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The Contribution of Raised Intraneuronal Chloride to Epileptic Network Activity

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Altered inhibitory function is an important facet of epileptic pathology. A key concept is that GABAergic activity can become excitatory if intraneuronal chloride rises. However, it has proved difficult to separate the role of raised chloride from other contributory factors in complex network phenomena, such as epileptic pathology. Therefore, we asked what patterns of activity are associated with chloride dysregulation by making novel use of HaloHodopsin to load clusters of mouse pyramidal cells artificially with Cl⁻. Brief (1–10 s) activation of HaloHodopsin caused substantial positive shifts in the GABAergic reversal potential that were proportional to the charge transfer during the illumination and in adult neocortical pyramidal neurons decayed with a time constant of τ = 8.0 ± 2.8 s. At the network level, these positive shifts in E_GAIN produced a transient rise in network excitability, with many distinctive features of epileptic foci, including high-frequency oscillations with evidence of out-of-phase firing (Ibarz et al., 2010). We show how such firing patterns can arise from quite small shifts in the mean intracellular Cl⁻ level, within heterogeneous neuronal populations. Notably, however, chloride loading by itself did not trigger full ictal events, even with additional electrical stimulation to the underlying white matter. In contrast, when performed in combination with low, subepileptic levels of 4-aminopyridine, HaloHodopsin activation rapidly induced full ictal activity. These results suggest that chloride loading has at most an adjunctive role in ictogenesis. Our simulations also show how chloride loading can affect the jitter of action potential timing associated with imminent recruitment to an ictal event (Netoff and Schiff, 2002).

Key words: chloride; fast spiking interneuron; high frequency oscillations; neocortex; pyramidal neuron; seizure

Introduction

Inhibitory dysfunction has long been considered to be a major factor in triggering epileptic seizures (Miles and Wong, 1983; Sloviter, 1987; Traub and Miles, 1991; Cossart et al., 2005; Pinto et al., 2005; Huberfeld et al., 2007; Zsiros and Maccaferri, 2008; Kaila et al., 2014; Pallud et al., 2014). In recent years, particular attention has focused on how positive shifts in the GABAergic reversal potential, caused by raised intracellular Cl⁻ levels in neurons, may contribute to this process. Recordings from human brain slices, resected to treat epilepsy, show evidence of excitatory GABAergic activity (Cohen et al., 2002; Pallud et al., 2014) associated with reduced expression of the potassium chloride cotransporter KCC2 (Huberfeld et al., 2007). Even without deficits in KCC2 expression, Cl⁻ levels may rise acutely during intense GABAergic activation (Alger and Nicoll, 1979; Fujiwara-Tsukamoto et al., 2003; Isomura et al., 2003; Kaila, 1994; Kaila et al., 1997; Staley et al., 1995; Thompson and Gähwiler, 1989a,b,c) or be influenced by the distribution of other permeant and impermeant anions (Glykys et al., 2014). Collectively, these suggest an important role for chloride dysregulation in epileptogenesis, but it has not actually been shown whether raising intracellular Cl⁻ generally within the neuronal population is sufficient by itself to trigger epileptiform events or what patterns of activity are associated with such Cl⁻ loading.

The largest group of cortical neurons is the pyramidal population, representing ~80% of all cortical neurons, and the timing of their firing is influenced strongly by the output of basket cells (Cobb et al., 1995). In epileptic cortical networks, there are particularly intense bursts of interneuronal activity that appear to provide a restraint on the propagation of epileptiform discharges (Prince and Wilder, 1967; Trevelyan et al., 2006; Schevon et al., 2012). We reasoned that chloride loading in the pyramidal population would strongly influence the pattern of any breakthrough firing during the high-frequency inhibitory volley. In particular, we asked whether it might explain certain distinctive high-frequency features (Foffani et al., 2007; Ibarz et al., 2010) and firing patterns (Netoff and Schiff, 2002) associated with epileptic
foci (Bragin et al., 2002a,b; Staba et al., 2002; Jiruska et al., 2010) and the onset of ictal activity. To test this, we made novel use of the enhanced optogenetic chloride pump derived from Natromonas, Halorhodospin (eNpHR; Gradinaru et al., 2008) to artificially load pyramidal cells with chloride (Zhang et al., 2007; Raimondo et al., 2012) to examine the specific contribution of chloride dysregulation to epileptiform activity.

Materials and Methods

**Dissociated neuronal culture recordings.** All procedures were performed according to the requirements of the United Kingdom Animals (Scientific Procedures) Act 1986. Assessment of the chloride-loading effect of eNpHR activation was performed using dissociated neuronal cultures. Primary neuronal cultures were prepared from rat pups at embryonic days 18–20 in the following way. A pregnant rat was killed by cervical dislocation, and a sagittal incision was made in the abdominal area to remove the pups. The neocortex and hippocampal tissue was isolated from the pups and digested using papain enzyme (Sigma-Aldrich) for 40 min. Cells were then dissociated into the growing medium (Neurobasal A, 2% B-27 supplement, 1% FBS, 0.5% glutamate, and 0.5% antibiotic–antimycotic; Invitrogen) by pipetting it up and down. Cells were plated (10^5 cells/ml) on coverslips coated with poly-l-lysine and laminin (Sigma-Aldrich). Cells were allowed to attach to the coverslips by incubating at 37°C for 2.5 h. Cells were transfected using virus [human synapsin–eNpHR3.0–enhanced yellow fluorescent protein (EYFP)] at 7 d in vitro, and experiments were conducted on 14–28 d in vitro.

**Optogenetic expression.** All animal handling and experimentation were done according to United Kingdom Home Office guidelines. Optogenetic proteins were expressed on pyramidal cells by doing injection of viral vectors or breeding. To achieve pyramidal expression of eNpHR, we injected adeno-associated virus (AAV–CaMKII–Cre–GFP (University of North Carolina Viral Vector Core) into homoyzogous floxed–eNpHR animals [Ai39; B6:129–Gt(Rosa)26Sortm1(CAGRO/EYFP)H2/H]; stock #014539 The Jackson Laboratory]. For pyramidal expression of Archaeorhodopsin (Arch), we injected AAV–CaMKII–ArChT30–EYFP (University of North Carolina Viral Vector Core) into wild-type C57BL/6J mice. Injections were made into either postnatal day 0–1 pups or young adult mice. The pups had EMLA cream (2.5% lidocaine and 2.5% prilocaine) applied to the left top of their head and were anesthetized subsequently using isoflurane inhalation. A single injection of virus was made using a 10 μl Hamilton syringe, with a beveled 36 gauge needle (World Precision Instruments) ~1 mm anterior to the lamda and 1 mm lateral to the midline into the left hemisphere, at 1.7–0.8 mm deep to the skin (four injections per hemisphere). Following injection, mice were terminally anesthetized by brief inhalation of isoflurane (33–36°C) perfused with this same ACSF. The concentration of divalent cations was used following the study by Sanchez-Vivez and McCormick (1999), because this allowed the reliable triggering of sustained bursts of activity, with a prominent high-frequency discharge of fast-spiking interneurons, by white matter stimulation (putative thalamic inputs).

**Perforated patch-clamp recordings.** Recordings were made using a laser spinning disc confocal microscope (Visitech) and Digidata acquisition boards connected to desktop computers (Dell Computer Company) running pClamp software (Molecular Devices). During the entire recording, cells were bathed in circulating oxygenated ACSF solution (perfusion at 1–3 ml/min), heated to 33–37°C by a sleeve warmer, and perfused with the ACSF for at least 1 h before being transferred to a recording interface chamber (33–36°C) perfused with the same ACSF. The concentration of divalent cations was used following the study by Sanchez-Vivez and McCormick (1999), because this allowed the reliable triggering of sustained bursts of activity, with a prominent high-frequency discharge of fast-spiking interneurons, by white matter stimulation (putative thalamic inputs).

**Fluorescence immunocytochemistry.** Adult male HaloGFP/CaMKIICre mice were terminally anesthetized by brief inhalation of isoflurane (0.05% in air), followed by an intramuscular injection of ketamine (≥100 mg/kg) and xylazine (≥10 mg/kg). Once anesthetized mice were perfused through the heart with 4% (w/v) paraformaldehyde (PFA) in PBS (0.1 M), pH 7.2. After perfusion, brains were removed and postfixed in 4% PFA in PBS overnight at 4°C. The brain was cut coronally (50-μm-thick sections) using a Leica VT1000S vibratome. Sections were collected in PBS and then incubated in 500 mM ammonium chloride in PBS for 20 min at room temperature. After washing in PBS, sections were incubated in 0.1% (w/v) gelatin and 0.1% (v/v) Triton X-100 in PBS. Sections were double labeled with antibodies to eNpHR (rabbit anti-Halorhodospin at 1:2000; catalogue #AS12 1851; Agrisera) and interneuronal markers (sheep anti-neuropeptide Y at 1:1000, catalogue #AB1583, Millipore; goat anti-parvalbumin at 1:1000, catalogue #PVG214, Swant; goat anti-somatostatin at 1:250, catalogue #sc7819, Santa Cruz Biotechnology). The antibodies were diluted in 0.1% (w/v) gelatin and 0.1% (v/v) Triton X-100 in PBS and incubated at 4°C for 48–72 h. After extensive washing with PBS, sections were incubated with fluorescein isothiocyanate donkey anti-rabbit and Cy3 donkey anti-sheep and anti-goat secondary antibodies (Jackson ImmunoResearch) at a dilution of 1:500 in PBS for 2 h at room temperature. Finally, sections were washed extensively in PBS and mounted in VECTASHIELD HardSet mounting medium (Vector Laboratories). Control experiments, in which the primary antibodies were omitted, resulted in no immunoreactivity. Double-labeling images were obtained by using a Leica TCS SP2 UV confocal microscope. For analysis, optical slices were reconstructed by using NIH Image. The photomicrographs used in the figures were produced by first generating an extended depth of field projection of the z-stack by using NIH Image. The brightness and contrast of each image were then optimized, and multipanel figures were composed and labeled by using Adobe Photoshop and Adobe Illustrator (Adobe Systems).

**Patch-Clamp recordings.** Cerebral brain slices (250-μm-thick slices for perforated patch recordings and analysis of cellular excitability: 400 μm for field recordings) were prepared from the injected animals (3–10 months) on ice-cold oxygenated (95% O2/5% CO2) artificial CSF (ACSF; in mM: 125 NaCl, 26 NaHCO3, 10 glucose, 3.5 KCl, 1.26 Na2HPO4, 3 MgCl2, 1 Na-kynurenate, and 0.3 Na-ascorbate). For the perforated patch recordings, slices were simply transferred to a submerged incubation chamber for at least 1 h before recording. For the field recordings, after cutting, the slices were washed twice for 10 min with the oxygenated ACSF (in mM: 125 NaCl, 26 NaHCO3, 10 glucose, 3.5 KCl, 1.26 Na2HPO4, 1.2 CaCl2, and 1 MgCl2) and transferred to an incubation, interface chamber (room temperature) perfused with the same ACSF for at least 1 h before being transferred to a recording interface chamber (33–36°C) perfused with the same ACSF. The concentration of divalent cations was used following the study by Sanchez-Vivez and McCormick (1999), because this allowed the reliable triggering of sustained bursts of activity, with a prominent high-frequency discharge of fast-spiking interneurons, by white matter stimulation (putative thalamic inputs).
pipette coupled to a Picospritzer II (Parker Instrumentation) delivering the illumination (optogenetic activation) using pClamp software coupled to a Digitizer box. Test voltage ramps (200 ms duration, a saw-tooth, up-down function; peak, −50 mV; trough, −90 mV; slope, ±400 mV/s) were applied at baseline and also at close to the peak of GABAergic current (\( I_{\text{GABA}} \)). The GABAergic current, \( I_{\text{GABA}} \), was derived from the difference between the ramp currents to allow a GABAergic current–voltage (\( I-V \)) curve to be plotted. \( I-V \) plots were derived with and without previous eNhPR activation, and the effect of eNhPR activation on \( E_{\text{GABA}} \) was estimated from the shift in the \( x \)-intercept of the \( I-V \) plot. The time course of the chloride loading was measured by doing 2.8 s eNhPR activation, followed by muscimol puffs with increasing delay. The change in \( E_{\text{GABA}} \) was calculated for each time delay and normalized to the value for the shortest delay, and a single exponential curve was fitted to the normalized data to give the time constant for chloride clearance.

**Recovery of cellular properties after optogenetic activation.** Whole-cell patch-clamp recordings were made by direct visualization of neocortical pyramidal neurons (differential interference contrast microscopy). Cells were recorded in current-clamp mode (Multiclamp 700B and 1440 Digi-data analog-to-digital board), and brief hyperpolarizing and depolarizing current pulses were applied before, during, and immediately after activation of eNhPR current (orange bar). D. Phase plots of the rate of voltage change (dV/dt) versus the voltage for AP doublets immediately before (red) and within 1 s after a 5 s eNhPR activation (black), illustrating that eNhPR activation had no lasting effect on the AP threshold or shape. E. Measures of input resistance (\( R_i \)) normalized to the pre-optogenetic activation value, showing a pronounced drop after the optogenetic activation but a rapid recovery. The post-optogenetic measures were taken within 1 s of the end of the illumination period.

**Figure 1.** eNhPR chloride-loading effect. A. **Ai**, \( E_{\text{GABA}} \) was measured by doing perforated gramicidin patch recording and applying muscimol with or without previous eNhPR activation. **Ai**, Sample traces from a single neuron, derived from different duration eNhPR activations (n cells) and five data points from five neurons (red) in adult brain slices, all from 2 s illumination. **Bii**, The recovery of \( E_{\text{GABA}} \) after eNhPR activation. **Bi**, Sample traces from one of the neurons recorded in a brain slice, showing the response to muscimol puffs at progressively longer latencies after a 2 s eNhPR activation. A single-exponential curve (black line) is fitted to the minima of the voltage-ramp responses. **Bi**, Pooled data showing the recovery of \( E_{\text{GABA}} \) after eNhPR activation in dissociated neurons (black) and neurons recorded in adult brain slices (red). Single-exponential fits of the mean data are shown, but note that the time constants reported in Results are the averages of fits made to each individual cell. C. Example trace showing the excitability, in response to somatic charge injection, of a layer 5 pyramidal cells recorded in an adult brain slice, before, during, and immediately after activation of eNhPR current (orange bar).
shown that a more stable optogenetic current could be obtained by coillumination of both 561 and 491 nm light, thought to be attributable to improved \textit{cis-to-trans} recovery of the activated retinal by blue light (Han and Boyden, 2007). Therefore, we provided coillumination of the brain slices, using the 561 nm laser light with blue epifluorescence light (460 \pm 20 nm excitation filter) through a 4\x air objective (0.28 numerical aperture; Nikon). We confirmed that this coillumination strategy provided an enhanced and sustained eNpHR current by assessing the suppressive effect on electrical stimulation at different times during a long illumination period.

\textbf{Spike-timing analyses.} Frequency band analysis and all the spectrograms were done on the raw data, whereas for single-unit and multunit analyses, bandpass filter of 300 – 5000 Hz was applied and spikes were detected by a simple thresholding algorithm. Spike analyses were only performed for periods when the tissue was not illuminated (that is to say, not during the periods of optogenetic activation). To analyze spike timing according to the dominant oscillation, detected spikes were plotted on the Hilbert transform of the corresponding dominant oscillation in the 75–300 Hz bandpass. Angle and linear histograms were derived from the Hilbert plot. Baseline data were centralized and fitted to a Gaussian curve to obtain the baseline half-width. To assess the AP jitter and out-of-phase firing across multiple experiments, we derived a “half-width index,” which was calculated using the following formula:

$$\text{Half-width index} = \frac{|\text{spikes outside baseline HW}|}{|\text{spikes within baseline HW}|}$$

where HW is the half-width measured on the baseline period. Note that the baseline HW is used for all analyses for a single experiment, including for the optogenetically primed datasets that have a different width modal index, which was calculated using the following formula:

$$\text{Half-width index} = \frac{|\text{spikes outside baseline HW}|}{|\text{spikes within baseline HW}|}$$

where HW is the half-width measured on the baseline period. Note that the baseline HW is used for all analyses for a single experiment, including for the optogenetically primed datasets that have a different width modal histogram peak.

\textbf{Computer simulations of firing patterns.} Modeling was performed to provide a simple, intuitive illustration of the phenomenon we describe in our experimental studies, showing the consequence of changing $E_{\text{GABA}}$ in a conductance-based model neuron. All the simulations presented here were run on personal computers (Dell Computer Company) using the NEURON simulation program (Hines and Carnevale, 2001). We will provide the main model files on request but will describe their features below.

All simulations were run on simple four-compartment models possessing a soma, an axon, and a single dendrite comprising a short proximal compartment and a long distal one. The passive properties were homogeneous throughout the cell (membrane capacitance, 1 $\mu$F/cm$^2$; axial resistivity, 160 $\Omega$ cm; leak conductance ($g_{\text{leak}}$), 0.66 pS/µm$^2$; $g_{\text{leak}}$ reversal potential, −80.6 mV; resting membrane potential, −76 mV). The input resistance in the passive structure (also zero synaptic conductance) was relatively high (≈1 GΩ), primarily because of the relatively small structure of the cell, but the membrane time constant (passive $\tau_m$ = 27 ms) was consistent with that measured in layer 2/3 pyramidal cells at physiological temperature (Trevelyan and Jack, 2002). All model cells had the same somatic and axonal active conductances: a Hodgkin–Huxley-type Na$^+$ conductance (peak conductance, 2000 pS/µm$^2$) and two non-inactivating K$^+$ conductances (peak conductance, 5 and 3.5 pS/µm$^2$, respectively).

Ten excitatory synaptic conductances ($E_{\text{syn}} = 0$ mV) were located on the distal compartment, spaced equally along its length, providing a persistent, noisy excitatory drive of mean 6.4 pS/µm$^2$ (calculated over the entire distal dendritic compartment) with a variance of 10%. This noisy excitatory drive was implemented as described by Destexhe et al. (2001) and reduced the input resistance to −170 MΩ. In the absence of inhibition, this generated a high rate of spiking, with no periodicity (see Fig. 4A). Threshold for AP generation was −57.7 mV.

Inhibitory synapses were located only on the proximal dendritic tree to simulate the powerful basket cell inputs that are known to dictate pyramidal spike timing (Cobb et al., 1995). Because our intention was only to explore the constraint of pyramidal firing by basket cells, we did not include other inhibitory synapses in our models. Basket cells were set to fire at 100 Hz and deliver a steady amplitude train of postsynaptic conductance events. Each synaptic event was modeled as a transient conductance, described by a rising time constant ($\tau_{\text{rise}}$ = 0.2 ms) and a decay time constant ($\tau_{\text{decay}}$ = 2.5 ms) as follows:

$$g = W_0 \exp(-t/\tau_{\text{decay}}) - \exp(-t/\tau_{\text{rise}}),$$

where g is the conductance, and W is the synaptic weight, giving a peak conductance of 15.9 pS/µm$^2$ over the whole proximal dendritic compartment. We ran separate simulations for a range of $E_{\text{GABA}}$ values from −45 to −85 mV. At 100 Hz, and the mean conductance was 4.86 pS/µm$^2$ across the whole compartment. Consistent with experimental models, our simulations showed that this high-frequency proximal dendritic inhibition imposed very clear phase relation on the times of pyramidal firing for all values of $E_{\text{GABA}}$, with respect to the rhythm of the basket cell firing. We will make the NEURON code available freely on request.
confirming previous work by Raimondo et al. (2012). The shift in $E_{GABA}$ decayed back to baseline levels with a time constant, $\tau_{\text{decay}}$, of 28.12 ± 9.26 s ($n = 14$; Fig. 1B). We repeated these experiments also on pyramidal cells in brain slices prepared from adult mice (5 months) and showed a qualitatively similar effect ($\tau_{\text{decay}} = 8.0 \pm 2.8$ s; $n = 5$). Notably, both groups showed large variance in the calculated time constants, indicating some degree of heterogeneity with respect to this cellular behavior, and consequently, although the averages suggested that the decay was faster in brain slices, the difference from the cultured neurons was not significant in our samples ($t = 1.36; 0.2 < p < 0.3$).

Importantly, other cellular parameters affecting neuronal excitability returned rapidly to normal after the end of eNpHR activation. In whole-cell current-clamp recordings of pyramidal cells recorded in brain slices prepared from adult mice, we induced eNpHR activation, causing an average hyperpolarizing shift of 23.0 ± 1.5 mV, strongly suppressing firing driven by current injection and also inducing a 27 ± 8% drop in input resistance. Measurements taken within the first second after the end of eNpHR activation showed no difference from the preillumination measures for resting membrane potential (pre-eNpHR $E_m = -74.8 \pm 2.5$ mV; post-eNpHR $E_m = -75.0 \pm 2.9$ mV; $n = 4$ cells), input resistance (Fig. 1C; change from baseline, 4.4 ± 1.4%), and AP threshold [pre-eNpHR threshold (above $E_m$), 33.0 ± 1.2 mV; post-eNpHR threshold, 33.3 ± 1.2 mV; $n = 4$], height (peak − threshold: pre, 53.9 ± 4.4 mV; post, 52.9 ± 4.5 mV; $n = 4$), or shape (Fig. 1D). At this same time point, $E_{GABA}$ was shifted positively by 5.9 ± 1.0 mV in cultures ($n = 12$ cells) and 3.9 ± 0.6 mV in adult pyramidal cells ($n = 4$ cells) in acutely prepared brain slices (culture vs slice comparison: $t = 1.09; 0.4 < p < 0.2$, nonsignificant). In short, the only apparent cellular change persisting beyond 1 s was the change in $E_{GABA}$.

We made similar measurements after Arch activation. This optogenetic protein also hyperpolarizes neurons but by pumping protons out of cells. As with eNpHR activation, Arch activation was associated with a suppression of firing and a drop in input resistance (normalized to preillumination, $R_n = 61 \pm 20\%$, $n = 4$), but these both corrected rapidly within a second of ending illumination (post-illumination, normalized $R_n = 106 \pm 4\%$), as did baseline $E_m$ (pre, −71.1 mV; post, −70.6 mV; $n = 4$; $t = 0.03$, n.s.), AP threshold relative to baseline $E_m$ (pre, 32.1 ± 4.2 mV; post, 32.5 ± 4.3 mV; $n = 4$; $t = 0.02$, n.s.), and AP height (peak − threshold: pre, 56.2 ± 6.7 mV; post, 54.9 ± 7.0 mV; $n = 4$; $t = 0.13$, n.s.). However, unlike after eNpHR activation, Arch activation produced no change in $E_{GABA}$ (measured at −1 s after illumination; pre, −64.5 ± 5.1 mV; post, −64.7 ± 5.5 mV; $n = 3$; $t$
Figure 4. Neuronal chloride loading triggers out-of-phase firing during spontaneous bursts of activity. A. Spectrograms of extracellular recordings of spontaneous bursts of activity in baseline (left) and eNpHR-primed (chloride-loaded) tissue. Note the prominent "double-frequency" signal in the chloride-loaded tissue. Detected spikes (dots, red (early) to green (late) during spontaneous bursts in baseline (left) and eNpHR-primed (chloride-loaded) tissue. B. Arch priming. C. Baseline, Half-width index = 0.4. eNpHR-primed, Half-width index = 1.1. D. p<0.001 p<0.02.
Thus, eNpHR activation can non-invasively induce a transient loading of Cl\(^-\) into many neurons simultaneously through broad illumination of networks expressing the protein. Therefore, we used this technique to modulate intraneuronal Cl\(^-\) levels in pyramidal cells in brain slices to investigate what patterns of activity ensue when neurons become loaded with Cl\(^-\), as is thought to happen in epileptic pathology.

We recorded neocortical activity patterns in brain slices taken from animals expressing either eNpHR or Arch (Chow et al., 2010), both under the CaMKII\(\alpha\) promoter (Fig. 2). We used a methodology developed by Sanchez-Vives and McCormick (2000), in which the extracellular divalent cation concentration is lowered slightly (1.2 mM Ca\(^{2+}\), 1 mM Mg\(^{2+}\)), because this allowed the reliable triggering of sustained bursts of activity, with a prominent discharge of fast-spiking interneurons (Shu et al., 2003), by white matter stimulation (putative thalamic inputs). We first confirmed that activation of the opticogenetic probe, by illumination through an optic fiber placed adjacent to the recording electrode, could suppress substantially the electrically evoked network response (Fig. 2Aii). Then we changed the timing of electrical stimulation so that it was delivered after the opticogenetic activation to investigate the chloride-loaded state induced by eNpHR priming. Prolonged eNpHR activation might also induce rebound firing simply as a reaction to the hyperpolarization. Therefore, we repeated these experiments in brain slices expressing instead Arch, which is another hyperpolarizing opticogenetic tool that pumps protons out and which thus allows us to separate the rebound-loading effects from the rebound activation.

**Distinctive network activity induced by chloride loading**

Chloride loading the pyramidal population, by previous eNpHR activation (“eNpHR priming”), reliably induced a large increase in evoked network activity with a prominent signal in high-frequency bandwidth between 150 and 600 Hz. This effect reversed rapidly, within five trials (150 s), once eNpHR priming was stopped (Figs. 2B, C, 3A). In contrast, after Arch priming, there was no change in the high-frequency power from baseline trials, confirming that the eNpHR-primed induced high-frequency activity is Cl\(^-\)-dependent and not merely attributable to rebound firing caused by synchronized hyperpolarization of pyramidal cells (Figs. 2C, 3A,B). We performed more detailed analysis of the high-frequency components of the extracellular signal between 75 and 600 Hz. The eNpHR-primed tissue showed increases above that shown by Arch-primed tissue at all frequencies, but this was only significantly different for the 300–600 Hz band [Fig. 3A; first priming, eNpHR (4.13 ± 0.87, n = 11) vs Arch (1.44 ± 0.26, n = 11), p < 0.01; second priming, eNpHR (3.67 ± 1.09, n = 7) vs Arch (1.56 ± 0.31, n = 10), p < 0.05, Student’s t test]. The 600–1200 Hz signal showed virtually no change for either eNpHR- or Arch-primed tissue, indicating that the eNpHR-priming effect at 300–600 Hz is unlikely to arise simply from a signal harmonic process, because that would have also produced increases at higher-order harmonics, too.

Previous studies of high-frequency oscillations in epileptic rats suggested that they arise from out-of-phase firing between subpopulations of pyramidal cells (Ibarz et al., 2007; Ibarz et al., 2010). Therefore, we conducted analysis of spontaneous spiking in the periods between optogenetic activation (Fig. 4). Analysis was conducted on the periods after the end of the evoked burst of activity and before the subsequent optogenetic illumination (10 4.5-s epochs, 45 s total per slice). After eNpHR priming, there was a significant increase in the multiunit firing rate (note the different scale bars in Fig. 4B, baseline multiunit rate, 6.7 ± 2.5 spikes/s; eNpHR-primed, 38.3 ± 6.6 spikes/s; n = 8, p < 0.001; eNpHR-priming increase, normalized to baseline, 12.7 ± 4.8). In contrast, Arch-priming produced only a small, but nonsignificant, trend toward increased firing over baseline (Arch baseline multiunit rate, 6.5 ± 1.9 spikes/s; Arch-primed, 10.2 ± 2.6 spikes/s; n = 6; n.s.; Arch-priming increase, 2.2 ± 0.7).

In both baseline and eNpHR-primed periods, there were episodes of bursting activity that showed up as prominent bands in the spectrograms between 150 and 300 Hz (Fig. 4A), with an additional band at approximately double this frequency after eNpHR priming (Fig. 4A, right). We speculated that the lower-frequency bandwidth (<300 Hz) may be dictated by bursts of fast-spiking interneurons, which are known to fire in this range, and because the output of these neurons can entain pyramidal firing, we analyzed the timing of the multiunit activity with respect to the dominant frequency band between 75 and 300 Hz. We did this by performing a Hilbert transform on the 75–300 Hz band, which allows the oscillating signal to be plotted as a continuous circular trajectory, and then plotting histograms of the spike times with respect to this cycle (Fig. 4A). Spiking during the baseline period was virtually entirely confined to the lower left quadrant (eNpHR-expressing tissue: mean phase angle, 3.58 ± 0.11 Rad, n = 7; Arch-expressing tissue: mean phase angle, 3.24 ± 0.14 Rad, n = 6), which represented spikes occurring close to the trough of the local field potential. In contrast, after eNpHR priming, the spiking was far more intense, spike timing showed a marked increase in jitter with respect to the dominant high gamma oscillations in all slices, and, in three of eight slices, there was a biphase distribution, with a prominent peak in the histogram exactly out-of-phase with the main peak. This feature was unique to the eNpHR-priming experiments and did not occur with Arch priming (Fig. 4B).

To pool data from different experiments, we derived an out-of-phase index by fitting a Gaussian curve centered on the circular mean to define the half-width of the modal peak and then calculating the ratio of the number of spikes outside of this half-width limit to the number of spikes within it (Fig. 4C). This showed a highly significant increase in out-of-phase spiking in the eNpHR-primed tissue (half-width index, 1.35 ± 0.15, n = 8) compared with both baseline periods (0.56 ± 0.04, n = 8, p < 0.001) and also to the control Arch-primed tissue (0.60 ± 0.20, n = 6, p < 0.02; Fig. 4D).

An important consideration when analyzing such spike-phase relationships, especially when relating units and field recordings made in the same electrode, is spectral leak, which is when a very
A high-frequency event will create a small oscillation also in lower bandpass filtered traces. To separate the spectral leak from the spike-independent field oscillation, we therefore performed two control analyses to examine potential confounding effects of spectral leak. First, we performed amputations of the unit spikes in the raw data before performing the 75–300 Hz filtering (Fig. 5). This reduced the amplitudes of the peaks (Fig. 5A, contrast the blue and black lines), and the spike-phase distributions showed slightly broader peaks than for the normal signal, both indicating that there was indeed a small spectral leak effect, but importantly, the distribution remained strongly skewed. We further tested the confounding influence of spectral leak with a second more ex-

Figure 5. Control analyses to examine the effect of spectral leak. A, A raw data trace, showing the spikes (blue) and the amputated version (black). The bottom traces show the 75–300 Hz bandpass filtered traces. B, 75–300 Hz bandpass filtered traces of the raw (blue) and the amputated traces. Red dots indicate the spike times. Note the reduced amplitude peaks for the amputated spikes but that these still occur at the same phase of the oscillation, as evidence by the same skewed orientation of the pooled data shown in the spike-phase plots (right columns). C, The same filtered traces but showing the new locations of APs for the time-shifted analysis. The spike-phase plots showed a shifted phase, reflecting the time shift, and a broader main peak, but importantly, they were still heavily skewed for both the raw and amputated data.

Figure 6. Spike-phase relationships were preserved for three different control analyses for spectral leak effects. A, Example phase histograms showing the shifted spike-phase distributions for the three control paradigms illustrated in Figure 5, but note that the key features, with baseline spike-phase plots having single peaks, and eNpHR-primed plots having double peaks, are maintained. B, Pooled data showing that there were highly significant differences between the eNpHR-primed and Arch-primed matched analyses for all control paradigms. *p < 0.05; **p < 0.01; ***p < 0.001. Halo, Halorhodopsin.
treme test by examining the phase relationship at a time ahead of the spikes. Our reasoning was that, if the spikes were indeed embedded within genuine oscillations, then these oscillations would extend sufficiently far in front and after the spike that the relationship would still exist for time-shifted points. We chose a forward time shift to avoid any postsynaptic influences. We tested increasing $\Delta t$ until the amputated and non-amputated spike-phase plots were identical, indicating a time point beyond the effect of the spikes. The $\Delta t$ was specific for each trace and ranged between 1.3 and 1.8 ms ahead of the spike. Importantly, for all three analyses [(1) amputated; (2) time-shifted; and (3) both amputated and shifted data], the baseline and Arch-primed datasets produced single-peak phase distribu-

Figure 7. Heterogeneity in levels of intracellular $\text{Cl}^-$ can explain the appearance of out-of-phase population firing. A, Simulation using NEURON of how a train of high-frequency IPSCs from a fast-spiking interneuron superimposed onto a noisy, desynchronized glutamatergic drive creates a patterned firing in the pyramidal cell. AP threshold was approximately $-48 \text{ mV}$. B, Simulations in the same model, at four different GABAergic reversal potentials. The firing probability is plotted with respect to the field oscillation, which is approximately $\pi/4$ phase shifted from the start of the IPSC, as judged by comparisons with the timing of fast-spiking interneuron APs (Hasenstaub et al., 2005). These different probability histograms are convolved with estimates of the distribution of $E_{\text{GABA}}$ in the pyramidal population (middle, black; the bottom histogram is taken from the study of Huberfeld et al. (2007), mean $E_{\text{GABA}} = -64 \text{ mV}$; the top is a simulated, normal distribution shifted to a slightly more hyperpolarized mean $E_{\text{GABA}} = -68 \text{ mV}$). The convolution is achieved by assuming that the various bins show the firing phase relations in the left plots, to yield estimates of the population firing for the “normal” and the “Huberfeld” populations.
tions, whereas the eNpHR-primed datasets produced double peaks (Fig. 6). We derived half-width indices for all three analyses, which all showed highly significant differences between eNpHR-primed and baseline periods and also between the eNpHR-primed and Arch-primed data (Fig. 6). Therefore, we concluded that, although there is indeed a small but demonstrable spectral leak effect, this could not explain the distinctive spike-phase plots found in the different experiments.

Activity patterns explained by heterogeneity within the Cl−-loaded pyramidal population

We hypothesized that a mechanism involving fast-spiking interneurons, which are known to influence the timing of pyramidal neurons and are also thought to discharge at high rates in response to surges in network activity (Cammarota et al., 2013), might explain these spiking patterns. In particular, we investigated how the influence of a high frequency inhibitory barrage might be distorted by changing the intracellular Cl− levels in the postsynaptic population, using compartmental neuronal modeling (Hines and Carnevale, 2001). We simulated an intense, desynchronized glutamatergic drive on to a pyramidal cell and then further delivered a high-frequency barrage of IPSCs on to the soma and proximal dendrites; Fig. 7A shows repeated simulations of postsynaptic spiking for a cell with a relatively hyperpolarizing ECl− of 68.5 mV, which are collated into spiking phase histograms based on the cycle of IPSCs. We next simulated how the effect of this same inhibitory barrage changed with ECABAr shifting toward more positive levels (Fig. 7B) to show that the window of opportunity for pyramidal spiking broadens as the effective inhibition diminishes (ECl− shifting from −75 to −65 to −55 mV). Eventually, when ECABAr exceeds AP threshold (more than −48 mV), there is a sudden 180° (π) phase shift in the spiking.

Previous studies have shown that the output of fast-spiking interneurons is synchronized precisely on to the multiple postsynaptic pyramidal cells (Miles et al., 1996; Trevelyan, 2009). If we consider that this postsynaptic population has a distribution of ECABAr values, as measured by Huberfeld et al. (2007) and also shown using Cl− imaging (Dzhala et al., 2010), we can use the simulations of firing patterns for the single cell at different ECABAr levels to simulate the multiunit spiking patterns in this heterogenous population. We derived the population response using two different distributions of ECABAr values, one distribution as described in resected human epileptic hippocampi (Huberfeld et al., 2007) and a negatively shifted one as the physiological distribution. The physiologically shifted distribution (mean ECABAr = −68.5 mV; Fig. 7B, bottom right) showed a unimodal spiking distribution with respect to the rhythm imposed by the basket cells. In contrast, only a small positive shift in ECABAr (mean ECABAr = −60.8 mV; Fig. 7B, bottom right) allowed a marked increase in spiking because of the broadening of the main peak but also the appearance of a prominent out-of-phase peak, reflecting the activity of the subpopulation of neurons with pathologically high levels of Cl− and a correspondingly high ECABAr, in excess of AP threshold. These distributions reproduced very well the histograms drawn from different eNpHR-primed brain slices (Fig. 4A).

Cl− loading only triggered full ictal events in conjunction with other pathological activity

This model thus provides a coherent explanation of how our eNpHR-priming experiments can give rise to activity patterns that have also been described in epileptic animals (Foffani et al., 2007; Ibáñez et al., 2010). Therefore, it was a surprise that, in none of these experiments, either with eNpHR or Arch priming, did repeated electrical stimulation to the network actually trigger ictal-like events, with hypersynchronous, rhythmic discharges. We next repeated whether Cl− loading altered the seizure threshold for other treatments. We examined the 4-aminopyridine (4-AP) model, because this model is known to trigger intense bursts of firing in the fast-spiking interneuronal population (Cammarota et al., 2013), therefore reasoning that such activity may escalate toward ictal activity if their postsynaptic output were imposed on a population of pyramidal cells with raised intracellular Cl− levels. Epileptiform activity can be readily induced by bath application of 50–100 μM 4-AP in brain slices. When we used only 20 μM 4-AP, 4 of 20 slices showed epileptiform activity very quickly (<10 min), and this activity was not in any way modulated by subsequent eNpHR activation (Fig. 8). The majority of slices were quiescent (80%), even when bathed in 20 μM for >1 h. However, when these quiescent slices were then eNpHR primed, full ictal activity was very rapidly induced, within a few minutes,
in all but a single slice (15 of 16 slices; latency to the first full ictal event after start of eNpHR priming, 8.9 ± 2.7 cycles (30 s cycles of 25 s illumination/5 s dark); latency from the start of the first cycle, 268 ± 80 s, n = 15; Fig. 8B). The ictal activity generally started immediately after the light was turned off, without the need of electrical stimulation, and often persisted into the next illumination cycle, resisting the inhibitory action of the eNpHR (see the expanded example traces in Fig. 8A).

**Discussion**

We explored a key hypothesis in epilepsy: that chloride disregulation in neurons is a major factor in triggering seizures. Surprisingly, Cl \(^{-}\) loading by itself did not trigger full ictal activity, even when electrical stimulation was delivered to the network. However, when Cl \(^{-}\) loading was associated with other pathological activation, by bathing in low levels of 4-AP, it did rapidly induce ictal activity. Thus, we make a clear distinction between how Cl \(^{-}\) loading creates a primed brain state and the requirement for some adjunct pathology to actually trigger a seizure.

Previous animal work has suggested that intense bursts of GABAergic activity can themselves be a direct trigger of seizures, with the proposed mechanism being a positive shift in E\(_{\text{GABA}}\) (Bennard et al., 2000; Gnatskiv et al., 2008). An important component of the pathology may be that this pattern of inhibitory discharge can synchronize the postsynaptic population of pyramidal cells (Klaassen et al., 2006). These issues are explored further in several review articles (Menendez de la Prida and Trevelyan, 2011; Jiruska et al., 2013). Our results suggest that the combination of such intense interneuronal discharges together with a progressive shifting of E\(_{\text{GABA}}\) may be particularly ictogenic and also give rise to certain previously unexplained features of electrophysiological recordings immediately before the seizure onset.

Our findings have a clear parallel with recent studies of human brain tissue resected during epilepsy surgery (Cohen et al., 2002; Huberfeld et al., 2007; Pallud et al., 2014), which show spontaneously occurring interictal events when bathed in conventional ACSF. These interictal events are sensitive to GABAergic blockade, suggestive of a Cl \(^{-}\) -loaded, excitatory GABAergic state, but to trigger full ictal events in these slices required excitability to be further enhanced by bathing in raised K\(^{+}\). Importantly, the subsequent ictal activity appeared to arise out of a fundamentally different type of transient discharge that was not sensitive to GABAergic blockade (Pallud et al., 2014). In other words, as with our data, we can distinguish between interictal activity associated with Cl \(^{-}\) dysregulation and a second pattern of pathological activity that is independent of Cl \(^{-}\) dysregulation; the combination of these is associated with full ictal activation, but the first alone does not predispose to full ictal events.

We contrasted the immediate changes in network excitability after eNpHR priming versus Arch priming. We attempted to achieve an approximately equivalent suppressive effect of network activation by Arch and eNpHR by adjusting the illumination intensity. Of course this is rather inexact, but the key issue is that, in both optogenetic paradigms, we clearly achieved some measurable network suppression, and then our data collection focused on the residual, post-illumination effects, of which the change in E\(_{\text{GABA}}\) appeared to be the most persistent. After activation of both eNpHR and Arch, there was a rise in excitability, suggesting that rebound activation may contribute, but the effect was far larger for eNpHR priming. Importantly, there were other, highly distinctive changes in firing patterns induced by eNpHR priming that are well captured by a model of a heterogeneous Cl \(^{-}\) loading in the population of pyramidal cells, causing them to react differently to the same high-frequency, GABAergic, synaptic barrage. The eNpHR-priming network changes also reversed with a timeframe similar to the recovery of E\(_{\text{GABA}}\) measured in single cells. Collectively, this strongly suggests that the eNpHR-priming effect identifies unique features of network excitability attributable to Cl \(^{-}\) loading.

These changes in activity in the Cl \(^{-}\) -loaded tissue correspond well with activity patterns recorded in epileptic animals (Foffani et al., 2007; Ibarz et al., 2010) in that both show particular high-frequency field oscillations with an apparent harmonic feature. The activity in the epileptic animals has been explained in terms of individual neurons firing at lower rates but with different subpopulations of neurons firing out-of-phase with each other. Our Cl \(^{-}\) -loading experiments support this view, and our model explains how this binary segregation could occur, depending on whether E\(_{\text{GABA}}\) is below or above the AP threshold in different cells. The important feature of this model is that there is a distribution of E\(_{\text{GABA}}\) values, and, in this situation, the eNpHR-priming effect can arise with relatively small (single figure millivolt changes) shifts in the mean E\(_{\text{GABA}}\).

There is an increasing body of evidence that links high-frequency oscillations to the focus of epileptic pathology in humans, too (Bragin et al., 2002b; Staba et al., 2002). There, of course, other mechanisms suggested for the origin of high-frequency oscillations, albeit without this “harmonic” feature. Any intense, focal activation of large numbers of neurons will generate a high-frequency signal, and there are several paradigms of epileptiform activity in which this occurs independent of any fast-spiking interneuron involvement. For instance, epileptic activity may arise from local loss of inhibition or even in preparations without synaptic function (Draguhn et al., 1998), in which ephaptic (Jiruska et al., 2010) or gap junction-mediated (Traub et al., 1999) spread has been implicated. However, our data provide the first evidence that there may be characteristic features of high-frequency activity that are pathognomonic for Cl \(^{-}\) dysregulation. It should further prompt us to look for other features that may also be used to subclassify pathological activity patterns, particularly if they also offer insights into the underlying pathology.

Our model also pertains to another long-time puzzle about epileptic spiking patterns, which is that, as the cortical network is recruited to a full ictal event, there is an increase in spiking jitter between neurons (Netoff and Schiff, 2002). This result had seemed to contradict the traditional concept of a progression toward hypersynchrony, but our data now offer an explanation, suggesting, instead, that the progressive trend is with the shift in E\(_{\text{GABA}}\) and a consequence of this is that the initial apparent effect is that the spiking window is made broader, before the critical stage is reached when E\(_{\text{GABA}}\) surpasses AP threshold.

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