Optical coherence tomography visualizes neurons in human entorhinal cortex

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Abstract. The cytoarchitecture of the human brain is of great interest in diverse fields: neuroanatomy, neurology, neuroscience, and neuropathology. Traditional histology is a method that has been historically used to assess cell and fiber content in the ex vivo human brain. However, this technique suffers from significant distortions. We used a previously demonstrated optical coherence microscopy technique to image individual neurons in several square millimeters of en-face tissue blocks from layer II of the human entorhinal cortex, over 50 μm in depth. The same slices were then sectioned and stained for Nissl substance. We registered the optical coherence tomography (OCT) images with the corresponding Nissl stained slices using a nonlinear transformation. The neurons were then segmented in both images and we quantified the overlap. We show that OCT images contain information about neurons that is comparable to what can be obtained from Nissl staining, and thus can be used to assess the cytoarchitecture of the ex vivo human brain with minimal distortion. With the future integration of a vibratome into the OCT imaging rig, this technique can be scaled up to obtain undistorted volumetric data of large sections of the human brain. Optical coherence tomography (OCT) is versatile and generates cytoarchitecture and myeloarchitecture-like images. In addition, current OCT techniques can only clear a few hundred microns or at most a millimeter or so of the myelinated human tissue, making it currently infeasible to image large sections of the human brain. Optical coherence tomography (OCT) is versatile and generates cytoarchitecture and myeloarchitecture-like images. OCT relies solely on the intrinsic optical properties of the tissues, mainly those of neurons and myelinated fibers. No dye is required and the tissue block is directly imaged, no sectioning is necessary prior to image acquisition. This last point is critical as imaging before cutting implies that the vast majority of distortions that plague standard histology are avoided in the OCT images. It is important to note that while optical coherence microscopy has been performed in vivo on fixed rodent tissue previously, fixed human tissue has been far less studied. In large part, this is due to the challenges one faces in analyzing human tissue that

tissue over several hundreds of microns in depth. In standard microscopy, fluorescent dyes label structures or proteins of interest. When coupled with a vibratome, larger volumes are imaged, such as a full mouse brain. To improve the depth penetration of light, tissue can be cleared using techniques such as CLARITY and rendered optically transparent while retaining the structural anatomy. The tissue can then be stained, imaged by two-photon microscopy, and repeated, which allows undistorted results, automatically registered with diverse and specific immunocytochemical information. All these methods, however, rely on staining that contains multiple steps and can be a long process for several cubic centimeters of tissue blocks. In addition, current CLARITY techniques can only clear a few hundred microns or at most a millimeter or so of the myelinated human tissue, making it currently infeasible to image large sections of the human brain. Optical coherence tomography (OCT) is versatile and generates cytoarchitecture and myeloarchitecture-like images. OCT relies solely on the intrinsic optical properties of the tissues, mainly those of neurons and myelinated fibers. No dye is required and the tissue block is directly imaged, no sectioning is necessary prior to image acquisition. This last point is critical as imaging before cutting implies that the vast majority of distortions that plague standard histology are avoided in the OCT images. It is important to note that while optical coherence microscopy has been performed in vivo on fixed rodent tissue previously, fixed human tissue has been far less studied. In large part, this is due to the challenges one faces in analyzing human tissue that

1 Introduction

Cellular features of the human brain are not fully observable with current in vivo imaging technologies [i.e., magnetic resonance imaging (MRI) and positron emission tomography]. The importance of these features is raised by the observation that many diseases are defined by tissue properties or neuronal loss that are not visible individually in living humans. The quantification of stereologic factors such as counting neurons typically performed in postmortem histology helps define disease diagnosis and severity such as Alzheimer’s disease. Traditional histology provides the ground truth for neuroanatomy and neuropathology, and remains by far the most common way to visualize neurons and axons. Recently, a three-dimensional (3-D) model of a human brain, called BigBrain, at a nearly cellular resolution of 20 μm and based on the reconstruction of 7404 histological sections was created. Each of the 7404 slices were beautifully stained and digitized. However, traditional methods are labor intensive and introduce irremediable distortions due to cutting, mounting, and staining. Distortions lead to challenges in registering histology slices back to minimally deformed volumetric data such as blockface images, ex vivo MRI, and in vivo MRI, and in registering each slice to its neighbors to generate undistorted 3-D volumes.

Optical imaging has emerged as a promising alternative to traditional histology. For example, two-photon microscopy provides undistorted high resolution volumetric imaging of brain

Keywords: histology; neuroanatomy; neurons; neuropathology; neurophotonics; optical coherence microscopy.

Paper 14063RR received Sep. 12, 2014; accepted for publication Jan. 12, 2015; published online Feb. 9, 2015.
pose significant hurdles. For example, the time period between death and fixation, called the postmortem interval (PMI) ranges from hours to days in humans, while in rodents it is essentially zero. During this interval, there are significant autolytic processes that degrade tissue quality, making subsequent analyses more difficult and variable. Similarly, rodent brains are typically perfusion fixed through their vascular system, providing rapid and homogeneous fixation of the tissue that is not possible in humans. In contrast, full fixation in humans requires a month or more of immersion in fixative, and fixation time is, therefore, variable across the brain, with deep white matter (WM) regions fixing last. Further, environmental and dietary variability is enormous in humans vis-à-vis rodents, in which living conditions and food supply are tightly controlled. Finally, rodents are typically sacrificed early in their normal lifespan, as opposed to most human tissue that comes to autopsy only at old age or as a consequence of some traumatic or pathological cause of death that can degrade tissue quality. Here, we demonstrate micron-resolution OCT to image individual neurons in human ex vivo tissue at various depths in the first 50 μm, and validate this imaging technique with Nissl staining of the same 50-μm thick tissue. We used layer II of entorhinal cortex (EC) samples of human brain. Images obtained by both modalities (OCT acquisition and digitized Nissl stained sections) were registered using a non-linear transform. Once registered, the degree of overlap was assessed to quantify the agreement between the neuronal content obtained with OCT and with the gold standard Nissl stain.

2 Optical Coherence Tomography

For this study, we used a spectral domain optical coherence tomography/optical coherence microscopy (OCM) that was described in Ref. 22. The broadband light source is provided by a superluminescent diode (LS2000B SLD, Thorlabs Inc., Newton, New Jersey) with a center wavelength of 1310 nm and a full width at half maximum of about 200 nm, which yielded an axial resolution of 4.7 μm in air (3.5 μm in tissue). The spectrometer consisted of a grating and a 1024 pixel InGaAs line scan camera (Thorlabs Inc.), which provides a depth of field of 2.2 mm in air (1.5 mm in tissue) and an axial pixel size of 2.9 μm. In the sample arm, two objectives were used: a 10× water immersion objective (Zeiss N-Achromplan 10× W, NA 0.3) that gives the global laminar structure of the cortex, and a 40× water immersion objective (Olympus LUMPLANFL/IR 40 W, NA 0.8) that allows the imaging of individual neurons. The lateral resolution of the two objectives was 2.5 and 1.25 μm, respectively. Each volume consisting of 1024 frames with 1024 axial scans per frame, was acquired over a field of view (FOV) of 1 mm × 1 mm and 400 μm × 400 μm, respectively, corresponding to a lateral pixel size of 0.97 and 0.39 μm for the 10× and 40×, respectively. To cover the whole sample, the tissue was placed on a manual xy translation stage (Optométrix, 1 in. displacement). The x and y displacements allow for overlap between the volumes in order to reconstruct the sample by stitching the images using a Fiji plug-in based on the Fourier shift theorem.

3 Tissue Samples

Three human brains were obtained from the Massachusetts General Hospital Autopsy Suite (Boston, Massachusetts). The demographics were as follows: mean age 50.3 ± 11.0 y.o., 1M/2F, PMI <24 h, neurologically normal. Each brain was immersed in 10% formalin for at least 2 months until thoroughly fixed. A subregion within the EC of each brain was then blocked in several square millimeters of en-face area (approximately 3 × 5 mm²). The samples were then embedded in melted oxidized agarose and covalent cross-linking between tissue and agarose was activated using borohydride borate solution.

4 Histology Protocol

A vibrotome (TissueCyte 1000, TissueVision), described in Ref. 18, was used first to flatface the samples and subsequently to section 50 μm slices after the imaging over this depth was performed. The agarose from the sections was removed by heating the phosphate buffer above the agarose melting temperature (above 50°C) and rinsing the slices for about 3 s. The sections were then mounted onto gelatin dipped glass slides and stained for Nissl substance, revealing mainly neuronal and glial cells (details in our previous paper). The stained slices were digitized with a camera mounted on an 80i Nikon Microscope (Microvideo Instruments, Avon, Massachusetts) with high magnification (20×), giving a pixel size of 0.37 μm. We used the image series workflow (“SRS Image workflow”) provided by Stereo Investigator (MBF Bioscience, Burlington, Vermont) to automatically mosaic the entire slice, and the tiles were then stitched with the same Fiji plug-in used for the OCT images. Figure 1(a) shows a Nissl stained section of EC where layers I–VI as well as WM were labeled. We confirmed that layer II exhibits large neurons in island formation in our samples.

5 OCT Acquisition

As shown in our previous publication, to obtain the overall laminar structure of the EC using the 10× objective, the average intensity projection over 300 μm in depth from the surface of each volume was performed, filtered and intensity adjusted to enhance the contrast. The images were then stitched together to obtain the full cortical ribbon of the sample. Figure 1 shows the cortical layers in EC that are easily observable with OCT imaging [Fig. 1(b)] and corresponding Nissl [Fig. 1(a)]. Layers exhibit different intensities related to their cellular architecture and myelin content. To evaluate the capability of OCT to accurately resolve individual neurons, we performed high-resolution imaging on layers I–III in EC. The 40× objective provides a depth of focus of about 10 μm. Therefore, at each xy position of the sample, we acquired five volumes, every 10 μm in depth, starting at 5 μm under the surface until the first 50 μm of the sample was imaged. The framed inset in Fig. 2 shows the depth schematic where the dashed lines represent the five different focus depths from which we acquired data. Contrary to what has been reported concerning freshly resected brain tissue or rodent brains, where neurons exhibit a lack of backscattered light (shown as black spots in the images), neurons in ex vivo fixed human brain highly backscatter the light compared to the surrounding tissue, resulting in high intensity (white spots). Thus, for each of the five volumes, the maximum intensity projection (MIP) over 10 μm around the focus plane (hatched regions around the focus planes) was performed to highlight the neurons. Each tile was then filtered and intensity adjusted. For each different depth, the tiles were acquired across the whole region of interest and were stitched using Fiji to provide the full FOV. Figure 1(d) shows an example of a high resolution OCT image corresponding to the Nissl stained slice and lower resolution OCT image. For the full cortical lamina, only one focus depth (25 μm, MIP over 10 μm around the focus) was imaged and presented. Neurons of layers...
Fig. 1 Cortical layers of entorhinal cortex (EC) observed using Nissl staining (a) and average intensity projection of lower resolution optical coherence microscopy obtained with a 10× objective (b). Layers I–IV have been labeled, as well as the white matter. The individual neurons can be observed by OCT using a higher magnification (c). The insets (d-i) show the different structures observed by OCT (e, g, i) and its correspondence in the Nissl stained slice (d, f, h): vessels found in layer III (d, e), neurons of layer II (f, g), and neurons of layer IV (h, i). Myelinated fibers are also visible in the OCT images. Scale bar: 500 μm.

Fig. 2 Images obtained of the same sample at five different focus depths (a-e). Dashed lines of the framed inset represent the different focus. The MIP was performed over the 10 μm around the focus (dashed regions of the inset). (f) Composite image representing the neuron content over 50 μm in depth and (g) the corresponding Nissl stained slice of 50 μm section thickness. Scale bar: 500 μm.
II and IV were visible, as shown in the inset of Figs. 1(g) and 1(i) and correspond to neurons observed on the Nissl stained slices [Figs. 1(f) and 1(h)]. Figures 2(a)–2(e) show the images obtained at the five different depths. Next, the five full images corresponding to each depth were stacked and the MIP in depth was performed to visualize all the neurons contained in the 50-μm thick slices of the Nissl stained image. The resulting OCT image is shown in Fig. 2(f) and the corresponding Nissl stained slice in Fig. 2(g). The modular organization of layer II is clearly visible both in the OCT and the Nissl stained images. The dendrites arising from the neurons are visible on both images. However, on the OCT image, tangential fibers in layer I can also be observed.

6 Registration and Segmentation of Neurons

To assess the colocalization of the neurons between histologically stained slices and OCT images, the images were registered. Histology protocols can suffer from irretrievable distortions such as tears due to the sectioning, shrinking occurring during the drying process, and geometric warping during mounting. In spite of precautions taken to avoid distortions during the tissue processing, artifacts cannot be avoided completely in large human tissue samples. (For animal histology, these distortions can be reduced significantly using perfusion fixation, shorter PMLs and small samples.) For example, in Fig. 1(a) no outright tearing was observed, but note the vessel on the lower left corner. After mounting and drying in the Nissl protocol, the vessel left a gap, which stretched and locally shifted the location of neurons within the slice. OCT acquisition on the blockface does not suffer from such distortions since imaging occurs on the blockface prior to sectioning. A rigid registration is not sufficient to account for these distortions, therefore, we developed an in-house nonlinear registration tool. Corresponding landmarks on the Nissl stained and OCT images were manually selected, and a nonlinear transformation between the landmark points was computed based on the thin-plate splines deformation model using the implementation in the freely available ITK library (National Library of Medicine Insight Segmentation and Registration Toolkit). The computed transformation was applied to the Nissl-stained images, and the warped Nissl images were resampled by linear interpolation into the new coordinates. The accuracy of the registration algorithm with respect to the number of landmarks was then tested. A large number of corresponding neurons found on both modalities of tissue from Figs. 2(f) to 2(g) were manually selected (\(N_{\text{tot}} = 957\)). Figure 3(a) shows the position of the cells on the OCT in green, and on the Nissl stain slice in blue, prior to nonlinear registration. In this example, the shrinkage due to drying and staining in Nissl is visible. \(N\) landmarks were randomly selected to register the remaining points and the mean distance between the corresponding neurons was calculated. For each \(N\) between 5 and 305, the procedure was repeated 100 times to estimate statistics on the distance error between the remaining corresponding landmarks. Figure 3(b) shows the registration results. The mean registration error is <6 μm with \(N = 5\) selected landmarks, and quickly asymptotes to just over 3 μm after 50 landmarks.

Next, the neurons were segmented. The neuronal segmentation in the Nissl-stained images was done using adaptive thresholding based on the implementation provided in the freely available OpenCV library. The thresholds used in the adaptive threshold at \((x, y)\) were the mean value of the \(S \times S\) neighborhood of \((x, y)\) minus a constant \(C (S = 105, C = 45)\) in Nissl-stained images. The segmentation was manually edited to add neurons whose contrast was insufficient to be identified automatically, and to remove glia and vessels that were segmented incorrectly. OCT images reveal more than just neurons. Vessels, dendrites, and possibly axons are also present in OCT images, for example, and exhibit the same kind of contrast. Moreover, the images are noisier than the Nissl stained slices. We first reduced the noise on each of the images obtained at the five different depths using a nonorthogonal wavelet algorithm optimized on a region of the image containing mainly noise (no visible neurons, fibers, or vessels). We then used adaptive thresholding \((S = 105, C = 25)\) to segment the neurons. As we did for the Nissl stained slices, we manually edited each segmentation to remove nonneuronal features and to add missing neurons. Finally, those five segmentations were overlaid to generate the final OCT segmentation corresponding to the 50 μm Nissl stained slice.

7 Results and Discussion

Figure 4 shows the registered Nissl stained slices (left panel), OCT images (center panel), and overlap of the segmented neurons found in layer II, delineated on the Nissl and OCT images by the lines (right panel) for the six tissue samples studied: green for Nissl, red for OCT, and yellow for the overlap. For each brain sample (A, B, and C), two slices were imaged, sectioned,
and stained (1 and 2: two different slices from the same case). The agreement between the cytoarchitecture observed by the traditional Nissl stain and the OCT imaging appears excellent in case A, slice 1 (A1) and 2 (A2), good for slice B1 and case C (C1 and C2), and finally fair for B2. The agreement was visually assessed by CM. Looking closely at the overlay (Fig. 4 (left)), we can see that the shapes of the corresponding neurons are overall visually the same. However, a slight shift is often
observed showing the limit of the registration between OCT images and histological slices that underwent multiple physical transformations (slicing, mounting, and drying).

At least two possible reasons may explain the varied qualitative colocalization across samples. The first explanation lies in the registration of the Nissl stained slice to the OCT image, which depends on tissue integrity, sectioning, and mounting on the glass slides. All the slices but one (B2) were well prepared, with no major distortions observed and homogeneous tissue thickness. We observed that slice B2 was thinner on its right side. As a result, the density of neurons was lower on this side (Fig. 4). This thickness discrepancy within tissue is due to mispositioning of the sample with regard to the vibratome blade plane. In our present protocol, the vibratome is not yet integrated with the OCT rig and is a separate apparatus. The sample, once flatfaced, is moved to the OCT system for the acquisition and returned to the vibratome for the sectioning. When the sample is not placed exactly in the same position relative to the blade (difference in height and/or tilt occurs), the thickness is not uniform. In the future, the vibratome will be integrated with the OCT rig to remove this complication.

The second explanation is the OCT imaging itself. Ideally, we focus the light at the depth of 5 μm for the first image and then move the sample in 10 μm increments until the first 50 μm of the tissue is imaged, corresponding to the stained tissue. At the beginning of each experiment, we visually position the sample so that the light is focused at the surface and then the sample is moved up 5 μm. Exact positioning of the sample is challenging since the axial pixel size is 2.9 μm. After visually noting the discrepancy in the neuronal content agreement between modalities, we decided to evaluate the position of the light focus with respect to the surface. The surface and the focus planes were fitted by a third degree polynomial surface on every volume acquired at the five different focus depths on the region of interest (layer II). For the first theoretical depth (5 μm), the focus plane and the surface plane are too close to clearly differentiate them. We used the four subsequent theoretical depths to evaluate the experimental depth of focus and use a linear regression to assess the first experimental depth. Table 1 shows the results in the last column. Moreover, we used a built-in algorithm (bwlabel) in the MATLAB programming environment to count the number of neurons on the OCT segmentation image (binary image) obtained at the five different depths (neurons on OCT) and the number of those segmented neurons that are corresponding to neurons on the Nissl segmentation image (overlap on OCT). Finally, we used the same algorithm to assess the total number of neurons segmented on the final OCT segmentation (after overlaying the segmentation of the different depths) and the Nissl segmentation. For each slice, Table 1 reports the intended focus depth, the number of neurons on the OCT at that depth, the number of overlapping neurons, and the percentage of overlapping neurons normalized by the total number of neurons in OCT at each depth. Finally, we also noted the total neurons on OCT and Nissl staining in Table 1. The total number of neurons observed on the final image of OCT (for the 50 μm) is underestimated due to the presence of different neurons at different depths. It does not reflect the number of neurons we would obtain by adding the numbers of neurons at each depth. However, this total number of neurons is in excellent agreement with the number of neurons found on the Nissl stained images (with the exception of case B1, which will be explained in the next paragraph). Contrary to OCT,

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<th>Depth (μm)</th>
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<th>Percentage (%)</th>
<th>Real first OCTNissl depth (μm)</th>
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9 Conclusion

In this study, we showed that OCT can discriminate healthy neurons in \textit{ex vivo} fixed human EC. This technique has been validated by the histological Nissl staining. The same 50 \(\mu\)m of tissue was imaged by OCT and then stained with Nissl. The modalities were registered using a nonlinear transformation, neuronomically segmented, and the overlap was quantified. The results showed good colocalization. Moreover, we demonstrate that OCT can discriminate the neurons in depth.

In conclusion, we demonstrated that OCT/OCM is a promising technique to image the postmortem human brain at the level of single neurons. One critical advantage of OCT over Nissl staining is the minimal distortion of tissue, since the block-face tissue is imaged prior to sectioning. OCT paves the way to undistorted, high resolution, 3-D visualization of the cytoarchitecture in the human cortex. We anticipated that OCT can have a far-reaching impact in both basic neuroscience and clinical neuropathology.

Acknowledgments

We acknowledge the National Center for Research Resources (P41-EB001386, U24 RR021382), the National Institute of Biomedical Imaging and Bioengineering (R01EB006758), the National Institute on Aging (AG022381, R01AG008122-22, K01AG028521, R01AG016945-11), the National Center for Alternative Medicine (RC1 AT005728-01), the National Institute for Neurological Disorders and Stroke (R01 NS005285-01, 1R21NS072652-01, 1R01NS070963), the Shared Instrumentation Grants (1S10RR023401, 1S10RR019307, 1S10RR023043) as well as The Autism & Dyslexia Project funded by the Ellison Medical Foundation, and by the NIH Blueprint for Neuroscience Research (5U01-MH093765), part of the multi-institutional Human Connectome Project. In addition, B.F. has a financial interest in CorticoMetrics, a company whose medical pursuits focus on brain imaging and measurement technologies. B.F. interests were reviewed and are managed by Massachusetts General Hospital and Partners HealthCare in accordance with their conflict of interest policies.

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