly scale with body mass. Our small animal study required 3.3MBq 111In in order to detect ligand receptor interaction. Similar doses ranging from 1.8MBq to 7.2MBq have been used in small animals with 111In, 18F, and 99mTc, for studies involving targeting of a radiolabel to the αvβ3 integrin via an RGD peptide conjugate.

Despite controlled fluorophore and isotope specific activities, the high fluorescent photon sensitivity of the ICCD system may be due to the theoretically larger number of fluorescent photons available for imaging over the available gamma rays emitted from the animal. Indeed, the number of radiative events per second emitted by a fluorescent dye can be several orders of magnitude greater than the total emitted by a radionuclide of the same quantity. For example, 111In has a specific activity of 4.2 * 10^6 [Bq/g]. To quantify this in terms of emitted disintegrations, the rationale is:

\[
\frac{4.2 \times 10^6 \text{Ci}}{g} \times \frac{3.7 \times 10^{10} \text{Bq}}{\text{Ci}} \times \frac{1 \text{ event/sec}}{\text{Bq}} = 1.5 \times 10^{19} \text{ event/sec} \cdot \text{g}^{-1}.
\]

A direct comparison of this activity to the fluorescent dye IRDye800, which has a molecular weight of 1166 g/mol and a fluorescent lifetime estimated at 0.5 ns, produces:
1 mole $\times 6.02 \times 10^{23}$ molecule/mole $\times 0.5 \times 10^9$ event/molecule/sec $\times 1166$g/mol $\div 0.16$ = 2.6 $\times 10^{29}$ event/sec.g$^{-1}$.

Thus, an equivalent “specific activity” for the fluorophore and radionuclide in theory provide significantly different amounts of photons emitted per second (10$^{10}$). Owing to this low sensitivity, nuclear scans require several minutes for data collection. Optical techniques may be favorable for the clinic because they require only a fraction of a second to resolve a fluorescent image.

The discrepancy between the theoretical difference in sensitivity and the actual observed image sensitivity can be explained by the comparative attenuation of NIR and gamma rays during tissue propagation. In tissue, gamma rays attenuate to a much lesser extent than NIR photons. For example, with a linear attenuation coefficient of $\mu_s=0.16$ cm$^{-1}$ for a gamma ray, and a reduced scattering and absorption coefficient of $\mu_s'=10$ cm$^{-1}$ and $\mu_e=0.1$ cm$^{-1}$ for NIR photons in thick tissue, a rough estimate of the attenuation of gamma and NIR intensity across 5 cm of tissue can be made:

$$I_{\text{gamma}} = I_0 e^{-\mu_s L} = 1.0 e^{-0.16 \times 5} = 0.45$$

$$I_{\text{optical}} = I_0 e^{-\mu_s' L} = 1.0 e^{-0.1 \times 10} = 1 \times 10^{-22}$$

showing that almost 50% of gamma rays are transmitted and much less than 0.01% of optical NIR photons are transmitted through an equivalent tissue depth. Attenuation unquestionably depends on the radiation energy; for example $\mu_s\approx0.16 - 0.35$ cm$^{-1}$ for photon energies of 30 to 100 keV and $\mu_e=100 - 300$ cm$^{-1}$ for NIR photons, where scattering dominates. Additionally, photons and gamma rays experience different types of scattering (i.e., Rayleigh versus Compton) in soft tissue.
A large part of NIR attenuation for fluorescence-enhanced imaging is due to scattering and absorption as well as the decay kinetics of the fluorophore. Excitation light attenuates before reaching the embedded fluorophore and the quantum efficiency, \( \phi \), of the fluorophore reduces the amount of fluorescent light that is re-emitted. Upon traveling back to the tissue surface, fluorescent photon attenuation occurs. Ultimately, the photon flux measured at the tissue surface is reduced and for the small animal imaging described herein, its attenuation must be predicted by the full radiative transport equation.

A last and important preface to nuclear and optical imager differences that may impact equal comparison is camera resolution. The CCD chip has fixed, 24-\( \mu \)m-size wells, thus the image resolution, or actual pixel size, may vary depending on the field of view chosen and achieved using any typical camera focusing lens. For the ICCD system, the resolution on an optical image is ultimately limited by the image intensifier; the CCD images a constant FOV at the phosphor screen, which likewise can be controlled by a simple zoom lens. Unlike the optical system however, the M.CAM has a fixed field of view, therefore the resolution is fixed for a given collimator size.

The comparative target-to-background ratios that are presented reflect some distinguishing qualities between the planar optical and nuclear image display. The M21:M21-L TBR values were analyzed and found to be similar; this similarity was anticipated because the contrast agent was dual-labeled causing the radionuclide and fluorophore to have an equal relative accumulation between the M21 and M21-L tumor. Yet upon viewing the ROI intensities in the actual optical and nuclear images (Fig. 1) this TBR similarity is less apparent. There is little signal delineating the gamma image M21 ROI over the M21-L ROI, which indicates a high variance in pixel intensity resulting from a raised background signal. The large background signal in the nuclear scintigram reflects the fact that gamma cameras detect scattered gamma rays that are emitted from deep tissue sections. Thus, any annihilated photons that emit from the liver, kidneys, and deeper tissue (>5 mm) are imaged and contribute to the total mean projected signal. Moreover, gamma rays that scatter into a single collimating well and emit from locations not adjacent to that well are weighted the same as those photons that travel in a straight trajectory from the tumor to the collimator. Thus, the deviation in pixel intensity values is much greater for the M.CAM than for the ICCD system.

Additionally, the similarities between the optical and nuclear M21:NT ratios and optical and nuclear M21:WB ratios further demonstrate that the dual-labeled agent was target-specific to the \( \alpha\beta \) integrin positive melanoma tumor. The nuclear and optical detectors equally discriminated between targeted signals and nontargeted signals from the background of the entire animal body and normal tissue regions. Yet, the optical and nuclear imaging systems appear to detect different background signals because (i) optical ICCD sensitivity is weighted toward shallow, subsurface photons; (ii) attenuated optical signals are often reported at suppressed intensities relative to nonattenuated signals owing to excessive excitation light leakage; (iii) conventional gamma camera systems detect gamma rays from any depth or location in the

### Table 3

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...with equal sensitivity; and (iv) gamma systems detect photons that travel in a mostly linear fashion through the body with little attenuation. Therefore the background signals from the optical M21-WB and M21-NT ratios are from fluorescence emitted by tissues and blood located a few millimeters below the skin’s surface as well as excitation light leakage through the optical filters. Alternatively, the gamma camera TBRs are indicative of the total radio-nuclide emissions from within the whole body relative to the gamma ray emission from the entire tumor volume.

**Fig. 2** The bar plots present (a) target-to-background ratios and (b) signal-to-noise ratios of the M21 tumor region of interest to the M21-L tumor background region of interest computed from NIR (dark-shaded) and gamma (light-shaded) images for all six animals studied. Error bars represent the relative standard deviation. The TBRs predicted from the ICCD and the MCAM yield statistically similar results (paired t-test with \( \alpha = 0.05 \)). On the contrary, the SNR values predicted from the ICCD and the MCAM yield statistically different results (paired t-test with \( P \text{-value} = 1.3 \times 10^{-6} \)), thus indicating that the ICCD imager produces on the average higher SNR values than does the M.CAM imager.

Unlike the TBR ratios, the statistically different SNR may be an indication of the reduced noise floor that an ICCD system provides. ICCDs are typically operated under shot-noise limited conditions, where photon noise is the limiting noise factor. Therefore, when high photon signals emitting from the tumor are detected, the Poisson distributed shot noise becomes relatively much smaller and this is reflected in the image output SNR. The SNR computed from the conventional gamma images is much lower than the optical owing also to the imaging system’s intrinsic signal-to-noise characteristics. The radioactive decay and gamma detection variance is closer to the gain, or sensitivity of the detector; this is reflected in the scintigram pixel values. However the favorable SNR of optical over nuclear may dramatically decrease when comparing deeply seated tumors as in the case of human imaging. Consider two targets 0.5 and 2 cm deep to be imaged using planar optical and nuclear imaging. The noise associated with each of the imaging modalities remains constant as the depth of the target increases. Assuming that the gamma emitting targets are of equal energy, the signal from the two centimeter depth \( (e^{-0.16 \times 2} = 0.7) \) will be attenuated by a difference of 0.2 in comparison to that emitted from the target 0.5 cm in depth \( (e^{-0.16 \times 0.5} = 0.9) \). For optical imaging, greater attenuation occurs with typical tissue scattering and absorption coefficients such as \( \mu_s' = 10 \text{ cm}^{-1} \) and \( \mu_a = 0.1 \text{ cm}^{-1} \). Using these optical properties for a total attenuation coefficient of \( \mu_t = 10.1 \), the comparative attenuation of photons from 0.5 cm to 2 cm is 0.006 and \( 2 \times 10^{-9} \), respectively, a magnitude difference of 10⁶. Of course, these calculations do not consider the half-life of the radiotracer, wherein the SNR of fluorescent targets at all depths may be significantly greater than that of a spent tracer at any depth.

Yet for detection of surface signals, plotting the SNR versus contrast further demonstrates improved sensitivity by the ICCD system. A high photon signal detected by the ICCD system yields a large SNR value despite the low contrast between the tumor and background. For the SNR contrast plots, the background value used was a true background ROI; a selected area outside of the animal body was used as the baseline noise signal. The change in optical SNR relative to the change in optical image contrast is more than 200-fold greater.
than the conventional gamma camera SNR-contrast gradient. Moreover, fluorescence-enhanced optical imaging has potential for improved SNR with better reduction in the excitation light noise. With the increase in intensified CCD sensitivity, excitation light is hard to eliminate from the total detected signal because of the fraction of excitation light (19%) passed through standard band rejection and optical bandpass filters.

Although this work demarcates favorable optical image quality relative to a conventional gamma camera, multimodality imaging combinations are truly complementary with neither imaging method providing completely autonomous diagnostic information. In this work however, the combined imaging methods provided a comparative tool for optical ICCD camera image quality assessment, which is essential for clinical translation of functional planar NIR imaging. Upon comparison to conventional nuclear imaging, it is evident that optical imaging with ICCD cameras is highly sensitive to photons emanating from superficial depths. Additionally, as optical tomography is developed for multipixel techniques on larger volumes, assessment of the target detection for small target sizes and greater target depths is increasingly important for locating the tumor position through a three-dimensional map in thick (>1 cm) human tissues.

Acknowledgments

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References


