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Cell-specific spatial profiling of targeted protein expression to characterize the impact of intracortical microelectrode implantation on neuronal health†

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Intracortical microelectrode arrays (MEAs) can record neuronal activity and advance brain-computer interface (BCI) devices. Implantation of the invasive MEA kills local neurons, which has been documented using immunohistochemistry (IHC). Neuronal nuclear protein (NeuN), a protein that lines the nuclei of exclusively neuronal cells, has been used as a marker for neuronal health and survival for decades in neuroscience and neural engineering. NeuN staining is often used to describe the neuronal response to intracortical microelectrode array (MEA) implantation. However, IHC is semiquantitative, relying on intensity readings rather than directly counting expressed proteins. To supplement previous IHC studies, we evaluated the expression of proteins representing different aspects of neuronal structure or function: microtubule-associated protein 2 (MAP2), neurofilament light (NFL), synaptophysin (SYP), myelin basic protein (MBP), and oligodendrocyte transcription factor 2 (OLIG2) following a neural injury caused by intracortical MEA implantation. Together, these five proteins evaluate the cytoskeletal structure, neurotransmitter release, and myelination of neurons. To fully evaluate neuronal health in NeuN-positive (NeuN+) regions, we only quantified protein expression in NeuN+ regions, making this the first-ever cell-specific spatial profiling evaluation of targeted proteins by multiplex immunocytochemistry following MEA implantation. We performed our protein quantification along with NeuN IHC to compare the results of the two techniques directly. We found that NeuN immunohistochemical analysis does not show the same trends as MAP2, NFL, SYP, MBP, and OLIG2 expression. Further, we found that all five quantified proteins show a decreased expression pattern that aligns more with historic intracortical MEA recording performance.

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Introduction

Implanted intracortical neural electrodes, such as microelectrode arrays (MEAs), can help characterize the underlying pathophysiology of various diseases or allow for interfacing between human patients and external systems.^{1–3} MEAs implanted directly into the cortex can record neurons with single-unit resolution, making them extremely valuable for brain-computer interfacing (BCI) applications.^{4–7} BCIs are devices that allow for communication between the brain and an external device,

including prosthetics or wheelchairs, to aid patients with loss of motor function.^{8,9} However, challenges in MEA reliability restrict the long-term clinical implementation of BCIs with MEA input. MEA recording quality is highly variable and generally decreases with time.^{10–15} MEA failures weeks to months post-implantation are mostly attributed to the tissue's biological response to the implant.¹⁶

MEAs are implanted in the cortex of the brain and, like many neural injuries, result in the activation of microglia and macrophages, the formation of a glial scar, the dieback of local neurons, and demyelination.^{15,17} Microglia are macrophage cells localized to the brain and are immediately activated by MEA implantation.¹⁸ Astrocytes are activated by microglia signaling and will migrate to the MEA surface, forming a tight glial scar that encapsulates the MEA and separates it from the neurons.¹⁹ Immune cells such as macrophages and T-cells will be recruited from the periphery to the MEA implant site.^{20,21} Collectively, the neuroinflammatory response to implanted

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MEAs contributes to the breakdown of electrode material, increased impedance between the neurons and the microelectrode contacts, dieback of local neurons, production of an impermeable glial scar, and therefore reduction in the electrode recording quality.^{15,22} Many different mechanisms have been investigated as major contributors to chronic MEA failure, including BBB permeability, material degradation, glial scar impedance, innate immune activation, and neuronal death.^{23–27} The complexity of device failure and the tissue response make it difficult to develop a therapeutic that adequately stabilizes recording quality.

Upon implantation, mechanical stress will rupture the cell bodies or the axons of the neurons in the path of the MEA.²⁸ The initial injury of MEA insertion is considered the primary injury.^{29,30} Neurons injured during the primary stage will release damage-associated molecular pattern molecules (DAMPs), activating local microglia and heightening the acute inflammatory response.^{31,32} Chronic neuroinflammatory responses to the MEAs result in a secondary neuronal injury. M1 microglia and A1 astrocytes will secrete pro-inflammatory factors that cause immune cell recruitment and proliferation.^{33,34} Reactive oxygen species (ROS) are highly reactive free radicals that are produced by inflammatory processes following inflammation. ROS are known to contribute to aging and neurodegeneration in injured states.³⁵ ROS will react with DNA, RNA, proteins, and lipid cell membranes in the neural environment, damaging local cells.^{35,36} The recording window required for an MEA to obtain single unit action potentials is approximately 50–150 μm from the implant,³⁷ and many neurons within this region are killed post-implantation.³⁸

The current standard for neuronal evaluation following MEA implantation is immunohistochemistry. The complex mechanisms of neuronal dieback are most commonly evaluated by immunohistochemistry using neuronal nuclear protein (NeuN).^{1,13–15,27,39–47} NeuN is a protein abundant in neuronal nuclei that binds to the nuclear membrane of mature neurons. Though the function of NeuN is not fully understood, it is hypothesized that NeuN is a transcription factor that regulates the general neuronal phenotype.⁴⁸ NeuN is present exclusively in neurons but is not detected in some neuronal cell types, such as Purkinje cells in the cerebral cortex and mitral cells in the olfactory bulb.^{49,50} Bedell *et al.* and Nguyen *et al.* have shown that NeuN expression returns to a healthy baseline level 50–100 μm from the MEA implant site.^{23,42} Essentially, NeuN expression initially declines and then replenishes with time post-MEA implantation. However, MEA neuronal recording performance decreases with time post-implantation.^{10,51} The stark difference between trends in neuronal recordings and NeuN expression could be interpreted to suggest that neuronal health is not a main contributor to chronic MEA failure. A more holistic understanding of the changes in targeted protein expression within neurons following MEA implantation is necessary to drive forward therapeutics to improve recording performance.

Our lab recently conducted a proteomic study of inflammation and neuronal health following MEA implantation in the motor and sensory cortices, evaluating the expression of several

key proteins for neuronal and oligodendrocyte health and functionality using the NanoString Digital Spatial Profiler (DSP).⁵² The NanoString DSP quantifies the proteins found in individual regions of interest (ROIs) using UV capture with <1-micron resolution, allowing for customizable and precise calculations of protein expression. Proteomic evaluation within 180 μm of the implant site revealed that several key proteins, including neurofilament light (NfL) and myelin basic protein (MBP), exhibit more than a 50% decrease in implanted animals compared to controls. The proteomic deficits did not include cell-specific or distance-specific capture of proteins. We are expanding on our previous proteomic analysis by adding segmentation to our multiplex immunochemistry collection process. In previous papers, we utilized much larger panels, which could be called proteomics. Here, due to the focus on a subset of the proteins examined in that panel, we will more accurately refer to this subset of the larger proteomics as multiplex immunochemistry. The present study is the first to segment multiplex immunochemistry quantification based on factors like distance from the MEA implant or cell type. Utilizing the DSP's functionality of ROI-specific collection, we split collection regions based on both distance from the implant (90 μm rings), as well as by cell type using NeuN immunohistochemistry (IHC). Essentially, we collected proteins only within NeuN+ regions in both implanted and naïve control samples. The additional segmentation methods will allow us to isolate neuronal proteins and quantify the damage based on distance from the MEA.

Here, we evaluated the expression of five proteins associated with neuronal and oligodendrocyte health: microtubule-associated protein 2 (MAP2), neurofilament light (NfL), synaptophysin (SYP), myelin basic protein (MBP), and oligodendrocyte transcription factor 2 (OLIG2). Taken together, this group of proteins can give an indication of a neuron's structure and ability to properly conduct an action potential or release neurotransmitters. MAP2, NfL, SYP, MBP, or OLIG2 are found in nearly all major regions of neurons: the dendrites, nucleus, cytoplasm, axon, and axon terminal (Fig. 1). MAP2 is a cytoskeletal protein abundant within neuronal dendrites that increases microtubule bundling and polymerization.⁵³ Downregulation of MAP2 could lead to increased microtubule collapse or depolymerization in the dendrites, reducing connections between neurons, and leading to neuronal hypoexcitability.⁵⁴ NfL is a cytoskeletal protein that makes up most of the proteins within an axon. NfL is an intermediate filament or cluster of fibers containing rod-like domains that comprise the cytoskeleton. NfL is also present in the cytoplasm of the cell body, but it is not as abundant in that region.^{55,56} SYP is a protein that forms the walls of synaptic vesicles that store and release neurotransmitters in the synaptic terminals. SYP is used as a marker for synaptic density.^{57,58} Neurotransmitter release is crucial to the function of a neuron because it allows for the excitation of the post-synaptic neuron in the neural circuit. Therefore, SYP downregulation causes neuronal hypoexcitability.⁵⁹ MBP is the second-most abundant protein in oligodendrocytes, making up 30% of the total proteins. Oligodendrocytes are the myelin cells of the central nervous system and are made up of proteolipid





Fig. 1 Simplified illustration of the general location of each protein on a motor neuron. OLIG2 is not illustrated, as it is a signaling protein rather than structural. Created with <https://BioRender.com>.

membranes. MBP binds to the lipid membranes and helps to form the structure of the myelin.^{60,61} Downregulation of MBP is likely indicative of demyelination of local neurons. OLIG2 is a transcription factor expressed highly in oligodendrocytes and oligodendrocyte precursor cells, while some expression has also been observed in astrocytes.⁶² OLIG2 is considered an important protein for myelin healing after disease or injury.⁶³

In addition to the 5-protein panel, we performed immunohistochemical analysis of NeuN, consistent with previous MEA studies.^{13,40,42,46} The direct comparison of NeuN immunohistochemical analysis to quantifying other neuronal health proteins could advance our understanding of neuronal health following MEA implantation. Under the assumption that NeuN represents neuronal health surrounding the MEA, other proteins involved in neuronal functionality would be expected to show similar trends. Evaluating additional proteins could also provide a more insightful understanding of neurodegenerative mechanisms and the extent of neuronal damage. We can call upon a vast collection of MEA recording literature evaluating chronic device performance,^{10,16} and neuronal proteins that align with MEA recording trends could be further evaluated as markers that indicate recording capabilities. Finally, adding distance and cell segmentation, we present new innovative techniques for quantifying protein expression following MEA implantation.

Methods

Animal preparation

Mice were housed at Case Western Reserve University and cared for in compliance with an IACUC-approved protocol. The mice used for this study were male C57BL/6J mice. For the experimental groups, mice underwent intracortical MEA implantation surgery once they reached 8 weeks of age. Surgical procedures were conducted according to established laboratory protocols approved by the Case Western Reserve University IACUC.^{64,65}

For each mouse in the experimental groups, four non-functional probes were implanted into the motor and sensory cortices on both the left and right sides of the brain, with coordinates 1.5 mm lateral and 1.0 mm anterior and posterior to the bregma. Mice were sacrificed at 4 or 8 weeks following MEA implantation (4 mice per group, each with 4 implants). The naïve control mice were sacrificed without any craniotomy or implant (4 mice).

Mice were sacrificed *via* cardiac perfusion. Animals were anesthetized and treated with ketamine and xylazine intraperitoneal injections. While the heart was still beating, incisions were made in the chest to expose the heart, and a butterfly needle attached to a peristaltic pump was inserted into the left ventricle. An incision was made in the right atrium to relieve the pressure on the heart and allow the outflow of blood and PBS. First, 30 ml of 1× phosphate-buffered saline (PBS) was circulated through the bloodstream. Next, 30 ml of 30% sucrose in 1× PBS was pumped through the mouse to cryoprotect the brain. Brains were extracted, flash frozen on dry ice suspended in optimum cutting temperature compound (OCT, Sakura Finetek USA, Item Number: 25608-930), and kept at -80° until cryo-slicing.

Microelectrode probes

The probes used in this study were non-functional silicon probes in the shape of single-shank, Michigan-style electrodes. Probes were 2 mm long, 150 μm wide at the widest section of the shank, and 15 μm thick. The Pancrazio Laboratory at the University of Texas Dallas generously provided these probes.

Analysis for spatially resolved protein expression

The full protocol for spatial protein expression analysis was provided by NanoString Technologies (<https://university.nanostring.com/geomx-dsp-manual-slide-preparation-user-manual>) using the FFPE protocol with amendments for fresh frozen tissue. The main steps are illustrated in Fig. 2. This protocol is





Fig. 2 Illustration of protein quantification methods. Intermediate steps such as washes or fixation are not illustrated but are described at length in the provided protocol.

described in depth in previous proteomic analyses.⁵² Isolated brains were cryo-sliced into five μm sections along the length of the cortex. These slices were fixed in 10% neutral buffered formalin (NBF) overnight at 4 °C. Slides were washed thrice for five minutes each in 1 \times tris-buffered saline with 0.1% tween 20 detergent (TBST). Following fixation and washing, slides were placed in 1 \times citrate buffer and underwent antigen retrieval in the TintoRetriever Pressure Cooker (Bio SB, Item Number: BSB 7008) on high temperature and pressure settings. Slides were then left at room temperature for 25 minutes and washed once again with 1 \times TBST. Following this, slides were stained with 1:10 anti-NeuN (Alexa Fluor[®] 532 GA-5, Item Number: NBP2-33184AF532) and 1:40 anti-GFAP antibodies (Alexa Fluor[®] 647 EPR12763, Item Number: ab190565), which were used to identify the implant site.

The protein panels purchased for this experiment include the Neural Cell Profiling Core (25 proteins, Item Number: 121300120), which was paired with the Glial Cell Subtyping Module (10 proteins, Item Number: 121300125), and Autophagy Module (10 proteins, Item Number: 121300124), which were analyzed previously. Our former proteomics analysis used the same protein panels but focused largely on immune cell activation and potential immunotherapy targets.⁵² For this study, we only included the expression of a subset of proteins that we thought would best represent the structure, health, and function of neurons and oligodendrocytes. The five proteins used in this study are: MAP2, NFL, SYP, MBP, and OLIG2. The slides were stained with custom antibodies that are each associated with a unique fluorescent identifier, or “barcode.” Each barcode uniquely corresponds to a protein that was quantified in this study. After staining with morphological and NanoString-specific panel antibodies, slides were incubated in a humidity chamber overnight at 4 °C.

The following day, slides were washed in 1 \times TBST three times for 10 minutes each, and then fixed with 10% NBF in a

humidity chamber for 30 minutes at room temperature. Next, the slides were washed twice with 1 \times TBST for five minutes each. Slides were then placed in a humidity chamber and stained with SYTO-13 (NanoString Technologies, Item Number:121300303) as a nuclear stain to better visualize the implant site. The slides were allowed to incubate for 15 minutes at room temperature, after which they were washed with 1 \times TBST two times. Finally, slides were loaded into the NanoString GeoMx Digital Spatial Profiler to image the implant site and select a region of interest (ROI), which was segmented into NeuN+ and GFAP+ regions. While NeuN and GFAP were both used to identify the implant sites, only NeuN was used to identify collection sites. Images were exported to FIJI, where custom segmentation was created to segment each NeuN+ region into three concentric rings 0–90 μm (inner), 90–180 μm (middle), and 180–270 μm (outer) from the implant site (Fig. 3). These rings will hereafter be referred to as “inner,” “middle,” and “outer” rings, respectively. The 270 μm collection region was selected to fit into the maximum ROI radius of the GeoMx DSP, which is 300 μm . The 90 μm interval was chosen for consistency with transcriptomic studies to be reported in future publications. A minimum area of tissue for the collection of at least 20 cells for transcriptomic analysis in the area closest to the MEAs. To later be able to correlate the protein expression with transcriptomics, we binned at 90 μm . Segmented images were imported back into the GeoMx and assigned separate areas of interest from which to collect protein counts. The GeoMx shined a UV light on NeuN+ area in each ring, cleaving the barcodes from the antibodies they were bound to. Barcodes were collected in a 96-well plate. GeoMx Hybridization Codes (NanoString Technologies, Item Number: 121300401) were added to the rows in the 96-well plate to allow for pooling each column and streamlining the 96-well plate into a 12-strip tube. Following the pooling of the samples, the 12-strip tube was loaded into the NanoString nCounter MAX/FLEX System, which





Fig. 3 A cross-section of an MEA implant site stained for neuronal nuclei (NeuN, red) and activated astrocytes (GFAP, green), showing the segmented NeuN+ inner ((A), 0–90 μm), middle ((B), 90–180 μm), and outer ((C), 180–270 μm) regions (yellow).

transferred samples into a clear cartridge and scanned the barcodes, generating the counts of each barcode, and thus each protein, present within each selected NeuN+ ring.

Protein expression statistics

The raw counts of each measured protein were exported from the NanoString nCounter MAX/FLEX system and analyzed for differential expression in MATLAB. A custom MATLAB script normalized the protein counts to three housekeeping proteins included in the protein panels: histone H3, S6, and glyceraldehyde 3-phosphate dehydrogenase (GAPDH). The MATLAB script also calculated the $\log_2(\text{fold change})$ (“ $\log_2\text{FC}$ ”) for each protein, which is the log of the average of the experimental group protein expression divided by the average of the control group protein expression. The control group, or baseline, for this study is always naïve controls. A positive $\log_2\text{FC}$ indicates that the ratio of experimental to control expression is greater than 1, which means the given protein is found in higher concentration in the experimental group. We denote higher experimental expression as “upregulation.” Likewise, a negative $\log_2\text{FC}$ value is “down-regulation” of the experimental group compared to naïve controls. The experimental group is always one of the implanted groups, either 4- or 8-weeks post-implantation. The MATLAB script also generated the p -values and the adjusted p -values (p_{adj}) for each protein. The p -values were adjusted using the Benjamini–Hochberg correction to filter out random significance. The R packages “tidyverse” and “readxl” were used to generate probability density functions (PDFs) for each protein using a kernel density estimation.^{66,67} A kernel density estimation is used to PDFs visualize the likelihood of a normalized protein count appearing at a specific value. PDFs are used to show the distribution, such as normal, bimodal, or skewed, of a protein within a specific comparison.⁶⁸ GraphPad Prism 10 (GraphPad Software, Boston, Massachusetts) was used to generate volcano plots and heat maps for data visualization. Within the volcano plots, all proteins on the left side of the x -axis (negative values) are downregulated in the experimental group, and all proteins on the right side of the x -axis (positive values) are upregulated in the experimental group. The volcano plots show the significance cutoff of $p_{\text{adj}} < 0.05$.

Analysis of neuronal nuclear counts using NeuN

Images from the ROI selection step of the experimental pipeline was exported from the GeoMx. The exact inner, middle,

and outer rings used in the protein expression analysis were used for the NeuN nuclear count analysis (Fig. 3). The NeuN stain was used to manually count NeuN+ cells in each ring separately for the experimental groups. Cell counts were uploaded to an Excel (Microsoft, Redmond, Washington) spreadsheet for statistical analysis. The NeuN+ cell count for each ring was normalized by the area of the individual ring. This gave us an area density of neuronal nuclei per square millimeter for each 90 μm ring of the experimental groups. The control groups did not consist of rings, as there was no implant, but instead, an equivalent area to the combined inner, middle, and outer rings (270 μm radius) was collected at the same coordinates as the implants in the experimental groups. The average nuclear density in the naïve control mice region is referred to as the control baseline. An unpaired, two-tailed t -test was performed on densities with a significance cutoff of $p < 0.05$. Data from the neuronal nuclear counts was visualized using GraphPad Prism 10.

Results

Analysis of neuronal and oligodendrocyte health using multiplex protein panel

Five proteins were quantified in NeuN+ regions using the NanoString GeoMx Digital Spatial Profiler: MAP2, NfL, SYP, MBP, and OLIG2. Six comparisons were made for protein expression: a combination of the two time points (4- and 8 weeks post-implantation) and three 90 μm concentric rings (inner, middle, outer) extending from the implant site. All comparisons have a baseline of healthy, unimplanted naïve control mice, meaning that the reported results represent changes from healthy tissue.

At 4 weeks post-MEA implantation, no proteins were found to be significantly differentially expressed. MAP2, NfL, SYP, and OLIG2 show insignificant upregulation in the implanted mice compared to naïve controls. OLIG2 exhibited a relatively high average $\log_2\text{FC}$ value of ~ 8.0 . Though insignificant, MBP was the only downregulated protein, with a $\log_2\text{FC}$ of ~ -0.8 (Fig. 4A–C and G). The protein expression 4 weeks post-implantation is highly variable, which resulted in a lack of significance at a threshold of $p_{\text{adj}} < 0.05$. Fig. 5 shows the probability density functions of all 5 measured proteins, illustrating the protein expression distribution within each comparison.





Fig. 4 Quantification of protein counts. (A)–(C) Volcano plots of protein expression within the NeuN+ region in 4-weeks post-implantation mice compared to the NeuN+ region in naïve control mice. The red line is the significance threshold $p_{\text{adj}} < 0.05$. All significant proteins are labeled (only MBP). (D)–(F) Volcano plot of protein expression within the NeuN+ region in 8-weeks post-implantation mice compared to the NeuN+ region in naïve controls. All significant proteins are labeled (all proteins were significant in the middle and outer regions). (A) and (D) Analysis of inner ring only. (B) and (E) Analysis of middle ring only. (C) and (F) Analysis of outer ring only. (G) Heat map of protein expression. This figure combines data from (A)–(F). Black asterisks represent significant differential expression between experimental groups compared to naïve controls. The log₂FC scale on the right represents comparisons that have significant differential expression, ranging from low magnitude log₂FC values (dark blue) to high magnitude log₂FC values (red). This figure was assembled with <https://BioRender.com>.

By 8 weeks post-implantation, MAP2, NfL, and MBP show significant downregulation in all three rings extending to 270 μm from the MEA-tissue interface. The average log₂FC of MAP2, MBP, and NfL between the three rings were ~ -0.6 , -2.0 , and -0.9 , respectively. In the inner ring, OLIG2 and SYP show insignificant downregulation with a log₂FC of ~ -0.4 and -0.2 , respectively (Fig. 4D–F and G). In the middle and outer rings, OLIG2 and SYP are significantly downregulated with log₂FC values ranging from ~ -0.6 to -0.7 and -0.5 to -0.6 , respectively.

Immunohistochemistry using NeuN as a marker for neuronal nuclei

The number of NeuN+ cells was counted on a per-region basis. The naïve control tissue was evaluated to create a baseline for

comparison to implanted tissue. Here, baseline NeuN+ cell density was determined to be ~ 1534 neuronal nuclei per square millimeter (NN mm^{-2}). Both 4-weeks and 8-weeks post-implantation groups showed significant decreases in nuclear density within the inner ring (0–90 μm). The 4-week post-implantation inner ring averaged ~ 860 NN mm^{-2} , while the 8-week post-implantation inner ring nuclear count was ~ 1124 NN mm^{-2} . Both inner ring comparisons had a $p < 0.05$, in the 4-week comparison $p = 0.00006$ and in the 8-week comparison $p = 0.0202$. The middle rings (90–180 μm) in the 4-week and 8-week post-implantation groups showed no significant deficits in neuronal nuclear density, averaging ~ 1370 and ~ 1540 NN mm^{-2} , respectively. In the 4-week middle ring comparison, $p = 0.2786$, and in the 8-week middle ring





Fig. 5 Probability density functions of measured proteins. The individual probability density functions show the distribution of protein expression between samples for the inner, middle, and outer rings of each protein within experimental and control groups.

comparison, $p = 0.9681$. Like the middle rings, the outer rings (180–270 μm) showed no significant neuronal deficits, with the 4-week group average of $\sim 1386 \text{ NN mm}^{-2}$ ($p = 0.2710$) and the 8-week group average of $\sim 1528 \text{ NN mm}^{-2}$ ($p = 0.9667$) (Fig. 6).

When comparing the 4-week group and the 8-week groups, the inner rings were not significantly different from one another ($p = 0.1193$), and neither were the middle or outer rings ($p = 0.2487$ and 0.2581 , respectively). When comparing rings within 4- and 8-week time points, for both groups, the inner rings were significantly different from both the middle and outer rings ($p < 0.02$ in all cases), but the middle and outer rings were not significantly different than one another ($p > 0.9$ in both cases).

Consistent with a previous study from our lab,⁶⁹ both the middle and outer rings show reduced nuclear count in the 4-week post-implantation group compared to baseline, though the effect is not statistically significant (Fig. 6). To determine what sample size would be required to reject the null hypothesis at a significance level of $p < 0.05$, we ran a power analysis. Our power analysis revealed that in the 4-week implanted vs. naïve control middle ring comparison, a sample size of ~ 100 per group would be required to yield a $p < 0.05$. The 4-week outer ring comparison would require ~ 96 samples per group to yield a $p < 0.05$. A distinction between the current and prior study was that the binning intervals used here were 90 μm while previous studies have used 50 μm intervals.



Fig. 6 Bar graph of neuronal nuclei density as a function of time and distance from the implant, showing inner (0–90 μm), middle (90–180 μm), and outer (180–270 μm) rings. Error bars represent the standard error of each group. Asterisks (*) indicate significance ($p < 0.02$) compared to a baseline of naïve control mice. The baseline neuronal nuclear density is $\sim 1534 \text{ NN mm}^{-2}$, indicated on this plot with a dotted line.

Discussion

Analysis of neuronal and oligodendrocyte health using multiplex immunocytochemistry protein panel

In the present study, we aimed to conduct a thorough evaluation of proteins that would provide insights into the effect of



site, of both the 4-week and 8-week post-implantation groups have neuronal damage or loss from the MEA implantation. The middle and outer rings for both time points showed no significant deficits compared to naïve controls (Fig. 6). Our power analysis, which resulted in a required sample size of ~100 for both 4 weeks middle and outer rings, confirmed that in both comparisons, NeuN neuronal nuclear counts were not sufficient to indicate damage within a reasonable sample size. However, the multiplex protein panel of MAP2, MBP, NfL, OLIG2, and SYP suggested a larger region of neurodegeneration over an extended time course. Together, our combined histological analysis suggests that certain elements of the neuron show decreased protein expression only near the implant site (decreased NeuN expression in the nucleus), while others have deficits that do not manifest until the chronic time points over the broader 270 μm region of interest explored here (MAP2, NfL, SYP, MBP and OLIG2 downregulation).

IHC and the cell-specific protein expression assay performed in this study represent two methods evaluating the same molecules: proteins, but with very different approaches. IHC is quicker, easier, and can generate images mapping the exact locations of each protein. IHC is extremely useful for proof-of-concept experiments or applications where knowing the location is vital. Other cell-specific proteomic methods, such as FACS-sorted mass spectrometry, can give the advantage of quantifying thousands of proteins at once, but typically are performed in bulk on isolated samples. The isolated samples cannot be reconstructed to determine the exact location of that protein in the brain.^{91,92} The NanoString DSP provides exact expression data within specified locations, depending on the user's ROIs. The advantages of the DSP methodology are preserving the sample *in situ* and the customizability of the ROIs. However, it is limited in the number of proteins it can quantify based on available panels. Each method has its own pros and cons, and researchers should explore different options to determine what best fits the needs of their experiment.

Implications for recording and neuronal health post-MEA implantation

Single-unit recordings from intracortical MEAs have been shown to decline in quality with time.⁹³ Optimizing MEA performance is crucial to clinical applications due to the invasiveness of the surgical implant procedure and the persisting need for BCIs to perform basic tasks. Traditional analyses of the MEA biological response have utilized immunohistochemistry to visualize and provide a semiquantitative assessment of the progression of both neuroinflammation and neurodegeneration.^{15,94} Based largely on histological reports, the general understanding in the field of neural interfacing is that the neuroinflammatory response to intracortical MEA implantation is a major contributor to the decline in signal quality.^{16,20,95,96} However, there is a disconnect between the documented timeline of neuronal "dieback" adjacent intracortical MEAs^{69,97} the decline in recording performance,⁹³ and the required proximity of neuronal nuclei to obtain single-unit recordings.⁹⁸ The density of neurons around the implant site has been described as the most accurate barometer for chronic

electrode performance.³⁸ Promoting neuronal growth and survival has been shown to improve cortical recordings, meaning that neuronal health is at least a partial contributor to chronic failure.

NeuN has historically been used to denote neuronal loss or "dieback".³⁸ However, injured, or damaged neurons can also temporarily stop expressing NeuN, while NeuN has also been shown to lose immunoreactivity while still being present in injured or damaged tissue.⁴⁸ NeuN IHC results can also vary between MEA studies. Some NeuN IHC results show a general trend of an initial decrease and steady increase toward a healthy baseline over time (Fig. 6),^{13,42} while others show variations from the standard trend, specifically a decline in NeuN over chronic time points.^{1,97}

The purpose of adding NeuN staining to this study was to determine if, within the same tissue, neuronal nuclei counts obtained from NeuN immunohistochemistry and multiplex immunochemistry protein quantification of other neuronal proteins would yield the same conclusion concerning neuronal health. Our findings suggest that they do not. Neurons may still have NeuN proteins present in their nuclei while exhibiting major defects in structure and functionality. To be useful to an organism, neurons need to be able to conduct bioelectric signals, which would be impeded by diminished MAP2 and NfL for maintaining the structure of the neuron and its connections, or SYP, a major structural protein in synaptic vesicles that allows the exocytosis of neurotransmitters. MBP and OLIG2 may not be expressed in neurons directly, but their loss would limit the conduction of action potentials down the length of the axon.⁸⁷

Our goal is to better understand the correlation, if any, between the histological response to MEA implantation and the time course of signal degradation. In the current study, NeuN expression increases from 4 weeks to 8 weeks post-implantation, while recording performance typically declines over the same interval – suggesting NeuN to be an unreliable marker for MEA recording performance (Fig. 6). MAP2, NfL, SYP, MBP, and OLIG2 are not significantly differentially expressed at 4 weeks after MEA implantation but are significantly downregulated in protein expression at 8 weeks post-implantation. The expression of the 5 proteins quantified in this study more closely correlates with documented trends in decreased MEA performance – suggesting that a combination of MAP2, NfL, SYP, MBP, and OLIG2 could represent more appropriate histological markers than NeuN alone to predict MEA performance. Finally, strategies to prevent the downregulation of MAP2, NfL, OLIG2, and SYP or approaches to increase the expression of MAP2, NfL, OLIG2, and SYP may prove valuable in improving the chronic recording performance of intracortical MEAs.

Ultimately, a major goal of our lab and others investigating the neuroinflammatory response to MEAs is to improve chronic recording performance due to the improved quality of life MEAs and BCIs could offer patients. However, neuronal health impacts more than just recording performance in live human patients. MEA implantation in the motor cortex of healthy rats causes fine motor deficits,⁹⁹ showing that the device causes



enough damage to noticeably affect brain function. MEAs have been applied beyond the motor cortex, to the greater frontal cortex, such as the sensory cortex or Broca's area, where neuronal damage could affect a person's senses or speech.^{100,101} Preserving neurons that cannot be regenerated is crucial to the safety of MEAs in clinical applications.

Conclusions

This study intended to utilize cell-specific, spatially resolved multiplex protein analysis techniques to perform a comprehensive analysis of neuronal health following MEA implantation. We found that MAP2, NfL, SYP, MBP, and OLIG2 significantly decreased expression at 8 weeks post-implantation but not 4 weeks post-implantation. We directly compared the multiplex protein panel results to the field standard, NeuN immunohistochemistry, to assess the utility of histological protocols to predict or correlate with the recording performance of intracortical MEAs. The expression profiles for MAP2, NfL, SYP, MBP, and OLIG2 more closely correlate with documented trends in decreased MEA performance – suggesting that a combination of MAP2, NfL, SYP, MBP, or OLIG2 could represent more appropriate histological markers than NeuN to predict MEA performance.

Author contributions

LD and NK contributed to the conception and design of the work. LD, NK, SS, and JW contributed to the methodology, software analysis, validation, formal analysis, investigation, and data curation. ERC guided the statistical analysis of the work. LD wrote the original draft. LD, NK, JW, and JRC prepared figures. All authors edited the final manuscript. JRC provided the funding and resources to conduct the study.

Disclosures

The contents do not represent the views of the U.S. Department of Veterans Affairs, the National Institutes of Health, or the United States Government.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no conflict of interest.

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