

Early changes in the properties of CA3 engram cells explored with a novel viral tool

Reviewed Preprint

v1 • February 24, 2025

Not revised

Dario Cupollilo, Noëlle Grosjean, Catherine Marneffe, Julio Viotti, Célia Reynaud, Séverine Deforges, Mario Carta, Christophe Mulle 

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France •

Neuroscience and Brain Technologies Department, Fondazione Istituto Italiano di Tecnologia, Genova, Italy

 https://en.wikipedia.org/wiki/Open_access

 Copyright information

eLife Assessment

This **important** study characterizes and validates a new activity marker - fast labelling of engram neurons (FLEN) - which is transiently active and driven by cFos, allowing the monitoring of intrinsic and synaptic properties of engram neurons shortly after the learning experience. The results **convincingly** demonstrate the utility of this novel viral tool for studying early changes in the properties of engram cells. However, the study would benefit from exploring how accurately FLEN reflects endogenous cFos activity, how this labelling technique compares to previous versions, and from careful consideration of alternative explanations such as changes in release probability.

<https://doi.org/10.7554/eLife.105452.1.sa2>

Abstract

Forming new memories after a one-time experience requires initial encoding then consolidation over time. During learning, multimodal information converges onto the hippocampus, activating sparse neuronal assemblies which are thought to form a memory representation through concerted activity and synaptic interconnectivity. In this work, we use a novel tool for fast-labeling of engram neurons (FLEN). FLEN is based on c-Fos activity-dependent transient expression of a destabilized fluorescent marker ZsGreen1 rapidly after one-trial learning. With FLEN, we explore the electrophysiological properties of c-Fos activated CA3 pyramidal neurons a few hours following one-trial learning of an episodic-like memory. In parallel, we employ the Robust Activity Marker (RAM) system, which provides activity-dependent labelling 24 hours following a novel experience. Comparing FLEN+ and RAM+ neurons allows to characterize how the properties of neuronal assemblies evolve during an initial phase of consolidation. Whereas no difference was observed in the excitability of FLEN+ vs. FLEN-neurons, RAM+ neurons were more excitable than RAM-neurons. This suggests that CA3 pyramidal neurons recruited in an engram progressively acquire increased excitability as compared to neurons which were not activated by the one-trial contextual memory task. In contrast, FLEN+ CA3 neurons show an increased number of excitatory inputs. Overall, with the FLEN strategy, we can show that both the intrinsic

excitability and the synaptic properties of CA3 pyramidal neurons undergo progressive plastic changes over the first day following a one-trial memory task.

Introduction

Animal survival depends on the ability to rapidly form new and durable episodic memories of one-time meaningful events which, when recalled, allow to adapt behavior (Tulving, 2002). Contextual memories of ongoing experiences are continuously and rapidly generated in the hippocampus and subsequently stored in an extended cortical network (Eichenbaum, 2000). Computational models of memory point to a crucial role of the hippocampal CA3 region in episodic memory encoding and retrieval (Kesner & Rolls, 2015). Based on its distinctive extensive recurrent architecture (CA3-CA3), CA3 may act as an auto-associative or attractor network, enabling pattern completion during recollection (Kesner & Rolls, 2015; McNaughton, 1987). Indeed, mice with CA3-specific synaptic plasticity impairment (CA3 NR1-KO mice) are unable to encode the novel location of a hidden platform in a delayed matching-to-place version of the Morris water maze (K. Nakazawa et al., 2003) and to retrieve the learned position of a submerged platform when partial cues are presented (K. Nakazawa et al., 2002). In addition to recurrent connections, the main excitatory synaptic drivers of CA3 pyramidal neurons (PNs) are inputs from medial and lateral entorhinal cortex (MEC and LEC) via the perforant pathway (PP), and indirectly from mossy fibers (Mfs) through the dentate gyrus (DG). Mf inputs display ‘conditional detonator’ properties, enabling individual Mf synapses to trigger postsynaptic firing in response to repetitive firing of presynaptic DG cells (Henze et al., 2002; Marneffe et al., 2024; Rebola et al., 2017; Sachidhanandam et al., 2009; Vandael & Jonas, 2024).

During encoding of novel information, the orchestrated activity patterns of selected neuronal groups distributed throughout the brain, called engrams, represent the initial memory trace (Josselyn & Tonegawa, 2020). Engrams undergo long-lasting synaptic modifications during consolidation and are selectively re-engaged when the memory is retrieved (Bessi eres et al., 2024; Goto et al., 2021; Josselyn & Tonegawa, 2020; Kitamura et al., 2017; Ryan et al., 2015; Tom   et al., 2024). Experimental manipulation of neuronal engrams in the hippocampus allows selective memory erasure (Han et al., 2009), artificial memory recall (Liu et al., 2012) and creation of a synthetic memory (Vetere et al., 2019), highlighting both their necessity and sufficiency for memory functions.

The investigation of the properties of engram neurons has taken advantage of immediate early genes (IEGs) such as c-Fos, Arc (activity-regulated cytoskeleton-associated protein), or *Npas4* (neuronal PAS domain protein 4) to visualize and label neurons active during a memory task (DeNardo et al., 2019; Kawashima et al., 2013; Reijmers et al., 2007; S orensen et al., 2016; X. Sun et al., 2020). Generally, IEGs are rapidly and transiently expressed; however different IEGs are triggered by different sets of extracellular events and possess specific expression kinetics. Temporal control of IEG-based systems is crucial to accurately link engram labeling to specific events that occur in a specific in time.

The activity of engram neurons can be tracked *in vivo* during their maturation from the process of encoding throughout consolidation in behaving mice, by means of functional indicators like GCaMP (Milczarek et al., 2018; Mocle et al., 2024; Tom   et al., 2024). However, *in vivo* Ca²⁺ imaging does not allow to address the synaptic and intrinsic properties of engram neurons. Labeling of engram neurons in order to characterize these properties *ex vivo* over time after an experience has occurred, often relies on labeling techniques using inducible transgenic systems (DeNardo et al., 2019; Reijmers et al., 2007). In several studies using transgenic mouse lines, engram labeling requires 48 hours of doxycycline (Dox) deprivation before a novel experience (Kitamura et al., 2017; Liu et al., 2012; Pignatelli et al., 2019; Ryan et al., 2015), or the use of tamoxifen injection to open a 12-hours time window for engram labeling (DeNardo et al.,

2019 [Cupolillo et al., 2019](#)). Therefore, these systems only offer temporal control over extended time windows, thus increasing the probability of compromising the experience specificity of labeled engram cells. In addition, most of these approaches drive a relatively late labeling, enabling investigation of learning-induced modifications more than 24 hours after a novel experience (e.g RAM system) (X. Sun et al., 2020 [Sun et al., 2020](#)) or shortly after recall for neurons which have been labeled several days before (Pignatelli et al., 2019 [Pignatelli et al., 2019](#)), when consolidation processes have already been engaged.

Little has been learned about how neuronal engrams in the hippocampus rapidly (in the time range of hours) adapt their synaptic and physiological properties following encoding of a memory event, and how these properties evolve over time. In this sense, it is unknown whether initial memory storage is enabled by engram cell modifications of intrinsic properties, synaptic plasticity or a combination of them. Here, we tackle these questions using a combination of viral strategies that allow us to identify engram neurons and compare *ex vivo* their properties shortly after learning (3-4 hours), and 24 hours after the experience has occurred (**Fig. 1A** [Cupolillo, 2021](#)). To track early engrams, we developed a virally delivered c-Fos-based genetic construct that provides physiological-like rapid expression and fast-decay of the bright fluorescent cytosolic marker ZsGreen1. This novel viral strategy for fast and efficient labeling of engram neurons (FLEN) allows to identify and characterize engram neurons within 3-4 hours following a novel experience, in reference to non-activated neurons. Then, to understand how initial engram cell properties evolve within a 24-hour time scale, we employ the more durable c-Fos-based artificial RAM system (Sørensen et al., 2016 [Sørensen et al., 2016](#)) to label neurons and characterize their properties 24 hours after the exposure to a one-trial memory task.

Results

Creation and validation of c-Fos-ZsGreen1-DR for fast-labeling of engram neurons (FLEN)

We generated a genetic cassette for adeno-associated virus-mediated delivery which allows fast labelling of engram neurons (FLEN). This construct includes the full-length, activity-dependent c-Fos promoter which controls the expression of a destabilized variant of the green fluorescent protein ZsGreen1 (Matz et al., 1999 [Matz et al., 1999](#)) (**Fig. 1B** [Cupolillo, 2021](#)). The complete sequence of the c-Fos promoter ensures a physiological-like gene expression of ZsGreen1, which follows c-Fos expression. ZsGreen1 provides a bright whole-cell somatic and dendritic labelling of activated neurons (**Fig. 1B** [Cupolillo, 2021](#)). To restrict the labelling time window and minimize labeling unrelated to the designed experience, ZsGreen1 was fused with a destabilizing domain (DR); when fused to EGFP, DR decreases the fluorescence half-time to 2 hours (Li et al., 1998 [Li et al., 1998](#)).

To determine the expression kinetics of FLEN in response to neuronal activity, P0 mouse cortical neurons were cultured (cellular density 3×10^5 cells/well) and then co-infected at DIV3 with AAV9-c-Fos/ZsGreen1-DR (FLEN) and a secondary AAV9-CB7/mCherry, which constitutively expresses mCherry in infected neurons (**Fig. 1B-C** [Cupolillo, 2021](#)). Neurons were cultured until DIV11 in regular neurobasal medium, before switching to an activating medium containing 4-aminopyridine (4-AP, 100 μ M) and bicuculline (Bic, 10 μ M) for 30 minutes. The drugs were then washed out, and tetrodotoxin (TTX, 0.5 μ M) was added to block further neuronal activity until cells were fixed (**Fig. 1C** [Cupolillo, 2021](#)). The percentage of activated neurons (ZsGreen+, further named FLEN+ neurons) over the total number of transduced neurons (mCherry+ neurons) was calculated at different time points within a 6-hour period after pharmacological stimulation. Only a limited number of FLEN+ neurons could be found in unstimulated conditions (5.4 ± 0.5 FLEN+/FLEN-, $n=3$ plates) (**Fig. 1D-E** [Cupolillo, 2021](#)). The activating medium triggered ZsGreen1 expression in neurons as early as 2 hours, reaching a 3-fold peak within 3 hours (1 hour: 6.7 ± 1.0 , $n=3$; 2 hours: 13.2 ± 5.0 , $n=3$; 3 hours: 15.7 ± 2.2 , $n=5$) (**Fig. 1D-E** [Cupolillo, 2021](#)). The number of FLEN+ neurons decreased after the peak, decaying to pre-

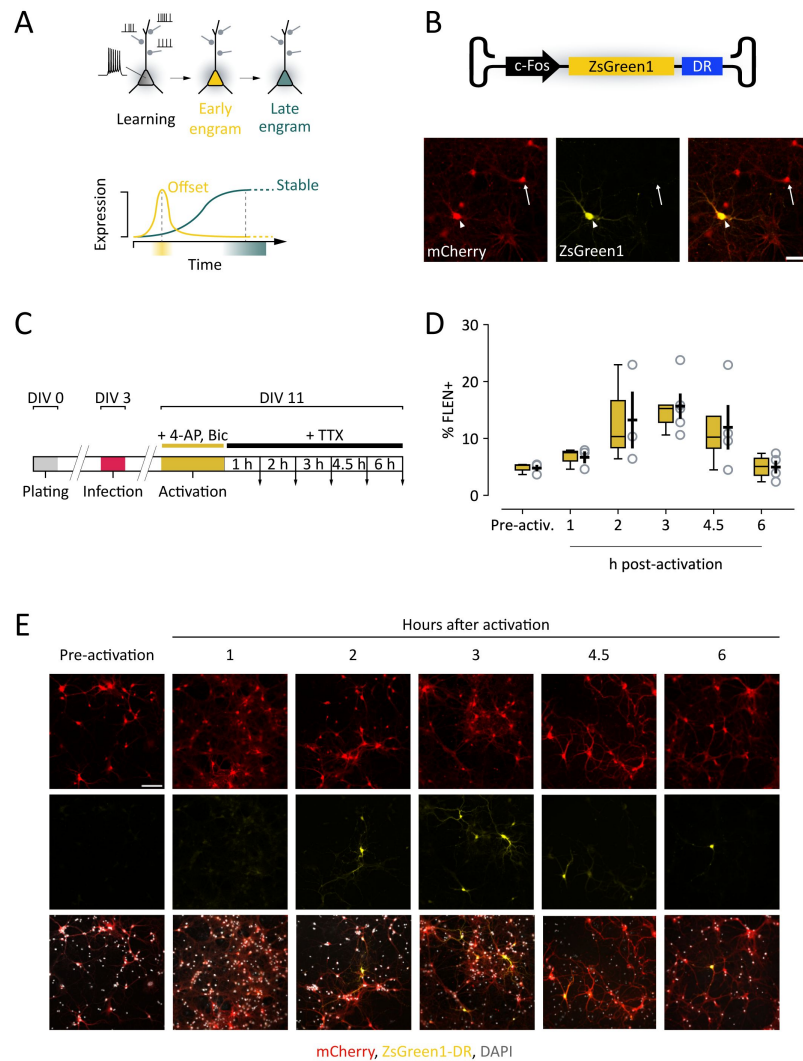


Figure 1.

Development of a novel viral tool to study early changes in the properties of CA3 engram cells.

(A) Scheme representing the maturation of engram neurons during memory formation. Top panel: during learning, a neuron receives numerous synaptic inputs, resulting in sequences of action potentials. This activity triggers the rapid expression of immediate early genes (IEGs), marking the neuron as part of an engram (yellow). As the engram matures (green), IEGs expression is not detectable anymore, yet the neuron retains established engram properties. Bottom panel: expression function of IEG-based construct to investigate and compare early and late engram properties. A construct with rapid offset (yellow) best captures engrams shortly after learning, whereas a progressively expressing construct best captures more mature engrams. **(B)** FLEN construct (c-Fos.ZsGreen1-DR) encapsulated in AAV2/9 vector. Bottom panel: representative images showing ZsGreen1 fluorescence in c-Fos+ neurons (arrowhead) and c-Fos-neurons (arrow). Scalebar = 50 μ m. **(C)** Timeline of *in vitro* experiments to determine FLEN expression and offset time course. After infection (DIV3) and activation (DIV11), samples are washed with a blocking medium and images are acquired at different intervals (downward arrows). **(D)** Percentage of ZsGreen+ vs. all transduced neurons (mCherry+ cells) at the different intervals following activation. **(E)** Representative images of activated cultured neurons at 1, 2, 3, 4.5 and 6 hours after activation. Scalebar = 100 μ m.

stimulation-like levels within 6 hours (4.5 hours: 12.0 ± 3.9 , $n=4$; 6 hours: 5.0 ± 1.1 , $n=4$) (**Fig. 1D-E**). Thus, FLEN showed a rapid induction (2-3 hours) and a fast decay (within 6 hours) of ZsGreen1-DR fluorescence (one-way ANOVA, $p=0.04$).

FLEN captures early neuronal activity following CFC

We next investigated the time course of FLEN expression *in vivo* in wild-type mice in response to exposure to a salient novel experience. We virally targeted bilaterally dorsal CA3 in adult mice (2–3-month-old) co-injecting AAV9-c-Fos/ZsGreen1-DR and the infection control AAV9-CB7/mCherry (**Fig. 2A**). Then, mice were subjected to contextual fear conditioning (CFC), a hippocampal-dependent one-trial memory task. During this task, mice learned to associate the delivery of 3 mild foot-shocks (0.4 pA, 2 seconds) with a specific environment (**Fig. 2B**). We found that CFC, a one-trial experience, can generate a strong context-specific fear memory, as mice responded by freezing when re-exposed to the conditioning environment 1 hour after training (HC: $14.7\% \pm 2.3\%$, 1-hour post-CFC: $47.4\% \pm 7.9\%$, $n=7$, unpaired t-test, $p=0.0018$), a response that outlasted 24 hours after training, but was not triggered in an unfamiliar novel environment, nor when they were left undisturbed in their home cage (HC) (HC: $18.7\% \pm 3.5\%$, 24 hours post-CFC: $61.5\% \pm 7.5\%$, neutral: $14.3\% \pm 4.5\%$, $n=8$, one-way ANOVA, Tukey's post-hoc test HC : 1 hour post-CFC $p=0.00004$, HC : Neutral $p=0.84116$, Neutral : 1 hour post-CFC $p=0.00001$) (**Fig. 2B**).

The percentage of FLEN+ CA3 neurons (i. e. that display FLEN+/mCherry+ labeling, **Fig. 2C**) among the overall population of infected neurons (FLEN-, i. e. that display mCherry+ labeling) was quantified 1.5, 3, 4.5, 6 and 12 hours following CFC and compared to that of untrained mice. Consistent with previous studies (Sørensen et al., 2016; Weng et al., 2018), less than 2% of infected CA3 neurons were reported to be labeled in the absence of a novel experience (**Fig. 2D-E**, HC, $0.84\% \pm 0.29\%$ FLEN+/mCherry+, $n=6$). One-trial CFC conditioning generated an engram, as indicated by an increase in FLEN+ neurons in CA3 in the following hours (**Fig 2D-E**). Specifically, CA3 displayed $1.11\% \pm 0.42\%$ FLEN+ neurons 1.5 hours after CFC ($n=5$), which increased to $3.11\% \pm 0.74\%$ 3 hours post-CFC ($n=4$) and $2.31 \pm 0.44\%$ at 4.5 hours post-CFC ($n=7$). The fraction of active cells returned to basal-like levels at 6 hours ($1.86 \pm 0.16\%$ FLEN+/ mCherry+, $n=5$) and further decreased 12 hours post-CFC ($0.73\% \pm 0.32\%$ FLEN+/ mCherry+, $n=4$) (**Fig. 2D-E**). Overall, the emergence of FLEN assembly peaks 3 hours following CFC and significantly decreases 12 hours later (one-way ANOVA $p=0.003$, *post hoc* Tukey's multiple comparisons test: HC vs. 3 h $p=0.008$, 1.5 h vs. 3 h $p=0.04$, 3h vs. 12h $p=0.01$), faithfully mimicking the progression and transient nature of native c-Fos expression (Holtmaat & Caroni, 2016).

We sought to probe CA3 engram formation 3 hours after different behavioral situations, namely context-only (CO) and immediate-shock (IS) conditions, which offer the possibility for spatial exploration only and no spatial exploration at all, respectively (**Fig. S1A**). In CO conditions, mice were allowed to freely explore a novel cue-enriched arena. Comparably to other exploratory tasks, we found an increase in the number of FLEN+ CA3 PNs 3 hours following CO ($3.50 \pm 1.18\%$ FLEN+/FLEN-, $n=5$) compared to control HC mice (Kruskal-Wallis non-parametric test, $p=0.03$; Dunn *post hoc* multiple comparisons test, $p=0.02$) (**Fig. S2B**). In the IS test, a 6-second-long foot-shock was delivered 2 seconds after the mouse was placed in the conditioning cage, preventing sensorial sampling of the novel context (**Fig. S1A**). In contrast, the IS test did not induce a significant increase in the number of FLEN+ in CA3 PNs as compared to HC ($1.84 \pm 0.77\%$ FLEN+/FLEN-, $n=4$, Dunn *post hoc* multiple comparisons test, HC: IS, $p=0.51$) and to context-only (CO: IS, $p=0.88$, $n=4$), possibly because of minimal spatial exploration.

Because of the reported functional heterogeneity of CA3 PNs in relation to memory encoding and recall, which is directly related to their position within the proximodistal axis of CA3 (Nakamura et al., 2013; Y. Nakazawa et al., 2016; Q. Sun et al., 2017), we analyzed the spatial distribution of FLEN+ PNs within CA3 pyramidal layer along the proximodistal and superficial-to-deep axis of CA3, following one-trial CFC (**Fig. 2F** and **Fig. S2C**). We found a clear skewedness in the location of FLEN+ neurons along the proximodistal axis toward CA3a at all time points following

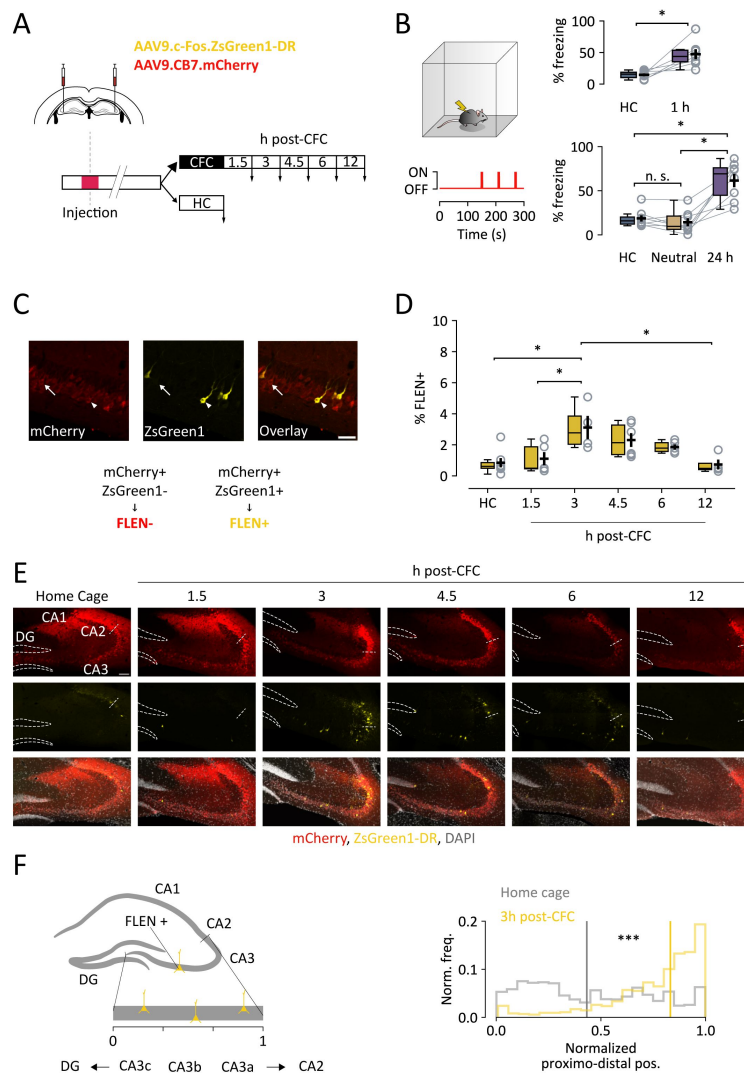


Figure 2.

In vivo FLEN labeling and decay time course following a salient experience.

(A) FLEN and an infection marker viral vectors are injected bilaterally into CA3. Mice undergo contextual fear conditioning (CFC), and sections are collected (downward arrows) 1.5, 3, 4.5, 6 and 12 hours following CFC. As a control, sections are obtained from mice left undisturbed in their home cage (HC). (B) Left: one-time CFC experiment scheme. Mice explore an arena for 5 minutes, 3 foot-shocks are given at 180, 210 and 270 seconds after the start of the test (red line). Top-right: paired comparison of the percentage of time mice spent freezing during training and when replaced in the conditioning arena 1 hour after CFC. Bottom-right: comparison of the percentage of time mice spent freezing during training, when placed in a different neutral environment and 24 hours after CFC. These experiments show that the CA3-dependent shock-context association is rapidly learned (within 1 hour), is highly environment-selective and it is retained for 24 hours. (C) Representative images of CA3 neurons expressing the infection marker (mCherry+, indicated by arrow) and neurons expressing both the infection marker and FLEN (ZsGreen1+/mCherry+, indicated by arrowhead). The latter are considered FLEN+ neurons. Scalebar = 50 μ m. (D) Percentage of ZsGreen+ CA3 neurons over the total number of mCherry+ neurons at different intervals following activation. (E) Representative sections of CFC-trained mice compared to HC untrained mice at different intervals following CFC. Dashed lines outline the dentate gyrus (DG) cellular layer, while the dashed segment indicates the separation between CA3 and CA2. Scale bar = 100 μ m. (F) Left panel: Schematic representation of neuronal position along the proximodistal axis analysis of CA3. The CA3 pyramidal layer was straightened and the x position of FLEN+ cells was normalized on a 0 (closer to DG) to 1 (closer to CA2) scale. Right panel: comparison of normalized proximodistal frequency distribution of CA3 FLEN+ cells placement in HC mice (gray) and 3 hours post-CFC mice (yellow). Bold lines represent the median of the respective distributions. HC median is displayed in the CFC plot to highlight distribution difference.

CFC and following CO compared to HC (Kolmogorov-Smirnov test: HC vs. 1.5 h post-CFC, $p < 0.001$; HC vs. 3 h post-CFC, $p < 0.001$; HC vs. 4.5 h post-CFC, $p < 0.001$; HC vs. 6 h post-CFC, $p < 0.001$; HC vs. 12 h post-CFC, $p < 0.001$; HC vs. CO, $p < 0.001$; HC vs. IS, $p < 0.001$) whilst they appeared to be more evenly distributed along the superficial-deep axis (Kolmogorov-Smirnov test: HC vs. 1.5 h post-CFC, $p = 0.013$; HC vs. 3 h post-CFC, $p < 0.001$; HC vs. 4.5 h post-CFC, $p < 0.001$; HC vs. 6 h post-CFC, $p < 0.001$; HC vs. 12 h post-CFC, $p = 0.61$; HC vs. CO, $p < 0.001$; HC vs. IS, $p = 0.02$) (**Fig. 2F** and **Fig. S2**). Overall, these data indicate that the FLEN labeling strategy faithfully identifies neurons activated during a one-trial exposure to a novel context and allows to characterize the morpho-functional properties of CA3 engram neurons shortly (< 6 hours) following the experience.

In parallel, we confirmed that the RAM system (Sørensen et al., 2016) is a powerful tool to tag neurons selectively activated by CFC in a temporally controlled manner (**Fig. S3A**) (Sørensen et al., 2016; Weng et al., 2018). To identify RAM+ neurons, we stereotactically injected AAV9-RAM/tTA::TRE/EGFP and AAV9-CB7/mCherry to label dorsal CA3 in wild-type mice (**Fig. S3A**). Mice were fed with doxycycline pellets starting one day before injection and then switched to a regular food diet for 48 hours before CFC (**Fig. S3A**). As reported, we found that the number of RAM+ neurons was notably higher 24 hours after CFC with respect to HC conditions (% RAM+ vs. mCherry+ neurons, HC: $1.13 \pm 0.18\%$, $n = 5$; CFC: $3.54 \pm 0.77\%$, $n = 5$; Welch's t-test $p = 0.03$) (**Fig. S3B-C**), although not to the same extent as in previous reports (Sørensen et al., 2016). The ratio of FLEN+ neurons 3 hours after CFC over HC (approximately 3:1) closely resembles that of RAM+ neurons 24 hours after CFC (approximately 3:1).

CA3 engram neurons evolve from low to higher firing frequency over 24 hours

To understand if rapid memory acquisition and early consolidation is supported by specific changes in electrophysiological intrinsic properties, we performed whole-cell patch-clamp recordings of CA3 PNs (mainly in CA3b) in acute hippocampal slices of young adult mice injected with either FLEN or RAM, respectively (**Fig. 3A**). AAV9-c-Fos/ZsGreen1-DR or AAV9-RAM-tTA::TRE-EGFP viral constructs were injected bilaterally in dorsal CA3, together with AAV9-CB7-mCherry (**Fig. 3A**). After CFC, mice were returned to their home cage for 3 hours (FLEN) or 24 hours (RAM), to yield peak expression of ZsGreen1-DR or EGFP and then sacrificed to collect parasagittal slices (**Fig. 3A**). Activity-dependent FLEN+ neurons remained detectable in acute slices, enabling specific targeting of engram CA3 PNs neurons for approximately 4 to 5 hours after slicing. Engram neurons (FLEN+ or RAM+) were compared to their respective neighboring non-engram neurons (FLEN- or RAM-) (**Fig. 3A**).

We first investigated the intrinsic physiological properties of FLEN+ and RAM+ CA3 PNs. Indeed, the probability for a neuron to be recruited in a memory engram is thought to depend on its inherent excitability level at the time of learning (see Josselyn & Tonegawa, 2020 for review). We assessed input resistance (R_i) by injecting a series of hyper- and depolarizing current pulses with the membrane potential held at -70 mV (**Fig. 3B**). R_i did not change significantly between FLEN+ and FLEN-neurons (FLEN+: 253 ± 35 M Ω , $n = 19$; FLEN-: 210 ± 19 M Ω , $n = 30$; non-parametric Mann-Whitney U test, p -value = 0.57) and was similar to RAM+ and RAM-neurons which were also not different between them (RAM+: 323.7 ± 14.1 M Ω , $n = 21$; RAM-: 396.1 ± 15.2 M Ω , $n = 30$, parametric t-test, $p = 0.68$) (**Fig. 3B**). Further depolarizing steps of current were injected until an action potential was fired. Action potential properties such as threshold, amplitude, width, upstroke and downstroke were not statistically different between FLEN+ and FLEN-neurons and between RAM+ and RAM-neurons (**Fig. S4B, C**; **Table 1**). To study the action potential firing pattern, we injected 1 s-long pulses of current just above rheobase (200 pA) (**Fig. 3C**). Overall, the firing frequency was not different in the FLEN groups (FLEN+: 2.64 ± 0.65 Hz, FLEN-: 3.59 ± 0.62 Hz, Mann-Whitney U test, $p = 0.51$) (**Fig. 3C**). More than half of the total number of action potentials occurred within the first 100 ms for all CA3 PNs, showing a consistent discharge pattern across both FLEN+ and FLEN-cells (Kolmogorov-Smirnov test, p -value = 1.0) (**Fig. 3D**). In contrast to

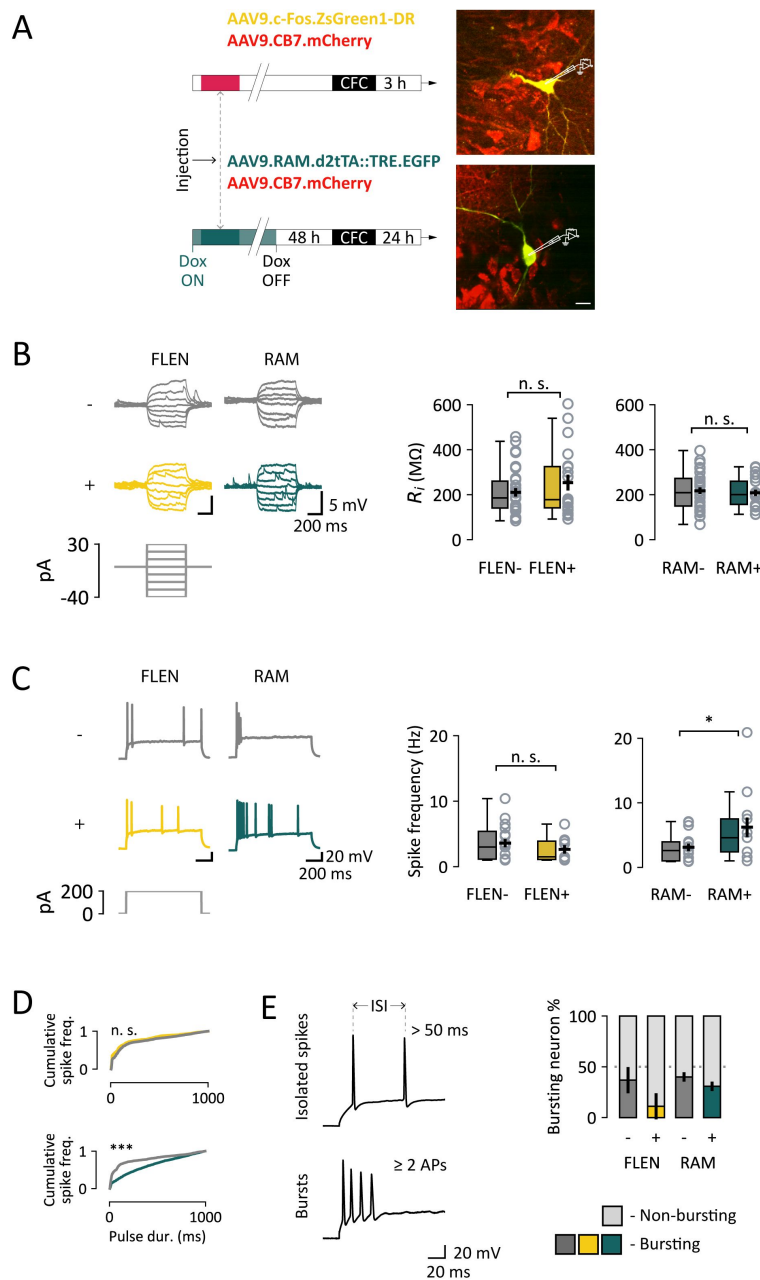


Figure 3.

Electrophysiological analysis of intrinsic properties of CA3 engram neurons.

(A) Experimental outline: mice were injected with either FLEN or RAM construct, subjected to CFC and then sacrificed at either 3- or 24-hours post-conditioning to collect acute hippocampal slices. Right: representative images of FLEN+ and RAM+ cells targeted for patch clamp recordings, with neighboring control FLEN- or RAM-neurons. Scale bar = 20 μ m. (B) Membrane potential response. Left panel: representative membrane potential trace in response to incremental injected current in control cells (gray) and engram cells (yellow for FLEN and green for RAM). Right panel: graphs of input resistance for comparison between engram cells and their corresponding control cells. (C) Action potential firing pattern. Left panel: sample traces of long (1 second) above-rheobase current step to analyze the pattern of action potential firing current in control cells (gray) and engram cells (yellow and green). Right panel: average spike frequency graphs. (D) Cumulative spike frequency curve over the 1 second current step shown in (C) showing a similar distribution between FLEN+ and FLEN - neurons (top), and a more sustained firing of RAM+ compared to their RAM-neurons (bottom). (E) Left panel: difference between isolated spikes and bursts. A group of action potential is considered a burst if two or more spikes occur within a 50 ms interval. Right panel: stacked bar plot showing the fraction of bursting cells in each group.

FLEN+ CA3 PNs, RAM+ CA3 PNs showed a prolonged spiking activity which outlasted the first 100 ms (**Fig. 3D**). This is reflected by an overall increased number of APs (RAM+: 20.9 ± 1.5 Hz, $n=13$; RAM-: 7.1 ± 0.6 Hz, $n=15$; Mann-Whitney U Test $p=0.04$) (**Fig. 3C**). CA3 PNs have been reported to have a distinctive burst-like firing pattern relying on the generation of dendritic Ca^{2+} spikes (Ding et al., 2020; Raus Balind et al., 2019). We defined bursts as a firing pattern where two or more spikes occur within a 50 ms interval (Zucca et al., 2017). Bursts were triggered by rheobase current pulses in 7 out of 19 FLEN-CA3 PNs (36.8 ± 12.9 %, with an average of 2.33 ± 0.33 spikes per burst), but only 1 out of 9 FLEN+ neurons displayed bursting behavior (11.1 ± 12.9 %, with 3.86 average spikes per burst). Bursts were induced in 6 out of 15 RAM-neurons (40.0 ± 4.6 %, with an average of 2.13 ± 0.07 spikes per burst). However, contrarily to FLEN+ neurons, 30.8 $\pm$ 4.6 % (4 out of 13 CA3-PN RAM+ neurons with an average spike/burst of 2.10 ± 0.10), indicating a 2.7-fold increase in the propensity of CA3 engram neurons to display bursting behavior over 24 hours after CFC (**Fig 3D**). Altogether, these results show that early after encoding, FLEN+ CA3 PNs do not show differences in excitability or patterns of bursting activity as compared to control neurons. However, after 24 hours, CA3 engram cells display an increase in excitability and in bursting behavior without changes in R_i .

Synaptic properties of FLEN+ neurons

To assess overall excitatory synaptic strength and overall number of active synaptic connections, we recorded spontaneous excitatory post-synaptic currents (sEPSCs) in FLEN+ CA3 PNs while blocking GABA_A-dependent inhibitory currents (Bicuculline, 10 μM). The average amplitude of sEPSCs was not different between FLEN+ and FLEN-CA3 PNs (FLEN+: 29.7 ± 5.3 pA, FLEN-: 32.2 ± 3.7 pA, Kolmogorov-Smirnov test, $p = 0.99$), whereas their frequency was increased (Inter-event interval (IEI), FLEN+: 788 ± 152 ms, FLEN-: 930 ± 91 ms, Kolmogorov-Smirnov test, $p < 0.005$) (**Fig. 4A**). Similarly, 24 hours after a novel experience, the average amplitude of sEPSCs was similar between RAM+ and RAM-CA3 PNs (RAM+: 26.22 ± 2.46 pA, $n=14$; RAM-: 21.31 ± 2.72 pA, $n=14$, Kolmogorov-Smirnov test, $p=0.99$) (**Fig. 4A**), but their frequency was significantly increased in RAM+ CA3 PNs (RAM+: 645 ± 88 ms, $n=17$; RAM -: 1342 ± 207 ms, $n=17$; Kolmogorov-Smirnov test, $p < 0.05$) (**Fig. 4A**).

Then, to investigate excitatory synaptic strength and release probability independent of network activity, we measured miniature EPSCs (mEPSCs) by further adding TTX (0.5 μM). The average amplitude was similar between FLEN+ and FLEN-neurons (FLEN+: 20.5 ± 2.5 pA, FLEN-: 18.7 ± 1.6 pA, Kolmogorov-Smirnov test, $p = 0.99$), whereas we observed a higher frequency of mEPSCs in FLEN+ neurons as compared to control neurons (IEI, FLEN+: 1722 ± 242 ms; FLEN-: 1228 ± 186 ms, Kolmogorov-Smirnov test, $p < 0.0005$) (**Fig. 4B**). Comparably, we found a higher frequency of mEPSCs in RAM+ neurons as compared to RAM-CA3 PNs (RAM+: 782 ± 91 ms, $n=14$; RAM-: 1244 ± 247 ms, $n=13$; Kolmogorov-Smirnov test, $p<0.005$), with unchanged amplitude (RAM+: 29.6 ± 4.8 pA, $n=14$; RAM-: 24.7 ± 4.2 pA, $n=13$; Kolmogorov-Smirnov test, $p=0.99$) (**Fig. 4B**). Thus, neurons recruited in an engram appear to show increased excitatory drive after a few hours that persist after 24 hours, which may be linked to a higher number of synaptic contacts.

Previous reports indicate that CA3 PNs activated during by contextual learning and identified 24 hours later by RAM had strengthened mossy fiber inputs (Weng et al., 2018). The existence of mEPSCs with larger amplitudes may be related to synaptic plasticity in a subset of synapses, or theoretically to an increased participation of Mf-driven mEPSCs (Henze et al., 2002). We thus calculated the proportion miniature excitatory events with an amplitude greater than 40 pA. During spontaneous activity recording, we found no difference in the fraction of putative Mf-driven inputs between FLEN+ and FLEN-PNs (FLEN+: 0.09 ± 0.03 %, $n=9$; FLEN-: 0.06 ± 0.02 %, $n=14$; Mann-Whitney U Test $p=0.33$) and between RAM+ and RAM-PNs 24 hours after learning (RAM+: 0.25 ± 0.08 %, $n=14$; RAM-: 0.18 ± 0.07 %, $n=13$; Mann Whitney U test, $p=0.42$) (**Fig. 4B**, inset). This later finding contrasts with earlier finding (Weng et al., 2018), which found a

	FLEN			RAM		
	FLEN-	FLEN+	Test	RAM-	RAM+	Test
Peak (mV)	45.19 ± 2.25, n=19	43.92 ± 4.62, n=9	Mann-Whitney U, p=1.00	47.93 ± 1.41, n=15	50.91 ± 1.57, n=13	T-test, p=0.17
Threshold (mV)	-40.41 ± 1.54, n=20	-42.47 ± 1.36, n=14	T-test, p=0.35	-41.53 ± 1.35, n=15	-39.78 ± 2.46, n=13	T-test, p=0.52
Amplitude (mV)	99.91 ± 4.63, n=19	98.67 ± 8.73, n=9	T-test, p=0.89	103.71 ± 3.93, n=15	101.18 ± 4.53, n=13	T-test, p=0.67
Width (ms)	0.002 ± 0.001, n=16	0.002 ± 0.001, n=6	Mann-Whitney U, p=0.51	0.0017 ± 0.0001, n=13	0.0016 ± 0.00003, n=7	Mann-Whitney U, p=0.36
Upstroke (mV/s)	197.06 ± 19.54, n=19	213.35 ± 35.48, n=9	T-test, p=0.67	229.89 ± 17.86, n=15	241.65 ± 23.2, n=13	T-test, p=0.69
Downstroke (mV/s)	-57.99 ± 7.91, n=19	-54.99 ± 6.18, n=9	Mann-Whitney U, p=0.84	-56.70 ± 4.43, n=15	-59.96 ± 3.62, n=13	T-test, p=0.58
Initial frequency (Hz)	21.58 ± 4.15, n=16	25.04 ± 12.83, n=7	Mann-Whitney U, p=0.62	41.74 ± 9.54, n=10	25.30 ± 6.43, n=11	Mann-Whitney U, p=0.25
ISI (ms)	0.19 ± 0.05, n=16	0.26 ± 0.08, n=7	Mann-Whitney U, p=0.28	0.14 ± 0.02, n=10	0.14 ± 0.02, n=11	T-test, p=0.98

Table 1.

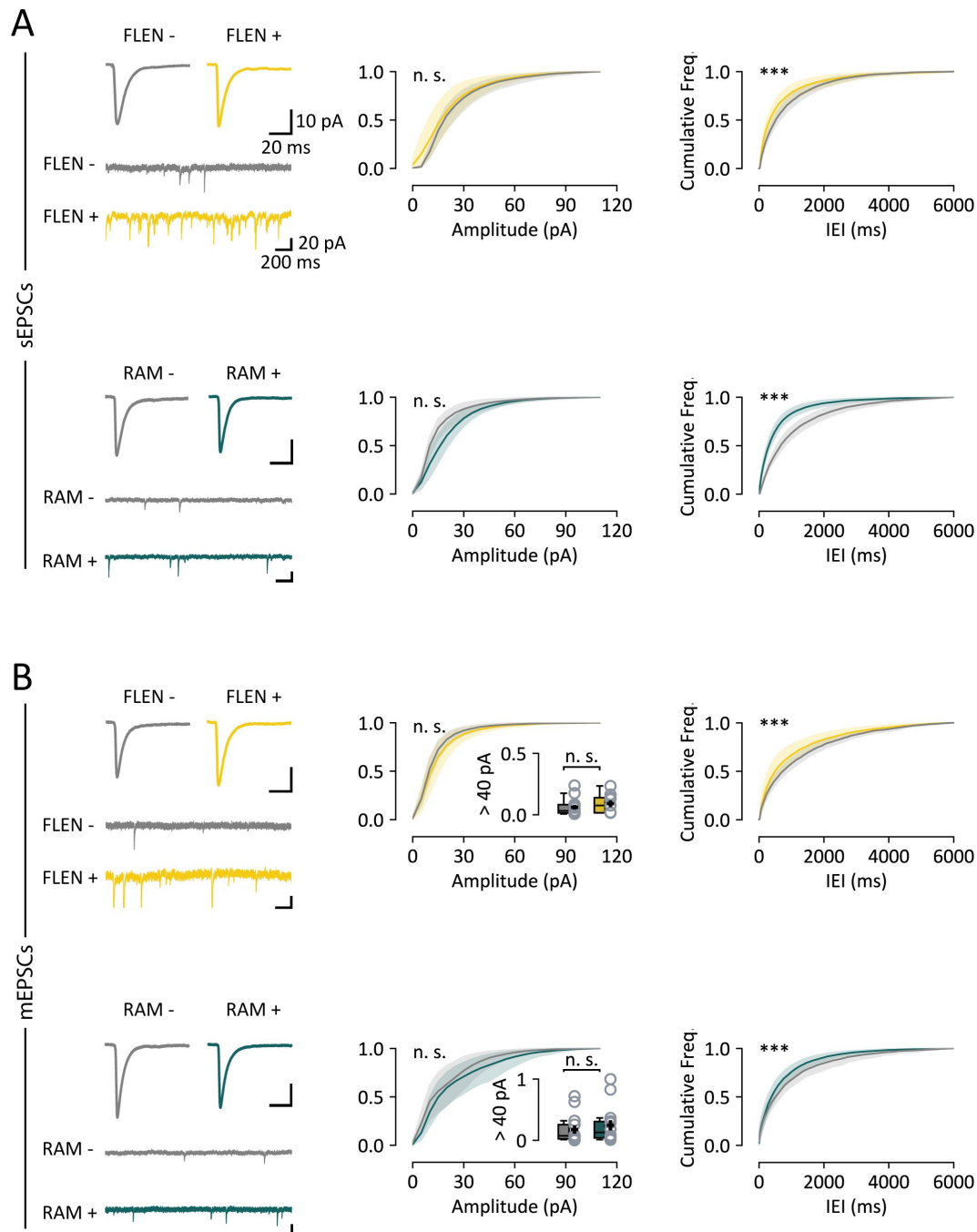


Figure 4.

Analysis of spontaneous EPSCs in CA3 engram neurons

(A) Sample traces and cumulative distributions of sEPSCs recorded from CA3 engram neurons (FLEN in yellow, RAM in green). Top-left: comparison of spontaneous EPSCs (sEPSCs) between FLEN+ and FLEN-neurons, showing no difference in average amplitude, and representative traces showing differences in the frequency of sEPSCs between FLEN- and FLEN+ neurons. Top-center and top-right: cumulative frequency distribution of amplitude and inter-event interval (IEI) as a readout of sEPSC frequency in FLEN+ and FLEN-neurons. Bottom-left: comparison of spontaneous EPSCs between RAM+ and RAM-neurons showing no difference in average amplitude and sample traces showing differences in the frequency of sEPSCs between RAM+ and RAM-neurons. Bottom-center and bottom-right: cumulative frequency distribution of amplitude and inter-event interval (IEI) of RAM+ and RAM-neurons. (B) The same as in (A) but for miniature EPSCs. Inset within cumulative distribution of amplitude plot represent the normalized fraction of mEPSC larger than 40 pA, likely due to giant mossy fiber activity.

selective strengthening of Mf inputs onto the ensemble of CA3 neurons >24 hours after activation by CFC. At variance with our analysis, this later study uses a mGluR2/3 agonist, LY354740, which inhibits evoked Mf-CA3 synaptic transmission (Kamiya & Ozawa, 1999 [\[1\]](#)).

Inhibitory neurons are important in controlling spike generation of CA3 PNs, shaping network activity and participation in neuronal assembly formation (Holtmaat & Caroni, 2016 [\[2\]](#)). We therefore probed miniature inhibitory post-synaptic currents (mIPSCs), comparing FLEN+ vs. FLEN-neurons and RAM+ vs. RAM-neurons, in the presence of NBQX (20 μ M), D-AP5 (50 μ M) and TTX (0.5 μ M). We found that mIPSCs were statistically more frequent (FLEN+: 340.1 $\pm$ 4.0 ms, FLEN-: 511.0 $\pm$ 40.5 ms, Kolmogorov-Smirnov test, $p < 0.005$), but their amplitude was on average comparable in FLEN+ vs. FLEN-neurons (FLEN+: 35.1 $\pm$ 2.3 pA, FLEN-: 31.7 $\pm$ 2.2 pA, Kolmogorov-Smirnov test, $p = 0.99$, Welch's t-test, $p = 0.30$) (**Fig 5A** [\[3\]](#)). At 24 hours after learning, RAM+ neurons showed an increase in the frequency of mIPSC (IEI 330.5 $\pm$ 51.9 ms, $n=6$) compared to RAM-neurons (IEI 437.1 $\pm$ 47.1 ms, $n=11$; Kolmogorov-Smirnov test, $p < 0.005$), but no difference in their amplitude (RAM+: 22.7 $\pm$ 1.8 pA, $n=11$; RAM-: 26.5 $\pm$ 5.2, $n=6$, Kolmogorov-Smirnov test, $p=0.89$) (**Fig. 5A** [\[3\]](#)). The frequency of all excitatory and inhibitory inputs was elevated after CFC, both after 3 hours and after 24 hours, but not their amplitude.

Mf-CA3 synapses and feedforward inhibition in FLEN+ neurons

Mf-CA3 connections between the DG and CA3 pyramidal cells provide powerful inputs via 'giant' Mf boutons to CA3 PCs, which are thought to assist CA3 in the encoding of memory (Marneffe et al., 2024 [\[4\]](#); Vandael & Jonas, 2024 [\[5\]](#)). We recorded EPSCs evoked by minimal stimulation of Mfs (Marchal & Mulle, 2004 [\[6\]](#)) in FLEN+ and FLEN-CA3 PNs using an extracellular electrode placed within the DG hilus to stimulate the Mf axonal bundle (**fig. 6A** [\[3\]](#)). Low-frequency stimulation (0.1 Hz) evoked EPSCs of variable amplitude that, on average, did not differ between FLEN+ (81.7 $\pm$ 17.5 pA) and FLEN-neurons (80.8 $\pm$ 13.5 pA, Welch's t-test $p = 0.97$) (**Fig. 6B** [\[3\]](#)). Increasing stimulation frequency to 1 Hz induced short-term presynaptic frequency facilitation, causing an approximatively 3-fold increase in evoked current amplitude in both FLEN+ (258.3 $\pm$ 49.5 pA) and FLEN-neurons (238.2 $\pm$ 39.0 pA, Welch's t-test $p = 0.76$) (**Fig. 6B** [\[3\]](#)). The EPSC facilitation ratio (1Hz vs. 0.1 Hz) was not different between FLEN+ and FLEN-neurons (FLEN+: 3.28 $\pm$ 0.40, FLEN-: 3.28 $\pm$ 0.57, Welch's t-test $p = 0.99$) (**Fig. 6B** [\[3\]](#)).

One-trial memory tasks reversibly induce a robust and rapid (within hours) increase in the number of filopodia emanating from Mf synaptic terminals which contact GABAergic interneurons (Ruediger et al., 2011 [\[7\]](#)), lending support to the hypothesis of an increase in feedforward inhibition in most CA3 PNs. Because of the rapid time course with which structural plasticity occurs at Mf-CA3 synapses following one-trial memory task, we took advantage of FLEN in order to explore a functional counterpart to structural plasticity, and to address whether the change in feedforward inhibition was differentially affected in FLEN+ vs. FLEN-CA3 PNs. We recorded excitatory and inhibitory synaptic currents in CA3 PNs while stimulating Mfs (**Fig. 6A** [\[3\]](#)). To calculate the excitatory to inhibitory (E/I) ratio at these synapses, we first recorded evoked EPSCs by holding cells at the experimentally determined GABA_A reversal potential ($E_{Cl^-} = -70$ mV), and then at the reversal potential for cations ($E_{Na^+} = +5$ mV) (**Fig. 6C** [\[3\]](#)) (Torborg et al., 2010 [\[8\]](#)). We measured the peak amplitude of evoked EPSCs and IPSCs at a stimulation rate of 0.1 Hz to prevent frequency facilitation (**Fig. 6C** [\[3\]](#)). We found that the EPSC/IPSC ratio was higher in FLEN+ vs FLEN-cells, although this not significantly different (FLEN+: 3.48 $\pm$ 0.96, FLEN-: 2.20 $\pm$ 0.42, Welch's t-test, $p = 0.27$) (**Fig. 6C** [\[3\]](#)). These data do not support the notion that shortly after one-trial learning, there is an increase in feedforward inhibition at Mf-CA3 synapses.

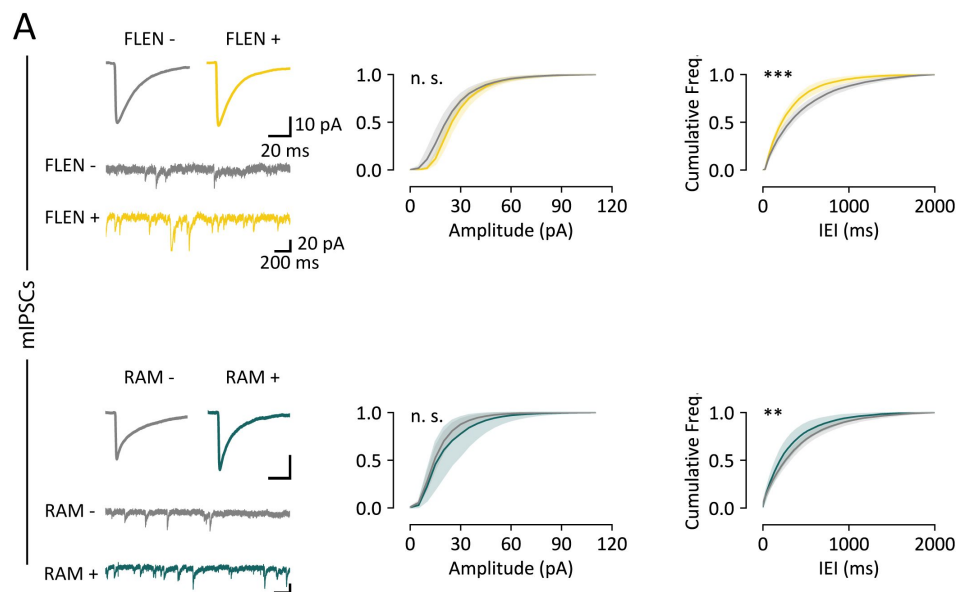


Figure 5.

Analysis of spontaneous IPSCs in CA3 engram neurons

(A) Sample traces and cumulative distributions of mIPSCs recorded from hippocampal CA3 engram (FLEN in yellow, RAM in green). Top-left: comparison of the average miniature IPSC between FLEN+ and FLEN-neurons showing no difference in amplitude. Below, sample traces of FLEN+ and FLEN-neurons showing differences in the frequency of mIPSCs. Top-center and top-right: cumulative distribution plots for amplitude and inter-event interval (IEI) in FLEN+ versus FLEN-neurons. Bottom-left: sample traces highlighting the difference in mIPSC frequency between RAM- and RAM+ neurons. Below, sample traces of FLEN+ and FLEN-neurons showing difference in the frequency of mIPSCs. Bottom-center and bottom-right: cumulative frequency distribution for amplitude and inter-event interval (IEI) of RAM+ and RAM-neurons.

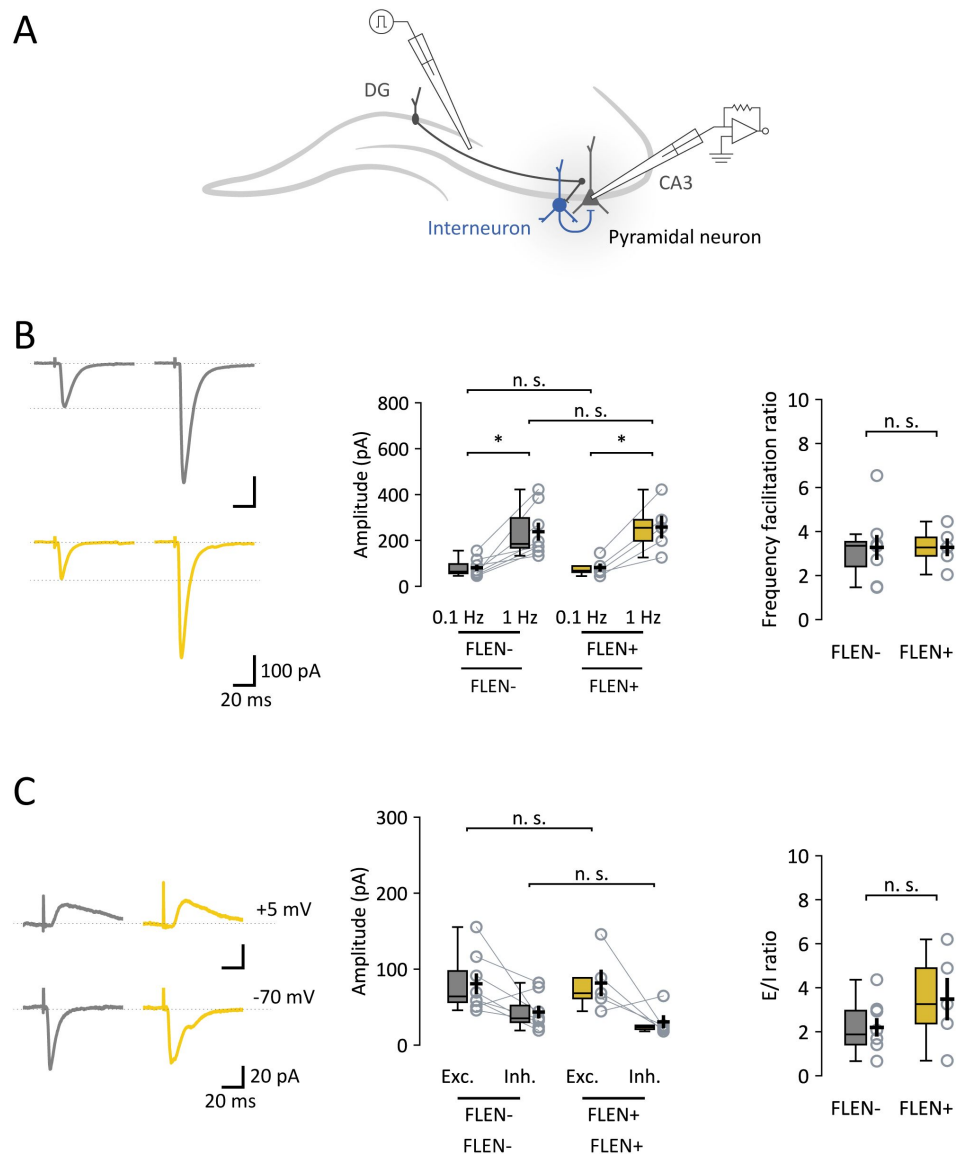


Figure 6.

Properties of Mf-CA3 synapses in CA3 engram neurons

(A) Schematic representation of Mf-CA3 EPSCs and feedforward inhibition recordings. **(B)** Left panel: representative traces showing EPSC amplitude in FLEN- (gray) and FLEN+ (yellow) cells following low frequency stimulation (0.1 Hz) and corresponding increase evoked with a higher stimulation frequency (1 Hz), due to a presynaptic form of synaptic plasticity characteristics of Mf-CA3 synapses (frequency facilitation). Center panel: amplitude plot for FLEN- and FLEN+ stimulated at 0.1 Hz and 1 Hz. Right panel: frequency facilitation ratio plot shows no difference between FLEN- and FLEN+ neurons. **(C)** Left panel: representative traces showing Mf-driven IPSCs (feedforward inhibition) and EPSCs (direct excitatory activity) recorded in individual FLEN- (gray) and FLEN+ (yellow) neurons. Center panel: amplitude plot of EPSCs and IPSCs for FLEN- and FLEN+ neurons. Right panel: graph showing excitation-to-inhibition ratio, comparing FLEN- and FLEN+ neurons.

Discussion

In the past decade, the search for engrams and the characterization of engram cells has largely benefited from new gene targeting approaches to label the cells and to control their reactivation (Josselyn & Tonegawa, 2020 [↗](#)). The primary goal of our work was to test the hypothesis that the functional properties of hippocampal cells activated during a one-trial contextual memory task were distinct from surrounding ones shortly (3-6 hours) after encoding and were subject to progressive plastic changes which could be linked to a first phase of memory consolidation.

Most techniques currently employed to study the properties of engram cells *ex vivo*, combine promoter expression based on IEGs, intersectional transgenic strategies, optogenetics and pharmacogenetics with behavioral paradigms (Denny et al., 2014 [↗](#); Pignatelli et al., 2019 [↗](#); X. Sun et al., 2020 [↗](#); Tonegawa et al., 2015 [↗](#)). A major caveat for using activity-dependent promoter/enhancers is the background basal expression which can result in the labeling of neurons unrelated to the triggering behavioral event. To limit this, fluorescent marker and/or actuator expression can be temporally controlled by directly coupling IEG promoters, such as c-Fos or synthetic promoters (RAM), to inducible systems such as the doxycycline (Dox) or the tamoxifen (TAM) systems (12/4/2024 5:31:00 PM). With these approaches, the physiological properties of engram vs. non-engram cells has been examined by *ex vivo* patch-clamp recordings 24 hours to several days after the initial contextual memory task (Pignatelli et al., 2019 [↗](#); Sørensen et al., 2016 [↗](#); Weng et al., 2018 [↗](#); Yap et al., 2021 [↗](#)).

We thus developed a viral strategy for transient neuronal expression of a fluorescent protein (FLEN) following the exposure of mice to a novel context making use of the full-length c-Fos promoter and of the brightly fluorescent ZsGreen1 protein engineered to be quickly destabilized. The time course of expression of ZsGreen1 in FLEN labeled neurons peaked at 3 hours both after pharmacological activation in cultured neurons and *in vivo* after a 5-minute exposure to a novel context. Neuron labeling persisted for a few hours, enabling *ex vivo* electrophysiological characterization of neurons thought to be involved in the initial encoding of contextual memory. As with all techniques based on IEG promoters, a small percentage of AAV-transduced neurons showed ZsGreen1 fluorescence under control conditions (HC or no activation of cultured neurons). However, following CFC, the number of FLEN labeled neurons increased by a factor of 3.12. The bright fluorescence of ZsGreen1 enabled the electrophysiological characterization of FLEN+ neurons, the majority of which (>60%) were activated in relation to the CFC memory task.

We have used FLEN to label neurons in CA3, which is involved in the rapid encoding of new spatial and contextual information (Kesner and Rolls, 2015 [↗](#)). One of the key features of CA3 is the presence of recurrent excitatory connections which are subject to NMDA-dependent synaptic plasticity (Rebola et al., 2017 [↗](#)), and this appears essential for memory of a one-trial experience (K. Nakazawa et al., 2003 [↗](#)). It has been proposed that the excitability state of neurons may contribute to memory formation (Zhang & Linden, 2003 [↗](#)). We first examined whether FLEN+ neurons showed distinctive intrinsic excitability properties. We observed no difference in the membrane potential, input resistance, threshold or bursting activity between FLEN+ and FLEN- neurons in slices prepared 3 hours following CFC. This suggests that the subpopulation of CA3 PNs engaged in a one-trial learning task are not initially more excitable than the general population of neurons, hence are not predetermined by their excitability state. Alternatively, engram neurons may be transiently more excitable, at the time of encoding, and lose this property within a few hours. In apparent contrast with our finding that FLEN+ CA3 PNs did not show increased excitability, a simple associative memory task, trace eye-blink conditioning, appears to increase the excitability of a large fraction (more than half) of CA3 neurons starting at 1 hour and decreasing after several days (Thompson et al., 1996 [↗](#)). Beyond the difference in the task and in the species, these contrasting results may be explained by an overall increased activity of CA1

neurons related to the learning task (Cai et al., 2016; Moyer et al., 1996), however not specific to engram neurons. This does not preclude however the possibility that relative neuronal excitability immediately before training could contribute to the selection of neurons to an engram as proposed in the amygdala (Yiu et al., 2014) and the hippocampus (Mocle et al., 2024). However, in our hands, a one-trial memory task does not per se trigger a selective increase in the excitability of engram neurons 3 hours after CFC as compared to the general population of CA3 neurons. In the DG, recall of a previously stored contextual memory (48 hours before), hence reactivation of engram neurons, rapidly increased the excitability of these engram neurons (within 1 hour) with respect to engram neurons which were not subject to reactivation in a separate group of mice (Pignatelli et al., 2019). A learning-related task may however induce plasticity of intrinsic excitability of engram neurons on a longer time scale. We thus compared the intrinsic excitability of FLEN+ neurons to those labeled more than 24 hours after CFC by using the previously described RAM (Sørensen et al., 2016). We found that RAM+ neurons showed prolonged spike firing in response to a depolarizing pulse of current, hence decreased accommodation in comparison with non-labeled neurons. Considering that the set of neurons labeled by these two strategies may be of the same set, based on the expression of a fluorescent marker under the control of the c-Fos promoter, our data strongly suggest that engram neurons in CA3 progressively increase excitability as compared to neurons which were not activated by the one-trial contextual memory task. The acquisition of increased excitability 24 hours after one-trial memory encoding suggests that intrinsic excitability is a key feature of an early consolidation of memory, which may even be strengthened upon reactivation of memory (Pignatelli et al., 2019). It would be interesting to assess whether increased intrinsic excitability persists throughout further consolidation of memory.

Increased neuronal activity in engram neurons at the time of encoding may result from more efficient or more synchronous excitatory synaptic inputs to these neurons. Here we observed that FLEN+ neurons (hence characterized >3 hours after initial encoding) showed an increased excitatory drive, which may be explained by a higher number of synaptic contacts. We cannot discriminate whether the increased number of synaptic inputs preexists to the formation of the engram, or whether it is a plastic mechanism taking place shortly following the contextual memory. The time course of appearance of additional synapses, by unmuting of silent synapses following contextual memory acquisition, is compatible with experimentally induced functional and structural synaptic plasticity (Holtmaat & Caroni, 2016). On the other hand, if FLEN+ neurons showed higher synaptic inputs at the time of memory encoding, this would increase the chance that these neurons show increased excitation (c-Fos activation) hence participate in the engram. The amplitude of mEPSCs in FLEN+ neurons does not appear to increase on average in comparison with FLEN-CA3 PNs. This finding may not favor the possibility that LTP-like synaptic plasticity occurs early during the process of memory formation. In this event, NMDA-dependent forms of synaptic plasticity are more likely to occur later on during the early phase of consolidation of the memory trace (Humeau & Choquet, 2019). Interestingly, we observed a comparable increase in the frequency of sEPSCs and mEPSCs in RAM+ CA3 neurons 24 hours after CFC, as previously reported (Weng et al., 2018), without any change in the amplitude. Using a pharmacological agent which selectively inhibits Mf-CA3 transmission, it was hypothesized that the increased mEPSC frequency is mainly due to more active Mf-CA3 synapses (Weng et al., 2018). However, if this was the case, we would also expect an increase in the amplitude of mEPSCs as well as an increase in the fraction of mEPSCs larger than 45 pA. In addition, we did not find evidence for an increase in Mf-CA3-evoked EPSCs (see below). Overall, our data indicate that CA3 engram neurons characterized a few hours after the one-trial memory task show a higher level of excitatory synaptic inputs, and this feature seems to extend to 24 hours. We found no evidence for increased amplitude of EPSCs in FLEN+ or RAM+ CA3 PNs which could have been indicative of postsynaptic plasticity through an increased number of synaptic AMPA receptors (Humeau & Choquet, 2019; Takeuchi et al., 2014). One reason may be that synaptic plasticity only occurs in a subset of synaptic contacts, and the recording of mEPSCs is not resolute enough to capture these modifications.

Local inhibition controls the spiking of CA3 PNs, hence their capacity to be engaged in an active ensemble of neurons at the time of contextual memory encoding. In CA1, there is a fine control of inhibition upon novel environment exploration (Yap et al., 2021 [DOI](#)) or fear learning (Lovett-Barron et al., 2014 [DOI](#)). We found an increase in the frequency of mIPSCs after CFC in engram CA3 neurons, both after 3 hours and after 24 hours. This suggests that the elevation of mEPSCs in the engram neurons is counterbalanced by increased inhibition, although this would need to be generally tested in conditions of hippocampal circuit activity *in vivo*. We have also provided information on the process of Mf-driven feedforward inhibition in CA3 following one-trial memory encoding. Mf-CA3 synapses have been proposed to play a major role in assisting CA3 in the encoding of memory (Kesner & Rolls, 2015 [DOI](#)). For these reasons, it is important to directly characterize the properties of Mf-CA3 synapses impinging on CA3 engram neurons. We observed no difference in the amplitude and short-term plasticity of Mf-CA3 EPSCs evoked in FLEN+ CA3 neurons vs. FLEN-neurons. It has been proposed that 24 hours after CFC, the increased amplitude of mEPSCs in RAM+ neurons can be attributed to modifications of Mf-CA3 synapses (Weng et al., 2018 [DOI](#)), which may suggest progressive plasticity of these inputs. There is abundant evidence that the structural properties of Mf-CA3 synapses are modified by experience (Ruediger et al., 2011 [DOI](#))(Galimberti et al., 2006 [DOI](#); Maruo et al., 2016 [DOI](#)). Feedforward inhibition combines with short-term plasticity of Mf-CA3 synapses to define the physiological conditions for efficient spike transfer between the DG and CA3 (Marneffe et al., 2024 [DOI](#); Vandael & Jonas, 2024 [DOI](#); Zucca et al., 2017 [DOI](#)). CFC induces a robust and global increase in the number of filopodia emanating from the giant Mf terminals, which peaks between 1 and 5 hours after the one-trial memory task and is related to the precision of memory and the size of engrams (Ruediger et al., 2011 [DOI](#)). Because filopodia contact GABAergic interneurons, this structural plasticity has been proposed to increase feedforward inhibition, albeit this has not yet been demonstrated. Here we compared the excitation/inhibition ratio of Mf-evoked synaptic responses between FLEN+ and FLEN-neurons. We did not observe a significant increase of feedforward inhibition onto CA3 engrams, as may be expected from an increase number of filopodia, but we rather observed a tendency towards less feedforward inhibition. This leads us to speculate that shortly following contextual memory formation, Mf-driven feedforward inhibition is increased in the general population of neurons but not in Mf inputs onto engram neurons, thus allowing for a comparatively stronger input from the DG to the engram neurons. It will be interesting to test whether this increased signal to noise ratio is maintained 24 hours after contextual memory encoding, given that the number of filopodia per Mf bouton has returned to basal level (Ruediger et al., 2011 [DOI](#)).

In conclusion, we used a viral tool allowing for the identification of neurons involved in a behavioral task within 3 hours, as a complement to the transgenic mouse strategies and viral vectors which have been used to assess changes in engram neuron properties in the time range of days. Although it is not possible to fully assess the electrophysiological properties of engram neurons at the time of the encoding, this work shows that the engram neurons are not different in their intrinsic membrane properties shortly after encoding. This set of properties may however follow delayed (progressive) plastic changes which are evident 24 hours after initial encoding. Overall, we propose that engram neurons do not show higher excitability preexisting to the selection of neurons to the engram during the memory encoding task. Hence the increased activation state of the FLEN+ neurons at the time of encoding may rather relate to a higher number of active synaptic inputs providing more efficient excitation. Interestingly, although more work is needed to confirm this tendency, increased excitation may also be linked to decreased Mf-driven feedforward inhibition, thus favoring spike transfer from the DG to CA3 at the time or shortly after memory encoding. The FLEN viral tool will be useful to assess how the properties of neurons in brain regions involved in the contextual memory encoding process, including the entorhinal cortex and the DG.

Methods

Animals

C57Bl6j female mice were obtained from Janvier and cared according to the regulations of the University of Bordeaux/CNRS Animal Care and Use Committee. Animals were housed with their littermates with ad libitum access to food and water. Cages were kept in a temperature-regulated room on a 12 h light/dark cycle. Prior to any behavioral test, all mice were single housed for 4-5 days and then handled for at least 3 consecutive days. Mice injected with the RAM system were kept on a doxycycline diet (4 mg/Kg) starting 1 day before the viral injection.

Viral vectors

Adeno-associated virus (AAV) serotype 2/9 vectors containing our custom-made c-Fos-ZsGreen1-DR (titer 4.00E13) were generated in our lab, whereas AAV2/9 vectors containing the RAM system (titer 3.39E13) were purchased from AddGene. Each vector was mixed with an infection marker virus AAV2/9-CB7/mCherry (titer 1.50E13) and diluted 1:10 for co-injection.

Cell Culture

P0 mouse cortical neurons were plated (300000 cells/well) and allowed to grow on glia cells. At DIV3 cells were infected with the viral mix of c-Fos-ZsGreen1-DR and CB7-mCherry. At DIV14, neurons were switched to an activating medium containing 4-AP (100 μ M) and Bicuculline (10 μ M) for 30 minutes. Then this activating medium was washed and tetrodotoxin was added (0.5 μ M) to prevent action potential firing and stop neuronal activation. Neurons were then fixed using 0.4% paraformaldehyde (PFA) in 0.1 mM PBS at the different intervals following pharmacological activation.

Stereotaxic injection

2-month-old C57Bl6j mice were bilaterally injected with either AAV9-c-Fos-ZsGreen1-DR or the AAV9-RAM virus mixed with the infection control virus AAV9-CB7-mCherry into dorsal hippocampus. Mice were anesthetized with 4% isoflurane and placed in a stereotaxic apparatus, where they were kept under 1.5-2% isoflurane anesthesia (0.8-1 L/min), and injected with buprenorphine (0.1 mg/kg) and carprofene (5 mg/kg) to prevent post-surgery pain. The injection volume and flow rate (300 nl at 60 nl/min) were controlled with an injection micro-pump (World Precision Instruments). Injection coordinates, using bregma as a reference point, were as follows: AP -1.78, ML $\pm$ 2.40, DV -2.35. This site targets dorsal CA3b, but virus spreading resulted the infection of CA3a, CA3b, CA3c, CA2 and in some cases few granule neurons and hilar cells of the DG. Mice were carefully monitored for 4 days following surgery, and behavioral manipulation started after a recovery period of approximately 2 weeks.

Behavioral tests

Once recovered, mice were single-housed to reduce uncontrolled experiences that could contaminate the ensemble of CFC-related labelled engram cells. Then, mice were handled to habituate to the experimenter for at least 3 consecutive days prior to CFC. Mice injected with the RAM virus were deprived of doxycycline, switching to a regular diet, 48 hours before CFC training. On the one-trial learning day, mice were transported in the experimental room and immediately placed in a novel conditioning chamber enclosed in a sound-attenuating cabinet (Ugo Basile), enriched with visual cues, and allowed to explore it for a total duration of 300s. After an initial exploration period, 3 mild foot-shocks (2s, 0.7 mA) were delivered at a 150s, 210s, and 270s, leaving 30s for additional context sampling at the end. Mice were recorded and freezing behavior was scored using ANY-Maze 6 software (Stoelting Europe). Freezing was detected as the absence of

movement apart from respiration, and freezing events were scored when immobility lasted more than a threshold of 300ms. Mice were then returned to their home cage. For context only (CO) exploration, mice were placed in a 50 cm x 50 cm custom-built open field, where some objects were distributed. Mice were allowed to explore for 5 minutes before returned to their home cage. The immediate shock test was carried out in the fear conditioning system. Mice explored the conditioning chamber for only 2 seconds before receiving a 6-second-long foot-shock and were then returned to their home cage.

Histology

Mice were anesthetized with intraperitoneal administration of a ketamine/xylazine mix (100 mg/kg/10 mg/kg) at different time intervals following CFC (1.5 h, 3 h, 4.5 h, 6 h, 12 h for c-Fos-ZsGreen1-DR, 24 h for RAM). Mice were then intracardially perfused with 0.9% NaCl solution followed by 4% paraformaldehyde (PFA) in 0.1 mM PBS. Brains were removed and postfixed in 4% PFA overnight at 4°C. Brains were transferred in sucrose 30% in 0.1 mM PBS at 4°C until they sank. Brains were cut on a cryostat into 25 µm-thick coronal sections and collected in PBS with 0.3% Triton X-100. Slices were immediately mounted on glass slides and covered in DAPI-based mounting medium. The number of fluorescence neurons was assessed directly without amplification.

Confocal microscope

Confocal acquisitions were performed using a Leica SP8 White Light Laser 2 on an inverted stand (Leica Microsystems, Mannheim, Germany) and with a x20 (NA 0.75) immersion objective. Images were analyzed using ImageJ. At least 5 slices per animal were acquired. Imaging of neuronal cultures was carried out as well using confocal Leica SP8 White Light Laser 2 (Leica Microsystems, Mannheim, Germany) and with a x20 (NA 0.75) immersion objective. In all experiments, the settings (laser power, gain, offset) were fixed between groups as well as the pixel size (120 nm).

Ex-vivo slice preparation

Slices were prepared 3 hours or 24 hours following CFC for c-Fos-ZsGreen1-DR-injected mice and RAM-injected mice respectively. To attempt to reduce stress-related unspecific c-Fos activation, mice were anesthetized with a ketamine/xylazine mix (100 mg/kg/10 mg/kg; i.p.) and intracardially perfused with an ice-cold oxygenated (95% O₂ and 5% CO₂) cutting solution containing (in mM): 87 NaCl, 2.5 KCl, 1.25 NaH₂PO₄, 25 NaHCO₃, 25 glucose, 75 sucrose, 0.5 CaCl₂, 7 MgCl₂, pH 7.4, 315 mOsm/kg. Brains were quickly dissected out and parasagittal slices from both hemispheres (300 µm thick) were cut on a vibratome (Leica VT1200S, Germany) in the same oxygenated ice-cold cutting solution. Slices were then incubated at 34°C in a resting chamber containing the same cutting solution for 20-30 minutes for recovery. Slices were then transferred in a resting chamber filled with an oxygenated (95% O₂ and 5% CO₂) artificial cerebrospinal fluid (aCSF) containing (mM): 125 NaCl, 2.5 KCl, 1.25 NaH₂PO₄, 26 NaHCO₃, 1.3 MgCl₂, 2.3 CaCl₂, 10 glucose, pH 7.4, 300 mOsm/kg at room temperature for the rest of the day.

Electrophysiology

When transferred to a recording chamber, slices were continuously superfused with aCSF. CA3 pyramidal cells were identified with an infrared differential interference contrast (IR-DIC) microscope with a water immersion 63X objective (N.A. 0.8). Neurons were then classified as engram neurons (simultaneous expression of ZsGreen1-DR/mCherry or EGFP/mCherry when RAM was used) or non-engram neuron (mCherry expression only) with a 2p microscope. Whole-cell recordings were made at room temperature using borosilicate glass capillaries with resistances between 4-6 MΩ. For current clamp recordings, the intracellular solution contained (in mM): 135 K-gluconate, 5 KCl, 2 NaCl, 10 Na₂-phosphocreatine, 0.1 EGTA, 10 HEPES, 5 Mg-ATP, 0.4 Na-GTP, pH 7.2 adjusted with KOH, 280-290 mOsm/Kg. For voltage clamp mode, the patch pipettes were filled with a solution containing (mM): 140 Cs-methanesulfonate, 2 MgCl₂, 4 NaCl, 5 Na-phosphocreatine,

0.2 EGTA, 10 HEPES, 3 Na₂-ATP, 0.3 GTP, pH 7.2 adjusted with CsOH, 280-290 mOsm/Kg. To record inhibitory currents, a high-chloride intracellular solution was used, containing (mM): 103 CsCl, 12 CsCH₃O₃S, 5 TEA-Cl, 10 HEPES, 4 Mg-ATP, 0.5 Na-GTP, 1 MgCl₂, pH adjusted to 7.2 with CsOH, 280-290 mOsm/Kg. No liquid-junction correction was used. Neurons were held at -70 mV. Leak current and series resistance were monitored throughout each experiment; neurons with a leak current >200 pA and a series resistance >25MΩ or if these parameters changed more than 20% were excluded from analysis. All recordings were performed 5-10 min after opening the membrane. Input resistance was quantified by linearly fitting the voltage change against injected current. Steps of hyper- and depolarizing current steps were injected to determine action potential threshold and rheobase. Action potential firing pattern was determined by injecting 200 pA steps of current. Spontaneous EPSCs (sEPSCs) were isolated by bath application of bicuculline (10 μM), while mEPSCs were recorded in presence of bicuculline (10 μM) and tetrodotoxin (0.5 μM). Spontaneous IPSCs (sIPSCs) were recorded in presence of D-AP5 (50 μM) and NBQX (20 μM), while mIPSCs were isolated in presence of D-AP5 (50 μM), NBQX (20 μM) and TTX (0.5 μM). Spontaneous and miniature synaptic events were detected using a deconvolution-based approach (Pernía-Andrade et al., 2012). To record evoked EPSCs (eEPSCs), stimulating glass pipettes (World Precision Instruments) were filled with aCSF and placed in the hilus, close to the granule cell layer to stimulate initial portion of Mf axons. Voltage pulses (200 μs) were delivered through a stimulus isolator (Digitimer, UK). Stimulation intensity was adjusted to obtain minimal Mf stimulation (Marchal and Mulle, 2004). Basal synaptic transmission was recorded while holding CA3 neurons at -70 mV with pulses delivered at a low frequency of 0.1 Hz. Frequency was then increased to 1 Hz to induce frequency facilitation. After 2-3 minutes, when facilitation was extinguished, the voltage was switched to +5 mV (reversal potential for cations) to record di-synaptic eIPSC. All recordings were obtained with a HEKA EPC10 amplifier, filtered at 3.3 kHz and digitized at 10 kHz via PatchMaster software (Lambrecht, Germany). Data was analyzed offline using custom-made Python scripts.

Statistical analysis and graphical representations

Statistical analyses and graphical representation were performed with custom-written Python codes. Data are presented as boxplots and scatter plots indicating the number of samples per group. Boxplots indicate median, first and third quartiles and whiskers represent the rest of the distribution. Bold horizontal and vertical lines on scattered data points represent mean and SEM respectively. Values were first tested for normality (D'Agostino and Pearson omnibus test or Shapiro-Wilk test). Normally distributed data were compared using a t-test (two-sided) while Mann-Whitney rank test (two-sided) was used for non-normal data set. Multiple group comparison was carried out using one-way ANOVA when data were normally distributed or Kruskalis-Wallis when normality did not apply. Tukey's *post hoc* comparison analysis or Dunn *post hoc* comparison analysis was applied. Cumulative frequency distributions and normalized distributions were tested using Kolgomorov-Smirnov test. Statistical differences were considered as significant at $p < 0.05$.

Supplementary Figures

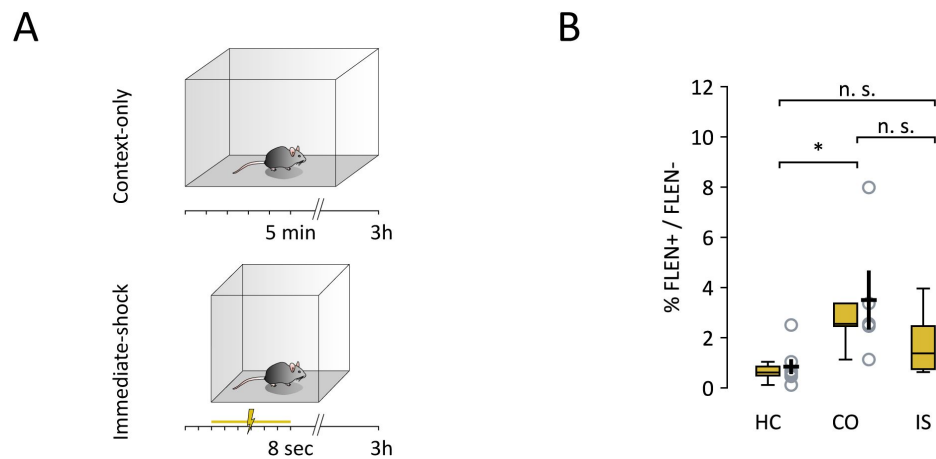


Figure S1.

(A) Left panel: context-only and immediate-shock behavioral test layout. In the context-only (CO) task, mice freely explore an arena without being conditioned; mice subjected to the immediate-shock (IS) experience a long (8 seconds) foot-shock immediately after being placed in the conditioning cage, preventing them exploring the environment and forming contextual memories. Right panel: plot comparing the percentage of FLEN+ cells in the home cage group (from **Figure 2D**) and the context-only and immediate-shock groups.

A

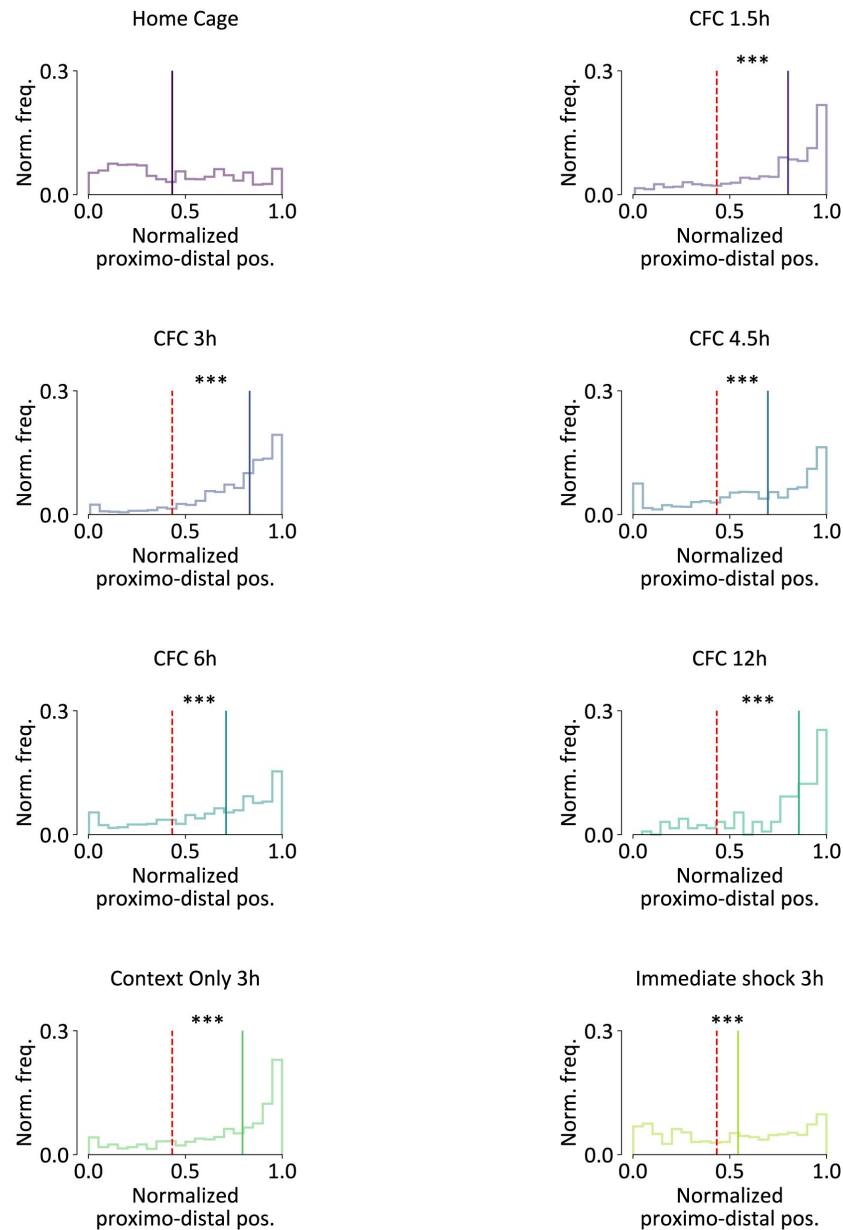


Figure S2.

(A) Normalized proximodistal (x) and superficial-to-deep (y) frequency distribution of the position of CA3 FLEN+ neurons in all behavioral groups. Black dashed lines indicate each group's median value, while red dashed line represents the home cage group's median value. Each plot includes the scattered x-y position of FLEN+ cells within the pyramidal layer, and the frequency distribution along the proximodistal (top) and superficial-to-deep (right) axis.

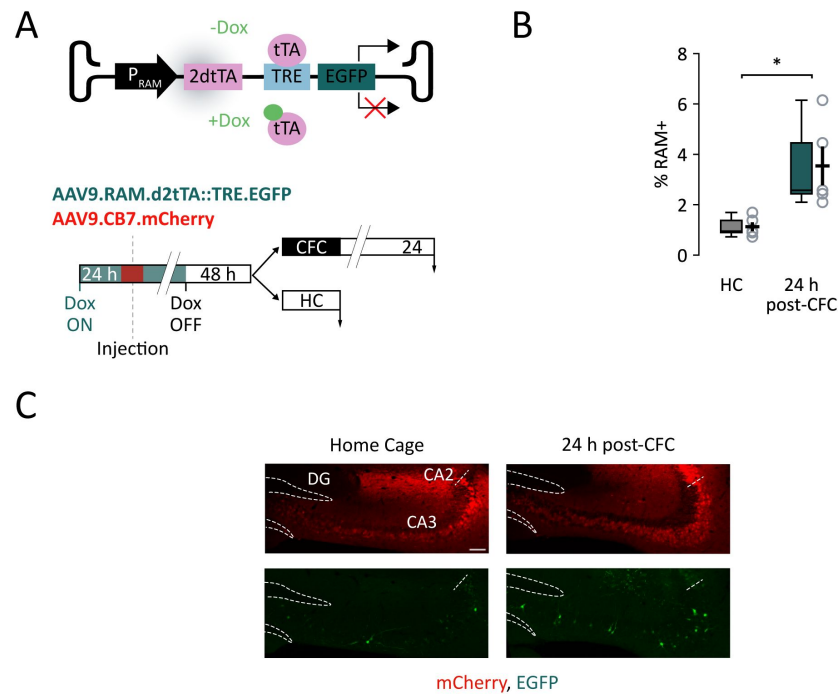


Figure S3.

(A) RAM construct outline and expression mechanism. Top panel: Dox administration prevents binding of tTA protein to the TRE promoter, whereas Dox deprivation allows downstream EGFP transcription. Bottom panel: scheme for *in vivo* experiments using the RAM system. Dox-fed mice are bilaterally injected in CA3 with viral constructs for RAM and the infection marker AAV-CB7-mCherry. The diet is switched to regular food chow 48 hours prior to the CFC. Then, mice undergo CFC and sections are collected (downward arrows) 24 hours later and compared to untrained mice that never left their home cage. (B) Percentage of RAM+ CA3 neurons over the total number of mCherry+ neurons in HC and 24 hours post-CFC. (C) Representative sections of CFC-trained mice compared to HC mice. Dashed lines outline the dentate gyrus (DG) cellular layer, while the dashed segment indicates the separation between CA3 and CA2. Scale bar = 100 μ m.

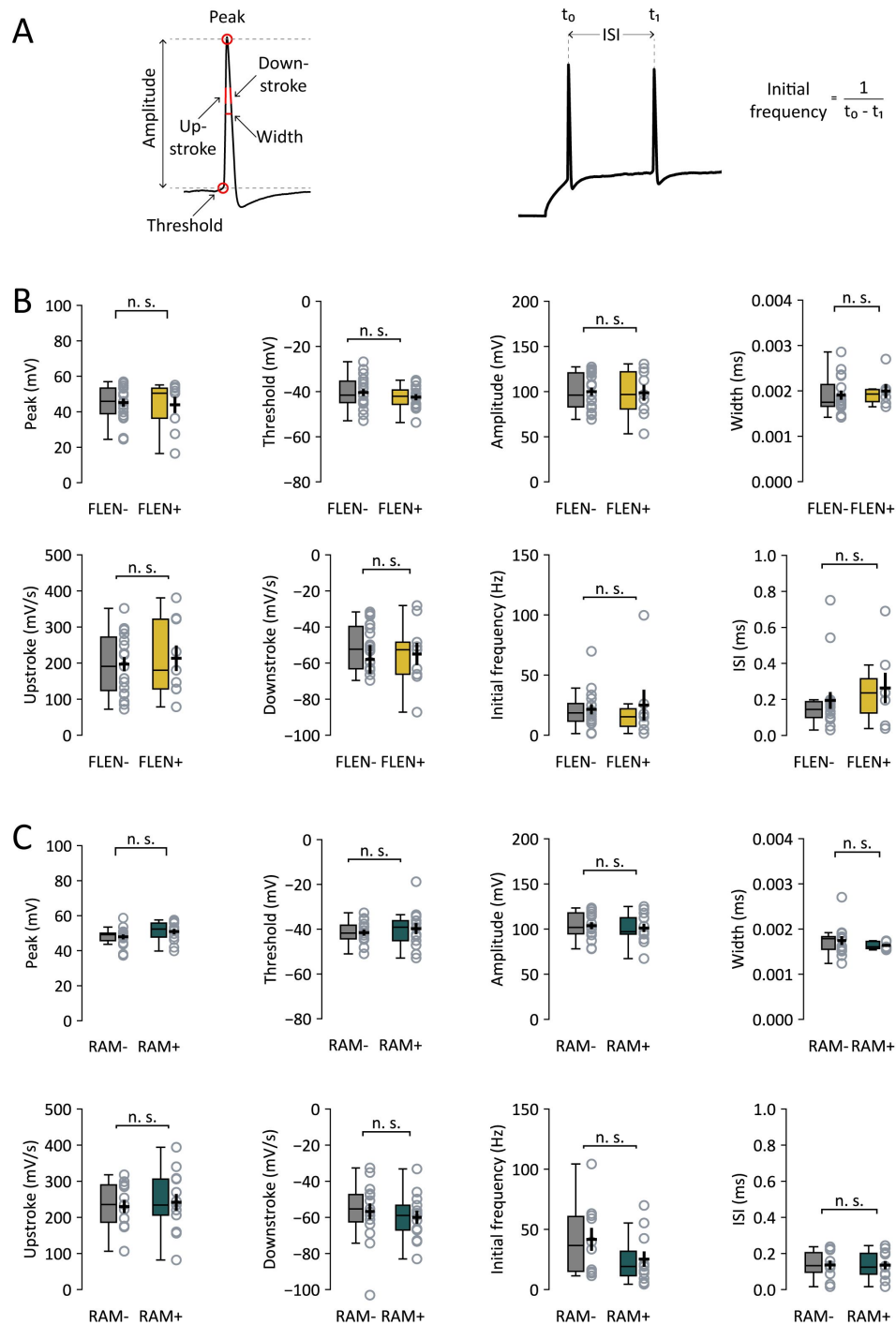


Figure S4.

(A) Left panel: overview of action potential features analyzed. Right panel: Initial frequency calculation. (B and C) Summary of analyzed action potential properties in FLEN+ vs. FLEN-(B) and RAM+ vs. RAM-(C).

References

- Bessi res B., Dupuis J., Groc L., Bontempi B., Nicole O. 2024) **Synaptic rearrangement of NMDA receptors controls memory engram formation and malleability in the cortex** *Science Advances* **10**:35 <https://doi.org/10.1126/sciadv.ado1148>
- Cupolillo D. 2021) **Comparing properties of CA3 engram neurons over initial periods of encoding and early phase of consolidation** PhD, Universit  de Bordeaux <https://theses.hal.science/tel-03336166v1>
- DeNardo L. A., Liu C. D., Allen W. E., Adams E. L., Friedmann D., Fu L., Guenther C. J., Tessier-Lavigne M., Luo L. 2019) **Temporal evolution of cortical ensembles promoting remote memory retrieval** *Nature Neuroscience* **22**:460–469 <https://doi.org/10.1038/s41593-018-0318-7>
- Denny C. A., Kheirbek M. A., Alba E. L., Tanaka K. F., Brachman R. A., Laughman K. B., Tomm N. K., Turi G. F., Losonczy A., Hen R. 2014) **Hippocampal Memory Traces Are Differentially Modulated by Experience, Time, and Adult Neurogenesis** *Neuron* **83**:189–201 <https://doi.org/10.1016/j.neuron.2014.05.018>
- Ding L., Chen H., Diamantaki M., Coletta S., Preston-Ferrer P., Burgalossi A. 2020) **Structural Correlates of CA2 and CA3 Pyramidal Cell Activity in Freely-Moving Mice** *The Journal of Neuroscience* **40**:5797–5806 <https://doi.org/10.1523/JNEUROSCI.0099-20.2020>
- Eichenbaum H. 2000) **A cortical-hippocampal system for declarative memory** *Nature Reviews Neuroscience* **1**:41–50 <https://doi.org/10.1038/35036213>
- Galimberti I., Gogolla N., Alberi S., Santos A. F., Muller D., Caroni P. 2006) **Long-Term Rearrangements of Hippocampal Mossy Fiber Terminal Connectivity in the Adult Regulated by Experience** *Neuron* **50**:749–763 <https://doi.org/10.1016/j.neuron.2006.04.026>
- Goto A., Bota A., Miya K., Wang J., Tsukamoto S., Jiang X., Hirai D., Murayama M., Matsuda T., McHugh T. J., Nagai T., Hayashi Y. 2021) **Stepwise synaptic plasticity events drive the early phase of memory consolidation** *Science* **374**:857–863 <https://doi.org/10.1126/science.abj9195>
- Han J.-H., Kushner S. A., Yiu A. P., Hsiang H.-L., Buch T., Waisman A., Bontempi B., Neve R. L., Frankland P. W., Josselyn S. A. 2009) **Selective Erasure of a Fear Memory** *Science* **323**:1492–1496 <https://doi.org/10.1126/science.1164139>
- Henze D. A., Wittner L., Buzs ki G. 2002) **Single granule cells reliably discharge targets in the hippocampal CA3 network in vivo** *Nature Neuroscience* **5**:790–795 <https://doi.org/10.1038/nn887>
- Holtmaat A., Caroni P. 2016) **Functional and structural underpinnings of neuronal assembly formation in learning** *Nature Neuroscience* **19**:1553–1562 <https://doi.org/10.1038/nn.4418>
- Humeau Y., Choquet D. 2019) **The next generation of approaches to investigate the link between synaptic plasticity and learning** *Nature Neuroscience* **22**:1536–1543 <https://doi.org/10.1038/s41593-019-0480-6>

- Josselyn S. A., Tonegawa S. 2020) **Memory engrams : Recalling the past and imagining the future** *Science* **367**:eaaw4325 <https://doi.org/10.1126/science.aaw4325>
- Kamiya H., Ozawa S. 1999) **Dual mechanism for presynaptic modulation by axonal metabotropic glutamate receptor at the mouse mossy fibre-CA3 synapse** *The Journal of Physiology* **518**:497–506 <https://doi.org/10.1111/j.1469-7793.1999.0497p.x>
- Kawashima T., Kitamura K., Suzuki K., Nonaka M., Kamijo S., Takemoto-Kimura S., Kano M., Okuno H., Ohki K., Bito H. 2013) **Functional labeling of neurons and their projections using the synthetic activity-dependent promoter E-SARE** *Nature Methods* **10**:889–895 <https://doi.org/10.1038/nmeth.2559>
- Kesner R. P., Rolls E. T. 2015) **A computational theory of hippocampal function, and tests of the theory : New developments** *Neuroscience & Biobehavioral Reviews* **48**:92–147 <https://doi.org/10.1016/j.neubiorev.2014.11.009>
- Kitamura T., Ogawa S. K., Roy D. S., Okuyama T., Morrissey M. D., Smith L. M., Redondo R. L., Tonegawa S. 2017) **Engrams and circuits crucial for systems consolidation of a memory** *Science* **356**:73–78 <https://doi.org/10.1126/science.aam6808>
- Li X., Zhao X., Fang Y., Jiang X., Duong T., Fan C., Huang C.-C., Kain S. R. 1998) **Generation of Destabilized Green Fluorescent Protein as a Transcription Reporter** *Journal of Biological Chemistry* **273**:34970–34975 <https://doi.org/10.1074/jbc.273.52.34970>
- Liu X., Ramirez S., Pang P. T., Puryear C. B., Govindarajan A., Deisseroth K., Tonegawa S. 2012) **Optogenetic stimulation of a hippocampal engram activates fear memory recall** *Nature* **484**:381–385 <https://doi.org/10.1038/nature11028>
- Lovett-Barron M., Kaifosh P., Kheirbek M. A., Danielson N., Zaremba J. D., Reardon T. R., Turi G. F., Hen R., Zeman B. V., Losonczy A. 2014) **Dendritic Inhibition in the Hippocampus Supports Fear Learning** *Science* **343**:857–863 <https://doi.org/10.1126/science.1247485>
- Marchal C., Mulle C. 2004) **Postnatal maturation of mossy fibre excitatory transmission in mouse CA3 pyramidal cells : A potential role for kainate receptors** *The Journal of Physiology* **561**:27–37 <https://doi.org/10.1113/jphysiol.2004.069922>
- Marneffe C., Moreira-de-Sá A., Lecomte S., Erhardt A., Mulle C. 2024) **Short term plasticity at hippocampal mossy fiber synapses.** *Neuroscience* **S 0306452224004974** <https://doi.org/10.1016/j.neuroscience.2024.09.044>
- Maruo T., Mandai K., Takai Y., Mori M. 2016) **Activity-dependent alteration of the morphology of a hippocampal giant synapse** *Molecular and Cellular Neuroscience* **71**:25–33 <https://doi.org/10.1016/j.mcn.2015.12.005>
- Matz M. V., Fradkov A. F., Labas Y. A., Savitsky A. P., Zaraisky A. G., Markelov M. L., Lukyanov S. A. 1999) **Fluorescent proteins from nonbioluminescent Anthozoa species** *Nature Biotechnology* **17**:969–973 <https://doi.org/10.1038/13657>
- McNaughton L., Morris G. M. 1987) **Hippocampal synaptic enhancement and information storage within a distributed memory system** *Trends in Neurosciences* **10**:10–415
- Milczarek M. M., Vann S. D., Sengpiel F. 2018) **Spatial Memory Engram in the Mouse Retrosplenial Cortex** *Current Biology* **28**:1975–1980 <https://doi.org/10.1016/j.cub.2018.05.002>

- Mocle A. J., Ramsaran A. I., Jacob A. D., Rashid A. J., Luchetti A., Tran L. M., Richards B. A., Frankland P. W., Josselyn S. A. 2024) **Excitability mediates allocation of pre-configured ensembles to a hippocampal engram supporting contextual conditioned threat in mice** *Neuron* **112**:1487–1497 <https://doi.org/10.1016/j.neuron.2024.02.007>
- Nakamura N. H., Flasbeck V., Maingret N., Kitsukawa T., Sauvage M. M. 2013) **Proximodistal Segregation of Nonspatial Information in CA3 : Preferential Recruitment of a Proximal CA3-Distal CA1 Network in Nonspatial Recognition Memory** *Journal of Neuroscience* **33**:11506–11514 <https://doi.org/10.1523/JNEUROSCI.4480-12.2013>
- Nakazawa K., Quirk M. C., Chitwood R. A., Watanabe M., Yeckel M. F., Sun L. D., Kato A., Carr C. A., Johnston D., Wilson M. A., Tonegawa S. 2002) **Requirement for Hippocampal CA3 NMDA Receptors in Associative Memory Recall** *Science* **297**:211–218 <https://doi.org/10.1126/science.1071795>
- Nakazawa K., Sun L. D., Quirk M. C., Rondi-Reig L., Wilson M. A., Tonegawa S. 2003) **Hippocampal CA3 NMDA Receptors Are Crucial for Memory Acquisition of One-Time Experience** *Neuron* **38**:305–315 [https://doi.org/10.1016/S0896-6273\(03\)00165-X](https://doi.org/10.1016/S0896-6273(03)00165-X)
- Nakazawa Y., Pevzner A., Tanaka K. Z., Wiltgen B. J. 2016) **Memory retrieval along the proximodistal axis of CA1 : PROXIMODISTAL CA1** *Hippocampus* **26**:1140–1148 <https://doi.org/10.1002/hipo.22596>
- Pernía-Andrade A. J., Goswami S. P., Stickler Y., Fröbe U., Schlögl A., Jonas P. 2012) **A Deconvolution-Based Method with High Sensitivity and Temporal Resolution for Detection of Spontaneous Synaptic Currents In Vitro and In Vivo** *Biophysical Journal* **103**:1429–1439 <https://doi.org/10.1016/j.bpj.2012.08.039>
- Pignatelli M., Ryan T. J., Roy D. S., Lovett C., Smith L. M., Muralidhar S., Tonegawa S. 2019) **Engram Cell Excitability State Determines the Efficacy of Memory Retrieval** *Neuron* **101**:274–284 <https://doi.org/10.1016/j.neuron.2018.11.029>
- Raus Balind S., Magó Á., Ahmadi M., Kis N., Varga-Németh Z., Lőrincz A., Makara J. K. 2019) **Diverse synaptic and dendritic mechanisms of complex spike burst generation in hippocampal CA3 pyramidal cells** *Nature Communications* **10**:1859 <https://doi.org/10.1038/s41467-019-09767-w>
- Rebola N., Carta M., Mulle C. 2017) **Operation and plasticity of hippocampal CA3 circuits : Implications for memory encoding** *Nature Reviews Neuroscience* **18**:208–220 <https://doi.org/10.1038/nrn.2017.10>
- Reijmers L. G., Perkins B. L., Matsuo N., Mayford M. 2007) **Localization of a Stable Neural Correlate of Associative Memory** *Science* **317**:1230–1233 <https://doi.org/10.1126/science.1143839>
- Ruediger S., Vittori C., Bednarek E., Genoud C., Strata P., Sacchetti B., Caroni P. 2011) **Learning-related feedforward inhibitory connectivity growth required for memory precision** *Nature* **473**:514–518 <https://doi.org/10.1038/nature09946>
- Ryan T. J., Roy D. S., Pignatelli M., Arons A., Tonegawa S. 2015) **Engram cells retain memory under retrograde amnesia** *Science* **348**:1007–1013 <https://doi.org/10.1126/science.aaa5542>
- Sachidhanandam S., Blanchet C., Jeantet Y., Cho Y. H., Mulle C. 2009) **Kainate Receptors Act as Conditional Amplifiers of Spike Transmission at Hippocampal Mossy Fiber Synapses** *The*

Journal of Neuroscience **29**:5000–5008 <https://doi.org/10.1523/JNEUROSCI.5807-08.2009>

Sørensen A. T., Cooper Y. A., Baratta M. V., Weng F.-J., Zhang Y., Ramamoorthi K., Fropf R., LaVerriere E., Xue J., Young A., Schneider C., Göttsche C. R., Hemberg M., Yin J. C., Maier S. F., Lin Y. 2016) **A robust activity marking system for exploring active neuronal ensembles** *eLife* **5**:e13918 <https://doi.org/10.7554/eLife.13918>

Sun Q., Sotayo A., Cazzulino A. S., Snyder A. M., Denny C. A., Siegelbaum S. A. 2017) **Proximodistal Heterogeneity of Hippocampal CA3 Pyramidal Neuron Intrinsic Properties, Connectivity, and Reactivation during Memory Recall** *Neuron* **95**:656–672 <https://doi.org/10.1016/j.neuron.2017.07.012>

Sun X., Bernstein M. J., Meng M., Rao S., Sørensen A. T., Yao L., Zhang X., Anikeeva P. O., Lin Y. 2020) **Functionally Distinct Neuronal Ensembles within the Memory Engram** *Cell* **181**:410–423 <https://doi.org/10.1016/j.cell.2020.02.055>

Takeuchi T., Duzskiewicz A. J., Morris R. G. M. 2014) **The synaptic plasticity and memory hypothesis : Encoding, storage and persistence** *Philosophical Transactions of the Royal Society B: Biological Sciences* **369**:20130288 <https://doi.org/10.1098/rstb.2013.0288>

Thompson L. T., Moyer J. R., Disterhoft J. F. 1996) **Transient changes in excitability of rabbit CA3 neurons with a time course appropriate to support memory consolidation** *Journal of Neurophysiology* **76**:1836–1849 <https://doi.org/10.1152/jn.1996.76.3.1836>

Tomé D. F., Zhang Y., Aida T., Mosto O., Lu Y., Chen M., Sadeh S., Roy D. S., Clopath C. 2024) **Dynamic and selective engrams emerge with memory consolidation** *Nature Neuroscience* **27**:561–572 <https://doi.org/10.1038/s41593-023-01551-w>

Tonegawa S., Liu X., Ramirez S., Redondo R. 2015) **Memory Engram Cells Have Come of Age** *Neuron* **87**:918–931 <https://doi.org/10.1016/j.neuron.2015.08.002>

Torborg C. L., Nakashiba T., Tonegawa S., McBain C. J. 2010) **Control of CA3 Output by Feedforward Inhibition Despite Developmental Changes in the Excitation–Inhibition Balance** *The Journal of Neuroscience* **30**:15628–15637 <https://doi.org/10.1523/JNEUROSCI.3099-10.2010>

Tulving E. 2002) **Episodic Memory : From Mind to Brain** *Annual Review of Psychology* **53**:1–25 <https://doi.org/10.1146/annurev.psych.53.100901.135114>

Vandael D., Jonas P. 2024) **Structure, biophysics, and circuit function of a “giant” cortical presynaptic terminal** *Science* **383**:eadg6757 <https://doi.org/10.1126/science.adg6757>

Vetere G., Tran L. M., Moberg S., Steadman P. E., Restivo L., Morrison F. G., Ressler K. J., Josselyn S. A., Frankland P. W. 2019) **Memory formation in the absence of experience** *Nature Neuroscience* **22**:933–940 <https://doi.org/10.1038/s41593-019-0389-0>

Weng F.-J., Garcia R. I., Lutz S., Alviña K., Zhang Y., Dushko M., Ku T., Zemoura K., Rich D., Garcia-Dominguez D., Hung M., Yelhekar T. D., Sørensen A. T., Xu W., Chung K., Castillo P. E., Lin Y. 2018) **Npas4 Is a Critical Regulator of Learning-Induced Plasticity at Mossy Fiber-CA3 Synapses during Contextual Memory Formation** *Neuron* **97**:1137–1152 <https://doi.org/10.1016/j.neuron.2018.01.026>

Yap E.-L., Pettit N. L., Davis C. P., Nagy M. A., Harmin D. A., Golden E., Dagliyan O., Lin C., Rudolph S., Sharma N., Griffith E. C., Harvey C. D., Greenberg M. E. 2021) **Bidirectional**

perisomatic inhibitory plasticity of a Fos neuronal network *Nature* **590**:115–121 <https://doi.org/10.1038/s41586-020-3031-0>

Yiu A. P., Mercaldo V., Yan C., Richards B., Rashid A. J., Hsiang H.-L. L., Pressey J., Mahadevan V., Tran M. M., Kushner S. A., Woodin M. A., Frankland P. W., Josselyn S. A. 2014) **Neurons Are Recruited to a Memory Trace Based on Relative Neuronal Excitability Immediately before Training** *Neuron* **83**:722–735 <https://doi.org/10.1016/j.neuron.2014.07.017>

Zhang W., Linden D. J. 2003) **The other side of the engram : Experience-driven changes in neuronal intrinsic excitability** *Nature Reviews Neuroscience* **4**:885–900 <https://doi.org/10.1038/nrn1248>

Zucca S., Griguoli M., Malézieux M., Grosjean N., Carta M., Mulle C. 2017) **Control of Spike Transfer at Hippocampal Mossy Fiber Synapses In Vivo by GABA A and GABA B Receptor-Mediated Inhibition** *The Journal of Neuroscience* **37**:587–598 <https://doi.org/10.1523/JNEUROSCI.2057-16.2016>

Author information

Dario Cupollilo

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France, Neuroscience and Brain Technologies Department, Fondazione Istituto Italiano di Tecnologia, Genova, Italy
ORCID iD: [0000-0002-1517-3815](https://orcid.org/0000-0002-1517-3815)

Noëlle Grosjean

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France

Catherine Marneffe

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France
ORCID iD: [0000-0002-4155-6369](https://orcid.org/0000-0002-4155-6369)

Julio Viotti

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France

Célia Reynaud

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France

Séverine Deforges

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France

Mario Carta

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux, Bordeaux, France
ORCID iD: [0000-0003-1225-2742](https://orcid.org/0000-0003-1225-2742)

Christophe Mulle*

Interdisciplinary Institute for Neuroscience, CNRS UMR 5297, University of Bordeaux,
Bordeaux, France

ORCID iD: [0000-0003-2709-6615](https://orcid.org/0000-0003-2709-6615)

For correspondence: christophe.mulle@u-bordeaux.fr

*Lead contact

Editors

Reviewing Editor

Katalin Toth

University of Ottawa, Ottawa, Canada

Senior Editor

Laura Colgin

University of Texas at Austin, Austin, United States of America

Reviewer #1 (Public review):

Summary:

The manuscript by Cupollilo et al describes the development, characterization, and application of a novel activity labeling system; fast labelling of engram neurons (FLEN). Several such systems already exist but this study adds additional capability by leveraging an activity marker that is destabilized (and thus temporally active) as well as being driven by the full-length promoter of cFos. The authors demonstrate the activity-dependent induction and time course of expression, first in cultured neurons and then in vivo in hippocampal CA3 neurons after one trial of contextual fear conditioning. In a series of ex vivo experiments, the authors perform patch clamp analysis of labeled neurons to determine if these putative engram neurons differ from non-labelled neurons using both the FLEN system as well as the previously characterized RAM system. Interestingly the early labelled neurons at 3 h post CFC (FLEN+) demonstrated no differences in excitability whereas the RAM-labelled neurons at 24h after CFC had increased excitability. Examination of synaptic properties demonstrated an increase in sEPSCs and mEPSC frequencies as well as those for sIPSCs and mIPSCs which was not due to a change in the mossy fiber input to these neurons.

Strengths:

Overall the data is of high quality and the study introduces a new tool while also reassessing some principles of circuit plasticity in the CA3 that have been the focus of prior studies.

Weaknesses:

No major weaknesses were noted.

<https://doi.org/10.7554/eLife.105452.1.sa1>

Reviewer #2 (Public review):

Summary:

Cupollilo et al. investigate the properties of hippocampal CA3 neurons that express the immediate early gene cFos in response to a single foot shock. They compare ex-vivo the electrophysiological properties of these "engram neurons" labeled with two different cFos promoter-driven green markers: Their new tool FLEN labels neurons 2-6 h after activity, while RAM contains additional enhancers and peaks considerably later (>24 h). Since the fraction of labeled CA3 cells is comparable with both constructs, it is assumed (but not tested) that they label the same population of activated neurons at different time points. Both FLEN+ and RAM+ neurons in CA3 receive more synaptic inputs compared to non-expressing control neurons, which could be a causal factor for cFos activation, or a very early consequence thereof. Frequency facilitation and E/I ratio of mossy fiber inputs were also tested, but are not different in both cFos+ groups of neurons. One day after foot shock, RAM+ neurons are more excitable than RAM- neurons, suggesting a slow increase in excitability as a major consequence of cFos activation.

Strengths:

The study is conducted to high standards and contributes significantly to our understanding of memory formation and consolidation in the hippocampus. Modifications of intrinsic neuronal properties seem to be more salient than overall changes in the total number of (excitatory and inhibitory) inputs, although a switch in the source of the synaptic inputs would not have been detected by the methods employed in this study

Weaknesses:

With regard to the new viral tool, a direct comparison between the new tool FLEN and existing cFos reporters is missing.

<https://doi.org/10.7554/eLife.105452.1.sa0>