Deprivation-induced synaptic depression by distinct mechanisms in different layers of mouse visual cortex

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Long-term depression (LTD) induced by low-frequency synaptic stimulation (LFS) was originally introduced as a model to probe potential mechanisms of deprivation-induced synaptic depression in visual cortex. In hippocampus, LTD requires activation of postsynaptic NMDA receptors, PKA, and the clathrin-dependent endocytosis of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors. It has long been assumed that LTD induced in visual cortical layer 2/3 by LFS of layer 4 uses similar mechanisms. Here we show in mouse visual cortex that this conclusion requires revision. We find that LTD induced in layer 2/3 by LFS is unaffected by inhibitors of PKA or α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor endocytosis but is reliably blocked by an endocannabinoid CB1 receptor antagonist. Conversely, LFS applied to synapses on layer 4 neurons produces LTD that appears mechanistically identical to that in CA1 and is insensitive to CB1 blockers. Oclusion experiments suggest that both mechanisms contribute to the loss of visual responsiveness after monocular deprivation.

Endocannabinoid | glutamate receptor | long-term depression | ocular dominance plasticity

For decades, neuroscientists have sought to understand how brief monocular deprivation (MD) in early life causes a loss of visual responsiveness in visual cortex that results in amblyopia. The laminar structure of the neocortex has allowed researchers to investigate the mechanisms of synaptic depression both in vivo and in vitro in a layer-specific manner. The feed-forward circuit is as follows: thalamus → layer 4 → layer 2/3 → layer 5 (1). Long-term depression (LTD) of synaptic responses in visual cortical layer 2/3 induced by low-frequency stimulation (LFS) of layer 4 is a widely studied phenomenon that is believed to share mechanisms with those that cause synaptic depression after MD (2). Based on similar frequency and NMDA receptor (NMDAR) dependence, it has long been assumed that mechanisms of layer 2/3 LTD are the same as those in area CA1 of the hippocampus (3, 4), which involve postsynaptic calcium flux through NMDARs, dephosphorylation of postsynaptic PKA substrates, and clathrin-dependent internalization of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors (AMPARs; ref. 5). Indeed, MD produces changes in AMPAR phosphorylation and surface expression in rat visual cortex that mimic CA1 LTD (6).

In the meantime, a protocol for the induction of LTD by precisely timed pre- and postsynaptic action potentials has gained popularity (7). That spike-timing-dependent LTD (STD-LTD), like LFS-dependent LTD (LFS-LTD), is also blocked by bath application of an NMDAR antagonist initially suggested that common mechanisms are engaged by the two types of induction protocol (8). However, this notion was soon challenged by evidence that STD-LTD in layer 5 neurons from neonatal visual cortex depends on endocannabinoid signaling and pre- rather than postsynaptic NMDAR activation (9).

These interesting findings compelled us to reexamine with greater precision the question of how LFS induces NMDAR-dependent LTD in the radial excitatory pathways of visual cortex. This issue is of considerable significance, because CA1 LTD has been used to inform hypotheses about the molecular basis of ocular dominance plasticity (2, 6). We performed this study in mouse visual cortex, because this has emerged as a favored preparation to study the mechanisms of ocular dominance plasticity in vivo. LTD-LTD induced in layer 4 by white-matter stimulation was compared with LFS-LTD induced in layer 2/3 by layer 4 stimulation. We found that identical LFS protocols induce comparable LTD in both locations by activation of postsynaptic NMDARs, as expected. To our surprise, however, we found that the mechanism of layer 2/3 LTD is qualitatively different from that of CA1 and requires activation of cannabinoid receptors. On the other hand, layer 4 LTD is very similar to that of CA1, notably including sensitivity to inhibitors of PKA and AMPAR endocytosis. We further found that both types of LTD are reduced in visual cortex after a period of MD in vivo.

Together, our results show that LTD induced by identical stimulation protocols in mouse visual cortex depends on distinct mechanisms in layers 3 and 4. Because deprivation-induced synaptic depression in vivo occludes both types of LTD, our findings strongly suggest that different mechanisms contribute to the effects of MD in different layers of mouse visual cortex.

Results

Reliable LTD Expression in Layers 3 and 4. We first sought to establish that LTD could be induced with an LFS protocol in both layers 2/3 and 4. Somatic whole-cell voltage-clamp recordings were obtained from pyramidal neurons in either layer in slices prepared from mice at postnatal day (P)21–28. Excitatory postsynaptic currents (EPSCs) were elicited in layer 2/3 by layer 4 stimulation or in layer 4 by white-matter stimulation (just below layer 6). After achieving a stable baseline, repeated brief postsynaptic depolarizations were paired with LFS to induce LTD (Fig. 1). This protocol reliably induced LTD in layer 2/3 (EPSC amplitude after LFS = 65.7 ± 2.6% of baseline, n = 5; Fig. 1a). In layer 4, previous reports suggested considerable variability in the ability to induce LTD with LFS (compare refs. 4 and 10). However, we found that our pairing protocol also reliably induced LTD in layer 4 (71.5 ± 8.1% of baseline, n = 5; Fig. 1b).

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Abbreviations: LFS, low-frequency stimulation; LTD, long-term depression; STD-LTD, spike-timing-dependent LTD; EPSC, excitatory postsynaptic current; EPSP, excitatory postsynaptic potential; AMPA, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid; AMPAR, AMPA receptor; MD, monocular deprivation; NMDAR, NMDA receptor; Pn, postnatal day n; APV, α-2-amino-5-phosphonovalerate.

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Mechanistic Similarities in LTD Between Layers 2/3 and 4. We then asked whether LTD in both layers was similarly NMDAR-dependent. As expected from previous work (5, 4, 11), LTD in layer 2/3 was prevented by bath-applied 100 μM dl-2-amino-5-phosphonovalerate (APV). EPSC amplitude after LFS was 106.7 ± 9.2% of baseline in the presence of APV (n = 5; Fig. 1a). Similarly, APV blocked LTD in layer 4 (97.3 ± 3.1% of baseline, n = 3; Fig. 1b). To disambiguate the role of pre- vs. postsynaptic NMDAR in layers 2/3 and 4, we loaded individual neurons with MK801 (500 μM), which can block postsynaptic NMDAR channels intracellularly (12–15). We found that LTD was almost entirely blocked by intracellular MK801 in layers 2/3 (91.2 ± 6.3% of baseline, n = 6; Fig. 1a) and 4 (94.3 ± 2.8% of baseline, n = 7; Fig. 1b). Interleaved control recordings yielded robust LTD in both layers (layer 2/3, 65.5 ± 6.2%; n = 5; layer 4, 73.8 ± 3.9%; n = 6).

To ensure that MK801 introduced postsynaptically does not leak out to affect presynaptic NMDAR, we studied STD-LTD in layer 3 of juvenile mice, which, like STD-LTD in layer 5 (9), is sensitive to blockade of pre-but not postsynaptic NMDARs (15). In contrast to LTD-LTD and in support of those prior studies, blocking postsynaptic NMDARs with 500 μM MK801 intracellularly did not prevent STD-LTD [excitatory postsynaptic potential (EPSP) amplitude, 62.4 ± 11.9%; slope, 62.4 ± 15.3%, n = 7; Fig. 2]. However, bath-applied APV prevented STD-LTD (EPSP amplitude, 95.5 ± 6.7%; slope, 99.1 ± 13.9%, n = 7). In addition to confirming that STD-LTD is insensitive to blockade of postsynaptic NMDARs, these experiments show that MK801 applied intracellularly does not leak out to affect presynaptic NMDARs, in which case the effect of MK801 on STD-LTD would be identical to bath-applied APV. Together, the data suggest that the induction of LTD by identical LFS protocols in both layers 2/3 and 4 requires postsynaptic NMDAR activation.

Mechanistic Differences in LTD Between Layers 2/3 and 4. The dependence on postsynaptic NMDAR activation for LTD-LTD in both layers 2/3 and 4 of mouse visual cortex allows specific predictions about the molecular cascades stimulated by calcium influx through NMDARs, based on work performed in CA1. Consistent findings in CA1 are inhibition of LTD by blockers of PKA and AMPAR endocytosis. Therefore, we next performed intracellular loading experiments, using either the selective PKA inhibitor PKI (6–22 amide; 10 μM) or a peptide (termed G2CT; 10 μM, KRMKLNIPSN) that competes with endogenous GluR2 for the AP2 clathrin adaptor complex and blocks AMPAR endocytosis (16).

We were very surprised to find that the magnitude of LTD in layer 2/3 cells loaded with PKI (66.6 ± 7.6%, n = 4; Fig. 3a) was indistinguishable from LTD in interleaved control experiments (67.7 ± 8.3%, n = 5). Similarly, loading neurons with the G2CT peptide had no effect on LTD in layer 2/3 (70.4 ± 4.8%, n = 7, vs. interleaved control LTD, 68.8 ± 2.4%, n = 6; Fig. 3a). However, the same reagents loaded into layer 4 neurons strongly blocked LTD induced by the same induction protocol. LTD in layer 4 was almost completely prevented by PKI (94.1 ± 5.1%, n = 5; control, 74.8 ± 5.3%, n = 7; Fig. 3b).

These results suggest that LTD induced in layer 4 by stimulating radial inputs involves the PKA-regulated endocytosis of GluR2-containing AMPARs, as has been proposed for CA1. However, what could account for the NMDAR-dependent LTD in layer 2/3? Studies of STD-LTD between pairs of visual cortex layer 5 neurons first revealed a potential mechanism involving retrograde endocannabinoid signaling by presynaptic CB1 receptors (9). The possibility that this mechanism could also contribute to LTD in layer 2/3 was also recently suggested by studies in rat barrel cortex (15). We therefore investigated the effect of the highly potent CB1 receptor antagonist AM251 (2...
Occlusion of LTD in Layers 3 and 4 by Previous MD. These findings, considered with an earlier report of differences in LTD induction requirements in the superficial and deep layers of rat visual cortex (4) potentially impact the conclusion that MD induces CA1-type LTD in visual cortex in vivo (6). That conclusion, from a study of rat visual cortex, was supported by two lines of evidence. First, it was found that MD mimics LTD with respect to depressed synaptic transmission, altered AMPAR phosphorylation, and decreased AMPAR surface expression. Second, it was found that synaptic depression caused by MD occludes LTD in layer 2/3 studied ex vivo. However, the current findings in mice suggest that occlusion of LTD in layer 2/3 may not adequately address the question of whether AMPAR internalization is a mechanism for ocular dominance plasticity. Therefore, we thought it was important to reexamine the effect of MD in vivo on the CA1-type LTD in layer 4 and the CB1-mediated LTD in layer 2/3 in mouse visual cortex ex vivo.

Mice were monocularly deprived at P28 for 3 days, a duration that produces maximal depression of visually evoked potentials recorded from layer 4 in vivo (17) without inducing metaplasticity, which could alter the induction requirements for LTD (18). On the third day, slices were prepared from visual cortex contralateral and ipsilateral to the deprived eye. Rodents show an interhemispheric asymmetry in the effect of MD; visual responses in the cortex contralateral to the deprived eye are depressed relative to the ipsilateral hemisphere (6). Thus, the contralateral hemisphere can be viewed as “deprived” relative to the same-animal ipsilateral “control” hemisphere.

With the experimenter blind to the visual history of the mice from which slices were obtained, whole-cell voltage-clamp recordings were performed from layer 2 neurons from the binocular zone of each hemisphere. LTD was induced with the same pairing protocol but for twice the duration as the previous recordings, to saturate the magnitude of LTD. That we observed no further depression after doubling the number of stimuli in control (nondeprived) slices confirmed that LTD is saturated and, further, that extended stimulation does not recruit alternate (e.g., CB1-dependent) mechanisms of LTD (compare Figs. 1 and 4). The results clearly showed that LTD at saturation is reduced in the hemisphere contralateral to the deprived eye (90.1 ± 3.6%, n = 6; Fig. 5b) compared with LTD in the ipsilateral hemisphere (72.6 ± 3.3%, n = 5).

We next asked whether LTD in layer 2/3 of mouse visual cortex is similarly disrupted by prior MD. LTD in layer 2/3 was significantly reduced (89.5 ± 3.6%, n = 7) in the visual cortex contralateral to the deprived eye compared with values obtained in the control ipsilateral hemispheres (73.8 ± 5.4%, n = 6; Fig. 5a) confirming previous findings in rats. Together, these data are consistent with the hypothesis that deprivation-induced synaptic depression in vivo utilizes, at least in part, the mechanisms of layer 4 (CA1-like) and layer 2/3 (CB1-mediated) LTD described here in brain slices.

Discussion

Modifications of synapses in the radial excitatory pathways afferent to layers 4 and 3 are responsible for the functional consequences of MD in the binocular region of visual cortex (19, 20). Motivated by a theoretical analysis of ocular dominance plasticity (21), LFS-LTD was introduced as an experimental paradigm that might help reveal the mechanisms (22). Here we show that, although activation of postsynaptic NMDARs by identical LFS protocols induces synaptic depression in both layers, there are significant laminar differences in the mechanism, and that LTD in layer 2/3 is mechanistically different than previously thought. These findings suggest new opportunities to dissect the individual contributions of plasticity in different layers to the ocular dominance shift in mice.

Crozier et al.
The first synaptic relay from the thalamus occurs in layer 4, which is a site of robust ocular dominance plasticity in mice (17, 23). Although extracellular stimulation of white matter activates these thalamic synapses (24), it is very likely that other sources of synaptic input are also recruited (e.g., recurrent collaterals). Considering this potential complication, it is remarkable that the LTD induced by our stimulation protocol was reliably blocked in all experiments by APV and postsynaptic inhibition of NMDAR channels, PKA, and AMPAR endocytosis. Thus, the large majority of synapses capable of expressing LTD on layer 4 neurons in response to LFS of the white matter use a mechanism that closely resembles what has been documented extensively in area CA1 of hippocampus (reviewed in ref. 5).

We hasten to point out that AMPAR endocytosis may not be the only mechanism for LTD in layer 4, particularly at intracortical synaptic connections. In synaptically coupled pairs of spiny stellate neurons in layer 4, LFS pairing of layer 4 inputs results in a more pronounced LTD than LFS pairing of layer 2/3 inputs. This is consistent with our previous findings and the observed differences in LTD expression between layers 2/3 and 4 (17, 23). The LTD protocol was administered at time 0 and lasted 5 min. The dashed horizontal line indicates no change from baseline responses. Sweep numbers (1 and 2) refer to averaged responses collected during the last 5 min of the baseline and postpairing periods. Stimulation artifacts were minimized for clarity. (Scale bars: 50 pA, 20 msec.) See Fig. 4 for statistical comparisons.

**Fig. 3.** Differences in pairing-induced LTD in layers 4 and 3 of visual cortex. (a and b) Displayed are grouped data time courses and representative sweeps from layers 2/3 (a) and 4 (b) recordings. LTD was challenged both extracellularly and intracellularly by a battery of antagonists, which highlight differences in LTD expression between layers. (a1 and b1) Schematic illustrations of recording configurations for layers 2/3 (a1) and 4 (b1). (a2 and b2) Intracellular loading of a PKA inhibitor PKI (6–22 amide, 10 μM) prevented LTD in layer 4 (b2) but not in layer 2/3 (a2). (a3 and b3) Intracellular loading of G2CT (10 μM), a peptide that interferes with regulated endocytosis of GluR2, also prevents LTD in layer 4 (b3) but not layer 2/3 (a3). (a4 and b4) Bath application of the CB1 antagonist AM251 (2 μM) had no effect on LTD in layer 4 (b4) but largely prevented LTD in layer 2/3 (a4). The y axis is EPSC amplitude normalized to a 15-min baseline period, with error bars indicating the SEM. The pairing protocol, denoted by the black bar, was administered at time 0 and lasted 5 min. The dashed horizontal line indicates no change from baseline responses. Sweep numbers (1 and 2) refer to averaged responses collected during the last 5 min of the baseline and postpairing periods. Stimulation artifacts were minimized for clarity. (Scale bars: 50 pA, 20 msec.) See Fig. 4 for statistical comparisons.

**Fig. 4.** Summary data comparing laminar differences in LTD. Filled bars represent mean ± SEM of EPSC 25–30 min after LFS (normalized to baseline) in neurons treated with the drugs indicated. Gray bars represent the effect of LFS in the interleaved vehicle-treated controls. The number of experiments in each group is indicated in the text. Note the consistency of LTD magnitude in all control experiments. APV and AM251 were bath-applied. MK801, PKI, and G2CT were loaded intracellularly. * P < 0.01.
neurons in layer 4 of barrel cortex, for example, Egger et al. (25) reported a form of LTD induced with a spike-timing protocol that required activation of postsynaptic group II metabotropic glutamate receptors but not NMDARs (25). If these findings generalize to layer 4 of visual cortex (where most spiny neurons are pyramidal), they suggest that the diversity of mechanisms for LTD could parallel the diversity of sources of excitatory synaptic input to layer 4 neurons. Nonetheless, that prior MD decreases the magnitude (occludes) LTD of the radial input to layer 4 strongly supports the conclusion that naturally occurring synaptic depression similarly utilizes a mechanism of NMDAR-dependent endocytosis of AMPAR that is reliably elicited with LFS. Thus, our data support the previous conclusions of Heynen et al. (6) in rats and extend them to layer 4 of mice.

We were initially surprised to find that LTD induced in layer 2/3 by the same LFS protocol was insensitive to inhibitors of PKA and AMPAR endocytosis, but sensitive instead to antagonists of CB1 receptors. However, there is certainly ample precedent for laminar diversity (4) and endocannabinoid-dependent LTD in the brain (15, 26, 27). Indeed, paired recordings of synaptically coupled layer 5 neurons have revealed a form of LTD that requires the calcium-dependent release of endocannabinoid from the postsynaptic neuron that acts on presynaptic CB1 receptors (34). Our finding that LTD within layer 4 is insensitive to CB1 antagonist is consistent with the fact that MK801 (1 mM) had no effect on STD-LTD induction, whereas it was blocked by antagonists of metabotropic glutamate receptor 5 (mGlur5). They conclude that the relevant source of postsynaptic calcium for induction of STD-LTD is release from intracellular stores. These findings are in clear contrast with what has been observed with LFS-LTD. First, there is no effect of either group 1 mGluR antagonists or genetic deletion of mGlur5 on LTD-LTD in layer 2/3 of rat or mouse visual cortex (4,32). Second, as we show here, LTD-LTD is reliably blocked by intracellular MK801 (500 μM). As a control for the possibility that MK801 might have leaked out of our recording pipette to affect presynaptic NMDARs implicated in STD-LTD, we repeated the experiments of Bender et al. (15,31) using their spike-timing protocol in visual cortex of juvenile mice (Fig. 2). Consistent with their results, intracellular MK801 (500 μM) had no effect on STD-LTD that was reliably blocked by APV. These experiments confirm that MK801 applied through our pipettes does not meaningfully affect presynaptic NMDARs. Thus, we propose that postsynaptic NMDAR activation triggers the CB1-dependent LTD induced by LFS in layer 2/3 of mouse visual cortex.

Ocular dominance plasticity occurs in both layers 2/3 and 4. There is evidence in cats that plasticity in layer 2/3 occurs independently from that in layer 4 (e.g., late in the critical period (33)), and, further, that it may precede and actually be required for plasticity in layer 4 (34). Our finding that LTD within layer 4 is insensitive to CB1 antagonist is consistent with the fact that there is very little CB1 protein in this layer (35). Our results support the general notion of laminar differences in the mechanisms of deprivation-induced response depression and suggest an opportunity to dissect their contributions to ocular dominance shifts. Because CB1 antagonists, like AM251, can completely block cannabinoid-dependent processes, including LTD in layers 3 and 5, they can potentially be used to pharmacologically isolate cannabinoid-independent mechanisms, like those used for LTD in layer 4, to assess their contributions to ocular dominance plasticity in vivo.

**Materials and Methods**

**Slice Preparation for Whole-Cell Recordings.** Slices of visual cortex (350 μm) from P21–P28 or P13–P15 C57BL/6 mice (Fig. 2) were
prepared as described (36). Briefly, mice were anesthetized, and the brains were rapidly removed and placed in ice-cold high-sucrose dissection solution and sectioned using a vibrating-blade microtome (Leica, Deerfield, IL; VT1000S). Slices containing primary visual cortex were transferred to a holding chamber with artificial cerebrospinal fluid containing 124 mM NaCl, 2.5 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 1.23 mM NaH₂PO₄, 26 mM NaHCO₃, and 10 mM glucose. Slices were left undisturbed at 32°C for 1 h and subsequently stored at room temperature for the remainder of the day. Recordings were performed at 28–30°C. Individual neurons were visualized with IR-DIC optics by using a Nikon (Florham Park, NJ) microscope (E600FN) and 60× water-immersion objective.

**Voltage-Clamp Recordings of STD-LTD.** Somatic whole-cell voltage-clamp recordings (37) were obtained from spiny pyramidal neurons in layer 2/3 and star pyramids in layer 4 by using patch pipettes filled with internal solution containing 107 mM D-glucosic acid, 107 mM CsOH, 5 mM QX-314-Cl, 0.2 mM EGTA, 20 mM Heps, 4 mM Mg-ATP, 0.3 mM Na-GTP, and 10 mM Na-phosphocreatine with pH adjusted to 7.2. When filled with internal solution, pipette resistances were ~5 MΩ. Piezo-electrically determined liquid-junction potential (approximately ~12 mV) was corrected (38). EPSCs were elicited with a two-contact cluster electrode (FHC, Bowdoin, ME) placed in layer 4 for layer 2/3 recordings or in white matter for layer 4 recordings. Test stimuli (100 μsec) were delivered at 0.05 Hz. The test holding potential was −65 mV. LFS-LTD was induced by pairing 1-Hz presynaptic stimulation with a brief (100-msec) postsynaptic-step depolarization from −65 to −45 mV for each of the 300 or 600 pulses (Fig. 5). Each presynaptic stimulation occurred midway (50 msec) into the step depolarization. Series resistance was estimated every fifth sweep by measuring the peak of the capacity transient elicited by a 5-mV hyperpolarizing pulse from −65 to −70 mV, and experiments were discarded if this value changed by >20% during the recording. Data acquisition and analysis were performed on a personal computer running pClamp 9.2 (Molecular Devices, Sunnyvale, CA).

**Current-Clamp Recordings of STD-LTD.** The internal solution consisted of 100 mM K-glucurate/20 mM KCl/4 mM Mg-ATP/0.3 mM GTP/10 mM phosphocreatine/10 mM Heps, pH 7.3. Pipette resistances were ~6 MΩ when filled with internal solution. For current-clamp recordings of STD-LTD (Fig. 2), test stimuli (0.3 msec) were delivered at 0.1 Hz, and STD-LTD was induced by pairing presynaptic stimulation ~25 msec after a postsynaptic spike repeated for 100 times at 0.2 Hz. The postsynaptic spikes were elicited by injecting a depolarizing current (0.2 msec in duration) and were of sufficient magnitude to yield an action potential for every trial. Picrotoxin (20 μM) was present for the duration of the STD-LTD experiment. Recordings were considered acceptable if membrane potentials were maintained between −55 and −70 mV. EPSPs were acquired and analyzed on a Macintosh (Apple, Cupertino, CA) computer by using custom routines written in Igor (WaveMetrics, Lake Oswego, OR).

**Reagents.** The following reagents were prepared as stock solutions in distilled water, stored at ~80°C, and diluted 1:1,000 on the day of use: 100 μM APV (Calbiochem, La Jolla, CA), 10 μM PKI (Calbiochem); 10 μM G2CT peptide (KRMKLNINPS; Massachusetts Institute of Technology, CCR Biopolymers Laboratory, Cambridge, MA); 500 μM MK801 (Sigma, St. Louis, MO). For experiments involving intracellular loading of MK801, a cell was patched, and then the slice was rinsed for 15–20 min such that a minimum of 25–30 min had elapsed before induction of LFS- or STD-LTD to ensure clearance of MK801 from the extracellular milieu. Biocytin 0.2–0.4% (wt/vol; Molecular Probes, Eugene, OR) was included in pipette solutions for post hoc examination. Biocytin was detected with streptavidin AlexaFluor 488 (Molecular Probes) and imaged on a confocal microscope (Olympus, Melville, NY). AM251 (2 μM; Tocris Cookson, Bristol, U.K.) was diluted 1:40,000 in DMSO and also contained BSA at a final concentration of 0.5 mg/ml to prevent AM251 from adhering to the perfusion line. Lid suture for MD was performed as described (6, 17).

**Stats.** The peak amplitude of the EPSC or EPSP was measured. A 15-min (LFS-LTD) or 10-min (STD-LTD) baseline period just before the respective conditioning protocol was averaged and compared with the average of the final 5 min of the recording. Data are displayed as percent baseline ± SEM. Statistics were computed by using StatView 5 (SAS Institute, Cary, NC). Statistical differences were confirmed by one-way analysis of variance with Fisher’s least-squares difference comparisons among groups, and data were considered statistically significant at P < 0.05.

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