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Visualizing and mapping the cerebellum with serial optical coherence scanner

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Abstract. We present the visualization of the mouse cerebellum and adjacent brainstem using a serial optical coherence scanner, which integrates a vibratome slicer and polarization-sensitive optical coherence tomography for ex vivo imaging. The scanner provides intrinsic optical contrasts to distinguish the cerebellar cortical layers and white matter. Images from serial scans reveal the large-scale anatomy in detail and map the nerve fiber pathways in the cerebellum and brainstem. By incorporating a water-immersion microscope objective, we also present high-resolution tiled images that delineate fine structures in the cerebellum and brainstem.

Keywords: cerebellum; brainstem; optical coherence tomography; polarization; fiber orientation.

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1 Introduction

In addition to coordination of movement, studies show that the cerebellum participates in other brain activities including certain cognitive function and experience-dependent adaptive process.1–3 Even though the cerebrum occupies the predominant volume in the brain, the cerebellum holds the majority of brain neurons across different species of mammals.4,5 The neurons in the cerebellum are mainly located in the tightly folded cerebellar cortex, which can be further divided into, from outside to inside, molecular, Purkinje cell, and granular layers. Abundant nerve fiber tracts connect the cerebellum with other brain regions through cerebellar peduncles. Diseases involving the cerebellum often lead to motor disturbance and loss of movement coordination.6

Optical coherence tomography (OCT) produces depth resolved images (cross sections) of tissue structures at the micrometer scale resolution.3 In recent years, OCT for brain imaging has drawn attention because of the increasing research interest in brain microstructure and circuitry. In addition to the visualizations of cortical layers and neurons,8–11 OCT has been used to study pathological changes in brain conditions such as brain tumors,12–14 edema,15 and seizures.16 OCT has also been shown to separate different layers in the cerebellar cortex17,18 and Purkinje cell bodies11 in the cerebellum.

Reflectivity, the conventional contrast for OCT and confocal microscopy, may not be descriptive enough to delineate different structures in complex tissue. Polarization-sensitive OCT (PS-OCT)19 provides additional contrasts that originate from tissue birefringence. It distinguishes white matter and gray matter, and visualizes fiber tract in the brain.20 PS-OCT highlights the myelinated and birefringent nerve fibers with the retardance contrast and indicates the in-plane fiber alignment with the axis orientation contrast. To facilitate large-scale imaging at microscopic resolution, a tissue slicer has been integrated with PS-OCT to form the serial optical coherence scanner (SOCS).21 Brain imaging with SOCS showed that the reflectivity contrast portrays morphology, and the polarization contrasts primarily probe myelinated nerve fibers in the white matter. Architecture of nerve fibers delineated by retardance and the in-plane fiber orientations quantified by optic axis orientation have been covalidated with diffusion MRI.22

In this study, we used SOCS to visualize and map the cerebellum and nearby brainstem areas of ex vivo mouse brains. Reflectivity- and polarization-based intrinsic contrasts are utilized for coronal or sagittal sections. Stacking several sections allows reconstruction of a large three-dimensional (3-D) volume, in which the global anatomy is visualized and the nerve fiber pathways are tracked. By incorporating a water immersion microscope objective, we also present higher resolution tiled images that delineate fine structures in the coronal and sagittal planes. The movements of stage for imaging or slicing steps are computer controlled, and the entire process is semiautomated. The results show that SOCS is an efficient tool to image rodent cerebellum at microscopic resolution and it offers a viable technique for studying cerebellar diseases.

2 Methods

2.1 Sample Preparation

Adult C57BL/6J wide type mice were euthanized and transcardially perfused by 10% buffered formalin. All animal treatments and experiments are in accordance with the Institutional Animal Care and Use Committee at the University of Minnesota approved protocols. Dissected brains were kept in 10% buffered formalin for 48 h prior to imaging. Only the cerebellum and adjacent brain stem region were scanned by SOCS. Samples were imaged in coronal and sagittal sections. After imaging the superficial region of a sample, a 200-μm slice was removed by the vibratome slicer to access the next section.
2.2 Optical Setup

SOCS\textsuperscript{21} integrates polarization maintaining fiber (PMF)-based spectral domain PS-OCT\textsuperscript{23} and a vibratome tissue slice. To obtain absolute optic axis orientation, calibration is included for dynamic removal of a phase offset.\textsuperscript{24} The light source is a 25-mW superluminescent diode with 840-nm center wavelength and 50-nm FWHM bandwidth, yielding an axial resolution of 5.5 μm in tissue (refractive index: ~1.4). Linearly polarized light is coupled into one of the PMF channels in a fiber bench and split into reference and sample arms by a 2 × 2 PMF coupler. Briefly, light in the reference arm (not shown) passes through a quarter-wave plate (QWP) whose optic axis is 22.5 deg with respect to the incoming polarization state. As a result, light returning from a reflector passes through the QWP one more time and becomes linearly polarized at 45 deg. This ensures equal powers propagating in the orthogonal channels of PMF toward the coupler, where interference of light returning from the reference and sample arms occurs. Details of the optical setup including the spectrometer design, operation, and performance can be found in Ref. \textsuperscript{23}.

Figure 1 shows the sample arm configuration. Linearly polarized light emerging from PMF is collimated by a lens (f = 11 mm). Light passes through a beam sampler, which diverts a small portion of the beam toward the calibration unit (dashed box) consisting of a polarizer and a mirror. After passing a QWP with optic axis set at 45 deg with respect to the incoming linear state, light becomes circularly polarized. Galvanometer controlled scanning mirrors provide beam scanning in two lateral dimensions. A beam expander (1.6×) consisting of two achromatic lenses is positioned between the scanning mirrors to pivot the light at the back focal plane of a scan lens or a microscope objective and to increase the effective numerical aperture. This system is capable of imaging samples using two different schemes. Solid boxes in Fig. 1 show switchable parts for selecting either a scan lens (LSM03-BB 5×, Thorlabs, Inc., Newton, New Jersey) or a water immersion objective (UMPLFLN 10× W, Olympus, Co., Tokyo, Japan) with another 1.6× beam expander, respectively. For the latter, the required space can be gained by removing the vibratome or by raising the height of the main platform. The FWHM lateral resolution is 10 μm with the scan lens or 4 μm with the objective. Imaging with high NA microscope objectives relates to optical coherence microcopy that can also utilize polarization sensitivity for high-resolution brain imaging.\textsuperscript{25} In our setup, the optical path of the reference arm is also adjustable to accommodate the path length change due to switching between the scan lens and the objective. Light reflected/backscattered within a birefringent sample can be represented by an elliptical polarization state that results in variable light powers coupled back into the PMF channels. In the coupler, light from the reference and sample arms interfere in the corresponding PMF channels.

2.3 Contrasts and Image Formation

Spectra from two orthogonal polarization channels are focused side-by-side onto a single line-scan camera in a custom spectrometer. Acquired spectra are resampled in linear k-space. Dispersion compensation is performed in software.\textsuperscript{26} Inverse Fourier transform of interference-related spectral oscillations yields complex depth profiles that are in the form of $A_{1,2}(z)\exp\{i\phi_{1,2}(z)\}$, where $A$ and $\phi$ denote the amplitude and phase as a function of depth $z$ and the subscripts represent the polarization channels.\textsuperscript{23} Reflectivity, $R(z)$, phase retardance, $\delta(z)$, and relative axis orientation, $\theta'(z)$, information of the depth profile (A-line), are obtained by

\begin{align*}
R(z) &\propto A_1(z)^2 + A_2(z)^2, \\
\delta(z) &\propto \arctan \frac{A_1(z)}{A_2(z)}, \\
\theta'(z) &\propto (\phi_1(z) - \phi_2(z))/2 + \phi_o,
\end{align*}

where $\phi_o$ is an arbitrary offset due to PMF and it is potentially time-varying with fiber movement and temperature change. $\phi_o$ is measured from the interference of the calibration paths (the path in reference arm is not shown), which is positioned below the imaging area dedicated to the tissue sample. Absolute optic axis orientation is computed by subtracting the offset from the orientation measurement, $\theta(z) = \theta'(z) - \phi_o$.\textsuperscript{24} Correspondingly, the in-plane orientation of white matter tracts is determined by dynamic calibration.

A cross-sectional image is formed by stacking several A-lines acquired during a lateral scan (x-axis). Multiple cross sections are acquired during raster scan to construct a 3-D volume. \textit{En-face} images are two-dimensional (2-D) projections of the 3-D datasets onto the $xy$-plane. The \textit{en-face} images can be generated from different contrasts in a number of ways. For example, the \textit{en-face} reflectivity image can get its pixel values from the average reflectivity values within the depth profiles. It is useful to present the anatomical structures in 2-D for the imaged section. Likewise, the \textit{en-face} retardance images are produced by averaging to delineate the nerve fiber tracts. A median filter is applied to the cross-sectional image prior to averaging in $z$-direction. For the \textit{en-face} orientation image, absolute optic axis orientation values within each A-line are binned into 5 deg intervals, and the mean value of Gaussian fitted histogram determines the pixel value. Axis orientation values are color coded in Hue-saturation-value (HSV) space as illustrated by a color wheel, while the brightness of colors is regulated by the \textit{en-face} retardance values that highlight the white matter regions.
Light intensity in tissue decays exponentially due to scattering and absorption events. Since cross-sectional reflectivity images are typically presented in logarithmic scale, the attenuation corresponds to a linear decay for uniform samples. Therefore, the values in an en-face attenuation image are calculated by a least-square first-order polynomial fitting of the logarithmic reflectivity signal within the depth profiles. The en-face birefringence image is produced by taking least-square first-order polynomial fitting of the retardance signal, as the derivative of measured retardance yields the apparent birefringence. Depth range used in calculations matches the physical slice thickness.

Open source image processing tool Fiji was used to produce composite images and stitch tiles for high-resolution imaging. Anatomical identifications are carefully compared with Allen mouse brain atlas. Abbreviations of anatomical structures also match with Allen mouse brain atlas.

### 2.4 Serial imaging

Serial imaging is semiautomated. A custom program written in LabView (National Instruments, Austin, Texas) controls linear actuators (T-NA series, Zaber Technologies Inc., Vancouver, British Columbia, Canada) of a one-inch three-axis translation stage, which accomplishes the repositioning of the sample. The actuators have 8 mm/s maximum speed, <1-μm repeatability and <4-μm backlash. As shown in Fig. 1, separate positions for imaging and slicing could be a requirement to avoid the slicer blade interfering with the sample arm optics. The vibratome is manually controlled. The workflow of the imaging/slicing procedure is described as follows.

1. User initially determines and stores the imaging and slicing locations.
2. The sample is transported to the imaging location.
3. After imaging, the sample is transported to the slicing position.
4. The vibratome removes the imaged portion to expose the deeper regions for imaging.
5. Steps 2 to 4 are repeated until the whole block is imaged.

The optical imaging for a single-volumetric scan takes 25 s when 500 cross sections each with 1000 A-lines are acquired with an A-line rate of 20 kHz. However, the imaging and slicing procedure for a section took 4 min due to a low speed of the vibratome blade to maintain slicing quality. The raster scans covered 10 mm × 8 mm for coronal sections and 7.5 mm × 8 mm for sagittal sections. Consequently, the A-line spacing was 10 μm and 7.5 μm for coronal and sagittal cross sections, respectively. Spacing between the cross sections was 16 μm, which is close to the ~17 μm 1/e² beam diameter in the field of focus that the scan lens provides.

After completing SOCS imaging for an entire sample, the scan lens and the vibratome were removed. Then, the water-immersion microscope objective as shown in Fig. 1 was incorporated for imaging selected slices at high lateral resolution. A raster scan in this case covered an area of 1.25 mm × 1.25 mm. In order to cover the entire lateral plane, several locations with 5% overlapping region had to be scanned. Each scan included 500 cross sections, each of which contained 500 A-lines. The computer controlled the procedure with the software moving the actuators between the optical scans. High-resolution smaller-scale images are then stitched together to create a tiled image of the tissue section.

### 3 Results

#### 3.1 Differentiation of Anatomy from Cross-Sectional Images

Figure 2 shows cross-sectional reflectivity and retardance images containing the cerebellum, the fourth ventricle (V4), and the brainstem, and compares the contrasts in anatomical regions. The scan for the reflectivity [Fig. 2(a)] and retardance [Fig. 2(d)] images was taken from the midline region of a coronal section which is at about Bregma −6.1 mm as compared with the stereotaxic coordinates of the mouse brain. Scale bars in Fig. 2(a) represent 1 mm in x-axis and 0.1 mm in z-axis. A color-coded bar is shown to label different anatomy. Molecular layer (ML, blue), granular layer (GL, green), and white matter (WM, red) regions are indicated by color-coded boxes above the images. The thin Purkinje cell layer is not visualized in these cross-sectional images. Comparisons in the subplots also share the same color-coding for these anatomical structures.

We calculated and compared the attenuation of the OCT signal in cerebellar ML, GL, and WM. Twenty A-lines were chosen from each of these regions, and the reflectivity signals in dB scale were averaged and fitted with a least-square first-order polynomial function. Figure 2(b) shows the averages (dots) and fitted lines as a function of depth. The attenuations of the OCT signal in the ML, GL, and WM region are found to be (3.3 ± 0.8) × 10⁻² dB/μm, (4.3 ± 1.1) × 10⁻² dB/μm, and (10 ± 1.1) × 10⁻² dB/μm, respectively. Clearly, light in the white matter region decayed faster than the layers. Attenuation for each A-line was also calculated. Figure 2(c) shows the mean and standard deviation of these attenuations in the aforementioned regions. Unpaired two-sample student t-tests were performed for every pair of the three regions. Attenuations in the cerebellar ML, GL, and WM are all significantly different from each other (p < 0.001). This suggests that the attenuation map can be useful for delineating the two cerebellar cortex layers, and at the same time differentiating the white matter from the gray matter.

We performed a similar comparison with the retardance contrast, which is represented as phase retardance [Eq. (2)]. Figure 2(e) shows the averaged retardance values as a function of depth. Averages were calculated for each region using the same twenty A-lines chosen earlier. Figure 2(e) also shows the slopes of retardance (apparent birefringence) by line fitting. The white matter region clearly exhibits the largest birefringence in the group. Calculated from each A-line, the apparent birefringence values are given in Fig. 2(f) as (2.6 ± 1.9) × 10⁻² deg/μm, (5 ± 2.9) × 10⁻² deg/μm, and (14 ± 3.1) × 10⁻² deg/μm for the ML, GL, and WM regions, respectively. Statistical comparison demonstrates the slope of retardance for the WM is significantly different from two cerebellar cortex layers (p < 0.001). The slopes of two cerebellar cortex layers were also significantly different from each other but with a lower p-value (p < 0.01). The difference between the attenuation and birefringence values in GL and ML can be attributed to the anatomy of the cerebellar cortex, as GL is densely packed with small granule cells and contains...
myelinated axons. WM is composed of bundles of myelinated axons, and therefore, the optical properties of myelin is associated with the results.

3.2 Visualizations with En-Face Coronal Images

Figure 3 shows en-face images of a coronal section. Comparing the images with the stereotaxic coordinates of the mouse brain, the section is located at about Bregma −6.3 mm. Images are derived from the reflectivity, retardance, and axis-orientation contrasts as described in Sec. 2.3. Figure 3(a) shows the en-face reflectivity image that shows the gross anatomy. The GL can be distinguished from the ML and white matter tracts. The scale bar represents 1 mm and the labels of anatomical structures are given only in this image. Figure 3(b) shows the en-face retardance image that highlights the white matter connections in the section. Figure 3(c) shows a composite image with the green and red channels representing the reflectivity and retardance, respectively. The brightness is regulated by retardance values. Abbreviations: arbor vitae (arb), medial longitudinal fascicle (mlf), interior cerebellar peduncle, (icp) and spinal tract of the trigeminal nerve (sptV). Scale bar: 1 mm.
Figures 3(a) and 3(b) show the attenuation and apparent birefringence maps, respectively. Figures 3(d) and 3(e) show the \textit{en-face} axis orientation image with orientation values given in the color wheel. The axis orientations of the branches of arbor vitae (arb) agree with the anatomical geometry.

The medial longitudinal fascicle (mlf) in the midline of medulla is visualized in Figs. 3(a), 3(b), and 3(f). The inner interior cerebellar peduncle (icp), and the outer spinal tract of the trigeminal nerve (sptV) are adjacent to each other in medulla. However, in the coronal section, the inclination angle of sptV is higher than icp. Measured reflectivity, retardance [Figs. 3(a) and 3(b)], and their slopes [Figs. 3(d) and 3(e)] for the sptV are lower than the ones for the icp region. The high inclination angle of the sptV also reduces the apparent birefringence and makes the axis orientation measurement unreliable.

By replacing the scan lens with the water immersion microscope, fine structures in the cerebellum and brainstem are delineated at high resolution. The coronal section, previously imaged with the scan lens (Fig. 3) and removed by the slicer, was positioned under the objective.

Figure 4 shows the tiled image consisting of smaller-scale high-resolution images and two selected regions to highlight structures with different contrasts. The composite image in Fig. 4(a) is created by merging the \textit{en-face} reflectivity (green) and retardance (red) images. Anatomical structures are presented in remarkable details. Two regions of interest (ROI) enclosed by white dashed lines are further investigated. Figures 4(b)–4(d) show fine structures in ROI-1 with the \textit{en-face} reflectivity, retardance, and optic axis orientation images. In Fig. 4(b), three layers in cerebellar cortex are clearly visualized with the Purkinje cell layer (arrow) appearing as an interface between GL and ML. Compared to the arb region, the cerebellar nuclei (CBN) region has less white matter and consequently appears with reduced reflectivity and retardance. Medulla oblongata in ROI-2 [Figs. 4(e)–4(g)] consists of abundant nuclei. Reflectivity contrast [Fig. 4(e)] shows a lattice pattern. The optical contrasts show the characteristics of icp and sptV labeled in Fig. 4(e)] in the high-resolution images.

### 3.3 Visualizations with En-Face Sagittal Images

Twenty sagittal sections were imaged serially with the scan lens. The sagittal section representing the \textit{en-face} plane starts at lateral $-2.8$ mm in the left hemisphere, and reaches lateral $1$ mm across the midline into the right hemisphere. A portion of the scan (10%, or 100 A-lines in $x$-direction) was removed from the images as it covered agarose gel supporting the tissue during slicing. As a result, the \textit{en-face} images represent an area of 6.75 $\times$ 8 mm$^2$. The \textit{en-face} images were stacked together to present the anatomy in the cerebellum and brainstem. The effective volume covered with the scans is 6.75 $\times$ 8 $\times$ 4 mm$^3$ in $xyz$-directions. Video 1 presents the composite \textit{en-face} images of reflectivity (green) and retardance (red). Video 2, on the other hand, shows the fiber axis in the plane with the axis orientation images. Each frame in the videos represents a 200-$\mu$m thick section. Scale bar is 1 mm in the videos and in Fig. 5, which shows two representative sections.

The sections shown in Fig. 5 are located at the left hemisphere at lateral $-2.2$ mm [Figs. 5(a) and 5(b)] and lateral $-0.6$ mm [Figs. 5(c) and 5(d)] from the midline. and lateral from the midline, respectively. Arrangement and axis orientation of the branching arb tracts are visualized in the cerebellum. Plentiful nerve tracts in the brainstem are observed since the retardance contrast highlights these connections. The spinal trigeminal nucleus [SPN in Fig. 5(a)] demonstrates lower retardance. In contrast, the surrounding fiber tracts (mainly vestibulocochlear nerve and sptV) appear red due to high birefringence. The facial nerve (vII) is clearly visualized in Fig. 5(c) and its position changes from interior to anterior in the videos. In Figs. 5(c) and 5(d), the sptV fibers run along the rostral edge of medulla oblongata. Several major fiber tracts such as middle cerebellar peduncle and pyramid in corticospinal tract, align with the ventral edge of medulla. Figures 5(b) and 5(d) show the in-plane axis orientations of the tracts.

After imaging the entire sample, the scan lens was replaced by a water-immersion microscope objective for high-resolution imaging of one of the sagittal section presented in Figs. 5(c) and 5(d). Figure 6 shows the tiled image and two ROIs for showing
the details. Figure 6(a) is the composite image of reflectivity and retardance contrasts. ROI-1 [Figs. 6(b)–6(d)] demonstrates the tightly folded cerebellar cortex and several arbor tracts with the en-face reflectivity, retardance, and axis orientation images. All three layers in the cerebellar cortex are clearly presented in Fig. 6(b). The arrow indicates the darker Purkinje cell layer [Fig. 6(b)]. ROI-2 [Fig. 6(e)–6(g)] visualizes the fiber tracts in the brainstem. Retardance image [Fig. 6(f)] captures vIIn, which is located between the pons and medulla, even though the high inclination angle reduces the reflectivity and retardance contrasts.

4 Discussion

Volumetric large-scale optical brain imaging at high resolution is challenging due to light scattering and limited field of view of focal lens. Recent techniques for mouse brain imaging include micro-optical sectioning tomography,32 two photon tomography,33,34 light sheet microscopy,35,36 and SOCS.21 Among these, SOCS is a label-free imaging technique and it is suitable for ex vivo imaging of the entire brain. One of the limitations is the depth of view, which may be improved to millimeter scale by using a light source operating in the longer wavelength region31 or by incorporating a tissue clearing method. The latter may reduce the contrasts. Enhancement in imaging depth could reduce the number of physical sections required for reconstructing the entire brain. The other limitation is the slicer speed to maintain good slicing quality. The system we report here is semiautomated and it only requires the user to press buttons to realize tissue slicing and data acquisition. Additional time can be saved with a fully automated system that controls the vibratome functions electronically. The current system is sufficient for a rodent brain. Imaging larger brains like primate brains is feasible, but it would require a bigger blade and longer-range stages and actuators.

We can incorporate microscope objectives into SOCS to enhance lateral resolution; however, reduction in the field of view requires tile scans to cover the entire sample. The short working distance under the objective may necessitate adjustments and modifications on the stages. Moreover, the tight focus reduces the confocal parameter. Its effect for 3-D reconstruction of the sample could be avoided by shifting the focus for consecutive imaging or reducing the physical slice thickness that would increase the recording time. Quantification of attenuation would also be challenging with the microscope objective since the rapidly changing beam profile needs to be decoupled from

Fig. 5 SOCS images of two representative sagittal sections. Composite images in (a) and (c) are with reflectivity (green) and retardance (red) contrasts. Axis orientation images in (b) and (d) are with colors in HSV space (see the color wheel) and the brightness of pixels is regulated by retardance values. Abbreviations: spinal trigeminal nucleus (SPN), facial nerve (vIIn). Scale bar: 1 mm. See Video 1 and Video 2 that present twenty sagittal sections with the composite and axis orientation images, respectively. Scale bars: 1 mm. [Video 1, MOV, 1.92 MB [URL: http://dx.doi.org/10.1117/1.NPh.4.1.011006.1]; Video 2, MOV, 2.54 MB [URL: http://dx.doi.org/10.1117/1.NPh.4.1.011006.2]]

Fig. 6 High-resolution en-face images of a sagittal section. (a) Composite image of reflectivity (green) and retardance (red). Dashed boxes represent ROIs. (b) and (e) Reflectivity, (c) and (f) retardance, and (d) and (g) axis orientation images show these contrasts for the fine structures in ROI-1 and ROI-2, respectively. Arrow in (b) indicates Purkinje cell layer. vIIn in (e) denotes the facial nerve. (d) and (g) Axis orientation is color coded in HSV space by the color wheel, and the brightness is regulated by retardance values. Scale bars in (a) and (b): 1 mm.
the measurement and the focal plane may be hard to define accurately in the images. Compared with sptV in sagittal sections, the sptV fibers in coronal sections may have misinterpretation in reflectivity, retardance, and axial orientation contrasts due to high inclination angle. Related to this, the axial orientation measurement in this study is defined in a plane (xy) orthogonal to the direction of light propagation (z). However, the description of nerve fiber axis orientation in 3-D requires both azimuthal and polar angles. Computational methods and experimental methods utilizing measurements with variable incident angle have been proposed to retrieve 3-D axis using PS-OCT. Quantification of the fiber inclination would allow calculation of the true birefringence Δn, since the measured apparent birefringence can be written as Δn′ = Δn cos² α, where α is the inclination angle. Combining the information from variable incident angles after registering the 3-D datasets has potential to improve the image quality and representation of tissue anisotropy.

5 Conclusion

SOCs provides an efficient approach to visualize and map the mouse cerebellum and nearby brainstem. The cerebellar layers, abundant nuclei in brainstem, and nerve fiber tracts are presented for coronal and sagittal sections at two different resolutions. Several contrasts within the anatomical structures are quantified and used in the visualizations. SOCs has the potential to study the diseases of the cerebellum and brainstem including ataxia and medulloblastoma and opens new perspectives to better understand the anatomy and pathology.

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References


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