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# Stereotyped large amplitude cortical LFP events can be clustered and reveal precisely ordered phase-locking in neuronal populations

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1 **Title Page**

2 **Abbreviated title:** Clustering Electrical Oscillations in Cortex Reveals Precise Single Neuron Spike Timing

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## 24 **Abstract**

25 During quiet wakefulness, slow-wave sleep and anesthesia, mammalian cortex exhibits a synchronised state  
26 during which transient changes in the local field potential (LFP) accompany periods of increased single  
27 neuron firing, known as UP-states. While UP-state genesis is still debated (Crunelli and Hughes, 2010) such  
28 transitions may constitute the default activity pattern of the entire cortex (Neske, 2016). Recent findings  
29 of preserved firing order between UP-state transitions and stimulus processing in high-firing rate ( $>2\text{Hz}$ )  
30 rat auditory and barrel cortex neurons (Luczak et al., 2015) support this hypothesis. Yet it is unknown  
31 whether UP-states are homogeneous and whether neurons with firing rates  $<2\text{Hz}$  in visual cortex or other  
32 species exhibit spiking order. Using extracellular recordings during anesthetized states in cat visual cortex  
33 and mouse visual, auditory and barrel cortex, we show that UP-states can be tracked and clustered based  
34 on the shape of the LFP waveform. We show that LFP event clusters (LECs) have current-source-density  
35 profiles that are common across different recordings or animals and using simultaneous electrophysiology  
36 and widefield voltage and calcium imaging in mouse we confirm that LEC transitions are cortex-wide  
37 phenomena. Individual LEC events can be resolved in time to within 1 – 4 ms and they elicit synchronous  
38 firing of over 75% of recorded neurons with most neurons synchronizing their firing to within  $\pm 5 - 15$  ms  
39 relative LECs. Firing order of different neurons during LEC events was preserved over periods of  $\sim 30$   
40 minutes enabling future studies of UP-state transitions and firing order with near millisecond precision.

## 41 **Significant Statement**

42 During sleep and anesthetic states mammalian cortex undergoes substantial changes from awake active  
43 states. Recent studies show that single neurons in some cortical areas in rats undergo increased spiking  
44 during sleep and anesthetic states (called UP-state transitions) with some neurons firing in an order similar  
45 to awake states. This suggests that sensory processing may be similar across all states and that firing order  
46 is important for stimulus processing. Yet UP-state transitions remain poorly understood and it is unclear  
47 whether firing order is present in other cortical areas or species. Here we describe multiple classes of UP-  
48 state transitions and show most neurons in visual cortex in cats and visual, barrel and auditory cortex in mice  
49 exhibit firing order during such transitions.

## 50 **Introduction**

51 An established finding of the past several decades is that the cortical neurons of mammals spike differentially  
52 in two different states. One of these is a desynchronized state which is present during awake and attending  
53 periods and during rapid-eye-movement sleep (REM). During this state neurons fire largely independently  
54 of each other ([Harris and Thiele, 2011](#)). The other is a synchronized state, present during slow-wave sleep  
55 (SWS), quiet waking and anesthesia, where individual neurons cycle, at rates of 0.2 Hz – 0.9 Hz, between a  
56 depolarized (spiking) state and a hyperpolarized (non-spiking) state. These are known as UP- and DOWN-  
57 states, respectively ([Steriade et al., 1993a](#); [Sanchez-Vives and McCormick, 2000](#); [McCormick and Yuste,](#)  
58 [2006](#); [Neske, 2016](#); [Sanchez-Vives et al., 2017](#)). UP- and DOWN-states are brain-wide phenomena which  
59 engage cortex, thalamus, hippocampus, striatum and cerebellum ([Neske, 2016](#)). They may facilitate flexible  
60 processing of information ([McCormick et al., 2004](#); [McCormick and Yuste, 2006](#); [Haider et al., 2006](#)) and  
61 may mediate changes in functional connectivity during waking states ([Neske, 2016](#)). They may also be  
62 involved in memory replay ([Wilson et al., 1994](#); [Sirota et al., 2003](#); [Sirota and Buzsaki, 2005](#)). While  
63 UP-state transitions can be evoked by sensory or thalamic activation ([Amzica and Steriade, 1998](#); [Steriade,](#)  
64 [2001](#)) transitions also occur spontaneously ([Amzica and Steriade, 1995](#); [Destexhe et al., 1999](#); [Volgushev](#)  
65 [et al., 2006](#)). Overall, slow oscillations and synchronized states might provide a “unifying paradigm for the  
66 study of cortical function” ([Sanchez-Vives et al., 2017](#)).

67 In addition to such broad scale functions the cortical “machinery” engaged by spontaneous UP-state transitions  
68 may be the same as used to represent stimuli during both awake, sleep or anesthetized states ([Amzica and](#)  
69 [Steriade, 1998](#)). A number of extracellular recording studies found that high firing rate (>2Hz) neurons  
70 in rat somatosensory and auditory cortex tend to fire in a similar order during UP-state transition (though  
71 some firing distributions were broad, e.g. 50-100ms full-width-half-max) as well as during the first 100ms  
72 following stimulus onset ([Luczak et al., 2007, 2009](#); [Bermudez-Contreras et al., 2013](#); [Luczak et al., 2015](#)).  
73 Firing order during UP-state transitions may therefore reveal a functional role for order in cortex, and  
74 tracking spiking order for all cells and in other cortical areas might help reveal a coding strategy used by  
75 cortex. While traditional methods for defining UP-state transitions rely on single-cell intracellular recordings  
76 ([Steriade et al., 1993a](#)), these methods can only be used to track UP-states for a few neurons at a time.

77 Additional methods for UP-state detection also have relied on peaks in synchronous firing (Luczak et al.,  
78 2007), but such methods exclude low-firing rate neurons (e.g. neurons firing <2Hz) and sparsely firing  
79 cortical areas (e.g. visual cortex).

80 Here we expand the analysis of UP-states by providing a method for detecting them using only extracellular  
81 recordings and show that most neurons (even those with low firing rates) have a preserved firing order.  
82 We show that the multi-channel local field potential (LFP) generated during UP-state transitions can be  
83 clustered and provides a more temporally precise definition of UP-states. Previous studies have shown  
84 that single-channel LFP events correlate with UP-state transitions (Saleem et al., 2010; Chauvette et al.,  
85 2010) and more recent work in rat hippocampal slices (Reichinnek et al., 2010) and anesthetized macaque  
86 hippocampus (Ramirez-Villegas et al., 2015) have shown that LFP events can have stereotyped shapes. Here  
87 we go further and show that in the synchronized state, large amplitude channel LFP events can be clustered  
88 (termed LFP event clusters - LECs) within a 1 – 4ms temporal precision enabling the accurate measurement  
89 of latencies of simultaneously recorded single units. CSD analyses of LECs revealed characteristic laminar  
90 profiles of sources and sinks for each LEC and potentially 3 groups of such clusters. We additionally used  
91 widefield voltage-sensitive dye (VSD) imaging of mouse cerebral cortex to show that LECs are associated  
92 with cortex-wide activations consistent with previous work (Amzica and Steriade, 1998). Most neurons  
93 synchronized their firing to individual events to within  $\pm 5 - 15$  ms. Consistent with this, the relative firing  
94 order of different units during LFP events was present in cat visual cortex and mouse visual, barrel and  
95 auditory cortex adding to the evidence that cortical neurons are capable of firing with high temporal precision  
96 relative to each other.

## 97 **Materials and Methods**

### 98 **Experimental Design**

99 **Cat Electrophysiological Recordings.** Experimental procedures are described in detail in previous work  
100 ([Swindale and Spacek, 2014](#)) and were carried out in accordance with guidelines established by the Canadian  
101 Council on Animal Care and the Animal Care Committee of the University of British Columbia. Data  
102 analysed here were obtained from 15 electrode penetration sites in 5 adult cats (animal IDs: C1-C5). The  
103 cats were anesthetized either with 0.5-1.5% isoflurane and 70% N<sub>2</sub>O + 30% O<sub>2</sub> (C1, C2 and C3) or with  
104 continuously infused propofol (6 – 9 mg/kg/hr) and fentanyl (4–6  $\mu$ g/kg/hr) (C4, C5). Following craniotomy  
105 surgery, a high-density polytrode was inserted perpendicularly into the cortex until the upper recording sites  
106 were 100 – 200  $\mu$ m below the surface. Polytrodes were either 2-column (C2,C4,C5) or 3-column (C1, C3)  
107 with electrode site spacing of 50-75  $\mu$ m. Voltage signals were analogue bandpass filtered between 0.5 and  
108 6 kHz, sampled at a rate of 25 kHz and digitized with 12-bit resolution (Blanche et al., 2005). A subset  
109 of 10 electrode sites were used to separately record the LFP, band pass filtered between 0.5-200 Hz, were  
110 fed in parallel to separate amplifiers. On the 3-column electrodes the channels were 130  $\mu$ m apart, with the  
111 exception of the bottom 2 channels, which were 65  $\mu$ m (C1) or 97  $\mu$ m (C3) apart. On the 2-column electrodes  
112 the channels were 150  $\mu$ m (C2 and C4) or 195  $\mu$ m (C5) apart, with the lower two channels being 100  $\mu$ m (C2  
113 and C4) or 195  $\mu$ m (C5) apart. Recording sites were in area 17 and receptive fields (not reported here) were  
114 typically within 10 degrees of the *area centralis*. In addition to recordings of spontaneous activity, visual  
115 stimuli, including moving bars, gratings, m-sequence stimuli and natural scene movies were presented on a  
116 CRT screen. Table 1 summarizes recording IDs, anesthetic methods and recording duration for individual  
117 experiments in cats.

118 **Mouse Electrophysiological Recordings.** Experimental protocols were established and carried out in  
119 accordance with guidelines established by the Canadian Council on Animal Care and the Animal Care  
120 Committee of the University of British Columbia. Data reported here were obtained from a total of 4  
121 electrode tracks in 4 mice (C57/BL6) anesthetized with isoflurane (1.5–2%) for surgery and with subsequent  
122 recording periods under reduced concentration of isoflurane (1.0–1.2%). The skull was fixed to a head-plate  
123 to stabilize recording and facilitate imaging in simultaneous acquisition sessions. Extracellular recordings



124 were made with 64-channel polytrodes (NeuroNexus A1x64-Poly2-6mm-23s-160-A64) with a 2-column  
125 (32 channels per column) staggered-format with vertical and horizontal (inter-column-distance) of 46  $\mu\text{m}$   
126 covering 1450  $\mu\text{m}$  of the probe. Voltage signals were acquired using a headstage amplifier (RHD2164,  
127 IntanTech, Los Angeles) and USB interface board (RHD2000, Intan) at a sampling rate of 25 kHz - 16 bit  
128 resolution. Electrodes were inserted perpendicular to the surface of the cortex using a micro-manipulator  
129 (MP-225, Sutter Instrument Company). Cortical penetration depth was tracked using micro-manipulator  
130 coordinates with the tip of the electrode being inserted between 900  $\mu\text{m}$  to 1450  $\mu\text{m}$  (mean of 1256  $\mu\text{m} \pm$   
131 157  $\mu\text{m}$ ) below the cortical surface. Further details for these recordings can be found in (Xiao et al., 2017).

132 **Mouse VSD Imaging.** To determine the cortex-wide correlates of LECs, widefield VSD imaging was  
133 carried out in anesthetized mice as previously described (Mohajerani et al., 2010, 2013; Vanni and Murphy,  
134 2014) while simultaneously recording LFP and single unit activity extracellularly. Either a unilateral craniotomy  
135 (1 wildtype C57/BL6 mouse, from bregma 2.5 to -4.5 mm anterior-posterior, and 0 to 6 mm lateral) or a  
136 bilateral craniotomy (2 wildtype C57/BL6 mice, from bregma 3.5 to -5.5 mm anterior-posterior, and -4.5  
137 to 4.5 mm lateral) was made and the underlying dura removed. RH1692 dye (Optical Imaging, New York,  
138 NY; (Shoham et al., 1999) dissolved in HEPES-buffered saline (1 mg/ml) was added to cortex for 60–90  
139 min. VSD imaging began  $\sim$ 30 minutes following washing of unbound dye with saline. VSD data (12 bit  
140 monochrome) was captured with 6.67 ms (150Hz) temporal resolution using a CCD camera (1M60 Pantera,  
141 Dalsa, Waterloo, ON) and EPIX E4DB frame grabber with XCAP 3.1 software (EPIX, Inc., Buffalo Grove  
142 IL).

143 Table 2 summarises recording IDs, anesthetic methods and recording duration for individual extracellular  
144 and VSD recording experiments in mice.

## 145 **Analysis**

146 Most of the analyses were carried out using custom Python code developed as part of an electrophysiology  
147 and optical physiology toolkit currently in development (<https://github.com/catubc/openneuron>). Methods  
148 for computing event triggered analysis for VSD imaging have been previously published (Xiao et al., 2017)  
149 and are also available online ([https://github.com/catubc/sta\\_maps](https://github.com/catubc/sta_maps)).

150 **Single Unit Spike Sorting.** Spike sorting of cat and mouse data was carried out primarily using SpikeSorter

151 (Swindale and Spacek, 2014, 2015) and selectively using Kilosort (Pachitariu et al., 2017) and JRClust (Jun  
152 et al., 2017). For recordings sorted using SpikeSorter, electrophysiological traces were high-pass filtered and  
153 spikes detected using a threshold of 5 times the median of the absolute voltage values of each channel was  
154 divided by 0.675 (Quiñero et al., 2004) followed by a dynamic-multiphasic event detection method  
155 (Swindale and Spacek, 2015). A summary of the sorting results is provided in Table 3 and examples of  
156 sorted spike waveforms are shown in Supplementary Figure S3.

157 **Clustering LFP Events.** High-amplitude LFP events were detected and clustered by converting LFP  
158 recordings to a data format similar to that of a high-pass spike recording. Existing spike sorting tools were  
159 then used for event detection, alignment, feature-extraction, clustering and review. Synchronized states  
160 (Figs 1A) were identified using the deepest LFP channel and on the basis of a synchrony index (SI) (Li et  
161 al., 2009; Saleem et al., 2010) which measures the ratio between power below 4 Hz and total power. Values  
162 of SI greater than 0.5 (i.e. periods where most of LFP power lies in the 0.1 – 4 Hz band) were used to define  
163 the synchronous state. In cat V1 recordings, synchronized state periods accounted for  $2.7 \pm 1.5$  hrs out of a  
164 total recording time of  $8.8 \pm 3.0$  hrs (Table 1) but varied substantially for each recording ranging from 4% to  
165 86% of the total recording period for each animal. This was likely due to variability of anesthetic depth and  
166 animal physiology. In anesthetized mouse sensory cortex recordings, synchronized state periods accounted  
167 for  $2.3 \pm 0.2$  hrs ranging from 84% to 100% of the total recording periods (Table 2).

168 Synchronized state LFP recordings were next high-pass filtered with a 4-pole Butterworth filter with a  
169 cutoff of 4 Hz to remove slower LFP fluctuations and improve signal-to-noise ratio (SNR) for subsequent  
170 event detection, alignment and clustering. Next, event-detection was performed using the same methods  
171 described above for spikes, with the same detection threshold and a temporal window of 50 ms (Swindale  
172 and Spacek, 2015). A temporal lockout of 150 ms and a spatial lockout of 2 mm were used to ensure that  
173 in a 150 ms period only a single LFP event could be identified. Following detection, events were initially  
174 aligned using a weighted center of gravity definition of the time of the event (Swindale and Spacek, 2014).  
175 Max peak alignment revealed similar results. Principal components were then calculated based on the  
176 covariance matrix obtained from the 50 points with the highest voltage variance taken across all channels.  
177 The observed principal component value distributions normally showed clear evidence of clustering (Figs.  
178 1E, 2). Clustering was done based on the first two principal component values. Following clustering, the

179 mean waveform of the events in the cluster was calculated and the individual waveforms were then further  
180 aligned to this mean using r.m.s. error minimization (Swindale and Spacek, 2014) computed over all the LFP  
181 channels. The mean waveform was then recalculated and the process was repeated until further realignments  
182 were of vanishingly small magnitude. The time in the aligned event waveform that corresponded to the  
183 center of gravity of the template (defined as above for individual waveforms) was then taken as the time of  
184 the event. Note that any other stable feature of the template could equally well have been used as an anchor  
185 as this would simply change the times of all the events by the same amount. We used the center of gravity  
186 measure in preference to peaks, troughs or zero-crossings as these features can be variable across different  
187 LECs and are occasionally ambiguous.

188 Clusters with  $< 20$  events or with peak-to-peak heights of  $< 100$   $\mu\text{V}$  were deleted and excluded from further  
189 analysis.

190 **Precision of Estimation of Event Times.** The accuracy with which individual LEC events can be aligned  
191 to the template in the presence of background variability in the LFP signal determines the accuracy of the  
192 estimate of the times at which individual events can be said to have occurred. This accuracy potentially  
193 limits the ability to determine the variability in the timing of spikes of individual units relative to the event.  
194 If the accuracy is low, the measured variability in timing will be larger than it actually is. We estimated the  
195 accuracy of the r.m.s. alignments by computing hybrid ground truth data (i.e. using real data with simulated  
196 shifts). We first calculated the covariance matrix of the noise in the LFP signal relative to a particular LEC  
197 template. This was normally based on the noise values for 100 points (1 ms) on the LFP channel for which  
198 the LEC peak-to-peak amplitude was a maximum. Alternatively, several different channels were included  
199 in the noise calculation. This matrix was then used to generate simulated noise samples with the same  
200 amplitude and covariance structure as the real noise. Simulated LEC waveforms were then generated by  
201 adding samples of simulated noise to the LEC template. Each waveform was then realigned to the template  
202 using r.m.s. minimisation and the resulting shift in position was taken as the error for that particular sample.  
203 The mean of the errors was normally close to zero and the standard deviation of the errors measured for  
204 1000 random samples was taken as an estimate of the accuracy.

205 **CSD Computation.** LEC CSDs were computed by calculating the second spatial derivative (Nicholson and

206 [Freeman, 1975](#)) of LEC templates using all available LFP channels. This calculation was implemented using  
207 the gradient function of the numpy Python library which provides “first or second order accurate one-sides  
208 (forward or backwards) differences at the boundaries” (<https://docs.scipy.org/doc/numpy-1.13.0/reference/generated/numpy.g>

209 **LEC-Triggered VSD Motifs.** VSD motifs were computed as described by ([Xiao et al., 2017](#)). A response,  
210  $dF/F_0$ , was computed for -3 s to +3 s around each LEC event, with  $F_0$  calculated as the average of the signal  
211 -6s to -3s before each event. Strong sensory stimulation resulted in VSD signals which generally peaked at  
212 0.5%  $dF/F_0$ . The LEC triggered VSD motifs had peaks of 0.1-0.2%. These were substantially larger than  
213 randomly generated motif peaks (See Fig 8A-control).

214 **Grouping of LECs using CSD shapes.** CSDs were equalized and then clustered using a generalized  
215 mixture model with 3 components (Fig 4). Because the recordings were made with different length electrodes,  
216 all CSDs shapes were clipped to represent only 0  $\mu\text{m}$  to 1200  $\mu\text{m}$  of cortical tissue. This allowed for a proper  
217 comparison to be made across all 24 selected recordings. Next, the 2D-shape CSDs were aligned to the mean  
218 of all 2D shapes and then converted to a 1D vector. The 1D vector array for all CSDs was then compressed  
219 using principal component analysis and a generalized mixture model with  $n=3$  components was fit to the  
220 resulting distributions (3 was chosen as qualitatively there appeared to be 3 different shapes considering  
221 both CSD shapes and PCA distributions).

## 222 **Results**

### 223 **Clustering Large Amplitude Multi-Laminar LFP Events Reveals Distinct Event Classes**

224 Fig 1 here

225 Multi-channel extracellular recordings were made during synchronized states in anesthetized cat visual  
226 cortex and mouse visual, barrel and auditory cortex (Fig 1A, B). Synchronized state recordings in anesthetized  
227 cats and mice contained large-amplitude, stereotypically shaped LFP events (Fig. 1C). These large amplitude  
228 events were mostly absent during desynchronized cortical states where only lower amplitude events were  
229 typically observed (Fig. 1D). Large amplitude LFP events correlated with peaks in multi-unit-activity (Fig  
230 1E) which are commonly known to be global indicators of UP-state transitions (e.g. (Luczak et al., 2007)).  
231 The LFP events generally had 1 – 3 peaks and troughs with varying relative heights and widths (Fig. 1F).  
232 Peak-to-peak amplitudes were typically 250 – 500  $\mu$ V (after high-pass filtering at 4 Hz) and the duration of  
233 the events was typically 50 – 100 ms. Clustering these events (see Methods) we identified 1 - 4 LECs per  
234 recording session in cat visual cortex and 1 - 2 LECs in mouse visual, auditory and barrel cortex recording  
235 sessions (Fig 1H, I; Tables 1 and 2; see also Methods).

236 Fig 2 here

237 The stability of LEC shapes could be tracked over time using the principal component values of the LEC  
238 event waveforms. We found that LEC shapes were stable and the principal component values of events  
239 within each cluster did not change substantially over periods of up to 3 hours in either cat or mouse cortical  
240 recordings (Fig. 2A, B).

### 241 **LECs Reflect Distinct CSD Patterns Common Within and Across Animals**

242 Fig 3 here

243 We used CSD analysis to further investigate the properties of LECs within and across animals (Figs. 3  
244 and 4). Both the raw unaveraged LFP traces (Fig. 3A, B, E, G) and the averaged LEC profile (Fig. 3C,  
245 D, F, H) revealed distinct CSD patterns present in cortex during synchronized states. CSD profiles from  
246 synchronized state cortical recordings showed periodic large amplitude events corresponding to LECs (Fig  
247 3A, E and G) whereas CSD profiles from desynchronized state recordings contained more frequent activity

248 distributed across multiple depths and with lower amplitude (Fig 3B; note amplitude were normalized within  
249 each session). Computing CSD profiles for different LEC templates in cat V1 recordings current-sink  
250 distributions showed they could be similar across different tracks within the same animal. For example,  
251 Fig. 3C shows 4 similarly shaped LEC CSD profiles from 4 different tracks in two hemispheres from a  
252 single cat. Some profiles were similar across different animals. Fig. 3D shows 5 similarly shaped CSDs, 4  
253 of which are from four tracks in one cat in both hemispheres and 1 from another cat. Figures 3E and G show  
254 CSD profiles for unaveraged LFP events recorded in mouse visual and auditory cortex respectively. Figures  
255 3F and H show CSD profiles obtained from the averaged LEC waveform for mouse visual and auditory  
256 cortex respectively.

### 257 **LECs Are Similar Within and Across Animals**

258 Fig 4 here

259 Across 14 V1 cortex tracks in 5 cats we identified 34 LECs (Table 1) with between 1 – 4 LEC per track.  
260 Using a gaussian mixture model with 3 components we grouped the 24 most common CSD shapes into  
261 three groups (Fig 4; see Methods). Thus rather than being unique or specific to a particular animal or track,  
262 LECs can be grouped on the basis of their CSD shape and can be the same within and across different cats.  
263 This suggests LECs may be generated by common neural circuits underlying UP-state transitions. Different  
264 types of UP-state transitions may thus be present across animals within a species (Table 4). Additionally,  
265 in a total of 8 mice recordings from visual, auditory and barrel cortex we found 13 LECs that could also be  
266 grouped (Table 7). The average frequency of the LECs (taken across all types) was 0.12 Hz across all cat  
267 V1 recordings, and 0.18 Hz across all mouse cortex recordings.

### 268 **LEC Events Can be Localised with 1 - 4ms Temporal Precision**

269 Fig 5 here

270 We estimated the accuracy with which the times of individual LEC events could be measured by taking  
271 individual LEC templates and creating synthetic events by adding artificial noise with the same temporal  
272 structure as the real noise measured from the individual, aligned, events in the LEC (Fig 5). We then  
273 measured the change in the time of the event required to minimise the r.m.s. difference between it and the

274 template, this change being the alignment error resulting solely from the added noise. The standard deviation  
275 of the resulting changes with repeated noise samples was taken as an estimate of the likely alignment error  
276 present with the real events in the sample. This was done for a number of different LECs in several different  
277 recordings. Error estimates ranged from 1.0 ms to 4.7 ms with a mean of 2.7 ms (n=11 in 6 different  
278 recordings). These estimates also put a lower bound on the accuracy with which spike times relative to  
279 individual LEC events can be measured.

280 The estimates were based on waveform data from single LFP channels (the one on which the peak-to-  
281 peak amplitude of the LEC template was a maximum). Adding additional channels improves the accuracy  
282 (though the improvements were found to be small, likely because signals tended to be highly correlated  
283 across adjacent channels). We also explored the temporal precision of LEC event detection by computing the  
284 stability of the full-width-half-max (FWHM) of the largest negative peak of each LEC event (Supplementary  
285 Figure 3). We found that the standard deviation of the FWHM were less than 10ms in most cases further  
286 suggesting that the overall shape of the LECs is relatively stable across time.

### 287 **Most Single Neurons Fire Precisely in Relation to LECs**

288 Fig 6 here

289 We next considered the timing relation between single-unit firing and LEC events. We computed Peri-LEC-  
290 Event-Time-Histograms (PLETHs) using 5 ms bins and triggering off the LEC event times (Figure 6; see  
291 also methods). We found that neurons had qualitatively different PLETH distributions with different peaks  
292 shifted in time (Fig 6A: cat visual cortex recording; Fig 6B mouse visual cortex recording). Plotting the  
293 PLETHs for each neuron in a recording and arranging neurons by depth (Fig 6C-E: left) revealed a gradual  
294 temporal difference in peaks across almost all units recorded. Plotting units by depth (Fig 6C-E: right  
295 panels) revealed a preference for deeper units to fire earlier in the recording. We further analyzed whether  
296 deeper layer units fired before upper layer units, as reported by others ([Sanchez-Vives and McCormick, 2000](#);  
297 [Volgushev et al., 2006](#); [Chauvette et al., 2010](#); [Beltramo et al., 2013](#)). PLETHs were ranked in order  
298 of depth (position of the unit) along the electrode (Figure 6C-E, rightmost panels). Units in the recordings  
299 were next divided into upper and lower halves (0 – 425 and 425 – 850  $\mu$ m respectively for the mouse; and  
300 0 – 750 and 750 – 1500  $\mu$ m respectively for the cat recordings). We then counted the number of times the

301 mean latency to the peak of the fitted gaussian was shorter for units in the upper vs. the lower halves of  
302 the recording (Supplementary Figure 5). In most cases latencies in the lower halves were shorter (10 of 14  
303 mouse recordings and 21 of 32 cat recordings). These differences were not significant when tested separately  
304 ( $p = 0.09$  and  $0.06$  respectively, binomial test) but were significant when both data sets were combined ( $p =$   
305  $0.013$ , binomial test).

306 We lastly considered whether the spiking distributions (i.e. not just the peaks) were statistically different  
307 across neurons. We found that despite sparse firing for many neurons, spiking distributions were almost  
308 always different between pairs of units recorded simultaneously (i.e. more than 90% of neuron pairs had  
309  $< 0.01$  pvalues; 2-sample Kolmogorov-Smirnov tests, Bonferroni corrected, Supplementary Figure 4). In  
310 other words, almost all neurons had unique firing distributions relative all other neurons even though their  
311 spiking distributions fell within a window of  $\sim 25$ -50 ms.

### 312 **Most Neurons Lock to LEC Defined UP-states with $\pm 5$ -15 ms Latencies**

313

Fig 7 here

314 In order to quantify the temporal precision with which different units could fire in relation to the LEC we  
315 selected units based on how well a gaussian fitted the PLETH (see Methods). Unit-PLETH pairs where the  
316 standard deviation of the fitted gaussian was  $> 50$  ms showed poor fits and were excluded from analysis, as  
317 were units where no histogram bin exceeded a count of 5 in any 5 ms wide window (i.e. when only a few  
318 spikes occurred during the recording period). These criteria resulted in the removal of  $\sim 25\%$  of recorded  
319 units). We found most neurons had gaussian fits with 1-5ms standard deviations for both mouse (e.g. Fig  
320 7A) and cat cortex (Fig 7B). Plotting the distribution of fits for all units recorded in different tracks and  
321 animals yielded an estimate of how 'precise' the firing of a unit could be determined relative the LEC. We  
322 found that across all unit-LEC pairs (3009) the mean precision was  $\pm 11.9$  ms - though many units had lower  
323 values, with the lowest being 1.4 ms - close to the limit of the accuracy with which LEC event times could  
324 be measured.

### 325 **LECs Have Broad Mesoscale Correlates in Mouse Dorsal Cortex**

326

Fig 8 here



327 We lastly sought to capture the meso-scale correlates of LECs by simultaneously recording widefield voltage-  
328 sensitive-dye (VSD) signals in mouse visual and auditory cortex (Fig 8A) and in GCaMP6s mice while  
329 recording extracellular potentials in mouse visual, barrel and auditory cortex (Fig 8B). LEC-triggered averages  
330 of widefield activity were computed as previously described (Xiao et al., 2017) for periods of 2 s relative  
331 to the LEC event time (i.e.  $\pm 2$  s relative to each LEC event). In VSD recordings, the spatio-temporal  
332 patterns (termed ‘motifs’ here) showed a peak at the electrode recording site and revealed gradual multi-area  
333 cortical activation preceding LEC events (Fig 8A,  $t=0$  s). This indicates that LECs are preceded by gradual  
334 membrane depolarizations and/or firing of many neurons as VSD activity represents both subthreshold  
335 and suprathreshold neural activity. This suggests that clustered LFP events are consistent with UP-state  
336 transition dynamics observed in intracellular recordings where near-simultaneous (10 - 100 ms) activation  
337 of neurons is observed during UP-state transitions across many cortical areas (Destexhe et al., 1999; Amzica  
338 and Steriade, 1995). The findings also suggest that LECs are the LFP-correlates of UP-state transitions in  
339 cortex. GCaMP6s mouse recordings also revealed a similar structure during LEC events (Figs 8B; see also  
340 Supplementary Videos 1 and 2).

## 341 **Discussion**

### 342 **Multiple LEC Types Suggests Multiple Sources of UP-state Genesis**

343 It has been previously shown that UP-state transitions in single neurons have LFP correlates and various  
344 recent studies have used peaks in multi-unit-activity as markers of UP-state transitions ([Amzica and Steriade,](#)  
345 [1998](#); [Chauvette et al., 2010](#); [Saleem et al., 2010](#); [Luczak et al., 2007](#)). We describe a method for grouping  
346 UP-state transitions based on stereotyped multi-channel LFP event shapes. We show that in some recordings  
347 multiple classes of such events can be defined and that most single neurons synchronize their firing to  
348 within a few milliseconds of such events. The finding of multiple classes (1 to 4) of LECs in different  
349 species and cortical areas is consistent with a growing body of work identifying stereotypy in LFP recordings  
350 ([Reichinnek et al., 2010](#); [Ramirez-Villegas et al., 2015](#)). Another study that used MUA to define UP-state  
351 transitions found two types of transitions in ketamine/xylazine-anesthetized rats based on UP-state duration  
352 ([Luczak and Bartho, 2012](#)). The findings of multiple classes of UP-state transitions is also supported by  
353 CSD correlates which have clearly different laminar patterns that are common across cortical areas and  
354 individuals of the same species. These findings of different patterns potentially occurring during UP-state  
355 transitions lend some support to the three cardinal oscillator hypothesis that UP-states can be caused and  
356 sustained by potentially independent cortico-thalamic-cortico populations ([Crunelli and Hughes, 2010](#)).

### 357 **Multi-channel LECs Capture Global UP-state Transitions**

358 The term UP-state has commonly referred to a single neuron's resting membrane potential transitioning from  
359 a hyperpolarized (i.e. non-spiking, e.g. -80 mV) to a depolarized (e.g. -60 mV) state ([Steriade et al., 1993a](#)).  
360 The term UP-state has also been used to refer to a global correlate of single neuron UP-state transitions  
361 where many (or possibly all) cortical neurons in a region depolarize and spike simultaneously ([Neske, 2016](#)).  
362 Defining an exact UP-state transition time faces certain problems. While most (or all) neurons depolarize  
363 simultaneously during an UP-state transition, not all neurons spike on every UP-state cycle ([Volgushev et](#)  
364 [al., 2006](#); [Chauvette et al., 2010](#)). Additionally, individual neurons depolarize at different times and rates  
365 based on measurements of their intracellular membrane potentials ([Lampl et al., 1999](#); [Petersen et al., 2003](#))  
366 and can have different UP-state dynamics ([Ros et al., 2009](#)).

367 These factors make it challenging to track UP-states globally solely by recording a few neurons intracellularly  
368 or tracking peaks in MUA. In the present study we have shown that stereotypy in the LFP waveform can be  
369 used to establish the time of each event with a precision of a few milliseconds. This marker can then serve  
370 as a physiologically relevant temporal reference for evaluating single unit spike timing. Previous methods  
371 of identifying the time of UP-state transitions include measures based on changes in the firing rate of  
372 simultaneously recorded neurons (Luczak et al., 2007, 2009, 2013) but these methods have the limitation that  
373 they can only be applied to high-firing rate neurons (i.e. because peaks in cumulative firing rate histograms  
374 have low-SNR and are unreliable) in cortical areas that are not sparsely firing. Combined intracellular and  
375 LFP recordings (Chauvette et al., 2010) can be used to define UP-state transitions by fitting a sigmoid to  
376 LFP traces and defining a transition point at 10% of the amplitude of the sigmoid. The limitation is that  
377 both intracellular and extracellular recordings are required and that a somewhat arbitrary point is chosen as  
378 the time of the UP-state transition. (Saleem et al., 2010) used a method based on the phase of the LFP at  
379 frequencies below 4 Hz combined with multi-unit activity and single neuron recordings. This also has the  
380 limitation that both LFP and multi-unit activity are needed to define the onset of UP-states. Overall, none of  
381 the previous work has demonstrated a particular degree of precision in defining the time of UP-state onset.  
382 While we claim to have found temporally precise markers of UP-state transitions our LEC times reflect a  
383 choice in event feature location (e.g. LEC  $t=0$  ms can be chosen at peak, trough or centre-of-gravity of LEC  
384 event). Yet this limitation is also present in defining a global UP-state transition time using intracellular  
385 membrane potentials given that individual neurons can transition to UP-states at different times (LampI et  
386 al., 1999; Petersen et al., 2003).

### 387 **Relation between LECs and K-complexes**

388 Like the LECs studied here, K-complexes are transient large amplitude events that occur in EEG or LFP  
389 recordings during the synchronised state in anesthesia and during stage 2 slow wave sleep (e.g. (Loomis  
390 et al., 1937; Amzica and Steriade, 1998). In humans, K-complexes typically last up to 1 second and occur  
391 every 1 – 2 minutes. They are often followed by sleep spindles - a burst of rapid oscillations at a frequency  
392 of 10 – 12 Hz. In recordings from ketamine-anesthetised rats (Luczak and Bartho, 2012) also describe  
393 large amplitude fluctuations in LFP recordings which mark transitions to UP-states where large numbers of

394 neurons start firing at about the same time. That study suggests that these fluctuations are homologous to  
395 K-complexes. However there are differences between all three sets of observations – classical sleep related  
396 K-complexes in humans, those of (Luczak and Bartho, 2012) and ours. Like Luczak and Bartho’s findings,  
397 ours differ from classical K-complexes in being much faster (lasting 100 – 200 ms compared to up to 1  
398 second) and occurring at much higher rates (several/second compared to every 1 – 2 minutes). Ours also  
399 differ from those observed by Luczak and Bartho in being simpler in structure, with typically a single large  
400 peak (positive or negative) flanked by two of opposite sign (see Fig 1G). We also did not find the traveling  
401 wave events described by Luczak and Bartho, possibly because these are reported to be of smaller amplitude  
402 and they perhaps fell below our detection threshold. Nor did we observe obvious sleep spindles following  
403 our events. Reasons for these various differences would include species (human K-complexes and spindles  
404 may be generally slower than in rats and cats), the fact that the animals were not naturally sleeping, types  
405 of anesthesia (Luczak and Bartho used ketamine, we used either isoflurane or propofol) and cortical area  
406 (we studied visual areas whereas Luczak and Bartho studied rat auditory cortex). A conservative hypothesis  
407 that might reconcile all of these findings is that K-complexes constitute a large and heterogeneous class of  
408 high amplitude transient activity in the LFP associated with UP-state transitions and widespread firing of  
409 neurons. Our findings of multiple types of LECs support such heterogeneity within single cortical areas and  
410 recording sessions, as well as suggesting that specific types of events may be identifiable within areas and  
411 across different individuals of the same species (Fig. 4).

#### 412 **LECs as temporally precise global markers of UP-state transitions**

413 As an alternative global-definition of UP-state transitions, LECs have advantages over single neuron patch  
414 clamp recordings in that they can be more rigorously defined using statistical clustering methods while  
415 also being more stable as they consist of spatially broad (i.e. 100  $\mu$ m to 1000  $\mu$ m) LFP contributions from  
416 multiple sources (Buzsáki et al., 2012) while being largely independent of any single neuron’s activity.  
417 While simultaneous intracellular recordings from many neurons might eventually be feasible, defining  
418 global UP-state transitions using such recordings stills requires averaging UP-state transition times leading  
419 to a definition that is dependent on the particular set of recorded neurons. Since the LFP represents the  
420 activity of a large population of neurons, transition times estimated from the stereotyped shapes of multi-  
421 channel LFP signals may provide a principled and non-circular methodology, i.e. it does not define UP-state

422 transition spiking based on the cumulative spiking of many neurons.

423 We propose that future work should focus on the implications of the narrowly defined UP-state transitions  
424 spiking (previous work showed histogram widths of 20-150 ms ([Luczak et al., 2007](#))). Such narrower spiking  
425 distributions lend support to spike timing and firing order being present in cortical processing ([Panzeri et al.,  
426 2001](#); [Gautrais and Thorpe, 1998](#)).

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## 527 **Legends**

528 **Figure 1. Clustering LFP Events During Synchronized Cortical States.** A. Example of a 200 s cat visual  
529 cortex extracellular recording with power spectrogram (top), synchrony index (middle), single neuron rasters  
530 (bottom) and 64 channel extracellular probe diagram (right). B. Same as (A) but from mouse visual cortex  
531 using a different type of extracellular probe containing 64 channels. C. Extracellular recording of 10-LFP  
532 channels obtained from a track in cat visual cortex during a synchronized state reveals infrequent large  
533 amplitude cross-laminar events (animal ID: C5). D. Same track as in (A) but recording acquired during a  
534 desynchronized cortical state shows higher frequency cross-laminar events that are much smaller. E. Large  
535 amplitude LFP events correlate with peaks in MUA histograms during synchronized state recordings (ID:  
536 C5). F. Large amplitude LFP events are detected (see text for details). G. detected events are aligned and  
537 features are clustered using PCA. H. LFP events are labeled (colors) and event times exported for analysis  
538 (see also Methods). I: left four panels: LFP templates for the 4 LECs identified in (F) show distinct multi-  
539 laminar LFP patterns. Middle and right panels: LFP templates for LECs identified in mouse visual, barrel  
540 and auditory cortex respectively.

541 **Figure 2. Stability of LECs over Time.** A: shows the principal component values (PC1 and PC2) for the  
542 4 LECs shown in Fig 1E, plotted over a period of 400 minutes (6.7 hours). This shows that individual LEC  
543 features are relatively stable over time. B: same as A but for 2 LECs in a mouse visual cortex recording.

544 **Figure 3. Current-Source-Density (CSD) Analysis of LECs.** A. CSDs for a 850 ms synchronized state  
545 recording (same track in animal C5 as shown in Fig 1A). B. CSD distributions for a 850 ms desynchronized  
546 state recording (same track as Fig 1B, 3A). C. CSD obtained from averaged LEC event profiles (i.e. templates)  
547 recorded in 4 different visual cortex tracks in animal C5. D. CSD distributions for averaged LEC profiles  
548 recorded in 5 different visual cortex tracks in two cats (1-4: animal C5; 5: animal C4). E. Same as panel A,  
549 but for a recording from mouse visual cortex. F. CSDs from two averaged LEC waveforms in a single mouse  
550 visual cortex recording. G. CSD profiles from unaveraged LFP events in a synchronised recording from  
551 mouse auditory cortex. H. Same as 3C, F but for a mouse auditory cortex recording. All CSD distributions  
552 were normalized to lie between -1 and +1 for visualization.

553 **Figure 4. LEC Grouping and LEC Shape Stability.** A. Principal component distributions for the most

554 common 24 CSDs detected in cat visual cortex recordings were fit with a 3-component generalized gaussian  
555 mixture model (see Methods). Shaded regions represent 1 standard deviation. B-D. CSD for each group  
556 clustered in A.

557 **Figure 5. Estimating the temporal precision of individual LEC events.** The solid black line shows the  
558 average (the template) of the events in a LEC. The 3 blue traces show the profiles of 3 individual events and  
559 the 3 red traces show the profiles of simulated events obtained by adding correlated noise to the template.  
560 Event times are estimated by finding the horizontal position of a profile that minimises the r.m.s. difference  
561 between it and the template. Time zero is defined as a weighted measure of the center of gravity of the  
562 template (see Methods).

563 **Figure 6. LECs Correlate with Single Neuron Activity.** A. Peri-LEC event time histograms (PLETHs) for  
564 3 example neurons and 4 LECs from a cat visual cortex recording (obtained as described in the Methods).  
565 Each dot represents a spike for a particular neuron (color) relative to the LEC event time ( $t=0$  ms). B. Same  
566 as A but from a recording in mouse visual cortex. C. PLETHs for all neurons in A ordered by latency of the  
567 peak of each histogram relative to the time ( $t = 0$ ) of each LEC-event (left; shortest latencies at the top) or  
568 by the depth of recorded neuron (right). D. Same as C but from a mouse visual cortex recording (MV1). E.  
569 Same as (C, D) but for a mouse auditory cortex recording (MA1)

570 **Figure 7. Single units fire precisely in relation to LEC onset.** A. PLETH firing rate histograms (green  
571 lines) from a mouse visual cortex recording (ID: MV4) were fit with a Gaussian function (dashed black  
572 lines) to determine the mean latency and the width of the distribution. Fits were confined to units which  
573 fired reliably in relation to the events (see Results for details). B. Same as A, but for a cat visual cortex  
574 recording (ID: C4). C. All gaussian fits (3009) for each neuron-LEC pair in every track for all animal  
575 recordings. The vertical axis shows the width (sigma) of the fitted peak for each of the selected units. Each  
576 vertical column of points is the data from a single LEC type. Columns are grouped horizontally by animal  
577 and then by track number in each animal. The narrowest widths (indicating the most precisely firing units)  
578 are below 10 ms. The mean and mode of the overall distribution were 11.9 and 11.0 ms respectively.

579 **Figure 8. LEC-Triggered Single- and Bi-hemispheric Wide-field VSD Motifs.** A. Voltage-sensitive-dye  
580 imaging of LEC-triggered dynamics in an auditory cortex recording reveals dynamics surrounding LEC

581 event time for two different LECs. The control panel shows imaging data from randomly generated trigger  
582 times. Controls for other experiments were similar. B: Same as A but from two different recordings in  
583 mouse visual cortex. C: A, B but from GCaMP6s mice recordings in auditory, visual and barrel cortex (see  
584 also main text, Methods). White dots show the position of the recording electrode on which the triggering  
585 LECs were detected.

586 **Figure 1-1: Examples of single unit clustered spikes.** Spikes on the maximum amplitude channels for  
587 nine randomly chosen single units from both cat and mouse recordings.

588 **Figure 3-1:** PETH distributions computed for wider temporal windows (-500 ms to +500 ms) than Figure  
589 6, for 3 LECs recorded in cat visual cortex, mouse visual cortex and mouse auditory cortex.

590 **Figure 5-1: Measuring the stability of LEC events.** A. The four LEC templates shown in Fig 1G. B.  
591 Measurement of the full-width-half-max (FWHM) from the first trough (i.e. negative peak) of each LEC  
592 event. C. FWHM means and standard deviations of the four LEC events shown in A reveal that most LEC's  
593 FWHM standard deviations are <10 ms. D. Same as C but for the first 6 LEC groups in Fig 4, reveal the  
594 vast majority of LECs have individual standard deviations < 10 ms.

595 **Figure 6-1.** LEC-triggered spiking distributions are significantly different across neurons. The significance  
596 of the difference in the distributions between pairs of neurons was assessed using 2 sample Kolmogorov-  
597 Smirnov tests with Bonferonni correction. Histograms show the distributions of  $p$  values for all the unit  
598 pairs in particular recordings A: recording C5.3; B recording MV1 and C recording MA1.

599 **Figure 6-2.** Deeper layer neurons are more likely to fire first during LEC-events. Top: histograms show  
600 the number of LECs in which superficial layer (blue) neurons (0-425  $\mu\text{m}$ : mouse; 0-750  $\mu\text{m}$ : cat) spiked  
601 before deeper layer (red) neurons (425-850  $\mu\text{m}$ : mouse; 750-1500  $\mu\text{m}$ : cat). Order was based on the means  
602 of Gaussian fits to LEC-triggered firing rate histograms. There is an overall bias for deeper layer neurons  
603 to spike first (see text for further description of statistical tests). Bottom: pooling all neuron relative firing  
604 times did not reveal substantial order differences between superficial layer neurons (blue) or deeper layer  
605 neurons (red) firing first. This was likely due to individual LECs eliciting different lag spiking which is  
606 averaged out when pooling all relative spiking times across all LECs.

607 **Illustrations and Tables**

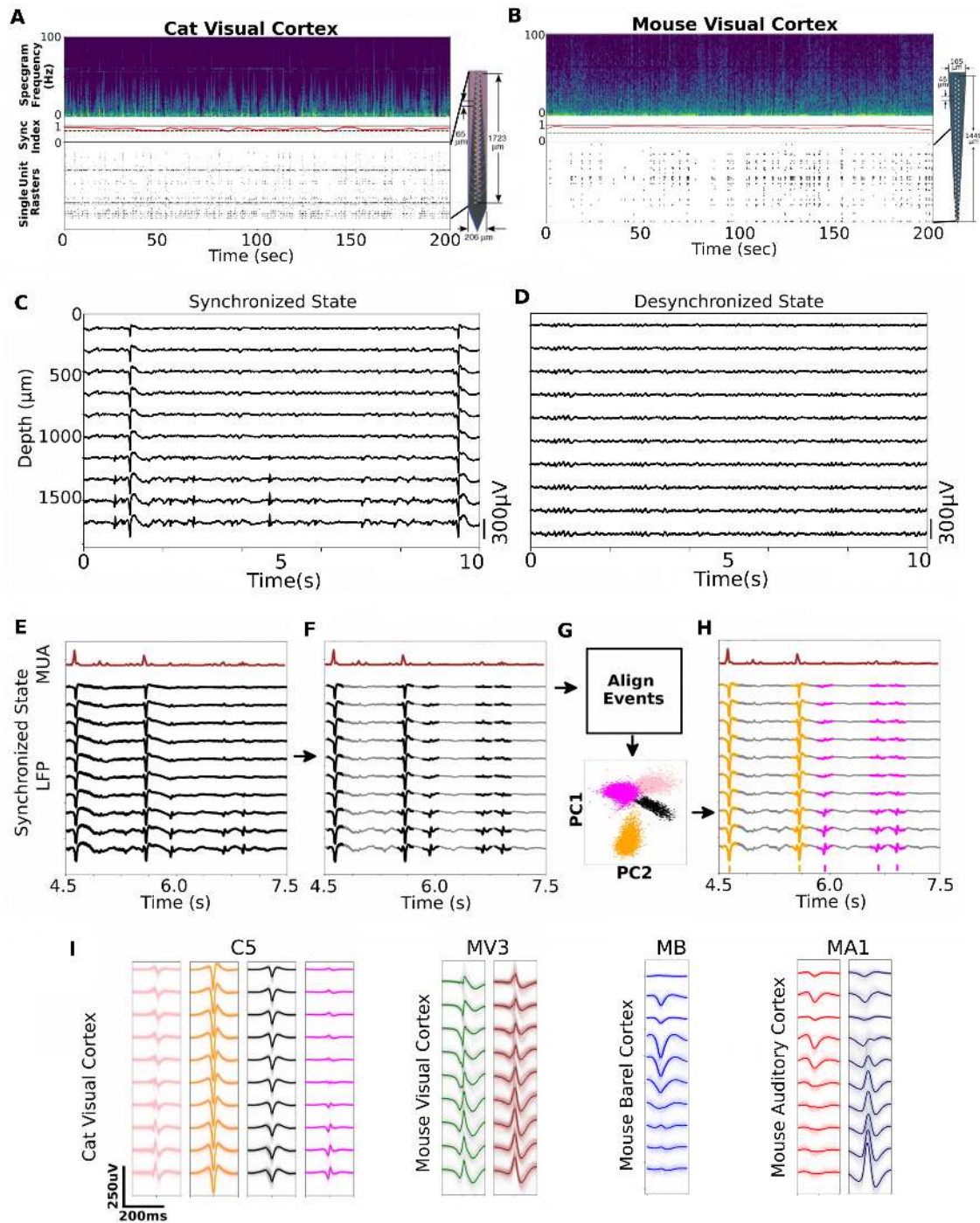


Figure 1

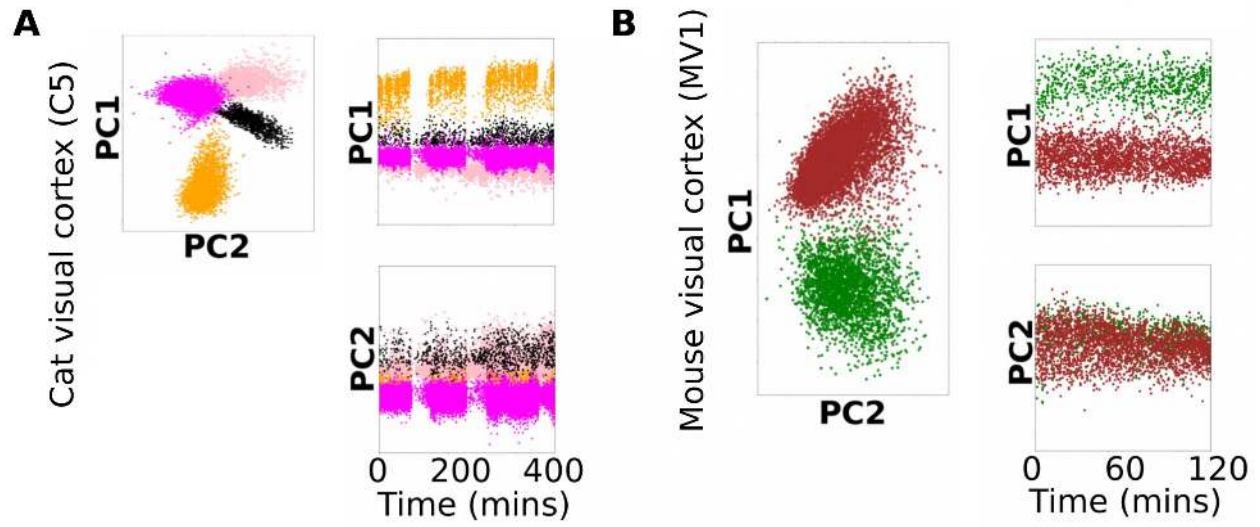


Figure 2

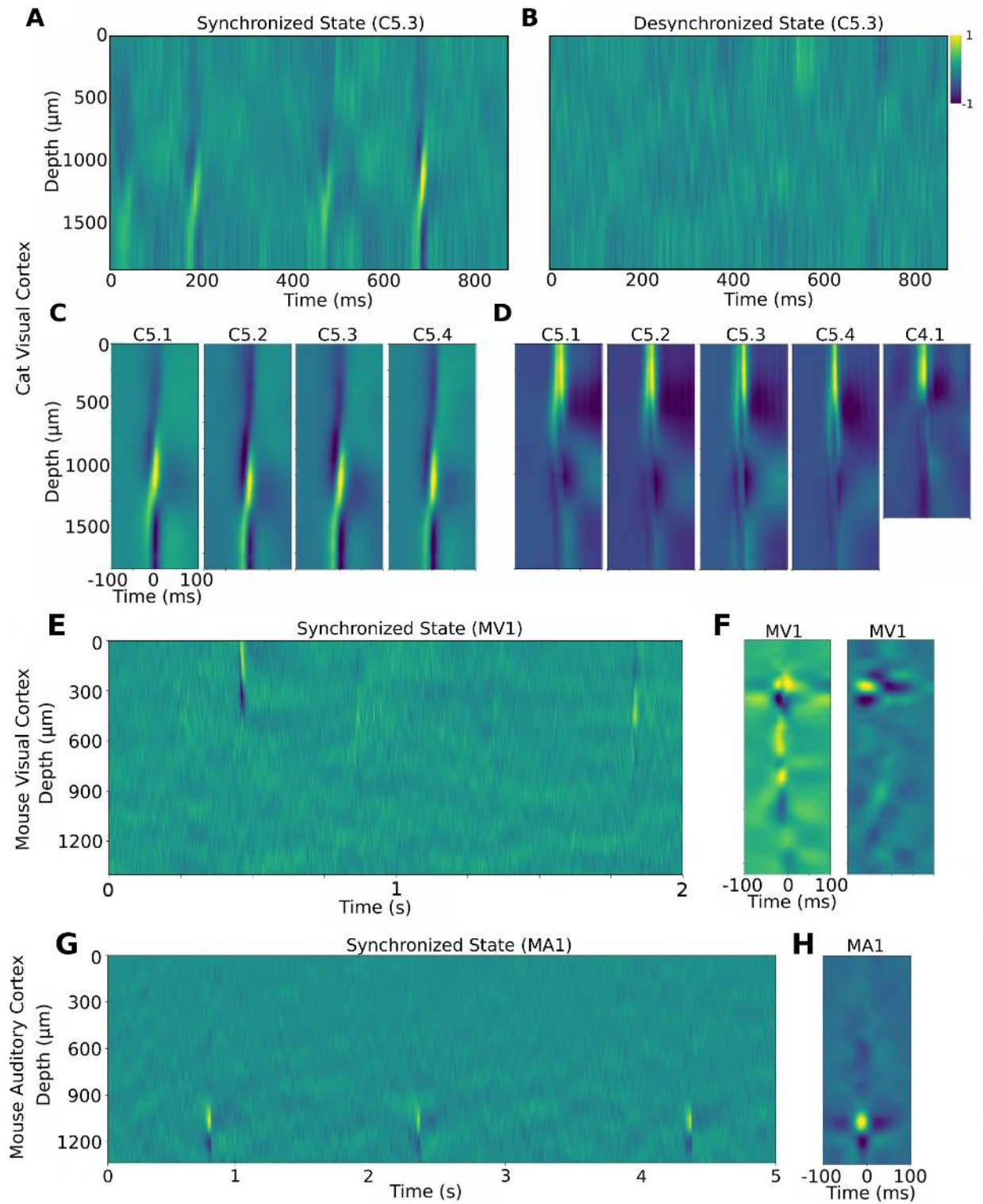


Figure 3



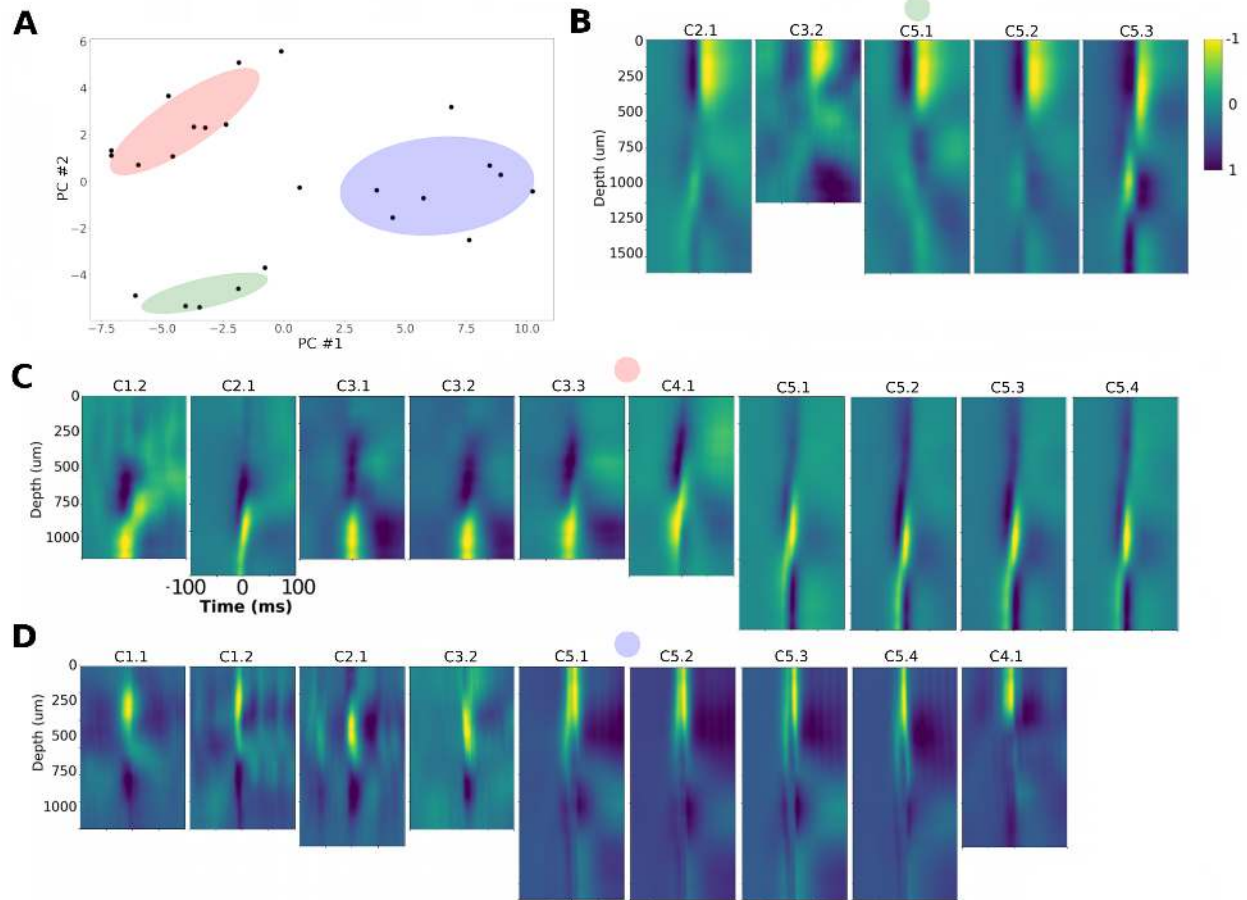


Figure 4

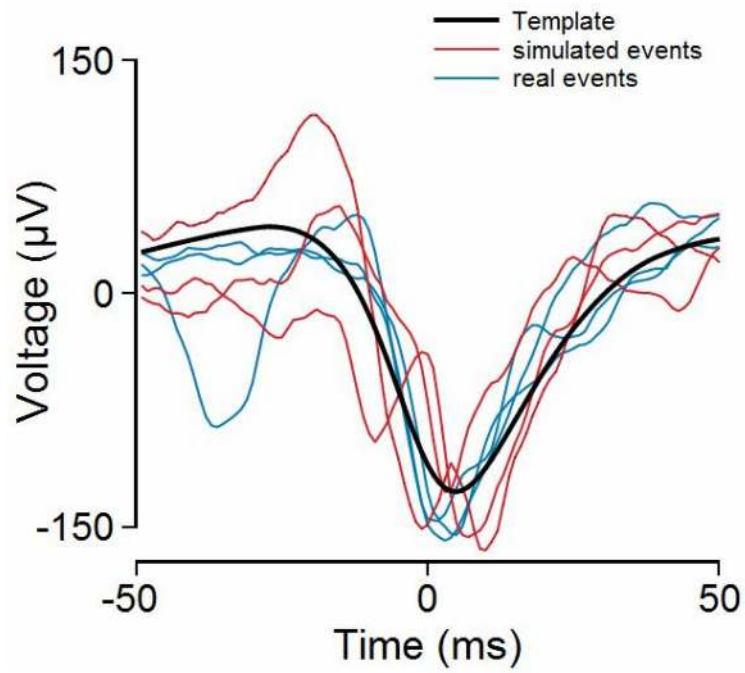


Figure 5

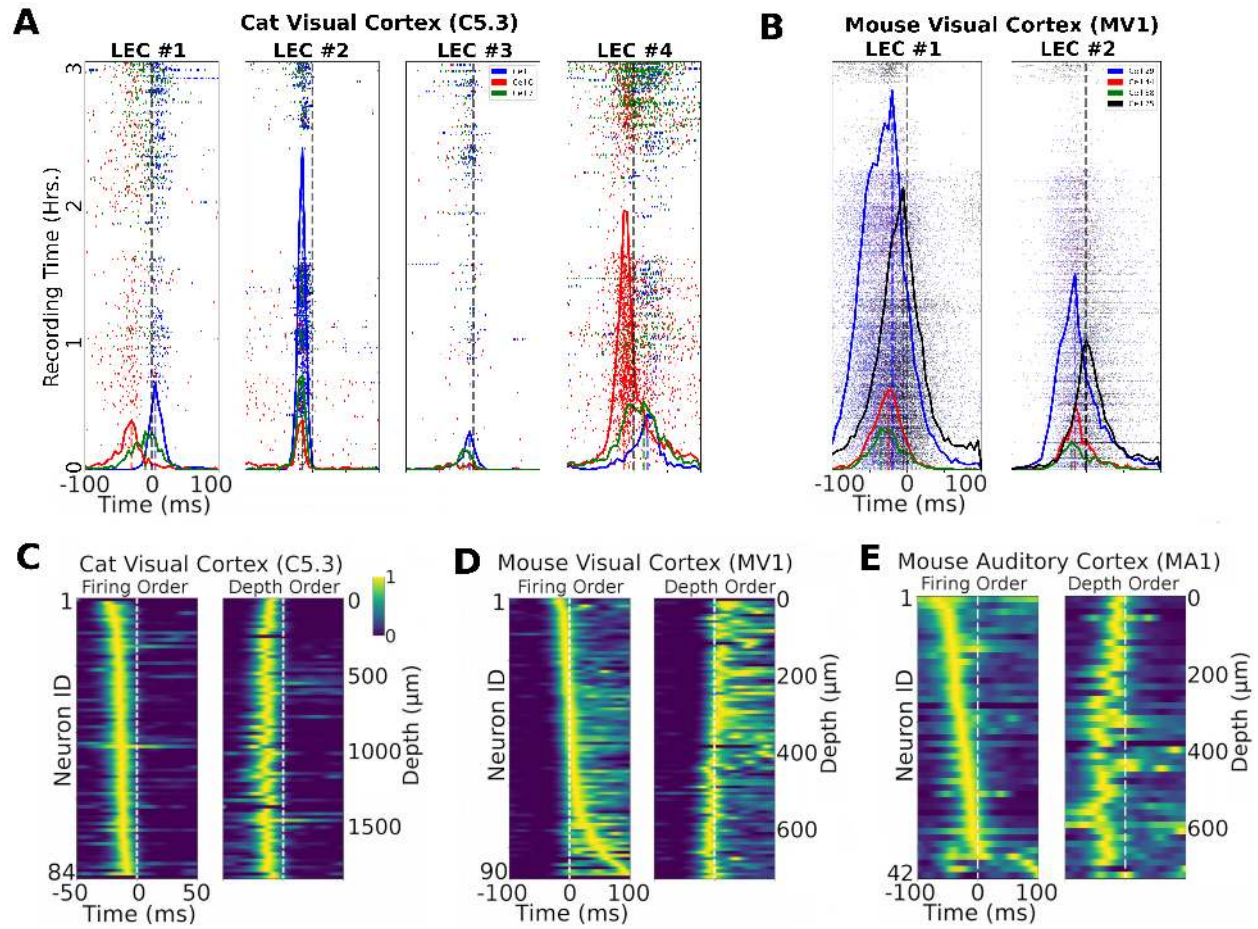


Figure 6

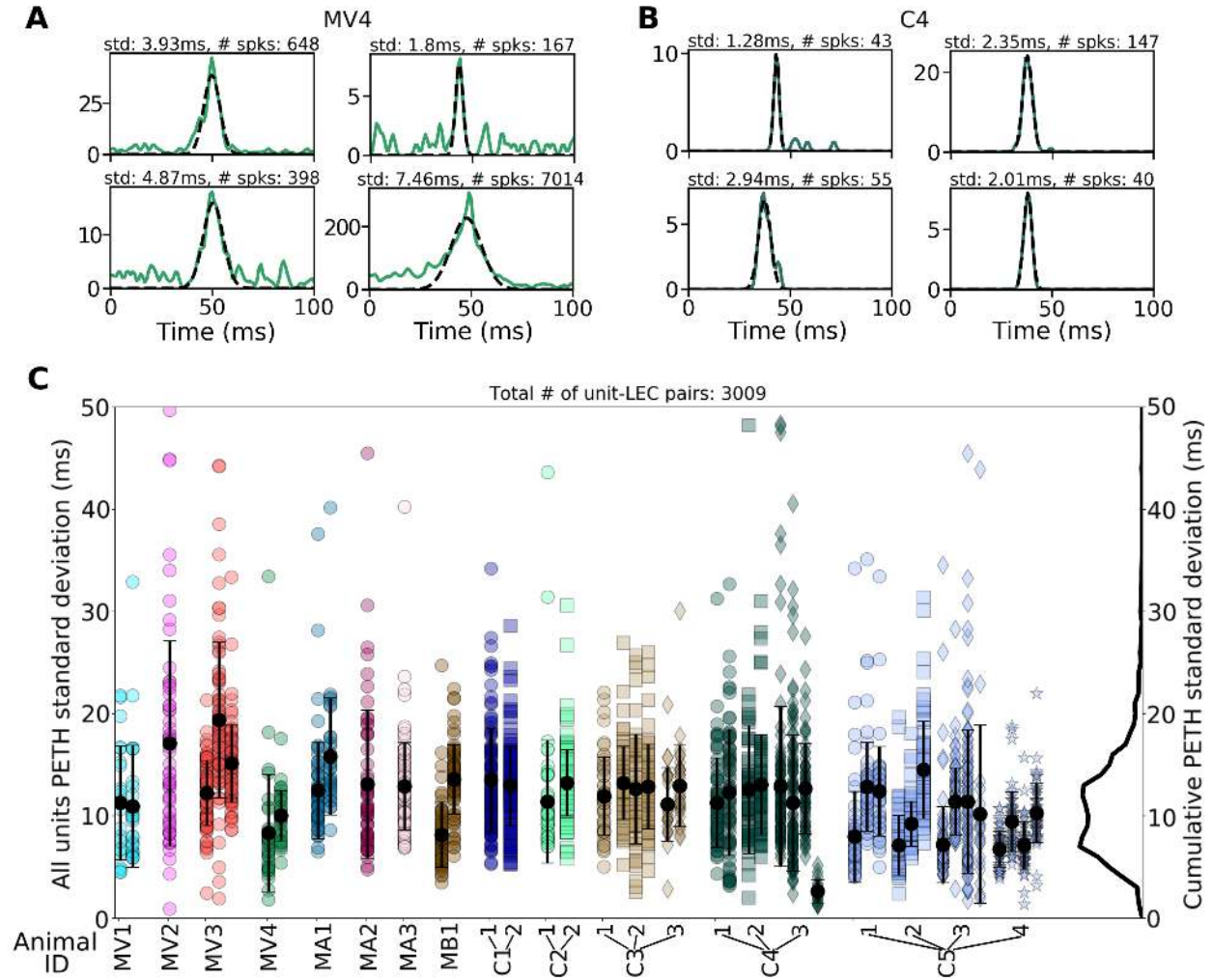


Figure 7



Figure 8

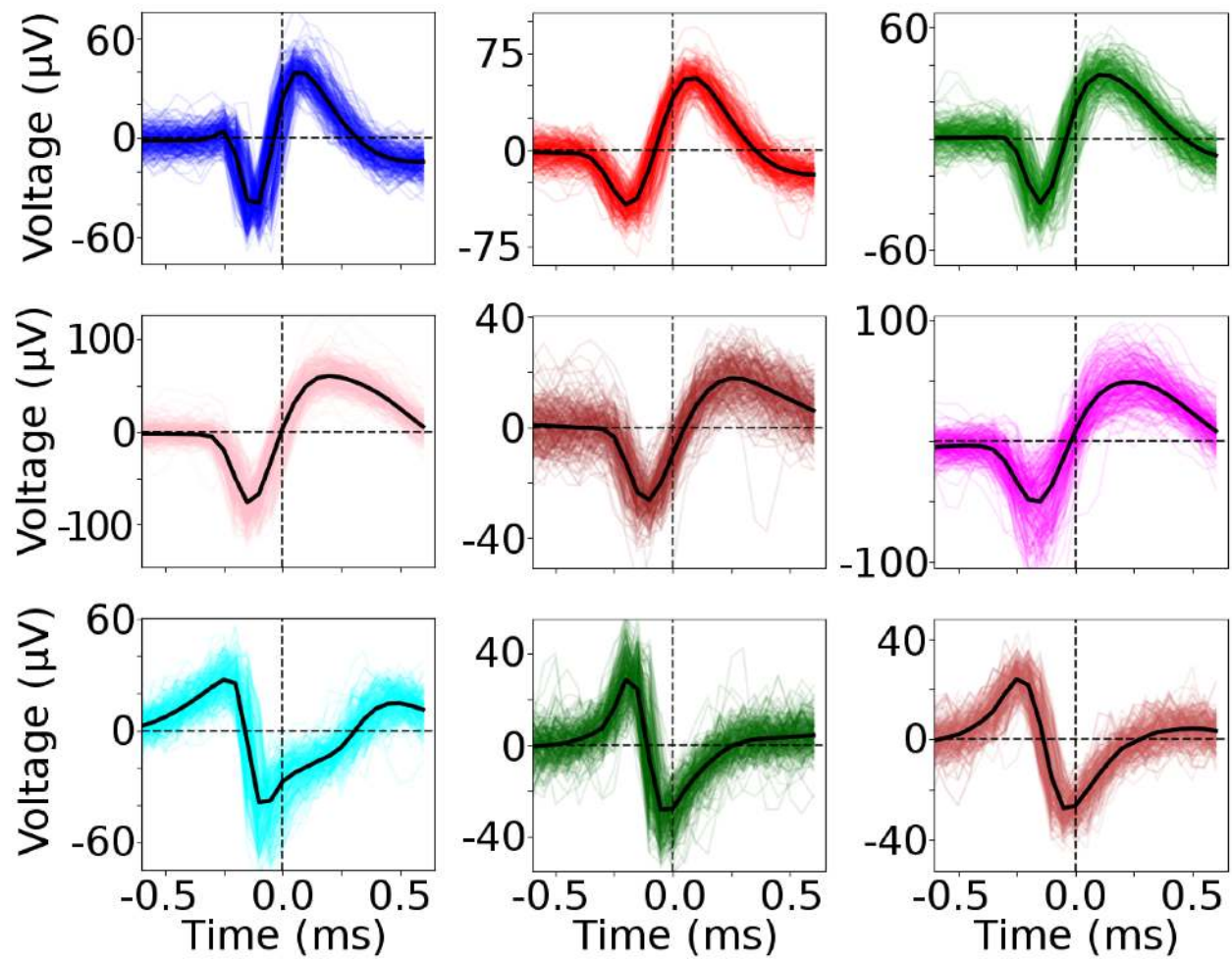


Figure 1-1

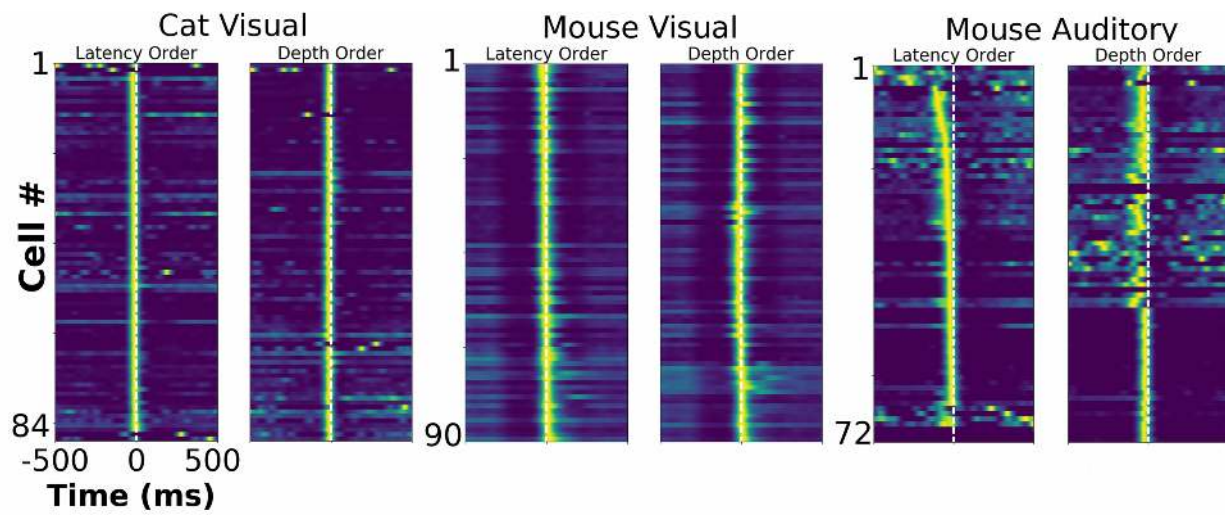


Figure 3-1

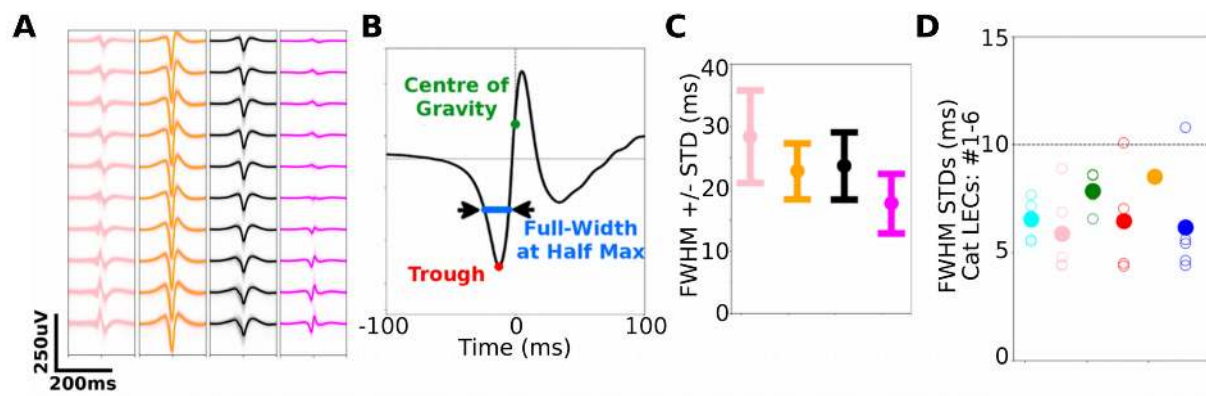


Figure 5-1



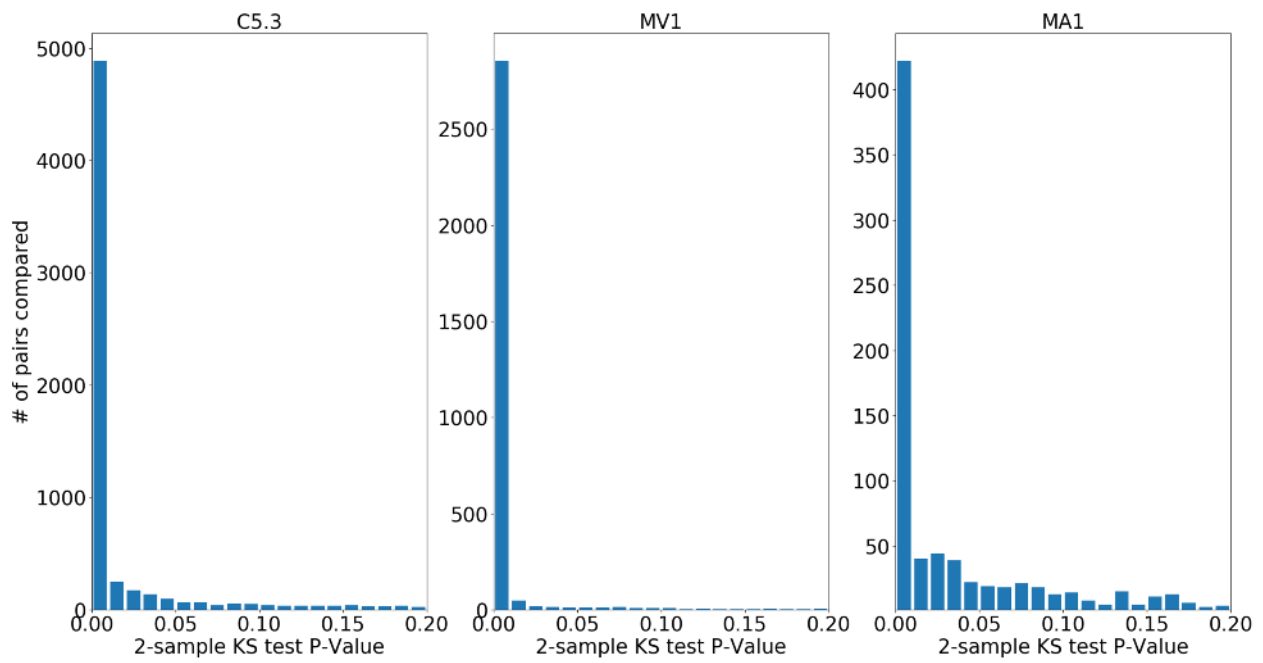


Figure 6-1

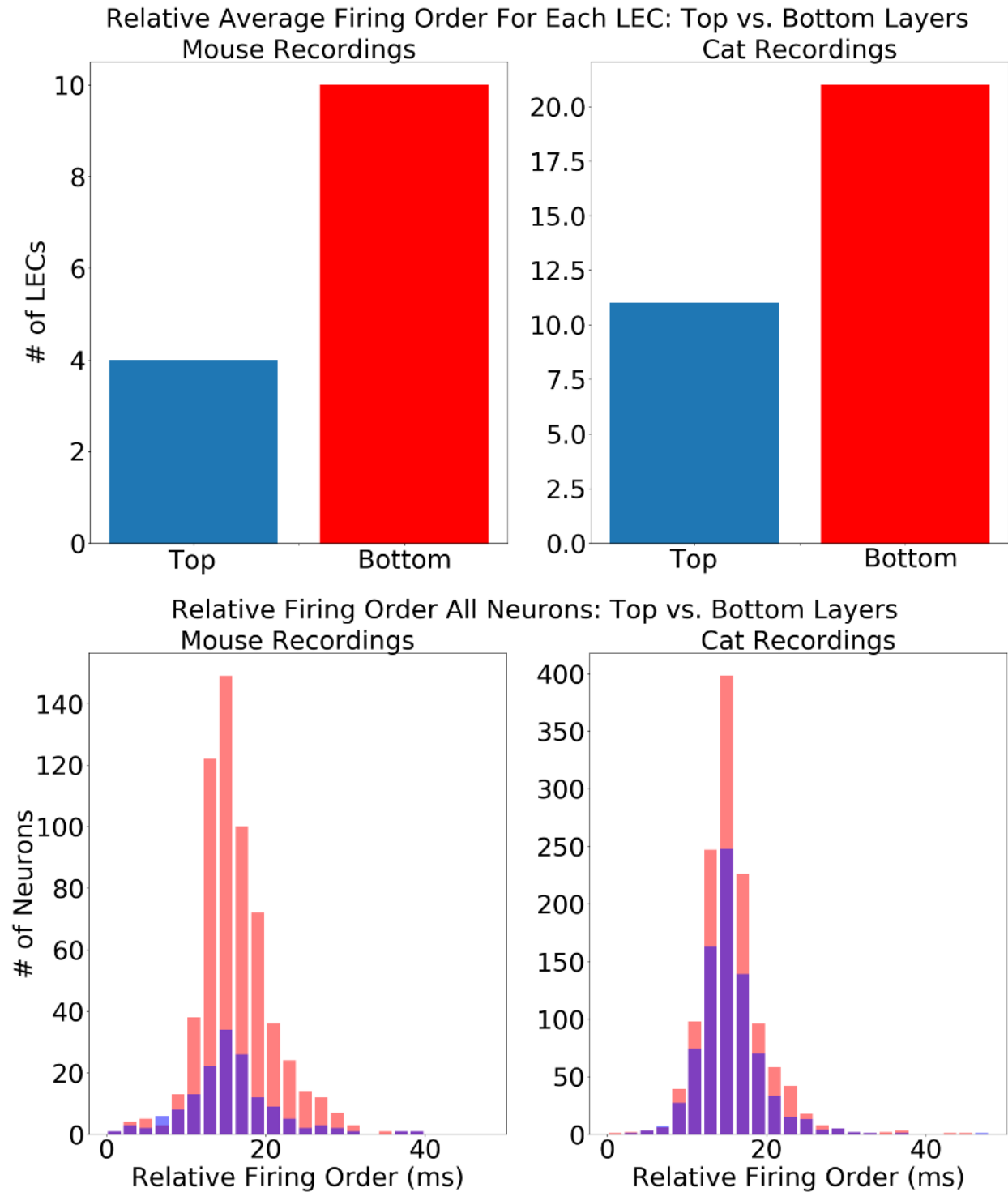


Figure 6-2

Table 1: Cat Visual Cortex – Recording Summary

Rec ID	Anesthetic	Track ID	Synchronized Rec Time <sup>1</sup> /Total Rec Time	No. of LECs
C1.1	Iso/N20	1	0.6 / 13.4hrs <sup>2</sup>	2
C1.2	Iso/N20	2	1.4 / 9.4hrs <sup>2</sup>	2
C2.1	Iso/N20	1	1.7 / 11.1hrs <sup>2</sup>	3
C2.2	Iso/N20	2	4.7 / 12.0hrs <sup>2</sup>	1
C3.1	Iso/N20	1	1.4 / 8.4hrs	1
C3.2	Iso/N20	2	4.2 / 8.3hrs	3
C3.3	Iso/N20	3	3.4 / 7.4hrs	2
C3.4	Iso/N20	4	3.0 / 5.3hrs	1
C4.1	Prop/Fent	1	3.3 / 11.9hrs	1
C4.2	Prop/Fent	2	0.5 / 11.5hrs	2 <sup>3</sup>
C4.3	Prop/Fent	3	0.9 / 4.6hrs	2
C5.1	Prop/Fent	1	2.5 / 2.9hrs	3
C5.2	Prop/Fent	2	4.9 / 10.6hrs	3
C5.3	Prop/Fent	3	3.1 / 8.5hrs	4
C5.4	Prop/Fent	4	4.6 / 6.3hrs <sup>2</sup>	4
Totals		15		34

1 – Synchronized recording time was defined as periods with synchrony index > 0.5.

2 – Synchrony index of 0.3 used.

3 – LECs had peak-to-peak amplitude and CSD maxima below threshold and were excluded from CSD grouping and further analysis.

Table 2: Mouse Cortex – Recording Summary

Rec ID	Anesthetic	Track ID	Area	Imaging	Sync Rec Time /Total Rec Time	No. of LECs
MV1	Isoflurane	1	Visual	VSD	2.6 / 3.1hrs	2
MV2	Isoflurane	1	Visual	VSD	2.2 / 2.2 hrs	1
MV3	Isoflurane	1	Visual	No	2.5 / 2.5 hrs	2
MV4	Isoflurane	1	Visual	GCaMP6s	1.0 / 1.0	1
MA1	Isoflurane	1	Auditory	VSD	2.1 / 2.1 hrs	1
MA2	Isoflurane	1	Auditory	GCaMP6s	3.1 / 3.1 hrs	1
MA3	Isoflurane	1	Auditory	No	2.0 / 2.0 hrs	1
MB1	Isoflurane	1	Barrel	GCaMP6s	2.7 / 2.7 hrs	1
Totals		8				9

Table 3: Single Unit Sorting Summary

Species	Ave. Track / Length( $\pm$ std)	Neuron Yield per Track( $\pm$ std)	Ave. Spikes per Unit( $\pm$ std)	Median Firing Rates	% Neurons Firing <2Hz
Cat (anesth)	8.8 $\pm$ 3.0hrs	99 $\pm$ 38	31807	0.31Hz	86.00%
Mouse (anesth)	2.5 $\pm$ 0.4hrs	85 $\pm$ 29	11974	0.26Hz	89.00%

Table 4: Cat V1 – LEC Groupings Summary

Rec ID	Anesthetic	#1	#2	#3	#4	#5	#6	Other
C1.1	IsoN20		X					X
C1.2	IsoN20	X	X					
C2.1	IsoN20	X	X					
C2.2	IsoN20						X	
C3.1	IsoN20	X						
C3.2	IsoN20	X	X	X				
C3.3	IsoN20	X				X		
C3.4	IsoN20				X			
C4.1	Prop/Fent	X	X					
C4.2	Prop/Fent						X	X
C5.1	Prop/Fent	X	X	X				
C5.2	Prop/Fent	X	X	X				
C5.3	Prop/Fent	X	X	X				X
C5.4	Prop/Fent	X	X	X				
Totals		10	9	5	1	1	2	3

1 – These 2 LECs are similar to Class 6 but may form separate class.

Table 5: Mouse Cortex – LEC Groupings Summary

Rec ID	Anesthetic	Area	#1	#2	#3	#4	#5	#6	#7	#8
MV1	Isoflurane	Visual	X	X						
MV2	Isoflurane	Visual	X							
MV3	Isoflurane	Visual	X		X	X				
MV4	Isoflurane	Visual	X			X				
MA1	Isoflurane	Auditory					X	X		
MA2	Isoflurane	Auditory						X		
MA3	Isoflurane	Auditory							X	
MB1	Isoflurane	Barrel								X
Totals	8		4	1	1	2	1	2	1	1

Table 6: Cat V1 – LEC Firing Rates (Hz)

Rec ID	Anesthetic	#1	#2	#3	#4	#5	#6	Other
C1.1	IsoN20		0.10					0.02
C1.2	IsoN20	0.04	0.52					
C2.1	IsoN20	0.05	0.11					
C2.2	IsoN20						0.25	
C3.1	IsoN20	0.16						
C3.2	IsoN20	0.08	0.08	0.35				
C3.3	IsoN20	0.04				0.32		
C3.4	IsoN20				0.03			
C4.1	Prop/Fent	0.29	1.4					
C4.2	Prop/Fent						0.4	0.90
C5.1	Prop/Fent	0.31	0.04	1.74				
C5.2	Prop/Fent	0.25	1.11	0.84				
C5.3	Prop/Fent	0.32	0.24	1.68				0.05
C5.4	Prop/Fent	0.21	0.28	2.57				



Table 7: Mouse Cortex – LEC Firing Rates (Hz)

Rec ID	Anesthetic	Area	#1	#2	#3	#4	#5	#6	#7	#8
MV1	Isoflurane	Visual	0.07	0.09						
MV2	Isoflurane	Visual	0.07							
MV3	Isoflurane	Visual	0.09		0.32	0.05				
MV4	Isoflurane	Visual	0.10			0.01				
MA1	Isoflurane	Auditory					0.06	0.01		
MA2	Isoflurane	Auditory						0.09		
MA3	Isoflurane	Auditory							0.37	
MB1	Isoflurane	Barrel								0.03
Totals	8		4	1	1	2	1	2	1	1