Acetylation of calmodulin regulates synaptic plasticity and fear learning

Synaptic plasticity is critical for brain function, including learning and memory. It is regulated by gene transcription and protein synthesis as well as posttranslational modifications at synapses. Although protein acetylation has been shown to be involved in the regulation of synaptic plasticity, this was mainly for histone protein acetylation. To investigate whether acetylation of nonhistone proteins is important for synaptic plasticity, we analyzed mouse brain acetylome and found that calmodulin (CaM), a ubiquitous Ca2+ sensor, was acetylated on three lysine residues, which were conserved across species. NMDA receptor-dependent long-term potentiation (LTP) is considered the most compelling form of synaptic plasticity. During LTP induction, activation of NMDA receptor triggers Ca2+ influx, and the Ca2+ binds with CaM and activates calcium/calmodulin-dependent protein kinase IIα (CaMKIIα), which is essential for LTP induction. By using home-generated and site-specific antibodies against acetylated CaM, we show that CaM acetylation is upregulated by neural activities in an NMDA receptor-dependent manner. Moreover, mutation of acetyllysines in CaM1 proteins disrupts synaptic plasticity and fear learning in a mouse model. We further demonstrate that acetylation of CaM reduces the binding free energy and increases the binding affinity toward CaMKIIα, a protein kinase pivotal to synaptic plasticity and learning. Taken together, our results demonstrate importance of CaM acetylation in regulating synaptic plasticity and learning.

Synaptic plasticity is critical for several brain functions including learning and memory. It is regulated by long-term mechanisms such as gene transcription (1, 2), as well as acute mechanisms at synapses including local protein synthesis (3), neurotransmitter receptor trafficking (4–6), and protein phosphorylation (7, 8). Calcium signaling is essential for synaptic plasticity and is initiated by activation of ion channels such as NMDA receptor and voltage-dependent Ca2+ channels (VDCC) and is mediated through several Ca2+-binding proteins such as calmodulin (CaM), S100 family proteins, synaptotagmin (SYT), and protein kinase C (PKC) (9–13).

CaM is a ubiquitous Ca2+ sensor and has more than 300 target proteins including protein kinases, enzymes, cytoskeleton proteins, ion and water channels (14–19). For instance, CaM interacts with and activates Ca2+/calmodulin-dependent protein kinase IIα (CaMKIIα) and protein phosphatase calcineurin (CaN), which are important for several cellular functions including plasticity and learning in the brain (14, 20–23). In addition, the conformational change of CaM upon binding with Ca2+ has been utilized to generate fluorescence proteins to monitor the activity of excitable cells (24).

Protein acetylation is regulated by lysine acetyltransferase (KAT) and lysine deacetylase (KDAC) and was discovered initially as a posttranslational modification of histone proteins to activate gene transcription (25). In accord, histone protein acetylation has been implicated in diverse physiological conditions including synaptic plasticity and learning (26–30). On the other hand, acetylation of nonhistone proteins has been shown to regulate a plethora of biological processes such as DNA repair, cytoskeleton dynamics, cell metabolism, and autophagy (31–36). However, the roles of nonhistone protein acetylation in synaptic plasticity and learning are less well understood (37).

In analyzing the mouse brain acetylome, we found that CaM was heavily acetylated. By using home-generated and site-specific antibodies against acetylated CaM, we show that CaM acetylation is increased within minutes of LTP (long-term potentiation) induction, but not LTD (long-term depression), in a manner dependent on NMDA receptor activation. Mutation of acetyllysines in CaM1 proteins impairs hippocampal LTP and contextual fear learning. The mechanistic study indicates that acetylation of CaM reduces the binding free energy and increases the binding affinity toward CaMKIIα, a protein kinase pivotal to synaptic plasticity and learning. Together, our results reveal important roles of CaM acetylation in synaptic plasticity and learning.
**Acetylation of CaM in synaptic plasticity**

**Results**

**Increase of CaM acetylation by neural activities**

CaM is a ubiquitous calcium sensor and plays important roles in calcium signaling and synaptic plasticity (38–40).

Previous proteomic studies have identified CaM acetylation in several tissues including the blood, liver, heart, lung, and brain (33, 36, 41, 42). However, the function and regulation of CaM acetylation remain largely unknown. To investigate whether CaM acetylation is regulated by neural activities, we analyzed two paradigms—chemical LTP (cLTP) stimulation in vitro and contextual fear learning in vivo, both of which can activate calcium signaling through NMDA receptor (43, 44). When hippocampal slices were stimulated with 50 μM forskolin and 100 nM rolipram for 15 min, LTP can be induced at Schaffer collateral (SC)-CA1 synapses (Fig. 1A) (43). Acetylated proteins were precipitated with anti-Ac-K antibody and probed by anti-CaM antibody. cLTP-induced CaM acetylation (Ac-CaM) occurred within minutes of the stimulation, but returned to basal levels by 30 min after washout of cLTP stimulation (Fig. 1, B and C).

The potentiation effect of cLTP on CaM acetylation was specific because it was not observed in hippocampal slices.
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To study whether CaM acetylation is regulated by hippocampus-dependent contextual fear learning, mice were subjected to training (i.e., pairing with the US—electric shock and CS—contextual box) and 1 day later, freezing or fear memory was tested when mice were returned to the contextual box (Fig. 1G). Of mice in the test box without electric shock (CS, contextual box), hippocampal Ac-CaM was similar to control (Fig. 1, H and I). However, their levels in the hippocampus were upregulated after pairing CS with US, almost as high as in mice injected with TSA, a KDAC inhibitor (Fig. 1, H and I). Ac-CaM remained high when mice were exposed to a new environment (i.e., unpaired tests) (Fig. 1, H and I). By contrast, Ac-CaM returned to control levels after paired tests (i.e., exposure to contextual box where they received electric shock) (Fig. 1, H and I). These results suggest that CaM acetylation is increased during contextual fear learning.

Increased CaM acetylation on K22, 95, and 116 by neural activities

By analyzing the previous proteomic studies (45, 46) and publicly available database (https://www.phosphosite.org/curatedInfoAction?action?record=9921012), we found that in mouse brain, CaM was acetylated on three lysine residues (K22, K95, and K116) that are conserved across species (Fig. 2A). To determine which of the three lysine residues in CaM could be acetylated, we generated site-specific antibodies against Ac-CaM at K22, K95, and K116. To demonstrate the specificity of the anti-Ac-CaM antibodies, we purified site-specifically acetylated recombinant CaM proteins using the strategy of genetic code expansion (Fig. 2B). This strategy uses an engineered pyrolysyl-tRNA synthetase specific for Ac-K and its cognate tRNA<sup>Pys</sup> to incorporate Ac-K at an assigned codon to produce site-specifically acetylated proteins (47). As shown in Figure 2C, the pan-anti-Ac-K antibody can detect CaM acetylated at any of the three lysine residues. However, the anti-Ac-K22-CaM antibody only recognized CaM acetylated at K22, but not WT-CaM or CaM acetylated at K95 or K116, and vice versa (Fig. 2C). These results demonstrate the specificity of the anti-Ac-CaM antibodies.

We probed the total lysates of hippocampal slices with these antibodies. As shown in Figure 3, A–D, CaM acetylation at K22, K95, and K116 was increased by cLTP stimulation for 10 min. In addition, the induced acetylation was diminished by AP5, an antagonist of NMDA receptor, suggesting its dependence on NMDA receptor activation (Fig. 3, A–D). We used purified Ac-CaM proteins as standards to analyze the stoichiometry levels for each lysine residue under control and cLTP conditions. The stoichiometry levels of K22, K95, and K116 in the total lysates are 2.80%, 2.76%, 2.87% under basal conditions and are increased to 6.77%, 7.33%, 5.84% after cLTP stimulation (Fig. 3, E–G). The stoichiometry results showed that cLTP upregulated acetylation at individual lysine residue to 2- to 3-fold, which are consistent with the western blot results using the site-specific antibodies (Fig. 3, A–D). Taken together, these results demonstrate that CaM acetylation is upregulated by cLTP in a manner dependent on NMDA receptor activation.

To study whether the protein levels of Ac-CaM are increased in the synapse or nucleus after cLTP stimulation, we purified postsynaptic density (PSD) and nuclear fractions from hippocampal slices. Ac-CaM was upregulated in the PSD fraction of hippocampal slices after cLTP stimulation for 10 min, which was prevented by AP5 (Fig. 3, H–K). By contrast, acetylation of CaM was not elevated in the nucleus of hippocampal slices after cLTP stimulation for 10 min (Fig. S1). These results indicate that Ac-CaM was increased in the synapse after cLTP stimulation in an NMDAR-dependent way.

Next, we investigate whether CaM acetylation at the three lysine residues could be regulated by contextual fear learning. Mice were subjected to training (i.e., pairing with the US—foot shocks and CS—contextual box) and 1 day later, freezing or fear memory was tested when mice were returned to the contextual box (Fig. 1G). Protein levels of Ac-CaM at K22, 95,
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and 116 in the hippocampus were upregulated after pairing CS with US, compared with control mice in the contextual box without foot shocks (Fig. 3, L–O). By contrast, protein levels of Ac-CaM at K22, 95, and 116 returned to control levels after paired tests (i.e., exposed to contextual box where they received foot shocks) (Fig. 3, L–O). These results suggest that contextual fear learning increases the acetylation of CaM at the three lysine residues.

Increase of CaMKIIα activity by acetylation-mimicking CaM

To determine whether acetylation alters the interaction between CaM and CaMKIIα, we generated 3KR-CaM and 3KQ-CaM where the three lysine residues were mutated to arginine and glutamine, respectively to block and mimic acetylation (41, 48). K to Q mutation neutralizes the positive charge of K and is commonly used to be a mimic of acetyllysine (41, 48). We used 1 μM GST-CaM and His-CaMKIIα for the GST-pull-down assay in the presence of 0.1 mM Ca2+. We finished the pull-down assay within 2 h but not overnight to avoid the saturation of CaMKIIα binding due to the long-time incubation. As shown in Figure 4A, GST-tagged WT-CaM was able to pull down His-CaMKIIα in a calcium-dependent manner. However, this interaction was inhibited by the KR mutation, but potentiated by the KQ mutation (Fig. 4, A and B). These results suggest that acetylation-resistant CaM reduces while acetylation-mimicking CaM enhances the interaction with CaMKIIα, compared with WT-CaM.

Next, we determine whether acetylation-mimicking CaM could enhance the kinase activity of CaMKIIα by an assay of in vitro CaMKIIα autophosphorylation—with puriﬁed His-CaMKIIα and GST-tagged WT or mutant CaM. We used 1 μM GST-CaM and His-CaMKIIα for the assay in the presence of 0, 0.004, or 0.1 mM Ca2+. Autophosphorylated CaMKIIα was detected with phospho-speciﬁc antibody against Thr286, whose phosphorylation is an indicator of CaMKIIα activation (49). CaMKIIα became activated by incubation with GST-CaM in a Ca2+-dependent manner (Fig. 4C). Remarkably, p-CaMKIIα was reduced by 3KR-CaM, but increased by 3KQ-CaM (Fig. 4, C–F). Note that acetylation-mimicking CaM had no effects on the interaction with CaN or its phosphatase activity (Fig. S2), indicating the speciﬁcity of CaM acetylation to

Figure 3. Increase of CaM acetylation on K22, 95, and 116 by neural activities. A, increased CaM acetylation at different lysine residues 10 min after cLTP stimulation in hippocampal slices without or with AP5. B–D, quantification of Ac-K22-CaM/CaM (B), Ac-K95-CaM/CaM (C), and Ac-K116-CaM/CaM (D) in panel A. Data were represented as mean ± SD. NS, not significant. ***p < 0.0001, compared with controls, one-way ANOVA, n = 6, data were normalized to control. E–G, stoichiometry levels of Ac-K22-CaM (E), Ac-K95-CaM (F), and Ac-K116-CaM (G) from control and cLTP-stimulated hippocampal slices. Data were represented as mean ± SD. ***p = 0.0008, **p = 0.0023, *p = 0.01, t test, n = 3. H, increased CaM acetylation at different lysine residues 10 min after cLTP stimulation in PSD fraction without or with AP5. The homogenates of hippocampal PSD fraction were probed with the indicated antibodies. I–K, quantification of Ac-K22-CaM/CaM (I), Ac-K95-CaM/CaM (J), and Ac-K116-CaM/CaM (K) in panel H. Data were represented as mean ± SD. NS, not significant, ***p < 0.0001, compared with controls, one-way ANOVA, n = 6, data were normalized to control. L, increased CaM acetylation at different lysine residues after pairing US and CS. The total lysates of hippocampus were probed with the indicated antibodies. M–O, quantification of Ac-K22-CaM/CaM (M), Ac-K95-CaM/CaM (N), and Ac-K116-CaM/CaM (O) in panel L. Data were represented as mean ± SD. NS, not significant, ***p < 0.0001, compared with controls, one-way ANOVA, n = 6, data were normalized to control.
activate CaMKIIα. Taken together, these results demonstrate that acetylation-resistant CaM reduces while acetylation-mimicking CaM increases the kinase activity of CaMKIIα, compared with WT-CaM.

To explore how acetylation-mimicking CaM affects the interaction with CaMKIIα, we performed bioluminescence resonance energy transfer (BRET) analysis. HEK293 cells were transfected with a BRET construct encoding Venus (a yellow fluorescent protein), CaM, C18 (a CaM binding peptide from CaMIIα), and RLuc8 (enhanced Renilla luciferase) (Fig. 4G) (modified from a previous report (50)). Compared with WT-CaM (Kd = 3.92 ± 0.08 μM), 3KR-CaM had a right-shifted binding curve and a higher Kd value (14.41 ± 0.98 μM) while 3KQ-CaM exhibited a left-shifted binding curve and a lower Kd value (1.39 ± 0.01 μM) (Fig. 4, H and I). These results indicate that 3KR-CaM decreases while 3KQ-CaM increases the binding affinity toward CaMKIIα, compared with WT-CaM. By contrast, 3KR and 3KQ-CaM show the similar binding affinity toward myosin light chain kinase (MLCK) in the BRET assay (Fig. S3), demonstrating the specificity on CaMKIIα interaction. Altogether, these data demonstrate that acetylation-mimicking CaM has a higher binding affinity with CaMKIIα, compared with WT-CaM.

Promotion of CaMKIIα activity by acetylated CaM

To further study whether CaM acetylation promotes the interaction with CaMKIIα, we generated site-specifically acetylated recombinant CaM proteins using the strategy of genetic code expansion (Fig. 2B). We purified His-tagged WT-CaM and acetylated CaM at the three lysine residues (Ac-3K-CaM) (Fig. 5A). Note that WT-CaM purified from bacteria could not be acetylated (Figs. 2C and 5A). His-tagged WT or Ac-3K-CaM was incubated with Flag-CaMKIIα purified from HEK293 cells for coimmunoprecipitation (Co-IP) experiments. Similar with the GST pull-down assay with 3KQ-CaM, we used 1 μM His-CaM and Flag-CaMKIIα for the Co-IP experiments in the presence of 0.1 mM Ca2+. We finished the Co-IP experiments within 2 h but not overnight to avoid the saturation of CaMKIIα binding due to the long-time incubation. As shown in Figure 5B, WT-CaM was able to interact with Flag-CaMKIIα in the presence of 0.1 mM Ca2+. However, this interaction was potentiated by Ac-3K-CaM (Fig. 5, B and C), suggesting that CaM acetylation promotes the CaM-CaMKIIα interaction.

We next determine whether acetylated CaM increases the kinase activity of CaMKIIα by an assay of in vitro CaMKIIα autophosphorylation—with purified Flag-CaMKIIα and different forms of His-CaM. Similar with the assay with 3KQ-CaM, we used 1 μM His-CaM and Flag-CaMKIIα for the assay in the presence of 0.004 or 0.1 mM Ca2+. Autophosphorylated CaMKIIα was detected with phospho-specific antibody against Thr286, whose phosphorylation is an indicator of CaMKIIα activation (49). CaMKIIα became activated by incubation with WT-CaM in the presence of Ca2+ (Fig. 5, D–G). Remarkably, p-CaMKIIα was significantly increased by Ac-3K-CaM (Fig. 5,
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**Figure 5.** Promotion of CaMKII activity by acetylated CaM. A, purification of His-tagged WT and acetylated CaM proteins from bacteria. Top, Coomassie blue staining of 10 µg His-tagged WT and acetylated CaM proteins, bottom, western blot of the purified proteins with the indicated Abs. B, increased CaMKIIα interaction by acetylated CaM. Flag-CaMKIIα was purified from HEK293 cells and was incubated with His-tagged WT or acetylated CaM to test the interaction in the presence of 0.1 mM Ca2+. CaM was immunoprecipitated with anti-His antibodies and then immunoblotted with anti-Flag antibodies. C, quantification of the interaction between WT or acetylated CaM and CaMKIIα in panel B. Data were represented as mean ± SD. **p < 0.0001, t test, n = 6, data were normalized to WT-CaM. D, enhanced CaMKIIα activity by acetylated CaM. Purified Flag-CaMKIIα and His-tagged WT or acetylated CaM were incubated in the presence of 0.1 mM Ca2+; CaMKIIα activity was revealed by anti-p-CaMKIIα. E, quantification of p-CaMKIIα/CaMKIIα in panel D. Data were represented as mean ± SD. **p < 0.0001, t test, n = 6, data were normalized to WT-CaM. F, p-CaMKIIα was significantly increased by Ac-3K-CaM compared with WT-CaM in the presence of 4 µM Ca2+. G, quantification of p-CaMKIIα/CaMKIIα in panel F. Data were represented as mean ± SD. **p < 0.0001, t test, n = 6, data were normalized to WT-CaM. H, structural comparison of the dominant conformations in CaM-CaMKIIα and Ac-CaM-CaMKIIα simulation system (adopted from PDB code 1CM4). CaM is colored in gray. Target peptide from CaMKIIα interacted with CaM and Ac-CaM, which are displayed in green and blue, respectively. Acetylated residues K21, K94, and K115 are highlighted with sticks. I, free energy (kcal/mol) analysis for the interaction between WT or acetylated CaM and C18 from CaMKIIα. Data were presented as mean (SEM). ΔE<sub>ele</sub>, energy contribution from electrostatic force; ΔE<sub>solv</sub>, energy contribution from van der Waals force; ΔG<sub>binding</sub>, total energy change for the protein–protein interaction process; ΔG<sub>solv</sub>, energy contribution from total solvation-free energy.

D–G), suggesting that CaM acetylation promotes CaMKIIα activation. Taken together, these results suggest that acetylated CaM increases the kinase activity of CaMKIIα, compared with WT-CaM.

In the following experiment, we study how acetylation of CaM promotes its interaction with CaMKIIα. Analysis of a 3-D CaM-CaMKIIα complex (Protein Data Bank, ID: 1CM4) from two simulation trajectories suggests that acetylation may increase the flexibility of CaM, enabling a tighter wrapping around CaMKIIα (Fig. 5H). Three lysine residues adopted different conformations after acetylation. K21 and K115 (no counting the first methionine) underwent structural reorientations and projected above the binding groove of CaM-KIIα. Given their close proximity and favorable topology, K21 and K115 might be closely implicated in the protein–protein interaction. For K94, allosteric effects might be involved in its impact on the CaM-CaMKIIα interaction upon acetylation, given its relatively distal location to the binding pocket.

To further study the impact of CaM acetylation on its interaction with CaMKIIα, we analyzed the free energy (ΔG<sub>binding</sub>) of the CaM-CaMKIIα interaction by using Amber package (51). The ΔG<sub>binding</sub> was calculated by molecular dynamics (MD) simulation of the CaM-CaMKIIα complex (see Experimental procedures). As shown in Figure 5I, upon acetylation of CaM, ΔG<sub>binding</sub> for CaMKIIα toward CaM decreased from ~75 ± 0.2 kcal/mol to ~89 ± 0.13 kcal/mol, rendering the binding event more favorable. ΔG<sub>binding</sub> is regulated by three parameters—ΔE<sub>ele</sub> (van der Waals force), ΔE<sub>solv</sub> (electrostatic force), and ΔG<sub>solv</sub> (solvation-free energy). Our result revealed a reduced ΔE<sub>ele</sub> by CaM acetylation, suggesting a role of the electrostatic force in promoting the CaM-CaMKIIα interaction (Fig. 5I).

**Generation and characterization of 3KR-Cam1 knockin mice**

The K to R mutation is commonly used as a dominant negative mutant for protein acetylation because R preserves the positive charge on the side chain (similar to K), but cannot be acetylated (41, 52). To investigate whether acetylation of CaM on the three lysine residues is important for LTP, we used CRISPR-Cas9 technique to generate mutant...
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Figure 6. Generation and characterization of 3KR-Cam1 knockin mice. A, schematic overview of generating 3KR-Cam1 knockin mice using Tild-CRISPR. The knockin fragment was composed of the DNA encoding the 22 to 149 amino acid of 3KR-CaM (green box) and the BGH polyA (red box) together with the homology arms. The knockin fragment was inserted in exon 3 of mouse Cam1 gene. HAL or HAR, left or right homology arm. B, sequence analysis of 3KR-Cam1 knockin mice. DNA of mouse tails from 3KR-Cam1 mouse was isolated. PCR products amplified from 5' and 3' junction sites were sequenced. C, transcriptional levels of Cam1 are higher than those of Cam2 and Cam3 in CA1 pyramidal neurons of mouse hippocampus (n = 9589 cells). Shown are transcripts of Cam genes per 100k total transcripts from single-cell RNA sequencing. D, lung hemorrhage in 6-week-old 3KR/3KR mice. E, normal brain weight from 5-week-old 3KR/3KR mice. Scale bar, 5 mm. NS, not significant, t-test, n = 6. F, normal global morphology of hippocampus for 5-week-old 3KR/3KR mice. The hippocampal slices were stained with anti-NeuN and DAPI. Scale bar, 400 μm. G, similar input–output (I/O) curve at SC-CA1 synapses between WT and 3KR/3KR mice. Data were represented as mean ± SD. F (1,23) = 0.0266, p = 0.8719, two-way ANOVA, n = 13 slices from four WT mice, n = 12 slices from four 3KR/3KR mice.

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Hippocampal slices from WT and 3KR/3KR mice were subjected to LTP stimulation. Autophosphorylated CaMKIIα was detected with phospho-specific antibody against Thr286, whose phosphorylation is an indicator of CaMKIIα activation (49). As shown in Figure 7, A–C, LTP-induced CaM acetylation and CaMKIIα activation was diminished in hippocampal slices from 3KR/3KR mice, compared with WT littermates. Note that Ac-CaM was not abolished in 3KR/3KR mice probably because of the intact of Cam2 and Cam3 genes. CaMKIIα could phosphorylate the AMPA receptor subunit GluR1 at Ser831 during LTP (55, 56). Consistent with the reduction of CaMKIIα activity, p-GluR1 Ser831 also decreased in LTP in 3KR/3KR mice (Fig. 7, A and D). These results suggest that acetylation of CaM is important for calcium signaling activated by LTP stimulation.

We further study whether acetylation of CaM is important for LTP. We performed field potential recording in hippocampal slices from 5-week-old WT and 3KR/3KR mice (Fig. 7E). The glutamate release reflected by the paired pulse facilitation (PPF) at SC-CA1 synapses was not altered in 3KR/3KR mice (Fig. 7F). However, LTP at SC-CA1 synapses in response to theta burst stimulation (TBS) was greatly attenuated in 3KR/3KR mice, compared with WT littermates (Fig. 7, G and H). To exclude any potential developmental effects mediated by 3KR/3KR mutant, we studied whether acute delivery of 3KR-CaM recombinant proteins in WT mice could impair LTP. To this end, we purified GST-tagged WT, 3KR, or 3KQ-CaM (mimic acetylation) proteins from bacteria (Fig. 7I). The GST-tagged CaM proteins were delivered into hippocampal CA1 neurons through recording peptide, and 30 min later, the high-frequency stimulation (HFS)-induced LTP was recorded (Fig. 7J). As indicated in Figure 7, K and L, 3KR-CaM but not 3KQ-CaM inhibited LTP in hippocampal CA1 neurons from whole-cell recording. Altogether, these results demonstrate an important role of CaM acetylation in LTP.
Importance of CaM acetylation in contextual fear learning

The LTP of SC-CA1 synapses is coupled to contextual fear learning (57, 58), which could increase CaM acetylation in the hippocampus (Fig. 3, L–O). Lastly, we study whether CaM acetylation plays an important role in contextual fear learning. Because 3KR/3KR mice died prematurely prior to adulthood, due to lung hemorrhage (Fig. 6D), we took a knockdown-plus-rescue strategy in adult mice for the behavioral study (Fig. 8A). Endogenous CaM was downregulated by lentivirus expressing shRNAs against Cam1, 2, and 3, and at the same time, the shRNA-resistant WT-CaM, 3KR-CaM, or 3KQ-CaM was expressed in the hippocampus (Fig. 8, B–D). No difference was observed in locomotor activity in open fields among these different groups (Fig. 8, E and F). Two weeks after injection of lentivirus into hippocampus, mice were subjected to contextual fear training and tests (Fig. 8G). In mice where Cam1, 2, and 3 were knocked down and WT-CaM was expressed (i.e., Camkd+WT-CaM), freezing was normal in exploring, training as well as tests, compared with untreated controls (Fig. 8H). In mice where Cam1, 2, and 3 were knocked down and 3KR-CaM was expressed (i.e., Camkd+3KR-CaM), freezing was normal in exploring and training (Fig. 8H). However, their freezing during tests was reduced, compared with controls (Fig. 8H), suggesting that CaM acetylation is important for contextual fear learning. On the other hand, Camkd+3KQ-CaM mice displayed freezing similar to that of control or Camkd+WT-CaM mice (Fig. 8H). These results provide evidence that CaM acetylation is important for contextual fear learning.

Figure 7. Importance of CaM acetylation in hippocampal LTP. A, reduced Ac-CaM, p-CaMKIIα, and p-GluR1 in 3KR/3KR hippocampal slices, in response to cLTP stimulation. B–D, quantification of Ac-CaM/CaM (B), p-CaMKIIα/CaMKIIα (C), and p-GluR1/GluR1 (D) in panel A. Data were represented as mean ± SD. ***p < 0.0001, two-way ANOVA followed by Tukey’s multiple comparisons test, n = 6, data were normalized to WT slices under control condition. E, diagram showing field EPSP recording at SC-CA1 synapses from WT and 3KR/3KR mice. F, comparable paired pulse facilitation (PPF) at SC-CA1 synapses between WT and 3KR/3KR mice. Data were represented as mean ± SD. F (1,19) = 0.304, p = 0.5876, two-way ANOVA, n = 10 slices from four WT, n = 11 slices from four 3KR/3KR mice. G, normalized fEPSP amplitudes were plotted every 1 min for hippocampal slices from WT and 3KR/3KR mice. H, reduced TBS-induced LTP at SC-CA1 synapses in 3KR/3KR hippocampal slices, compared with WT. Data in panel G were quantified. Data were represented as mean ± SD. **p = 0.0021, t test, n = 11 slices from six WT mice, n = 15 slices from eight 3KR/3KR mice. I, Coomassie blue staining of 10 μg GST-tagged WT, 3KR, and 3KQ-CaM proteins purified from bacteria. J, diagram showing whole-cell recording of eEPSC in hippocampal CA1 pyramidal neurons. Recording pipettes were infused with GST-WT, 3KR, or 3KQ-CaM. K, normalized eEPSC amplitudes were plotted every 1 min for CA1 pyramidal neurons infused with 100 nM GST-WT, 3KR, or 3KQ-CaM. L, quantification of LTP in panel K. Data were represented as mean ± SD. NS, not significant, ***p = 0.0001, one-way ANOVA, n = 9 cells from nine mice for WT and 3KQ-CaM, n = 10 cells from ten mice for 3KR-CaM.
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Discussion

In this study, we provide evidence that acetylation of CaM can be regulated by neural activities and is important for synaptic plasticity and fear learning. Acetylation of CaM occurred within minutes of cLTP stimulation in hippocampal slices, in a time course similar to that of CaMKIIa activation during cLTP stimulation in cultured neurons (59). Mutation of the lysine residues to prevent CaM acetylation impairs hippocampal LTP and contextual fear learning. A parsimonious explanation of these results is that CaM acetylation plays a role in regulating synaptic plasticity.

The stoichiometry level (6%–7%) of CaM acetylation in the total lysates of hippocampus was similar to those of bona fide acetylated proteins in mammalian cells (60). The calcium elevation during LTP induction is localized to stimulated spines, or a region called calcium nanodomain, which is near the inner mouth of postsynaptic NMDA receptor (21). The data presented here and in the accompanying paper (61) demonstrated that neural activities increased CaM acetylation through an NMDA receptor and calcium–dependent manner. For these reasons, one could speculate that the increased acetylation of CaM during LTP induction mainly occurred in the stimulated spines or calcium nanodomain. Note that 6% to 7% of CaM acetylation in the stimulated neurons was from the total lysates rather than from the stimulated spines or calcium nanodomain. One could argue that the stoichiometry level of CaM acetylation in the stimulated spines or calcium nanodomain might be much higher than that in the total lysates. Future studies are warranted to generate new tools to study CaM acetylation in the stimulated spines or calcium nanodomain.

How neural activities increase CaM acetylation is an intriguing question. Here we showed that neural activities enhance CaM acetylation through NMDAR activation. Identification of the CaM acetyltransferase will be important for understanding how neural activities increase CaM acetylation. Regardless, our results indicate that CaM acetylation is important for the full activation of CaMKIIa and phosphorylation of GluR1 during LTP. In addition to the GluR1 subunit of AMPA receptor, CaMKIIa can interact with and phosphorylate a variety of synaptic proteins, which contribute to synaptic plasticity and learning (62–65). It will be interesting to study whether CaM acetylation regulates phosphorylation of other downstream target proteins of CaMKIIa.

The results presented here addressed the roles of nonhistone protein acetylation in synaptic plasticity. NMDAR-dependent LTP is arguably the most compelling form of synaptic plasticity (4, 20, 66). NMDAR-dependent LTP operates widely in the brain, but is most extensively studied at Schaffer collateral (SC)-CA1 synapses in the hippocampus (4, 20, 66). The LTP at SC-CA1 synapses in the hippocampus is linked with contextual fear memory (57, 58). These are the reasons why we choose hippocampus in this study. In addition to hippocampus, other brain regions such as amygdala and the hippocampus–amygdala circuit can also regulate contextual fear memory (67–69). Likewise, the LTP in the amygdala also plays important roles in fear memory (69, 70). Future studies are required to investigate whether CaM acetylation can...
regulate synaptic plasticity in other brain regions besides hippocampus.

Ideally, a loss-of-function approach could be informative for studying CaM acetylation. However, mice have three Cam genes—Cam1, Cam2 and Cam3—that encode completely identical proteins, in addition to more distantly related genes (53). Such genetic complexity hinders genetic targeting for all Cam genes at the same time. Recent study from single-cell RNA sequencing in adult mouse hippocampus revealed that the gene expression levels of Cam1 were higher than that of Cam2 and Cam3 in CA1 pyramidal neurons (54). Due to these reasons, we generated 3KR/3KR mutant in Cam1 gene to study the roles of CaM acetylation in LTP at SC-CA1 synapses. Nonetheless, acetylation of Cam2 and Cam3 may remain and account for the residual acetylation in the 3KR/3KR mutant mice. In addition to CaM, the recent studies suggest that LTP is also regulated by other Ca2+ sensors such as synaptotagmins (5), which might explain why LTP is not completely abolished in 3KR/3KR mice. K to Q mutation is commonly used to be a mimic of acetyllysine (41, 48), and thus 3KQ-CaM displayed potent activity to stimulate CaMKIIα. However, the CaM acetylation is already very high after cLTP stimulation, which might explain why 3KQ-CaM could not further increase LTP.

Acetylation of CaM in synaptic plasticity

CaM, as a calcium sensor, regulates synaptic plasticity by activating kinases such as CaMKIIα and phosphatases including CaN. We provided evidence that acetylated CaM promotes the interaction and activation of CaMKIIα, but not CaN, compared with WT-CaM. In addition to CaMKIIα and CaN, CaM has many other target proteins, several of which have been implicated in synaptic plasticity, neurotransmission, or neuronal excitability (14). For example, CaM regulates NR1 subunit (15), L-type calcium channel (16), cGMP-gated cation channel (71), and other ion channels (17). We observed lung hemorrhage in the 3KR/3KR mice. It might be possible that CaM acetylation affected the interaction and/or function of proteins critical for the integrity of endothelial cells in the pulmonary blood vessels. Future work is warranted to investigate whether CaM acetylation alters the function of these proteins.

Experimental procedures

Animals

C57BL/6N male mice at age of 7 to 8 weeks were used in experiments unless otherwise described. Animals were housed in rooms at 23 °C and 50% humidity in a 12 h light/dark cycle and with food and water available ad libitum. All experimental procedures were approved by the Institutional Animal Care and Use Committees of East China Normal University.

K22, 95, 116R (3KR) knockin mice were generated on C57BL/6 background using Tild-CRISPR (targeted integration with linearized dsDNA-CRISPR) (72). The Tild donor vector for 3KR-Cam1 contained 35 bp HAL-3KR-Cam1(22–149 amino acid)-BGH polyA-35 bp HAR. The exon3 of mouse Cam1 gene was replaced by the Tild donor vector. Two gRNAs targeting Cam1 exon 3 (gRNA-L: ttctccctattcgataaa-gatgg, gRNA-R: ttgagaaactgagtgcaccaggg) were transcribed in vitro. A mixture of Cas9 mRNA (100 ng/ml), sgRNAs (50 ng/ml), and donor vectors (50 ng/ml) was injected into fertilized eggs. A total of 694 injected zygotes were transferred into recipients, and 124 pups were obtained and genotyped. The PCR amplification of the targeting loci and DNA sequencing validated the correct gene targeting. The 3KR-Cam1 founder mouse was crossed with WT C57BL/6 mice for three generations, and the F4 homozygous 3KR-Cam1 knockin (3KR/3KR) mice were used for experiments. Since most 3KR/3KR mice died at 6 weeks old due to the lung hemorrhage, 5-week-old 3KR/3KR male mice were used in LTP experiments. The genotyping primers for WT and 3KR-Cam1 knockin mice are as follows: forward: 5’TTTAT TGTTAC CTGGT AAATC GG 3’, reverse: 5’ACCAA TCCAG GAAAT GCTCT 3’. The genotyping products for WT and 3KR-Cam1 are 646 and 1115 bp, respectively.

Detection of CaM acetylation

Because the CaM monoclonal antibody was not suitable for IP, we opted to purify acetylated proteins. Lysates were incubated with acetyllysine antibody-conjugated agarose and transferred to nitrocellulose membranes, which were incubated in the TBS containing 0.1% Tween-20 and 5% milk for 1 h at room temperature before incubation with a primary antibody overnight at 4 °C. After wash, the membranes were incubated with HRP-conjugated secondary antibody in the same TBS buffer for 1 h at room temperature. Immunoreactive bands were visualized by ChemiDocTM XRS + Imaging System (BIO-RAD) using enhanced chemiluminescence (Pierce) and analyzed with Image J (NIH). The primary antibodies used were as follows: anti-Histone H3, Cell Signaling (9715); anti-Calmodulin, Millipore (05-173); anti-CaMKIIα, Cell Signaling (11945); anti-p-CaMKIIα Thr286, Sigma (SAB4300228); anti-Acetyllsine, Cell Signaling (9441); anti-p-GluR1 Ser 831, Abcam (ab109464); anti-GluR1, Abcam (ab109450); anti-GST, Abmart (12G8); and anti-His, Abmart (10E2).

Western blot

Homogenates of hippocampal tissue were prepared in RIPA buffer containing 50 mM Tris–HCl, pH 7.4, 150 mM NaCl, 2 mM EDTA, 1% sodium deoxycholate, 1% Triton X-00, 1 mM PMSF, 50 mM sodium fluoride, 1 mM sodium vanadate, 1 mM DTT, and protease inhibitors cocktails. PSD fractions were directly dissolved in 1x SDS-PAGE sample buffer. All the protein samples were boiled in 100 °C water bath for 10 min before western blot. The homogenates were resolved on SDS-PAGE and transferred to nitrocellulose membranes, which were incubated in the TBS containing 0.1% Tween-20 and 5% milk for 1 h at room temperature before incubation with a primary antibody overnight at 4 °C. After wash, the membranes were incubated with an HRP-conjugated secondary antibody in the same TBS buffer for 1 h at room temperature. Immunoreactive bands were visualized by ChemiDocTM XRS + Imaging System (BIO-RAD) using enhanced chemiluminescence (Pierce) and analyzed with Image J (NIH). The primary antibodies used were as follows: anti-Histone H3, Cell Signaling (9715); anti-Calmodulin, Millipore (05-173); anti-CaMKIIα, Cell Signaling (11945); anti-p-CaMKIIα Thr286, Sigma (SAB4300228); anti-Acetyllsine, Cell Signaling (9441); anti-p-GluR1 Ser831, Abcam (ab109464); anti-GluR1, Abcam (ab109450); anti-GST, Abmart (12G8); and anti-His, Abmart (10E2).
immunized with 500 µg of peptides C-FSLFD(Ac-K)DGDGT, C-FRVFD(Ac-K)DGNGY and C-TNLGE(Ac-K)LTDEE with complete Freund’s adjuvant three times, and boosted with 250 µg peptides with incomplete Freund’s adjuvant additional four times. Rabbit anti-serum was collected and purified by respective nonacetylated peptides.

**Subcellular fractions**

Mouse brain tissues were homogenized in Buffer A (0.32 M sucrose, 1 mM MgCl₂, 1 mM PMSF and a protease inhibitor cocktail). Homogenates were passed through a filter to remove cell debris and centrifuged at 500g for 5 min in a fixed angle rotor to yield P1 and S1 fractions. P1 fractions were washed in Buffer B containing 10 mM KCl, 1.5 mM MgCl₂, 10 mM Tris-HCl (pH 7.4) and centrifuged at 500g for 5 min. Pellets were dissolved in Buffer C containing 20 mM HEPES (pH 7.9), 25% glycerol, 1.5 mM MgCl₂, 1.4 M KCl, 0.2 mM EDTA, 0.2 mM PMSF, 0.5 mM DTT and incubated on a shaker at 4 °C for 30 min. After centrifugation at 12,000 g for 10 min, the supernatant of P1 was collected as nuclear proteins. The S1 fraction was centrifuged at 10,000g for 10 min to yield P2 that contains membranes and synaptosomes and the cytoplasmic S2.

P2 fractions were resuspended in 0.32 M sucrose, which was then layered onto 0.8 M sucrose. After being centrifuged at 9100g for 15 min in a swinging bucket rotor, synaptosomes were collected from 0.8 M sucrose layer and concentrated by centrifugation at 20,800g for 1 h. To further purify the post-synaptic (PSD) fractions, synaptosomes in the 0.8 M sucrose solution were mixed with 1/19 volume of Buffer D containing 200 mM HEPES (pH 7.0), 20% Triton X-100, and 1.5 M KCl. Samples were centrifuged at 20,800g for 45 min using a fixed angle rotor. The resulting pellets were resuspended in Buffer E containing 1% Triton X-100 and 75 mM KCl using a Dounce mini-homogenizer and centrifuged again at 20,800g for 30 min to yield final pellets (PSD fraction), which were washed with 20 mM HEPES (pH 7.9) and dissolved in 1× SDS-PAGE sample buffer.

**Chemical LTP and LTD**

cLTP stimulation in hippocampal slices was performed as previously described (43). Briefly, cLTP was induced by incubating slices for 15 min in aCSF lacking MgCl₂ and containing 4 mM CaCl₂, 100 µM picrotoxin, 50 µM forskolin, and 100 nM rolipram. For western blots, the brain slices were collected 1 or 10 min after cLTP stimulation. cLTD in hippocampal slices was induced by submerging the slices in 30 µM NMND A for 3 min, as previously described (73). Slices were transferred to a well containing standard aCSF solution.

**Immunofluorescence**

Brain slices were fixed in 4% PFA, permeabilized with 0.3% Triton-X 100 and 5% BSA in PBS, and incubated with primary antibodies at 4 °C overnight. After washing with PBS for three times, samples were incubated with Alexa Fluor-594 secondary antibodies (1:1000, Invitrogen) for 1 h at room temperature. Samples were mounted with Vectashield mounting medium (Vector) and images were taken by Leica TCS SP8 confocal microscope. The following primary antibody was used: rabbit anti-NeuN (Abcam, ab177487).

**Acetylation of CaM in synaptic plasticity**

Samples were mounted with Vectashield mounting medium (Vector) and images were taken by Leica TCS SP8 confocal microscope. The following primary antibody was used: rabbit anti-NeuN (Abcam, ab177487).

**Purification of site-specifically acetylated CaM recombinant proteins**

The site-specifically acetylated CaM recombinant proteins were synthesized according to a previous report (47). In brief, *Escherichia coli* strain, BL21 (DE3), was transformed with plasmids pAcKRS-3 and pCDF PyrT-1 carrying the ORF for Cam with an amber codon at the desired site. The cells were first grown overnight in LB medium supplemented with 50 mg/ml kanamycin and 50 mg/ml spectinomycin (LB-KS) at 37 °C. Two-milliliter bacteria were cultured overnight and then inoculated into 200 ml LB-KS for further culturing. When the OD600 reached 0.4–0.6, 20 mM nicotinamide (NAM) and 10 mM acetyl-lysine were added, and 30 min later, the protein expression was induced at 18 °C overnight by adding 0.5 mM IPTG. Cells were harvested after induction and were washed with ice-cold PBS containing 20 mM NAM, the proteins were purified with HisTrap FF (GE Healthcare, 17-5319-01) according to the manufacturer’s protocol.

**Stoichiometry of CaM acetylation at K22, 95, and 116**

We used purified acetylated CaM proteins as standards to analyze the stoichiometry levels for each acetylated lysine residue of CaM under control and cLTP conditions. The standard samples containing 0%, 0.1%, 0.3%, 1%, 3%, and 10% acetylated His-CaM proteins in 1 µg total His-CaM proteins were subjected to ELISA assay in 96-well microplates, which were coated with 100 µl site-specific anti-Ac-CaM antibodies (0.01 µg/ml) overnight. The hippocampal lysates with 100 µg proteins were used for the same ELISA assay considering that the endogenous CaM proteins were about 1% of total proteins in the hippocampus (74). The standard samples and hippocampal lysates with the volume of 100 µl were added to each well and were incubated for 2 h at room temperature, and then each well was washed with PBS containing 0.1% Tween-20 for three times. After washing, 100 µl secondary antibodies (HRP-conjugated goat-anti-rabbit IgG, 1 µg/ml) was added to each well and incubated for 2 h at room temperature. After that, each well was washed with PBS containing 0.1% Tween-20 for three times before adding 100 µl HRP substrate solution (1:1 mixture of Color Reagent A H₂O₂ and Color Reagent B Tetramethylbenzidine, R&D Systems, Catalog # DY999) into each well. After incubation with the HRP substrate solution for 20 min at room temperature, 50 µl of stop solution (2 N H₂SO₄, R&D Systems, Catalog # DY994) was applied to each well. The optical density of each well was determined using a microplate reader set to 450 nm. We first generated a standard curve for stoichiometry of standard samples containing different concentration of Ac-CaM and then determined the stoichiometry levels of endogenous Ac-CaM proteins in the hippocampal lysates.
Acetylation of CaM in synaptic plasticity

**GST pull-down**

GST-tagged WT-CaM or CaM with K to R mutation was expressed in *E. coli* BL21 cells and purified using Glutathione Sepharose 4 Fast Flow (GE Health) according to the manufacturer’s instructions. His-tagged CaMIIα and acetylated CaM proteins were purified using Ni-NTA agarose beads (QIAGEN) following the manufacturer’s protocols. For binding assays, purified His-CaMIIα proteins (1 μM) were incubated with immobilized GST-CaM (1 μM) for 2 h at 4°C with 1 mM EGTA (no Ca²⁺, as a negative control), 0.1 mM, or 1 mM CaCl₂. The mixture was then washed, eluted, and subjected to western blot with anti-His and anti-GST antibodies (Abmart).

**In vitro CaMIIα autophosphorylation**

In brief, the purified His-CaMIIα proteins (1 μM) and GST-tagged WT, 3KR, or 3KRQ-CaM proteins (1 μM) were incubated at 31°C for 5 min in the reaction buffer (500 μM ATP, 50 mM HEPES, pH 7.5, 1 mM DTT, 0.1% Tween-20, and 10 mM MgCl₂) with 1 mM EGTA (no Ca²⁺, as a negative control), 0.1 mM, or 4 μM CaCl₂ with constant shaking. Reactions were stopped by an addition of 2× SDS sampling buffer, followed by western blot with anti-p-CaMIIα and anti-CaMIIα antibodies (Cell Signaling, 11945).

**Calcineurin activity assay**

The phosphatase activity of calcineurin was assayed by a colorimetric kit (Abcam, ab139461) according to the manufacturer’s instructions. The kit provided both the A and B subunit of calcineurin. Dephosphorylation of the RII phosphopeptide substrate in the presence of CaM was detected by the phosphatase activity, was read on a microplate reader (Thermo Fisher Scientific).

**BRET assay**

We applied bioluminescence resonance energy transfer (BRET) assay (50) to study CaM-CaMIIα interaction systems in HEK293 cells. We first transfected one dish (10 cm diameter) of cells with 10 μg BRET plasmids containing WT, 3KR, or 3KRQ-CaM and the target peptide from CaMIIα (C18). The cells were harvested with 1 ml KCl (100 mM) 48 h after transfection and were separated into ten 15 ml centrifuge tubes. After spin-down (500 rpm × 5 min), the pellet was resuspended with calcium calibration buffer (Invitrogen, C-3008MP) plus 1000 units/ml α-hemolysin (Sigma, H9395). We added 200 μl cell resuspension solution plus 50 μl luciferase substrate (Promega, S2011) in each well of 96-well plate and then read the emission ratio of 530 nm to 480 nm. The calcium concentration was adjusted to 0, 0.1, 0.3, 1, 3, 10, 30, 100 μM for Ca²⁺ binding assay using the calcium calibration buffer kit. A Ca²⁺ titration curve was used to calculate Kd value by nonlinear regression analysis. The averaged data from four independent experiments were fitted to the Hill equation using Prism 8 software.

**Molecular dynamics simulation**

We carried out 200 ns MD simulation for CaM-CaMIIα interaction systems in the presence or absence of acetylation with the help of Amber package (51). The CaM-CaMIIα complex structure was extracted from Protein Data Bank (PDB ID: 1CM4), and for the acetylated system, modifications were introduced and optimized through Discovery Studio. After setting up, Amber ff14SB force field was applied to the parameter calculations and TIP3P water model was used for the system solvation. NaCl ions were added to the complex systems for the charge neutralization and the simulation of the physiological conditions. After the primary preparations, two rounds of energy minimization were carried out. In the first step, the overall structural scaffolds were held rigid and water and counterions underwent a 5000-step maximum minimization cycles. After that, the whole systems were relaxed, and minimizations with no restriction followed. Then, under NVT conditions, two simulation systems were heated up from 0 to 300 K within 300 ps. They were further equilibrated at 300 K for another 700 ps, after which, 200 ns of MD simulations were performed for these two systems respectively. Analyses of the simulation trajectories were carried out with the cpptraj and MMTPBSA.py plugin within Amber package. In total, 200 representative snapshots from the last 180 ns trajectories, when the simulations had already reached equilibrium, were extracted for detailed investigations.

For binding free energy evaluation, we followed the Equation 1 below, with the complex systems, the receptors (CaM/ acetylated CaM), and the ligands (CaMIIα) calculated respectively.

\[
\Delta G_{\text{binding}} = \Delta G_{\text{complex}} - \Delta G_{\text{receptor}} - \Delta G_{\text{ligand}}
\]  

Within Equation 1, the free energy terms were given by the following Equation 2:

\[
\Delta G = \Delta E_{\text{vdW}} + \Delta E_{\text{ele}} + \Delta G_{\text{solv}}
\]  

in which \(\Delta E_{\text{vdW}}\) denoted the energy contribution from van der Waals force, \(\Delta E_{\text{ele}}\) represented the energy contribution from electrostatic force, and \(\Delta G_{\text{solv}}\) referred to the energy contribution from total solvation-free energy.

**LTP recording and quantification**

Hippocampal slices from 5-week-old WT and 3KR/3KR mice were placed in a recording chamber continuously superfused with prewarmed (32 °C ± 1 deg. C) aCSF at a rate of 3 ml/min. IEPSPs were evoked (0.033 Hz, 0.1 ms current pulses) in the CA1 stratum radiatum by stimulating Schaffer collateral (SC) with a two-concentric bipolar stimulating electrode (FHC) and recorded in current-clamp by a HEKA EPC 10 (HEKA Elektronik) amplifier with aCSF-filled glass pipettes (1–5 MΩ). LTP was induced using three trains of theta burst stimulation (ten bursts at 5 Hz, each having 50 pulses, at 100 Hz) with intertrain intervals of 30 s (75, 76).
Whole-cell LTP recording was performed as previously described (5). Three-week-old male mice were used for whole-cell LTP recording with 20 μM BIC in the aCSF. Briefly, EPSCs of hippocampal CA1 pyramidal neurons were evoked (0.1 Hz, 0.1 ms current pulses) by stimulating SC with electrode placed 200 μm away from recorded neurons at −70 mV. LTP was induced by three trains of high-frequency stimulation (HFS, 100 Hz, 1 s) separated by 20 s with the patched cells depolarized to −20 mV. To avoid “wash-out” of LTP, the HFS was applied within 10 min after achieving whole cell configuration.

All data were acquired at a 10 kHz sampling rate of using PATCHMASTER version 2 x 90.1 software (HEKA Elektronik) and filtered offline at 2 kHz. Analysis was performed with Neuromatic version 3.0 (http://www.neuromatic.thinkrandom.com). Each EPSP or EPSC trace was normalized to baseline. Two consecutive EPSP or six consecutive EPSC traces were averaged to generate 1-min bin, which generated LTP summary time course graphs. For the LTP of field potential, the magnitude of LTP was calculated at an average of normalized EPSP amplitudes 50 to 60 min after TBS. For whole-cell LTP, the magnitude of LTP was calculated at the averaged of normalized EPSC amplitudes 45 to 50 min after HFS.

Generation and stereotaxic injection of lentivirus (LV)

The lentivirus vectors expressing Cam shRNA and rescue genes were kindly provided by Dr Thomas C. Südofer (53). We changed the sequence of IRES in the original vector to p2A to increase the expression level of EGFP. All the AAV and lentivirus were generated in Obio Technology Corp, Ltd. For virus injection, WT male mice at age of 7 to 8 weeks were anesthetized with 1% pentobarbital sodium (100 mg/kg, i.p.) and were placed in a stereotaxic apparatus (RWD Life Science). Viruses were injected bilaterally in the CA1 regions of hippocampus with the coordinates: anteroposterior (AP) −2.7 mm, mediolateral (ML) ± 2.25 mm, dorsoventral (DV) −1.625 mm relative to bregma. Each injection used 1 μl LV and took 10 min. After injection, the glass pipette was left in place for 5 min in order to facilitate diffusion of the virus. The injection sites were examined at the end of the experiments, and animals with incorrect injection site were excluded from the data analysis. Two weeks after LV injection, mice were subjected to experiments. All surgery was conducted with aseptic technique.

Contextual fear conditioning

The investigators who performed behavioral analysis were blind to the genotype or treatment of the mice. Eight-week-old mice were first habituated to the behavioral room and apparatus (Fear Conditioning System, Panlab) for 5 min. During training, mice were placed in the conditioning chamber and exposed to three foot shocks (2 s, 0.5 mA) with an interval of 30 s. One day after training, mice were returned to the chamber to evaluate contextual fear learning. Freezing during training and testing was scored using PACKWIN software.

Data were expressed as percent freezing in 180-s epochs, with each epoch divided into 12 bins.

Statistics

All the data were shown as mean ± SD. Comparisons between two groups were made using unpaired t test. Comparisons between three or more groups were made using one-way ANOVA analysis followed by Tukey’s post-hoc test. Data on the binding curve between CaM and C18, CaM and M13, I/O curve, and PPF at SC-CA1 synapse were analyzed by two-way ANOVA. Statistically significant difference was indicated as follows: ***p < 0.001, **p < 0.01, and *p < 0.05. The statistical analysis was performed with the software of GraphPad Prism 8.

Data availability

All data supporting the results presented herein are available from the article paper.

Supporting information—This article contains supporting information.

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Conflict of interest—The authors declare no competing interests.

Abbreviations—The abbreviations used are: CaM, calmodulin; CaMKI/IIα, calcium/calmodulin-dependent protein kinase IIα; cLTP, chemical LTP; LTP, long-term potentiation; MD, molecular dynamics; NAM, nicotinamide; PPF, paired pulse facilitation; SC, Schaffer collateral.

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