

Natural forgetting reversibly modulates engram expression

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Abstract

Summary

Memories are stored as ensembles of engram neurons and their successful recall involves the reactivation of these cellular networks. However, significant gaps remain in connecting these cell ensembles with the process of forgetting. Here, we utilized a mouse model of object memory and investigated the conditions in which a memory could be preserved, retrieved, or forgotten. Direct modulation of engram activity via optogenetic stimulation or inhibition either facilitated or prevented the recall of an object memory. In addition, through behavioral and pharmacological interventions, we successfully prevented or accelerated forgetting of an object memory. Finally, we showed that these results can be explained by a computational model in which engrams that are subjectively less relevant for adaptive behavior are more likely to be forgotten. Together, these findings suggest that forgetting may be an adaptive form of engram plasticity which allows engrams to switch from an accessible state to an inaccessible state.

eLife assessment

This is an **important** paper on the role of engrams and relevant conditions that influence memory and forgetting. The variety of methods used, namely, behavioural, labeling, interrogation, immunohistochemistry, microscopy, pharmacology, computational, are exemplary and provide **convincing** evidence for the role of engrams in the dentate gyrus in memory retrieval and forgetting. This examination will be of interest broadly across behavioural and neural science communities.

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Introduction

For an organism to adapt to its environment there must be processes by which learned information is updated or replaced with new more relevant information (1, 2). For example, learning that a specific food source is no longer present in a previous location, or that a given action stops yielding the same outcome (1, 3, 4). Without a process to update or forget irrelevant information, adaptability would be impaired with frequent instances of correct but outdated memory recall, and a subsequently biased behavioral response (2, 4, 5–7). Forgetting, that is the loss of learned behavioral responses, may therefore function adaptively to improve learning, and promote survival (8).

Memories are stored as ensembles of engram cells that undergo specific forms of plasticity during learning, and their successful recall involves the reactivation of these cellular networks (9–13). However, despite advancements in understanding the biology of engram cells, more work is still needed to fully elucidate how these cell ensembles contribute to the process of forgetting (8, 13, 14). Forgetting is often considered a deficit of memory function. The standard view in the field of memory is that this kind of active forgetting is due to the loss or dissipation of the engram/trace (8). We argue that a new conceptualization is necessitated by recent behavioral and physiological findings that emphasize retrieval deficits as a key characteristic of memory impairment, supporting the idea that memory recall or accessibility may be driven by learning feedback from the environment (7, 8, 14–18). Little is understood about the role experience plays in altering forgetting rates. Recently it has been suggested that different forms of forgetting may exist along a gradient of engram activity or expression (8). Indeed, numerous studies have suggested that memory recall (15, 17, 19–22), and most recently forgetting, require engram activity (18, 23). Long-term memories of salient experiences can last a lifetime and must involve significant and specific changes to brain structure, while the successful recall of learned information requires the reactivation of these cell ensembles (11). In contrast, forgetting may occur when engrams are not or cannot be reactivated (8, 18, 24). In severe amnesic states, this may be due to the destruction of the engram itself (8, 25–28). However, in other pathological and non-pathological states, forgetting may be due to reduced accessibility of engrams that otherwise endure (28). Altered engram accessibility may be caused by structural plasticity in the engram or indeed by competing engrams of similar or recent experiences (8, 18, 29–33).

We hypothesize that natural forgetting represents a reversible suppression of engram ensembles due to experience and perceptual feedback, prompting cellular plasticity processes that modulate memory access adaptively. To investigate this idea, we utilized a forgetting paradigm based on an object recognition task and interrogated the conditions in which a memory could be preserved, retrieved, or forgotten to elucidate how experience and learning govern engram reactivation and memory recall. Based on our experimental results, we used the Rescorla-Wagner model to explain our findings across studies in a formal framework. This model describes forgetting as a learning process through which engram accessibility is dynamically updated in response to perceptual feedback, and engrams that are subjectively less relevant for adaptive behavior are more likely to be ‘forgotten’.

Results

Natural forgetting of an object memory

Natural forgetting was assessed using an object recognition task. This task utilizes the natural tendency of mice to explore novel stimuli to assess recognition memory over time (**Figure 1a**) (34, 35). During object memory recall, mice tested 24 hours (h) after training spent significantly more time exploring the novel object compared to the familiar object, $t(9) = 5.12$, $p < 0.001$, 95% CI [11.18 to 28.68]. Mice that were tested 2 weeks (wk) after training spent a similar amount of time exploring both novel and familiar objects, $t(11) = 0.93$, $p = 0.36$, 95% CI [-3.88 to 9.64] (**Figure 1b**). In order to compare performance in object memory, a discrimination index was calculated. The discrimination index was calculated as the time spent exploring the novel object minus the time spent exploring the familiar object divided by the total time spent exploring both objects (novel – familiar / novel + familiar). There was a significant difference between the discrimination index of the 24 h and 2 wk group, $t(20) = 4.30$, $p < 0.001$, 95% CI [-0.59 to -0.20] (**Figure 1c**). This finding was further supported by a minute-by-minute analysis of object exploration (Supplementary Figure 1). Together, these data indicate that mice successfully recall an object memory 24 hours after acquisition, while at 2 weeks there was no preference for the novel object suggesting the original familiar object had been forgotten.

Engram reactivation is associated with object memory recall

In order to assess engram reactivation on object memory retrievability an AAV_g-TRE- ChR2-eYFP virus was injected into the dentate gyrus of c-fos-tTA mice (36, 37). The immediate early gene *c-fos* becomes upregulated following neural activity and as such the c-fos-tTA transgene selectively expresses tTA in active cells (37, 38). The activity-dependent tTA induces expression of ChR2-eYFP following a learning event (**Figure 1d**, e). To restrict activity-dependent labelling to a single learning experience, mice were maintained on a diet containing doxycycline (DOX) which prevents binding of tTA to the TRE elements of the TetO promoter, thus controlling the activity-dependent labelling to a specific time window of the object acquisition training (**Figure 1d-f**). Prior to training (36 h) doxycycline was removed from the animal's diet, thereby allowing for activity-dependent labelling during the subsequent acquisition training (**Figure 1f**). During the object recall test, c-fos-tTA mice tested 24 hours after acquisition training spent significantly more time exploring the novel object compared to the familiar object, $t(10) = 6.00$, $p < 0.001$, 95% CI [10.80 to 23.54]. Mice that were tested 2 weeks after training spent a similar amount of time exploring both novel and familiar objects, $t(7) = 0.67$, $p = 0.51$, 95% CI [-5.69 to 10.27] (**Figure 1g**). In order to compare performance in object memory, a discrimination index was calculated, as described earlier. There was a significant difference between the discrimination index of the 24 h and 2 wk group, $t(17) = 4.30$, $p < 0.001$, 95% CI [-0.47 to -0.16] (**Figure 1h**). When comparing the engram ensemble between the two cohorts, we observed no significant difference in the number of engram cells between the 24 h and 2 wk tested groups (eYFP⁺) $t(6) = 0.18$, $p = 0.86$, 95% CI [-2.24 to 2.60] (**Figure 1j**, k). Similarly, there was no significant difference in c-Fos activity within the dentate gyrus between the 24 h tested or 2 wk tested mice, $t(6) = 0.84$, $p = 0.42$, 95% CI [-1.37 to 0.66] (**Figure 1l**). However, when comparing engram reactivation, that is the number of engram cells (eYFP⁺) that also express c-Fos during the recall test, which indicates these cells were active during encoding and then became active again during the memory recall (**Figure 1j**), there was a significant decrease in engram reactivation in mice that were tested 2 weeks after training and that had forgotten the object memory, $t(6) = 4.56$, $p = 0.003$, 95% CI [-14.96 to -4.52] (**Figure 1m**). Furthermore, the level of engram reactivation correlated with the performance in object discrimination, $r(1,6) = 0.61$, $p = 0.02$ (**Figure 1i**). Together these data suggest that natural forgetting may be driven by a reduction in engram ensemble activity.

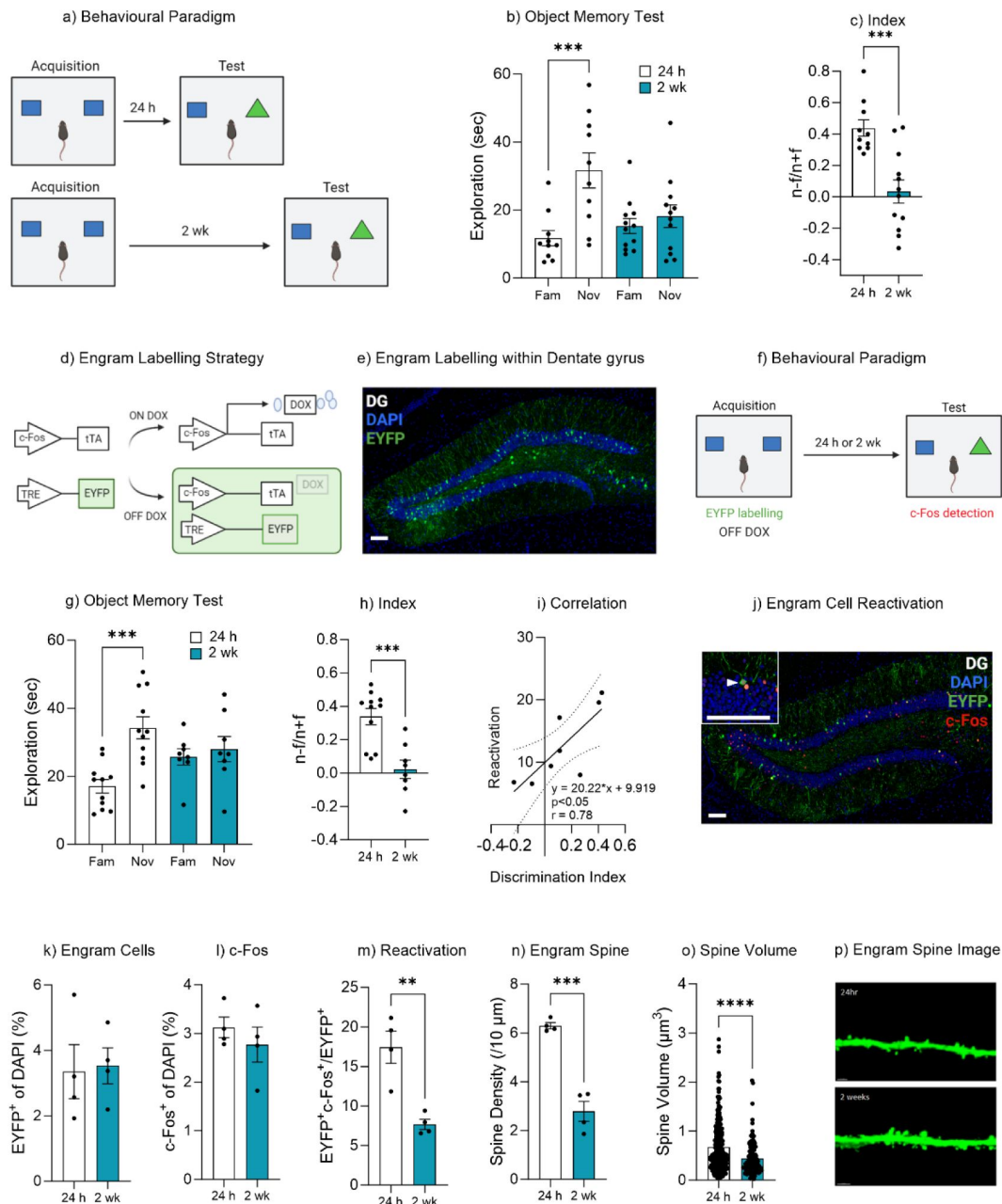


Figure 1

Natural Forgetting of an Object Memory

(a) Object recognition paradigm **(b)** Object memory test for recall at 24 h or 2 wk **(c)** Discrimination index **(d)** Schematic of c-Fos-tTA engram labelling system **(e)** Representative image of engram labeling within the dentate gyrus **(f)** Engram labeling of object memory and c-Fos detection following recall **(g)** Object memory test for recall at 24 h or 2 wk **(h)** Discrimination index **(i)** Correlation between Discrimination Index and Engram reactivation **(j)** Representative image eYFP⁺ cells, c-Fos⁺ cells and Merged eYFP⁺ and c-Fos⁺ for both 24 h and 2 wk test **(k)** Engram cells **(l)** c-Fos⁺ cells **(m)** Engram reactivation **(n)** Engram spine density average per mouse, **(o)** Engram spine volume average per dendrite **(p)** Representative image of engram dendrite for morphological analysis. Bar graphs indicates average values in $n = 4-11$ per group (** $p < 0.01$, *** $p < 0.001$). Data graphed as means $\pm$ SEM. Scale bar 100 μm .

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Engram spine density decreased following natural forgetting

In order to gain further insight into the changes that occur within the memory engram following natural forgetting we performed morphological analysis of engram dendritic spines following successful memory recall at 24 hours or forgetting at 2 weeks. Following natural forgetting mice displayed a significant reduction in spine density, $t(6) = 8.23$, $p < 0.001$, 95% CI [-4.55 to -2.46] (**Figure 1n**). Moreover, there was also a significant decrease in spine volume, $t(372) = 4.50$, $p < 0.001$, 95% CI [-0.33 to -0.13] (**Figure 1o**, p). Together, these findings suggest that in addition to the reduced engram activity, there was also a reduction in engram spine density.

Dentate gyrus engrams are necessary and sufficient for recall of an object memory

To test the hypothesis that the retrieval of an object memory requires the activity of engram cells within the dentate gyrus, we labelled engrams with the virus AAV_g-TRE- ArchT-GFP and inhibited engram activity during memory retrieval 24 hours following acquisition training (**Figure 2a**). During the object recall test the No Light control mice spent significantly more time with the novel object compared to the familiar object, indicating successful memory retrieval, $t(9) = 2.91$, $p = 0.01$, 95% CI [3.43 to 27.09] (**Figure 2b**). Whereas mice in which the engram was inhibited spent equal time exploring both the familiar and novel objects, indicating mice failed to distinguish the novel from the familiar object, $t(8) = 0.58$, $p = 0.57$, 95% CI [-4.21 to 7.09] (**Figure 2b**). In order to compare performance in object memory, a discrimination index was calculated, as described earlier. There was a significant difference between the discrimination index of the No Light control and Light-induced inhibition groups, $t(17) = 2.18$, $p = 0.04$, 95% CI [-0.49 to -0.008] (**Figure 2c**). Together, these data demonstrate that the engram cells within the dentate gyrus are required for successful retrieval of an object memory.

Since the data suggested that forgetting correlated with a reduction in engram activity (**Figure 1c**) and that inhibition of the ensemble was sufficient to impair successful recall of an object memory (**Figure 2b, c**), we next asked if the retrieval of an object memory could be induced via activation of the engram after natural forgetting. We labelled engrams cells within the dentate gyrus with the virus AAV_g-TRE-ChR2-eYFP and optogenetically activated the ensemble after natural forgetting. Specifically, optogenetic stimulation was induced just prior to memory recall (3min), but not during the recall test itself (**Figure 2d**). During the object recall test the No Light control mice displayed equal exploration of both the novel and familiar objects suggesting mice had forgotten the familiar object, $t(10) = 1.81$, $p = 0.10$, 95% CI [-0.90 to 8.84] (**Figure 2e**). Whereas mice that underwent optogenetic activation spent more time exploring the novel object compared to the familiar object, suggesting the original object memory had been retrieved, $t(7) = 3.41$, $p = 0.01$, 95% CI [3.71 to 20.43] (**Figure 2e**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference between the discrimination index of the No Light control and Light-induced activation group, $t(17) = 2.14$, $p = 0.04$, 95% CI [0.002 to 0.34] (**Figure 2f**). These data suggest that artificial activation of the original engram was sufficient to induce recall of the forgotten object memory, and that the forgotten engram was still intact, otherwise activation of those cells would no longer trigger memory recall. Together, the results of **Figure 2** suggest that engram activity can modulate memory retrieval, where activation of the engram ensemble is both necessary for successful memory retrieval as well as sufficient to induced recall despite natural forgetting.

Environmental enrichment reduces the rate of forgetting by increasing engram activation

Environmental enrichment has been shown to enhance cognitive function and improve memory retention (40–44). A key component of enriched environments is the additional perceptual and tactile feedback provided by extra toys and objects within the home-cage for the mice to

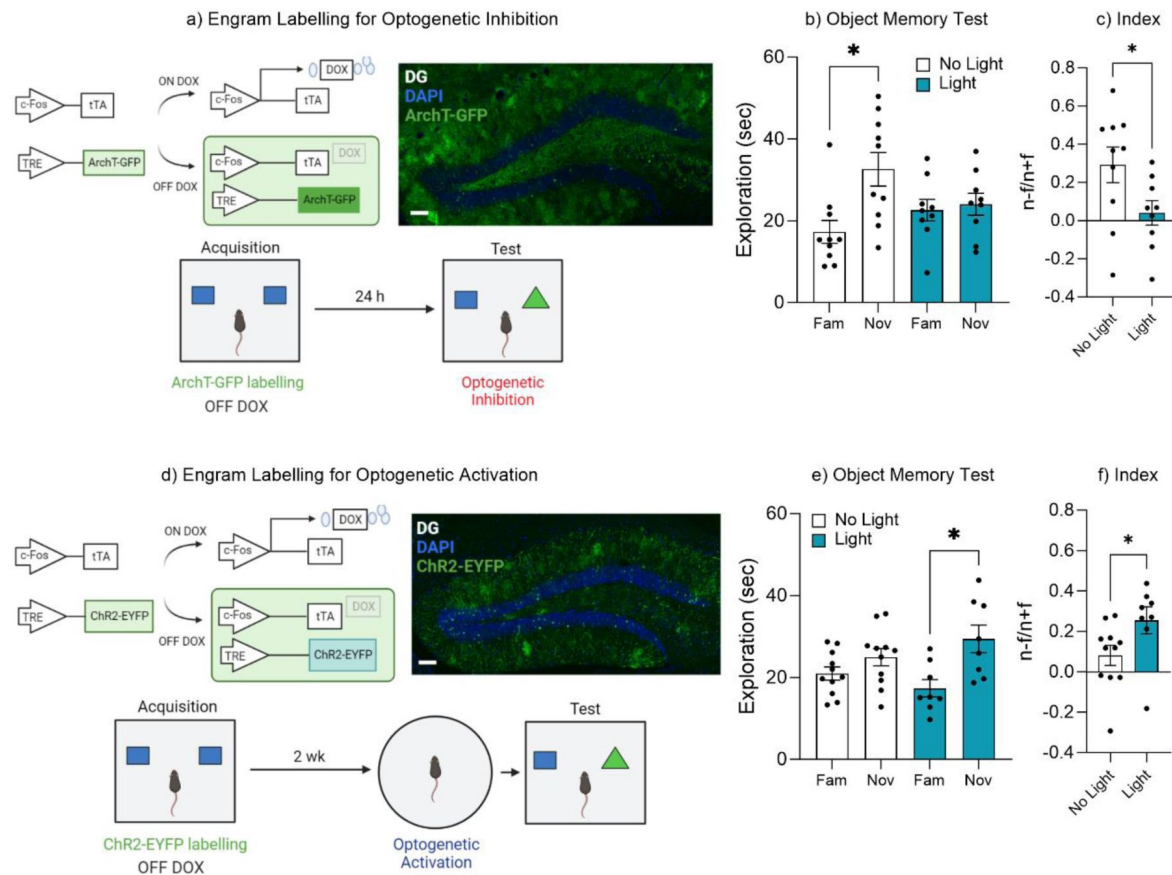


Figure 2

Dentate gyrus engrams are necessary and sufficient for recall of an object memory

(a) Engram labelling for optogenetic inhibition and behavioral timeline **(b)** Object memory test **(c)** Discrimination index **(d)** Engram labelling for optogenetic activation and behavioral timeline **(e)** Object memory test **(f)** Discrimination index Bar graphs indicate average values in $n = 9-12$ per group (* $p < 0.05$). Data graphed as means $\pm$ SEM. Scale bar 100 μ m. © 2024, BioRender Inc. Any parts of this image created with *BioRender* are not made available under the same license as the Reviewed Preprint, and are © 2024, BioRender Inc.

interact with and explore. We, therefore sought to investigate how enrichment might alter the rate of forgetting through changes in the engram given this increase in tactile and perceptual experience. First, we characterized the forgetting curve of mice housed under either standard or enriched conditions (**Figure 3a**). There was a significant effect of enriched housing on object memory performance $F(3, 53) = 3.26$, $p = 0.028$, 95% CI [-0.18 to 0.01] (**Figure 3b**). Bonferroni post hoc comparisons indicated there was no difference in object memory performance between the standard housed (Std) or Environmental Enrichment housed (EE) mice when tested 24 hours or 1 week after acquisition training, indicating no change in memory recall (**Figure 3b**). However, there was a significant difference in discrimination index between standard housed and enriched mice when tested 2 after training, ($p = 0.047$), suggesting the enriched housed mice were still able to recall the familiar object, whereas the standard housed mice had forgotten. This finding was further supported by a minute-by minute analysis of object exploration (Supplementary Figure 2).

In order to assess the impact of environmental enrichment on engram activity we again injected an AAV₉-TRE-ChR2-eYFP virus into c-fos-tTA mice to label engram cells within the dentate gyrus. Following surgery, mice were housed in either standard or enriched housing for 3 weeks before behavioral testing and remained in the housing condition for the duration of the experiment. Again, we observed that mice housed in enrichment maintained the object memory 2 weeks after training, as they spent more time exploring the novel object compared to the familiar object, $t(11) = 5.05$, $p < 0.001$, 95% CI [8.59 to 21.83]. Mice housed in standard housing spent a similar amount of time exploring both the novel and familiar objects, $t(11) = 1.77$, $p = 0.10$, 95% CI [-1.34 to 12.57] (**Figure 3c**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference between the discrimination index of standard housed (Std) and enriched (EE) mice, $t(22) = 2.75$, $p = 0.01$, 95% CI [0.054 to 0.38] (**Figure 3d**). When comparing the engram size between standard and enriched mice, we observed no significant difference in the number of engram cells following enrichment, $t(12) = 0.56$, $p = 0.58$, 95% CI [-3.09 to 1.82] (**Figure 3e & f**). There was a marginal, but non-significant increase in the number of labelled cells within the standard housed mice in **Figure 3f**. The cell counting criteria was the same across experimental groups and conditions, where the entire dorsal and ventral blade of the dorsal DG was counted for each animal. Similarly, there was also no significant difference in c-Fos activity, $t(12) = 1.45$, $p = 0.17$, 95% CI [-0.62 to 3.13] (**Figure 3g**). However, there was a significant increase in engram reactivation in mice that underwent enrichment, $t(12) = 3.52$, $p = 0.004$, 95% CI [2.84 to 12.06] (**Figure 3h**). These data suggest that environmental enrichment reduces the rate of forgetting, by increasing or maintaining engram reactivation. To confirm the effectiveness of the enrichment paradigm to enhance neural plasticity, we measured hippocampal neurogenesis following environmental enrichment. Enrichment increased hippocampal neurogenesis, with both the number of immature neurons (doublecortin⁺ cells) (Supplementary Figure S3a & b) as well as neuronal survival (BrdU-NeuN positive cells) (Supplementary Figure S3c & d). The effects of enrichment on neural plasticity are well established and it likely contributes to the enhanced memory recall. In addition, the enriched environment, as it contains extra items such as objects and toys, may also help to reinforce the original object memory. Resulting in the preservation of an object's memory through multiple engram retrievals, and updating plasticity. In this framework, the additional reminding experience of enrichment toys might help maintain or nudge mice to infer a higher engram relevancy that is more robust against forgetting, leading to higher engram expression.

Exposure to the original stimuli facilitates memory recall

We next sought to investigate conditions in which more nuanced learning from experience may alter engram activity and subsequently switch a forgotten memory from an inaccessible state to an accessible state. Previous work from both human experimental psychology studies and rodent behavioral paradigms have shown that a brief exposure to reminder cues can aid memory recall (45–48). Here we modified our object-based paradigm to include a brief exposure to the original encoding environment and objects (**Figure 4a**). A control experiment was performed to

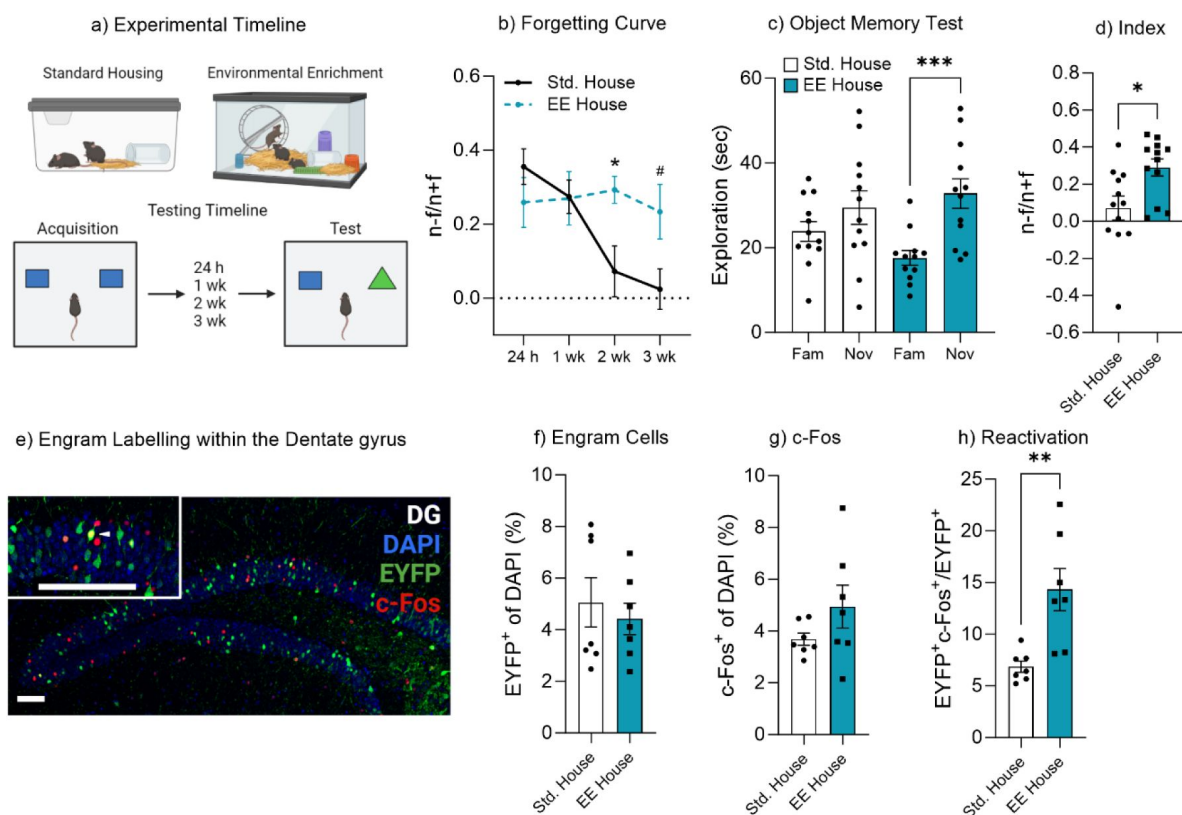


Figure 3

Environmental enrichment reduces the rate of forgetting by increasing engram activation

(a) Enrichment behavioral paradigm **(b)** Natural forgetting curve of an object memory at 24 h, 1 wk, 2 wk and 3 wk **(c)** Object memory test **(d)** Discrimination index **(e)** Engram labelling within the dentate gyrus **(f)** Engram cells **(g)** c-Fos⁺ cells **(h)** Engram reactivation. Bar graphs indicate average values in $n = 4-12$ per group (* $p < 0.05$, *** $p < 0.001$). Data graphed as means $\pm$ SEM. Scale Bar 100 μ m.

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demonstrate that a brief reminder exposure of 5 mins on its own was insufficient to induce new learning that formed a lasting memory (Supplementary Figure S4a). Mice given only a brief acquisition period of 5 mins exhibited no preference for the novel object when tested 1 hour after training, suggesting the absence of a lasting object memory (Supplementary Figure S4b & c). We therefore used the 1-hour time point for the brief reminder experiment in **Figure 4a**. An AAV₉-TRE-ChR2-eYFP virus was injected into c-fos-tTA mice to label engram cells in the dentate gyrus. During the object recall test, mice within the control group spent a similar amount of time exploring both the novel and familiar object, $t(10) = 1.62$, $p = 0.13$, 95% CI [-1.13 to 7.22], again suggesting the original object memory was forgotten (**Figure 4b**). However, mice given a brief reminder session 1 hour prior to the recall test spent significantly more time exploring the novel object compared to the familiar object, $t(4) = 4.63$, $p < 0.01$, 95% CI [7.27 to 29.00] (**Figure 4b**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference between the discrimination index of the Control and Reminder group, $t(14) = 4.57$, $p = 0.001$, 95% CI [0.19 to 0.54] (**Figure 4c**). Furthermore, the mice that underwent the reminder session and displayed intact memory recall also exhibited an increase in original engram reactivation, $t(8) = 3.13$, $p = 0.014$, 95% CI [2.66 to 17.57] (Figure d & g), while there was no difference in the engram size, $t(8) = 0.14$, $p = 0.88$, 95% CI [-1.76 to 1.55] (**Figure 4e**) or the level of neuronal activity within the dentate gyrus, $t(8) = 0.11$, $p = 0.91$, 95% CI [-1.16 to 1.05] (**Figure 4f**). We next conducted an experiment to determine if repeated object reminder sessions would also reduce forgetting. In this paradigm, mice were repeatedly exposed to the original object pair (Supplementary Figure S5a). The results indicated that repeated reminder trials spaced over several days was also sufficient to facilitate persistent object memory recall (Supplementary Figure 5b&c). Together, these data indicate that a brief reminder or subsequent object reminders over time facilitates the transition of a forgotten memory to an accessible memory via modulating engram activity.

Repeated context-only exposure to the training environment facilitates forgetting

Given that memory recall could be facilitated by optogenetic activation (**Figure 2**), maintained by enrichment (**Figure 3**), or even retrieved after forgetting by exposure to natural “reminder” cues (**Figure 4**), we wanted to test that forgetting was driven by an adaptive process which updates memory engrams according to perceptual and environmental feedback. We therefore developed an altered version of the object memory task where mice were repeatedly reintroduced to the training context in the absence of objects (**Figure 5a**). We hypothesized that the repeated exposure to the training context without objects would update the original memory engram signaling that the objects were no longer relevant to the environment. During the recall test 1 week after training control mice spent significantly more time exploring the novel object compared to the familiar object, indicating a retrievable object memory, $t(7) = 4.14$, $p = 0.004$, 95% CI [5.45 to 19.92] (**Figure 5b**). Whereas mice that had been repeatedly exposed to the training environment devoid of objects spent a similar amount of time exploring both the novel and familiar objects, $t(8) = 1.26$, $p = 0.24$, 95% CI [-5.29 to 18.18], suggesting an updated and now inaccessible engram. In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference in the discrimination index between the control and empty box exposed mice, $t(15) = 2.83$, $p = 0.012$, 95% CI [-0.55 to -0.077] (**Figure 5c**). As the mice exhibit relatively high exploration of both the novel and familiar objects. An alternative explanation could be that the mice have not truly forgotten the familiar object, but rather as the mouse has not seen the familiar object in the previous 6 context only sessions, its reappearance makes it somewhat novel again. Therefore, this change in the object’s reappearance might trigger the animal’s curiosity, and in turn drives exploration by the animal. Indeed, cross groups comparison reveals mice in the context-only group showed increased object exploration, $F(3, 30) = 11.30$, $p = 0.005$. A post hoc Tukey test showed that context only mice explore the novel object, $p = 0.028$, and object familiar, $p = 0.001$ more than control mice. In order to confirm that the

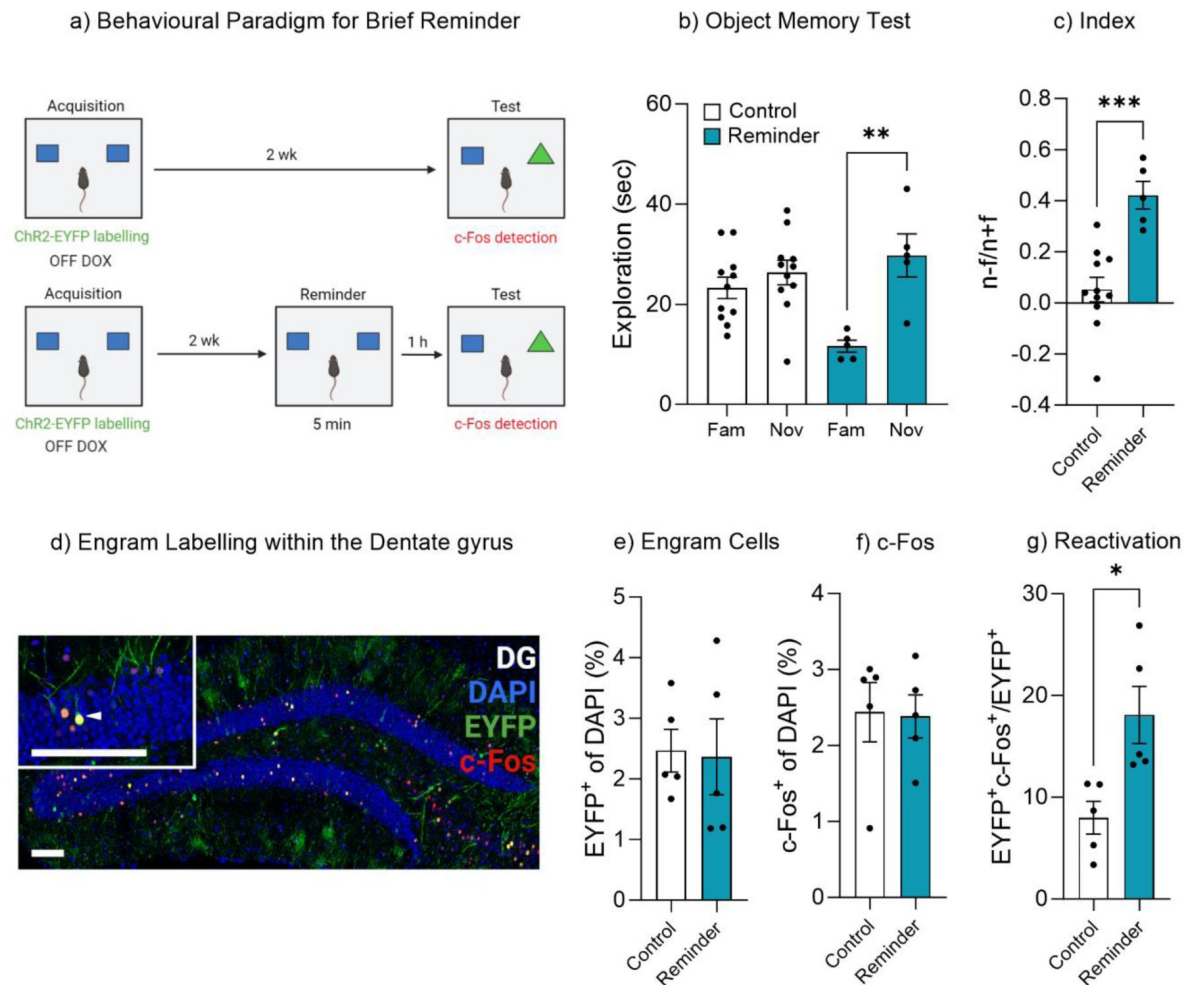


Figure 4

Exposure to the original stimuli facilitates memory recall

(a) Behavioral timeline for a brief reminder exposure **(b)** Object memory test **(c)** Discrimination index **(d)** Engram labelling within the dentate gyrus **(e)** Engram cells **(f)** c-Fos⁺ cells **(g)** Engram reactivation. Bar graphs indicate average values in $n = 5-11$ per group (* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$). Data graphed as means $\pm$ SEM. Scale Bars 100 μ m.

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experience during the repeated exposure was indeed updating the original engram resulting in accelerated forgetting, we ran a second cohort of c-fos-tTA mice which had been implanted with an optogenetic fiber and expressed the inhibitory opsin ArchT (**Figure 5d**). Following acquisition training the engram was optogenetically silenced during the repeated context-only exposures (**Figure 5d**). During the recall test, the No Light control group spent a similar amount of time exploring both the novel and familiar objects, $t(9) = 1.36$, $p = 0.20$, 95% CI [-2.04 to 8.27] (**Figure 5e**), while the Light-induced inhibition mice, in which the engram was silenced, spent significantly more time exploring the novel object compared to the familiar object, $t(11) = 3.60$, $p = 0.004$, 95% CI [4.70 to 19.46] (**Figure 5e**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference in discrimination index between No light control and Light-induced inhibition group, $t(20) = 2.12$, $p = 0.046$, 95% CI [0.003 to 0.33] (**Figure 5f**). Combined these data suggest that experience following the encoding of an object memory can update the original memory engram accelerating the rate at which the information is forgotten. Further, this adaptive process requires the activity of the original engram ensemble.

Rac1 mediates forgetting and engram reactivation

Rac1 has previously been shown to be involved in memory encoding and consolidation as well as active forgetting of contextual and social memory (29, 49–54). Rac1 has also recently been shown to be involved in memory retrieval (21). We therefore sought to investigate the role of Rac1 signaling in mediating natural forgetting within the object memory paradigm employed in this study. An AAV_g-TRE- ChR2-eYFP virus was injected into c-fos-tTA mice to label engram cells within the dentate gyrus. Immediately after acquisition training, mice were administered a Rac1 inhibitor (**Figure 6a**). Mice that were administered saline displayed typical forgetting at 2 weeks, as indicated by similar object exploration between the novel and familiar objects, $t(8) = 1.07$, $p = 0.31$, 95% CI [-2.61 to 7.20] (**Figure 6b**), while mice treated with a Rac1 inhibitor displayed intact memory recall, spending significantly more time with the novel object compared to the familiar object, $t(8) = 4.27$, $p = 0.002$, 95% CI [8.10 to 27.10] (**Figure 6b**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference in discrimination index between the Saline and Rac1 inhibitor group, $t(17) = 3.53$, $p = 0.002$, 95% CI [0.12 to 0.49] (**Figure 6c**). Furthermore, there was no difference in engram size between the two groups, $t(10) = 1.09$, $p = 0.29$, 95% CI [-0.45 to 1.33] (**Figure 6d**) or the level of neuronal activity within the dentate gyrus, $t(10) = 0.86$, $p = 0.40$, 95% CI [-1.31 to 0.57] (**Figure 6e**). However, there was a significant decrease in engram reactivation, $t(10) = 5.87$, $p < 0.001$, 95% CI [3.39 to 7.54] (**Figure 6f & g**). We also conducted a control experiment to demonstrate that Rac1 inhibition did not enhance memory recall at 24 h (Supplementary Figure S6a-f).

Given Rac1 inhibition prevented forgetting, it stands to reason that by activating Rac1 forgetting maybe accelerated. We next injected a Rac1 activator into a second cohort of c-fos-tTA mice following memory encoding and again daily for 6 days (**Figure 6h**). During the object recall test, mice within the control group spent more time exploring the novel object compared to the familiar object, suggesting intact memory recall, $t(8) = 4.98$, $p = 0.001$, 95% CI [9.11 to 24.78] (**Figure 6i**). Whereas mice with Rac1 activator spent a similar amount of time exploring both the novel and familiar object, suggesting the original object memory was forgotten, $t(7) = 1.19$, $p = 0.27$, 95% CI [-2.33 to 7.05] (**Figure 6i**). In order to compare performance in object memory, a discrimination index was calculated. There was a significant difference between the discrimination index of the Control and Rac1 activator group, $t(15) = 3.68$, $p = 0.002$, 95% CI [-0.39 to -0.10] (**Figure 6j**). Furthermore, there was no difference in the engram size, $t(12) = 0.29$, $p = 0.77$, 95% CI [-1.35 to 1.77] (**Figure 6k**) or the level of neuronal activity within the dentate gyrus, $t(12) = 0.23$, $p = 0.81$, 95% CI [-1.09 to 1.36] (**Figure 6l**). However, there was a decrease in engram reactivation, $t(12) = 3.15$, $p = 0.008$, 95% CI [-13.98 to -2.56] (**Figure 6m & n**). We also conducted a control experiment which indicated a single dose of Rac1 activator was sufficient to promote

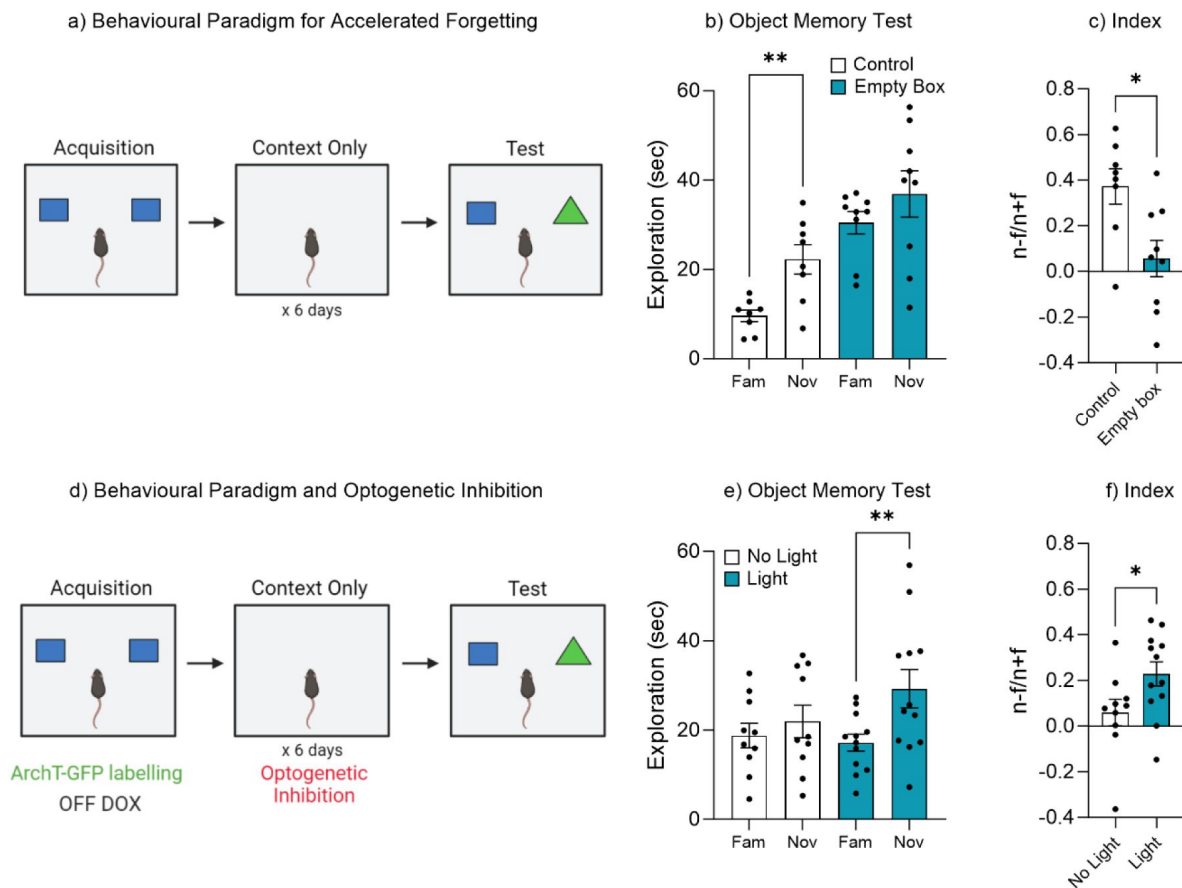


Figure 5

Repeated exposure to the training environment facilitates forgetting

(a) Behavioral paradigm **(b)** Object memory test **(c)** Discrimination index **(d)** Behavioral paradigm for optogenetic inhibition during repeated context exposure **(e)** Object memory test **(f)** Discrimination index. Bar graphs indicate average values in n = 8-12 per group (*p < 0.05, **p < 0.01). Data graphed as means ± SEM.

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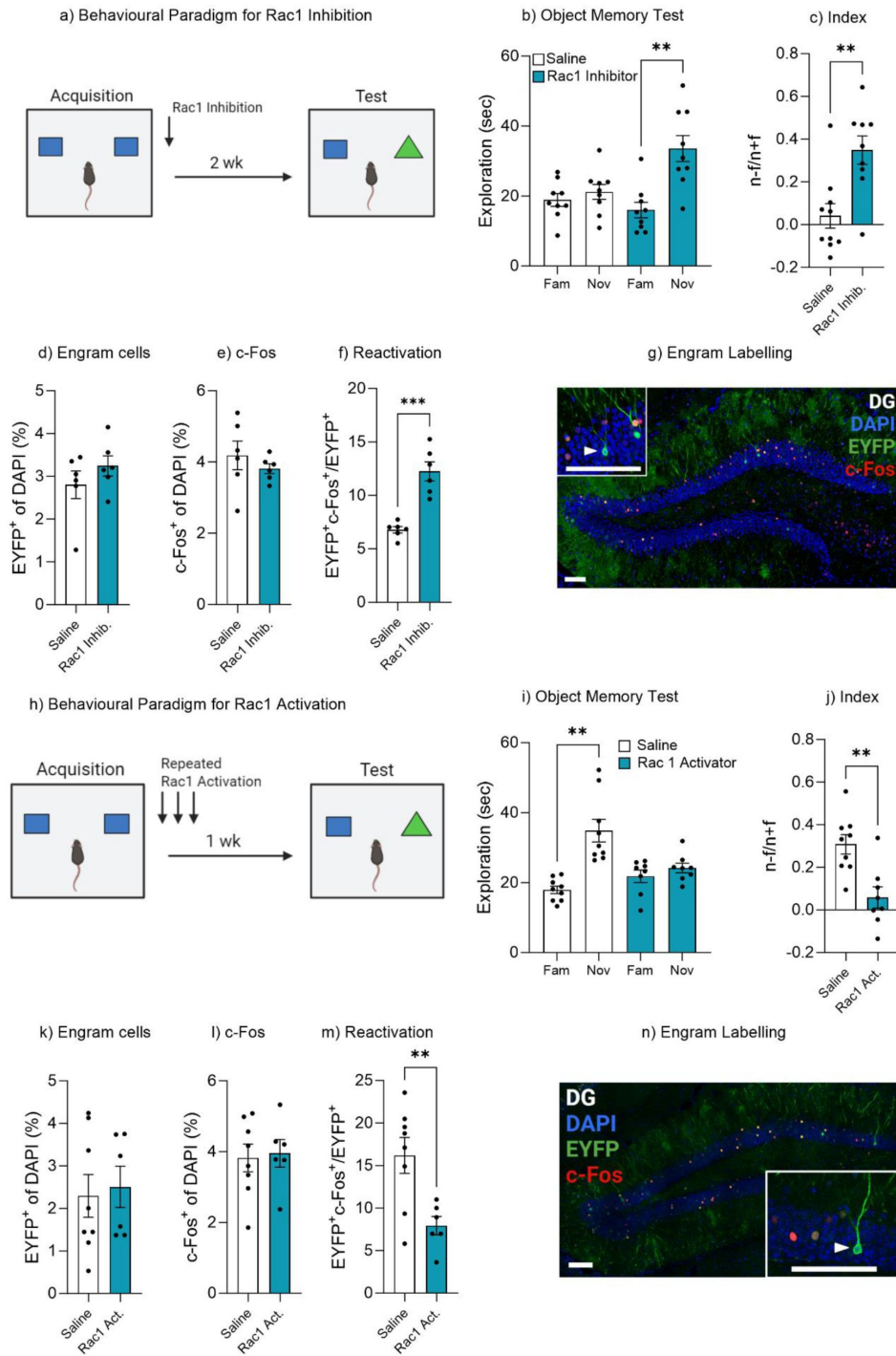


Figure 6

Rac1 modulates forgetting through engram activation

(a) Experimental timeline and drug administration for Rac1 inhibition **(b)** Object memory test **(c)** Discrimination index **(d)** Engram cells **(e)** c-Fos⁺ cells **(f)** Engram reactivation **(h)** Engram labelling within the dentate gyrus **(i)** Experimental timeline and drug administration for Rac1 inhibition **(j)** Object memory test **(k)** Discrimination index **(l)** Engram cells **(m)** c-Fos⁺ cells **(n)** Engram reactivation. Bar graphs indicate average values in $n = 8-9$ per group (** $p < 0.01$, *** $p < 0.001$). Data graphed as means $\pm$ SEM. Scale Bar 100 μ m.

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forgetting (Supplementary Figure S7a-f). Combined these data demonstrate the Rac1 signaling is involved in memory encoding and consolidation and may function as a plasticity enabling molecule to mediate retrievability. Rac1 may therefore be a signaling mechanism involved in adaptive forgetting.

Learning Model Explains Forgetting Dynamics

Utilizing a mouse model of object memory, we have investigated the conditions in which a memory could be preserved, retrieved, or forgotten. The data indicate that forgetting is a dynamic process that can be explained as a gradient of engram activity, where the level of memory expression is experience-dependent and influenced by the animal's rate at which it learns from new experiences. Based on these findings, we sought to utilize the Rescorla-Wagner model to formalize the idea that forgetting is a form of learning and to integrate the empirical findings into a coherent, mechanistic, and quantitative framework (8, 55, 56). Our proposed Rescorla-Wagner model dynamically updates object-context associations in response to prediction errors (Figure 7a). Accordingly, engrams that are subjectively less relevant for adaptive behavior (representing weaker object-context associations) are more likely to be forgotten (56). As illustrated in Figure 7b, "engram strength" modeling the current object-context association is a function of the prediction error (difference between experienced objects and expected engram strength) and the "learning rate" governing the influence of prediction errors on learning and forgetting. Positive prediction errors strengthen the engram, and negative errors yield weaker object-context associations (Figure 7d). Therefore, faster forgetting is driven by higher rates at which mice learn from negative prediction errors. In summary, this mechanism could help the animal prioritize relevant information in memory (Figure 7c). The animal predicts what it will encounter in its environment (e.g., what objects) and adjusts these predictions as a function of its experiences (e.g., the presence or absence of objects). Within this conceptual framework, the animal learns what features of the environment are currently relevant and thus important for remembering. In order to further support a link between our empirical data and computational modeling, we demonstrated that the modulation of engram cells within the dentate gyrus can indeed regulate object-context associations (see Supplementary Figure 12a-f).

Supporting the idea that forgetting can be cast as an adaptive learning process, model simulations showed similar forgetting curves as mice in our experiment. We first applied the Rescorla-Wagner model to exploration behavior of mice in the standard housing condition that included data across four test time points (Figure 3). To obtain an empirical estimate of the subjects' learning rate governing the speed of forgetting (i.e., how quickly the object-context association degrades over time due to inhibitory object-context associations), we fitted the model to the data using a maximum-likelihood approach (see Methods). Here, the learning rate indicating the weight given to negative prediction errors for updating engram strength was $\alpha^- = 0.07$, explaining the increase in forgetting across time (see Supplementary Table 1 for the other parameter estimates). Simulated forgetting curves that were based on the estimated parameters showed a close similarity to mice in the standard housing condition (Figure 7e; see Supplementary Figure S8 and S9 for a direct comparison of model predictions and empirical data).

The model also offers an explanation for the key results of the different experimental interventions. Experiencing objects during environmental enrichment that resemble the familiar objects in the memory test might nudge subjects to infer a higher engram relevancy that is more robust against prediction errors leading to higher engram expression (Figure 8a & 8b, d, f). Supporting the perspective of forgetting as learning, the estimated learning rate in response to negative errors driving forgetting $\alpha^- = 0$. Thereby, negative prediction errors did not induce forgetting so that memory performance was constant across 4 weeks (see also Supplementary Figure S8). From a computational perspective, a low learning rate may also explain the high memory performance of the Rac1-inhibition group. For this group, model fitting revealed a similar learning rate to the enrichment group (learning rate for negative prediction errors $\alpha^- = 0.01$), suggesting that the inhibition of Rac1 made subjects largely ignore negative prediction errors that

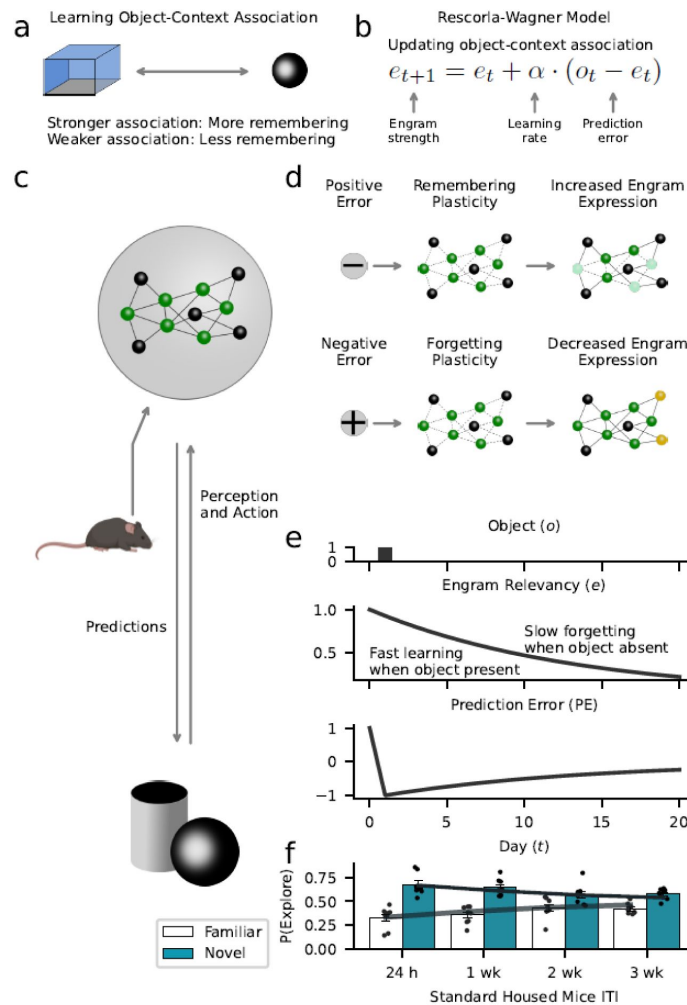


Figure 7

Forgetting as adaptive learning.

Our model assumes that animals create and update memory engrams to flexibly adjust their behavior to their environment. **(a)** We used a Rescorla-Wagner model that learns object-context associations in the object-based memory task. The key assumption is that stronger associations lead to higher memory performance, while weaker associations lead to more forgetting. **(b)** The Rescorla-Wagner model updated the strength of object-context associations ("engram strength") as a function of the prediction error (difference between experienced object and engram strength) and the learning rates governing the influence of the prediction error. **(c)** Based on learned representations, animals constantly predict what happens in the environment (e.g., the occurrence of objects), and if predictions are violated (prediction errors), engrams are updated to improve the accuracy of future predictions. Here, established engram cells are shown in green; non-engram cells in gray. **(d)** Positive prediction errors signaling the occurrence of an unexpected event (e.g., new object) induce a learning process that increases the probability of remembering. This might rely on the recruitment of new engram cells (shown in yellow). In contrast, negative prediction errors signaling the absence of an expected event (e.g., predicted object did not appear) induce forgetting. This might rely on "forgetting" plasticity reducing access to engrams (light green cells). **(e)** Our model formalizes this perspective based on the notion of "engram relevancy", i.e., the strength of the object-context association. Higher engram relevancy makes it more likely that an engram is behaviorally expressed, e.g., through exploration behavior. The presentation of a novel object (upper panel) leads to a high engram relevancy (middle panel) in response to a positive prediction error (lower panel). The absence of an expected object decreases engram relevancy through negative prediction errors. **(f)** Model simulations corroborate the behavioral effects of our data (Figure 3a). Gray lines and bars show the average exploration probability for the familiar and novel object according to the model; markers show simulated mice.

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would normally drive forgetting, thereby avoiding the formation of inhibitory object-context associations. In contrast to reduced learning preventing forgetting due to Rac1 inhibition, Rac1 activation might speed forgetting through a higher learning rate. Similarly, we found accelerated forgetting following repeated exposure to the context-only training environment. Model simulations (**Figure 8c**) assuming a higher learning rate and, therefore, reduced engram expression, reproduced these findings (**Figure 5** & 6).

A crucial assumption of our model is that seemingly forgotten engrams are not necessarily lost but rather inaccessible when the inferred object relevance is low. This is consistent with the experimental results above, where we presented evidence for the reinstatement of memory representations based on real or artificial reminder cues (**Figure 2** and **4**). Here, we modeled memory reinstatement through reminder cues by inducing an artificial, positive prediction error in response to the reminder that boosted the inferred object relevance (strengthening object-context associations). Similar to the experimental data, the modeled reminder cues increased engram expression and yielded reduced forgetting rates (**Figure 8e**).

Finally, the model fits the data better than a baseline model predicting random forgetting rates, suggesting that it describes the data accurately and above chance-level (Supplementary Figure S10). We also performed a parameter-recovery analysis suggesting that parameters can sufficiently be estimated despite the limited data for model fitting (Supplementary Figure S11). Our modeling results support the perspective that natural forgetting is a form of adaptive learning that alters engram accessibility in response to environmental feedback. Accordingly, environmental, optogenetic, and pharmacological manipulations might change the learning rate, suggesting that miscalibrated learning rates could indeed give rise to pathological forgetting. In summary, our Rescorla-Wagner model provides a mechanistic interpretation of our own data, a parsimonious explanation for the dynamics of natural forgetting, and could inform future studies on pathological forgetting.

Discussion

For an organism to adapt to its environment, there must be processes by which learned information can be updated (1, 8, 57). Thus forgetting, or the letting go of information when it is appropriate, may be a key characteristic underlying learning and memory. It is possible that forgetting may be an adaptive process that is driven by feedback from the environment (8). This experience-dependent process could help an animal to prioritize relevant information in memory. Previous work has demonstrated that hippocampal engram cells are differently modulated by experience (21, 58).

Here, we investigated the conditions in which a memory could be preserved, retrieved, or forgotten through changes in experience and learning. Altered memory recall was associated with corresponding changes in engram activation, indicating that altered memory engram expression is a reversible feature of natural forgetting. Under baseline conditions mice were able to recall an object memory 24-hour post-training but exhibited impaired recall after two weeks, indicating the initial memory was forgotten. The results demonstrated that engram activity correlated with the rate of forgetting. Moreover, these reactivated cells were shown to be a functional component of the initially encoded memory as the direct modulation of engram activity via optogenetic stimulation or inhibition either facilitated or prevented the recall of the object memory. These findings are in support of previous work which demonstrated hippocampal engram cells could be differently modulated by experience (58). These findings are also in agreement with the current literature which has shown that engram activation is associated with the level of memory recall (1, 47, 58–61). In addition, they also support a role for the hippocampus as a fundamental structure for object memory (62, 63). It is important to consider that forgetting may also occur when an experience or an engram is not properly encoded during the learning

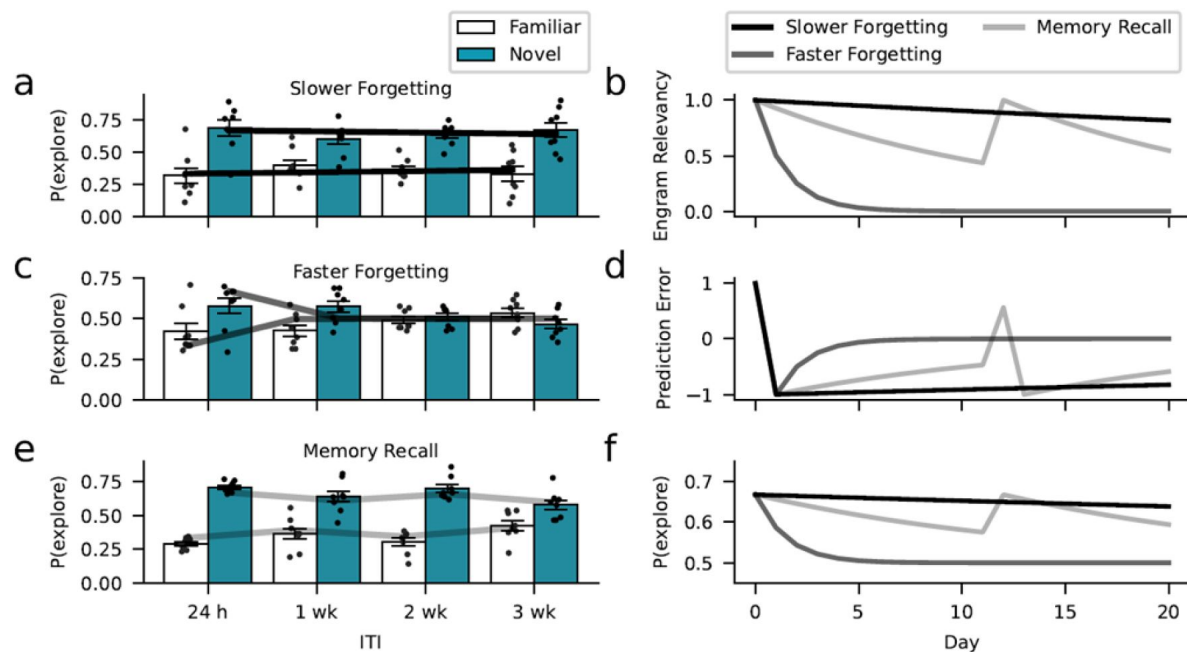


Figure 8

Learning model captures the dynamics of forgetting.

Environmental, optogenetic, and pharmacological manipulations might modulate the speed of forgetting by altering key parameters of our model. Simulations with different learning-rate parameters explain the forgetting dynamics of the different experimental conditions. **(a)** The enrichment and Rac1-inhibition conditions were successfully captured using a low learning rate (0.01, similar to the empirical estimates). **(c)** In contrast, assuming a larger learning rate (0.5), we could capture faster forgetting as observed in the Rac1-activator and context-only conditions. **(e)** Moreover, improved memory performance after reminder cues can be explained by assuming that these interventions induce a positive prediction error boosting object relevancy. Here, we assumed a learning rate of 0.07 (based on the empirical estimate). **(b)** Development of engram relevancy and **(d)** prediction errors across conditions. **(f)** Probability of exploring the novel object plotted separately for each condition.

phase, or not properly consolidated (64–66). In the present study forgetting was also associated with a reduction in spine density relative to a 24 h recall timepoint. This observation suggested that the memory was properly consolidated one day post training (67). We also observed intact memory recall at 1 week post training, which further supports the idea that the initial object memory was indeed properly encoded and consolidated. Forgetting may encompass a broad spectrum of types of memory loss or apparent memory loss, some of which may involve the actual loss or damage of the engram, whilst others may reflect suppressed engram expression or simply altered conditions of engram activation. Here we investigated spine density within engram cells following recall at 24 hour or 2 weeks. Future work could investigate if spine density or spine type is reduced in reactivated and non-reactivated engram cells within the ensemble and across engram and non-engram cells following different learning and forgetting conditions. This avenue of research could provide evidence to whether there are functionally distinct aspects of engram cells that are altered following forgetting (39, 68).

Our data showing memory retrieval following optogenetic reactivation demonstrate that the forgotten engram is indeed still intact, otherwise activation of those cells would no longer trigger memory recall. Interestingly, previous work has shown that an engram remains intact even when memory consolidation is disrupted by protein synthesis inhibition or transgenic induction of Alzheimer's disease (15, 17, 69). In those studies, the engram persists in a latent state despite lower dendritic spine density (15, 17, 69). Moreover, optogenetic stimulation of amnesic engram cells was also sufficient to restore access to the amnesic memory (15, 17, 69). Taken together, these studies suggest the engram remains intact, whilst not properly retrievable (15, 17, 69).

We showed that the rate of forgetting, and the level of engram activation could be modulated through enrichment. This enhanced memory recall was associated with increased engram reactivation. These findings are supported by earlier work, which showed exposure to an enriched environment prior to learning improved engram reactivation and rescued memory recall in a mouse model of Fragile X syndrome (70). Previous literature has also shown the benefits of environmental enrichment in alleviating other pathological forms of memory impairment, such as Alzheimer's disease (71–73). The effects of enrichment on learning and neural plasticity are well established and it is likely the contributing factor to the enhanced memory recall (40, 42, 43, 72, 74, 75). A key component of enrichment is the increased perceptual and tactile feedback provided by extra items such as toys, objects and running wheels, within the home cage for the mice to interact with and explore (76, 77). The added opportunity to experience these extra cage items, at an algorithmic level, may help to reinforce the original object memory, resulting in the preservation of an object's memory through multiple engram retrievals, and updating plasticity. Under this framework, the additional reminding experience of enrichment toys might help maintain or nudge mice to infer a higher engram relevancy that is more robust against forgetting, leading to higher engram expression. However, this line of investigation requires further empirical evidence to separate the impact of similar percepts on object memory recall from the broadly cognitive enhancing effects of environmental enrichment. Based on the computational analyses using the Rescorla-Wagner model we suggest that this form of natural forgetting is governed by the strength of the object- context associations, referred to here as engram relevancy. At the algorithmic level, stronger associations signal that the memory or engram representation is important or relevant and should not be forgotten. Accordingly, due to the experiences with extra objects in the enriched environment, mice might learn that objects are common in their environment and potentially relevant in the future. Thus, the object-context association is strong, preventing forgetting. It is important to emphasize that this process of updating is not necessarily separate from enrichment-induced plasticity at an implementational level, but part of the learning experience within an environment containing multiple objects. The enrichment or, more generally, experience, may therefore enhance memory through the modification of activity of specific engram ensembles. The idea of enrichment facilitating memory updating is consistent with the results obtained by the reminder experiments and further

supported by our analysis with the Rescorla-Wagner computational model, where experience updates the accessibility of existing memories, possibly through reactivation of the original engram ensemble.

We validated the effectiveness of the enrichment paradigm to enhance neural plasticity by measuring adult hippocampal neurogenesis. The hippocampus has been identified as one of the only regions where postnatal neurogenesis continues throughout life (75 [↗](#)). Levels of adult hippocampal neurogenesis do not remain constant and can be altered by experience (42 [↗](#)–44 [↗](#), 58 [↗](#)). In addition, adult born neurons have been also shown to contribute to the process of forgetting (74 [↗](#), 78 [↗](#), 79 [↗](#)). Although the contribution of adult born neurons to cognition and the memory engram is not fully understood (80 [↗](#), 81 [↗](#)). Mishra et al, showed that immature neurons were actively recruited into the engram following a hippocampal-dependent task (39 [↗](#)). Increasing the level of neurogenesis rescued memory deficits by restoring engram activity, in a mouse model of Alzheimer's disease (39 [↗](#)). Specifically, augmenting neurogenesis rescued the deficits in spine density in both immature and mature engram neurons (39 [↗](#)). Whether neurogenesis alters spine density differentially for reactivated or non- reactivation engrams cells remains to be investigated (39 [↗](#), 68 [↗](#)). This avenue of research would help to illuminate the morphological changes following forgetting and provide evidence if there is a functionally distinct aspect of engram cells that is altered in forgetting (39 [↗](#), 68 [↗](#)). Future work could utilize a different engram preparation based on a stable genetic labelling strategy could investigate the contribution of new-born neurons to the engram ensemble.

Our results showed that forgetting could be reversed following a brief reminder experience, which was associated with a corresponding increase in engram activity. Previous work from both human and rodent behavioral studies have shown that a brief exposure to reminder cues can aid memory recall (45 [↗](#)–48 [↗](#)). We further demonstrated that forgetting could be accelerated following repeated exposure to the training environment, therefore signaling the updated irrelevance of the object memory. This updating through experience was impaired when the original memory was inhibited. In line with the Rescorla-Wagner computational model, this suggests that in the present study, when mice are repeatedly exposed to the training context, the association between the context and the objects is reduced, resulting in a disrupted ability of the context to activate the object engram. In fact, earlier work by Bouton and colleagues has demonstrated that contextual changes from the passage of time can lead to retrieval failure (48 [↗](#), 82 [↗](#), 83 [↗](#)). Future work addressing how changes in internal and external contextual cues alter memory expression will expand our understanding forgetting processes of multiple memory engrams, and their interactions.

Our findings indicate that the degree of learning and memory specificity corresponded with engram activity (84 [↗](#)). The learning rate may therefore offer insight into the mechanism of forgetting. Here the learning rate was altered following environmental enrichment as well as following repeated context exposure, which in turn affected the rate of forgetting. Together, this suggests the possibility of an 'optimal' learning rate that yields environmentally adaptive, natural forgetting and that learning rates that are too high may be linked to amnesia and learning rates that are too low to some sort of hypermnesia.

These findings raise an important question regarding the interpretation of forgetting (85 [↗](#), 86 [↗](#)). Previous work has emphasized retrieval deficits as a key characteristic of memory impairment, supporting the idea that memory recall or accessibility may be driven by learning through perceptual feedback from the environment (7 [↗](#), 8 [↗](#), 14 [↗](#)–18 [↗](#)). Within our behavioral paradigm, a lack of memory expression would still constitute forgetting due to the loss of learned behavioral response in the presence of natural retrieval cues. The changes in memory expression may therefore underlie the adaptive nature of forgetting. Here we studied natural forgetting, and our data showing memory retrieval following optogenetic reactivation demonstrates that the engram endures even when not expressed.

The Rescorla-Wagner model can be used to describe forgetting as a process of learning, where experience alters the degree of association between retrieval cues and memory (56, 87). Based upon this idea, we proposed a parsimonious computational model that integrates our findings into a cognitively plausible framework in which memories more relevant for adaptive behavior are more likely to be accessible than memories representing irrelevant, outdated information. As such, our modeling analyses suggest that the different experimental manipulations altered the learning rate determining how rapidly engrams switch from accessible to inaccessible states in response to environmental feedback, such as the presence or absence of objects. This model suggests that an emergent property underlying forgetting is the prediction error. Within this framework, forgetting is a form of learning whereby experience drives changes in the learning rate of an organism, and the expectation of one's environment can alter the efficiency of recall through forgetting plasticity (8). It therefore follows that prediction errors may determine whether an engram is strengthened, leading to higher engram expression, or weakened, leading to lower engram expression (**Figure 7d**). Positive prediction errors indicate that an unexpected event has taken place (e.g., some unexpected object is present). We assume that positive errors induce plasticity processes that alter the engram and increase the likelihood of engram expression. In contrast, negative errors indicate that something that was expected has not taken place (e.g., object absent). Consequently, negative errors induce forgetting plasticity, and yield decreased engram expression.

Dopamine is known to play a role in prediction error and learning rates (88, 89). Moreover, dopamine has been shown to differently regulate Rac1 to modulate behavioral plasticity (29, 90). Here, we showed that the inhibition of Rac1 prevented forgetting, while its activation following memory encoding accelerated the rate of forgetting. This finding is in agreement with previous work where Rac1 impaired memory recall by driving forgetting-induced plasticity (29, 49–52, 54). Rac1 has been shown to function during the memory encoding and consolidation phases, and thereby to contribute to active forgetting (29, 49–52). Recently, Rac1 has also been shown to be involved in memory retrieval (21). Lei et al., demonstrated that a social reward enabled latent engrams to switch to active engrams via suppression of Rac1 activity in CA1 neurons of the hippocampus (21). It was further demonstrated that this effect was bidirectional as a social stressor switched active memory engrams back to latent engrams (21). This social stress-induced forgetting was modulated through the activation of Rac1 signaling (21). Overall, the activation of Rac1 has been shown to accelerate the rate of forgetting, while its inhibition prevented forgetting (50). Here, our work advanced this knowledge by demonstrating that Rac1-induced changes in memory recall alter engram reactivation. These works support our data suggesting that Rac1 is involved in altering the rate of forgetting at multiple time intervals after learning, not just immediately after learning. Our broader view is that Rac1 may function like a ubiquitous plasticity enabling molecule that is involved in memory encoding, consolidation, forgetting, and retrieval, depending on the experimental conditions and behavioral settings (8, 29).

Our Rescorla-Wagner model could be a starting point for more comprehensive models that account for forgetting across different experimental paradigms. Recent work suggests that retroactive interference emerges from the interplay of multiple engrams competing for expression (18). Further, there is the possibility of an interplay or connection between memory relevance and context within the process of extinction (91). Lacagnina et al., demonstrated that extinction training suppressed the reactivation of a fear engram, while activating a second putative extinction ensemble (60). In another study, these extinction engram cells and reward cells were shown to be functionally interchangeable (92). Moreover, in a study conducted by Lay et al., the balance between extinction and acquisition was disrupted by inhibiting the extinction recruited neurons in the BLA and central nucleus of the amygdala (93). These results suggested that decision making after extinction was governed by a balance between acquisition and extinction-specific ensembles (93).

Future models could explicitly incorporate multiple engrams and their competition, explaining a broader range of forgetting effects. Several approaches to modeling extinction (94 [↗](#)–96 [↗](#)), memory interference (97 [↗](#)), or the creation and updating of motor memories (98 [↗](#)) that more explicitly assumed multiple memory representations could inform such a generalized model of natural forgetting. Furthermore, future work could also include other variables in addition to prediction errors to the model framework to better capture the dynamic conditions of memory recall. These may include boundary conditions of destabilization and reconsolidation, the salience or strength of the memory as well as the timing of retrieval cues or updating experience. Future models could focus on understanding the specific boundary conditions in which a memory becomes retrievable and the degree to which it is sufficiently destabilized and liable to updating and forgetting. The role of perceptual learning in memory retrieval and forgetting may also be an avenue for future investigation. Understanding how experience alters object familiarity versus object retrieval and its impact on learning would also help to develop better models of object memory and forgetting.

Learning and memory allow humans and animals to maintain and update predictions about future outcomes. While having access to a large number of memories is adaptive in vast environments, it is equally important to prioritize the accessibility of the most relevant memories and to forget outdated information. Suppressing stored information through natural forgetting might therefore promote adaptive behavior. However, forgetting too much (e.g., amnesia) or too little (hypermnesia) under pathological conditions is maladaptive. Here, we utilized a mouse model of object memory and investigated the conditions in which a memory could be preserved, retrieved, or forgotten. Moreover, through pharmacological and behavioral interventions, we successfully prevented or accelerated forgetting. Based on these findings and our computational model in which memories subjectively less relevant to adaptive behavior are more likely to be forgotten we conclude that natural forgetting may be considered a form of adaptive learning where engrams that are subjectively less relevant for behavior are more likely to be forgotten.

Methods

Animals

All wild-type behavior was conducted with male C57BL/6J mice aged 7–12 weeks, bred in-house from Charles River breeding pairs. c-fos-tTA mice were generated in-house by breeding TetTag mice with C57BL/6J mice and selecting offspring carrying the c-fos-tTA (38 [↗](#)). All mice used for the experiments were male between 8–12 weeks old at the time of surgery and had been raised on food containing 40 mg/kg doxycycline for at least five days prior to surgery. c-fos-tTA mice remained on doxycycline diet for the duration of the experiments. Doxycycline was removed from the diet for 36 hours prior to object acquisition to allow for engram labelling, following completion of the object acquisition period c-fos-tTA mice were placed back on the doxycycline diet.

All mice were grouped housed in standard housing conditions, with temperature $22 \pm 1^\circ\text{C}$, relative humidity 50% and a 12:12 hour light-dark cycle (lights on 0730h) and had ad libitum access to food and water. All experiments were conducted in accordance with the European Directive 2010/63/EU and under an authorization issued by the Health Products Regulatory Authority Ireland and approved by the Animal Ethics Committee of Trinity College Dublin.

Virus-mediated gene expression

The recombinant AAV vectors used for viral production were AAV-TRE-ChR2-eYFP and AAV-TRE-ArchT-GFP. Plasmids were serotyped with AAV9 coat proteins and packaged commercially by Vigene. The recombinant AAV vectors were injected with viral titers were 1×10^{13} genome copy (GC) ml⁻¹ for AAV₉-TRE-ChR2-eYFP and AAV₉-TRE-ArchT-GFP.

Stereotactic viral injection and Optic Fiber implantation

Mice were anaesthetized with Avertin 500 mg/kg and placed into a stereotactic frame (World Precision Instruments). Bilateral craniotomies were performed using 0.5 mm diameter drill and the virus was injected using a 10 µl Hamilton microsyringe filled with mineral oil. A microsyringe pump and its controller were used to maintain the speed of the injection. The needle was slowly lowered to the target site and remained for 5 min before the beginning of the injection. For engram labeling and activation studies (AAV₉-TRE-ChR2-eYFP) or inhibition studies (AAV₉-TRE-ArchT-GFP) (Volume; 300nl) virus was injected bilaterally into the DG using the coordinates AP: -2 mm, ML: ± 1.35 mm, DV: -2 mm relative to Bregma at a rate of 60nl/min. Followed by a 10 min diffusion period. For optogenetic experiments, a Doric optical fiber (200 µm core diameter; Doric Lenses) was implanted above the injection site (-2 mm AP; ± 1.35 mm ML; -1.85 mm DV). A layer of adhesive cement (C&B Metabond) was applied followed by dental cement to secure the optic fiber implant to the skull. Mice were given 1.5 mg/kg meloxicam as analgesic. Animals were allowed two weeks to recover before behavioral testing. For optogenetic experiments. Control No Light mice underwent the identical surgery procedure with virus and optic fiber implantation. However, no light was delivered to excite or inhibit the respective opsin during the experiment.

Engram labelling strategy

The AAV₉-TRE-ChR2-eYFP virus was injected into the dentate gyrus of *c-fos*-tTA mice, under the control of a *c-fos* promoter (37, 38). The immediate early gene *c-fos* becomes upregulated following neural activity and as such the *c-fos*-tTA transgene selectively expresses tTA in active cells (36, 99). The activity-dependent tTA induces expression of ChR2-eYFP. In order to restrict activity-dependent labelling to a single learning experience, mice were maintained on diet containing doxycycline (DOX).

Object Recognition

The novel object recognition task was based on the protocol previously described by Bevins and Besheer (2006). Mice were first habituated to the testing arena (30cm L x 20cm W x 30cm H) for a 10-minute exploration period on two consecutive days. On day three, two identical objects were positioned on adjacent sides of the arena and each animal was introduced for a 10-minute exploration period. Mice were then placed directly back into their home cages. After a 24-hour or two-week inter-trial interval, one familiar object was replaced with a novel object, and each animal was introduced for a five-minute exploration period. Objects and locations were counterbalanced across groups. Object exploration was defined when the animal's nose came in contact with the object. The testing arena and objects were cleaned with a disinfectant, TriGene, between each animal and training session. Video recordings were made to allow for manual scoring of object exploration. The object discrimination index was calculated as the time spent exploring the novel object minus the time spent exploring the familiar object divided by the total time spent exploring both objects (novel – familiar / novel + familiar).

Environmental Enrichment

Mice were placed in the enriched housing for three weeks prior to behavioral testing and remained in the housing for the duration of the experiment. The enriched environment consisted of a larger home cage (50cm L x 20cm W x 40cm H). The cage was included a running wheel, tunnels, extra nesting material as well as an assortment of Lego bricks. The configuration of tunnels, Lego bricks and housing enrichment changed each week.

Pharmacological modification of Rac1

Rac1 activator (CN04) was reconstituted and prepared in saline at 2ug/mL solution. CN04 was administered via intraperitoneal injection at a dose of 0.0125 mg/kg immediately following completion of the acquisition training. Mice were administered CN04 once per day for the following 5 days, for a total of 6 injections. Rac1 inhibitor (Ehop016) was reconstituted and prepared in PBS with 1% Tween-80, 1% DMSO and 30% PEG. Ehop016 was administered via intraperitoneal injection at a dose of 20 mg/kg immediately following completion of the acquisition training.

Immunohistochemistry

On completion of the behavioral testing, mice were deeply anaesthetized with sodium pentobarbital and then transcardially perfused with 4% paraformaldehyde. Brains were post-fixed for 24 hours in 4% paraformaldehyde, transferred to PBS, and stored at 4°C. Coronal sections through the DG were collected onto slides at 50 µm thickness in a 1:4 series. Coronal sections were immunostained for eYFP and c-Fos. Non-specific antibody binding was blocked using 10% normal goat serum (NGS) in a solution of PBS with 0.2% Triton-X100 and tissue sections were incubated with goat anti-eYFP (Anti-GFP chicken IgY fraction 1:1000, Invitrogen) and anti-c-Fos (Anti-c-Fos rabbit, 1:500, Synaptic System). Sections were then incubated in the appropriate AlexaFluor secondary antibody (AF488 for eYFP and AF596 for c-Fos) and then with Dapi (1:1000; Sigma) to stain nuclei. Lastly, sections were washed, mounted, and coverslipped with anti-fade mounting media (Vectashield-DAPI). Images were obtained using a Leica SP8 gated STED confocal microscope at 40x magnification.

To measure adult born neurons. Mice were administered BrdU (50mg/kg) 3 weeks before behavioral testing. For BrdU/NeuN immunohistochemistry, hippocampal sections were washed, denatured in HCl (2 N) for 45 min at 37 °C and renatured in 0.1 M sodium tetraborate. Sections were then washed in PBS and blocked in 10% normal goat serum (NGS; Sigma) diluted in 0.1% Triton-X 100 PBS to prevent non-specific binding. Sections were incubated with rat anti-BrdU antibody (1:100, Abcam) in 1% NGS diluted in 0.1% Triton-X 100 PBS, washed, and then incubated in AlexaFluor secondary antibody (AF598 for BrdU and AF88 for NeuN) and then with Dapi (1:1000; Sigma) to stain nuclei. Lastly, sections were washed, mounted, and coverslipped with anti-fade mounting media (Vectashield-DAPI). To quantify the cell counts, images were obtained using a Leica SP8 gated STED confocal microscope at 40x magnification. The number of engram cells (eYFP+ cells) and c-Fos+ cells were counted. These were divided by the total number of DAPI-positive cells to estimate the percentage of c-Fos positive cells and eYFP positive cells in the DG (both dorsal and ventral blade). To quantify the reactivation of engram cells, the number of cells positive for both eYFP and c-Fos was divided by the number of eYFP+ cells (18 [18](#)). The reactivation value for each mouse was calculated by averaging the reactivation for each slice (4 x sections per mouse) (18 [18](#)). To measure immature neurons, the total number of DCX positive cells were counted in the dentate gyrus and averaged for each section (4 x sections per mouse). Cells expressing BrdU/NeuN were manually counted through the DG. The area of the DG for each animal was obtained using ImageJ software (National Institutes of Health, USA) and data for BrdU/NeuN or DCX- positive cells were expressed as the number of cells per µm² of DG (100 [100](#)).

Morphological Analysis

To obtain engram dendritic spine density, the dentate gyrus was imaged using the Leica SP8 gated STED confocal microscope and images were collected with the Leica Application Suite X (LasX) software. Z-stacks were taken bidirectional under a 40x lens. Dendritic spine analysis was carried out using the Imaris software (Oxford Instruments, Imaris v9.5). The dendrites of eYFP-positive cells were traced using a semiautomated neurofilament tracer tool and dendritic spines were individually highlighted and manually traced with the software. The image processing feature of

Imaris was used to apply the Gaussian and Median filters to Z-stack images to remove background eYFP staining and allow for better resolution and visualisation of dendritic fragments and associated spines. Following the labelling of spines on traced dendritic fragments, parameters for spine volume, spine head volume and dendritic spine density (/10 μ m) were collected.

Statistical analyses

All data were analysed using SPSS statistical software (SPSS, Chicago, IL). Behavioral data and the number of Chr2-eYFP-positive and c-Fos cells were graphed as means $\pm$ SEM. Data were analysed by Student's t-test or ANOVA where appropriate. An alpha level of 0.05 was used as a criterion for statistical significance, and probability levels were quoted for non-significance.

Computational Modelling

We used an error-driven learning rule based on the Rescorla-Wagner model (101 [↗](#), 102 [↗](#)). The model offers a mechanistic explanation for the key behavioral results of our experiments (Figure 7 [↗](#) and 8 [↗](#)). In line with the idea of the Rescorla-Wagner model that learning is an error-driven updating process, the model computes the relevancy of an engram e_t on each day t of the experiment, indicating the strength of the current object-context association. The engram relevancy is updated as a function of the prediction error ($o_t - e_t$; where o_t denotes the object observation). The degree to which the prediction error changes the computed engram relevancy is determined by the learning rate α_t . To update the engram relevancy, the model relies on a two-speeded mechanism through which the relevancy quickly increases when an object is present (higher learning rate) and slowly decays when the object is absent (lower learning rate). In particular, the appearance of an object yields a positive prediction error and triggers the fast-updating process, while the absence of an object is associated with negative prediction errors and slow decay in the engram relevancy. Depending on the time between the acquisition phase and the retrieval test, the model predicts different levels of memory performance, where a longer time interval between acquisition and test is associated with less object recognition.

Task Model

To model mouse behavior in the object-based memory task, we first formulated a model of the experimental paradigm.

- T : = 21 denotes the maximum number of days, which are indexed as $t = 0, 1, \dots, T$.
- $x \in \{0, 1\}$ denotes the objects of the task. $x = 0$ refers to the novel object; $x = 1$ refers to the acquisition object that is presented together with the novel object.
- $O \in \{0, 1\}$ indicates whether an object is presented. When the acquisition (and therefore familiar) object $x = 1$ is presented in the acquisition phase, $o_t^{x=1} = 1$. Moreover, $o_t^{x=1} = 1$ when the retrieval test takes place, which depending on the task condition is after one day, one week, two weeks or three weeks. In all other cases, $o_t^{x=1} = 0$.

Learning Model

To formalize the learning and memory processes of mice in the object-based memory task, we developed a simple reinforcement-learning model based on the Rescorla-Wagner model. Here,

- $e_t \in [0, 1]$ denotes the relevancy of an engram, which is updated on each trial.

Higher values indicate higher object relevancy (stronger object-context association).

The key assumption of the model is that once engrams are formed, they endure but might be inaccessible when they cannot be reactivated following natural forgetting. The expressibility of the engram depends on the inferred relevancy of the respective object, where objects that are

subjectively more likely to be encountered in the current environment (higher relevancy), are more easily accessible. The inferred relevancy of the (familiar) acquisition object $x = 1$ governing the accessibility of the engram was computed according to the delta rule

$$e_{t+1} = e_t + \alpha_t(o_t - e_t) \quad (\text{eq. 1})$$

where e_t denotes the relevancy of the engram, α_t the learning rate, and $o_t - e_t =: \delta_t$ the prediction error. For clarity, we omit the dependency on x here. Moreover, we used separate learning rates for positive and negative prediction errors

$$\alpha_t := \begin{cases} 1, & \delta_t \geq 0 \\ \alpha^-, & \delta_t < 0 \end{cases} \quad (\text{eq. 2})$$

That is, positive prediction errors indicating the presence of an object lead to a rapid increase in the relevancy of the engram. Negative prediction errors indicating the absence of an object lead to a slower decrease in the relevancy (when $\alpha^- < 1$).

Exploration model

To translate the modeled engram relevancy into exploration behavior, we used the softmax and beta functions. The softmax function was used to compute the average exploration probability for the familiar object

$$\mu = \frac{\exp(\beta e_t^{x=1})}{\sum_x \exp(\beta e_t^x)} \quad (\text{eq. 3})$$

where the inverse temperature parameter β determines the slope of the function. $e_t^{x=1}$ denotes the engram relevancy of the familiar object that is dynamically updated according to the delta rule (eq. 1). $e_t^{x=0}$ corresponds to the relevancy of the novel object, where, since the object has not been experienced before, we assume an engram relevancy of 0. Subsequently, we utilized the beta distribution to model exploration variability across individuals. The beta distribution for $\mu \in [0,1]$ is defined by

$$\text{Beta}(\mu; a, b) := \frac{1}{B(a,b)} \mu^{a-1} (1-\mu)^{b-1} \quad (\text{eq. 4})$$

with the shape parameters a and b . In order to compute the distribution over exploration probability conditional on the mean μ , we exploited that the sum of a and b indicates the concentration κ of the distribution. That is, when $\kappa = a + b$ gets larger, the beta distribution is more concentrated, and the predicted exploration variability is lower. When κ is lower, the distribution is wider and the model predicts greater exploration variability (for more details, see (103)).

Therefore, to compute the exploration probabilities for object $x = 1$ using the beta distribution conditional on the computed average exploration probability μ and concentration κ , we computed the shape parameters a and b in the following way:

$$a = \mu\kappa \text{ and } b = (1 - \mu)\kappa. \quad (\text{eq. 5})$$

Parameter estimation

We implemented the following free parameters. The learning rate conditional on negative prediction errors α^- (see eq. 2), the inverse temperature parameter β of the softmax distribution (see eq. 3), and the κ parameter of the beta distribution (see eq. 4 and 5). To estimate these parameters, we computed the probability of the observed exploration probabilities μ for the

familiar object $x = 1$ conditional on a and b on the test day using the beta distribution (eq. 4 [↗](#)). Based upon this, we computed the model fit across subjects of the experimental and control group by summing the log-likelihoods

$$\ell = \sum_s \log p(\hat{\mu}_s | a_s, b_s) \quad (\text{eq. 6})$$

where S denotes the subject. The free parameters were estimated using the bound- constrained optimization algorithm L-BFGS-B of the SciPy library in Python 3.10. The parameter boundaries were $[0, 0.25]$ for α^- , $[-10, 0]$ for β , and $[1, 100]$ for κ .

Model comparison

We systematically compared our model to a baseline model that explored the two objects with the same probability ($\mu = 0.5$), where we computed the log-likelihood based on $a = 1$ and $b = 1$. We compared the Bayesian information criterion (BIC; (104 [↗](#))) of the two models defined by

$$BIC := \ell - \frac{\kappa}{2} \log S \quad (\text{eq. 7})$$

where κ denotes the number of free parameters and S the number of subjects of the respective group.

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Author Contributions

J.O.L. and T.R. conceived the original scientific design. J.O.L. and L.A. conducted the experiments and analysed the data. J.O.L., R.B., L.A. and T.R. interpreted the results. R.B. conducted all modelling work. J.O.L., R.B. and T.R. wrote the manuscript draft. J.O.L., R.B., L.A. and T.R. reviewed and edited the manuscript.

Declaration of Interests

The authors declare no competing interests.

References

1. Pignatelli M., et al. (2019) **Engram Cell Excitability State Determines the Efficacy of Memory Retrieval** *Neuron* **101**:274–284
2. Cowan E. T., Schapiro A. C., Dunsmoor J. E., Murty V. P. (2021) **Memory consolidation as an adaptive process** *Psychonomic Bulletin & Review* **28**:1796–1810
3. Hughes R. N., Kaiser M. J., Mackney P. A., Warburton K. (1992) **Optimizing foraging behaviour through learning** *Journal of Fish Biology* **41**:77–91
4. Eliassen S., Jørgensen C., Mangel M., Giske J. (2007) **Exploration or exploitation: life expectancy changes the value of learning in foraging strategies** *Oikos* **116**:513–523
5. Marvin C. B., Shohamy D. (2016) **Curiosity and reward: Valence predicts choice and information prediction errors enhance learning** *Journal of Experimental Psychology: General* **145**
6. den Ouden H. E. M., Friston K. J., Daw N. D., McIntosh A. R., Stephan K. E. (2009) **A Dual Role for Prediction Error in Associative Learning** *Cerebral Cortex* **19**:1175–1185
7. Miller R. R. (2021) **Failures of memory and the fate of forgotten memories** *Neurobiology of Learning and Memory* **181**
8. Ryan T. J., Frankland P. W. (2022) **Forgetting as a form of adaptive engram cell plasticity** *Nature Reviews Neuroscience* **23**:173–186
9. Josselyn S. A., Tonegawa S. (2020) **Memory engrams: Recalling the past and imagining the future** *Science* **367**
10. Ortega-de San Luis C., Ryan T. J. (2022) **Understanding the physical basis of memory: Molecular mechanisms of the engram** *Journal of Biological Chemistry* **298**
11. Tonegawa S., Pignatelli M., Roy D. S., Ryan T. J. (2015) **Memory engram storage and retrieval** *Current Opinion in Neurobiology* **35**:101–109
12. Josselyn S. A., Köhler S., Frankland P. W. (2015) **Finding the engram** *Nature Reviews Neuroscience* **16**:521–534
13. Denny C. A., Lebois E., Ramirez S. (2017) **From Engrams to Pathologies of the Brain** *Front Neural Circuits* **11**
14. Richards B. A., Frankland P. W. (2017) **The Persistence and Transience of Memory** *Neuron* **94**:1071–1084
15. Perusini J. N., et al. (2017) **Optogenetic stimulation of dentate gyrus engrams restores memory in Alzheimer’s disease mice** *Hippocampus* **27**:1110–1122
16. Guskjolen A., et al. (2018) **Recovery of “Lost” Infant Memories in Mice** *Current Biology* **28**:2283–2290

17. Roy D. S., et al. (2016) **Memory retrieval by activating engram cells in mouse models of early Alzheimer's disease** *Nature* **531**:508–512
18. Autore L., O'Leary J. D., Ortega-de San Luis C., Ryan T. J. (2023) **Adaptive expression of engrams by retroactive interference** *Cell Rep* **42**
19. Frankland P. W., Josselyn S. A., Köhler S. (2019) **The neurobiological foundation of memory retrieval** *Nat Neurosci* **22**:1576–1585
20. Josselyn S. A., Frankland P. W. (2018) **Memory Allocation: Mechanisms and Function** *Annu. Rev. Neurosci.* **41**:389–413
21. Lei B., Lv L., Hu S., Tang Y., Zhong Y. (2022) **Social experiences switch states of memory engrams through regulating hippocampal Rac1 activity** *Proc Natl Acad Sci U S A* **119**
22. Lv L., et al. (2019) **Interplay between α 2-chimaerin and Rac1 activity determines dynamic maintenance of long-term memory** *Nature Communications* **10**
23. Khalaf O., et al. (2018) **Reactivation of recall-induced neurons contributes to remote fear memory attenuation** *Science* **360**:1239–1242
24. Poo M., et al. (2016) **What is memory? The present state of the engram** *BMC Biology* **14**
25. Washington P. M., Villapol S., Burns M. P. (2016) **Polypathology and dementia after brain trauma: Does brain injury trigger distinct neurodegenerative diseases, or should they be classified together as traumatic encephalopathy?** *Experimental Neurology* **275**:381–388
26. Chapman D. P., Sloley S. S., Caccavano A. P., Vicini S., Burns M. P. (2021) **High-Frequency Head Impact Disrupts Hippocampal Neural Ensemble Dynamics** *Front Cell Neurosci* **15**
27. Sloley S. S., et al. (2021) **High-frequency head impact causes chronic synaptic adaptation and long-term cognitive impairment in mice** *Nat Commun* **12**
28. McTighe S. M., Cowell R. A., Winters B. D., Bussey T. J., Saksida L. M. (2010) **Paradoxical false memory for objects after brain damage** *Science* **330**:1408–1410
29. Davis R. L., Zhong Y. (2017) **The Biology of Forgetting-A Perspective** *Neuron* **95**:490–503
30. Wang C., et al. (2020) **Microglia mediate forgetting via complement-dependent synaptic elimination** *Science* **367**:688–694
31. Rao-Ruiz P., Yu J., Kushner S. A., Josselyn S. A. (2019) **Neuronal competition: microcircuit mechanisms define the sparsity of the engram** *Current Opinion in Neurobiology* **54**:163–170
32. Rashid A. J., et al. (2016) **Competition between engrams influences fear memory formation and recall** *Science* **353**:383–387
33. Poll S., et al. (2020) **Memory trace interference impairs recall in a mouse model of Alzheimer's disease** *Nature neuroscience* **23**:952–958
34. Bevins R. A., Besheer J. (2006) **Object recognition in rats and mice: a one-trial non-matching-to-sample learning task to study "recognition memory"** *Nat Protoc* **1**:1306–1311
35. Leger M., et al. (2013) **Object recognition test in mice** *Nature Protocols* **8**:2531–2537

36. Reijmers L. G., Perkins B. L., Matsuo N., Mayford M. (2007) **Localization of a stable neural correlate of associative memory** *Science* **317**:1230–1233
37. Ramirez S., et al. (2013) **Creating a false memory in the hippocampus** *Science* **341**:387–391
38. Ramirez S., et al. (2015) **Activating positive memory engrams suppresses depression-like behaviour** *Nature* **522**:335–339
39. Mishra R., et al. (2022) **Augmenting neurogenesis rescues memory impairments in Alzheimer’s disease by restoring the memory-storing neurons** *J Exp Med* **219**
40. Bekinschtein P., Oomen C. A., Saksida L. M., Bussey T. J. (2011) **Effects of environmental enrichment and voluntary exercise on neurogenesis, learning and memory, and pattern separation: BDNF as a critical variable?** *Seminars in Cell & Developmental Biology* **22**:536–542
41. Gubert C., Hannan A. J. (2019) **Environmental enrichment as an experience-dependent modulator of social plasticity and cognition** *Brain Research* **1717**:1–14
42. Clemenson G. D., Deng W., Gage F. H. (2015) **Environmental enrichment and neurogenesis: from mice to humans** *Current Opinion in Behavioral Sciences* **4**:56–62
43. O’Leary J. D., Hoban A. E., Cryan J. F., O’Leary O. F., Nolan Y. M. (2019) **Differential effects of adolescent and adult-initiated voluntary exercise on context and cued fear conditioning** *Neuropharmacology* **145**:49–58
44. O’Leary J. D., et al. (2019) **Differential effects of adolescent and adult-initiated exercise on cognition and hippocampal neurogenesis** *Hippocampus* **29**:352–365
45. Wahlheim C. N., Smith W. G., Delaney P. F. (2019) **Reminders can enhance or impair episodic memory updating: a memory-for-change perspective** *Memory* **27**:849–867
46. Tambini A., Berners-Lee A., Davachi L. (2017) **Brief targeted memory reactivation during the awake state enhances memory stability and benefits the weakest memories** *Scientific Reports* **7**
47. Finkelstein A. B., et al. (2022) **Social reactivation of fear engrams enhances memory recall** *Proceedings of the National Academy of Sciences* **119**
48. Bouton M. E. (1993) **Context, time, and memory retrieval in the interference paradigms of Pavlovian learning** *Psychological bulletin* **114**
49. Liu Y., Lv L., Wang L., Zhong Y. (2018) **Social Isolation Induces Rac1-Dependent Forgetting of Social Memory** *Cell Reports* **25**:288–295
50. Liu Y., et al. (2016) **Hippocampal Activation of Rac1 Regulates the Forgetting of Object Recognition Memory** *Current Biology* **26**:2351–2357
51. Shuai Y., et al. (2010) **Forgetting Is Regulated through Rac Activity in Drosophila** *Cell* **140**:579–589
52. Cervantes-Sandoval I., Davis R. L., Berry J. A. (2020) **Rac1 Impairs Forgetting-Induced Cellular Plasticity in Mushroom Body Output Neurons** *Front Cell Neurosci* **14**

53. Berry J. A., Phan A., Davis R. L. (2018) **Dopamine Neurons Mediate Learning and Forgetting through Bidirectional Modulation of a Memory Trace** *Cell Reports* **25**:651–662
54. Wu W., et al. (2019) **Inhibition of Rac1-dependent forgetting alleviates memory deficits in animal models of Alzheimer's disease** *Protein & Cell* **10**:745–759
55. Wagner A. R., Rescorla R. A. (1972) **Inhibition in Pavlovian conditioning: Application of a theory** *Inhibition and learning* :301–336
56. Todd T. P., Holmes N. M. (2022) **How common is a common error term? The rules that govern associative learning in sensory preconditioning and second-order conditioning** *Frontiers in Behavioral Neuroscience* **16**
57. Yassa M. A., Stark C. E. L. (2011) **Pattern separation in the hippocampus** *Trends Neurosci* **34**:515–525
58. Denny C. A., et al. (2014) **Hippocampal Memory Traces Are Differentially Modulated by Experience, Time, and Adult Neurogenesis** *Neuron* **83**:189–201
59. Chen B. K., et al. (2019) **Artificially Enhancing and Suppressing Hippocampus-Mediated Memories** *Current Biology* **29**:1885–1894
60. Lacagnina A. F., et al. (2019) **Distinct hippocampal engrams control extinction and relapse of fear memory** *Nat Neurosci* **22**:753–761
61. Karaca K. G., Brito D. V., Kupke J., Zeuch B., Oliveira A. M. (2021) **Engram reactivation during memory retrieval predicts long-term memory performance in aged mice** *Neurobiology of Aging* **101**:256–261
62. Cohen S. J., Stackman R. W. J. (2015) **Assessing rodent hippocampal involvement in the novel object recognition task. A review** *Behav Brain Res* **285**:105–117
63. Cohen S. J., et al. (2013) **The rodent hippocampus is essential for nonspatial object memory** *Curr Biol* **23**:1685–1690
64. McGaugh J. L. (2000) **Memory--a Century of Consolidation** *Science* **287**:248–251
65. Dudai Y., Karni A., Born J. (2015) **The Consolidation and Transformation of Memory** *Neuron* **88**:20–32
66. Osorio-Gómez D., Saldivar-Mares K. S., Perera-López A., McGaugh J. L., Bermúdez-Rattoni F. (2019) **Early memory consolidation window enables drug induced state-dependent memory** *Neuropharmacology* **146**:84–94
67. Tonegawa S., Morrissey M. D., Kitamura T. (2018) **The role of engram cells in the systems consolidation of memory** *Nature Reviews Neuroscience* **19**:485–498
68. Kastellakis G., Tasciotti S., Pandi I., Poirazi P. (2023) **The dendritic engram** *Front Behav Neurosci* **17**
69. Ryan T. J., Roy D. S., Pignatelli M., Arons A., Tonegawa S. (2015) **Engram cells retain memory under retrograde amnesia** *Science* **348**:1007–1013

70. Li J., et al. (2020) **Defective memory engram reactivation underlies impaired fear memory recall in Fragile X syndrome** *eLife* **9**
71. Barak B., et al. (2013) **Opposing actions of environmental enrichment and Alzheimer's disease on the expression of hippocampal microRNAs in mouse models** *Translational Psychiatry* **3**:e304–e304
72. Griñán-Ferré C., et al. (2018) **Environmental Enrichment Improves Cognitive Deficits, AD Hallmarks and Epigenetic Alterations Presented in 5xFAD Mouse Model** *Front Cell Neurosci* **12**
73. Jankowsky J. L., et al. (2005) **Environmental enrichment mitigates cognitive deficits in a mouse model of Alzheimer's disease** *J Neurosci* **25**:5217–5224
74. Epp J. R., Botly L. C. P., Josselyn S. A., Frankland P. W. (2021) **Voluntary Exercise Increases Neurogenesis and Mediates Forgetting of Complex Paired Associates Memories** *Neuroscience* **475**:1–9
75. Snyder J. S., Drew M. R. (2020) **Functional neurogenesis over the years** *Behavioural Brain Research* **382**
76. Sztainberg Y., Chen A. (2010) **An environmental enrichment model for mice** *Nature protocols* **5**:1535–1539
77. Wolfer D. P., et al. (2004) **Cage enrichment and mouse behaviour** *Nature* **432**:821–822
78. Frankland P. W., Köhler S., Josselyn S. A. (2013) **Hippocampal neurogenesis and forgetting** *Trends in Neurosciences* **36**:497–503
79. Akers K. G., et al. (2014) **Hippocampal neurogenesis regulates forgetting during adulthood and infancy** *Science* **344**:598–602
80. Ko S. Y., Frankland P. W. (2021) **Neurogenesis-dependent transformation of hippocampal engrams** *Neuroscience Letters* **762**
81. Tran L. M., et al. (2022) **Adult neurogenesis acts as a neural regularizer** *Proceedings of the National Academy of Sciences* **119**
82. Bouton M. E., Nelson J. B., Rosas J. M. (1999) **Stimulus generalization, context change, and forgetting** *Psychological Bulletin* **125**
83. Rosas J. M., Bouton M. E. (1997) **Additivity of the effects of retention interval and context change on latent inhibition: toward resolution of the context forgetting paradox** *Journal of Experimental Psychology: Animal Behavior Processes* **23**
84. Leake J., Zinn R., Corbit L. H., Fanselow M. S., Vissel B. (2021) **Engram Size Varies with Learning and Reflects Memory Content and Precision** *J Neurosci* **41**:4120–4130
85. Rosiles Tania, et al. (2020) **Registered Report: Transcriptional Analysis of Savings Memory Suggests Forgetting is Due to Retrieval Failure** *eNeuro* **7**
86. Riccio D. C., Richardson R. (1984) **The status of memory following experimentally induced amnesias: Gone, but not forgotten** *Physiological Psychology* **12**:59–72

87. Yau J. O.-Y., McNally G. P. (2023) **The Rescorla-Wagner model, prediction error, and fear learning** *Neurobiol Learn Mem* **203**
88. Cai L. X., et al. (2020) **Distinct signals in medial and lateral VTA dopamine neurons modulate fear extinction at different times** *Elife* **9**
89. Kalisch R., Gerlicher A. M. V., Duvarci S. (2019) **A Dopaminergic Basis for Fear Extinction** *Trends Cogn Sci* **23**:274–277
90. Tu G., et al. (2019) **Dopamine D1 and D2 Receptors Differentially Regulate Rac1 and Cdc42 Signaling in the Nucleus Accumbens to Modulate Behavioral and Structural Plasticity After Repeated Methamphetamine Treatment** *Biological Psychiatry* **86**:820–835
91. Bouton M. E. (2004) **Context and behavioral processes in extinction** *Learning & memory* **11**:485–494
92. Zhang X., Kim J., Tonegawa S. (2020) **Amygdala Reward Neurons Form and Store Fear Extinction Memory** *Neuron* **105**:1077–1093
93. Lay B. P., Koya E., Hope B. T., Esber G. R., Iordanova M. D. (2023) **The recruitment of a neuronal ensemble in the central nucleus of the amygdala during the first extinction episode has persistent effects on extinction expression** *Biological Psychiatry* **93**:300–308
94. Redish A. D., Jensen S., Johnson A., Kurth-Nelson Z. (2007) **Reconciling reinforcement learning models with behavioral extinction and renewal: implications for addiction, relapse, and problem gambling** *Psychol Rev* **114**:784–805
95. Gershman S. J., Blei D. M., Niv Y. (2010) **Context, learning, and extinction** *Psychol Rev* **117**:197–209
96. Maren S., Phan K. L., Liberzon I. (2013) **The contextual brain: implications for fear conditioning, extinction and psychopathology** *Nature Reviews Neuroscience* **14**:417–428
97. Gershman S. J., Radulescu A., Norman K. A., Niv Y. (2014) **Statistical computations underlying the dynamics of memory updating** *PLoS Comput Biol* **10**
98. Heald J. B., Lengyel M., Wolpert D. M. (2021) **Contextual inference underlies the learning of sensorimotor repertoires** *Nature* **600**:489–493
99. Reijmers L., Mayford M. (2009) **Genetic control of active neural circuits** *Front Mol Neurosci* **2**
100. Hueston C. M., et al. (2018) **Chronic interleukin-1 β in the dorsal hippocampus impairs behavioural pattern separation** *Brain Behav Immun* **74**:252–264
101. 101., Daw N. D., Tobler P. N. (2014) **Value learning through reinforcement: the basics of dopamine and reinforcement learning** *Neuroeconomics* :283–298
102. Glimcher P. W., Fehr E. (2013) **Neuroeconomics: Decision making and the brain**
103. Krushke J. (2014) **Doing Bayesian Data Analysis: A Tutorial with R, JAGS, and Stan**

104. Stephan K. E., Penny W. D., Daunizeau J., Moran R. J., Friston K. J. (2009) **Bayesian model selection for group studies** *Neuroimage* **46**:1004–1017

Editors

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Reviewer #1 (Public Review):

O'Leary and colleagues sought to understand the factors that underlie memory processes, including formation, retrieval, and forgetting. The present data identify time, environmental enrichment, Rac-1, context reexposure, and brief reminders of the familiar object as factors that alter discrimination between novel and familiar objects. This is complimented with an engram approach to quantify cells that are active during learning to examine how their activation is impacted with each of the above factors at test. There are many strengths in the manuscript, including systematic testing of several factors that contribute to poor discrimination between novel and familiar objects. These results are interesting and outline essential boundaries of incidental, nonaversive memory. With this behavioral data, authors apply a modeling approach to understand the factors that contribute to good and poor object memory recall.

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Reviewer #2 (Public Review):

Summary:

The manuscript examines an important question about how an inaccessible, natural forgotten memory can be retrieved through engram ensemble reactivation. It uses a variety of strategies including optogenetics, behavioral and pharmacological interventions to modulate engram accessibility. The data characterize the time course of natural forgetting using an object recognition task, in which animals can retrieve 1 day and 1 week after learning, but not 2 weeks later. Forgetting is correlated with lower levels of cell reactivation (c-fos expression during learning compared to retrieval) and reduction in spine density and volume in the engram cells. Artificial activation of the original engram was sufficient to induce recall of the forgotten object memory while artificial inhibition of the engram cells precluded memory retrieval. Mice housed in an enriched environment had a slower rate of forgetting, and a brief reminder before the retrieval session promoted retrieval of a forgotten memory. Repeated reintroduction to the training context in the absence of objects accelerated forgetting. Additionally, activation of Rac1-mediated plasticity mechanisms enhanced forgetting, while its inhibition prolonged memory retrieval. Authors also reproduce the behavioral findings using a computational model inspired by Rescorla-Wagner model. In essence, the model proposes that forgetting is a form of adaptive learning that can be updated based on prediction error rules in which engram relevancy is altered in response to environmental feedback.

Strengths:

- (1) The data presented in the current paper are consistent with the authors claim that seemingly forgotten engrams are, in fact, accessible. This suggests that retrieval deficits can lead to memory impairments rather than a loss of the original engram (at least in some cases).
- (2) The experimental procedures and statistics are appropriate, and the behavioral effects appear to be very robust. Several key effects are replicated multiple times in the manuscript.

Comments on revised version:

The authors have adequately addressed my prior concerns.

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Reviewer #3 (Public Review):

Summary:

The manuscript by Ryan and colleagues uses a well-established object recognition task to examine memory retrieval and forgetting. They show that memory retrieval requires activation of the acquisition engram in the dentate gyrus and failure to do so leads to forgetting. Using a variety of clever behavioural methods, the authors show that memories can be maintained and retrieval slowed when animals are reared in environmental enrichment and that normally retrieved memories can be forgotten if exposed to the environment in which the expected objects are no longer presented. Using a series of neural methods, the authors also show that activation or inhibition of the acquisition engram is key to memory expression and that forgetting is due to Rac1.

Strengths:

This is an exemplary examination of different conditions that affect successful retrieval vs forgetting of object memory. Furthermore, the computational modelling that captures in a formal way how certain parameters may influence memory provides an important and testable approach to understanding forgetting.

The use of the Rescorla-Wanger model in the context of object recognition and the idea of relevance being captured in negative prediction error are novel (but see below).

The use of gain and loss of function approaches are a considerable strength and the dissociable effects on behaviour eliminate the possibility of extraneous variables such as light artifacts as potential explanations for the effects.

Weaknesses:

A closer examination of the process that governs the behavioural effect in the present investigation would have been of even greater significance. The authors acknowledge the distinction between object familiarity vs object recognition, but a direct assessment would benefit the field's understanding of the current role of engrams on behaviour.

Relatedly, while relevance is an interesting concept that has been operationalized in the paper, it is unclear how distinct it is from extinction. Specifically, in the case where the animals are exposed to the context in the absence of the object, the paper currently expresses this as a process of relevance - the objects are no longer relevant in that context. Another way to think about this is in terms of extinction - the association between the context and the objects is reduced resulting in a disrupted ability of the context to activate the object engram. The authors have noted the potential role of extinction in their studies.

The impact of the paper is somewhat limited by the use of only one sex. Do the authors expect an identical process to be engaged in females in the present set of studies?

<https://doi.org/10.7554/eLife.92860.2.sa1>

Author response:

The following is the authors' response to the original reviews.

Public Reviews:

Reviewer #1 (Public Review):

Summary:

O'Leary and colleagues present data identifying several procedures that alter discrimination between novel and familiar objects, including time, environmental enrichment, Rac-1, context reexposure, and brief reminders of the familiar object. This is complimented with an engram approach to quantify cells that are active during learning to examine how their activation is impacted following each of the above procedures at test. With this behavioral data, authors apply a modeling approach to understand the factors that contribute to good and poor object memory recall.

We thank the Reviewer for summarizing the scope and depth of our manuscript, and indeed for recognizing our efforts. We engage below with the Reviewer's specific criticisms.

Strengths:

Authors systematically test several factors that contribute to poor discrimination between novel and familiar objects. These results are extremely interesting and outline essential boundaries of incidental, nonaversive memory.

These results are further supported by engram-focused approaches to examine engram cells that are reactivated in states with poor and good object recognition recall.

We thank the Reviewer for these positive comments.

Weaknesses:

For the environmental enrichment, authors seem to suggest objects in the homecage are similar to (or reminiscent of) the familiar object. Thus, the effect of improved memory may not be related to enrichment per se as much as it may be related to the preservation of an object's memory through multiple retrievals, not the enriching experiences of the environment itself. This would be consistent with the brief retrieval figure. Authors should include a more thorough discussion of this.

This is one of the main issues highlighted by the Editor and the Reviewers. We agree that these results dove-tail with the reminder experiments. We have included additional discussion, see line 510-546.

Authors should justify the marginally increased number of engram cells in the non-enrichment group that did not show object discrimination at test, especially relative to other figures. More specific cell counting criteria may be helpful for this. For example, was the DG region counted for engram and cfos cells or only a subsection?

There was a marginal, but non-significant increase in the number of labelled cells within the standard housed mice in Figure 3f. The cell counting criteria was the same across experimental groups and conditions, where the entire dorsal and ventral blade of the dorsal DG was counted for each animal. This non-statistically significant variance may be due to surgical and viral spread difference between mice. We have clarified this in the manuscript, see line 229-232.

It is unclear why the authors chose a reactivation time point of 1hr prior to testing. While this may be outside of the effective time window for pharmacological interference with reconsolidation for most compounds, it is not necessarily outside of the structural and functional neuronal changes accompanied by reconsolidation-related manipulations.

A control experiment was performed to demonstrate that a brief reminder exposure of 5 mins on its own was insufficient to induce new learning that formed a lasting memory (Supplementary Figure S4a). Mice given only a brief acquisition period of 5 mins, exhibited no preference for the novel object when tested 1 hour after training, suggesting the absence of a lasting object memory (Supplementary Figure S4b & c). We therefore used the 1-hour time point for the brief reminder experiment in Figure 4a. We have clarified this within the manuscript and supplementary data see line 258-264.

Figure 5: Levels of exploration at test are inconsistent between manipulations. This is problematic, as context-only reexposures seem to increase exploration for objects overall in a manner that I'm unsure resembles 'forgetting'. Instead, cross-group comparisons would likely reveal increased exploration time for familiar and novel objects. While I understand 'forgetting' may be accompanied by greater exploration towards objects, this is inconsistent across and within the same figure. Further, this effect is within the period of time that rodents should show intact recognition. Instead, context-only exposures may form a competing (empty context) memory for the familiar object in that particular context.

The Reviewer raises an important question, and we agree with the Reviewer that there should be caution and qualification around interpreting these results as “forgetting”. Indeed, for the context-only reexposures, cross-group comparisons show increased exploration time for familiar and novel objects. As the mice exhibit relatively high exploration of both the novel and familiar objects. An alternative explanation would be that the mice have not truly forgotten the familiar object, but rather as the mouse has not seen the familiar object in the last 6 context only sessions, its reappearance makes it somewhat novel again. Therefore, this change in the object's reappearance triggers the animal's curiosity, and in turn drives exploration by the animal. In addition, the context-only exposures may form a competing memory for the familiar object in that particular context. We have highlighted this in the results and also included greater discussion. See lines 306-315.

I am concerned at the interpretation that a memory is 'forgotten' across figures, especially considering the brief reminder experiments. Typically, if a reminder session can trigger the original memory or there is rapid reacquisition, then this implies there is some savings for the original content of the memory. For instance, multiple context retrievals in the absence of an object reminder may be more consistent with procedures that create a distinct memory and subsequently recruit a distinct engram.

These findings raise an important question regarding the interpretation of ‘forgetting’. If a reminder trial or experience can trigger the original memory, or there is rapid reacquisition, then this would suggest there is a degree of savings for the original memory content (85, 86). Previous work has emphasized retrieval deficits as a key characteristic of memory

impairment, supporting the idea that memory recall or accessibility may be driven by learning feedback from the environment (7, 8, 14–18). Within our behavioral paradigm, a lack of memory expression would still constitute forgetting due to the loss of learned behavioral response in the presence of natural retrieval cues. The changes in memory expression may therefore underlie the adaptive nature of forgetting. This is consistent with the idea that the engram is intact and available, but not accessible. Here we studied natural forgetting, and our data showing memory retrieval following optogenetic reactivation demonstrates that the original engram persists at a cellular level, otherwise activation of those cells would no longer trigger memory recall. We also agree with the reviewer that multiple context retrievals may indeed lead to the formation of a second distinct engram that competes with the original. Recent work suggests that retroactive interference emerges from the interplay of multiple engrams competing for accessibility (18). We have added clarification and included extra discussion of this interpretation. See lines 589-598.

Authors state that spine density decreases over time. While that may be generally true, there is no evidence that mature mushroom spines are altered or that this is consistent across figures. Additionally, it's unclear if spine volume is consistently reduced in reactivated and non-reactivated engram cells across groups. This would provide evidence that there is a functionally distinct aspect of engram cells that is altered consistently in procedures resulting in poor recognition memory (e.g. increased spine density relative to spine density of non-reactivated engram cells and non-engram cells)

We thank the Reviewer for their helpful comments on explaining our engram dendritic spine data. We agree with the Reviewer that an analysis of the changes in spine type, as well as the difference between engram and non-engram spines as well and reactivation and non-reactivated engram spines would be interesting and may help to further illuminate the morphological changes of forgetting and memory retrieval. Indeed, future analysis could determine if spine density is reduced in reactivated and non-reactivated engram cells or indeed across engram non-engram cells within different learning conditions. This avenue of investigation could determine if there is a functionally distinct aspect of engram cells that are altered following forgetting (67). However, such analysis is beyond the scope of this study. We have highlighted this limitation and included its discussion. See lines 493-499.

Authors should discuss how the enrichment-neurogenesis results here are compatible with other neurogenesis work that supports forgetting.

We validated the effectiveness of the enrichment paradigm to enhance neural plasticity by measuring adult hippocampal neurogenesis. The hippocampus has been identified as one of the only regions where postnatal neurogenesis continues throughout life (75). Levels of adult hippocampal neurogenesis do not remain constant throughout life and can be altered by experience (41–43, 57). In addition, adult born neurons have been shown to contribute to the process of forgetting (74, 78, 79). Although the contribution of adult born neurons to cognition and the memory engram is not fully understood (80, 81). Mishra et al, showed that immature neurons were actively recruited into the engram following a hippocampal-dependent task (67). Moreover, increasing the level of neurogenesis rescued memory deficits by restoring engram activity (67). Augmenting neurogenesis further rescued the deficits in spine density in both immature and mature engram neurons in a mouse model of Alzheimer's disease (67). Whether neurogenesis alters spine density on differentially for reactivated or non-reactivation engrams cells remains to be investigated (67, 68). This avenue of research would help to illuminate the morphological changes following forgetting and provide evidence if there is a functionally distinct aspect of engram cells that is altered in forgetting (67, 68). Our engram labelling strategy which utilized c-fos-tTA transgenic mice combined with an AAV9-TRE-ChR2-eYFP virus does not necessarily label sufficient immature neurons. Future work could utilize a different engram preparation, such as a genetic labelling strategy (TRAP2) or

using a different immediate early gene promoter such as Arc to investigate the contribution of new-born neurons to the engram ensemble. We have added additional discussion of how our work fits with previous literature investigating neurogenesis and forgetting. See lines 547-565.

Reviewer #2 (Public Review):

Summary:

The manuscript examines an important question about how an inaccessible, natural forgotten memory can be retrieved through engram ensemble reactivation. It uses a variety of strategies including optogenetics, behavioral and pharmacological interventions to modulate engram accessibility. The data characterize the time course of natural forgetting using an object recognition task, in which animals can retrieve 1 day and 1 week after learning, but not 2 weeks later. Forgetting is correlated with lower levels of cell reactivation (c-fos expression during learning compared to retrieval) and reduction in spine density and volume in the engram cells. Artificial activation of the original engram was sufficient to induce recall of the forgotten object memory while artificial inhibition of the engram cells precluded memory retrieval. Mice housed in an enriched environment had a slower rate of forgetting, and a brief reminder before the retrieval session promoted retrieval of a forgotten memory. Repeated reintroduction to the training context in the absence of objects accelerated forgetting. Additionally, activation of Rac1-mediated plasticity mechanisms enhanced forgetting, while its inhibition prolonged memory retrieval. The authors also reproduce the behavioral findings using a computational model inspired by Rescorla-Wagner model. In essence, the model proposes that forgetting is a form of adaptive learning that can be updated based on prediction error rules in which engram relevancy is altered in response to environmental feedback.

We thank the Reviewer for summarizing the scope and depth of our manuscript, and for recognizing our efforts. We engage below the Reviewer's specific criticisms of our interpretations.

Strengths:

(1) The data presented in the current paper are consistent with the authors claim that seemingly forgotten engrams sometimes remain accessible. This suggests that retrieval deficits can lead to memory impairments rather than a loss of the original engram (at least in some cases).

We thank the Reviewer for their positive summary.

(2) The experimental procedures and statistics are appropriate, and the behavioral effects appear to be very robust. Several key effects are replicated multiple times in the manuscript.

We thank the Reviewer for their positive comments.

Weaknesses:

(1) My major issue with the paper is the forgetting model proposed in Figure 7. Prior work has shown that neutral stimuli become associated in a manner similar to conditioned and unconditioned stimuli. As a result, the Rescorla-Wagner model can be used to describe this learning (Todd & Homes, 2022). In the current experiments, the neutral context will become associated with the unpredicted objects during training (due

to a positive prediction error). Consequently, the context will activate a memory for the objects during the test, which should facilitate performance. Conversely, any manipulation that degrades the association between the context and object should disrupt performance. An example of this can be found in Figure 5A. Exposing the mice to the context in the absence of the objects should violate their expectations and create a negative prediction error. According to the Rescorla-Wagner model, this error will create an inhibitory association between the context and the objects, which should make it harder for the former to activate a memory of the latter (Rescorla & Wagner, 1972). As a result, performance should be impaired, and this is what the authors find. However, if the cells encoding the context and objects were inhibited during the context-alone sessions (Figure 5D) then no prediction error should occur, and inhibitory associations would not be formed. As a result, performance should be intact, which is what the authors observe.

What about forgetting of the objects that occurs over time? Bouton and others have demonstrated that retrieval failure is often due to contextual changes that occur with the passage of time (Bouton, 1993; Rosas & Bouton, 1997, Bouton, Nelson & Rosas, 1999). That is, both internal (e.g. state of the animal) and external (e.g. testing room, chambers, experimenter) contextual cues change over time. This shift makes it difficult for the context to activate memories with which it was once associated (in the current paper, objects). To overcome this deficit, one can simply re-expose animals to the original context, which facilitates memory retrieval (Bouton, 1993). In Figure 2D, the authors do something similar. They activate the engram cells encoding the original context and objects, which enhances retrieval.

Therefore, the forgetting effects presented in the current paper can be explained by changes in the context and the associations it has formed with the objects (excitatory or inhibitory). The results are perfectly predicted by the Rescorla-Wagner model and the context-change findings of Bouton and others. As a result, the authors do not need to propose the existence of a new "forgetting" variable that is driven by negative prediction errors. This does not add anything novel to the paper as it is not necessary to explain the data (Figures 7 and 8).

We thank the reviewer for clearly explaining their concern about our model. We are very sorry that we did not sufficiently explain that our model is, in fact, based on the classic Rescorla-Wagner model. The key equation of the model that updates "engram strength" is equivalent to the canonical Rescorla-Wagner model that is commonly used in research on reinforcement learning and decision-making (105). One potential minor difference is that we crucially assume different learning rates for positive and negative prediction errors. However, this variant of the Rescorla-Wagner model is common in the computational literature and is generally not regarded as a qualitatively different kind of model. In fact, it allows us to capture that establishing an object-context association (after a positive prediction error) is faster than the forgetting process (through negative errors).

The other equations that are explained in detail in the Methods are necessary to simulate exploration behavior and render the model suitable for model fitting. Concerning exploration behavior, we use the softmax function, which is commonly used in combination with the Rescorla-Wagner model, in order to translate the learned quantity (in our case, engram strength) into behavior (here exploration). The other equations are necessary to fit the model to the data (learning rate α and behavioral variability in exploration behavior).

Therefore, we fully agree with the reviewer that the Rescorla-Wagner can explain our empirical results, in particular by assuming that the different manipulations affect the strength of object-context associations, which, in turn, governs forgetting as behaviorally observed.

In our previous version of the manuscript, we only referred to the Rescorla-Wagner model directly in the Methods. But to make this important point clearer, we now refer to the origin of the model multiple times in the Results section as well. See lines 81, 386-393.

We also agree with the reviewer that the learning/forgetting process can be described in terms of changes in object-context associations (e.g., inhibitory associations after a negative prediction error). Therefore, we now explicitly refer to the relationship between updated object-context associations and forgetting and highlight that we believe that stronger associations signal higher engram “relevancy”. See lines 386-393.

We have extended Figure 7 (new panels a and b), where we illustrate the idea that (a) object-context associations govern forgetting and (b) show the key Rescorla-Wagner equation, including a simple explanation of the main terms (engram strength, prediction error, and learning rate). Finally, we have also extended our discussion of the model, where we now directly state that the Rescorla-Wagner model captures the key results of our experiments. See lines 573-580.

In order to further support a link between our empirical data and computational modeling, we also added extra experiments that showed the modulation of engram cells within the dentate gyrus can regulate these object-context associations. See Supplementary Figure 12a-f and lines 401-404.

To summarize our reply, we agree with the reviewer’s comment and hope that we have clarified the direct relationship to the Rescorla-Wagner model.

(2) I also have an issue with the conclusions drawn from the enriched environment experiment (Figure 3). The authors hypothesize that this manipulation alleviates forgetting because "Experiencing extra toys and objects during environmental enrichment that are reminiscent of the previously learned familiar object might help maintain or nudge mice to infer a higher engram relevancy that is more robust against forgetting." This statement is completely speculative. A much simpler explanation (based on the existing literature) is that enrichment enhances synaptic plasticity, spine growth, etc., which in turn reduces forgetting. If the authors want to make their claim, then they need to test it experimentally. For example, the enriched environment could be filled with objects that are similar or dissimilar to those used in the memory experiments. If their hypothesis is correct, only the similar condition should prevent forgetting.

We thank the Reviewer for this alternative perspective on our findings. First of all, we agree that this statement is speculative. The effects of enrichment on neural plasticity are well established and it likely contributes to the enhanced memory recall. It is important to emphasize that this process of updating is not necessarily separate from enrichment-induced plasticity at an implementational level, but part of the learning experience within an environment containing multiple objects. The enrichment or, more generally, experience, may therefore enhance memory through the modification of activity of specific engram ensembles. The idea of enrichment facilitating memory updating is consistent with the results obtained by the reminder experiments and further supported by our analysis with the Rescorla-Wagner computational model, where experience updates the accessibility of existing memories, possibly through reactivation of the original engram ensemble.

We would like to further clarify that our explanation concerns the algorithmic level, in contrast to the neural level. Based on the computational analyses using the Rescorla-Wagner model and in line with the reviewer’s previous comment on the model, we believe that forgetting is governed by the strength of object-context associations (or engram relevancy). Our interpretation is that stronger associations signal that the memory or engram representation is important (“relevant”) and should not be forgotten. Accordingly, due to a

vast majority of experiences with extra cage objects in the enriched environment, mice might generally learn that such objects are common in their environment and potentially relevant in the future (i.e., the object-context association is strong, preventing forgetting). Our speculation of these results is to help unify our empirical data with the computational model.

We believe that the Reviewer's alternative explanation in terms of synaptic plasticity, spine growth is not mutually exclusive with the modelling work. It is possible that the computational mechanisms that we explore based on the Rescorla-Wagner model are neurally related to the biological mechanisms that the reviewer suggests at the implementational level. Therefore, ultimately, the two perspectives might even complement each other. We have included additional discussion to clarify this point. See lines 510-546.

(3) It is well-known that updating can both weaken or strengthen memory. The authors suggest that memory is updated when animals are exposed to the context in the absence of the objects. If the engram is artificially inhibited (opto) during context-only re-exposures, memory cannot be updated. To further support this updating idea, it would be good to run experiments that investigate whether multiple short re-exposures to the training context (in the presence of the objects or during optogenetic activation of the engram) could prevent forgetting. It would also be good to know the levels of neuronal reactivation during multiple re-exposures to the context in the absence versus context in the presence of the objects.

We thank the Reviewer for their comments. We agree that additional experiments would be helpful to further support the idea of updating. We have performed additional experiments to test the idea that multiple short re-exposures to the training context, in the presence of objects prevents forgetting. In this paradigm, mice were repeatedly exposed to the original object pair (Supplementary Figure S5a). The results indicate that repeated reminder trials facilitate object memory recall (Supplementary Figure 5b&c). These data indicated that subsequent object reminders over time facilitates the transition of a forgotten memory to an accessible memory. See Supplementary Figure S5 and Lines 279-287.

(4) There are a number of studies that show boundary conditions for memory destabilization/reconsolidation. Is there any evidence that similar boundary conditions exist to make an inaccessible engram accessible?

The Reviewer asks an interesting question about boundary conditions and engram accessibility. Boundary conditions could indeed affect the degree of destabilization and reconsolidation, the salience or strength of the memory, as well as the timing of retrieval cues. Future models could focus on understanding the specific boundary conditions in which a memory becomes retrievable and the degree to which it is sufficiently destabilized and liable for updating and forgetting. We have included additional discussion on the potential role of boundary conditions for engram accessibility. See lines 661-666.

(5) More details about how the quantification of immunohistochemistry (c-fos, BrdU, DAPI) was performed should be provided (which software and parameters were used to consider a fos positive neurons, for example).

We have added additional information for the parameters of quantification of immunohistochemistry. See lines 796-809.

(6) Duration of the enrichment environment was not detailed.

We have highlighted the details for the environmental enrichment duration. See lines 756.

Reviewer #3 (Public Review):

Summary:

The manuscript by Ryan and colleagues uses a well-established object recognition task to examine memory retrieval and forgetting. They show that memory retrieval requires activation of the acquisition engram in the dentate gyrus and failure to do so leads to forgetting. Using a variety of clever behavioural methods, the authors show that memories can be maintained and retrieval slowed when animals are reared in environmental enrichment and that normally retrieved memories can be forgotten if exposed to the environment in which the expected objects are no longer presented. Using a series of neural methods, the authors also show that activation or inhibition of the acquisition engram is key to memory expression and that forgetting is due to Rac1.

We thank the Reviewer for summarizing the scope and depth of our manuscript, and indeed for recognizing our efforts. We engage below the Reviewer's specific criticisms of our interpretations.

Strengths:

This is an exemplary examination of different conditions that affect successful retrieval vs forgetting of object memory. Furthermore, the computational modelling that captures in a formal way how certain parameters may influence memory provides an important and testable approach to understanding forgetting.

The use of the Rescorla-Wagner model in the context of object recognition and the idea of relevance being captured in negative prediction error are novel (but see below).

The use of gain and loss of function approaches are a considerable strength and the dissociable effects on behaviour eliminate the possibility of extraneous variables such as light artifacts as potential explanations for the effects.

We thank the Reviewer for their positive comments.

Weaknesses:

Knowing what process (object retrieval vs familiarity) governed the behavioural effect in the present investigation would have been of even greater significance.

The Reviewer touches on an important issue of the object recognition task. Understanding how experience alters object familiarity versus object retrieval and its impact on learning would help to develop better models of object memory and forgetting. We have added additional discussion. See lines 666-669.

The impact of the paper is somewhat limited by the use of only one sex.

We agree that using only male mice limits the impact of the paper. Indeed, the field of behavioural neuroscience is moving to include sex as a variable. Future experiments should include both male and female mice.

While relevance is an interesting concept that has been operationalized in the paper, it is unclear how distinct it is from extinction. Specifically, in the case where the animals are exposed to the context in the absence of the object, the paper currently expresses this as a process of relevance - the objects are no longer relevant in that context. Another way to think about this is in terms of extinction - the association between the context and the

objects is reduced results in a disrupted ability of the context to activate the object engram.

We thank the reviewer for their insightful comment on the connection between engram relevance and memory extinction. Lacagnina et al., demonstrated that extinction training suppressed the reactivation of a fear engram, while activating a second putative extinction ensemble (59). In another study, these extinction engram cells and reward cells were shown to be functionally interchangeable (92). Moreover, in a study conducted by Lay et al., the balance between extinction and acquisition was disrupted by inhibiting the extinction recruited neurons in the BLA and CN (93). These results suggested that decision making after extinction can be governed by a balance between acquisition and extinction specific ensembles (93). Together, this may suggest that in the present study, when mice are repeatedly exposed to the training context, the association between the context and the objects is reduced, resulting in a disrupted ability of the context to activate the object engram. Therefore, memory relevance and extinction may operate similarly to effect engram accessibility, and in essence ‘forgetting’ of object memories may be due to neurobiological mechanisms similar to that of extinction learning (4). We have included additional discussion on the link between our results and the extinction literature. See lines 642-654.

Recommendations for the authors:

Reviewer #1 (Recommendations For The Authors):

Additional measures that may help interpretation of and clarify data are:

A minute-by-minute analysis for training and testing may provide insight about the learning rate and testing temporal dynamics that may shed light substantially on differential levels of exploration. This should be applied across figures and would support conclusions from models in Figures 7-8 as well.

Locomotion/distance travelled measures.

We have included additional analysis for a minute-by-minute analysis of training and testing of the object memory test at 24 hr, 2 weeks as well as under the standard housing and enrichment conditions. The results further support the initial finding that novel object recognition is increased in mice that recall the object at 24 hr. Similarly, mice housed in the enriched housing initially explore the novel object more compared to the familiar object. See Supplementary Figure 1 and 2, as well as lines 103-105 and 211-213.

The appropriate control for the context exposure figure would be to expose to a novel context in one group and the acquisition/testing context for the other.

We agree with the reviewer that an additional control of a novel context would further support our findings. Indeed, this line of investigate may dovetail with the other reviewer comments on the role of competing engrams and interference. Future work could investigate the degree to which novel contexts and multiple memories can affect the rate of forgetting through engram updating. We have included additional discussion. See lines 643 and 655. However, in our experience it is necessary to pre-expose mice to different contexts before object exposure (e.g. Autore et al '23), in order to form discriminate object/context associations. Establishing such a paradigm for this study would be at odds with the established paradigms and schedules in this current study. Moreover, the possibility that the effect of object displacement on forgetting requires the familiar context, or not, does not impact the main conclusions of this study. However, we agree that it is a point for expansion in the future.

A control virus+light group vs simply a no-light condition.

For optogenetic experiments. Control mice underwent the same surgery procedure with virus and optic fibre implantation. However, no light was delivered to excite or inhibit the respective opsin. Previous papers have shown laser light delivered to tissue expressing an AAV-TRE-EYFP lacking an light-opsin does cause cellular excitation. We have clarified this in the text. See lines 726-729.

Reviewer #2 (Recommendations For The Authors):

Minor details:

(1) In the pharmacological modification of Rac 1, please specify what percentage of DMSO was used to dissolve Rac1 inhibitor and correct the typo 'DSMO'

Rac1 inhibitor (Ehop016) was reconstituted and prepared in PBS with 1% Tween-80, 1% DMSO and 30% PEG. We have clarified this in the text and corrected the typo, thank you. See lines 767.

(2) In the penultimate paragraph there is a typo 'predication error'

This is now corrected. Thankyou.

Reviewer #3 (Recommendations For The Authors):

I was unable to find information on what the No Light group consisted of. Was there a control virus infused, were the animals implanted with optical fibres (in the presence or absence of a virus), were they surgical controls, etc?

For optogenetic experiments. No Light Control mice underwent the same surgery procedure with virus and optic fibre implantation. However, no light was delivered to excite or inhibit the respective opsin. We have clarified this in the text. See lines 726-729.

The discussion lacked specificity in places. For example, the idea of eluding to 'other variables' is somewhat vague (p. 21, middle paragraph). Some examples of what other variables could be relevant would be helpful in capturing what direction or relevance the model may have going forward.

We have expanded the discussion of other variables which might impact engram relevance and how the model might be developed moving forward. These may include, boundary conditions of destabilization and reconsolidation, the salience or strength of the memory as well as the timing of retrieval cues or updating experience. Future models could focus on understanding the specific boundary conditions in which a memory becomes retrievable and the degree to which it is sufficiently destabilized and liable for updating and forgetting. The role of perceptual learning on memory retrieval and forgetting may also be an avenue of future investigation. Understanding how experience alters object familiarity versus object retrieval and its impact on learning would also help to develop better models of object memory and forgetting. In the current study, only male mice were utilized. Therefore, future work could also include sex as a variable to fully elucidate the impact of experience on the processes of forgetting. See lines 642-669.

In the same paragraph (p. 21, middle paragraph) there is mention of multiple engrams and how they can compete. The authors reference Autore et al (2023), but I thought Lacagina did this really beautifully also in an experimental setting. This idea is also

expressed in Lay et al. (2022). So additional references would further strengthen the authors argument here.

We thank the reviewer for the additional references for discussing engram competition. We have included these papers in the discission. See lines 642-654.

Relatedly, environmental enrichment was considered in terms of object relevance. I wonder if the authors may want to consider thinking about their results in terms of effects on perceptual learning.

Indeed, perceptual learning maybe playing a role in environmental enrichment. We have included additional discussion. See lines 666-669.

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