

Norepinephrine Controls Astroglial Responsiveness to Local Circuit Activity

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<http://dx.doi.org/10.1016/j.neuron.2014.04.038>

SUMMARY

Astrocytes perform crucial supportive functions, including neurotransmitter clearance, ion buffering, and metabolite delivery. They can also influence blood flow and neuronal activity by releasing gliotransmitters in response to intracellular Ca^{2+} transients. However, little is known about how astrocytes are engaged during different behaviors *in vivo*. Here we demonstrate that norepinephrine primes astrocytes to detect changes in cortical network activity. We show in mice that locomotion triggers simultaneous activation of astrocyte networks in multiple brain regions. This global stimulation of astrocytes was inhibited by alpha-adrenoceptor antagonists and abolished by depletion of norepinephrine from the brain. Although astrocytes in visual cortex of awake mice were rarely engaged when neurons were activated by light stimulation alone, pairing norepinephrine release with light stimulation markedly enhanced astrocyte Ca^{2+} signaling. Our findings indicate that norepinephrine shifts the gain of astrocyte networks according to behavioral state, enabling astrocytes to respond to local changes in neuronal activity.

INTRODUCTION

Astrocytes are an essential component of neural circuits *in vivo*. They form highly interconnected networks, in which individuals occupy distinct domains that are extensively coupled through gap junctions. Each cell extends highly ramified processes that ensheathe synapses, make contact with nodes of Ranvier, and form endfeet specializations on blood vessels, placing these cells in an ideal position to both control the extracellular milieu and influence neuronal activity. Indeed, astrocytes have been shown to participate in diverse functions, including neurotransmitter clearance, ion homeostasis (Djukic et al., 2007), hemo-

dynamic control (Mulligan and MacVicar, 2004), and synaptic plasticity (Min and Nevian, 2012). However, there is still uncertainty about how astrocyte networks are controlled *in vivo* and when they engage in these distinct behaviors.

Astrocytes express an extensive complement of Gq-coupled neurotransmitter receptors that liberate Ca^{2+} from intracellular stores, providing a means to adjust their behavior in response to changes in neural activity. Ca^{2+} signaling in astrocytes has been linked to diverse phenomena, including changes in blood vessel diameter (Attwell et al., 2010; Mulligan and MacVicar, 2004) and synaptic plasticity (Di Castro et al., 2011; Min and Nevian, 2012; Jourdain et al., 2007), suggesting that the impact of astrocytes on various aspects of brain physiology is controlled by these metabotropic receptors. Nevertheless, the role of Ca^{2+} signaling in astrocytes *in vivo* remains uncertain, and mice that lack IP3R2 Ca^{2+} release channels that are responsible for receptor-evoked Ca^{2+} transients are overtly normal (Petravicz et al., 2008). Our lack of understanding about the interaction of astrocytes with neural circuits reflects our limited knowledge about the behavioral contexts in which astrocyte networks are activated. Despite evidence that astrocytes are responsive to multiple neurotransmitters, the pathways used to activate astrocytes *in vivo* and the patterns of activity that they exhibit during different behaviors remain to be defined.

In *vivo* two-photon imaging using Ca^{2+} -sensitive dyes has revealed that astrocyte network activity can be enhanced by local glutamatergic signaling (Nimmerjahn et al., 2009; Schummers et al., 2008) or by stimulation of long-range cholinergic (Takata et al., 2011; Chen et al., 2012) or noradrenergic (Bekar et al., 2008; Ding et al., 2013) neuromodulatory projections. How these local and global neuronal pathways interact to control the activity of astrocyte networks in awake, behaving animals has not been determined. Here we developed mice that express the genetically encoded Ca^{2+} indicator GCaMP3 in astrocytes and used *in vivo* two-photon imaging to define the activity patterns of cortical and cerebellar astrocytes during locomotion. Our results indicate that the increase in arousal that accompanies locomotion promotes widespread activation of astrocyte networks in the cortex and enhances their responsiveness to local changes in neuronal activity.

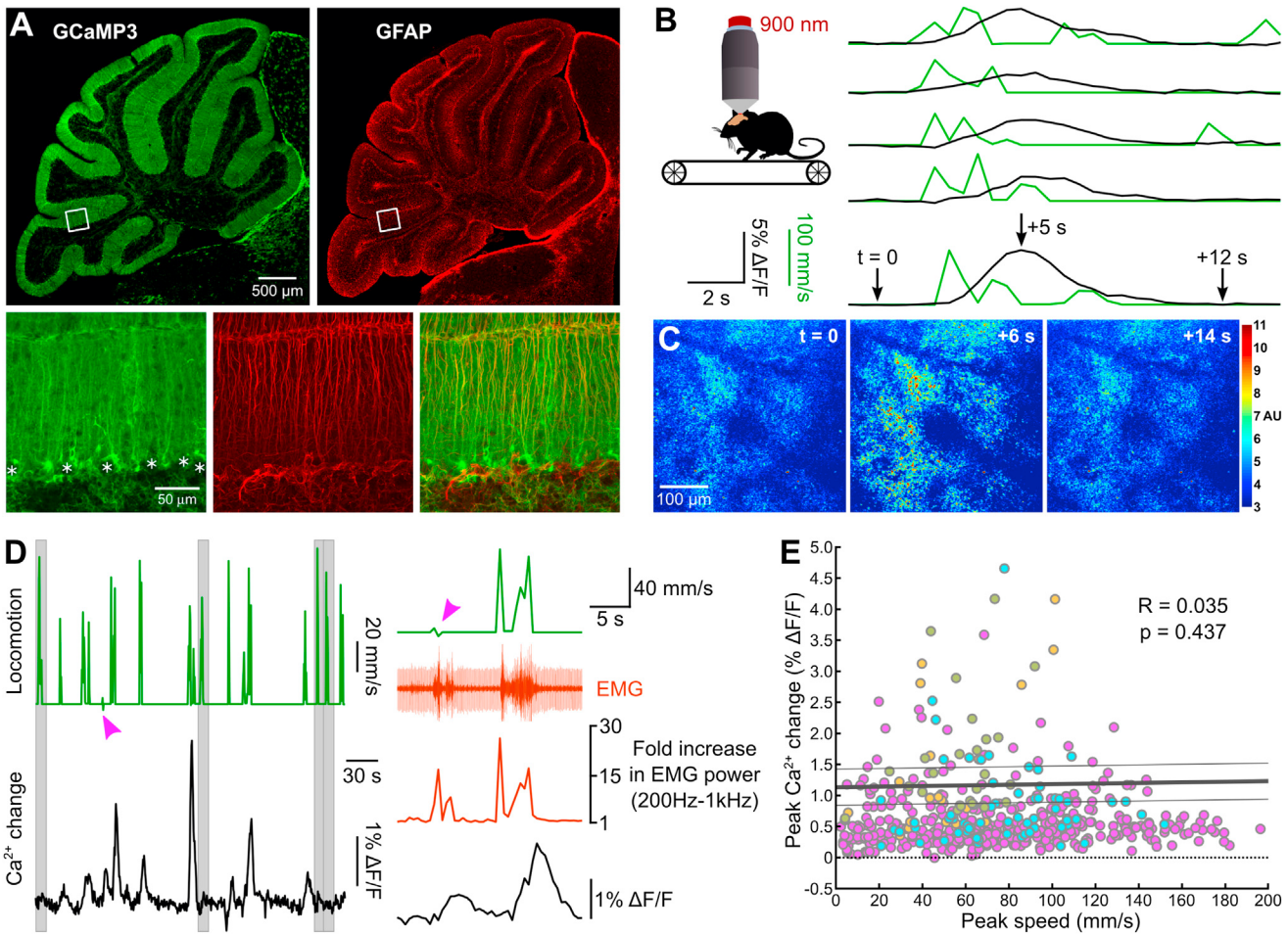


Figure 1. Weak Correlation between Voluntary Locomotion and Ca²⁺ Elevation in Bergmann Glia

(A) Cerebellar section from a 5-week-old *GLAST-CreER;R26-*lsl*-GCaMP3* mouse immunostained for GCaMP3 and glial fibrillary acidic protein (GFAP). Boxed areas are shown at higher magnification below. Asterisks highlight Purkinje cell somata devoid of fluorescence. (B) Left: schematic of imaging configuration. Right: five representative trials showing Bergmann glia Ca²⁺ increase (GCaMP3 fluorescence, black traces) relative to mouse locomotion (optical encoder, green traces). (C) Images of GCaMP3 fluorescence in Bergmann glial processes at times indicated by arrows in (B). (D) Left: continuous record of locomotion (green trace) and Bergmann glia Ca²⁺ levels (black trace, GCaMP3 fluorescence). Gray bars highlight periods when locomotion was not associated with Bergmann glia Ca²⁺ elevations. Arrowhead highlights a Bergmann glia Ca²⁺ transient that was not associated with locomotion. Right: expanded portion of trial including electromyography (EMG) signal. Arrowhead indicates timing of Bergmann glia Ca²⁺ elevation. (E) Plot of locomotion speed and Bergmann glia Ca²⁺ change (GCaMP3 fluorescence) for 707 locomotion periods from 4 mice. Colors represent trials from different individuals. Black lines represent mean ± SEM of 4 regression lines.

RESULTS

Ca²⁺ Transients in Bergmann Glia during Locomotion

Depend on Animal State of Arousal

To define the mechanisms that control astrocyte activity in vivo, we developed transgenic knockin mice (*R26-*lsl*-GCaMP3*) in which the genetically encoded Ca²⁺ indicator GCaMP3 can be expressed in a cell-specific manner (Figure S1 available online). After breeding to *GLAST-CreER* mice, tamoxifen administration induced GCaMP3 expression in 35% ± 2% of cortical astrocytes (n = 20 mice) (Figure S1C) and 100% of Bergmann glia (n = 17 mice), a distinct group of astroglial cells found in the cerebellar cortex (Figure 1A; Figure S1C), which could be visualized in vivo for weeks to months using two-photon imaging through a cranial window (Movie S1). Apart from neurons (dentate gyrus granule

cells, olfactory bulb interneurons) derived from SVZ/SGZ progenitors that express GLAST, no neuronal expression was detected in these mice.

Locomotion has been shown to trigger a transient rise in intracellular Ca²⁺ in Bergmann glia (Nimmerjahn et al., 2009). This activity, visualized acutely with a Ca²⁺ indicator dye, extended over large areas of the cerebellum and required local activation of glutamate receptors. To define the mechanisms required to engage this glial network, we trained GCaMP3-expressing mice to walk on a treadmill and monitored locomotion-induced Ca²⁺ levels in Bergmann glia. In accordance with previous findings (Nimmerjahn et al., 2009), brief bouts of locomotion were often associated with widespread elevation of Ca²⁺ in Bergmann glia that persisted for many seconds after cessation of movement (Figure 1B; Movie S2). However, the magnitude of the

Bergmann glia Ca^{2+} response was not correlated with locomotion speed ($R = 0.0352$, $p = 0.4365$, 495 events from 4 mice), and locomotion often did not trigger activation of Bergmann glia (212 failures in 707 locomotion events). As comparable bouts of locomotion should produce similar activity in glutamatergic afferents, these results suggest that other signaling pathways are involved in recruitment of these glial cells. Indeed, Bergmann glia often exhibited widespread activity in the absence of locomotion (Figures 1D and 1E) (83 events, 4 mice).

To provide an independent measure of motor activity in these mice, we monitored muscle contraction during imaging trials. Post hoc analysis of electromyogram (EMG) recordings revealed that “spontaneous” Ca^{2+} transients that occurred in the absence of locomotion were often associated with an increase in EMG power (73/83 events, 3 mice), suggestive of startle behavior. This observation raised the possibility that activity in Bergmann glia may correspond to an increase in arousal. To facilitate pharmacological analysis of locomotion-induced Bergmann glial Ca^{2+} responses, we standardized their motor activity by subjecting mice to brief periods (5 s) of forced locomotion. Strikingly, enforced locomotion consistently evoked large Ca^{2+} transients throughout the Bergmann glia network (Figure 2A). Ca^{2+} transients induced by enforced locomotion had a similar time course to events that occurred during voluntary locomotion (enforced: half-maximum width, 8.30 ± 0.60 s, $n = 159$ events; voluntary: 8.26 ± 1.24 s, $n = 13$ events, 5 mice, $p = 0.975$) (Figure 2B) and enforced events that immediately followed voluntary events were depressed by $47\% \pm 5\%$ ($n = 11$ events, 7 mice, $p < 0.001$) (Figure 2C), suggesting that similar pathways are activated during both voluntary and enforced locomotion.

Norepinephrine Induces Ca^{2+} Elevations in Bergmann Glia during Locomotion

We assessed whether antagonists of different neuromodulatory receptors attenuated Bergmann glia Ca^{2+} changes induced by locomotion. Although Bergmann glia responses were not affected by peripheral administration of CNS accessible antagonists of serotonergic, muscarinic, metabotropic glutamate, or cannabinoid receptors, administration of trazodone, a broad-spectrum inhibitor of adrenergic, serotonergic, and histaminergic receptors (Cusack et al., 1994), dramatically reduced locomotion-induced Ca^{2+} transients in these cells without altering the intensity of their motor response (EMGs) (Figures 2D–2F). Increases in arousal are often associated with the release of norepinephrine (Foote et al., 1980), raising the possibility that the trazodone-induced block of Ca^{2+} signaling in Bergmann glia is due to an inhibition of the noradrenergic signaling. Indeed, chemical depletion of norepinephrine with the neurotoxin DSP4 (Jonsson et al., 1981) abolished Bergmann glia Ca^{2+} transients induced by either voluntary or enforced locomotion (Figure S2), indicating that norepinephrine plays a key role in activating these astroglial cells during this behavior. The histaminergic H1 receptor antagonist diphenhydramine (DPH) also partially inhibited Bergmann glia responses, possibly due to its antagonism at $\alpha 1$ -adrenoceptors (Kester et al., 2003). To define the receptor subtypes responsible, we locally applied selective adrenoceptor antagonists to the cerebellar surface between imaging sessions. Locomotion-induced Ca^{2+} transients were

strongly inhibited by the $\alpha 1$ -adrenergic receptor antagonist terazosin but not by the β -adrenoceptor antagonist metoprolol (Figure 2G). In contrast, AMPA and NMDA receptor antagonists attenuated but did not block this activity, suggesting that local glutamatergic and noradrenergic signaling act synergistically in the cerebellar cortex to promote Bergmann glia activity.

Locomotion-Induced Norepinephrine Release Activates Astrocytes in Visual Cortex

Noradrenergic neurons in the brainstem nucleus locus coeruleus (LC) extend axon collaterals diffusely throughout the brain, providing the means to exert control over brain activity states (Steindler, 1981). To determine whether locomotion-induced engagement of the noradrenergic system results in global activation of astrocytes throughout the brain, we monitored astrocyte Ca^{2+} levels in primary visual cortex (V1) during locomotion (Figures 3A–3C). Remarkably, in complete darkness, enforced locomotion reliably elicited Ca^{2+} transients in V1 astrocytes that had a similar time course to Bergmann glia responses in the cerebellum (Figures 3D and 3E; Movie S3). Locomotion-induced astrocyte activity in V1 was similarly blocked by trazodone and abolished by depletion of norepinephrine with DSP4 (Figure S3). To assess the relative timing and spatial extent of changes in astrocyte activity, we developed a dual fiber-optic imaging system to monitor GCaMP3 fluorescence simultaneously in different brain regions (Figure 3F). When fiber-optic probes were positioned over cranial windows implanted above the visual cortex (V1) and cerebellum (lobulus simplex), coincident Ca^{2+} elevations were detected in both astrocytes and Bergmann glia in response to voluntary and enforced locomotion (Figure 3G). The magnitude of Ca^{2+} changes in these two regions covaried (correlation coefficient = 0.756, $p < 0.001$; $n = 348$ events from 6 mice), suggesting that amplitude fluctuations arise primarily from different levels of activity in noradrenergic neurons. Although events in these regions began at the same time (cerebellum onset delay = 1.93 ± 0.13 s; V1 onset delay = 1.91 ± 0.10 s, 348 events from 6 mice, $p = 0.8982$), consistent with the minimal delay predicted to arise solely from differences in axonal length (~ 40 ms, assuming a conduction velocity of 0.5 m/s and an additional distance of 20 mm to V1), Ca^{2+} transients in visual cortex reached their peak 1.6 ± 0.2 s later than events in the cerebellum ($p < 0.001$), raising the possibility that local circuit activity modifies the timing of astrocyte recruitment.

Norepinephrine Enhances the Sensitivity of Astrocytes to Local Circuit Activity

Locomotion induces release of norepinephrine in the visual cortex in complete darkness, providing a means to determine whether activation of this endogenous neuromodulatory input influences the response of astrocytes to local network activity. To assess the effects of norepinephrine, we monitored the behavior of groups of V1 astrocytes to locomotion, light stimulation alone, and locomotion paired with light stimulation. Enforced locomotion reliably elicited Ca^{2+} transients in V1 astrocytes, although the response of individual cells was variable, with some cells responding to each trial and others responding to only a subset of trials (Figures 4A and 4B). In contrast, light stimulation alone rarely activated V1 astrocytes, despite triggering large evoked

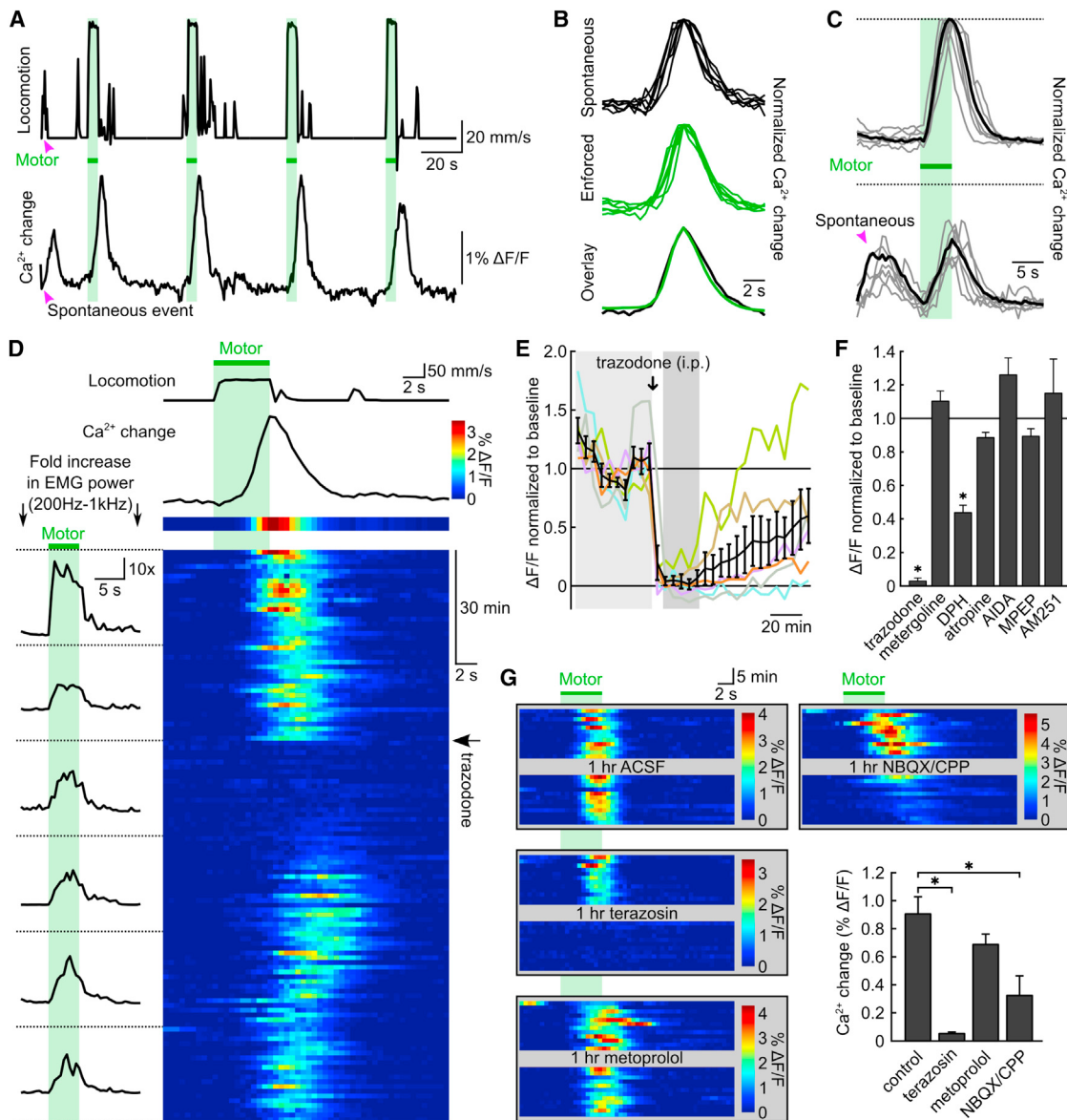


Figure 2. Noradrenergic Signaling Is Required for Locomotion-Induced Activation of Bergmann Glia

(A) Spontaneous (arrowhead) and enforced (green bars) locomotion and corresponding Bergmann glia Ca^{2+} levels (GCaMP3 fluorescence). (B) Normalized Bergmann glia Ca^{2+} changes associated with spontaneous (6 of 13 events) and enforced locomotion (6 of 159 events) from experiment in (A). (C) Individual (gray traces) and average (black traces) Bergmann glia Ca^{2+} transients resulting from enforced locomotion with (bottom traces) or without (top traces) a preceding spontaneous event. (D) Consecutive trials of enforced locomotion (green bar) pseudocolored according to the magnitude of Ca^{2+} change. Arrow indicates time of trazodone injection (10 mg/kg i.p.). Left: average EMG power during 20 consecutive trials at right. (E) Normalized amplitude of Bergmann glia Ca^{2+} elevations elicited by enforced locomotion. Each point is average of four consecutive trials normalized to baseline. Regions highlighted by light gray bar used to determine baseline and dark gray bar the maximal response to drug. Black symbols represent mean \pm SEM. (F) Effect of neuromodulatory receptor antagonists on Bergmann glia Ca^{2+} response to enforced locomotion. Columns represent mean \pm SEM. Concentrations from left: 10, 10, 20, 20, 20, 20, and 10 mg/kg i.p. Number of experiments from left: 6, 5, 5, 5, 5, 4, and 3. Asterisks indicate significant reduction relative to baseline in one-way ANOVA followed by Bonferroni post hoc test. (G) Bergmann glia Ca^{2+} transients in consecutive trials of enforced locomotion before and after local application of antagonists. Summary graph represents mean \pm SEM. Concentrations are the following: 100 μM terazosin, 3 μM metoprolol, 300/200 μM NBQX/CPP. Number of experiments from left: 8, 5, and 5. Asterisks indicate significant difference in one-way ANOVA followed by Bonferroni post hoc test.

potentials and inducing hemodynamic changes in this area (Figures S4A and S4B). Occasional coincident astrocyte activity that occurred in response to light stimulation was always associated with enhanced EMG power (Figure 4A), reflecting voluntary locomotion of the animals or possible startle behavior. Despite the weak activation of V1 astrocytes by visual input, pairing locomotion with light stimulation markedly enhanced astrocyte activity above that produced by locomotion alone, without altering

motion of the animals or possible startle behavior. Despite the weak activation of V1 astrocytes by visual input, pairing locomotion with light stimulation markedly enhanced astrocyte activity above that produced by locomotion alone, without altering

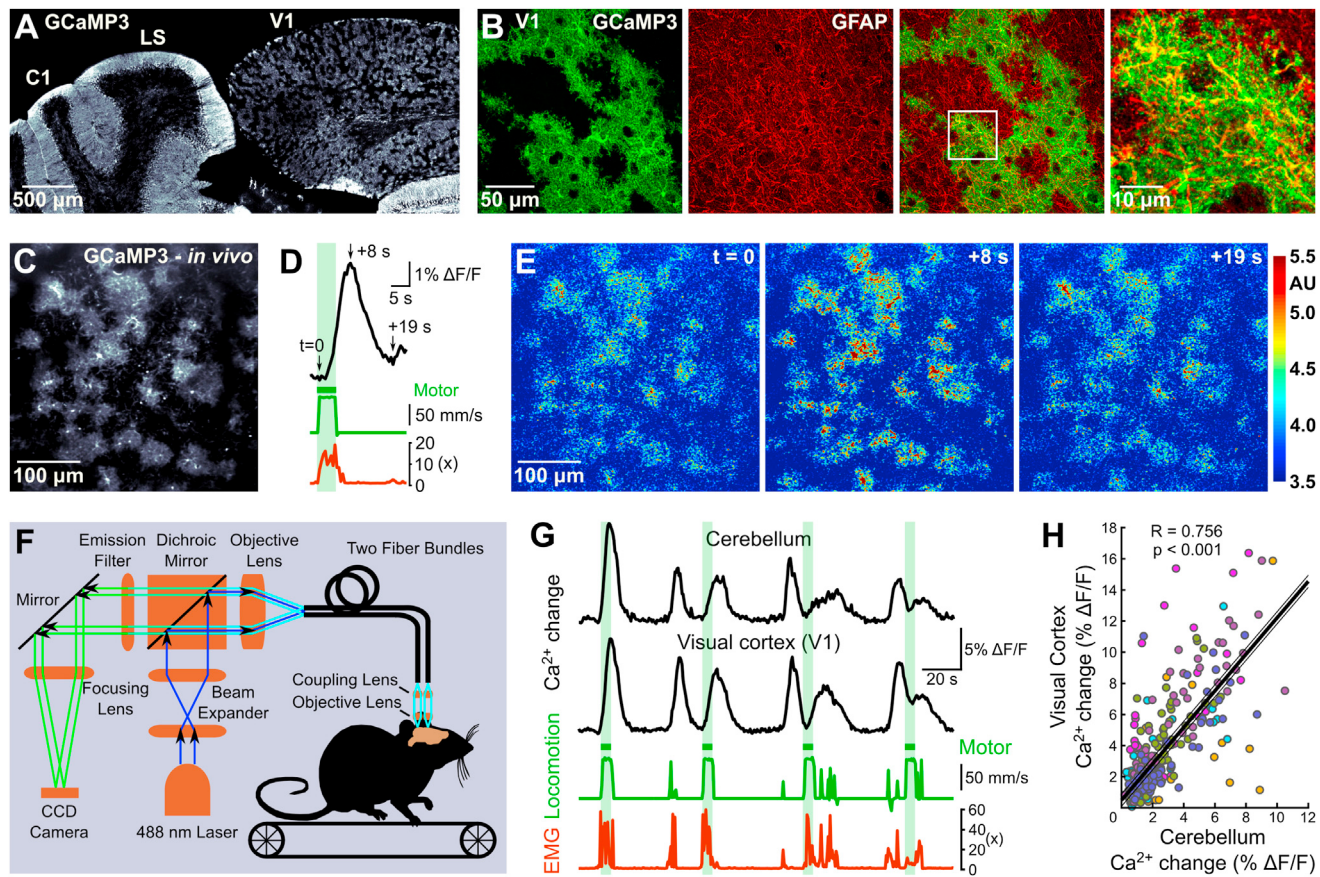


Figure 3. Locomotion Induces Simultaneous Activation of Astroglia in Different Regions of the Brain

(A) Paraxial section of *GLAST-CreER;R26-*Isl*-GCaMP3* mouse (P57) immunostained for GCaMP3. C1, crus1; LS, lobulus simplex; V1, primary visual cortex. (B) Higher-magnification image of GCaMP3 (green) and GFAP (red) in V1. (C) In vivo image of V1, layer 1 (70 μ m below pial surface) cortical astrocytes expressing GCaMP3 in 3-month-old *GLAST-CreER;R26-*Isl*-GCaMP3* mouse. (D) Mean change in cytosolic Ca^{2+} (GCaMP3) in V1 cortical astrocytes (black trace) induced by enforced locomotion (green trace). Green bar: period of enforced locomotion; green trace: locomotion; red trace: fold increase in EMG activity. (E) Images of GCaMP3 fluorescence in V1 astrocytes at times indicated in (D). (F) Schematic of dual fiber-optic photometry configuration. (G) Ca^{2+} changes in Bergmann glia and V1 astrocytes visualized simultaneously during enforced locomotion (green bars) and corresponding EMG activity. (H) Covariance between Bergmann glia and V1 astrocyte Ca^{2+} changes during spontaneous locomotion. $n = 348$ spontaneous locomotion events from 6 mice. Black lines represent mean \pm SEM of 6 regression lines.

motor output as measured by EMG (Figures 4A–4D). Analysis of individual astrocytes revealed that light stimulation increased both the amplitude of Ca^{2+} transients in each cell and the proportion of cells that responded to each trial (Figures 4B and 4C). To assess whether the highly ramified processes of astrocytes, which are comprised of distinct microdomains (Grosche et al., 1999), exhibit Ca^{2+} transients with thresholds and kinetics distinct from those observed at the soma, we analyzed Ca^{2+} events in astrocyte processes and somata independently by masking individual astrocyte somata within the field; these non-somatic regions yielded signals from both the main processes and the highly ramified segments within the neuropil. Although it is possible that some microdomain activity may go undetected using our imaging method, astrocyte processes responded very similarly to their somata (Figures S4C–S4H), suggesting that $\alpha 1$ adrenoceptors are widely distributed in astrocyte membranes.

DISCUSSION

These findings indicate that astrocytes throughout the brain are activated by norepinephrine during periods of heightened vigilance and that this modulation plays a critical role in enabling these glial cells to respond to local network activity. Unlike fast excitatory and inhibitory neurotransmitters, where transmission occurs at synapses defined by directly apposed pre- and post-synaptic elements, norepinephrine is released from axonal varicosities into the surrounding neuropil, a “volume transmission” that enables multiple targets in the surrounding area to be affected. Astroglia appear to be a direct target of the noradrenergic system, as gene expression profiling indicates that they express a variety of Gq-coupled alpha adrenergic receptors (Cahoy et al., 2008); exogenous norepinephrine triggers a rise in Ca^{2+} in cultured astrocytes (Salm and McCarthy, 1990) as well as astrocytes and Bergmann glia in acute brain slices

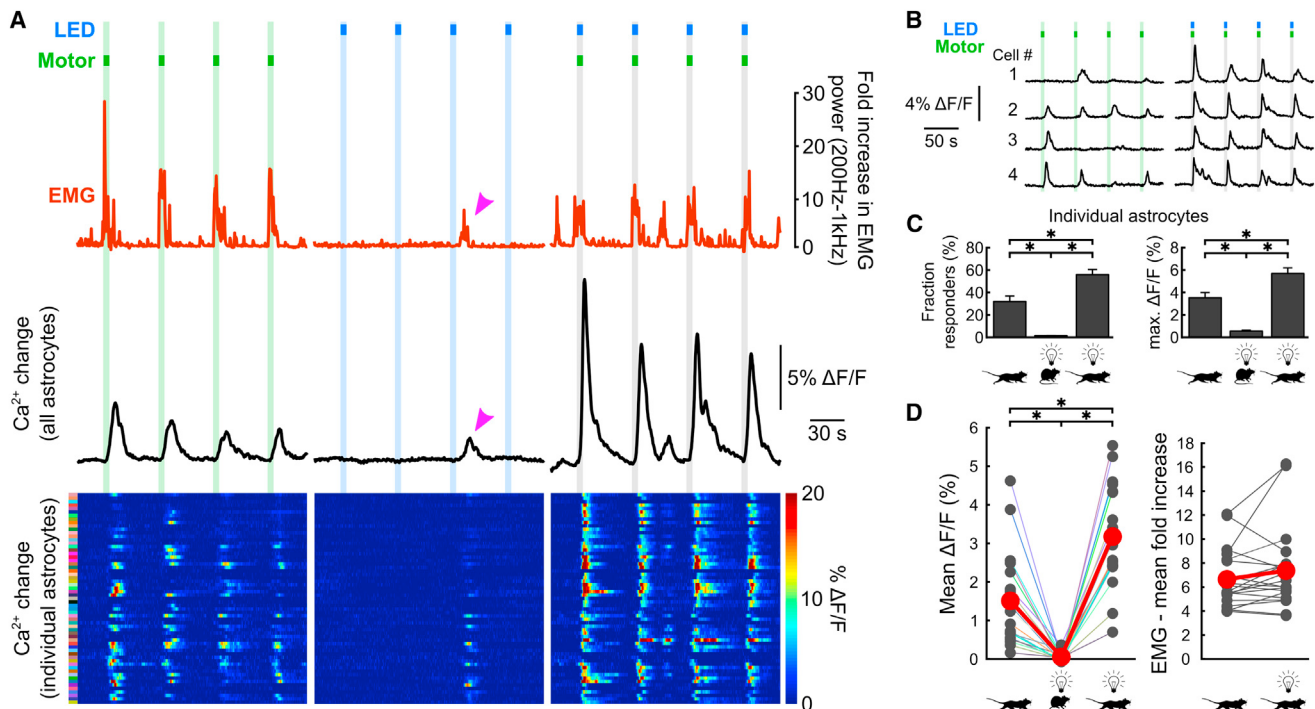


Figure 4. Norepinephrine Enhances the Sensitivity of Astrocytes to Local Circuit Activity

(A) V1 astrocyte Ca^{2+} responses to enforced locomotion (green bars), visual stimulation (blue bars), or simultaneous enforced locomotion and visual stimulation (gray bars). Red traces represent EMG activity; black traces represent mean Ca^{2+} change in all astrocytes. Bottom: Ca^{2+} changes in individual astrocytes pseudocolored according to amplitude. Arrowhead highlights Ca^{2+} elevation associated with spontaneous locomotion. (B) Ca^{2+} transients in four representative cells during four consecutive trials of enforced locomotion (green bars) or simultaneous enforced locomotion and visual stimulation (gray bars). (C) Left: average fraction of astrocytes responding with a mean Ca^{2+} elevation exceeding 1% $\Delta F/F$. Right: average maximum Ca^{2+} responses in individual astrocytes (16–20 trials). Error bars represent mean \pm SEM; $n = 20$ mice; asterisks indicate significant difference (one-way ANOVA with Bonferroni post hoc test). (D) Left: mean Ca^{2+} elevations in all astrocytes (16–20 trials). Colored lines connect data points from individual mice. Asterisks indicate significant difference (one-way ANOVA with Bonferroni post hoc test). Right: change in EMG power during enforced locomotion or simultaneous enforced locomotion and visual stimulation. Each point represents average response in 16–20 trials. Red circles indicate mean of all trials.

when neuronal activity is blocked (Duffy and MacVicar, 1995; Shao and McCarthy, 1997), and aversive stimuli or direct electrical stimulation of the LC induces Ca^{2+} transients in astrocytes in the somatosensory cortex (Bekar et al., 2008; Ding et al., 2013). The mechanisms that enhance the responsiveness of astrocytes to local increases in neuronal activity are unknown but may involve intracellular changes in signaling intermediates that enhance the ability of other Gq-coupled metabotropic receptors to induce Ca^{2+} release from intracellular stores (Ryzhov et al., 2006). It is also possible that synergy occurs by enhancing the release of norepinephrine in V1, leading to larger, more prolonged Ca^{2+} transients in astrocytes. This effect could be initiated locally in the cortex or by enhancing the activity of these neurons at the level of the LC. Direct assessment of the activity of noradrenergic projections to V1 during this behavior would help to define whether their activity is modulated in this form of plasticity.

Reduced Responsiveness of Astrocyte Networks in Unanesthetized Mice

The close association of astrocytes with synapses allows detection of neuronal activity and the potential for feedback modulation (Parpura et al., 1994; Di Castro et al., 2011; Jourdain et al.,

2007; Saab et al., 2012). Thus, it was unexpected that exposure to strong visual stimulation alone failed to engage astrocyte networks in V1, particularly given previous studies demonstrating robust activation of V1 astrocytes with light in anesthetized mice (Chen et al., 2012). However, anesthesia has been shown to dramatically expand active regions in mouse V1 and prolong responses to visual stimulation (Haider et al., 2013), which may lower the threshold for astrocyte activation. Moreover, astrocytes have been implicated in the regulation of sleep-wake cycles (Halassa et al., 2009) and their functional state differs considerably between sleep/anesthesia and wakefulness (Xie et al., 2013). It has also been reported that astrocyte Ca^{2+} signaling in vivo is inhibited by various anesthetics (Nimmerjahn et al., 2009; Thrane et al., 2012), raising concern that the depth of anesthesia and type of anesthetic can dramatically alter their responsiveness. Together, these findings highlight the importance of defining astrocyte activity in awake animals during different behavioral states.

Norepinephrine Enhances the Gain of Both Astrocyte and Neuronal Networks

These studies indicate that astrocyte networks have a high threshold for activation in periods of quiescence and suggest

that recruitment of the noradrenergic system is required to enhance the gain of these networks to enable local interactions between astrocytes and neurons in circuits involved in processing sensory information. The activation of astrocytes in V1 by norepinephrine during locomotion parallels recent *in vivo* electrophysiological studies indicating that locomotion triggers norepinephrine-dependent depolarization of neurons in V1 (Polack et al., 2013). In contrast to neuronal activation, which began within 50 ms of locomotion onset, Ca^{2+} transients in astrocytes began more than a second later (1.4 ± 0.1 s, $n = 106$ events from 13 mice), presumably reflecting the additional biochemical steps required to release Ca^{2+} from intracellular stores. These results suggest that astrocyte Ca^{2+} transients are unlikely to be causal for the initial depolarization of cortical neurons, for example, by stimulating the release of gliotransmitters; however, Ca^{2+} is not the only signaling intermediate produced by activation of α adrenoceptors and astrocytes may express adrenoceptors coupled to other G proteins (e.g., Gi) (Cahoy et al., 2008) that could trigger responses on a timescale coincident with neuronal depolarization. The ability of astroglia to detect the coincidence of arousal and local network activity may allow these cells to contribute to sensory-specific attentional shifts (Cohen and Maunsell, 2011), promoted by the slow onset and prolonged nature of their Ca^{2+} responses. Future analysis of mice in which alpha adrenoceptors have been selectively removed from astrocytes will help to define their role in noradrenergic modulation of cortical networks and arousal states.

Astrocytes are well positioned to amplify the effects of diffuse neuromodulatory projections; their highly ramified processes not only increase the probability of interaction with low concentrations of neurotransmitters but also shield neurons by increasing the diffusional distance from varicosities to neuronal membranes. Although the exact complement of neurotransmitter receptors expressed by astrocytes in different brain regions has not been defined, cortical astrocytes have also been shown to respond to glutamate, ATP, and acetylcholine (Verkhratsky and Kettenmann, 1996; Takata et al., 2011). The ability to stably express Ca^{2+} indicators selectively in astrocytes *in vivo* using conditional GCaMP3 mice and viral-based delivery methods (Shigetomi et al., 2013) will help define the behavioral contexts in which these different receptors become activated and reveal how they influence interactions between neurons and astrocytes in distinct neural circuits.

EXPERIMENTAL PROCEDURES

Animals

Cre recombinase-conditional GCaMP3 mice were generated using a knockin strategy into the ROSA26 locus and crossed to *GLAST-CreER* mice to enable GCaMP3 expression in astrocytes and Bergmann glia. Cranial windows were implanted above the cerebellum or V1.

In Vivo Two-Photon Imaging

Fluorescence images were collected using a Movable Objective Microscope (MOM) (Sutter Instruments) with a 20 \times , 1.0 NA objective (Zeiss). Image acquisition rate was 2 frames/s.

Dual Optic Fiber Bundle Imaging

For simultaneous imaging of astroglial Ca^{2+} changes in cerebellum and visual cortex, cranial windows were implanted and two multicore optical fibers

aligned above these regions. The light path is illustrated in Figure 3F. No optical crosstalk between fiber bundles was detected: the objective lens pairs had a diameter of 2.4 mm and a working distance of 0.88 mm.

Locomotion Behavioral Paradigm

Mice were placed on a custom-designed linear treadmill and their heads immobilized. The treadmill was either freely movable or placed under motor control and the motion of the treadmill belt was monitored with an optical encoder.

Electromyography

Body surface potentials were recorded as the voltage between two silver wires placed subcutaneously at the right shoulder and left hip.

Visual Stimulation

A UV-LED with a Lambertian emission profile was used as a light source at a distance of 40 mm, centered between the eyes to achieve uniform light exposure.

Data Analysis

Data were processed and analyzed in MATLAB. $\Delta F/F$ fluorescence intensity traces represent $(F - F_{\text{median}})/F_{\text{median}}$, with F representing mean fluorescence value of all pixels within a region of interest (ROI) of one image frame and F_{median} representing median F of all image frames.

Statistical Analysis

The following statistical analyses were used: Figures 1E and 3H, analysis of covariance for correlation within subjects; Figures 2B, 2C, and 4D (right), difference in onset of response in Figures 3F–3H, Figures S2 and S3, one-tailed paired t test; Figures 2F, 2G, 4C, and 4D (left), one-way ANOVA followed by Bonferroni post hoc test; text, difference in peak of response in Figures 3F–3H, one-sample t test.

Additional information about the procedures used in this study is described in the *Supplemental Experimental Procedures*.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, four figures, and three movies and can be found with this article online at <http://dx.doi.org/10.1016/j.neuron.2014.04.038>.

AUTHOR CONTRIBUTIONS

M.P., A.A., and D.E.B. conceived the project. M.P. and D.E.B. designed the experiments and wrote the manuscript with help from the other authors. M.P. conducted experiments, developed software, and analyzed data. A.A. generated and characterized the *R26-*lsl*-GCaMP3* mice, J.C. and J.K. developed the dual fiber imaging probe and assisted with fiber-optic imaging, and V.A.D. provided assistance with adrenergic receptor pharmacology.

ACKNOWLEDGMENTS

We thank L. Looger for providing the GCaMP3 construct, J. Nathans for providing *GLAST-CreER* mice and assistance with Rosa26 targeting, M. Fukaya for providing antibodies, Terry Shelley for his machining expertise, and Naiqing Ye for assistance with animal colony maintenance. This work was supported by grants from the NIAAA (AA022239) to M.P., the NIMH (Conte Center for Neuroscience, MH084020), the Brain Science Institute and the Science of Learning Institute at Johns Hopkins to D.E.B. A.A. was supported by a postdoctoral fellowship from the National Multiple Sclerosis Society.

Accepted: April 11, 2014

Published: June 18, 2014

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