

Published in final edited form as:

Sci Signal.; 5(218): ra26. doi:10.1126/scisignal.2002334.

Astrocytes Modulate Neural Network Activity by Ca2+-Dependent Uptake of Extracellular K+

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Abstract

Astrocytes are electrically nonexcitable cells that display increases in cytosolic calcium ion (Ca²⁺) in response to various neurotransmitters and neuromodulators. However, the physiological role of astrocytic Ca²⁺ signaling remains controversial. We show here that astrocytic Ca²⁺ signaling ex vivo and in vivo stimulated the Na⁺, K⁺-ATPase (Na⁺- and K⁺-dependent adenosine triphosphatase), leading to a transient decrease in the extracellular potassium ion (K^+) concentration. This in turn led to neuronal hyperpolarization and suppressed baseline excitatory synaptic activity, detected as a reduced frequency of excitatory postsynaptic currents. Synaptic failures decreased in parallel, leading to an increase in synaptic fidelity. The net result was that astrocytes, through active uptake of K+, improved the signal-to-noise ratio of synaptic transmission. Active control of the extracellular K⁺ concentration thus provides astrocytes with a simple yet powerful mechanism to rapidly modulate network activity.

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Author contributions: F.W., N.A.S., O.X., and T.T. performed the experiments and analyzed the data, A.B. and T.M. synthesized reagents. F.W., N.A.S., T.T., L.B., and M.N. designed the experiments. F.W., N.A.S., T.F., T.T., L.B., and M.N. wrote the paper.

Competing interests: The authors declare that they have no competing interests. SEA0400 is a proprietary compound of Taisho Pharmaceutical Co. Ltd. (Japan) and was used while under development as a drug candidate; the amount available by request from the authors is limited; however, it can be obtained by request from Taisho Pharmaceutical Co.

SUPPLEMENTARY MATERIALS

www.sciencesignaling.org/cgi/content/full/5/218/ra26/DC1 Fig. S1. Ouabain-insensitive ⁸⁶Rb⁺ uptake.

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Fig. S2. Doxycycline suppresses eGFP expression in cultured astrocytes prepared from MrgA1^{+/-} mice. Fig. S3. GPCR-activated increases in ⁸⁶Rb⁺ uptake require increases in cytosolic Ca²⁺ in cultured astrocytes.

Fig. S4. Comparison of ⁸⁶Rb⁺ uptake in rat versus mouse and rat cortical versus hippocampal astrocytes.

Fig. S5. Detection of GPCR-activated changes in EPSPs is depth- and [K⁺]_{Bath}-dependent.

Fig. S6. GPCR activation triggers a transient minor increase in intra-astrocytic Na+ concentration and a minor decrease in extracellular Ca²⁺ concentration in hippocampal slices.

Fig. S7. Activation of GPCR-induced changes in PKA and PKC activity does not mediate ATP-induced increases in 86Rb+ uptake in

Fig. S8. Combined recordings of extracellular K⁺ with mEPSCs or mIPSCs in hippocampal slices.

Fig. S9. Effects of GPCR activation or reduction of bath K+ concentration on extracellular K+, neuronal membrane potential, and evoked EPSP in hippocampal slices.

INTRODUCTION

Extracellular K⁺ is a critical determinant of the resting membrane potential and is normally maintained within a range of 3.0 to 4.0 mM (1). Even minor increases in the extracellular K⁺ concentration can alter the probability of a neuron firing an action potential by affecting K⁺ channel gating, slowing action potential recovery (2), modulating synaptic transmission (3), or narrowing the gap between resting membrane potential and the threshold for activation of voltage-gated Na⁺ channels (4). Astrocytes, which function as sinks for the rapid uptake of extracellular K⁺ (5), have traditionally been regarded as supportive cells of the central nervous system (CNS) whose major housekeeping functions include the maintenance of extracellular K⁺ homeostasis (6). The highly negative resting membrane potential (~-85 mV), set by basal Na⁺- and K⁺-dependent adenosine tri-phosphatase (Na⁺,K⁺-ATPase) activity, and their large number of inwardly rectifying K⁺ channels drive excess K⁺ into astrocytes (7, 8). Although inwardly rectifying K⁺ channels appear to be primarily responsible for K⁺ homeostasis at rest, the Na⁺,K⁺-ATPase plays the predominant role in normalizing the increase in extracellular K⁺ that occurs during neural activity (7–11).

Work in tissues outside the CNS has demonstrated that activity of the Na⁺,K⁺-ATPase is regulated by the second messengers Ca²⁺ and adenosine 3',5'-monophosphate (cAMP) (5, 7). Moreover, the intracellular Na⁺ concentrations are rate-limiting for Na⁺,K⁺-ATPase activity in many cell types including astrocytes (8, 9). Thus, increases in cytosolic Na⁺ concentration in astrocytes could potentially lead to increased activity of the Na⁺, K⁺-ATPase and thereby to a decrease in extracellular K⁺.

The observations that astrocytes propagate Ca²⁺ signals over considerable distances (12), signal to neurons (13, 14), and modulate neural network activity (15, 16) have sparked interest in defining broader roles for astrocytes in more complex CNS function. Although passive astrocytic buffering of excess K⁺ is well established (17), whether astrocytes actively control resting extracellular K⁺ concentration by transiently changing the activity of the Na⁺,K⁺-ATPase is unclear. Astrocytic Ca²⁺ transients play a central role in signaling between neurons and glia (18). Thus, we wondered whether activation of Gq-linked heterotrimeric guanosine triphosphate (GTP)–binding protein (G protein)–coupled receptors (GPCRs) could, through production of inositol 1,4,5-trisphosphate (IP₃), activation of its receptor (IP₃R), and the subsequent increase in cytosolic Ca²⁺, control the astrocytic Na⁺,K⁺-ATPase activity and thereby modulate neuronal excitability by active uptake of K⁺.

Here, we show that a rise in cytosolic Ca^{2+} triggers an increase in ouabain-sensitive K^+ uptake in cultured astrocytes and a transient decrease in extracellular K^+ in hippocampal slices. Ca^{2+} -mediated K^+ uptake into astrocytes was primarily driven by Na^+ influx mediated by astrocyte Na^+ , Ca^{2+} exchange. Combined, these findings show that Ca^{2+} -dependent activation of the astrocytic Na^+ , K^+ -ATPase through GPCR-activated pathways enables astrocytes to dynamically regulate the extracellular K^+ concentration. Because extracellular K^+ concentration is a key determinant of neuronal excitability, its active regulation by astrocytes has numerous implications for basic CNS functions.

RESULTS

GPCR-activated Ca²⁺ signaling increases K⁺ uptake in cultured astrocytes

Astrocytes propagate intercellular Ca²⁺ signals through release of adenosine 5'-triphosphate (ATP) and the consequent activation of purinergic receptors (19, 20); therefore, to evaluate the potential role of astrocytic Ca²⁺ signaling in K⁺ uptake, we first assessed the role of purinergic receptors on Na⁺,K⁺-ATPase activity by quantifying the ouabain-sensitive fraction of rubidium-86 uptake (⁸⁶Rb⁺; used as a K⁺ analog) (9, 10). ATP stimulated an

increase in ouabain-sensitive $^{86}\text{Rb}^+$ uptake averaging $35.6 \pm 7.1\%$ (n = 27) in cultured cortical rat astrocytes (Fig. 1A) as well as an increase in cytosolic Ca²⁺. Similarly, uridine 5′-triphosphate (UTP, a P2Y2 and P2Y4 receptor agonist) and 2-methylthioadenosine 5′-diphosphate (2MeSADP, a P2Y1 receptor agonist) increased not only cytosolic Ca²⁺ but also ouabain-sensitive $^{86}\text{Rb}^+$ uptake by $24.1 \pm 3.6\%$ (n = 39) and $10.7 \pm 3.9\%$ (n = 8), respectively (Fig. 1A). Exposure to the protease-activated receptor-1 (PAR1)–selective agonist Thr-Phe-Leu-Leu-Arg-NH₂ (TFLLR-NH₂) or to prostaglandin E₂ (PGE₂) also increased cytosolic Ca²⁺ and K⁺ uptake, whereas the metabotropic glutamate receptor (mGluR) agonist *trans*-(1S,3R)-1-amino-1,3-cyclopentanedicarboxylic acid (t-ACPD) and the P1R agonist adenosine had no effect (Fig. 1A). The ouabain-insensitive fraction of t-ACPD and the pathways, such as channel-mediated K⁺ influx, were not affected by changes in cytosolic Ca²⁺.

We prepared astrocyte cultures from transgenic mice expressing an exogenous Gq-coupled receptor (MrgA1) under the control of the inducible human glial fibrillary acidic protein (hGFAP) promoter (21) to further evaluate the Ca²⁺ dependency of GPCR-activated K⁺ uptake (fig. S2). The MrgA1 agonist peptide Phe-Met-Arg-Phe amide (FMRF) increased intracellular Ca²⁺ and ⁸⁶Rb⁺ uptake in MrgA1^{+/-} astrocytes (Fig. 1A). ⁸⁶Rb⁺ uptake increased as a linear function of Ca²⁺ concentration regardless of the ligand used (Fig. 1B, left, $t^2 = 0.94$). Moreover, pretreatment with the endoplasmic reticulum (ER) Ca²⁺ pump inhibitor cyclopiazonic acid (CPA) blocked GPCR-activated increases in cytosolic Ca²⁺ and in ouabain-sensitive ⁸⁶Rb⁺ uptake as well as nonstimulated ouabain-sensitive ⁸⁶Rb⁺ uptake (Fig. 1B, right, and fig. S3). In contrast to astrocyte cultures, neuronal cultures failed to show increases in 86Rb+ uptake in response to activation of GPCR pathways. 2MeSADP and glutamate reduced ⁸⁶Rb⁺ uptake in neurons despite triggering increases in cytosolic Ca²⁺ (Fig. 1C). Thus, increases in the cytosolic Ca²⁺ concentration selectively increased Na⁺,K⁺-ATPase activity in astrocytic, but not neuronal cultures (compare Fig. 1, B and D). Whereas these experiments were performed in neo-cortical cultures, UTP-induced K⁺ uptake appeared to be comparable in astrocytic cultures prepared from hippocampus or cortex from either rat or mouse pups (fig. S4).

To determine whether cytosolic Ca^{2+} increased Na^+,K^+ -ATPase activity independently of GPCR activation, we exposed astrocytic cultures to increasing concentrations of the Ca^{2+} ionophore ionomycin (22). Ionomycin elicited a dose-dependent increase of ouabainsensitive $^{86}Rb^+$ uptake and Ca^{2+} responses in astrocyte cultures at $0.05~\mu M$ (Fig. 1E). Ionomycin also increased cytosolic Ca^{2+} in neuronal cultures; however, in neuronal cultures, it did not increase $^{86}Rb^+$ uptake (Fig. 1F). Thus, we concluded that increased cytosolic Ca^{2+} can stimulate Na^+,K^+ -ATPase activity in astrocytes, but not neurons.

GPCR-mediated astrocytic Ca²⁺ signals decrease extracellular K⁺ in hippocampal slices

We next investigated whether GPCR-mediated increases in astrocytic Ca^{2+} could regulate extracellular K^+ . We loaded hippocampal slices with the Ca^{2+} indicator rhod-2 AM (rhod-2 acetoxymethyl ester) to assess GPCR-activated increases in Ca^{2+} and confirmed that FMRF induced robust increases in rhod-2 signal in MrgA1^{+/-} mice, but not in MrgA1^{-/-} littermates (Fig. 2A). To determine whether GPCR-activated decreases in the extracellular K^+ concentration were secondary to an increase in astrocyte cytosolic Ca^{2+} concentration, we used mice in which astrocytes are unable to mobilize intracellular Ca^{2+} stores as a consequence of global deletion of IP₃R2 (IP₃R2^{-/-} mice) (23, 24). Application of ATP, UTP, or TFLLR-NH₂ increased cytosolic Ca^{2+} in slices from wild-type (IP₃R2^{+/+}) mice, whereas, consistent with previous studies (21, 24), they failed to do so in IP₃R2^{-/-} mice (Fig. 2B). Using K^+ -sensitive microelectrodes to measure extracellular K^+ (25) 80 μ m below the surface, we found that in the presence of a bath concentration of 4 mM K^+ ,

exposure to FMRF, ATP, UTP, or TFLLR-NH₂ elicited a transient decrease in the extracellular K⁺ concentration in wild-type mice (Fig. 2, C and D). This decrease in extracellular K⁺ concentration was markedly attenuated in IP₃R2^{-/-} mice (Fig. 2, C and D). For instance, whereas UTP induced a transient decline in extracellular K⁺ concentration averaging -0.43 ± 0.05 mM (n = 6) in slices prepared from wild-type mice, the K⁺ decline was only -0.13 ± 0.08 mM (n = 6) in IP₃R2^{-/-} mice. Similarly, the PAR1 agonist TFLLR-NH₂ decreased extracellular K⁺ by 0.33 ± 0.04 mM (n = 8) in IP₃R2^{+/+} mice, but not in IP₃R2^{-/-} mice (0.05 ± 0.08 mM, n = 5). Thus, experiments in hippocampal slices were consistent with the hypothesis that GPCR-activated increases in astrocyte cytosolic Ca²⁺ lead to decreased extracellular K⁺. These GPCR-activated decreases in extracellular K⁺ concentration were not apparent, however, when K⁺ concentration was recorded <30 μ m below the surface of the slice or at a subphysiological bath concentration of K⁺ (2.5 mM) (fig. S5).

Astrocytic Na⁺,K⁺-ATPase activity is stimulated by small increases in extracellular K⁺, but decreased at subphysiological extracellular K⁺ concentrations (26, 27). This enables astrocytes to transiently buffer the minor increases in extracellular K⁺ concentration that occur during neural activity and then slowly release K⁺ during the period of decreased extracellular K⁺ that follows (28). We wondered whether the increase in K⁺ uptake that occurs in response to increases in extracellular K⁺ concentration depended on cytosolic Ca²⁺ signals in astrocytes. We found that ouabain-sensitive ⁸⁶Rb⁺ uptake increased with increasing extracellular K⁺ concentrations and that preloading cultured astrocytes with CPA blocked the increase in ⁸⁶Rb⁺ uptake when the extracellular K⁺ concentration was increased from 2.5 to 5 mM, but not when extracellular K⁺ was increased to 10 or 20 mM (Fig. 3A). We therefore concluded that astrocytic buffering of modest increases of extracellular K⁺ is potentiated by cytosolic Ca²⁺ increases, whereas buffering of pathophysiological increases of extracellular K⁺ (10 mM or higher) is Ca²⁺-independent.

To assess whether Ca^{2+} -dependent K^+ uptake can contribute to buffering of increases in extracellular K^+ in response to more physiological stimuli, we compared high-frequency stimulation (HFS)-induced changes in extracellular K^+ in hippocampal slices from $IP_3R2^{+/+}$ and $IP_3R2^{-/-}$ mice. HFS consistently induced a higher peak increase in extracellular K^+ in $IP_3R2^{-/-}$ mice. Moreover, the undershoot during the K^+ recovery was reduced in $IP_3R2^{-/-}$ mice (Fig. 3, B and C), consistent with the lack of Ca^{2+} responses in slices prepared from $IP_3R2^{-/-}$ mice (Fig. 3C, left). Thus, we concluded that the ability of astrocytes to mobilize intracellular Ca^{2+} is a determinant of astrocytic K^+ uptake in hippocampal slices.

Na+ influx drives GPCR-mediated increases in K+ uptake

In many cell types, including astrocytes, the intracellular Na⁺ concentration provides the primary mechanism for regulating Na⁺,K⁺-ATPase activity (8, 9), which increases with increasing intracellular Na⁺ concentrations. For example, glutamate uptake stimulates Na⁺,K⁺-ATPase activity by increasing cytosolic Na⁺ (9). To define the role of cytosolic Na⁺ in GPCR-activated K⁺ uptake, we used several approaches to manipulate intra-astrocytic Na⁺. These included removal of extracellular Na⁺, application of the Na⁺ ionophore monensin, inhibition of the Na⁺,Ca²⁺ exchanger (NCX), and inhibition of store-operated Ca²⁺ channels (SOCs) (Fig. 4). We first assessed whether Na⁺ was necessary for GPCR-activated K⁺ uptake. Removal of extracellular Na⁺ reduced baseline ouabain-sensitive ⁸⁶Rb⁺ uptake and eliminated GPCR-activated increases, supporting the notion that Na⁺ is required for both baseline and Ca²⁺-induced increases in Na⁺,K⁺-ATPase activity (Fig. 4A). One mechanism whereby increases in intracellular Ca²⁺ could increase Na⁺ availability in astrocytes involves the NCX, which uses the energy stored in the Na⁺ electrochemical gradient to extrude Ca²⁺ (exchanging three Na⁺ for one Ca²⁺) (29, 30). We found that the NCX inhibitors SN-6 and SEA0400 (31) decreased TFLLR-NH₂-induced ⁸⁶Rb⁺ uptake

(Fig. 4B). Consistent with cytosolic Na⁺ being rate-limiting for Na⁺,K⁺-ATPase, the Na⁺ ionophore monensin triggered a dose-dependent increase of ⁸⁶Rb⁺ uptake in both astrocytic and neuronal cultures (Fig. 4C). Monensin-induced ⁸⁶Rb⁺ uptake was of larger amplitude than GPCR-activated uptake in astrocytes and severalfold higher in astrocytic than neuronal cultures (compare Figs. 1A and 4C), suggesting that the Na⁺,K⁺-ATPase activity is not maximally activated by GPCR-activated increases in cytosolic Ca²⁺. As previously reported (32), however, monensin also induced Ca²⁺ increases in cultured astrocytes. To dissect the relative contribution of Na⁺ influx versus Ca²⁺ increases, we attenuated monensin-induced cytosolic Ca²⁺ increases by preloading them with CPA. CPA reduced monensininduced ⁸⁶Rb⁺ uptake (by ~30%), indicating that the predominant action of monensin on Na⁺,K⁺-ATPase activity involves the increase in intracellular Na⁺ rather than that in Ca²⁺. The NCX inhibitors SN-6 and SEA0400 reduced ⁸⁶Rb⁺ uptake without decreasing the intracellular Ca²⁺ concentration, suggesting that SN-6 and SEA0400 not only inhibit NCX, but may have other effects, or that the NCX may contribute to ⁸⁶Rb⁺ uptake through some alternative mechanism. Thus, this analysis indicates that the NCX contributes to GPCRactivated increases in ouabain-sensitive ⁸⁶Rb⁺ uptake and that it is possible to bypass the NCX by directly increasing intracellular Na⁺ concentration (Fig. 4D).

Depletion of intracellular Ca^{2+} stores during Ca^{2+} signaling in astrocytes has previously been shown to result in opening of SOCs (33). These Ca^{2+} channels, which in astrocytes include transient receptor potential C1 (TRPC1) channels (34), represent a major pathway for Ca^{2+} entry in nonexcitable cells (35, 36). Indeed, using Ca^{2+} -sensitive micro-electrodes, we found that GPCR-activated Ca^{2+} signaling in astrocytes induced a transient decrease in extracellular Ca^{2+} averaging ~0.1 mM, from a resting concentration ranging from 1.2 to 1.4 mM (fig. S6) (33, 37). SKF96365, a nonspecific inhibitor of TRPC1 (38), shortened GPCR-activated Ca^{2+} increases and suppressed ${}^{86}Rb^{+}$ uptake (Fig. 4, E and F).

We used K⁺-sensitive microelectrodes to determine whether the NCX plays a role in GPCR-mediated K⁺ uptake in hippocampal slices. Na⁺ in the bath was replaced with *N*-methyl-D-glucamine (NMDG⁺). Na⁺ suppressed GPCR-activated decreases in extracellular K⁺ (Fig. 4, G and H). Moreover, the NCX inhibitors SN-6 and SEA0400 attenuated the transient decline in extracellular K⁺ triggered by TFLLR-NH₂ (Fig. 4, G and H). The TRPC1 inhibitor SKF96365 also attenuated GPCR-activated decreases in extracellular K⁺ and suppressed TFLLR-NH₂-induced Ca²⁺ increases (Fig. 4I). Together, these observations suggest that high cytosolic Ca²⁺ activates the NCX. In turn, the inflow of Na⁺ ions activates the Na⁺,K⁺-ATPase, resulting in a transient decrease in extracellular K⁺ (Fig. 4J).

Support for this model was obtained by an analysis of cytosolic Na⁺ concentration in cultured astrocytes. Astrocytes loaded with the fluorescent Na⁺ indicator sodium-binding benzofuran isophthalate (SBFI) (26, 27) showed a small increase in SBFI fluorescence when exposed to either ATP or FMRF in hippocampal slices from MrgA1^{+/-} mice (n = 5; P < 0.01 compared to before, t test) that paralleled in increase in cytosolic Ca²⁺ (fig. S6). Thus, several lines of analysis show that increases in cytosolic Na⁺ may be a prerequisite for GPCR-activated reductions in extracellular K⁺.

Na⁺,K⁺-ATPase activity is regulated by both protein kinase C (PKC)–and PKA-mediated phosphorylation in a broad range of cell types (39, 40). However, an analysis of cultured astrocytes showed that pharmacological activation or inhibition of PKA or PKC activity consistently suppressed GPCR-activated ⁸⁶Rb⁺ uptake (fig. S7). This suggests that PKC- or PKA-mediated modulation of the Na⁺,K⁺-ATPase activity is optimal under resting in cultured astrocytes. However, kinase regulation of the Na⁺,K⁺-ATPase activity is complex and dependent on the ambient concentrations of cytosolic Ca²⁺ and Na⁺ (39). Thus, we

cannot eliminate the possibility that kinase modulation regulates the activity of the Na⁺,K⁺-ATPase activity in vivo.

Astrocytic Ca²⁺ signaling suppresses basal excitatory synaptic activity

Given that the Na⁺,K⁺-ATPase activity in astrocytes is modulated by changes in the cytosolic Ca²⁺ concentration, we wondered whether astrocytes might modulate synaptic activity through dynamic changes in extracellular K⁺. Although several studies have addressed the role of pathological increases in extracellular K⁺ on synaptic activity (17, 41, 42), the effects of lowering extracellular K⁺ are less clear. We achieved reductions in extracellular K⁺ by either exposing hippocampal slices from MrgA1^{+/-} mice to FMRF for 1 min (Fig. 5, A and B and E and F, and fig. S8) or reducing bath K⁺ from 4 to 2 mM for 2 min (Fig. 5, C and D and G and H, and fig. S8). Extracellular K^+ fell from 3.81 \pm 0.07 to 3.14 ± 0.06 mM (n = 13) when bath K⁺ was lowered from 4 to 2 mM, similar to the drop in extracellular K⁺ evoked by GPCR-activated astrocytic Ca²⁺ signaling (see Figs. 2D and 4I). CA1 hippocampal neurons typically underwent a small hyperpolarization in response to decreasing the bath extracellular K⁺ concentration (-3.44 ± 0.37 mV, n = 13). Similarly, FMRF triggered a small hyperpolarization in CA1 neurons in MrgA1^{+/-} mice (-1.78 ± 0.33) mV, n = 6), whereas it did not affect the membrane potential $(0.22 \pm 0.12 \text{ mV}, n = 6)$ of CA1 neurons in littermates (MrgA1^{-/-}). Thus, Ca²⁺ signaling in hippocampal astrocytes is linked to a minor hyperpolarization of the membrane potential of CA1 neurons.

Neuronal hyperpolarization was associated with a decrease in the frequency of miniature excitatory postsynaptic currents (mEPSCs) (Fig. 5, A to D). The frequency of mEPSCs fell from 1.37 ± 0.17 to 0.80 ± 0.10 Hz during FMRF application (n = 9, Fig. 5B) and from 1.61 ± 0.27 to 0.68 ± 0.13 Hz when bath K⁺ concentration was lowered to 2 mM K⁺ (n = 8, Fig. 5D). Moreover, repolarization of mEPSCs occurred more rapidly when the extracellular K⁺ concentration was reduced (Fig. 5, B and D), likely reflecting the increase in K+ driving force. mEPSC recovery (assessed as the time constant of decay, τ) fell from 97.2 ± 13.6 to 66.5 ± 7.9 ms during FMRF application (n = 9, Fig. 5B) and from 95.4 ± 16.6 to 63.6 ± 14.5 ms when bath K^+ concentration was lowered to 2 mM K^+ (n = 8, Fig. 5D). Lowering of extracellular K⁺ may therefore suppress excitatory synaptic activity by reducing both the probability of glutamate release from presynaptic terminals and the duration of the depolarization of the postsynaptic membrane. In contrast, miniature inhibitory postsynaptic currents (mIPSCs) were largely unaffected by lowering of extracellular K⁺ (Fig. 5, E to H). mIPSC frequency before and after FMRF application was 0.56 ± 0.08 to 0.45 ± 0.10 Hz (n =7, Fig. 5, E to F) and 0.66 ± 0.03 to 0.62 ± 0.06 Hz (n = 7) before and after reduction of bath K⁺ concentration from 4 to 2 mM (Fig. 5, G and H). Moreover, lowering of bath K⁺ concentration or exposure to FMRF had no effects on mIPSC recovery (τ) (Fig. 5H). The lack of effects with manipulation of extracellular K⁺ on inhibitory transmission has been previously reported (43, 44). Postsynaptically, the IPSCs are mediated by a transient increase in Cl⁻ conductance, which is insensitive to minor changes in extracellular K⁺ (45). Moreover, presynaptically, inhibitory neurons express Kv3 potassium channels in their terminals. Kv3 channels are inactive at resting membrane potential, and their open probability is not altered in response to minor voltage changes (46–48). Thus, the net effect of lowering extracellular K⁺ is a selective reduction of spontaneous excitatory transmission, which can be considered a suppression of background synaptic noise.

Astrocytic Ca²⁺ signaling enhances the signal-to-noise ratio of excitatory synaptic transmission

Given that astrocytic Ca²⁺ signaling decreased basal excitatory activity, we next asked what effect it had on synaptic transmission. We used evoked excitatory postsynaptic potentials (eEPSPs) as signal and spontaneous EPSPs (sEPSPs) as a measurement of background

afferent synaptic transmission (noise) by obtaining whole-cell recordings from CA1 hippocampal neurons while stimulating the Schaeffer collaterals with a stimulation electrode placed in the stratum radiatum (stimulation interval, 2 s) (Fig. 6A). FMRF decreased the frequency of sEPSPs in slices from MrgA1^{+/-} mice, but not the amplitude of eEPSP concurrently with a transient decrease in the extracellular K⁺ concentration (Fig. 6A). In contrast, FMRF failed to alter either the frequency of sEPSPs, the amplitude of the eEPSP, or the concentration of extracellular K⁺ in MrgA1^{-/-} littermates (Fig. 6A). To more broadly assess the effect of astrocytic Ca²⁺ signaling on excitatory synaptic transmission, we next exposed slices to an array of GPCR agonists, including ATP, UTP, 2MeSADP, and TFLLR-NH₂. Combined whole-cell recordings of CA1 neurons with measurements of extracellular K⁺ showed that GPCR-activated pathways triggered a transient neuronal hyperpolarization (fig. S9) concomitantly with a decrease in the frequency of sEPSPs (Fig. 6B). The frequency of sEPSPs decreased as a direct function of the reduction of extracellular K⁺ (Fig. 6B). To address the effect of GPCR-activated Ca²⁺ signaling in astrocytes on hippocampal network function, we calculated the signal-to-noise ratio (SNR): eEPSPs were taken as signal and sEPSPs were taken as noise (Fig. 6C). The integrated area of all the eEPSPs was divided by the integrated area of all the sEPSPs within the same time period (49–51). We found that the SNR increased in response to GPCR-activated Ca²⁺ signaling in astrocytes, primarily a reflection of the fact that the eEPSP amplitude was insensitive to reductions in the extracellular K⁺ concentration (fig. S9), whereas the frequency of sEPSPs decreased in response to astrocytic K⁺ uptake (Fig. 6B).

We next asked whether a reduction in bath K⁺ concentration was sufficient to reproduce the effects of GPCR-activated astrocytic Ca²⁺ signaling on excitatory synaptic transmission. We first assessed the impact of minor transient reductions of bath K⁺ on sEPSPs and eEPSPs. In these experiments, whole-cell recordings of CA1 pyramidal neurons were combined with measurements of extracellular K⁺, because the actual changes in interstitial K⁺ concentration were of smaller amplitude than in the bath solution (fig. S9A). For example, reducing bath K⁺ concentration from 4 to 2 mM for 2 min only lowered interstitial K⁺ concentration by \sim 0.6 mM (from 3.84 \pm 0.06 to 3.24 \pm 0.06 mM, n = 5) (fig. S9D). CA1 neurons typically exhibited a minor hyperpolarization in response to lowering the extracellular K⁺ concentration, and the changes in membrane potential were a linear function of bath K⁺ concentration ($t^2 = 0.61$, fig. S9E). This hyper-polarization was associated with a suppression of basal excitatory activity detected as a reduction of the frequency of sEPSPs (Fig. 6, D and E). Plotting sEPSP as a function of the decrease in extracellular K⁺ concentration showed that the frequency (Fig. 6E) of sEPSPs was inversely related to extracellular K^+ ($t^2 = 0.36$). Thus, lowering the extracellular K^+ concentration reduced spontaneous excitatory activity. Recordings of eEPSP in CA1 neurons showed that eEPSP amplitude was largely insensitive to changes in extracellular K⁺ over the concentration range studied. Consequently, the ratio of integrated areas for eEPSPs and sEPSPs over the same time period increased as an inverse function of extracellular K^+ ($r^2 = 0.54$) (Fig. 6F). We next compared the effect of activation of GPCRs in astrocytes upon sEPSP and eEPSPs in hippocampal slices from IP₃R2^{-/-} mice. UTP (100 μ M) failed to change the frequency of sEPSP (after/before = 1.02 ± 0.03 , n = 5) in slices prepared from IP₃R2^{-/-} mice (Fig. 6G). In contrast, eEPSP amplitude was insensitive to UTP. Thus, mobilization of intracellular Ca²⁺ store in astrocytes was essential for UTP-induced suppression of sEPSPs. In contrast, direct reduction of bath K⁺ concentration triggered membrane hyperpolarization and a decrease in the frequency of sEPSP in IP₃R2^{-/-} mice (Fig. 6G).

These observations indicate that the reduction in extracellular K^+ concentration induced by astrocytic Ca^{2+} signaling leads to an increase in eEPSPs versus sEPSPs.

Astrocytic Ca²⁺ signaling reduces the frequency of synaptic failures

The stimulation protocol used above for eEPSPs will recruit essentially all fibers in the Schaffer collaterals and may not detect minor modulation of synaptic transmission triggered by decreased extracellular K⁺ concentration. To investigate the possibility that astrocytic Ca²⁺ signaling enhances the evoked response directly, we used a minimal stimulation protocol (52, 53) in which we placed a glass electrode in the stratum radiatum in hippocampal CA1 and adjusted the stimulus to approximate the threshold for getting synaptic failures (~50%). By stimulation of a small number of Schaffer collaterals (potentially a single fiber), the effects of minor reductions in extracellular K⁺ on synaptic fidelity can be quantified. At 4 mM extracellular K⁺, the rate of synaptic failures detected by whole-cell recording from CA1 neurons averaged 0.46 ± 0.03 (n = 33). FMRF application in MrgA1^{+/-} slices (Fig. 7A) or directly lowering bath K⁺ concentration of these slices (Fig. 7B) consistently reduced conduction failures (Fig. 7C). No effects were observed on synaptic potency (mean amplitude of successful stimuli) (Fig. 7D), indicating that the decrease in conduction failure was associated with an increase in synaptic efficacy (mean amplitude of all stimuli) (Fig. 7E). Because mGluRs reduce astrocyte-mediated increases in synaptic efficacy (52, 53), we administered FMRF to MrgA1^{+/-} slices in the presence of the mGluR antagonists 6-methyl-2-(phenylethynyl)pyridine hydrochloride (MPEP) (50 μM) and LY367385 (100 μ M) and found that synaptic failures still decreased from 0.46 \pm 0.07 to 0.24 ± 0.04 (n = 6, P < 0.01, paired t test). Similarly, synaptic efficacy increased from 1.02 ± 0.04 0.13 mV to $1.5 \pm 0.19 \text{ mV}$ (n = 6, P < 0.01, paired t test), whereas synaptic potency remained unchanged, 1.96 ± 0.19 mV to 1.97 ± 0.08 mV (n = 6, P = 0.38). Thus, MPEP and LY367385 did not interfere with suppression of synaptic failures evoked by FMRF-induced astrocytic Ca²⁺ signaling. The minimal stimulation protocol provides a more sensitive dissection of the effect of astrocytic Ca²⁺ signaling on eEPSPs. The net effect of astrocytic activation is therefore a transient increase in circuit SNR.

GPCR-activated astrocytic Ca²⁺ signaling mediates a decrease in extracellular K⁺ concentration in vivo

We next assessed the effects of astrocytic Ca^{2+} signaling on extracellular K^+ in live adult mice, using topical application of agonists in an open cranial window preparation to trigger astrocytic Ca^{2+} signaling. We used this approach, rather than evoking Ca^{2+} signaling by sensory stimulation, because interpretation of sensory-evoked changes in extracellular K^+ concentration is complicated by the direct effect of synaptic activity upon extracellular K^+ (17, 42, 54, 55). We first evaluated the efficacy of different agonists in triggering cytosolic Ca^{2+} increases in cortical astrocytes. The exposed cortex was loaded with the Ca^{2+} indicator rhod-2 AM (56–59) and superfused with either ATP (100 μ M) in Glt1-eGFP (enhanced green fluorescent protein) reporter mice (60) or FMRF in MrgA1+/- mice (21). Both ATP and FMRF evoked robust increases in cytosolic Ca^{2+} in astrocytes located 100 μ m below the pial surface (Fig. 8, A to D). Moreover, extracellular K^+ recorded at a depth of ~100 μ m exhibited a transient decline in response to either ATP or FMRF superfusion (Fig. 8, E and F). Amplitudes of Ca^{2+} increases ($\Delta F/F$) and reductions in extracellular K^+ were similar to those observed in acute slice preparations from mouse pups. Thus, astrocytic Ca^{2+} signaling in live adult mice can also trigger a transient reduction in the extracellular K^+ concentration.

DISCUSSION

We show here that GPCR-mediated Ca^{2+} signaling in astrocytes is linked to an increase in K^+ uptake resulting in a transient decrease in the extracellular concentration of K^+ (Fig. 1). The average decrease in the extracellular K^+ concentration was in the range of 0.3 to 0.6 mM and lasted 20 to 80 s in hippocampal slice preparations as well as in live adult mouse cortex (Figs. 2 and 8). A detailed analysis suggested that the transient increase in astrocytic

K⁺ uptake was mediated primarily by the Na⁺,K⁺-ATPase (Fig. 4 and fig. S1). The intracellular Na⁺ concentration has been identified as a factor that limits the rate of Na⁺,K⁺-ATPase activity in astrocytes as in other cell types (8–11). We confirmed this observation and showed that Ca²⁺-mediated activation of the Na⁺,K⁺-ATPase, at least in part, is the result of Na⁺ influx through the Na⁺,Ca²⁺ exchanger, because removal of extracellular Na⁺ or addition of inhibitors of the Na⁺,Ca²⁺ exchanger reduced or eliminated GPCR-activated astrocytic K⁺ uptake (Fig. 4). In support of the proposed mechanism (Fig. 4J), we found that cytosolic Na⁺ concentration in hippocampal astrocytes increased slightly in response to activation of GPCR pathways (fig. S7). The classical role of astrocytes is to buffer excess K⁺ released during synaptic transmission. Previous analyses have shown that astrocytes are specialized for rapid removal of K⁺, because the astrocytic Na⁺,K⁺-ATPase exhibits an unusual sensitivity to minor increases in extracellular K⁺ (26, 27). We confirmed these original observation and extended them by showing that Ca²⁺ signaling potentiated K⁺ uptake in response to modest increases of extracellular K⁺ (5 mM), but not when K⁺ is raised to pathological concentrations (10 mM or higher) (Fig. 3). Moreover, we showed that GPCR-activated Ca²⁺ signaling in astrocytes reduces baseline extracellular K⁺ concentration in slices (Figs. 2D and 4H) and in vivo (Fig. 8). As expected, lowering the extracellular K⁺ concentration led to a neuronal hyperpolarization averaging -2 to -4 mV. The hyperpolarization was observed in response to direct lowering of bath K⁺ concentration, or indirectly by stimulating astrocytic Ca²⁺ signaling. As previously described, lowering the extracellular K⁺ concentration decreased the frequency and the duration of mEPSCs (61, 62) in hippocampal slices, but had little effect on eEPSP amplitude (Figs. 5 and 6) (43, 44). However, when a minimal stimulation protocol was used, we found that astrocytic Ca²⁺ signaling triggered a small but significant decrease in synaptic failures ($P \le 0.01$ compared to "Before," Bonferroni-Dunn test, n = 7, Fig. 7). Thus, the transient reduction of extracellular K⁺ evoked by astrocytic Ca²⁺ signaling decreased synaptic noise (concomitant with an increase in synaptic fidelity detected as a decrease in synaptic failures). The net effect was that astrocytic Ca²⁺ signaling improved the SNR of hippocampal neurotransmission. Thus, Ca²⁺-mediated K⁺ uptake endows astrocytes with a simple yet powerful mechanism for rapid modulation of neural activity. We have observed Ca²⁺dependent K⁺ uptake in cortex, hippocampus, and cerebellum, suggesting that this signaling pathway is active in many areas of CNS.

Potassium homeostasis is unquestionably an essential housekeeping duty of astrocytes. The concept of astrocytic "K+ spatial buffering" in which K+ is taken up in regions of high extracellular K+ concentration and released at distant sites was first advanced in a leech CNS preparation (63) and later confirmed in isolated retinal Muller cells (64). It was subsequently established that astrocytic K+ uptake is accomplished through multiple mechanisms (10). One study found that the astrocytic Na+,K+-ATPase constituted the dominant mechanism for K+ recovery after high-frequency neuronal activity (9), whereas other studies provided evidence suggesting that inwardly rectifying K+ channels play a predominant role in maintenance of resting extracellular K+ concentration and recovery from mild neuronal activity (1, 8, 9). However, the relative importance of the Na+,K+-ATPase and inward rectifying potassium channels in K+ buffering may exhibit regional differences. For example, the Na+,K+-ATPase is solely responsible for K+ buffering in rat optic nerve (1), whereas K+ siphoning through inwardly rectifying K channels is important in the retina (64). Thus, additional studies are needed to definitively establish whether the mechanism described here is active in regions outside the hippocampus and cortex.

Although our in vitro analysis indicates that the Na⁺,Ca²⁺ exchanger, at least in part, provides Na⁺ ions to fuel the Na⁺,K⁺-ATPase, the catalytic activity of the Na⁺,K⁺-ATPase is regulated by multiple pathways, including phosphorylation of the catalytic α subunit (65). The astrocytic Na⁺,K⁺-ATPase is also activated by minor increases in extracellular K⁺

concentration, consistent with the role in astrocytic buffering of excess K^+ released during neuronal activity (26, 27). Additional studies are required to establish the relative contribution of the many potential regulatory pathways that determine the activity of the Na^+, K^+ -ATPase.

Through strict regulation of the extracellular K⁺ concentration, astrocytes exert powerful control over the membrane potential of both excitatory and inhibitory synapses. Why then, do astrocyte-mediated Ca²⁺-dependent decreases in extracellular K⁺ modulate excitatory and not inhibitory transmission? This question can be addressed by considering the differences between these two modes of synaptic communication in light of the mechanisms by which lowering the extracellular K⁺ concentration modulates synaptic transmission: (i) the decay kinetics of the mEPSC versus those of the mIPSC, (ii) the voltage dependence of presynaptic ion channels in excitatory and inhibitory nerve terminals, and (iii) the probability of axonal conductance failure. First, the shortening of mEPSCs we observed in response to lowering of extracellular K⁺ has been reported before (66) and likely reflects that cation currents drive excitatory currents. Low K⁺ will tend to shorten mEPSCs, because the decay of mEPSCs in part is driven by the outward K⁺ current. Simply, a reduction of extracellular K⁺ will increase the driving force for the outward K⁺ currents and therefore shorten the decay of mEPSCs (61). In contrast, the decay of mIPSCs is primarily mediated by desensitization of GABA_A (γ -aminobutyric acid type A) receptors (45) and is therefore insensitive to minor decreases in the extracellular K⁺ concentration. The decrease in the frequency of mEPSCs, but not that of mIPSCs, in response to low extracellular K⁺ likely might result from differences in the properties of potassium channels found in nerve terminals of excitatory (67) versus inhibitory neurons (47, 48). Potassium channels in excitatory neurons exhibit rapid depolarization-induced inactivation with slow reactivation kinetics upon repolarization (67). In contrast, potassium channels in inhibitory neurons are activated only at relatively depolarized potentials, and their open probability is not altered in response to neuronal hyperpolarization (68). The net result of astrocytic Ca²⁺-mediated K⁺ uptake is therefore a decrease in the frequency and duration of mEPSCs, whereas mIPSCs are largely insensitive. Lastly, the increase in synaptic fidelity associated with low extracellular K⁺ is most likely a consequence of the direct effect of extracellular K⁺ on axonal conduction (44). High extracellular K⁺ concentration increases the rate of action potential conduction failure in various types of neurons (44, 69), an observation that has been attributed to the slowing of Na⁺ channel recovery from inactivation. Conversely, a decrease in K⁺ concentration will hasten Na⁺ channel recovery and improve action potential conduction reliability. The consequence of the transient lowering of extracellular K+ concentration during GPCR-activated astrocytic Ca²⁺ signaling is thereby a selective reduction in mEPSCs (background noise) concomitant with enhancement of conduction reliability (signal). Thus, astrocytic Ca²⁺ signaling will enhance network SNR and thereby improve information processing. The ability of astrocytes to decrease extracellular K⁺ concentration provides an elegantly simple and powerful means for rapid astrocytic modulation of network activity. Previous studies on neuroglial signaling have primarily focused on gliotransmitters rather than measurements of extracellular K⁺ or neuronal membrane potential. In addition, recordings in this study were obtained in neurons located deeper than 80 µm from the surface of the slice. We found that GPCR-activated changes in K⁺ were of smaller amplitude near the surface of the slice, where the bath solution rapidly buffers the response (fig. S5). Lastly, many previous studies of neuroglial signaling used bath solution containing 2.5 mM K⁺, an artificially low concentration of K⁺, given extracellular K⁺ in vivo is in the range of 3.0 to 4.0 mM in rat brain cortex (58). GPCRactivated astrocytic modulation of EPSPs was not detectable when we used a bath solution containing 2.5 mM K⁺ (fig. S5). Thus, it is possible that these and other experimental details have attenuated the effects of Ca²⁺-dependent K⁺ uptake in astrocytes and obscured this mechanism of neuroglial signaling.

Our findings support the existence of a previously unappreciated mechanism by which astrocytes can modulate neuronal function. Hence, astrocytic control of extracellular K^+ concentration may, at least in part, be responsible for effects on neural networks that hitherto have been ascribed to Ca^{2+} -dependent release of gliotransmitters. Neuromodulators, many of which potently increase astrocytic Ca^{2+} signals (59), are likewise well suited to mediate large-scale network modulation through astrocytic control of extracellular K^+ .

MATERIALS AND METHODS

Cultures

Cultured neocortical astrocytes were prepared from 1- to 2-day-old Wistar rat pups (Taconic Farms Inc.) or C57BL/6 mouse pups (Taconic Farms) as previously described (70). Briefly, cerebral cortices were dissected and meninges were removed. The tissue was washed three times in Hank's balanced salt solution without Ca²⁺. Once washed, the tissue was triturated, filtered through 70-µm nylon mesh, and centrifuged. The pellet was resuspended in 10% fetal bovine serum in Dulbecco's modified Eagle's medium (DMEM)/F12 containing penicillin (100 IU ml⁻¹) and streptomycin (100 µg ml⁻¹) and transferred to culture flasks. Cells were maintained at 37°C in an incubator containing humidified air and 5% CO₂. Medium was changed after 24 hours and twice a week thereafter. More than 95% of the cells stained positive for GFAP. When the monolayers became confluent, they were rinsed two times in Hank's balanced salt solution without Ca²⁺, suspended in 0.05% trypsin-containing solution for 1 min, resuspended in DMEM/F12, centrifuged to remove trypsin, and then plated in 24-well plates. Experiments were performed when the cells were 95% confluent.

A comparison of ⁸⁶Rb⁺ uptake in astrocytic cultures prepared from rat and mouse astrocytes as well as cultures of rat cortical and hippocampal astrocytes showed no differences in basal or GPCR-activated ⁸⁶Rb⁺ uptake, except for a higher sensitivity to ouabain of cultures prepared from rat (fig. S4). Unless otherwise noted, rat neocortical cultures were used for ⁸⁶Rb⁺ uptake experiments. Cultured neocortical neurons were prepared from 1- to 3-dayold Wistar rat pups as previously described (42), with slight modification. Briefly, cerebral cortices were dissected and meninges were removed. The tissues were minced and then suspended in Hibernate A solution minus calcium (BrainBits, LLC) containing papain (Worthington) at 37°C for 30 min. The suspension was then triturated with a 9-inch siliconized pipette. The cell suspension was then separated by density gradient centrifugation. The neuronal cells were then seeded onto poly-D-lysine-coated 24-well plates and maintained in Neurobasal A medium (Invitrogen) with 0.5 ml of B27, 0.5 mM GlutaMAX, gentamycin (10 µg ml⁻¹), and human fibroblast growth factor 2 (FGF2) (5 ng ml⁻¹) at 37°C in an incubator containing humidified air and 5% CO₂. Immunohistochemistry showed that more than 90% cells in the cultures [identified by DAPI (4',6-diamidino-2-phenylindole)-stained nuclei] were MAP2-positive. Neuronal cultures were used 8 days after plating. All animal experiments were conducted in accordance with the guidelines of the Animal Care and Use Committee of the University of Rochester.

⁸⁶Rb⁺ uptake assay and Ca²⁺ imaging of cultured cells

On the day of the experiment, culture medium was replaced with oxygenated artificial cerebrospinal fluid (aCSF) (pH 7.4, osmolarity = 310 mosM). After a 30-min recovery period, the cells were incubated for 10 min in the presence or absence of ouabain. The potassium analog $^{86}Rb^+$ was added to each of the 24 wells for 15 min (1 μ Ci, Perkin Elmer). The reaction was stopped by placing the cells on ice and washing them three times with ice-cold aCSF. The remaining aCSF was removed, the cells were lysed with NaOH (100 μ l of 0.5 N), and $^{86}Rb^+$ uptake was quantified by liquid scintillation counting (Beckman Coulter). Ouabain-sensitive $^{86}Rb^+$ uptake was calculated by subtracting $^{86}Rb^+$ uptake during the same

conditions, but in the presence of ouabain (1 mM). For experiments where 5, 10, and 20 mM K^+ were used to stimulate $^{86}Rb^+$ uptake, the $^{86}Rb^+$ concentration was increased proportionally with K^+ concentration.

For imaging of GPCR-activated increases in Ca²⁺, cultures were loaded with rhod-2 AM (4.5 μ M), and rhod-2 signal was visualized by confocal microscopy (FV500, Olympus) every 3 s before and after application of agonists. The peak fluorescence intensity was calculated as percentage of baseline intensity before stimulation (Δ *FI*). Pharmacological agents used in cultures included adenosine (100 μ M), ATP (100 μ M), CPA (20 μ M), glutamate (100 μ M), H89 (10 μ M), ionomycin (0.01 to 0.05 μ M), monensin (5 nM to 3 μ M), *t*-ACPD (100 μ M), 2MeSADP (100 μ M, Tocris), N^6 ,2′-O-dibutyryladenosine cAMP (DBcAMP, 1 mM), PGE₂ (20 μ M, Tocris), ouabain (1 mM), FMRF (15 μ M), rhod-2 AM (4.5 μ M, Invitrogen), phorbol 12-myristate 13-acetate (PMA, 1 μ M), SEA0400 (10 μ M, synthesized by Taisyo Pharmaceutical Co. Ltd.), potassium chloride (KCl, 5, 10, 20 mM), 1-[2-(4-methoxyphenyl)-2-[3-(4-methoxyphenyl)propoxy]ethyl]imidazole (SKF96365, 25 μ M), (2-[[4-[(4nitrophenyl) methoxy]phenyl]methyl]-4-thiazolidinecarboxylic acid ethyl ester (SN6, 10 μ M, Tocris), staurosporine (2 μ M), UTP (100 μ M), TFLLR-NH₂ (30 μ M, Tocris), MPEP (50 μ M), LY367385 (100 μ M), and DL-*threo*- β -benzyloxyaspartic acid (TBOA, 100 μ M, Tocris). All chemicals were from Sigma unless otherwise noted.

Fabrication and calibration of K+- and Ca²⁺-sensitive microelectrodes

To detect K⁺ changes in the extracellular space, we fabricated ion-sensitive microelectrodes from double-barreled pipette glass (PB150F-6, WPI) pulled to a tip of <3 μ m with a puller (Sutter P-97). Pipettes were silanized by dimethylsilane (Fluka, Sigma) loaded with a 100-to 150- μ m column of valinomycin-based K⁺ ion-exchange resin (Potassium Ionophore 1-Cocktail B, Fluka, Sigma), and backfilled with 150 mM KCl. The reference barrel was filled with Hepes-buffered 150 mM NaCl solution (pH 7.4) (25, 71, 72). The pipette was connected to headstages of an Axoprobe 700B amplifier (Axon Instruments). Electrodes were calibrated in 1 to 6 mM K⁺ in aCSF before and after experiments. The recording pipette was located ~100 μ m below the surface of the slice. The electrodes used in experiments typically displayed voltage responses of 5 to 6 mV per millimolar increase in K⁺ concentrations in the concentration range studied. In selected experiments, single-barrel pipettes were used in combination with a reference electrode located ~10 μ m away.

Fabrication and use of Ca²⁺-sensitive microelectrodes was similar to that of K⁺-sensitive microelectrodes, except that the Ca²⁺ ion-exchange resin (Calcium Ionophore 1-Cocktail A, Fluka, Sigma) was used and the pipettes were backfilled with 100 mM CaCl₂ (73).

Slice preparation, electrophysiology, and Ca²⁺ and Na⁺ imaging

Unless otherwise noted, 15- to 21-day-old C57BL/6 (Charles River), MrgA1^{+/-}, MrgA1^{-/-}, $IP_3R2^{+/+}$, and $IP_3R2^{-/-}$ (21, 24) pups were used for preparation of hippocampal slices as previously described (74). The pups were anesthetized with isoflurane in a closed chamber (1.5%) and decapitated. The brains were rapidly removed and immersed in ice-cold cutting solution that contained 230 mM sucrose, 2.5 mM KCl, 0.5 mM CaCl₂, 10 mM MgCl₂, 26 mM NaHCO₃, 1.25 mM NaH₂PO₄, and 10 mM glucose (pH 7.2 to 7.4). Coronal slices (400 μ m) were cut with a vibratome (Vibratome Company) and transferred to oxygenated aCSF that contained 126 mM NaCl, 4 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 26 mM NaHCO₃, 1.25 mM NaH₂PO₄, and 10 mM glucose (pH 7.2 to 7.4, osmolarity = 310 mosM). Slices were incubated in aCSF for 1 to 5 hours at room temperature before recording. Experiments were performed at room temperature (21° to 23°C). During the recordings, the slices were placed in a perfusion chamber and superfused with aCSF gassed with 5% CO₂ and 95% O₂

at room temperature. Cells were visualized with a 40x water-immersion objective and differential inference contrast (DIC) optics (BX51 upright microscope, Olympus Optical).

Patch electrodes were fabricated from filament thin-wall glass (World Precision Instruments) on a vertical puller; the resistance of the pipette was about 6 to 9 megohms with intracellular pipette solution added. Membrane potentials were recorded under current clamp (I= 0) with an Axopatch MultiClamp 700B amplifier (Axon Instruments). The pipette solution contained 140 mM K-gluconate, 5 mM Na-phosphocreatine, 2 mM MgCl₂, 10 mM Hepes, 4 mM Mg-ATP, and 0.3 mM Na-GTP (pH adjusted to 7.2 with KOH). With these solutions, the IPSCs reversed at -70 mV, and EPSCs reversed at about 0 mV. The junction potential between the patch pipette and the bath solution was zeroed before forming a gigaseal. Patches with seal resistances of less than 1 gigohm were rejected. Data were low pass–filtered at 2 kHz and digitized at 10 kHz with a Digidata 1440 interface controlled by pClamp Software (Molecular Devices).

In experiments recording GPCR-activated changes in K⁺ or EPSCs, agonists were delivered by superperfusion for a period of 1 min unless otherwise noted. mEPSCs were isolated by adding 0.5 µM tetrodotoxin (TTX) and bicuculline methylchloride (50 µM) to the bath solution and maintaining a holding potential of -70 mV. Miniature IPSCs were isolated with 0.5 µM TTX, CNQX (6-cyano-7-nitroquinoxaline-2,3-dione) (25 µM), and APV (D,L-2amino-5-phosphonovaleric acid) (50 μ M) and a holding potential of -50 mV. HFS was delivered as a train of 300 square pulses (200 µs, 50 to 100 µA) at 100 Hz applied in the stratum radiatum. Minimal stimulation was used to activate single or few presynaptic fibers (52, 53). Briefly, a glass pipette filled with aCSF (3 µm tip) was positioned in the stratum radiatum so that 100-µs current pulse evoked postsynaptic potentials in a patched CA1 neuron. The stimulus strength (0.5-1 mA) was titrated to obtain ~50% synaptic failures and was not changed during the experiment. Each condition was tested by delivering 20 stimulations at 1 Hz. Synapses with unstable postsynaptic potential amplitudes were rejected. We analyzed (i) synaptic efficacy defined as mean peak amplitude of all responses including failures, (ii) synaptic potency defined as mean peak amplitude of all successful responses, and (iii) probability of release or the ratio of the success versus total number of stimuli (n = 20). For analysis of paired-pulse facilitation, an electrode (FHC, CBARC75) was placed in the stratum radiatum at a distance of 100 µm from the patched CA1 neurons. eEPSPs were induced by delivering a 50- to 100-μA, 200-μs current. Analysis of data was performed with Clampfit 10.0 software (Axon Instruments) and SigmaPlot (Systat Software). For SNR analysis, eEPSPs were taken as signal, and sEPSPs were taken as noise. The integrated area of all the eEPSPs divided by the integrated area of all the sEPSPs within the same time period was used as SNR. The integrated area was calculated by the mean area multiplied by averaged frequency with Event Statistics collected in Clampfit 10.0 software. All results are reported as means ± SEM, and significance was determined by paired or unpaired t tests or Tukey-Kramer post hoc multiple comparison tests.

For Ca^{2+} imaging experiments, the slices were loaded with rhod-2 AM (4.5 μ M, 1 hour). A custom-built microscope attached to Tsunami/Millennium laser (10 W, SpectraPhysics) and scanning box (FV 300, Olympus) with FluoView software and 40× objective (0.9 numerical aperture, Olympus) was used in all experiments (56). The excitation wavelength was fixed at 820 nm, whereas two-channel detection of emission was achieved by a 565-nm dichroic mirror (Chroma) and two external photomultiplier tubes. A 525/50 band-pass filter (Chroma) was used to detect GFP emission wavelength, and a 620/60 band-pass filter (Chroma) to detect rhod-2 signals. Relative changes in rhod-2 signaling were quantified with FluoView software. Time-lapse images of astrocytic Ca^{2+} signaling were recorded every second. For Ca^{2+} imaging experiments, the agonists were dissolved in aCSF containing fluorescein (1%) to visualize picospritzer-delivered solution. In experiments recording

GPCR-activated changes in K^+ or EPSPs, the agonists were delivered by perfusion for a period of 1 min unless otherwise noted. SBFI, tetraammonium salt was used to measure intracellular Na⁺. SBFI was dissolved to 20 μ M in intracellular solution and loaded into astrocytes through whole-cell patch clamp. The same parameters of two-photon imaging were used to image SBFI.

In vivo K⁺ recordings and Ca²⁺ imaging

Adult mice (8 to 12 weeks old) were anesthetized with ketamine (60 to 80 mg kg⁻¹) and xylazine (10 to 20 mg kg $^{-1}$). Body temperature was monitored by a rectal probe and maintained at 37°C by a heating blanket (BS4, Harvard Apparatus). The mice were incubated and artificially ventilated with a small animal ventilator (SAAR-830, CWE) (56). Blood was taken through a femoral artery catheter, and PCO₂ (partial pressure of CO₂), PO₂ (partial pressure of O₂), and pH were analyzed in microsamples (Rapidlab 248, Bayer, sample size 40 ml). Experiments were completed only if physiological variables remained within normal limits. The normal limits for PCO₂ were set at 35 to 45 mmHg, PO₂ at 80 to 115 mmHg, and arterial blood pH at 7.35 to 7.45; body temperature was maintained at 37°C. A stainless steel frame with one opening was glued to the skull. A cranial borehole (~1.0 mm) was prepared below the hole in the metal plate over the parietal cortex, and the dura was carefully removed. The exposed cortex was loaded with rhod-2 AM (0.4 mM, 1 hour) (56). The cranial window was superfused with aCSF gassed with 5% CO₂ and 95% O₂ (57). A potassium-sensitive microelectrode was inserted 100 µm (cortical layer II) below the pial surface microelectrode. After a stable baseline recording was obtained, ATP (100 µM), FMRF (15 μM), or vehicle (aCSF) was superfused for 1 min. K⁺-sensitive microelectrodes were calibrated before and after each recording. In selected experiments, the exposed cortex was loaded with rhod-2 AM (50 μM, 1 hour, 15 washes in aCSF). Time-lapse images of astrocytic Ca²⁺ signaling were recorded every second by FluoView with the custom-built two-photon laser-scanning setup.

Statistics

All histograms are expressed as means \pm SEM. Student's t test or analysis of variance (ANOVA) with Bonferroni-Dunn post hoc tests was used, where normality of the data was assumed by Shapiro-Wilk's test. Otherwise, nonparametric Kruskal-Wallis test with Mann-Whitney pairwise test with Bonferroni-adjusted α level was used. For repeated measures, paired t test or repeated-measures ANOVA with Bonferroni-Dunn post hoc test was used, where normality and sphericity conditions were met.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank C. Nicholson and S. Hrabetova for valuable comments on the manuscript and for sharing their technical expertise regarding fabrication of ion-selective sensitive microelectrodes. We thank J. Chen, K. McCarthy, and J. Rothstein for sharing genetically modified mice.

Funding: Supported by G. Harold and Leila Y. Mathers Charitable Foundation, U.S. NIH (NS075177 and NS078304 to M.N.), and F31 National Research Service Awards (F31NS073390 to N.A.S.).

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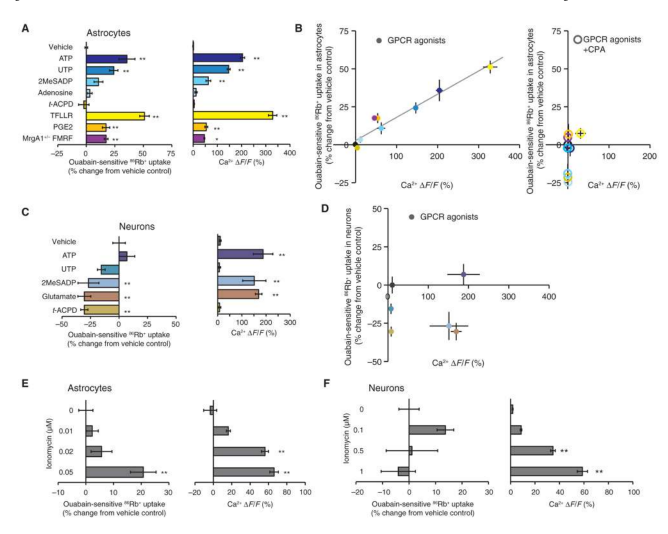
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Receptor-mediated Ca²⁺ signaling enhances astrocytic K⁺ uptake in vitro. (A) The effect of GPCR agonists ATP (100 μ M, n = 27), UTP (100 μ M, n = 39), 2MeSADP (100 μ M, n = 8), adenosine (100 μ M, n = 4), t-ACPD (100 μ M, n = 8), TFLLR-NH₂ (30 μ M, n = 37), PGE₂ $(50 \,\mu\text{M}, n = 10)$, or FMRF (15 μM in MrgA1^{+/-} mice, n = 12) on ouabain-sensitive $^{86}\text{Rb}^+$ uptake in cultured astrocytes compared with vehicle (left). **P< 0.01, Mann-Whitney Utest. Cytosolic Ca²⁺ increases in response to the same agonists (right). The cultured astrocytes were loaded with the Ca²⁺ indicator rhod-2 AM (4.5 μ M). n = 3 to 5. *P < 0.05; **P< 0.01, Bonferroni-Dunn test. (**B**) Relative increases in 86 Rb⁺ uptake plotted as a function of agonist-induced Ca²⁺ increases in astrocytes (y = 0.153x + 2.90, $t^2 = 0.94$). CPA $(20 \,\mu\text{M}, \text{ right})$ eliminated agonist-induced enhancement of $^{86}\text{Rb}^+$ uptake (n = 9). (C) The effect of ATP, UTP, 2MeSADP, glutamate, and t-ACPD (all 100 μ M, n = 4 to 8) on 86 Rb⁺ uptake in cultured neurons (left) with corresponding agonist-evoked increases in Ca²⁺ (right). Cultured neurons were loaded with the Ca²⁺ indicator rhod-2 AM (4.5 μ M). n = 3 to 5. **P< 0.01, Bonferroni-Dunn test. (**D**) Relative increases in 86 Rb⁺ uptake plotted as a function of agonist-induced Ca²⁺ increases in neurons. (E and F) Effect of the Ca²⁺ ionophore, ionomycin, on 86 Rb⁺ uptake and Ca²⁺ increases in astrocytes (E) (n = 4 to 12, **P < 0.01, Bonferroni-Dunn test) and neurons (F) (n = 4 to 8, **P < 0.01, Bonferroni-Dunn test).

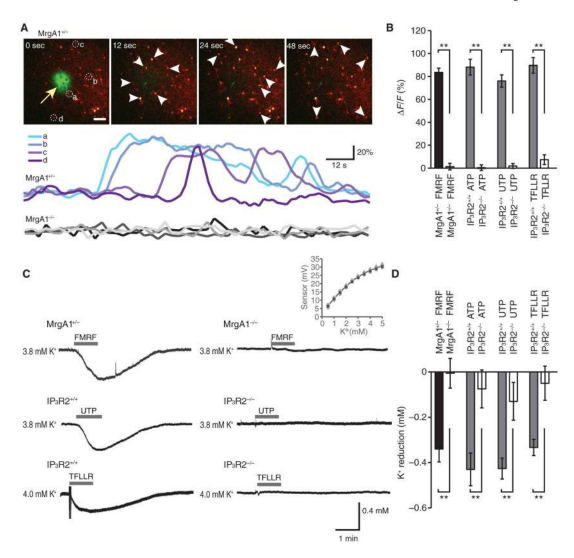


Fig. 2. Receptor-mediated astrocytic Ca²⁺ signaling enhances K⁺ uptake in hippocampal slices. (A) Time-lapse images of FMRF-induced increases in Ca²⁺ in a hippocampal slice prepared from a MrgA1 $^{+/-}$ mouse loaded with rhod-2 AM (4.5 μ M). The pipette solution contained, in addition to FMRF (15 µM), fluorescein isothiocyanate (FITC)-dextran (1%, 10 kD) to visualize delivery of agonist (arrow). Colored traces indicate relative changes in rhod-2 signals of astrocytes (a to d) labeled in the first panel, whereas gray traces at the bottom show the effect of FMRF in a MrgA1^{-/-}littermate. Scale bar, 50 µm. (**B**) Bar graph comparing relative increases in Ca^{2+} (rhod-2 signal, ΔFIF) in astrocytes in MrgA1^{+/-} and MrgA1^{-/-} mice (n = 3 to 6 slices) and the effects of agonist exposure, ATP (100 μ M), UTP (100 μ M), and TFLLR-NH₂ (30 μ M), in slices prepared from wild-type and IP₃R2^{-/-} mice. **P < 0.01, t test. (C) Recordings of extracellular K⁺ concentration in hippocampal slices exposed to FMRF, UTP, or TFLLR-NH2. The slices were prepared from wild-type (IP₃R2^{+/+}), IP₃R2^{-/-}, or MrgA1^{+/-} and MrgA1^{-/-} mice as indicated. Inset: average calibration of K⁺-sensitive microelectrodes. (**D**) Bar graph comparing relative changes in extracellular K⁺ in hippocampal slices exposed to ATP (n = 6 for both IP₃R2^{+/+} and $IP_3R2^{-/-}$), UTP (n = 6 for both $IP_3R2^{+/+}$ and $IP_3R2^{-/-}$), and TFLLR-NH₂ (n = 8 for $IP_3R2^{+/+}$ and n = 5 for $IP_3R2^{-/-}$) in slices prepared from $IP_3R2^{+/+}$ and $IP_3R2^{-/-}$ mice, as

well as exposure to FMRF in slices prepared from MrgA1^{+/-} (n = 6) and MrgA1^{-/-} mice (n = 5). **P < 0.01, t test.

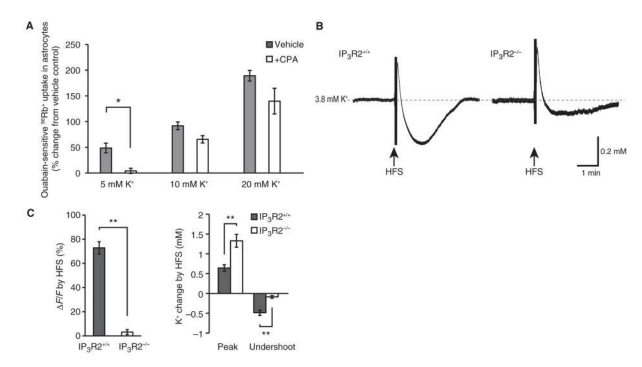


Fig. 3. Astrocytic calcium transients contribute to K⁺ clearance in response to moderate, but not pathophysiological, changes in extracellular K⁺ in cultured astrocytes and in hippocampal slices. (**A**) Bar graph illustrating the calcium dependence of ouabain-sensitive ⁸⁶Rb⁺ uptake in cultured astrocytes to physiological and pathophysiological changes in extracellular K⁺ concentration (n = 4 slices). *P < 0.05, t test. (**B**) HFS-induced changes in extracellular K⁺ in wild-type (IP₃R2^{+/+}) and IP₃R2^{-/-} mice. Dashed line indicates baseline K⁺ concentration. (**C**) Bar graphs comparing relative increase in Ca²⁺ (rhod-2 signal, n = 12 slices, ΔFIF , left panel) and changes in extracellular K⁺ (peak increase and undershoot) in IP₃R2^{+/+} and IP₃R2^{-/-} mice (right panel, n = 7 slices). **P < 0.01, t test.

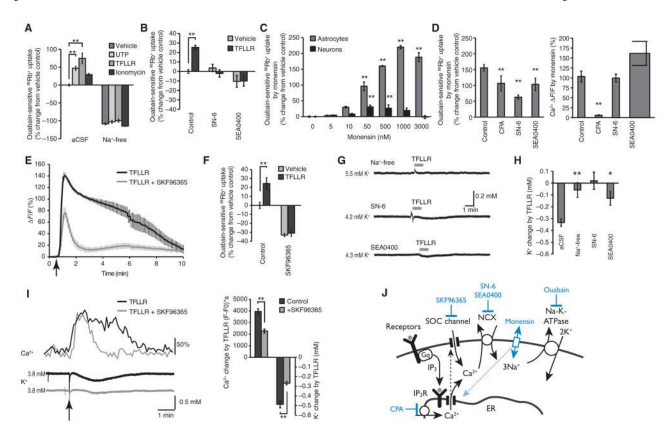


Fig. 4. Agonist-induced astrocytic Ca²⁺ signaling drives K⁺ uptake by increasing Na⁺ availability in cultured astrocytes and in hippocampal slices. (A) Effects of replacement of extracellular Na⁺ on 86 Rb⁺ uptake under baseline conditions and in response to UTP (100 μ M, n = 7), TFLLR-NH₂ (30 μ M, n = 7), or ionomycin (0.05 μ M) in cultured astrocytes. n = 3 to 7 wells. **P < 0.01, Mann-Whitney U test. (B) Effects of the Na⁺,Ca²⁺ exchange inhibitors, SN-6 (10 μ M) and SEA0400 (10 μ M) on baseline, and agonist (30 μ M TFLLR-NH₂)– induced $^{86}\text{Rb}^+$ uptake in cultured astrocytes. n = 3 to 6 wells. **P < 0.01, Mann-Whitney Utest. (C) Effects of increasing concentration of monensin, a Na⁺ ionophore, on ⁸⁶Rb⁺ uptake in cultured astrocytes and neurons. n = 4 to 8 wells. **P < 0.01 compared to before, Bonferroni-Dunn test. (**D**) Effects of CPA (20 μM), SN-6 (10 μM), and SEA0400 (10 μM) on monensin (3 μ M)-induced ⁸⁶Rb⁺ uptake in cultured astrocytes (n = 6 to 15 wells; **P <0.01 compared to before, Bonferroni-Dunn test) with corresponding agonist-evoked Ca²⁺ increases (right panel; n = 4 wells, **P < 0.01 compared to before, Bonferroni-Dunn test). (E and F) Store-operated calcium channel blocker SKF96365 (25 μM) reduced both the increase in cytosolic Ca²⁺ evoked by ATP (n = 4 wells) and the $^{86}\text{Rb}^+$ uptake by 30 μ M TFLLR-NH₂ (n = 4 wells) in cultured astrocytes. Arrow indicates the time ATP was applied to the cells. **P < 0.01, t test. (G) Effects of replacing extracellular Na⁺ with NMDG⁺, or addition of NCX blockers SN-6 (10 µM) and SEA0400 (15 µM) on TFLLR-NH2-induced (100 µM) changes in extracellular K⁺ concentration in hippocampal slices. (H) Comparison of TFLLR-NH₂-induced changes in extracellular K+ under Na+-free (n = 6) and control conditions in the presence of SN-6 (n = 5) or SEA0400 (n = 5) in hippocampal slices. *P< 0.05; **P< 0.01, Bonferroni-Dunn test. (I) Typical trace illustrating Ca²⁺ waves and K⁺ recordings activated by TFLLR-NH₂ (30 µM) in hippocampal slices with or without SKF96365 (25 μ M) with corresponding bar graphs compared on the right. n = 8 to 13, **P

< 0.01, t test. (**J**) Diagram of proposed pathways involved in agonist-induced K⁺ uptake in astrocytes.

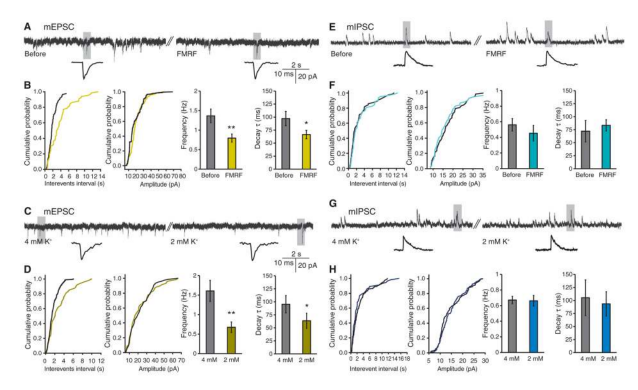


Fig. 5. Astrocytic Ca²⁺ signaling decreases mEPSC frequency but not mIPSC frequency in hippocampal slices. (A) Representative recordings of mEPSCs before and after superfusion of FMRF (15 μ M) in slices prepared from MrgA1^{+/-} mice. (**B**) Cumulative analysis of the inter-event interval and amplitude of mEPSCs in a 1-min representative recording before and during application of FMRF. Bar graphs compare frequency and time constant of decay (τ) . n = 9 slices. *P < 0.05; **P < 0.01 compared to before FMRF treatment, paired t test. (C) Representative recordings of mEPSCs when bath K⁺ concentration was decreased from 4 to 2 mM; shaded area is shown below at extended time scale. (D) Cumulative analyses of the inter-event interval and amplitude of mEPSCs. Bar graphs compare frequency and decay τ of mEPSCs. n = 8 slices. *P < 0.05; **P < 0.01 compared to 4 mM K⁺, paired t test. (E) Representative recordings of mIPSCs before and after perfusion of FMRF in slices prepared from MrgA1^{+/-} mice. (**F**) Cumulative analyses of the inter-event interval and amplitude of mIPSCs from representative traces. Comparison of the effect of FMRF on frequency and decay τ (n = 7 slices, paired ttest). (G) Representative recordings of mIPSCs when bath K⁺ concentration was decreased from 4 to 2 mM. (H) Cumulative analyses of the inter-event interval and amplitude of mIPSCs. Histograms compare frequency and decay τ of mIPSCs (n = 7 slices, paired t test).

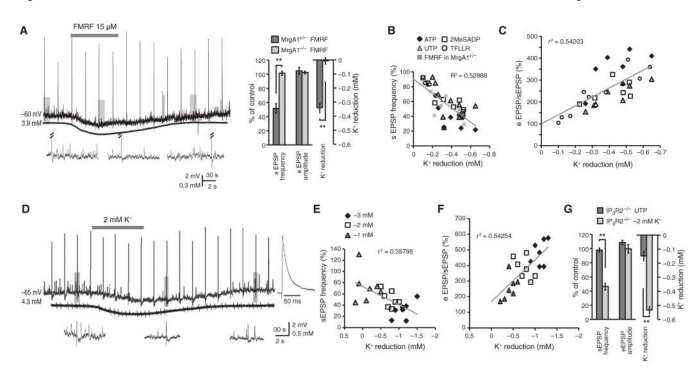


Fig. 6. Suppression of spontaneous excitatory activity in hippocampal slices in response to Ca²⁺ signaling in astrocytes. (A) Left: whole-cell recording of CA1 pyramidal neurons combined with measurements of extracellular K⁺ in a hippocampal slice. Representative recording of sEPSPs and eEPSPs in a CA1 hippocampal neuron in response to FMRF (15 µM) in a slice prepared from a MrgA1^{+/-} mouse. Enlarged time scales of EPSPs for the indicated periods are shown at the bottom. Right: bar graph comparing the effect of FMRF on the frequency of sEPSPs and amplitude of eEPSPs in CA1 neurons in slices prepared from MrgA1 $^{+/-}$ (n =6 slices) or MrgA1^{-/-} (n = 5 slices) mice. Concomitant changes in extracellular K⁺ are also shown. **P < 0.01, t test. (B) The frequency of sEPSPs plotted as a function of changes in extracellular K⁺ concentration in response to agonist exposure (ATP, 2MeSADP, UTP, TFLLR-NH₂, or FMRF) in MrgA1^{+/-} mice. (C) The ratio of integrated areas of eEPSPs and sEPSPs increased as a direct function of the decrease in extracellular K⁺ induced by agonist exposure ($r^2 = 0.54$). (**D**) Representative recording of sEPSPs together with eEPSPs from five CA1 hippocampal slices when K⁺ in the bath solution was transiently reduced from 4 to 2 mM. Enlarged time scales of the indicated periods are shown at the bottom. Trace on the right shows enlarged time scale for eEPSP recorded in 4 mM K⁺. (E) Relative changes in the frequency of sEPSP plotted as a function of the decrease in extracellular K⁺ induced by lowering bath K⁺ concentration. (F) The ratio of integrated area of eEPSPs and sEPSPs during the same time period increases as a direct function of lowering extracellular K^+ ($r^2 =$ 0.54). (G) Effects of UTP or reduction of extracellular K⁺ (from 4 to 2 mM) on the frequency of sEPSPs and amplitude of eEPSPs in CA1 hippocampal neurons in slices prepared from IP_3R_2 —mice (n = 5 slices). Concomitant changes in extracellular K^+ are also plotted. **P < 0.01, t test.

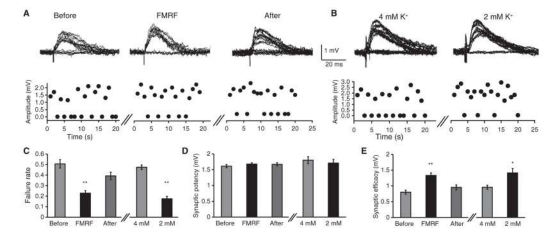


Fig. 7. Astrocytic Ca²⁺ signaling increases synaptic efficacy in hippocampal slices. (**A**) Failure rate from a representative recording in response to minimal stimulation before, during, or after application of FMRF in MrgA1^{+/-} mice. Traces show eEPSP amplitude. (**B**) Failure rate to minimal stimulation in 4 or 2 mM K⁺ perfusate. (**C**) Comparison of the effects of FMRF (n = 7 slices, **P < 0.01 compared to before FMRF treatment, Bonferroni-Dunn test) and extracellular K⁺ (n = 8 slices, **P < 0.01 compared to 4 mM K⁺, paired t test) on failure rate. (**D** and **E**) Effects of FMRF and K⁺ on (D) synaptic potency (mean amplitude of EPSP successes) and (E) synaptic efficacy (mean amplitude of all trials). n = 7 slices, repeated-measures ANOVA for FMRF; n = 8 slices, paired t test for K⁺, **P < 0.01, *P < 0.05).

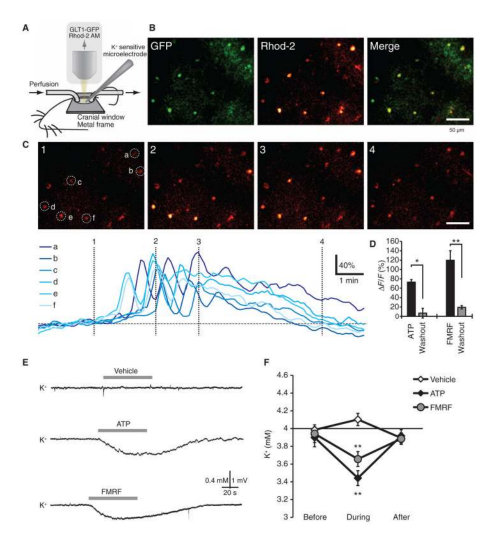


Fig. 8. Agonist-induced increases in astrocytic Ca²⁺ reduce extracellular K⁺ concentration in vivo. (**A**) Experimental setup for in vivo recordings. (**B**) Glt1-eGFP reporter mouse loaded with the Ca²⁺ indicator rhod-2 AM. Rhod-2 only–labeled eGFP⁺ astrocytes. (**C**) Time-lapse images of ATP (100 μM)–evoked increases in Ca²⁺ in astrocytes located 95 μm below the pial surface. Colored traces indicate relative changes in rhod-2 signals of astrocytes (a to f) labeled in the first panel. (**D**) Comparison of ATP (100 μM)– and FMRF (15 μM)–induced increases in rhod-2 signal (ΔPF). n = 4 to 5 mice. *P < 0.05; **P < 0.01, paired t test. (**E**) Representative recordings of extracellular K⁺ concentration in mouse cortex exposed to vehicle (aCSF), ATP (100 μM), or FMRF (15 μM). (**F**) Graph comparing the effects of vehicle, ATP, and FMRF on extracellular K⁺. All recordings were obtained 100 μm below the cortical surface. n = 7 mice. **P < 0.01, Bonferroni-Dunn test compared to baseline.