Conjunctive input processing drives feature selectivity in hippocampal CA1 neurons

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Feature-selective firing allows networks to produce representations of the external and internal environments. Despite its importance, the mechanisms generating neuronal feature selectivity are incompletely understood. In many cortical microcircuits, the integration of functionally distinct inputs occurs nonlinearly through generation of active dendritic signals that drive burst firing and robust plasticity. To examine the role of this processing in feature selectivity, we recorded CA1 pyramidal neuron membrane potential and local field potential in mice running on a linear treadmill. We found that dendritic plateau potentials were produced by an interaction between properly timed input from entorhinal cortex and hippocampal CA3. These conjunctive signals positively modulated the firing of previously established place fields and rapidly induced new place field formation to produce feature selectivity in CA1 that is a function of both entorhinal cortex and CA3 input. Such selectivity could allow mixed network level representations that support context-dependent spatial maps.

The neuronal microcircuit is a fundamental processing unit of nervous systems and as such is intricately involved in producing the network computations that drive behavior. Many cortical microcircuits show a stereotyped organization that may allow them to implement a common set of core operations1. One such core operation is derived from the fact that most cortical microcircuits receive at least two functionally distinct afferent inputs from different brain areas1. In pyramidal neuron–based microcircuits, these different inputs are spatially segregated, with long-range inputs mainly innervating the most distal apical tuft dendrite regions and more local inputs usually forming synapses onto the proximal perisomatic dendrites2–5. The electrical isolation present in most pyramidal neurons means that each functional class of inputs is initially processed in relative independence of the other6–9. However, active dendritic mechanisms, such as backpropagating action potentials (APs) and dendritic Ca2+ plateau potentials, allow the different afferent input streams to interact nonlinearly when they arrive with appropriate coincident timing6–13. Additionally, the initiation of dendritic plateau potentials is known to robustly induce rapid and long-lasting changes in synaptic strength and dendritic excitability10,14. This form of input interaction or integration could support a nonlinear combination of the input streams to produce neurons with a feature selectivity that is a function of both input paths6,9. Such a population will, in turn, generate mixed or multimodal network representations with advantageous computational properties15,16.

In the CA1 region of the hippocampus, long-range external information from entorhinal cortex (EC) arrives onto the most distal dendritic regions of principal neurons while more local, internal information from hippocampal area CA3 arrives onto proximal perisomatic dendrites2,3,17,18. In vitro studies have shown that these inputs can be integrated supralinearly, with the exact level of amplification being dependent on the coincidence of the two input pathways. A voltage-gated Ca2+ channel and NMDA receptor-dependent dendritic plateau potential generated in the distal apical dendritic regions mediates this nonlinear form of dendritic integration9–11. Recently, several in vivo studies have provided strong evidence that dendritic plateau potentials and the complex burst firing they produce are present during both anesthetized and awake conditions13,19–21. Such active dendritic processing could mediate a nonlinear interaction between EC3 and CA3 inputs that allows the CA1 circuit to associate contextual and spatial information, thus generating and maintaining context-dependent spatial maps10,22–25.

RESULTS

Intracellular characterization of place-specific firing

To investigate dendritic plateau potentials in awake, behaving mice, we trained water-restricted mice to run head-fixed on a linear track treadmill with spatial cues for a 10% sucrose water reward delivered each trial (lap) at the same location on a ~2 m track (Fig. 1a; see Online Methods)26. Extracellular recordings under these conditions have demonstrated that CA1 pyramidal neurons fire at specific locations on the track and that these place cells have similar properties to those recorded from freely moving animals26. We obtained simultaneous whole-cell intracellular and local field potential (LFP) extracellular recordings in area CA1 of the dorsal hippocampus of navigating mice (Fig. 1b). The activity profiles of recorded neurons generally fell into three categories: silent cells, non-place cells and place cells (Supplementary Fig. 1; see Online Methods for criteria). At this point

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we will restrict our analysis to place cells, which exhibited location-specific AP firing rates that were similar to those measured using extracellular methods (mean peak rate, 14.3 ± 1.5 Hz, N = 22 neurons from 21 mice; Fig. 1 and Supplementary Fig. 2). In accord with previous intracellular recordings of place cells (19, 20), we observed a ramp-like depolarization in the subthreshold membrane potential (ramp) (mean ramp amplitude, 7.9 ± 1.0 mV, N = 22; Fig. 1 and Supplementary Fig. 2), an increase in the amplitude of intrinsic theta oscillation (thetain) (mean Δtheta, 2.8 ± 0.26 mV, N = 22; Fig. 1 and Supplementary Fig. 2) and phase precession of both APs and thetan in the preferred firing location (Supplementary Fig. 2). The entire population of place fields spanned the length of the belt (Fig. 1d).

Detection and properties of dendritic plateau–driven complex spiking and burst firing

We also observed bursts of APs that were driven by characteristic membrane potential (V_m) depolarizations that are the signature of dendritic plateau potential initiation (Fig. 1e–g and Supplementary Fig. 3). Plateau events were detected as a threshold crossing (~35 mV) of the smoothed intracellular voltage (Supplementary Fig. 3). The threshold value was determined from dual dendritic and somatic in vitro recordings with simultaneous calcium imaging (Supplementary Fig. 3f–j). The average duration of plateau events in vivo, measured as the time spent above threshold, was 56 ± 4 ms and the amplitude of these events from baseline V_m was 41 ± 0.3 mV (Fig. 1h, i, N = 17 cells, 226 events), similar to that recorded in vitro (Supplementary Fig. 3) and from anesthetized mice (21). We also found that the probability of plateau initiation varied between laps and was positively correlated with ramp amplitude, Δtheta and peak firing rate (Supplementary Fig. 4). NMDA receptor blockade using intracellular MK801 reduced plateau probability and duration, which is consistent with the pharmacology of plateau potentials recorded in vitro and in anesthetized mice (Supplementary Fig. 5).

Spatio-temporal properties of plateaus

To examine the microcircuit mechanisms of plateau potential initiation, we measured the extracellular theta phase preference of plateaus and APs in all place cells. In vitro studies predict that plateaus should be initiated when EC3 and CA3 input are coincidently active. Extracellular studies, however, have shown that EC3 and CA3 inputs tend to be active at different phases of theta, with the phase preference of EC3 being near the peak of theta while CA3 inputs are most active near the trough of extracellular theta (27, 28). Consistent with previous reports for CA1 cells, the phase preference of APs recorded here was near the trough of extracellular theta (173 ± 16°; 12,784 events in 21 neurons). The phase preference of plateaus, however, was more advanced than that of APs (56 ± 7°; 223 events in 16 neurons; two-tailed unpaired t-test, P = 1.8 × 10^{-6}; Fig. 2b), making their phase preference more similar to that previously reported for EC3 input. Plateaus also displayed a significantly higher level of theta modulation (APs, 0.43 ± 0.04; plateaus, 0.92 ± 0.03; two-tailed unpaired t-test, P = 4.9 × 10^{-4}). Interestingly, we observed a relationship between

Figure 1 Intracellular recordings of CA1 place cell firing. (a) Schematic of recording set up (left) and confocal stack of a biocytin filled cell (right). Scale bar, 50 μm. (b) Representative intracellular V_m (black) and extracellular theta (green) for laps 12–17 for a representative place cell (cell 3 in d). (c) AP firing rate for all single laps as a function of distance for cell shown in b. (d) Normalized firing rates of all place cells (N = 22 from 21 mice) ordered by place field position. (e, f) Representative extracellular theta (green), intracellular V_m (black), intracellular ramp (red) and thetan (blue) as a function of time for lap 12 (e) and lap 16 (f) for cell in b. (g) Representative complex spikes (plateaus) from traces shown in e and f. (h, i) Amplitude (h) and duration (i) distributions for all plateaus (N = 226 events from 22 neurons from 21 mice).
plateau probability and AP theta phase preference of individual neurons: plateau probability was higher in those cells showing the most advanced theta phase preference (Fig. 2c). Plateau durations also had a phasic relationship with the LFP, such that the duration of plateau events decreased as the theta phase of their initiation moved from the peak of extracellular theta to the trough, farther from the preferred phase of EC3 input (Fig. 2d). Together these results are consistent with previous in vitro data showing that dendritic Ca2+ plateaus occur when CA3 input is appropriately timed to drive AP output and back-propagation to coincide with EC3 input.

Experimental manipulation of perisomatic $V_m$ and AP firing phase

We further examined this temporal relationship by manipulating the phase of perisomatic $V_m$ fluctuations and resultant CA1 firing, thereby mimicking the impact of either advanced or delayed CA3 input. For this we used a closed loop experiment in which we online filtered and variably delayed the extracellular LFP and then used the resulting waveform to drive current injection into the recorded cell (Fig. 2 and Supplementary Fig. 6). In the first condition we minimized the delay to produce advanced $V_m$ fluctuations and spiking biased toward the peak of extracellular theta (AP peak near 45°; Fig. 2e–g). In the second condition we increased the delay on the injected current to produce later phase $V_m$ fluctuations, and the cells preferentially spiked near the trough of extracellular theta (~200°; ~95 ms delay; Fig. 2f–g). Plateau probability and normalized duration were significantly higher when spiking was biased toward the peak of extracellular theta (probability, 4.1 ± 1.2 plateaus per 100 APs; duration 183.2 ± 41 ms per 100 APs; $N = 6$; Fig. 2i–j) than when spiking was near the trough (probability, 1.3 ± 0.4 plateaus per 100 APs; duration; 63.5 ± 33.5 ms per 100 APs; $N = 6$; Fig. 2i–j) even though the average firing rates were similar under the two conditions (peak, 8.6 ± 1.4 Hz; trough, 7.8 ± 1.7 Hz; $N = 6$). The phase preference of plateaus under these conditions was similar to that of the control (Fig. 2h). This manipulation provides further evidence that plateau potentials in CA1 pyramidal neurons are preferentially generated when CA3 input drives AP output to coarsely overlap with synaptic input from EC3.

Experimental reduction of EC3 input to CA1

To elucidate the role of the distal long-range input, we recorded from mice expressing archaerhodopsin (Arch) in EC3 pyramidal cells (see Online Methods and Fig. 3a)17. We inhibited synaptic input from EC3 neurons by activating Arch in the axons and terminals of these cells using an optical fiber fitted inside the LFP electrode that was located in the stratum radiatum near the recorded neuron (3–5 s duration 594-nm laser pulses; Supplementary Fig. 7). A similar manipulation reduced the time integral of synthetically evoked excitatory postsynaptic potential (EPSP) (as measured by area under the curve) ~30% in hippocampal brain slices from these transgenic mice (Supplementary Fig. 8). Activation of Arch influenced our measures of burst firing (Fig. 3b,e,f) and significantly reduced plateau probability and normalized duration by ~50% (no light versus light: probability, 3.8 ± 0.8 versus 1.7 ± 0.5 plateaus per 100 APs; duration, 63.4 ± 6.8 versus 27.8 ± 6.0 ms per 100 APs; $N = 8$ neurons; Supplementary Fig. 9c–e) whereas running speed was comparable for laps with and without light stimuli (no light, 29.6 ± 1.6 cm/s; light, 28.5 ± 1.3 cm/s; $N = 8$). This manipulation of the distal synaptic input stream indicates that EC3 input is critical in generating plateau potentials in awake, behaving mice.

EC3 and CA3 pathways in the generation of plateaus

We next examined the relative contribution of each individual pathway in plateau initiation by performing an in vitro experiment...
that allows independent pathway activation. Here we paired separate stimulation of either the EC3 or CA3 input pathways with long-duration current injections (300–600 pA, 400 ms) into CA1 pyramidal somata (Fig. 3–n). Pairing of current injection with EC3 input pathway stimulation increased the occurrence of dendritic plateau potentials, quantified as the associated tuft Ca\textsuperscript{2+} signal area (measured as time integral of the change in fluorescence signal over baseline fluorescence, ΔF/F) (600 pA alone, 50 ± 11 ΔF/F ms; plus EC3, 151 ± 16 ΔF/F ms; N = 6 neurons; Fig. 3j,n). This was not the case, however, when the same currents were injected together with similarly sized CA3 input pathway stimulation. Instead, Ca\textsuperscript{2+} plateau initiation was rare and the tuft Ca\textsuperscript{2+} signal was not different from that observed for current injection alone (600 pA alone, 50 ± 11 ΔF/F ms; plus CA3, 60 ± 10 ΔF/F ms; N = 6; Fig. 3m,n). This result highlights the special, perhaps necessary, role of the EC3 pathway in the production of plateau potentials in CA1 pyramidal neurons. For its part, CA3 input appears to provide apical dendritic depolarization and, perhaps most importantly, to produce AP output that backpropagates into the distant regions of the apical dendrite to interact with the EC3 input\textsuperscript{7–10}.

Gain modulation of place-field firing by plateau initiation

To determine the impact of these events on CA1 output, we calculated plateau probability as a function of the animal's location. We found when examining place field traversals that plateau probability was highest within the field and peaked near the place field center, consistent with previous reports of intracellularly recorded complex spiking\textsuperscript{19} (bootstrapped permutation test, \(P = 1 \times 10^{-4}\); Fig. 4a–c). Notably, plateau potentials positively modulated the frequency of in-field AP firing (~30% increase in peak firing rate) in a manner consistent with an increase in firing rate gain and heavily increased the incidence of high-frequency firing and short inter-spike interval (ISI) events (Fig. 4d–f). These data indicate that the generation of plateau potentials elevates in-field AP firing frequency by promoting
high-frequency burst firing output from CA1 pyramidal neurons. In addition to this effect on firing rate, burst firing output should also be particularly effective at transmitting information to downstream regions owing to the facilitating nature of CA1 pyramidal neuron synapses.\(^\text{29,30}\) Thus, one function of dendritic plateau potential initiation appears to be a distinctive gain modulation of place-field firing rate via the induction of high-frequency bursts of AP output within the field.

Considering all of the above data, dendritic plateau potentials appear to operate as coincidence detectors that signal the presence of correlated or conjunctive EC3 and CA3 input by modulating the rate and mode of AP output in CA1 pyramidal neurons. Such a nonlinear input interaction would allow the association of internal and external features through a burst-firing-related output modulation mechanism that could produce a mixed representation of space and context useful for encoding episodic memory.\(^\text{15,16,22–25}\).

**Spontaneous plateaus are associated with new place field formation**

In addition to the above described ability of plateau potentials to produce a robust gain modulation of CA1 output, plateaus were also observed to be associated with the rapid formation of new place fields. A subset of pyramidal neurons abruptly transitioned from either silent or non-place firing into position-specific firing at a particular time during the recording session (\(N = 6\) neurons; Fig. 5 and Supplementary Fig. 10). Inspection of the \(V_m\) traces during this transition revealed the presence of long-duration, plateau-driven complex spiking that consisted of bursts of high-frequency APs riding on top of large (~40 mV) and long-lasting (138 ± 23 ms, see Fig. 5).

**Figure 4** Plateaus drive burst firing output within place fields. (a) AP firing rate for single laps as a function of distance for a representative cell. (b) Mean intracellular ramp for all laps as a function of distance (red). Spatial locations of plateaus are indicated by blue dots. (c) Spatial distribution of all plateaus in all neurons as a function of animal position. Place field (PF) centers were normalized within trials. \(P = 1 \times 10^{-4}\), bootstrapped permutation test (\(N = 226\) events from 17 neurons; see Online Methods). (d) Normalized (norm) firing rate as a function of animal position aligned to place field centers for trials with and without plateaus. (e) Frequency distribution for all laps with and without plateaus. Bin size is 140 ms. (f) Interspike interval (ISI) distribution for laps with and without plateaus. For d-f, \(N = 5,645\) APs, 78 laps with plateau; \(N = 5,754\) APs, 141 laps without plateau; from 17 neurons. * \(P < 0.05\), paired two-tailed t-tests. Exact \(P\) values (from left to right): d, 0.034, 0.024, 0.041; e, 0.005, 0.044, 0.041, 0.032, 0.002; f, 0.042, 0.038, 0.043, 0.027, 0.040.

**Figure 5** Place fields form after appearance of a large plateau potential. (a) Representative intracellular \(V_m\) as a function of time for laps (numbers at left) before, during and after appearance of a spontaneous plateau. Red box indicates portion of trace shown in c. (b) AP firing rate for single laps as a function of distance for representative cell from a. (c) Top, intracellular \(V_m\) from lap 8 on an expanded time base to show plateau characteristics. Red line is smoothed trace. Bottom, phase preference of plateaus for all place cells induced by a spontaneous plateau (\(N = 6\) cells from 6 mice). Gray line is a sine fit. (d) Mean intracellular ramp amplitude (top) and \(\Delta \theta\) (bottom) as a function of time for cell in a. The traces at right are averages across laps within the time window indicated by the gray or black bar on left plot to show the average ramp and \(\Delta \theta\) for laps before and after plateau. (e) Plateau duration as a function of phase for all place cells induced by a spontaneous plateau. (f,g) Average intracellular ramp amplitude (f) and \(\Delta \theta\) (g) for all place cells induced by a spontaneous plateau as a function of time. Data are shown as mean ± s.e.m.; see Online Methods for number of neurons in each data point.
N = 22 events) depolarizations in all neurons that formed new place fields (Fig. 5a,c and Supplementary Fig. 10). $V_m$ traces from laps following the dendritic plateau potential displayed a consistent location-specific increase in firing rate (mean peak firing rate $12.7 \pm 2.2$ Hz; $N = 6$) that was driven by large-amplitude, slow $V_m$ depolarizations (ramp; before, $0.8 \pm 0.2$ mV; after, $6.9 \pm 1.0$ mV; $N = 6$; $P = 2.3 \times 10^{-9}$, paired two-tailed $t$-tests) and substantial intracellular $V_m$ theta oscillations ($\Delta$theta; before, $0.6 \pm 0.2$; after, $1.7 \pm 0.4$ mV; $N = 6$; $P = 1.6 \times 10^{-4}$, paired two-tailed $t$-tests) at the location on the track where the plateau had appeared for the remaining duration of the recordings ($19.8 \pm 4.2$ laps; $9.6 \pm 1.2$ min; $N = 6$) (Fig. 5 and Supplementary Fig. 10). The above data suggest a role for dendritic plateau potentials in the rapid formation of new place fields in CA1 pyramidal neurons.

Figure 6 Plateaus are sufficient to drive new place field formation. (a) Representative intracellular $V_m$ as a function of time for laps (numbers at left) before, during and after plateau induction. (b) AP firing rate for single laps as a function of distance for cell from a. Arrow indicates current injection locations. (c) Intracellular $V_m$ (black) and extracellular theta (green) expanded from box in a. Red line is smoothed trace. (d) Mean intracellular ramp amplitude (top) and $\Delta$theta (bottom) as a function of time for cell in a. (e) Average intracellular ramp (top) and theta (bottom) as a function of position for laps indicated by gray or black bars in d. (f) Plateau duration as a function of theta phase. (g,h) Average intracellular ramp amplitude (g) and $\Delta$theta (h) for cells with plateau-inducing (black) or AP-inducing (red) current injections as a function of time. Data are shown as mean ± s.e.m.; see Online Methods for number of neurons in each data point. (i,j) AP firing rate for single laps as a function of distance for place fields induced at locations 1 (i) and 3 (j). (k) Average ramp amplitude after place field induction for all place cells induced with current injection (black circles) and for spontaneous plateaus (open squares) as a function of track position. Averages across cells at the three positions are shown as large open circles (mean ± s.e.m.).
Induced plateaus drive new place field formation

To more directly test whether plateau potentials are sufficient to induce place field formation, we developed a closed-loop experiment to evoke plateau potentials at specific locations along the track. Although naturally occurring plateaus require coincident input from CA3 and EC3, we were able to induce plateau potential initiation through intracellular current injection (300 ms, ~600 pA, Fig. 6) at the same location along the track for repeated laps (~5 laps; Fig. 6 and Supplementary Fig. 11). The induced plateau potentials had similar properties to naturally occurring plateaus (phase preference, 25°±15°, N = 102 events; phase modulation index, 0.83 ± 0.04, N = 102 events; duration, 56 ± 10 ms, N = 141 events). Such repeated plateau initiation was sufficient to induce spatially selective AP firing at the location of the current injection (mean peak firing rate 15.4 ± 2.1 Hz; N = 14 neurons) in 93% of cells tested (14 of 15 neurons). This position-specific firing was driven by the rapid appearance of a large-amplitude $V_m$ depolarization (ramp; before, 0.8 ± 0.2 mV; after, 7.2 ± 0.6 mV; N = 14; $P = 2.3 \times 10^{-11}$, paired two-tailed $t$-tests; Fig. 6d,e,g) and $V_m$ theta oscillation ($\Delta$theta; before, 0.6 ± 0.2 mV; after, 1.7 ± 0.2 mV; N = 14; $P = 7.4 \times 10^{-8}$, paired two-tailed $t$-tests; Fig. 6d,e,h). The induced place fields were stable for the remaining duration of the recording (15.1 ± 1.8 min; N = 14) and for many laps (39 ± 7; N = 14). In most respects, the induced place fields were similar to those of non-induced place cells: the mean peak firing rate, ramp amplitude, phase precession of APs, phase precession of intracellular theta and AP theta phase preference were not significantly different from those of standard place cells (Supplementary Fig. 11d–f; two-tailed unpaired $t$-tests, $P = 0.22, 0.46, 0.067, 0.09$ and 0.28, respectively). However, $\Delta$theta was slightly, but significantly, lower in the induced place fields when compared to standard place cells (Supplementary Fig. 11e; two-tailed unpaired $t$-test, $P = 0.001$).

Input resistance did not change following place field induction (before, 69.5 ± 2.8 MΩ; after, 73.1 ± 3.5 MΩ; N = 14 neurons, $P = 0.48$; paired two-tailed $t$-test; Supplementary Fig. 12).

The amplitude of the resulting position-specific increase in $V_m$ ramp depolarization and firing rate also did not vary as a function of position for induced plateau initiation (N = 14; $F = 0.2425$, ANOVA; Fig. 6k). This indicates that there was no spatial bias to place-field generation and that the only determinant was the position of repeated plateau potential initiation. These observations suggest that CA1 pyramidal neurons receive proportional amounts of input from presynaptic cells that are tuned to all possible spatial locations and as a result are equally capable of expressing place-specific firing at any potential position.

High-frequency single spiking is not effective

To determine whether the plateau potentials or the associated high-frequency AP firing was the main driver of place field formation, we next used a current-injection waveform that generated a similar number of high-frequency AP trains but that did not contain any plateau potentials (train of 30 APs at 100 Hz, ~1-nA, 2-ms-duration pulses). The trains of APs induced only a small increase in the ramp amplitude (ramp; before, 1.1 ± 0.3 mV; after, 1.9 ± 0.4 mV; N = 6...
neurons; \( P = 0.007 \), paired two-tailed \(-t\)-tests) that was both smaller than that induced by plateau initiation (\( P = 4.7 \times 10^{-3} \); unpaired two-tailed \(-t\)-test) and insufficient to produce a lasting increase in position-specific firing rate \((0.8 \pm 0.1 \text{ Hz}; N = 6; \text{ Supplementary Fig. 13})\). There was no significant effect on intracellular theta oscillations \((\Delta \text{theta}; \text{ before, } 0.7 \pm 0.1 \text{ mV}; \text{ after, } 0.8 \pm 0.1 \text{ mV}; N = 6 \text{ neurons}; \ P = 0.41 \), paired two-tailed \(-t\)-tests). These data indicate that dendritic plateau potentials, but not high-frequency AP firing, are sufficient for rapid place field generation in CA1 pyramidal neurons during our behavioral task.

**Plateaus increase the amplitude of incoming input**

We next sought to determine the mechanism of the plateau-induced formation of new place fields. Given the previously described ability of plateau potentials to induce synaptic plasticity\(^{10,14} \), a straightforward prediction is that synaptic potentiation at the position of the plateaus drives subsequent place-specific firing. Alternatively, plateaus could produce an increase in the frequency of inputs present at a given position of the track. An analysis of \( V_m \) fluctuations can distinguish between these two possibilities, as \( V_m \) variance is a linear function of the frequency of inputs but increases as the square of EPSP amplitude (\textit{Supplementary Fig. 14})\(^{31,32} \). The analysis showed that both spontaneous and induced place field generation caused the variance of \( V_m \) fluctuations within the fields to increase nearly fourfold over that of laps without place field firing \((\text{before, } 2.4 \pm 0.1 \text{ mV}^2; \text{ after, } 8.0 \pm 0.8 \text{ mV}^2; N = 19 \text{ neurons}; \ P = 8.5 \times 10^{-9} \), paired two-tailed \(-t\)-tests; \ Fig. 7\), and variance was found to increase as approximately the square of the mean \( V_m \) \((\text{variance versus mean plots were best fit with power } = 1.7 \pm 0.2; N = 19 \text{ neurons})\). Additionally, place field generation caused an increase in the higher order moments \((\text{particularly excess kurtosis})\) of the \( V_m \) distribution \((\text{before, } 0.36 \pm 0.08; \text{ after, } 1.0 \pm 0.2; N = 19 \text{ neurons}; \ P = 3.2 \times 10^{-5} \), paired two-tailed \(-t\)-tests; \ Fig. 7f,h\). The observed large increase in \( V_m \) variance within the induced place fields and the supralinear relationship to the mean \( V_m \) is consistent with a position-specific increase in EPSP amplitude rather than an increase in the frequency of the synaptic inputs (\textit{Supplementary Fig. 14}).

We next performed the \( V_m \) fluctuation analysis on place cell and silent cell populations. This analysis showed that place cells had within-field \( V_m \) variance that was approximately four times that of silent cells \((\text{silent cell, } 2.0 \pm 0.2 \text{ mV}^2; n = 10 \text{; place cell, } 8.8 \pm 1.4 \text{ mV}^2; n = 12; \ P = 4.8 \times 10^{-7} \), unpaired two-tailed \(-t\)-tests; \ Fig. 7g\) and was not different from that in neurons with induced place fields \((\text{unpaired two-tailed } -t\text{-test; } P = 0.42; \text{ Fig. 7g})\). The degree of nonlinearity in the relationship between \( V_m \) variance and mean was also not different from that of induced place fields \((\text{power } = 1.6 \pm 0.2; N = 15 \text{ neurons}; \text{ unpaired two-tailed } -t\text{-test; } P = 0.30)\). Together these analyses support the idea that an elevation in position-specific synaptic weight is an important mechanism underlying place-field firing in hippocampal CA1 pyramidal neurons.

**Impact on sharp wave/ripple-related activity**

Finally, we observed that the induction of new place fields by plateau potential initiation during active running was associated with an alteration in the activity of neurons during subsequent sharp wave/ripple (SPW/R) network states that occurred within nonexploratory periods. When rodents stop actively exploring an environment, the hippocampus transiently enters a distinct network state that is characterized by short-duration \((<100 \text{ ms})\), high-frequency \((\sim 200 \text{ Hz})\) oscillations in the extracellular field potential (\textit{Fig. 8})\(^{33,34} \). Intracellular \( V_m \) depolarization produced during ripple activity was significantly enhanced following the induction of place fields during active exploration \((\text{before, } 2.0 \pm 0.3 \text{ mV}; \text{ after, } 4.3 \pm 0.2 \text{ mV}; N = 16; \text{ Fig. 8c})\).
Furthermore, this enhanced $V_{in}$ depolarization also increased the number of APs produced during ripple activity (before, $0.01 \pm 0.004$ APs; after, $0.10 \pm 0.025$ APs; $N = 16$; Fig. 8d). The effect, particularly on AP firing, was most pronounced when ripples occurred at positions on the track that were near the location of the induced place field (after inside place field, $0.154 \pm 0.034$ AP; after outside place field, $0.047 \pm 0.026$ AP; $N = 16$; Fig. 8d). These data indicate that a common mechanism, plateau potential–mediated plasticity, underlies neuronal feature selectivity during both distinct hippocampal network states (theta versus SPW/R states), further strengthening the idea that the neuronal activity associated with ripples is a reactivation of the activity that occurred during active exploration.

**DISCUSSION**

We show that dendritic plateau potentials enhance place field firing rates and the incidence of complex spike burst firing in CA1 pyramidal neurons. The timing of plateau potential initiation coincides with EC3 input to CA1 cells, and plateau probability is increased when perisomatic $V_{in}$ is manipulated to drive CA1 spiking near the theta phase preference of EC3 input. Furthermore, optogenetic inhibition of EC3 input in the behaving animal, as well as independent pathway activation in the slice, suggest a unique role for EC3 input in the generation of dendritic plateau potentials. These data are consistent with previous in vitro studies showing that dendritic plateau potentials produce a nonlinear integration of CA3 and EC3 input streams, the size of which is related to the degree of correlation.

In addition, the data show that dendritic plateau potentials are sufficient to induce place field formation in CA1 pyramidal cells and that the underlying mechanism is likely to be a strengthening of synaptic inputs active around the time of plateau generation. The ability of plateaus to induce fields at any position along the track indicates that a given CA1 pyramidal cell receives input tuned to all spatial locations and is therefore equally able to express location–specific firing at any position. We did not find any evidence of a position–specific bias in the number or frequency of inputs, suggesting that CA1 cells receive an approximately proportional amount of inputs for all regions of the linear track and that the generation of a long-duration plateau potential chooses the particular feature for which the neuron will be selective (Supplementary Fig. 15). The plateau–potential–driven increase in synaptic input also appears to result in an enhanced reactivation during SPW/R activity, implicating a common mechanism for position–specific firing during both active exploration and non-exploratory network states. Furthermore, since CA3 is considered to be the source of SPW/R activity$^{35,36}$, the plasticity evoked by dendritic plateaus should involve CA3 inputs to CA1 pyramidal neurons.

The two mechanisms currently thought to be involved in the formation of place fields in hippocampal neurons are a linear combination of different periodic grid cell inputs$^{37,38}$ and various Hebbian synaptic plasticity mechanisms$^{39–41}$. The data presented here indicate that input-potentiating plasticity mechanisms rather than location–dependent increases in input numbers are the most prominent determinant of position–selective firing in CA1 place cells. However, there remains the possibility that variations in EC3 input, mediated by overlapping grid–cell input, may regulate the probability of plateau potential initiation in a position–dependent manner. This could potentially connect the two mechanisms. The observation that dendritic plateau potentials were much more effective than high–frequency APs at rapidly producing long–lasting place–specific input potentiation has implications for the rules governing place field formation in CA1 pyramidal neurons. Since plateau potentials are induced by the conjunction of two input pathways (EC3 and CA3), it follows that CA1 neurons may use nonstandard associative learning rules$^{42–43}$ to form place fields with properties that are a function of both CA3 and EC3 input. More experiments are needed to determine the exact synaptic mechanisms and induction rules involved in the generation of CA1 place fields.

The spontaneous plateaus that were found to induce place fields were approximately three times as long as those plateaus produced within existing place fields or those induced by current injections. This suggests that there may be some special condition associated with their initiation. Such a condition could be subcortical input driving both the dendritic disinhibition and neuromodulation that are likely required for such long-duration events. Future experiments manipulating these elements should reveal the detailed circuit mechanisms involved in spontaneous plateau generation. In the end, the plasticity produced by plateau potentials is capable of generating a place field in a single trial, and such rapid field induction is a known property of CA1 pyramidal neuron activity during exploration$^{44–46}$. While it is not entirely clear how long newly generated place fields will persist, the continued generation of plateau potentials within the new field could enhance their stability and this could be another important function for plateau generation within place fields.

Although we have above mainly focused on the role of excitatory input in dendritic plateau potential generation, inhibitory inputs are also likely to be important in regulating these signals$^{36,47–49}$. In particular, dendritically targeting interneurons that display a late theta phase firing preference are well suited to control the initiation and duration of dendritic plateau potentials. Along these lines, oriens–lacunosum–moleculare (OLM) interneurons are known to reduce complex spike probability and duration as well as influence spatial learning$^{46,49}$, underscoring the role of inhibitory elements in this microcircuit operation.

This and other studies present evidence that active dendritic integration in pyramidal neuron based cortical microcircuits supports a fundamental operation, the nonlinear interaction of long-range and local inputs that leads to firing rate modulation and neuronal plasticity$^{6–11,15,16,50}$. Indeed, evidence of a similar nonlinear modulation of tuned responses by active dendritic processing has been reported in layer 5 pyramidal neurons of the barrel cortex of behaving mice$^{6,7}$, and a comparable form of dendritic plateau–driven plasticity has been observed in neocortical neurons both in vitro and in anesthetized mice$^{14,50}$. Additional experiments are needed to fully explore the roles of various cortical microcircuit elements in what appears to be a candidate general microcircuit operation or low–level computation.

**METHODS**

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

K.C.B. and J.C.M. designed experiments; K.C.B. and C.G. performed in vivo recordings; S.P.V. and A.D.M. performed in vitro recordings; J.I.M. designed and...
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ONLINE METHODS

Animal experimentation. All experiments were performed according to methods approved by the Janelia Farm Institutional Animal Care and Use Committee (protocol 12-84).

Surgery and training schedule. Custom-made titanium head plates with an opening over the right hippocampus were affixed to the skulls of 5- to 6-week-old mice using Locite brand cyanoacrylate glue and dental acrylic while under either isoflurane or ketamine/xylazine anesthesia. The skull underneath the head implant opening was covered with cyanoacrylate glue and future locations for the craniotomies for the whole cell and field recordings were marked using stereotactic coordinates (2.0 caudal, 1.7 lateral for whole cell and 3.6–3.7 caudal, 1.7 lateral for field). After 5–7 d of recovery from surgery, running wheels were added to the cages and animals were placed on a water schedule protocol receiving 1.5 ml/d. After 5–7 d of water scheduling, animals were placed on the head fixation apparatus on either a training rig or the experimental rig for 20- to 40-min sessions per day for 5–14 d. Mice were trained to run for a 10% sucrose/water reward delivered once for every rotation of the belt (approximately 2 m) at the same location through alicking port. Mice were supplemented with additional water after training sessions to ensure they received 1.5 ml/d. Mice were used for recording after the number of trials (laps) completed per day was similar for 2–3 d. Animals that started training on the training rig were trained for at least two sessions on the experimental rig. The belt the animal used for training was the same belt used during the experimental session. The day before recording, animals were anesthetized using either isoflurane or ketamine/xylazine and two small (approximately 0.5 mm in diameter) craniotomies, one for the patch electrode and one for the field electrode, were drilled at previously marked locations. The dura was left intact and craniotomies were covered with silicone elastomer (Kwik-cast, World Precision Instruments).

Behavioral set-up and control. The linear track treadmill contains a belt made from velvet fabric (McMaster Carr) coated with visual and tactile cues as previously described26. There were three types of cues covering the length of the belt: glue spines, Velcro tabs and white spots. The belt is self-propelled and the mouse was used for recording after the number of trials (laps) completed per day was similar for 2–3 d. Animals that started training on the training rig were trained for at least two sessions on the experimental rig. The belt the animal used for training was the same belt used during the experimental session. The day before recording, animals were anesthetized using either isoflurane or ketamine/xylazine and two small (approximately 0.5 mm in diameter) craniotomies, one for the patch electrode and one for the field electrode, were drilled at previously marked locations. The dura was left intact and craniotomies were covered with silicone elastomer (Kwik-cast, World Precision Instruments).

In vivo electrophysiology. For each animal before whole cell patching, the depth of the pyramidal cell layer was determined using an extracellular recording. Extracellular electrodes were glass patch electrodes broken so that the resistance when filled with 0.9% NaCl was 1.5–3 MΩ. The extracellular electrode used to determine the pyramidal cell depth for whole-cell recording was mounted vertically on a micromanipulator (Luigs and Neumann). The extracellular signal was monitored using an audio amplifier (Grass Technologies). As the electrode was advanced through the cortex, the pyramidal cell layer was identified when theta modulated spikes were prevalent and the amplitude of ripples increased, typically 1.1–1.3 mm below the surface of the brain. The extracellular electrode to be used for LFP recordings during whole cell patching was mounted to a second micromanipulator (Narishige) on a 40–45° angle relative to the bregma, unless otherwise indicated, and advanced through the more caudal craniotomy. For most experiments, this extracellular electrode was advanced until detection of the pyramidal cell layer but not through it. For EC3 inactivation experiments, the fiber-containing electrode was advanced approximately 200–300 μm beyond the pyramidal cell layer to position the light near EC3 axons. To correct for any location differences in theta phase between the LFP electrode location and the location of whole cell patch recordings, we recorded the two LFP signals simultaneously. Long-taper whole-cell patch electrodes (7–12 MΩ) were filled with (in mM) 134 potassium gluconate, 6 KCl, 10 HEPES, 0.3 Mg-GTP, 4 Mg-ATP and 14 Tris-phosphocreatine. Whole-cell patch electrodes were advanced through the cortex with 6–8 psi of pressure to prevent the electrode tip from clogging. Upon entry into the hippocampus (approximately 100–200 μm from depth measured using the extracellular electrode), the pressure was reduced to 0.3–0.4 psi. Cells were identified by reproducible increases in electrode resistance. All recordings were made in current clamp using a Dagan BVC-700A amplifier and digitized at 20 kHz using an Instrutech ITC-18 with HEKA Patchmaster acquisition software. For the experiments in Figure 2, the LFP was filtered using a single-pole high-pass filter with 1-Hz cut-off frequency followed by a second-order Butterworth low-pass filter with a 12-Hz cutoff, delayed, and amplified using a custom-built FPGA device controlled by a Matlab graphical user interface. The filtering procedure produced an ~25 ms lag in the peak of the filtered response as compared with the unfiltered LFP. During whole-cell recording, the amount of amplification was adjusted until the current injection was sufficient to elicit spiking in either of the two delay conditions. To bias spiking toward the peak of extracellular theta, the filtered LFP was used with either 0 or 10 ms additional delay over that introduced by the instrument (25–35 ms total delay). To produce spiking on the trough of extracellular theta, the LFP was filtered and inverted before injection (producing the equivalent of ~95 ms total delay). The exact timing of the current injection relative to the LFP was subject to any theta phase offsets between the local LFP and that at the recording location. All data were grouped on the basis of the actual timing of the spiking relative to the LFP phase as measured during data analysis.

Optical fiber-containing LFP electrode preparation. To prepare an optical fiber-containing electrode, 9 μm core fiber (1550 BHP, Thorlabs) with one end connectorized was loaded through the perfusion port of an electrode holder, stripped, cleaved, and etched in hydrofluoric acid in two steps: 5 mm of fiber etched to 30 μm diameter, then the last 1 mm of fiber etched to 8–10 μm. The etched fiber was loaded into a quartz (1.0/0.7 mm or 1.2/0.9 mm OD/ID, Sutter Instruments) electrode with a tip size of 12–20 μm. The fiber positioning inside the pipette was adjusted until the fiber tip was recessed 100–500 μm from the pipette tip. These optical fiber-containing electrodes (LFP/fiber) were filled with saline the day of the experiment and used on subsequent days until the power out of the device dropped by more than 30% or the electrical resistance doubled. For laser light delivery, the output of a 594-nm laser (Mambo, Cobolt) was fiber-coupled into a single-mode fiber patch cable using a 3.3-mm focal length aspheric lens and the fiber mounted on a fiber-launch translation stage (MBT616D, Thorlabs) with associated hardware for FC/APC connectors. APC angled connectors were necessary to prevent back reflections into the laser head. The coupling fiber was connected to the LFP/fiber electrode using a FC/APC-FC/APC coupler. Optical throughput of a freshly prepared LFP/fiber electrode was 60–70%, and laser power out of the LFP/fiber electrode was between 7 and 12 mW for silencing experiments.

Photodetector for in vivo intensity measurements. To determine the light intensity profile for LFP/fiber electrodes, we constructed a photodetector device that could measure local light intensities in vivo. A plug of 645-nm-emitting quantum dots (QDP-645, Ocean Nanotech) was imbedded in epoxy at the tip of a pipette. To collect quantum dot fluorescence excited by laser light, this photodetector pipette also contained a 200-μm-core, 0.22 NA fiber, etched at one end to 40 μm diameter and fully inserted and glued into the photodetector pipette (Supplementary Fig. 7a). This fiber was connected to a bandpass filter (641/75, Semrock) and fluorescence was measured using a fiber-coupled spectrometer (QE65000, Ocean Optics). For the in vivo work, the photodetector was mounted onto the same manipulator used for patch experiments and inserted in a behaving animal at the same location as a patch electrode, to record the light intensity as a function of depth from the cortical surface. The laser light intensity was also measured at several distances from the LFP/fiber electrode tip, to generate an in vivo laser intensity heat map (Supplementary Fig. 7b). The photodetector was calibrated in saline using a 200-μm-core fiber coupled to a 561-nm laser as a light source of spatially defined light intensity using the same geometry as that used in vivo. EC3 inactivation. To inactivate EC3 inputs, we crossed mice expressing Cre in EC3 (pOx1-I-cre)17 with mice expressing Rosa-CAG-LSL-Arch-GFP-WPRE
The firing fields of single neurons were determined by first band-pass filtering solution containing (in mM) 210 sucrose, 25 NaHCO3, 2.5 KCl, 1.25 Na2HPO4, 0.75 CaCl2, 7 MgCl2, and 7 glucose. Cells were incubated for 30 min at 35 °C, and then recorded from at 35 °C, in a solution containing (in mM) 125 NaCl, 25 NaHCO3, 3 KCl, 1.25 Na2HPO4, 1 MgCl2, 1.3 CaCl2, and 25 glucose. All solutions contained fresh sodium pyruvate (3 mM) and ascorbic acid (1 mM), and were bubbled with 85% O2 and 5% CO2. Cells were visualized using a Leica BX-61 microscope using a water-immersion lens (60x; 0.9 NA, Olympus, Melville, NY). Whole-cell current-clamp recordings were performed using a Dagan BVC-700 (Minneapolis, MN) in active ‘bridge’ mode and were analog-filtered at 10 kHz before being digitized at 50 kHz using Prairie View acquisition software (Bruker Technologies, Middleton, WI). The pipette solution was the same as described for the in vivo electrophysiology but also included 100 µM OGB-6f and 50 µM Alexa Fluor 594 (Invitrogen). Dual recordings were performed by first filling the soma and then target-patching the dendrite using a galvanometer-based two-photon laser scanning system (Ultima; Bruker Technologies, Middleton, WI; using Chameleon Ultra II; Coherent, Auburn, CA). Line scans for Ca2+ imaging were performed with excitation at 920 nm using the photodetector (above and Supplementary Fig. 5). To quantify the average baseline Vm, APs were removed by deleting all points 0.26 ms before and 3.4 ms after a threshold value (dVm/dt = 50 V/s) and baseline Vm was defined as the peak of Vm histograms (Vm mode) from the resulting traces using a bin size of 0.2 mV. All traces were corrected for baseline drift on a lap-by-lap basis by determining the average deviation of the most hyperpolarized 5% of AP voltage thresholds measured from an expected value of –50 mV. To determine the properties of plateau potentials underlying complex spiking, we used a signal detection protocol that began by smoothing the above trace with a running average (401 point, ~20 ms) boxcar procedure, following linear interpolation of the deleted AP points. Event On and Off were detected from this trace by threshold crossing (~35 mV). This value was determined as a conservative estimate of where Vm distributions deviate from normal (Gaussian) (Supplementary Fig. 3). A second threshold level was evaluated (~30 mV) and, other than decreasing the total number of events detected by 12%, no quantitative differences were observed in any parameters. For the in vitro dual recordings, a burst of APs was classified as a plateau potential if the dendritic membrane potential corresponding to a somatic AP failed to repolarize below –30 mV. Plateau amplitude was calculated as the difference of event peak, and the baseline Vm and plateau duration was the time between the positive-going and negative-going threshold crossings. Plateau probability was determined by dividing the total number of plateau events by the total number of APs per cell. To analyze ramps of depolarization, we low-pass filtered (<3 Hz) the raw AP-deleted traces using an FIR filter with a 200 ms Hamming window. Ramp amplitude was quantified as the difference between peak (10 cm average around the most depolarized value) and the baseline (10 cm average around the most hyperpolarized value). To analyze subthreshold Vm oscillations, we band-pass filtered (4–11 Hz) the raw AP-deleted traces using an FIR filter with a 200 ms Hamming window and derived the amplitude vector using the Hilbert transform. The change in theta envelope (Δtheta) was calculated in the same manner as the ramp amplitude. The theta phase of APs and theta,dep were determined by first band-pass filtering the extracellular LFP in the theta range (4–11 Hz) and then deriving a vector of instantaneous phase using the Hilbert transform. The theta phase of each AP threshold crossing and theta,dep peak depolarization was found by interpolating from the vector of instantaneous phase. Theta-phase histograms of APs and plateaus were produced by dividing the theta cycle into 36 or 10 equal phase bins, respectively. The phase histogram of the AP and plateau events were computed for each individual neuron. Each AP phase histogram was fit by a sine function and the phase preference and modulation index were determined from the peak and modulation depth ((peak – trough)/peak) determined from this fit. These same values were determined for the plateau from smoothed plateau phase histograms (three-bin Gaussian window). For the population averages in Figure 2c, histograms of each neuron were first normalized by dividing by the total number of events found in each neuron. To determine whether there were any offsets in the phase of the recorded theta oscillation, in all recordings a reference LFP recording was performed at the site of the whole-cell recording simultaneously with the standard LFP recording and the theta phase peaks of the two simultaneously recorded LFP signals were compared. Most phase peak differences were less than 10° and no corrections were performed. However, differences between 10° and 40° were found in five neurons and we corrected for these differences during analysis. For the determination of phase precession of APs and theta,dep the phase values of these events were binned into 10-cm spatial bins for each individual neuron. For the population, the average phase of the first 10 cm of the place field

In vitro electrophysiology (Fig. 3 and Supplementary Fig. 3). 400-µm transverse hippocampal slices were cut from 8- to 10-week-old male C57/B6 mice using a Leica Vibratome VT1200S after perfusing an isoflurane-anesthetized animal with ice-cold solution containing (in mM) 197.2 sucrose, 2.5 KCl, 1.25 Na2HPO4, 0.75 CaCl2, 7 MgCl2, and 7 glucose. Slices were incubated at 30 min at 35 °C, and then recorded from at 35 °C, in a solution containing (in mM) 125 NaCl, 25 NaHCO3, 3 KCl, 1.25 Na2HPO4, 1 MgCl2, 1.3 CaCl2, and 25 glucose. All solutions contained fresh sodium pyruvate (3 mM) and ascorbic acid (1 mM), and were bubbled with 85% O2 and 5% CO2. Cells were visualized using a Leica Vibratome VT1200S after perfusing an isoflurane-anesthetized animal with ice-cold solution containing (in mM) 197.2 sucrose, 2.5 KCl, 1.25 Na2HPO4, 0.75 CaCl2, 7 MgCl2, and 7 glucose. 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input rate does not increase significantly (that is, fourfold; Supplementary Fig. 14g) it is impossible to achieve the levels of \( V_m \) variance (elevations of approximately fourfold) observed in experimentally recorded place fields (with \(-10 \) mV ramp) for this simple type of input integration model. Furthermore, even when extreme rate changes are used (sixfold excitation increase paired with fourfold inhibition increase) the relationship between \( V_m \) variance and mean is approximately linear (Supplementary Fig. 14g). By contrast, the relationship between \( V_m \) variance and mean was found to be greater than linear (power = 1.7) when the forcing function was applied to unit amplitude (Supplementary Fig. 14f).

Ripples were detected using the following algorithm. (1) LFP recordings from periods during which the animal did not run were concatenated and subsequently band-pass filtered between 100 and 250 Hz using an FIR filter. (2) The envelope of the filtered signal was calculated using the Hilbert transformation and ripples were subsequently detected as signals increasing >7 s.d. above baseline. (3) The onset and the end of the ripple were marked as the time points at which the envelope decreased below 3 s.d. of the baseline period; the time of the ripple peak was determined by peak detection between ripple onset and end. (4) The successful identification of the ripples and their features (peak, onset and end) was confirmed through visual inspection of the filtered signals along with the raw LFP signal. \( V_m \) periods during ripples were subsequently further analyzed. The number of APs during a ripple was determined by the number of threshold crossings in the corresponding \( dV_m/ dt \) trace (threshold = 50 V/s). Smoothed \( V_m \) traces (as above) were used to analyze ripple-associated subthreshold \( V_m \Delta V_m \) was calculated as the difference between the mean \( V_m \) values centered ±10 ms around the ripple peak and the baseline \( V_m \). For each lap, the baseline \( V_m \) value was determined by the peak of the \( V_m \) histogram (see above).

For the analysis in Figure 8 (\( N = 16 \) neurons), the neuron's place field was determined using the subthreshold slow \( V_m \) depolarization (or ramp), which is a typical intracellular feature of place fields\(^{20}\). To quantify the ramp, the linear track was divided into 100 spatial bins (each 1.9 cm wide) and an average \( V_m \) value was calculated for each bin using the previously generated AP-deleted \( V_m \) trace. After calculating the mean ramp depolarization across all laps, the place field was defined as locations on the linear track where the average of the sub-threshold ramp continuously exceeded a baseline period (determined by visual inspection) of about 10 neighboring bins (>19 cm) for at least 5 adjacent bins (9 cm). The remainder of the linear track was considered to be outside the place field. These place fields covered on average 74 ± 2% (mean ± s.e.m.) of the linear track. Ripple location was calculated relative to the midpoint (place field center) between place field start and end. Locations before and after the place field center were pooled.

**Statistical methods.** No statistical methods were used to predetermine sample sizes, but our sample sizes are similar to or larger than those reported in previous publications\(^{19–21}\). Where appropriate, data were analyzed using two-tailed paired or unpaired \( t \)-tests. Plateau duration, amplitude and probability are shown in text to be normally distributed. We assumed the same for ramp and \( \theta \) amplitudes. If normality was uncertain, we used non-parametric test as stated in the main text or figure legends. All manipulations were done within single neurons, with manipulations interleaved trial-by-trial (light versus no light) or in blocks of five trials (current injections). Data were analyzed automatically without consideration of trial condition. To determine whether plateau counts displayed location dependence (Fig. 4c), locations were shuffled 10,000 times, resulting histograms were fit with a Gaussian and \( R^2 \) values were compared between data histogram and location-shuffled data (bootstrapped permutation test). Data are shown as mean ± s.e.m. Data collection and analysis were not performed blind to the experimental conditions.

A Supplementary Methods Checklist is available.