Clear cell renal cell carcinoma detection by multimodal photoacoustic tomography

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ABSTRACT

There is a need for accurate and rapid detection of renal cancer in clinic. Here, we integrated photoacoustic tomography (PAT) with ultrasound imaging in a single system, which achieved tissue imaging depth about 3 mm and imaging speed about 3.5 cm/s/min. We used the wavelength at 1197 nm to map lipid distribution in normal renal tissues and clear cell renal cell carcinoma (ccRCC) tissues collected from 19 patients undergone nephrectomy. Our results indicated that the photoacoustic signal from lipids was significantly higher in ccRCC tissues than that in normal tissues. Moreover, based on the quantification of lipid area ratio, we were able to differentiate normal and ccRCC with 100 % sensitivity, 80 % specificity, and area under receiver operating characteristic curve of 0.95. Our findings demonstrate that multimodal PAT can differentiate normal and ccRCC by integrating the morphologic information from ultrasound and lipid amount information from vibrational PAT.

1. Introduction

Among the current methods for renal cancer treatment, surgical resection (radical nephrectomy or partial nephrectomy) is still the most effective way [1, 2]. Partial nephrectomy can maximize the preservation of kidney function while ensuring the therapeutic effect. However, there is an increased risk of local cancer recurrence and a lower survival rate for having cancer residues left inside the kidney during partial nephrectomy [3, 4]. The gold standard of tumor assessment for partial nephrectomy is postoperative histopathology examination. However, its turnaround time is 3–5 days.

Currently, there have been several existing or emerging techniques for intraoperative renal cancer diagnosis. Frozen section analysis has been applied in clinic, and it usually takes 20–30 min. However, the accuracy of frozen section analysis is argued because of the limited sampling of the excised tumor tissue [5–7]. Intraoperative ultrasound imaging has been used for the detection of renal masses, but it lacks chemical selectivity to identify the pathological lesions [8–10]. Structured light illumination microscopy was shown to image the edge of the renal parenchyma at a high speed, but it required the use of fluorescent contrast agent and had very limited imaging depth [11]. Raman spectroscopy and fluorescence diffuse reflectance spectroscopy were used to classify tumor and normal tissues, but could only focus on a few spots of interest [12, 13]. More recently, some new techniques have been developed for intraoperative histological diagnosis, such as stimulated Raman scattering microscopy [14], microtomy-assisted photoacoustic microscopy [15], spatial light interference microscopy [16], optical coherence microscopy [17], microscopy with ultraviolet surface excitation [18], but these technologies are limited by either slow imaging speed or small field-of-view. Taken together, these current techniques for renal cancer detection are either limited to low sensitivity (<90 %) or long procedure time to image the entire resected tissue surface with deep tissue penetration. Considering that nephrectomy requires temporary artery block-off no longer than 30 min., a label-free imaging technique for renal cancer diagnosis with high sensitivity and fast speed is desirable.

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Recently, photoacoustic tomography (PAT) has been emerging as an unique noninvasive biomedical imaging tool with strong molecular selectivity and deep tissue penetration [19–22]. In particular, melanoma can be detected based on photoacoustic signals from melanin [23, 24]; prostate cancer [25], ovarian cancer [26], colorectal cancer [27], skin cancer [28] and breast cancer [29, 30], can be detected based on photoacoustic signals from blood. As for vibrational PAT, a series of previous work has demonstrated that neutral lipids in biological tissues produce a strong photoacoustic signal at around 1200 nm due to the 2nd overtone absorption of CH stretching vibrations [31–34]. Specifically, the photoacoustic signal from adipose tissues is about 7 times higher than that from blood at around 1200 nm [31]. Based on vibrational PAT of neutral lipid accumulation, breast cancer can be identified, because normal breast tissues contain significantly higher amount of neutral lipids than breast cancer tissues [35, 36]. Interestingly, an early study has reported that clear cell renal cell carcinoma (ccRCC), which is the most common and aggressive form of renal cancer [37], contains significantly more neutral lipids in the form of cholesteryl ester than normal renal tissues [38]. Thus, it is promising of using PAT for ccRCC detection based on lipid accumulation.

Herein, we developed a multimodal PAT system that could provide both ultrasound images and 1197 nm wavelength PAT images. PAT images at 1197 nm specifically showed lipid accumulation in 32 intact human tissues collected from 19 patients undergone nephrectomy. Due to the big difference in lipid accumulation between normal and ccRCC tissues, we have achieved a 100% sensitivity and 80% specificity for ccRCC detection. This result demonstrated the capability of multimodal PAT to distinguish ccRCC tissues from normal tissues.

2. Materials and methods

2.1. Multimodal PAT system

Our multimodal PAT system was developed based on a customized ultrasound system with 128-channel data acquisition board and a customized all-solid-state Raman laser, which generated 10 Hz pulses with wavelengths at 1197 nm (Fig. 1a). The Raman laser was built based on stimulated Raman scattering process in a gain medium. The output wavelength of Raman laser was determined by the pump wavelength and Raman shift of the medium. With 1064 nm laser light as pump, a Ba(NO3)2 crystal was used to produce a laser output at 1197 nm. The technical details about the Raman laser can be found in Ref [36]. The performance of the Raman laser, including laser pulse duration, pulse energy stability and laser output power at 1197 nm, was shown in Fig. A.1. The 128-element ultrasound transducer array sends and receives acoustic wave at 18.5 MHz center frequency and a bandwidth of 14 MHz–22 MHz (L22–14vX, Verasonics Inc.). The laser and ultrasound system were connected to a computer for synchronizing time trigger. The excitation laser beam was focused by a convex lens and then coupled into a multimode fiber bundle, which transmitted the light to the sample. As shown in Fig. 1b, Two 45-degree transparent glass slides were placed inside the probe holder to reflect the acoustic waves, making the light and the acoustic wave collinear for better adaption to the unevenness of the tissue surface. The imaging probe was mounted on a 2D stage for X-Z scanning. The ultrasonic and PAT signals were detected by a 128-element ultrasound transducer array, and then were transmitted to the ultrasound system for post-processing and image reconstruction. To characterize the imaging depth, polyethylene tubes with 1.27 mm outer diameter were imaged in a 2.5% agarose gel phantom. The SNR of polyethylene tubes was calculated from 20 log10 ratio of the mean of PA signal to the standard deviation of the background noise.

2.2. Tissue imaging and processing

This study was approved by an institutional review board. Frozen specimens of human renal tissues were obtained from the tissue bank of Department of Urology in Peking University First Hospital. Before imaging, frozen tissues were first fixed with 4% paraformaldehyde, and then embedded in 2.5% agarose gel. During imaging, embedded tissues were placed in a bath containing phosphate buffered saline solution, and the front end of the probe was immersed into the solution. The 1197 nm laser with energy density of 82 mJ/cm2 were utilized for PAT signal excitation, which were below the American National Standards Institute (ANSI) safety standard (100 mJ/cm2 for nanosecond laser at 1197 nm) [32].

As shown in Fig. 1b, a series of two-dimensional ultrasound and PAT images in X–Y direction were acquired for each tissue by moving the
probe at a step size of 150 μm. Then, three-dimensional ultrasound and PAT images were reconstructed by ImageJ (NIH, USA). A single image section from the tissue surface in the X–Z direction was selected for comparison with the histopathology image of the adjacent tissue slice.

2.3. Histopathology

Tissues were peeled from the agarose gel after imaging and dehydrated with a gradient of 30 % sucrose solution overnight at 4 degrees. Then, the tissues were sectioned along the X–Z direction and stained with hematoxylin and eosin (H&E). Finally, two professional pathologists examined the H&E slides independently. According to Fuhrman grading system, renal cancers are given a grade from 1 to 4. Grade 1 and 2 are considered low-grade, and Grade 3 and 4 are considered high-grade.

2.4. Data quantification and statistical analysis

The lipid amount was represented by area ratios between the corresponding areas covered by PAT signals and ultrasound signals. First, we defined the threshold to pick valid signal. The threshold is three times of the noise level, which is estimated as the standard deviation of the background. Then, we calculated the number of pixels from signal to represent the signal area. The calculation of the signal area was done by using ImageJ. To compare the lipid amount between different groups, two-sample, Welch’s t-test was used. p < 0.05 was considered statistically significant. The exact p values are shown in the corresponding figure captions. To illustrate the ability of using the lipid area ratio to differentiate normal and ccRCC, a receiver operating characteristic (ROC) curve was generated by plotting sensitivity versus (1-specificity) and the area under curve (AUC) was calculated.

3. Results

3.1. Characterization of multimodal PAT system

Our multimodal PAT system, which integrated ultrasound and PAT, was established based on a customized ultrasound system with 128-channel data acquisition and a customized all-solid-state Raman laser. The imaging depth of the system was first measured. The polyethylene tube, rich in C–H bonds, was used to characterize PAT imaging depth at 1197 nm excitation. As shown in Fig. 2a–b, both of the upper and lower walls of the polyethylene tubes at different depths can be clearly visualized by ultrasound and PAT. The PAT imaging depth could reach up to 1 cm. The abundant absorption of laser light by the upper wall of the polyethylene tube resulted in the loss of laser energy, and so less laser energy transmitted to the lower wall of the tube. This further led to weaker signals from the lower wall of the tube and showed a thinner tube wall at the bottom. Signal to noise ratio of the PAT signals were then measured at different depths. As shown in Fig. 2c, the signal-to-noise ratio was around 38 dB at 2 mm depth and gradually decreased to 28 dB around 9 mm depth, presumably because the laser power reduced with depth. The spatial resolution was further characterized by imaging a 50 μm diameter tungsten wire placed at ~1 mm beneath the ultrasound transducer. Based on the PAT image of the tungsten wire (Fig. 2d), the lateral and axial resolution was 372 μm and 230 μm, respectively, as shown in Fig. 2e–f.

Before multimodal PAT imaging of human renal tissues, we used a beef tissue to test the imaging capabilities. As shown in Fig. A.2, by using laser excitation at 1197 nm, we were able to detect strong signals from lipid deposition with ~3 mm depth and reconstruct 3D multimodal PAT image, which was consistent with the photograph of the beef tissue. Moreover, with 12.8 mm * 40 mm field of view and 450 μm/second scanning speed along the z-direction at 150 μm translational step size, the system could assess ~3.5 cm² tissue area per minute.

3.2. Multimodal PAT of normal and ccRCC tissues

In order to demonstrate the capability of multimodal PAT system for
ccRCC detection, we compared the multimodal PAT images with the histological images at the same location. In total, 32 human renal tissues (15 normal, 12 low-grade ccRCC, 4 high-grade ccRCC and 1 sample containing both cancer and normal tissue) from 19 patients were imaged. The patients information was listed in Table A.1. The PAT image at 1197 nm showed the distribution of lipids. Diagnosis was confirmed by histopathology. As shown in Fig. 3, we found abundant lipid signals in the ccRCC tissue, but very little or no detectable lipid signals in the normal tissue. This finding was consistent throughout most of the patient tissues. An enlarged histopathological image of the ccRCC is shown in Fig. A.3. More representative images are shown in Fig. A.4. Moreover, we assessed the tissue from patient #19, which contained both normal and ccRCC tissue, and found that the lipid signal in the ccRCC area was higher than that in the normal area, consistent with previous results. This demonstrated that multimodal PAT system was able to differentiate normal and ccRCC on the same tissue sample (Fig. A.5).

3.3. Differentiation between normal and ccRCC by multimodal PAT

The lipid amount was quantified by the area ratios between the corresponding areas covered by PAT signals and those covered by ultrasound signals. The lipid area ratio was found to be significantly higher (~9 times) in ccRCC tissues than that in normal tissues (Fig. 4a). Although the lipid area ratio was higher in high-grade ccRCC compared to low-grade ccRCC, there was no statistical difference (Fig. A.6). Detailed information about lipid area ratios are listed in Table 1 and Table A.2.

To test the ability of using the lipid area ratio to differentiate ccRCC from normal tissues, the ROC curve was generated by plotting sensitivity versus (1-specificity), as shown in Fig. 4b. The large area under curve (AUC = 0.95) demonstrated that the lipid area ratio can accurately differentiate between normal and ccRCC tissues. Moreover, the ROC curve provided a way to obtain the desired degree of sensitivity at the cost of specificity. Because detection sensitivity is more important for ccRCC detection, the threshold of lipid area ratio was chosen to be 7.62 % to achieve 100 % sensitivity and 80 % specificity.

4. Discussion

There is a need for accurate and rapid detection of renal cancer in clinic. In this work, we developed a customized multimodal PAT system at 1197 nm to detect lipid distribution respectively, at an imaging speed of 3.5 cm² tissue area per minute. Our data from 32 intact human tissues demonstrated that the PAT signal of lipids was significantly higher in ccRCC tissues than that in normal tissues. By collecting the morphologic information from ultrasound imaging and lipid area ratio information from vibrational PAT, multimodal PAT can identify ccRCC from normal renal tissues with 100 % sensitivity, 80 % specificity, and the AUC of 0.95. As discussed below, these results demonstrated the potential of using multimodal PAT system for intraoperative ccRCC detection.

First, lipid accumulation can serve as a marker for ccRCC detection. Our data has shown significantly stronger vibrational photoacoustic signals from neutral lipids in ccRCC compared to the normal counterparts, although there is indeed a variation of lipid signal level among different patients. As for the ccRCC tissues with small amount of lipid accumulation, photoacoustic microscopy could help accurate detection owing to high spatial resolution [39-43].

Second, high sensitivity makes our multimodal PAT system a promising way for sensitive ccRCC detection. Tumor residue in partial nephrectomy increases the risk of local cancer recurrence and results in a lower survival rate [3,4]. Therefore, it is essential to get high sensitivity of renal cancer detection. In this study, based on the morphological information provided by ultrasound imaging and the lipid amount information provided by PAT imaging, we were able to differentiate normal and ccRCC tissues with 100 % sensitivity and 80 % specificity. Such high sensitivity could help surgeons to remove ccRCC, the most common and aggressive form of renal cancer, as much as possible and leave a healthy kidney.

Third, high speed of our multimodal PAT system fulfills the requirement for intraoperative cancer detection. Since renal tumor masses smaller than 4 cm in diameter can be usually treated by partial nephrectomy, the maximal surface area of the resected renal tumor tissue is about 50 cm². With an imaging speed at 3.5 cm²/min of our system, the whole surface area of the resected tissue can be assessed within 15 min. Such high imaging speed meets the current clinical need.
for rapid intraoperative renal cancer detection.

Finally, we utilized a customized Raman laser as the excitation light source, which showed greater efficiency to generate 1197 nm laser light output for photoacoustic imaging of lipids, compared with traditional optical parametric oscillator. Due to its excellent performance, the Raman laser-based photoacoustic imaging has been reported in many previous works [36, 46, 47]. Owing to the compact size and cost-effective feature, the Raman laser shows great promise for clinical translation.

5. Conclusions

In this study, we developed a multimodal PAT system with 1197 nm excitation to map both lipid distribution in intact human renal tissues collected from nephrectomy. Our data indicated that the PAT signal from lipids was significantly higher in ccRCC tissues than that in normal tissues. Based on the lipid amount information provided by ultrasound imaging, we were able to differentiate normal and ccRCC tissues with 100% sensitivity and 80% specificity. Our results show promise of using multimodal PAT system for intraoperative ccRCC detection.

Table 1
Lipid area ratios in normal and ccRCC tissues.

| Tissue Type       | Number of Tissues | Lipid Area Ratio (%) (mean ± SEM) |
|-------------------|-------------------|-----------------------------------|
| Normal            | 15                | 7.15 ± 3.33                       |
| ccRCC*            | 16                | 47.40 ± 6.38                      |
| Low-grade ccRCC   | 12                | 44.22 ± 6.60                      |
| High-grade ccRCC  | 4                 | 56.93 ± 17.19                     |

* N represents the number of tissues; 
ccRCC: clear cell renal cell carcinoma; 
Standard error of the mean. 

The results of each sample keep two decimals.

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