Size-tunable ICG-based contrast agent platform for targeted near-infrared photoacoustic imaging.

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ABSTRACT

Near-infrared photoacoustic imaging (NIR-PAI) combines the advantages of optical and ultrasound imaging to provide anatomical and functional information of tissues with high resolution. Although NIR-PAI is promising, its widespread use is hindered by the limited availability of NIR contrast agents. J-aggregates (JA) made of indocyanine green dye (ICG) represent an attractive class of biocompatible contrast agents for PAI. Here, we present a facile synthesis method that combines ICG and ICG-azole dyes for producing contrast agents with tunable size down to 230 nm and direct functionalization with targeting moieties. The ICG-JA platform has a detectable PA signal in vitro that is two times stronger than whole blood and high photostability. The targeting ability of ICG-JA was measured in vitro using HeLa cells. The ICG-JA platform was then injected into mice and in vivo NIR-PAI showed enhanced visualization of liver and spleen for 90 min post-injection with a contrast-to-noise ratio of 2.42.

1. Introduction

Photoacoustic imaging (PAI) [1-3] is a relatively low-cost non-invasive biomedical imaging modality, which combines the advantages of optical imaging (e.g., high contrast and molecular specificity) and ultrasound imaging (e.g., deep penetration in tissue). Near-infrared I (NIR-I) illumination (620–950 nm) can be used with PAI to reduce photon scattering in biological tissues and improve imaging depth [4]. To this end, several exogenous contrast agents (CA) such as organic particles, metallic nanoparticles, [5,6] or polymer-based nanomaterials [7,8] have been designed for NIR-PAI to enhance the image contrast. Hence, NIR-PAI has found use in various pre-clinical applications, including imaging tumor lesions, [9] detecting metastatic lymph nodes, [10,11] delineation of tumor margins, [12,13] identification of pathological tissues, [14] and measurement of blood oxygenation level [15]. However, even with the current catalogue of available CAs, NIR-PAI still requires improved CAs that combine critical features such as stability in vivo, solubility, biocompatibility, photostability and targeting capabilities [16,17]. In addition, there is also a need to develop robust and simple methods that reproducibly synthesize different sized CAs depending on the biological system and applications [18,19]. Thus, it is essential to engineer a stable and biocompatible CA platform that can enable prescribed size tuning and easy functionalization with targeting moieties [16].

One of the main approaches taken for producing CA of a specific size and functionalization is to produce aggregates of dyes [20]. Indocyanine green (ICG) is an attractive candidate for this purpose since it is a biocompatible FDA-approved NIR dye currently used for PAI [9]. However, monomeric ICG dye has variation in its optical properties depending on its surrounding environment [21], has low photostability and its targeting chemistry is difficult [22]. On incubation in aqueous solutions at temperatures above 50 °C [23], ICG can self-assemble to

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form J-aggregates (JA) \[24,25\]. J-aggregates are stable head-tail arrangements of monomeric dye held together by non-covalent interactions with a strong red (bathochromic) shift of the absorbance peak coupled with higher absorbance \[26,27\]. Compared to monomeric ICG, ICG-JA have advantages like quenched fluorescence \[28\], a red-shifted absorption peak at a wavelength of 895 nm \[29\], stronger PAI signal, \[30\] and greater photothermal stability \[31\]. However, the formation of ICG-JA is a process of self-assembly of monomeric ICG, which often yields micron-sized, polydisperse aggregates. Therefore, most current methods of synthesizing monodisperse, nanometer-sized ICG-JA rely on syringe filtration \[28,29\] or encapsulation in nanocarriers \[24,32,33\]. The addition of targeting moieties also requires encapsulation in nanocarriers. These factors limit the flexibility of ICG-JA with respect to size, targeting and photoacoustic properties.

Hence, in this study, we report the scalable preparation of azide-modified ICG J-aggregates (JAAZ) that have controllable mean sizes ranging from 230 to 1200 nm, are amenable to direct functionalization, with NIR absorption, and a strong PA signal. As a proof of principle, we functionalize RGD peptide as a model of targeting moiety to enable targeting of cells that overexpress the integrin receptor such as HeLa cells. We determined the intensity of the PA signal in an agarose phantom and upon binding to HeLa cells. We further evaluated the biodistribution, PA properties and capability to visualize blood-rich organs of RGD functionalized JAAZ with preliminary experiments in vivo using tomographic PAI.

2. Results and discussion

2.1. Synthesis, characterization and stability of azide-modified J-aggregates

To synthesize azide-modified ICG-JA (JAAZ), we first incubated different ratios of ICG and ICG-azide dye in water using a procedure previously described for ICG only \(\text{Fig. 1A} \) \[29\]. After incubation for 20 and 40 h at 60 °C in water, we observed the typical JA peak at 895 nm for 1:10 ICG-azide:ICG solution but not for 100% ICG-azide dye or for ICG-azide:ICG molar ratios of 3:4, 1:2 and 1:4 (Fig. S1). However, for the JA formed using the 1:10 ratio, the Fourier-transform Infrared (FT-IR) spectrum did not show a peak between 2141 and 2100 cm⁻¹ (Fig. S2), which is a characteristic range for the azide group due to the strong asymmetric N = N = N stretching, \[34\] indicating that the formed JA did not contain ICG-azide but only ICG. To promote the incorporation of ICG-azide into the JA, we modified our initial protocol to include cations, which are known to influence aggregate formation \[35\]. Indeed, monovalent metal cations like Na⁺ and K⁺ are known to facilitate J-aggregate formation of anionic cyanine dyes such as ICG and pseu-doisylocyanine (PIC) due to ionic interactions between cations and the dye molecules that limit electrostatic repulsion between dyes \[36,37\]. Thus, we tested different concentrations of potassium chloride (KCl) (1, 5, 10 and 20 mM) with 1:10 ICG-azide:ICG molar ratio solutions at a total dye concentration of 1 mM. After 20 h of incubation at 60 °C, these conditions showed the presence of a red-shifted absorption peak at 895 nm and the disappearance of both the monomer and the dimer peaks (780 and 715 nm, respectively) \(\text{Fig. 1B, Fig. S3}\). The size of the JAAZ particles (Fig. S4A) formed with 1 mM, 5 mM, 10 mM and 20 mM of KCl were 771 ± 41 nm, 571 ± 120 nm, 602 ± 396 nm, and 1326 ± 73 nm, respectively. JAAZ particles formed at 5 mM and 10 mM were too polydisperse (Fig. S4B); hence, these conditions were not utilized further for synthesis. Unless explicitly stated, 20 mM of KCl was selected for the synthesis of JAAZ particles for all further experiments because it resulted in the formation of monodisperse micron-sized particles. We then tested formation of JAAZ particles with different molar ratios of ICG-azide to ICG (1:10, 1:5 and 1:3) using 20 mM KCl to detect the incorporated azide groups using FT-IR. FT-IR spectra of all synthesized JAAZ particles with KCl show a peak at 2100 cm⁻¹ due to incorporated azide groups, which is absent from the control ICG-JA (Figs. S5B-D). We further studied temperature as an experimental parameter effecting the formation of JAAZ particles. The formation of JAAZ particles (1:10 ICG-azide:ICG molar ratio, 20 mM KCl dye concentration of 1 mM) was quantified by the intensity of the 895 nm absorption peak (Fig. S6) at 50, 60, 60 °C and 70 °C. At 50 °C, the intensity of the 895 nm was lower than 60 and 70 °C at 20 h of incubation. At 70 °C, the intensity decreased after 14 h indicating the slight decomposition of ICG dye. Hence, incubation at 60 °C for 20 h was appropriate for JAAZ formation. Hence, all micron-sized JAAZ particles were synthesized using 1:10 ICG-azide:ICG molar ratio solutions incubated in 20 mM KCl for 20 h at 60 °C at a total dye concentration of 1 mM, unless stated otherwise.

Before further functionalization with targeting moieties, excess free dye and residual salt were efficiently removed from the prepared ICG-JA and JAAZ using spin column filtration (Fig. S7A). There was no fluorescence emission at 800 nm for both the ICG-JA and JAAZ particles as previously described in literature \[28\] (Fig. S7B), which is an important characteristic since it ensures maximum conversion of the absorbed photons to acoustic signal. Therefore, the experimental parameters

Fig. 1. Synthesis of ICG J-aggregates (ICG-JA) and azide-modified ICG J-aggregates (JAAZ) particles. (A) Schematic of the procedure used to synthesize ICG-JA and JAAZ particles. (B) (top) Representative absorbance spectra of ICG-JA and JAAZ particles at 50 μM of total dye concentration. A change in the color of the solution is observable by the naked eye between monomeric ICG solution and the ICG-JA and JAAZ particle solutions due to the shift toward longer wavelengths.
studied for the formation of micron-sized J-aggregates with incorporated azide groups were KCl concentration and temperature.

After confirming formation of JAAZ particles, we examined the presence and accessibility of the azide group using measurements of size and surface charge. We first measured the surface zeta (ζ) potential of both ICG-JA and JAAZ, knowing that the difference in charge between ICG-azole dye and regular ICG dye should modify the overall surface charge of the JAAZ particles if ICG-azole dyes were to be displayed on the surface. The average ζ potential measured was −50.32 mV and −60.11 mV for the JAAZ and the ICG-JA particles, respectively (Fig. 2A). The less-negative zeta potential of JAAZ particles indicates the presence of azide groups on the surface of the JAAZ particles. Moreover, the narrow surface charge distribution confirms the presence of a single J-aggregate population (Fig. S8). Using dynamic light scattering (DLS), we measured the average hydrodynamic diameter of ICG-JA and JAAZ particles as 1.17 ± 0.26 and 1.34 ± 0.18 μm, respectively (Fig. 2B). The slightly larger size of JAAZ particles could be attributed to a less densely-packed or organized arrangement of ICG molecules due to the presence of azide groups. Scanning electron microscopy (SEM) reveals that JAAZ particles have a uniform morphology with an average diameter of 1.1 μm, in good agreement with the DLS data (Fig. 2C and S9A). Energy-dispersive X-ray (EDX) spectroscopy confirmed the elemental composition of JAAZ particles (Fig. S9B).

In order to determine the stability of the physical properties of JAAZ particles, we lyophilized and resuspended the JAAZ particles in water. There was no significant change in the absorption spectra, size and ζ potential, suggesting that JAAZ particles can be stably stored in lyophilized form at room temperature protected from light (Fig. S10 and Table S1). Moreover, they are easily dispersed in different media, such as 1X phosphate buffered saline (PBS), Dulbecco’s Modified Eagle Medium (DMEM), DMEM + 10% fetal bovine serum and 100% fetal bovine serum (FBS, mimicking biological conditions). JAAZ particles were stable in 1X PBS at 37 °C and different pH (5, 7 and 9) for up to 126 h (Fig. 2D). They were also stable in DMEM, DMEM + 10% FBS and 100% FBS for 48 h as observed by the lack of intensity difference in the 895 nm peak (Fig. 2E), indicating JAAZ particles were stable in in-vitro physiological conditions.

Since J-aggregates are an assembly of dye molecules, it was important to determine the concentration of dye within the cluster before functionalization. Hence, we disassembled the J-aggregates into free dye using Triton X-100 [29] to determine the equivalent dye concentration (or total dye concentration) and compare it from one sample to another. A similar concentration of dye was determined using a standard curve (Fig. S11) for both ICG-JA and JAAZ particles, indicating that the efficiency of azide-modified J-aggregate formation is similar to ICG-JA.

Fig. 2. Characterization of JAAZ particles prepared in 20 mM KCl at 1 mM dye concentration. (A) Zeta potential measurements of ICG-JA and JAAZ particles. (B) Dynamic light scattering measurements of ICG-JA and JAAZ particles. (C) SEM images of six selected JAAZ particles prepared in 20 mM KCl with 1 mM dye. (Scale bar: 2 μm, the complete SEM image is provided in Fig. S9). (D) UV-Vis-NIR absorbance of JAAZ particles incubated in solutions of pH 5 (red), 7 (blue), and 9 (green) at 37 °C for 126 h. (E) Percentage stability of JAAZ particles in water as control (grey), DMEM (blue), DMEM + 10% FBS (orange) and 100% FBS (black). Stability over time was quantified as difference in the intensity of the 895 nm absorption peak when compared to control at 0 min. All JAAZ samples were made with 1:10 molar ratio of ICG-azole/ICG in 20 mM KCl at dye concentration of 1 mM (n = 5 for A and B, n = 3 for D and E, all results are mean±SD with p value calculated using unpaired t-test; * and *** = p < 0.05, all P values provided in Table S3).
2.2. Functionalization of the azide-modified J-aggregates

To enable further functionalization of JAAZ with streptavidin, we performed copper-free click chemistry to graft DBCO-PEG-biotin linker onto the azide groups available on their surface to generate biotinylated JAAZ particles (Bio-JAAZ). After removal of the excess of DBCO-biotin linker, we incubated the particles with streptavidin to prepare Strep-JAAZ. Biotinylated RGD was then attached to available streptavidin to create RGD-JAAZ, as shown in Fig. 3 A. A summary of the concentration of biomolecules used to functionalize JAAZ to create Bio-JAAZ, Strep-JAAZ and RGD-JAAZ is provided in Table S2. Conjugation of the aggregates with biotin, streptavidin and RGD did not affect the aggregates’ stability as demonstrated with the absorbance spectra of Bio-JAAZ, Strep-JAAZ and RGD-JAAZ showing the typical J-aggregate absorption peak at 895 nm (Fig. 3B). Bio-JAAZ particles exhibit an average \( \zeta \) potential of \( -54.1 \) mV, a non-significant change compared to JAAZ particles. In contrast, the attachment of streptavidin changed the average \( \zeta \) potential of the particles to \( -15.8 \) mV, making them significantly less charged than JAAZ (Fig. 3C) as expected given the charge of streptavidin (38). DLS measurements show the size of Bio-JAAZ, Strep-JAAZ and RGD-JAAZ particles were \( 1.26 \pm 0.34 \) \( \mu \)m, \( 1.51 \pm 0.35 \) \( \mu \)m and \( 1.72 \pm 0.36 \) \( \mu \)m, respectively (Fig. S12A). Attachment of streptavidin caused no statistically significant increase in the size of JAAZ particles, confirming the absence of aggregation of Strep-JAAZ particles. Both Strep-JAAZ and RGD-JAAZ particles have a PDI of 0.26 when compared to JAAZ particles with a polydispersity index (PDI) of 0.37 (Fig. S12B). SEM images and EDX analysis (Fig. 3D) show the average size of Strep-JAAZ particles as 917.11 nm and confirm the coating of streptavidin present when compared to JAAZ samples. EDX analysis of three uncoated JAAZ particles is shown for comparison. All JAAZ samples were made were made with 1:10 molar ratio of ICG-azide:ICG without affecting the stability of the aggregates. We observed that modification of JAAZ particles with a soluble protein like streptavidin increased their solubility, preventing sedimentation. To test this, JAAZ and Strep-JAAZ particles were left to sediment at room temperature on a benchtop. No sedimentation was observed for up to 8 h with Strep-JAAZ particles, whereas a visible pellet was formed for the JAAZ particles. At 24 h, Strep-JAAZ particles settled down at the bottom of the centrifuge tube. However, upon slight mixing, they returned into solution, whereas a visible pellet was observed for JAAZ particles at 24 h, which required vortex mixing to be properly dispersed (Fig. S14A and S14B).

The results presented in this section demonstrate our capability to conjugate biomolecules to the surface of JAAZ particles.

2.3. Synthesis and characterization of nanosized azide-modified J-aggregates

Because the micrometer size of JAAZ particles might restrict their use in applications that require contrast agents to pass through biological
bars, such as the blood-brain barrier, we adapted our synthesis protocol to enable control over the size of JAAZ particles. According to previous literature, [23,35,39] J-aggregate formation is dependent on time, [40,41] salt and dye concentration; hence, we explored the impact of these three parameters and their combination on the size of these JAAZ particles. Solutions of 250, 500, 750 and 1000 µM total dye with 1:10 ICG-azide:ICG molar ratio were incubated at 60 °C for 20 h under KCl concentrations of 0.1, 1 and 20 mM. At 0.1 mM KCl, the 895 nm absorbance peak appeared after incubation for 6 h for 1000 and 750 µM of total dye. For 500 and 250 µM concentrations, 8 and 14 h were required for the formation of J-aggregates, respectively (Fig. S1A-D). For 1 and 20 mM KCl, a J-aggregate peak formed after 6 h for all dye concentrations (Fig. S16 and S17). The size of the formed JAAZ particles was measured using DLS for the above-mentioned conditions after 20 h. At higher total dye concentrations (1000 and 750 µM) for all salt molarities, the size of the formed JAAZ particles ranged from 700 nm to 1.4 µm. Hence, we determined that micron sized JAAZ particles can be synthesized using total dye concentrations of 1000 and 750 µM with 0.1, 1 or 20 mM KCl at 20 h (Fig. S1A-B and Fig. S19). On reducing the initial concentration of total dye to 500 and 250 µM, at lower salt molarities of 0.1 mM and 1 mM KCl, nanoscale JAAZ (N-JAAZ) of average size ranging from 230 to 600 nm were formed (Fig. S18C-D). Using total dye concentrations of 500 and 250 µM in 0.1 mM KCl at 14 h of incubation, we demonstrated the synthesis of 350 nm and 225 nm N-JAAZ particles, respectively (Fig. 4A). These results indicate that varying the concentration of total dye, concentration of KCl and incubation time can be used to tune the size of JAAZ particles. Specifically, decreased total dye concentration and decreased concentration of KCl at 14 h formed N-JAAZ particles. We were able to synthesize N-JAAZ particles with average size of 225 nm (Fig. 4B) and a PDI of 0.20 with 250 µM total dye and 0.1 mM KCl at 14 h. Using a higher concentration of KCl and total dye (1 mM KCl, 750 µM dye, 20 h), JAAZ particles of 640 nm were obtained (Fig. 4B). JAAZ particles of 1.3 µm, as shown in Figs. 2B and 4A, were formed in a solution of 20 mM KCl, 1000 µM total dye at 20 h. A detailed summary of the size of JAAZ particles formed varying time, KCl and total dye concentrations is given in Fig. S19. To demonstrate the difference in sizes between the JAAZ and N-JAAZ, the two samples were loaded onto an agarose gel where the negatively charged JAAZ and N-JAAZ particles were expected to move toward the positive electrode. N-JAAZ particles of ~230 nm moved through the gel and formed a distinct green band after traversing the gel as seen in Fig. 4C, whereas the size of JAAZ particles (~1 µm) prevents them from migrating through the gel. The size of N-JAAZ particles was further confirmed by SEM images (Fig. 4D) which showed particles with an average diameter of 175 nm. EDX analysis (Fig. S20) confirmed the composition of N-JAAZ particles.

Following the same biomolecule functionalization strategy used for JAAZ particles (Fig. 3A), we attached biotin and streptavidin onto the surface of N-JAAZ. After attaching DBCO-Biotin and streptavidin, the average hydrodynamic diameter of N-Bio-JAAZ and N-Strep-JAAZ particles were 210 ± 30 nm and 255 ± 27 nm, respectively. After RGD attachment, the average size of N-RGD-JAAZ particles was 313 ± 37 nm (Fig. 4E), a statistically non-significant increase compared to N-JAAZ particles. Hence, the synthesized N-JAAZ particles can be functionalized with RGD yielding nanosized and targeted J-aggregates. As in the case of JAAZ particles, attachment of the slightly negatively charged streptavidin to N-JAAZ particles makes their zeta potential more positive. The average ζ potentials of N-JAAZ, N-Bio-JAAZ, N-Strep-JAAZ and N-RGD-JAAZ particles were −52 mV ± 5.1, −53 mV ± 5.7, −21 mV ± 3.3 and −18 ± 2.4 mV respectively (Fig. 4F), comparable to those obtained for JAAZ particles. To determine if the reduction in size resulted in changes in the stability of N-JAAZ particles, we performed stability studies in DMEM, DMEM + 10% FBS and 100% FBS. When incubated in DMEM and DMEM + 10% FBS for 24 h at 37 °C, no significant differences were

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**Fig. 4. Characterization of N-JAAZ.** (A) Absorbance spectrum of N-JAAZ particles formed in 0.1 mM KCl using 1:10 ICG-azide:ICG solutions at 250 and 500 µM of total dye. Representative absorbance spectrum is for particles formed at 14 h of incubation at 60 °C. Inset: Size of N-JAAZ after formation for 8, 14 and 20 h. (B) Size of N-JAAZ and JAAZ particles. The total concentration of dye used for 0.1 mM KCl samples was 250 µM and for 1 mM and 20 mM KCl samples, the dye concentration was 1 mM. The time of incubation was 14 h for 0.1 mM and 20 h for both 1 mM and 20 mM KCl. (C) Representative agarose gel electrophoresis of JAAZ (1 mM KCl, 1 mM dye, 20 h) and N-JAAZ (0.1 mM KCl, 250 µM KCl, 14 h) particles. (D) SEM images of ten selected N-JAAZ particles prepared in 0.1 mM KCl with 250 µM of dye. (Scale bar: 250 nm, complete image is provided in Fig. S20). (E) DLS measurements of functionalized N-JAAZ particles show average hydrodynamic diameters of 217.5 ± 39 nm for bare N-JAAZ, 206.6 ± 30 nm for Bio-N-JAAZ, 256.5 ± 27 for Strept-N-JAAZ and 313.0 ± 37 nm for RGD-N-JAAZ, respectively. (F) Zeta potential values of functionalized N-JAAZ particles are −52 mV, −53.1 mV, −20.6 and −18 mV respectively. (n = 4 for A, E and F, n = 7 for N-JAAZ in E, all data presented as mean±SD, p-values are calculated using one-way ANOVA, *p < 0.05, **p < 0.01 and ***p < 0.001, ns, not significant; all P values are provided in Table S3).
observed in the intensity of the absorbance peak at 895 nm. However, there was a 20% decrease in stability of N-JAAZ particles in 100% FBS at 24 h of incubation (Fig. S21). After vacuum drying and resuspension, we observed a ~26% decrease in the absorbance peak (Fig. S10) and an 80 nm increase in the size of the N-JAAZ particles and no significant changes in the ζ potential of the particles, (Table S1). We also studied the shelf stability of N-JAAZ particles for two weeks in water at room temperature. As seen in Fig. S22, the size of the particles increased by 3% after 14 days, a statistically non-significant increase, which showed that the N-JAAZ particles are stable. Thus, we have designed an experimental protocol to control the size of JAAZ particles synthesizing stable N-JAAZ particles, which can be modified with biomolecules making targeted nanoscale J-aggregates.

2.4. Stability and cytotoxicity of the JAAZ and N-JAAZ particles in vitro

We also tested the hemotoxicity of JAAZ, N-JAAZ and RGD-JAAZ particles (Fig. S23A). Optical microscopy images of smeared red blood cells stained with all three types of JAAZ particles showed no difference in morphology compared to control blood indicating no adverse reaction of all combinations of JAAZ particles. To assess the cytotoxicity of different JAAZ particles, we performed an MTT assay using HeLa cells (Fig. S23B). HeLa cells were incubated with JAAZ, N-JAAZ and RGD-JAAZ at an equivalent dye concentration of 20 µM. With all three samples at high concentration, cells did not show any cytotoxicity after 24 h of incubation. These data indicate the in-vitro safety and efficacy of JAAZ particles.

Before testing the in vitro PA properties of JAAZ and N-JAAZ particles, we determined their stability in whole blood (Fig. S24). JAAZ particles were ~60% stable when incubated in whole blood at 37 °C for 24 h. N-JAAZ particles were 90% stable at the end of the 24-hour incubation. RGD-JAAZ particles were stable for 24 h, an increased stability when compared to uncoated particles.

2.5. Characterization of the photoacoustic properties of JAAZ particles in vitro

To quantify their use for PAI applications, we characterized the in vitro photoacoustic properties of JAAZ and N-JAAZ along with RGD-JAAZ. The photoacoustic setup shown in Fig. 5A includes the different J-aggregate samples embedded in agarose submerged in a water reservoir and illuminated obliquely by a pulsed laser. PA signal intensities were recorded for JAAZ, N-JAAZ, and RGD-JAAZ particles at 25, 12 and 5 µM equivalent free dye concentration in 0.5% (w/v) agarose. All PA signals were normalized with respect to 1:100 dilution of India Ink in water. The normalized PA signal magnitude of RGD-JAAZ particles was comparable to the PA magnitude of the non-functionalized particles, demonstrating that attaching a protein like streptavidin does not substantially affect the PA conversion efficiency of the J-aggregates (Fig. 5B). To ascertain whether the size of the JAAZ particles has any influence on the PA signal intensity, the PA signal of different sized JAAZ particles (930–250 nm) at the same absorbance intensity at 895 nm was recorded (Fig. S25). The PA signal intensities of the 363 ± 90 nm, 570 ± 52 nm and 930 ± 27 were not significantly different and in the same range. The intensity of the 256 ± 20 nm N-JAAZ particles was ~16% higher than the 930 ± 27 nm JAAZ particles which

![Photoacoustic properties of various JAAZ particles.](image-url)

Fig. 5. Photoacoustic properties of various JAAZ particles. (A) In vitro PAI set-up including (i) laser source (ii) transducer (iii) and reservoir containing deionized water to submerge the (iv) 96-well sample plate. (B) PA signal of the different JAAZ particles at different concentrations. The normalization is performed with respect to 1:100 (v/v) dilution of India Ink. Size measurements of all particles were taken at 25, 12 and 5 µM dye concentration to validate that there are no changes in size with concentration. (C) Normalized absorbance (dashed line) and PA spectra (continuous line) for different JAAZ particles. (D) PA signal intensity of free ICG dye, and various JAAZ particles measured in sheep’s blood. Samples concentration was set at 20 µM and all values were normalized with respect to whole blood PA signal. Individual data points are not shown for simplicity. Inset shows the PA spectra of RGD-JAAZ particles in whole blood. (E) 2D photoacoustic image of HeLa cells stained with 10 µM JAAZ and RGD-JAAZ particles showing photoacoustic signal normalized to maximum value. An area of the glass slide was scraped with a razor blade to remove cells before staining and the scraped region is indicated by an arrow in both figures. The entire scraped regions are highlighted in Fig. S29A-B. Representative scale of PA signal amplitude is also given in the figure. (n = 100 for B and D with data presented as mean±SD, p-values are calculated using one-way ANOVA; ****=p < 0.0001; all P values are provided in Table S6).
suggests a non-linear relationship between the particle size and PA signal intensity. To explain this observation, we hypothesize that the smallest N-JAAZ may absorb the laser light more efficiently because of their higher surface-area-to-volume ratio, leading to more thermoelastic expansion and stronger PA signal [42,43]. An increase in PA signal magnitude was observed with increasing equivalent free dye concentration (5, 12 and 25 µM, Fig. 5B) of the J-aggregates, which establishes the correlation between the total free dye concentration and PA signal intensity. The individual absorbance curves of all samples (JAAZ, N-JAAZ and RGD-JAAZ) at the above-mentioned concentrations are provided in Fig. S26. The normalized absorbance of JAAZ and N-JAAZ shows a peak at 895 nm (Fig. 5C, dashed lines) and the normalized PA profile matches the absorbance profile, with the PA signal peak at 890 nm (Fig. 5C, solid lines). The slight shift of the PA signal maxima from the absorbance maximum can be attributed to slight offset between the wavelength setting and output wavelength of the tunable pulsed laser. The photoacoustic spectrum of RGD-JAAZ has a PA signal maximum at 890 nm, which is in the NIR-I region (Fig. 5C), and optimally suited for deep-tissue imaging. We also tested the change in absorbance peak and size stability of different J-aggregates (JAAZ, N-JAAZ, RGD-JAAZ and RGD-N-JAAZ) under pulsed laser irradiation for up to 300 s (3000 shots). N-JAAZ, RGD-JAAZ and RGD-N-JAAZ exhibited neither a decrease in their absorbance peak at 895 nm nor a change in size (Fig. S27A-B). Micron-sized JAAZ particles demonstrated a 20% decrease in the absorbance peak after 200 s of irradiation and a 12% decrease in size, which was not significant (Fig. S27A-B), indicating that these particles are stable upon laser irradiation.

To explore the possibility of using JAAZ particles as a contrast agent in vivo, their photoacoustic signal amplitude was compared to that of whole blood. Short segments of 24-gauge PTFE tubing (containing approximately 100 µl of sample) were filled with 20 µM of ICG dye, JAAZ, N-JAAZ, RGD- JAAZ particles mixed in whole sheep’s blood. PA signals were recorded from each sample and the signal magnitude was normalized to the whole-blood PA signal magnitude (Fig. 5D). The PA amplitude of 20 µM RGD-JAAZ was ~2.1x that of blood excited at 895 nm, while 20 µM of free ICG dye was nearly ~0.03x the intensity of whole blood at this wavelength. The PA signal of RGD-JAAZ particles did not change while delivering pulsed laser light at ~58 mJ/cm² per 5 nanosecond pulse in over 200 s when subjected to 2000 laser pulses (10 Hz) at 895 nm, which indicates that RGD-JAAZ particles are photostable (Fig. S28A). Short segments of PTFE tubing were filled with 10 µM RGD-JAAZ and arranged in an ‘M’ pattern. The ‘M’ pattern was scanned to create a 2D PA image (Fig. S28B). As a demonstration of cell labeling using JAAZ, 2D cultures of HeLa cells were stained with 10 µM JAAZ and RGD-JAAZ particles on a custom glass slide with a red silicone insert. Before staining with JAAZ and RGD-JAAZ, a specific area of cells was scraped away using a razor blade as shown in Fig. S29A-B and the PA signal magnitude of the scraped region was used as a background to determine the level of non-specific binding of both particles. After washing the stained cells, a 35-MHz transducer was raster-scanned to produce a 2D PA image of the stained cells. The PA-color map indicates that the region where the cells were scraped away produced a weak PA signal that was less than 10% of the peak PA signal for cells stained with the particles (Fig. 5E). When comparing regions with similar density of seeded (100,000 cells/ml) cells on the slide, RGD-JAAZ labeling resulted in a more uniform labeling of the cells as well as significantly stronger PA signals when compared to cells labeled with JAAZ (Fig. 5E). Whereas the PA signal magnitude of the JAAZ stained cells decreased as the raster scan was completed (~3 h), for the RGD-JAAZ stained cell culture, there was minimal to no diminution of the PA signal magnitude as the raster scan was completed (~3 h). The direct maximum PA signal from each stained cell culture was compared for different laser intensities for RGD- JAAZ vs JAAZ with the signal from JAAZ stained cells being only half that of cells stained with RGD-JAAZ (Fig. S29C). Our results indicate that the RGD-JAAZ particles can efficiently target and label integrin expressing HeLa cells taking advantage of the well-known binding affinity between RGD and the integrin receptors [44], as opposed to JAAZ particles without RGD, which do not bind to or label HeLa cells efficiently.

2.6. Pilot in-vivo imaging of two mice using RGD functionalized J-aggregates

To further demonstrate the clinical translation potential of the JAAZ particles, an in vivo pilot study was conducted on the commercially available small animal PAI system TriTom™ (Fig. 6 A) [45-47]. We first acquired multiwavelength images of microcuvette tubes containing 0.4 mM of ICG dye or RGD-JAAZ particles to evaluate the PA spectra of both contrast agents. The peak PA signal in the RGD-JAAZ tube was measured at 895 nm while the ICG dye tube had peaks at 740 nm and 780 nm (Fig. S30). For in vivo imaging experiments, multispectral PAI were acquired in two female n/nu nude mice at various timepoints (0, 10, 90 min) following tail-vein administration of ICG dye and RGD-JAAZ particles (Fig. 6). A pre-injection baseline PA signal at 895 nm was also acquired for reference (Fig. 6B). We found that while RGD-JAAZ and ICG dye demonstrate a similar biodistribution in vivo, intravenous administration of RGD-JAAZ particles improved visualization of blood-rich tissues such as the liver and spleen for up to 90-minutes post-injection (Fig. 6C and Fig. S31A for PA images of the second mouse) compared to ICG. The improved visualization with RGD-JAAZ particles continued for up to 120-minutes post-injection as seen in Fig. S31B. The achievable imaging depth was also improved by RGD-JAAZ particles demonstrated by the increased contrast-to-noise ratio (CNR) in blood vessels as deep as 5 mm from the surface of the skin compared to ICG dye 90-minutes post-injection (Fig. 6D and Table 1). The circulation half-life of RGD-JAAZ particles was 10 times greater than ICG dye (Table 1). Additionally, the molecular maps of oxyhemoglobin, deoxyhemoglobin, and RGD-JAAZ particles show the biodistribution of the contrast agent is restricted to the vasculature and blood-rich tissues such as the liver and spleen (Fig. 6E). Molecular maps were generated from a linear spectral unmixing of the multiwavelength PA images using previously reported spectra for hemoglobin and the measured spectra of the RGD-JAAZ particles at their injection concentration [47,48].

RGD-modified ICG [9,49] or RGD-modified polymeric particles [50] or micelles [51] encapsulating ICG have been previously used to improve the visualization and targeting of ICG in vivo for both fluorescence and PAI. While comparing non-RGD-labelled JAAZ particles with RGD-JAAZ particles in vivo was beyond the scope of this pilot study, based on the studies referenced above, we can hypothesize that the enhanced circulation time and increased CNR might be attributed to the presence of the RGD peptide. This needs to be confirmed in future systematic studies comparing various JAAZ formulations.

3. Conclusion

In this study, we designed and validated a simple and robust protocol for the synthesis of functionalized NIR-PAI contrast agent with tunable sizes ranging from 1.2 µm to 230 nm, which allows photoacoustic imaging at an excitation wavelength of 895 nm. As an added advantage over previously reported ICG J-aggregates [29], the modified J-aggregates synthesized in our study do not require size selection or further downstream processing. The particles are also stable in physiological conditions and can be easily dried and resuspended for long-term storage.

We successfully demonstrated direct functionalization of the azide groups using efficient and reliable copper-free click chemistry, which allow for further conjugation with biomolecules and targeting moieties such as streptavidin and RGD onto the surface of JAAZ and N-JAAZ. Our functionalized particles exhibited a strong PA signal even at concentrations as low as 5 µM equivalent free dye and showed nearly 2.1x greater PA magnitude in comparison with whole blood. RGD-JAAZ
demonstrated the cell targeting capabilities of our platform with a significantly higher PA magnitude compared to JAAZ stained HeLa cells. The in vivo 3D PAI images showed that RGD-JAAZ particles can be visualized with a CNR of 2.42 in blood vessels as deep as 5 mm from the surface of the skin and generate strong and quantifiable PA signals in blood-rich tissues such as the liver and spleen for up to 90-minutes post-injection in comparison with ICG. At 90 min of injection, in the RGD-JAAZ unmixed images, high intensity signals of these particles are clearly visible in liver and spleen over hemoglobin and deoxyhemoglobin. Hence, these results demonstrate the potential use of RGD-

Fig. 6. In vivo imaging using RGD-JAAZ particles. (A) Schematic and photograph of the 3D PAI acquisition on the TriTom™ imaging system. (B) Coronal and axial slabs of the baseline PA signal acquired at 895 nm. (C) Representative coronal and axial slabs of the ICG (left) and RGD-JAAZ particle (right) injections acquired at the peak excitation wavelength (i.e., 800 and 895 nm, respectively) over time. The displayed dynamic range for all images was set based on the baseline scan acquired at the same wavelength. (Legend): a. superficial thoracic vessels, b. liver c. spleen d. intestines e. thoracic vertebrae f. CuSO₄ fiducial tube. (D) Axial slabs encompassing the iliac arteries (white ROI) used to calculate the achievable imaging depth for ICG and RGD-JAAZ particles 90-minutes post-injection. (E) Coronal slabs of oxyhemoglobin (red), deoxyhemoglobin (blue), and RGD-JAAZ particles (green) spectral unmixing overlaid on the 800 nm excitation scan of anatomy (white) at 0- and 90-minutes post-injection. Scale bars for all images are 10 mm. (n = 2 female nu/nu nude mice).
**Table 1**
Contrast-to-noise ratio and circulation half-life of ICG and RGD-JAAZ particles.

| Injection       | Contrast-to-noise ratio | Circulation half-life (mins) |
|----------------|-------------------------|------------------------------|
| ICG            | 1.59                    | 1.19 ± 0.45                  |
| RGD-JAAZ       | 2.42                    | 10.26 ± 1.24                 |

n = 2. Data is represented as mean ± SD.

JAAZ as a promising contrast agent platform for future applications such as tumor margin staining, [29] imaging diseased tissue [28] and lymph node biopsy. [52].

3.1. Experimental section

3.1.1. Materials

Indocyanine Green (ICG) and ICG-Azide were purchased from AAT Bioquest. Dibenzylooctyne-PEG-Biotin (DBCO-PEG₄-Biotin) was acquired from Jena Biosciences. The streptavidin and the Cyclo-Arg-Gly-Asp-D-Phe-Lys(Biotin) (RGD-Biotin) were purchased from Peptides International. The Amicon Ultra-0.5 centrifugal filters, potassium chloride (KCl), agarose, molecular biology grade water, penicillin, and Bicinchoninic Acid (BCA) Kits were acquired from Sigma-Aldrich. Triton X-100, Tween-20, Dimethylsulfoxide (DMSO) and glycerol were purchased from Fisher Scientific. Phosphate-Buffered Saline (PBS, 10X solution at pH 7.4) was acquired from ThermoScientific. Low melt Agarose was purchased from IBI Scientific. Molecular biology grade water was purchased from Quality Biological. The low-volume quartz glass cuvettes and the folded capillary zeta cells were purchased from Malvern Panalytical. 1X Dulbecco’s Modified Eagle’s Medium was purchased from Corning Inc. and Fetal Bovine Serum (FBS) were purchased from VWR Life Sciences. India Ink and whole sheep blood containing anti-coagulant were purchased from Hardy Diagnostics. Eagle’s Minimum Essential Media (EMEM) was purchased from ATCC. Cell Proliferation Kit I (MTT) was purchased from Millipore Sigma with active ingredient 3-(4,5-dimethylthiazol-2-yl-2,5-diphenyltetrazoliumbromide (MTT).

3.2. Methods

3.2.1. Synthesis of ICG-JA, JAAZ and N-JAAZ particles

ICG and ICG-azide stock solutions were respectively prepared at a concentration of 5 mg/ml (6 mM) in water and 10 mg/ml (12 mM) in DMSO and stored at −20 °C. To assemble the ICG-JA, ICG dye was prepared at 1 mM in water complemented with 20 mM of KCl and incubated at 60 °C for 20 h. For greater than 1 µm sized JAAZ particles, ICG and ICG stock solutions at 1:10 ratio was mixed to reach a final ICG concentration of 1 mM in water with 20 mM KCl. The mixture was incubated for 20 h at 60 °C. Micron-sized JAAZ particles could also be prepared by incubating 1:10 molar ratio of ICG-azide:ICG at a final dye concentration of 1 mM in water with 1 mM KCl. For JAAZ particles in the 600–700 nm range, the reaction solution of 1:10 ICG-azide:ICG molar ratio was incubated in 1 mM KCl for 20 h at the same temperature as mentioned above to a final concentration of 750 µM total dye. To make N-JAAZ particles of size ~200 nm, 250 µM total dye solution was incubated in 0.1 mM KCl for 14 h at 60 °C. A complete summary of the total dye concentration at 1:10 ICG-azide:ICG molar ratio, KCl molarity and time required for the formation of JAAZ particles is given in Fig. S19. Before further use, ICG-JA, JAAZ, and N-JAAZ samples were filtered using 0.5 ml Amicon centrifugal filters (100 kDa cutoff) to remove the non-aggregated free monomers of ICG and exchange the buffer. Filtration was carried out by centrifugation at 4000 g for 3 min at room temperature. Three cycles of filtration in water were used to ensure complete removal of unreacted dye. The percentage of samples recovered after filtration was calculated by:

\[
\% \text{recovery} = \left( \frac{A_{894(BF)} - A_{894(AF)}}{A_{894(BF)}} \right) \times 100
\]

Where, \(A_{894(BF)}\) is the absorbance intensity at 894 nm before filtration (BF) and \(A_{894(AF)}\) is the absorbance intensity after filtration.

3.2.2. Characterization of JAAZ particles

Samples were scanned from 440 nm to 990 nm with 2 nm step size using a PowerWave™ microplate spectrophotometer by Bio-Tek Instruments Inc. The amount of free dye in all J-aggregate samples was determined by incubating the formed J-aggregate particles in 1% Triton X-100 for 10 min at 37 °C and scanning the samples from 400 to 990 nm. A standard curve of a 1:10 molar solution of ICG-azole:ICG dye was created to determine the free dye concentration in the dissociated J-aggregate samples. Fluorescence intensity of different concentrations of free ICG dye and J-aggregates was recorded using a Tecan Safire® fluorescence microplate reader. Both free ICG dye and J-aggregate particles were excited at 720 nm with an emission at 770 nm. The fluorescence emission spectrum was recorded from 770 to 850 nm with a step size of 2 nm. The hydrodynamic diameter of the formed ICG-JA and JAAZ particles was measured using a Malvern Zetasizer Nano ZS instrument. The size measurements were taken using low-volume quartz glass cuvettes at 25 µM of equivalent free dye concentration. The zeta potential (surface charge) of all samples was determined using the Malvern Zetasizer Nano ZS using DTS-1070 folded capillary zeta cells. The measurements for zeta potential were taken at an equivalent free dye concentration of 15 µM and a volume of 700 µl in 0.1X PBS at pH 7.4. A 2% (w/v) gel was prepared using high strength agarose in 1X TAE buffer for electrophoresis. A square window of 6 × 6 cm was cut in the 2% gel. A 0.1% (w/v) low melting agarose gel prepared in 1X TAE buffer was poured in the square window and let to polymerize at 4 °C overnight. 30 µl of 500 µM JAAZ and N-JAAZ samples were mixed with an equal volume of glycerol and loaded into the wells. The gel was run at 120 V for approximately 10 min. For scanning electron microscopy, 2 µl of the JAAZ or the N-JAAZ samples were deposited on a clean mica surface and dried overnight under vacuum. SEM images were collected using a JEOL JSM-IT500HR InTouchScope™ at an accelerating voltage of 15 kV and a working distance of 10 nm. EDX spectra were collected using Octane Elect EDS System at an accelerating voltage of 15 kV and a resolution of 127 eV. ImageJ [53] was used to determine the size distribution of the particles in the obtained SEM images.

3.2.3. Functionalization of JAAZ particles

500 µM of JAAZ or N-JAAZ particles with 1:10 ICG-azole:ICG molar ratio were labelled with DBCO-PEG₄-Biotin at a concentration 10 times greater than the estimated number of azide groups available (Table S2). The samples were incubated and mixed in an incubator shaker overnight at room temperature protected from light. To remove excess DBCO-Biotin, the samples were filtered as mentioned above. These biotin functionalized samples (Bio-JAAZ) were diluted to an equivalent of 100 µM of free dye and incubated with streptavidin at 5X concentration of the available biotin sites (Strep-JAAZ). The samples were kept at 4 °C for two hours to ensure attachment of streptavidin to the available biotin sites. Excess streptavidin was removed by filtration. The Bio-JAAZ and the Strep-JAAZ samples were characterized using the procedures mentioned in the characterization section. The Strep-JAAZ samples were labelled with RGD-Biotin at 10X molar concentration and incubated for two hours at 4 °C. Excess RGD-Biotin was removed by filtration using the same procedure as mentioned before. The labelled samples were named RGD-JAAZ. The RGD-JAAZ samples were characterized using the procedures mentioned in the characterization section. Purified Strep-JAAZ samples were diluted to an equivalent of 5 µM of equivalent free dye concentration. BCA working reagent was made following the instructions provide by the supplier. 25 µl of 5 µM samples are incubated in 200 µl of BCA working solution and the amount of streptavidin attached is calculated using a BSA standard curve.
3.2.4. Hemotoxicity, Cytotoxicity and stability in whole blood of JAAZ particles

Whole sheep’s blood was centrifuged at 1000 g for 2 min at 4 °C to remove any coagulated blood. 80 µl of the centrifuged blood was mixed with 20 µl of samples by gently pipetting up and down and incubated at 37 °C for 30 min. 10 µl of treated blood was pipetted onto a clean glass slide and another glass slide was snapped across its entire length to create a blood smear. Microscopy images were acquired using Zeiss AxioVert 200 microscope. Hela cells were used for the cytotoxicity experiment of RGD-JAAZ, N-JAAZ, and JAAZ. The cells were seeded at 5 × 10^3 cells/well in a 96-well plate incubated in augmented EMEM at 37 °C and 5% CO2 for 24 h. Cells were then cultured with 10 µM of RGD-JAAZ, N-JAAZ, or JAAZ, 10 µl (final concentration of 0.5 mg/ml) of the standard 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazoliumbromide (MTT) was added to each well of the 96-well plate and incubated for 4 h. Then 100 µl of solubilization solution was added to each well and the plate was incubated overnight. The absorbance of each well was tested for a microplate (ELISA) reader (accurSkan FC FisherBrand, Waltham, MA, USA) measuring the absorbance of the formazan product between 550 and 600 nm. A positive control group was tested with an equivalent mixture of saline mixed with culture medium.

80 µl of whole blood was incubated with JAAZ, N-JAAZ and RGD-JAAZ samples respectively, at 37 °C. The final concentration of all samples in whole blood was 10 µM. At 0, 2, 4, 6, 8, 16 and 24 h, 60 µl of the incubated sample for each condition was scanned from 440 to 990 nm with 2 nm steps using a microplate spectrophotometer.

3.2.5. Characterisation of the photoacoustic properties of JAAZ particles in-vitro

The 96-well plate containing samples embedded in agarose was secured to the bottom of a plastic water reservoir. The reservoir was filled with deionized water, completely submerging the samples. For comparison of smaller volumes, 2-inch segments of 24-gauge PTFE tubing (~100 µl) were filled with various samples. The tubes were placed in a clear acrylic chamber submerging the samples in deionized water. A single element piezoelectric 35 MHz (PI35, Olympus, Massachusetts, USA) or 5 MHz (V326, Olympus, Massachusetts, USA) focused transducer was fixed at 12 mm or 45 mm above each sample with their functional element submerged in deionized water. Optimization of positioning and photoacoustic signal from each sample was done through movement of a fine x-y axis stage, moving above each sample with a fixed transducer and pulsed laser. Excitation of the sample was done obliquely by a wavelength-tunable (690–950 nm) pulsed laser designed for photoacoustic imaging (Phoicus Mobile, Opotek) at the peak absorption wavelength for JAAZ particles (895 nm) for magnitude comparison. The photoacoustic signal recorded from the transducer was sent to a 20/40 db amplifier (HVA-200 M, Femto) from there the signal is recorded through a lock-in amplifier (Zurich Instruments, Zurich, Switzerland) triggered on each laser pulse using a photodiode placed near the laser source. The sample was then raster scanned through movement of the stage at 100 µm steps (with fixed laser source and transducer) mounted on motorized translation stages (Thorlabs). Absolute signal magnitude was then plotted in 2D space to create the image. A simple Matlab image gaussian filter was then used to smooth the image.

3.2.6. In-vivo photoacoustic imaging using RGD-JAAZ particles

The phantom and in vivo photoacoustic imaging studies were performed using the TriTom™ imaging platform (PhotoSound Technologies, Inc.) and 20 Hz nanosecond-pulsed OPO laser (PhotoSonus, EKSPLA) located at MD Anderson Cancer Research Center House in Houston, TX. The TriTom is a commercially available multimodal imaging technology capable of high-resolution, 3D multispectral photoacoustic imaging in the whole body of small animal models[45]. To prepare for imaging, the TriTom™ imaging chamber was filled with degassed, deionized water at room temperature for phantom experiments or heated to 36.0 ± 0.5 °C for animal studies. In both studies, free ICG dye was diluted to 0.4 mM from a stock solution and the same concentration of RGD-JAAZ particles was prepared using the methods mentioned in the above sections. The size of the RGD-JAAZ particles was 1.24 ± 0.24 µm.

The sample phantom was constructed using 810 µm inner diameter PTFE microcuvette tubes filled with a 50 µl volume of water, PBS, free ICG dye, or RGD-JAAZ particles. The four sample tubes were secured in a phantom holder that was then mounted in the TriTom and lowered into the imaging chamber. A series of single-wavelength 3D PA scans were acquired using excitation wavelengths from 700 to 900 nm in 10 nm steps with an additional scan at 895 nm. During each scan, the sample phantom was rotated 360° while acquiring 760 ± 5 frames of PA data. The acquired PA data were reconstructed into 30×30×30 mm volumes with a voxel size of 0.1 mm using a filtered back-projection method [54]. All visualization and analysis of the reconstructed volumes was performed in 3D Slicer. To determine the PA spectra of the ICG and RGD-JAAZ samples measured with the TriTom™, a 5 mm vertical section in the center of each microcuvette tube was manually segmented and the maximum PA signal intensity was calculated for all wavelengths.

A pilot study consisting of two female nu/nu nude mice (Charles River Laboratories) was conducted on the TriTom to compare the RGD-JAAZ particles of size 1.24 ± 0.24 µm to ICG dye in-vivo. During each imaging session, the animal was anesthetized with isoflurane and a saline-filled catheter was inserted in the tail vein and secured. The animal was then transferred to a mouse holder and mounted in the TriTom imaging chamber so that the liver, spleen, and abdominal region of the mouse were visible. First, a baseline series of 3D PAI data was acquired with 700, 750, 800, 850, 875, 885, and 895 nm excitation wavelengths. A 100 µl bolus of 0.4 mM free ICG dye was then injected via the tail vein catheter followed by a 50 µl saline flush and the same set of PAI data was collected at 0, 10, 30, 60, and 90-minutes post-injection. After the final imaging time point, the PA scan acquired at 800 nm was reconstructed and compared to the pre-injection scan to confirm that the ICG dye had completely cleared circulation. A matching 100 µl bolus of 0.4 mM RGD-JAAZ particles and 50 µl saline flush was then administered and the 3D PAI imaging sequence was repeated with an additional time point at 120 min post-injection. Each 3D PAI scan was then reconstructed into a
30 × 30 × 30 mm volume with a voxel size of 0.1 mm using a filtered back-projection method. The displayed dynamic range was determined using the baseline scans and set for all 3D reconstructions and axial slices.

To estimate the circulation half-life of RGD-JAAZ particles compared to monomeric ICG, four superficial blood vessels in the thorax region were manually segmented for the 0, 10, 30, and 60-minute post-injection volumes. Half-life estimation was performed using the 800 nm and 895 nm excitation scans (i.e., peak absorbance wavelength) for ICG and RGD-JAAZ particles, respectively. The baseline PA signal in each vessel was subtracted and then averaged for each time point. An exponential function fit to the data in each animal was then used to calculate the circulation half-life of ICG or RGD-JAAZ [24].

The achievable imaging depth of RGD-JAAZ particles and monomeric ICG was assessed by comparing the CNR in the iliac arteries from the baseline and 90-minute post-injection scans. Three axial slices containing both blood vessels at varying depths from the skin were identified using the 3D reconstructed volumes. The vessels and background signal (i.e., the area surrounding the animal) were manually segmented and the shortest distance from the center of each vessel to the skin was measured. CNR was then calculated for each artery as:

\[
\text{CNR} = \frac{\mu_i - \mu_b}{\sqrt{\sigma_i^2 + \sigma_b^2}}
\]

Where \(\mu\) and \(\sigma\) are the averages and standard deviation of the vessel and background, respectively. The blood vessel was considered detectable at the measured imaging depth if the calculated CNR was \(\geq 2\). Finally, multispectral PAI (700, 750, 800, 850 and 895 nm) was performed to spectrally unmix the endogenous PA signal from oxyhemoglobin, deoxyhemoglobin from the RGD-JAAZ signal. The unmixed volumes were then overlaid on the reconstructed 800 nm scan for anatomical reference.

**Statistical Analysis**

All bar graphs are expressed as the mean ± standard deviation. Statistical analyses were performed using the unpaired, two-tailed Student’s t-test and One-Way Analysis of Variance (ANOVA) with a Tukey post-hoc test using the software GraphPad Prism 9.0.2. Data was deemed statistically significant if the p-value was < 0.05. Absolute PA signal amplitude is defined as the sum of the absolute minimum and maximum peak PA signal.

**Ethical Statement**

All animal studies were approved by the Institutional Animal Care and Use Committee at MD Anderson Cancer Center. IUCAC protocol number 00001779-RN01, “Evaluations of Tumor Models with Multispectral Optoacoustic Tomography.”

**Author contributions**

S.S. and G.G. contributed equally to the manuscript. S.S. synthesized all the J-aggregate samples, performed the characterization and the conjugation experiments, and analyzed the data. G.G., P.C., helped by S.S., and R.V. designed the photoacoustic experiments and analyzed the data. C.H.H., J.B. and L.S.C performed additional characterization experiments and data analysis. D.J.L. performed the phantom and in-vivo experiments along with the associated data analysis. J.L.M. performed the SEM imaging and assisted with data analysis. R.V. designed and supervised the study and interpreted the results. S.S., G.G., D.J.L., and R.V. wrote the manuscript. All authors commented on and edited the manuscript.

**Declaration of Competing Interest**

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Shrishti Singh has patent Provisional Patent pending to George Mason University. Parag V. Chitnis has patent Provisional patent pending to George Mason University. Remi Veneziano has patent Provisional patent pending to George Mason University.

**Data Availability**

Data will be made available on request.

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**Appendix A. Supporting information**

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.pacs.2022.100437.

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