Photoacoustic imaging depth comparison at 532-, 800-, and 1064-nm wavelengths: Monte Carlo simulation and experimental validation

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Abstract. Photoacoustic imaging (PAI) provides high-resolution and high-optical-contrast imaging beyond optical diffusion limit. Further improvement in imaging depth has been achieved by using near-infrared window-I (NIR-I, 700 to 900 nm) for illumination, due to lower scattering and absorption by tissues in this wavelength range. Recently, near-infrared window-II (NIR-II, 900 to 1700 nm) has been explored for PAI. We studied the imaging depths in biological tissues for different illumination wavelengths in visible, NIR-I, and NIR-II regions using Monte Carlo (MC) simulations and validated with experimental results. MC simulations were done to compute fluence in tissue, absorbance in blood vessel, and in a spherical absorber (mimicking sentinel lymph node) embedded at different depths in breast tissue. Photoacoustic tomography and acoustic resolution photoacoustic microscopy experiments were conducted to validate the MC results. We demonstrate that maximum imaging depth is achieved by wavelengths in NIR-I window (~800 nm) when the energy density deposited is same for all wavelengths. However, illumination using wavelengths around 1064 nm (NIR-II window) gives the maximum imaging depth when the energy density deposited is proportional to maximum permissible exposure (MPE) at corresponding wavelength. These results show that it is the higher MPE of NIR-II window that helps in increasing the PAI depth for chromophores embedded in breast tissue. © The Authors. Published by SPIE under a Creative Commons Attribution 4.0 Unported License. Distribution or reproduction in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.24.12.121904]

Keywords: photoacoustic imaging; photoacoustic tomography; acoustic resolution photoacoustic microscopy; near-infrared window; imaging depth; Monte Carlo simulation with embedded object.

1 Introduction

Photoacoustic imaging (PAI) is a rapidly growing biomedical imaging technique used in various clinical and preclinical applications. PAI is based on photoacoustic (PA) effect, wherein ultrasound (US) signals are generated on absorption of light by an absorber. In PAI, the target samples or tissues generate US signals after being irradiated by laser pulses. Being a hybrid imaging modality, PAI provides strong optical contrast and high US resolution, making it a promising tool for breast tumor imaging, angiogenesis monitoring, sentinel lymph node (SLN) mapping, urinary bladder imaging, monitoring of hemoglobin concentration, etc.

Depending upon system configuration, PAI can be broadly classified as PA computed tomography/photoacoustic tomography (PAT) and photoacoustic microscopy (PAM). In PAT, the sample is homogeneously illuminated by the laser, and the acoustic signal is collected by single or multiple ultrasound transducers (USTs) from different locations around the sample. Reconstruction algorithms are applied on the acquired signals to form images of the internal structure of the object. On the other hand, PAM generally employs confocal alignment of optical illumination and US detection, and by raster scanning of the sample, three-dimensional volumetric information of the object is obtained. Such alignment provides PAM the advantage of better imaging resolution, but at a cost of relatively poor imaging depth.

Unlike pure optical imaging modalities such as optical coherence tomography, two-photon microscopy, confocal microscopy, or Raman imaging, which have a maximum imaging depth of a few millimeters, PAI has an imaging depth up to a few centimeters. This is because PAI is based on acquisition of US signals generated by the absorbers inside the sample. The PA signal generated depends on the illumination light wavelength, the optical properties of the target sample, and propagating background medium. Light within the visible window (400 to 600 nm) is generally used to image vasculature, as blood is a strong optical absorber in the visible window. However, surrounding tissues also absorb visible light, leading to poor imaging depth. The imaging depth can be increased by selecting suitable wavelengths for illumination, wherein scattering and absorption in background tissue are minimal. Therefore, first near-infrared window (NIR-I) (700 to 900 nm) is widely used for sample illumination in PAI rather than visible (400 to 600 nm) region for deep tissue imaging. Although imaging depth is increased in NIR-I, it comes with a cost of a bulky, inefficient, expensive two-stage [Ti:sapphire, dye, or optical parametric oscillator (OPO)] laser needed to generate the light in this window. Recently, a second window of near-infrared region (NIR-II, 900 to 1200 nm) has been used for PAI. This increased interest in NIR-II imaging has been attributed to further decrease in scattering of light by tissue in this region, and to the increased maximum permissible exposure (MPE) of light...
illumination in accordance with the American National Standards
Institute (ANSI) safety limit. However, as compared to the NIR-
I window, absorption of light by surrounding tissues is higher in
NIR-II. The combined effect of decreased scattering, increased
absorption, and increased MPE on depth of imaging in biological
tissues has not been compared systematically.

In this work, we have done a comparative analysis of PAI
depth in breast tissue in various optical wavelength ranges.
For the first time, we have systematically compared and ana-
lyzed the penetration depth in tissue using Monte Carlo (MC)
simulations and have further validated them experimentally. We
compared different scenarios to account for change in imaging
depth due to change in target sample, MPE at different wave-
lengths, and imaging modality. Since vasculature imaging and
SLN imaging using external contrast agent are two common
applications of PAI, environments mimicking these scenarios
were considered. Different chromophores present in the two
environments helped evaluate the effects of absorption coeffi-
cient ($\mu_a$) of target sample on imaging depth. Further, by ana-
lizing the results at both constant energy and MPE, relative
effect on imaging depth due to change in MPE and change in
tissue optical properties at different wavelengths has been
examined.

MC simulations were first performed to simulate light propa-
gation in human breast tissue with embedded targets (mimicking
blood vessel or SLN) at wavelengths 532, 800, and 1064 nm.
Owing to ease of availability, 532 and 1064 nm are extensively
used for PAI. No considerable change in optical properties is
present within NIR-I; therefore, the penetration depth remains
similar in the entire window. $\mu_s$ of 800 nm has been used for
breast vasculature imaging, it was considered an ideal choice
for performing simulations. Another consideration for selecting
the wavelengths was availability of tissue optical prop-
eties in a single publication in all the three optical windows.
Therefore, for comparing penetration depths in visible, NIR-I,
and NIR-II regions, wavelengths of 532, 800, and 1064 nm were
selected, respectively.

The simulation results were then validated by performing
experiments on both PAT and acoustic resolution photoacoustic
microscopy (ARPAM) systems using chicken breast tissue sample
and blood/ink targets. Wavelengths of 532, 740, and 1064 nm were employed for sample illumination in PAT.
Wavelengths used for experiments were slightly different from
those used for simulations, due to system restrictions. However,
since changes in optical properties of blood and tissue between
740 and 800 nm are minimal, simulation and experimental
results were expected to align. For ARPAM, samples were illu-
minated with 570- and 1064-nm wavelengths. The depth of im-
ing and the signal-to-noise ratio (SNR) were studied for all the
experiments.

2 Materials and Methods
2.1 Monte Carlo Simulations
MC simulation for light propagation in multilayered tissue
(MCML) with embedded objects (MCEO) was modified to
compute the fluence in tissue across depth and the absorbance
in objects embedded in tissue at different depths. The light
propagation in the simulation geometry is determined by the
optical properties of the medium such as $\mu_a$, scattering coeffi-
cient ($\mu_s$), scattering anisotropy ($g$), and refractive index ($n$).
Three different simulation geometries were used to compare the
propagation of photons at different wavelengths. Figures 1(a–
1(c) illustrate the cross-sectional view of simulation geometries
at $y = 0$ plane, across $xz$ axis. The first simulation geometry
(Fig. 1(a)) mimics human breast tissue of thickness 30 cm. The
fluence across the depth of the medium was recorded for a pencil
(an infinitely narrow collimated) beam. Simulations were
executed for wavelengths 532, 800, and 1064 nm.

Figure 1(b) demonstrates the second geometry mimicking
absorption of light in a blood vessel embedded in human breast
tissue. A cylinder of 1-mm diameter and infinite length was
placed at different depths in the tissue medium, which was sub-
ject to pencil beam illumination. Absorbance in cylinder was
computed at different wavelengths by varying optical proper-
ties of the medium and the cylinder to match the properties
of human breast tissue and blood, respectively. Distance
between the tissue surface and the cylinder was increased to
compute the absorption of light by the blood vessel at different
depths.

MCEO was further performed on the geometry as illustrated
in Fig. 1(c). Here, absorbance was calculated in a sphere of
diameter 1 cm placed at different depths in tissue. The $\mu_a$ of
the sphere was constant (1 cm$^{-1}$) across all wavelengths, while its
$\mu_s$ was equal to the $\mu_s$ of surrounding tissue. Such geometry
mimics imaging of organs such as urinary bladder or SLN,
which generally require injecting contrast agent such as methyl-
ene blue, indocyanine green, gold nanoparticles, and black ink,
during imaging. The sphere, therefore, mimics the organ filled
with contrast agent. Similar to the previous case, the position of
the sphere and the properties of the sphere and the tissue were
modified to get absorption at different depths and at different
wavelengths, respectively, Table 1 gives the optical properties
of medium, cylinder, and sphere, used for simulations.
10$^8$ photons were launched for each simulation to ensure
high computational accuracy. The cylinder and sphere were
embedded at depths 1 to 6 cm from the tissue surface in steps
of 1 cm. All simulations were performed on a desktop computer
with an Intel Xeon 3.7 GHz, 64-bit processor, and 16-GB RAM
running windows 10 operating system.

2.2 Acoustic Resolution Photoacoustic Microscopy
Imaging System Configuration
ARPAM system was used for comparing the images obtained
using laser illumination of 570 and 1064 nm. A nanosecond
diode-pumped solid-state Nd:YAG laser (INNOSLAB,
Edgewave, Wurselen, Germany) generated laser pulses of wave-
lengths 532 and 1064 nm at a pulse repetition rate (PRR) of
5000 Hz. A harmonic beam splitter (HBSY11) reflected a
1064-nm beam toward the 1064-nm ARPAM system and the
transmitted 532-nm beam was allowed to pass through a dye
laser (Credo-DYE-N, Sirah dye laser, Spectra Physics, Santa
Clara, California) toward the 570-nm ARPAM system. More
detailed system descriptions (and performance) of the 570-
nm ARPAM system and the 1064-nm ARPAM system have
been reported earlier. The PA signal generated by the sample
after irradiation with the 570-nm beam was collected by a
50-MHz UST (V214-BB-RM, Olympus NDT, Waltham,
Massachusetts) housed in the center of an optical condenser.
Before acquisition, the PA signal was focused with an acoustic
lens (LC 4573, Thorlabs) of diameter 1 cm placed at different
depths with contrast agent such as methyl-
ene blue, indocyanine green, gold nanoparticles, and black ink,
during imaging. The sphere, therefore, mimics the organ filled
with contrast agent. Similar to the previous case, the position of
the sphere and the properties of the sphere and the tissue were
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of 1 cm. All simulations were performed on a desktop computer
with an Intel Xeon 3.7 GHz, 64-bit processor, and 16-GB RAM
running windows 10 operating system.
of 12.4 mm was used to acquire the generated PA signal. The UST was positioned at the center of the optical condenser to achieve confocal alignment of optical and acoustic foci.

In both the ARPAM systems, the scanning setup was mounted on a three-axis motorized stage (PLS 85 for $x$ and $y$ axes, VT 80 for $z$ axis, PI-Physik Instrumente, Karlsruhe, Germany) which was controlled by a three-axis controller (SMC corvus eco, PI miCos). The scanning head was immersed in a water tank (13 x 30 cm) having an imaging window with diameter of 10 cm sealed with polyethylene membrane.

### 2.3 Photoacoustic Tomography Imaging System Configuration

The system configuration (and performance) for PAT system has been described in detail earlier. A Q-switched Nd:YAG laser

<table>
<thead>
<tr>
<th>Layer</th>
<th>Medium (human breast tissue)</th>
<th>Cylinder (blood vessel)</th>
<th>Sphere (constant absorber)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavelength (nm)</td>
<td>532</td>
<td>800</td>
<td>1064</td>
</tr>
<tr>
<td>$\mu_a$ (cm$^{-1}$)</td>
<td>0.45</td>
<td>0.011</td>
<td>0.033</td>
</tr>
<tr>
<td>$\mu_s$ (cm$^{-1}$)</td>
<td>120</td>
<td>116</td>
<td>77</td>
</tr>
<tr>
<td>$g$</td>
<td>0.9</td>
<td>0.9</td>
<td>0.9</td>
</tr>
</tbody>
</table>
delivered laser pulses of wavelengths 532 and 1064 nm at a PRR of 10 Hz. These pulses were split using a harmonic beam splitter (HBSY11) before being guided toward the imaging area. For generating pulses of wavelength 740 nm, the beam from Nd: YAG laser was pumped into a tunable OPO (Continuum, Surelite OPO) laser. The desired wavelength was guided to the sample using three right-angled prisms (PS911, Thorlabs). The setup was arranged such that the sample was irradiated homogeneously by laser pulses from above. An optical diffuser was used to homogenize the beam.

The PA signal generated by the sample was collected using a 5-MHz single-element UST (V309-SU, Olympus NDT) having a 13-mm diameter active area and a 70% nominal bandwidth. UST was connected to a stepper motor (Lin Engineering, Silverpak 23C) which rotated it around continuously to collect signal from different angles. Both the sample and the UST were placed in a water tank to enable better coupling of US signal.

2.4 Sample Preparation

Two transparent, low-density polyethylene (LDPE) tubes having an inner diameter of 0.59 mm were filled with black ink and fresh blood, respectively. Blood was chosen owing to its vast use as an endogenous light absorber in the body. Black ink, due to its high coefficient of absorption across the visible and NIR spectrum, mimics the properties of any exogenous contrast agent used during PAI.

For ARPAM, the tubes were placed under the imaging window of the water tank, while for PAT, they were directly placed inside the tank. Slices of different thicknesses of commercially available chicken breast tissue were stacked on the LDPE tubes to illustrate imaging at different depths. ARPAM data were acquired by raster scanning over the tubes in an area of 8 mm x 10 mm. Maximum amplitude projection (MAP) of the acquired data was computed using MATLAB (MathWorks, Natick, Massachusetts). PAT data were acquired by rotating the UST around the sample at a speed of 1.5 deg /s. The collected data were averaged into 240 A-lines to compensate for pulse-to-pulse variation of laser power. A simple delay-and-sum algorithm was implemented for data reconstruction.

United States Air Force 1951 test target (RIDSIP, Thorlabs) was further imaged to validate the depth of imaging in ARPAM systems. The target was placed below the imaging window, and slices of chicken breast were stacked on it. The scanning head was raster scanned in an area of 20 mm x 20 mm across the test target. Transparent US gel was used as a coupling medium between the sample and the imaging window. MAP of the acquired data was computed to form the final image.

2.5 Laser Power

Laser energy density on the surface of biological tissues is governed by ANSI safety limits. The safety limit depends on incident light wavelength and time of exposure on the sample. To analyze the effect on imaging depth due to change in optical properties of tissue (which is dependent on the wavelength of illumination) and change in MPE of light independently, experiments were conducted in two ways. Initially, imaging depth was compared using constant laser fluence across all wavelengths. Later, the fluence was changed proportionally to the MPE at that wavelength.

For ARPAM systems used here, the allowable per-pulse energy density for the 570 and 1064 nm are 1 and 2.96 mJ/cm², respectively. To analyze images at constant laser power, experiments were conducted at pulse energy of 0.30 mJ/cm² for both the wavelengths. Later, pulse energy of the 1064-nm system was increased to 1 mJ/cm² and the acquired images were compared with the images obtained previously using the 570-nm system, to compare the results when energy is proportional to MPE.

Owing to the change in exposure time, the allowable per-pulse energy density was different for PAT. In this case, initial experiments were conducted by maintaining a constant energy density of 5.1 mJ/cm² across all wavelengths. Later, pulse energy of the 532- and 1064-nm pulses was altered such that it became proportional to MPE at respective wavelengths.

For MCEO results, MPE was calculated for exposure duration between 10 s and 3 x 10⁴ s. Simulations were performed to give values of absorbance and fluence at different wavelengths and different depths. These values were multiplied with a multiplication factor (k) proportional to MPE, to give an estimate of relative absorbance at MPE. Values of MPE and k used during simulations are provided in Table 2.

3 Results

Figures 1(d)–1(i) show the results of MC simulations when tissue [Figs. 1(d) and 1(g)] and tissue with embedded objects [Figs. 1(e), 1(f), 1(h), and 1(j)] were illuminated by a pencil beam at different wavelengths. Fluence across depth in human breast tissue for wavelengths 532, 800, and 1064 nm has been shown in Fig. 1(d). Figures 1(e) and 1(f) show absorbance in cylinder and sphere, respectively, when they were embedded at different depths in breast tissue. Corresponding results of Figs. 1(d), 1(e), and 1(f) at MPE are shown in Figs. 1(g), 1(h), and 1(i), respectively. The results obtained show a drastic decrease in tissue fluence with an increase in depth for illumination with wavelength of 532 nm. The decrease in fluence resulted in a corresponding decrease in absorbance in both cylinder and sphere. This decrease can be attributed to the high μs and high μs of breast tissue in visible region compared to NIR window.

The change in absorption with depth is much lower in NIR region, as compared to visible region. It can be observed in Figs. 1(d)–1(f) that there is a higher decrease in absorption at 1064-nm illumination than at 800-nm illumination when the energy deposited for both the wavelengths is same. Although the μs decreases with increase in wavelength, the μs in breast
tissue is higher at 1064 nm than at 800 nm. This increase in $\mu_a$ results in a decrease in the amount of light reaching higher depths. Conversely, it was observed that when the tissue was subjected to exposure by all three wavelengths at MPE, fluence was highest for 1064 nm till an approximate depth of 4.5 cm, after which fluence for 1064 nm reduces compared to 800 nm. The absorbance of 1064 nm is higher than the absorbance of 800 nm, in both the cylinder and the sphere at MPE till a depth of 6 cm. Between 4.5 and 6 cm, the fluence in tissue for the 1064-nm illumination reduces with depth compared to the 800 nm, while the absorbance in the objects for 1064 nm remain higher than 800 nm. This observation can be attributed to the different properties of embedded objects at different wavelengths. Higher $\mu_s$ of embedded objects at 800 nm (compared to 1064 nm) might result in decreased absorbance of light in some parts of sphere, even though fluence in the given $xy$ plane is larger.

It can be concluded from simulations that the fluence decreases by almost 96% per cm in visible (532 nm) region, while the average decrease in NIR-I (800 nm) is $\sim$30% per cm and the average decrease in NIR-II (1064 nm) is 45% per cm. Thus, the absorbance changes by a factor of $\sim$100/cm in visible region and by a factor $<10$ in NIR region. Further, till $\sim$6 cm depth, NIR-I has higher absorbance in embedded object when laser power is same, while NIR-II has higher absorbance in embedded objects when laser power is proportional to MPE.

Experiments were conducted to validate the simulation results. Figure 2 shows the results obtained when LDPE tubes filled with black ink and blood were subjected to ARPAM experiments. Figures 2(a)–2(e) correspond to images obtained using (a)–(e) 570-nm ARPAM and (f)–(o) 1064-nm ARPAM system at (a)–(k) same fluence and (a)–(e), (l)–(o) fluence proportional to MPE. Left and right tubes were filled with black ink and blood, respectively. Scale bar is same for all images as given in (a). Numbers on top of images indicate the thickness of tissue placed on the tubes. (p) The SNR for (a)–(k). Solid lines indicate ink and dashed lines indicate blood.
using the 570-nm ARPAM system, while Figs. 2(f)–2(o) correspond to images obtained using the 1064-nm ARPAM system. In the images, the tube on the left was filled with ink, while that on the right [only visible in Figs. 2(a)–2(c)] was filled with blood. Laser fluence per pulse was maintained at 0.30 mJ/cm² for Figs. 2(a)–2(k) and at 1 mJ/cm² for Figs. 2(l)–2(o).

At same fluence, the tube containing ink was visible till a depth of 4 mm in the 570-nm ARPAM system and 5 mm in the 1064-nm ARPAM system. Further, on increasing the surface fluence of the 1064-nm ARPAM system such that the surface fluence for both the wavelengths were proportional to MPE at respective wavelength, the imaging depth increased to almost 10 mm. This corresponds to more than 100% increase in imaging depth between 570 and 1064 nm ARPAM systems for black ink. However, 570 nm showed a much stronger blood absorbance as compared to 1064 nm. LDPE tube filled with blood was visible till a depth of 3 mm in the 570-nm system, while it was not at all visible in the 1064-nm system. Visibility of blood tubes in these images is poor due to a high relative contrast of black ink. Tubes were clearly visible in Figs. 2(a)–2(c) on adjusting the color bar (images not shown). For quantification of results, SNR at different depths was computed as

\[
\text{SNR (dB)} = 10 \times \log \left( \frac{V}{N} \right),
\]

where \(V\) is the mean of PA signal (above a threshold) from an area where sample was placed and \(N\) is the standard deviation of the background noise.

Figure 2(p) shows SNR for ink (solid lines) and blood (dotted lines) for the 1064- (black triangles) and 570-nm (green circles and squares) systems corresponding to images in Figs. 2(a)–2(k). The average change in SNR of black ink was more (8 dB/mm) for the 570-nm system, as compared to the 1064-nm system (6 dB/mm). SNR of blood was 25–30% less than the SNR of ink for the 570 nm ARPAM system. Owing to low SNR, blood was not visible beyond a depth of 3 mm.

Experiments were further conducted on test targets to assess the images obtained at different imaging depths. Figures 3(a), 3(b) and 3(c)–3(e) show images obtained in the 570- and 1064-nm systems, respectively. Chicken tissue of thicknesses 0, 3, and 8 mm were stacked on targets in Figs. 3(a), 3(c); 3(b), 3(d); and 3(e), respectively. It was observed that when no tissue was placed on the sample, the images obtained using the 570-nm system showed better image quality, as compared to images obtained on using the 1064-nm system. This may be attributed to the different USTs used in the two imaging systems. Since the imaging depth of the 1064-nm system is higher, a transducer with lower central frequency was used for signal detection. This resulted in a decrease in system resolution. It can also be observed from Figs. 3(c)–3(e) that an increase in tissue thickness results in an increase in the image artifacts. Presence of chicken tissue results in refraction of PA signal at multiple surfaces. Owing to the nonuniformity of tissue, refraction at different points across the image is different, resulting in slight changes in acoustic focus and acoustic impedance. Therefore, some parts of the image appear out of focus. With an increase in tissue thickness, slight degradation in image quality was observed. However, the structures of test targets were distinct and intact. This confirms that imaging of bigger structures such as lymph node and urinary bladder can be accurately performed at higher depths using the 1064-nm ARPAM system; however, smaller structures with low absorbance might be difficult to discern at such depths.

Following this, experiments at wavelengths of 532, 740, and 1064 nm were conducted using PAT system. The images of LDPE tubes filled with blood and black ink are shown in Figs. 4 and 5. The left tube in the image was filled with black ink, while the right one was filled with blood. Figure 4 shows images acquired when the tubes were irradiated with constant surface fluence for all wavelengths. SNR values, computed in the same way as explained earlier, for blood (dotted lines) and black ink (solid lines) at different depths, for all three wavelengths are shown in Fig. 4(o). The graph shows that PA signal from blood was maximum at 532 nm in the absence of chicken tissue. This is expected due to high \(\mu_s\) of blood at 532 nm. An increase in depth of chicken tissue resulted in a decrease in SNR.
for all three wavelengths. The decrease in SNR per unit length was similar for 532 and 1064 nm and was slightly lower for 740 nm. No particular trend in SNR variation was observed. For 532 and 1064 nm, the LDPE tubes containing black ink were discernible only till imaging depth of 15 mm, while for 740 nm, the tube was visible till 25 mm. Thus, the experimental results validated the simulation results, which suggested that at constant fluence, absorbance in constant absorber at higher depths will be maximum for NIR-I region.

Figures 5(a)–5(l) show the images acquired for LDPE tubes when the surface fluence at different wavelengths was as given in Table 2. Corresponding SNR values for blood and ink at all three wavelengths are shown in Fig. 5(m). Owing to a relatively low MPE in visible region, the SNR for blood and ink for 532 nm was lower than that observed in the previous case. Black ink was still visible to a depth of 15 mm, but image quality degraded due to reduced SNR. However, due to much higher MPE of NIR-II, the SNR values at 1064 nm were at least 30% higher than the SNR values at 740 nm for imaging depths between 10 and 25 mm. The imaging depth for 1064 nm increased to 35 mm at this surface fluence. For depths beyond 25 mm, the LDPE tubes were not discernible with 740 nm. Imaging of tissue <10 mm was not performed with 1064 nm in this case, as such high power could have resulted in boiling of black ink, thus damaging the tube.

As can be seen in Table 2, MPE for all wavelengths was higher than the surface fluence used for the experiments. The laser power was limited due to the use of OPO, which was used for generating 740-nm pulses. Energy of Nd:YAG laser could further be increased to achieve an imaging depth of 45 mm at 1064-nm wavelength, as shown in Figs. 5(n) and 5(o). For all the images obtained using PAT, other image characteristics, such as resolution and spatial accuracy, remained same irrespective of the illumination wavelength used.

4 Discussion and Conclusion

In this work, imaging depths in breast tissue for different wavelengths have been compared using both MC simulations and
experiments. Two environments, one mimicking a blood vessel inside human breast tissue and another mimicking imaging of SLN following injection of a contrast agent, were considered for simulations. Tissue properties corresponding to three wavelengths, one each in visible, NIR-I, and NIR-II regions were used. Simulation results showed a severe decrease in laser fluence for visible region (532 nm) with an increase in depth of breast tissue. It was also observed that the decrease in fluence with depth was more for NIR-II (1064 nm) as compared to NIR-I (800 nm), when both were irradiated with same number of photons. At MPE, the fluence for 1064 nm was higher till an approximate depth of 4.5 cm, following which fluence of 800 and 1064 nm became equal and then fluence at 800 nm became larger. MCML simulations may further be theoretically convolved to compare penetration depth for different beam profiles; however, the trend in imaging depth for different wavelengths is expected to remain similar for all profiles.

Following simulations, experiments were conducted on both ARPAM and PAT systems. ARPAM experiments showed an increase in imaging depth for 1064 nm, as compared to 570 nm. The increment in depth of chicken breast was from 4 to 5 mm, when both the systems had same fluence, and from 4 to 10 mm, when fluence was proportional to MPE. To validate that the imaging depth of the 570-nm system was low due to the wavelength and not due to the UST used, experiments were conducted by irradiating the sample at 570 nm but by using a 30-MHz UST instead of a 50-MHz UST (results not shown). The imaging depth using this setup was similar to what was observed in a conventional 570-nm ARPAM system, as described above. Thus, confirming that the comparatively lower imaging depth was due to high $\mu_a$ and $\mu_s$ in tissue at 570 nm.

PAT experimental results validated the simulation results that the decrease in SNR with imaging depth was more for NIR-II (1064 nm) as compared to NIR-I (740 nm) when samples were
irradiated with constant surface fluence. This resulted in higher achievable imaging depth in chicken breast tissue for 740 nm. Nevertheless, analogous to simulation results, the SNR for NIR-II was higher than the SNR for NIR-I, when pulse energy in both cases was proportional to MPE at that wavelength. Therefore, in this case, the imaging depth was higher for NIR-II, as compared to NIR-I. For our experiments, imaging depths in chromophores embedded in chicken breast were 15, 25, and 15 mm, respectively, for wavelengths 532, 740, and 1064 nm when irradiated by same surface fluence. The imaging depths changed to 15, 25, and 35 mm, respectively, for 532-, 740-, and 1064-nm wavelengths, when surface fluence was proportional to MPE. Although the simulation and experimental results follow similar trend, slight differences could still be observed. This is due to the differences in optical properties of human breast tissues used in simulations versus the chicken breast tissues used in the experiments. Further variation in penetration depth for all wavelengths may be observed on changing the type of tissue (e.g., muscles and liver). In addition, variation can be observed on using excitation wavelengths different from the ones that have been used in this study, even though they might be within the same optical window. One needs to use appropriate simulation or experiments to find out the imaging depth for different combination of tissue types or wavelengths.

Our results show that the imaging depth in breast tissue for 1064 nm was higher than 740 nm due to its higher MPE. Compared to visible region, NIR regions have higher imaging depths, owing to the decreased scattering. Penetration depth of visible region may further decrease during in-vivo imaging due to presence of skin layer; however, this decrease is nominal for NIR regions and therefore skin layer has not been considered in this work. Nevertheless, between NIR-I (740 nm) and NIR-II (1064 nm), the decrease in scattering is accompanied with an increase in absorption of surrounding tissues, which results in an overall decrease in fluence at higher depths. However, since the maximum skin exposure allowed for NIR-II is higher than NIR-I, the tissue can be irradiated with higher laser energy. Therefore, deep tissue imaging possible at NIR-II can only be attributed to higher illumination energy. Simulation results also showed that absorbance in sphere becomes similar for 800 and 1064 nm at an approximate depth of 6 cm. Further, absorbance in cylinder and sphere was higher in the NIR-I region than in the NIR-II region, even when illumination was proportional to MPE. Owing to the system restrictions, experimental validation of this trend was not possible as it was difficult to get enough energy density to achieve such imaging depth experimentally.

Another reason of growing popularity of NIR-II window, specifically 1064 nm, is the availability of high laser power at this wavelength. Since traditional Nd:YAG laser can directly deliver pulses of wavelength 1064 nm, there is no need of a dye laser or OPO, which is generally needed for generating wavelengths in NIR-I region. Owing to low efficiency (15% to 25% efficiency) of the second-stage laser, their presence not only increases the cost of the system but also results in a decrease in available power. For the PAT experiments illustrated above, maximum laser power of OPO could provide an imaging depth of 25 mm at 740-nm illumination, whereas maximum laser power of Nd:YAG could provide an imaging depth of 45 mm at 1064-nm illumination.

In summary, we have compared the PAI depth in breast tissue for illumination of target samples at different wavelengths.


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