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1 Tracking neural activity from the same cells during the entire adult life of mice

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12 Abstract

Recording the activity of the same neurons over the adult life of an animal is important to 13 14 neuroscience research and biomedical applications. Current implantable devices cannot provide 15 stable recording on this time scale. Here, we introduce a method to precisely implant nanoelectronics with an open, unfolded mesh structure across multiple brain regions in the mouse. 16 17 The open mesh structure forms a stable interwoven structure with the neural network, preventing probe drifting and showing no immune response and neuron loss during the yearlong implantation. 18 Using the implanted nanoelectronics, we can track single-unit action potentials from the same 19 neurons over the entire adult life of mice. Leveraging the stable recordings, we build machine 20 21 learning algorithms that enable automated spike sorting, noise rejection, stability validation, and 22 generate pseudotime analysis, revealing aging-associated evolution of the single-neuron activities.

23 Main

Long-term stable recording¹⁻⁴ of the same neuron at single-cell and single-spike resolution over 24 25 the entire adult stage of life of behaving animals is important to understand how neural activity changes with learning and age^{4,5,6,7}, to improve current brain-machine interface performance by 26 reliably interpreting the brain's behavioral and internal states^{4,6}, and to study neurodegenerative 27 28 diseases, aging-associated neurological disorders and cognitive decline^{7,8}. Current implantable electronic and optical tools can record neural activity at single-cell and single-spike resolution but 29 suffer from immune response and recording drift due to the mechanical and structural disparities 30 between rigid electronic or optical devices and brain tissue^{9,10}. Relative shear and repeat motion at 31 the implanted interface keep changing the relative position between recording devices and 32 33 recorded neurons. The proliferation of astrocytes and microglia form a $\sim 100 \,\mu m$ thick glial sheath that cause the death of neurons and isolate recording devices from neurons. Together, they lead to 34 chronic instability of recordings. Optical imaging techniques are further limited by the light 35 36 penetration depth and three-dimensional (3D) volumetric scanning across the 3D tissue due to optical aberration and attenuation¹¹. 37

38 Miniaturized flexible electronics such as mesh nanoelectronics and thin-film probes have been utilized in *in vivo* electrophysiology¹²⁻¹⁷ given their unique mechanical properties. Mesh 39 nanoelectronics provide a chronically stable, gliosis-free implantation over a few months through 40 the incorporation of tissue-like structural and mechanical properties into nanoelectronics¹⁸⁻²⁰. 41 However, due to their mechanical flexibility, invasive methods such as syringe injection are 42 required to implant the tissue-like electronics into the brain¹⁸⁻²⁰. The relatively large mechanical 43 damage from the implantation causes permanent damage to the neural network. In addition, 44 45 implanted mesh nanoelectronics can only unfold in the cavities of the brain such as the

subventricular zone, not in tissue-dense brain regions¹⁸⁻²¹. As a result, the bundled mesh 46 nanoelectronics do not have the optimized mechanical flexibility required to interface with the 47 brain tissue over long time periods¹⁸⁻²¹. On the other hand, flexible thin film brain probes lack the 48 open mesh structures that allow for 3D integration with the neural network²²⁻²⁴. They also need to 49 maintain mechanical strength to prevent damage of the probe during the implantation²²⁻²⁵, which 50 51 can cause an immune response and probe drifting during long-term implantation. As a result, none of the existing technologies has demonstrated long-term stable tracking of the same neuron over 52 the entire adult stage of life of a behaving animal. 53

Here, we solved this issue by implanting a fully unfolded tissue-like mesh nanoelectronics into the 54 brain of a mouse. The fully unfolded mesh can form an interwoven structure within the neural 55 network and eliminate the immune response and probe drifting, maintaining a long-term stable 56 electrode-to-neuron interface at the single-cell level, thus enabling the same neuron to be recorded 57 over the entire adult life of animals (Fig. 1a). To enable this method, we developed mesh 58 59 nanoelectronics monolithically integrated with ultra-thin and releasable polymer shuttles through lithographic fabrication. We also incorporated unique polymer anchors and water-releasable 60 structures, allowing for controllable, precise, and minimally invasive delivery of mesh 61 62 nanoelectronics in the mouse brain. This implantation method can keep the designed open mesh structure of the device in the brain across multiple brain regions, including cell-dense regions. The 63 open mesh allows the neural network to form seamlessly interwoven structures, provides a tissue-64 level flexible interface, and prevents the repeated micromotion and drift between recording 65 electrodes and surrounding neural tissue during yearlong recording, thus allowing for a highly 66 67 stable recording of neuron activities across multiple brain regions.

By optimizing the size of the implanted nanoelectronics, we achieved stable tracking of the same 68 neuron over the entire adult life of mice until their natural death (*i.e.*, 5-18 months for mouse #1, 69 70 5-20 months for mouse #2, and 5-19 months for mouse #3), confirmed by vigorous statistical tests^{3,18-21,26}. Tracking the whisker-stimulation-evoked single spikes from the barrel cortex 71 indicated that the electrode embedded in a single whisker barrel does not drift over the animal's 72 73 adult life. Using the first several months' recording data to train an autoencoder²⁷, a machine learning (ML) tool for representation learning, we further confirmed the stable recording, which 74 also allowed for fully automatic spike sorting, noise rejection, and stability analysis over the adult 75 life of the mice. Finally, ML-based pseudotime analysis of single-unit waveforms identified 76 several neurons with age-dependent changes in electrical activities. 77

78

79 **Results**

80 Monolithically integrated mesh nanoelectronics and ultra-thin shuttle

To implant completely open and unfolded mesh nanoelectronics into the brain, we integrated the 81 mesh nanoelectronics monolithically with a releasable, ultra-thin polymeric shuttle using standard 82 photolithography procedures (Fig. 1b, Extended Data Fig. 1, 2a-k, Methods). The mesh 83 nanoelectronics were fabricated as described in previous reports¹⁸⁻²⁰. Briefly, 16 or 32 15-µm-84 85 diameter electrodes were connected by SU-8 encapsulated Cr/Au interconnects to Cr/Au input/output (I/O) pads. The encapsulated interconnects were 10-µm wide and <1-µm thick, 86 forming a mesh network with a 2D filling ratio at 73.3%, which yielded an effective bending 87 stiffness of 1.26×10⁻¹⁵ N·m². The mesh nanoelectronics were partially fabricated on the top of a 88 Ni sacrificial layer. Next, a 25-µm-thick polymer shuttle was defined on the top of the mesh 89

nanoelectronics with a 3-µm-thick water-soluble dextran and 20-µm-thick polymer anchors. These 90 polymer anchors (Fig. 1b, inset) connected the mesh nanoelectronics with the polymer shuttle 91 92 through the dextran layer (Fig. 1b, red box). After the integrated device was released from the 93 substrate (Fig. 1c, top) by removing the Ni sacrificial layer, the anchor kept the pattern of the mesh nanoelectronics on the polymer shuttle. Then, a few drops of 10 wt% PEG (35 kDa) was coated to 94 reinforce the bonding between the mesh nanoelectronics and polymer shuttle (Fig. 1c, bottom), as 95 well as enhanced the temporary stiffness and provided protection to the mesh nanoelectronics 96 during implantation. The biodegradable PEG adhesion layer has a sub-micron thickness; thus, the 97 surgical footprint is mostly affected by the thin polymer shuttle. After removing the anchor 98 connection (Fig. 1c, bottom), the mesh nanoelectronics can be readily released from the polymer 99 shuttle by dissolving the PEG in an aqueous solution (Fig. 1d). 100

The polymer shuttle was used to guide the implantation of the mesh nanoelectronics into the brain 101 tissue (Fig. 1e, left). After the integrated brain probe reached the targeted position, saline was 102 103 applied to quickly dissolve the PEG and to release the mesh nanoelectronics from the polymer shuttle, which was subsequently withdrawn from the brain tissue (Fig. 1e, right). We characterized 104 the implantation procedure in a transparent 0.6 wt% agarose gel-based brain phantom with 105 mechanical properties comparable to that of the brain tissue²⁸. At a typical implantation speed of 106 100 μ we did not observe any buckles on the probes (Fig. 1f. left). After insertion, 1× 107 phosphate buffered saline (PBS) solution was applied to dissolve the PEG/dextran adhesive layer. 108 The shuttle was then withdrawn at a speed of 10 µm/s. After withdrawing the shuttle, the mesh 109 nanoelectronics still maintained its implantation location and open mesh structure without any 110 111 deformations (Fig. 1f, right). We tested the yield of the implantation in brain phantoms with various speeds. We achieved a 93.3% yield for insertion and extraction of 16-channel mesh 112

113 nanoelectronics at 100 μ m/s (Extended Data Fig. 2l, m). There was no significant change of 114 implantation yield with different size mesh electronics of 32 channels (n = 3, p > 0.05, Extended 115 Data Fig. 2l, m).

The optimized implantation procedure was then used for mouse brain implantation. Figure 1g 116 shows typical implantation of 300-µm-wide 16-channel mesh nanoelectronics in the anesthetized 117 118 mouse brain (Methods). The probe (Fig. 1g, inset) can be easily implanted with the polymer shuttle withdrawn by the same conditions tested for the phantom gel. Damages to the blood vessels were 119 minimal throughout the imaging-guided implantation. To evaluate acute tissue damage, cell loss, 120 and mesh nanoelectronics distribution, we imaged the post hoc fixed and stained tissue slices 121 immediately after implantation (Fig. 1h). Acute damaged area with the thin-shuttle was 122 approximately $0.0068 \pm 0.0016 \text{ mm}^2$ (mean \pm SD, n = 5, Fig. 1h), which is significantly smaller 123 than those from previous reported implantations using syringe-injection^{18,19} $(0.0164 \pm 0.0033 \text{ mm}^2)$, 124 mean \pm SD, p < 0.01, n = 5, Fig. 1h) and biodegradable shuttles^{21,29} (0.0182 \pm 0.0039 mm², mean 125 \pm SD, p < 0.01, n = 5, Fig. 1h). While the tissue damage showed no significant difference compared 126 with samples implanted by 50-µm diameter tungsten wire²⁵ (0.0062 \pm 0.0014 mm², mean \pm SD, n 127 = 5, Fig. 1h), the cross-sectional images of brain slices with implants showed the clear unfolded 128 mesh structures vs. bundled ribbons by tungsten probe-based delivery (Fig. 1i). Compared with a 129 previously demonstrated implantation method for flexible neural probes^{18-21,25,29,30}, the integrated 130 25-um-thick polymer shuttle drastically reduces tissue displacement during implantation as well 131 as maintains the designed open structure with nearly 90% implantation yield. Moreover, the 132 presented method involves minimal manual manipulations since the ultra-flexible nanoelectronics 133 was pre-attached to the thin-shuttle with the lithography process. On average, it took less than 3 134 min to assemble one mesh nanoelectronics (more than 20 nanoelectronics per hour), including sub-135

micro-thick PEG coating, anchor dicing, and additional packaging with a success rate approaching100%.

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139 Unfolded mesh nanoelectronics structure 3D interwoven with the neural network.

140 We implanted mesh nanoelectronics with different sizes across multiple brain regions. Each mesh structure has an ultra-small cross-section of $10 \times 1 \ \mu m^2$. The longitudinal bending stiffness of each 141 individual mesh structure reached 1.26×10^{-15} N·m², which is comparable to that of brain tissue and 142 orders of magnitude lower than state-of-the-art probes (i.e., ultrasmall carbon³¹⁻³³, polyimide³⁴ and 143 elastomer-based 'e-dura' probes³⁵). Rhodamine 6G was added to the SU-8 encapsulation layer, 144 enabling the imaging of the mesh structure in the brain. To explore the potential capability of the 145 implantation, Figure 2a shows a 2-mm-wide, 3-mm-long mesh nanoelectronics implanted into a 146 mouse brain across cortex, hippocampus, and thalamus regions. The size of this device can 147 potentially include 1,024 recording sites through 3D stacking of electrodes³⁶ (Fig. 2a-d, Methods). 148 A representative 3D reconstructed image of the mesh nanoelectronics in the brain tissue at 6-week 149 post-implantation (Fig. 2a) showed the fully unfolded, open mesh structure interweaving with 150 151 neurons and astrocytes across multiple brain regions (i.e., cortex, hippocampus, thalamus, etc.). A slight bending of the mesh suggested that the tissue-like nanoelectronics were flexible within the 152 tissue. A zoomed-in view of the hippocampus CA1 region (Fig. 2b) shows a smooth distribution 153 of neurons and astrocytes across the mesh. Notably, neurons in the cell-dense region (hippocampus) 154 can still penetrate the open mesh structure (Fig. 2c), forming an intertwined tissue-nanoelectronics 155 interface. Figure 2d shows that the size of the recording electrode (white dashed circles) is 156 comparable to the size of the soma. The subcellular feature size, tissue-level flexibility, and 3D 157 interwoven network collectively eliminated the micromotion between the functional electrode and 158

recorded neurons³⁷. Additional replications of mesh nanoelectronics with different sizes were
implanted in mouse brains and are shown in Extended Data Fig. 3.

161 Next, we performed longitudinal immunostaining characterizations to assess the distribution of key cell types around mesh nanoelectronics over the time course of implantation. To demonstrate 162 that the open mesh structure reduces immune responses during chronic implantation (Fig. 2e, top), 163 164 thin-film nanoelectronics with the same dimensions as mesh nanoelectronics were used (Fig. 2e, bottom) as control and contralaterally implanted in the same mouse brain. The bending stiffness 165 of the thin film nanoelectronics is only slightly higher than that of the mesh nanoelectronics ($39.8 \times$ 166 10⁻¹⁵ N·m² vs. 1.26×10⁻¹⁵ N·m², Methods). The brain tissue was harvested and sliced for 167 immunostaining 2-, 6-and 12-week, and 1-year post-implantation. Horizontal slices were stained 168 with cell-type-specific protein markers for imaging of neurons, astrocytes, and microglia 169 (Extended Data Fig. 4). We quantitatively analyzed horizontal brain slices implanted with 16-170 channel, 300-um-wide film/mesh nanoelectronics (Fig. 2f-i, Methods). Protein marker signals 171 172 were calculated by normalizing the fluorescence intensity around the implantation site with the baseline value defined as the average fluorescence intensity over a range of 525-550 µm away 173 from the nanoelectronics. Statistical analysis demonstrated a significant degradation of neuron 174 density (NeuN) and an enhancement of astrocytes and microglia intensity (GFAP and Iba-1, 175 respectively) near the thin-film nanoelectronics at all time points (p < 0.05, n = 5, Fig. 2f). These 176 results proved that the thin-film nanoelectronics can still trigger the proliferation of 177 astrocytes/microglia and reduced the neuron density at the nanoelectronics-brain interface. 178 Meanwhile, the open mesh nanoelectronics introduced minimal damage to the surrounding 179 180 neurons and negligible immune response. Importantly, the result demonstrated that no significant neuron loss was detected at 2-week post-implantation for mesh samples (Fig. 2f), suggesting 181

minimal acute damage from the thin-polymer shuttle. In addition, the neuron density near the meshsurface remained the same at one-year post-implantation (Fig. 2f).

184 We further calculated the normalized intensity of neural cell fluorescence signals within regions 100-µm away from the nanoelectronics to assess neuron loss and inflammation reaction at the 185 different post-implantation periods (Fig. 2g-i). The mesh nanoelectronics samples showed a 186 neuron density of $83.9 \pm 13.0\%$ (mean \pm SD, n = 5, Fig. 2g) at 2 weeks, which is greater (p < 0.001, 187 n = 5, Fig. 2g) than that from the thin film nanoelectronics (66.9 \pm 7.8%, mean \pm SD, Fig. 2g) in 188 the contralateral brain slices. Compared to the non-implanted regions, neuron intensity of mesh 189 nanoelectronics samples increased to $93.1 \pm 10.2\%$, $99.1 \pm 8.0\%$, and $102.4 \pm 11.2\%$ (mean \pm SD, 190 n = 5, Fig. 2g) 6 weeks, 12 weeks, and 1-year after implantation, respectively. On the contrary, 191 192 thin-film nanoelectronics samples showed significant neuron loss for the same periods (70.2 \pm 4.6%, 84.8 \pm 3.3%, and 85.2 \pm 2.3% at 6 weeks, 12 weeks, and 1-year after implantation, 193 respectively. p < 0.05, n = 5, Fig. 2g). The intensity of astrocytes and microglia around the mesh 194 195 slightly increased at 2 weeks ($115.1 \pm 9.1\%$ at microglia, $132.6 \pm 20.5\%$ at astrocytes, mean \pm SD, n = 5, Fig. 2h, i) and then reduced at 6 weeks (113.6 \pm 5.8% at microglia, 110.1 \pm 10.0% at 196 astrocytes, mean \pm SD, n = 5, Fig. 2h, i). Moreover, continuous monitoring of the inflammation 197 around the mesh nanoelectronics revealed nearly normal immune cell distribution at 12 weeks 198 $(95.3 \pm 13.1\%$ at microglia, $105.5 \pm 12.7\%$ at astrocytes, mean \pm SD, n = 5, Fig. 2h, i), and even 199 up to one year (104.9 \pm 9.0% at microglia, 109.5 \pm 17.9% at astrocytes, mean \pm SD, n = 5, Fig. 2h, 200 i). We attribute the little-to-no immune response of the mesh nanoelectronics to the ultra-flexible 201 open structure that is imperceptive to surrounding brain tissue, neurons, and the cells involved in 202 203 inflammation. Compared with the open mesh structure, the thin-film nanoelectronics implantation introduced significantly higher levels of astrocytes ($181.9 \pm 17.9\%$, $272.6 \pm 40.3\%$, $174.4 \pm 22.8\%$ 204

and $182.1 \pm 32.7\%$ at 2-, 6-, 12-and 1-year post-implantation, respectively, mean \pm SD, n = 5, Fig. 205 2h, i) and microglia aggregation (177.2 \pm 14.5%, 162.3 \pm 18.4%, and 125.1 \pm 12.6% and 154.9 \pm 206 207 30.0% at 2-, 6-, 12-and 1-year post-implantation, respectively. Mean \pm SD, n = 5, Fig. 2h, i) over the same period (p < 0.05, significant enhancement compared with the open mesh at all time points, 208 Fig. 2h, i). Notably, we can still observe the proliferation of astrocytes and microglia around the 209 210 thin-film nanoelectronics at one year post-implantation. Together, these results demonstrate that open mesh nanoelectronics introduce little-to-no inflammation and mechanical damage to the 211 surrounding tissues as compared with thin-film nanoelectronics over yearlong implantation. 212

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214 Long-term stable recording at single-cell resolution across multiple brain regions.

215 To test the stability of the recording, we implanted 600-µm-wide mesh nanoelectronics with 32 channels and 300-µm-wide mesh nanoelectronics with 16 channels into multiple mouse brain 216 regions for head-fixed behaving electrophysiology (Methods). Electrodes were implanted into 217 different brain regions including the somatosensory cortex and striatum (32-channel, mesh 218 nanoelectronics#1, Fig. 3a); red nucleus, interstitial nucleus, and ventral tegmental area in 219 220 midbrain (32-channel mesh nanoelectronics#2, Fig. 3a); and visual cortex and hippocampus (16channel mesh nanoelectronics#3, Fig. 3a). Putative individual neurons were isolated using 221 *Waveclus3*³⁸ (Methods). Intrinsic spike waveform variability from the superficial (mesh 222 223 nanoelectronics #1, #3, Fig. 3b, d) and deep (mesh nanoelectronics #2, Fig. 3c) brain regions is consistent with different putative neuron types in each brain region^{39,40}. Moreover, the 224 hippocampus recordings show higher neuron yield per electrode (2.0 ± 0.4 neurons, median ± 1.5 225 interquartile range, n = 8, Fig. 3e), spike amplitude (148.146 ± 77.4 μ V, median ± 1.5 interquartile 226 range, n = 8, Fig. 3f) and firing rate $(17.3 \pm 7.6 \text{ spike/s}, \text{median} \pm 1.5 \text{ interquartile range}, n = 8,$ 227

Fig. 3g) as compared to other recorded brain regions (primary somatosensory cortex, striatum,
midbrain, primary visual cortex, Fig. 3e-g).

230 Next, we evaluated the long-term stability of recordings from 32-channel and 16-channel mesh nanoelectronics from 5 independent animals (n = 43 electrodes from two 32-channel and n = 29231 electrodes from three 16-channel mesh nanoelectronics). 72 putative individual neurons from 232 233 multiple regions were recorded 10 days post-implantation, which increased to 115 putative individual neurons after 60 days. Both 16-channel and 32-channel mesh nanoelectronics show low 234 noise level and high signal to noise ratio (SNR) at 60 days post-implantation in behaving animals 235 (16-channel: 9.97 \pm 1.72 μ V at noise level, 12.67 \pm 6.34 at SNR, *n* = 29 electrodes; 32-channel: 236 $9.27 \pm 2.10 \text{ }\mu\text{V}$ at noise level, 13.25 ± 5.94 at SNR, n = 43 electrodes, mean \pm SD). The statistical 237 results (Fig. 3h-j) showed that the normalized neuron count per electrode, average amplitude, and 238 SNR of 300-µm-wide,16-channel mesh nanoelectronics increased over the first 30 days of 239 implantation and then stabilized (n = 29 electrodes from three 16-channel mesh nanoelectronics). 240 241 These parameters from the 600-um-wide, 32-channel mesh nanoelectronics stabilized at 50-day post-implantation, suggesting the potential device size-related effect on the signal stability (n = 43242 electrodes from two 32-channel mesh nanoelectronics). These results contrast with reports from 243 previous brain probes where amplitudes, SNR, and neuron counts degrade weeks after 244 implantation^{3,4,37}, suggesting that the unfolded, open mesh nanoelectronics formed a long-term 245 stable interface with neurons and tissue. 246

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248 Tracking the same neuron's activity over the entire adult life of mice

Two mice implanted with 16-channel mesh nanoelectronics and one mouse with 32-channel mesh 249 nanoelectronics were recorded monthly until their natural death (5-18 months for mouse #1, 5-20 250 251 months for mouse #2, and 5-19 months for mouse #3). 82.8 ± 6.2 % neurons are stably recorded (mean \pm SD, compared to the first recording session). Compared to mice of 5 months with glossy 252 brown fur, the aged mice of 18-months with mesh electronics implanted exhibited normal and 253 254 healthy aging, including weight gain, barbering around the eyes, and thinning and grey fur in the dorsal back skin⁴¹ (Extended Data Fig. 5a-e). Statistical analysis revealed that there was a 255 significant increase of gray hairs and a decrease of black hairs in aged mice when compared to the 256 mature adult mice (p < 0.05, n = 3, Extended Data Fig. 5f). The electrode interfacial impedances 257 exhibited relatively constant values of 920.2 \pm 107.2 k Ω vs. 857.2 \pm 85.7 k Ω at months 6 vs.18 258 (mean \pm SD, n = 30, Extended Data Fig. 6a), indicating stable electrical and mechanical properties 259 of the mesh nanoelectronics⁴². 260

We first assessed the stability of the signals by spike sorting and statistical analyses. Spike 261 waveforms were projected to a 2D embedding space for stability validation by using UMAP-²⁶ 262 and PCA-based dimension reduction algorithms. PCA is commonly used to define the number and 263 stability of recorded single-neuron signals over time^{19,20,31}. UMAP is a non-linear ML-based 264 dimension reduction algorithm that can learn a low-dimensional embedding space to preserve as 265 much of the local and more of the global data structure than linear dimension reduction algorithms 266 such as PCA. 26 neurons were isolated across all recording sessions starting from 5 months and 267 lasting until natural death. The clusters for each sorted spike in both UMAP and PCA embeddings 268 show nearly constant positions and well separated from each other in the first and second 269 component plane (UMAP1-UMAP2 and PC1-PC2) through >1-year recordings (Fig. 4a, Extended 270 Data Fig. 6b, Methods). In addition, the corresponding single-unit waveforms' shapes (Fig. 4b), 271

as well as their firing dynamics (i.e., interspike interval) were stable (Extended Data Fig. 6c, 272 Methods). Auto-correlation analysis showed that single-unit waveforms were highly similar and 273 274 almost indistinguishable to themselves $(0.90 \pm 0.14 \text{ across all recording sessions from 3 mice},$ mean \pm SD, Fig 4c, Methods). L-ratio⁴³ and silhouette score⁴⁴ analysis (Fig. 4d, Methods) 275 confirmed good unit separation and accurate identification of individual neurons, demonstrating 276 that the signals were sufficiently separated to permit isolation of single units. Statistical analysis 277 on single-unit recording stability examined by five waveform features (amplitude, duration, peak-278 trough ratio, repolarization slope, and recovery slope)⁴⁰ and signal-to-noise ratio (SNR) showed 279 that their average values were nearly constant and the majority (79% from 3 mice) of neuron 280 waveform features did not change significantly over time (p > 0.05 two-sided t test, Fig. 4e-h, 281 Methods), demonstrating that the neuron spikes showed consistent characteristic features over time. 282 Notably, the consistent signal-to-noise ratio (SNR) demonstrated that the electrode-to-cell 283 interface was not degraded during the entire period (Fig. 4f). Collectively, all these results indicate 284 285 that the waveforms were stably recorded from the same neuron over the entire recording period.

In addition to recording of spontaneous activity, we validated the stability of recording by 286 examining the stable recording of whisker stimulation-elicited neuron activities^{32,45}. Specifically, 287 we identified one electrode (electrode A) on the mesh nanoelectronics as being close to a D2 barrel 288 neuron in primary somatosensory cortex (S1 cortex) by successful recording of the whisker-289 stimulation-elicited single-unit spikes (Fig. 4i-l). Recording from stimulation of other whiskers 290 (e.g., C3) or another electrode (e.g., electrode B that is close to electrode A) away from the D2 291 barrel field were used as control (Extended Data Fig. 7). We performed the whisker deflection 292 with a galvanometer-driven stimulation contralateral to the implant (Fig. 4i). A 1 Hz, 900 deg/s 293 deflection was applied to the targeted whisker (Methods). The raster plot and peri-stimulus time 294

histogram (PSTH) of this single-unit recording showed that the electrode A of the mesh 295 nanoelectronics can record strong and rapid neuron firing in response to the principal whisker D2 296 297 deflection (Fig. 4j). We can record well defined neuron activity and waveforms from 8 months to 16 months (Fig. 4j). Spikes observed from the electrode A with C3 whisker deflection (Extended 298 Data Fig. 7a-c) or from electrode B with D2 whisker stimulation (Extended Data Fig. 7d-e) showed 299 300 no correlation with the whisker stimulation. Notably, Figure 4k and I showed that the evoked spike count and the time delay to the stimulation exhibited no significant change over time (p > 0.05, n)301 = 5). These behavior-triggered electrical activities further demonstrated the capability of this 302 method to track the same neuron during the adult lifetime of mice. 303

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305 ML-based validation and analysis

We further applied an unbiased, autoencoder-based²⁷ automated neuronal signal processing and 306 analysis to benchmark the stability of the signal. Autoencoders is a self-supervised learning 307 algorithm, which is only able to meaningfully reconstruct data similar to what they have been 308 trained on, thus providing an unbiased way to examine the stability of the recording. We trained 309 310 three two-headed autoencoders by using detected spike waveforms, corresponding electrode information and sorted neuron labels from the first 6-month recording data of the three mice. Each 311 autoencoder i) learned nonlinear dimension reduction transformations compressing spike 312 waveforms into a 2D embedding space, ii) classified the spike as a specific neuron in the training 313 data, and iii) reconstructed the embeddings back to the input data space (Fig. 5a). The classification 314 and reconstruction were simultaneously optimized during the training of the autoencoder, enabling 315 the autoencoder the capabilities of spike sorting, postprocessing, and stability verification 316 (Extended Data Fig. 8a, Methods) at the same time. Notably, the autoencoder showed much faster 317

classification speed and higher reconstruction accuracy compared with UMAP and random forest⁴⁶ 318 classifier-based spike sorting (Extended Data Fig. 8b, Methods). While we used the autoencoder 319 320 trained with the first six-month recording to achieve the best performance, we found that a twomonth recording dataset is sufficient to train the model with only about 4% accuracy decrease 321 (Extended Data Fig. 8c). The waveforms of the remaining 8-month recording data can still be 322 classified and reconstructed (Fig. 5b and Extended Data Fig. 8d) with the classification accuracy 323 (Fig. 5c, Extended Data Fig. 8e) reaching $89 \pm 4\%$ (mean \pm SD, n = 3 mice). An anomaly dataset 324 was constructed to simulate the drift of the recording using spikes gathered from a fourth 325 independent mouse to test the drift detection ability of the autoencoder. The results showed that 326 the mean squared error (MSE) between reconstructed and original waveforms (Fig. 5d) was higher 327 328 for spikes detected from simulated drifting neurons, compared to the stable neurons used in the 329 training and testing dataset. This significant difference allowed for drift detection based on 330 reconstruction accuracy. A threshold could be used to distinguish the testing spikes and drifting 331 spikes, which eliminated the majority of spikes from the drifting dataset ($83 \pm 12\%$, mean \pm SD, n = 3 mice) and kept most spikes from the testing dataset ($86 \pm 7\%$, mean \pm SD, n = 3 mice, Extended 332 333 Data Fig. 8f). By visualizing the autoencoders' embedding space in the bottleneck (Fig. 5e), cluster 334 embeddings of the neuron spike waveforms from the same mouse showed higher separability than 335 drifting spikes from a different mouse (Extended Data Fig. 8g-k). Furthermore, the training 336 manifold convex hull was used as a stability verification tool for spike processing to quantify the 337 within-boundary subset of testing dataset spikes (Fig. 5f and Extended Data Fig. 8h-k). Low 338 amounts (8 \pm 11%, mean \pm SD, n = 3 mice) of out-of-manifold testing spike embeddings further demonstrated recording stability as a similarity in waveform shape between the first and last 339 recording months, meaning the later spikes' embeddings lay within the embedding space created 340

by the first recorded spikes (Fig. 5f and Extended Data Fig. 8h-k). Collectively, high classification 341 accuracy and low out-of-manifold percentage confirmed the long-term stable single-unit spike 342 343 detected from the same neurons over time. This result also suggests that the stable recording data can be used to build the ML model to perform automated spike sorting based on the first few 344 months of recording from a given mouse by capturing single-neuron waveform salient 345 characteristics used as the input to the classification head. Additionally, the ML model successfully 346 detected the spikes from a different mouse, highlighting the model's ability to detect the drift of 347 348 recording.

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350 Entire adult life study of brain aging at the single-neuron level

The adult mice life recording offers an opportunity to observe aging-associated electrical behavior 351 changes at single-neuron resolution. We performed both qualitative and quantitative analyses of 352 353 aging-associated changes at the single-neuron level over the mouse adult life. Analysis of clusters 354 using PCA showed the stability of spikes from a group of neurons with largely overlapping clusters (purple and red, respectively) from each electrode, while the other spikes (green and blue) from 355 356 the same electrode showed a slight change over the time course of recording (Fig. 6a). We quantitatively assessed the multivariate spread of cluster centroid positions by comparing average 357 position shifts between consecutive cluster centroid positions (Fig. 6a, Methods) to average cluster 358 distribution spreads. This assessment supported the stability (0.47 σ_{purple} , 0.58 σ_{red}) and variability 359 $(1.96 \sigma_{\text{green}}, 2.63 \sigma_{\text{blue}})$ described previously. We analyzed the time-evolution of average neuron 360 waveforms in a representative 3D feature space while calculating trajectories with a B-spline 361 interpolation of successive positions (Fig. 6b, Methods). Non-correlated features were chosen after 362 performing correlation analysis (duration, peak-trough ratio, and repolarization slope in Extended 363

Data Fig. 9a). 21% of neurons (*i.e.*, green and blue) showed a clear trajectories trend while others 364 (*i.e.*, purple and red) remained the same over the aging of the mice (Fig. 6b, Methods). We 365 366 analyzed the time-evolution of the UMAP embeddings of spike waveforms (Fig. 6c, Extended Data Fig. 9b) in real time and ML-defined pseudotime⁴⁷. To study the continuous and gradual 367 transition of the neuron waveforms instead of the discrete real time label, we constructed a pseudo-368 369 temporal path termed as pseudotime to order spikes in the latent space using *monocle3*⁴⁷, an ML tool originally for exploring the dynamics of gene expression within cell types and trajectories 370 over time (Methods). The pseudotime of stable neurons (red and purple) remained the same value 371 as the real time varied, which further validated the stability in the waveform during the aging-long 372 recording. Similarly, the pseudotime of previously defined slow aging-associated neurons 373 increased as the mouse got old. These qualitative and quantitative results suggest that the long-374 375 term stable recording from these open mesh nanoelectronics can track the aging-related electrical activity evolution from the same or same group of neurons in mice at single-cell level. 376

377

378 Conclusion

379 We demonstrated that the ultra-thin shuttle monolithically integrated mesh nanoelectronics can be implanted across multiple brain regions with an open mesh structure with minimal tissue damage. 380 The open mesh structure is interwoven with the neural network in the brain of the animal, enabling 381 382 immune response-free implantation and long-term stable 3D electrode-to-neuron integration. This structural stability allowed us to track the activity of the same neuron over the entire adult life of 383 mice until natural death as supported by our extensive statistical data analyses showing stable 384 impedance, waveform, firing dynamics, and recording performance, something not achieved by 385 other *state-of-the-art* electrodes. We leveraged the high recording stability of this method, capable 386

of successfully training an autoencoder using the first month's recording, which further validate 387 the stability of recording. Combining the stable recording and autoencoder, we can automate spike 388 389 processing, sorting and stability verification on the remaining months' recording. The unique 390 ability to successfully track individual neurons in a chronically stable manner over such a long timespan provides a continuous view of aging-associated changes in neural activity. Combining 391 392 the evolution of spiking activity at both real time and ML-calculated pseudotime, we observed potential aging-associated waveform changes at the single-neuron level. We believe long-term 393 stable tracking of single neuron activity patterns across a stably recorded population of cells 394 combined with automated data analysis tools will open new opportunities for the next-generation 395 brain-machine interface and bioelectronic medicine. This technology also promises to inform our 396 397 understanding of many long-term processes, including development, learning, recovery from 398 injury, neurodegeneration and age-related cognitive decline. In the future, we envision that further integration of stretchability into our current device design, which can further adapt to the large 399 400 volume change during early brain development, could further allow us to achieve the long-term stable recording over the entire lifespan of animals. 401

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403 Figures and Figure Legends

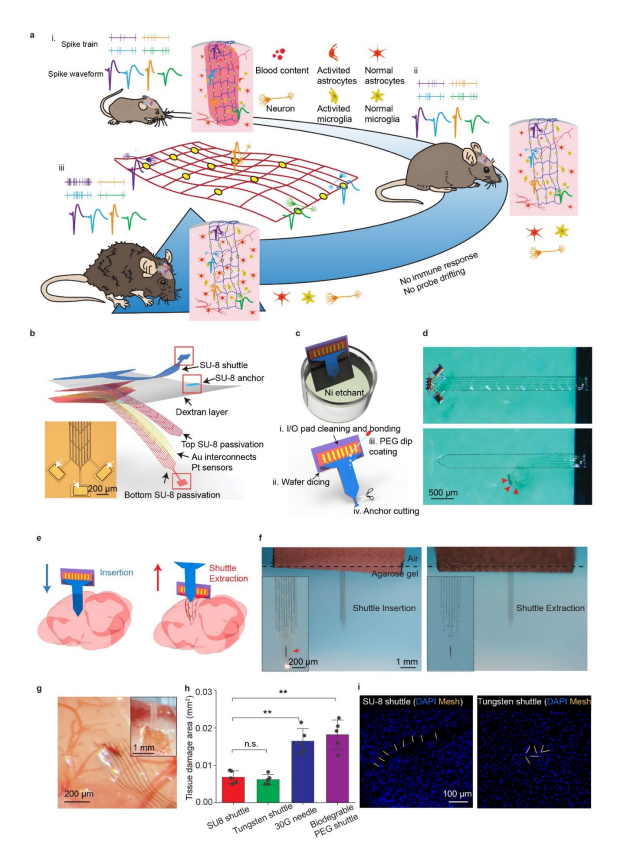


Fig. 1 | Minimally invasive implantation of tissue-level flexible mesh nanoelectronics in the 405 brain. a, Schematics showing the long-term stable electrical recording of the same neuron over 406 the entire adult life of mice enabled by the minimally invasively implanted and fully unfolded 407 tissue-level flexible mesh nanoelectronics. Mesh nanoelectronics seamlessly integrate with neural 408 networks without immune response. Single-cell electrophysiology from the same animal at (i) 409 mature adult (3-6 months), (ii) middle (10-14 months), and (iii) old (18-24 months) stages are 410 recorded. Colored waveforms represent different neurons that are stably recorded over the entire 411 adult life of mice. b, Exploded view of the integrated mesh nanoelectronics showing the distinct 412 material layers. The fully assembled nanoelectronics consisting of (from top to bottom) a 25-um-413 thick polymer shuttle, a 20-µm-thick polymer anchor, a 3-µm-thick dextran dissolvable layer, a 414 415 450-nm-thick top SU-8 encapsulation layer, 50-nm-thick platinum electrodes and 70-nm-thick gold (Au) interconnects, and a 450-nm-thick bottom SU-8 encapsulation layer. Inset: bright-field 416 (BF) microscopic image of mesh nanoelectronics connected with polymer shuttle through anchors 417 (white arrows). c, Schematics showing stepwise releasing of mesh nanoelectronics from the 418 substrate and shuttle. Mesh nanoelectronics was released from the fabrication substrate after 419 removing the Ni sacrificial layer while connected with the polymer shuttle by the anchors (top). 420 421 The released shuttle/nanoelectronics were cleaned for bonding and dicing, and then coated by the biodegradable PEG through dip-coating. After cutting the polymer anchors, the mesh 422 nanoelectronics was released from the shuttle by dissolving PEG (bottom). d, Photographs 423 424 showing the released polymer shuttle/mesh nanoelectronics hybrid from the substrate (top) and released mesh nanoelectronics from the shuttle (bottom). Red dashed lines and arrows highlight 425 the cutting lines of the anchors and the released mesh nanoelectronics, respectively. e, Schematics 426 showing the brain implantation process. f, In vitro images of mesh nanoelectronics implantation 427 in a 0.6% agarose gel. Insets: zoom-in images showing the released mesh nanoelectronics maintain 428 the unfolded structure and implantation location after withdrawing the polymer shuttle. g, 429 430 Photograph showing the representative brain implantation with minimal tissue damage. Inset: ultrathin polymer shuttle-enabled implantation. h, Statistical analysis of tissue acute damage zone 431 of different implantation methods. Data represented as mean \pm SD, individual data points are 432 overlaid (**p < 0.01, two-tailed unpaired t test, n = 5). i, Representative images of 20-µm-thick 433 horizontal brain slices showing the acute mechanical injuries by the ultrathin-polymer shuttle (left) 434 and 50-µm diameter tungsten shuttle (right). Yellow and blue represent mesh nanoelectronics and 435 DAPI, respectively. 436 437

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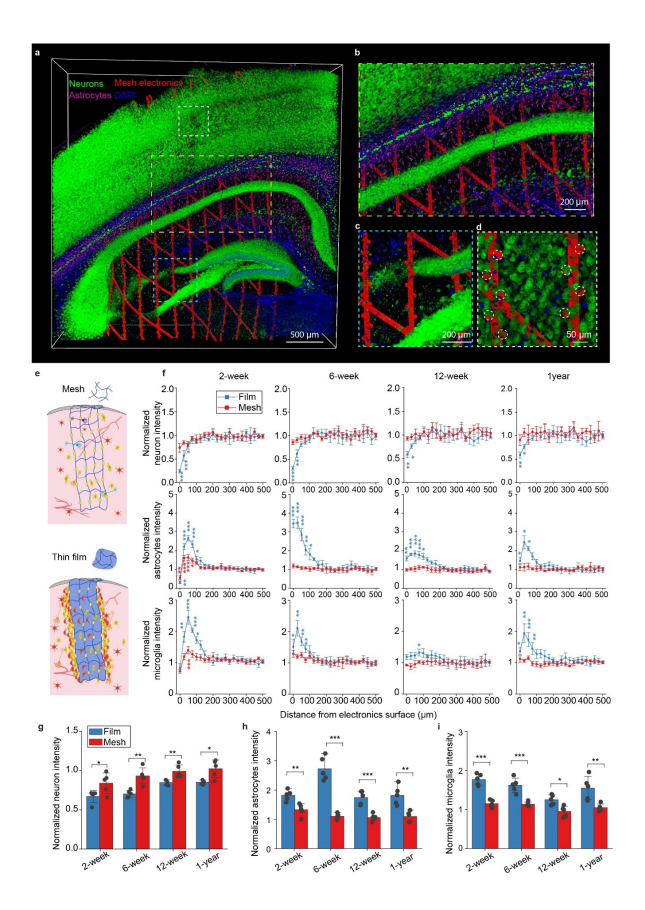


Fig. 2 | Unfolded mesh nanoelectronics seamlessly integrating with the neuron network 440 across multiple brain regions. a, Representative 3D reconstructed confocal fluorescence imaging 441 of 600-µm-thick brain tissue implanted with 2-mm wide mesh nanoelectronics for 6 weeks. Green, 442 purple, blue, and red label neurons, astrocytes, nuclei, and mesh nanoelectronics. b-d, Zoom-in 443 views of the regions highlighted by white (b), cvan (c), and vellow (d) dashed boxes in (a), 444 showing the seamless integration of the mesh with the neural network. Neuron-like electrodes are 445 highlighted by white dashed circles in (d). e. Schematics illustrating 1-um-thick mesh (top) and 446 thin-film (bottom) nanoelectronics implanted and unfolded inside brain tissue for the long-term 447 immune response characterization. f, Normalized fluorescence intensity as a function of distance 448 from the mesh/thin-film electronic and tissue boundary at 2 weeks, 6 weeks, 12 weeks, and 1-year 449 450 post-implantation. The relative signal was obtained by normalizing the fluorescence intensity with the baseline value defined as the fluorescence intensity over a range of 525–550 µm away from 451 the electronics. Data represented as mean \pm SEM (intensity compared with that of distance at 500-452 μ m, *p < 0.05, **p < 0.01, ***p < 0.001, two-tailed unpaired t test, n = 5). g-i, Normalized neuron 453 (g), astrocytes (h), and microglia (i) intensity and neuron density within 100-µm from the 454 electronic surface. Data represented as mean \pm SD, individual data points are overlaid (*p < 0.05, 455 456 **p < 0.01, ***p < 0.001, two-tailed unpaired t test, n = 5). 457

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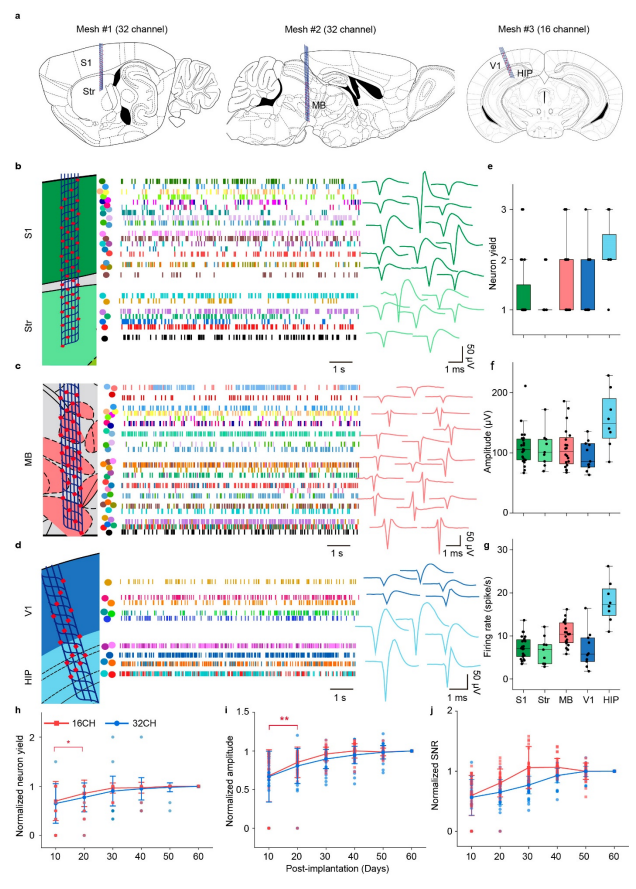


Fig. 3 | Chronically stable recording across multiple brain regions. a, Schematics illustrating 459 the representative brain regions implanted with mesh nanoelectronics for chronic recording. S1, 460 primary somatosensory cortex; Str, striatum; MB, midbrain; V1, primary visual cortex; HIP, 461 hippocampus. b-d, Approximate mesh locations overlaid on the Allen Mouse Brain Atlas (left) 462 and representative spiking raster (middle) and single-unit waveform (right) from two 32-channel 463 (b, c) and one 16-channel (d) mesh nanoelectronics in head-fixed behaving animals at 60 days 464 post-implantation. Colored dots and blocks indicate individual neurons and spike times, 465 respectively (middle). Representative average single-unit action potential waveforms were 466 extracted from a 2-min recording session (right). e-g, Quantification of sorted neuron yield per 467 electrode (e), waveforms amplitude (f), and firing rate (g) across 5 brain regions in a 2 min 468 469 recording session. Box plots show median and quartile range (whiskers denote $1.5 \times$ the interquartile range). Individual data points are overlaid (n = 29 electrodes from three 16-channel 470 mesh nanoelectronics and n = 43 electrodes from two 32-channel mesh). h-j, Normalized sorted 471 neuron yield (h), amplitude (i), and signal-to-noise ratio (SNR) (j) over the time course of 60 days. 472 Data represented mean \pm SD, individual data points are overlaid (n = 29 electrodes from three 16-473 channel mesh nanoelectronics and n = 43 electrodes from two 32-channel mesh nanoelectronics. 474 475 *p < 0.05, **p < 0.01, comparison of different days within 16-,32-channel mesh electronics, twotailed unpaired t test). 476 477

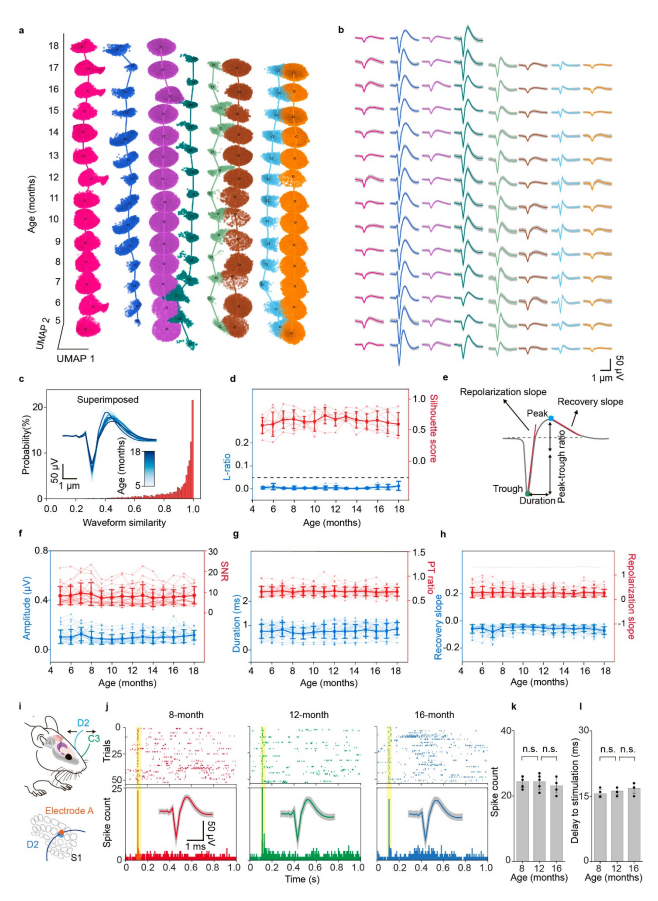
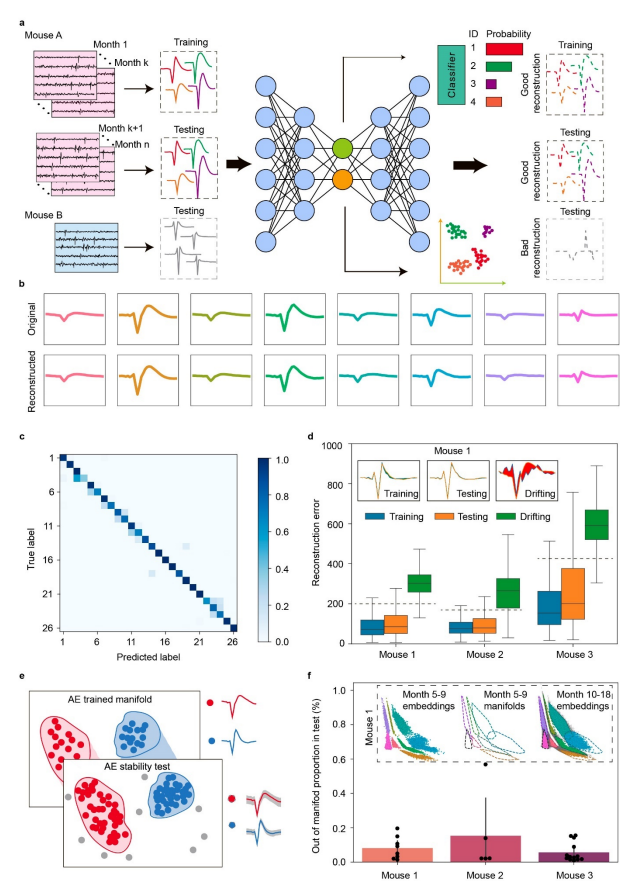


Fig. 4 | Stably tracking the single-unit action potential from the same group of neurons over 479 the entire adult life of mice. a, Time evolution of single-units clustered by Leiden over the entire 480 adult life of mice from the mature adult (5 months) to the aged (18 months) stage. The x- and y-481 axes denote the first and second uniform manifold approximation and projection (UMAP) 482 dimensions, respectively, and the z-axis denotes mouse age in months. Color scheme maintained 483 throughout. **b**, Time course analysis of the average waveforms of single-unit action potential in 484 the *Leiden* clustering results in (a). Waveform represented mean \pm SD. c, Waveform correlations 485 for every single-unit across all recording sessions (n = 26 units from 3 mice). Inset: Overlaid 486 average waveforms across all recording session for the representative putative same neuron. **d**, 487 Likelihood-ratio (L-ratio) and silhouette score showing the clustering quality for the single unit 488 489 action potentials from identified neurons. Data represented mean \pm SD, individual data points are overlaid. The constant dashed line is 0.05 L-ratio, commonly taken as a threshold of high cluster 490 quality. e, Schematic showing features extracted from the single-unit action potential waveform 491 used for the analysis in (f-h). f-h, mean \pm SD (individual data points are overlaid) of features 492 illustrated in (e) and signal-to-noise ratio (SNR) over three mice life recording. In order, amplitude, 493 SNR, duration, peak-trough ratio (PT ratio), recovery slope, and repolarization slope are shown. i-494 495 I, Long-term stable recording of the behavior-associated neuron. i, Schematic diagram of whisker deflection. An individual vibrissa was deflected in the rostral-caudal plane using a computer-496 controlled galvanometer system. D2 and C3 whiskers are labeled in blue and green, respectively 497 498 (top). Schematic diagram of whisker barrel arrangement in S1 (bottom). Evoked spike from electrode A was associated with D2 whisker deflection. Control experiments were shown in 499 Extended Data Fig. 7. i, Raster plot and peri-stimulus time histogram (PSTH, 1 ms bin size) of the 500 single unit identified from electrode A response to D2 whisker deflection from 8- to 16-month 501 recording. Inset: average single-unit waveforms from recordings in the S1 in response to D2 502 whisker deflection over time, Waveform represented mean \pm SD. **k**, Population data showing the 503 number of spikes (out of 55 trials) evoked by whisker deflection over the time course of 504 implantation. I, Time delay of the evoked spikes to the stimulation over the time course of 505 implantation (n.s.: not significant, two-tailed unpaired t test, n = 5). Individual data points are 506 overlaid. 507

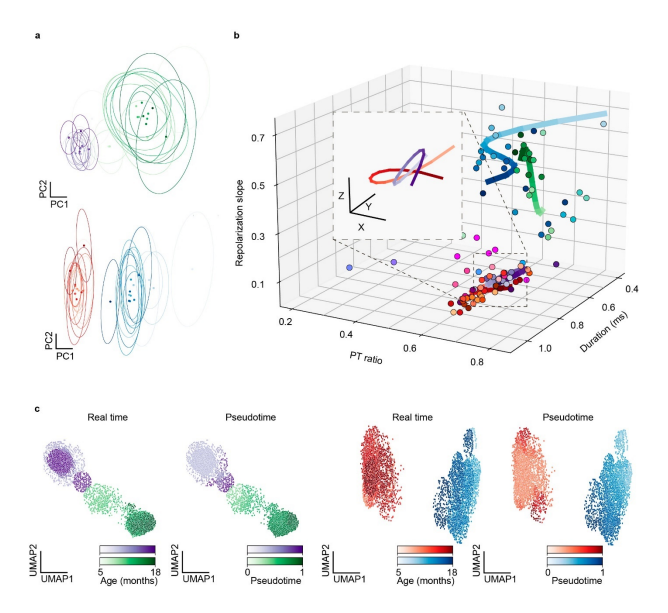
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509 Fig. 5 | Long-term stable single-cell recording enabled autoencoder-based spike processing.

a, Schematics of an autoencoder-based model which consists of an encoder, a classification head, 510 and a decoder. Single-unit action potential waveforms are first extracted from the recording 511 voltage traces and encoded with channel number to a lower-dimensional latent space. The latent 512 representation is then used for classification and decoding. Spikes from the first 6 recording 513 months including 5,490 spikes were used for training. The subsequent 8 months of data including 514 158.317 spikes were used for testing. Spikes from different groups of neurons collected from a 515 different mouse were also used as simulated drifting data for testing. b, Original and reconstructed 516 average representative single-unit waveforms from the testing data. Original waveforms are 517 colored by neuron labels. Reconstructed waveforms are colored by the classification output. c, 518 519 Confusion matrix comparing testing dataset ground-truth neuron labels with autoencoder classification output across all three mice. d, Boxplot of median and quartile range of mean-520 squared error (MSE) distribution for each of the training, testing, and drift datasets. Whiskers 521 denote 1.5× interquartile range. Dashed lines indicate thresholds corresponding to mean + SD of 522 mouse-specific MSE training distribution, eliminating 83 ± 4 % of training and 86 ± 7 % of testing 523 (Extended Data Fig. 8f). Inset: representative average waveforms and their corresponding 524 525 reconstructions by the mouse-specifically trained autoencoder. The colored areas correspond to the difference between the reconstructed and original waveform. e, Schematics showing the 526 autoencoder can be used to quantify the stability of the recording. The trained manifold 527 528 (highlighted by red and blue) was used to test the stability of the recording by quantifying the percentage of the neuron single-unit action potentials from the testing data falling inside the 529 training manifold. **f**, Bar plot illustrating the percentage of the testing dataset for each neuron 530 falling outside of the training manifold as per the process illustrated in (e). Data represented mean 531 \pm SD, individual data points are overlaid. Low levels of detected noise suggest recording stability 532 through train-test waveform resemblance. Inset: applying noise rejection process to mouse 1 data, 533 534 in order: training embeddings, training manifold boundaries, testing embeddings with out-ofmanifold spikes colored in grey. 535

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536

Fig. 6 | Adult life study of brain aging at the single-neuron level. a, Principal component 537 analysis (PCA) of representative waveforms of neurons showing stable (purple and red) and 538 progressive changing (green and blue) electrophysiological behaviors. Dots and ovals represent 539 the centers and $\pm 2 \sigma$ of PC clusters, respectively. Average cluster centroid successive position 540 shifts: 0.47 opurple, 0.58 ored, 1.96 ogreen, 2.63 oblue. Low values indicate stable cluster centroid 541 position across time relative to single cluster whereas high values suggest time-evolution of 542 corresponding cluster centroids and associated single-unit action potential waveforms. In both 543 cases, a single electrode channel tracked stable (< 1σ) and time-evolving single-unit action 544 potential waveform. **b**, Trajectory analysis conducted with a B-spline interpolation of 545 representative waveform features over the time course of implantation for the same neurons. Dots 546 with the same color represented mean values of features calculated by the waveforms associated 547 with the same color-coded neurons from data shown in (a). Trajectories highlight the 548 representative neurons shown in (a) showing slow progressive and aging-associated 549 electrophysiological properties. Inset: zoom-in view of purple and red trajectory. Scale bar, x = 550

- 551 0.01, y = 0.025, z = 0.01. c, Comparison of the electrophysiology Monocle3-based pseudotime and
- real time evolution over the adult life of mice of overall UMAP representation of representative individual single-unit action potential waveforms. Pairs of neurons recorded by the same electrode
- 554 are compared.

555 Methods

556 1. Fabrication of ultra-thin shuttle monolithically integrated mesh nanoelectronics

(1) Cleaning a silicon wafer grown with thermal oxide (500-nm thickness) with acetone, isopropyl 557 alcohol, and deionized water. (2) Depositing 100-nm-thick nickel (Ni) using a Sharon thermal 558 559 evaporator as a sacrificial layer. (3) Spin-coating SU-8 precursor (SU-8 2000.5, MicroChem) at 3000 rpm, which was pre-baked at (65 °C, 95 °C) for 2 min each, exposed to 365 nm ultra-violet 560 (UV) for 200 mJ/cm², post-baked at (65 °C, 95 °C) for 2 min each, developed using SU-8 561 562 developer (MicroChem) for 60 s, and hard-baked at 180 °C for 40 min to define mesh SU-8 patterns for bottom encapsulation. (4) Spin-coating LOR3A photoresist (MicroChem) at 4000 rpm, 563 followed by pre-baking at 180 °C for 5 min; spin-coating S1805 photoresist (MicroChem) at 4000 564 rpm, followed by pre-backing at 115 °C for 1 min; the sample was then exposed to 405 nm UV 565 for 40 mJ/cm², and developed using CD-26 developer (Microposit) for 70 s to define interconnects 566 patterns. (5) Depositing 5/70/5-nm-thick chromium/gold/chromium (Cr/Au/Cr) by a Denton 567 electron-beam evaporator, followed by a standard lift-off procedure in remover PG (MicroChem) 568 to define the Au interconnects. (6) Repeating Step (4) to define electrode array patterns in 569 570 LOR3A/S1805 bilayer photoresists. (7) Depositing 5/50-nm-thick chromium/platinum (Cr/Pt) by a Denton electron-beam evaporator, followed by a standard lift-off procedure in remover PG 571 572 (MicroChem) to define the electrode array. (8) Repeating Step (3) for top SU-8 encapsulation. (9) 573 Spin-coating SU-8 precursor (SU-8 2025, MicroChem) at 4000 rpm, which was pre-baked at 65 °C for 2 min and 95 °C for 8 min, exposed to 365 nm ultra-violet (UV) for 200 mJ/cm², post-baked 574 at 65 °C for 2 min and 95 °C for 6 min, developed using SU-8 developer (MicroChem) for 6 min, 575 576 and hard-baked at 180 °C for 1 hour to define SU-8 anchors patterns to connect the mesh and SU-577 8 shuttle. (10) Spin-coating 20 wt% dextran solution at 1000 rpm for 20 s. which was at 80 °C for

1 min and 180 °C for 30 min. (11) Spin-coating SU-8 precursor (SU-8 2025, MicroChem) at 3000 578 rpm, which was pre-baked at 65 °C for 2 min and 95 °C for 8 min, exposed to 365 nm ultra-violet 579 580 (UV) for 200 mJ/cm², post-baked at 65 °C for 2 min and 95 °C for 6 min, developed using SU-8 developer (MicroChem) for 6 min, and hard-baked at 180 °C for 1 hour to define the SU-8 shuttle 581 pattern. (12) Cleaning the input/output with water and soldering a 32-channel flexible flat cable 582 (Molex) onto the input/output pads using a flip-chip bonder (Finetech Fineplacer). (13) Soaking 583 the mesh nanoelectronics in nickel etchant for 2 to 4 hours to completely release the mesh 584 nanoelectronics from the substrate wafer. (14) Rinsing the mesh nanoelectronics with deionized 585 water and PBS three times each. (15) Dicing the substrate to the desired length. Dip-coating 10 586 wt% PEG solution to attach the mesh nanoelectronics and SU-8 polymer shuttle. The 587 monolithically integrated mesh nanoelectronics was allowed to dry in the air. (16) After cutting 588 the anchor, the monolithically integrated mesh nanoelectronics was ready for implantation. 589

590

591 **2. Bending stiffness calculations**

592 We estimated and compared the bending stiffness values (D) of mesh and thin-film nanoelectronics 593 using a beam model. The bending stiffness of mesh nanoelectronics with three-layer 594 polymer/mesh/polymer structure can be calculated as

595
$$D = \frac{E_S}{12} (wh^3 - w_m h_m^3) + \frac{E_m}{12} w_m h_m^3$$
(1)

where E_s and E_m are the young's moduli of SU8 and gold, 2 and 79 GPa, respectively, *w* is the width of SU-8 scaffolds, *w_m* is the width of gold interconnects, $h_{mesh} = 0.91 \mu m$ and $h_{film} = 0.92 \mu m$ is the measured total thickness of SU-8, $h_{metal} = 80 \text{ nm}$ is the thickness of Cr/Au interconnects. The calculated bending stiffness of mesh and thin-film nanoelectronics, $1.26 \times 10^{-15} \,\mathrm{N \cdot m^2}$ and $39.8 \times 10^{-15} \,\mathrm{N \cdot m^2}$, respectively.

601

602 **3. Brain implantation**

603 All the animal experiments were approved by the Institutional Animal Care and Use Committee 604 of Harvard University. The implantations were carried out on the male C57BJ/6 mice (~16 weeks of age). The animals were housed in a regular 12 h/12 h light/dark cycle. Animals were 605 anesthetized with 2-3% isoflurane and maintained under anesthesia with 0.75-1% isoflurane 606 during the intracranial implantation surgery. Stainless-steel screws were implanted in the 607 cerebellum and used as ground electrodes. A craniotomy $(2 \times 2 \text{ mm}^2)$ was performed on the brain, 608 and the cortical surface was exposed upon removal of the dura mater. The mesh nanoelectronics 609 with the releasable shuttle was attached onto a micromanipulator on the stereotaxic frame. The 610 micromanipulator was manually controlled to insert the nanoelectronics into the mouse brain at 611 the targeted depth at the tip. Sterile PBS was applied on the rear end of the nanoelectronics to 612 dissolve the PEG/dextran and release the SU-8 shuttle from mesh nanoelectronics. After 613 614 PEG/dextran was fully dissolved on both ends, the SU-8 shuttle was extracted with the second manipulator, leaving ultra-flexible mesh nanoelectronics implanted at the target positions. 615 Craniotomies were sealed with a silicone elastomer (World Precision Instruments, USA). Ceramic 616 617 bone anchor screws, together with dental methacrylate, were used to fix the FFC and electrode set onto the mice's skull. 618

619

620 4. Immunohistochemistry

621 The following procedures were performed according to our previous reports 18,48

Histology sample preparation: At each time point (2-, 6-, 12-week and 1-year post-implantation), 622 623 mice were anesthetized with 40-50 mg/kg sodium pentobarbital and then transcardially perfused with ~40 ml PBS pre-wash, and ~40 ml 4 % paraformaldehyde (PFA) in PBS, followed by 624 decapitation. The scalp skin was removed, and the exposed skull/dental cement were ground for 625 626 10-20 min at 15,000 r.p.m. using a high-speed micro motor tool. The brain with the mesh/thin film nanoelectronics undisturbed was removed from the cranium and postfixed in PFA for 24 h at 4 °C. 627 The brain was transferred to incrementally increasing sucrose solutions (10-30%, w/v) until sunk 628 to the bottom for the thin tissue preparation. 629

Immunohistochemical staining of 20-µm thin tissue: After cryostat sectioning, brain slices were 630 incubated PBST (1 × PBS with 0.2% Triton X-100, Thermo Fisher Scientific) for 30 min, and then 631 blocked with 5 % (w/v) normal donkey serum for 2 hours. After three rinses with PBST for 30 632 min each, slices were then incubated at 4 °C overnight in the primary antibodies: chicken anti-glial 633 fibrillary protein GFAP (targeting astrocytes, 1:200, Abcam #ab4674, USA), goat anti-ionized 634 calcium binding adaptor molecule 1 (Iba1) (targeting microglia, 1:100, Abcam #ab5076, USA), 635 636 and rabbit anti-neuronal nuclear NeuN (targeting nuclei of neurons, 1:200, Abcam #ab177487, USA), followed by the slices being washed three times for 30 min each with PBST. Slices were 637 then incubated in a secondary antibody solution at room temperature for 2 h with protection from 638 639 light (1:500, Alexa Fluor 647 donkey anti-chicken, Jackson Immunoresearch, USA; 1:500, Alexa 640 Fluor 594 donkey anti-goat; 1:500, Alexa Fluor 488 donkey anti-rabbit, Invitrogen, USA). After being washed three times for 30 min each with PBST, brain slices were also stained by incubating 641 with 4',6-diamidino-2-phenylindole (DAPI, Sigma-Aldrich, USA) to mark all cell nuclei for 30 642 643 min. After being washed, slices were mounted on glass slides with coverslips using Prolong Gold 644 (Invitrogen, USA) mounting media. The slides remained covered (protected from light) at room
645 temperature, allowing for 12 h of clearance before imaging.

646 Tissue clearing and staining for thick tissues: After vibratome sectioning, brain slices were placed in 1 × PBS containing 4% (w/v) acrylamide (Sigma-Aldrich) and 0.25% (w/v) VA-044 647 thermal polymerization initiator (Fisher Scientific) at 4 °C for 3 days. The solution was replaced 648 649 with fresh solution immediately before placing the brain slices in X-CLARITY polymerization system (Logos Biosystems) for 3 h at 37 °C. After polymerization, any remaining gel from the 650 tissue surface was removed and the slices were rinsed with PBST before placing them in 651 electrophoretic tissue clearing solution (Logos Biosystems) at 37 °C for 3–5 days until the samples 652 were translucent. Brain slices were incubated with PBST overnight, followed by three washes with 653 PBST, and then blocked with 5 % (w/v) normal donkey serum for 2 days. After three rinses with 654 PBST, slices were then incubated at 4 °C for 5-7 days in the primary antibodies containing: chicken 655 anti-glial fibrillary acidic protein (GFAP) (targeting astrocytes, 1:200, Abcam #ab4674, USA) 656 657 and/or and NeuN (targeting nuclei of neurons, 1:200, Abcam #ab177487, USA), followed by the slices being washed three times with PBST. Slices were then incubated in a secondary antibody 658 solution at 4 °C for 5-7 days with protection from light (1:500, Alexa Fluor 647 donkey anti-659 chicken, Jackson Immunoresearch, USA; 1:500 and/or Alexa Fluor 488 donkey anti-rabbit, 660 Invitrogen, USA). After being washed three times with PBST, brain slices were also stained by 661 incubating with 4',6-diamidino-2-phenylindole (DAPI, Sigma-Aldrich, USA) to mark all cell 662 nuclei for 2 days. Brain slices were glued at their edge to the bottom of Petri dishes with 1% (w/v) 663 agarose in optical clearing solution (Lifecanvas Technologies) 24 h before microscopy imaging. 664

665

666 5. Microscope imaging and image data analysis

Confocal fluorescent images were acquired using a Leica SP8 confocal system (Leica, USA). 667 Images were collected using a 25×, 0.95 NA water-immersion or 40×, 1.3 NA oil-immersion 668 669 objective lens. 488 nm, 591 nm, and 633 nm lasers as the excitation sources for Alexa Fluor 488, Alexa Fluor 594, and Alexa Fluor 647, respectively. Standard TIFF files were exported and 670 colorized using LAS X Software. ImageJ software and custom code were used for image analysis. 671 672 The distance of each pixel to mesh/film nanoelectronics was defined as its shortest distance from the mesh/film boundary. Baseline fluorescence intensity is defined as the average fluorescence 673 intensity of all pixels 525–550 µm away from the boundary. Intensity values with distances binned 674 over an interval of 25 µm were averaged and normalized against the baseline intensity. 675

676

677 6. Electrophysical measurement

Mice with implanted mesh nanoelectronics and FFC connector were recorded chronically monthly 678 679 using CerePlex Direct recording system (Blackrock Microsystem, USA), starting from 1-month post-implantation. Mice were anesthetized with 1 % isoflurane in medical-grade O₂ or head-fixed 680 during the measurement. Homemade PCB for connecting the FFC and head stage. Two animals 681 682 are excluded from the long-term study (Fig. 4) since the poor body condition and connectors failed at an early stage. The electrophysiological recording was made with a 30-kHz sampling rate and a 683 60-Hz notch filter. Whiskers of interest were trimmed at \sim 15 mm from the face and inserted into 684 a polyimide tube fixed to the galvanometer system (PT-30K, SpaceLas, China) positioned ~10 mm 685 from the vibrissal pad to yield high-fidelity sensory stimuli. Stimulation was always delivered 686 along the rostro-caudal axis. Voltage command and output for the actuator were programmed by 687 Axon Digidata 1550B (Molecular Devices, USA). 688

689 7. Data analysis

Spike sorting and clustering: All recording data was analyzed offline. 6 min continuous 690 691 recordings were used for analysis of each month. Spike detection was performed using the WaveClus3 software package (https://github.com/csn-le/wave_clus). In brief, raw recording data 692 was filtered using four poles Butterworth filters in the 300-3000 Hz frequency range before spikes 693 694 were detected using an amplitude threshold 5 times the estimated standard deviation of the noise. After spike alignment, 30 data points with the sampling rate of 10 kHz were kept for each detected 695 spike representing 3 ms. No normalization procedure was applied to the spike waveforms due to 696 the stable nature of recorded amplitudes (Fig. 4f) and experiments (Extended Data Fig. 10a-f) 697 showing normalization decreased putative neuron cluster separability as well as meaningfulness 698 of PCA embeddings. Spike sorting results of WaveClus3 obtained through superparamagnetic 699 clustering were kept and used for comparison with the results of our chosen spike sorting approach. 700 Additional quality metrics calculated for individual neurons demonstrate that the individual 701 702 neurons were clearly defined (Extended Data Fig. 10g-i). Results reported in Figure 4 and 6 used clusters determined by *Leiden*⁴⁹(https://github.com/vtraag/leidenalg) clustering performed on the 703 graph constructed by Uniform Manifold Approximation and Projection (UMAP, 704 705 https://github.com/lmcinnes/umap) for each individual month and over individual channels. Geometric considerations ensured that no neuron was being recorded by two separate channels. 706 Cluster labels were then aligned by choosing a label matching scheme which minimized the mean-707 squared error between the average cluster waveforms of a given month and a chosen template. 708

709 Stability analysis of single-unit action potentials: After spike detection, sorting and clustering,710 stability analysis was performed. A custom computational pipeline for assessing the stability711 analysis, which was built using Python v3.9.4. Performing prior spike extraction using third-party

software or using the pipeline to perform spike extraction and sorting directly in python are bothpossible.

714 UMAP analysis was used to confirm the recording stability. UMAP embeddings' coordinates were calculated for all spikes of a given channel before plotting the corresponding points per recording 715 month producing a visualization over time. The representations originating from different channels 716 were plotted next to each other after manual curation for quality control to ensure only the best 717 quality spikes were included in the analysis. Additionally (Fig. 6a), average cluster centroid 718 position shifts were compared to average cluster distributions spread. Concretely, let us name $X \in$ 719 \Re^{dx^2} a vector of cluster centroid positions over d days in a given 2D (x, y) embedding. For each 720 identified neuron, we calculated: 721

722
$$\sqrt{\left(\frac{u_x}{\sigma^{av}x}\right)^2 + \left(\frac{u_y}{\sigma^{av}y}\right)^2} \tag{2}$$

with $u_x = \frac{1}{d-1} \sum_{i=i}^{d-1} |x_{i+1} - x_i|$, $u_y = \frac{1}{d-1} \sum_{i=i}^{d-1} |y_{i+1} - y_i|$ the average successive cluster centroid absolute position shifts along each axis, and $\sigma^{av}{}_x = \frac{1}{d} \sum_{i=1}^{d} \sigma^{x}{}_i$, $\sigma^{av}{}_y = \frac{1}{d} \sum_{i=1}^{d} \sigma^{y}{}_i$ the average cluster distribution SD along each axis. This allowed us to compare centroid position shifts with average cluster SD along each axis to examine the embedded waveform stability. Results of correlation analysis using Pearson correlation coefficients and associated two-sided *t* tests were carried out using *scipy* v1.6.3 (http://www.scipy.org) stats module.

730 Cluster quality metrics shown in Figure 4d were obtained by calculating silhouette score using

rikit-learn v0.24.2's (<u>http://scikit-learn.org</u>) implementation over individual channels and months.

732 L-ratio for an individual cluster was calculated as

$$L_{ratio}(C) = \frac{L(C)}{n_c},$$
(3)

734

where,
$$L(C) = \sum_{i \notin C} 1 - CDF_{\chi^2_{df}}(D_{i,C}^2)$$
 (4)

with n_c the number of spikes belonging to cluster C, $CDF_{\chi^2_{df}}$ the cumulative distribution function 735 of the χ^2 distribution, df = 2 in our case and $D_{i,C}^2$ being the mahalanobis distance of spike i from 736 the center of cluster C. Linear regressions were performed using scikit-learn. 737 Within unit correlation analysis in Figure. 4c was calculated based on the Pearson product-moment 738 correlation coefficient. The corrcoef function in numpy v1.18.5 (http://www.numpy.org) was used 739 740 between all pairs of average cluster waveforms and recording days to provide the auto and cross correlation coefficients. Feature extraction was performed by using functions adapted from 741 AllenSDK (https://github.com/AllenInstitute/AllenSDK). 742

Analysis of interspike intervals: The spiking times of each sorted neuron were used to calculate
the interspike interval (ISI) histograms for individual months per cluster with a bin size of 2 ms.

745 **Data analysis of whisker-stimulus provoked recording:** For whisker responsive recording, 746 firing timing of each detected single-unit spike was presented and aligned by trial number in the 747 raster plot. For the peristimulus time histogram (PSTH), 1 ms bin size was used and the spike 748 counts were accumulated from all trials within each recording session.

Autoencoder-based automatic spike processing: All autoencoders were built using *tensorflow* 2.5 (https://www.tensorflow.org). The encoder consisted of two fully connected (FC) layers with 100 nodes each before the bottleneck 2D layer. Inputs to the network were concatenations of individual spike waveforms and their corresponding one-hot encoded channel (geometric considerations of the sparsity of the mesh recording device ensure that no single neuron can be

recorded on two separate channels). From the bottleneck, the classification head, which consists 754 of a single FC layer with *softmax* activation, outputs probabilities of belonging to a particular 755 756 neuron class. The decoder's symmetric structure with respect to the encoder allows for reconstruction of the original waveform. Activations for encoder and decoder layer were set as 757 LeakyReLU with alpha = 0.3 as well as an added L_2 regularizer on the encoder's last layer, 758 enforcing lower latent embedding values. Adam optimizer with default parameters was used. For 759 a given dataset, $X \in R^{nx30}$, $n \in \Re$ of n spikes, each composed of 30 data points recorded on c 760 channels, we denote the input batch to the autoencoder $\hat{X} \in R^{px(30+c)}$. The autoencoder 761 considered here is composed of an encoder, f_e , decoder f_d and classifying head f_c parametrized by 762 $\theta = (\theta_e, \theta_d, \theta_c)$. The network's (slightly simplified, as the encoder's last layer regularization is 763 not explicit here) loss function over a batch of p spikes is thus: 764

765
$$L(\widehat{X},\theta) = \sum_{i=1}^{p} \left[\alpha | |x_i - f_d(f_e(x_i))| |_2^2 + \beta CE(f_c(f_e(x_i))) \right]$$
(5)

where CE is the usual categorical cross-entropy loss used for multi-class classification problems. 766 767 The loss function is clearly defined by two components: reconstruction and classification, each 768 with their own weight to regulate their overall contribution to the loss function (reconstruction 769 error is also on a different scale than the cross-entropy loss so the relative weighting terms α and 770 β are necessary for balancing contributions of the losses). Training the network consists of learning the correct parametrization θ (network weights) which minimizes the loss function over the 771 772 training dataset. Early Stopping was applied on an evaluation dataset to avoid overfitting of the training data. One autoencoder was trained per mouse using a subset of that mouse's first months 773 recording data. The subset was chosen using conservative thresholds on the within-cluster centroid 774 distance of UMAP embedded spike waveforms. This allows to train the autoencoder in a very short 775 time on high-quality spikes without hurting classification results (Extended Data Fig. 8b). It also 776

means the autoencoder's training manifold can be used as a stability verification tool (Fig. 5f and
Extended Data Fig. 8h-k). Indeed, waveforms recorded in the later parts of the recordings with
similar shapes as the high-quality training waveforms will fall inside the training manifold.

780 Aging analysis at the single-neuron level: UMAP coordinates of the representation in Fig. 4 were used to visualize the different neuron trajectories shown in Figure 6 defined by their 781 782 electrophysiological properties across the adult mouse life recording. Indeed, these UMAP embeddings were used to compute a trajectory graph with the corresponding "pseudo temporal" 783 scale of evolution, with *Monocle 3* (https://cole-trapnell-lab.github.io/monocle3). The convex hull 784 delimiting regions spanned by individual neuron time-evolution was calculated using Scipy; 785 Principal Component Analysis (PCA) was used as another form of dimensionality reduction and 786 787 was calculated per channel and month to view the evolution of spike waveforms per cluster (Extended Data Fig. 6b). Clusters obtained from the same channel were used to obtain cluster 788 centroids per channel for individual recording months along with 2 SD confidence ellipses of 789 790 corresponding covariance. Confidence ellipses were centered on cluster means while radii were calculated without explicit computation of eigenvalues by using Pearson correlation coefficients 791 and equations (6) and (7): 792

793

794

$$\sqrt{\lambda_1} = \sqrt{1 + p} \tag{6}$$

$$\sqrt{\lambda_2} = \sqrt{1-p} \tag{7}$$

where λ_1 and λ_2 are the eigenvalues of the covariance matrix and p is the Pearson correlation coefficient. These equations hold for normalized data; thus, ellipses were scaled by twice the standard deviation along each axis to render the final plots. 799 Feature trajectory analysis was performed by fitting trajectories, from feature embeddings in a selected 3D feature space of mean feature values per cluster over real recording months. The 800 801 features chosen for the representation were repolarization slope, peak trough ratio and duration. These were selected based on prior correlation analysis (Extended Data Fig. 9a) to make sure the 802 trajectory was not a degenerate representation of feature evolution because of highly correlated 803 804 features. Trajectory fittings were constructed using a B-spline interpolation on previously discussed points by first finding the parametric definition of the curve using Scipy splrep function 805 before evaluating the spline using *splev* function. A cubic (k = 3) spline was used with s=2. as the 806 807 smoothing condition in *splrep*.

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Author contributions J.L. and S.Z. conceived and designed the experiments. S.Z., R. L, J. Lee.
fabricated and characterized the electrodes. S.Z., Z.L. performed the brain implantation, *in vivo*

recording and histology study. S.Z., X.T., S.P. conducted the data analysis. The manuscript was

815 written by J.L., S.Z., X.T., and S.P.

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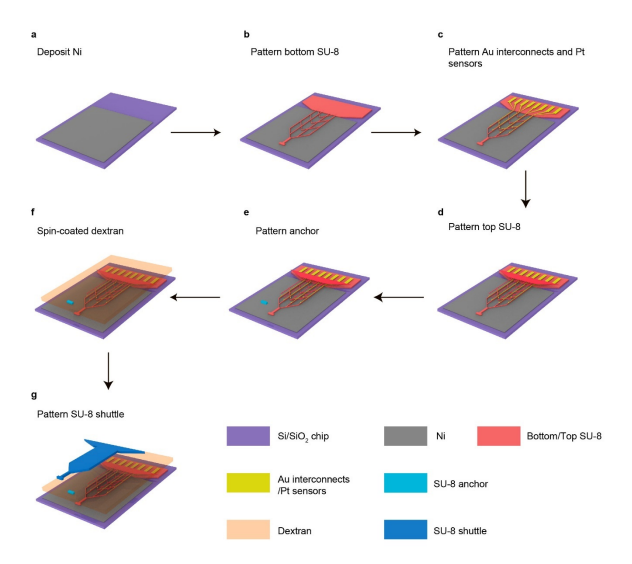
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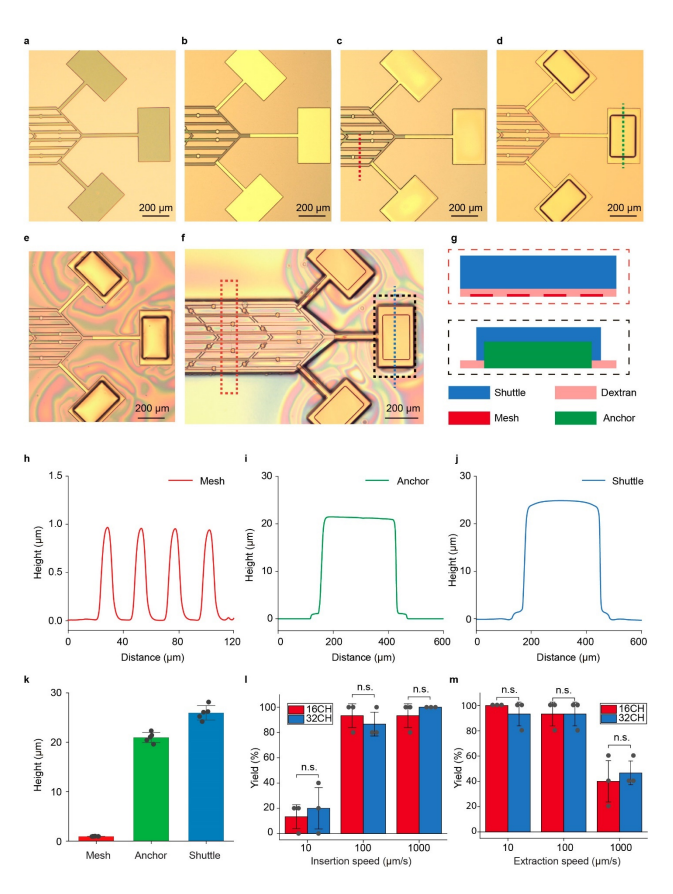
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928 Extended Data Figures



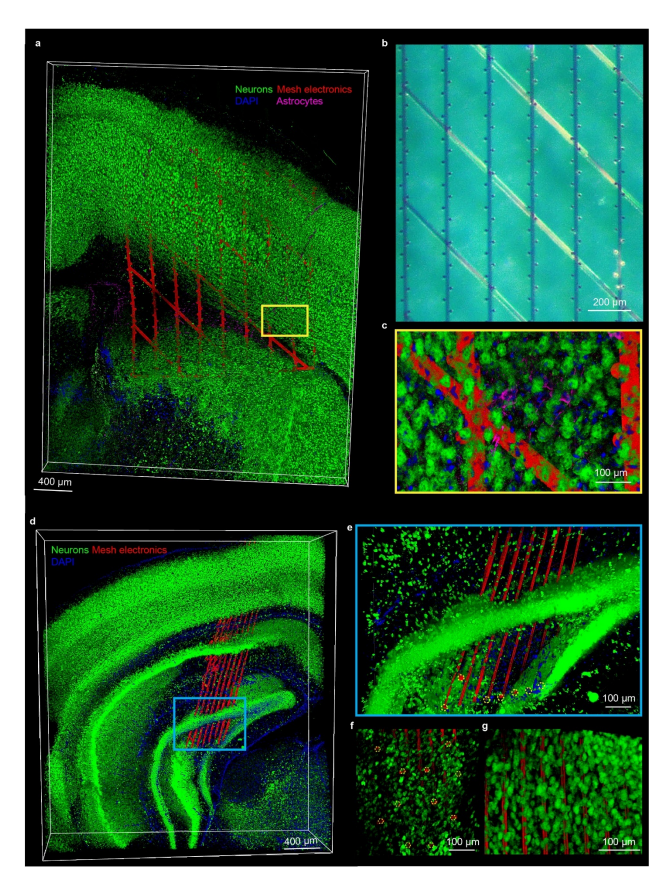
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Extended Data Fig. 1 | Schematics show the stepwise fabrication of monolithically integrated 930 931 tissue-level flexible mesh nanoelectronics. a, A Ni sacrificial layer (grey) was defined by photolithography and deposited through thermal evaporation on the Si/SiO₂ wafer (purple). **b**, SU-932 8 2000.5 bottom passivation layer (red) was defined by photolithography. c, Cr/Au interconnects 933 and Pt microelectrodes (yellow) were sequentially defined by photolithography and deposited 934 through electron beam (e-beam) evaporation on the top of the SU-8 passivation layer. d, SU-8 935 2000.5 top passivation was defined by photolithography (red). e, SU-8 2025 anchor was defined 936 937 by photolithography (cyan). f, Dextran sacrificial layer (pink) was spin coated. g, SU-8 2025 shuttle was defined by photolithography (navy). 938

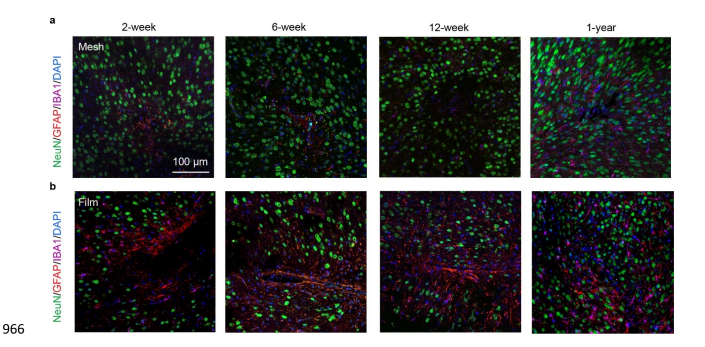


Extended Data Fig. 2 | Anchor structures for controllable implantation. a-f, Optical images 940 941 illustrating each step of the fabrication corresponding with Extended Data Fig. 1b-g, respectively. g, Schematics showing the cross-section of the monolithically integrated mesh nanoelectronics at 942 the red and black dashed boxes highlighted regions in (f), respectively. h-j, Contact profilometer 943 measurements along with the open mesh structure in (c, red dashed line), anchor structure in (d, 944 green dashed line), and shuttle structure in (f, blue dashed line). k, Statistical summary of the 945 thickness of the open mesh, anchor, and shuttle layer structures (n = 5). I, m, Insertion (I) and 946 extraction (m) yield of 16-channel and 32-channel mesh nanoelectronics with different speeds (n.s.: 947 not significant, two-tailed unpaired t test, n = 3 times experiment, each time include 5 948

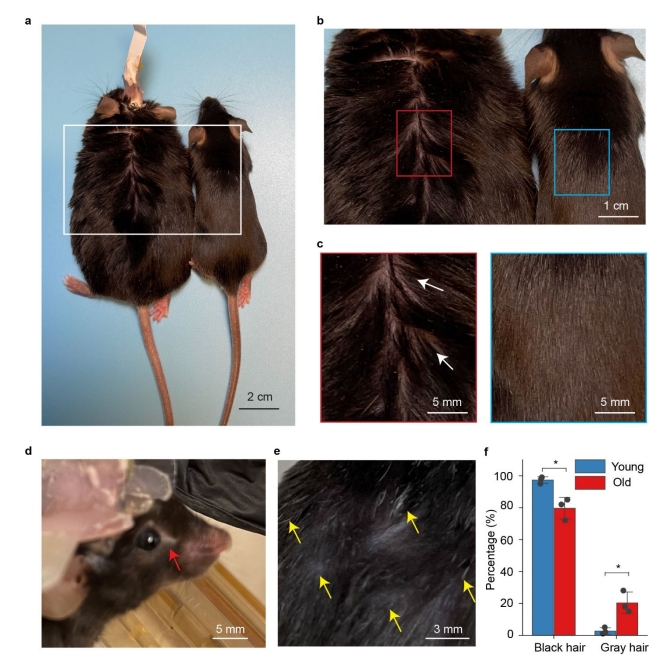
949 insertion/extraction).



Extended Data Fig. 3 | Independent replicates of mesh nanoelectronics with different sizes 951 952 were implanted in mice brains. a, 3D reconstructed confocal fluorescence imaging of neuron (green), nuclei (blue), and astrocytes (purple) with 1024-channel mesh nanoelectronics (red) 953 954 sustaining their open mesh structure across multiple brain regions at 6-week post-implantation. **b**, Representative photograph illustrates the high density, 1024-channel mesh nanoelectronics. c, 955 Zoom-in views of the regions highlighted by yellow (c) box in (a). d, A 3D reconstructed interface 956 of neurons (green), nuclei (blue) with shape maintained 16-channel mesh electronic (red) at 6-957 week post-implantation. The mesh electronic was across the cortex and hippocampus with a 958 designed 30-degree angle corresponding to the dorsal-ventral direction. e, A zoom-in view of the 959 hippocampus region highlighted by the cyan box in (a). f, g, Neuron interpenetration inside the 960 961 subcellular electrode, individual electrodes are indicated by vellow dashed circles in (e) and (f)and opening mesh structure (g). These results show neuron interpenetration inside the opening 962 mesh and minimal astrocyte increases at the surface and interior of mesh, and thus demonstrate 963 the reproducibility of these results in Fig. 2. 964

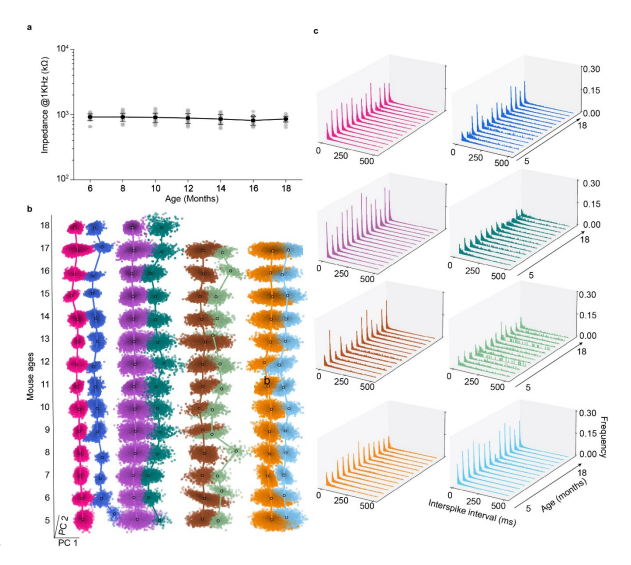


967 Extended Data Fig. 4 | Time-dependent histology studies of brain tissue reaction to ~1 μ m-968 thick open mesh/thin-film nanoelectronics. a, b, Representative immunofluorescence images of 969 brain tissue reaction following 2-week, 6-week, 12-week, and 1-year post-implantation of a mesh 970 (a) and a thin-film (b) nanoelectronics from the contralateral hemisphere. The tissue was labeled 971 for astrocytes (red), microglia (purple), neurons (green), and nuclei (blue). Time-dependent 972 histology studies have been repeated on n = 5 independent samples for each time point, with 973 statistical analyses shown in Fig. 2f-i.

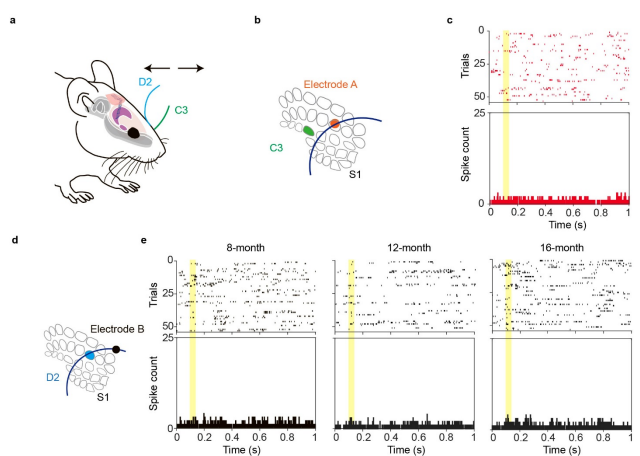


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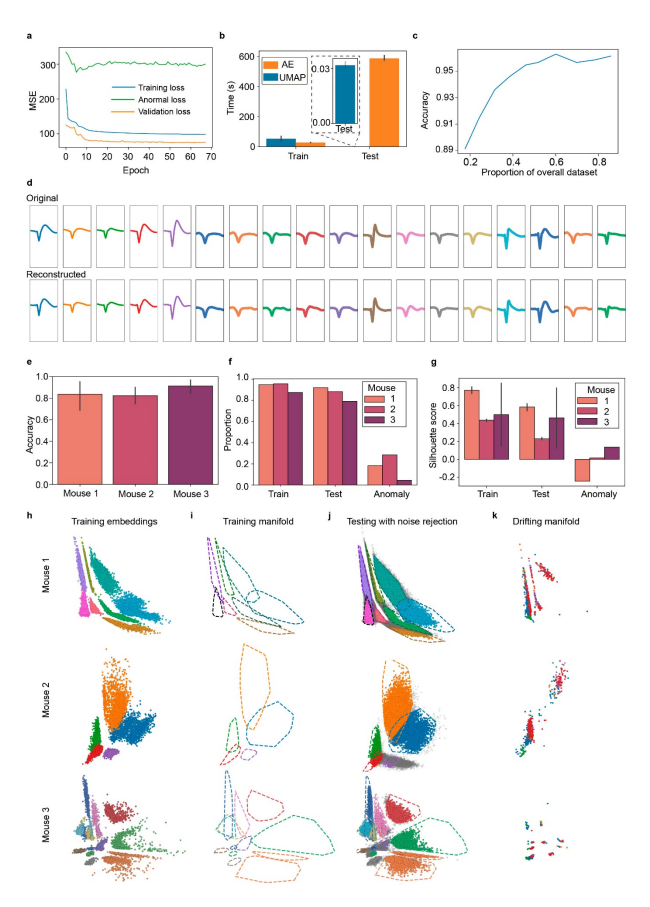
Extended Data Fig. 5 | Aged mice characterization. a, Representative photograph showing the 975 old mouse of weight gain (18 months) with tissue-like mesh nanoelectronics implant (left) 976 compared with the mature adult mouse (5 months, right). b, Zoom-in views of the regions 977 highlighted by white boxes in (a). c, Zoom-in views of the thinning hair (white arrows) of aged 978 mouse and glossy brown fur of mature adult mouse highlighted by red and blue boxes in (b). d-e, 979 Representative photograph showing the barbering around eyes (d, red arrow), grey and thinning 980 fur in the dorsal back skin (e, yellow arrows) of the aged mouse (18 months) with mesh 981 nanoelectronics implant. Statistical analysis reveals that significantly increased gray hairs in dorsal 982 back skin in old-aged mice (*p < 0.05, two-tailed unpaired t test, n = 3). 983



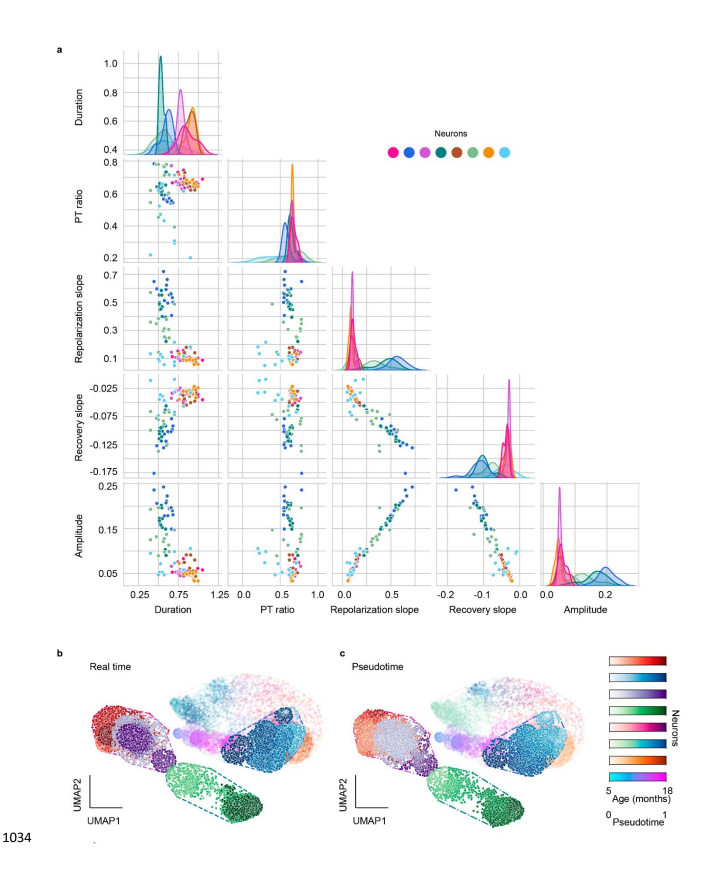
Extended Data Fig. 6 | Long-term stable recording characterization. a, Time-dependent 985 electrode interfacial impedance at 1 kHz measured by the Cereplex Direct (Blackrock 986 987 Microsystems, USA). Data represented mean \pm SD, individual data points are overlaid. The electrode interfacial impedances exhibited relatively constant values of $920.2 \pm 107.2 \text{ k}\Omega \text{ vs. } 857.2$ 988 989 \pm 85.7 k Ω (mean \pm SD, n = 30) at months 6 vs.18. **b**, Time evolution of representative single-unit spikes clustered by Leiden clustering. The x- and y-axes denote the first and second PC dimensions, 990 respectively, and the z-axis denotes mouse age in months. Dimension-reduced clusters associated 991 with a single unit are shown (Fig. 4a) using the same color. These data show stable clusters with 992 993 nearly constant positions in the first and second principal component plane (PC1-PC2) over the entire recording period from 5 months to 18 months. c, Time evolution of interspike interval (ISI) 994 histograms of representative neurons identified in Fig. 4 from 5 months to 18 months. The x- and 995 996 y-axes denote the time between subsequent action potentials of spontaneous firing neuron, and mouse age in months, respectively, and the z-axis denotes frequency. Bin size, 2 ms. 997 998



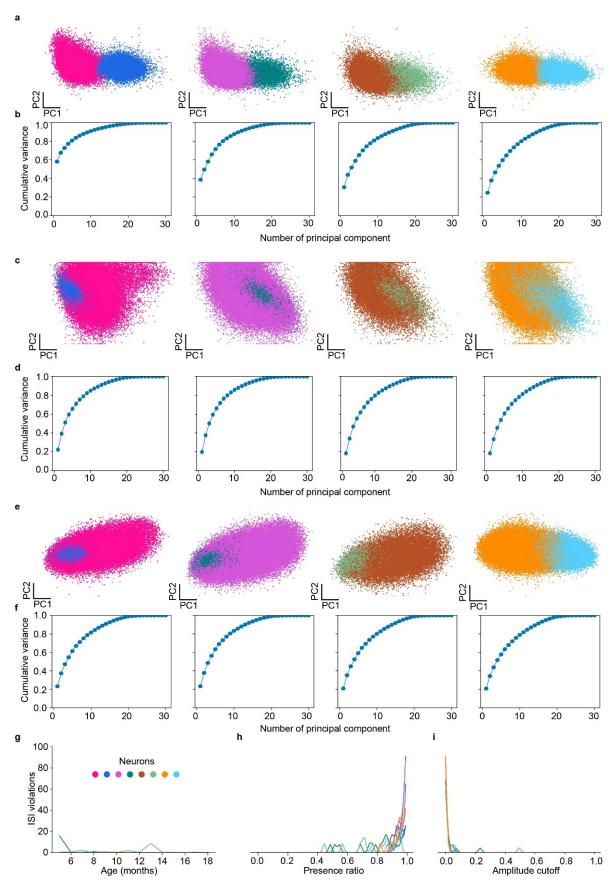
999 Extended Data Fig. 7 | Validation of whisker-related activity using sham stimulation. a, 1000 Schematic diagram of whisker deflection. Individual vibrissa was deflected in the rostral-caudal 1001 1002 plane using a computer-controlled piezoelectric bending actuator. D2 and C3 whiskers are labeled in blue and green, respectively. b, Schematic diagram of whisker barrel and electrode arrangement 1003 in S1. C3 whisker barrel and electrode A position is labeled in green and orange, respectively. c, 1004 Raster plot and peri-stimulus time histogram (PSTH, 1 ms bin size) of electrode A show no 1005 observable spiking activities when applied to the C3 whisker at 8 months. d, D2 whisker and 1006 electrode B position are labeled in blue and black in S1 barrel field, respectively. e, Raster plot 1007 1008 and PSTH (1 ms bin size) of electrode B show no observable spiking activities when applied to the D2 whisker from 8 months to 16 months. 1009



Extended Data Fig. 8 | Autoencoder-based spike sorting and stability analysis a, 1011 Representative training, validation, and drift loss curves obtained during the training process of 1012 one of the autoencoders. b, Time performance comparison of autoencoder vs computational 1013 1014 pipeline allowing for dimensionality reduction and classification (UMAP and a Random Forest classifier trained on UMAP embeddings). Tests were performed using a single CPU with Intel i5. 1015 8 cores @ 4.1GHz. The autoencoder has a significant time advantage for inference. c, Overall 1016 accuracy on the testing dataset as a function of the proportion of the overall dataset used for 1017 training the network. d, Original and reconstructed average neuron waveforms for the second and 1018 third mice data. Original waveforms on the top row are colored by within-mouse neuron labels 1019 (the first 5 neurons from the left are from mouse 2) while reconstructed waveforms on the bottom 1020 1021 row are colored by autoencoder classifier predicted labels. Reconstruction and classification are 1022 near-perfect. e, Bar plot of per-mouse classification accuracy calculated for each neuron label class. f, Proportion of spikes kept for each dataset when using MSE-based thresholds for drift detection 1023 (Fig. 5d). g, Silhouette score calculated using autoencoder embeddings of training, testing and drift 1024 1025 datasets with associated true neuron labels. The training and testing scores reflect the emergent latent space cluster separability for observed neurons while the anomalous scores highlight the 1026 1027 poor separability of previously unseen embedded neuron waveforms. h, i, j, Stability verification process illustrated for all three mice: visualizing training latent space embeddings (dots colored by 1028 their true neuron label), calculating training manifold boundaries, and finally applying these 1029 1030 boundaries to quantify the proportion of testing dataset spikes latent embeddings lying inside the predicted neuron's training manifold. k, Visualization of the latent embeddings of drift spikes. 1031 Dots are colored by their true neuron labels; mixed colors within clusters showed poor ability to 1032 separate different neurons from the drift dataset, in accordance with (g). 1033



Extended Data Fig. 9 | Feature selection and pseudo time analysis for potential aging-1035 1036 associated analysis a, Pairplot of 5 AllenSDK selected features shown in Fig. 4e-h. Dots represent mean values of paired features calculated over neuron clusters over the entire recording period. 1037 1038 Diagonal subplots show neuron feature univariate distributions using kernel density estimators. **b**, c, Real time (b) and pseudotime (c) comparison of UMAP embedding time-evolution for neurons 1039 in Fig. 4. The x- and y axes denote the first and second UMAP dimensions, respectively. Each 1040 gradient color-coded cluster represents a distinct neuron in Fig. 4. The color bars show the mouse 1041 age time points from 5 to 18 months and the corresponding pseudo time from 0 to 1. Highlighted 1042 dots correspond to the representative neurons used in the aging analysis of Fig. 6. Delimiting lines 1043 are the convex hull of a neuron's UMAP embedding time-evolution. 1044



Extended Data Fig. 10 | Quality assessment for PCA embeddings and neuron clusters. a, 1046 1047 Principal Component (PC) embeddings of all recorded spikes without any normalization colored by neuron. **b**, Cumulative proportion of variance explained by top principal components of (**a**). **c**, 1048 1049 d, Same as (a, b) with prior min-max normalization to scale spikes to (-1, 1) range. e, f, Same as (a, b) with prior standard scaling normalization per spike. Normalization decreases cluster 1050 separability in PC representation and lowers proportion of variance explained by the top two 1051 components by removing amplitude information from spikes. g, Interspike interval (ISI) violations 1052 in the absolute number of spikes calculated over all recordings for each recording month. h, 1053 Presence ratio smoothed density plot over all recording sessions per unit. i, Amplitude cutoff 1054 smoothed density plot over all recording sessions per unit. 1055