Optical clearing for luminal organ imaging with ultrahigh-resolution optical coherence tomography

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Abstract. The imaging depth of optical coherence tomography (OCT) in highly scattering biological tissues (such as luminal organs) is limited, particularly for OCT operating at shorter wavelength regions (such as around 800 nm). For the first time, the optical clearing effect of the mixture of liquid paraffin and glycerol on luminal organs was explored with ultrahigh-resolution spectral domain OCT at 800 nm. Ex vivo studies were performed on pig esophagus and bronchus, and guinea pig esophagus with different volume ratios of the mixture. We found that the mixture of 40% liquid paraffin had the best optical clearing effect on esophageal tissues with a short effective time of ∼10 min, which means the clearing effect occurs about 10 min after the application of the clearing agent. In contrast, no obvious optical clearing effect was identified on bronchus tissues.

Keywords: optical coherence tomography; optical clearing technique; luminal organs.

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1 Introduction

Optical coherence tomography (OCT) affords high-resolution imaging of tissue microanatomy in vivo. However, the limited imaging depth due to the high scattering of light in biological tissues hinders its potential applications, where a deeper imaging depth is desired. To achieve a better imaging depth, optical clearing is regarded as the most effective method. It is generally considered that optical clearing results from collective effects including the refractive index matching effect, the cell dehydration effect, and the increased collagen solubility due to the osmotic property of optical clearing agents (OCAs), which leads to a deeper penetration of light in biological tissues.1,2 Recently, increasing research interest has been witnessed on the efforts to develop and validate new and more effective OCAs.

New OCAs, mixtures of two or more agents, capable of extending imaging depth or even making tissue transparent, such as SeeDB,3 benzyl alcohol and benzyl benzoate,4 Scale,5 and the mixture of tetrahydrofuran and dibenzyl ether,6 were developed and successfully demonstrated in several ex vivo mouse brain tissue studies. With these new OCAs, a few days’ process for dehydration and maceration of biological tissues was needed for optical clearing to take effect. In contrast, a new method for three-dimensional imaging of solvent-cleared organs successfully reduced the process time to several hours.7 However, all these new OCAs are toxic and suitable only for ex vivo applications.

For in vivo applications, an ideal OCA should be safe to use and has a short effective time (which is defined as the time for the optical clearing effect to take place after the application of the agent). To this end, some biocompatible hyperosmotic and hydrophilic chemical agents, e.g., polyethylene glycol,8 glucose,9 glycerol,10,11 and propylene glycol,12 have been tested. The hyperosmotic property leads to an efficient diffusion of these biocompatible OCAs into biological tissues. For example, an effective time of about 50 min was reported in an in vivo study on human skin by using glycerol, which is one of the most commonly used biocompatible agents for optical clearing.1 However, the limited optical clearing effect of glycerol and other biocompatible OCAs was achieved.30 To further shorten the effective time, some physical methods, such as sonophoresis with ultrasound,21 rubbing with sandpaper,22 or puncturing with microneedle roller,23 were developed to aid the diffusion of glycerol through the dense stratum corneum (SC) of skin. However, these methods cannot be directly or conveniently applied to luminal organs in vivo. Alternatively, it was proposed to optimize the penetration efficiency of glycerol in biological tissues by incorporating a permeation enhancer, such as oleic acid,24 dimethyl sulfoxide,25 thiazione,26 azone,27 thiazone,28,29 or liquid paraffin.30,31,32 The enhanced optical clearing effect of liquid paraffin and glycerol mixture was validated on human skin, where a much shorter effective time of ∼10 min was achieved.30,32 Furthermore, as a lipophilic agent, it was found that liquid paraffin can greatly relieve the severe dehydration effect caused by glycerol,30,31 which induces tissue warping and might not be safe to use for certain clinical applications. It is noteworthy that liquid paraffin has been used in cosmetics and food applications for >100 years.34 Liquid paraffin has also been approved by the Food and Drug Administration as a safe pharmaceutical vehicle and is listed in many different pharmacopeias.34

In this paper, we studied the optical clearing effect of liquid paraffin and glycerol mixture on luminal organs ex vivo with a spectral domain optical coherence tomography (SD-OCT) system operating at 800 nm. Our experimental results verified that the mixture of 40% liquid paraffin had the optimum optical clearing effect on both guinea pig and pig esophagus tissues with an effective time as short as only about 10 min, whereas no obvious optical clearing effect was found on pig bronchus tissues.
2 Materials and Methods

2.1 Materials

Esophagus and bronchus tissues from four pigs and esophagus tissues from one guinea pig were harvested for this study right after the animals were euthanized for other purposes approved by the Animal Care and Use Committee at the Johns Hopkins University. The luminal tissues were cut into small pieces of ~1 x 1 cm immediately after tissue harvest. Fresh specimens were submerged in phosphate-buffered saline (PBS) solution and kept in a 4°C refrigerator for no more than 5 h before use. After being well mixed by stirring for at least 10 min, about 40 μl of the liquid paraffin and glycerol mixture with a volume ratio of 20% to 60% were immediately applied on the inner surface of the luminal tissues using a syringe needle. It was found that a well-mixed liquid paraffin/glycerol mixture took over 4 h to have a complete phase separation, while in our experiments each test lasted for at most 50 min so as to greatly mitigate the influence of phase separation. Herein, the volume ratio was calculated as \( V_{lp} / (V_{lp} + V_{gl}) \), where \( V_{lp} \) is the volume of liquid paraffin and \( V_{gl} \) is the volume of glycerol.

2.2 Imaging System

As shown in Fig. 1, our SD-OCT system adopted a customized broadband linear-in-wavenumber spectrometer to afford an imaging depth of ~1.2 mm, the detail of which was reported previously.35,36 A homemade Ti:Sapphire laser with a central wavelength of ~825 nm and a 3-dB bandwidth of ~150 nm was used to achieve an axial resolution of ~2.8 μm (in air). Two galvanometer scanning mirrors were employed in the sample arm for performing X – Y two-dimensional beam scanning. A scan lens was used to afford a lateral resolution of ~16 μm. Sample imaging was performed at 70 k A-scans/s.

![Schematic of the ultrahigh-resolution spectral-domain OCT imaging system.](image)

\( \text{Ti:Sapphire Laser} \quad \lambda_c = 825 \text{ nm} \)
\( \Delta \lambda_{\text{FWHM}} = 150 \text{ nm} \)

\( \text{Linear K Spectrometer} \)

\( \text{BS, beam splitter; C, multielement achromatic collimator; charge coupled device (CCD), line scan CCD; CG, cover glass; G, grating; GSM, galvanometer scanning mirrors; M, mirror; MESL, multielement scan lens; P, linear K mapping prism; PP, prism pair; S, specimen; PBS, phosphate-buffered saline; PD, Petri dish; SL, scan lens; and TF, tunable filter.} \)

2.3 Experiment Design

All imaging experiments were performed at room temperature. As shown in Fig. 1, the lower part of the specimen was immersed in a small volume of PBS solution during imaging to prevent the specimen from dehydration. The liquid paraffin/glycerol mixture was applied evenly on the tissue surface and sealed between a cover glass and the specimen surface. In our control experiments, PBS instead of the liquid paraffin/glycerol mixture was applied on the tissue surface. The diffusion process of the mixture was studied by B-scan (cross-sectional) imaging of the tissue at a given place every 5 min for a total of 50-min duration to understand the efficacy of the optical clearing effect. Each B-scan image consisted of 2048 x 2048 pixels (lateral x axial) corresponding to a physical image size of 2 mm x 1.2 mm. Each dataset consisted of 20 B-scan images, which were used for averaging to reduce speckle noise.

2.4 Merit Metrics

B-scan images provided direct and qualitative visual assessment of the optical clearing effect. To quantify such effect, some metrics have been proposed previously.17,30,31,37-40 Considering the heterogeneous nature of tissue structures, tissue optical property, and their potential changes during the process of OCA application and diffusion, evaluating the optical property change at one fixed tissue depth alone would not be sufficiently accurate. Therefore, in addition to optical property, we proposed a merit metric called intensity ratio of regions (RIR),30,31 which calculates the ratio of average intensities of a designated region inside the tissue and a region nearby the tissue surface, and it is defined as

\[
\text{RIR} = \frac{1}{N_1} \sum_{x,y \in \text{Internal}} I(x,y) \quad \frac{1}{N_2} \sum_{x,y \in \text{Surface}} I(x,y),
\]

where \( N_1 \) and \( N_2 \) are the number of pixels of the two regions of interest, respectively, and \( I(x,y) \) represents the OCT intensity at position \((x,y)\). The time-dependent optical clearing effect was assessed with RIR and benchmarked with the RIR value a minute after the application of the clearing agent to obtain the relative RIRs. Generally, a larger relative RIR value means a better optical clearing effect.

3 Experimental Results

As shown in Fig. 2, a series of representative B-scan images at a fixed location on the guinea pig esophagus were displayed over a time window of about 50 min after the application of the liquid paraffin (at 30% volume concentration) and glycerol mixture. Since it was difficult to ensure the same B-scan position before and after the application of the clearing agent, the B-scan image used as a benchmark was acquired at the first minute after the application of the mixture. As shown in Fig. 2(b), the optical clearing effect of the mixture on the esophagus tissue was evident. The increased transparency of upper layers, such as epithelium and lamina propria, led to a pronounced muscle layer at only 10 min after the application of the paraffin–glycerol mixture.

To quantitatively study the optical clearing effect and find the optimum paraffin–glycerol volume ratio, mixtures of 30%, 40%, and 50% of liquid paraffin were used on pig esophagus tissues.
Small image regions of 400 × 100 pixels (lateral × axial) near the tissue surface and at different depths were used to calculate the RIRs. In this study, regions starting from pixel 1, 600, and 1000 below the tissue surface along the imaging depth were selected, which correspond to 0, ∼360, and ∼600 μm below the tissue surface, respectively, as shown in Fig. 2.

As shown in Fig. 3, the relative RIR value [calculated by using Eq. (1) and then dividing by the RIR benchmark value] was studied at different tissue depths and plotted along with time after the application of the mixture. As mentioned earlier, the RIR benchmark value was the one calculated over the same image region but at the first minute after the mixture application. The standard deviations were calculated with three pig esophagus tissues and displayed as error bars in Fig. 3. It is found that optical clearing by the mixture occurred on the esophagus tissue with the relative RIRs > 1 for all paraffin volume ratios used in this study. The bigger the relative RIR is, the higher the improvement is for the intensity value at the internal position relative to that at the surface, which means a better optical clearing effect. The best optical clearing effect (represented by the highest relative RIR) was achieved with a paraffin volume ratio of 40% by comparing the relative RIR values at different tissue depths (in particular at depths of ∼360 μm), which is consistent with the observation reported in Ref. 30. As seen from Fig. 3, the optical clearing effect took place very rapidly during the first 10 min for all tested mixtures on the esophagus tissue. This is a remarkable property, which may be very useful for the in vivo applications, where a fast optical clearing effect is needed. Validation experiments using guinea pig esophagus tissues revealed the similar results (results not shown).

To further scrutinize the consequences of optical clearing, depth-resolved optical properties (i.e., the attenuation coefficient) of the esophagus tissue have been studied versus time after the application of the liquid paraffin/glycerol mixture. The algorithm used for quantifying the optical attenuation coefficient is similar to the one reported in Ref. 41. In essence, the depth-resolved attenuation coefficients for each A-scan were calculated with the following equation:

$$\mu[i] \approx \frac{I[i]}{\Delta \sum_{i'=-1}^{i+1} I[i']}.$$

(2)

where $\mu[i]$ is the attenuation coefficient at the $i$’th pixel of a given A-scan, which is defined as the averaged attenuation over the pixel size of $\Delta$. $I[i]$ is the OCT intensity at the $i$’th pixel of that A-scan. It is noteworthy that Eq. (2) is slightly different from the one reported in Ref. 41 by assuming the OCT...
A representative attenuation image of the pig esophagus tissue at 10 min after the application of a 40% paraffin/glycerol mixture is shown in Fig. 4(a). As highlighted with a dashed line rectangle in Fig. 4(a), a region of interest with 400×1300 pixels (lateral × axial) was selected to calculate the averaged attenuation coefficient along the A-scan. A significant time-dependent change on the attenuation coefficient was observed as shown in Fig. 4(b). It was found that the attenuation coefficient decreased gradually at the region near the tissue surface and increased at the deeper tissue regions with time. This change eventually led to a decreased attenuation inhomogeneity of the esophagus tissue with the most dynamic change happening during the first 10 min after the application of the optical clearing mixture.

To study the optical clearing effect of liquid paraffin and glycerol mixture on other internal organs, we further tested the pig bronchus tissues. No significant optical clearing effect was observed for all tested mixtures with different volume ratios. As shown in Fig. 5 (for a representative 30% paraffin/glycerol volume ratio), the relative RIR values changed only slightly with time at different tissue depths. The error bars were calculated with three pig bronchus tissues. It is also noteworthy that in the control experiments no optical clearing effect of PBS was observed for either esophagus or bronchus tissues.

4 Discussions and Conclusion

It has been proved that lipophilic agents, such as liquid paraffin, are capable of penetrating epithelium tissue and can serve as a carrier for drug delivery.\textsuperscript{34,42,44} Our experimental results verified that liquid paraffin can facilitate the penetration of glycerol through the dense stratified squamous epithelium of esophagus tissue into deeper layers and work collaboratively with glycerol for tissue index matching. This effect was also confirmed in our previous studies on human skin, where a significantly enhanced optical clearing effect was found by using liquid paraffin to aid the penetration of glycerol through the SC layer of skin.\textsuperscript{30–32} The reason that this facilitation effect does not work for bronchus tissue may be due to the difficulty for the mixture to penetrate the cartilage rich in pig bronchus.

Compared with our previous studies, where a 1500-nm light source was used, higher relative RIR values (i.e., more pronounced optical clearing effect) were found in this study with an 800-nm SD-OCT system. This finding coincides with many previous publications reporting a better optical clearing effect at shorter wavelengths.\textsuperscript{20,26,28} Furthermore, it was found that a fine balance between the dehydration effect of glycerol and the water retention capability of the liquid paraffin helped avoid tissue deformation when using the mixture of 40% liquid paraffin. This feature is very useful for potential in vivo applications. In contrast, it was also noted that with the mixture of a low-volume ratio, such as 20%, the esophagus tissues showed some obvious warping of tissue surface due to the dominant dehydration effect of glycerol.

The phenomena of a decrease in scattering in upper layers and an increase in scattering in lower layers with time in Fig. 4 prove the attenuation coefficient tends to become homogeneous. From Fig. 4, we can see the attenuation coefficients that have the most dynamic change in the first 10 min after the application of the optical clearing mixture, which means

\[
I[i] \propto e^{-\mu(z) \cdot z}, \quad \text{where } \mu(z) \text{ is the attenuation coefficient at medium depth of } z.
\]
glycerol rapidly penetrates into inner tissue with the help of liquid paraffin. After 20 min, they change slowly with time until they become homogeneous. This change can be attributed to the index matching effect of the liquid paraffin/glycerol mixture, which leads to more homogeneous tissue refractive index that in turn reduces tissue attenuation (mainly scattering) and consequently enhances light penetration in the tissue.

In summary, the optical clearing effect of the liquid paraffin/glycerol mixture on luminal organs, such as the esophagus and bronchus, was studied for the first time at 800-nm wavelength region with an ultrahigh-resolution SD-OCT system. A significant optical clearing effect on esophagus tissues was observed with a short effective time of ~10 min, whereas no obvious optical clearing effect was found on bronchus tissues. We validated that the mixture of 40% liquid paraffin afforded the optimum optical clearing effect on the esophagus tissues. These findings will facilitate the use of the liquid paraffin/glycerol mixture in the in vivo endoscopic imaging of lumon organs in the future.

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References


Biographies for the authors are not available.