Preexisting hippocampal network dynamics constrain optogenetically induced place fields

Highlights
- μLED optogenetic stimulation induces remapping in CA1 neurons
- Stimulated and non-stimulated neurons show comparable place field reorganization
- Fields emerge in places with weak preexisting drive, not at site of depolarization
- Stimulation changes coupling between pyramidal cells and neighboring interneurons

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In Brief
McKenzie et al. show that optogenetic depolarization of CA1 pyramidal neurons induces place fields to emerge in locations with weak preexisting drive. Stimulation also alters synaptic coupling strength between pyramidal cells and interneurons. Reorganization of feedback/lateral inhibition may be one mechanism for long-lasting changes in hippocampal neural computation.

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Preexisting hippocampal network dynamics constrain optogenetically induced place fields

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SUMMARY

Memory models often emphasize the need to encode novel patterns of neural activity imposed by sensory drive. Prior learning and innate architecture likely restrict neural plasticity, however. Here, we test how the incorporation of synthetic hippocampal signals is constrained by preexisting circuit dynamics. We optogenetically stimulated small groups of CA1 neurons as mice traversed a chosen segment of a linear track, mimicking the emergence of place fields. Stimulation induced persistent place field remapping in stimulated and non-stimulated neurons. The emergence of place fields could be predicted from sporadic firing in the new place field location and the temporal relationship to peer neurons before the optogenetic perturbation. Circuit modification was reflected by altered spike transmission between connected pyramidal cells and inhibitory interneurons, which persisted during post-experience sleep. We hypothesize that optogenetic perturbation unmasked sub-threshold place fields. Plasticity in recurrent/lateral inhibition may drive learning through the rapid association of existing states.

INTRODUCTION

Neural systems face a fundamental tension between maintaining existing structure and changing to accommodate new information. Several potential solutions have been raised to deal with this stability-plasticity dilemma. At one extreme, novel patterns of neural activity are imposed solely by “outside-in” sensory drive. In such models, the storage of these new activity patterns occurs through synaptic plasticity, a process which potentially interferes with the stability of previously stored knowledge (Bittner et al., 2017; Caillo-Reid et al., 2016; McClelland, 2013). At the other extreme, neuronal circuits could generate and maintain a preconfigured stable dynamic, sometimes referred to as an attractor, a manifold, or a neural schema (Hopfield, 1982; McKenzie et al., 2014; Tse et al., 2007), with a large reservoir of patterns available for matching with novel experiences (Battaglia et al., 2005; Dragoi and Tonegawa, 2011; Edelman, 1993; Golub et al., 2018; Luczak et al., 2009; Mizuseki and Buzsáki, 2013). Where biological memory systems fall along this “inside-out/ outside-in” continuum is a subject of debate.

The hippocampus is thought to contribute to episodic memory through rapid association of the constellation of sensory inputs that define a moment (Alme et al., 2014; Hasselmo and Wyble, 1997; Kumaran and McClelland, 2012; Liu et al., 2013; McNaghton and Nadel, 1990; Rolls, 1990; Tayler et al., 2013). According to this learning framework, the pattern of hippocampal neurons active at the time of learning generates an index for that episode (Teyler and DiScenna, 1986) and effectively binds the cortical regions carrying the multimodal episodic content (Tanaka et al., 2014). In support of models emphasizing rapid encoding of inputs, when single neurons are depolarized in a predetermined spatial position, robust firing persists in that location in the absence of external drive (Bittner et al., 2015, 2017; Diamantaki et al., 2016, 2018). This new spatial “code” is thought to be due to the association of that neuron with excitatory afferent signals naturally tuned to the stimulated location (Bittner et al., 2017; Dragoi et al., 2003), thus suggesting that any hippocampal neuron can be associated with any spatial location.

Neural communication in general, and the hippocampal index model in particular (Buzsáki and Tingley, 2018; Tayler and
DiScenna, 1986), are thought to depend upon the synchronous activity of groups of neurons. Little is known about how preexisting connectivity and self-organized dynamics affect the encoding of novel experiences. Here, we address these problems by pairing focal optogenetic stimulation of small groups of CA1 pyramidal (PYR) neurons with a segment of familiar space. Stimulation induced persistent place field remapping. Novel place fields emerged in both stimulated and non-stimulated neurons, and could appear outside the stimulation zone. The new place field location could be predicted by the correlation structure of neural activity before the optogenetic perturbation, suggesting that the incorporation of arbitrary signals was constrained by preexisting circuit dynamics. Circuit modification was reflected by altered coupling between connected PYR cell-inhibitory interneuron (INT) pairs and by changes in “synapsembles” (Buzsáki, 2010), which persisted during post-experience sleep. We hypothesize that optogenetic perturbation caused plasticity in recurrent/lateral inhibition to convert sub-threshold inputs into bona fide place fields.

RESULTS

Microscopic light-emitting diode (µLED) stimulation drives spiking in nearby CA1 PYR cells and INTs

The postulated ability of hippocampal circuits to imprint a random, novel pattern (Rolls, 1990) was tested in transgenic mice in which channelrhodopsin2 (ChR2) was expressed in excitatory PYR neurons (n = 5 CaMKIIα-Cre::Ai32 mice). µLED silicon probes were implanted in dorsal CA1 to record neural activity during sleep and behavior as mice ran on a linear track for a water reward. Before any stimulation, neurons were recorded for at least 1 h while mice stayed in their homecage (REST1), followed by running laps on a linear track for 10 baseline trials (pre-stim), stimulation (stim = 1 s half-sine
wave) was given for 2–10 trials (see Table S1) at a fixed position and running direction that changed daily. Immediately following stimulation, at least another 10 trials were given (post-stim) to test place field stability. During “control” sessions, identical test conditions were given; however, no μLED stimulation was provided on the linear track. During stimulation sessions, μLED optogenetic stimulation induced highly focal drive in CA1 PYR neurons recorded on the stimulated shank (Figures 1C–1E and S1; n = 631; median increase above expected rate = 4.67 ± 0.37 Hz, Wilcoxon signed rank test p < 10^-10; effect size = 1.14, 95% highest density interval (HDI) effect size that excludes 0; 84.0% neurons on stimulated shank numerically increased rate above expected baseline). Induced firing was calibrated, before track running, to be within the physiological range normally seen at the center of a place field (McClain et al., 2019). PYR cells on the neighboring shanks (≥ 250 μm away; n = 529) showed a small but significant decrease in expected firing (Figures 1C–1E; median rate difference = 0.02 ± 0.09 Hz, Wilcoxon signed rank test p < 10^-14; effect size = 0.29, 95% HDI effect size that excludes 0; 61.2% PYR cells on non-stimulated shank numerically decreased rate below expected baseline). The small decrease in the non-stimulated shanks was potentially due to the recruitment of local lateral inhibition. Inhibitory INTs recorded on the stimulated shank were also driven (Figure 1F; n = 126; median rate difference = 2.29 ± 2.83 Hz, Wilcoxon signed rank test p < 10^-10; effect size = 0.60, 95% HDI effect size that excludes 0; 69.1% INTs on stimulated shank numerically increased rate above expected baseline), likely through monosynaptic drive from the optogenetically stimulated PYR cells (English et al., 2017). The overall increase in excitatory firing was higher than that observed for inhibitory cells. Considering the multi-unit activity of all recorded PYR cells and all INTs, the excitatory:inhibitory (E:I) ratio increased transiently during stimulation before returning to baseline (Figure 1G). As shown by previous studies (English et al., 2017; Stark et al., 2015), we found that focal optogenetic stimulation engaged small networks of PYR cells and INTs during track running.

**Focal optogenetic stimulation induces partial remapping of place fields**

We next tested for the stability of optogenetically induced place fields (Rickgauer et al., 2014; Stark et al., 2012). In contrast to prior studies that manipulated single neurons with current injection (Bittner et al., 2015, 2017; Diamantaki et al., 2016, 2018; Zhao et al., 2020), we found that novel place fields emerged and disappeared both within and outside the stimulation zone (Figures 2 and S2). Furthermore, for CA1 neurons that remapped (see STAR methods), the locations of the place fields post-stimulation were not clustered around the stimulation zone on the track (Figures 2C–2F), showing a reorganization of the place representation, at least within the small volume of the recorded CA1 circuit.

Two approaches were taken to quantify the illustrated remapping. The log-likelihood (LL) of an observed spike train was assessed given two conditional intensity functions—one derived from that neuron’s place field recorded before stimulation trials (pre-template LL) and another from place fields recorded after the last stimulation trial (post-template LL). The degree to which the LLs differed, the LL ratio (post-template LL/pre-template LL; Harris et al., 2003), was used to quantify remapping. Using this LL ratio metric, spatial tuning of neurons recorded on both stimulated and non-stimulated shanks (neighboring shanks for which the μLED was not activated) showed greater reorganization immediately after the stimulation trials as compared to the spatial tuning observed during control sessions that lacked track stimulation (stim versus control: Mann-Whitney U test, p < 10^-5, effect size = 0.37, 95% HDI effect size that excludes 0; non-stim versus control: Mann-Whitney U test, p < 10^-7, effect size = 0.40, 95% HDI); remapping did not differ between stimulated and non-stimulated shanks on stimulation sessions (Figures 3A and 3B). Similar results were obtained with a different remapping metric that uses Pearson correlation to compare trial-by-trial rate maps with the average place field maps recorded before and after stimulation (Figure S3). Trial-by-trial estimates of place field drift also revealed greater remapping after optogenetic stimulation, regardless of whether the trials were separated by short or long intervals (Figure S3).

The LL approach provides a moment-to-moment estimate of whether the observed spike train is better described by the pre- or post-stimulation fire rate maps. We used this metric to quantify how quickly new spatial firing fields emerged after stimulation. For stimulated neurons that remapped (n = 84, 12.3% of all stimulated PYR cells; see STAR methods for remapping criteria), the spiking immediately after stimulation was better predicted by post-stim place fields (Figure 3C), suggesting that the place fields in these neurons emerged immediately. However, spiking of simultaneously recorded remapping neurons on the non-stimulated shanks (n = 44, 10.3% of all non-stimulated PYR cells) was equally well described by the pre- and post-stimulation place fields (Figure 3C) in the moments around stimulation, suggesting a distinct, slower process driving this firing field reorganization.

For those neurons that remapped, post-stimulation place fields signaled more spatial information (median information score = 4.94 ± 0.12 bits) than simultaneously recorded neurons with stable fields (median information score = 4.53 ± 0.10 bits; Mann-Whitney U test, p = 0.002; effect size = 0.35, 95% HDI effect size that excludes 0), and comparable information as neurons recorded in control sessions (median information score = 5.09 ± 0.17 bits, Mann-Whitney U test, p = 0.49). Consistent with this observation, Bayesian decoding of the position of the animal during pre-stimulation trials using pre-stimulation place fields (median decoding error = 15.1 ± 0.92 cm) was as accurate as location decoding during post-stimulation trials using post-stimulation place fields (median decoding error = 15.4 ± 1.0 cm; n = 27 sessions, Mann-Whitney U test, p = 0.87).

A second recording session (RUN2) revealed that the remapping could be stable after 1–4 h in the homecage (Figures 1C and 3D). Cells that remapped showed post-stimulation place fields that were more similar to those recorded during the second recording session (median corr<sub>POSTvsRUN2</sub> = 0.59 ± 0.03) than to those place fields recorded pre-stimulation (Mann-Whitney U test corr<sub>PREvsRUN</sub> versus corr<sub>POSTvsRUN2</sub>, p = 0.0003; effect size = 0.40, 95% HDI that excludes 0), despite the shorter time difference between the first and last stimulations (median = 6 min), as compared to the interval between RUN1 and RUN2 (median = 3 h 7 min). This stability after the homecage interval...
suggests that the observed remapping was not due to drift in place field coding or systematic recording instability. However, remapped place fields were not as stable as those observed in control sessions (median corr_{POST\text{vs\,RUN2}} = 0.75 ± 0.03; Mann-Whitney U test corr_{PRE\text{vs\,RUN2}} versus corr_{POST\text{vs\,RUN2}}, p<10^{-5}; effect size = 0.63, 95% HDI that excludes 0). We also found no difference in spike sorting quality between the remapping and non-remapping sub-populations, as measured by the L-ratio (p = 0.85) and isolation distance (Mann-Whitney U test p = 0.24). These results show moderate long-term stability of optogenetically induced remapping in CA1, which was less stable than that observed during control sessions.

Next, we examined potential physiological features that would reveal insights into why some neurons remapped and others did not. Comparing stimulated remapping cells versus stable cells, we did not find any differences in the radial position of the neuron (i.e., deep versus superficial, p = 0.23), nor in overall firing rate or burstiness during pre-rest or pre-stim RUN periods (Figure 3E, all p > 0.20). Remapping neurons recorded tended to fire at a higher rate during stimulation and during subsequent post-stim RUN trials (Figures 3D, 3E, and S3). Higher firing persisted during REST2 and was marginally higher during RUN2 (Figures 3D and 3E). Therefore, remapping was preferentially observed in neurons that responded to light with early
spiking and a high firing rate, while relative stability was observed in neighboring neurons.

Prior studies have found that back-propagating action potentials and dendritic plateau potentials cause new place fields to emerge in the position in which those strong dendritic events occurred (Bittner et al., 2015, 2017; Milstein et al., 2020). It is possible that the complex spike burst is an extracellular correlate of the plateau potential. We found that the first pass through a new place field (first trial with >3 Hz firing) was associated with bursting (spikes with ~5-ms inter-spike interval; Figure S3F), a defining feature of a complex spike burst, although bursts were not more prominent in the first trial (12.2% ± 0.02% spikes in a burst) as compared to later trials (11.1% ± 0.01% spikes in a burst), irrespective of the rate threshold used to define the first place field pass. The geometry of the implanted silicon probes (Figure 1A) also allowed for an analysis of the backpropagating spikes over the radial axis of CA1, albeit to a limited extent (<200 μm). We therefore tested whether the spikes that appeared on the first pass of a new place field differed from the mean waveform recorded for each unit in which a new field emerged. This analysis revealed no differences in spike waveforms (Figure S3). However, because of the limitations in detecting spike features from more distal apical dendrites with extracellular methods, these results do not exclude the possibility of important distal dendritic events. We next tested whether the running speed on the first trial predicted subsequent place field width (width at half the peak firing rate), as has been observed with spontaneous and artificial plateau potentials.
We found no correlation (Pearson correlation coefficient = 0.10, $p = 0.27$) between first trial running speed and subsequent place field width, irrespective of the firing rate criteria used to define a new field (1–10 Hz tested).

**Pre-stimulation dynamics predict post-stimulation remapping**

Since place fields did not emerge preferentially around the stimulation site on the track, we next examined whether any systematic firing patterns can explain the findings. We tested whether, before stimulation, neurons fired differently in the location that would later become the center of the remapped place field. During pre-stim trials, firing rates in the future place field were already elevated relative to the median rate recorded outside the future place field for both neurons on the stimulated shank (Figures 4A and 4C) ($n = 84$, Mann-Whitney $U$ test, $p < 0.05$ from trials 2–10; effect size 0.41–0.7, 95% HDI that excludes 0), and those recorded on the non-stimulated shanks (Figures 4B and 4C) ($n = 42$ neurons, Mann-Whitney $U$ test, $p < 0.05$ from trials 8–10; effect size 0.39–0.53, 95% HDI that excludes 0), even if such sporadic spiking did not translate to a quantitatively detectable place field. The pre-existing elevated spiking suggested that remapping unveiled the presence of preexisting weak drive to these neurons at the location of the future place field.

Next, we tested whether we could invert the analysis and predict where the place field would be located given the trial-by-trial pattern within and outside the future place field. Since the majority of data recorded before stimulation occurred outside the new place field, the classification labels (“future field pattern” versus “not future field pattern”) were heavily unbalanced. To confront this issue, we adopted the RUSBoost binary classification algorithm (Seiffert et al., 2009) to predict whether each location would become a new place field based on a feature space defined by the trial-by-trial firing rate of each neuron within that location, and the proportion of spikes emitted in bursts (inter-spike interval [ISI] $< 7$ ms) in that location. Half of the pre-stimulation trials were used to train the classifier, which was able to classify ripples activity for stim and control sessions) showed an increase in SPW-R recruitment as compared to those seen in control sessions in which light stimulation was not given. Stimulated PYR neurons showed slightly higher rate increases within ripples as compared to changes seen in neurons ($n = 291$) recorded in control sessions ($p = 0.02$; effect size $= 0.13$, 95% HDI that excludes 0) but not from neurons recorded on non-stimulated shanks (Figure 5A; $n = 220$; Mann-Whitney $U$ test, $p = 0.46$; effect size $= 0.04$). In contrast, INTs driven synaptically on the stimulated shank ($n = 63$ interneurons) showed a larger increase in SPW-R recruitment as compared to those ($n = 77$ interneurons) recorded on the non-stimulated shank (Figures 5B–5D) (Mann-Whitney $U$ test, $p = 0.005$; effect size $= 0.49$, 95% HDI that excludes 0), and compared to control sessions ($n = 70$ sessions; Mann-Whitney $U$ test, $p = 0.005$; effect size $= 0.45$, 95% HDI that excludes 0).

Typically, INT firing is excluded from SPW-R replay analyses. Therefore, we tested whether INTs that are naturally recruited during the running task persist in their firing during the following homecage recording session. We also tested PYR cell reactivation as a positive control. Since INTs lack well-defined firing fields, and typically have higher rates overall, we focused on the firing observed during waking SPW-Rs recorded on the track. As expected, CA1 PYR cells that were more strongly recruited into waking SPW-Rs ($n = 703$ neurons, threshold for ripple recruitment $= 2.5$ Hz, derived from median split of waking ripple activity for stim and control sessions) showed an increase in firing from REST1 to REST2 (Figure 5E) (paired Mann-Whitney
Figure 4. Relationship between pre-stimulation neural activity and post-stimulation remapping

(A) For remapping neurons recorded on the stimulated shank, the median firing rate in the center of the induced place field (blue) was already higher before optogenetic stimulation than in other regions of the track (red).

(B) Firing rate in the future place field before stimulation was also higher for neurons recorded on the non-stimulated shank. In-field versus out-of-field rate: \#p < 0.1, *p < 0.05, **p < 0.01.

(C) The distribution across remapping neurons of the difference-over-sum of the in-future-field firing rates versus the out-of-field firing rates measured before stimulation. In-future-field rates were defined as spikes measured when the mouse was ±5 cm around the future place field peak; out-of-field rates were the mean firing rates measured at all of the other locations on the track.

(D) The classification performance of the RUSBoost algorithm across neurons (points = neuron) to predict whether the pattern of neural activity in a given place on a given trial was recorded in the location of the future place field, or within another location on the track. Classifier performance compared against place-shuffled data.

(E) Method for relating pre-stimulation activity to subsequent place field distribution. Using data recorded during REST1, the binned firing rate of each target cell \( f_t \) was predicted using the weighted (\( w_a \)) binned rate \( f_{st_a} \) of other simultaneously recorded PYR cells. These weights were then related to the distance between place fields of the target cell and each peer predictor.

(F) Left: on average, for all pairs of neurons with overlapping pre-stimulation place fields (peaks within 5 cm), higher peer prediction weights were observed during REST1 as compared to pairs whose fields were further apart (spikes binned at 44 ms). Right: higher peer prediction strengths for PYR pairs with overlapping place fields were observed across a range of binning timescales. Bar indicates \( p < 0.01 \) comparing peer prediction weights for pairs of neurons with overlapping versus non-overlapping place fields. Inset, difference between peer prediction weights for neurons with overlapping versus non-overlapping place fields at each time bin.

(G) Same as (D), although comparing post-stimulation place fields and limiting the analysis to only neuron pairs in which either the target or peer predictor remapped. Shaded error bars, SEMs.
U test, p < 10^{-8}; effect size = 0.18, 95% HDI that excludes 0). This increase was greater than that observed for PYR cells (n = 672 neurons) that were less recruited into waking SPW-Rs (Figure 5E; unpaired Mann-Whitney U test, p < 10^{-8}; effect size = 0.21, 95% HDI). Similarly, INTs that fired more during waking SPW-R (n = 127 interneurons; threshold for ripple recruitment = 15.8 Hz) also showed a REST2 versus REST1 gain in homecage SPW-R firing (Figure 5F; paired Mann-Whitney U test, p = 0.0002; effect size = 0.44, 95% HDI that excludes 0), which was greater than that observed in INTs that were only weakly recruited during waking SPW-R (Figure 5F; n = 119 interneurons; unpaired Mann-Whitney U test, p = 0.003; effect size = 0.34, 95% HDI that excludes 0). Therefore, the synaptic drive of INTs during both optogenetic stimulation and natural drive during waking SPW-Rs was predictive of subsequent sleep SPW-R recruitment.

In addition, we tested for the reactivation of the optogenetically induced remapped spatial representations. We focused on those same pairs in which one or both neurons shifted place fields after stimulations. The degree of co-fluctuation (Spearman ρ) during SPW-Rs before track running strongly correlated with the pairwise correlation in ripple activity for ripples recorded both before and after track running. Only pairs of neurons with fields within 20 cm showed a significant (p = 0.007) gain in ripple co-modulation. 

These results demonstrate that the optogenetic perturbation can induce long-lasting changes in both PYR cell and INT firing during post-RUN SPW-Rs. Overall, these findings are consistent with the observation that the sequential order of place fields in both novel experiences (Dragoi and Tonegawa, 2011) and after artificial perturbation predicts prior internal circuit dynamics.
Optogenetic stimulation alters excitatory → inhibitory synaptic coupling

We hypothesized that the changes in SPW-R participation in synaptically activated INTs and the place field remapping, particularly of non-stimulated cells, was due to a reorganization of local CA1 circuitry, mainly affecting lateral inhibition. To explore this hypothesis, we examined spike transmission between pairs of monosynaptically connected PYR cells and INTs (PYR-INT) (English et al., 2017; Fujisawa et al., 2008). To track how spike transmission changed over time, we developed a generalized linear model (GLM; see STAR methods) to measure long timescale changes in the influence of the presynaptic drive to the postsynaptic INT while regressing out changes in the postsynaptic firing rate (Figures 6A and S4). We found that putative synaptic coupling strength, as approximated by our spike transmission measure (see STAR methods), varied 64.7% ± 0.5% around the mean over the recording session (Figures 6A, 6B, and S4). These temporal fluctuations were independent of the presynaptic rate (Spearman ρ = 0.0124, p = 0.077, n = 1,771 pairs), but correlated with the postsynaptic rate (ρ = 0.38, p < 0.001, n = 1,771 pairs), potentially reflecting a confounding effect of postsynaptic excitability (Figures S5A and S5B). However, spike transmissions across neuron pairs sharing the same postsynaptic INT were only moderately correlated with each other (ρ = 0.24, p < 0.05; Mann-Whitney U test) that were included in putative monosynaptic connections. *p < 0.05, ***p < 0.001.

See also Figures S4–S6.
coordinated and changes as a result of optogenetic stimulation, we extracted patterns of significant coactivity using a combination of principal- and independent-component analyses (PCA, ICA) (see STAR methods) to define a vector of factors for the contribution of each PYR-INT pair to each synaptic state (Figure S4), referred to here as synapsembles (Buzsáki, 2010) (Figures 6B–6E). Each synapsemble was detected before or after stimulation, and its expression strength was tracked throughout the entire daily session (see STAR methods). Compared to synapsembles in control sessions (n = 153), optogenetic perturbation was associated with significant changes in synapsemble expression strengths (n = 543, p < 0.001, Mann-Whitney U test; effect size = 0.44, 95% HDI that excludes 0; Figure 6F).

These results also held when controlling for differences in baseline synapsemble expression strengths between optogenetic perturbation and control conditions (Figure S5D).

Based on these analyses, we reasoned that changes in synapsemble expression ought to be accompanied by changes in cell ensembles defined by spiking. To investigate this possibility, we repeated the above analysis for presynaptic PYRs and postsynaptic INTs whose spike trains were smoothed to fluctuate at timescales equivalent to those of spike transmission (see STAR methods). While we found no differences in PYR ensemble expression (p = 0.128, Mann-Whitney U test; Figure 6F), INT ensembles were significantly reorganized following optogenetic stimulation (p = 0.012, Mann-Whitney U test; effect size = 0.48, 95% HDI that excludes 0; Figure 6F).

To further investigate the nature of this plasticity, we tested an alternative statistical model to capture changes in spike transmission (see STAR methods). In particular, spike transmission was constrained to fluctuate according to the Tsodyks-Markram dynamical model for short-term plasticity (Tsodyks and Markram, 1997), while long-term coupling strengths were held constant. The synapsemble analysis was repeated for this short-term plasticity model (Ghanbari et al., 2020), and as observed in the long-term model, synapsembles reorganized more in the optogenetic stimulation sessions as compared to controls (Figure S6). Therefore, multiple mechanisms could account for the observed changes in the coupling of PYR cells to INTs in the minutes and hours after stimulation.

**Discussion**

We report that optogenetic activation of a small group of PYR neurons resulted in the remapping of place fields both at and away from the stimulation site and in both directly stimulated and non-stimulated neurons. New place fields tended to emerge in places already associated with sparse preexisting activity. In addition, neurons with newly expressed place fields had preexisting correlated firing with partner cells whose place fields were co-tuned following optogenetic stimulation. Optogenetically induced place field remapping was accompanied by a reorganization of PYR-INT coupling and altered SPW-R membership of both PYR cells and INTs, which persisted in the hours after perturbation. We hypothesize that reorganization of hippocampal circuits is both constrained and guided by preexisting connectivity and internally organized physiological states.

**Constraints on place field remapping**

Interpretations of previous experiments on artificially induced place fields fall into two extremes. At one end is the assumption that the association of any arbitrary environmental input with the appropriate plasticity manipulation in the hippocampus can give rise to new place fields (omnipotent blank slate model). Interpretations at the other extreme assume that perturbations simply uncover preexisting or dormant place fields, and the success of artificial perturbations is, therefore, limited (preconfigured dynamical view). In support of the arbitrary place field induction interpretation, studies have shown that single neurons reliably form new place fields at any position on the running belt where those cells are injected with strong intracelluar current, resulting in a large dendritic plateau potential (Bittner et al., 2015, 2017). These experiments (see also Sheffield et al., 2017) gave rise to a model in which CA1 PYR cells receive inputs from an array of upstream neurons (presumably in CA3) that represent all parts of the covered space evenly, and the plateau-related “instructive signal” is broadcast to all input synapses. Any input that occurs within a multi-second time window around the plateau potential can get strengthened, but more so when the synapses are active before a plateau rather than during or after a plateau (Bittner et al., 2017; Milstein et al., 2020; Zhao et al., 2020). In support of the plateau potential experiments, juxtacellular stimulation of hippocampal PYR neurons (Diamantaki et al., 2018) and granule cells (Diamantaki et al., 2016) in freely moving subjects also induced place cell remapping to the stimulation site in a fraction of cases, especially when stimulation drove intense spiking (Diamantaki et al., 2018).

In contrast, our optogenetic perturbation experiments, along with previous studies on natural place field dynamics (Dragoi and Tonegawa, 2011; Grosmark and Buzsáki, 2016; Samsonovich and McNaughton, 1997), support the so-called preconfigured framework. Under our experimental conditions, neurons with induced place fields had dormant place fields and were weakly associated with preexisting neuronal assemblies. These relationships were strengthened after optogenetic stimulation, as was revealed by their gain in SPW-R-related firing. What appeared to be a freshly learned novel place field (i.e., newly added information to hippocampal circuits) may be more appropriately described as a matching process between existing internally organized neuronal trajectories and constellations of external inputs (Dragoi and Tonegawa, 2011; Farooq et al., 2019; Guo et al., 2020; Mizuseki and Buzsáki, 2013; Villette et al., 2015; Buzsáki 2010).

Other lines of evidence suggest that new place fields emerge upon a background of weak prior drive. The first place field perturbation study used long-term potentiation (LTP) of the CA3-CA1 Schaffer collaterals under the hypothesis that each CA1 neuron receives input from an array of CA3 neurons that covered the available space equally (Dragoi et al., 2003). LTP perturbation induced novel place fields and made existing place fields disappear, analogous to plateau potential studies in single neurons (Milstein et al., 2020). However, new fields tended to emerge in low firing rate neurons in locations associated with preexisting dormant activity; strong, existing place fields could not be modified. In line with these findings, spatially uniform depolarization of silent CA1 PYR cells led to the reversible...
emergence of a spatially tuned subthreshold response and place field spiking (Lee et al., 2012), indicating that a neuron driven by a preconfigured network in which it is embedded can selectively match to a particular external event. A related study showed that in a novel environment, subthreshold fields preexisted in the locations where place fields subsequently emerged without prior plateau potentials (Cohen et al., 2017).

Place fields can emerge rapidly in familiar and novel environments (Frank et al., 2004), precluding a scenario in which receptive fields emerge due to incremental plasticity with repetitive pairings of pre- and postsynaptic activity (Isaac et al., 2009). A recent behavioral study made the remarkable observation that before spontaneous emergence of a novel place field in a familiar environment, the exploring rat displayed prominent head scanning at that very place on the preceding trial (Monaco et al., 2014). Although plateau potentials may have occurred during such head scanning, the authors also found dormant place field firing before head scanning (their Figure 2), allowing the interpretation that head scanning, like optogenetic perturbation, strengthened the membership with an assembly that already signified that place.

There are important differences between our study and others that have linked plateau potentials to arbitrary remapping. In our study, groups of neurons were depolarized with focal optogenetic stimulation, rather than single neurons driven with strong intrasomatic current injection. Our optogenetic stimulation excited and synchronized a small group of PYR cells (an estimated 10–50 neurons by μLED) (Wu et al., 2015) and effectively discharged local inhibitory cells (English et al., 2017). In turn, the resulting recruitment of feedback inhibition could have prevented regenerative dendritic depolarization, underlying plateau potentials (Grienberger et al., 2017; Lovett-Barron et al., 2012; Sheffield et al., 2017). Single-neuron versus multi-neuron drive may or may not be important because remapping to the stimulation site was not observed when single CA1 neurons were depolarized by holographic two-photon stimulation in a head-fixed situation (Rickgauer et al., 2014) that was similar to that used in the plateau potential studies (Bittner et al., 2015, 2017; Milstein et al., 2020). In all of the optical preparations, targeted stimulation of specific neurons causes the place field remapping in both the targeted and non-targeted populations, with place fields appearing at locations not associated with the artificial drive (Rickgauer et al., 2014; Robinson et al., 2020), similar to what was observed here and in previous reports (Stark et al., 2012). Finally, we have observed that a newly induced place field can persist alongside an existing field of the same neuron, in line with findings that a minority of PYR neurons can have two or multiple place fields (Mizuseki and Buzsáki, 2013). In contrast, a plateau-induced novel place field makes the spiking at the original place field disappear (Milstein et al., 2020).

We hypothesize that the magnitude of artificial perturbation in single neurons may explain the differences among these seemingly contradicting studies. Firing rates and synaptic strengths of neurons upstream to the recorded CA1 cells are strongly skewed (Mizuseki and Buzsáki, 2013); thus, the assumption of equal representation of each position on the track may not hold. Such preconfigured, skewed synaptic weight distributions may explain why the dormancy of place cells predicted the expression of novel place fields. We submit, however, that if postsynaptic perturbation is strong, even very weak, preexisting connections can be strengthened sufficiently and give rise to a novel place field. Thus, while the dormant-to-overt place field conversion may be the physiological mechanism for novel place field induction, strong dendritic events may also trigger arbitrary position-related place field expression. Importantly, this interpretation does not negate the possibility that plateau potentials reflect a physiological mechanism by which place fields are induced in the intact brain (Bittner et al., 2015). However, our findings demonstrate that this process in vivo may be more one of unmasking preexisting connections and strengthened partnership with assembly peers. This unmasking mechanism of place field emergence may be analogous to the rate remapping observation, where “ghosts” of place fields in one apparatus can often be recognized, which predict bona fide place fields in another apparatus (Leutgeb et al., 2005).

**Mechanisms of perturbation-induced remapping**

Dormant activity in the induced place field recorded before optogenetic perturbation predicted the pattern of remapping, irrespective of whether the neuron was directly stimulated or not. The immediate changes during stimulation were likely brought about by the rapid redistribution of recurrent and lateral inhibition (Trouche et al., 2016), and some of the immediate changes persisted several hours post-stimulation. Alongside place field remapping, optogenetic stimulation also recruited INTs into ripples and affected the correlation structure among INTs and the pattern of coupling between monosynaptically connected PYR→INT pairs. We hypothesize that the primary component of this local circuit reorganization is long-term plasticity in inhibitory signaling.

Although plastic changes may take place at multiple levels of CA1 local connectivity, one potential locus of this plasticity is the PYR to INT synapse, which is known to undergo activity-dependent plasticity (Buzsáki and Eidelberg, 1982; Kullmann and Lamsa, 2011; Lau et al., 2017; Le Roux et al., 2013; Lu et al., 2007; Nissen et al., 2010). To study PYR→INT plasticity, we quantified the change in spike transmission probability (Csicsvari et al., 1998; English et al., 2017), a measure shown to be highly correlated with single PYR neuron-induced excitatory postsynaptic potentials (EPSPs) in parvalbumin (PV)-expressing INTs in vivo (Galarreta and Hestrin, 2001; Jouhanneau et al., 2018). A population measure of PYR→INT spike transmission (synapsembles) showed a significant rearrangement following optogenetic stimulation, possibly driving the remapping in the place fields of PYR cells. We also observed significant synapsemble reorganization in a statistical model lacking slow timescale fluctuations, where changes in spike transmission only depended on the recent (seconds) history of presynaptic spiking (Ghanbari et al., 2020). We acknowledge that these indirect measures of synaptic plasticity need to be confirmed by future intracellular experiments. However, the current data are compatible with the hypothesis that INTs played an important role (Dupret et al., 2013; Schoenenberger et al., 2016). As further support for the involvement of inhibition, we also demonstrated that INT firing rate ensembles, although not PYR ensembles, were reorganized following optogenetic stimulation. In addition, INTs on the stimulated shank showed a large gain in
SPW-R-related firing, compared to INTs on non-stimulated shanks or in control sessions. Finally, INTs that were highly active during waking SPW-Rs on the track showed a larger gain in SPW-Rs from pre-RUN sleep to post-RUN sleep in the home cage compared to INTs with low waking SPW-R activity. Overall, our analysis of ensemble activity points to the critical role of inhibitory neurons in both short-term and long-term remapping of place fields. The molecular identity of the INT type(s) that played a critical role in our experiments need to be explored in future experiments.

In addition, the relationship between induced and spontaneous reorganization of hippocampal circuits needs to also be addressed in future studies. A growing body of literature has shown that, even when task demands and sensory stimuli remain fixed, the spatial code of the hippocampus “drifts” (Bladon et al., 2019; Cai et al., 2016; Mankin et al., 2012; Ziv et al., 2013), as reflected by different neural firing patterns observed in the same location at different times. It is possible that some of this drift may be related to the reorganization of local feedback circuits. However, other mechanisms have also been postulated, such as fluctuations in intrinsic excitability (Cai et al., 2016) and varying readout of decaying cortical activity signaling some event that happened in the past (Bright et al., 2020). Rapid, task-dependent changes in the place coding have also been described (Jackson and Redish, 2007; Jezek et al., 2011; Kelemen and Fenton, 2010), which likely require inhibitory neurons to coordinate groups of PYR neurons and allow for rapid transitions to other ensembles when task demands change or when learning about a novel environment (Cohen et al., 2017; Frank et al., 2004; Sheffield et al., 2017). We hypothesize that our artificial stimulation affected this inhibitory recurrent circuitry to produce place field reorganization.

Conclusions
In summary, we hypothesize that moderate perturbations can unmask subthreshold, preexisting place fields. Overall, perturbation studies are compatible with the general framework that incorporation of novel information within brain circuits is constrained and guided by a backbone of a preexisting repertoire of states (Battaglia et al., 2005; Dragoi and Tonegawa, 2011; Farooq et al., 2019; Golub et al., 2018; Luczak et al., 2009; Mizuseki and Buzsáki, 2013). Such preconfigured dynamics may reflect regularities captured across prior learning experience—the neural schema (McKenzie et al., 2014; Sanders et al., 2020; Tse et al., 2007)—in combination with the self-organized scaffold assembled across development (Farooq and Dragoi, 2019; Li et al., 2012; Xu et al., 2014).

STAR Methods
Detailed methods are provided in the online version of this paper and include the following:

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**Supplemental Information**
Supplemental Information can be found online at https://doi.org/10.1016/j.neuron.2021.01.011.

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**Author Contributions**
S.M. and R.H. collected and analyzed the data. D.F.E prepared the experimental subjects. K.K. and E.Y. provided the surgical probes. G.B. supervised all aspects of the experiment. F.C. assisted with the statistical analyses. S.M., R.H., and G.B designed the experiments and wrote the manuscript.

**Declaration of Interests**
The authors declare no competing interests.

**References**


**STAR METHODS**

**KEY RESOURCES TABLE**

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**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources and data should be directed to and will be fulfilled by the Lead Contact, György Buzsáki (Gyorgy.Buzsaki@nyumc.org)

**Materials availability**
No new materials were generated in these studies.

**Data and code availability**
The datasets generated during the current study are available in the Buzsaki lab repository, https://buzsakilab.nyumc.org/datasets/McKenzieS/.

Custom lab software is available at https://github.com/buzsakilab/buzcode and MATLAB scripts for the current project will be made available upon reasonable request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

Homozygous CaMKIIα-Cre line T29-1 transgenic mice (Jackson Laboratory #005359) were crossed with homozygous Ai32 mice (Jackson Laboratory #012569) to express channelrhodopsin2 (ChR2) in neurons expressing male and female CaMKIIα in F1 hybrid mice (n = 5; 3 male; 25-40 g, 30-50 weeks of age). After implantation, animals were housed individually on a reversed 12/12 h day/night schedule. Following one week of recovery, mice were recorded 5-7 days/week for two months before being euthanized with pentobarbital cocktail (Euthasol®, intraperitoneal 300 mg/kg) and perfused with formalin (10%). All experiments were conducted in accordance with the Institutional Animal Care and Use Committee of New York University Medical Center.
METHOD DETAILS

Task
Mice were trained to run laps on a linear track (120 cm long, 4 cm wide) to retrieve water reward (5–10 μL) at each end. Before implantation, water access was restricted and was only available as reward on a linear track and ad libitum for 30 minutes at the end of each day. After mice reliably ran 50 trials in under an hour (~1 week daily training), free water access was restored for at least two days, and surgery was scheduled.

For a typical recording session, a one hour baseline recording was conducted in the mouse’s home cage (REST1), followed by a calibration of light intensity for optogenetic stimulation. Then mice ran 25–30 trials in a morning session (RUN1), followed by 1–4 hours of homocage recording (REST2) and another 25 trials in an afternoon/evening session (RUN2), followed by redelivery of the calibrating light pulses in the homecage and ad libitum water access. On Control sessions, the calibration pulses were given though no stimulation was delivered on the track.

To deliver optogenetic stimulation at a fixed location and running direction, an infrared sensor was placed at a random location on the track. Sensors were also placed at each end to control water delivery. An Arduino circuit detected beam breaks to activate solenoids to deliver water and to send a TTL pulse to a DAQ (CED Power 1401 Cambridge, UK) which delivered voltage control signals to the integrated μLEDs.

In every subject tested we noticed that there would be days in which mice would stop running past the unmarked stimulation site after stimulation and defecate despite being very familiar with the recording protocol. This occurred in 10%–15% of recording sessions and these sessions were excluded. In the remaining sessions, mouse behavior did not appear to be affected by the stimulation (Figure 1).

Surgery
Mice were anesthetized with 1.5%–2% isoflurane (2 L/min) and provided with a local anesthetic to the incision site (bupivacaine at 0.05 mg/kg, 2.5 mg/ml, S.C.). The skull was cleaned with saline and hydrogen peroxide and ground wires (bare stainless steel) were positioned intracranially over the cerebellum. The skull was then coated with Optibond (Kerr Dental, Brea, CA) and a craniotomy (~1.5 × 0.5 mm) was performed at AP ~2.2, ML ~2.0 (left hemisphere), 45° angle from the midline. The dura was removed and the probe was implanted ~0.5mm into the cortex. The probe and custom driver were cemented to the skull with C & B Metabond Quick Adhesive Cement (Parkell) and Unifast Trad acrylic (GC America). The craniotomy was capped with a mixture of mineral oil (one part) and dental wax (three parts), and a Faraday cage was constructed using copper mesh and connected to the cerebellar ground wire. Following surgery, an opioid analgesic was injected (Buprenex at 0.06 mg/kg, 0.015 mg/ml, IM) and given as needed for the next 1–3 days.

Recording and stimulation
Neural data was acquired using 32 site, 4-shank μLED probes (Wu et al., 2015; Neurolight, MI). Data were amplified and digitized at 30 kHz with Intan amplifier boards (RHD2132/RHD2000 Evaluation System, Intan). μLEDs were controlled with voltage (2–3.5 V generating 1–20 μW of total light power) provided by a CED Power 1401 programmed with Spike2 (CED) which delivered light pulses (pre and post run) or 1 s long sine waves (when the track IR beam was crossed). The animal’s position was monitored with a Basler camera (acA1300-60 gmNIR, Graftek Imaging) sampling at 30 Hz to detect head-mounted blue and red LEDs. Position was synchronized with neural data with TTLs signaling shutter position as well as a blinking LED (0.5 Hz) mounted 1 m above the maze.

Blue light (centered emission at 460 nm) was delivered on one or two μLEDs (always 1 μLED/shank, emission surface area = 150 μm²). To minimize artifact, the control voltage was held just under the forward voltage (2V). For calibrating light intensity, pulses (100 ms, 1–1.5 s variable inter-stimulus intervals) were delivered at 5 amplitudes (20 pulses/amplitude). The total radiant flux of the minimum pulse amplitude was typically around 1 - 5 μW, and the radiant flux of maximum amplitude, that was also used during track stimulation, was around 15-20 μW. This maximum intensity in the homecage was calibrated to be just subthreshold of a population oscillation, which was never observed on the track, likely since inhibitory tone is higher, as this highest stimulation level evoked a weaker response during the theta state (Figure S1). On the track, 1 s half sine waves were delivered when mice crossed an IR beam. Track stimulation only occurred in one running direction and, unless otherwise noted, was most frequently given for a block of 5 trials (see Table S1).

QUANTIFICATION AND STATISTICAL ANALYSIS

Unit isolation
Spikes were extracted and classified using Kilosort (Pachitariu et al., 2016). Global principal components were calculated (three per channel,8 channels/shank) and spikes were extracted from the highpass filtered wideband signal (3rd order Butterworth filter, passband: 0.5 – 15 kHz). Manual unit curation was done using Klusters. Spike sorting quality was assessed with L-Ratio (Schmitzer-Torbert et al., 2005), Isolation distance (Schmitzer-Torbert et al., 2005), inter-spike interval violation, and visual inspection of cross-correlations suggestive of erroneous splitting of single units.
Cell type classification
Spike waveform (width and asymmetry), autocorrelation properties, and mean firing rate (mean inter-spike interval) were used to classify neurons into excitatory cells and interneurons. The autocorrelation was parameterized with a double exponential model (Platkiewicz et al., 2021):

\[ \text{ACG}_\text{BASE} = (\beta_1 + \beta_2) \cdot e^{\frac{-x^2}{\sigma_1^2}}, \text{ for } x \leq 0 \text{ otherwise } \text{ACG}_\text{BASE} = 0 \]  

\[ \text{ACG}_\text{BURST} = \beta_1 \cdot e^{\frac{-x^2}{\sigma_2^2}} + \beta_2, \text{ for } x > 0 \text{ otherwise } \text{ACG}_\text{BURST} = 0, \text{ and } 0.9 < \beta_4 < 2.1 \]  

\[ \text{ACG} = \text{ACG}_\text{BASE} + \text{ACG}_\text{BURST} \]  

Units were defined by: rate, autocorrelation peak above baseline \( \beta_1 \), spike width, and spike asymmetry. Then k-means clustering (k = 2) was performed on the z-score normalized feature matrix which separated excitatory cells from putative interneurons (including fast spiking and regular spiking interneurons). The validity of the cluster labels was confirmed through the cross-correlation (CCG) analysis, revealing increased synchrony at synaptic time-scales (see Synapsemble analysis).

Ripple detection
The local field potential (LFP) was extracted by low-pass filtering the 30 kHz raw data (sinc filter with a 450 Hz cut-off band) and then downsampling to 1250 Hz. For ripple detection, the LFP was bandpass-filtered (3rd order Butterworth, passband: 130-200 Hz), squared, and z-score normalized. Events with peak power > 5 standard deviation (SD), sustained power > 2 SD, and duration between 30-200 ms were detected. When available, such events were also detected on a non-hippocampal, ‘noise’ channels and events common to both (e.g., EMG artifacts) were excluded. Stimulation periods were also excluded. Ripple onset was the first moment when the bandpassed signal increased > 2 SD.

Ripple co-fluctuation analysis
Ripple start and stop times were taken as the moments when ripple-band power rose above and fell below 2 SD of baseline. Rate per ripple was simply the number of spikes per neuron divided by the duration of the ripple event. Pairwise co-fluctuations were quantified through analysis of the Spearman correlation of ripple rates pre- \( \text{Corr}_\text{PreRUN} \) and postRUN \( \text{Corr}_\text{PostRUN} \). Multiple regression analyses were used to predict:

Post-RUN ripple correlations

\[ \text{Corr}_\text{PostRUN} = \beta_3 + \beta_1 \cdot \text{Corr}_\text{PreRUN} + \beta_2 \cdot \text{PlaceFieldSeparation}_\text{PostSTIM} \]  

while accounting for pre-stimulation place fields

\[ \text{Corr}_\text{PreRUN} = \beta_0 + \beta_1 \cdot \text{Corr}_\text{PreRUN} + \beta_2 \cdot \text{PlaceFieldSeparation}_\text{PostSTIM} + \beta_3 \cdot \text{PlaceFieldSeparation}_\text{PreSTIM} \]  

and preRUN ripple correlations,

\[ \text{Corr}_\text{PreRUN} = \beta_0 + \beta_1 \cdot \text{PlaceFieldSeparation}_\text{PostSTIM} \]  

while accounting for pre-stimulation place fields

\[ \text{Corr}_\text{PreRUN} = \beta_0 + \beta_1 \cdot \text{PlaceFieldSeparation}_\text{PreSTIM} + \beta_2 \cdot \text{PlaceFieldSeparation}_\text{PreSTIM} \]  

Place field remapping analysis
The two-dimensional location of the mouse was linearized by projecting actual position onto the straight line of the track. A Kalman filter (2nd order; locally quadratic) was used to derive a Bayesian MAP estimate of instantaneous speed and only moments with speed > 1.5 cm/sec were considered for place field analysis. Position was binned (100 bins, each 1.2 cm) and the spike count and occupancy at each binned position was convolved with a Gaussian kernel. Place maps were calculated as the smoothed spike counts divided by the smoothed occupancy. Rate maps were calculated separately for inbound and outbound journeys and for each of four conditions: \( f_{\text{pre-stim}}(x) \), \( f_{\text{stim-trial}}(x) \), \( f_{\text{post-stim}}(x) \), \( f_{\text{RUN2}}(x) \).

To calculate a trial-by-trial estimate of remapping, two methods were adopted. First, a statistical modeling approach was taken to assess the likelihood of the observed spike train derived from a Poisson process with a conditional intensity function defined by place field templates observed either before or after stimulation. The place maps were constructed from a training set of all trials except for the one for which the log-likelihood of the spike train was evaluated. The position \( x(t) \) at each moment during this test trial, and the place fields \( f_{\text{pre-stim}}(x) \), \( f_{\text{post-stim}}(x) \) constructed from the training set were used to produce an estimated intensity at each time point.
according to each place field template, $f_{\text{pre-stim}}(t) = f_{\text{pre-stim}}(x(t))$ and $f_{\text{post-stim}}(t) = f_{\text{post-stim}}(x(t))$. The log-likelihood of the spike train $t_s$ was calculated following Harris et al. (2003), under two place field models:

$$LL_{f_{\text{pre-stim}}} = - \int f_{\text{pre-stim}}(t)dt + \sum_{s} \log(f_{\text{pre-stim}}(t_s)),$$

(Equation 8)

$$LL_{f_{\text{post-stim}}} = - \int f_{\text{post-stim}}(t)dt + \sum_{s} \log(f_{\text{post-stim}}(t_s))$$

(Equation 9)

where $t_s$ is the time of each spike and $dt = \frac{1}{30}$ s.

The degree to which remapping occurred was quantified with the log-likelihood ratio normalized by the duration of the trial for the evaluated spike train $T$,

$$LLratio = \frac{LL_{f_{\text{post-stim}}} - LL_{f_{\text{pre-stim}}}}{T}$$

(Equation 10)

$LLratio$ gives how much information is conveyed by the spike train about the post-stimulation rate map relative to the pre-stimulation rate map in bits per second. The cumulative sum of the $LLratio$ without the normalization provides a running estimate of the total amount of information conveyed by the spike train with respect to one place field map or the other.

There were three classes of remapping neurons, those for which fields emerged, those for which fields disappeared, and those for which fields shifted. In all cases remapping neurons must have had average post-stim $LLratio > 0.75$ bit/sec. Shifting neurons must have had a spatial information score (Olypher et al., 2002) of more than 2 bits/spike for place fields before and after stimulation and the peak firing must have been at least 10 cm apart; there was no rate threshold. For disappearing fields, the peak rate before stimulation must have been $> 5$ Hz and the peak rate after stimulation must have been $< 5$ Hz with a pre-stim spatial information score $> 2$ bits/spike. Similarly, for appearing fields, the peak rate after stimulation must have been $> 5$ Hz and the peak rate before stimulation must have been $< 5$ Hz with a post-stim spatial information score $> 2$ bits/spike. Stable neurons were defined as those with an $LLratio < 0.75$ bit/sec, place field peaks that stayed within 10 cm, and fields either before or after stimulation with $> 2$ bits/spike; no peak rate criteria were adopted. The main findings were robust against selection of spatial information score and $LLratio$, as well as for a correlation based remapping metric described below.

A second method to quantify remapping was also adopted that uses more standard correlation of place field templates. Here, two templates were calculated, one for place field maps prior to stimulation (premap) and another for post-stimulation place field maps (postmap). Both templates were correlated (Pearson’s correlation coefficient) with single trial rate maps (premap corr/postmap corr), always excluding the correlated trial from the data from which the template was defined (i.e., the premap template is derived from data from trials 2-10 when correlated with trial 1). The mean difference between the template matches prior to stimulation, $\Delta\text{corr}_\text{prestim}$, gives a measure for predictiveness of poststimulation spatial tuning in generating the prestimulation rate maps. The main difference between the template matches post stimulation, $\Delta\text{corr}_\text{poststim}$ provides the converse measure, the predictability of prestimulation place field tuning for post stimulation activity. To avoid firing rate thresholds, we opted to include any place field map that showed trial-to-trial spatial reliability (mean premap corr. $> 0.25$ for trials before stimulation and mean postmap corr. $> 0.25$ for trials after stimulation). Trial-by-trial correlations were done in a similar manner where each trial’s firing rate map was correlated with each other and the rate of decorrelation (drift) was measured with or without intervening optogenetic simulation (Figure S4). Inhibitory cells were excluded from place field analyses.

**Bayesian decoding**

Bayesian decoding was conducted following Zhang et al. (1998). The decoding time bin was 100 ms. Cross validation was accomplished by constructing place fields that excluded activity recorded on the trial for which position was estimated. Only sessions in which more than 20 pyramidal cells were simultaneously recorded were considered.

**Predicting new field locations with RUSBoost**

We adopted the RUSBoost binary classification algorithm (Seiffert et al., 2008) to predict whether neural activity observed at each location on the track was recorded in the future place location. RUSBoost is designed to handle the class imbalance problem in data with discrete class labels, which we faced, as we defined latent place field locations as 10 cm segments of the track centered on the post-stim place field peak. The RUSBoost algorithm uses a combination of RUS (random under-sampling) and the standard boosting procedure AdaBoost, to better model the minority class by removing majority class samples. RUSBoost was implemented with MATLAB 2020a’s fitensemble algorithm to create 100 classification trees (without shrinkage) from the training set. The learned mapping was then used to predict whether withheld data was recorded in a future place field or not. The features that defined a location were the cell’s single-pass firing rate as observed for each 10 cm segment of the track, and the proportion of spikes observed in a burst. Ten random 50/50 splits of the pre-stimulation data were performed for the cross validation and the reported hit and false positive rates reflect the mean across independent cross-validations. A control bootstrap distribution was generated by shuffling
the position labels for the held-out data (100 shuffles). The positive likelihood ratio (hits divided by false alarms) was used to compare whether neurons did better than the bootstrap distribution (> 95% bootstrap) and whether bursts or firing rates were more predictive.

**Peer prediction analysis**

To predict the activity of one target cell from the population of peer cells, we used the generalized linear model approach of Harris et al. (2003). Spike trains of the target and peers were binned with different resolutions (20 bin widths, logarithmically spaced from 5-5000 ms). Ten-fold cross validation was used to build a training set with 90% of the data and a test set with the remaining 10%. The predicted intensity at a time \( t \) was given by

\[
\hat{f}_t = g \left( \sum_a S_{\alpha} w_a \right) \tag{Equation 11}
\]

The link function \( g(\eta) \) was defined as

\[
g(\eta) = \begin{cases} 
e^\eta, & \eta < 0 \\ \eta + 1, & \eta \geq 0 \end{cases} \tag{Equation 12}
\]

The prediction weights \( w_a \) were chosen to maximize the regularized log-likelihood on the training set:

\[
LL_{\text{peer}} = \sum_t -f_t dt + n_t \log (f_t) - \frac{1}{2} \sum_a w_a^2 \tag{Equation 13}
\]

where \( n_t \) is the number of spikes observed in that time bin.

The minimization of \( LL_{\text{peer}} \) was carried out using MATLAB 2018b’s fmincon function using the default Interior Point Algorithm and upper and lower weight bound of ± 100.

**Synapsemble analysis**

A wealth of *in-vitro* data suggests that the strength of synaptic coupling (e.g., magnitude of the evoked PSP) changes following different pairing protocols. Our goal was to capture related changes in our dataset by estimating long-term changes in spike transmission between monosynaptically connected pyramidal cells and interneurons (as defined by English et al., 2017; Figure S4). In order to do this, we model the postsynaptic spike train using a generalized linear model (GLM) with the following two features: (1) a coarsened, slowly-evolving version of the postsynaptic rate (baseline term) and (2) a transient boost following the presynaptic spike whose magnitude varies with time on a long timescale (coupling term). The features are summed and passed through an exponential nonlinearity to yield the instantaneous postsynaptic rate \( \lambda_{\text{post}}(t) \). The exponentiated coupling term can therefore be interpreted as a multiplicative, presynaptically-induced gain acting on an otherwise slowly evolving postsynaptic rate (the exponentiated baseline term). This separation of timescales assumption conveniently dodges the issue of capturing all parameters modulating the postsynaptic rate (e.g., theta, behavioral state). The conditional intensity function takes the following form:

\[
\lambda_{\text{post}}(t) = C(\text{Post}(t)) e^{\text{Pre}(t - \Delta) X_C(t) \cdot k} \tag{Equation 14}
\]

where Pre and Post are the pre- and postsynaptic spike trains binned at \( dt = 0.8 \) ms, ensuring that each bin has at most one spike. \( \Delta \) is the mode time lag between the pre- and postsynaptic spikes estimated from the raw CCG. \( C(\text{Post}) \) is the coarsened baseline rate obtained by counting postsynaptic spikes in 15 ms wide bins, expressing these as rates, and linearly interpolating at times corresponding to the centers of the dt-sized bins that were used to bin spikes. Note that in the absence of a presynaptic spike immediately preceding \( t \) (i.e., \( \text{Pre}(t - \Delta) = 0 \)), the predicted rate is equal to \( C(\text{Post}(t)) \). We model the slow changes in synaptic coupling \( X_C \cdot k \) with a linear combination of cubic B-splines with equally spaced knots. For each pair of neurons, the spacing of the knots (every 400 - 1000 s, in increments of 100) is selected by cross-validation using 100 s even/odd data splits. More specifically, the data are divided into 100 s data segments, such that odd segments are used for training, and even segments for testing. Because the parameters \( k \) are only constrained when \( \text{Pre}(t - \Delta) = 1 \), we fit the model in these bins exclusively. We use an LBFGS algorithm to minimize the convex negative log-likelihood with analytically computed gradients.

In order to explore the specific dynamics that govern changes in spike transmission, we made use of an extended GLM in which the coupling term takes the form of a dynamical model for short-term plasticity (Ghanbari et al., 2020; Tsodyks and Markram, 1997):

\[
\lambda_{\text{post}}(t_i + \Delta) = C(\text{Post}(t_i + \Delta)) e^{A \cdot w_i} \tag{Equation 15}
\]

where \( \lambda_{\text{post}}(t_i + \Delta) \) is the postsynaptic rate at monosynaptic lag \( \Delta \) following the \( i \)-th presynaptic spike. At all other times, \( \lambda_{\text{post}} = C(\text{Post}) \). \( A \) is the constant coupling strength, and \( w_i \) is a scale factor that depends on the history of presynaptic spiking through the Tsodyks-Markram (TM) model (Tsodyks and Markram, 1997), a nonlinear dynamical system to capture short term plasticity:

\[
w_i = R_i u_i \tag{Equation 16}
\]
was computed as,

\[ R_i = 1 - (1 - R_{i-1}(1 - u_{i-1})) e^{\frac{-t_i}{\tau_d}} \]  

(Equation 17)

\[ u_i = U + (u_{i-1} + f(1 - u_{i-1}) - U)e^{\frac{t_i}{\tau_f}} \]  

(Equation 18)

\[ \lambda_{\text{diff}} = \lambda_{\text{post}} - C(\text{Post}) \]  

(Equation 19)

In order to obtain a smoothly varying estimate of spike transmission (i.e., even at times when the presynaptic neuron is not spiking), \( \lambda_{\text{diff}} \) was convolved using a Gaussian kernel with a standard deviation of 120 s. In order to get a bin-by-bin estimate of spike transmission (in Hz) per presynaptic spike, we normalized the resulting time series by the presynaptic spike train convolved in the same manner.

In order to find ensembles reflecting higher order coactivations among the spike transmission time series (i.e., synapsemblies), we performed an unsupervised statistical analysis based on ICA (Lopes-dos-Santos et al., 2013; Peyrache et al., 2009; van de Ven et al., 2016). Briefly, spike transmission time series were represented as a matrix \( z \), which was \( z \)-scored and downscaled to 10 Hz. The number of synapsemblies in each session was based on the N principal components of the correlation matrix whose variances exceeded an analytical threshold based on the Marcenko-Pastur distribution describing variances expected for uncorrelated data. The high-dimensional activity matrix was projected onto the subspace spanned by these N principal components, and ICA was performed to extract synapsemblies (each corresponds to an independent component). The expression strength of each synapsemble was computed as,

\[ A(t) = z(t)^T P_i z(t) \]  

(Equation 20)

where \( P_i \) is the projection matrix (outer product, diagonal set to zero) of the i-th independent component. As such, \( A(t) \) quantifies the moment-to-moment similarity between an independent component (synapsemble template) and the instantaneous spike transmission pattern across all PYR-INT pairs.

In order to quantify synapsemble rearrangements across a relevant time point \( T \), we defined the change in synapsemble expression as the difference between mean expression strength around \( T \). For stimulation sessions, \( T \) was the first stimulation event on the track, while for controls, we selected the halfway point of the track running session. In each case, differences in mean expression strengths were based on time intervals that included entire homecage periods flanking the track period. Synapsemblies were detected either in the period before or following \( T \). A negative value for the change in expression strength reflects that the synapsemble is re-expressed less prominently following \( T \) (if it was detected prior) or before \( T \) (if it was detected after).

This analysis pipeline was also conducted for ensembles of presynaptic pyramidal neurons (PYR ensembles) or postsynaptic interneurons (INT ensembles). For each session, spike trains of neurons that figured in PYR-INT connections were binned at \( dt = 0.8 \) ms, and convolved with a Gaussian whose full-width at half maximum matched the timescale of the corresponding spike transmission time series as estimated from cross-validation in the long-term model. The resulting firing rate time series were \( z \)-scored, and the above described ensemble analysis was performed to estimate changes in rate ensembles around \( T \).

**GFP control**

Two adult male Tg(Camk2α-cre)T29-1Stl mice (The Jackson Laboratory; #005359; Tsien et al., 1996) were anesthetized and injected with a Cre-dependent adeno-associated virus (AAV) expressing EYFP (Addgene viral prep # 27056-AAV5; http://addgene.org/27056; RRID:Addgene_27056) delivered through a glass pipette (Sutter Instruments). Injections targeted right dorsal CA1 (–2 mm AP, 1.7 mm ML w.r.t bregma; 1.2 mm depth), and 300 nL of AAV-Ef1a-DIO EYFP (titer \( \geq 1 \times 10^{13} \) vg/mL) was injected at a flow rate of 50 nL/min (World Precision Instruments, UMP3 UltraMicroPump). Three weeks following virus injection, animals were re-anesthetized for head plate implantation. The skull was exposed, and a custom designed 3D printed head plate was attached with C & B Metabond Quick Adhesive Cement (Parkell). A single steel screw was implanted over the cerebellum as ground and reference. Lastly, a craniotomy was drilled over the site of the virus injection, and sealed with silicone adhesive (World Precision Instruments, Kwik-Sil) until the day of recording.

Over the course of one week, mice were habituated to head fixation while running on a custom designed running wheel. On the day of recording, Kwik-Sil was removed to expose the craniotomy, and a μLED-equipped silicon probe was slowly lowered into the brain until sharp-wave ripples were detectable on all four shanks. Following 10-20 minutes, a 1h recording session was performed, during
which light was delivered, one μLED at a time, to shanks with high single unit activity. 100ms square pulses with 1 s inter-pulse inter-
vals with 5 driving currents chosen to span the full range of light intensities used in mice expressing ChR2. μLEDs were driven using
an open-source μLED current controller (https://github.com/YoonGroupUmich/osc1lite). Following the acute experiment, mice were
perfused as described, and EYFP expression was verified using light microscopy (Olympus).

**Effect size estimates**
We follow the recommendation of Calin-Jageman and Cumming (2019) and the methods developed by Kruschke (2013) to report
effect sizes. We include an additional Bayesian statistical test for each comparison and report the highest density interval (HDI) of
the effect size that lies outside of zero. For comparisons with underlying skewed distributions, we model the data, y, as having derived
from a skewed, non-centered Student’s t-distribution.

**In JAGS notation**

\[
Y \sim \text{dt}(\mu_{\text{COND}} + \mu_{\text{MOUSE}}, \tau_{\text{CON}}, \nu_{\text{CON}}) \text{ for } \tau > 0 \text{ and } \nu > 0
\]  
\text{(Equation 21)}

where,

\[
Y = - U / \sqrt{V}
\]  
\text{(Equation 22)}

\[
U \sim \text{dnorm}(\mu_{\text{COND}} + \mu_{\text{MOUSE}}, \tau_{\text{CON}})
\]  
\text{(Equation 23)}

\[
V \sim \text{dgamma}(\nu_{\text{CON}} / 2, \nu_{\text{CON}} / 2)
\]  
\text{(Equation 24)}

The choice of \(\tau, \mu, \text{ and } \nu\) that best fits the data is found through Markov Chain Monte Carlo (MCMC) with weakly informed priors
as in (Kruschke, 2013). Inter-subject differences were modeled by fitting means that were a linear combination of effects driven by
random-effects for individual mice, \(\mu_{\text{MOUSE}}\), and the condition-specific effect of interest \(\mu_{\text{COND}}\) (e.g., \(\text{CON} = \text{stim} \text{ versus Control})).

For normal data, a similar approach was taken, though assuming

\[
y \sim \text{dnorm}(\mu_{\text{COND}} + \mu_{\text{MOUSE}}, \tau_{\text{CON}})
\]  
\text{(Equation 25)}

The distribution of credible parameters was identified for each group under comparison and the effect size calculated with three
MCMC chains, 1000 burn-in steps and five thinning steps (to avoid auto-correlations in the Markov process).