Serial Section Scanning Electron Microscopy of Adult Brain Tissue Using Focused Ion Beam Milling

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Introduction

Analyzing the synaptic basis of neuronal circuits within a volume of brain tissue requires electron microscopy. With a resolution capable of seeing the smallest synaptic contacts, this method uses different sectioning techniques to produce serial images suitable for seeing the ultrastructure within significant volumes of brain tissue. Sections are cut from a block of tissue, leaving either the block face to be imaged with a scanning electron microscope or the section itself to be imaged with a transmission electron microscope. Here, we have explored the use of focused ion beam milling combined with blockface scanning electron microscopy to serially cut and image a volume of neuropil in the adult rodent brain. We show that a conventional transmission electron microscopy (TEM) fixation, staining, and embedding procedure gives sufficient membrane contrast to see all the axons and dendrites and their synaptic contacts. Milling and imaging of the block face provided serial images with sufficient resolution to allow the observer to follow all the neurites and their connections throughout the volume. This technology provides a viable alternative to conventional tissue slicing and electron microscopy (EM) imaging approaches for the acquisition of large datasets that could be used for the reconstruction and analysis of neural connectivity.

For several decades, EM has been used to study neuronal ultrastructure, from the first study using TEM to classify synapses in the cerebral cortex (Gray, 1959) to detailed serial section analyses on large parts of identified neurons (White and Rock, 1980; Anderson et al., 1994). Despite advances in light microscopy, only EM is capable of seeing every axon, dendrite, and synaptic connection within a volume of neuropil. However, imaging large volumes of tissue with serial section transmission EM is one approach that, for the moment, relies on considerable manual labor by skilled persons. Studies thus far have analyzed either the connectivity within tiny portions of complex nervous systems (Shepherd and Harris, 1998) or complete systems of much simpler ones (White et al., 1986). Successful acquisition of large sets of aligned TEM images is challenging for even the experienced microscopist. A number of factors can contribute to interrupted series or imperfect image stacks: section folds during the placement onto support films, electron beam damage that distorts the sections, and stain contamination.

Recently, an alternative to the serial section TEM approach for imaging tissue was described using scanning electron microscopy. A resin-embedded tissue sample was serially sectioned, using a microtome integrated inside a scanning electron microscope (SEM) (Denk and Horstmann, 2004), and the milled face was imaged with the scanning electron beam. The method therefore relies on the mechanical microtomy approach, with a diamond knife cutting thin sections from the block face. Unlike TEM, in which the contrast in the images is generated by the elastic scattering of electrons in regions of higher density, the block-face scanning approach captures electrons backscattered from below the surface of the block itself. More backscattering occurs in regions of higher electron-dense stain, making it possible to image conventionally prepared biological tissue. This technique can be highly automated, removing many of the problems associated with manual serial section manipulation and imaging in the TEM, automatically generating aligned serial images.

We present here an alternative sectioning approach using a focused ion beam (FIB) directed parallel to a block face for removing (or milling) thin layers of embedded tissue, and a scanning electron beam integrated into the same microscope for imaging this milled region (Heymann et al., 2006). We used this technology to serially image conventionally prepared adult brain tissue at a resolution that allowed us to follow the axons and dendrites in a volume of 286 µm³ and identify their synaptic connections. This milling approach can remove layers as thin as 15 nm from the block face. The study demonstrates the potential of this method for the automated collection of serial images through volumes of brain tissue that can be used for the analysis of neural connectivity.

Materials and Methods

Tissue preparation

An adult mouse (female, C57BL/6; 10 weeks of age) was deeply anesthetized using an intraperitoneal injection of sodium pentobarbitone. The animal then underwent chemical fixation by cardiac perfusion of 10 ml of isotonic PBS followed immediately with 200 ml of 2.5% glutaraldehyde and 2% paraformaldehyde in phosphate buffer (0.1 M, pH 7.4). Two hours after perfusion, the brain was then removed, and 60-µm-thick vibratome (Leica VT100; Leica Microsystems, Wetzlar, Germany) sections were cut through the somatosensory cortex and the cerebellum. The sections were then washed in cacodylate buffer (0.1 M, pH 7.4) and postfixed for 40 min in 1.5% potassium ferrocyanide and 1% osmium tetroxide, followed by 1 h in 1% osmium tetroxide alone, each in the same buffer, and then 40 min in 1% uranyl acetate in water. They were then dehydrated

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Figure 1. *a*, Three-dimensional drawing of the resin block showing the arrangement of the gallium ion beam (white) that scans parallel to the block (indicated by double-headed arrow) and mills away a layer of resin, creating an imaging face. The electron imaging beam (gray) at an angle of 52° from this vertical face is then used to image the embedded tissue at this surface. *b*, Scanning electron micrograph of a tissue block in the same orientation as *a* with a dotted line indicating the milled region, which is shown in *c* in reverse contrast. Scale bar, 20 µm.

in alcohol and finally infiltrated with Durcupan resin (Fluka, Buchs, Switzerland). The sections were flatembedded between glass slides in fresh resin and left for 48 h at 60°C for the resin to harden. This procedure was in accordance with the Swiss Federal Laws on Animal Experimentation administered by the Véterinaire Cantonale Lausanne.

Block preparation

Once the resin was hard, a 1 \times 1 mm square block was cut from a section that contained layer IV of the somatosensory cortex. This block was stuck to a blank resin support, and using a microtome and glass knife, trimmed down to a block face measuring 600 \times 200 µm (Fig. 1). These are typically the block dimensions used to make serial sections for TEM. The final block was then mounted, using conductive silver paint (Electron Microscopy Sciences, Hatfield, PA), on a standard SEM stub and then coated with a layer of platinum/palladium (thickness of \sim 30 nm) using a sputter coater (Cressington Scientific, Watford, UK) to remove any charge that may build up on the surface.

The block was placed inside a Dual-Beam microscope (FEI Helios NanoLab; FEI Company, Acht, Eindhoven, The Netherlands), which combines a scanning electron beam produced by a field emission gun, with a focused beam of gallium ions. This ion beam was used in conjunction with a gas injection system to deposit a thick layer (~ 1 µm) of platinum on the top surface of the sample, above the region of interest. This system releases a precursor gas, methylcyclopentadienyl (trimethyl) platinum (IV) (FEI Company), close to the sample. When the gallium ions interact with the sample, in the presence of this gas, decomposition takes place, causing platinum solid to be deposited on the surface of the sample where the ion beam was directed.

This additional deposition of a platinum layer immediately above the region of interest reduced FIB milling artifacts. The coating ensured that the resin face was milled evenly and remained parallel to the direction of the ion beam. Without it, excessive nonuniform milling occurred because of sample topography or material differences in the face. This can lead to excessive streaking or vertical stripes down the block face known as curtaining (Fig. 2*a*) (Heymann et al., 2006).

Sectioning and imaging

To prepare the initial imaging face, a region measuring $\sim 10,000 \ \mu\text{m}^2$ (Figs. 1c, 2a) was milled. Essentially, the beam skimmed the surface, and when



Figure 2. *a*, Reverse-contrast backscattered electron micrograph of an imaging face milled to expose a field of view $\sim 120 \times 115 \mu$ m. For this initial preparation phase, a small amount of "curtaining" can be seen in the bottom right corner. This imaging artifact can be removed once serial milling and imaging begins. The image was taken with a beam voltage of 5 keV, beam current of 0.40 nA, and pixel dwell time of 10 µs. The resolution of the image is 30 nm/pixel. Within this image, cell bodies (cb) and blood vessels (bv) are easily visible. *b*, Reverse-contrast backscattered scanning electron micrograph taken from the imaging face shown in *a*, of the neuronal cell body indicated as cb, using the same beam parameters, but a resolution of 10 nm/pixel. With these settings, the microscope can generate images that clearly show the stained membranes and many features within the neuropil, e.g., dendrites (de). *c*, The same image of the region enclosed in the white square in *b*, which has been enlarged to illustrate the detail that can be seen with these imaging parameters. The image shows an axonal bouton (ab) and dendritic spine head (sp). Scale bars: *a*, 20 µm; *b*, 5 µm.

ions collided with molecules in the material, they were ejected away. As the beam scanned parallel to the block face, it was slowly moved closer and closer until a $100 \times 100 \,\mu\text{m}$ face had been milled and all the resin in the path of the ion beam was removed. This created a smooth, polished face parallel to the plane of milling (Fig. 1). The progress was monitored using images formed from the electron beam, simultaneous with the ion beam milling (simultaneous patterning and imaging [SPI] mode). Next, the ion beam was set to mill layers from this region. We milled 40 nm layers, but also tested milling as little as 15 nm. After each layer was removed, a backscattered electron image was collected. The electron imaging beam was directed at an angle of 52° to the block face (Fig. 1*a*), the image being generated through the electrons interacting with the sample and being backscattered out again. These electrons were detected via the in-lens detector, using a negative potential on the tube through which the electrons were collected. This negative potential meant that the lower-energy secondary electrons were repulsed with only the high-energy backscattered electrons contributing to image formation.

Image interference can occur when milled material redeposits around the imaged area. When this occurs, there is a risk that it can build up and obscure the field of view. For this reason, the milled face was always much larger than the imaged area in both the x and y directions, to ensure that redeposition did not occur nearby.

From the initial $100 \times 100 \,\mu\text{m}$ face, a smaller region was selected, measuring $\sim 30 \times 30 \,\mu\text{m}$, free from blood vessels and not completely obscured by cell somata, and therefore containing mostly axons, dendrites, and astrocytic processes. This smaller face was milled, using an acceleration voltage of 30 keV and a current of 1000 pA. Each mill took 75 s to complete. After the removal of each layer, images were collected from a smaller subregion at a magnification of 4 nm/pixel, and with a total image size of 2048×1768 pixels (\sim 4 MB/image). This gave a horizontal field width of 8.2 um. The acceleration voltage

of the imaging beam was 2 keV, with a current of between 340 and 400 pA and dwell time of 100 μ s/ pixel (unless otherwise indicated). For each image collected, the contrast was adjusted automatically. Throughout the entire procedure, the ion and electron beam–coincident position remained at 5 mm from the final lens of the SEM column.

Although SPI mode was used to monitor the creation of the block face, the acquisition of the higher-resolution images was performed after the milling had finished. Milling and acquisition was fully automated, producing a series of nearly aligned images. Final alignment was achieved manually using Photoshop software (Adobe Systems, San Jose, CA). Individual images were overlaid on the preceding one, as a separate image, in a single file. Optimal alignment was achieved by making the top image (layer) partially transparent and moving it in the x and y directions until it was aligned with the visible image (layer) immediately below. Realignment of the serial images was never >40 pixels in the x or y direction, corresponding to 0.16 µm. The contrast of the images was also reversed to give the final images the more familiar TEM look. NOTES

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Results

Because the images were generated from the backscattered electrons, we wanted to estimate the depth that they were penetrating. This volume of interaction is a function of the density of the sample and the accelerating voltage. The lower the accelerating voltage, the less penetrating the electron beam, and therefore, the greater the information that is obtained close to the surface (Denk and Horstmann, 2004). This will have a significant effect on the imaging resolution, such that lower accelerating voltages will provide higher effective resolution, which may be critical for resolving small structures such as membranes and vesicles. Therefore, different acceleration voltages were tested to identify the appropriate voltage for identifying the cell membranes and synapses. Because we made 40 nm slices, an electron beam energy was selected that would give backscattered electrons at a depth that is less than this slice thickness (Fig. 3).

The acceleration voltage of 2 keV allowed us to see membranes, including those of vesicles within axonal boutons. At higher voltages (5 keV), these features were less distinct, and deeper underlying structures blurred the images. This is particularly evident when a myelinated axon, lying beneath the We tried to estimate the depth from which the electrons were being backscattered using 2 keV beam energy. Using a resin block of cerebellar tissue, we imaged the molecular layer, examining a single asymmetric synapse and its vesicles. We used the same imaging parameters as previously. The face was milled at depths of 15 nm, and in each image, vesicles could clearly be identified. Because these structures have diameters of between 40 and 50 nm, when section thickness is 15 nm, they can be sectioned a maximum of four times. We saw in our image series that each vesicle was present in three to four images as a circular membrane with a clear core (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). However, they first appear in a single preceding image, but without a clear center, suggesting that the imaging beam penetrates to a depth below the surface at which backscattering occurs from the vesicle. Because this occurs only once before it appears as a hollow structure, presumably when it is imaged at the surface, it would indicate a penetration depth of <30 nm.

In our larger imaging series taken from the neocortex, we milled 120 layers, each 40 nm thick, from the block face. This entire process of repeated milling and imaging occurred without any intervention from



Figure 3. a, Reverse-contrast backscattered electron micrograph of a region of neuropil with a resolution of 4 nm/pixel. The electron beam energy was 5 keV. The image shows limited contrast between the stained membranes and intracellular space. This effect appears to blur the image. b, Reverse-contrast micrograph of the same region of the imaging face that was imaged in a. All the imaging parameters are the same, except that the electron beam energy has been lowered to 2 keV. In this image, the membrane contrast is improved and the depth of imaging is decreased. Arrows in both images indicate a myelinated axon. The inset images in both panels show a dendritic spine head (sp), but only in **b** can the cluster of vesicles be seen along with the presynaptic and postsynaptic densities (arrowheads).

surface, appears as a shadow in part of the image (Fig. 3a). This shadowing does not appear when a lower voltage (2 keV) is used (Fig. 3b), reducing the interaction volume to what would appear to be less than the slice thickness.

 2048×1768 pixels, with a pixel dwell time of 100 µs; therefore, each image took \sim 4 min to acquire. A subset of the image series is shown as supplemental data (supplemental movie 1, available at www.jneurosci.org as supplemental material) in which a portion of the field of view has been included (Fig. 4c, region indicated by dotted line). This shows the milling and imaging process, enabling the viewer to follow each individual neurite through the tissue.

To independently check the milling thickness, we used the method of Fiala and Harris (2001). Measuring the diameter of several mitochondria lying longitudinal to the imaging plane and

counting the number of times this region was cut allow us to make an estimate of section thickness, presuming these structure are cylindrical. For the series cut at 40 nm intervals, this gave a milling

thickness of 48 ± 8.6 nm (mean \pm SD); for the series cut at 15 nm intervals, the milling thickness was measured as 14.1 ± 3.1 nm (mean \pm SD). It is evident in the serial images (supplemental movie 1, available at www.jneurosci.org as supplemental material) that milling thickness is not entirely consistent throughout the stack, although it is not clear why this occurred.

An analysis of neural connectivity requires the identification of the synapses. The parameters that we chose in order to collect this image series allowed for the identification of single vesicles, making it possible to see the neuronal connections. Moreover, morphological differences between identified synapses indicated that both asymmetric (presumed glutamatergic) and symmetric (presumed GABAergic) synapses could also be distinguished (Fig. 4; supplemental movie 1, available at www.jneurosci. org as supplemental material). For asymmetric synapses, vesicles were round, and their lumens appeared clear and unstained (Fig. 4). These were present in axonal boutons making contacts that were asymmetric, with the postsynaptic density being thicker than the presynaptic. This was in contrast to symmetric synapses, where the vesicles were ovoid, smaller than the asymmetric type, with the axonal boutons displaying presynaptic and postsynaptic densities that were of a similar thickness (Fig. 4). In our other studies in this region of the neocortex, all these symmetric synapses were immunopositive for GABA (Knott et al., 2002).

Discussion

Visualizing the complete wiring and connections within a neural circuit is a realistic challenge with today's EM technology. Using TEM, serial sections can be collected, stained, and imaged, and the sections revisited numerous times at any magnification (Harris et al., 2006). For the moment, much of this process needs to be done by hand and is not without pitfalls. The serial block-face scanning SEM approach (Denk and Horstmann, 2004), although removing sections that are discarded, is capable of collecting aligned series of images automatically, and with large fields of view that are not possible with currently available TEMs. The diamond knife is well suited to sectioning these large block faces. However, imaging this surface at a resolution suitable to see synapses means that significant energy is delivered to the block face, resulting in changes in the resin's structure (i.e., cross-linking); the increased hardness makes it difficult to continue the sectioning process with the knife (Denk and Horstmann, 2004).



Figure 4. a-c, Serial scanning electron micrographs taken from the imaging face each time that 40 nm of resin had been milled away. The beam voltage was 2 keV, the beam current was 0.4 nA, and the dwell time was 100 µs. The magnification was set at 4 nm/pixel, giving a field width of 8.4 µm. In each image, the inset shows a digitally magnified region that is indicated with the white square showing a bouton making an asymmetric synapse (white arrowhead) and a bouton making a symmetric synapse (black arrowhead). The dotted line in cindicates a portion of the field of view that was selected from the stack of 120 images that is shown in supplemental movie 1 (available at www.jneurosci.org as supplemental material).

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These technical limitations led us to explore the FIB milling technology with the aim of producing a series of images in which all the neurites and their connections could be identified. The resulting images required only limited alignment. Moreover, we found that imaging at a resolution that allowed us to see the axons, dendrites, and synapses did not change the properties of the resin that would interfere with the milling process. This meant that cutting with the ion beam, although limited in area, is possible regardless of the parameters of the imaging beam. Excessive cross-linking of the embedding resin does not appear to affect the milling beam and is not noticeable in the final image series.

We were able to mill a surface area of >10,000 μ m², which we then reduced to 900 µm² when we collected the final image series. The maximum possible area that can be milled, however, is limited. The largest length along which the ion beam can scan is 800 µm. This theoretical width to the imaging window means that the beam would be deflected far from the vertical. Ions at the outer edges would not be localized to a circular spot but an ellipse, with a larger surface area, resulting in fewer ions impacting the resin per unit area and decreasing their milling effect. Here we milled a region 120 um wide without seeing any detrimental effects at the windows' edges. The milling depth is also limited. The beam is focused on the upper edge of the block face, giving the best imaging surface in the upper regions. Further down the face, the beam is less focused, reducing the milling effect and decreasing the chance that ions will be able to eject molecules away from the block.

Milling the final 900 μ m² face took 75 s, which was insignificant for the short series of acquired images. When images are required over much larger surfaces, however, the milling time would become significant, as well as the possibility of charge buildup over the imaged surface. Although the initial block is coated with platinum, providing a conductive layer surrounding the imaged regions, this may become less effective when large areas are prepared.

Despite the milling time, image acquisition time is the main limiting factor for large volumes. Each image is generated as a result of the electron beam scanning the block surface, and a balance needs to be reached between fast acquisition time, high resolution, and adequate signal to generate images with the suitable contrast. Using a 4 nm/pixel magnification and dwell time of 100 μ s/pixel, each image was acquired in 4 min, and the total stack in >12 h. Although significant improvements in the dwell time

could reduce the total imaging time, we were not able to achieve this and also maintain an image quality with sufficient contrast.

The technique allowed us to see and follow all the neurites in the image series and recognize their synaptic connections. However, there was certain variation in milling depth from image to image, through the stack. This was considerably reduced when milling thinner layers, but it is not clear why, so this needs further investigation.¹ For the series of images taken here, the inconsistency in milling depth does not appear to cause problems in following the neurites through the neuropil. We could also identify both the symmetric and asymmetric synapses. However, because some of the neuronal processes, such as dendritic spine necks, can be as thin as 40 nm (Sorra and Harris, 2000), it may be necessary to mill thinner layers. Milling layers of 15 nm, although possible, and increasing the ability to automatically segment all of the structures, would also increase the number of images needed.²

In conclusion, this study shows that ion beam milling, combined with block-face scanning, can be used to generate serial images through substantial volumes of neuropil in the adult mammalian brain. These image series had sufficient contrast and resolution to identify all the axons and dendrites and their synaptic contacts. However, as an approach to collecting images on a large scale through volumes of nervous tissue that may even contain complete neural circuits, this block-face scanning technique needs further improvements. For the moment, scanning beam technology is limited in its ability to mill large surfaces, and imaging across these areas at high resolution would require considerable time. Nevertheless, the method is able to identify synapses, and the whole process of milling and image acquisition can be completely automated without relying on manual collection, staining, and imaging of large series of sections.

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¹Since the publication of this paper, we were able to achieve consistent milling thickness of ~5 nm for each section for up to 3000 images. By using beam energy of 1.5 kV, and in-lens filtering (ESB detector in the Zeiss NVision 40) (Zeiss, Oberkochen, Germany), the backscattered electrons that make up the image emerge only from within the top 5 nm of block face.

²Using image data sets with near isotropic voxels of 5–6 nm in each dimension, software is now available that can automatically segment neurites and detect symmetric synapses (www.ilastik.org).