Imaging ex vivo and in vitro brain morphology in animal models with ultrahigh resolution optical coherence tomography

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Abstract. The feasibility of ultrahigh resolution optical coherence tomography (UHR OCT) to image ex vivo and in vitro brain tissue morphology on a scale from single neuron cells to a whole animal brain was investigated using a number of animal models. Sub-2-μm axial resolution OCT in biological tissue was achieved at different central wavelengths by separately interfacing two state-of-the-art broad bandwidth light sources (titanium:sapphire, Ti:Al2O3 laser, λc=800 nm, Δλ = 260 nm, Pout = 50 mW and a fiber laser light source, λc =1350 nm, Δλ =470 nm, Pout = 4 mW) to free-space or fiber-based OCT systems, designed for optimal performance in the appropriate wavelength regions. The ability of sub-2-μm axial resolution OCT to visualize intracellular morphology was demonstrated by imaging living ganglion cells in cultures. The feasibility of UHR OCT to image the globular structure of an entire animal brain as well as to resolve fine morphological features at various depths in it was tested by imaging a fixed honeybee brain. Possible degradation of OCT axial resolution with depth in optically dense brain tissue was examined by depositing microspheres through the blood stream to various depths in the brain of a living rabbit. It was determined that in the 1100 to 1600-nm wavelength range, OCT axial resolution was well preserved, even at depths greater than 500 μm, and permitted distinct visualization of microspheres 15 μm in diameter. In addition, the OCT image penetration depth and the scattering properties of gray and white brain matter were evaluated in tissue samples from the visual cortex of a fixed monkey brain. © 2004 American Institute of Physics.

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1 Introduction

In recent years there has been a constant demand for development of noninvasive or minimally invasive imaging techniques that can be applied in neurosurgery as guiding tools and as an alternative to standard excisional biopsy. So far a vast variety of imaging methods ranging from magnetic resonance imaging (MRI) to optical microscopy have been used at different stages in the diagnostics, treatment, and postoperative monitoring of various neuropathologies. Despite all their advantages, each of these methods possesses some inherent limitation related to image resolution, acquisition time, specificity and accuracy of the acquired image or information, etc., which motivates the constant search for new more precise and less invasive imaging modalities.

In neurosurgery the goal of any intracranial intervention is to provide accurate localization, diagnosis, and appropriate treatment of intracranial abnormalities, while causing minimal damage to the intact brain. During open skull surgery, the boundaries of some lesions, in particular low-grade gliomas, as well as the presence and location of small metastases of various brain tumors, may be difficult to determine by visual inspection or microscopic imaging of the tissue surface. Since brain pathologies are characterized by alterations in the morphology and physiology of healthy brain tissue, an imaging method with sufficient resolution and penetration depth that is able to discriminate between healthy and pathological brain tissue can find applications in neurosurgery as a diagnostics tool.

Optical coherence tomography (OCT) is a noncontact optical imaging method that combines micrometer-scale resolution with millimeter image penetration depth and as such may have a potential as a guiding and diagnostics tool in neurosurgery. Although OCT has been successfully applied to many biomedical fields, such as ophthalmology, dermatology, gastroenterology, etc., since its invention about a decade ago, so far only a few attempts have been made to image brain
tissue morphology and function with standard resolution OCT.\textsuperscript{9–12} However, the spatial resolution in these cases (axial resolution ~10 to 15 $\mu$m) was insufficient to permit visualization of fine morphological details such as microcalcifications and displasias that are characteristic of various types of brain tumors. Recent advances in laser technology have led to the development of UHR OCT,\textsuperscript{13–19} (axial resolution <2 $\mu$m), and current research in ophthalmology has clearly demonstrated the feasibility of UHR OCT to image intraretinal morphology, as well as to view small structural features in intraocular pathologies.\textsuperscript{20–22}

The objective of this project was to investigate the feasibility of using UHR OCT to visualize small morphological features in brain tissue by using animal models and to establish a correlation between structural details present both in the OCT tomograms and in the eosin-stained histological cross-sections. In addition, this study aimed to determine the wavefront effect was compensated for by acquiring and subsequently fusing multiple tomograms obtained at different depths at the same transverse location. The image acquisition rates for the free-space and the fiber-based OCT systems were 10 A-scans/s and 250 A-scans/s, respectively. The OCT systems were evaluated to provide 0.9×2 $\mu$m (axial×lateral) resolution in biological tissue, and a sensitivity of 110 dB for 5 mW at the sample surface for the case of the Ti:Al$_2$O$_3$ laser ($\lambda_c$ = 800 nm) and 1.4×4.5 $\mu$m spatial resolution and 98 dB sensitivity at 0.5 mW for the case of the MenloSystems fiber laser-based light source ($\lambda_c$ = 1350 nm). Full fringe detection was realized by use of a high-speed (10 Ms/s), 16-bit A/D converter to digitize the fringe data, thus permitting extraction of functional and spectroscopic information in addition to the high-resolution morphological imaging.

For the purpose of this study, tissue samples were prepared from the brains of honeybees, rabbits, and monkeys. The sophisticated structural organization of the honeybee brain, along with its small size, have made it an ideal object for testing the ability of UHR OCT to image small morphological details and to discriminate among tissue layers in optically dense brain tissue. The monkey brain was chosen because its structural organization and tissue optical properties most closely resemble the human brain. To test the ability of UHR OCT to image subcellular features in living neuronal cells, cell cultures of sympathetic ganglion cells were obtained from rat superior cervical ganglia.

Because postmortem brain tissue quickly loses optical quality as a result of cell degradation, all brain issue samples were fixed in 4% paraformaldehyde solution. Depending on the animal model, a brain tissue slice or the whole brain was placed in a custom-designed chamber with an optical window (~150 $\mu$m thick glass coverslip) through which the tissue was imaged. During the imaging procedure, care was taken to properly compensate for the dispersion mismatch introduced by the glass coverslip and the excess fixation solution between the glass window and the tissue surface in order to preserve the high OCT axial resolution in all imaged tissue samples.

For the case of the honeybee, eosin-stained histological cross-sections (5 $\mu$m thick) of the whole brain were prepared.
and imaged with a regular microscope for comparison with the OCT tomograms. To examine the degradation of OCT axial resolution with imaging depth in optically dense brain tissue, resulting from dispersion mismatch and detection of multiply scattered light, 15-μm large microspheres were injected into the blood stream of a living rabbit. After euthanasia, the rabbit brain was fixed in 4% paraformaldehyde and cross-sections containing microspheres trapped in small blood vessels were identified using a regular microscope. Tissue slices were also obtained from the visual cortex of a fixed monkey brain. For the case of imaging cell morphology, ganglion cell cultures were prepared on protein-coated glass coverslips and placed in a custom-designed perfusion chamber filled with nutritious solution to preserve the normal condition of the cells.

3 Results and Discussion

To test the suitability of UHR OCT for imaging morphology of neuron cells, a series of images were obtained from ganglion cell cultures using the fiber-based OCT system. Figures 2(b) to 2(d) show tomograms of single cells and groups of cells acquired at \( \lambda \sim 800 \) nm with 110-dB sensitivity compared with a representative image of the cells obtained with a regular microscope [Fig. 2(a)]. Considering the average size of the cells (<50 μm) and the depth of focus of the imaging optics (~80 μm), the imaging beam was focused slightly below the glass/solution interface and single tomograms of the cells were acquired (no fusing of images acquired at different focal depths was necessary). The thick black line in the upper part of all OCT images corresponds to a reflection from the glass coverslip, to which the cells were attached. The highly reflective (black) spots most likely correspond to various cell organelles, judging from the high optical density and the size of the objects. Although the cell membrane is not clearly visualized, the boundaries of the cell cytoplasm (pale gray color) are distinctly visible. Furthermore, the high spatial OCT resolution permits imaging of the thin axonal extensions of the ganglion cells (marked with black arrows in Fig. 2). Previous research on isolated neurons has demonstrated that the propagation of action potentials is accompanied by cell membrane depolarization, which further results in changes in the light-scattering properties of the cell, which are most pronounced at the base of the axon. Distinct visualization of neuronal axonal extensions in cell cultures containing multiple cells may provide useful information on the transmission of signals in neuronal networks.

To examine the feasibility of UHR OCT for imaging fine morphological details in depth in optically dense brain tissue, tomograms were acquired from the fixed whole brain of a honeybee. Figure 3(a) shows a schematic of the bee brain, where the dotted white line marks the location of a representative OCT tomogram. Although the bee brain is very small (~1 mm³), it has a sophisticated globular structure and functional organization that correspond to a vast repertoire of behavioral patterns. Furthermore, neurons in the bee brain...
are spatially organized to form layers for example, the visual center of the brain [medulla, Fig. 3(a), ME] contains alternating layers composed of neuron somas and axonal extensions, while the neurons in the antenna lobes [Fig. 3(a), AL] are oriented in such a way that the somas create a spherical shell inside which the axonal extensions are tightly packed. Approximately 20 OCT tomograms (3 × 1 mm), separated by steps of 25 μm were obtained from the bee brain with both the Ti:Al₂O₃ laser (coupled to the fiber-based OCT system) and the fiber-based light source (interfaced to the free-space OCT system). Figure 3 shows representative images acquired at λc~800 nm [Fig. 3(c)] and λc~1350 nm [Fig. 3(d)] through the medulla and the antenna lobes at a location marked with a dotted white line in Fig. 3(a). Since the fiber-based OCT system has a limited depth of focus, to minimize loss of transverse image resolution, three tomograms were acquired at different focal depths and fused together to produce the image in Fig. 3(c). Figure 3(b) shows the corresponding histological cross-section. Boundaries between different cell layers are distinctly outlined on both OCT tomograms (marked with solid black arrows) and correspond well to the layered structure observed in the histological image. A part of the lobula [Fig. 3(a), LO] positioned below the medulla is clearly visible on the right side of the image acquired at longer wavelengths [Fig. 3(d), dotted black arrow] and corresponds well to the structure observed in the histological cross-section. Note that the central parts of the antenna lobes appear dark (highly backscattering), surrounded by a ring of low backscattering tissue. This appearance can be explained by the fact that the large number of tightly packed axon-dendritic cell extensions tend to scatter light more than the neuron somas. Additional OCT tomograms acquired at various locations in the bee brain and not shown here also demonstrated the ability of UHR OCT to clearly visualize the nerve fiber bundles connecting the medulla with the honeybee eyes, as well as the honeycomb structure of the bee’s eyes consisting of small compartments 30 to 50 μm in diameter. Although the tomogram in Fig. 3(d) obtained with the fiber-based light source was acquired with significantly lower sensitivity (98 dB versus 105 dB for Ti:Al₂O₃), the penetration depth and the overall image quality were comparable to that of the tomogram acquired with the Ti:Al₂O₃ laser. In general, light scattering in optically dense tissue decreases monotonically with wavelength, which accounts for the better image penetration in all tomograms acquired at λc~1350 nm compared with images obtained at λc~800 nm.

Figure 4 summarizes representative results from the test of OCT axial resolution degradation in brain tissue. Figure 4(a) shows a regular microscope image of a fixed rabbit brain slice. The white dashed line through it marks the position of a transverse OCT scan acquired through the gray and white brain matter using the fiber-based light source coupled to the free-space OCT system. Figure 4(b) shows the OCT image (6 × 2 mm), the left edge of which corresponds to the brain surface. The tomogram clearly demonstrates the difference in optical scattering properties of gray and white brain matter, resulting in varying image depth penetrations through the tomogram. The penetration depth is approximately 1 mm in gray matter (left part of the image) and diminishes to less than 500 μm in white matter (close to the central part of the image). The OCT scan was acquired at a location near the brainstem where nerve fiber bundles are abundant. The central part of the OCT tomogram shows a group of densely packed, almost elliptically shaped nerve fiber bundles. Figure 4(c) presents an enlarged view of the region and demonstrates the ability of UHR OCT to distinctly outline single fiber bundles ~50 μm in size in a transverse direction.

The right part of the OCT tomogram shows a number of highly reflective (dark) spots, almost spherical in shape and positioned on a straight line [Fig. 4(e), enlarged view of the region]. A comparison with regular microscope images [Fig. 4(e)] revealed that the spots correspond to 15-μm-diameter microspheres embedded in a small blood vessel. Considering that both water dispersion and detection of multiply scattered light can cause degradation of OCT image axial resolution as a function of depth in biological tissue, it is important to note that in this case the deterioration of image reso-

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**Figure 4** (a) A microscope image of a rabbit brain slice in which the white dashed line marks the location of a transverse OCT scan. (b) Corresponding OCT tomogram (6 × 2 mm) acquired at λc ~1350 nm with 98-dB sensitivity. Insets: (c) enlarged view of the nerve fiber bundles, (d) enlarged view of the embedded microspheres, and (e) a regular microscope image of the microspheres.
lution was not sufficient to prevent distinct visualization of a 15-μm large microsphere positioned at a depth of ~500 μm below the brain tissue surface.

To evaluate the OCT image penetration depth in fixed brain tissue, tomograms were obtained from a brain slice from the visual cortex of a monkey brain with both the Ti:Al₂O₃ laser and the fiber-based laser. Figure 5 shows two representative OCT images (1×0.63 mm) acquired at λ ≈ 800 nm with 110-dB sensitivity [Fig. 5(a)] and at λ ≈ 1350 nm with 98-dB sensitivity [Fig. 5(b)]. In both tomograms the brain tissue appears almost homogeneous, with no distinct layered structure, except for the clearly visualized dura. In general, the monkey visual cortex (thickness ~0.5 mm) consists of six layers characterized by different thicknesses, cell types, and cell densities. The lack of a distinctly visible layered structure in the OCT tomograms may be caused by two factors: first, the spatial distribution and density of neuron somas varies slowly and gradually from one layer to the next, so the layers’ boundaries are not distinctly outlined; and second, the fixation in 4% paraformaldehyde changes the optical properties of brain tissue and may reduce the naturally existing contrast between neuron somas and the extracellular milieu.

To determine the scattering properties of fixed brain matter in the 670 to 930-nm wavelength range, OCT tomograms of gray and white brain matter were acquired [Figs. 5(c) and 5(d)]. Reflectivity profiles as functions of imaging depth were extracted by averaging over twenty adjacent A-scans [Fig. 5(e)]. The scattering coefficient averaged over the spectral bandwidth of the Ti:Al₂O₃ laser was evaluated to be μ_s ~ 9 mm⁻¹ in the case of gray matter, which compared well with previously published data on the optical properties of human brain tissue. However, for white matter, the measured scattering coefficient μ_s ~ 22 mm⁻¹ was about 30% smaller relative to the value obtained for fresh human white matter. This discrepancy may be attributed to two factors. It is possible that monkey white brain matter is not as highly scattering as human white matter, or that the difference arises from the fixation of the monkey brain in 4% formaldehyde. Since neither fresh monkey brain slices nor literature values for the scattering properties of fresh monkey white matter were available during this study, this discrepancy remains to be investigated in the future.

4 Conclusion

The feasibility of UHR OCT for imaging in vitro brain tissue morphology on the scale from single neuron cells to the structural organization of an entire animal brain was investigated with the intention of determining the suitability of this imaging technique as a potential diagnostics and research tool in neurology and neurosurgery. The results presented in this paper have demonstrated that although the UHR OCT penetration depth is limited to about 1 to 2 mm in optically dense brain tissue, the micrometer-scale image resolution is well preserved, even at greater imaging depths, to permit clear visualization of morphological features the size of 10 to 50 μm. Furthermore, tomograms acquired in various animal brain tissue samples have shown that in cases when the neuron cell density and spatial distribution are such as to provide sufficient optical contrast, UHR OCT is able to spatially resolve layered or globular structures within the highly scattering brain matter. Considering that development of any neurological disease is accompanied by morphological and physiological changes on the cellular level or a larger scale, UHR OCT may be able to discriminate, not only between healthy and pathological brain tissue, but also between various neuro pathologies. Provided that future investigations confirm this hypothesis, UHR OCT may be successfully used as a diagnostics tool in neurology and neurosurgery.
In addition, the sub-2-μm OCT axial resolution proved sufficient to distinguish intracellular components such as nuclei in cultures of living electrically active ganglion cells, as well as to image thin axonal cell extensions. These results are of significant importance, suggesting that UHR OCT may find applications as an investigative technique in basic neurological research.

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References