

Syringe-injectable mesh electronics integrate seamlessly with minimal chronic immune response in the brain

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Implantation of electrical probes into the brain has been central to both neuroscience research and biomedical applications, although conventional probes induce gliosis in surrounding tissue. We recently reported ultraflexible open mesh electronics implanted into rodent brains by syringe injection that exhibit promising chronic tissue response and recording stability. Here we report time-dependent histology studies of the mesh electronics/brain-tissue interface obtained from sections perpendicular and parallel to probe long axis, as well as studies of conventional flexible thin-film probes. Confocal fluorescence microscopy images of the perpendicular and parallel brain slices containing mesh electronics showed that the distribution of astrocytes, microglia, and neurons became uniform from 2–12 wk, whereas flexible thin-film probes yield a marked accumulation of astrocytes and microglia and decrease of neurons for the same period. Quantitative analyses of 4- and 12-wk data showed that the signals for neurons, axons, astrocytes, and microglia are nearly the same from the mesh electronics surface to the baseline far from the probes, in contrast to flexible polymer probes, which show decreases in neuron and increases in astrocyte and microglia signals. Notably, images of sagittal brain slices containing nearly the entire mesh electronics probe showed that the tissue interface was uniform and neurons and neurofilaments penetrated through the mesh by 3 mo post-implantation. The minimal immune response and seamless interface with brain tissue postimplantation achieved by ultraflexible open mesh electronics probes provide substantial advantages and could enable a wide range of opportunities for in vivo chronic recording and modulation of brain activity in the future.

neural probes | ultraflexible macroporous probes | in vivo implants | minimal neuroinflammation | brain-machine interface

Stable chronic mapping of single neurons with action potential temporal resolution in the central neural system could have significant impact on research focused on both fundamental questions in neuroscience and biomedical applications (1–3). For example, fMRI (4, 5) has the ability to map whole-brain activity, although low spatiotemporal resolution (6) precludes monitoring neural circuits at the cellular level. Optical imaging (7, 8) is capable of mapping at single neuron spatial resolution, although applications have been limited by penetration depth, temporal resolution, specimen heating, and incorporation of genetically encoded reporters (9). Flexible surface electrode arrays (10–12) are also capable of mapping activity at the cellular level but are not capable of accessing deeper brain regions and generally have spatial resolution inferior to that of optical imaging. Implantable electrical probes can provide advantages for high-spatiotemporal-resolution neural recordings independent of probing depth compared with other techniques (6). However, conventional implanted electrical probes, such as silicon and microwire probes (13, 14), generally exhibit immune responses that lead to glial scar formation and neuronal cell depletion at the interfacial tissue/probe region (15–18).

Recent studies have investigated the potential advantages of reducing mechanical stiffness (19–21), feature sizes (22), as well as density of the material used for fabrication of probes (15) in terms of reduced gliosis. For example, polymer fiber-based neural probes with bending stiffness values *ca.* 8% of a rigid microwire showed reduced accumulation of astrocytes (*ca.* 50%) around probes compared with microwires 1 and 2 wk post-implantation (20). However, accumulations of astrocytes and microglia have still been observed around the fiber probe surfaces, presumably due to the bending stiffness mismatch with soft neural tissue.

To address the issues of conventional rigid devices we have introduced a paradigm for implantable multielectrode probes based upon an ultraflexible open mesh structure (23–26), where the mesh has a unique combination of structural and mechanical features. Specifically, the mesh has all size features comparable to or smaller than neuron soma and bending stiffness values of ~ 0.1 nN-m, comparable to a 150- μ m-thick brain tissue slice (i.e., the same as the overall implanted open mesh diameter), ~ 0.4 nN-m (27, 28) and orders of magnitude smaller (more flexible) than conventional probes (14, 29, 30). Herein, we present systematic time-dependent histology studies of the mesh electronics/brain-tissue interface obtained from tissue sections perpendicular (horizontal slices) and parallel (sagittal slices) to the long axis of probes, as well as time-dependent measurements obtained from horizontal tissue sections containing the cross-sections

Significance

Seamless integration of electrical probes within neural tissue could substantially enhance their impact and open up new opportunities in neuroscience research through electronic therapeutics. This paper describes systematic studies of brain tissue behavior following implantation of a design for probes that can be precisely targeted to specific brain regions by syringe injection as in many biological species and have an ultraflexible open mesh structure similar to brain tissue itself. Studies of the chronic tissue response postimplantation demonstrate that these tissue-like probes do not elicit inflammation or scarring, in contrast to more conventional probes. Moreover, neurons were found to penetrate through the probes' open mesh structure, thus demonstrating an unprecedented level of integration and compatibility with the brain circuitry.

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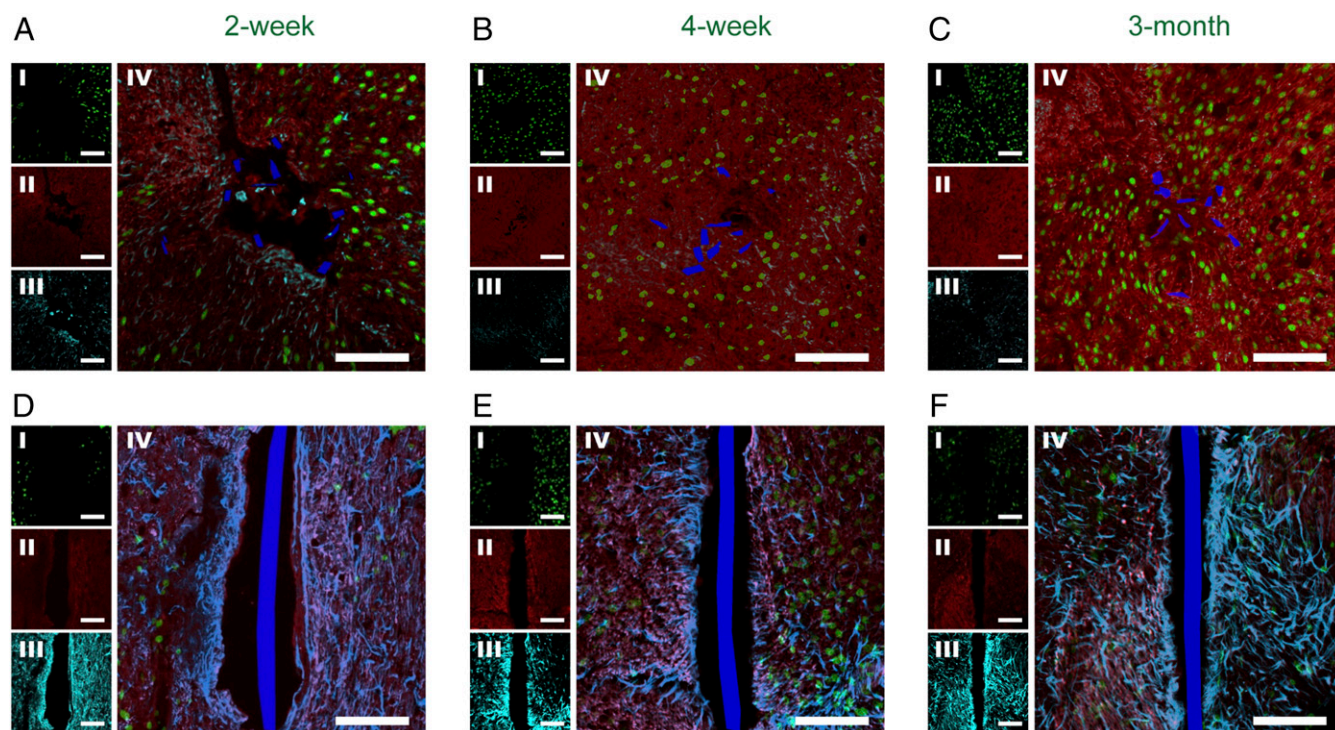


Fig. 2. Time-dependent histology of horizontal tissue slices containing implanted mesh electronics and flexible thin-film probes. Confocal fluorescence microscopy images of horizontal tissue slices containing mesh electronics/flexible thin-film probes at 2 wk (A and D), 4 wk (B and E), and 3 mo (C and F) postimplantation. In all of the panels the image labels were NeuN (I, green), NF (II, red), GFAP (III, cyan), and NeuN, NF, GFAP composite (IV). The mesh electronics and flexible thin-film cross-sections are pseudocolored blue. (Scale bars in all images, 100 μ m.)

flexible thin-film probes at 2 wk (Fig. 2D), as well as substantial accumulation of microglia (Fig. S2D).

Second, images from the intermediate 4-wk mesh electronics-implanted samples (Fig. 2B and Fig. S2B) show that (i) there is no astrocyte and little or no microglia accumulation near the mesh elements, (ii) axons interpenetrated into the interior region and the tissue is continuous within and across the boundaries of the mesh, and (iii) there are neuron somata near and/or close to the interior boundary of mesh elements. Significantly, images from 3-mo mesh electronics-implanted samples (Fig. 2C and Fig. S2C) exhibit axons and neuron somata within the mesh electronics interior at levels close to the signals hundreds of microns away, and, additionally, background levels of astrocytes and microglia around the mesh elements. It is also worthwhile to note that the penetration of axons and neuron somata into mesh interior is not correlated with statistically significant contraction of the mesh. Specifically, acute microcomputed tomography (micro-CT) images of mesh electronics implanted into mouse brains (Fig. S3) show mesh diameters similar to those estimated from the optical microscopy studies of horizontal sections from samples at 2, 4, and 12 wk (Fig. 2A–C). In comparison, images of 4-wk and 3-mo flexible thin-film probe-implanted samples (Fig. 2E and F) maintain the substantial accumulation of astrocytes and depletion of neuron somata around the thin film that is consistent with glial scarring (30, 32).

To quantify the above observations we have analyzed confocal microscopy images of the different antibody cell marker distributions as a function of distance from the surface of mesh and thin-film probes at 2 wk, 4 wk, and 3 mo, as shown in Fig. 3. In these plots, the relative specific cell marker signal was obtained by normalizing the fluorescence intensity with the baseline value defined as the fluorescence intensity averaged over a range of 500–520 μ m away from the probe (*Materials and Methods* and *SI Text*). Several key findings are evident from these plots. First, at

all times the mesh electronics show lower increases in GFAP (astrocytes) and Iba-1 (microglia) signals near the probe surfaces than the flexible thin films. For 2-wk samples, the GFAP (Fig. 3C) and Iba-1 signals (Fig. 3D) at the surface of flexible thin-film probes was 1.3 and 1.7 times the maximum of the mesh electronics, respectively. Importantly, the 4-wk and 3-mo mesh electronics-implanted samples show that the GFAP (Fig. 3G and K) and Iba-1 (Fig. 3H and L) signals return to baseline at the surface of the mesh electronics, while remaining elevated by *ca.* 2 and 1.3 times, respectively, at the flexible thin-film probe surfaces. In addition, these data show that both astrocytes and microglia penetrate to the interior of the mesh electronics reaching baseline levels at 3 mo; that is, there is near normal distribution of these cells across the entire sample.

Second, the NeuN (neuron somata) signals showed little or no decrease when approaching the mesh electronics probe surface for 4-wk (Fig. 3E) and 3-mo (Fig. 3I) samples, whereas there were 90% and 50% decreases at the same times near the surfaces of the flexible thin-film probes. Notably, these data also show that the NeuN signal is *ca.* 95% of baseline within the mesh electronics interior for 3-mo samples (Fig. 3I). In addition, at all times the NF (axons) signals for the mesh electronics showed little or no decrease near the probe surfaces, whereas NF signals are reduced from 90 to 10% from 2 wk to 3 mo near the flexible thin-film surfaces (Fig. 3B, F, and J). These data further showed NF signals within the mesh electronics interior at all times, and that the interior NF signal is the same as baseline at 3 mo.

The horizontal brain slices described above provide detailed information about the local tissue/probe interfaces for the mesh electronics and flexible thin-film implants and also allow comparison with previous studies of more rigid conventional probes (29, 30, 32, 33). These horizontal sections cannot, however, provide a global view of the probe/tissue interface, which typically extends millimeters in depth in the mouse brain. As a step

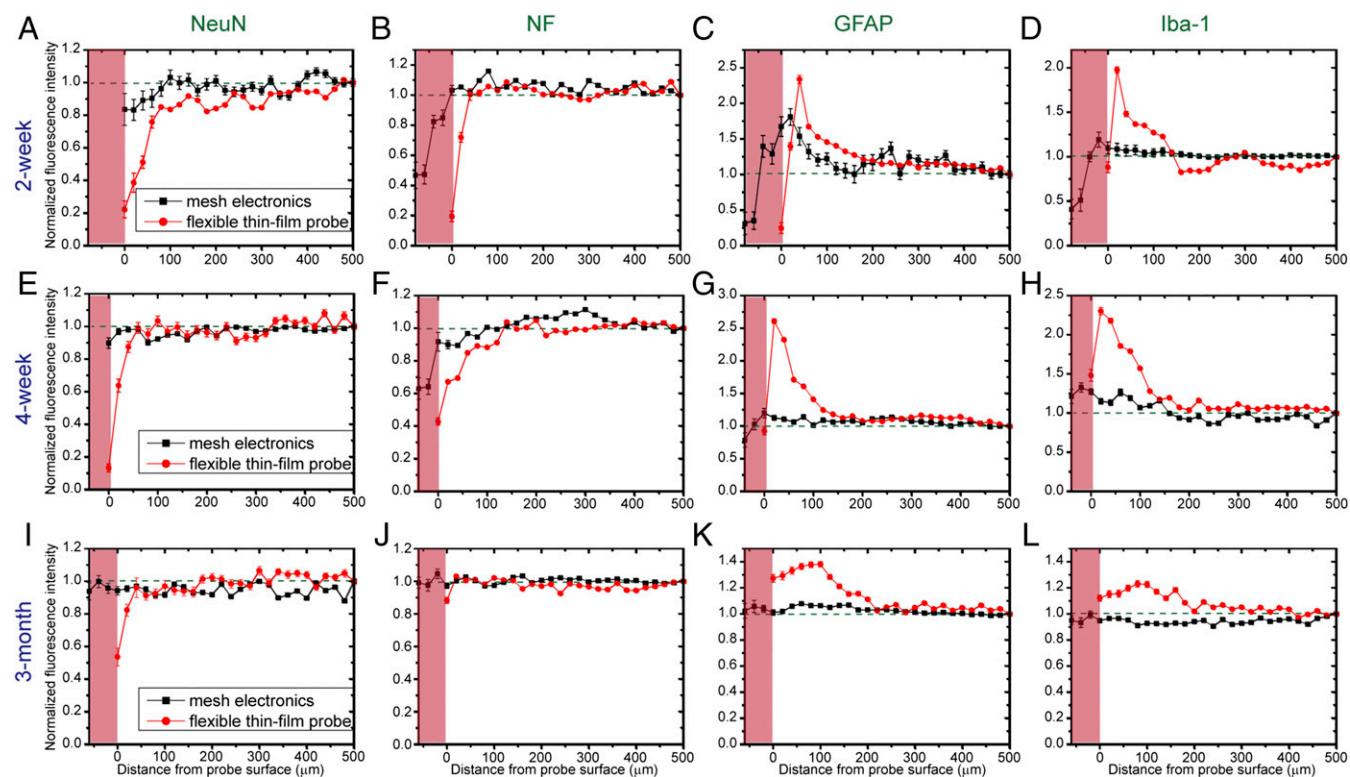


Fig. 3. Fluorescence intensity as a function of distance from the probe boundary for mesh electronics (black) and flexible thin-film probe (red) analyzed based on zoomed-out images of those shown in Fig. 2 and Fig. S2 with a field of view of $1,275 \mu\text{m} \times 1,275 \mu\text{m}$. (A–D) Fluorescence intensity of NeuN, NF, GFAP, and Iba-1, respectively, for 2-wk postimplanted samples. (E–H) Fluorescence intensity of NeuN, NF, GFAP, and Iba-1, respectively, for 4-wk postimplanted samples. (I–L) Fluorescence intensity of NeuN, NF, GFAP, and Iba-1, respectively, for 3-mo postimplanted samples. The pink shaded regions indicate interior of mesh electronics. Details of methods that were used to determine the boundary of mesh electronics and to average and normalize fluorescence intensity values of pixels in the images are described in *SI Text* and Fig. S4. Error bars represent the SEM.

toward addressing this key issue we carried out histology studies on a sagittal brain section parallel to the mesh electronics injection axes and containing nearly the entire length of the probe within the tissue slice at 3 mo postimplantation (*Materials and Methods* and *SI Text*). The confocal microscopy images (Fig. 4 A and B and Fig. S5), which were recorded from both sides of the tissue slice at a depth of *ca.* $5 \mu\text{m}$ due to limited diffusion of the antibody labels through the *ca.* $200\text{-}\mu\text{m}$ -thick sagittal slice (*Materials and Methods* and *SI Text*), highlight several key points. First, there is no evidence for proliferation of astrocytes (GFAP) or depletion of axons (NF) adjacent to the mesh electronics elements, which span $>1 \text{ mm}$ in length for these *ca.* $1.3 \times 1.3\text{-mm}$ images. Second, there seems to be nearly uniform penetration of axons through and density within the mesh probe elements, although there are some variations across the images as a whole. Third, although there are image-wide variations in the density of neuron somata (NeuN), examination of similar density regions indicate that neuron somata density is similar in the mesh electronics interior and adjacent exterior regions over length scales of at least 0.5 mm (e.g., yellow dashed box, Fig. 4B). Analysis of the normalized fluorescence intensity for NeuN, NF, and GFAP (Fig. 4C) allows the above observations to be quantified (*Materials and Methods* and *SI Text*). For example, the GFAP and NF normalized signals analyzed for each side of the sagittal slice demonstrate a uniform distribution of astrocytes and axons, respectively, from *ca.* 0.5 mm away from the probe surface through its interior along the $\sim 1.3\text{-mm}$ length of the probe. In addition, analysis of NeuN signals from regions of relatively uniform distribution on both sides of the tissue slice clearly show that the density of neuron somata is approximately the same within the mesh interior as far away as over at least the 0.4- to 0.5-mm lengths analyzed.

These histology results for both horizontal and sagittal sections highlight several unique features of the implanted mesh electronics compared with conventional probes. First, the mesh electronics produces little inflammation at short times (2 wk) postimplantation, and, moreover, there is essentially no evidence for immune response—no elevation of astrocytes or microglia, and no glial scar formation—at longer times. In contrast, conventional silicon, microwire, and flexible thin-film probes, which have orders of magnitude greater bending stiffness (i.e., are more rigid) elicit a substantial inflammatory response postimplantation as evidenced by the accumulation of astrocytes and microglia and of glial scar formation around these implanted probes (refs. 15–19, Figs. 2 D–F and 3, and Fig. S2 D–F). Second, and perhaps most important, the mesh electronics do not adversely affect the distributions of neuron somata and axons adjacent to mesh electronics implant elements. Moreover, the macroporous open structure of the mesh electronics enables axons and neuron somata to penetrate into its interior with natural cell distributions over at least *ca.* millimeter length scales at longer times. However, conventional rigid and flexible thin-film probes usually introduce a 20- to $50\text{-}\mu\text{m}$ depletion region of neurons and thus yield interfaces distinct from natural tissue (refs. 15–19, Figs. 2 D–F and 3, and Fig. S2 D–F). Hence, the mesh electronics yield a seamless integration of probes with neural tissue not possible with more rigid conventional probes, and moreover the penetration of neurons through the open mesh structure can yield a relatively large electrophysiological mapping area with only a very small cost ($<1\%$ vs. a solid probe) of occupation volume.

Conclusion

We have conducted systematic time-dependent histology studies of the interface of mesh electronics and flexible thin-film probes

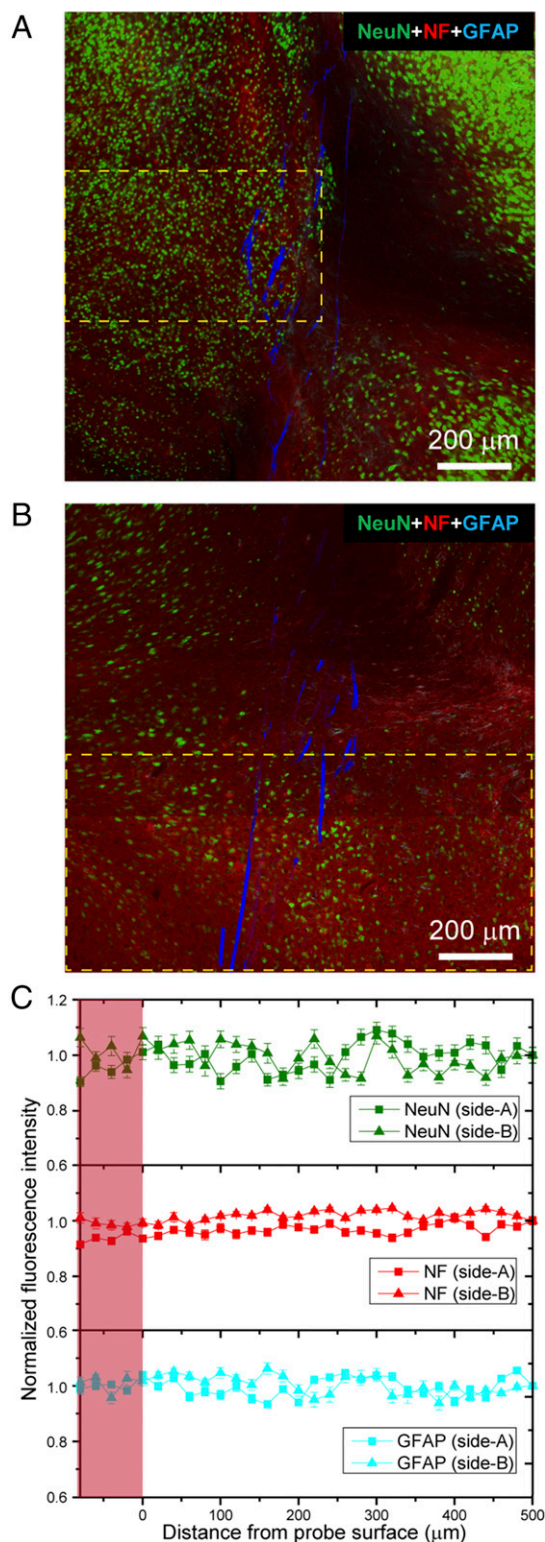


Fig. 4. Histology of a sagittal tissue slice containing nearly the full implanted mesh electronics probe. (A and B) Confocal fluorescence microscopy images of a sagittal tissue slice including the mesh electronics probe at 3 mo post-implantation. Each of the images are 3×3 composite images recorded directly in the Tile Scan mode, where each component image of the Tile Scan had a field of view of 425 μm × 425 μm. The tissue slice was stained with antibodies for NeuN (green), NF (red), and GFAP (cyan); the mesh is shown as pseudocolored blue. The images were recorded at an optical focal plane ca. 5 μm below the surfaces of side-A (A) and side-B (B). (C) Fluorescence intensity as a function of distance from the boundary of the mesh electronics

with brain tissue. The results show that mesh electronics lead to much less inflammation and damage to surrounding neurons than flexible thin-film probes at short times, and, moreover, the mesh electronics exhibit essentially no evidence for chronic immune response and does not adversely affect the natural distribution of neurons at longer times. The small amount of tissue damage and astrocyte proliferation observed in mesh electronics/tissue samples at 2 wk postimplantation, which recovers by 4 wk, can be attributed to acute damage during the initial implantation. Thus, it should be possible to reduce this acute damage by using smaller-diameter needles for mesh injection and possibly speed the recovery time by coinjection of drugs that reduce inflammation (34, 35) and/or biochemical modification of the mesh electronics (36–39). More importantly, interpenetration of axons and neuron somata into the interior of mesh electronics allows for the formation of a seamless interface between mesh electronics and neural tissue in a manner not previously achieved with more conventional probes. This observation of seamless integration and the ultraflexibility of mesh electronics suggests that the mesh electronics could affect related areas such as implants to, for example, the spinal cord and neuromuscular junctions (40, 41), as well as opening a new window for brain–machine interfaces and cyborg animals (42–44).

Materials and Methods

Fabrication of Syringe-Injectable Mesh Electronics and Flexible Thin Film. Mesh electronics were fabricated following procedures reported previously (23, 25, 26). Key steps in mesh electronics fabrication are described in *SI Text*. In brief, (i) an Ni sacrificial layer with a thickness of 100 nm was thermally evaporated (Sharon Vacuum) onto a 3-inch Si wafer, (ii) a 420-nm layer of negative photoresist SU-8 (SU-8 2000.5; MicroChem Corp.) was spin-coated on the Si wafer then patterned by PL, (iii) PL and thermal deposition were used to pattern and deposit metal interconnects Cr/Au (5/100 nm) and Pt sensing electrodes Cr/Pt (5/50 nm) on the SU-8 layer, (iv) and step ii was repeated to pattern an SU-8 passivation layer, serving as the top insulating layer of the metal interconnects. Flexible thin film was prepared by cutting polyimide film (Kapton, 25 μm in thickness; CS Hyde Company) into small probes with a width of ca. 500 μm and length of ca. 4 mm.

Stereotaxic Surgery and Probe Implantation into Mice Brains. Key steps for stereotaxic surgery and implantation of mesh electronics and flexible thin-film probes are described in *SI Text*. In brief, all tools and probes were sterilized before using. An anesthetized mouse (adult male C57BL/6J mice; Jackson Laboratory) was fixed on a stereotaxic frame (Lab Standard Stereotaxic Instrument) to perform surgery and implantation of probes.

All procedures performed on the mice were approved by the Animal Care and Use Committee of Harvard University. The animal care and use programs at Harvard University meet the requirements of the federal law (89-544 and 91-579) and NIH regulations and are also accredited by the American Association for Accreditation of Laboratory Animal Care.

Histology Sample Preparation. Key steps for sample preparation are described in *SI Text*. In brief, mice with implanted mesh electronics or flexible thin film at postimplantation times of 2 wk, 4 wk, and 3 mo were anesthetized and then transcardially perfused with 1× PBS, followed with 4% formaldehyde (Sigma-Aldrich) before decapitation. The brain was resected from the cranium and placed in 4% formaldehyde for 24 h and then transferred to 1× PBS for 24 h at 4 °C to remove remaining formaldehyde. For horizontal sectioning, the brain was cryoprotected, embedded, and then sectioned into 10-μm-thick horizontal slices using Leica CM1950 cryosectioning instrument (Leica Microsystems). For sagittal sectioning, the brain was embedded in 3% agarose (SeaPlaque Lonza Group Ltd.) hydrogel after being fixed in 4% formaldehyde, imaged by micro-CT to determine mesh probe orientation, and sectioned into 200-μm slices through vibratome stage (VT1000 S vibrating blade microtome; Leica).

in images of side-A (A) and side-B (B). The pink shaded regions indicate interior of mesh electronics on each side. The NF and GFAP fluorescence intensity was analyzed based on the entire images, and the NeuN fluorescence intensity was analyzed based on the regions shown in yellow dashed boxes in A and B. Error bars represent SEM.

Immunohistochemistry. Horizontal brain tissue sections were rinsed three times in 1× PBS and blocked using a blocking solution consisting of 0.3% Triton X-100 (Life Technologies) and 5% goat serum (Life Technologies) in 1× PBS for 1 h at room temperature. Slices were then incubated with the primary antibodies, rabbit anti-NeuN (1:200 dilution; Abcam), mouse anti-neurofilament (1:400 dilution; Abcam), rat anti-GFAP (1:500 dilution; Thermo Fisher Scientific), or rabbit anti-Iba1 (1:400 dilution; Wako Chemicals) containing 0.3% Triton X-100 and 3% goat serum overnight at 4 °C. After incubation, slices were rinsed nine times for total of 45 min with 1× PBS, before they were incubated with the secondary antibodies, Alexa Fluor 488 goat anti-rabbit (1:200 dilution; Abcam), Alexa Fluor 568 goat anti-mouse (1:200 dilution; Abcam), and Alexa Fluor 647 goat anti-rat (1:200 dilution; Abcam), for 1 h at room temperature. Slices were then rinsed nine times for a total of 30 min before they were mounted on glass slides with coverslips using ProLong Gold Antifade Mountant (Life Technologies). The slides remained in the dark at room temperature for at least 24 h before microscopic imaging. Sagittal brain tissue sections were rinsed three times in 1× PBS and blocked using a solution consisting of 0.3% Triton X-100 (Life Technologies) and 5% goat serum (Life Technologies) in 1× PBS for 6 h at room temperature. Slices were then incubated with the primary antibodies, rabbit anti-NeuN (1:200 dilution; Abcam), mouse anti-neurofilament (1:400 dilution; Abcam), and rat anti-GFAP (1:500 dilution; Thermo Fisher Scientific Inc) containing 0.3% Triton X-100 and 3% goat serum for 12 h at 4 °C and then flipped over for another 12 h at 4 °C. After incubation, slices were rinsed three times every hour for a total of 6 h. Then they were incubated with the secondary antibodies, Alexa Fluor 488 goat anti-rabbit (1:200 dilution; Abcam), Alexa Fluor 568 goat anti-mouse (1:200 dilution; Abcam), and Alexa Fluor 647 goat anti-rat (1:200 dilution; Abcam) for 6 h and flipped over for another 6 h at room temperature. Slices were then rinsed three times every hour for a total of 6 h before they were mounted on glass slides with coverslips using ProLong Gold Antifade Mountant (Life

Technologies). The slides remained in the dark at room temperature for at least 24 h before microscopic imaging.

Imaging and Image Data Analysis. Confocal fluorescence imaging of the samples was acquired on a Zeiss LSM 880 confocal microscope (Carl Zeiss Microscopy GmbH). Confocal images of antibody-labeled horizontal slices in Fig. 2 and Fig. S2 were acquired using 488-, 561-, and 633-nm lasers as the excitation sources for Alexa Fluor 488, Alexa Fluor 568, and Alexa Fluor 647, respectively, and a 1-Airy unit (AU) pinhole. The mesh electronics in each slice was imaged with DIC on the same microscope (Fig. S1) and was assigned as a false blue color in the composite images (SI Text). Images of flexible thin-film probes were acquired using the same method. Confocal images from both sides of the antibody-labeled sagittal slice in Fig. 4 were acquired using the same excitation sources as above, a 3-AU pinhole, which yields an optical section thickness of ca. 9 μ m, and image planes centered at ca. 5 μ m below surfaces of the sample. The mesh electronics in Fig. 4 were imaged in reflection mode using the same confocal microscope and imaging conditions (Fig. S5 and SI Text).

ImageJ software and custom MATLAB software were used for image analysis of both horizontal and sagittal slices (SI Text). In short, the distance of each pixel from mesh electronics in a given image was defined as its shortest distance from mesh boundary and the intensity values for all pixels with distances binned over an interval of 20 μ m were averaged and normalized against the baseline intensity, which is defined as the average fluorescence intensity of all pixels 500–520 μ m away from the mesh boundary.

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