Recoding a cocaine-place memory engram to a neutral engram in the hippocampus

Stéphanie Trouche¹, Pavel V Perestenko¹, Gido M van de Ven¹, Claire T Bratley¹, Colin G McNamara¹, Natalia Campo-Urriza¹, S Lucas Black¹, Leon G Reijmers² & David Dupret¹

The hippocampus provides the brain’s memory system with a subset of neurons holding a map-like representation of each environment experienced. We found in mice that optogenetic silencing those neurons active in an environment unmasked a subset of quiet neurons, enabling the emergence of an alternative map. When applied in a cocaine-paired environment, this intervention neutralized an otherwise long-lasting drug-place preference, showing that recoding a spatial memory engram can alleviate associated maladaptive behavior.

Memory is central to behavior, allowing individuals to respond to their environment using past experiences. The spatially tuned and temporally coordinated activity of hippocampal CA1 principal cells contributes to adaptive memories by providing the brain with neuronal engrams that represent the spatial context of life events. Nonetheless, the recall of certain representations, such as those associating an environment with a drug of abuse, can impair the individual’s ability to make adaptive choices, including an uncontrolled drive to revisit drug-paired locations. Manipulating hippocampal neuronal representations could redress undesirable spatio-contextual behaviors. However, a given hippocampal ‘place cell’ can contribute to not one but multiple representations. Thus, it remains unknown how the hippocampal neuronal representation of a particular place could be selectively edited. In a given environment, the hippocampus exhibits, alongside the active place cells, a subset of quiet neurons without clear spatial selectivity. We used a c-fos–based optogenetic approach to silence hippocampal neurons active in a particular environment and examined whether quiet neurons could implement an alternative representation that neutralizes a drug-place memory.

To label and manipulate hippocampal neurons recruited during exploration of an environment, we bilaterally injected the dorsal CA1 of c-fos–tTA transgenic mice with a TRE3G-ArchT (Archaerhodopsin-T)-GFP viral construct (Fig. 1a). In these mice, neuronal activation leads to the expression of the tetracycline transactivator (tTA) through the promoter of the c-fos immediate early gene. The transient removal of doxycycline (Dox) from the mouse diet allows the tTA to interact with the tetracycline-responsive element (TRE3G), thereby tagging active neurons with the light-driven silencer ArchT. To test this tagging procedure, a first group of mice explored a circular-walled enclosure (the circle) while transiently off Dox (Fig. 1b). In these tagged mice, 6.28 ± 0.38% of neurons in the CA1 pyramidal cell layer expressed ArchT-GFP, of which 99.70 ± 0.10% were pyramidal cells (tagged neurons, n = 4 mice; Fig. 1c). This neuronal tagging lasted at least 6 d (n = 2 mice; Supplementary Fig. 1b) and was virtually absent in c-fos::ArchT control mice (n = 3) that were always fed Dox (0.09 ± 0.02%; P < 0.001 tagged versus control; Fig. 1a–c).

We next optogenetically silenced ArchT-GFP–tagged neurons while monitoring CA1 ensemble activity from c-fos::ArchT–tagged mice (n = 5; Fig. 1a,b). Tagged mice were transiently taken off Dox to tag CA1 neurons in the circle (day 0; Fig. 1b), and recordings were performed on the following days during re-exposure to the circle with CA1 light delivery (561-nm, 30-s light-ON pulses with 70-s light-OFF intervals; Fig. 1b). We found that ArchT photo-silencing substantially decreased the firing rate of 133 neurons (putative tagged neurons; OFF = 1.08 ± 0.08 Hz, ON = 0.06 ± 0.005 Hz) of the 1,083 principal neurons recorded (range: 161–231 daily recorded neurons over 6 d) and increased the firing rate of 332 neurons (alternative neurons’ OFF = 0.84 ± 0.04 Hz, ON = 1.97 ± 0.08 Hz; Fig. 1d,e). We did not observe such a light-induced firing switch amongst the 358 CA1 principal neurons recorded from c-fos::ArchT control mice continuously fed on Dox (n = 2; Fig. 1b and Supplementary Fig. 2a,b).

We then sought to identify functional differences between tagged and alternative neurons. We found that the baseline (light-OFF) firing rate of alternative neurons was initially lower in the circle than that of tagged and unchanged neurons (days 1–2; Fig. 2a and Supplementary Fig. 2d,e). In contrast, their firing rate matched that of the tagged and unchanged neurons during exploration of a square-walled enclosure mice also experienced each day (the square; Supplementary Fig. 2d,e). This suggests that the preferred recruitment of a selected subset of neurons in a given environment is associated with tight inhibitory control over other principal cells. Accordingly, the photo-evoked switch in activity of tagged and alternative neurons (Fig. 1e) was accompanied by a fine redistribution of interneuron firing activity, without a change in the interneuron population mean rate (n = 189 interneurons recorded; Supplementary Fig. 3).

During the following days, the baseline (light-OFF) activity of alternative neurons increased in the circle while that of tagged neurons concomitantly decreased (Fig. 2a), as revealed by the slope of the linear regression (firing rate = β0 + β1 × days; with β1 = 0.15 and −19 alternative and tagged neurons, respectively; P < 0.0001 for both). In these tagged mice, there was no similar rate shift of light-unresponsive neurons in the circle (unchanged neurons; Supplementary Fig. 4a,b), nor for tagged and alternative neurons in the square (P > 0.19 for both). Moreover, no shift in neuronal activity was observed when mice with a subset of neurons holding a map-like representation that neutralizes a drug-place memory.

1Medical Research Council Brain Network Dynamics Unit, Department of Pharmacology, University of Oxford, Oxford, UK. 2Department of Neuroscience, School of Medicine, Tufts University, Boston, Massachusetts, USA. Correspondence should be addressed to D.D. (david.dupret@pharm.ox.ac.uk) or S.T. (stephanie.trouche@pharm.ox.ac.uk).

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The emergence of spatially structured firing delivery strongly disrupted the place fields according to their light-modulation (as in $\not$ observed in $c$-fos $:\not$ArchT–tagged mice exploring the circle without CA1 light delivery ($n = 3$; Supplementary Fig. 4c,d). Thus, the repetitive photo-silencing of neuronal activity in the circle, but not context re-exposure per se, caused the tagged/alternative firing rate shift.

We next computed place firing rate maps in light-OFF and light-ON epochs. During the first days of re-exposure to the circle, tagged neurons, but not alternative neurons, exhibited clear light-OFF spatial firing fields (Fig. 2b,c and Supplementary Fig. 5). Although light delivery strongly disrupted the place fields of tagged neurons, it simultaneously led to the emergence of spatially structured firing of alternative neurons (Fig. 2c and Supplementary Figs. 5b,c and 6). This photo-evoked switch from one hippocampal map to another was not observed in $c$-fos $:\not$ArchT control mice (Supplementary Figs. 7 and 8), ruling out nonspecific remapping caused by light delivery. Moreover, the hippocampal map photo-switch was associated with a transient increase in locomotor reactivity, which diminished over the following days of light delivery, suggesting that the expression of the alternative representation caused tagged mice to initially treat the circle as a new enclosure (Supplementary Fig. 8). Notably, concurrent with this behavioral shift, the baseline spatial tuning of the alternative neurons increased, whereas that of tagged neurons decreased ($\beta_1 = 0.08$ and $-0.07$, respectively; $P < 0.00001$ for both; Fig. 2b,d and Supplementary Figs. 5 and 6), but no such change was observed in light-unresponsive neurons (Supplementary Fig. 4b).

Information processing in cortical circuits is thought to rely on spiking activity coordinated over short intervals in distributed groups of neurons forming assemblies$^5$. We tested whether our intervention changed the contribution of tagged and alternative neurons to the temporal expression of hippocampal assemblies. We detected assembly patterns from the co-fluctuation of (light-OFF) neuronal discharges in 25-ms windows, and described them by weight vectors representing the contribution of each neuron to each assembly pattern (Fig. 2e and Supplementary Fig. 9a–f). We found that, although tagged neurons predominantly contributed to assembly-pattern expression in the first 3-d block, this was reversed in the next 3-d block, which was marked by the dominant contribution of their alternative peers (Fig. 2f) without a change in the average strength of assembly activations (Supplementary Fig. 9g,h).

Finally, we investigated whether recoding a hippocampal map could reset the memory of an environment in which a drug had been previously experienced. Drugs of abuse, through their association with environmental cues and contexts, produce powerful and long-lasting memories that precipitate relapse$^6$. Concordantly, we found that more CA1 neurons recruited in a cocaine-paired environment were re-activated during drug-free re-exposure to that environment than for a saline-paired environment (Supplementary Fig. 10). We therefore tested whether our optogenetic intervention could re-wake the memory of a cocaine-paired environment. We trained mice in a conditioned place preference task to learn the association of one compartment with cocaine and the other with saline, such that they expressed a cocaine-place preference (test 1; Fig. 3). This cocaine-place association was long-lasting for both
Figure 2  Photo-silencing of tagged neurons enabled alternative neurons to emerge and provide an alternative map. (a,b) Baseline (light-OFF) firing rate (a) and spatial coherence (b) of tagged and alternative neurons from tagged mice exploring the circle (mean ± s.e.m.). *P < 0.05, **P < 0.01, ***P < 0.001; comparison to day 1. (c,d) Example raster plots and spatial rate maps of simultaneously recorded neurons on days 2 (c) and 6 (d). Distinct neurons were recorded on each day; one cell per row. Spatial rate maps are scaled to the peak firing rate (Hz; top right of each map) or 1 Hz for low-firing cells. Warm colors (red) correspond to the cell’s place field. (e) Example weight vectors representing simultaneously recorded assembly patterns detected from 25-ms co-fluctuation of (light-OFF) spike discharge in the circle. (f) Average single-neuron contribution to the temporal expression of assembly patterns during the first and the second 3-d blocks using 25-ms time windows (mean ± s.e.m.). **P < 0.01, ***P < 0.001.

c-fos::ArchT control mice (n = 8) and wild-type control littermates (n = 8) (test 2; Fig. 3). However, c-fos::ArchT–tagged mice (n = 13) no longer exhibited a cocaine-place preference (Fig. 3 and Supplementary Fig. 11) after the repetitive CA1 photo-silencing in the cocaine-paired compartment had shifted neuronal activity from the tagged to the alternative subset (Supplementary Fig. 11c). The cocaine-place preference of tagged mice could not be reinstated in response to cocaine-priming injection, suggesting that the shift to the alternative hippocampal map had neutralized the cocaine-place memory (Supplementary Fig. 11d). However, both tagged and control mice were able to express a place preference for a never before experienced compartment (Supplementary Fig. 11e,f). We therefore propose that the absence of spatial preference after hippocampal map recoding in tagged mice indicates that the compartment previously paired with cocaine was no longer associated with the drug experience and yet was perceived as familiar again.

Figure 3  CA1 photo-recording revoked an otherwise long-lasting cocaine-place memory. (a) Cocaine-conditioned place preference procedure. Mice explored the entire apparatus (pre-test) before the preferred compartment was paired with saline (+S) and the other with cocaine (+C) administration, followed by a place-preference test (test 1). Then tagged, but not control, c-fos::ArchT mice were transiently taken off Dox to tag active CA1 neurons and all mice received cocaine in the cocaine-paired compartment. CA1 light was delivered to all c-fos::ArchT mice during subsequent (drug-free) re-exposure to the cocaine-paired compartment, followed by a second place-preference test (test 2). (b) Example paths for c-fos::ArchT–tagged and control mice during pre-test, test 1 and test 2. (c) Cocaine-place preference score (mean ± s.e.m.). During each test, drug-free mice had access to the entire apparatus and their place preference was scored as the difference between the time spent in the cocaine-paired compartment minus that in the saline-paired compartment over their sum. Within-group comparison to pre-test: ***P < 0.001. Test 2, between-group comparison: ###P < 0.001.
Previous studies using c-fos–based optogenetic technology have influenced the behavioral output of a spatio-contextual memory by altering the interaction the hippocampus has with extra-hippocampal circuits that attach valence to context\(^{19,20}\). We developed an alternative approach that directly changed the hippocampal ensemble representation of a spatial context by disengaging the initially recruited neurons while enabling previously quiet neurons to emerge and provide an alternative representation. Drug-paired spatio-contextual memories increase the propensity for relapse in cocaine users and prompt drug-seeking behavior in laboratory rodents\(^6\). Our approach for editing the hippocampal ensemble representing an environment associated with cocaine was able to revoke drug-induced spatial preference in mice. In contrast with regular extinction, the neutralizing effect of this intervention was not overturned by a priming cocaine experience. Together, our data demonstrate that recoding a selective hippocampal engram represents a potent strategy to reset spatial memories and neutralize maladaptive behavior.

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

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**AUTHOR CONTRIBUTIONS**
S.T. and D.D. designed the experiments. P.V.P. cloned and tested the construct. S.T., N.C.-U., C.T.B. and P.V.P. carried out the experiments. S.T. and S.L.B. performed the cell counting. S.T., G.M.v.d.V., C.T.B., P.V.P., C.G.M. and D.D. analyzed the data. L.G.R. provided the mice and helped with the behavioral protocols. S.T. and D.D. wrote the manuscript. D.D. supervised the project. All of the authors discussed the results and commented on the manuscript.

**COMPETING FINANCIAL INTERESTS**
The authors declare no competing financial interests.

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ONLINE METHODS

Subjects. c-fos-tTA animals were adult transgenic mice heterozygous for the transgene carrying the c-fos promoter–driven tetracycline transactivator (tTA)1. Mice were bred from c-fos-tTA mice crossed with C57Bl/6j mice, and housed with their littersmates until used in the experimental procedure. Food containing doxycycline (the ‘regular Dox’, 40 mg per kg of body weight Dox chow; Bio-Serv) was provided for at least two weeks before the injection surgery; c-fos-tTA-LacZ animals were adult male double transgenic mice carrying the tetracycline transactivator (tTA) transgene under the control of the c-fos promoter, as well as a second transgene linking the teto promoter to the tau-LacZ reporter and the tetracycline transactivator mutant tTA1026Q (tTA′) in order to maintain the teto-linked gene expression after the initial neuronal activation. In the absence of doxycycline the active teto promoter forms a positive feedback loop with tTA′ that then becomes doxycycline insensitive. The c-fos-tTA-LacZ mice were backcrossed to a C57Bl/6j background and were raised on regular Dox. These two c-fos-tTA mouse lines were generated at The Scripps Research Institute and maintained at Tufts University until shipment to the MRC BNUU at the University of Oxford. Mice had free access to food and water ad libitum in a dedicated housing room with a 12/12-h light/dark cycle (7 a.m. to 7 p.m.). Behavioral experiments were performed during daylight. All experiments involving animals were conducted according to the UK Animals (Scientific Procedures) Act 1986 under personal and project licenses issued by the Home Office following ethical review.

Viral construct cloning, packaging and testing. The pAAV-TREG-ArchT-GFP plasmid was constructed using the pAAV-CamKII-ArchT-GFP backbone vector (Addgene plasmid #37807)23 where the original CamKII promoter was substituted, using PacI and the BamHI restriction enzyme recognition sites, with the third generation of tetracycline responsive element containing promoter (TRE3G, Clontech Laboratories). The recombinant AAV vector was serotyped and packaged with AAV2 coat proteins (University of North Carolina). Dox-controlled transcriptional activation of ArchT-GFP expression was first assessed in cultured HEK-293 cells (Supplementary Fig. 1a) co-transfected with a pTet-DualOFF vector (Clontech Laboratories Inc.; IRES2-ZsGreen1 cassette removed using EcoRI and HindIII restriction sites followed by Klenow fragment blunting and self-ligation of the vector). The ArchT-GFP expression was assessed in c-fos-tTA mice injected in the CA1 region of the dorsal hippocampus with the TREG3, Clontech Laboratories vector was delivered at a rate of 100 nl min−1 using a glass micropipette24. After 2 weeks of recovery mice were implanted with a ten-independently movable tetrode microdrive combined with two optic fibers (Doric Lenses) targeting the dorsal CA1 bilaterally24.

Recording procedures. Following the implant surgery, mice recovered for at least one week before familiarization to the recording procedure commenced. Mice were handled 5–10 min in a dedicated towel, twice a day, and connected to the recording system for 1–2 h a day for at least a week. During this time, tetrodes were gradually lowered toward the pyramidal cell layer. All cell ensemble recordings were performed from the CA1 hippocampus of adult male mice. On recording days, each morning tetrodes were lowered into the pyramidal cell layer in search of multi-unit spiking activity as previously described24. All recordings were performed under dim light conditions (20–40 lux) with no rewards provided. At the end of each recording day, tetrodes were raised to avoid damaging the pyramidal cell layer overnight. The primary recordings completed in this study were from drug-free c-fos:–ArchT tagged mice exploring the circle paired with CA1 light delivery (see below) and a square-walled enclosure (the square, 46 cm × 46 cm × 38 cm; ~30 min) without light delivery (n = 5 tagged light-IN mice). Wooden pieces were added on some of the square walls on each day to make it novel. In these drug-free experiments recordings were also performed from two control groups. The first control group consisted of c-fos:–ArchT control (non-tagged) mice exploring the circle paired with light delivery and the square without light delivery (n = 2 control light-IN mice) and the second control group consisted of c-fos:–ArchT mice exploring the circle without light delivery and the square with light delivery (n = 3 tagged light-OUT mice). The purpose of light delivery in tagged light-OUT mice was to allow the identification of tagged and alternative neurons without affecting their activity during re-exposure to the circle. Locomotor behavior of tagged light-IN mice was similar to that of both non-tagged light-IN and tagged light-OUT mice in the circle during baseline condition (distance traveled = 110.54 ± 7.90 m, 103.23 ± 5.78 m and 113.19 ± 3.01 m, respectively; animal speed = 5.26 ± 0.38, 4.91 ± 0.27 and 5.39 ± 0.14; P = 0.78). CA1 light delivery in tagged light-IN mice was however initially associated with an increased locomotor reactivity, indicative of a behavioral novelty response to the expression of the alternative hippocampal map (Supplementary Fig. 8). In experiments involving cocaine, recordings were performed in mice (n = 4 on C57Bl6 background) exploring an enclosure paired with a saline administration (0.9% saline, i.p.) followed by the exploration of another enclosure paired with a cocaine administration (10 mg per kg, i.p.; Sigma) (Supplementary Fig. 11a). In addition, to determine whether the shift in neuronal activity caused by the repetitive photo-silencing in the drug-free circle enclosure (Fig. 2) also occurred in a cocaine-paired environment recordings were performed in c-fos:–ArchT tagged mice (n = 2) during the conditioned place preference task (Supplementary Fig. 11c).

Doxycycline treatment. Tagged, but not control, c-fos:–ArchT mice were taken off regular Dox 3 d before the tagging day (Day 0), a period during which familiarization to the recording procedure was temporarily discontinued for both tagged and control mice. On the tagging day of the drug-free open-field experiment both tagged and control c-fos:–ArchT mice explored the circle for 45 min. For the CPP experiment, the least preferred enclosure was identified for each mouse during the pre-test and that enclosure was paired with cocaine and explored for 45 min on the tagging day (see below). All mice went back to their home cage immediately after the tagging and were fed with high dose of Dox. The following day mice returned to the regular Dox diet.

Multichannel data acquisition. The signals from the electrodes were buffered on the head of the animal (unity gain op-amps, Axona) and transmitted over a single strand of litz wire to a dual stage amplifier and band pass filter (gain 1,000, pass band 0.1 Hz to 5 kHz; Sensorium)23, or (in a second setup) the electrode signals were amplified, multiplexed, and digitized using a single integrated circuit located on the head of the animal (RHD2164, Intan Technologies; pass band 0.09 Hz to 7.60 kHz). The amplified and filtered electrophysiological signals were digitized at 20 kHz and saved to disk along with the synchronization signals from the position tracking and the laser activation. To track the location of the animal the three LED clusters were attached to the electrode casing and captured at 25 frames per s by an overhead color camera.

CA1 light delivery. A 561-nm diode pumped solid-state laser (Laser 2000, Ringstead) was used to deliver light to the dorsal CA1 (~15 mW) of c-fos:–ArchT–tagged and control mice through two lengths of optic fiber and a rotary joint with splitter (Doric Lenses)24. For the drug-free open-field experiment, light was delivered in the circle (light-IN) using trains of 30 pulses 30 s in duration (light-ON) with 70 s between pulses (light-OFF). As these experiments relied on the propensity of the mice to spontaneously explore, this light delivery protocol was chosen to ensure the mice evenly covered the circle in both light-OFF and light-ON epochs (that is, ~35-min and 15-min total exploration, respectively). CA1 photo-silencing was also performed in c-fos:–ArchT–tagged mice during the exploration of the square, that is after that of the circle (light-OUT); 1 s light pulses with 9 s between pulses, in order to optogenetically identify tagged and alternative neurons. For the CPP experiment, trains of 30 light pulses of 30-s duration were delivered during the 20-min re-exposure sessions to the cocaine-paired

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To quantify the extent to which hippocampal principal cells contribute to the formation of neuronal assemblies by synchronizing their firing activity with peer neurons, significant co-firing patterns were detected using an unsupervised statistical method based on independent component analysis (ICA) (Fig. 2ef and Supplementary Fig. 9)\textsuperscript{33}. Light-Off epochs during exploration of the circle were divided into 25-ms time bins\textsuperscript{34}, and for every principal cell the number of spikes per bin was counted. Time windows matching theta-band (4–12 Hz) oscillatory waves (~100 ms) were also used as they represent another natural temporal division for the expression of hippocampal cell assemblies (Supplementary Fig. 9f). To restrict the analysis to the temporal coordination of spiking activity without bias due to differences in the average firing rates, the binned spike-counts were normalized for each neuron by a z-score transform

\[ z_{i,b} = \frac{x_{i,b} - \mu_{i}}{\sigma_{i}} \]

where for neuron \( i \), \( z_{i,b} \) is its z-scored spike-count in bin \( b \), \( x_{i,b} \) its spike-count in bin \( b \), \( \mu_{i} \), its mean spike-count across all bins and \( \sigma_{i} \), the s.d. of its spike-counts. This set each neuron to have a null mean rate with unitary variance. With the number of principal cells denoted by \( n \) and the number of time-bins denoted by \( B \), let \( Z \) be the \( n \times B \) binned and z-scored spike-count matrix with element \((i, b)\) equal to \( z_{i,b} \). From this matrix \( Z \) assembly patterns were identified in two steps: (1) the number of significant patterns in the data was estimated based on random matrix theory, after which (2) the corresponding number of assembly patterns was extracted using ICA. Note that here a pattern is described by a weight vector over the \( n \) principal cells recorded that day. In the data set used here the average number of principal cells was 30.11 ± 2.51 per recording day.

**Determination of the number of significant assembly patterns.** A principal component analysis was first applied to matrix \( Z \). The pairwise correlation matrix of \( Z \) is given by \( C = \frac{1}{n}ZZ^{T} \) (where \( Z^{T} \) is the transpose of \( Z \)) and the eigenvalue-decomposition of \( C \) is given by

\[ \sum_{j=1}^{n} \lambda_{p_{j}p_{j}}^{T} = C \]

where \( \lambda_{j} \) is the \( j \)th eigenvalue of \( C \) and \( p_{i} \) its corresponding eigenvector (that is, principal component of \( Z \)) for \( j = 1, \ldots, n \). To estimate the number of significant patterns embedded within \( Z \), the Marčenko-Pastur distribution was used\textsuperscript{35}, which states that the eigenvalues of the correlation matrix obtained from a \( Z \) matrix whose elements are independent and identically distributed random variables (with zero mean and unit variance) are asymptotically (that is, when \( n, B \rightarrow \infty \) such that \( B/n \) converges to a finite positive value) bounded to the interval

\[ \left[ 1 - \frac{1}{nB}, 1 + \frac{1}{nB} \right] \]

This implies that none of the eigenvalues of the correlation matrix of a \( Z \) matrix constructed from the spike trains of neurons with independent firing activity is expected to exceed \( \lambda_{\text{max}} = \left( 1 + \frac{1}{nB} \right)^{2} \), as demonstrated elsewhere (for \( B > n \))\textsuperscript{36,37}. An eigenvalue above \( \lambda_{\text{max}} \) thus indicates that the pattern given by the corresponding principal component captures more correlation in the data than any pattern could if the firing activity of all neurons was independent. The number of eigenvalues above \( \lambda_{\text{max}} \) (denoted by \( N_{A} \)) therefore represents the minimum number of distinct significant patterns in the data\textsuperscript{33,38}.

**Identification of assembly-pattern composition using ICA.** The first \( N_{A} \) principal components each capture a significant amount of correlation between the firing patterns of the neurons. However, principal components are restricted to be orthogonal to each other while cell assemblies do not need to be (for example, they can contain overlapping neurons). Principal components are also extracted from the data sequentially, which usually causes the first principal component to seemingly be a mixture of multiple assembly patterns\textsuperscript{36,39} (our own observation not shown). Moreover, principal component analysis is solely based on pairwise correlations while higher-order correlations can also inform assembly identification. To avoid these issues, ICA was used instead. The ICA extracts patterns such that the linear projections of the data onto these patterns are as independent as possible.
from each other as possible. However ICA directly applied to Z without prior dimension reduction would extract as many patterns as there are neurons. To restrict the number of patterns identified by ICA to N_H, the data was first projected onto the subspace spanned by the first N_H principal components

\[
\mathbf{Z}_{\text{PROJ}} = \mathbf{P}_{\text{SIGN}} \mathbf{T} \mathbf{Z}
\]

where \( \mathbf{P}_{\text{SIGN}} \) is the \( n \times N_H \) matrix with the first \( N_H \) principal components as columns. ICA was then applied to the matrix \( \mathbf{Z}_{\text{PROJ}} \). That is, an \( N_H \times N_H \) un-mixing matrix \( \mathbf{W} \) was found such that the rows of the matrix \( \mathbf{X} = \mathbf{W}^{\top} \mathbf{Z}_{\text{PROJ}} \) were as independent as possible. This optimization-problem was solved using the fast ICA algorithm. The resulting un-mixing matrix \( \mathbf{W} \) was then expressed in the original basis spanned by all the neurons

\[
\mathbf{V} = \mathbf{P}_{\text{SIGN}} \mathbf{W}
\]

where the columns of \( \mathbf{V} \) (that is, \( v_1, \ldots, v_{N_H} \)) are the assembly patterns. As both the sign and the scale of the ICA output is arbitrary, all assembly patterns were scaled to unit length (that is, \( \sum_i v_i^T v_i(i) = 1 \)) and their sign set such that the highest absolute weight of each pattern was always positive. The contribution of each neuron toward the temporal expression of an assembly pattern was taken as its squared weight (25-ms time bins: days 1–3: 436 and 1,312 patterns; days 4–6: 162 and 411 patterns; theta-cycle time-bins: days 1–3: 380 and 1,128 patterns; days 4–6: 123 and 438 patterns; with tagged and alternative neurons, respectively).

**Tracking the activation-strength of assembly patterns over time.** To determine whether the optogenetic manipulation altered the strength of the expressed assemblies, we tracked each assembly pattern \( v_k \) over time by

\[
R_k(t) = z(t)^\top \mathbf{P}_k z(t)
\]

where \( z(t) \) is a smooth vector-function containing for each neuron its z-scored instantaneous firing-rate and \( \mathbf{P}_k \) is the matrix projecting \( z(t) \) to the activation-strength of assembly pattern \( k \) at time \( t \). \( \mathbf{P}_k \) was taken as the outer product of \( v_k \) with the main diagonal entries set to zero to prevent high activation-strength caused by the isolated activity of a single neuron with high weight to that pattern. With this approach, only interactions between neurons could contribute toward the activation-strength of an assembly pattern. To increase the temporal resolution beyond the bin-size used to identify the assembly patterns, \( z(t) \) was obtained by convolving the spike-train of each neuron with a kernel-function after which the resulting smooth curve was normalized by a

\[
\text{instantaneous firing-rate and } \mathbf{w} \text{-scored } z(t)
\]

**Conditioned place preference (CPP).** The CPP apparatus consisted of a circle (46-cm diameter, 38-cm height) and a square (46 cm × 46 cm × 38 cm) compartment connected via a bridge (8-cm length, 5-cm width), which was present during both conditioning sessions each day. Conditioning was performed with respect to the initial preference for one of the two compartments (pre-test). Then, mice were conditioned for three days (days 1–3) with two pairing sessions each day. Conditioning was performed with respect to the initial preference of each animal (as identified during the pre-test) for one of the two compartments. In the first session, mice received saline (0.9% saline, i.p.) before exploring the preferred compartment for 20 min (saline-paired compartment). In the second session, 5 h later, mice received cocaine (10 mg per kg, i.p.; Sigma) before exploring the least-preferred compartment for 20 min (cocaine-paired compartment). On the day after (22 h after) the last conditioning session the cocaine place memory was assessed by allowing (drug-free) mice to explore the entire apparatus for 15 min (test 1; day –3). Dox was subsequently removed from the diet of tagged mice, but not control mice. Mice were injected with cocaine (10 mg per kg, i.p.; Sigma) before re-exposure to the cocaine-paired compartment for the tagging procedure (45 min; day 0). Both tagged and control mice were put on high Dox immediately after, and on regular Dox the next day. During each one of the three following days (days 1–3) mice were successively re-exposed to the saline-paired compartment without photo-silencing (20-min exploration) and to the cocaine-paired compartment 5 h later with ArchT photo-silencing (20-min exploration). The locomotor behavior of control and tagged c-fos::ArchT mice during re-exposure to the cocaine-paired enclosure was similar (distance traveled = 48.51 ± 3.32 m versus 54.26 ± 2.74 m, animal speed = 4.04 ± 0.27 cm s⁻¹ versus 4.52 ± 0.23 cm s⁻¹; \( P = 0.18 \)). The cocaine place memory was re-assessed by allowing mice to explore the entire apparatus for 15 min (test 2, day 4). For each mouse the time spent in each compartment was measured during the pre-test and the two tests sessions and the cocaine place preference score calculated as the difference between the time spent in the cocaine-paired compartment minus that in the saline-paired compartment over their sum. Two additional experiments using the CPP apparatus were performed. The first was to test the ability of the mice to subsequently reactivate a cocaine place preference when cocaine-primed (10 mg per kg, i.p.; Sigma; Supplementary Fig. 11d) and the second was to test their ability to detect spatial novelty (Supplementary Fig. 11e). Because control mice exhibited a cocaine place preference in test 2 (see Fig. 3c), we used a trial-to-criterion approach to extinguish their place preference before the drug-primed reinstatement test by repeatedly testing their place preference until their CPP score was no more than 0.04 (number of additional tests: 3.50 ± 0.88).** The drug-priming test was performed 24 h after the extinction criterion was reached for control mice or 24 h after test 2 for tagged mice, then followed by the spatial novelty test. For each mouse the time spent in each compartment was measured during both the drug-priming and the novelty tests. The cocaine place preference score was calculated as mentioned above, and the novel place preference was scored by the time spent in the new compartment minus that in the other (familiar) compartment over their sum.

**Cocaine place memory retrieval in c-fos-tTA-LacZ mice.** A group of c-fos-tTA-LacZ adult male mice was used to evaluate the extent to which dorsal CA1 neurons that were activated during the exploration of the tagged open-field enclosure associated with a cocaine (\( n = 7 \) mice) or a saline (\( n = 4 \) mice) administration were subsequently re-activated during a (drug-free) spatial context re-exposure (Supplementary Fig. 10). Mice were individually housed for at least two weeks (and regular Dox was removed 3 d) before the tagging day. The tagging day consisted of injecting mice with either saline (0.9% saline, i.p.) or cocaine (10 mg per kg, i.p.; Sigma) before they were allowed to explore the open-field enclosure for 30 min. A high dose of Dox (1 g per kg) was provided immediately after the exploration session before returning to regular Dox 1 d after. Drug-free mice were re-exposed to the tagged enclosure 3 d later and perfused 90 min after. Brains were extracted for immunohistochemistry procedures (Supplementary Fig. 10).

**Tissue processing and immunohistochemistry.** At the completion of the experiments all mice were deeply anesthetized with pentobarbital and transcardially perfused with 0.1 M phosphate buffer saline (PBS) followed by cold 4% paraformaldehyde (PFA) dissolved in PBS. Brains were extracted and kept in 4% PFA for at least 24 h before slicing and sections were stored in PBS-azide. For immunostaining, free-floating sections were rinsed extensively in PBS with 0.25% Triton X-100 (PBS-T) and were blocked for 1 h at −20 °C in PBS-T with 10% normal donkey serum (NDS). Sections were then incubated with primary antibodies diluted in 3% NDS blocking solution and incubated at 4 °C for 72 h (rabbit polyclonal anti-neuropod peptide tyrosine (NPY) 1:1,500, Abcam), goat polyclonal anti-parvalbumin (PV; 1:4,000, Swant), rat monoclonal anti-somatostatin (1:1,000), mouse monoclonal anti-beta-galactosidase (1:10,000, Promega), rabbit polyclonal anti-c-Fos (1:10,000, Santa Cruz). All sections were rinsed three times for 15 min in PBS-T and incubated for 4 h at −20 °C in secondary antibodies (Jackson ImmunoResearch; donkey anti-rat AMCA (1:250), donkey anti-rabbit Cy3 (1:1,000), donkey anti-mouse 647 (1:500), donkey anti-goat 647 (1:500)) diluted in the blocking solution. This step was followed by three rinses for 15 min in PBS-T. Sections were then incubated for 1 min with 4’, 6-diamidino-2-phenylinodole (DAPI, 0.5 µg ml⁻¹, Sigma, D8417) diluted in PBS to label cell nuclei before undergoing three additional rinse steps of 10 min each.
in PBS. Sections were mounted on slides, coverslipped with Vectashield mounting medium (Vector Laboratories) and stored at 4 °C.

**Image acquisition and quantification.** Images shown in Figure 1c and Supplementary Figure 10c were acquired using a laser-scanning confocal microscope (LSM 710; Zeiss, 40× /1.3 objective) in sequential scanning mode. A single 1-μm confocal plane was used for DAPI pictures with flattened z-stacks (2-μm step) for the GFP, LacZ or Fos signals. Series of tiled single-plan GFP and DAPI images shown in Supplementary Figure 1b were acquired using an epifluorescence microscope (AxioImager M2; Zeiss, 10× /0.45 objective) and Stereoinvestigator software (Virtual Tissue 2D module, MBF Bioscience). Counting was performed on z-stacks using Stereoinvestigator (MBF Biosciences) and ImageJ (http://rsb.info.nih.gov/) software. An experimenter blind to each condition outlined the pyramidal cell layer of the dorsal CA1 according to the DAPI signal in each brain section. The experimenter estimated the total number of DAPI-positive cells in the CA1 pyramidal cell layer by randomly outlining nine CA1 sample areas on sections from a subset of c-fos::ArchT mice (n = 3). In Figure 1, quantification of ArchT-GFP positive neurons in the pyramidal cell layer was conducted from one-in-three coronal sections (30-μm thickness, average of n = 11.14 ± 1.31 CA1 hippocampi per mouse) throughout the dorsal CA1 (−1.46 to −2.46 mm from bregma) from tagged (n = 4) and control (n = 3) mice perfused on day 3. In each hippocampal section, the percentage of ArchT-GFP cells expressing the different interneuron markers (PV, NPY and somatostatin) was evaluated. Putative pyramidal cells were required to exhibit dendritic spines (Fig. 1c) and be immunonegative for PV, NPY and somatostatin, markers known to capture interneuron.27,28,47,48 In Supplementary Figure 10, quantification of LacZ-positive and Fos-positive neurons in the pyramidal cell layer was conducted from one-in-seven coronal sections (30-μm thickness, average of n = 11.91 ± 0.45 CA1 hippocampi per mouse) throughout the dorsal CA1 (−1.31 to −2.53 from bregma) from saline (n = 4) and cocaine (n = 7) mice.

**Statistical analysis.** Data are presented as mean ± s.e.m. All P values were calculated in R (https://www.r-project.org/) using a two-sided t test or an ANOVA followed by a post hoc Tukey’s test for multiple comparisons, unless specified otherwise.

A **Supplementary Methods Checklist** is available.

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