Electrophysiological Evidence for Functional Astrocytic P2X$_3$ Receptors in the Mouse Trigeminal Caudal Nucleus

Jaekwang Lee$^{1,2}$, Jin Young Bae$^3$, C. Justin Lee$^{2,*}$ and Yong Chul Bae$^3*$

$^1$Division of Functional Food Research, Korea Food Research Institute, Jeonju 55365,
$^2$Center for Neuroscience and Functional Connectomics, Korea Institute of Science and Technology, Seoul 02792,
$^3$Department of Anatomy and Neurobiology, School of Dentistry, Kyungpook National University, Daegu 41940, Korea

Recently, we reported that astrocytes in the trigeminal caudal nucleus (Vc) of the brain stem express a purinergic receptor P2X$_3$, which is involved in the craniofacial pathologic pain. Although we observed protein expression of P2X$_3$ receptors (P2X$_3$Rs) in the astrocyte of the Vc, it is still unclear that astrocyte has functional P2X$_3$Rs in Vc. To address this issue, we recorded astrocytic P2X$_3$Rs by using whole cell voltage-clamp recording in the Vc of the GFAP-GFP mice, which was used as a guide to astrocytes with green fluorescence. While measuring voltage ramp-induced astrocytic membrane current, we found the amplitude of the current was increased when we applied P2-purinoreceptor agonist, α,β-meATP. This increase was blocked by co-application of A317491, P2X$_3$R antagonist. These results demonstrate that astrocytes in the Vc express functional P2X$_3$Rs, which might be critical in craniofacial pathologic pain.

Key words: Electrophysiology, Trigeminal Caudal Nucleus, Astrocytes, Purinergic P2X$_3$, Pain

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*To whom correspondence should be addressed.
C. Justin Lee, TEL: 82-2-958-6940, FAX: 82-2-958-6937
e-mail: cj@kist.re.kr
Yong Chul Bae, TEL: 82-53-660-6860, FAX: 82-53-425-6025
e-mail: ycbae@knu.ac.kr

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neuropathic pain, there is a lack of direct evidence for functional P2X3R in the Vc.

To address this issue, we investigated the activity of P2X3R in astrocytes in the Vc using whole cell voltage-clamp recordings while applying voltage ramp protocol. In this study, we report, for the first time, a functional expression of P2X3R in the astrocyte of the Vc, which may be involved in the regulation of neuronal activity in pathologic pain condition.

MATERIALS AND METHODS

Animals and tissue preparation

Three male Sprague-Dawley rats, weighing 290–310 g, were used for this study. All animal procedures were reviewed and approved by the Kyungpook National University Intramural Animal Care and Use Committee. The rats were deeply anesthetized with sodium pentobarbital (80 mg/kg, i.p.) and perfused intracardially with 100 ml of heparinized normal saline (0.9% NaCl solution), followed by 500 ml of freshly prepared fixative, a mixture of 4% paraformaldehyde and 0.01% glutaraldehyde in 0.1 M phosphate buffer (PB), pH 7.4 (PB). Brainstem including trigeminal caudal nucleus (Vc) was removed, postfixed in the same fixative for 2 hours at 4°C, and rinsed in PB. Sections were transversely cut on a Vibratome at 60 μm and cryoprotected in 30% sucrose in PB overnight at 4°C.

Electron microscopic preembedding immunohistochemistry

For double immunostaining for P2X3 and GFAP, sections of Vc were frozen on dry ice for 20 minutes, thawed in phosphate-buffered saline (PBS; 0.01 M, pH 7.4) to enhance penetration, and pretreated with 1% sodium borohydrate for 30 minutes to remove glutaraldehyde. Sections were then blocked with 3% hydrogen peroxide for 10 minutes, to suppress endogenous peroxidase, and with 10% normal donkey serum (NDS, Jackson ImmunoResearch, West Grove, PA) for 30 minutes, to quench secondary antibody binding sites. Sections were incubated overnight in a mixture of rabbit anti-P2X3 (1:200; Alomone Labs Ltd., Cat. no. APR-016) antibody and mouse GFAP (1:5000, Chemicon, Cat. no. MAB360) antibody in PBS. After rinsing in PBS, sections were incubated with a mixture of biotinylated donkey anti-mouse (1:200, Jackson ImmunoResearch) and 1 nm gold-conjugated donkey anti-rabbit (1:50, EMS, Hatfield, PA) antibodies for 2 hours. The sections were postfixed with 1% glutaraldehyde in PB for 10 minutes, rinsed in PB several times, incubated for 4 minutes with HQ silver enhancement solution (Nanoprobes, Yaphank, NY), and rinsed in 0.1 M sodium acetate and PB. After rinsing, sections were incubated with ExtrAvidin peroxidase (1:5000, Sigma-Aldrich) for 1 hour and the immunoperoxidase was visualized with nickel-intensified 3,3’-diaminobenzidine tetrahydrochloride (DAB). Then, sections were further rinsed in PB, osmicated in 1% osmium tetroxide (in PB) for 1 hour. Sections were further dehydrated in graded alcohols, flat-embedded in Durcupan ACM (Fluka, Buchs, Switzerland) between strips of Aclar plastic film (EMS, Hatfield, PA), and cured for 48 hours at 59°C. Small pieces containing immunostaining for P2X3, and GFAP in Vc were cut out of wafers and glued onto blank resin blocks with cyanoacrylate. Thin sections were cut with a diamond knife, mounted on formvar-coated single slot nickel grids, and stained with uranyl acetate and lead citrate. Grids were examined on a Hitachi H 7500 electron microscope (Hitachi, Tokyo, Japan) at 80 kV accelerating voltage. Images were captured with Digital Micrograph software driving a cooled CCD camera (SC1000; Gatan, Pleasanton, CA) attached to the electron microscope, and saved as TIFF files.

For the quantitative analysis of the P2X3 expression in the astrocytic soma and process, 60 electron micrographs (at 25,000 original magnification) were taken in each section of the Vc from each of three rats. Gold particle density (number of gold particles/µm²) for P2X3 in the GFAP⁺ astrocytic soma and process in each rat was measured using a digitizing tablet and Image J software (v.1.45; NIH, Bethesda, MD). Statistical analysis of differences between astrocytic soma and process was performed with Student’s t-test.

Slice preparation and electrophysiology

Coronal mouse brain slices (200–300 μm) containing trigeminal caudal nucleus region were acutely prepared from adult GFAP-GFP mice (age 4–8 weeks). Following decapitation the brain was rapidly removed and placed in cold artificial cerebrospinal fluid (ACSF) having the following composition (in mM): 130 NaCl, 24 NaHCO3, 3.5 KCl, 1.25 NaH 2PO4, 1 CaCl2, 3 MgCl2 and 10 glucose, pH 7.4; room temperature with oxygenation (95% O2 and 5% CO2). The slices were made using an oscillating tissue slicer (Linear Slicer) at 4°C and stored in room temperature with oxygenation (95% O2, 5% CO2) and prepared slices were left to recover for at least 1 hour before recording. Each slice that was studied was transferred from a recovery/holding reservoir to the recording chamber of a fixed-stage upright microscope (Zeiss Axio Examiner) and submerged in oxygenated ACSF that was supplied to the chamber at a rate of 1.5–2 ml/min. The submerged slice was visualized either directly via the microscope’s optics, or indirectly via a high resolution CCD camera system (Orca Flash 2.1, Hamamatsu) that received the output of a CCD camera attached to the microscope’s video port. Experiments with a holding current of more than -100 pA or in which there was a change in input resistance >30% of the control were rejected. Recordings were obtained
using Multiclamp 700A (Axon Instruments) and were filtered at 1~2 kHz. Current recordings under ramp protocol were digitized at 5 kHz and analyzed using pCLAMP 10 software (Axon Instruments). Whole-cell recordings from trigeminal astrocyte were carried out with KCl based internal solution composed of (mM): 140 KCl, 2 MgCl₂, 10 EGTA, 10 HEPES, 0.3 Na₃-GTP and pH 7.2 (OSM=304).

**Light microscopic immunohistochemistry**

Animals were deeply anesthetized using 2% avertin and perfused with 0.1 M PBS, followed by 4% paraformaldehyde. Brains were postfixed in 4% paraformaldehyde at 4°C for 24 hr and 30% sucrose 4°C for 48 hr. Brains were then cut in coronal sections of 30 μm on a cryosection. Sections were blocked in 0.1 M PBS containing 0.3% Triton X-100 (Sigma) and 2% serum (Donkey and Goat mixture, 1:1, Abcam) for 1 hr at room temperature. Primary antibody was then applied at appropriate dilution [Chicken anti-GFAP at 1:1000 (ab4674, Abcam), Rabbit anti-S100β at 1:500 (ab868, Abcam)] and incubated overnight at 4°C. After this, the sections were washed three times in 0.1 M PBS and incubated in secondary antibody [Alexa 555 donkey anti-chicken IgG at 1:500 (ab150170, Abcam), Alexa 488 goat anti-rabbit IgG at 1:500 (ab150077, Abcam) for 2h]. After three rinses in 0.1 M PBS and the sections were mounted on polysine microscopic slide glass (Thermo Scientific) with fluorosheild with DAPI (F6057, Sigma).

Images were acquired using a Nikon A1R confocal microscope and analysed with Imagel software.

In all statistical comparisons, significance was set at *p<0.05 and **p<0.01. All data are presented as the mean±SEM.

**RESULTS AND DISCUSSION**

We firstly confirmed the expression of P2X₃R in astrocytes in the trigeminal caudal nucleus (Vc) of the brain stem as reported previously [12]. Under the electron microscopic observation, the immunostaining for P2X₃R in the superficial lamina of the Vc was identified by discrete gold-silver particles, easily distinguishable from that for GFAP, which was in the form of amorphous, electron-dense patches of reaction product (Fig. 1). Immunostaining for P2X₃R was observed in the somata and processes of the GFAP-immunopositive (GFAP⁺) astrocytes (Soma, Fig. 1A, B and process, Fig. 1C, D) in the Vc. Gold particle density for P2X₃, representing P2X₃ expression level was significantly higher in the astrocytic process than in the astrocytic soma suggesting that expression level of P2X₃ is significantly higher in the astrocytic process than in the astrocytic soma (Fig. 1E).

To test whether the P2X₃Rs in the astrocytes in the Vc are functional, we performed whole cell patch clamp recordings from Vc slices. The soma of the astrocytes in the Vc (trigeminal astrocytes) were relatively smaller (approximately 7~8 μm of diameter of

![Fig. 1](image-url)
Evidence for Functional Expression of P2X3 in Trigeminal Astrocytes

soma) than that of hippocampal astrocytes when we measured soma area of double positive cells with GFAP and S100β antibodies, both of which are astrocytic marker in CNS (area of soma, size of pixel on S100β signals, μm²: hippocampal astrocyte, 78.41±6.71, n=10; trigeminal astrocyte, 30.30±0.97, n=15; p<0.001, Fig. 2A and 2B). This physical aspect of Vc astrocytes prompted us to use GFAP-GFP mice (mice overexpressing green fluorescent protein under the control of astrocyte-specific GFAP promoter) to visually identify astrocytes in acutely prepared slices. Simultaneously, to enhance chance for tight seal, we used recording electrode with higher pipette resistance (11~14 MΩ) compared with conventional recording electrode (pipette resistance is 4~8 MΩ, which would be ≈ 2 μm pore size of glass electrode). Visual guidance with GFP and smaller tip size of electrode allowed us to identify and successfully whole-cell patch-clamp the astrocytes in Vc (Fig. 3A). To observe ATP-mediated astrocytic current, we monitored the change of membrane conductance stimulated by a periodic voltage ramps from +100 mV to -100 mV and examined both the amplitude of ramp-currents and reversal potentials to verify that recorded-current is mediated by purinergic receptor activity (Fig. 3B). During recordings from astrocyte in the Vc, we found that application of a P2X agonist, α,β-meATP (M6517, Sigma, USA), increased the membrane conductance, and that this effect was blocked by co-application of A317491 (A2979, Sigma, USA), a P2X3 specific antagonist (Fig. 3B and 3C, black and green line). Application of α,β-meATP induced not only increase of amplitude of ramp current, but also induced slow inward current during recordings which was partially blocked by co-application of A317491 (Fig. 3B). The voltage dependence of P2X3-mediated current was determined by subtracting ramp current traces recorded during co-application of α,β-meATP and A-317491 from those recorded during application of α,β-meATP only (Fig. 3C, blue line). The reversal potential of the P2X3-mediated current fell at around 0 mV, which is consistent with a current mediated by a nonselective cation channel. Summary bar graph showed that application of α,β-meATP increased current density of ramp current in astrocyte and its increase was almost completely blocked by A-317491 (average of pA/pF at maximal peak: Control 29.74±3.75; α,β-meATP, 46.65±5.79; α,β-meATP+A-317491, 32.57±5.28; n=9; p=0.0227 , Control vs α,β-meATP; p=0.0029, α,β-meATP vs α,β-meATP+A-317491, paired sample t-test, Fig. 3D). These results indicate that astrocytes in the Vc express functional purinergic receptors, including P2X3Rs.

In conclusion, we demonstrate for the first time that astrocytes in the Vc express a functional P2X3Rs using whole-cell voltage-clamp recording from trigeminal astrocytes. Together with our previous report that P2X3Rs are upregulated in the fine astrocytic process following CCI-ION [12], this result strongly suggests the existence of astrocytic P2X3-R mediated mechanism for craniofacial neuropathic pain which was initially attributed to neuronal P2X3R on pain sensation. Future investigations on the astrocytic P2X3-R mediated mechanism for the modulation of the primary nociceptive neurons and their postsynaptic neurons in the Vc will be needed.

Fig. 2. Comparison of size of soma for astrocytes in hippocampus and trigeminal nucleus. (A) Pictures show result of immunolabeling for GFAP (red) and S100β (green) in hippocampal CA1 and trigeminal nucleus. Upper panel shows low magnification. Lower panel shows high magnification of dotted line box in upper panel. Scale bar indicates 60 μm (upper panel) and 10 μm (lower panel) respectively. (B) Summary bar graph shows area of soma measured pixel size of soma on S100β positive cells in hippocampus and trigeminal astrocyte. *** indicates p<0.001 by student t-test.
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