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Nucleus basalis-enabled stimulus-specific plasticity in the visual cortex is mediated by astrocytes

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Although cholinergic innervation of the cortex by the nucleus basalis (NB) is known to modulate cortical neuronal responses and instruct cortical plasticity, little is known about the underlying cellular mechanisms. Using cell-attached recordings in vivo, we demonstrate that electrical stimulation of the NB, paired with visual stimulation, can induce significant potentiation of visual responses in excitatory neurons of the primary visual cortex in mice. We further show with in vivo two-photon calcium imaging, ex vivo calcium imaging, and whole-cell recordings that this pairing-induced potentiation is mediated by direct cholinergic activation of primary visual cortex astrocytes via muscarinic AChR. The potentiation is absent in conditional inositol 1,4,5 trisphosphate receptor type 2 KO mice, which lack astrocyte calcium activation, and is stimulus-specific, because pairing NB stimulation with a specific visual orientation reveals a highly selective potentiation of responses to the paired orientation compared with unpaired orientations. Collectively, these findings reveal a unique and surprising role for astrocytes in NB-induced stimulus-specific plasticity in the cerebral cortex.

acetylcholine | response potentiation | gial calcium | basal forebrain | astrocyte-neuron interactions

Sensory experience associated with nucleus basalis (NB)-driven, cholinergic activation of the cortex (1) has been shown to induce cortical plasticity at both single-cell and cortical map levels (2–6). To understand how cortical responses and representations can be altered by experience during cholinergic modulation, it is critical to identify the circuit elements involved and define how their interactions can contribute to the restructuring of cortical network dynamics.

Previous studies have shown that multiple cortical cell types, including neurons (7–9) and astrocytes (10–12), can be responsive to ACh. Among these cell types, astrocytes are a promising candidate for contributing to NB-mediated cortical plasticity. Ex vivo studies have implicated hippocampal astrocytes in synaptic potentiation [(13–15) compare with (16)], demonstrating that they can potentially provide a powerful means of altering the state of neuronal networks to induce plasticity. More recently, studies using combined somatosensory and cholinergic stimulation have revealed that NB-induced astrocytic activation can induce potentiation of local field potentials recorded in somatosensory cortex (17, 18). These findings open up several key questions. Does the NB-mediated potentiation manifest at the level of single neurons and astrocytes? If so, does the potentiation influence specific features of single neuronal responses and representations? In particular, is the potentiation a nonspecific increase in responses independent of sensory stimulus features, or does it selectively facilitate responses to stimuli that have been paired with NB stimulation?

The primary visual cortex (V1) provides an excellent model system to address these issues. Modulation by ACh in general and cholinergic drive from the NB in particular are known to influence V1 circuits in multiple ways, resulting in an enhancement of direction and orientation selectivity in V1 neurons (19–22), increase in attentional modulation of V1 neurons in behaving monkeys (23), and alteration in the reliability and synchrony of stimulus-evoked spikes in V1 neurons (24). Responses of mature V1 neurons to specific visual stimulus features have also been shown to be plastic and depend on the history of visual stimulation (25). Moreover, cortical astrocytes have previously been demonstrated to be an integral component of V1 circuits because they are visually responsive and can be capable of modulating visually driven neuronal responses (26). We therefore examined the influence of NB-mediated cholinergic activation of astrocytes and neurons in mouse V1 in vivo, and the mechanisms of their interactions in V1 slices ex vivo, using both calcium imaging and electrophysiological recording. We show that pairing electrical stimulation of the NB with visual stimuli can induce potentiation of visual responses in V1 excitatory neurons. The potentiation is facilitated by astrocytes, which are activated by cholinergic inputs from the NB and, in turn, directly influence neuronal responses, and is abolished in mice that lack astrocyte calcium increases due to deletion of astrocytic inositol 1,4,5 trisphosphate receptor type 2 (IP-3R2).

This astrocyte-mediated response potentiation is stimulus-specific, because pairing one stimulus orientation with NB stimulation selectively potentiates the visual response of the paired orientation over other unpaired orientations.

Results

Pairing NB and Visual Stimulation Potentiates Visual Responses in Excitatory Neurons in Vivo. We first investigated if paired NB and visual stimulation can induce potentiation of identified excitatory neuron responses. The NB was stimulated with an implanted bipolar electrode (SI Materials and Methods, In Vivo Surgery) (24), whereas responses of single visual cortical neurons in the supragranular layers were recorded with in vivo cell-attached recordings (Fig. 1 A and D, Upper). The stereotaxic accuracy of the implantation was determined by (i) localization of the electrode tip within the NB as assessed by acetylcholinesterase staining (24) (Fig. 1B) and (ii) the effect of stimulation on desynchronization of the interhemispheric electroencephalogram (24, 27) (Fig. 1C). Excitatory neurons were characterized by their “regular spiking” properties (28), including spike half-widths and peak-to-valley ratios (Fig. S1 A and B), and their responses before, during, and after the pairing paradigm were recorded at single-spike resolution for long durations. The visual stimuli consisted of gratings of

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random orientation designed to evoke robust responses and were presented alternately with blank gray screens (Fig. 1D, Lower and SI Materials and Methods, Visual Stimulation). Stimulating the NB with multiple trains of pulses, where each train (50 pulses at 100 Hz; SI Materials and Methods, NB Stimulation) was paired with a cycle of visual stimulus (Fig. 1D, Lower), induced a prominent sustained slow potentiation of visual responses in excitatory neurons lasting over 40 min (Fig. 1E and F). The firing rates during the alternate visual stimulus and blank gray screen presentations were quantified as “ON” and “OFF” responses, respectively, whereas the ON responses relative to the preceding OFF responses were quantified as “ON-OFF.” The increase was pronounced in both visual ON and ON-OFF responses (Fig. 1F: n = 6 neurons in 6 animals; P << 0.001). Thus, paired NB and visual stimulation reliably potentiates visually driven responses of regular spiking, presumably pyramidal V1 neurons.

In Vivo NB Stimulation Evokes Robust Calcium Responses in Visual Cortical Astrocytes via Muscarinic Receptors. Next, we use in vivo two-photon imaging (Fig. 2A, Left) to investigate if NB stimulation can activate V1 astrocytes. V1 astrocytes and neurons were loaded with the fluorescent calcium indicator Oregon Green 488 Bapta-1-AM (OGB1-AM), whereas astrocytes were loaded with the selective astrocytic marker sulforhodamine101 (SR101; Molecular Probes) (29) (Fig. 2A, Right). OGB1-AM fluorescence in astrocytes was monitored continuously with two-photon imaging during NB stimulation. First, we examined the
effect of a brief train of pulses applied to the NB (50 pulses at 100 Hz) on the spontaneous activity in astrocytes. Indeed, we observed robust increases in the calcium responses of astrocytes (Fig. 2B, Left; n = 30 of 44 astrocytes in 5 animals; P < 0.0001, paired t test, comparing population, trial-averaged responses before and after NB stimulation). When atropine, a muscarinic AChR (mAChr) antagonist, was delivered by a visualized pipette (Fig. 2A, Right), NB-evoked responses were reduced to pre-NB levels (Fig. 2B, Right, and C (n = 17 NB stimulation-facilitated astrocytes in 3 animals; P = 0.001, paired t test, comparing responses before and after NB stimulation; P < 0.001, paired t test, comparing NB evoked responses before and after atropine application) and Fig. S2A). Thus, responses of cortical astrocytes to NB stimulation are evoked by cholinergic modulation via mAChRs.

To examine how NB activation influences visual responses of astrocytes, we presented randomly oriented visual gratings synchronized to a single, brief train of pulses applied to the NB (Fig. 2D) while imaging astrocyte responses. Indeed, visual responses of...
astrocytes were enhanced by NB stimulation [Fig. 2 E and F (n = 47 visually responsive astrocytes in 5 animals; P < 0.01, paired t test, comparing population average of visual responses pre- and post-NB) and Fig. S2B], showing that V1 cortical astrocytes are capable of integrating visual and cholinergic inputs.

**ACH Stimulation in V1 Slices Evokes Calcium Responses in Astrocytes via Muscarinic Receptors.** To examine the mechanisms underlying NB-evoked responses in astrocytes, we performed calcium imaging of astrocytes in slices of V1. Responses of OGB1-AM-loaded layer 2/3 astrocytes were imaged (Fig. 2G) before and after ACH application. Imaged astrocytes had small round somas with thin radiating processes revealed by OGB1-AM loading (Fig. 2H, Upper Left). The identity of the imaged astrocytes was further confirmed by colocalization of their dye-filled processes (Fig. 2H, Upper Right) with anti-GFAP immunohistochemistry (Fig. 2H, Lower Left) and with the astrocyte-selective marker SR101 (29) in selected experiments (Fig. 2H, Lower Right). To mimic brief NB stimulation, we applied a brief pulse (0.2–1 s) of ACH, which evoked robust calcium transients in astrocytes (Fig. 2I, Upper). These responses were TTX-insensitive (Fig. 2J) and were abolished by the mAChR antagonists scopolamine and atropine (Fig. 2 I, Lower, and J; n = 8 astrocytes in 3 animals; P = 0.0001, paired t test). Immunohistochemistry confirmed that mAChRs were expressed on GFAP-expressing astrocytes (Fig. S3 A and B). To mimic prolonged NB stimulation, we bath-applied ACH, which caused an increase in the frequency and duration of calcium transients lasting for several minutes (Fig. S3 C–F). These findings, together with in vivo results (Fig. 2 A–F), indicate that V1 astrocytes are direct targets of NB stimulation-evoked cholinergic modulation via mAChRs (10, 11) and that the time scale of their calcium responses depend on the mode of ACH application and not on neuronal action potentials.

**Cholinergic Activation of Astrocytes Contributes to Prolonged Depolarizing Responses in Excitatory Neurons via Calcium-Mediated Processes.** The NB- and ACh-induced astrocyte responses suggested that the sustained slow potentiation of excitatory neuronal responses following paired NB and visual stimulation (Fig. 1 D–F) could be a contribution by astrocytes to neuronal plasticity, based on previous evidence that hippocampal astrocytes play a role in synaptic plasticity (13–15) and facilitate neuronal responses via gliotransmitter release (30, 31) or regulation of glutamate uptake (32).

We first tested the hypothesis that cholinergic activation of astrocytes can contribute to cholinergic facilitation of neuronal responses by performing whole-cell patch recordings in V1 slices (Fig. S4). Excitatory neurons, identified by their electrophysiological regular spiking and morphological characteristics, responded to a brief pulse of ACH with slow, prolonged depolarization (Fig. 3 E–G, Upper; n = 50 of 50 neurons), which was TTX-insensitive (Fig. S4; n = 10; P > 0.9, paired t test). We examined the mechanisms behind the ACh-induced slow depolarization further by blocking calcium responses in astrocytes (Fig. 3A). Astrocytic calcium was chelated through patch-loading electrophysiologically characterized astrocytes (Fig. 3B) with the cell-impermeable calcium chelator 1,2-Bis(2-aminophenoxy)ethane-N,N,N′,N′-tetraacetic acid (BAPTA) and allowing BAPTA to travel within the local syncytium of astrocytes via gap junctions (33, 34). The spread of BAPTA was assessed by Alexa Fluor 594 (A594; Molecular Probes) dye included in the patch pipette and determined to be ~150 µm from the patched astrocyte within 30–45 min of dialysis (Fig. 3C). BAPTA effectively blocked ACh-induced calcium transients in the dialed astrocytes (Fig. 3D). Excitatory neurons within 100 µm of the patched astrocytes (Fig. 3C) were recorded after BAPTA dialysis. The amplitude of the ACh-induced slow depolarization in neurons within the dialed astrocytic network was reduced compared with that in control conditions (Fig. 3E), either without astrocyte patch (n = 9 neurons; P < 0.001, t test) or in the presence of astrocytes dialed with A594 without BAPTA (n = 9 neurons; P < 0.0001, t test) (Fig. 3H). The slow neuronal depolarizations were mediated by mAChRs as they were abolished by atropine (Fig. 3 F and J; atropine, n = 18 neurons in 13 animals; P < 0.001, paired t test), consistent with our previous demonstration that ACh-induced astrocyte calcium transients were mAChR-mediated (Fig. 2 B, C, and F–J). These data therefore demonstrate that cholinergic activation of astrocytes can contribute to cholinergic facilitation of mAChR-mediated neuronal responses via increase of intracellular calcium.

**Cholinergic Activation of Astrocytes Evokes Slow NMDA Receptor-Mediated Currents in Neurons.** Earlier work has demonstrated that gliotransmitters released by astrocytes can act on the NRI/NR2B subunits of extrasynaptic NMDA receptors (NMDARs) in neurons (35, 36). These subunits have slow kinetics (37–39) and have been proposed to give rise to slow currents observed in neurons when adjacent astrocytes are activated (12, 36, 40). To investigate if NMDA-mediated currents underlie the astrocyte-evoked cholinergic responses in neurons, we performed whole-cell voltage-clamp recordings in which slow currents were defined and discriminated from miniature excitatory postsynaptic currents by their differential time courses (Fig. 4 A–C, Fig. S5 A and B, and SI Materials and Methods, Analysis of Slow Currents). Indeed, ACh induced an increase in the frequency of TTX-insensitive slow currents (Fig. 4 D–F, Upper, and G; Table S1 and Fig. SSC; n = 18 neurons in 9 animals; P < 0.0001, paired t test). These ACh-induced slow currents were atropine-sensitive (Fig. 4 E and G and Table S1: n = 10 neurons in 5 animals; P < 0.02, paired t test comparing ACh-induced slow current frequency before and after atropine application). To investigate if these currents have an astrocytic origin, double-patch experiments were performed in which a pyramidal neuron was patched and a neighboring astrocyte was contacted with a BAPTA-containing patch pipette in a cell-attached configuration, keeping the membrane intact to prevent BAPTA diffusion into the astrocyte. ACh-induced slow currents in the pyramidal neurons were first recorded before the seal between the astrocyte and BAPTA-containing patch pipette was broken to allow BAPTA dialysis of the astrocyte synecium. After 30–45 min of BAPTA dialysis, a reduction of ACh-induced slow currents was observed (Fig. 4 D, Lower, and G and Table S1: n = 5 neurons in 4 animals; P < 0.006, paired t test comparing ACh-induced slow current frequency before and after BAPTA dialysis of astrocytes). In a similar set of experiments in which excitatory neurons were patched after astrocytic BAPTA dialysis, we also observed a reduction in the frequency of ACh-induced slow currents [Fig. SSC; P < 0.001, t test comparing ACh-induced slow current frequency without (n = 18 neurons) and with (n = 11 neurons) BAPTA dialysis of astrocytes]. The reduction of slow currents was further confirmed not to be due to the extra-cellular action of BAPTA (Fig. SSD). Both the ACh-induced slow currents and slow depolarizations in neurons were reduced in the presence of D-2-Amino-5-phosphonovaleric acid (D-APV), indicating that they are mediated by NMDARs (Slow currents in Fig. 4 F and G and Table S1: n = 10 in 7 animals; P = 0.0001, paired t test comparing ACh-induced slow current frequency before and after D-APV application; slow depolarization in Fig. 3 G and J: n = 8 neurons in 8 animals; P < 0.003, paired t test). Although we cannot determine the causal relationship between the ACh-induced slow currents and slow depolarization, their similar insensitivity to TTX (Fig. 4 D–F and Fig. S4) and similar sensitivity to BAPTA dialysis, D-APV, and atropine (Figs. 3 E–I and 4 D–G), as well as comparable durations (mean duration of slow depolarization = 76.6 ± 10.25 s, n = 24; duration of increase in slow current frequency = 60–120 s, n = 13; Fig. 4 D–F, Upper) and long peak and valley latency (Figs. 3 E and 4 A, B, and D), suggest a correlation between the two phenomena. Considering the low slow-current frequency, it is possible that the depolarization of the cellular membrane induced by these slow currents favors the...
development of ACh-induced prolonged depolarization. Collectively, these findings indicate that cholinergic excitation of astrocytes via mAChRs leads to calcium-mediated processes that, in turn, evoke NMDAR-mediated facilitatory responses, including the neuronal slow currents and slow depolarization.

Astrocytic IP$_2$R2-Mediated Calcium Mediates NB Stimulation-Evoked Potentiation of Visual Responses in Excitatory Neurons. We next investigated if cholinergic activation of astrocytes can contribute to the NB-mediated potentiation of visual responses in excitatory neurons observed in vivo (Fig. 1 E and F). For this purpose, we used conditional IP$_2$R2 KO mice (IP$_2$R2-cKO) (Fig. S6 and SI Materials and Methods, Mice), where the astrocytic IP$_2$R2, previously shown to be the only IP$_2$R (41) that mediates agonist-induced calcium responses in astrocytes (33, 42), is specifically knocked out in GFAP-expressing astrocytes (Fig. S4). In WT mice 77 of 80 GFAP-expressing cortical astrocytes (96.3%) colocalized with IP$_2$R2; in IP$_2$R2-cKO mice the fraction was 7 of 90 (7.8%). Because the evidence for IP$_2$R2 expression in neurons remains inconclusive (41, 42), the IP$_2$R2-cKO mice were used instead of the...
full IP3R2 KO mice]. We first performed calcium imaging (Fig. 2G) in V1 slices of adult IP3R2-cKO and WT (control) animals. Although ACh induced calcium responses in astrocytes of WT mice, no astrocytic calcium responses were observed in the IP3R2-cKO mice (Fig. 5B). We next performed whole-cell patch recordings in V1 slices of adult IP3R2-cKO and WT animals. The amplitude of the ACh-induced slow depolarization in IP3R2-cKO regular spiking neurons was indeed drastically reduced compared with that in WT (Fig. 5C; n = 5 WT and n = 5 IP3R2-cKO neurons in 2 animals each; P < 0.001, t test). When the NB was stimulated with multiple trains of pulses, where each train was paired with visual stimulation in the IP3R2-cKO animals in vivo using the same protocol/analysis as in WT animals (Fig. 1D), no sustained potentiation was observed (Fig. 5D; n = 5 neurons in 4 animals; P > 0.1, paired t test comparing pre-NB responses with post-NB responses pooled across all 400-s time segments for ON, ON–OFF, and OFF responses). The neurons in the IP3R2-cKO animals had similar electrophysiological properties as those in the WT animals (Fig. S7); therefore, the lack of sustained potentiation in the IP3R2-cKO animals cannot be attributed to unintended secondary effects due to abnormal electrophysiology in the transgenic mice. These data therefore confirm that astrocytic IP3R2-mediated calcium plays a significant role in the potentiation of visual responses in excitatory neurons following prolonged pairing of NB stimulation with visual stimulation.

**NB-Induced Cholinergic Activation of Astrocytes Contributes to Potentiation of Stimulus-Specific Responses.** The NB stimulation-evoked, astrocyte-mediated potentiation of visual responses could be a general increase in neuronal responsiveness, such that responses to any visual stimulus are indiscriminately enhanced, or it could reflect a specific facilitation of particular visual stimuli.
To discriminate between these possibilities, we performed single-unit recordings from neuronal populations (Fig. 6A) in both WT and IP$_2$R2-cKO mice while simultaneously stimulating NB electrically and presenting gratings of a specific orientation as extrinsic visual stimulation. Responses to nine orientations each 20° apart, including an arbitrarily chosen orientation for the paired NB and visual orientation stimulation (conditioned orientation), were measured before and after the pairing protocol. Blank gray screens were interleaved between each orientation presentation. We observed an increase in the firing rates (Fig. 6B) as well as in normalized postresponses minus preresp onses at the conditioned orientation in WT animals (ON–OFF responses are shown in Fig. 6 C, Left, and D, Upper. n = 41 neurons; P < 0.001, Wilcoxon rank-sum test comparing responses before and after pairing protocol). Interestingly, this increase was specific to the conditioned orientation because the facilitation at the conditioned orientation was significantly greater than at unconditioned orientations, with the latter showing no change in response (Fig. 6 B, C, Left, and D and Fig. S8A and C). Recovery of this facilitation was observed earlier (Fig. S8D) than when NB stimulation was paired with random orientations (Fig. 1F), possibly due to reversal of synaptic modifications (43) induced by exposure to orientations other than the conditioned orientation after the pairing protocol (during assays of the postconditioning effects). In sum, these findings demonstrate that pairing NB stimulation with a specific visual orientation induces a highly specific potentiation of the conditioned orientation over unconditioned orientations in WT animals.

In the IP$_2$R2-cKO animals, no potentiation at the conditioned orientation was observed after the pairing protocol. Instead, a slight response depression was surprisingly revealed (Fig. 6 B,
C, Left, and D, Upper and Fig. S8 A and C, Upper). The population mean conditioned response minus the unconditioned response in the IP<sub>R2-cKO</sub> mice showed a reversal in signs of conditioned responses, in marked contrast to the findings in the WT experiments (Fig. 6C, Right and Fig. S8B; P < 0.0001 comparing mean differential ON–OFF and ON responses in WT and IP<sub>R2-cKO</sub> paired experiments: n = 41 WT neurons and 45 IP<sub>R2-cKO</sub> neurons).

In additional control experiments in which the same visual stimuli were presented without NB stimulation (visual-only experiments), a similar reversal in signs of conditioned responses was observed (Fig. 6C and D and Fig. S8 A–C; P < 0.0001 comparing mean differential responses, both ON–OFF and ON, between WT paired and visual-only experiments). Visual-only responses in WT and IP<sub>R2-cKO</sub> animals were similar (Fig. S8E). Conditioned responses were depressed to similar extents in the visual-only experiments, as well as in the IP<sub>R2 KO</sub> paired experiments (Fig. 6C, Right and Fig. S8 B and E; P > 0.05 for each comparison). Collectively, these findings show that the potentiation observed at the conditioned orientation requires both NB activation and intact astrocytic IP<sub>R2</sub>-mediated calcium elevation. The absence of either of these two factors changes the sign of conditioned responses, indicating that, by itself, the prolonged orientation-specific visual stimulation in the training protocol induces cortical response suppression, likely via adaptation mechanisms that have previously been shown to be reliably evoked by similar prolonged orientation-specific stimulation (25). This phenomenon is absent in the IP<sub>R2-cKO</sub> responses measured in Fig. 5D, likely because the random orientation gratings used for pairing constitute a rapid presentation of drifting gratings of multiple orientations, and thus do not induce orientation-
specific adaptation.) The presence of NB stimulation paired with single-orientation stimulation, however, decisively counters this adaptation-induced suppression in WT animals but not in IP,R2-KO animals. In sum, these data show that astrocytic calcium can mediate an NB-evoked stimulus-specific potentiation of V1 neuron responses that overrides adaptation-induced response suppression in these neurons.

Discussion

We have demonstrated that repeated NB stimulation paired with visual stimulation in vivo leads to prolonged potentiation of visual responses in V1 excitatory neurons. Previous work in rat auditory cortex has shown that NB stimulation paired with an auditory stimulus leads to significant cortical plasticity exemplified by changes in receptive fields (2, 6) and reorganization of cortical maps (3–5). Our work extends these findings by revealing the critical role of astrocytes in mediating this NB-induced cortical plasticity through their direct activation by cholinergic modulation via muscarinic receptors. Although we cannot rule out a direct facilitatory effect of NB stimulation on pyramidal neurons, the predominant contribution to the potentiation in neurons seems to be from an astrocytic calcium-driven mechanism, because removal of IP,R2-mediated astrocytic calcium blocks NB-induced potentiation in vivo, whereas chelation of intracellular astrocytic calcium abolishes ACh-induced neuronal depolarization and slow currents ex vivo.

Recent reports using combined somatosensory and cholinergic stimulation have shown that cholinergic-activated astrocytes can induce synaptic plasticity (17, 18). However, it remained unknown if this plasticity reflects a general increase of cortical responsiveness or a more specific potentiation at the stimulus and circuit level. Using visual cortex as a model system, together with cell-attached and single-unit recordings, we were able to apply a more refined stimulus protocol and address the question at a single-cell level. We find that the astrocyte-mediated potentiation is indeed stimulus-specific, because pairing a grating of a particular orientation with NB stimulation induces a highly specific increase in response to the paired orientation but not to unpaired orientations. Because individual synapses on superficial layer V1 neurons convey specific orientation information (44), the influence of astrocyte-mediated plasticity has to be more precise and even synapse-specific, and this suggests an intimate organization of astrocytes or their domains with respect to the synapses that convey and generate orientation-specific responses. Indeed, such organization is suggested by matched response features of neurons and adjacent astrocytes in V1 (26).

The generation and plasticity of orientation-selective responses in V1 requires both feed-forward inputs from the thalamus and recurrent inputs within the cortex (45, 46) and can be powerfully altered by top-down influences (47). NB inputs have also been implicated in top-down processes, including attention (23). Our findings indicate that astrocyte-mediated mechanisms, potentially through astrocytic calcium-dependent release of gliotransmitters that act on neuronal NMDARs, can induce a plasticity cascade in synapses and alter orientation-specific responses. Candidate ligands include glutamate (36) and D-serine [(15), compare with (48)]. Alternatively, the potentiation can be mediated by astrocytic calcium-dependent regulation of extracellular glutamate (26) or extracellular potassium (49), which may indirectly lead to NMDAR-mediated responses. Further investigation, however, is required to understand them in the context of NB-mediated potentiation. Regardless of the mechanism, our findings reveal an important role of astrocytes in NB-induced cortical plasticity and highlight their role as partners with neurons in restructuring specific circuits that govern stimulus-specific cortical responses.

Materials and Methods

C57BL/6 mice and IP,R2-KO mice older than 2 wk of age and older than 6 wk of age were used for ex vivo and in vivo experiments, respectively. All experiments were performed under protocols approved by the Animal Care and Use Committee, Massachusetts Institute of Technology, and conformed to National Institutes of Health guidelines. For in vivo two-photon calcium imaging, as well as cell-attached and single-unit experiments, mice were anesthetized with urethane or a fentanyl/midazolam/medetomidine mixture before craniotomy and NB electrode and EEG probe implantation. The NB was stimulated with either a single train or multiple trains of pulses, paired with presentation of either random or single-orientation visual gratings. EEG recordings during the experiment and acetylcholinesterase histochemistry after the experiment were performed to verify the stereotaxic accuracy of NB stimulation. Ex vivo slice experiments were performed for calcium imaging and whole-cell recording. Anti-GFAP, neuronal nuclei (NeuN), anti-mAChR, and DAPI immunohistochemistry was performed to verify the identity of astrocytes and localization of mAChRs on them. Student’s t test and Wilcoxon rank-sum test were used for statistical analyses as appropriate. Full details are provided in SI Materials and Methods.

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