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Protease activated receptor 1-induced glutamate release in cultured astrocytes is mediated by Bestrophin-1 channel but not by vesicular exocytosis

Soo-Jin Oh¹, Kyung-Seok Han^{1,2}, Hyungju Park¹, Dong ho Woo^{1,2}, Hye Yun Kim¹, Stephen F Traynelis³ and C Justin Lee^{1,2*}

Abstract

Background: Glutamate is the major transmitter that mediates the principal form of excitatory synaptic transmission in the brain. It has been well established that glutamate is released via Ca²⁺-dependent exocytosis of glutamate-containing vesicles in neurons. However, whether astrocytes exocytose to release glutamate under physiological condition is still unclear.

Findings: We report a novel form of glutamate release in astrocytes via the recently characterized Ca²⁺-activated anion channel, Bestrophin-1 (Best1) by Ca²⁺ dependent mechanism through the channel pore. We demonstrate that upon activation of protease activated receptor 1 (PAR1), an increase in intracellular Ca²⁺ concentration leads to an opening of Best1 channels and subsequent release of glutamate in cultured astrocytes.

Conclusions: These results provide strong molecular evidence for potential astrocyte-neuron interaction via Best1-mediated glutamate release.

Keywords: Astrocyte, Bestrophin-1, Glutamate

Background

Astrocytes and neurons are intimate partners in the brain. Astrocytes respond to neuronal activity by an increase in intracellular Ca^{2+} ($[Ca^{2+}]_i$) [1,2]. The synaptically released neurotransmitters such as glutamate and ATP activate corresponding $G_{\alpha q}$ protein-coupled receptors expressed in astrocytes, resulting in increase in $[Ca^{2+}]_i$ and Ca^{2+} -dependent release of gliotransmitters from astrocytes [2,3]. These gliotransmitters in turn influence the neuronal excitability and synaptic activities [3-8].

At the heart of this neuron-glia interaction there is glutamate, which is released from neurons by synaptic vesicles to mediate neuron-to-glia communication and from astrocytes by various Ca^{2+} -dependent mechanisms

¹Center for Neural Science and Center for Functional Connectomics, Korea Institute of Science and Technology (KIST), Seoul, South Korea ²Neuroscience Program, University of Science and Technology (UST),

Daejeon, South Korea

to mediate glia-to-neuron communication [9,10]. Therefore, understanding how astrocytes release glutamate in Ca²⁺-dependent manner and how astrocytically-released glutamate regulates neuronal synaptic activity has been a major challenge. One line of accumulating evidence supports that astrocytes release glutamate by Ca^{2+} and SNARE-dependent exocytosis mechanism [11], although this mechanism has been recently challenged [12]. It has been reported that glutamate released by SNAREdependent mechanism regulates hippocampal synapse between perforant path afferents and granule cells [13] or Schaffer Collateral afferent and CA1 pyramidal neurons [6] through N-methyl-D-aspartic acid receptor (NMDAR) activation. Yet, astrocytes also express a combination of anion-permeable channels to move Cl⁻ and large anions such as glutamate across membranes in response to specific physiological stimulations [14,15], raising the possibility that astrocytes have a multiple glutamate release mechanisms. Although astrocytes under



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^{*} Correspondence: cjl@kist.re.kr

Full list of author information is available at the end of the article

physiological conditions appear to possess a Ca^{2+} and anion channel-dependent glutamate release mechanism [10,16], until now this idea has not been directly demonstrated due to the lack of molecular evidence.

Bestrophin is the gene identified as the gene responsible for Best's vitelliform macular dystrophy and has been shown to encode a functional Ca²⁺-activated anion channel (CAAC) in nonneuronal tissue and peripheral neurons [17]. This Bestrophin-1 channel (Best1) is directly activated by submicromolar intracellular Ca²⁺ concentration and has an anion selective pore with single channel activities [17-24]. Recently, we have discovered that astrocytes express CAAC and that Best1 encodes most of the CAAC in astrocytes [25]. This astrocytic CAAC showed a considerable permeability to large anions such as isethionate and glutamate [25]. In addition, we have recently demonstrated that Best1 channel mediates tonic GABA release from cerebellar glia by a direct permeation [26]. Therefore, we hypothesized that Best1 is the molecular identity of Ca²⁺-dependent anion channel that mediates glutamate release from astrocytes. Here we tested whether Best1 is an alternative Ca²⁺-dependent glutamate release mechanism in cultured astrocytes. To stimulate astrocytes more physiologically, we activated endogenous GPCRs by application of TFLLR [27] a selective peptide agonist of the protease activated receptor 1 (PAR1). We demonstrate that PAR1-induced glutamate release is mediated not by conventional vesicular exocytosis but by an activation of glutamate permeable anion channel Best1 in cultured astrocytes.

Results

PAR1-induced glutamate release is mediated by CAAC

It has been suggested that hippocampal astrocytes utilize an anion channel-dependent mechanism for Ca^{2+} -induced glutamate release [10,16]. To confirm the anion channelmediated glutamate release from astrocytes, we performed an *in vitro* glutamate release assay using radioactive isotope-labeled glutamate (³H-glutamate) in cultured hippocampal astrocytes (Figure 1A) [7]. To elicit intracellular Ca^{2+} increase and the subsequent release of glutamate in astrocytes, we used TFLLR to activate PAR1. We found that an application of TFLLR induced a significant glutamate release from cultured astrocytes (Figure 1B), which was inhibited by a pre-incubation of anion channel blockers, such as niflumic acid and flufenamic acid (Figure 1B) [28]. These results suggested that astrocytes utilize anion channel-dependent glutamate release mechanism.

Best1 mediates a Ca²⁺-dependent glutamate release in cultured astrocytes

We have recently demonstrated that Best1 channel is activated by intracellular Ca^{2+} (EC₅₀ = 150 nM) with considerable permeability to GABA [26]. To test whether Ca^{2+} induced glutamate release is also mediated by Best1 in native astrocytes, we adopted a gene silencing tool using the short hairpin RNA (shRNA) specifically targeted to mouse Best1 transcript [25,26]. We recorded whole cell currents in cultured astrocytes under gramicidin-D perforated patch configuration (Figure 2B). We found the significant Ca^{2+} -induced conductance carried by the efflux





indicates amino acids position of best 1 pore mutant (w95C), ked square indicates target amino acids for best 1 shrive. Right; indicated and amino acids for best 1 shrive. Right; indicated and amino acids sequences of Best 1 shrive. Right; indicated and amino acids sequences of Best 1 shrive. Right; indicated and amino acids is pore mutant (w95C), ked square differences from wild type Best 1 (Best 1 WT) and shrNA. Right; indicated and (Best 1 (sh insens)) were highlightened. **B**) Schematic diagram for perforated-patch-current recording from cultured astrocytes. **C**) Representative perforated-patch-current recording from Best 1-shrNA and Best 1 (sh insens) or Best 1-shrNA and shrNA insensitive pore mutant Best 1 (Best 1-W93C (sh insens)) expressing cultured astrocytes. The current responses were recorded in response to a voltage ramp command (from -100 to +100 mV, 1 s duration, 0.2 Hz; V_h of -70 mV) before and after 30 µM TFLLR treatment. **D**) The bar graph summarizes the mean current amplitude at V_h = -70 mV ± s.e.m (*p < 0.05, **p < 0.01, unpaired t-test). Among 13 recorded astrocytes which expressed Best 1-shrNA and Best 1 (sh insens), only 5 cells showed comparable currents. And a big current recorded in an astrocyte expressing Best 1-shrNA and Best 1-W93C (sh insens) was eliminated in the bar graph. Numbers of determinations are indicated on the bar graph.

of glutamate at -70 mV in astrocyte with scrambledshRNA. Silencing the Best1 gene via specific shRNA for Best1 significantly eliminated PAR1 induced whole-cell current in cultured astrocytes. This effect was fully rescued by the cotransfection of Best1-shRNA, along with a shRNA-insensitive form of Best1 (Figure 2C and D). However, the addition of a pore mutation at position 93, from tryptophan to cystein (Best1-W93C sh insens) [29] failed to rescue the current (Figure 2C and D). These results indicate that pore of Best1 channel is responsible for the permeation of anions such as Cl⁻ and glutamate. We further tested the effect of Best1-shRNA on glutamate release using conventional high-performance liquid chromatography (HPLC) from the solution samples collected after 5 min application of TFLLR in cultured astrocytes. In this experiment, we firstly prepared astrocytes with the same cell number of cells that were transfected with Sc-shRNA or Best1-shRNA. Since we found that Best1-shRNA did

not affect astrocytic cell growth or survival (data not shown), we assumed that total number of astrocytes in each sample after transfection of Sc-shRNA or Best1-shRNA should be similar. We found a decrease in glutamate release by Best1-shRNA using HPLC method (Figure 3). These results indicate that the Best1 channel is the molecular identity of glutamate permeable CAAC and suggest that the Best1 channel might be the molecular machinery for Ca²⁺-dependent glutamate release from astrocytes.

To independently confirm the glutamate release by HPLC method, we tested the effect of Best1-shRNA on glutamate release using fluorescence resonance energy transfer (FRET) based glutamate sensor. The released glutamate was monitored using a glutamate-sensing fluorescent reporter, GluSnFR, which senses glutamate by changing the ratio of FRET from CFP to YFP upon glutamate binding [30] (Figure 4A). The concentration-response curve for glutamate on the sensor was within the range of estimated glutamate release from a single astrocyte in culture (~10⁻⁶ M) [7] (Figure 4C and 4D). The calculated EC_{50} of glutamate was ~25.57 μ M (Figure 4D). We observed that a brief pressure application of TFLLR caused a long lasting glutamate release as indicated by the relative change in FRET ratio (Figure 4B and 4E). The amount of glutamate release was on average significantly lower in the astrocytes expressing Best1-shRNA (Figure 4E, red symbols) than those expressing control scrambled shRNA (Figure 4E, black symbols). We found a similar degree of decrease in glutamate release by Best1-shRNA using the FRET based glutamate imaging, as compared to HPLC method (Figures 3C and 4F). Co-application of TFLLR and niflumic acid or treating the cells with BAPTA-AM, significantly reduced the glutamate release from astrocytes (Figure 4E and 4F).

PAR1 induced glutamate release is not due to vesicular release

We noted that the amount of released glutamate was not completely inhibited by gene-silencing of Best1 (Figure 4E and 4F), raising a possibility that astrocytes have other Best1-independent mechanisms for glutamate release such as vesicular exocytosis. It is possible that Best1-shRNA affects the expression level of other genes involved in vesicular release. To address this possibility, we performed semi-quantitative RT-PCR analysis of cultured astrocytes with Best1-shRNA. Best1-shRNA showed a knock down efficiency for Best1 of over 90% while not affecting the expression level of other known genes such as Syt4 (Synaptotagmin4), Munc18-1, Vamp2 (Vesicle-associated membrane protein 2) involved in vesicular release (Figure 5A). We also tested whether inhibiting vesicular release machinery affects the glutamate release by using FRET based glutamate imaging. We found that the pretreatment with Conconomycin A (preventing vesicular glutamate release by inhibiting vesicular hydrogen ATPase, [31]) and Tetanus toxin (interfering all vesicular releases by inhibiting fusion of vesicles, [32]) slightly but not significantly reduced TFLLR-induced glutamate release from cultured astrocytes (Figure 5B and 5C). Consistent with these results, hyperosmotic challenge, which is known to cause a Ca^{2+} independent exocytosis of glutamate containing releasable vesicles, did not show any significant increase in glutamate release (Figure 5B and 5C). Taken together, PAR1-induced glutamate release is not mediated by vesicular exocytosis in cultured astrocytes.

Discussion

In this study we report the direct evidence of anion channel-mediated glutamate release mechanism. We provide a series of evidence for Ca^{2+} -activated, Best1-mediated glutamate release from astrocytes. For example, the selective gene silencing of astrocytic Best1 channels significantly reduced GPCR-induced and Ca^{2+} -dependent glutamate release from astrocytes as measured by HPLC detection of glutamate from cultured astrocytes (Figure 3), and by FRET glutamate sensor in cultured astrocytes (Figure 4). The FRET based glutamate sensor has been wellcharacterized in the previous report [30]. In that report it has been demonstrated that FRET change is quite specific



for glutamate because the sensor shows binding kinetics and high selectivity toward glutamate with no apparent change in ratio with APV, NBQX, NMDA, KA, AMPA, etc. [30]. We performed a calibration experiment by measuring glutamate EC_{50} in our imaging system and experimental conditions. Our calibration results showed that the FRET-based glutamate sensor in cultured astrocyte can be useful in detecting micromolar and submicromolar concentrations of released extracellular glutamate from a single astrocyte. Until now, it has been proposed that glutamate could be released from astrocytes through multiple routes, including Ca^{2+} and SNARE-dependent vesicular exocytosis, the reversal action of glutamate transporters, transportation by cystine-glutamate antiporter, and permeation through channels or receptors, such as P2X7 receptor, volume regulated anion channel (VRAC), or gap junction hemichannel [33]. Among these, the molecular mechanism of Ca^{2+} -dependent glutamate release has been extensively studied mainly because the







generation of astrocytic Ca²⁺ transient responding to neuronal activity is one of the most important physiological readout of neural activity [3,5,7,10]. Numerous studies have proposed the SNARE-dependent exocytosis of glutamate as a potential route for astrocytic Ca²⁺dependent glutamate release. These studies demonstrated that astrocytic glutamate release was sensitive to exocytosis blockers or the expression of the essential machineries for regulating Ca²⁺-dependent exocytosis [33]. In addition to the vesicular mechanism, there is multiple lines of evidence suggesting that astrocytes express a Ca²⁺-dependent but non-vesicular glutamate release machinery [16]. Despite its unclear molecular identity, it has been reported that an increase in Ca^{2+} by GPCR activation induces glutamate release from astrocytes through osmolyte-permeable anion channels [10,28], which are activated by intracellular Ca^{2+} or increase in cell volume [16]. In our previous reports, we observed that Best1-mediated Ca2+-activated anion current could be induced by a host of GPCR agonists such as mGluR1/5, P2Y, B2, S1P, and LPAR in astrocytes [25], and that activation of those receptors elicits Ca^{2+} dependent glutamate release [7]. Finally, Best1 was demonstrated to be permeable to GABA and mediate tonic GABA release in cerebellar glial cells [26]. Recent

study showed that GABA is abundant in cerebellar glial cells to be released tonically via Best1 to cause tonic inhibition [34]. However, in hippocampus, GABA is not present in glial cells; rather, Best1 might release glutamate near synapse. Therefore, it is plausible that Best1, at least in part, could be the downstream target of various G_{q} -coupled GPCRs that signal through intracellular Ca²⁺ to release glutamate in hippocampal astrocytes.

In addition to Best1 channel, other anion channels might participate in anion channel-mediated glutamate release as previously suggested, such as Ca²⁺-induced but volume regulated osmolyte-permeable channels [16]. In fact, there are several studies showing that bestrophin channel could be opened by both cytosolic Ca²⁺ and cell volume increase [17,35], raising a possibility that Best1-mediated glutamate release is also triggered by Ca²⁺-dependent cell swelling and subsequent activation of VRAC. However, in our previous study we demonstrated that PAR-1 activation does not induce significant swelling [36] indicating that Ca²⁺ directly initiates Best1 channel-mediated glutamate release from astrocytes. The PAR1-induced astrocytic glutamate release was not completely inhibited by Best1 knock-down (Figure 4). Still, vesicular release mechanism was not involved because Best1 silencing did not affect the expression level

of known genes involved in vesicular release (Figure 5A) and glutamate release from astrocytes was not reduced by treatment with Conconomycin A or Tetanus toxin (Figure 5B). There is a possibility that other ion channels that are independent of Best1 channel might participate in PAR1-induced glutamate release from astrocytes.

We utilized the native GPCR, PAR1, which has been extensively used in numerous studies to selectively activate astrocytes. Even though PAR1 is expressed in a subset of dentate granule cells [37], PAR1 has been shown to be expressed exclusively in astrocytes in human and rodent CA1 hippocampus [7,38,39], as well as in the nucleus of solitary tract [40] to mediate neuron-glia interaction. Therefore, the source of PAR1-induced glutamate is most likely astrocyte, as a direct consequence of increase $[Ca^{2+}]_i$. Although there is no direct evidence of how PAR1 is activated in the physiological situation up to now, the recent study demonstrated that tPA-plasmin pathway is an endogenous PAR1 agonist [41], suggesting that physiological PAR1 activation is initiated by the activation of tPA-plasmin pathway in physiological condition such as synaptic plasticity [42,43].

In summary, we reveal a novel anion channel-mediated glutamate release mechanism in cultured astrocytes. The ideas and tools developed in this study should prove to be helpful in understanding the physiological role of glutamate release mechanism and its functional significances.

Methods

Primary astrocyte culture

Cell culture of mouse astrocytes was performed as previously described [7]. The cerebral cortex from P0 ~ P3 postnatal mice was dissected free of adherent meninges, minced and dissociated into single cell suspension by trituration. Dissociated cells were plated onto 12 mm glass coverslips coated with 0.1 mg/ml poly D-lysine. Cells were grown in DMEM supplemented with 25 mM glucose, 10% heat-inactivated horse serum, 10% heat-inactivated fetal bovine serum, 2 mM glutamine, and 1000 units ml-1 penicillin-streptomycin. Cultures were maintained at 37°C in humidified 5% CO₂-containing atmosphere.

Glutamate FRET imaging

In FRET-based glutamate imaging experiments, pDisplay-GluSnFR vector [30] was electroporated alone or with pSicoR-scrambled-shRNA or pSicoR-Best1-shRNA vector into cultured astrocytes (MicroPorator; Digital Bio, Korea). After 48 ~ 72 hour expression, FRET imaging was performed under a microscope (BX50WI; Olympus) equipped with xenon lamp with 436/20 excitation filter (D436/20x filter; Chroma). The emission beam was split with a DualView (Optical Insights) with a CFP/YFP filter set (OI-05-EX), recorded by EM-CCD camera (ANDOR IXON). Imaging Workbench software (INDEC BioSystems) was used for image acquisition and offline image analysis. TFLLR puff (TFLLR-NH₂; Peptron, Korea; 500 μ M) was made by using picospritzer-assisted positive pressure (~100 ms). The amount of released extracellular glutamate was described as ratio between the emission intensity of CFP and YFP (CFP/ YFP), which was divided by baseline CFP/YFP ratio (relative CFP/YFP ratio).

Radioactivity glutamate release assay

In vitro glutamate assay using ³H-labelled glutamate was performed as described previously [7]. Astrocytes were loaded with 0.5 µm l-[³H]glutamate for 60 min by adding 1 µm of 1 mCi ml⁻¹ l-[³H]glutamate stock solution to 2 ml of culture medium. The cultures were preincubated for 30 min with 1 mm amino-oxyacetic acid and 0.5 mm methionine sulfoximine before adding $[^{3}H]$ glutamate, and during the loading to inhibit the metabolism of glutamate to glutamine and other metabolites [44]. Cells were washed with external solution 3 times. In some experiments, the external solution was supplemented with 50 µm l-transpyrrolidine-2,4-dicarboxylic acid (trans-PDC) to block glutamate transporter, a maximally effective concentration ($6 \times IC_{50}$ of $4-8 \mu m$; [45]) that is well below that suggested to stimulate heteroexchange (0.2 mm; [46,47]). Agonists were added to external solution for 6 min and the experiment was terminated by collection of the solution. Each experimental run included the control condition in which no agonist was added. Six replicates were obtained for each drug condition. For analysis, the average radioactivity count was obtained from six replicates for each condition and compared to the average of control.

HPLC analysis for glutamate release

The Cortical primary astrocytes were cultured in 60 mm dishes for HPLC analysis. TFLLR (30 µM) was used to induce glutamate. Prior to TFLLR treatment, astrocytes were washed with PBS three times. For gene silencing experiment, Lentivirus containing Scrambled or Best1-shRNA was treated to cells and incubated for 48 hrs. The amino acid content was derivatized with o-phthaldialdehyde (OPA) and detected using UV (DAD) detection [48]. The OPA derivatized samples by programmed autosampler were injected on Zorbax Eclipse Plus C₁₈ column with detection at 338 nm (a reference = 390 nm). Mobile phase A was 40 mM Na_2HPO_4 pH 7.8, and B was acetonitrile/methanol/water (45:45:10, v/v/v). The flow rate was 2 mL/min with a gradient condition that allowed for 1.9 min at 0% B and raised to 26% B at a 12.5 min-step. Then washing at 100% B and equilibration at 0% B was performed in a total retention time of 15 min. Reagents of OPA derivatization and all equipments for HPLC analysis were obtained from Agilent Technologies.

Lentivirus with Best1-shRNA and Best1 mutagenesis

Best1-shRNA and lentivirus production were performed as previously described ([49], Virus facility, KIST). Best1 shRNA-insensitive pore mutant (Best1-shRNA insens W93C) was generated by using PCR-based site-directed mutagenesis kit (Stratagene, Cedar Creek, TX, USA).

Electrophysiology

The extracellular recording solution for perforated patch clamp recording was comprised of (in mM) 150 NaCl, 10 HEPES, 3 KCl, 2 CaCl₂, 2 MgCl₂, 5.5 glucose, at pH 7.3. The internal solution contained 25 μ g/ml gramicidin D and (in mM) 75 Cs₂SO₄, 10 NaCl, 0.1 CaCl₂, and 10 HEPES, at pH 7.1. Pipette resistances ranged from 5 to 8 MΩ. It took 20 to 30 min to achieve acceptable perforation, with final series resistances ranging from 15 to 40 MΩ. Current voltage curves were established by applying 100-, 200-, or 1000 ms duration voltage ramps from -100 to +100 mV. Data were acquired by an Axopatch 200A amplifier controlled by Clampex 9.2 via Digidata 1322A data acquisition system (Axon Instruments, Union City, CA, USA). Experiments were conducted at room temperature (20 ~ 24°C).

Reverse transcription-PCR

Total RNA was prepared from cultured astrocyte from postnatal day 0–3 mice using Trizol reagent (Invitrogen). cDNA was synthesized using Super Script III reverse transcriptase (Invitrogen). The reverse transcription (RT)-PCR primers used to check expression of Best1, Syt4 (Synaptotagmin4), Munc18-1, Vamp2 (Vesicle-associated membrane protein 2) GAPDH and glial fibrillary acidic protein (GFAP) were as followings

- ; Best1 forward, 5'-AGGACGATGATGATTTTGAG- 3'
- ; Best1 reverse, 5'-CTTTCTGGTTTTTCTGGTTG- 3'
- ; Syt4 forward, 5'-AGGCCAATTCCCCTGAGAGC- 3'
- ; Syt4 reverse, 5'-ACCGCTTCCTTCTGCTGTGG- 3'
- ; Munc18-1 forward, 5'-

ACTCCGCTGACTCTTTCCAA- 3'

- ; Munc18-1 reverse, 5'-
- GTCGGCTTTATAGGCATCCA- 3'
- ; Vamp2 forward, 5'-GTCACTGCCTCTGCCAAGTC- 3'
- ; Vamp2 reverse, 5'-GGCAGACTCCTCAGGGATTT- 3'
- ; GAPDH forward, 5'-
- AGCTGAACGGGAAGCTCACT- 3'
- ; GAPDH reverse, 5'-TGCTGTAGCCAAATTCGTTG- 3'
- ; GFAP forward, 5'-AGAACAACCTGGCTGCGTAT-3'
- ; GFAP reverse, 5'-CGGCGATAGTCGTTAGCTTC-3'

Primers for Syt4, Munc18-1 and Vamp2 were designed as previous reports [50,51]. All reactions were performed in a programmable thermocycler (Eppendorf).

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

SJO performed patch clamp recording and wrote the manuscript. HP and KSH carried out FRET imaging experiments. DW and HYK performed HPLC analysis. SFT designed the radioactivity glutamate release assay experiment. CJL carried out radioactivity glutamate release assay and wrote the manuscript, and supervised entire project. All authors read and approved the final manuscript.

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Author details

¹Center for Neural Science and Center for Functional Connectomics, Korea Institute of Science and Technology (KIST), Seoul, South Korea. ²Neuroscience Program, University of Science and Technology (UST), Daejeon, South Korea. ³Department of Pharmacology, School of Medicine, Emory University, Atlanta, GA, USA.

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