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**ABSTRACT**

The purpose of this grant is to build cancer-specific contrast agents for photoacoustic imaging, using which one could estimate the change in molecular expression of various breast-cancer-specific proteins undergoing chemotherapy treatment. We've made significant progress towards obtaining this goal: 1) we created the first-ever photoacoustic imaging agent (which is based on carbon nanotube nanoparticle) and showed it can specifically target tumors in tumor-bearing mice (paper published in Nature Nanotechnology); 2) We created 2 additional molecular imaging agents for photoacoustic imaging which exhibit 300-times higher sensitivity and for the first allow imaging photoacoustic molecular probes at sub-nanomolar concentrations (paper submitted to Nano Letters). We've shown that such sensitivity improvement results in the ability to image smaller tumors. Beyond higher sensitivity, the 3 imaging agents developed in this grant thus far have different optical spectra. We used this fact and have shown the ability to simultaneously image these agents (multiplexing). This ability is particularly powerful and important for this grant as we plan to progress to characterizing the response to chemotherapy of multiple cancer-specific proteins in the same tumor simultaneously.
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1. INTRODUCTION

The purpose of this grant is to build cancer-specific contrast agents for photoacoustic imaging, using which one could estimate the change in molecular expression of various breast-cancer-specific proteins undergoing chemotherapy treatment. We’ve made significant progress towards obtaining this goal: 1) we created the first-ever photoacoustic imaging agent (which is based on carbon nanotube nanoparticle) and showed it can specifically target tumors in tumor-bearing mice (paper published in Nature Nanotechnology); 2) We created 2 additional molecular imaging agents for photoacoustic imaging which exhibit 300-times higher sensitivity and for the first allow imaging photoacoustic molecular probes at sub-nanomolar concentrations (paper submitted to Nano Letters). We’ve shown that such sensitivity improvement results in the ability to image smaller tumors. Beyond higher sensitivity, the 3 imaging agents developed in this grant thus far have different optical spectra. We used this fact and have shown the ability to simultaneously image these agents (multiplexing). This ability is particularly powerful and important for this grant as we plan to progress to characterizing the response to chemotherapy of multiple cancer-specific proteins in the same tumor simultaneously.

2. BODY

2.1 Creation of 2 new Photoacoustic imaging agents

We have recently reported on the conjugation of cyclic Arg-Gly-Asp (RGD) peptides to pegylated SWNTs¹ and their use as photoacoustic imaging agents². In order to enhance the photoacoustic signal of the SWNTs, we attached Indocyanine Green (ICG) and QSY-21 dyes to the surface of the SWNTs through pi-pi stacking interactions³ (see Methods section for more details). The ultra-high surface area of SWNTs allows highly efficient loading of aromatic molecules such as ICG and QSY-21 on the nanotube surface. This created two new kinds of photoacoustic agents; SWNT-ICG and SWNT-QSY (Fig. 1a). The particles were targeted using the RGD-peptide to α₅β₃ integrins, which are over-expressed in tumor vasculature, while control untargeted particles were synthesized using a non-targeted peptide, RAD.

The optical absorbance spectra of the two new particles suggest that 710 nm and 780 nm are the preferable wavelengths for scanning SWNT-QSY and SWNT-ICG respectively (Fig. 1b). At their respective absorbance peaks, the SWNT-QSY and SWNT-ICG particles exhibit a 17 and 20-fold higher absorbance respectively as compared with plain SWNTs. Since blood absorption is significantly reduced at 780 nm compared to 710 nm, SWNT-ICG was the particle of choice for the small animal experiments for this study. Importantly, the attachment of RGD or RAD peptides to SWNT-ICG had little effect on the particles’ absorbance. We constructed a non-absorbing and non-scattering agarose phantom with inclusions of SWNT-ICG-RGD at increasing concentrations from 0.5 nM to 121.5 nM in multiples of 3 (n = 3 samples of each concentration). The photoacoustic signal produced by the SWNT-ICG-RGD particles correlated well with the nanoparticle concentration (R²=0.983) (Fig. 1c).
Figure 1. Characterization of the dye-enhanced SWNT. a, Illustration of SWNT-ICG and SWNT-QSY. ICG and QSY-21 (red molecules) are attached to the SWNT surface through non-covalent pi-pi stacking bonds. Polyethylene glycol-5000 (blue molecules) is conjugated to a targeting peptide in one end and to the SWNT surface on the other end through phospholipids. b, Optical spectra of plain SWNT (green), SWNT-ICG-RGD (red), SWNT-ICG-RAD (blue) and SWNT-QSY-RGD (black). The similarity of SWNT-ICG-RAD and SWNT-ICG-RGD spectra suggests that the peptide conjugation does not notably perturb the photoacoustic signal. c, The photoacoustic signal produced by SWNT-ICG was observed to be linearly dependent on the concentration ($R^2 = 0.9833$).

2.2 Sensitivity of the imaging agents in living mice

We then tested the particle’s sensitivity in living subjects by subcutaneously injecting the lower back of mice ($n = 3$) with 30 µl of SWNT-ICG-RAD mixed with matrigel at increasing concentrations of 820 pM to 200 nM in multiples of 3. Matrigel alone produced no significant photoacoustic signal (data not shown). Upon injection, the matrigel solidified, fixing the SWNT-ICG-RAD in place and three-dimensional (3D) ultrasound and photoacoustic images of the inclusions were acquired (Fig. 2a). While the ultrasound images visualized the
mouse anatomy (e.g., skin and inclusion edges), the photoacoustic images revealed the SWNT-ICG-RAD contrast in the mouse. The photoacoustic signal from each inclusion was quantified using a three dimensional region of interest (ROI) drawn over the inclusion. We observed a linear correlation \((R^2 = 0.97)\) between the SWNT-ICG-RAD concentration and the corresponding photoacoustic signal (Fig. 2b). Tissue background signal was calculated as the average photoacoustic signal in areas where no contrast agent was injected. Extrapolation of the signal-concentration graph reveals that 170 pM of SWNT-ICG-RAD gives the equivalent photoacoustic signal as the tissue background (i.e., signal to background ratio = 1). This value represents over 300-times improvement in sensitivity compared to plain SWNTs.

**Figure 2. Photoacoustic detection of SWNT-ICG in living mice.** a, Mice were injected subcutaneously with SWNT-ICG at concentrations of 0.82-200 nM. The images represent ultrasound (gray) and photoacoustic (green) vertical slices through the subcutaneous injections (dotted black line). The skin is visualized in the ultrasound images, while the photoacoustic images show the SWNT-ICG distribution. The white dotted lines on the images illustrate the approximate edges of each inclusion. b, The photoacoustic signal from each inclusion was calculated using 3D regions of interest and the ‘background’ represents the endogenous signal measured from tissues. The error bars represent standard error \((n = 3\) mice). Linear regression \((R^2 = 0.97)\) of the photoacoustic signal curve estimates that a concentration of 170 pM of SWNT-ICG will give the equivalent background signal of tissues.
2.3 Targeting of the imaging agents to tumors

Finally, we tested the nanoparticles targeting ability in living mice. Mice bearing tumor xenografts (150 mm³ in size) were injected through the tail vein (IV) with 200 μl of either targeted SWNT-ICG-RGD or untargeted SWNT-ICG-RAD particles (n = 4 mice per group) at a concentration of 1.2 μM. We acquired 3D photoacoustic and ultrasound images of the entire tumor area before and up to 4 hours after the injection. Mice injected with the targeted SWNT-ICG-RGD particles show significantly higher photoacoustic signal in the tumor compared with the control group (Fig. 3a). The ultrasound images were used for visualizing the boundaries of the tumor as well as to validate that no significant movement (beyond 100 μm) had occurred throughout the scan. While the pre-injection photoacoustic signal is primarily due to the tumor’s blood content, post-injection photoacoustic signal consists of both blood and SWNT-ICG. To subtract out the blood signal from the images, a subtraction image calculated as the 2 hour post-injection minus the pre-injection image was calculated. Measurement of the photoacoustic signal from a 3D ROI around the tumor (Fig. 3b) showed that the photoacoustic signal in the tumor was significantly higher in mice injected with SWNT-ICG-RGD as compared with the control particles SWNT-ICG-RAD (p < 0.001). For example, at 2 hours post-injection, mice injected with SWNT-ICG-RGD showed over 100% higher photoacoustic signal in the tumor than mice injected with the control SWNT-ICG-RAD.
Figure 3. SWNT-ICG-RGD tumor targeting in living mice. a, Ultrasound (gray) and photoacoustic (green) images of one vertical slice through the tumor (dotted black line). The ultrasound images show the skin and the tumor boundaries. Subtraction photoacoustic images were calculated as 2 hr post-injection minus pre-injection images. As can be seen in the subtraction images, SWNT-ICG-RGD accumulates in higher amount in the tumor as compared to the control SWNT-ICG-RAD. b, Mice injected with SWNT-ICG-RGD showed significantly higher photoacoustic signal than mice injected with the untargeted control SWNT-ICG-RAD (p < 0.001). The error bars represent standard error (n = 4 mice).

2.4 Imaging the two contrast agents simultaneously (Multiplexing)

Finally, we show that the two kinds of photoacoustic imaging agents we synthesized, SWNT-ICG and SWNT-QSY can be imaged simultaneously due to their unique, though overlapping, absorbance spectra (Fig. 1b). We created an agarose gel phantom containing increasing concentrations of SWNT-ICG and decreasing concentrations of SWNT-QSY (starting from 100nM:0nM up to 0nM:100nM respectively). Photoacoustic images of the phantom were taken at wavelengths of 700, 730, 760, 780, and 800 nm and a spectral un-mixing algorithm was then used to separate each particle’s signal to an individual image (Fig. 4).

Figure 5. Multiplexing of SWNT-ICG with SWNT-QSY particles in a phantom. A phantom with various concentrations of SWNT-ICG and SWNT-QSY was scanned under the photoacoustic instrument at wavelengths of 700, 730, 760, 780, and 800 nm. A spectral un-mixing algorithm based on least-squares was used to separate the signals of SWNT-ICG particles (green) from SWNT-QSY particles (red). Notice that no SWNT-QSY signal is seen in the well with pure SWNT-ICG and vice versa, despite the fact that the two particles have overlapping spectra.
KEY RESEARCH ACCOMPLISHMENTS

- Developed two more photoacoustic imaging agents
- Characterized the particles and optimized them for tumor targeting upon intra-venous administration to tumor-bearing mice
- Optimized the imaging system to allow imaging the two imaging agents simultaneously

REPORTABLE OUTCOMES


  *Abstract poster presentation was awarded the best poster presentation at the Photoacoustic session at the conference – the biggest photoacoustic conference.*

CONCLUSION

The main achievement over this past year was the development of 2 new photoacoustic imaging agents which allow reaching unprecedented sensitivities. This development will be the basis for measuring breast cancer response to chemotherapy and will allow investigating biomarkers which weren’t been able to investigate before due to insufficient sensitivity of photoacoustic instruments.

REFERENCES

APPENDICES

1. Paper published in Nature Nanotechnology:

Carbon nanotubes as photoacoustic molecular imaging agents in living mice

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Photoacoustic imaging of living subjects offers higher spatial resolution and allows deeper tissues to be imaged compared with most optical imaging techniques. As many diseases do not exhibit a natural photoacoustic contrast, especially in their early stages, it is necessary to administer a photoacoustic contrast agent. A number of contrast agents for photoacoustic imaging have been suggested previously1-3, but most were not shown to target a diseased site in living subjects. Here we show that single-walled carbon nanotubes conjugated with cyclic Arg-Gly-Asp (RGD) peptides can be used as a contrast agent for photoacoustic imaging of tumors. Intravenous administration of these targeted nanotubes to mice bearing tumors showed eight times greater photoacoustic signal in the tumor than mice injected with non-targeted nanotubes. These results were verified ex vivo using Raman microscopy. Photoacoustic imaging of targeted single-walled carbon nanotubes may contribute to non-invasive cancer imaging and monitoring of nanotherapeutic effects.

Recently, we reported on the conjugation of cyclic RGD-containing peptides to single-walled carbon nanotubes (SWNT-RGD) that is stable in serum. The single-walled carbon nanotubes, which were 1-2 nm in diameter and 50-300 nm in length were coupled to the RGD peptide through polyethylene glycol-5000-grafted phosphonate (PL-PGs5000). These SWNT-RGD conjugates bind with high affinity to α5β1 integrin, which is over-expressed in tumor microvasculature, and to other integrins expressed by tumors but with lower affinity4,5. We also synthesized non-targeted single-walled carbon nanotubes (that is, plain single-walled carbon nanotubes) by conjugating them solely to PL-PGs5000 (Fig. 4a). Our photoacoustic instrument6 used a single-element focused transducer to raster scan the object under study, which was illuminated through a fiber head (see Methods and Supplementary Information, Fig. S1). In a phantom study we measured the photoacoustic signal of plain single-walled carbon nanotubes and SWNT-RGD at wavelengths of 690-800 nm (Fig. 1b; shorter wavelengths are less desirable as the depth of penetration through the tissues is reduced6). These photoacoustic spectra suggest that 690 nm is the preferred wavelength, because the photoacoustic signal of the single-walled carbon nanotubes is higher at that wavelength. Furthermore, the ratio of single-walled carbon nanotubes to hemoglobin signal is higher at this wavelength when compared with other wavelengths. Importantly, the photoacoustic signal of single-walled carbon nanotubes was found to be unaffected by the RGD peptide conjugation. This finding was validated through measurements of the optical absorbance of the two single-walled carbon nanotubes conjugates (see Supplementary Information, Fig. S2). In a separate non-absorbing and non-scattering phantom study, we also validated that the photoacoustic signal produced by single-walled carbon nanotubes is in linear relationship with their concentration (Fig. 1c) with ρ = 0.9997.

We then subcutaneously injected the lower back of a mouse with 0.3 μl mixture of single-walled carbon nanotubes and murepate at concentrations between 50 and 600 nM (n = 3 for each concentration). Murepate alone produced no photoacoustic signal (data not shown). Upon injection, the murepate solidified, fixing the single-walled carbon nanotubes in place. Three-dimensional (3D) ultrasound and photoacoustic images of the inclusions were then acquired (Fig. 2a). The photoacoustic images showed the mouse anatomy (for example, skin and inclusion edges), and the photoacoustic image revealed the single-walled carbon nanotubes contrast in the mouse. The photoacoustic signal from each inclusion was quantified using a 3D region of interest drawn over the inclusion. We observed a linear correlation (R² = 0.9929) between the single-walled carbon nanotubes concentration and the corresponding photoacoustic signal (Fig. 2b). Importantly, this linear relation can only be expected in spatial cases where the die concentration does not perturb the tissue light distribution significantly. We concluded that the photoacoustic signal produced by tissues/background was equivalent to the photoacoustic signal produced by 50 nM of single-walled carbon nanotubes (that is, a signal-to-background ratio of 1). The experimental result correlates well with the theoretical analysis (see Supplementary Information), which predicts a background signal equal to 7-70 nM of single-walled carbon nanotubes, depending on the location of the nanotubes in the body.
Three-dimensional ultrasound and photoacoustic images of the tumour and its surroundings were acquired before and up to 4 h after injection. We found that mice injected with SWNT-RGD showed a significant increase of photoacoustic signal in the tumour compared with control mice injected with plain single-walled carbon nanotubes (Fig. 3a). The images from the different time points were aligned with one another using simple vertical translations to account for small vertical movements in the transducer positioning. This alignment allowed quantification of the photoacoustic signal at all time points using a single region of interest. We then calculated a subtraction image between the photoacoustic image taken at 4 h post-injection and the photoacoustic image taken before injection. The subtraction image better visualizes the real distribution of the single-walled carbon nanotubes as it removes, to a large extent, the background signal. For example, in the mice injected with plain single-walled carbon nanotubes (Fig. 3a), a high photoacoustic signal, likely produced by a large blood vessel, was seen in the pre-injection and post-injection images. However, the subtraction image showed a much lower signal from this area, reflecting the likely low concentration of plain single-walled carbon nanotubes there. We calculated the photoacoustic signal by drawing a 3D region of interest around the tumour (tumour boundaries were clearly visualized in the ultrasound images). The photoacoustic signal increase was quantified as a function of time.
Figure 3: Single-walled carbon nanotube targets tumour in living mice. a. Ultrasound (green) and photoacoustic (red) images of the vertical slice of the tumour in situ. The ultrasound images show the internal structure and boundary. The photoacoustic images were calculated as the 4th harmonic of the ultrasound image minus the pre-injection image. The high photoacoustic signal in the image indicates that the single-walled carbon nanotubes (yellow in the ultrasound image) are present in the tumour, suggesting that it is due to a large blood vessel and not single-walled carbon nanotubes. b. Mice injected with SWNT-RGD showed a significantly higher photoacoustic signal than mice injected with plain single-walled carbon nanotubes (P < 0.05). The error bars represent standard error (n = 4, *P < 0.05).

(Fig. 3b). Although SWNT-RGD led to a consistent higher photoacoustic signal, plain single-walled carbon nanotubes led only to a temporary increase in the photoacoustic signal of the tumour (P < 0.05) when comparing the signals at each time point independently. The temporary photoacoustic signal observed for plain single-walled carbon nanotubes is likely caused by circulating nanoparticles that are eventually cleared from the bloodstream. Conversely, SWNT-RGD bind to the tumour vasculature, creating a consistent photoacoustic signal from the tumour. On average, at 4 h post-injection, the SWNT-RGD resulted in ~3 times greater increase in photoacoustic signal compared with plain single-walled carbon nanotubes. The percentage injected dose per gram of tissue was calculated to be ~14%ID/g (see Supplementary Information).

Figure 4: Validation of the in vivo photoacoustic images by Raman ex vivo microscopy. a. Photograph of the tumour in vivo and the corresponding photoacoustic subtractions images (green). b. The Raman images showed both the photoacoustic and Raman signals on the same side of the tumour. c. Comparison between the photoacoustic signal of the tumour injected with SWNT-RGD and the Raman signal acquired from the excised tumour (right). *P < 0.05.

We further validated our photoacoustic results using a Raman microscope as an independent method for detection of single-walled carbon nanotubes. At the conclusion of the photoacoustic results, 4 h post-injection, the mice were sacrificed, the tumours were surgically removed and scanned as in vivo under a Raman microscope. The two-dimensional Raman images of the excised tumours were found to match the photoacoustic images (Fig. 4a). The mean Raman signal from the tumours was calculated from the Raman images. Similar to the photoacoustic results, the Raman signal from the tumours was ~4 times higher in mice injected with SWNT-RGD than in mice injected with plain single-walled carbon nanotubes (Fig. 4b).
Unlike photoacoustic imaging, optical imaging suffers from relatively poor spatial resolution as well as exponentially degraded sensitivity as tissue depth increases. We showed the superiority of our photoacoustic strategy by comparing it with fluorescence imaging of tumour-targeted quantum dots. The quantum dots were conjugated to RGD peptides (QD–RGD) and visualised with a fluorescence imaging instrument (Fig. 5a). Although the quantum dot and single-walled carbon nanotube conjugates might have different biodistributions, the photoacoustic images of single-walled carbon nanotubes from the tumour illustrate the depth information and the greater spatial resolution achieved by photoacoustic imaging compared with fluorescence imaging (Fig. 5a–d). The measured signal from the tumour in the photoacoustic image is due to light scattering. However, the photoacoustic images showed the 3D distribution of SWNT–RGD in the tumour with high spatial resolution. Similar results were also observed in a phantom study (see Supplementary Information, Fig. 5a).

We have demonstrated that single-walled carbon nanotubes can be exploited as photoacoustic contrast agents to non-invasively image tumours. Intravenous injection of targeted single-walled carbon nanotubes in mice led to a 10-fold higher photoacoustic signal in the tumour compared with mice injected with nontargeted single-walled carbon nanotubes. Our photoacoustic images were verified using Raman microscopy on the surgically excised tumours. Furthermore, our results agreed with a previous study9 where radiolabelled SWNT–RGD were monitored using small animal positron emission tomography (microPET). In the study SWNT–RGD were found to accumulate ~3–5 times more in tumours than pristine single-walled carbon nanotubes. That study also showed that the SWNT–RGD did not accumulate in the tissue surrounding the tumour.

Most previous work on photoacoustic contrast agents in vivo is limited to some targeted agents such as gold nanoparticles used for highlighting the blood vessels in a rat's brain10. A recent preliminary study11 showed that an indocyanine green derivative (IR780–RGD) was highly effective for photoacoustic spectroscopic imaging of U87MG tumours; however, the study was carried out on a single mouse and the validation of the agent has yet to be shown. Various gold nanoparticles have been previously suggested, primarily for their high absorption characteristics and the ability to control their spectra, which allows multiplexing studies. However, their main limitation is their relatively large size, which will lead to their rapid clearance by the reticuloendothelial system (RES) upon intravenous injection. It is possible that single-walled carbon nanotubes, due to their unique high aspect ratio (~100) and high surface area to volume ratio, are capable of minimizing RES uptake while having an increased affinity for molecular targets due to multivalency effects12. A concentration of 50 nM of single-walled carbon nanotubes was found to produce a photoacoustic signal equivalent to mouse tissue (background); however, the minimum detectable concentration of single-walled carbon nanotubes is likely to be less than 50 nM. This is because photoacoustic images were acquired before and after the administration of the contrast agent, thus making it possible to separate the contrast agent signal from the background signal. Further background reduction can be achieved by performing photoacoustic spectral imaging, improving hardware/reconstruction software, or by enhancing the single-walled carbon nanotubes' photoacoustic signal. With respect to acquisition time, our current instrument acquires a single photoacoustic image in ~20–30 minutes for a tumour ~100 mm³ in size. However, by using lasers with higher repetition rates, scan duration can be greatly reduced.

We are currently investigating the potential of single-walled carbon nanotubes to extravasate out of the leaky vasculature of tumours. Single-walled carbon nanotube extravasation is of particular interest, because upon exiting the vasculature the nanotubes would have access to many more molecular targets that exist only on the cancer cell's membrane. Future work should optimise the partial extravasation as well as bring new technologies to help quantify the degree of nanotube extravasation. Moreover, future studies can monitor various nanotherapeutic applications such as drug-eluting single-walled
carbon nanotube using photoacoustic imaging. Such nanotherapeutic and cancer imaging applications would gain further clinical interest as single-walled carbon nanotubes continue to show low toxicity effects. Although single-walled carbon nanotubes have the ability to efficiently bind to molecular targets, their high photoacoustic signal allows for high-resolution 3D plaque orientation, with substantial depth of penetration. None of the other molecular imaging modalities compare with the precise depth information and submillimeter resolution at nanomolar sensitivity that is achieved by photoacoustic imaging. We expect this work to stimulate further studies of biologically relevant problems using photoacoustic molecular imaging.

METHODS

SYNTHESIS OF SINGLE-WALLED CARBON NANO TUBE COLLOIDS

A complete description of the synthesis of SWNT-RGO is found elsewhere. The single-walled carbon nanotubes used in this study were dispersed in 0.1 N HCl for 2 h. The dispersion was then filtered and washed with water to remove excess acid, and then dried under vacuum.

STATISTICAL METHODS

For the single-walled carbon nanotube targeting experiments, we used a random-effects model to test the hypothesis that the mice treated with SWNT-RGO showed an increased photoacoustic signal over time in the tumor. The control group was analyzed separately with a single-walled carbon nanotube and a protein cocktail. The tumor size was measured at each time point independent of treatment and the photoacoustic signal was collected at each time point independent of treatment.

PHOTOACOUSTIC IMAGING

Our in-house photoacoustic system is illustrated in the Supplementary Information. Fig. 1. Amonolax pulsed laser with a repetition rate of 1 kHz and a wavelength of 1064 nm was used. The 1064 nm laser pulse was delivered through a 15-cm-long fiber-optic probe. The tumor was placed in a water bath to maintain an isothermal environment. The output power level of the laser was 200 mW, and the laser pulse duration was 10 ns. The laser was focused on the tumor surface using a 1.5-m focal length lens. The tumor was illuminated with 600 mW of light for 1 minute at each time point. The photoacoustic signal was detected by an acoustic transducer and recorded using an oscilloscope. The data were processed offline using a custom-built software program. The image was then analyzed to determine the size and shape of the tumor.

RESULTS

Animal studies showed that the SWNT-RGO treated group had a statistically significant increase in the tumor area compared to the control group. The tumor area increased by 20% in the SWNT-RGO treated group and by 5% in the control group. The increase in tumor area was observed in all animals treated with SWNT-RGO, regardless of the tumor size.

DISCUSSION

These results demonstrate the potential of using photoacoustic imaging to monitor the growth and regression of tumors. Photoacoustic imaging can provide real-time information about the size, shape, and location of tumors, and can be used to guide surgical interventions. The results also suggest that SWNT-RGO may be a promising therapeutic agent for the treatment of cancer.

REFERENCES


For more information, please visit our website at: www.nature.com/nature.

Supplementary Information: Please visit our website for additional information and data.

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Ultra-High Sensitivity Carbon Nanotube Agents for Photoacoustic Molecular Imaging in Living Mice

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Introduction

Photoacoustic imaging is an emerging modality that overcomes to a great extent the resolution and depth limitations of optical imaging while maintaining high-contrast\textsuperscript{1-6}. However, since many diseases will not manifest an endogenous photoacoustic contrast, it is essential to develop exogenous photoacoustic contrast agents that can target diseased area(s). Recently, we showed that single-walled carbon nanotubes (SWNTs) have utility as targeted photoacoustic contrast agents\textsuperscript{7}. Here we present a dye-enhanced SWNT agent that markedly increases the photoacoustic contrast in living tissues by 300-times compared to plain SWNTs, leading to sub-nanomolar sensitivities. By attaching two different dyes onto the SWNT surface, we show that the two resulting particles can be imaged simultaneously (multiplexing). Intravenous administration of targeted dye-enhanced SWNTs to tumor-bearing mice showed significantly higher signal in the tumor than mice injected with untargeted particles. Finally, we show that the new dye-enhanced SWNTs can detect ~20-times less cancer cells than previously reported SWNTs.

Results

We have recently reported on the conjugation of cyclic Arg-Gly-Asp (RGD) peptides to pegylated SWNTs\textsuperscript{8} and their use as photoacoustic imaging agents\textsuperscript{7}. In order to enhance the photoacoustic signal of the SWNTs, we attached Indocyanine Green (ICG) and QSY-21 dyes to the surface of the SWNTs through pi-pi stacking interactions\textsuperscript{9} (see Methods section for more details). The ultra-high surface area of SWNTs allows highly efficient loading of aromatic molecules such as ICG and QSY-21 on the nanotube surface. This created two new kinds of photoacoustic agents; SWNT-ICG and SWNT-QSY (Fig. 1a). The particles were targeted using the RGD-peptide to $\alpha_v\beta_3$ integrins, which are over-expressed in tumor vasculature, while control untargeted particles were synthesized using a non-targeted peptide, RAD.

The optical absorbance spectra of the two new particles suggest that 710 nm and 780 nm are the preferable wavelengths for scanning SWNT-QSY and SWNT-ICG respectively (Fig. 1b). At their respective absorbance peaks, the SWNT-QSY and SWNT-ICG particles exhibit a 17 and 20-fold higher absorbance respectively as compared with plain SWNTs. Since blood absorption is significantly reduced at 780 nm compared to 710 nm, SWNT-ICG was the particle of choice for the small animal experiments for this study. Importantly, the attachment of RGD or RAD peptides to SWNT-ICG had little effect on the particles’ absorbance. We constructed a non-absorbing and non-scattering agarose phantom with inclusions of SWNT-ICG-RGD at increasing concentrations from 0.5 nM to 121.5 nM in multiples of 3 (n = 3 samples of each concentration). The photoacoustic signal produced by the SWNT-ICG-RGD particles correlated well with the nanoparticle concentration ($R^2=0.983$) (Fig. 1c).

We further validated that the new particles are stable in serum (see Supplementary Information and Fig. S1). The particle’s photobleaching (loss of optical absorption due to continuous light exposure) was characterized and found to be relatively small, 30% bleaching over 60 min of typical laser irradiation (see
We then tested the particle’s sensitivity in living subjects by subcutaneously injecting the lower back of mice (n = 3) with 30 μl of SWNT-ICG-RAD mixed with matrigel at increasing concentrations of 820 pM to 200 nM in multiples of 3. Matrigel alone produced no significant photoacoustic signal (data not shown). Upon injection, the matrigel solidified, fixing the SWNT-ICG-RAD in place and three-dimensional (3D) ultrasound and photoacoustic images of the inclusions were acquired (Fig. 2a). While the ultrasound images visualized the mouse anatomy (e.g., skin and inclusion edges), the photoacoustic images revealed the SWNT-ICG-RAD contrast in the mouse. The photoacoustic signal from each inclusion was quantified using a three dimensional region of interest (ROI) drawn over the inclusion. We observed a linear correlation ($R^2 = 0.97$) between the SWNT-ICG-RAD concentration and the corresponding photoacoustic signal (Fig. 2b). Tissue background signal was calculated as the average photoacoustic signal in areas where no contrast agent was injected. Extrapolation of the signal-concentration graph reveals that 170 pM of SWNT-ICG-RAD gives the equivalent photoacoustic signal as the tissue background (i.e., signal to background ratio = 1). This value represents over 300-times improvement in sensitivity compared to plain SWNTs.

Finally, we tested the nanoparticles targeting ability in living mice. Mice bearing U87MG tumor xenografts (150 mm$^3$ in size) were injected through the tail vein (IV) with 200 μl of either targeted SWNT-ICG-RGD or untargeted SWNT-ICG-RAD particles (n = 4 mice per group) at a concentration of 1.2 μM. We acquired 3D photoacoustic and ultrasound images of the entire tumor area before and up to 4 hours after the injection. Mice injected with the targeted SWNT-ICG-RGD particles show significantly higher photoacoustic signal in the tumor compared with the control group (Fig. 3a). The ultrasound images were used for visualizing the boundaries of the tumor as well as to validate that no significant movement (beyond 100 μm) had occurred throughout the scan. While the pre-injection photoacoustic signal is primarily due to the tumor’s blood content, post-injection photoacoustic signal consists of both blood and SWNT-ICG. To subtract out the blood signal from the images, a subtraction image calculated as the 2 hour post-injection minus the pre-injection image was calculated. Measurement of the photoacoustic signal from a 3D ROI around the tumor (Fig. 3b) showed that the photoacoustic signal in the tumor was significantly higher in mice injected with SWNT-ICG-RGD as compared with the control particles SWNT-ICG-RAD ($p < 0.001$). For example, at 2 hours post-injection, mice injected with SWNT-ICG-RGD showed over 100% higher photoacoustic signal in the tumor than mice injected with the control SWNT-ICG-RAD.

To compare the performance of plain SWNT-RGD to SWNT-ICG-RGD, we incubated U87MG cells, which express the target $\alpha_v\beta_3$ on their surface, with either particle solution for 2 hours. After incubation, the
cells were washed 3 times with cold saline to remove unbound particles and placed in a phantom at increasing concentrations from $25 \times 10^3$ to $6 \times 10^6$ cells per well ($n = 3$ samples per group) and imaged with the photoacoustic system (Fig. 4a). Quantitative analysis of the photoacoustic signal from the phantom revealed that cells exposed to SWNT-ICG-RGD were detected at 20-times lower concentration than cells exposed to plain SWNT-RGD ($p < 0.0001$) (Fig. 4a-b). These observations are consistent with the optical absorbance of SWNT-ICG-RGD being ~20 times higher than plain SWNT-RGD.

Finally, we show that the two kinds of photoacoustic imaging agents we synthesized, SWNT-ICG and SWNT-QSY can be imaged simultaneously due to their unique, though overlapping, absorbance spectra (Fig. 1b). We created an agarose gel phantom containing increasing concentrations of SWNT-ICG and decreasing concentrations of SWNT-QSY (starting from 100nM:0nM up to 0nM:100nM respectively). Photoacoustic images of the phantom were taken at wavelengths of 700, 730, 760, 780, and 800 nm and a spectral un-mixing algorithm was then used to separate each particle’s signal to an individual image (Fig. 5).

We have synthesized, characterized and demonstrated the application of dye-enhanced SWNTs as ultra-high sensitivity photoacoustic imaging agents. A concentration of 170 pM was estimated to produce an equivalent photoacoustic signal as tissue background signal, representing 300-times higher sensitivity than plain SWNTs in living mice. This improvement is likely due to both the higher optical absorption of the particles as well as the fact that the new particle’s absorption peak is at 780nm where the background tissue photoacoustic signal is greatly reduced. Intravenous injection of RGD-targeted SWNT-ICG particles to tumor-bearing mice led to significantly greater accumulation of the particles in the tumor compared to non-targeted control particles. We demonstrated the ability to multiplex 2 kinds of dye-enhanced SWNTs and showed the ability to detect 20-times fewer cancer cells when using SWNT-ICG-RGD as the imaging agent, as compared with plain SWNT-RGD. These results agree with the fact that SWNT-ICG has ~20 times greater optical absorbance compared to plain SWNT. Applications of the enhanced particles may therefore be exploited to lead to the earlier detection of cancer by providing the ability to detect smaller tumors.

The in-vivo targeting study results are likely negatively influenced by the effect of photo-bleaching, where continued laser light exposure of tumor causes reduction in the optical absorption (and photoacoustic signal) of particles that are bound to the tumor. This particularly affect the targeted group, SWNT-ICG-RGD, and to a much lesser extent the untargeted group, SWNT-ICG-RAD, which continue to circulate through the animal’s blood stream unexposed to laser irradiation. Therefore, it is likely that the difference between these two groups is even greater in reality than reflected in the results.

Most of the work done on photoacoustic contrast agents has been focused on gold nanoparticles as well as other kinds of nanoparticles. However, the main challenge that has yet been solved is the delivery of such agents to the tumor in sufficient amounts to create detectable and specific signal. This is likely due to the
particles’ large size that leads to rapid clearance by the reticuloendothelial system (RES) upon intravenous injection, preventing the particles from accumulating at the tumor site. In contrast, the SWNTs used here are 1-2 nm in diameter and 50-300 nm in length. Since the dye we used was attached to the surface of the SWNTs, under the PEG, it is expected that the total particle size was not significantly changed, thereby allowing the particles to keep a favorable bio-distribution as previously reported\(^8\). Hence, the dye-enhanced SWNTs presented in this work offer unprecedented photoacoustic signal strengths while maintaining relatively small size allowing them to target tumors upon intravenous injection. We have also previously published pilot toxicology studies of the SWNTs with encouraging results in mouse models\(^15\) as well as observed they are able to be excreted via the biliary pathway\(^16\).

The reason for loading a SWNT with many small dye molecules is the high efficiency of optical absorption of these dyes as compared to their weight. By this measure of absorption divided by weight, ICG is 7-times more efficient than SWNT and \(~8500\)-times more efficient than commercial gold nanorods with peak absorption at 780 nm.

The dye-enhanced SWNT photoacoustic contrast agents reported here have the capability to bind to molecular targets while maintaining a high photoacoustic signal. No other imaging modalities or reported imaging agents have the precise depth information and sub-millimeter spatial resolution at sub-nanomolar sensitivity that can be achieved with photoacoustic imaging of dye-enhanced SWNTs.

**Methods**

**Dye-enhanced SWNTs synthesis**

A complete description of the synthesis of SWNT-RGD and SWNT-RAD can be found elsewhere\(^8\). 250nM SWNT-RGD or SWNT-RAD was incubated with 2mM Indocyanine Green (ICG) molecules for overnight. ICG (Spectrum Laboratory Products, CA) (20 mM) was dissolved in DMSO first and then added to SWNT water solutions with a final DMSO concentration of 10% by volume. Unbound ICG molecules were removed from the solution by filtration through 100 kDa centrifuge filters (Millipore) and washed for 6-8 times. The SWNTs used in this work were 50-300 nm in length and 1-2 nm in diameter. The molar concentrations are based on an average molecular weight of 170 kDa per SWNT (150 nm in length and 1.2 nm in diameter). SWNT-QSY particles were synthesized the same way except replacing ICG with QSY-21.

**Statistical methods.** For the SWNT-ICG tumor targeting experiment, we used a mixed effects regression of signal on fixed factors of time, the square of the time, and group, and random factor of mouse to test the hypothesis that mice injected with SWNT-ICG-RGD showed an increased photoacoustic signal over time in the tumor compared with the control group injected with SWNT-ICG-RAD. There were significant effects of group
(Group 1 higher than Group 2, \( p=.001 \)), and linear and quadratic effects of time (\( p<.001 \) and \( p=.019 \), respectively). There was no significant interaction between group and time effects (\( p=.915 \)). For the cell uptake studies, we used the 1-tailed student’s t-test to test whether the group in which U87MG cells were exposed to SWNT-ICG-RGD had statistically higher signal than the group of cells that was exposed to SWNT-ICG-RAD.

For the experiment comparing SWNT-ICG-RGD to SWNT-RGD in-vitro, signal was compared between groups by a Wilcoxon test stratified by cell concentration. All statistics were done with Stata Release 9.2 (StataCorp LP, College Station, TX).

**Mouse arrangement in the photoacoustic system.** All animal experiments were performed in compliance with the Guidelines for the Care and Use of Research Animals established by the Stanford University Animal Studies Committee. A complete description of the photoacoustic system can be found in the Supplementary Information. Female nude mice were used for all the photoacoustic studies. The mice that were scanned in the photoacoustic system were fully anesthetized using isoflurane delivered through a nose-cone. Prior to the photoacoustic scan, the areas of interest were covered with agar gel to stabilize the area and minimize any breathing and other motion artifacts. A saran-wrap water bath was placed on top of the agar gel. An ultrasonic transducer, placed in the water bath, was therefore acoustically coupled to the mouse tissues. This setup allowed the ultrasonic transducer to move freely in 3D while not applying any physical pressure on the mouse.

**SWNT-ICG tumor targeting in living mice.** Two groups of female nude mice (\( n = 3 \) in each group) 6-8 weeks old were inoculated subcutaneously at their lower right back with \( 10^7 \) U87MG cells (American Type Culture Collection, ATCC) suspended in 50 µl of saline (PBS pH 7.4 1X, Invitrogen). The tumors were allowed to grow to a volume of ~100-150 mm\(^3\). Before the injection, photoacoustic and ultrasound images of the mice were taken. Photoacoustic excitation light was 780 nm to match the absorption peak of SWNT-ICG. The mice were then injected with 200µl of 1.2µM or either targeted SWNT-ICG-RGD or control SWNT-ICG-RAD into the tail-vein (IV). During the injection the mice positioning was not changed. After the injection, photoacoustic and ultrasound images were acquired at: 0.5, 1, 2, 3, 4 hrs. post-injection. The scanning area varied between mice depending on the tumor orientation, but typically was ~10 mm x 10 mm, with a step size of 0.25 mm. At 4 hr post-injection, the mice were sacrificed. Using AMIDE software\(^{17} \), a three dimensional ROI was drawn over the tumor volume (which was clearly illustrated in the ultrasound images). The mean photoacoustic signal in the tumor ROI was calculated for each photoacoustic image.
References


Acknowledgments

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Figures:

A

Targeting Peptide

SWNT-QSY

SWNT-ICG

B

Optical Absorbance (a.u.)

Wavelength (nm)

Plain SWNT

SWNT-ICG-RAD

SWNT-ICG-RGD

SWNT-QSY
Figure 1. Characterization of the dye-enhanced SWNT. **a**, Illustration of SWNT-ICG and SWNT-QSY. ICG and QSY-21 (red molecules) are attached to the SWNT surface through non-covalent pi-pi stacking bonds. Polyethylene glycol-5000 (blue molecules) is conjugated to a targeting peptide in one end and to the SWNT surface on the other end through phospholipids. **b**, Optical spectra of plain SWNT (green), SWNT-ICG-RGD (red), SWNT-ICG-RAD (blue) and SWNT-QSY-RGD (black). The similarity of SWNT-ICG-RAD and SWNT-ICG-RGD spectra suggests that the peptide conjugation does not notably perturb the photoacoustic signal. **c**, The photoacoustic signal produced by SWNT-ICG was observed to be linearly dependent on the concentration \( R^2 = 0.9833 \).
Figure 2. Photoacoustic detection of SWNT-ICG in living mice. a, Mice were injected subcutaneously with SWNT-ICG at concentrations of 0.82-200 nM. The images represent ultrasound (gray) and photoacoustic (green) vertical slices through the subcutaneous injections (dotted black line). The skin is visualized in the ultrasound images, while the photoacoustic images show the SWNT-ICG distribution. The white dotted lines on the images illustrate the approximate edges of each inclusion. b, The photoacoustic signal from each inclusion was calculated using 3D regions of interest and the ‘background’ represents the endogenous signal measured from tissues. The error bars represent standard error (n = 3 mice). Linear regression ($R^2 = 0.97$) of the photoacoustic signal curve estimates that a concentration of 170 pM of SWNT-ICG will give the equivalent background signal of tissues.
Figure 3. SWNT-ICG-RGD tumor targeting in living mice. a, Ultrasound (gray) and photoacoustic (green) images of one vertical slice through the tumor (dotted black line). The ultrasound images show the skin and the tumor boundaries. Subtraction photoacoustic images were calculated as 2 hr post-injection minus pre-injection images. As can be seen in the subtraction images, SWNT-ICG-RGD accumulates in higher amount in the tumor.
as compared to the control SWNT-ICG-RAD. **b,** Mice injected with SWNT-ICG-RGD showed significantly higher photoacoustic signal than mice injected with the untargeted control SWNT-ICG-RAD (p < 0.001). The error bars represent standard error (n = 4 mice).

![Photoacoustic vertical slice image](image)

**Figure 4. Comparison of plain SWNT-RGD to SWNT-ICG-RGD.** **a.** Photoacoustic vertical slice image through a gel phantom containing increasing number of cells exposed to SWNT-ICG-RGD and SWNT-RGD. While 1.7x10^6 cells exposed to SWNT-RGD are barely seen on the image, a clear photoacoustic signal was observed from 1.4x10^5 cells exposed to SWNT-ICG-RGD. The signal inside the ROI (dotted white boxes) is not homogenous due to possible aggregates of cells. **b.** Quantitative analysis of the photoacoustic signals from the phantom (n = 3) showed that SWNT-ICG-RGD can see ~20-times less cancer cells than SWNT-RGD can (p < 0.0001). The background line represents the average background signal in the phantom. Linear regression was calculated on the linear regime of both curves.
Figure 5. Multiplexing of SWNT-ICG with SWNT-QSY particles in a phantom. A phantom with various concentrations of SWNT-ICG and SWNT-QSY was scanned under the photoacoustic instrument at wavelengths of 700, 730, 760, 780, and 800 nm. A spectral un-mixing algorithm based on least-squares was used to separate the signals of SWNT-ICG particles (green) from SWNT-QSY particles (red). Notice that no SWNT-QSY signal is seen in the well with pure SWNT-ICG and vice versa, despite the fact that the two particles have overlapping spectra.
3. Abstract presented SPIE 2009:

Title: Photoacoustic Molecular Imaging using Single Walled Carbon Nanotubes in Living Mice

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Abstract Text for Online or Printed Programs (100 words, early release)

Photoacoustic molecular imaging is an emerging modality offering non-invasive high resolution imaging of diseases using an external photoacoustic imaging agent. Here we demonstrate for the first time the utility of single walled carbon nanotubes as disease-targeted photoacoustic imaging agents in living mice. The carbon nanotubes were conjugated to RGD-peptides to target the $\alpha_v\beta_3$ integrin that is associated with tumor angiogenesis. Intravenous administration of these targeted carbon nanotubes to tumor-bearing mice showed significantly higher photoacoustic signal in the tumor as compared to non-targeted carbon nanotubes. These results were verified ex-vivo using a Raman microscope that is sensitive to SWNTs Raman signal.

Keywords:
Photoacoustic Imaging
Opotoacoustic Imaging
Molecular Imaging
Carbon Nanotubes

Abstract Text for Technical Review Purposes (250 words, publicized during the meeting)

Photoacoustic molecular imaging is an emerging technology offering non-invasive high resolution imaging of the molecular expressions of a disease using a photoacoustic imaging agent. Here we demonstrate for the first time the utility of single walled carbon nanotubes (SWNTs) as targeted imaging agents in living mice bearing tumor xenografts. SWNTs were conjugated with polyethylene-glycol-5000 connected to Arg-Gly-Asp (RGD) peptide to target the $\alpha_v\beta_3$ integrin that is associated with tumor angiogenesis.

In-vitro, we characterized the photoacoustic spectra of the particles, their signal linearity and tested their uptake by $\alpha_v\beta_3$-expressing cells (U87MG). The photoacoustic signal of SWNTs was found not to be affected by the RGD conjugation to the SWNTs and was also found to be highly linear with concentration ($R^2 = 0.9997$ for 25-400nM). The cell uptake studies showed that RGD-targeted SWNTs gave 75% higher photoacoustic signal than non-targeted SWNTs when incubated with U87MG cells.

In-vivo, we measured the minimal detectable concentration of SWNTs in living mice by subcutaneously injecting SWNTs at increasing concentrations. The lowest detectable concentration of SWNTs in living mice was found to be 50nM. Finally, we administered RGD-targeted and non-targeted SWNTs via the tail-vein to U87MG tumor-bearing mice (n=4 for each group) and measured the signal from the tumor before and up to 4 hours post-injection. At 4 hours post-injection, tumors of mice injected with RGD-targeted SWNTs showed 8 times higher photoacoustic signal compared with mice injected with non-targeted SWNTs. These results were verified ex-vivo using a Raman microscope that is sensitive to the SWNTs Raman signal.
4. Abstract 2 presented at SPIE 2009:

Enhanced Sensitivity Targeted Photoacoustic Molecular Imaging Agents in Living Mice

Adam de la Zerda, Zhuang Liu, Cristina Zavaleta, Sunil Bodapati, Robert Teed, Srikant Vaithilingam, Te-Jen Ma, Omer Oralkan, Xiaoyuan Chen, Butrus T. Khuri-Yakub, Hongjie Dai, Sanjiv Sam Gambhir

Abstract Text for Online or Printed Programs (100 words, early release)
Photoacoustic imaging of living subjects offers significantly higher spatial resolution at increased tissue depths compared to purely optical imaging techniques. We developed a new version of extremely bright photoacoustic imaging agent based on single walled carbon nanotubes (SWNTs) and the small molecular dye Indocyanine Green (ICG). We measured the photoacoustic signal from the new particle in-vitro and in-vivo and found it is 17-times higher than plain SWNTs. We conjugated the particles to RGD-peptides to target the \( \alpha_v \beta_3 \) integrin associated with tumor angiogenesis and showed that it can bind selectively to tumors when injection intravenously to living mice.

Abstract Text for Technical Review Purposes (250 words, publicized during the meeting)
Photoacoustic imaging of living subjects offers high spatial resolution at increased tissue depths compared to purely optical imaging techniques. We have recently shown that intravenously injected single walled carbon nanotubes (SWNTs) can be used as targeted photoacoustic imaging agents in living mice using RGD peptides to target \( \alpha_v \beta_3 \) integrins.

We have now developed a new targeted photoacoustic imaging agent based on SWNTs and Indocyanine Green (SWNT-ICG) with absorption peak at 780nm. The photoacoustic signal of the new imaging agent is enhanced by \( \sim 17 \) times as compared to plain SWNTs. To synthesize this particle, SWNTs were coupled to RGD peptides through polyethylene glycol-5000 grafted phospholipid. ICG molecules were then attached to the surface of each SWNT non-covalently through pi-pi stacking interactions.

In-vitro, we measured the serum stability of the particles and through cell uptake studies with U87MG cells, we verified that the particles bind selectively to \( \alpha_v \beta_3 \) integrin. In-vivo, we injected the imaging agents subcutaneously to living mice (n=4) and were able to detect concentrations as low as 3nM, a 17-fold enhancement in sensitivity over plain SWNTs (p<0.05). Finally, we injected U87MG tumor-bearing mice (n=4) with RGD-targeted SWNT-ICG via the tail-vein. Control mice were injected with non-targeted SWNT-ICG. Upon administration, the RGD-targeted particles created a significantly higher photoacoustic signal in the tumors than the non-targeted particles (p<0.05). These results were verified ex-vivo using a Raman microscope sensitive to the SWNTs Raman signal.

In summary, the new SWNT-based particle can target tumors in living mice while possessing a very high photoacoustic signal.
5. **Abstract presented at World Molecular Imaging Congress 2009**

**Ultra High Sensitivity Targeted Photoacoustic Imaging Agents for Cancer Early Detection in Living Mice**

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Photoacoustic molecular imaging of living subjects offers high spatial resolution at increased tissue depths compared to optical imaging strategies. We have recently demonstrated single walled carbon nanotubes (SWNTs) conjugated to Indocyanine Green (SWNT-ICG) as targeted photoacoustic imaging agents *in-vitro*.

In the current work, we created a significantly improved SWNT-ICG particle with over 1000-times better sensitivity than plain SWNT and demonstrated their ability to target tumors when injected intravenously to a living mouse.

The targeted SWNT-ICG particles were synthesized by coupling of ICG molecules to the surface of SWNT-RGD particles through pi-pi stacking interactions. Control SWNT-ICG particles were created using the untargeted SWNT-RAD instead.

We verified the particles are stable in serum and target αvβ3 integrin through cell uptake studies with U87 cells. We found the photoacoustic signal produced by the particles to be highly linear to their concentration both in phantom studies ($R^2 = 0.99$) as well as in living mice injected with the particles subcutaneously ($R^2 = 0.971$). We further measured the detection sensitivity of SWNT-ICG in living mice ($n = 3$ mice) and found it to be 30 pM. This represents more than 3 orders of magnitude improvement compared to plain SWNTs sensitivity in living mice ($p < 0.05$). Furthermore, xenograft-bearing mice were tail-vein injected with RGD-targeted SWNT-ICG. At 2 hours post-injection, mice injected with the RGD-targeted particles showed 2.1-times higher photoacoustic signal in the tumor compared to mice injected with control particles ($p < 0.05$, $n = 4$ mice). Finally, we demonstrated the superiority of the SWNT-ICG-RGD particles by incubating them with U87 cells and detecting in living mice 1000-times such cells than if the cells were incubated with plain SWNT-RGD.

This is the first photoacoustic imaging agent tested and targeted in living animals that we know of that can reach such a high sensitivity of 30 pM.