A 3-D 160-Site Microelectrode Array for Cochlear Nucleus Mapping

Mary Elizabeth Merriam, Susanne Dehmel, Onnop Srivannavit, Susan E. Shore, and Kensall D. Wise*, Life Fellow, IEEE

Abstract—A 3-D application-specific microelectrode array has been developed for physiological studies in guinea pig cochlear nucleus (CN). The batch-fabricated silicon probes contain integrated parylene cables and use a boron etch-stop to define 15 μm-thick shanks and limit tissue displacement. Targeting the ventral (three probes) and dorsal (two probes) subnuclei, the custom four-shank 32-site probes are combined in a slotted block platform having a 1.18-mm² footprint. The device has permitted, for the first time, high-density 3-D in vivo studies of ventral CN to dorsal CN connections, stimulating with 1000 μm² sites in one subnucleus while recording with 177 μm² sites in the other. Through these experiments, it has demonstrated the efficacy of bimodal silicon arrays to better understand the central nervous system at the circuit level. The 160 electrode sites also provide a high-density neural interface, which is an essential aspect of auditory prosthesis prototypes.

Index Terms—Auditory brainstem prosthesis, cochlear nucleus, microelectrode array, neural mapping, silicon probe, 3-D microassembly of microelectromechanical devices.

I. INTRODUCTION

Advances in neuroscience depend on the availability of supporting technology for the high-density stimulation and recording of neural activity. While electrode arrays can be highly versatile, individual neuroscience applications may require particular array specifications for optimal performance [1].

As alternatives to cochlear implants, research on a central auditory prosthesis is targeting the auditory nerve [2], the cochlear nucleus (CN) [3], [4], and the inferior colliculus [5] to lower thresholds, increase dynamic range, and improve frequency discrimination [2], [6]. Placement within the CN may be advantageous since, although, sound coding is divided into parallel channels, the more complex signal-processing functions are left to the higher portions of the auditory system [4]. While research on animal models and clinical trials has shown the potential for such devices, significant technological and neuroscience questions must be addressed before high-quality auditory brainstem implants (ABIs) become a reality. Increasing the number of implanted electrode sites and properly tailoring their 3-D locations to the targeted neural structures is one essential aspect [1], [4]. Once the proper supporting technology exists, the neuroscience experiments needed to determine the sound encoding strategies that will be most successful can be pursued.

The array presented here is an application-specific design with 160 electrode sites to enable the mapping of neural connections between the ventral CN (VCN) and dorsal CN (DCN) in guinea pig [7], [8]. It can also be used as an acute prototype for a central auditory prosthesis.

II. ARRAY DESIGN

A. Anatomical and Surgical Constraints

While the basic anatomical structure of the guinea pig CN and its placement within the brain is well known [9], the numerical details required for a single 3-D array to simultaneously reach the antero-ventral portion (AVCN) and the DCN are not readily available from current literature. In order to acquire the necessary information for probe and platform specifications, mock arrays with fluorescent fluorogold (FG) dye applied to the shanks were inserted and 3-D reconstructions of serial sections were assembled.

The CN is located in the auditory brainstem. The operative procedure involves removing part of the skull and overlying cerebellum to reach the DCN, as shown in Fig. 1. The access hole is made as small as possible to minimize the surgical impact. While the surgery and cerebellum removal directly exposes the surface of the DCN, the AVCN is located farther ventral, lateral, and rostral. Thus, the VCN shanks pass through the cerebellum to reach the AVCN. The small diameter of the access hole and the angle at which the array must be inserted leads to considerable size and positioning constraints on the electrode array, particularly on the platform and any required cabling. The difference between the mediolateral location of the DCN and the AVCN necessitates offset between the platform positions of the probes designated for the respective regions. The relative dorsoventral and rostrocaudal positions of the CN subdivisions determine the insertion angle and relative probe lengths.

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B. 2-D VCN–DCN Probes for Use in the 3-D Array

For a high-density interface, the VCN–DCN array consists of five 32-site four-shank probes secured in a mounting platform to form a 160-site device. Two probe designs were developed: one for VCN and the other for DCN; specifications are listed in Table I. Both designs had a maximum shank width of 61 μm in order to minimize any disruption to the neural circuitry during implantation. This was achieved by using an interconnect line width of 2 μm when necessary. The shanks also taper in the region, where the sites are located to keep the width at a minimum as the interconnect lines reach their designated endpoints.

1) Ventral Cochlear Nucleus Probes: Three of the five probes in the array are targeted at VCN. The VCN probe design has four shanks at a pitch of 200 μm, as shown in Fig. 2. The total length from the bottom of the platform to the center of the tip-most site is 5 mm. The merged shank region at the base of the array remains outside the cerebellum after implantation and spans the distance to the platform required by the insertion angle and insertion depth. By merging this region, the strength of the probe was increased; the impedances of the interconnect lines were also decreased since their line widths could be increased there.

Recording and stimulation offer conflicting constraints on electrode site area. Although single units may be more easily isolated with smaller sites, the maximum safe stimulating current levels would be substantially less with smaller area higher impedance sites. In order to both record and stimulate effectively with the VCN probe, the sites were arranged in pairs. The smaller site of each pair was designated for recording and had a site area of 177 μm². The larger site of each pair had an area of 1000 μm² and was intended for stimulation. There are four-site pairs per shank at a pitch of 300 μm for a total of eight sites per shank and thirty-two sites per probe. The site designations for recording and stimulation are purely based on site area; beside the aforementioned discussion, any of the sites are suitable for recording or stimulation.

2) Dorsal Cochlear Nucleus Probes: The remaining two probes were designated for the DCN, with four shanks at a pitch of 125 μm, as illustrated in Fig. 2. The desired recording locations in the DCN are within the top layers, up to a few hundred micrometers deep. Since the surface of the DCN is dome shaped, the shanks were designed with staggered lengths to account for the surface curvature and maintain recordings within the top layers across a given mediolateral probe position. The length of the lateral-most shank from the bottom of the platform to the center of the tip-most site is 3400 μm; subsequent shanks each decrease in length by 125 μm. There are eight equally spaced sites per shank at a pitch of 100 μm. All of the sites have an area of 177 μm²; although this site size is more appropriate for recording, these sites may also be used for stimulation.

C. 3-D Structure and Block Platform

The 3-D microassembly of boron-doped silicon probes has been previously achieved using a thin platform through which 2-D probes were placed and secured with spacers [10]–[13].
However, for large numbers of leads, this approach results in substantial wings on the ends of the probes, where the probe-platform lead transfers are made, increasing the size of the implant. More recently, an improved structure has been developed, in which the probes are inserted in slots formed using deep reactive ion etching (DRIE) in a platform formed from full-thickness (500 μm) 4-in silicon wafers [14]. The probes are held parallel by designing the vertical slots only slightly larger than the probes themselves, eliminating the need for any wings or platform spacers. A parylene overlay cable is bonded to the lead tabs coming from the probes [14], taking the neural signals to an adjacent signal-processing chip or directly to the external setup. The low profile of this design is especially important in chronic applications, minimizing overall implant size, and allowing the dura to be replaced over it. However, for acute applications, such as neural mapping, the structure can be further simplified by integrating the output cables directly into the probes themselves, completely eliminating any bonds or lead transfers at the implant site.

This paper employs a slotted 500-μm-thick platform with probes containing 10-mm-long integrated parylene cables. The cables route the leads from the back ends of the probes embedded in the platform out of the surgical access hole. The five cables from the probes may be stacked to form a space-saving “multilevel” interconnect, which is important for high-density designs, as in this paper. Each cable is less than three-fourths of a millimeter wide at its widest point, carrying 32 interconnect lines at a 20-μm pitch. Platform size constraints prevent expanding the platform to fit wider leads.

The platform slots mechanically secure the individual probes. Top (300-μm deep) and bottom (200-μm deep) DRIE etches form the slots and the overall platform shape. The top slot is longer than the bottom slot so that a ledge is formed, on which the probe rests when fully inserted, as illustrated in Fig. 3. The back ends of the VCN and DCN probes both employ this same design for countersinking them into the platform. The width of the platform slots is based on the fabrication thicknesses of the various probe layers. The top portion of the probe back end is 25-μm thick; the top slots are 29-μm thick, resulting in a 4-μm tolerance. The bottom slot width extends past the top by 1 μm on each side.

Several constraints determined the platform footprint for the VCN–DCN arrays. The exposed area for viewing and probe placement is extremely limited for the surgical and experimental reasons previously described. Therefore, the mediolateral and rostrocaudal dimensions of the platform must be no larger than necessary to support the probes. The narrow platform design, with a 1.18-mm² footprint, suspends probes with slots only 50 μm from the edge of the platform in one direction and 59 μm in the other. Since the AVCN is farther lateral than the DCN, the VCN probe slots are offset to the left of center in relation to the DCN slots, as presented in Fig. 4. Table II lists the platform specifications. The VCN probe slot pitch is 300 μm and the DCN probe slot pitch is 200 μm. The distance between the closest DCN and VCN probe slots is 628 μm. The VCN and DCN platform slots are perpendicular to each other in order to span the tonotopically organized subdivisions of the CN. The notched shape of the platform offers a reduced footprint over a rectangular design, allowing greater visibility of the probe shanks during surgical implantation and a lower weight for a chronic prosthesis. The 3-D 160-site array is also illustrated conceptually in Fig. 4.

III. Fabrication and Assembly

The 2-D probe process flow was based on the well-established passive-probe technology developed at the University of Michigan, which employs boron etch-stops to enable thin (~12-μm substrate) devices to be batch fabricated with integrated polymer cables [13], [15]. Table III lists the major process steps and layer thicknesses for the 2-D probes. The block platforms were fabricated using DRIE following a process similar to [14]. A photograph of fabricated devices is shown in Fig. 5.

A custom micromachined jig was used to hold the platform during assembly. In this 500-μm-thick frame, 250-μm-deep
TABLE II
PLATFORM SPECIFICATIONS FOR THE VCN–DCN 3-D ARRAY

<table>
<thead>
<tr>
<th></th>
<th>VCN Area</th>
<th>DCN Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (rostro-caudal)</td>
<td>947µm</td>
<td>335µm</td>
</tr>
<tr>
<td>Width (medio-lateral)</td>
<td>735µm</td>
<td>948µm</td>
</tr>
<tr>
<td>Max Probes per Array</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Probe Pitch</td>
<td>300µm</td>
<td>200µm</td>
</tr>
<tr>
<td></td>
<td>Overall</td>
<td></td>
</tr>
<tr>
<td>Total Length</td>
<td>1800µm</td>
<td></td>
</tr>
<tr>
<td>Total Width</td>
<td>1354µm</td>
<td></td>
</tr>
<tr>
<td>Thickness (dorso-ventral)</td>
<td>500µm</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Top (Major)</td>
<td>Bottom (Minor)</td>
</tr>
<tr>
<td>Slot Thickness</td>
<td>29µm</td>
<td>31µm</td>
</tr>
<tr>
<td>Slot Width</td>
<td>830µm</td>
<td>700µm</td>
</tr>
<tr>
<td>Slot Depth</td>
<td>300µm</td>
<td>200µm</td>
</tr>
<tr>
<td>Rim Etch</td>
<td>16µm</td>
<td>17µm</td>
</tr>
</tbody>
</table>

TABLE III
PROBE FABRICATION STEPS AND TARGET PARAMETERS

<table>
<thead>
<tr>
<th>Material</th>
<th>Thickness</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deep &amp; Shallow Boron Substrate</td>
<td>12µm</td>
<td>Diffusion</td>
</tr>
<tr>
<td>Pad Oxide</td>
<td>1500Å</td>
<td>Thermal</td>
</tr>
<tr>
<td>Oxide/Nitride/Oxide</td>
<td>3000Å/1500Å/3000Å</td>
<td>LPCVD&lt;sup&gt;4&lt;/sup&gt;</td>
</tr>
<tr>
<td>Doped Polysilicon Leads</td>
<td>6000Å</td>
<td>LPCVD / Etch</td>
</tr>
<tr>
<td>Pad Oxide</td>
<td>1500Å</td>
<td>Thermal</td>
</tr>
<tr>
<td>Oxide/Nitride</td>
<td>3000Å/1500Å</td>
<td>LPCVD</td>
</tr>
<tr>
<td>Large Contact</td>
<td>--</td>
<td>RIE&lt;sup&gt;3&lt;/sup&gt;</td>
</tr>
<tr>
<td>Oxide</td>
<td>3000Å</td>
<td>LPCVD</td>
</tr>
<tr>
<td>Small Contact</td>
<td>--</td>
<td>RIE</td>
</tr>
<tr>
<td>Titanium (Ti)/Iridium Sites (Ir)</td>
<td>500Å/3000Å</td>
<td>Sputtered / Lift off&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Chromium (Cr)/Gold Pads (Au)</td>
<td>500Å/300Å</td>
<td>Sputtered / Lift off&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Dielectric Etch</td>
<td>--</td>
<td>RIE</td>
</tr>
<tr>
<td>Bottom Parylene</td>
<td>2µm</td>
<td>Deposition / Etch</td>
</tr>
<tr>
<td>Chromium (Cr)/Gold Leads (Au)</td>
<td>250Å/1000Å</td>
<td>Sputtered / Lift off&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Silicon Anchor</td>
<td>--</td>
<td>DRIE&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Top Parylene</td>
<td>3µm</td>
<td>Deposition / Etch</td>
</tr>
<tr>
<td>Thinning</td>
<td>--</td>
<td>DRIE</td>
</tr>
<tr>
<td>Release</td>
<td>--</td>
<td>TMABH&lt;sup&gt;4&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Totals</th>
<th>Thickness</th>
<th>Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>Back-end Top</td>
<td>24.875µm</td>
<td>All except 6</td>
</tr>
<tr>
<td>Back-end Bottom &amp; Shanks</td>
<td>14.75µm</td>
<td>Through step 8</td>
</tr>
</tbody>
</table>

<sup>1</sup>Low-Pressure Chemical Vapor Deposition  
<sup>2</sup>Reactive Ion Etching  
<sup>3</sup>Deep Reactive Ion Etching  
<sup>4</sup>Tetramethylammonium Hydroxide

Fig. 5. Photograph of fabricated probes and platforms. Five platforms are shown; two are standing on their edge.

recesses were etched by DRIE for multiple platforms to enable the assembly of several arrays at once. The probes were positioned in the platform slots with a vacuum wand attached to a micromanipulator. A small amount of medical grade silicone (Med-4211, NuSil Technology LLC) was used to secure the probes in place and to seal the top and bottom ends of the platform slots.

Fig. 6. Photograph of a five-probe array in the Plexiglas workholder after wire bonding between the integrated parylene cables and hybrid polyimide cables.

Fig. 7. Photograph of a 160-site 3-D VCN–DCN mapping array on the back of a U.S. dime.

The parylene cables terminate in an integrated silicon bonding region. Ultrasonic wire bonding was used to connect them to flexible 25-cm-long polyimide printed circuit board (PCB) cables (manufactured by SunTech Circuits, Inc.), which attach to a site-selection board. A custom Plexiglas work holder, shown in Fig. 6, restrained the array to enable bonding between the integrated parylene and hybrid polyimide cables. Fig. 7 shows an assembled five-probe array.

IV. In Vivo Results

A. Surgical Approach and Verification of Probe Placement

The typical surgical preparation followed that of previously reported work with other electrodes [16]. The work was in accord with the National Institutes of Health (NIH) Guidelines for the Use and Care of Laboratory Animals (NIH publication
No. 80–23) as well as with the guidelines by the University of Michigan Committee on the Use and Care of Animals. Pigmented guinea pigs from Cady Ridge Farms (Chelmsford, MA, USA) of young adult size/age (300–800 g) were anesthetized using ketamine (40 mg/kg) and xylazine (10 mg/kg). A stereotactic frame stabilized the subject during surgery and recording, which took place in an electrically shielded and double-walled sound booth. The surgery included a craniotomy and partial aspiration of the cerebellum overlying the DCN. Hollow earbars joined to speakers (Beyer DT770pro) provided the acoustic stimulus.

The VCN–DCN 3-D array was stereotactically implanted, as shown in Fig. 8. The location of the array in the tissue was marked by dipping the shanks in FG dye prior to insertion and with an electrolytic lesion at the end of the experiment. Placement of electrode sites in DCN and AVCN were confirmed in serial brain sections. Animal ground was defined by connecting the neck muscles to the ground and reference pins of the head stage (RA16AC, Tucker Davis Technologies).

B. Single and Multunit Recordings

The primary purpose of the VCN–DCN array is to simultaneously record from both CN subdivisions. This was achieved repeatedly during in vivo experiments. Fig. 9 presents the clips of neural spikes for one of the recorded units. The complexity of the CN is underscored by its diverse cytoarchitecture. Over ten cell types are grouped in part by location [9]; the peri-stimulus time histogram (PSTH) of a CN neuron is one indicator of cell type. A spread of response patterns has been observed from recordings with the VCN–DCN array; a selection of these responses is presented in Fig. 10 for both subnuclei.

C. Tonotopic Mapping

It is well known that the CN is tonotopically organized, representing a spatial array of all frequencies that are coded by the incoming auditory nerve. This tonotopic pattern is repeated within each subdivision. In some experiments, such as those investigating aspects required for auditory prostheses, it is desirable to simultaneously record (or stimulate) from neurons with similar characteristic frequencies (CFs) in both subnuclei. The high-density 3-D array enables one to plot the CFs with respect to instrumented location. The recorded frequency map aids the researcher in intuitively visualizing corresponding CFs and their anatomical location. Fig. 11 shows recorded CF maps for two probes.
work to date demonstrates that the application-specific design permits high-density recording in vivo, enabling studies not heretofore feasible. Experiments exploring circuits. DCN array developed here for studying the underlying neural anatomical structures and in particular the 160-site 3-D VCN–DCN channels, the bimodal (stimulate/record) approach to anterograde and retrograde tracing studies, enables the study of the living network from the perspective of particular connections. Previous studies have investigated intrinsic pathways within the CN [17]–[21]. To expand on this understanding, in this paper, the bimodal (stimulate/recording) approach was used to explore the connections between the AVCN and DCN. The stimulus configuration consisted of 50 pulse trains at 20 Hz of 10 current pulses each. The pulses were charge balanced, biphasic, negative first, and of 10-μA amplitude. Fig. 12 shows elicited responses of a selected DCN channel before and during electrical stimulation on one channel in VCN; the response pattern shows a decrease in spike count immediately after the stimulus followed by significant excitation and then further inhibition. While further experiments are necessary before conclusions can be made about the connectivity between VCN and DCN channels, the in vivo work to date demonstrates the value of high-density electrode arrays tailored to specific anatomical structures and in particular the 160-site 3-D VCN–DCN array developed here for studying the underlying neural circuits.

D. Intersubnuclei Mapping

The neural architecture is composed of complex circuits and subcircuits. Electrical stimulation, as a complementary approach to anterograde and retrograde tracing studies, enables the study of the living network from the perspective of particular connections. Previous studies have investigated intrinsic pathways within the CN [17]–[21]. To expand on this understanding, in this paper, the bimodal (stimulate/recording) approach was used to explore the connections between the AVCN and DCN. The stimulus configuration consisted of 50 pulse trains at 20 Hz of 10 current pulses each. The pulses were charge balanced, biphasic, negative first, and of 10-μA amplitude. Fig. 12 shows elicited responses of a selected DCN channel before and during electrical stimulation on one channel in VCN; the response pattern shows a decrease in spike count immediately after the stimulus followed by significant excitation and then further inhibition. While further experiments are necessary before conclusions can be made about the connectivity between VCN and DCN channels, the in vivo work to date demonstrates the value of high-density electrode arrays tailored to specific anatomical structures and in particular the 160-site 3-D VCN–DCN array developed here for studying the underlying neural circuits.

V. CONCLUSION

The device presented in this paper represents one of the most advanced high-density neural interfaces ever reported and is the first 3-D array with over 100 sites for mapping in the guinea pig VCN–DCN. The VCN–DCN array has not only been used during in vivo experimentation, which proved its functionality, but it has also contributed to neuroscience research enabling studies not heretofore feasible. Experiments exploring auditory-somatosensory integration continue to use this device; the application-specific design permits high-density recording in two CN subnuclei with a commercial stimulation probe in the spinal trigeminal nucleus. On-going studies will focus on the intrinsic connections between the VCN and DCN subregions using bimodal functionality to simultaneously stimulate electrically and record the elicited waveforms. Additional experiments are also being planned using this array as a prototype for an auditory prosthesis. This application-specific device is not only a high-density interface for the guinea pig CN, but also stands as the forerunner for future silicon-based electrode arrays, which focus on target-specific neuroscience and medical needs, including neural mapping and prosthetic system prototypes.

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REFERENCES


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Dr. Wise organized and was as the first chairman of the Technical Subcommittee on Solid-State Sensors of the IEEE Electron Devices Society (EDS). He was the General Chairman of the 1984 IEEE Solid-State Sensor Conference, was an IEEE-EDS National Lecturer (1986), and was a Technical Program Chairman (1985) and General Chairman (1997) of the IEEE International Conference on Solid-State Sensors, Actuators, and Microsystems. He was the recipient of the Paul Rappaport Award from the EDS (1990), a Distinguished Faculty Achievement Award from the University of Michigan (1995), the Columbus Prize from the Christopher Columbus Fellowship Foundation (1996), the SRC Aristotle Award (1997), and the 1999 IEEE Solid-State Circuits Field Award. In 2002, he was named the William Gould Dow Distinguished University Professor at the University of Michigan. He held the 2007 Henry Russel Lectureship at the University and is a Fellow of the American Institute of Medical and Biological Engineering, and a member of the United States National Academy of Engineering.