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Multi-modal imaging of a single mouse brain over five orders of magnitude of resolution

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Abstract

Mammalian neurons operate at length scales spanning six orders of magnitude; they project millimeters to centimeters across brain regions, are composed of micrometer-scale-diameter myelinated axons, and ultimately form nanometer scale synapses. Capturing these anatomical features across that breadth of scale has required imaging samples with multiple independent imaging modalities. Translating between the different modalities, however, requires imaging the *same* brain with each. Here, we imaged the same postmortem mouse brain over five orders of spatial resolution using MRI, whole brain micrometer-scale synchrotron x-ray tomography (μ CT), and large volume automated serial electron microscopy. Using this pipeline, we can track individual myelinated axons previously relegated to axon bundles in diffusion tensor MRI or arbitrarily trace neurons and their processes brain-wide and identify individual synapses on them. This pipeline provides both an unprecedented look across a single brain's multi-scaled organization as well as a vehicle for studying the brain's multi-scale pathologies.

Declaration of Competing Interest

The authors declare no competing or conflicts of interest.

Ethics statement

The following ethics statement is in the 2. Materials and Methods subsection, 2.1 Sample Preparation:

All procedures performed on animals followed protocols approved by the Institutional Animal Care and Use Committee and were in compliance with the Animal Welfare Act and the NIH Guide for the Care and Use of Laboratory Animals.

Code availability

All image processing was performed with commonly used freeware tools, noted throughout.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2021.118250.

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Authors' contribution

S.F., P.LR., and N.K. conceived and designed the imaging pipeline. S.F. acquired and processed MRI data. V.DA. designed the μ -CT system used for data collection and contributed in reconstruction of μ -CT data. V.S. collected all EM data. S.T. registered MRI, μ CT, and EM data. S.F., A.S., and K.N. traced/segmented μ CT data and S.F. registered results to MRI data. S.F. and N.K. wrote the final manuscript and all authors contributed to its revision.

Keywords

MRI; Synchrotron source x-ray tomography; Electron microscopy; Whole mouse brain; Connectomics; Multi-scale imaging; Echo planar spectroscopic imaging

1. Introduction

The architecture of the mammalian brain is complicated, operating on multiple scales spanning orders of spatial magnitude (Lichtman and Denk, 2011). Neurons in the brain project macroscopic distances(millimeters to centimeters) (S.R., 1899), while simultaneously producing microscopic connections and plasticity-related changes at the nanoscale (Grutzendler et al., 2002).

No current single existing imaging modality spans the six orders of length-scale (mm to nm) required to map the whole brain while simultaneously resolving individual synapses. Magnetic resonance imaging (MRI) allows in vivo (Basser et al., 2000) and postmortem (Dyrby et al., 2007; Miller et al., 2011) mapping of neuronal tracts spanning macroscopic distances but cannot easily observe individual neurons much less identify individual neuronal connections. At the other end of the resolution scale, advances in automated electron microscopy (EM) provide synapse-level resolution (3 nm) (Helmstaedter, 2013) but remain limited by serious computational challenges; a single voxel of MRI data (~1mm³) is almost 2 million terabytes of raw EM image data (Bouchard et al., 2016). Thus, EM remains best suited for neuron 'connectomics' — reconstructing connections of individual neurons over limited ranges of 100s of micrometers. An imaging pipeline capable of bridging these approaches would have to solve the problem of 'contex' by providing sufficient information and volume to help register an EM dataset to the relative low-resolution framework of an MRI volume. An ideal intermediate imaging modality would (a) achieve micrometer-scale resolution throughout an entire large volume tissue sample and (b) use a sample preparation compatible with a pipeline including MRI and EM.

Synchrotron-based x-ray tomography (μ CT) nondestructively provides micrometer-scale resolution over ~1cm³ whole tissue volumes (the size of a small mammalian brain), bridging the resolution disparity between MRI and EM imaging techniques (Fig. 1). Sample preparation for μ CT uses aldehyde fixed material, compatible with postmortem MRI (Shepherd et al., 2009), followed by osmium staining and plastic embedding, compatible with large volume EM (Dyer et al., 2017). Thus, MRI followed by μ CT and EM provides a viable pipeline for multi-scale whole brain imaging. In this work, we demonstrate the feasibility of such a pipeline by imaging a fixed postmortem mouse brain with MRI, preparing the whole brain for subsequent μ CT and EM (Mikula et al., 2012; Mikula and Denk, 2015), re-imaging the entire brain with micrometer-resolution μ CT (Dyer et al., 2017; Vescovi et al., 2018), and, finally, imaging isolated sub-volumes with automated serial EM (Kasthuri et al., 2015).

2. Materials and methods

2.1. Sample preparation

All procedures performed on animals followed protocols approved by the Institutional Animal Care and Use Committee and were in compliance with the Animal Welfare Act and the NIH Guide for the Care and Use of Laboratory Animals. An eight week old, female C57bl6 mouse was deeply anesthetized with 60 mg/kg pentobarbital. The mouse was transcardially perfused with a solution (pH 7.4) of 0.1 M sodium cacodylate and heparin (15 units/ml) immediately followed by a solution of 2% paraformaldehyde, 2.5% glutaraldehyde, and 0.1 M sodium cacodylate (pH 7.4). The brain was carefully removed from the skull and post-fixed in the same fixative for 48 hours at 4°C. The order of serial imaging was MRI, μ CT, and then EM.

- **2.1.1. MRI**—The resected and aldehyde-fixed brain was washed in PBS for 72 hours to remove fixative from the tissue, which can be deleterious to MRI imaging (Dyrby et al., 2007; Shepherd et al., 2009). Just prior to imaging, brains were dried of excess PBS and placed in 10 ml falcon tubes. Tubes were filled with Fluorinert (FC-3283, 3M Electronics) for susceptibility matching and to improve shimming.
- **2.1.2. µCT and electron microscopy**—After MRI, the whole brain was transferred to 0.1 M cacodylate buffer. Whole brain staining was performed with multiple rounds of osmium and reduced osmium followed by dehydration and plastic embedding (Mikula et al., 2012; Mikula and Denk, 2015).

2.2. Acquisition protocols

2.2.1. MRI—Data were acquired at 9.4 Tesla (20 cm internal diameter, horizontal bore, Bruker BioSpec Small Animal MR System, Bruker Biospin, Billerica, MA) using a 6 cm high performance gradient insert (maximum gradient strength: 1000 mT/m, Bruker Biospin) and a 35 mm internal diameter quadrature volume coil (Rapid MR International, Columbus, Ohio). The brain was aligned such that the anterior/posterior portion of the olfactory limb of the anterior commissure (AC) was approximately parallel to the magnetic field of the scanner and the hemispheric midline was parallel to the scanner YZ plane.

Using the Paravision mapshim protocol, third order shimming was iteratively performed over an ellipse that encompassed the entire brain, but did not extend beyond the boundaries of the falcon tube/Fluorinert interface. B0 maps were produced by recording the voxel-wise frequency of the peak of the resonance, including additional sub-spectral resolution frequency produced by estimating the maximum peak amplitude of the resonance, described below. This was consistent with previously reported work (Foxley et al., 2015; Foxley et al., 2018), which described a high degree of field homogeneity across samples.

DTI was performed using a standard diffusion-weighted 3D spin echo sequence (TR = 400 ms, TE = 18.5 ms, b value = 3000 s/mm^2 , $\delta = 5 \text{ ms}$, = 11.04 ms, spatial resolution = 150 um isotropic, number of b0s = 16, number of non-collinear directions = 30, receiver bandwidth = 200 kHz, partial Fourier along first phase encoding direction = 7/8, duration = 55 hrs 19 min 40 sec).

Structural MRI data were acquired using a multi-echo gradient echo sequence. This produces a 4D dataset, where data are sampled along time in each spatial voxel to acquire the free induction decay (FID). Sequence parameters were chosen so that the entire voxel-wise FID was sampled to the noise floor (TR = 1000 ms, TE of first echo = 2.74 ms, echo spacing = 2.74 ms, number of echoes = 192, receiver bandwidth = 75 kHz, flip angle = 68° , $100 \mu \text{m}$ isotropic resolution, 4 averages, duration = 12 hours).

2.2.2. μ CT—The μ CT data were acquired at the 32-ID beamline at the Advanced Photon Source, Argonne National Laboratory. The setup consists of a 1.8 cm-period undulator operated at a low deflection parameter value of K=0.26. This yields a single quasi-monochomatic peak of energy 25 keV without the losses incurred by use of a crystal monochromator. For a sample 68 m from the undulator, this produces a photon fluence rate of about 1.8×10^7 photons s⁻¹ μ m⁻².

The x-rays were imaged using a 10 μ m thick thin-film LuAG:Ce scintillator producing visible-light images then magnified using a 5X Mitutoyo long working distance microscope objective onto a 1920 \times 1200 pixel CMOS camera (Point Gray GS3-U3–51S5M-C). The effective object space pixel size is 1.17 μ m 2.25 \times 1.41 mm² field of view. The thickness of the thin-film scintillator matched the depth of focus of the objective lens, achieving a spatial resolution equivalent to the resolving power of the lens (1.3 μ m for a NA of 0.21). Since the camera field of view is substantially smaller than the mouse brain, a mosaic strategy was employed (Vescovi et al., 2018).

The sample was mounted on an air-bearing rotary stage (PI-Micos UPR-160 AIR) with motorized x/y translation stages located underneath and x/y piezo stages on top. Typical exposure time for a single projection image at one mosaic grid point and one rotation angle was 30 ms. 3600 rotation angles were used at each grid point. The sample was translated through a 6 ×18 tomosaic grid.

2.2.3. Electron microscopy—Regions of interest (ROI) in the brainstem and cerebellum were identified in the μ CT data. The sample block was trimmed to each ROI and 50 nm thick serial sections were collected on aluminium tape, attached to a wafer (Kasthuri et al., 2015), and imaged in an Apreo scanning electron microscope (SEM) (ThermoFisher Scientific). Images were obtained with either 135 nm or 3 nm in-plane resolution with 3 μ s dwell times.

2.3. Data processing and analysis

2.3.1. MRI—Data were reconstructed, processed, and analyzed with IDL 8.2 (ITT Visual Information Solutions, Boulder CO), Matlab 2014b (The Math-Works Inc., Natick, MA, 2012), and FSL 5.0.9 (FMRIB Software Library, FMRIB, Oxford, UK).

3D multiple-gradient echo data were reconstructed to produce voxel-wise water spectra. Each complex 4D dataset $(x \times y \times z \times t)$ was Fourier transformed along time to produce three spatial dimensions and one spectral dimension $(x \times y \times z \times v)$. A T2*-weighted structural image was constructed from the spectral data, where image contrast was produced by isolating the maximum voxel-wise signal amplitude of the water spectrum (Al-Hallaq

et al., 2002). This was achieved by identifying and storing the voxel-wise maximum peak signal amplitude in a 3D array with the same spatial dimensions as the acquisition. The mean $b=0 \text{ s/mm}^2$ dataset from the diffusion acquisition was registered to the structural image via affine transformation using the FSL linear registration tool (FLIRT) (Jenkinson and Smith, 2001). Subsequent diffusion data were identically registered using the resultant transformation matrix. Principal, secondary, and tertiary diffusion directions were estimated using BEDPOSTX (Behrens et al., 2003).

2.3.2. μ CT—The acquired sub-sinograms were registered with sub-pixel shifting and stitched through pyramid blending into a complete sinogram (Adelson et al., 1984). The center of rotation of the composite sinogram was estimated using an entropy-based approach (Donath et al., 2006). Reconstruction of the entire dataset in whole-block mode was performed using the gridrec implementation of direct Fourier reconstruction (Dowd et al., 1999; Tsai et al., 2019) implemented in the TomoPy package (Gursoy et al., 2014) resulting in a $11.7 \times 11.7 \times 17.7$ mm³ reconstructed image volume with 1.17μ m isotropic voxels. The reconstructed volume was trimmed and stored in 8-bit format.

Following reconstruction, µCT data were visualized using Neuroglancer (https://github.com/google/neuroglancer), a high-performance WebGL based viewer for large volumetric datasets. Neuroglancer allows for memory-efficient loading of cross-sectional displays across arbitrary planes in the data. This allowed for simple, fine-tuned manual localization of the visible structures in the EM sections within the µCT dataset, using large blood vessels and neuronal somas as landmarks. Our µCT data can be publicly viewed online through Neuroglancer (http://neuroglancer-demo.appspot.com/) using the tinyurl link http://tinyurl.com/cxmbjy6b as a significantly shortened, precomputed source link URL to the general 'karturilab' Neuroglancer reconstruction of the data.

All manually labeling of somas and myelinated axons was performed using Knossos (Helmstaedter et al., 2011).

3. Results

3.1. Whole brain imaging with µCT

Previous work has shown the compatibility of MRI and EM in postmortem tissue samples (Liu et al., 2011). In order to assess the feasibility of utilizing μ CT as a compatible bridge between the two, we developed a protocol for imaging entire mouse brains (1 cm³) at micrometer resolution with μ CT. This included advances in whole mouse brain osmium staining and tissue embedding (Mikula et al., 2012; Mikula and Denk, 2015), extending the field of view of the μ CT system with tomographic imaging (Vescovi et al., 2018), and parallelization of the μ CT reconstruction algorithms using national lab supercomputers (Argonne National Laboratory) for creating contiguous image stacks from the ~10 terabyte sized raw 'sinogram' data. These developments allowed for both imaging and reconstructing an entire mouse brain at 1 μ m resolution (Fig. 2a). The resulting series of images formed a well-aligned isotropic dataset with sufficient resolution and contrast to capture both individual anatomic regions (Fig. 2b; the full dataset can be explored in Supplemental Movie 1) as well as underlying brain-wide cytoarchitecture. For example, both

neuronal somas (Fig. 2c) and long distance myelinated axon bundles (Fig. 2d) could be clearly visualized in the same brain. Data produced sufficient detail and contrast to trace cells, their morphologies, and individual myelinated axons several millimeters (Fig. 2e–g; Supplemental Movie 2).

3.2. Imaging a single sample through the complete pipeline

As described in the Methods section, a perfusion fixed mouse brain was imaged with MRI, prepared for microscopy, and imaged with both μ CT and automatic tape-collecting ultramicrotome (ATUM) serial EM (Kasthuri et al., 2015). Thus, resolutions from each modality spanned 5 orders of magnitude on the same brain: 150 μ m DTI and 50 μ m structural MRI data, 1.2 μ m μ CT data, and 3 nm EM data. Typical results (Fig. 3a–d) show similar regions from the same brain imaged with both 50 μ m resolution T2*-weighted structural MRI and 1.2 μ m resolution μ CT datasets centered on a coronal slice of brainstem and cerebellum. Using vasculature and neuronal somas as fiducials to guide the subsequent EM imaging (yellow arrows, Fig. 3d–e), we found individual synapses (white arrows, Fig. 3f) on neuronal somas in the EM data. Thus, we traversed from MRI data of the entire coronal cerebellum/brainstem to the underlying cellular architecture of the spinal vestibular nucleus with micrometer-resolution μ CT data and ultra-structural details of individual synapses with nanometer-resolution EM data, all from the same brain.

3.3. Comparing results of µCT with MRI

We next explored how whole brain μ CT could indicate underlying cellular correlates of MRI signals. Since both approaches produce isotropic 3D renderings of whole volumes (Figs. 2a and 4b), this allowed for simple co-registration between the two datasets. For example, Fig. 4a shows 3D somas/dendrites traced from μ CT data from the medial vestibular nucleus. While individually smaller than an MRI voxel, resolving the cytoarchitecture within a voxel suggests a potential source of underlying contrast variability within an MRI dataset. Similarly, the somas in the dentate gyrus (DG) of the hippocampus are easily seen in the μ CT data (Fig. 4c). Ten thousand individual cells were counted and traced within the DG over a 50 μ m slab of data. Again, while the individual somas are much smaller than the resolution of the MRI data, the high density of somas contribute to an anatomically dependent local variation in the MRI signal, rendering it resolvable from the surrounding structures (yellow box, Fig. 4b).

Finally, we investigated underlying axonal correlates from μ CT data with DTI, which is explicitly sensitive to underlying axon distributions. Fig. 4a and b show linearly coregistered sagittal cross-sections of the mouse cerebellum and brain stem in both structural MRI and μ CT, respectively. Overlain on the MRI structural scan are estimates of crossing fibers in each voxel. Specifically, the MRI data supported estimates of two fiber populations (Fig. 5b): one population traversing the anterior/posterior (A/P) direction and another traversing the left/right (L/R) direction (green and red cylinders, respectively). The yellow boxes in Fig. 5a and c indicate the location of 9 MRI voxels in which 47 individual axons (27 L/R, 20 A/P) were traced in the μ CT data (Fig. 5d). The individually traced axons from μ CT were similarly color coded for comparison. Fig. 5b shows the MRI diffusion direction estimates overlain on top of the axons. The populations of axons traced in the μ CT data

clearly follow similar directions to the MRI diffusion direction estimates while negligible numbers of axons in the μ CT data traverse directions other than two estimated with the DTI analyses. This supports the accuracy of the MRI diffusion model used to estimate directions of underlying fiber populations.

4. Discussion

We describe a pipeline using three imaging modalities, providing unprecedented views across orders of magnitude of a single mouse brain sample. This will allow for potentially novel understandings and/or interpretations of results within each modality, currently constrained by scale-dependent differences in each individual approach. For example, for axon bundles, MRI data will provide low-resolution, whole-brain estimates of neural tracts (Fig. 1a), μ CT will provide long range trajectories of individual myelinated axons in those tracts (Fig. 1b), validating and extending the MRI data, and EM would provide reconstructions of individual wrappings of myelin sheaths (Fig. 1c) or the eventual synaptic targets of those myelinated axons.

Because of the contrast mechanisms of each imaging modality, this pipeline works on brains (or any other organ type) of any species. MRI contrast is explicitly sensitive to tissue water, regardless of tissue type. Further, osmium is used as a contrast stain for both μ CT and EM because it stains membranes of any type of cell in any type of tissue (Palay et al., 1962). Therefore, in a single brain sample, our pipeline reveals the relative locations of nearly every soma, blood vessel, myelinated axon and synapse. Other ultra-high resolution whole brain imaging approaches often rely on genetic expression of fluorescent proteins to label a subset (often < 1%) of neurons, limiting these approaches to a small range of tractable animals (i.e. mouse, fly, and worm Chung and Deisseroth, 2013a). Subset labeling, by necessity, precludes detailing how different brain structures, like neuronal somas, vasculature, or myelinated tracts, relate to each other. Finally, whole brain optical approaches, whether based on expansion (Chen et al., 2015) or optical index matching (Chung and Deisseroth, 2013b), often remove cell membranes, making them essentially incompatible with subsequent large volume electron microscopy.

Other clear advantages of using whole brain μ CT as an intermediate approach is that datasets are: (1) acquired with isotropic resolution, facilitating unbiased tracing in any direction, (2) imaged through intact brains with minimal deformation, obviating computationally difficult large scale alignments (Chung and Deisseroth, 2013a; Saalfeld et al., 2012), and (3) compatible with sample preparation for downstream electron microscopy.

Such pipelines will likely be useful in studying brain pathology. Pathological alterations in neural function and behavior are well documented, but our understanding of their cellular underpinnings remains fragmentary. The complexity of brain circuitry and the marked heterogeneity of neuronal sizes and shapes across brains could mask even dramatic changes in specific neuronal subsets, making structural analyses difficult. Using μ CT and EM to inform contrast variability in MRI data over entire brains would significantly narrow the search for pathological structural change and lead to improved specificity in MRI. This would also lead to potential *in vivo* MRI biomarkers of disease. The next generation of

synchrotrons currently under construction (Schmidt et al., 2018) should allow for even deeper penetration of samples with the possibility of imaging even larger brains in their entirety. Finally, by instantiating our imaging pipeline in a national laboratory, we provide a path for future investigations at a no-cost facility open to researchers worldwide.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data availability

The data are freely available for viewing, as noted throughout the manuscript:

Our μ CT data can be publicly viewed online through Neuroglancer: (http://neuroglancerdemo.appspot.com/) with (http://tinyurl.com/cxmbjy6b) as the precomputed source link. Raw data from any of the three modalities will be made available from the corresponding author upon reasonable request.

Abbreviations:

MRI Magnetic resonance imaging
 DTI Diffusion tensor imaging
 EM automated electron microscopy
 μCT synchrotron-based x-ray tomography

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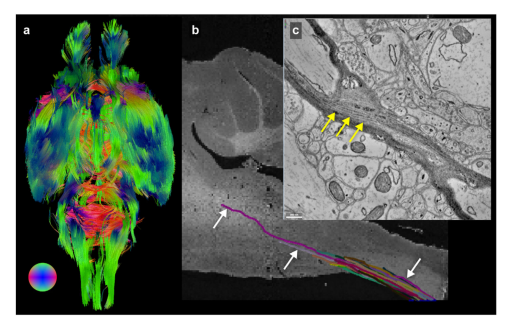


Fig. 1. Imaging modality scale differences capture scale dependent information. By using an imaging pipeline of MRI, μ CT, and EM, we can simultaneously resolve brain structures, like the white matter, at macro-, meso-, and microscopic-scales in the same brain. (a) We visualized whole brain interconnectivity by inferring white matter tracts from MRI diffusion tensor imaging. The colors denote tract directions - see inset color sphere. (b) We can trace individual axons (white arrows) within the same myelinated tracts over long distances with μ CT, directly comparing axon paths with tracts produced in MRI. With the addition of EM, we can re-inspect the ultra-structure of individually traced axons from (b), such as (c) the individual wraps of the myelin sheath (yellow arrows).

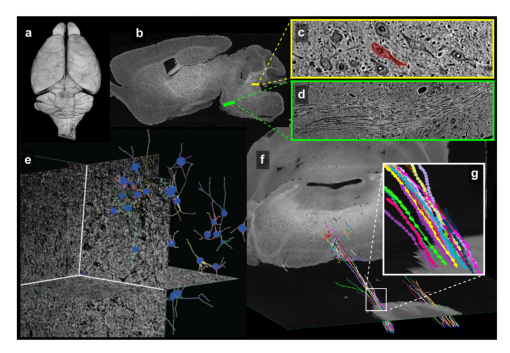


Fig. 2. An entire mouse brain imaged at 1 μ m resolution with μ CT. We used high energy x-rays (25 kEV) for tomographic imaging of (a) a whole, intact mouse brain at \sim 1 μ m resolution, collecting \sim 10 terabytes of data in \sim 7 hours. A single down-sampled sagittal slice (b) reveals obvious brain regions and myelinated tracts while the full resolution data contains enough detail and contrast to identify (c) individual neuronal somas, nuclei, and nucleoli as well as (d) individual myelinated axons. Since μ CT data are collected non-destructively, over large volumes, and with isotropic resolution, we can trace (e) somas and their neuronal processes and individual myelinated axons (g) over long distances (f).

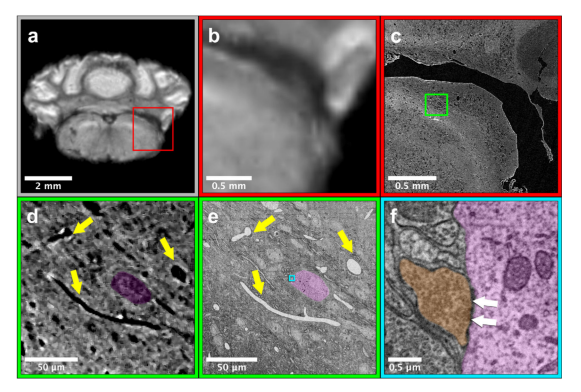


Fig. 3. A multi-scale, multimodal pipeline for imaging the same brain from MRI to EM. The same brain was imaged using diffusion (a, b) MRI (50 μ m isotropic voxels), (c, d) μ CT (\sim 1.2 um/voxel resolution), and (e, f) large volume serial electron microscopy (EM, \sim 3 nm/voxel resolution). (b, c) shows the same FOV from the whole brain MRI and μ CT imaging, corresponding to the red ROI in (a). (d - e) show a smaller FOV in both the μ CT and EM data, corresponding to the green ROI in (c). Yellow arrows indicate corresponding blood vessels, and a single neuron is labelled purple. Panel (f) highlights an individual somatic synapse (white arrows) on that soma, colored orange. The FOV of (f) is indicated by the blue ROI in (e). This pipeline demonstrates the ability to identify corresponding structures in a single brain imaged across four orders of magnitude of spatial resolution.

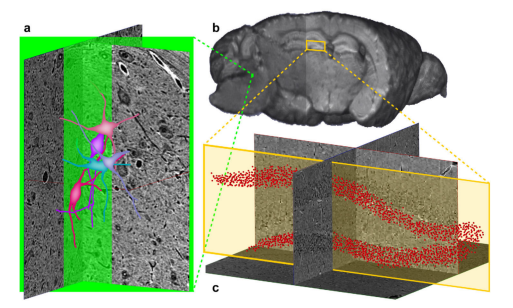


Fig. 4. Whole brain μ CT provides cellular correlates of MRI contrast. (b) 3D rendering of structural MRI data used for coregistration with similar μ CT data (Fig. 2a). (a) Somas and dendrites of neurons in the medial vestibular nucleus and somas in the (c) dentate gyrus were traced from the μ CT data. While these somas are much smaller than the resolution of the MRI data, they provide possible underlying sources of contrast in the T2* weighted images.

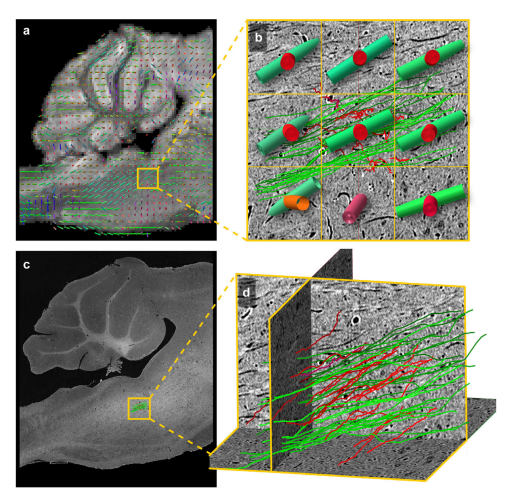


Fig. 5. Whole brain μ CT provides axonal correlates of MRI contrast) (a) DTI data of the same brain were also acquired. Principal and secondary fiber population estimates are shown overlaid on a sagittal slice of a corresponding structural MRI scan (green indicates fiber orientation along the rostral/caudal axis, red indicates left/right) as well as in (b) the magnified 9 voxels. (c) The μ CT data from the same brain has been linearly registered to the MRI structural data and (d) individually traced axons from the mCT data are shown with the same color scheme as (a) and (b). Both principal and secondary fiber estimates from DTI data indicate quite similar orientations as the underlying population of individual axons traced from the μ CT data, as seen by (b) the agreement of the DTI direction estimates superimposed on the individual traced individual axons.