Microglia mediate forgetting via complement-dependent synaptic elimination

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Synapses between engram cells are believed to be substrates for memory storage, and the weakening or loss of these synapses leads to the forgetting of related memories. We found engulfment of synaptic components by microglia in the hippocampi of healthy adult mice. Depletion of microglia or inhibition of microglial phagocytosis prevented forgetting and the dissociation of engram cells. By introducing CD55 to inhibit complement pathways, specifically in engram cells, we further demonstrated that microglia regulated forgetting in a complement- and activity-dependent manner. Additionally, microglia were involved in both neurogenesis-related and neurogenesis-unrelated memory degradation. Together, our findings revealed complement-dependent synapse elimination by microglia as a mechanism underlying the forgetting of remote memories.

Memory is coded and allocated to engrams within related brain regions (1, 2). Reactivation of engram cells is essential for memory recall, whereas failure in reactivation of engram cells leads to the forgetting of related memories (3). Synaptic connections between engram cells are believed to be substrates for memory storage (4, 5). Circuit rewiring and synaptic reorganization may lead to loss or weakening of synaptic connections between engram cells, resulting in the forgetting of previously existing memories. For example, massive synaptic reorganization takes place in the dentate gyrus (DG) as continuously generated newborn neurons integrate into the hippocampal neural circuit, which leads to the forgetting of hippocampus-dependent memories (6–8). Even in mature neurons, experience- and learning-dependent, dynamic remodeling of synapses occurs constantly throughout life (9–13), providing a potential mechanism for the erasure of stored memories in the synaptic connections of these cells. Microglia are not only important for pruning excessive synapses during postnatal brain development but are also involved in the dynamics of synapses in the adult brain (14–17). Because they survey the brain and play crucial roles in monitoring synapses and determining the wiring of the brain (15, 18, 19), microglia may affect the stability of synaptic connections within the neural circuits where memories are allocated.

First, we used contextual fear conditioning (CFC) to assess the memory retention in C57BL/6 mice. We measured the freezing behavior of the animals during a test performed 5 or 35 days after three training sessions, each training session consisting of three weak foot shocks (Fig. 1A). We observed a significant decrease in the freezing of animals at 35 days compared with 5 days after training (Fig. 1B). We then carried out CFC using CD11b-DTR mice, in which diphtheria toxin receptor (DTR) is specifically expressed in CD11b-expressing microglia (20). By intracerebroventricularly administering diphtheria toxin (DT) after training, we depleted microglia in these CD11b-DTR mice until the test (Fig. S1). Thirty-five days later, CD11b-DTR mice treated with DT showed significantly higher freezing levels than those in the saline group (Fig. S1). To avoid the effect of daily injection on animal behavior, we depleted microglia in C57BL/6 mice with PLX3397 (PLX), a CSF1R/c-kit antagonist (21), via mouse diet after CFC training (Fig. 1C).

To exclude the possibility that depleting microglia may affect formation or retrieval of memories, we tested the freezing of mice a short time (5 days) after training (Fig. S2A). We found that PLX treatment did not alter the freezing of animals (Fig. S2B). Furthermore, we started administration of PLX to deplete microglia before the training and tested 24 hours later (Fig. S2C). No significant difference was observed between control and PLX-treated animals (Fig. S2D). Further behavioral tests showed that PLX treatment for 35 days did not significantly change the behavior of animals in an elevated plus maze or an open field (Fig. S3).

Memory retrieval requires reactivation of engram cells (3), whereas dissociation of engram cells—i.e., engram cells being unable to reactivate at the same time—leads to forgetting. To test whether the microglia-mediated forgetting of already-formed memory correlates with dissociation of engram cells, we used a FosTRAP strategy for tagging activated neurons during CFC training (22). We trained c-FosCreERT2::Ai4 mice for contextual fear memory and administered tamoxifen (TAM) before the last training session to induce permanent expression of dTomato in activated engram neurons. Immunofluorescent staining for c-Fos was performed after the test, and the reactivation rate of engram cells was assessed by analyzing c-Fos·dTomato colocalization in the DG (Fig. 1, G and H). Under physiological conditions, the reactivation rate of engram cells 35 days after training significantly decreased compared with that measured at 5 days, which correlates with the forgetting of related memory (Fig. II). Thirty-five days but not 5 days after training, PLX treatment significantly increased the reactivation rate of the engram cells (Fig. II), without altering the number of dTomato-engram cells in the DG (Fig. S4). The freezing of animals during the test positively correlates with the reactivation rate of engram cells (Fig. I).

During postnatal development, microglia are involved in synaptic reorganization and circuitry refinement by synaptic pruning (15). We imaged microglia in the DG of adult CX3CR1GFP+ mice, in which microglia were labeled with green fluorescent protein (GFP). When coadministered with synaptophysin or PSD95, markers for pre- or postsynaptic components, we found synaptophysin and PSD95 puncta were present in GFP+ microglia, colocalizing with lysosome marker Lamp1 (Fig. 2, A and B, and movies S1 and S2).

To test whether synaptic elimination by microglial phagocytosis may mediate forgetting, we systematically administered minocycline (Mino)—which has been shown to inhibit microglial engulfment of synapses in vitro and in vivo (15, 23)—after CFC training until the test (fig. S5A). Thirty-five days later, Mino-treated animals showed significantly longer freezing time (fig. S5B). Immunostaining showed that microglia in Mino-treated CX3CR1GFP+ animals

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Complement cascades are important for tagging synapses to be eliminated by microglia during brain development. C1q, the initiating protein of the classical complement cascade, localizes to synapses during developmental circuit refinement (24). C1q-tagging of the synapses leads to deposition of C3, which activates C3 receptors on microglia and triggers microglial phagocytosis (15, 24). We found that C1q was present within microglia, colocalizing with PSD95 and CD68, a microglial lysosomal marker (Fig. 2C and movie S3). Furthermore, using brain sections from c-Fos-CreERT2::Ai14 mice, in which engram cells were labeled with dTomato, we found colocalization of C1q with ~1.193 ± 0.335% of the dendritic spines of engram cells (Fig. 2, D and E, and movie S4) as well as colocalization of dTomato, PSD95, and CD68 within microglia (Fig. 2F and movie S5). Correspondingly, engram cells showed higher spine density in PLX-treated animals (fig. S6).

To test whether complement pathways are responsible for microglia-mediated engram dissociation and forgetting, we constructed a Cre-dependent adeno-associated virus (AAV) vector expressing CD55 (also known as decay-accelerating factor, or DAF), which is a synthetic elimination by microglial phagocytosis (15, 24). We found that C1q was present within microglia, colocalizing with PSD95 and CD68, a microglial lysosomal marker (Fig. 2C and movie S3). Furthermore, using brain sections from c-Fos-CreERT2::Ai14 mice, in which engram cells were labeled with dTomato, we found colocalization of C1q with ~1.193 ± 0.335% of the dendritic spines of engram cells (Fig. 2, D and E, and movie S4) as well as colocalization of dTomato, PSD95, and CD68 within microglia (Fig. 2F and movie S5). Correspondingly, engram cells showed higher spine density in PLX-treated animals (fig. S6).

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Fig. 2. Microglia mediate forgetting through complement system. (A and B) Superresolution microscopy images and three-dimensional (3D) reconstructions showing the presence of synaptophysin (Syn) or PSD95 in microglia (GFP+), colocalizing with Lamp1, in the DG of CX3CR1<sup>GFP</sup>+ mice. Scale bars, 5 μm; white arrows indicate Syn+Lamp1+ (A) or PSD95+Lamp1+ (B) puncta within microglia. (C) Images and 3D reconstruction showing the presence of C1q and PSD95 in microglia, colocalizing with CD68. Scale bars, 5 μm; white arrows indicate C1q+PSD95+CD68+ puncta within microglia. (D) Images and 3D reconstruction showing colocalization of C1q with a dendritic spine of an engram cell (dT<sup>Tomato</sup>) in the DG. Scale bars, 1 μm; white arrows indicate colocalization of C1q with a dendritic spine of an engram cell. (E) Percentage of engram cell dendritic spines showing colocalization with C1q. n = 3 mice, N = 76 dendritic segments. (F) Images and 3D reconstruction showing the presence of engram cell components (dT<sup>Tomato</sup>) in the microglia (Iba-1<sup>-</sup>), colocalizing with PSD95 and CD68. Scale bars, 5 μm; white arrows indicate dTomato+PSD95+CD68+ puncta within microglia. (G) Diagram of AAV vectors. (H) Experimental scheme for expressing CD55 in engram cells in the DG of c-Fos-Cre<sup>ERT2</sup> mice. (I) CD55 animals showed higher freezing level. mCherry n = 21 mice, CD55 n = 17 mice; t = 5.033, df = 36, ***P < 0.0001. (J) Images showing engram cells labeled by mCherry or CD55 AAV vectors (red) in the DG and neurons activated during the test expressed by c-Fos (green). White arrows indicate reactivated engram cells (mCherry+c-Fos<sup>+</sup>). Insets show the colocalization of mCherry and c-Fos. Scale bar, 20 μm. (K) Reactivation rate of engram cells (c-Fos/mCherry<sup>+</sup>) showed an increase in CD55 animals. n = 8 mice, CD55 n = 9 mice; t = 5.916, df = 15, ****P < 0.0001. (L) Images showing engram cell components (mCherry<sup>+</sup>) colocalizing with CD68 in microglia (Iba-1<sup>-</sup>) in mCherry mice, but not in CD55 animals. White arrows indicate mCherry<sup>+</sup>/CD68<sup>+</sup> puncta in microglia. Scale bars, 5 μm; white arrows indicate mCherry<sup>+</sup>/CD68<sup>+</sup> (mCherry) puncta or mCherry<sup>-</sup>/CD68<sup>+</sup> (CD55) puncta in microglia. (M) Percentage of microglia containing mCherry<sup>+</sup> puncta decreased in CD55 mice. mCherry n = 3 mice, N = 36 cells; CD55 n = 4 mice, N = 34 cells; t = 24.16, df = 5, ****P < 0.0001. (N) CD55 expression in engram cells decreased the volume of mCherry<sup>+</sup> puncta in microglia containing mCherry<sup>+</sup> signals. mCherry n = 3 mice, N = 34 cells; CD55 n = 4 mice, N = 9 cells; t = 3.755, df = 41, ***P = 0.0005.
known inhibitor of both classical and alternative complement pathways (26). We injected AAV-hSyn-DIO-CD55-p2A-mCherry (CD55) or AAV-hSyn-DIO-mCherry (mCherry) viruses into the DG of c-Fos-CreER T2 mice 10 days before CFC training, and TAM was administered before the last training to induce the expression of CD55 in mCherry-only DG engram neurons (Fig. 2, G and H, and fig. S7). Thirty-five days after training, mice in the CD55 group showed higher freezing (Fig. 2D) and a higher reactivation rate (Fig. 2E) of engram cells (Fig. 2, J and K). Post hoc staining showed significantly fewer microglia containing mCherry+ puncta (Fig. 2, I and M) and smaller mCherry+ puncta within Iba-1+ microglia in the CD55 group of animals (Fig. 2N).

Connectivity between engram cells is essential for memories (26), whereas microglia-dependent synaptic elimination preferentially targets weak or less-active synapses (25). We next examined whether microglia-mediated forgetting depends on the activity of engram neurons. We trained c-Fos-CreER T2::hM4Di mice for contextual fear memory and administered TAM to induce the expression of inhibitory DREADD (designer receptors exclusively activated by designer drugs) receptor hM4Di in tagged engram cells. DREADD ligand clozapine-N-oxide (CNO) was administered every other day after CFC training to repetitively suppress the activity of tagged engram cells (Fig. 3F). Thirty-five days after training, mice treated with CNO alone exhibited significantly decreased freezing (Fig. 3B), whereas administration of PLX after training prevented the facilitated forgetting in CNO-treated animals (Fig. 3B).

To confirm this result, we injected AAV-hSyn-DIO-CD55-mCherry or AAV-hSyn-DIO-mCherry into the DG of c-Fos-CreER T2::hM4Di mice and trained them for conditioned contextual fear memory. TAM was administered to express both hM4Di and CD55/mCherry in the engram cells (Fig. 3C). The animals treated with CNO alone showed reduced freezing (Fig. 3D) and decreased reactivation rate of labeled engram cells (Fig. 3, E and F). Expression of CD55 in engram cells prevented the decrease of reactivation rate (CNO+mCherry versus CNO+CD55 t = 2.895, df = 6, P = 0.0275; CNO+mCherry versus CNO+CD55 t = 2.435, df = 7, P = 0.0451; CNO+CD55 versus CNO+CD55 t = 0.4375, df = 7, P = 0.6749).
MEM-treatment significantly increased the number of DCX+ cells in the DG, indicating enhanced neurogenesis (Fig. S8). We found significantly larger volumes of PSD95+Lamp1+ puncta within microglia in MEM-treated animals (Fig. 4, A and B). Additionally, MEM treatment facilitated forgetting after CFC training (Fig. 4, C and D), whereas administering PLX blocked MEM-facilitated forgetting (Fig. 4D) without altering the enhanced neurogenesis by MEM (fig. S8).

To further investigate whether microglia also contribute to neurogenesis-unrelated forgetting, we used a GFAP-TK transgenic mouse line, which expresses herpes simplex virus thymidine kinase (TK) under the control of the glial fibrillary acidic protein (GFAP) promoter. Administration of the antiviral drug ganciclovir (GCV) ablates only mitotic GFAP+ cells that express TK, thus depleting neurogenesis (29). To completely deplete integration of new neurons, we started treating TK+/− mice and their wild-type littermates (TK−/−) with GCV 4 weeks before CFC training and

Fig. 4. Microglia contribute to both neurogenesis-mediated and non–neurogenesis-mediated forgetting. 
(A) Superresolution images and 3D reconstruction showing the PSD95 “Lamp1” puncta in microglia in Ctrl and MEM-treated CX3CR1GFP/+ mice. Scale bars, 5 μm; white arrows indicate PSD95+Lamp1+ puncta in microglia. Insets are enlarged 3D reconstructions of PSD95+Lamp1+ puncta in microglia. Scale bars, 2 μm. (B) MEM-treatment increased the volume of PSD95+Lamp1+ puncta in each microglia. Ctrl n = 3 mice, MEM n = 3 mice; t = 2.774, df = 57, **P = 0.0075. (C) PLX and MEM treatment administered to mice. (D) Freezing of animals during the test 35 days after training. MEM+PLX− n = 10 mice, versus MEM−PLX+ n = 10 mice, t = 3.511, df = 18, **P = 0.0025; MEM−PLX− versus MEM+PLX+ n = 10 mice, t = 3.341, df = 18, **P = 0.0036; MEM+PLX+ versus MEM−PLX+ n = 10 mice, t = 2.277, df = 18, **P < 0.0001; MEM−PLX− versus MEM−PLX+ t = 2.072, df = 18, P = 0.0529. (E) GCV and PLX treatment administered to GFAP-TK+/− or GFAP-TK−/− mice. (F) Freezing during the test 35 days after training. TK−/− PLX+ n = 14 mice versus TK−/− PLX− n = 14 mice, t = 5.181, df = 26, ****P < 0.0001; TK−/− PLX− versus TK+/− PLX− n = 14 mice, t = 4.895, df = 26, ****P < 0.0001; TK+/− PLX+ versus TK+/− PLX− n = 13 mice, t = 2.333, df = 25, *P = 0.0280; TK+/− PLX− versus TK+/− PLX+ t = 2.2426, df = 25, *P = 0.0341. (G) c-Fos-CreERT2 mice received AAV injection into CA1 and recovered for 10 days before CFC training. TAM was administered before the last training, and freezing was tested 35 days later. (H) Confocal image showing engram cells (mCherry+) in CA1 but not in the DG. Scale bar, 100 μm. (I) CD55 animals showed higher freezing level. mCherry n = 11 mice, CD55 n = 11 mice; t = 2.728, df = 20, *P = 0.0130. (J) Confocal images showing the reactivation of engram cells in CA1. White arrows indicate a reactivated engram cell (mCherry+c-Fos+) in CA1. Inset shows the colocalization of mCherry and c-Fos; Scale bars, 20 μm. (K) Reactivation rate of engram cells in CA1 was increased in CD55 animals. mCherry n = 5 mice, CD55 n = 5 mice; t = 3.268, df = 8, *P = 0.0114.
continued until the test (Fig. 4E and fig. S9). PLX was administered after training to deplete microglia. GCV treatment in TK−/− mice prevented forgetting, whereas TK−/− mice treated with both PLX and GCV showed significantly higher levels of memory retention than TK−/− mice treated with GCV only (Fig. 4F).

To confirm that microglia-mediated forgetting contributes to neurogenesis-unrelated forgetting, we injected AAV-DIO-CD55-mCherry or AAV-DIO-mCherry into e-Fos-CreERT2 mice to label engram cells in hippocampal CA1 (Fig. 4, G and H), which is not a neurogenic region. We found that expression of CD55 in CA1 engram cells also prevented forgetting (Fig. 4I) and dissociation of engram cells (Fig. 4, J and K).

Synaptic connections in the brain are highly dynamic and variable in strength and connectivity (14). Our study shows that microglia eliminate synaptic components in the adult hippocampus, whereas depleting microglia or inhibiting phagocytosis of microglia prevents forgetting. This suggests that synapse elimination by microglia leads to dissociation of engrams and the forgetting of previously learned contextual fear memory. In the developing brain, microglial engulfment of synapses depends on the classical complement cascade (15). Disruption of the microglia-specific phagocytic pathway by knocking out complement components, such as C1q, C3, or CR3, results in sustained deficits in synaptic connectivity (15, 24). C1q levels in the brain increase during aging, whereas C1q-deficient mice exhibit enhanced synaptic plasticity and less cognitive and memory decline when aged (30). Notably, our study showed that the C1q-dependent complement pathway is actively involved in synapse elimination by microglia in the healthy adult hippocampus. CD55 is a known inhibitor of complement pathways in the immune system and is expressed in neurons in response to chronic inflammation (33). We overexpressed CD55 to inhibit the complement pathways, specifically in engram cells, without affecting microglia or other neurons in the circuits, and we found that forgetting was prevented. This indicates that the elimination of synaptic structures by microglia in the DG of the healthy adult brain occurs in a complement-dependent manner. Moreover, inhibiting the activity of engram cells facilitates the forgetting of related memory, which could be blocked by depleting microglia or inhibiting complement pathways in engram cells. This indicates that synapse elimination by microglia is also activity-dependent, following similar rules in the developing brain (15), thus resulting in the erasure of less-active memories. Besides eliminating synapses, microglia have also been reported to be able to trigger long-term synaptic depression via AMPA receptor internalization, through activation of CR3 (32), which may also contribute to forgetting.

New neurons are continuously generated in the DG, providing a substrate for massive synaptic reorganization and circuit rewiring in this region. Newborn dentate granule neurons integrate into hippocampal neural circuits by competitively replacing existing synaptic connections formed by mature granule neurons (7, 27), thus leading to the forgetting of hippocampal-contextual fearful memory (6). Our study shows that MEM-induced enhanced neurogenesis leads to increased synaptic engulfment by microglia, whereas depletion of microglia blocks facilitated memory forgetting induced by enhanced neurogenesis, suggesting that microglia contribute to neurogenesis-induced synaptic reorganization. Besides the rewiring of neural circuits caused by the continuous integration of newborn neurons, mature neurons are also able to reorganize their connectivity. We found that depletion of microglia in the DG without neurogenesis or inhibition of complement pathways in CA1 engram cells prevents forgetting. This indicates that microglia-mediated synaptic reorganization is also happening in mature hippocampal neurons, thus leading to weakening or loss of connections between engram cells and the forgetting of encoded memories. This also suggests that, in species lacking adult neurogenesis, or in non-neurogenic brain regions such as the cortex, microglia could be one major force contributing to synaptic loss and forgetting.

REFERENCES AND NOTES

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SUPPLEMENTARY MATERIALS
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Microglia modulate memories
Synaptic reorganization and circuit rewiring leads to loss or weakening of connections between neurons and may result in the erasure of previously formed memories. Microglia eliminate excessive synapses in the developing brain and regulate the dynamics of synaptic connections between neurons throughout life. However, it is still unclear whether forgetting is related to microglia activity and how microglia regulate memory erasure in the adult brain. Wang et al. discovered that microglia eliminated synaptic components in the adult hippocampus and that depleting microglia or inhibiting phagocytosis of microglia prevented forgetting. Synapse elimination by microglia may thus lead to degradation of memory engrams and forgetting of previously learned contextual fear memory.

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