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Metaplastic effect of apamin on LTP and paired-pulse facilitation

Laurence Ris,1 Brigitte Capron,1,2 Coralie Sclavons,1 Jean-François Liégeois,3 Vincent Seutin,4 and Emile Godaux1,5

1Laboratory of Neurosciences, University of Mons-Hainaut, 7000 Mons, Belgium; 2Department of Neurology, UCL, 1000 Brussels, Belgium; 3Laboratory of Medicinal Chemistry, University of Liège, 4000 Liège, Belgium; 4Laboratory of Pharmacology and Center for Cellular and Molecular Neurobiology, University of Liège, 4000 Liège, Belgium

In area CA1 of hippocampal slices, a single 1-ssec train of 100-Hz stimulation generally triggers a short-lasting long-term potentiation (S-LTP) of 1–2 h. Here, we found that when such a train was applied 45 min after application of the small conductance Ca2+-activated K+ (SK) channel blocker apamin, it induced a long-lasting LTP (L-LTP) of several hours, instead of an S-LTP. Apamin-induced SK channel blockage is known to resist washing. Nevertheless, the aforementioned effect is not a mere delayed effect; it is metaplastic. Indeed, when a single train was delivered to the Schaffer’s collaterals during apamin application, it induced an S-LTP, like in the control situation. At the moment of this LTP induction (15th min of apamin application), the SK channel blockage was nevertheless complete. Indeed, at that time, under the influence of apamin, the amplitude of the series of field excitatory postsynaptic potentials (fEPSPs) triggered by a stimulation train was increased. We found that the metaplastic effect of apamin on LTP was crucially dependent on the NO-synthase pathway, whereas the efficacy of the NMDA receptors was not modified at the time of its occurrence. We also found that apamin produced an increase in paired-pulse facilitation not during, but after, the application of the drug. Finally, we found that the induction of each of these two metaplastic phenomena was mediated by NMDA receptors. A speculative unitary hypothesis to explain these phenomena is proposed.

Results

Delayed facilitation of L-LTP induction by apamin applied during the recovery period

In all the experiments carried out here, the slices were left undisturbed after slicing for 1.5 h (recovery period) before starting to collect the data (recording period). An interface chamber was used for both recovery and recording. When the experiments
were performed in artificial cerebrospinal fluid (aCSF) containing Mg$^{2+}$ at a concentration of 1 mM, application of apamin (100 nM), an SK channel blocker, during the recovery period had a dramatic influence on the duration of the LTP induced by a single train of high-frequency stimulation (100 Hz, 1 sec) (Fig. 1A). In the control situation, a single train elicited an initial potentiation of the slope of the field excitatory postsynaptic potential (fEPSP) reaching 177% ± 9% (n = 5, mean ± SEM). Four hours later, it was reduced to 111% ± 7% of baseline, a level not significantly different from preinduction level (paired Student’s t-test, P = 0.09). In contrast, when apamin was present during recovery, a single train applied 45 min after the end of apamin application induced a strong and long-lasting LTP. The initial potentiation was 249% ± 7% (n = 6). Over the next 20 min it diminished to 176% ± 3% and then remained nearly stable, being as high as 169% ± 7% 4 h after LTP induction, a value significantly higher than that measured in the absence of apamin (Student’s t-test, 111% ± 7%, P < 0.05).

Apamin-triggered delayed facilitation of L-LTP induction is a metaplastic phenomenon

The fact that L-LTP induction was facilitated 45 min after stopping apamin application suggested that this effect was a metaplastic phenomenon. However, because apamin is known to induce a long-lasting block of SK channels (Scuée-Moreau et al. 2002), the aforementioned apamin-triggered facilitation of L-LTP induction could have been due to the persistence of the SK channel block at the moment of LTP induction. We ruled out this possibility. In a first experiment, apamin (100 nM) was applied not before but around the moment of LTP induction (Fig. 1B).

Like in the control situation, in presence of apamin, a single train induced a short-lasting LTP. Immediately after the high-frequency stimulation, the increase in the slope of the fEPSP was greater in the presence of apamin (n = 7) than in its absence (230% ± 11% vs. 177% ± 9%, P < 0.05). However, this facilitation was only temporary. Four hours after induction, the increase in the fEPSP slope was not different in the presence or in the absence of the drug (114% ± 7% vs. 111% ± 7%, P = 0.94). Given that recording (as well as recovery) occurred in an interface chamber with aCSF perfused at a relatively low perfusion rate (1 mL/min), we were concerned that drug wash-in could require more time than we thought. In this case, the apparent lack of action of apamin (15 min after the beginning of its application) could not be due to the fact that apamin was not fixed to its receptors at the time of LTP induction. To rule out this possibility, we carried out a second control experiment. We allowed the slices to recover in normal aCSF, but switched to apamin-containing aCSF for the entire duration of the field recording (Fig. 2A). Here too, 4 h after induction, the increase in the fEPSP slope was not different in the presence or absence of the drug (114% ± 10% vs. 111% ± 7%, P = 0.8). In a third experiment, we blocked SK channels during the recovery period using (+)-methyl-laudanosine, a blocker whose action is fully reversible after 10 min of washout (Scuée-Moreau, J.-F. Liégeois, and V. Seutin, unpubl.). When (+)-methyl-laudanosine (300 μM) was applied during the recovery period, it had an effect similar to that of apamin on the LTP induced 45 min after the end of the recovery (Fig. 2B). In these conditions, the initial potentiation of the slope of the fEPSP reached 259% ± 16% (n = 5). Over the next 20 min it diminished to 182% ± 10%. It then remained at such a high level until the end of the experiment, being as high as 181% ± 12% 4 h after LTP induction, a value significantly greater than that observed in the absence of (+)-methyl-laudanosine (111% ± 7%, P < 0.001).

Thus, apamin applied during the recovery period has a metaplastic effect on LTP: In this situation, a single train of stimulation triggers an L-LTP instead of an S-LTP.

Delayed facilitation of L-LTP induction by apamin applied during the recording period

As a next step, we planned to apply apamin not while the slice was still recovering from the trauma of slicing but during the recording period. To work in conditions close to those used in the initial experiment, where apamin was present during recovery (Fig. 1A), apamin had to be applied for 1 h 15 min after the baseline had been recorded and 45 min before LTP induction (Fig. 3A). However, the long duration (2 h 30 min) of the period prior to LTP induction associated with the usage of a concentration of Mg$^{2+}$ at 1 mM modified the properties of the control LTP. In these conditions, a single train induced a long-lasting LTP instead of a short-lasting one (n = 8). Three hours after induction, the increase in the slope of the fEPSP was still as high as 142% ± 8% (Fig. 3A). It would have been surprising if apamin had improved such a robust LTP. It did not. Three hours after induction, the fEPSP was as high as 149% ± 16%, a level not statistically different from that recorded in the control situation (P = 0.67).

To investigate potential effects of apamin on the induction of...
of L-LTP, we had to find conditions where a single train triggered an S-LTP in the absence of apamin. Because we showed in a previous work (Capron et al. 2006) that a small change in Mg$^{2+}$ concentration (from 1.0 mM to 1.3 mM) could have a surprisingly dramatic influence on metaplasticity, we repeated the experiment, this time using slices bathed in aCSF containing 1.3 mM Mg$^{2+}$ (Fig. 3B). In these conditions, the LTP triggered by a single train 2 h 30 min after the end of the recovery period was much less robust than when the Mg$^{2+}$ concentration was 1.0 mM (Fig. 3A). Three hours after LTP induction, the slope of the fEPSP was as low as 125% ± 6% (n = 14). In contrast, when the apamin perfusion lasted from 2 h to 45 min before LTP induction, a single train caused a long-lasting LTP (Fig. 3B). Immediately after the stimulus application, the slope of the fEPSP reached 248% ± 11% (n = 8). Afterward, it diminished rapidly to a level of 174% ± 5% 20 min later. It then ceased to decrease and was still as high as 167% ± 5% 3 h after the train application, a value statistically different from the control (125% ± 6%, P < 0.001).

The facilitating effect of prior application of apamin on induction of long-lasting LTP occurred within a specific time window. When apamin was applied 1 h before LTP induction it had no facilitating influence (Fig. 3C). Three hours after the train delivery the increase in the slope of the fEPSP was similar whether apamin was perfused or not (131% ± 9% [n = 5] vs. 125% ± 6% [n = 5]).

### Influence of apamin on the depolarization induced by the stimulation train.

In the CA1 region, the field depolarization recorded during a train of high-frequency stimulation (100 Hz, 1 sec) consists of a succession of fEPSPs, whose amplitudes decrease rapidly to −10% of their peak.
of the first response at the tenth electrical shock (Fig. 4A). SK2 channels have recently been demonstrated to be located in the dendritic spines of the hippocampal pyramidal neurons (Ngo-Anh et al. 2005), where they are in close vicinity to the NMDA receptors. Therefore, it can be anticipated that, when a synapse is stimulated repetitively at high frequency, the Ca\(^{2+}\) influx associated with an EPSP would trigger the opening of the SK channels, which would shunt the next EPSP and hence decrease its amplitude. Consequently, apamin should increase the global depolarization induced by a train of stimulation.

Here, we studied the influence of apamin on the field depolarization induced by a high-frequency stimulation (HFS) train. As illustrated in Figure 4A, the profile of the global depolarization induced by an HFS train varied from one slice to another (A) but not when the stimulus was delivered thrice at 30-min intervals on the same slice (B,C). (A) Superimposition of the first 10 fEPSPs of two series of fEPSPs recorded in two distinct slices in response to the delivery of one train of HFS. The intensity was adjusted so that the amplitudes of the first fEPSP were the same in slice number 1 (thick trace) as in slice number 2 (thin trace). (B) Superimposition of the global depolarizations induced in a same slice by three HFS trains separated by 30 min. The intensity of stimulation was adjusted to obtain the same amplitude for the first fEPSP in each of the three global depolarizations. (C) In each slice from a group of 10, the amplitude of each of the first 10 fEPSPs elicited by three HFS trains at 30-min intervals was measured. The means ± SEM of the measurements were plotted in function of the rank of the corresponding fEPSP in the series.}

during and after apamin application with that obtained in the absence of the drug. In a control experiment, we ascertained that, in absence of the drug, the three global depolarizations induced in the same slice by three trains of HFS delivered >30 min apart were the same (Fig. 4B). For relevant comparisons, the intensity of stimulation was adjusted so that the amplitudes of the first fEPSPs of the three-fEPSP series recorded from the same slice were the same. In a control group of 10 slices, the amplitudes of each of the first 10 fEPSPs were measured in the response to each of the three HFS trains (Fig. 4C). The total depolarization induced was the same in the three situations (two-way ANOVA with one repeated measure, \(F_{2,27} = 0.05, P = 0.95\)).

The effect of apamin, applied for 1 h 15 min, was then tested on two other groups of slices. In a first group of eight slices, an HFS train was delivered before apamin application, at the 45th min of the drug perfusion and 45 min after apamin treatment (Figs. 5B, 6A). In a second group of six slices, an HFS train was delivered before, during (at the 15th min), and after (60 min) apamin application (Figs. 5A,C, 6B). As illustrated in Figure 5, the global depolarization induced by a stimulation train was increased not only during (Fig. 5A) but also after (Fig. 5B) apamin
Apamin causes a long-lasting enhancement of the global depolarization elicited by a train of high-frequency stimulation, and this phenomenon is NMDA receptor-dependent. (A, B, C) The amplitude of each of the first 10 fEPSPs within the series of fEPSPs is plotted in function of its rank in the series. (A) Amplitudes of the fEPSPs induced by a stimulation train in a same group of six slices before apamin application, at the 45th min of the drug application, and 45 min after the end of apamin application. (B) Same display as in A, but the stimulation trains were applied on another set of eight slices before, 15 min after the start, and 60 min after the end of apamin perfusion. (C) Same display and same pattern of train delivery on another set of five slices, as in A, but when APV was present throughout the experiment. (D) Illustration of the lack of effect of apamin on the global depolarization induced by a high-frequency stimulation train when APV was coapplied with apamin. The first seven fEPSPs of two series of fEPSPs elicited before (thin trace) and 45 min after apamin application are superimposed.

These results show that the blockage of SK channels induced by apamin was already fully developed 15 min after the beginning of apamin application and persisted ~45 min after the washout of the drug. We interpreted the difference between the experimental results displayed in Figure 1, A and B, as a piece of evidence in favor of the metaplastic nature of the action of apamin on LTP. As aforementioned, the possibility that the apamin molecules had not yet reached their sites on the SK channels after only 15 min of apamin application could cast doubt on this interpretation. This doubt was not justified. Indeed, the fact that the depolarization induced by the stimulation train was already increased only 15 min after the start of the drug application proved that, at that time, the apamin molecules were already attached to their receptor sites.

Apamin did not have any delayed influence on the NMDA receptor-mediated fEPSP

A C

Figure 6. Apamin causes a long-lasting enhancement of the global depolarization elicited by a train of high-frequency stimulation, and this phenomenon is NMDA receptor-dependent. (A, B, C) The amplitude of each of the first 10 fEPSPs within the series of fEPSPs is plotted in function of its rank in the series. (A) Amplitudes of the fEPSPs induced by a stimulation train in a same group of six slices before apamin application, at the 45th min of the drug application, and 45 min after the end of apamin application. (B) Same display as in A, but the stimulation trains were applied on another set of eight slices before, 15 min after the start, and 60 min after the end of apamin perfusion. (C) Same display and same pattern of train delivery on another set of five slices, as in A, but when APV was present throughout the experiment. (D) Illustration of the lack of effect of apamin on the global depolarization induced by a high-frequency stimulation train when APV was coapplied with apamin. The first seven fEPSPs of two series of fEPSPs elicited before (thin trace) and 45 min after apamin application are superimposed.

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The dependence of the apamin-induced increase in depolar-
ization on the influx of Ca<sup>2+</sup> though the NMDA receptors was confirmed in experiments where APV was applied throughout the experiment, before, during, and after apamin application. As APV prevented any Ca<sup>2+</sup> influx through the NMDA receptors, there was no opening of the SK channels during a stimulation train, no matter when it was applied—before, during, or after apamin application. This phenomenon clearly emerges from the comparison of Figure 6C with Figure 6A. Under APV (n = 5), there was no difference between the total depolarizations recorded before, during (45th min), and after the end of (45 min) apamin treatment (Fig. 6C, two-way ANOVA with one repeated measure, F<sub>(2,12)</sub> = 1.8, P = 0.2).

Delayed facilitation of L-LTP induction by apamin is NMDA receptor-dependent

When APV, a blocker of NMDA receptors, was applied together with apamin, the delayed effect of apamin on LTP was suppressed (Fig. 7A). In these conditions, the LTP induced by a single train of stimulation was an S-LTP, as was the case where no drug was applied. Three hours after induction, the slopes of the fEPSP were similar in the control situation (125% ± 6%, n = 14) and when apamin and APV were previously co-applied (109% ± 10%, n = 5, Mann-Whitney rank sum test, P = 0.18). We ascertained that 45 min after its application, APV on its own had no effect (Fig. 7B). Three hours after induction, the slopes of fEPSP were not different, whether APV had been applied (117% ± 6%, n = 4) or not (125% ± 6%, n = 14, P = 0.15). Thus, we found that the metaplastic effect of apamin on L-LTP induction was NMDA receptor-dependent.

Apamin did not have any delayed influence on the NMDA receptor-mediated fEPSP

Prior application of apamin led to facilitated L-LTP induction. One possibility that could explain these results was an increased

Figure 7. Suppression of the metaplastic effect of apamin on LTP by APV. (A, B) Time course of fEPSP 2 h 15 min before and 3 h after one train of HFS. (A) When APV was coapplied with apamin, the LTP induced after prior application of apamin (C) was not different from the control LTP (●). (B) When APV was applied alone using the same timing as that used for apamin application, the LTP induced after prior application of APV (●) was not different from the control LTP (●).
efficacy of the NMDA receptors at the moment of LTP induction. To put this hypothesis to the test, we compared the fEPSP mediated by the NMDA receptors when apamin was previously applied and when it was not. The fEPSP recorded in the CA1 region in response to a stimulation of the Schaffer collaterals consists of a large, AMPA-receptor-mediated, and a small, NMDA receptor-mediated, component. At rest, the opening of NMDA receptors under the action of glutamate is hindered by the presence of Mg$^{2+}$ ions in their ionic channels. It is thus possible to record the NMDA receptor-mediated component of the fEPSP by blocking the AMPA receptors using CNQX (20 µM). It is also possible to measure an enhanced NMDA receptor-mediated component of the fEPSP by lifting the Mg$^{2+}$ blockage using aCSF free of Mg$^{2+}$ and containing CNQX. The fEPSPs of two groups of eight slices each were compared, one submitted to apamin, the other not. CNQX was applied for 20 min after apamin perfusion (where applied). At the end of this period the input/output relationship of the NMDA-mediated fEPSP [f(NMDA) EPSP] was established (Fig. 8A). At every tested stimulation intensity, the f(NMDA) EPSP amplitudes were not different whether apamin was applied previously or not (Student’s t-test, P > 0.05). After the end of the 20-min application of CNQX, aCSF free of Mg$^{2+}$ containing CNQX was perfused. Around 40 min after the end of prior application of apamin (where applied), the input–output relationships of the enhanced f(NMDA) EPSPs were established (Fig. 8B). At every tested stimulation intensity, the enhanced f(NMDA) EPSPs were not statistically different, either (P > 0.05). This suggests that there was no major delayed modification of the NMDA receptors induced by prior application of apamin.

The delayed effect of apamin on L-LTP induction is mediated by the NO-synthase signaling pathway LTP depends on activation by Ca$^{2+}$/calmodulin of several signaling pathways, among which is the NO-synthase pathway. Here, we tested whether or not the delayed action of apamin on LTP was dependent on this pathway. When L-NAME, an inhibitor of NO-synthase, was applied around the moment of LTP induction, it prevented prior application of apamin from transforming the S-LTP usually triggered by a single stimulation train into an L-LTP. In this case, the time course of the increase in the slope of the fEPSP triggered by one train of HFS delivered after prior application of apamin was not different from that observed when no drug was applied (Fig. 9A). When L-NAME was coapplied with apamin, it did not prevent apamin from having a metaplastic effect on LTP. In this case, the time course of the increase in the slope of the fEPSP triggered by a single stimulation train corresponded to the profile of an L-LTP whether apamin was applied 45 min before LTP induction with L-NAME (Fig. 9B). Three hours after induction, the increase of the fEPSP was still as high as 176% (n = 5), a level not different from that reached in the control situation (Fig. 3B, 125% ± 6, P = 0.59). Interestingly, L-NAME had no effect when coapplied with apamin (Fig. 9B). Three hours after induction, the slope of the fEPSP was still as high as 176% ± 5% (n = 5), a level not different from that observed when apamin was applied alone (Fig. 3B, 167% ± 5%, P = 0.5), but significantly greater than that reached in the control situation (P < 0.001).
Delayed action of apamin on paired-pulse facilitation: A second metaplastic effect of apamin

We next explored how paired-pulse facilitation, known to be due to a presynaptic mechanism, was affected by apamin. Thirty minutes after the end of the recovery period, six slices of a first group were submitted to apamin for 1 h 15 min, whereas five slices of a control group were perfused with plain aCSF. Schaffer collaterals were stimulated using a paired-pulse with a 25-msec interval before apamin application (5 min), during apamin perfusion (at the 60th min), and after the end of apamin treatment (at the 16th min, 36th min, and 56th min). The same paired-pulses were delivered at the corresponding times on the control slices. In the control situation, the initial paired-pulse facilitation in the slope of the fEPSP was 164% ± 5% (n = 6) and remained unchanged over time (one-way ANOVA with one repeated measure, F = 0.39, P = 0.81, Fig. 10A, black columns). In the group of slices treated with apamin, the initial paired-pulse facilitation in the slope of the fEPSP was 175% ± 5% (n = 6). Under the influence of apamin it increased (Fig. 10A, white columns, one-way ANOVA with one repeated measure, F = 3.96, P < 0.05). Paired comparisons using the Student-Newman-Keuls method showed that there was no significant increase during apamin perfusion (at the 60th min), whereas there was indeed a significant increase after apamin application (196% ± 14% at the 16th min and 200% ± 12% at the 36th min). The fact that apamin had no immediate action on paired-pulse facilitation but that it did have a delayed one indicated that we were facing another metaplastic phenomenon. Interestingly, 56 min after apamin application, at a time when apamin did not influence LTP any more, paired-pulse facilitation fell back to a level not statistically different from that measured before apamin application.

Like the metaplastic effect of apamin on LTP, the metaplastic action of apamin on paired-pulse facilitation was NMDA receptor-dependent. APV on its own did not modify the paired-pulse facilitation (152% ± 3% [n = 8] vs. 149% ± 2% in plain cerebrospinal fluid [aCSF], paired Student’s t-test, P = 0.46). When apamin was added in aCSF containing APV, the paired-pulse facilitation did not change under the influence of apamin (Fig. 10B, gray columns) (one-way ANOVA with one repeated measure, F = 1.67, P = 0.15).

Moreover, other stimulation intervals were also tested: Paired shocks with an interpulse interval of 50 msec, 100 msec, and 200 msec were delivered, respectively, 1 min, 2 min, and 3 min after the paired-pulses with an interval of 25 msec. Paired-pulse facilitation was also increased after apamin application when the interpulse interval was 50 msec but not when it was longer (Fig. 10C, Student’s t-test, P < 0.05). During apamin treatment, paired-pulse facilitation was not significantly changed for any tested interpulse intervals (Fig. 10D, Student’s t-test, P < 0.05).

Discussion

The main findings of this work are the following: (1) Apamin, an SK channel blocker, has a metaplastic effect on LTP (see the summary bar graph of Fig. 11A). Forty-five minutes after the end of apamin application, a single train of high-frequency stimulation (HSF) induces a long-lasting LTP (L-LTP) instead of a short-lasting one (S-LTP). In contrast, the presence of apamin around the time of LTP induction does not induce any change in the duration of the triggered LTP. (2) The metaplastic effect of apamin on LTP is mediated by NMDA receptors and is crucially dependent on the NO-synthase pathway (see the summary bar graph of Fig. 11B). (3) Apamin also has a metaplastic effect on paired-pulse facilitation (PPF). PPF is increased after drug application but not during it. This metaplastic effect is also mediated by NMDA receptors. (4) Apamin causes an increase in the global depolarization induced by a train of high-frequency stimulation (100 Hz, 1 sec). This effect is not a metaplastic one. It is fully developed during apamin application (as soon as 15 min, if not sooner, after the start of apamin treatment) and persists ~45 min after the beginning of the washout of the drug.

The delayed actions of apamin are metaplastic effects

The term metaplasticity refers to the fact that synaptic plasticity is sensitive not only to the “state” imposed by extra factors at the moment of induction, but also to the “state” created by certain factors prior to the induction (Abraham and Bear 1996; Abraham and Tate 1997). The starting point of this work consisted of a pair of experiments. In Experiment 1, 45 min after apamin application, a single HFS train induced an L-LTP instead of an S-LTP (Fig. 1A). In contrast, in Experiment 2, when a single HFS train was
Measurements were made 3 h after LTP induction. Another piece of evidence comes from the fact that (+)-apamin-induced transformation of an S-LTP into an L-LTP was already present at the 15th min of apamin perfusion, which was not the case. However, the negative result of the experiment 2 was not due to the fact that the drug had not yet reached its target 15 min after beginning the apamin perfusion. This was ruled out in two ways. First, the negative result of the experiment was confirmed when apamin was perfused during the whole recording period, and thus for 45 min before LTP induction. Second, the fact that the global depolarization triggered by an HFS train was already well established at the 15th min of apamin perfusion proved that SK channels were already blocked at the time of LTP induction in Experiment 2.

The mechanism of the positive result of Experiment 1 could also be questioned. Indeed, apamin is known to induce a long-lasting blockage of SK channels. In this work, we assessed the blockage of SK channels by measuring a consequence of it, the increase in the paired-pulse facilitation elicited by an HFS train. This effect was already present at the 15th min of apamin perfusion, unlike the control situation, it triggered an S-LTP (Fig. 1B).

Because recordings were made in an interface chamber while the slices were perfused at a relatively low perfusion rate (1 mL/min), one had to ascertain that the negative result of Experiment 2 was not due to the fact that the drug had not yet reached its target 15 min after beginning the apamin perfusion. This was confirmed when apamin was perfused during the whole recording period, and thus for 45 min before LTP induction. Second, the fact that the global depolarization triggered by an HFS train was already well established at the 15th min of apamin perfusion proved that SK channels were already blocked at the time of LTP induction in Experiment 2.

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Therefore, the results of Experiments 1 and 2 strongly suggest that the delayed action of apamin on LTP is a novel form of metaplasticity. This is not the first time that metaplastic phenomena concerning the duration of LTP have been reported. Woo and Nguyen (2002) found that when four trains of HFS were preceded 7 min earlier by the application of a train of low-frequency stimulation (3 min, 5 Hz), multiple trains induced an S-LTP instead of an L-LTP. Conversely, we recently reported that a single train of HFS induced an L-LTP instead of an S-LTP when the recovery of the slices occurred in submersion instead of in interface (Capron et al. 2006).

Similarly, the fact that apamin caused an increase in paired-pulse facilitation 16 min and 36 min after its application, whereas it left PPF unchanged during its application, shows that the action of apamin on PPF is also a metaplastic phenomenon.

**Potential molecular mechanisms of apamin-induced metaplasticity**

As far as the underlying molecular mechanisms are concerned, two questions arise. First, how can the action of apamin on SK channels induce, 45 min later, an enhancement of the maintenance of an LTP and an increase in the paired-pulse facilitation? Second, what is the role of the NMDA receptors in triggering this phenomenon?

The mechanisms of LTP are complex (Collingridge et al. 1983; Lynch et al. 1983; Frey et al. 1993; Nguyen et al. 1994; Abel et al. 1997; Giese et al. 1998; Kandel 2001). Basically, during LTP induction, Ca\(^{2+}\) enters postsynaptic spines through the NMDA receptors. Next, the newly entered Ca\(^{2+}\) links to calmodulin (CaM), and the complex Ca\(^{2+}\)/CaM activates several enzymes, among which are CaM kinase II, adenylate cyclase, and NO synthase (O’Dell et al. 1991; Haley et al. 1992; Williams et al. 1993; Schuman and Madison 1994; Arancio et al. 1996; Ko and Kelly 1999). Here, we demonstrated that the NO-synthase pathway was crucial for the expression of the metaplastic effect of apamin on LTP (Fig. 9). The fact that the NMDA-mediated fEPSP was unchanged (Fig. 8) strongly suggests that the Ca\(^{2+}\) influx occurring during LTP induction was not changed by prior application of apamin. Therefore, the modified element in the Ca\(^{2+}\)/CaM-NOS-synthetic signaling pathway must be at least one step beyond the influx of Ca\(^{2+}\). One possibility is that the affinity of calmodulin for Ca\(^{2+}\) would be increased. This is not impossible. For instance, it is known that the affinity of the CaM molecules constitutively bound to SK channels for Ca\(^{2+}\) (Fanger et al. 1999; Keen et al. 1999; Xia et al. 1998) vary in function of their state of phosphorylation at threonine 80 (Bild et al. 2004). Moreover, this state of phosphorylation is regulated by a protein kinase (CK2) and a protein phosphatase (PP2A) attached to SK channels. We speculate that it is not impossible that apamin would induce a modification of the CaM molecules attached to SK channels. Over time, this modified CaM could dissociate from the SK channels and play an enhanced role in the Ca\(^{2+}\)/CaM-NOS-synthetic signaling pathway.

Paired-pulse facilitation (PPF) of central synaptic transmission is primarily due to increased presynaptic glutamate release under the influence of residual calcium (Creager et al. 1980; McNaughton 1982; Muller and Lynch 1989; Manabe et al. 1993; Christie and Abraham 1994; Schulz et al. 1994). However, postsynaptic Ca\(^{2+}\)/CaM signaling pathways can modulate PPF (Wang and Kelly 1996). This involves a Ca\(^{2+}\)/CaM-activated CaMKII pathway leading to an increase in the desensitization of AMPA receptors and likely a Ca\(^{2+}\)/CaM phosphatase pathway causing a decrease in the desensitization of AMPA receptors (Wang and Kelly 1996).
Kelly 1996). Activation of kinases necessitates higher concentrations of Ca\(^{2+}\)/CaM than activation of phosphatases. For example, the protein phosphatase calcineurin is activated by 450-fold lower concentrations of Ca\(^{2+}\)/CaM relative to CaMKII (Mulkey et al. 1994). In the control situation (in absence of apamin), the postsynaptic influx of Ca\(^{2+}\) resulting from a single pulse is probably not sufficient to activate either CaMKII or phosphatases. After prior application of apamin, the hypothesized apamin-induced enhanced affinity of CaM for Ca\(^{2+}\) could lead to an increase in the concentration of Ca\(^{2+}\)/CaM reached after the first of the paired-pulses. This could be sufficient to activate phosphatases and hence decrease the desensitization of AMPA receptors. Consequently, paired-pulse facilitation could be increased.

Another possible mechanism is worth considering. Apamin, being a blocker of a synaptically located ion channel, might be exerting its meta-plastic effects on the synaptogenesis that occurs as a result of slicing (Kirov et al. 1999). However, this is unlikely. Slicing-induced synaptogenesis occurs during 2 h after preparing the slices and ceases afterward. In contrast, the meta-plastic effects of apamin were observed not only when apamin was applied just after slicing (Fig. 1A) but also 2 h after it (Fig. 3B).

Like a classic meta-plastic effect (i.e., stimulation of a synapse at frequencies ranging from 5 to 30 Hz causes a dramatic reduction in subsequently evoked LTP) (Huang et al. 1992; O’Dell and Kandel 1994; Zhang et al. 2005), the two meta-plastic effects of apamin reported here are also dependent on NMDA receptors. This implies that an NMDA receptor-mediated Ca\(^{2+}\) influx is necessary for the induction of the meta-plastic phenomena induced by apamin. During apamin application, NMDA receptors were weakly activated due to the glutamate released by test stimulations (when apamin was applied in interface during the recording period) and to that released as a result of the slicing trauma (when apamin was applied during the recovery period). This small Ca\(^{2+}\) influx could activate phosphatases, like PP2A—associated with the SK channels—and so lead to an enhanced affinity of CaM for Ca\(^{2+}\).

Because of this dependence of the meta-plastic effects of apamin on NMDA receptors, we have hypothesized that a post-synaptic mechanism explains the phenomenon. However, as SK3 channels have been found to be located on the presynaptic terminals of cultured hippocampal pyramidal neurons (Obernair et al. 2003), we cannot exclude the possibility that apamin might exert its effect via a presynaptic mechanism.

**Influence of apamin on the depolarization induced by a single train**

There is histochemical (Ngo-Anh et al. 2005) and electrophysiological (Faber et al. 2005) evidence showing that SK channels are colocalized with NMDA receptors in dendritic spines of glutamatergic synapses. It can be anticipated that during tetanic stimulation of glutamatergic synapses, the Ca\(^{2+}\) that entered through the NMDA receptors would open SK channels and consequently shut the next EPSP. As a result, the summation of EPSPs during tetanic stimulation is expected to be enhanced when SK channels are blocked by apamin. This was shown to be the case in the lateral amygdala (Faber et al. 2005). In this work, we demonstrated that such a phenomenon occurred in the synapses between the Schaffer collaterals and the CA1 pyramidal neurons. We also confirmed that it was NMDA receptor-dependent, as its development was blocked by APV.

**Materials and Methods**

**Animals**

Male C57BL/6 mice, aged 6–10 wk (Charles River) were used for all the experiments, which were carried out in accordance with National Institutes of Health regulations for the care and use of animals in research and with local ethics committee guidelines.

**Electrophysiology**

Transverse hippocampal slices (400-μm thickness) were prepared as described by Nguyen and Kandel (1997). The hippocampus was isolated and sliced with a McIlwain chopper. Slices were allowed to recover for 1.5 h at 28°C in interface. After this recovery period all the recordings were made in an interface chamber (IST) at 28°C. The rate of flow of the perfused liquid was 1 mL/min. Baseline recording started 15 min later and lasted 30 min. Then LTP was induced electically by applying a single 1-sec train (100 Hz, at test strength). Slices were perfused with ACSF of the following composition: 124 mM NaCl, 5 mM KCl, 26 mM NaHCO\(_3\), 1.0 mM NaH\(_2\)PO\(_4\), 2.5 mM CaCl\(_2\), either 1.3 or 1 mM MgSO\(_4\), 10 mM glucose. The ACSF was aerated with 95% O\(_2\) and 5% CO\(_2\). Extracellular fEPSPs were recorded with a glass micro-electrode (2–5 Mohm, filled with ACSF) positioned in the stratum radiatum of area CA1. A bipolar nickel–chromium stimulating electrode was used to elicit fEPSPs by stimulation of the Schaffer collateral fibers. Stimulation intensity (0.08-msec pulse duration) was adjusted to elicit fEPSP amplitudes that were –40% of maximum size. Basal synaptic transmission was assessed by stimulating Schaffer collaterals once per minute at this test stimulation intensity. Slices that showed maximal fEPSPs <2 mV were rejected.

Apamin (Tocris), 2-amino-5-phosphonopentanoic acid (APV, Acros Organics), and L-NAME (Sigma) were prepared as a concentrated stock solution in H\(_2\)O. (+)-Methyl-laudanosine was prepared in DMSO and subsequently diluted at 0.1% to obtain the final concentration.

**Data analysis**

For each slice, the fEPSP slopes were normalized against the average slope over the 30 min before LTP induction. The statistical significance (P<0.05) of the difference in increase of fEPSP measured 3 or 4 h after two different treatments applied to two distinct groups of slices was assessed using a Student’s t-test. When another test was used, it is indicated in the text.

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**References**


