

# Multiscale Cloud-based Pipeline for Neuronal Electrophysiology Analysis and Visualization

Jinghui Geng<sup>1,4,\*</sup>, Kateryna Voitiuk<sup>1,2,4</sup>, David F. Parks<sup>1,2,4</sup>, Ash Robbins<sup>1,4</sup>, Alex Spaeth<sup>1,4</sup>, Jessica L. Severson<sup>1,4</sup>, Sebastian Hernandez<sup>1,4</sup>, Hunter E. Schweiger<sup>1,4</sup>, John P. Andrews<sup>1,4</sup>, Spencer T. Seiler<sup>1,4</sup>, Matthew A.T. Elliott<sup>1,4</sup>, Edward F. Chang<sup>5,6</sup>, Tomasz J. Nowakowski<sup>5,6,7,8,9</sup>, Rob Currie<sup>4</sup>, Mohammed A. Mostajo-Radji<sup>4</sup>, David Haussler<sup>1,2,4</sup>, Tal Sharf<sup>1,2,4</sup>, Sofie R. Salama<sup>1,2,4</sup>, Mircea Teodorescu<sup>1,2,4,\*</sup>

<sup>1</sup>Department of Electrical and Computer Engineering, University of California Santa Cruz, Santa Cruz, CA, USA

<sup>2</sup>Department of Biomolecular Engineering, University of California Santa Cruz, Santa Cruz, CA, USA

<sup>3</sup>Department of Molecular, Cell, and Developmental Biology, University of California Santa Cruz, Santa Cruz, CA, USA

<sup>4</sup>Genomics Institute, University of California Santa Cruz, Santa Cruz, CA, USA

<sup>5</sup>Department of Neurological Surgery, University of California San Francisco, San Francisco, CA, USA

<sup>6</sup>Weill Institute for Neurosciences, University of California San Francisco, San Francisco, CA, USA

<sup>7</sup>The Eli and Edythe Broad Center of Regeneration Medicine and Stem Cell Research, University of California San Francisco, San Francisco, CA, USA

<sup>8</sup>Department of Psychiatry and Behavioral Sciences, University of California San Francisco, San Francisco, CA, USA

<sup>9</sup>Department of Anatomy, University of California San Francisco, San Francisco, CA, USA

\*Correspondence Author. Email: [jgeng2@ucsc.edu](mailto:jgeng2@ucsc.edu), [mteodorescu@ucsc.edu](mailto:mteodorescu@ucsc.edu)

## ABSTRACT

Electrophysiology offers a high-resolution method for real-time measurement of neural activity. The vast amount of data generated requires efficient storage and sophisticated processing to extract neural function and network dynamics. However, analysis is often challenging due to the need for multiple software tools with different runtime dependencies. Longitudinal recordings from high-density microelectrode arrays (HD-MEAs) can be of considerable size for local storage, complicating data management, sharing, and backup. To address these challenges, we developed an open-source cloud-based pipeline to store, analyze, and visualize neuronal electrophysiology recordings from HD-MEAs. This pipeline is dependency agnostic by utilizing cloud storage, cloud computing resources, and an Internet of Things messaging protocol. We containerized the analysis algorithms to serve as scalable and flexible building blocks within the pipeline. We designed graphical user interfaces and command line tools to remove the requirement of programming skills. The interactive visualizations provide multi-modality information on various neuronal features. This cloud-based pipeline is an efficient solution for electrophysiology data processing, the limitations of local software tools, and storage constraints. It simplifies the electrophysiology data analysis process and facilitates understanding neuronal activity. In this paper, we applied this pipeline on two types of cultures, cortical organoids and *ex vivo* brain slice recordings.

## INTRODUCTION

Recent advances in hardware and software platforms for neuronal recordings have enabled simultaneous recording of neuronal activity with high spatial and temporal resolution across var-

ious samples, including brain slices<sup>112</sup>, 2D cultures<sup>3-5</sup>, and 3D cerebral organoids<sup>67</sup>. These technologies facilitate comprehensive studies of brain function, neurodevelopment, and network topology<sup>8-10</sup>. However, the exponential growth in data volume and complexity<sup>11-13</sup> presents significant challenges in data storage, processing, and analysis. Recordings, images, and analysis results can consume substantial storage on computers and hard drives. Interpreting this multi-dimensional data requires specialized algorithms and tools to extract single neuronal unit activity, visualize firing patterns, and understand neuronal network-level information<sup>14-16</sup>. While efforts have been made to unify standards in electrophysiology, biologists still face difficulties performing comprehensive analyses.

Spike sorting algorithms are crucial for analyzing multi-electrode array (MEA) recordings<sup>17-21</sup>, identifying and categorizing individual neuronal spikes from raw voltage traces to analyze neuronal features<sup>22-25</sup> and network dynamics<sup>26,27</sup>. While various software tools have been developed to process MEA recordings and visualize neuronal features<sup>23,28-33</sup>, challenges persist due to differing programming languages, limited user support, and compatibility issues. Although integrated platforms offer end-to-end analysis capabilities, they may restrict custom data manipulation, requiring researchers to develop their own workflows and navigate steep learning curves for effective data interpretation.

Cloud computing enables processing a large amount of data in parallel by utilizing abundant resources while still being a cost-effective solution<sup>34-36</sup>. Cloud-based storage can address the issue of massive experimental data filling up local disks. It also provides extensive data sharing ability for collaborations across research labs. Infrastructures and web platforms have been developed to store and analyze various types of data, including electrophysiology, neuroimaging, and sequencing<sup>37-42</sup>. These platforms are designed to benefit the broader neuroscience community, emphasizing data publication and sharing<sup>43,44</sup>. A research laboratory-oriented data platform is needed to support consistent experiments and data processing.

The Internet of Things (IoT) has made a significant impact in many fields, including healthcare<sup>45,46</sup> and, in recent years, has been applied to cellular biology<sup>47,48</sup> and in vitro electrophysiology experiments<sup>49-51</sup>. Its resource efficiency enables the messaging protocol to work across different hardware, allowing networks to grow from a few devices to a large number without compromising performance.

We developed a cloud-based pipeline for electrophysiology data storage, processing, and sharing to facilitate the day-to-day research. We used containerization as the minimum building block. The IoT messaging services and data analysis algorithms are packaged into individual containers. The IoT services run on a web server to stream data, monitor processing tasks, and communicate with researchers through user interfaces. We applied Kubernetes<sup>52</sup> to orchestrate the analysis containers on the cloud computing clusters. By using cloud computing resources, the pipeline can process a large number of datasets with different algorithms in parallel, optimizing resource utilization, scalability, and flexibility. Moreover, we lower costs by replacing local computing hardware, such as CPUs and GPUs, with cloud-based technology. We also remove the barriers to data analysis by providing user interfaces, minimizing the software setup process, and making the Python code open source. The pipeline provides a suite of algorithms, including spike sorting, autocuration of putative neural units, visualization, and downstream analyses for specific goals using the curated data. We tested this pipeline with two applications. First, we analyzed mouse cortical organoid longitudinal recordings, 10 minutes long, one hour apart, over a 7-day period. This demonstrated the utility of our approach for neuron tracking. Second, we applied the pipeline to study optogenetic modulation of epileptiform activity in human hippocampus slices, contributing to our understanding and potential treatment of neurological diseases.

## 69 RESULTS

70 Our platform allows users to upload recordings from electrophysiology devices directly to cloud  
71 storage. The data is organized by experiment date and is annotated with automatically extracted  
72 as well as user-specified metadata. The pipeline can be scaled up as algorithms and services  
73 are containerized, making it easy to integrate new analytical tools as they become available. The  
74 pipeline supports multiple data processing paradigms to accommodate diverse research require-  
75 ments. The graphical interface allows users to initiate, monitor, and visualize data processing  
76 after upload, offering multimodal analysis and result downloads. An integrated IoT messaging  
77 service connects users, local recording devices, and the cloud, streamlining workflow.

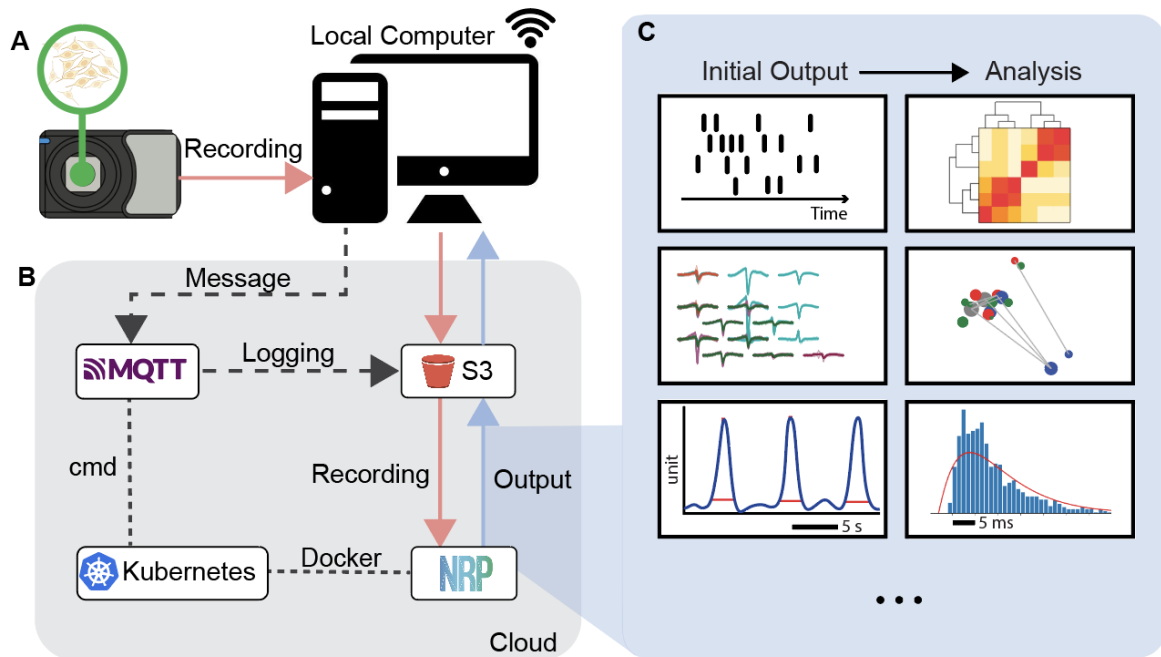
### 78 Framework Design

79 The pipeline is generic and capable of processing data from any electrophysiology platform that  
80 uses HDF5 and NWB<sup>53-55</sup> formats. In this paper, we tested it with data generated by a MaxOne  
81 HD-MEA (MaxWell Biosystems)<sup>56</sup>. The system has 26,400 electrodes in a 2.10x3.85mm<sup>2</sup> area.  
82 It supports data collection from 1,020 channels and can simulate 32 channels simultaneously at  
83 a 20kHz sample rate. Together with a small inter-electrode pitch (17.5µm), the system provides  
84 high temporal and spatial resolution, where the activity of a typical neuron will be recorded on  
85 multiple pads. We utilize Ceph S3 and the National Research Platform (NRP) computing clusters  
86 for data storage and processing.

87 The overview of the platform is shown in Figure 1. Neuronal tissue culture activity data is col-  
88 lected on a MaxWell MEA headstage, connected to a local computer running MaxLab software  
89 (Figure 1A). After recording, datasets are streamed to S3 and the data uploader generates cor-  
90 responding metadata and maintains the applicable S3 file structure for these datasets (Figure  
91 1B). Upon completion, an MQTT message is sent from the data uploader to the processing ser-  
92 vice – the job listener. This message contains the experiment identifiers and the image of the  
93 dockerized algorithm. The listener parses the message to gather the S3 paths for each dataset  
94 and calls the Kubernetes-Python API to deploy data processing jobs to the NRP computing clus-  
95 ter 15. The pipeline provides several containerized data processing applications, including spike  
96 sorting, data curation, and visualization. Once a job is completed on the NRP, the result is saved  
97 to S3 (Figure 1C). Researchers can access and download these results through the user inter-  
98 face.

99 To make the pipeline accessible to non-programmers, we have developed user interfaces for  
100 managing and interacting with both local and remote data processes (Figure 2A). Through these  
101 interfaces, researchers can have complete control over their data while bridging the gap between  
102 running complex algorithms and requiring extensive programming knowledge or technical exper-  
103 tise.

104 These interfaces include a data uploader, a Dashboard webpage, and a Slack channel, each  
105 serving distinct purposes while bridging local data collection, cloud-based data manipulation,  
106 and user notifications. The data uploader, installable on local laptops, enables users to upload  
107 electrophysiology recordings to S3 storage and initiate batch processing jobs with predefined  
108 parameters after the experiment is finished. The web Dashboard, accessible from any internet-  
109 connected device, provides access to existing S3 data for both batch and chained jobs. The data  
110 uploader and the Dashboard support downloading files from S3 to local directories. Addition-  
111 ally, the Dashboard features a visualization page displaying post-processing figures of selected  
112 recordings. A Slack channel is used to post status notifications for data processing jobs. Detailed  
113 descriptions of these user interfaces are provided in the Methods section. Screenshots of the ap-  
114 plications are shown in Supplementary Figure S2, S3.



**Figure 1: Cloud-based electrophysiology data processing pipeline architecture.** (A) Electrophysiology data from neuronal cultures is recorded on a local computer. Different neuronal cultures and their recordings are shown in Figures 4 and 7. (B) Once the dataset is saved, it is uploaded to a uniquely identified data bucket AWS S3 for permanent storage using the Uploader. An MQTT message is simultaneously sent to the job listener service to initiate data processing jobs. These jobs run containerized algorithms and are launched on the National Research Platform (NRP) computing cluster using Kubernetes. Results, including post-processed data and figures, are saved back to AWS S3. (C) The analysis outputs various interactive analytical figures for each dataset's network features and single-unit activity.

115 For cloud integration, we used the Message Queuing Telemetry Transport (MQTT) messaging  
116 protocol, a lightweight publish-subscribe protocol designed for Internet-of-Things (IoT) appli-  
117 cations. This approach reduces the dependency requirements for edge devices to run cloud-  
118 computing jobs. A local computer can utilize the pipeline as long as it can run a Python envi-  
119 ronment and has a network connection. We have designed job listener and scanner services to  
120 run and monitor jobs on the cloud (Figure 2B,C). For cloud computing, we used the National Re-  
121 search Platform (NRP), a distributed commodity compute cluster based on Kubernetes and the  
122 Ceph distributed file system. It has special CPUs and GPUs for data science, simulations, and  
123 machine learning. This setup allows for parallel data processing and can help reduce the com-  
124 puting infrastructure cost of individual labs.

125 We have a job scanner (Figure 2B) that checks on data processing jobs in the cloud every 30  
126 minutes. It updates a list of current job statuses using the Kubernetes Python API. The scanner  
127 reads job names and information, which are named based on the dataset or a job list. This helps  
128 the scanner find the correct information in the NRP. The scanner then updates the listener and  
129 the user about how jobs are progressing.

130 To keep the flexibility of data processing, we implemented two types of jobs: batch processing  
131 and chained jobs. Chained jobs run through several steps on different data, with subsequent  
132 processing dependent on prior results. When the scanner detects a status change in a chained  
133 job, it sends a message to the listener to update the corresponding job look-up table and initiate  
134 the next processing step. Concurrently, it notifies the user about completing the prior job and the  
135 start of the next. This notification is done through the “slack-bridge” service. For completed batch  
136 processing jobs, the scanner sends only a user notification. After a message is sent, finished  
137 jobs are removed from the scanner’s memory to prevent duplicate notifications.

138 The job listener (Figure 2C) receives messages from both the user interface and the scanner. It  
139 also sends user notifications to the “ephys-pipeline” Slack channel. The primary function of the  
140 job listener is to initiate cloud computing jobs. Upon receiving a run job message, the listener  
141 parses it to extract the data path, data format, parameter setting, and job type (analysis algo-  
142 rithms). The listener then calls functions from a Python Kubernetes object (Figure 2) to allocate  
143 computing resources on NRP and the appropriate analysis docker image for each dataset. This  
144 object creates a job on the NRP and a pod within each job. Finally, it sends a “job created” no-  
145 tification to the Slack channel. Both the scanner and listener services maintain logs on S3 for  
146 historical tracking and ease of maintenance. These logs are updated after each new message is  
147 received or sent.

148 The data organization on S3 (Figure 2D) is structured based on data types and characteristics.  
149 Electrophysiology recordings are grouped by experiment batch, assigned a universally unique  
150 identifier (UUID), and paired with a “metadata.json” file for overall content description and ex-  
151 periment notes. We create sub-buckets: “original/data” for raw recordings and “derived/algo” for  
152 analysis output, where “algo” represents the algorithm used to analyze the data. Additionally, we  
153 maintain a “service” bucket for the chained job scheduler and logging of listener and scanner ac-  
154 tivities. Since the computing clusters are designed to run containerized data processing jobs, we  
155 have created docker images for electrophysiology algorithms with minimum software dependen-  
156 cies.

157 As illustrated in Figure 2E, when the listener deploys a job to NRP, the platform assigns a node  
158 with all the requested resources. The node creates a pod, pulls the docker image from Docker-  
159 Hub, and retrieves data from S3 to run the analysis. The processing results are then uploaded  
160 back to S3 from the container. Figure 2F demonstrates an example of a containerized batch pro-



cessing algorithm. In this container, a Python script reads an electrophysiology recording, performs spike sorting on the raw data to identify putative firing neurons (single units), applies autocuration to preserve high-quality units, and generates both visualization figures and spike data for the recording. The spike data is stored as a NumPy data structure with temporal and spatial information of the single units.

Figure 2G shows the Python Kubernetes object configuration in the listener for job execution. This configuration specifies the number of CPUs and GPUs and the amount of memory and storage required to run a specific container. These resource allocations are calculated based on the algorithm workload and data size, optimized for efficient utilization of cloud computing resources. To execute a specific container, the configuration is provided with the corresponding docker image, input data (such as the recording or derived results from the recording), and metadata (including data format or parameter settings). Examples of Kubernetes configurations can be found in Supplementary Table S2.

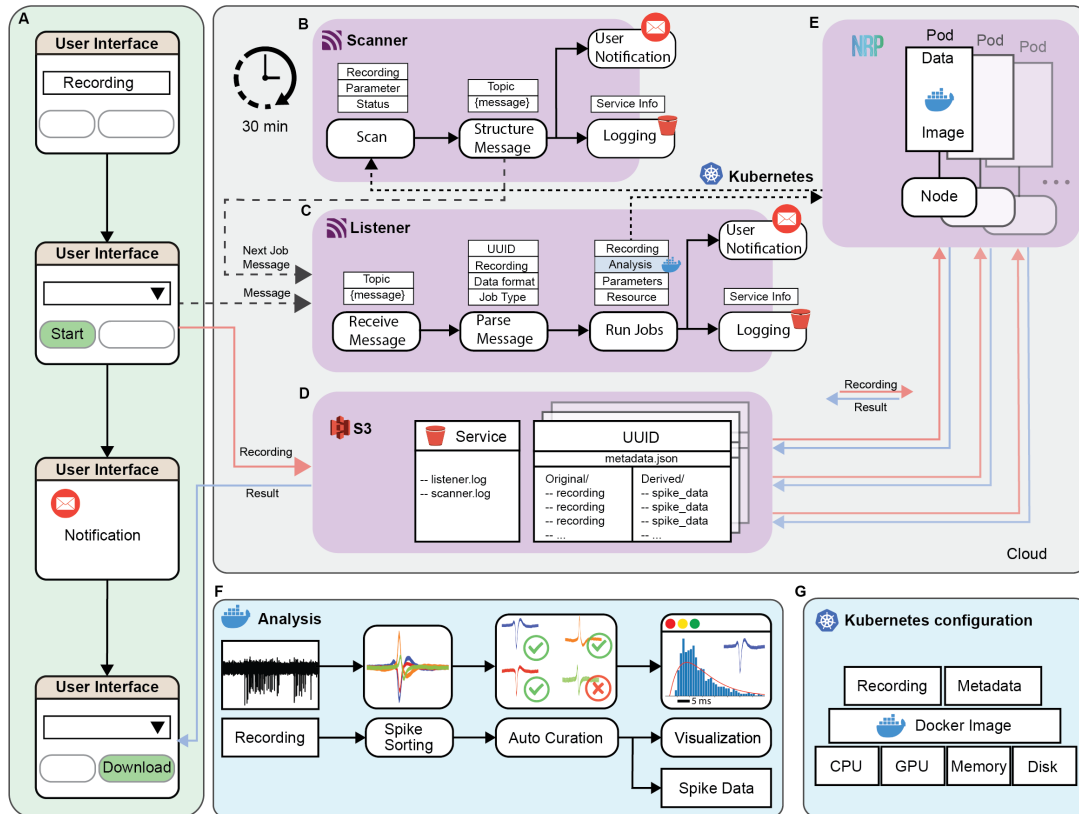
### **This Pipeline Enables Versatile Jobs**

Data processing and analysis often require multiple iterations for new experiments due to changes in recording hardware, biological samples, and data requirements. To ensure versatility in data processing jobs, we developed a minimum building block for the pipeline and designed various job execution paradigms.

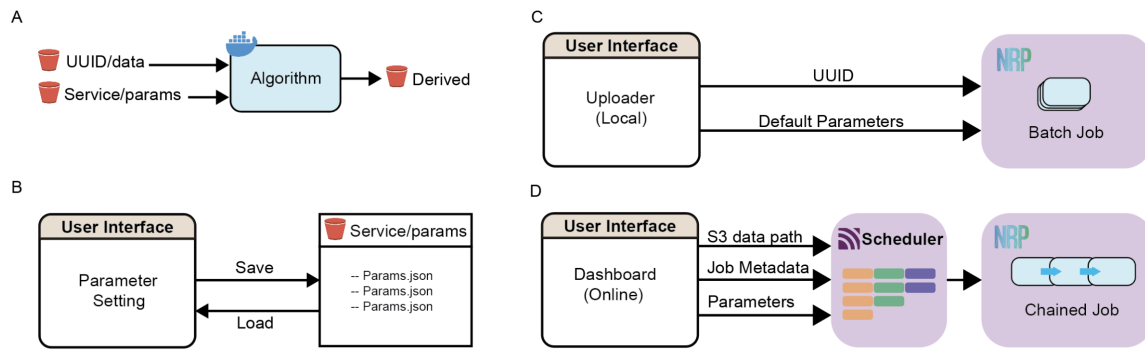
Figure 3A shows the minimum building blocks of our pipeline. It includes paths to S3 data storage and a containerized algorithm. Each algorithm needs two inputs (data and parameters) and produces one output file with results (processed data, visualization figures, and logs). We store input data and outputs in designated buckets on S3 under each UUID. We keep these parameters in designated sub-buckets (“service/params/algo”) on S3, named after each algorithm. Users can pick existing parameters or make new ones on the Dashboard’s “Job Center” page (Figure 3B, Supplementary Figure S3).

The pipeline supports both batch processing and chained jobs (Figure 3C,D). Batch processing enables the analysis of numerous recordings using identical parameter settings. All jobs can be processed in parallel on NRP. Users can initiate a batch job from the local data uploader after the experiment. In batch processing, each recording undergoes spike sorting, autocuration, and visualization. Detailed descriptions of these three steps can be found in the Methods section.

As algorithms are packaged in individual docker containers as minimum building blocks, multiple analysis jobs can be chained for a recording, with stage results passed to subsequent jobs upon completion of the previous job (Figure 3D). To implement this functionality, we designed a CSV job scheduler integrated into the Dashboard, Listener, and Scanner services. When users select recordings and a list of analysis jobs from the Dashboard, a CSV file is generated, with each row representing an analysis job. Columns contain sufficient information to initiate the job, including the S3 data path, computing resource requirements (job metadata), and parameter settings. We use the “next\_job” column to index the row of the job to run after the current row, allowing for multiple indices. After saving this CSV file to S3, the Dashboard sends a message to the Listener to start the first stage jobs by indexing them in the message body. We create the NRP job name using the CSV file name, enabling the Scanner to differentiate chained jobs from batch jobs by simply parsing the name. Upon completing the first stage jobs, the Scanner sends the Listener an “update” message. The Listener then checks for any available “next\_job” in the CSV file and launches the second-stage jobs. Detailed information on job chaining can be found in the Methods section.



**Figure 2: Pipeline components and workflow.** (A) The user interface allows researchers to upload their electrophysiology recordings to cloud storage, initiate data processing jobs, receive notification upon completion, and download results to local computers. (B) MQTT-based job scanner service monitors job status on the NRP, sends a message to the listener for the next job, and notifies users. (C) MQTT-based job listener service that subscribes to specific topics to run data processing jobs. When the service receives a message, it parses the JSON format to extract experiment identifiers and computing requirements, then deploys jobs to NRP through Python-Kubernetes API. Both scanner and listener services update their status to S3 log files on a scheduled basis. (D) S3 file structure for service logging and experiment data. Log files are human-readable text files that track service status. Experiment data is stored in batches, each with a unique identifier (UUID), metadata file, “original” bucket for experiment data, and “derived” bucket for analysis outputs. (E) Computing cluster (NRP) for running containerized jobs using Kubernetes. (F) An analysis container for batch processing is capable of loading electrophysiology recordings, running spike sorting and autocuration algorithms, producing visualization figures, and generating Numpy files for single units. (G) Kubernetes configuration for job deployment to a computing cluster.



**Figure 3: Minimum building block and job types.** (A) The minimum pipeline building block utilizing dockerized algorithms and S3 data storage. Data and parameter settings are retrieved from S3, processed by containerized algorithms on NRP, and results are uploaded back to S3. (B) Users can save and load parameter settings to and from the S3 “service” bucket through the Dashboard. (C) Batch processing of numerous recordings is achieved by providing UUID and default parameter settings to the pipeline. Users can initiate this process through the local data uploader. (D) Chained jobs are implemented using a CSV job scheduler containing S3 data paths, job metadata, and parameter settings. Users can initiate job chaining from the online Dashboard.

## 206 Pipeline Output for Individual Recordings

207 The pipeline output is designed to be comprehensive, structured, and accessible so the data can  
 208 be reproduced and distributed easily. Using batch processing algorithms, for example, each pro-  
 209 cessing step produces one compressed file (zip format). For spike sorting, the compressed file  
 210 is compatible with Phy GUI<sup>57</sup>. Users can download the file, uncompress it, and open it in Phy to  
 211 check the sorting result and perform manual curation. We also developed a function to load the  
 212 data directly into a Python object, enabling automated downstream analysis of the single-unit  
 213 features. Autocuration, the second step, outputs a compressed file (zip format) containing a spike  
 214 data object in NumPy array and Python dictionary. This object consists of a spike train list, a neu-  
 215 ron data dictionary, the recording’s sample rate, and electrode configuration. The neuron data  
 216 dictionary has spatial information such as the channel’s coordinates, neighbor channels, and  
 217 spike features such as waveform and amplitude. The spike train list and the neuron data dictio-  
 218 nary index match each other. The size of the autocuration file is approximately 10 times smaller  
 219 than the spike sorting output by re-constructing the data. For the final step, data visualization, the  
 220 pipeline generates interactive HTML format figures for the recording and a PNG format figure for  
 221 each single unit. All of the output files have a log to keep track of the actions and decisions made  
 222 by the algorithm. To make the data structure consistent, other algorithms’ outputs that are pro-  
 223 duced by this pipeline are also sorted into NumPy arrays and dictionaries. These outputs can be  
 224 easily converted to Pandas DataFrame and distributed as tabular data.

225 Figure 4 illustrates the visualization output for a 10-minute recording from a mouse cortical organoid  
 226 on day 42 in culture. Figure 4A is a photograph of the mouse cortical organoid on the HD-MEA.  
 227 Initially, two organoids were plated on the same HD-MEA for this experiment. As the majority  
 228 of activity originated from the right organoid, our analysis was focused on this organoid. The  
 229 pipeline’s interactive HTML overview figure includes a footprint map showing the spiking wave-  
 230 forms on the corresponding electrode locations (Figure 4B). The HD-MEA can detect a unit’s  
 231 footprint by multiple electrodes and potentially show the neuron’s orientation. Since a single elec-  
 232 trode can record activity from many neurons, different colors are used to label the units. Along-



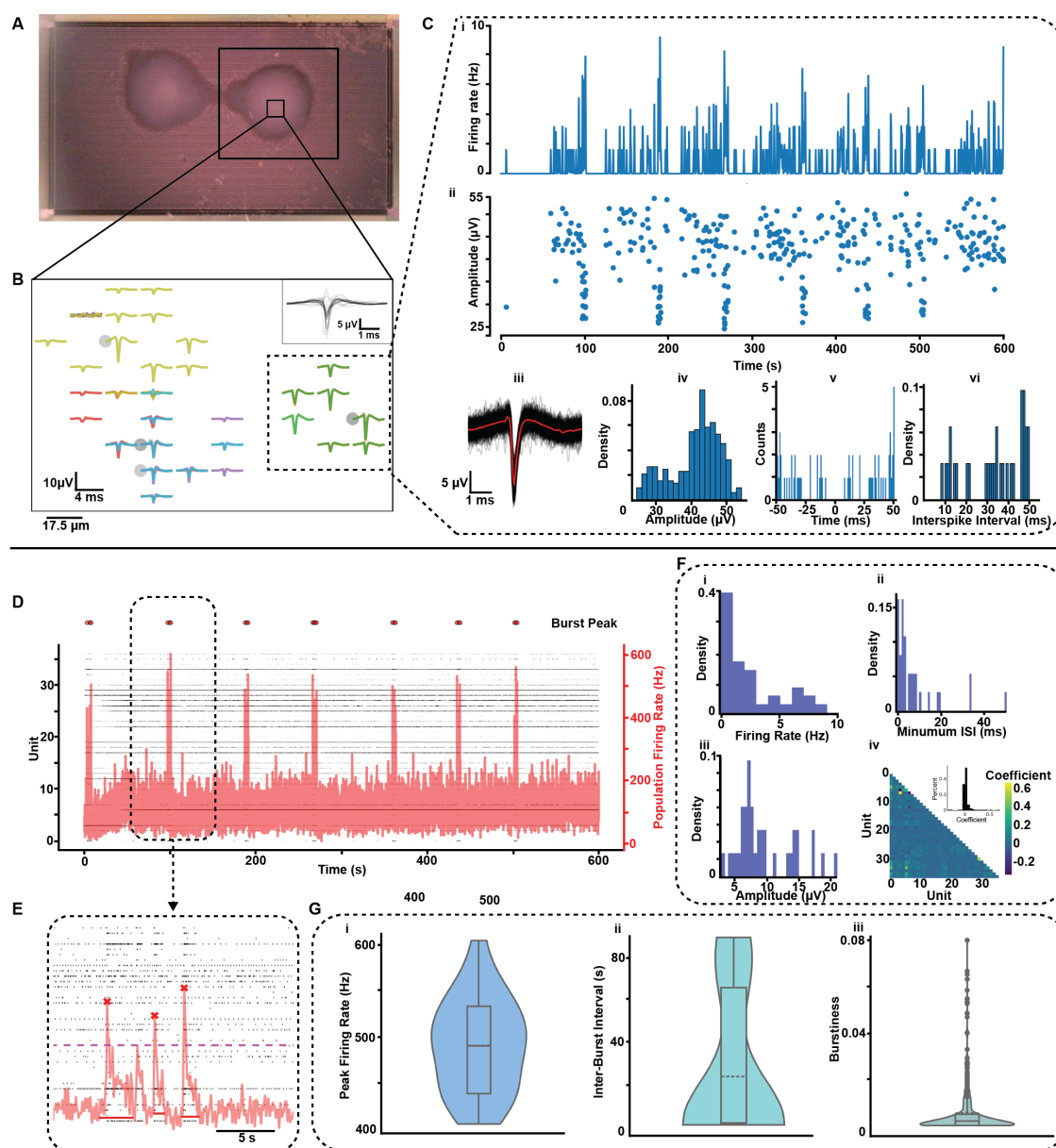
side every single unit's colored footprint (Figure 4B) we provide descriptive electrophysiology features (Figure 4C). We present the unit's temporal firing rate using 50 ms binning of the spike times over the course of the recording (Figure 4C-i). The result is smoothed by a Gaussian kernel with a sigma of 5. We also provide the amplitude of each spike and a histogram of the amplitude distribution (Figure 4C-ii,iv). Raw spikes and the averaged waveform are also displayed (Figure 4C-iii). Both the amplitudes and raw spikes are from the best channel which recorded the highest mean amplitude of the unit. Interspike interval (ISI) is a crucial feature for neurons, as it is associated with firing patterns and cell types<sup>23,24,58,59</sup>. We show this information through an auto-correlogram in the range of -50 to 50 ms and a histogram of ISI values in the range of 0 to 50ms (Figure 4C-v,C-vi).

In addition to the footprint map, the interactive HTML overview figure includes a spike raster and several statistical plots for population features for the organoid. The spike raster shows each unit's spike times and the population firing rate with labeled burst peaks (Figure 4D,E). Bursts are detected by thresholding the population firing rate. We show burst features such as the distributions of peak firing rate, interburst interval, and each unit's burstiness index in violin plots (Figure 4G). Furthermore, we display the distribution of firing rates, minimum ISI values, and mean spike amplitudes for all single units in the recording (Figure 4F-i,ii,iii). We also illustrate the pairwise correlation of units' firing activity by calculating the Spike Time Tiling Coefficient (STTC)<sup>60</sup> value of each unit relative to the others. We designed the overview figures to be interactive, allowing users to zoom in for a closer examination of the data. The figures for individual units are high-resolution. These figures can give users useful information to evaluate the recording object and perform cross-comparisons. Detailed descriptions of data visualization can be found in the Methods section. The complete figures are available in the Supplementary Figure S4, S5.

## Longitudinal Organoid Electrophysiology Properties

Longitudinal neuronal recordings provide invaluable data to study how neuron activity patterns change over time. The cortical organoid shown in Figure 4A was subjected to hourly ten minute recordings on the HD-MEA over seven days (see Voitiuk et al., 2024<sup>50</sup>). During this experiment, recordings were automatically scheduled at the beginning of each hour, uploaded to S3, and processed by the pipeline. Data processing included spike sorting using Kilosort2 and autocuration with quality metrics. Detailed descriptions of the data processing can be found in the Methods section.

Over time, we observed an increasing number of single units and intensified spiking activity. Figure 5A illustrates the time-lapse images of the units' locations and their action potential amplitudes on the HD-MEA. With a grayscale color bar, the darker color denotes a higher amplitude. The scale ranges from 0μV (white) to 30μV (black). There is a noticeable increase in the activity intensity and clustering of active areas as days progress, especially prominent between days 3 to 5. To visualize the development of the organoid neuronal network we plotted the number of detectable units in each recording (Figure 5C,D) and the individual unit firing rate (Figure 5E,F) over the recording time course. Figure 5C shows the distribution of the unit count across recordings for each day, while figure 5D shows the average number of units each day. There is a substantial growth in the number of units from day 0 to day 2 and decreased variability among the recordings. From day 2 to day 7, the number of units is relatively stable, with an increased variability across the samples. There is a clear upward trend for the firing rate for the individual neural units from each recording (Figure 5E) and the average firing rate for each day (Figure 5F). As the days progress, the firing rate distribution of individual units becomes wider with some units showing higher firing rates while other units have a firing rate between 0 and 10 Hz.



**Figure 4: Pipeline Output for an Electrophysiology Recording.** (A) Photograph of a mouse organoid on HD-MEA. (B) Zoomed-in view of spiking activities in the mouse organoid. Each color represents a single unit. Waveforms from all single units are shown in the top right corner. (C) Spiking features for the single unit labeled in B: i) Firing rate distribution over the recording time, calculated by binning spike train with a 50 ms time window. ii) Amplitude of each spike over the recording time. iii) Raw spike waveforms (black) and the averaged waveform (red). iv) Amplitude distribution. v) Auto-correlogram from -50 ms to 50 ms. vi) Interspike interval distribution for intervals in the 0 - 50 ms range. (D) Spike raster (black) overlaid with population firing rate (red) for the recording. Dots above the plot label population burst peaks. (E) Zoomed-in view of a population burst. (F) Distribution of i) unit firing rates, ii) minimum ISIs, iii) mean amplitudes, and iv) spike time tiling coefficients. (G) Violin plots showing the distribution of i) population burst peak firing rates, ii) interburst intervals, and iii) burstiness of each single unit.

We also found changes in neuron firing patterns over time. The neural unit firing patterns are represented by the coefficient of variation (CV) of interspike intervals (ISI)<sup>61,62</sup>. We show the evolution of CV by plotting the standard deviation of ISI to the mean of ISI for each unit over the 7 days. The stacked bar charts represent the proportion of neurons with different CV values, where the red portion indicates neurons with  $CV < 1$ , and the blue portion represents neurons with  $CV \geq 1$  (Figure 5B). Over the 7 days, we observe a trend of an increasing number of units showing a more regular pattern with  $CV < 1$ , implying the maturity of the neural network. Day 0 to day 2 starts with a fairly even split, with slightly more neurons having  $CV \geq 1$  (44%, 54%, and 53%) and  $CV < 1$  (56%, 46%, and 47%). As time progresses, there is a clear shift towards neurons with  $CV < 1$ . By day 6, the majority of neurons have  $CV < 1$  (72%), and a small proportion of neurons have  $CV \geq 1$  (28%).

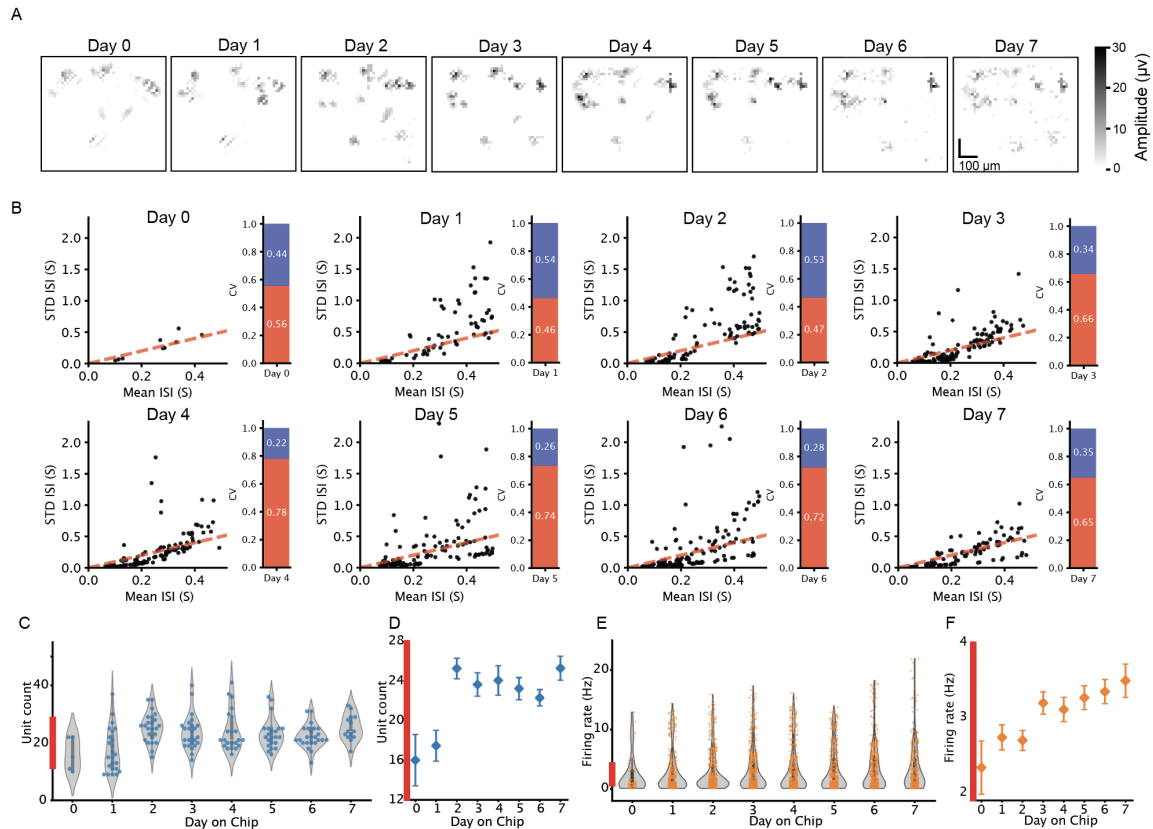
Overall, this analysis suggests maturation of the mouse cortical organoid neuronal activity over the 7 days with increases in both the number of units detected and their firing rate. The increased firing rate variability could indicate the emergence of more complex and heterogeneous neural circuits within the organoid.

## Neuron Tracking for Longitudinal Recordings

The consistency of the pipeline enables tracking putative neurons throughout the longitudinal experiment, as the same processing steps and parameters are applied to all datasets. A trackable unit can be identified by its consistent spike waveform and location on the HD-MEA. After spike sorting a recording, we gathered the average waveform (2.5ms), the best channel's location (x, y coordinate on the HD-MEA), footprint, and firing rate for each single unit. We used a waveform clustering algorithm (WaveMap)<sup>424</sup> to label the units and observed the change of electrophysiological features across multiple days. We ran WaveMap using both the waveform and the best channel's location. The best channel is defined as the one that recorded the unit's highest mean amplitude. For each unit, we concatenated the best channel's location to the end of the waveform. Then, we aggregated units from all recordings. The waveforms and the locations were normalized separately. As a result, WaveMap yielded 20 distinct clusters for the mouse organoid, as shown in Figure 6A. For each cluster, we characterized the waveform features by measuring the trough-to-peak width and Full Width at Half Maximum (FWHM) of the amplitude. The violin plots (Figure 6C) show significant differences in the waveform features among clusters, indicating potentially different cell types in the organoid. Details of running the algorithm can be found in Methods: Waveform Clustering for Cell Tracking.

For a trackable unit at a static location on the organoid, the unit's waveforms sampled across recordings should be in the same cluster and appear on adjacent recording channels. Using HD-MEA, we can locate a unit within a small area with an electrode pitch of tens of micrometers (17.5 $\mu$ m for MaxOne HD-MEA). We labeled each footprint by the color of the corresponding waveform cluster and observed the duration and change of its best channel throughout the recordings. For each cluster, we summarized the best channels for each recording and the frequency of each channel (Supplementary Figure S7) that shows activity.

Among these clusters, we selected Cluster 4 as our primary focus (Figure 6B, D). Figure 6B shows this cluster on the UMAP and the waveforms across recordings. We observed the channel locations of the units in this cluster and arranged the footprints from the three adjacent channels that showed the most activity. These activities are highly likely to be from an individual neuron. We labeled the best channels as L1, L2, and L3 and overlaid corresponding footprints for each channel (Figure 6D,E). On an HD-MEA, the electrical signal from a neuron can be picked up by nearby electrodes, which can be beneficial in identifying a neuron's orientation and movement.



**Figure 5: Single neuron features from hourly recordings over days.** (A) Spatial area of spiking activity in the mouse organoid on the HD-MEA over the recording time course. Color intensity corresponds to the amplitude of the neuron's action potential. (B) Changes in the Coefficient of Variation (CV) of interspike interval distribution over time. The bar plot shows the percentage of units with CV < 1 (red) and CV  $\geq$  1 (blue). (C) Distribution of the total number of single units for each day. (D) Average unit count with standard error of the mean (SEM) over time (Day 0:  $16 \pm 2.58$ , Day 1:  $17.45 \pm 1.55$ , Day 2:  $25.25 \pm 1.03$ , Day 3:  $23.64 \pm 1.18$ , Day 4:  $24.04 \pm 1.49$ , Day 5:  $23.22 \pm 1.12$ , Day 6:  $22.29 \pm 0.81$ , Day 7:  $25.28 \pm 1.20$ ). (E) Single unit firing rate distribution over the 7 days. (F) Average firing rate (Hz) with SEM over time (Day 0:  $2.33 \pm 0.35$ , Day 1:  $2.74 \pm 0.16$ , Day 2:  $2.7 \pm 0.13$ , Day 3:  $3.19 \pm 0.14$ , Day 4:  $3.11 \pm 0.16$ , Day 5:  $3.27 \pm 0.15$ , Day 6:  $3.35 \pm 0.16$ , Day 7:  $3.49 \pm 0.22$ )

During the experiment, this unit initially showed activity on L3. Then its signals were sampled mostly between the two main locations L1 and L2 (Figure 6F). We calculated the firing rate for each sample across recordings and locations (Figure 6F), and grouped the firing rates for each location in Figure 6G. Interestingly, while distributions of firing rates between L1 and L2 did not differ significantly (two-sample Kolmogorov–Smirnov test,  $p=0.11$ ), there was a significant difference between L2 and L3 distributions (two-sample Kolmogorov–Smirnov test,  $p=0.019$ ). This finding suggests that L3 may represent a subset of activity of L2 based on differences in their respective footprints.

Using this study, we show the pipeline provides stable, consistent and reproducible data analysis. The neuron tracking function can improve our understanding of an individual neuron's long-term activity by monitoring its electrophysiological features. Thus, this pipeline offers new possibilities to investigate neural dynamics, plasticity, and neural circuit development.

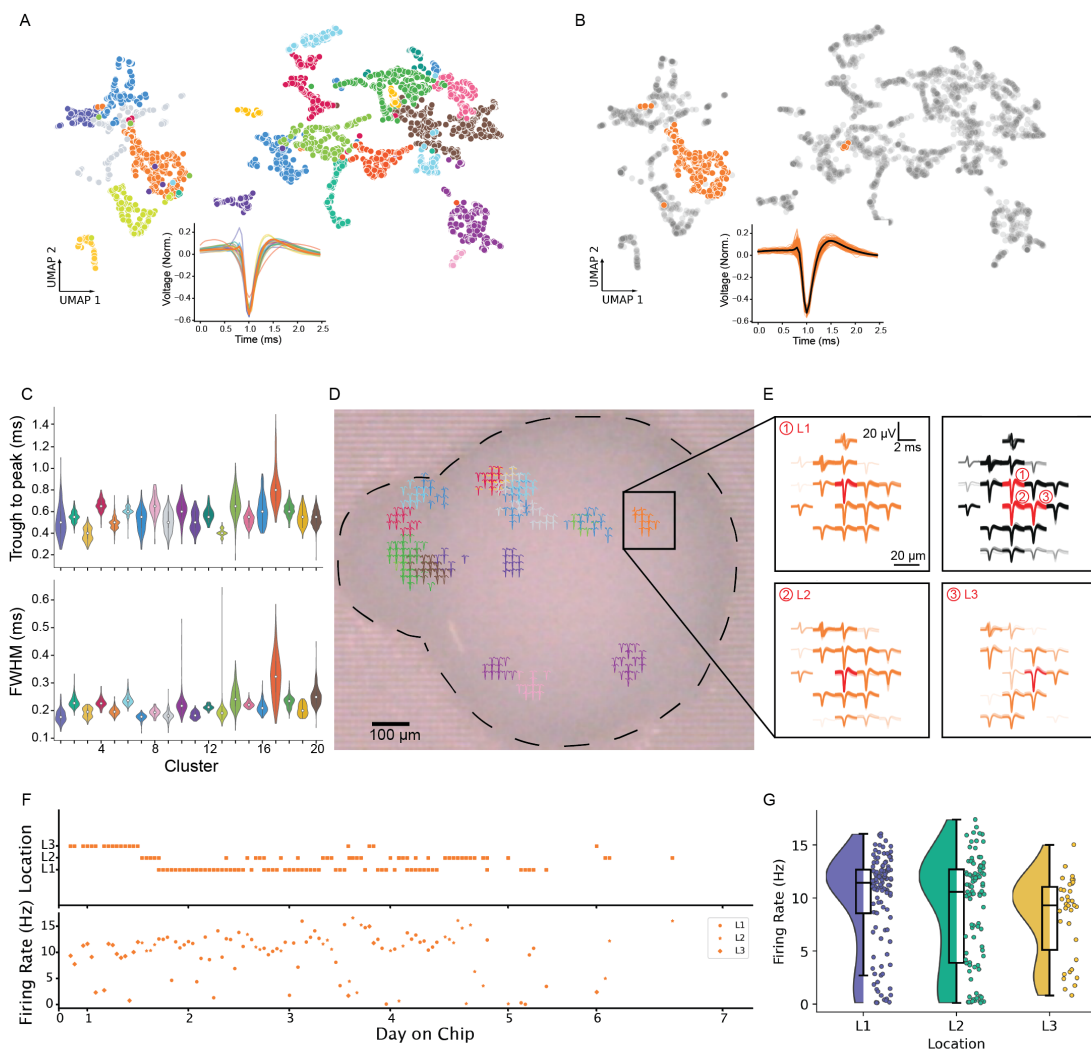
### **Pipeline Applied to Optogenetics Modulation of Epileptiform Activity from Human Hippocampus Slices**

Epilepsy is a neurological disorder characterized by abnormal brain activity resulting from an imbalance between excitatory and inhibitory processes<sup>63</sup>. Light-responsive channelrhodopsins enable optogenetic interventions to modulate the neuronal activity of brain tissues. We applied this pipeline's data processing and analysis functionality to study the optogenetic modulation of neural circuits from human hippocampus slices.

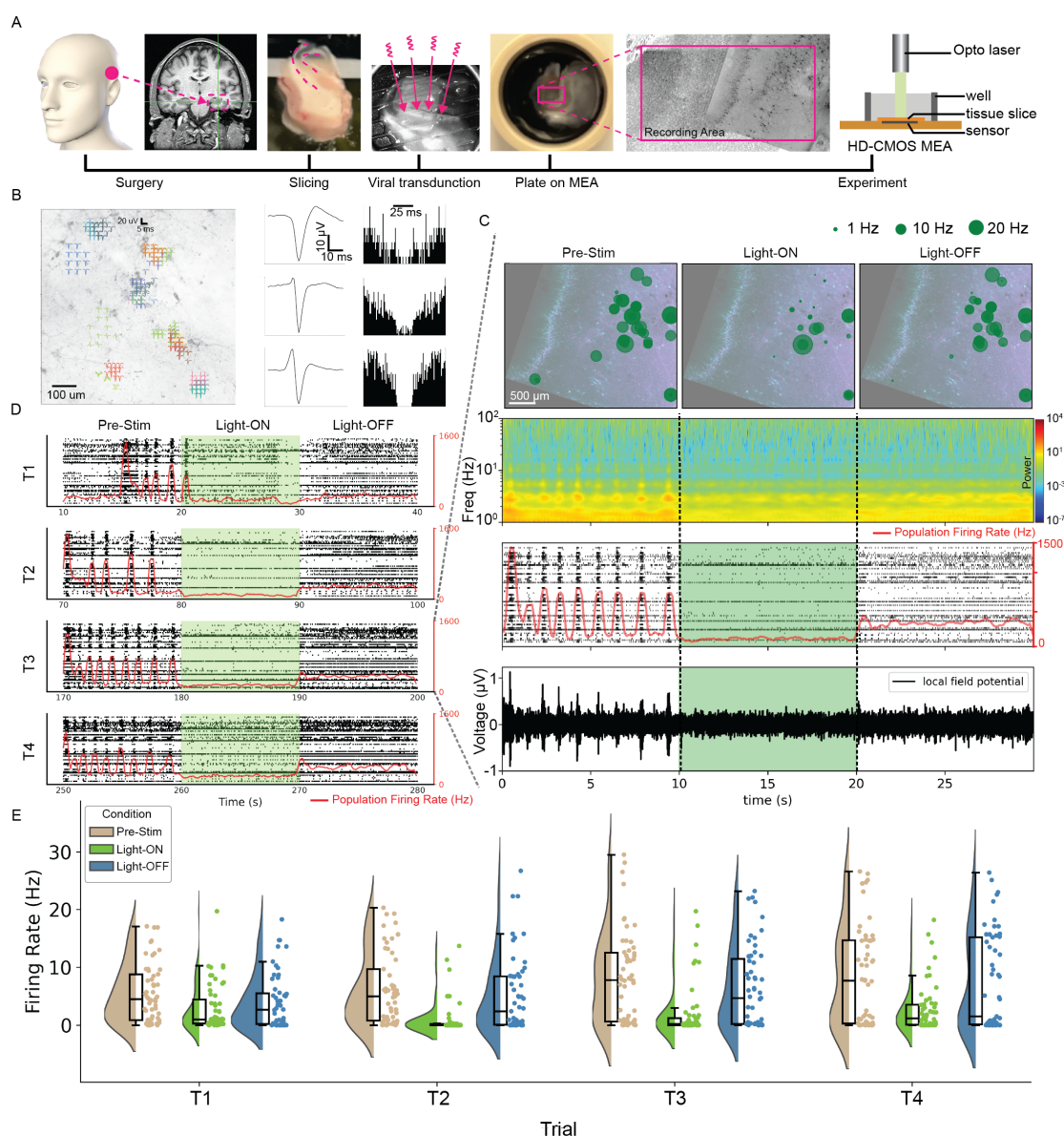
Before optogenetics illumination, human organotypic tissue slices from hippocampus tissue were established. The hippocampus tissue was obtained from patients with drug-refractory temporal lobe epilepsy, sliced to 300 $\mu$ m, and cultured at an air-liquid interface. Slices were transduced with AAV9 carrying an HcKCR1 transgene driven by a CaMKII $\alpha$  promoter and a fluorescent tag (eYFP) (see Andrews et al., 2024<sup>64</sup>). A hippocampus slice was plated on an HD-MEA (MaxOne) for electrophysiology recording, with a fiber-coupled LED to illuminate the slice from the top. Since HcKCR1 encodes a kaliun channelrhodopsin, a light-gated potassium channel that hyperpolarizes the neuronal membrane, the probability of neuronal spiking is reduced when activated by 530nm light. Bicuculline was applied to the slice after plating to increase neuron firing rates, inducing epileptiform activity. During the experiment, we illuminated the slice for 10 seconds at 0.6 light intensity (35.8 mW/mm<sup>2</sup>) of the LED driver, and observed the neuronal population firing activity for 10s prior to illumination (Pre-Stim), 10s during illumination (Light-On) and 10s following the end of illumination (Light-OFF). The experimental setup is shown in Figure 7A. Each HD-MEA recording was processed by the described pipeline using spike sorting and autocuration algorithms. Neuronal activities were aligned to optogenetics stimulation timestamps that were synchronized with the recording. As illustrated in Figure 7B, the units' footprints were overlaid with NeuN staining of the slice on the HD-MEA recording area, showing the physical location of the spiking activity. Examples of spike waveforms and auto-correlograms are shown.

The optogenetics modulation of population firing is shown in Figure 7D and Supplementary Figure S8. The bicuculline-provoked recurrent burst activity was rapidly suppressed by the illumination. Interestingly, the burst activity didn't completely recover when the illumination was off. The firing rate suppression of individual neuronal units was consistent among trials (Figure 7E). This pipeline is capable of providing multiple perspectives of an individual neuron's firing activity (Figure 7C). In addition to firing rate, the suppression of activity is visualized on the hippocampus slice through the electrodes on the HD-MEA. The pipeline can extract local field potential (LFP) data by applying a 5th-order Butterworth bandpass (0.1-100Hz) filter to the raw voltage data. During the 10 seconds "Light-ON" period, activities in the LFP frequency bands were also





**Figure 6: Neuron tracking in a mouse organoid over seven days of recording** Pipeline output demonstrating the capability of neuron tracking for longitudinal recordings. (A) UMAP of waveform clusters with location coordinates. The inset shows the mean waveforms of each cluster from a total of 20 clusters superimposed. (B) The cluster of interest (orange) is highlighted on the UMAP, with other clusters in gray. Inset displays the individual waveforms from this cluster, with the mean waveform in black obtained by averaging all the waveforms. (C) Distribution of waveform features for each cluster. The features include trough-to-peak width and Full Width Half Maximum (FWHM) of the amplitude. (D) Footprint projection on the organoid recorded on the MEA. The footprint is color-coded according to the UMAP cluster. (E) Footprints for the cluster of interest overlaid across the recording time course. L1, L2 and L3 are the best channels on the footprints. (F) Temporal tracking of location and firing rate change for the units in the orange cluster. (G) Firing rate distribution for the three locations.



**Figure 7: Pipeline facilitates seizure study by analyzing electrophysiology data from human hippocampus brain slice with optogenetic stimulation.** (A) Experimental setup for hippocampus slice HD-MEA recording<sup>[64]</sup>. Brain tissue from a seizure patient in 300 $\mu$ m thick slices cultured expressed channelrhodopsin delivered through adeno-associated virus (AAV) delivery. The slice is placed on HD-MEA for simultaneous optogenetic stimulation and recording detailed in Andrews et al., 2024<sup>[64]</sup> (B) NeuN-stained hippocampus slice overlaid with an image of the slice on HD-MEA and footprint of spiking activities on the slice. Example spike waveforms and auto-correlograms from representative neurons. (C) Hippocampus slice overlaid with single units' firing on the HD-MEA. The change in firing activity is shown for the three steps of Trial 3 (T3). From top to bottom, the panels display the single unit's location overlaid with GFP-stained slice, firing rate, local field potential spectrum, spike raster, and voltage data from all recording channels. (D) Spike raster with population firing rate for four experimental trials under Pre-Stim, Light-On, and Light-Off conditions. The population firing rate shows epileptiform activity suppressed by optogenetics illumination, with the firing rate remaining low afterwards. (E) The single unit's firing rate distribution for each Trial in (D).

attenuated.

This application showcases the pipeline's adaptability to diverse experimental paradigms, extending its utility beyond basic neural activity analysis to more complex neurological disease studies. This advancement opens new avenues for studying neurological disease mechanisms and potentially developing therapeutic approaches, highlighting the pipeline's significance in translational neuroscience research.

## DISCUSSION

The cloud-based electrophysiology data pipeline presented in this study represents an advancement in the processing and analysis of HD-MEA recordings, which are enabled by IoT and cloud computing technology. The flexible and modular architecture can meet different data processing goals, enabling high data quality and comprehensive electrophysiology feature extraction. The integration of the MQTT messaging protocol provides remote access to the pipeline as well as communications between various components of the pipeline. The cloud-based infrastructure addresses the challenge of storing and processing large volumes of long-term, high-throughput experiments. The parallelized processing capabilities allow for rapid analysis of multiple datasets simultaneously. In addition, the ability to process recordings consistently and without human intervention saves time and reduces the potential for human error and bias in data analysis.

The user interface allows easy access to the pipeline, and open source makes the pipeline adaptable to different computing environments and infrastructure setups. Cloud data storage and computing contribute to the scalability of the pipeline. The pipeline output data structure is straightforward and size efficient, making it easy for computational tasks.

### Consistent and Reliable Data Processing

By using the same parameter settings for spike sorting and curation across all recordings, we ensure that data is processed uniformly without human intervention. This consistency is crucial for longitudinal studies, where tracking changes in neural activity over time requires a stable and reproducible processing framework. The use of Kilosort2 for spike sorting, combined with autocuration algorithms, allows us to accurately identify and classify single-unit activity, even in complex datasets with overlapping spikes. Recent studies have highlighted the importance of such consistent processing in large-scale electrophysiology data analysis.

One of the most critical aspects of our pipeline is its ability to process data consistently and reliably. By using the same parameter settings for spike sorting and curation across all recordings, we ensure that data is processed uniformly without human intervention. This consistency is crucial for longitudinal studies, where tracking changes in neural activity over time requires a stable and reproducible processing framework. The use of Kilosort2 for spike sorting, combined with auto-curation algorithms, allows us to accurately identify and classify single-unit activity, even in complex datasets with overlapping spikes. Recent studies have highlighted the importance of such consistent processing in large-scale electrophysiology data analysis<sup>30,65</sup>.

### Data Management and Visualization

Our pipeline's data management capabilities are enhanced by the use of a hierarchical structure with strategically named buckets on AWS S3. This structure, combined with metadata files that store detailed experiment-related information, ensures that data is organized efficiently and can be accessed quickly. The integration of user interfaces, such as the data uploader and Dashboard, further empowers researchers by providing tools for data management, algorithm parameter configuration, and result visualization. The Dashboard, built using the Plotly Dash library,

offers interactive features that allow users to explore and analyze their data in depth. Similar approaches have been successfully implemented in recent neuroscience data management and visualization systems<sup>66</sup>.

### Code Availability

This electrophysiology data pipeline is an open-source project. The code will be released on GitHub upon manuscript publication and is currently available upon request.

### Data Availability

No new data was generated for this paper. All datasets described were obtained from Voitiuk et al., 2024<sup>50</sup>, Andrews et al., 2024<sup>64</sup>.

## METHODS AND MATERIALS

### Mouse Cortical Organoids

The data presented in this manuscript was collected using an integrated system for neuronal cell culture<sup>50</sup>. Mouse cortical organoids were made using a protocol described in Park et al., 2024<sup>67</sup>. Cortical organoid recordings were performed on a MaxWell Biosystems MaxOne CMOS HD-MEA chip. The system captured 10-minute recordings every hour for seven days. The recording configuration remained consistent throughout the experiment. Details can be found in Voitiuk et al., 2024<sup>50</sup>.

### User Interfaces, Data uploader, and Dashboard

To give researchers full control over their experimental data, we designed user interfaces that enable data management, data processing, algorithm parameter configuration, and result visualization. We developed the data uploader installed on a local computer of the recording device as well as an online Dashboard for remote data access.

The data uploader, created using the Python PyQt library, facilitates the uploading of experiment recordings from a local laptop to S3. Upon opening this application, users can navigate to a folder where recordings are stored. An initial Universally Unique Identifier (UUID) is generated using the date of the recordings, and users can add more descriptive labels to this UUID. Before uploading, users must generate a metadata file by loading a template and inputting any experiment-related information such as notes, cell lines, media used for culture, and recording features for each dataset. Recording features such as recording length, number of active channels, and data path are automatically populated in the metadata template. When users press the upload button, all recordings in the selected local folder are reorganized according to the S3 file structure and are uploaded to the UUID folder on S3. Users can start the data analysis pipeline after uploading by sending a request a message to the job listener.

The Dashboard was created using the Python Plotly Dash library. This library uses callback functions to achieve user-interactive features like dropdown lists, buttons, and tables. We built a multi-page website, with each page serving a different purpose.

On the “Data Processing Center” page, users can choose recordings from the S3 dropdown list, select preferred data processing jobs, set parameters, and start NRP jobs. It allows users to perform batch processing or chained tasks for chosen recordings. Batch processing is the most commonly used case since all parameters and algorithms are usually the same for an experiment. Chained tasks are practical for testing parameter and algorithm combinations for new experiment setups. Supplementary Figure S3 shows the job center webpage.



On the “Status” page, users can monitor the job status of current tasks on the NRP cluster. By using the “Show Status” button, users can check jobs labeled with prefix “edp-”. This function parses the information returned by Kubernetes Python API for all jobs in the namespace. Upon refreshing with the button, the NRP job name, running status, and data summary will be displayed on the webpage.

The “Visualization” page is designed to display figures of post-processed results. Users can select a processed recording from the dropdown menu to display an interactive raster plot and electrode map. Clicking a unit on the electrode map highlights its spike times in light red on the raster plot and shows its waveform and interspike interval histogram. This page allows users to evaluate MEA recordings effectively.

## **Cloud Data Storage and Organization**

Efficient cloud data organization is crucial for optimizing access performance and storage management. In this pipeline, we employ a hierarchical structure with strategically named buckets. We use a UUID that reflects the experiment date and key information. Upon data uploading, the metadata file is automatically generated and uploaded with the raw data. For each UUID, we keep the raw data in “/original/data”, and the processed result files in “/derived/algo”, where “algo” can be “kilosort2”, “connectivity” and others that are named after the specific algorithms. We store logging files on S3 for MQTT services and data analysis jobs. Detailed logs provide a comprehensive record of each pipeline component by capturing essential information. These logs enable researchers to track the progression of data processing, identify potential bottlenecks, and troubleshoot issues effectively. Service logs are generated when the MQTT broker sends or receives a message and updated to the S3 “/service” bucket on a schedule. Logs from the data analysis jobs include processing steps, quantities, and malfunctions. These log files are kept in the algorithm output directory.

## **Cloud Orchestration**

Each analysis algorithm is packaged into a Docker container with the minimum required dependencies, enabling parallel processing of a large volume of electrophysiology recordings in a cloud-agnostic environment. This approach simplifies the addition of new analysis algorithms to the pipeline.

We use Kubernetes to deploy and monitor data processing jobs on NRP. For each container, based on the input data size and algorithm requirements, we request computing resources from NRP, such as the number of CPUs, GPUs, memory, and storage. Supplementary Table S2 summarizes the computing resource requirements for each algorithm on a 10-minute HD-MEA recording with 1000 active channels. When a job is deployed to NRP, a pod with a job is created to run the data in a container. To get the status of the pod, we extract metadata from the return of the Kubernetes “list\_namespace\_pod” function. From the metadata, we provide the status of the job, such as “running” or “succeeded,” and the timestamps for running this job.

## **MQTT Messaging Application**

To enable remote job execution for a large number of recordings, we implemented services using MQTT messaging. This infrastructure has been previously described<sup>47,50</sup>. All messaging services are hosted on the Braingeneers UCSC Genomics Institute server. We package these services into Docker containers and manage them using Docker Compose.

## **Job Listener**

We designed a centralized MQTT service to parse analysis job run messages. This service subscribes to specific topics and responds by running the corresponding Docker container on the



502 given data. We assign the topics “experiments/upload” for batch processing or “services/csv\_job”  
503 for chained tasks.

504 The message body is designed according to the different topics. For “experiments/upload”, we  
505 use the UUID and recording file name from the metadata.json file. The service can assemble the  
506 S3 file path for each recording from this information. The computing requirements for running  
507 batch processing jobs are written to a look-up table in the listener service.

508 For “services/csv\_job”, we first create a CSV file where each row contains the UUID, recording  
509 file name, job type, and computing requirements for running the analysis. We name the CSV  
510 file using the current timestamp, upload it to the S3 services bucket, and create a message con-  
511 taining the path to the CSV file and the indices of the CSV rows. When the listener receives this  
512 message, it pulls the information from the CSV file and deploys jobs using the given indices. Ex-  
513 amples of the messages are included in Supplementary Materials.

#### 514 **Job Scanner**

515 When running analysis jobs on the NRP, we use the prefix label “edp-” in the job name. We name  
516 batch processing and chained jobs differently. For batch processing, we name the job using a  
517 prefix and the recording file name. For chained jobs, we name the job using a prefix, the CSV file  
518 name, and the index of the CSV row. This naming convention allows us to parse the job name to  
519 determine which analysis algorithm is running on which data.

520 The job scanner is designed to scan the “edp-” jobs on a schedule with two main aims. First, it  
521 notifies the job listener when the current step in a chained job is finished. This message body  
522 contains the keyword “update”. When the listener receives this message, it checks the corre-  
523 sponding CSV file to launch any pending jobs related to the current job. The scanner scans NRP  
524 every 2 minutes to minimize delays in running chained jobs. Second, it notifies users of their job  
525 status via a Slack channel using the messaging bridge service 18. These notifications are sent  
526 every 30 minutes.

527 Job information is pulled from NRP using Python-Kubernetes functions. We use the “list\_namespaced\_pod”  
528 function to get all “edp-” jobs. We loop through them, extracting job name, data file name, job  
529 type, and timestamps. This information is stored in a Python dictionary and updated when the  
530 scanner service scans NRP on schedule.

531 The scanner identifies job status and sends messages to other MQTT services. For batch pro-  
532 cessing jobs, the scanner sends a user notification message when the job status is “pending”,  
533 “running”, “failed”, or “succeeded”. Since “failed” and “succeeded” jobs are finished, the scanner  
534 removes these jobs from NRP and the dictionary after sending the message. For chained jobs,  
535 when a job is finished as “succeeded”, in addition to sending a user notification, the scanner also  
536 sends an “update” message to the listener to run the next job.

#### 537 **User Notification to a Slack Channel**

538 Both the job listener and scanner can send user notifications. When a run job message is sent to  
539 the listener, and the listener successfully deploys jobs to NRP, a notification is sent to the Slack  
540 channel with the S3 data path, job type, and “start” status.

541 To make user notifications human-readable and clear, when the scanner sends messages to the  
542 Slack channel, it groups the jobs by UUID and lists the recordings in each UUID.

#### 543 **Spike Sorting**

544 Spike sorting is fundamental in analyzing extracellular recordings for assigning action potentials  
545 picked up by electrode channels to neurons in an ensemble. For the HD-MEA recordings, Kilo-  
546 sort2 was used to sort the raw voltage data into single unit activity. Since HD-MEAs can record

one neuron from tens of channels, it is common for spikes from many neurons to overlap in time on a single channel. The template matching and clustering algorithm in Kilosort2 can distinguish spikes between different neurons based on their waveform. Before spike sorting, the raw data is bandpass filtered using an IIR filter with a 300 - 6000 Hz bandwidth. The data type is converted to int16 for running Kilosort2. The voltage detection threshold of Kilosort2 is set to 6 RMS over the baseline. Parameter settings for Kilosort2 are shown in Supplementary Table S1. Spike sorting was performed on the NationalResearch Platform computing cluster with an NVIDIA A10 GPU. The sorting output is saved to a compressed file (zip format) and uploaded to S3. The output file structure is compatible with the software Phy for performing manual curation. An autocuration process is built on top of the sorting result.

## Autocuration

The autocuration process is applied after spike sorting for each single unit. To assess data quality and retain good units for downstream analysis, we evaluate each unit by calculating the Inter-spike Interval (ISI) violation ratio, Signal-to-Noise ratio (SNR), firing rate, and spike waveform. We use the curation module from SpikeInterface API<sup>[30]</sup> in our Python script. For ISI violation, we apply the Hill method<sup>[68]</sup> of false positive errors with an absolute refractory period of 1.5 ms. We set the maximum violation rate to 20%. The SNR is calculated using the spike amplitude of a unit and the baseline voltage, with a minimum SNR threshold of 5 RMS. The unit's firing rate is defined as the total number of firing events divided by the recording length in seconds. In our default autocuration algorithm, this threshold is set to 0.1 Hz.

To check the spiking waveform for a unit, we run the WaveformExtractor class across all active channels and take the average of a maximum of 500 spikes. We then find the best channel and a maximum of eight neighboring locations on a 3x3 grid by sorting their waveform amplitudes on each channel from highest to lowest. The best channel is defined as the channel that captures the neuron's highest mean amplitude among all recording channels. Since HD-MEAs can record one neuron across multiple channels simultaneously, we expect the waveform distribution to have an adequate layout. This layout is defined as the unit's footprint. Thus, we save units that show a waveform on the best channel and at least one neighboring channel within a distance of 17.5 $\mu$ m for further analysis.

## Visualization of Electrophysiology Features

For each recording, the pipeline generates an interactive overview figure in HTML format that includes the activity map of the MEA, the neuron's footprint at its physical location on the map, a spike raster with population burst detection<sup>[7]</sup>, and a summary of electrophysiology features for all single units. The population firing rate is smoothed using moving average (20 ms window size) applied to aggregated spike trains, then further smoothed using a Gaussian kernel with sigma = 20. The population burst detection threshold is set to 2 RMS of the population's baseline firing rate. Burst detection is performed using `scipy.signal.find_peaks` with a minimum peak distance of 800 ms. Burst edges are defined as points where the firing rate drops by 90% from the peak on both sides. The "burstiness index"<sup>[69]</sup> of a single unit is represented by a number from 0 to 1 that measures the synchronization in spiking activity by binning (40 ms bin size) the spike train. Electrophysiology features include interspike interval (ISI), minimum ISI, firing rate, amplitude, spike time tiling coefficient (STTC)<sup>[60]</sup> and average spike waveforms. Distributions of these features for all single units are provided in the interactive figure. Each single unit is also paired with a PNG format figure showing the unit's footprint, raw and average spike waveform, auto-correlogram (ACG), ISI distribution, instantaneous firing rate and amplitudes. Both the interactive figure and single unit figures are created using the Plotly Python graphing library. Parameters for the visual-

ization are adjustable in the source code.

## Waveform Clustering for Cell Tracking

Neuronal cell types and their spiking waveforms are known to be correlated. To demonstrate the capability of tracking units in longitudinal recordings, we performed waveform clustering using the WaveMAP Python package<sup>424</sup>. This package combines non-linear dimensionality reduction (UMAP) with the Louvain clustering method.

To prepare the waveforms, we first extracted the spike times for each single unit through spike sorting. Since the spike time represents the peak of each spike, we initially took a 5 ms window of the complete waveform from the best channel, then averaged across up to 500 spikes. Before input into the WaveMAP algorithm, we centered the waveforms at their peak and truncated them to 1 ms before and 1.5 ms after the peak. Units with positive spikes were not included in this clustering due to the high possibility of axonal signals. We extracted waveforms for each recording, stacked them into a NumPy array, and pre-processed them using l2 normalization. The total number of waveforms was 3526 from 160 recordings.

Given that the mouse organoid recordings were taken hourly across seven days, and neurons can migrate during development, we appended the corresponding electrode location (x, y) to the end of each waveform for clustering. The location was normalized as a percentage of the maximum x and y coordinates, respectively, to ensure the data range was within [0, 1], comparable to the normalized waveform.

The UMAP parameters were set with 20 neighbors and a minimum distance of 0.1, while the Louvain clustering resolution was set to 1.5. As a result, the algorithm identified 20 distinct clusters and assigned a color to each. Based on the clustering results, we plotted the footprint of each unit on the electrode map using the assigned color. Throughout the recordings, we were able to track changes in a neuron's location and firing rate.

## Local Field Potential

Local field potentials (LFPs) are low frequency signals up to about 500 Hz that are generated by multiple signal processes in a neural population<sup>70</sup>. These signals are traditionally decomposed into frequency domain. In this pipeline, we focused on LFPs in the range of 0.1 to 100 Hz, and subband frequencies as delta (0.5 - 4 Hz), theta (4 - 8 Hz), alpha (8 - 13 Hz), beta (13 - 30 Hz) and gamma (30 - 50 Hz).

To get LFPs and subband frequencies, we first bandpass filter the raw voltage signal from all recording channels with 0.1-100 Hz 5-order Butterworth filter. Then, these signals are downsampled 1 kHz. A second bandpass filter is applied to separate subband frequencies. We use spectrograms to show the signal strength of different subbands. A spectrogram is the time-frequency spectrum of the local field potential signal, based on the power values, over the given time and frequency range. We applied a continuous wavelet transform (CWT) on the local field potentials to obtain wavelet coefficients and corresponding frequencies using the complex Morlet wavelet ('cmor1-1') in PyCWT library. Signal strength is computed as the magnitude squared of the wavelet coefficients and smoothed using a Gaussian filter with sigma of 2.

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## AUTHOR CONTRIBUTIONS

J.G., K.V., D.F.P., and A.R. conceived the project and established the cloud storage organization, cloud computing, and IoT codebase. J.G. designed the pipeline architecture, built the pipeline components, contributed to data collection, analyzed and interpreted the data, created figures and wrote the manuscript. A.R. developed the data uploader and contributed to MEA recordings and spike sorting. A.S. assisted with data analysis code. J.L.S. provided insights into pipeline design, interpreted the data, and contributed to data visualizations. R.C. contributed to the conceptualization of the cloud-based data architecture. S.H. and H.E.S. cultured and provided mouse cortical organoids for MEA recording. K.V. and S.T.S. designed and conducted mouse cortical organoid experiments, provided data, and interpreted the data. E.F.C. provided human tissue samples. J.P.A. obtained human tissue samples, designed and conducted experiments, provided data, interpreted the data, and performed histology. J.P.A., J.G., K.V., A.R., and M.A.T.E. conducted optogenetics experiments, gathered data, and interpreted the results. T.J.N., M.A.M.-R., D.H., T.S., S.R.S., and M.T. provided mentorship, intellectual consultation, input on experimental design and analytic methods, discussed the results. M.A.M.-R., D.H., S.R.S., and M.T. provided funding for the project. All authors commented, edited, and approved the manuscript.

## AUTHOR COMPETING INTERESTS

K.V. and S.T.S. are co-founders and D.H., S.R.S, M.T. are advisory board members of Open Culture Science, Inc., a company that may be affected by the research reported in the enclosed paper. All other authors declare no competing interests.

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