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Delayed Intrinsic Activation of an NMDA-Independent CaM-kinase II in a Critical Time Window Is Necessary for Late Consolidation of an Associative Memory

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Calcium/calmodulin-dependent kinases (CaM-kinases) are central to various forms of long-term memory (LTM) in a number of evolutionarily diverse organisms. However, it is still largely unknown what contributions specific CaM-kinases make to different phases of the same specific type of memory, such as acquisition, or early, intermediate, and late consolidation of associative LTM after classical conditioning. Here, we investigated the involvement of CaM-kinase II (CaMKII) in different phases of associative LTM induced by single-trial reward classical conditioning in Lymnaea, a well established invertebrate experimental system for studying molecular mechanisms of learning and memory. First, by using a general CaM-kinase inhibitor, KN-62, we found that CaM-kinase activation was necessary for acquisition and late consolidation, but not early or intermediate consolidation or retrieval of LTM. Then, we used Western blot-based phosphorylation assays and treatment with CaMKIINtide to identify CaMKII as the main CaM-kinase, the intrinsic activation of which, in a critical time window (~24 h after learning), is central to late consolidation of LTM. Additionally, using MK-801 and CaMKIINtide we found that acquisition was dependent on both NMDA receptor and CaMKII activation. However, unlike acquisition, CaMKII-dependent late memory consolidation does not require the activation of NMDA receptors. Our new findings support the notion that even apparently stable memory traces may undergo further molecular changes and identify NMDA-independent intrinsic activation of CaMKII as a mechanism underlying this “lingering consolidation.” This process may facilitate the preservation of LTM in the face of protein turnover or active molecular processes that underlie forgetting.

Introduction
Calcium/calmodulin-dependent kinases (CaM-kinases) are highly suitable molecular substrates for long-term memory storage due to the unique ability of this group of enzymes to maintain an active autophosphorylated state even after the decay of external stimuli (i.e., when Ca\(^{2+}\) influx stops) (Hook and Means, 2001; Hudmon and Schulman, 2002a). CaMKII, like other CaM-kinases, can be activated by transient influx of Ca\(^{2+}\) (e.g., through NMDA receptors) and plays a role in subsequent transcriptional and translational processes involving the cAMP response element binding protein (CREB), a key transcription factor for learning and memory (for review, see Silva et al., 1998; Abel and Lattal, 2001; Cammarota et al., 2002; Hudmon and Schulman, 2002b; Wang et al., 2006).

The involvement of CaMKII in memory acquisition has been well documented (for review, see Cammarota et al., 2002; Silva, 2003; Elgersma et al., 2004). However, the role of, and especially the temporal pattern for, the involvement of CaMKII in memory consolidation still remains largely unclear, with only a few studies directly showing that activation of CaMKII is necessary for the consolidation of behavioral long-term memory (Wolfman et al., 1994; Frankland et al., 2001; Miller et al., 2002; Wang et al., 2003; Bevilaqua et al., 2005).

Here, we sought to investigate the temporal involvement of CaMKII in various different stages of associative learning and memory in a single, well defined experimental system. Previous work in the pond snail Lymnaea stagnalis has provided important insights into the molecular mechanisms of associative memory after single-trial classical reward conditioning (for review, see Kemenes, 2008), and here we used the many advantages offered by this system and paradigm to study the role of CaMKII in the acquisition, early, intermediate, and late consolidation and retrieval of long-term associative memory.

Here, for the first time, we identify a critical time window (occurring at ~24 h after training) during which intrinsic activation of CaMKII is required for late consolidation of associative memory. Additionally, we demonstrate that CaMKII activation is required for memory acquisition, but not for early or intermediate consolidation or memory retrieval. We also show that NMDA receptors are involved in the acquisition but not late consolidation of long-term memory (LTM), indicating a dissociation of NMDA receptor function and CaMKII activation between these two different phases of memory formation.
Materials and Methods

Experimental animals. *Lymnaea stagnalis* pond snails were bred at the University of Sussex. The snails were maintained in large holding tanks using Ca$^{2+}$-free water (also used throughout the experiments) at 18–20°C with a 12 h light-dark cycle, and fed *ad libitum* on lettuce and a vegetable-based fish food (TETTRA Werke). The animals were food-deprived for two d before the beginning of the conditioning procedure.

Single-trial conditioning protocol and pharmacological treatments. A membrane-permeable inhibitor of the activation of CaM-kinases, 1-[[N,O-bis(5-isouquinolinesulfonyl)]-N-methyl-L-tyrosyl]-4-phenylpyridine (KN-62) (Sigma-Aldrich) was applied to the nervous system of intact animals for the first set of experiments. KN-62 has been shown to inhibit the activation of CaM-KII (Davies et al., 2000), but also other CaM-kinases such as CaMKI, CaMKIV, and CaMK kinase (Hidaka and Yokokura, 1996) and thus any effects observed after administering KN-62 can indicate whether or not there is an involvement of at least one of these CaM-kinases in the memory task used. When KN-62 was found to affect memory, we further investigated whether the observed effects were due to the inhibition of the activation of CaMKI. In these experiments we used myristoylated CaMKII(N) (Merck Chemicals), a specific CaMKII inhibitor that is membrane permeable and does not inhibit other CaM-kinases or CaMK kinases (Chang et al., 1998; Pereda et al., 1998; Soderling et al., 2001). We then performed further experiments administering an NMDA receptor blocker MK-801 (Sigma-Aldrich) at time points when inhibiting CaMKII activation resulted in anterograde amnesia. The purpose of this third experiment was to establish whether the involvement of CaMKII in acquisition or consolidation of associative memory is related to the activation of NMDA receptors; CaMKII activation has been linked to NMDA receptor activation in previous studies of both memory and synaptic plasticity (Davis et al., 1992; Giese et al., 1998; Shimizu et al., 2000; Miller et al., 2002; Bevilacqua et al., 2005).

Snails were trained using an established single-trial reward classical conditioning protocol (Alexander et al., 1984; Kemenes et al., 2002). Before training, the snails were placed individually into Petri dishes containing 90 ml of water for a 10 min acclimatization period, so that a constant low level of spontaneous rasping (stereotyped feeding movements of the mouth) was reached in the novel environment (Kemenes and Benjamin, 1994). Then four types of experiment were performed: “acquisition,” “retrieval,” “early consolidation,” and “intermediate and late consolidation” (Fig. 1). Three groups of animals were used in each type of experiment. To test the effects of general CaM-kinase inhibition on memory, one group was injected in the hemolymph with KN-62 (final concentration 20 μM in 0.67% dimethyl sulfoxide (DMSO, Sigma-Aldrich)) at different time points: 30 min before memory acquisition (Fig. 1A), 30 min before retrieval (Fig. 1B), 15 min after acquisition (Fig. 1C), or at a range of longer intervals after conditioning, with a further 18 h allowed before testing (Fig. 1D). A second group (Veh) was also trained in each experiment, but in this group the animals were injected with the vehicle (0.67% DMSO in the hemolymph). Finally, a naive group (not trained or injected) was also used in each experiment to identify baseline responses to the conditioned stimulus (CS). The set up of the groups of snails used in the experiments testing the specific effects of CaMKII(N) and MK-801 was similar to that used in the experiments with KN-62. Three groups were used: the drug-treated group (“Ntide” or “MK-801”) was injected with CaMKII(N) (final concentration 4 μM in 0.17% DMSO in the hemolymph) or MK-801 (final concentration 2 μM in 0.067% DMSO in the hemolymph), respectively; the “Veh” group was injected with the vehicle alone, i.e., 0.17% or 0.067% DMSO, respectively, and the “Naive” group was neither trained nor injected.

For the CS + unconditioned stimulus (US) paired protocol training, 5 ml of amyl acetate solution (0.08% in water, the CS) was delivered into the dish using a plastic syringe and was followed 15 s later by 5 ml of sucrose solution (13.4% in water, the US). Snails remained in the CS + US mixture for a further 105 s (thus the total time exposed to stimuli was 2 min). After training, snails were placed into a tank of clean water and after 10 min transferred back to their home tanks. For testing, individual snails were taken from their home tanks using a blind procedure and placed in Petri dishes. After a 10 min acclimatization period, rasps were counted for 2 min (i.e., spontaneous rasping). 5 ml of the CS was then applied to the dish, and rasps were counted for a further 2 min (i.e., the feeding response to the CS). The “feeding score” was defined as the feeding response to the CS minus spontaneous rasping.

Cloning of a partial *Lymnaea* αCaMKII cDNA sequence. To determine whether a homolog of vertebrate αCaMKII exists in *Lymnaea stagnalis*, we cloned the functional domains of the autoregulatory region of αCaMKII in the nervous system of this species. Total RNA samples were isolated from the pooled central ganglia of 10 snails with Absolutely RNA Miniprep Kit (Stratagene) according to the manufacturer’s instructions. To synthesize first strand cDNA, reverse transcription (RT) using SuperScript II Rnase Reverse Transcription (Invitrogen) was carried out, according to the manufacturers’ instructions, in 20 μl of mixture containing 1–5 μg of total RNA sample, 1 μl of random primer (Roche Diagnostics), 1 μl of 10 μM dNTP Mix (10 mM each dATP, dGTP, dCTP, dTTP), Roche Diagnostics), 4 μl of 5× First Strand Buffer, and 2 μl of 0.1 M DTT (both from Invitrogen), 1 μl (200 units) of SuperScript II Reverse Transcriptase (Invitrogen), 1 μl of RNase inhibitor (Ambion Applied Biosystems), and sterile distilled H₂O. The first strand cDNA produced was then used as a template for PCR amplification. Two pairs of degenerate nested oligonucleotides, directed to the conserved region among the cDNA sequence for human αCaMKII (708–921), were used to amplify the partial αCaMKII sequence of *Lymnaea*. These were as follows: 5’-GT(GC)(GG)(AC)/TA(CT)CC(AT)TT(C/T)TGG-3’ (forward outside), 5’-TT(C/T)TGGGA(CT)/GA(AG)(GA)/CT(GA)(C)-3’ (forward inside), 5’-CAACAT(TG)GTG(TG)GTG(AG)(AG) AT(AG)/GG(CT)CC-3’ (reverse inside), 5’-CGAAA(AG)/TGT(CT)(GT) GT(AG)/GC(CT)AACAT-3’ (reverse outside). PCR was performed in a GeneAmp PCR system (PerkinElmer Life and Analytical Sciences), for 40 cycles, denaturing at 95°C for 15 s, annealing at 48°C for 30 s and extending at 72°C for 30 s using Hotstar Plus Taq (Qiagen). Amplified cDNAs were subcloned into a pCR2.1 vector (25 ng/μl) with TA Cloning Kit (Invitrogen). The ligation products were used to transform TOP10® cells (Invitrogen). Colonies were selected and plasmid DNA was isolated and analyzed by restriction enzyme EcoRI and purified with Miniprep Kit (Qiagen) and finally was sequenced by MWG-Biotech AG.
Western blot analysis of pCaMKII levels after classical conditioning. Phosphorylated CaMKII levels were compared among five different groups of animals. One group was trained using the paired CS/US protocol 24.5 h before dissection. Two further groups were subjected to paired training, then injected with KN-62 or vehicle 24 h later (in the same concentrations that were used in the behavioral experiments), and dissected 30 min after injection. A fourth group contained animals that were subjected to an explicitly unpaired CS/US protocol (CS/US interval, 1 h) and a fifth group consisted of naive animals. Each group contained 5 animals and each experiment was repeated 5 times. Previous work has shown that the cerebral and buccal ganglia of the snail are two major brain regions which are involved in food-reward classical conditioning (Kemenes, 2008), thus these ganglia were dissected and prepared as protein samples for Western blotting. After homogenization in ice-cold Tris lysis buffer (125 mM TRIS, pH 6.8), the samples were put on ice and then centrifuged at 4°C. Supernatants were removed into a clean tube and then boiled at 95°C for 5 min with 50 mM Tris, pH 6.8, 2% (w/v) SDS, 10% glycerol, 10% β-mercaptoethanol (BME), 0.01% Bromophenol Blue (BFB), 2 mM Na3VO4, 10 mM NaF, 10 mM MgCl2, 50 μM ATP, 8 mM EGTA and 2% protease inhibitor cocktail (Sigma-Aldrich Company) and then centrifuged at 13,000 X g at 4°C to remove debris. Approximately 20 μg of each sample was run by SDS-PAGE and blotted according to standard methods. The blotted PVDF membrane was blocked for 1 h at RT or overnight at 4°C in Tris Buffer Saline (TBS) with 0.15% Tween 20 with 5% nonfat dry milk. The membrane was incubated for 2 h at RT or overnight at 4°C with either mouse monoclonal IgG anti-α subunit CaMKII-II, or mouse monoclonal IgG anti-phospho-CaMK-II (T286) (both from Millipore) diluted 1:500 in TBS/Tween 20/milk and BSA, then washed in TBS/Tween three times, and incubated for 1 h at RT with secondary antibody (1:1000 horseradish peroxidase (HRP)-linked anti-mouse IgG antibody, New England Biolabs ) diluted in TBS/Tween 20/milk and BSA. The membrane was then washed in TBS/Tween twice and once in TBS. Protein bands were visualized using enhanced chemiluminescence (Millipore). Control samples were only probed with the secondary antibody and showed no bands. Densitometric analysis was performed using ImageJ.

### Results

#### Acquisition, but not retrieval of memory, requires CaMK-kinase activation

In the first set of experiments we injected KN-62 into intact snails at different time points, either before CS + US training (for memory acquisition, Fig. 1A) or postlearning (for memory retrieval, Fig. 1B) to establish whether or not the activation of any CaMK-type enzyme was involved in these processes. When KN-62 was administered 30 min before training and memory was tested 24 h after training (Fig. 1A, acquisition), significant differences were found in the feeding scores among the three groups of snails (Fig. 2A). The significantly higher level of feeding responses to the CS in the vehicle-injected trained group (Veh) versus the Naive group (Fig. 2A) indicated that the single-trial classical conditioning was successful. The KN-62-injected trained group showed a significantly lower score than the Veh group, and was not significantly different from the Naive group (Fig. 2A). This indicates that administration of KN-62 prevents the increase in feeding response to the CS that is normally acquired during training, and further implies that the blocking of the activation of CaMK-kinases by KN-62 at around the time of training causes impairment to the animals’ long-term memory tested at 24 h after training.

In contrast, there was no significant difference between the KN-62 and Veh groups when animals were injected 30 min before the CS test at 24 h after training (Figs. 1B, retrieval, 2B). Both groups were able to remember what they had learned 24 h earlier, as shown by a significant increase in feeding response to the CS compared to the Naive group (Fig. 2B). In an additional experiment we found that 30 min after injection with KN-62, the animals’ ability to respond to their unconditioned food stimulus, sucrose, was not affected (Fig. 2C). Thus, blocking the activation of CaMK-kinases with KN-62 30 min before a food stimulus did not interfere with the animals’ ability to produce either the conditioned or unconditioned feeding response. This is in sharp contrast to the strong amnestic effect seen when KN-62 was injected 30 min before training. These results demonstrate that the blocking of the activation of CaMK-kinases produces impairment in memory acquisition, but not in memory retrieval, implying that there are different roles for the activation of CaMK-kinases in these two fundamental processes of memory.

A distinct critical time window of intrinsic CaMK-kinase activation is required for late memory consolidation

Systemic injection of KN-62 at 15 min (Fig. 1C, early consolidation) or 6, 12 or 20 h after training (Fig. 1D, intermediate consolidation) did not cause loss of long-term memory (Fig. 3A, B, left panel); the feeding responses of KN-62-injected trained animals were elevated compared to those of naive animals, and indistinguishable from the trained group injected with the vehicle alone. However, injection of KN-62 at 24 h after training (Fig. 1D, late consolidation) produced a significant reduction of the feeding response tested 18 h later (Fig. 3B, middle panel). This impairment of memory was quantitatively similar to the effects seen when KN-62 was injected 30 min before single-trial classical conditioning to block training-induced CaMK-kinase activation (Fig. 2A). As KN-62 injected at 24 h after training caused amnesia in the apparent absence of any external stimuli, we conclude that KN-62 at this late posttraining time point acts by preventing intrinsically triggered activation of a CaMK-kinase type enzyme. Together, the above results suggest that at around the 20 h point after training, memory seems to be consolidated without a require-
ment for ongoing activation of CaM-kinases. However, at ~24 h after training, there is a period where one or more CaM-kinases must become activated in order for associative memory to undergo a late consolidation process.

To determine whether this late consolidation process required a continued activation of a CaM-kinase type enzyme, KN-62 was injected at 30 h after training and memory was tested 18 h later (Fig. 1D). In this experiment, the increase in feeding response to the test CS was preserved (Fig. 3B, right panel). Thus, a requirement for intrinsic activation of a CaM-kinase type enzyme during late memory consolidation appears to exist in a time window between 20 and 30 h after training.

Identification of CaMKII as the main CaM-kinase involved in memory acquisition and late consolidation

CaMKII has been postulated as one of the most likely molecular substrates for LTM (Hook and Means, 2001; Hudmon and Schuman, 2002a). We therefore investigated whether the observed memory impairments by KN-62 resulted from the inhibition of this specific CaM-kinase during memory acquisition, and in particular when KN-62 was applied during the late consolidation phase.

We first confirmed the existence of a bona fide CaMKII in the Lymnaea CNS by cloning the core domains of the autoregulatory region of LymCaMKII, including the autophosphorylation domain (GenBank Accession Number: EF428260). The cloned sections are highly conserved (86% sequence homology with the mammalian α subtype of CaMKII) (Fig. 4) suggesting functional homology and justifying the use of inhibitors and antibodies originally developed against mammalian CaMKII.

To address the possible role of CaMKII as an important CaM-kinase involved in late memory consolidation, the main focus of this paper, we tested whether CaMKII actually showed intrinsic phosphorylation at 24 h after training. Five replicate experiments were conducted using the Western blotting technique (see Materials and Methods). In these experiments we also tested whether, if such phosphorylation occurred, it could be prevented by treatment with KN-62, the broad-spectrum CaM-kinase inhibitor used in the behavioral experiments. These experiments showed that, relative to total CaMKII levels, there was a significantly higher level of phosphorylated CaMKII present in the buccal and cerebral ganglia at ~24 h after contingent training (CS/US interval, 30 s) than in naive and explicitly unpaired control animals (CS/US interval, 1 h). Importantly, the same experiments also showed that the increased phosphorylation induced by contingent training was significantly blocked by KN-62 (Fig. 5A,B). The biochemical findings at ~24 h after training (Fig. 5B) corresponded strongly with the results of behavioral tests conducted 18 h later on groups of trained and control animals in the same experiment (Fig. 5C). These results both demonstrate that CaMKII is activated ~24 h after training and suggest that the activation of this specific CaM-kinase is necessary for the consolidation of associative memory.

In the next set of experiments, we tested whether inhibition of CaMKII alone was sufficient to lead to the significant memory impairments seen in the experiments where KN-62 was used. In these tests we used CaMKIIN tide, a specific inhibitor of CaMKII (Chang et al., 1998). The application of this drug 30 min before training significantly impaired the snails’ performance at the memory test 24 h after training (Fig. 6A), indicating that the inhibition of CaMKII alone is capable of impairing acquisition of associative memory. The impairment caused by the administration of CaMKIIN tide is quantitatively similar to that produced by the administration of KN-62 (compare Fig. 6A and Fig. 2A), demonstrating that the effects observed in the experiments using KN-62 (injected 30 min before training) were mainly due to the inhibition of CaMKII activation. This experiment therefore provides specific evidence for the important role of CaMKII in this stage of LTM.

More crucially, administration of CaMKIIN tide 24 h after training also significantly reduced the conditioned feeding score of the CaMKIIN tide-treated animals compared with the vehicle-treated animals when they were tested 18 h after injection (Fig. 6B). The effects caused by the administration of CaMKIIN tide were again similar to those produced by KN-62 in the same time window (compare Fig. 6B and Fig. 3B, middle panel). This demonstrates further that the effects observed in the experiment using KN-62 (injected 24 h after training) were due specifically to the inhibition CaMKII activation at ~1 d after training. These results, together with the results of the Western blot experiments, provide the first experimental evidence to show definitively that CaMKII plays a crucial role in late consolidation of LTM after single-trial classical food-reward conditioning in Lymnaea.

CaMKIIN tide recently has been shown to inhibit both calcium-dependent CaMKII activity and the autonomously active kinase itself (Vest et al., 2007). It was therefore also important to test the effect of CaMKIIN tide at a time point after training when KN-62 had no effect, such as 20 h after training (see Fig. 3),
to distinguish between these two mechanisms. A significant amnestic effect arising from treatment with CaMKIINtide at this time point would indicate that this peptide inhibitor impairs memory after single-trial conditioning in Lymnaea mainly by blocking autonomously active CaMKII. On the other hand, a lack of amnestic effect in response to CaMKIINtide treatment at 20 h after training would suggest that the amnesia observed at 24 h after training in both the KN-62 and CaMKIINtide experiments resulted from an inhibition of calcium-dependent intrinsic activation of CaMKII.

When we injected CaMKIINtide into a group of animals 20 h after training, these animals showed no memory impairment 18 h later compared to a vehicle-injected trained group \( n = 22 \) for each group. Both groups showed significantly higher levels of feeding response to the CS \((7.0 \pm 2.1\) and \(7.2 \pm 2.2\) bits/2 min, respectively) compared to a Naive control group \((n = 20, 2.0 \pm 1.2\) bits/2 min; ANOVA, \( F_{(2,63)} = 4.7, p < 0.01\); Tukey’s, \( p < 0.05\)) and they were not significantly different from one another (Tukey’s, \( p > 0.05\)). The findings from this experiment lend strong support to the notion that, similar to KN-62, the amnestic responses to CaMKIINtide treatment at 24 h after training are due to its inhibitory effect on calcium-dependent kinase activity rather than on autonomously active CaMKII.

**NMDA receptors are involved in CaMKII-dependent acquisition, but not late consolidation, of associative memory**

Previous work in other systems has demonstrated a direct link between NMDA receptors and activation of CaMKII (Bayer et al., 2001), and recently, NMDA-like receptors have been characterized at the molecular level in both Aplysia and Lymnaea (Ha et al., 2006). Moreover, glutamate has been shown to be an important transmitter in the Lymnaea feeding system (Brierley et al., 1997; Hatakeyama et al., 2007). The effects of glutamate on neurons of the feeding network have been shown to be modulated by nitric oxide (D’Yakonova and D’Yakonova, 2008), a transmitter necessary for memory consolidation after single-trial learning in Lymnaea (Kemenes et al., 2002). Importantly, a close link between NMDA receptor function and CaMKII activation has been found in various different forms of synaptic plasticity and learning and memory (Izquierdo et al., 1999; Moriya et al., 2000; Bevilaqua et al., 2001; Hatakeyama et al., 2007). The effects of glutamate on NMDA receptor function and CaMKII activation has been shown to be strongly supported by the findings from this experiment. When we injected CaMKIINtide into animals 24 h after single-trial training, these animals showed no memory impairment 24 h later compared to a vehicle-injected trained group \((n = 22\) for each group). Both groups showed significantly higher levels of feeding response to the CS \((7.0 \pm 2.1\) and \(7.2 \pm 2.2\) bits/2 min, respectively) compared to a Naive control group \((n = 20, 2.0 \pm 1.2\) bits/2 min; ANOVA, \( F_{(2,63)} = 4.7, p < 0.01\); Tukey’s, \( p < 0.05\)) and they were not significantly different from one another (Tukey’s, \( p > 0.05\)). The findings from this experiment lend strong support to the notion that, similar to KN-62, the amnestic responses to CaMKIINtide treatment at 24 h after training are due to its inhibitory effect on calcium-dependent kinase activity rather than on autonomously active CaMKII.

**Figure 4.** Amino acid (AA) sequence predicted from the partially cloned Lymnaea αCaMKII cDNA and the corresponding AA sequence of human αCaMKII. The homology of these sequences between the two species is 86%. The cloned region contains the αCaMKII autoregulatory (including autophosphorylation) domain (Lymnaea position 89–101) and part of the catalytic domain (position 1–88). Asterisks indicate identical amino acids in the Lymnaea and human sequences.

**Figure 5.** Increased intrinsic phosphorylation of CaMKII 24 h after single-trial classical reward conditioning and late memory consolidation. A, B, Western blot experiments. Three main experimental groups were used, one injected with KN-62 24 h after paired training, one injected with vehicle and one un.injected. Cerebral and buccal ganglia were dissected from all trained animals at 24.5 h after training and also from animals of an unpaired control group and a naive control group. Homogenates from five replicates of five pooled tissue samples per group (see Materials and Methods) were loaded onto gels and the resulting blots probed with either a pCaMKII antibody (example in A, top) or with a total CaMKII antibody (example in A, bottom). The total CaMKII signal was used as an internal standard for densitometric comparisons. Relative integrated density values \((p\text{CaMKII}/\text{CaMKII})\) in samples taken from naive animals were assigned a value of 1. All other relative integrated density values were normalized to this naive baseline (dashed line) and used for multiple statistical comparisons (shown in B, significance compared to unpaired group indicated by asterisks). Test statistics: ANOVA: \( F_{(2,63)} = 11.2, p < 0.003\); Tukey’s test: No inj. (paired) versus Unpaired, \( p < 0.05\); Vehicle (paired) versus Unpaired, \( p < 0.001\); KN-62 (paired) versus No inj. (paired) and Vehicle (paired), \( p < 0.05\) and 0.01, respectively; KN-62 (paired) versus Unpaired and Naive, \( p < 0.01\). C, Memory tests. Three groups of trained (paired stimulus) animals from the experiment described in A and B were kept for behavioral tests performed 18 h after injection. Naive and unpaired control animals from the same experiment were also saved for behavioral tests. Note that for an easier comparison with the densitometric data in B, for the Naive group, only the mean value is shown (dashed line; \( SE \pm 1.6\)). Test statistics: ANOVA: \( F_{(2,63)} = 6.6, p < 0.001\); Tukey’s test: No inj. (paired) versus Unpaired and Naive, \( p < 0.01\) and 0.05, respectively; Vehicle (paired) versus Unpaired and Naive, \( p < 0.01\) and 0.05, respectively; KN-62 (paired) versus No inj. (paired) and Vehicle (paired), \( p < 0.05\); KN-62 (paired) versus Unpaired and Naive, \( p > 0.05\) (n.s.); No inj. (paired) versus Vehicle (paired), \( p > 0.05\) (n.s.). Significance compared to unpaired group is indicated by asterisks.
The treatment of groups (Fig. 6) showed that MK-801 was injected 24 h after training did not affect LTM tested 24 h after injection. These results, in contrast to those found when MK-801 was injected 30 min before training and tested 24 h after training, indicate a dissociation between the activation of CaMKII and the activation of NMDA receptors, suggesting that the intrinsic activation of CaMKII 24 h after training is NMDA-independent.

**Discussion**

Here, we show first that activation of CaMKII, a major subtype of the CaM-kinases, is required for the acquisition of long-term associative memory in *Lymnaea*. Furthermore, we show that activation of NMDA receptors is also crucial for acquisition of this type of classical conditioning. This finding is consistent with the vertebrate learning literature (Silva et al., 1992; Davis et al., 1992; Giese et al., 1998; Shimizu et al., 2000; Rodrigues et al., 2004; Irvine et al., 2005; Bevilaqua et al., 2005), implying that the role of CaMKII and its close correlation with the activation of NMDA receptors in learning is conserved between invertebrates and vertebrates. NMDA-like receptors are present in both *Aplysia* and *Lymnaea* (Ha et al., 2006), providing a molecular foundation for functional conservation.

Importantly, we have identified for the first time a distinct critical time window (~24 h after training) during which intrinsic activation of CaMKII occurs and CaMKII is specifically required for late consolidation of associative memory. Moreover, in contrast to its role in acquisition and late consolidation, we found no evidence for a role of the activation of CaMKII (or other CaM-kinases) in early or intermediate consolidation, for up to 20 h after training.

Both KN-62 and CaMKIINtide affect the interaction between the Ca\(^{2+}\)/CaM complex by directly binding to the CaM-binding site of CaMKII. However, KN-62, unlike CaMKIINtide, does not affect the Ca\(^{2+}\)/CaM-independent activity of the already autophosphorylated (activated) enzyme (Tokumitsu et al., 1990; Vest et al., 2007). Thus, both of these drugs inhibit functions that require fresh activation of CaMKII, but in addition, CaMKIINtide may also act by inhibiting autonomously active CaMKII (Vest et al., 2007). The 20 h postraining treatment with either KN-62 or CaMKIINtide was equally ineffective at producing amnesia, showing that neither fresh activation of CaMKII nor autonomous activity of CaMKII are necessary for memory maintenance or late consolidation at this time point. However, the possibility remains that autonomous CaMKII activity plays a functional role in certain other phases of memory. Further studies are necessary to examine this possibility, for example by measuring the level of autophosphorylated CaMKII at different phases of memory acquisition, consolidation, or at the time of memory retrieval.

As treatment with KN-62 and CaMKIINtide at 24 h after training caused loss of memory at a much later time point, we conclude that an ongoing activation of CaMKII occurs at ~24 h and that this plays a crucial role in a distinct late memory consolidation process. Indeed, CaMKII activation may signal the beginning of this late consolidation phase. This observation suggests the interesting possibility that similar to mammalian systems, there is a late process of synaptic reentry reinforcement in *Lymnaea*, which is dependent on the reactivation of CaMKII and is necessary for the long-term storage of old memories (Wang et al., 2003). Whether or not this also indicates the existence of systems level memory consolidation in *Lymnaea*, which follows the previously described molecular consolidation processes taking place between 0 and 6 h after training (Kemenes, 2008), remains to be investigated.

We also established that unlike during acquisition, the activation of CaMKII at ~24 h after training is not dependent upon the activation of NMDA receptors. This finding is also different from what has been described in NMDA receptor and CaMKII knockout mice, where it was suggested that during memory consolidation activation of CaMKII results from the upstream activation of NMDA receptors (Wang et al., 2003). The question arises then: what triggers the intrinsic activation of CaMKII in the *Lymnaea* CNS if not an NMDA-dependent mechanism? This question may be answered in part by the recent finding that single-trial classical reward conditioning leads to a delayed somal membrane potential depolarization in identified modulatory neurons [the Cere-
bral Giant Cells (CGCs)] of the Lymnaea CNS, which plays a key role in long-term memory (I. Kemenes et al., 2006). This depolarization emerges between 20 and 24 h after training, and it is sufficient to lead to a prolonged increase in Ca\(^{2+}\) levels in proximal CGC axon terminals, which in turn is involved in the enhanced response to the CS that occurs after learning (I. Kemenes et al., 2006). A learning-induced delayed rise in Ca\(^{2+}\) levels may well account for the delayed activation of CaMKII involved in late consolidation of memory; our experiments have shown that the time windows for these two events coincide (between ~20 and 24 h after training). The exact cellular mechanisms of the depolarization-induced rise in Ca\(^{2+}\) levels in the CGCs are as yet unknown. As the behavioral pharmacological experiments with MK-801 suggested that this rise is not mediated by NMDA receptors, it is likely to be based on the activation of non-NMDA type voltage-gated calcium channels or intracellular calcium stores in the CGC axon terminals, as proposed previously by I. Kemenes et al. (2006).

CaMKII seems to play a distinct but complementary role to other identified signaling molecules in the processes of memory consolidation in Lymnaea. CaMKII is not involved in consolidation ~15 min after learning, but nitric oxide (NO) and protein kinase A (PKA) are (Kemenes et al., 2002; Michel et al., 2008). However, NO and PKA are not required for late memory consolidation at ~24 h after training (Kemenes et al., 2002; G. Kemenes, 2006; Michel et al., 2008). The complementary roles played by CaMKII and these other molecules also suggest that the consolidation process is dynamic, and may involve both parallel and sequential activation of different signaling cascades in different phases of the consolidation of long-term memory even after a single training trial.

The delayed intrinsic activation of CaMKII shown here also strongly supports the notion of “lingering consolidation” (Dudai and Eisenberg, 2004). This dynamic feature can be a general characteristic of the memory consolidation process and supports the hypothesis that even a long time after learning, memory may be considered to be consolidated with respect to certain molecules, but not to others.

Our finding that there is no requirement for the activation of CaMKII for memory consolidation >30 h after training also supports the general idea of dynamically organized time windows for the posttraining activation of different signaling cascades (Schwärl and Müller, 2006). It is possible that activation of CaMKII at ~24 h after training leads to a sustained high level of autophosphorylated CaMKII, and this is required for memory to be maintained in the even longer term [for up to 3 weeks (Alexander et al., 1984)]. However, it is yet to be shown that autophosphorylation of CaMKII in the Lymnaea CNS lasts long enough for this maintained effect and we cannot rule out either that there is recurrent intrinsic activation of CaMKII or other signaling molecules at even later stages of memory, as suggested by the synaptic reentry reinforcement hypothesis formulated in mammalian models of memory consolidation (Wang et al., 2003).

By cloning the functional domains of CaMKII in Lymnaea, we have directly demonstrated the existence of this ubiquitous and crucial molecule in this well established molluscan model system of learning and memory. Our behavioral results demonstrate not only that CaMKII exists in Lymnaea, but also that it shares with its mammalian counterpart important functional roles in learning and memory. Therefore, our findings further strengthen the comparability of the results obtained from invertebrate and vertebrate systems, implicating that there are important conserved molecular mechanisms that underlie fundamental cognitive tasks such as associative learning and memory. Most importantly, however, our work identified a novel NMDA-independent role for CaMKII in late memory consolidation and it has yet to be established whether or not this function too has been conserved in evolutionarily more advanced organisms.

References


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