Black Mesoporous Silicon as a Contrast Agent for LED-Based 3D Photoacoustic Tomography

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Abstract: Mesoporous silicon (PSi) nanoparticles have been widely studied in different biomedical imaging modalities due to their several beneficial material properties. However, they have not been found to be suitable for photoacoustic imaging due to their poor photothermal conversion performance. In the present study, biodegradable black mesoporous silicon (BPSi) nanoparticles with strong light absorbance were developed as superior image contrast agents for photoacoustic tomography (PAT), which was realized with a light-emitting diode (LED) instead of the commonly used laser. LED-based PAT offers the advantages of low cost, compactness, good mobility, and easy operation as compared to the traditional laser-based PAT modality. Nevertheless, the poor imaging sensitivity of the LED-PAT systems has been the main barrier to prevent their wide biomedical application. The LED light has low optical energy. The present study demonstrated that the imaging sensitivity of the LED-PAT system was significantly enhanced with the PEGylated BPSi (PEG–BPSi) nanoparticles. The PEG–BPSi nanoparticles were clearly detectable with a low concentration of 0.05 mg/mL in vitro and with an LED radiation energy of 5.2 μJ. The required concentration of the PEG–BPSi nanoparticles was 10 times lesser than that of the reference gold nanoparticles to reach the corresponding level of the imaging contrast. The ex vivo studies demonstrated that the submillimeter BPSi nanoparticle-based absorbers were distinguishable in chicken breast tissues. The strong contrast provided by the BPSi particles indicated that these particles can be utilized as novel contrast agents in PAT, especially in LED-based systems with low light intensity.

Keywords: silicon nanoparticles, porous materials, photothermal conversion, imaging contrast, sensitivity, photoacoustic imaging

Introduction

Photoacoustic tomography (PAT) has lately been investigated as one of the most promising photonic imaging and diagnostic modalities. Unlike other optical imaging modalities, PAT converts light into ultrasound via thermoelastic expansion, and the generated ultrasound signal is monitored for imaging. Thus, PAT integrates the unique optical contrast and the high spatial resolution of ultrasound. A wide range of nanomaterials such as organic dye nanoparticles, gold nanoparticles, carbon nanotubes, and copper sulfide nanoparticles have been extensively studied as potential contrast agents for PAT imaging due to their strong photothermal conversion effect. However, each of these nanomaterials has its own disadvantages. For instance, organic dyes usually encounter the issue of poor photostability. Inorganic gold and carbon nanoparticles have a very slow biodegradation rate. Copper sulfide increases the risk of heavy-metal accumulation in the body. In general, the nanomaterials used in vivo should be biodegradable and safe.

Mesoporous silicon (PSi) nanoparticles have a high surface area, large pore volume, and good biocompatibility. The degradation product of porous silicon is orthosilicic acid, which is nontoxic and is essential for optimal bone and collagen growth. They have been widely studied in different biomedical imaging systems like in fluorescence and ultrasound imaging. Even though there are some recent studies on the development of the photoacoustic imaging contrast agents based on PSi, PEG–BPSi nanoparticles have only been utilized as carriers for different contrast agents such as indocyanine green and carbon nanomaterials. In these studies, PSi itself did not play any functional role in the imaging due to its poor light absorption performance. To the best of our knowledge, there are no studies published on the PAT contrast agents solely based on PSi nanoparticles. In our previous study, a new type of black mesoporous silicon (BPSi) nanoparticles was developed via the chemical oxidation method. The BPSi nanoparticles possessed better photothermal performance than gold nanoparticles, carbon nanotubes, and Fe3O4, which have widely been studied as contrast agents in photoacoustic imaging. Therefore, due to their outstanding photothermal performance, we hypothesize that the BPSi nanoparticles can serve as novel contrast agents to enhance the imaging sensitivity of PAT.

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Until now, PAT systems have been commonly based on the light source of class-IV lasers such as Nd:YAG lasers. Although these lasers can meet the demands of PAT, they are bulky, expensive, poorly movable, and suffer from energy fluctuations. LEDs have the advantages of compactness, low cost, good mobility, and stable light sources. Moreover, LED light can be easily operated in a typical indoor environment, while special safety restrictions and requirements (e.g., light shielding and water cooling) are necessary for the operation of the laser-based PAT. Nevertheless, due to the much weaker optical energy of the LEDs (μJ) than traditional lasers (μJ), the poor imaging sensitivity is the main issue in the current LED-PAT system. The development of exogenous image contrast agents is an effective way to overcome such technical barriers. Up till now, a few contrast agents such as free organic dyes have been tested to enhance the imaging sensitivity of LED-PAT. The dye molecules were dissolved in a suitable solvent and then used as the contrast agent directly. However, even though different nanoparticles have been studied as the contrast agents in laser-based PAT, the application of nanoparticles as the contrast agent of LED-PAT has been rarely investigated. The aim of the present study is to develop novel contrast agents for PAT based on BPSi nanoparticles. These particles could provide a new way to overcome the challenge in the imaging sensitivity of LED-PAT. Tomographic imaging was performed with the prototype of the LED-PAT system.

RESULTS AND DISCUSSION

The Raman scattering peak of silicon from the BPSi sample was observed at 515.6 cm\(^{-1}\). Compared with the Raman peak of 521.5 cm\(^{-1}\) for the silicon wafer, there was a clear blue shift attributed to the nanocrystalline structure of silicon in BPSi (Figure S1, Supporting information). The average size of the Si clusters was about 4.2 nm. Two poly(ethylene glycol) (PEG) molecules with different lengths of the carbon chains (molecular weights 0.5 and 2.0 kDa) were conjugated on the surface of BPSi (PEG–BPSi). According to our previous study, the dual-PEGylation with two PEGs was more efficient to enhance the colloidal stability of the nanoparticles for biomedical applications than the common mono-PEGylation. The colloidal stability of the nanoparticles before and after the PEGylation was studied in PBS at 37 °C. The BPSi nanoparticles without PEGylation aggregated quickly, and the particle diameter increased to 1.0 and 4.0 μm after 1 h and 24 incubation, respectively (Figure S2, Supporting information).

In contrast, the PEG–BPSi nanoparticles had excellent colloidal stability within 24 h. The hydrophilic PEG coating inhibits effectively the aggregation of the nanoparticles and enhances the colloidal stability. The functional groups on the BPSi nanoparticles were further analyzed with Fourier transform infrared (FT-IR) spectroscopy. The absorbance peak at 1730 cm\(^{-1}\), attributed to the –C=O stretching in the 2.0 kDa PEG-silane, was observed after the dual-PEGylation (Figure S3, Supporting information). In addition, intensive peaks of C–H stretching in the range of 2800–3000 cm\(^{-1}\) also appeared in the spectrum of PEG–BPSi. Based on the thermogravimetric analysis (TGA) measurements, about 10.8% of PEGs were grafted on the surface of BPSi nanoparticles. The γ-potential of the nanoparticles was changed from −24 to +2.0 mV (Figure S5, Supporting information). The surface charge close to neutral is beneficial to biological stability for a long systemic circulation. All of these evidences indicated that the PEGs were successfully grafted on the surface of BPSi nanoparticles. The biocompatibility of the nanoparticles was evaluated with the noncancerous cells of mouse embryonic fibroblasts and macrophages RAW 264.7 in vitro (Figure S6, Supporting Information). Generally, the BPSi nanoparticles did not show a cytotoxicity effect on either the embryonic fibroblasts or macrophages after 24 h of exposure if the nanoparticle concentration was below 0.2 mg/mL. The BPSi nanoparticles showed moderate cytotoxicity with both types of cells when the concentrations were higher than 0.5 mg/mL. However, the biocompatibility of the BPSi nanoparticles was significantly enhanced because of the surface PEGylation. The PEG–BPSi nanoparticles did not exhibit cytotoxicity toward embryonic fibroblasts and macrophages even at a concentration of 0.5 mg/mL, indicating that the nanoparticles had better compatibility as compared with other inorganic imaging contrast agents like gold and carbon nanotubes.

In the present study, the Au nanoparticles were used as the reference material because (1) they have been widely studied as contrast agents in PAT, (2) the BPSi nanoparticles have a particle diameter similar to that of Au nanoparticles, and (3) the light-absorption peaks of both BPSi and Au nanoparticles are close to the wavelength of LED (617 nm) in the photoacoustic imaging system. The mean hydrodynamic diameters of the PEG–BPSi and PEG–Au nanoparticles were 174 and 165 nm in deionized H\(_2\)O, which were consistent with the results observed with transmission electron microscopy.

**Figure 1.** (a) UV–vis spectra of PEG–BPSi and PEG–Au nanoparticles at a concentration of 0.1 mg/mL. The light source in the LED-PAT system has a wavelength of 617 nm. (b) Heating curves of PEG–BPSi and PEG–Au nanoparticles with different concentrations under the illumination of LED (mean ± standard deviation (SD), n = 3). The concentrations of PEG–BPSi were 1.0, 0.5, 0.2, 0.1, and 0.05 mg/mL. The concentration of PEG–Au was 0.5 mg/mL.
microscopy (TEM) (Figure S7, Supporting Information). The PEG−BPSi nanoparticles presented an irregular shape, while the shape of the PEG−Au nanoparticles was spherical. Figure 1a shows UV−vis absorption spectra of PEG−BPSi and PEG−Au nanoparticles in an aqueous solution with a concentration of 0.1 mg/mL. The BPSi and Au nanoparticles have light-absorbance peaks at 550 and 629 nm, respectively. In the present study, the light source in the PAT system has a wavelength of 617 nm, which is close to the peak absorbance of BPSi and Au nanoparticles. The mass extinction coefficients of PEG−BPSi and PEG−Au at 617 nm were 24.3 and 5.6 L g⁻¹ cm⁻¹ based on the Lambert−Beer law, respectively. The BPSi nanoparticles have a low level of Na impurity (0.6%), lowering the bandgap down to 1.34 eV, which corresponds to the wavelength of 925 nm. This improves the light absorbance of BPSi at the wavelengths used in the present study. The strong light absorption of PEG−BPSi indicates that the nanoparticles have great potential to be applied as imaging contrasts in PAT where the produced image corresponds to the reconstructed initial pressure $p_0$ caused by the photoacoustic effect. This pressure is directly dependent on light absorption through the relation described by

$$p_0 = \Gamma \mu_a \Phi(\mu_a, \mu_s)$$

where $\mu_a$ is the absorption coefficient, and $\Phi(\mu_a, \mu_s)$ is the light fluence that depends on the intensity of the light source and absorption and scattering $\mu_s$ parameters, and can be solved using a light transport model such as the radiative transfer equation. Further, $\Gamma$ is the Grüneisen parameter that is used to identify photoacoustic efficiency.

To test and compare the heating tendency of nanoparticles, the temperature rise after 5 min of LED illumination was measured using a thermocouple immersed in the sample dispersion. The temperature increased as a function of the concentration (Figure 1b), and the highest temperature rise was measured with the concentration of 0.5 mg/mL (19.2 ± 0.4 °C), followed by the concentration of 1 mg/mL (19.0 ± 0.2 °C). The increased values of temperature are basically identical, indicating heating saturation above the concentration of 0.5 mg/mL due to the long optical path inside the sample.
measurement vessel, resulting in nearly complete light absorption. With 0.05 mg/ml PEG−BPSi, the temperature rise (9.5 ± 0.1 °C) was comparable to that generated by a 10 times higher concentration (0.5 mg/mL) of PEG−Au particles (9.6 ± 0.3 °C). In a sample of blank water, the temperature elevated by less than 1 °C. Based on the temperature measurements, the PEG−BPSi particles generated roughly twice as high temperatures compared to the PEG−Au particles with the same concentration. When the LED power was increased from 1 to 2 W and the 0.05 mg/mL sample was exposed, the temperature rise was doubled to 19 ± 0 °C (Figure S8, Supporting information). Accordingly, the LED-induced temperature rise of the PEG−BPSi particles with different concentrations can be controlled by adjusting the LED power. Furthermore, it can be expected that the PEG−BPSi particles can serve as good contrast agents for PAT where the initial acoustic pressure is generated by thermoelastic expansion.

To compare the photoacoustic signals from the PEG−BPSi and PEG−Au particles, single capillary tubes were filled with nanoparticle dispersions of different concentrations and measured with the LED-PAT configuration developed earlier (Figure 2a).24 The photoacoustic signal amplitude was calculated through a frequency spectrum analysis and averaged over the measurement angles (Figures S9–S11, Supporting information). The signal level from the PEG−BPSi particles was observed to follow a linear trend starting from 0.05 mg/mL concentration (Figure 2b). The lowest concentration of 0.05 mg/mL was detectable, and the signal was above the levels of the background. The signal from the PEG−Au particles was barely above the background values but showed a slightly positive slope as a function of the concentration.

To detect and image nanoparticles through tomographic reconstructions, three identical capillary tubes were filled with different nanoparticle concentrations, measured using the PAT setup, and reconstructed applying the Bayesian approach (Figure 3).24,35 The reconstructed volumetric three-dimen-

Figure 4. Reconstructions of three tubes having different nanoparticles or concentrations: (a) Illustration of three sample tubes illuminated with LED light. (b–d) Comparison of PEG−BPSi and PEG−Au with different concentrations. The reconstructed images visualize the maximum-intensity projections (MIPs) in the z-direction (xy-plane) and x-direction (yz-plane) and the volumetric images, respectively.

Figure 5. Reconstructions of three tubes placed inside a 4–6 mm-thick chicken tissue and having different PEG−BPSi concentrations: (a) 0.05 mg/mL, (b) 0.1 mg/mL, (c) 0.2 mg/mL, (d) 0.5 mg/mL, and (e) 1 mg/mL. The figures visualize the maximum-intensity projections (MIP) in the z-direction (xy-plane) and x-direction (yz-plane), and volumetric images. Dimensions of chicken tissue samples were 1 cm (x-direction), 3 cm (y-direction), and 5 cm (z-direction), that is, the reconstructions visualize the area and volume inside a tissue.
sional (3D) images and the maximum-intensity projections in the z-direction (xy-plane) are shown in Figure 3a,b, respectively. Based on these experiments, the reconstructed maximum amplitude at the highest concentration (1 mg/mL) of PEG–Au was approximately at the same level as compared to the amplitude from the smallest concentration (0.05 mg/mL) of PEG–BPSi. More specifically, the contrast value for the reconstructions was calculated from Figure 3a. The tube contrast using PEG–BPSi particles at 0.1 mg/mL was 0.84 ± 0.12 and exceeded the contrast of PEG–Au particles at 10 times higher concentrations (1 mg/mL: 0.60 ± 0.15) (Table S1, Supporting information).

Next, the tubes were filled with dispersions having different nanoparticle concentrations and different particles (Figure 4). In the first experiment, the tubes contained both PEG–Au (0.2 mg/mL) and PEG–BPSi (0.1 and 0.2 mg/mL) nanoparticles (Figure 4a). Both low and high PEG–BPSi concentrations could be detected from the background and discriminated from each other. The tube containing PEG–Au particles was not detectable. In the second experiment, the lowest concentration of PEG–BPSi particles (0.05 mg/mL) was placed between 0.1 and 0.2 mg/mL of PEG–BPSi (Figure 4b). The signal from the 0.05 mg/mL tube was small but detectable, while the signals from the two higher PEG–BPSi concentrations were similar to the measurement in Figure 4b. In the third experiment, 0.1 mg/mL PEG–BPSi concentration was placed between the tubes having the highest PEG–BPSi concentrations (0.5 and 1 mg/mL) (Figure 4c). The 0.1 mg/mL tube was detectable between the high contrast tubes and had an amplitude similar to that in Figure 4a,b, indicating systematic detection and a repeatable signal level.

To test the imaging contrast for LED-PAT ex vivo, tubes filled with different nanoparticle dispersions were imaged inside chicken tissue (Figure 5). The capillary tubes were placed at the center of 1 cm-thick (x-direction) tissue blocks so that the tubes were approximately 4–6 mm deep in the tissue. The height (z-direction) and width (y-direction) of these blocks were 5 and 3 cm, respectively. The qualitative estimate for a minimum detectable concentration, so that all three tubes were clearly distinguishable in all three repeats, was 0.2 mg/mL. The tubes with 0.1 mg/mL nanoparticles were still detectable in most repeated experiments (Figures 5 and S12 and S13, Supporting information). In addition to heterogeneities of chicken tissue pieces, the spacing between the tubes and the tube orientation varied as the flexible tubes were misaligned when they were pushed through the tissue block, causing variations of the received acoustic signal in the LED-PAT experiment.

**CONCLUSIONS**

The feasibility of black porous silicon nanoparticles as a contrast agent for 3D PAT was shown. They were observed to provide excellent photoacoustic contrast that was significantly stronger than that of the reference Au nanoparticles. More precisely, the PEG–BPSi particles were found to provide a higher contrast at 0.1 mg/mL than the PEG–Au particles at 1 mg/mL in the LED-PAT system. Furthermore, submillimeter absorbers with a concentration of 0.2 mg/mL PEG–BPSi could be clearly observed inside approximately 5 mm-thick chicken tissue ex vivo. This indicates that the BPSi nanoparticles can be utilized as efficient contrast agents for PAT, where they could be used, for example, to increase the sensitivity of the technique for imaging certain targeted tissue types and to increase the imaging depth. This can be especially useful in LED-based systems with low light intensity. Overall, versatile LED illumination combined with the highly efficient contrast agent of BPSi nanoparticles can offer an alternative for large solid-state lasers for certain applications of biomedical imaging: for example, the diagnosis of skin cancer, subcutaneous tumors such as breast cancer, and ocular choroidal melanoma.

**EXPERIMENTAL SECTION**

The BPSi nanoparticles were prepared via the reaction between NaSi and NH₄Br at 250 °C. Subsequently, the BPSi nanoparticles were functionalized with 0.5 and 2.0 kDa PEG-silanes to improve biocompatibility. More experimental details are provided in the Supporting Information.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.9b18844.

Experimental materials and methods; results of the Raman spectrum; colloidal stability, FT-IR spectra, TGA curves, ζ-potential, particle size distribution, TEM, biocompatibility, and temperature rise graphs; the repeated photoacoustic tomography measurements of chicken tissue samples; the sinograms for nanoparticles, water, and the reference measurement using 0.5% ink solution; frequency spectra as a function of the measurement angle calculated using the waveforms; and image contrast values (PDF)

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Notes

The authors declare no competing financial interest.

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