INTRODUCTION

Astrocytes are increasingly recognized as an important cell type in the central nervous system and its importance has been increasingly recognized in the brain pathophysiology. To study in vivo function of astrocyte, astrocyte-specific gene-targeting is regarded as a powerful approach. Especially, hGFAP-CreERT2, which expresses tamoxifen-inducible Cre recombinase under the human GFAP promoter, has been developed and characterized from several research groups. However, one of these mouse lines, [Tg(GFAP-Cre/ERT2)13Kdmc] from Ken McCarthy group has not been quantitatively analyzed, despite its frequent use. Here, we performed comprehensive characterization of this mouse line with quantitative analysis. By crossing this mouse line with Ai14 (RCL-tdTomato), a very sensitive Cre reporter mouse line, we visualized the Cre-expressing cells in various brain regions. For quantitative analysis, we immunostained S100β as an astrocytic marker and NeuN, tyrosine hydroxylase or calbindin as a neuronal marker in different brain regions. We calculated ‘astrocyte specificity’ as the proportion of co-labelled S100β and tdTomato positive cells in the total number of tdTomato positive cells and the ‘astrocyte coverage’ as the proportion of co-labelled S100β and tdTomato positive cells in the total number of S100β positive cells. Interestingly, we found varying degree of astrocyte specificity and coverage in each brain region. In cortex, hypothalamus, substantia nigra pars compacta and cerebellar Purkinje layer, we observed high astrocyte specificity (over 89%) and relatively high astrocyte coverage (over 70%). In striatum, hippocampal CA1 layer, dentate gyrus and cerebellar granule layer, we observed high astrocyte specificity (over 80%), but relatively low astrocyte coverage (50-60%). However, thalamus and amygdala showed low astrocyte specificity (about 65%) and significant neuron specificity (over 30%). This hGFAP-CreERT2 mouse line can be useful for genetic modulations of target gene either in gain-of-function or loss-of-function studies in the brain regions with high astrocyte specificity and coverage. However, the use of this mouse line should be restricted to gain-of-function studies in the brain regions with high astrocyte specificity but low coverage. In conclusion, hGFAP-CreERT2 mouse line could be a powerful tool for gene-targeting of astrocytes in cortex, striatum, hippocampus, hypothalamus, substantia nigra pars compacta and cerebellum, but not in thalamus and amygdala.

Key words: Astrocytes, Glial fibrillary astrocytic protein, Cre recombinase, Tamoxifen
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in the brain pathophysiology. Physiologically, astrocytes not only function at developmental stages by guiding neuronal migration [1] but also contribute to neuronal functions including taking up neurotransmitters through their transporters [2], sensing neuronal activity through many neurotransmitter receptors [3] and releasing gliotransmitters such as glutamate, GABA, ATP and D-serine [4-7]. Additionally, they also buffer extracellular potassium for proper neuronal firing [8]. At pathological conditions, including neurodegenerative disease, astrocytic functions are known to be altered [9]. For example, it was reported that glutamate transporter T1 (GLT1) levels are reduced [10] and tonic γ-aminobutyric acid (GABA) release is increased in animal models of Alzheimer’s disease [11], which cause excitotoxicity and neuronal inhibition, respectively.

During these processes, astrocytic gene expression is also altered and involved in these functional alternation. For studying in vivo functions, astrocytic gene manipulation is required. For this, promoters of human glial fibrillary acidic protein (hGFAP), mouse aldehyde dehydrogenase family 1 member L1 (mALDH1L1) and human ALDH1L1 (hALDH1L1) were introduced to modulate gene expression in astrocytes by expressing Cre recombinase (Cre) cell type-specific manner through transgenic mouse or virus injection [12-14]. However, these promoters showed leaky Cre expression in other brain cell types such as neuron, microglia or oligodendrocytes [13, 15]. Therefore, their astrocyte specificity has been questioned. In particular, it has been reported that transient GFAP expression in neuronal progenitor cells causes significant proportion of neuronal Cre expression in hGFAP-Cre mouse line [15]. Therefore, utilizing hGFAP-Cre mouse line is not suitable for targeting mature astrocytes. Based on this issue, inducible Cre driver transgenic mouse lines such as hGFAP-CreERT2, GLAST-CreERT2 and ALDH1L1-CreERT2 have been developed [16-21]. However, GLAST-CreERT2 also shows a significant neuronal expression, whereas ALDH1L1-CreERT2 does not [20]. Although mALDH1L1 promoter has high astrocyte specificity in the brain, it has significant limitations to utilize in brain-specific manner by the fact that ALDH1L1 mRNA is also highly expressed in other parts of the body [22]. Due to these limitations, until now there is no perfect option for genetically targeting astrocyte in the brain.

hGFAP-CreERT2 mouse lines, which are the most popular transgenic mouse line for genetically targeting astrocyte, were made by several groups [16-18, 21]. Two mouse lines which were initially developed each from Vaccarino group [Tg(GFAP-cre/ERT2) 505Fmvr] (MGI: 3774167), Kirchhoff group [Tg(GFAP-cre/ERT2) 1Fki] (MGI: 4418665), and Baker group were very well characterized with detailed quantitative analysis of astrocyte specificity. However, the other mouse line which is from Ken McCarthy group [Tg(GFAP-cre/ERT2) 13Kdmc] (MGI:3712447) has not been well characterized in various brain regions, despite its frequent use with over 75 citations of the original paper [23-25]. In this regard, it is necessary to characterize the cell types and its exact proportion in various brain regions for accurate interpretation of results.

Based on this need, we set out to characterize [Tg(GFAP-cre/ ERT2) 13Kdmc], which we refer to as hGFAP-CreERT2, for assessing its efficiency of targeting astrocyte in various brain regions. To quantitatively analyze Cre expression pattern by crossing with Ai14 mouse line whose floxed tdTomato fluorescence gene can be sensitively and brightly activated by Cre. In this double transgenic mouse line, we analyzed the population of tdTomato positive cells to examine the ‘astrocyte specificity’, indicating the proportion of astrocyte in the total number of Cre expressing cells, and ‘astrocyte coverage’ indicating the proportion of Cre expressing cells in the total number of astrocytes, of the hGFAP-CreERT2 mouse line depending on brain regions.

MATERIALS AND METHODS

Animals

To generate reporter mice for visualizing Cre expression, we made double transgenic mouse line using C57BL/6 strain by crossing Ai14 (Rosa-CAG-LSL-tdTomato-WPRE::deltaNeo) mouse with hGFAP-CreERT2 mouse (hGFAP-CreERT2×Ai14). The mice have been bred in suitable space mice cage (Thoren, Hazleton, USA) with freely accessing to food and water and kept on a 12-hour-light-dark cycle. All experimental steps described in this paper were performed in accordance with the institutional guidelines for experimental animal care and use of the Korea Institute of Science and Technology (KIST; Seoul, Korea).

Genotyping

Mice tails were digested overnight at 60°C, using 1 mg/ml Proteinase K (21560025-2, bioWORLD, Dublin, USA) in lysis reagent for genotyping (102-T, Viagenbiotech, Los Angeles, USA). On the following day, after inactivating the protein kinase K, the PCR reaction mixture contained Taq DNA Polymerase, MgCl2, dNTP mix (#QM13531, Bioquest, Seoul, Korea) and 1 μl genomic DNA template, 0.1 μM forward and reverse primer pairs and ddH2O. PCR was performed using the specific primer pairs as the detailed protocol is described in Fig. 1B. The reaction products were run on 1.5% agarose gels (HB0100500, E&S Bio Electronics, Daejeon, Korea) in TAE buffer (40 mM Tris-pH 7.6, 20 mM acetic acid and 1 mM EDTA) at 100 V and visualized using Nobel view (NOV001, www.enjournal.org 509
Fig. 1. Genetic confirmation of hGFAP-CreERT2 mouse line and its tamoxifen inducible Cre expression. (A) Genetic map of hGFAP-CreERT2. (B) The primer sequences and PCR protocol of hGFAP-CreERT2 transgenic mice. (C) PCR results of genotyping. TG: Transgenic mouse, WT: Wild-Type, DW: ddH2O. (D) Schematic diagram of generation of double transgenic mouse. (E) Sagittal brain section images from tamoxifen or vehicle (sunflower oil) treated hGFAP-CreERT2 mouse. Red fluorescence indicates tdTomato, and green fluorescence indicates S100β. OB=Olfactory bulb, CTX=Cortex, STR=Striatum, HIP=Hippocampus, TH=thalamus, HY=Hypothalamus, CB=Cerebellum, MB=Midbrain, P=Pons, and MY=Medulla. (F) S100β and GFAP localization images of hippocampal CA1 layer (CA1, left) and dentate gyrus (DG, middle). Green fluorescence indicates S100β, and magenta fluorescence indicates GFAP. (G) Summary table of co-localization ratio of S100β+ & GFAP+ cells among S100β+ cells in each images, (CA1: n=4 sections, DG: n=2 sections from 2 mice) (Mean±SEM).
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Tamoxifen administration

Stock solution of tamoxifen was made at concentration of 20 mg/ml in sunflower seed oil (Sigma, St. Louis, USA) mixture with 1/10 ethanol (64-17-5, Millipore, Burlington, USA). 100 μl of tamoxifen were administered into each hGFAP-CreERT2×Ai14 double transgenic mice (8–12 weeks, male and female) by intraperitoneal (i.p.) injection per day for consecutive 5 days.

Immunohistochemistry

Mice were anesthetized with 2% avertin (20 μl/g, T48402, Sigma) by intraperitoneal injection and perfused with 4% paraformaldehyde accompanied by 0.1 M phosphate buffered saline (PBS) at room temperature. Extracted whole brains were post-fixed in 4% paraformaldehyde at 4°C overnight. After that, post-fixed brains were incubated in 30% sucrose at 4°C for more than 24 hours. Both sagittal and coronal sections were sliced with 30 μm thickness in cryostat microtome (Thermo Scientific, Waltham, USA). Sliced brain sections were washed three times with PBS and blocked with blocking solution (0.3% Triton X-100 (X-100, Sigma), 2% donkey serum (GTX27475, Genetex, Irvine, USA) and 2% goat serum (ab7481, Abcam, Cambridge, UK) in 0.1 M PBS for 1.5 hours. And then, these sections were incubated in suitable primary antibodies (rabbit anti-S100β antibody (ab41548, Abcam) and mouse anti-NeuN antibody (MAB377, Abcam) or chicken anti-tyrosine hydroxylase (P40101-0 TH, Abcam) or mouse anti-Calbindin antibody (C9848, Sigma) mixture with shaking at 4°C overnight. After overnight incubation, brain sections were washed three times with PBS and incubated in matched secondary antibodies (Jackson, Bar Harbor, USA), they were also stained with 4’,6’-diamidino-2-phenylindole (DAPI, 62248, Pierce, Waltham, USA). 100 μl of tamoxifen were administered into each hGFAP-CreERT2×Ai14 double transgenic mice (8–12 weeks, male and female) by intraperitoneal (i.p.) injection per day for consecutive 5 days.

RESULTS

Genotypic and phenotypic confirmation of hGFAP-CreERT2 mouse line

According to the original description of hGFAP-CreERT2 mouse line, this mouse line was generated by micro injection of plasmid form of the transgene (2.2 kb human GFAP promoter and CreERT2) into mouse embryonic stem cell [16]. For genetic confirmation, we designed two unique sets of primers, each targeting the promoter and transgene, and performed polymerase chain reaction (PCR) of tail biopsy genomic DNA (Fig. 1A and B). Gel electrophoresis of the PCR products showed two bands at 439 bp and 179 bp (Fig. 1C), indicating hGFAP promoter and CreERT2, respectively.

To characterize cell type specificity of tamoxifen-inducible Cre-expressing cells in hGFAP-CreERT2×Ai14 double transgenic mouse line, we administered tamoxifen or sunflower oil in these mice by i.p. injection for five consecutive days (Fig. 1D). Upon tamoxifen administration, we observed the tdTomato fluorescence in Cre-expressing cells. To scan and validate the Cre-expressing cells as an astrocyte, we performed immunohistochemistry by using an antibody against S100β as an astrocytic marker in sagittal brain sections (Fig. 1E, top). These sections showed universal tdTomato fluorescence over the whole brain region, whereas in thalamus tdTomato expressing cells were rarely merged with S100β showing segregated two distinct colors (Fig. 1E, top). These results imply that the degree of astrocyte specificity is different in various brain regions.
These regional differences in the degree of astrocyte specificity could be originated from either the lack of specificity of S100β antibody to astrocytes or the region-dependent differential activity of human GFAP promoter in non-astrocytes. In fact, it has been reported previously that some S100β antibodies labels oligodendrocyte progenitor cells (OPCs) in the developing brain [27-29]. However, there has been no information on the proportion of S100β-positive astrocytes in adult brain. Therefore, to directly determine what proportions of S100β-positive cells are GFAP-positive astrocytes or GFAP-negative (non-astrocytic, possibly OPCs) cells, we co-immunostained the adult mouse brain with GFAP and S100β antibodies and observed that about 96% of S100β-positive cells were GFAP-positive astrocytes in hippocampal CA1 and DG regions (Fig. 1F and 1G). These results indicate that the S100β antibody is an excellent astrocyte marker in adult brain. Therefore, by using this S100β antibody as a reliable astrocyte marker, we quantitatively analyzed cell type-specific expression of Cre in various brain regions where astrocytic functions are extensively researched.

In cortex (somatomotor area) and striatum (caudoputamen), hGFAP-CreERT2 mouse shows high astrocyte specificity but differential coverage

To investigate tamoxifen-inducible Cre-expressing cells in cortex, we performed immunohistochemistry by utilizing S100β as an astrocytic marker rather than GFAP, which is known to be differentially expressed in various brain regions [30]. We used NeuN (Neuronal nuclear antigen) as a neuronal marker for quantitative analysis (Fig. 2A). We found that tdTomato expressing cells in each image were predominantly co-localized with S100β in cortex (Fig. 2B and C, green, 87.37±1.45%). However, these tdTomato expressing cells were rarely co-localized with NeuN (Fig. 2B and C, cyan, 8.04±1.03%). Based on these high magnification images, we calculated ‘astrocyte specificity’ which is the ratio of the total counted number of tdTomato and S100β double positive (tdTomato+&S100β+) cells to that of tdTomato positive (tdTomato+) cells. Moreover, we also calculated ‘astrocyte coverage’ which is the ratio of the total counted number of tdTomato+&S100β+ cells to that of S100β+ cells. Based on these calculations, astrocyte specificity was 87.69% and coverage was 74.13% in cortex (Fig. 2D and E). These results indicate that hGFAP-CreERT2 mouse line has high astrocyte specificity with over 80% of specificity and high astrocyte coverage with over 70% of coverage in cortex.

In striatum (caudoputamen) (Fig. 3A), tdTomato expressing cells in each image also showed pre-dominant S100β co-localization which was comparable or higher than that of cortex (Fig. 3B and C, green, 90.04±3.42%) and rare NeuN co-localization (Fig. 3B and C, cyan, 2.71±1.25%). Astrocyte specificity was 89.02% and coverage was 50.69% (Fig. 3D and E). These results indicate that hGFAP-CreERT2 mouse line has high astrocyte specificity with over 80% of specificity, but low astrocyte coverage with under 70% of coverage in striatum.

In hippocampus, hGFAP-CreERT2 mouse shows high astrocyte specificity but low coverage

Next, we examined hippocampal CA1 layer (CA1) and Dentate Gyrus (DG). In hippocampal CA1 (Fig. 4A), tdTomato expressing cells in each image were highly co-localized with S100β (Fig. 4B and D, green, 92.61±1.49%) but not with NeuN (Fig. 4B and D, cyan, 5.21±2.10%). Astrocyte specificity was 92.83% and coverage was 56.28% (Fig. 4E and F, left).

In DG area (Fig. 4B), tdTomato expressing cells in each image were highly co-localized with S100β (Fig. 4C and D, green, 85.91±1.08%) than that of NeuN (Fig. 4C and D, cyan, 10.59±1.13%). Although, a few tdTomato expressing cells were found in subgranular zone where GFAP positive neuronal progenitor cells are known to be present [31], the total number was negligible (<5%). Astrocyte specificity was 86.13% and coverage was 61.44% (Fig. 4E and F, right). These results indicate that hGFAP-CreERT2 mouse line has high astrocyte specificity with over 80% of specificity, but relatively low astrocyte coverage with under 70% of coverage in both hippocampus sub-regions.

In LHA and SNpC, hGFAP-CreERT2 mouse shows high astrocyte specificity and coverage

We further tested Cre expression pattern in lateral hypothalamus (LHA) (Fig. 5A). In LHA, tdTomato expressing cells were mainly in S100β+ in each image (Fig. 5B and C, green, 92.05±2.39%), but NeuN+ cells were little (Fig. 5B and C, cyan, 5.48±2.61%). Astrocyte specificity was 92.20% and coverage was 79.24% in LHA (Fig. 5D and E). These results indicate hGFAP-CreERT2 mouse line has high astrocyte specificity with over 80% of specificity and high astrocyte coverage with over 70% of coverage in LHA.

Similar with LHA, we also tested in substantia nigra pars compacta (SNpC) by using S100β and tyrosine hydroxylase (TH) which marks dopaminergic neuron (Fig. 6A). Almost all of tdTomato expressing cells in SNpC were also S100β+ in each image (Fig. 6B and C, green, 97.74±0.59%), but no TH+ cells (Fig. 6B and C, cyan, 0.00%). In SNpC, astrocyte specificity was 97.64% and coverage was 72.94% (Fig. 6D and E). These results indicate that hGFAP-CreERT2 mouse line has the highest astrocyte specificity and high astrocyte coverage with over 70% of coverage in SNpC.
Fig. 2. In cortex (somatomotor area), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of cortex. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in (A). (C) Quantification of tdTomato population with S100β (green), NeuN (cyan) and S100β & NeuN (gray) cells in cortex (Mean±SEM, n=5 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato & S100β/tdTomato) and coverage (tdTomato & S100β/S100β). (E) Summary table of above analysis.
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Fig. 3. In striatum (Caudoputamen), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of striatum. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in (A). (C) Quantification of tdTomato+ population with S100β+ (green), NeuN (cyan) and S100β & NeuN (gray) cells in striatum (Mean±SEM, n=5 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato+&S100β+/tdTomato+) and coverage (tdTomato+&S100β+/S100β+). (E) Summary table of above analysis.
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**Fig. 4.** In hippocampal CA1 layer (CA1) and Dentate gyrus (DG), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of hippocampal CA1 (left) and DG (right). Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in CA1. (C) High magnification images from the box in DG. (D) Quantification of tdTomato population with S100β (green), NeuN (cyan) and S100β & NeuN (gray) cells in hippocampal CA1 and DG (Mean±SEM, n=6 sections from 3 mice). (E) Summary graph of astrocyte specificity (tdTomato & S100β/tdTomato) and coverage (tdTomato & S100β/S100β) from each region. (F) Summary table of above analysis (CA1: left, DG: right).
**Fig. 5.** In Lateral hypothalamic area (LHA), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of LHA. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in (A). (C) Quantification of tdTomato population with S100β (green), NeuN (cyan) and S100β & NeuN (gray) cells in LHA (Mean±SEM, n=7 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato & S100β/tdTomato) and coverage (tdTomato & S100β/S100β). (E) Summary table of above analysis.
**Fig. 6.** In substantia nigra pars compacta (SNpC), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of SNpC. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=TH. (B) High magnification images from the box in (A). (C) Quantification of tdTomato⁺ population with S100β⁺ (green), TH⁺ (cyan) and S100β⁺ & TH⁺ (gray) cells in SNpC (Mean±SEM, n=6 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato⁺ & S100β⁺ / tdTomato⁺) and coverage (tdTomato⁺ & S100β⁺ / S100β⁺). (E) Summary table of above analysis.
Fig. 7. In cerebellar Purkinje cell layer (PL) and granule cell layer (GL), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of cerebellar PL (left) and GL (right). Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=Calbindin. (B) High magnification images from the box in PL. (C) High magnification images from the box in GL. (D) Quantification of tdTomato+ population with S100β+ (green), Calbindin+ (cyan) and S100β- & Calbindin- (gray) cells in cerebellar PL and GL. (Mean±SEM, n=6 sections from 3 mice). (E) Summary graph of astrocyte specificity (tdTomato+ & S100β+ / tdTomato+ ) and coverage (tdTomato+ & S100β+ / S100β+ ) from each region. (F) Summary table of above analysis (PL: left, GL: right).
Fig. 8. In basolateral amygdala (BLA), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of BLA. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in (A). (C) Quantification of tdTomato+ population with S100β+ (green), NeuN+ (cyan) and S100β−&NeuN− (gray) cells in BLA (Mean±SEM, n=6 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato+&S100β+/tdTomato−) and coverage (tdTomato+&S100β+/S100β−). (E) Summary table of above analysis.
Fig. 9. In thalamic ventro-basal complex (VB), tdTomato expression driven by hGFAP-CreERT2. (A) Low magnification images of VB. Each color represents as following description. Green=S100β; Red=tdTomato; Cyan=NeuN. (B) High magnification images from the box in (A). (C) Quantification of tdTomato+ population with S100β (green), NeuN (cyan) and S100β−&NeuN− (gray) cells in VB (Mean±SEM, n=6 sections from 3 mice). (D) Summary graph of astrocyte specificity (tdTomato+&S100β+/tdTomato+&S100β−) and coverage (tdTomato+&S100β+/S100β−). (E) Summary table of above analysis.
In cerebellum, hGFAP-CreERT2 mouse shows high astrocyte specificity but differential coverage

In addition, we also scrutinized Cre expression pattern in cerebellar Purkinje cell layer (PL) and granule cell layer (GL). Therefore, we stained with S100β for marking cerebellar astrocyte and Bergmann glia, and with Calbindin for marking Purkinje cell (Fig. 7A). In cerebellar PL, tdTomato expressing cells in each image were highly co-localized with S100β (Fig. 7B and D, green, 94.65±1.44%)
but not Calbindin (Fig. 7B and D, cyan, 0.00%). Therefore, astrocyte specificity was 94.96% and coverage was 91.25% (Fig. 8E and F, left).

In GL area (Fig. 7B), tdTomato expressing cells in each image were much more co-localized with S100β<sup>+</sup> cells (Fig. 7C and D, green, 83.46±1.91%) than that of Calbindin<sup>+</sup> cells (Fig. 7C and D, cyan, 0.00%). Accordingly, astrocyte specificity was 83.29% and coverage was 57.69% in GL (Fig. 7E and F, right). These results indicate that hGFAP-CreERT2 mouse shows high astrocyte specificity with over 80% of specificity in both sub-regions of cerebellum. However, astrocyte coverage is different between these two sub-regions. Astrocyte coverage is high in PL with over 70% but low in GL with under 70%.

**In BLA and thalamic VB, hGFAP-CreERT2 mouse shows low astrocyte specificity and high coverage**

Finally, we characterized Cre expression in basolateral amygdala (BLA) and thalamic ventro-basal complex (VB) (Fig. 8A and 9A). Even though tdTomato expressing cells in each image were much more co-localized with S100β<sup>+</sup> the ratio of co-localization with NeuN was also high (S100β<sup>+</sup> & tdTomato<sup>+</sup>: 66.28±2.80%, NeuN<sup>+</sup> & tdTomato<sup>+</sup>: 32.64±2.70%) (Fig. 8B and C, S100β<sup>+</sup>; green, NeuN: cyan). As a result, astrocyte specificity was 65.34%, and coverage was 84.33% in BLA (Fig. 8D and E). These results indicate that hGFAP-CreERT2 mouse line has low astrocyte specificity with under 80% of specificity and high astrocyte coverage with over 70% of coverage in BLA.

Similarly, in thalamic VB, tdTomato expressing cells in each image showed non-specific expression between S100β<sup>+</sup> and NeuN<sup>+</sup> (S100β<sup>+</sup> & tdTomato<sup>+</sup>: 67.42±5.66%, NeuN<sup>+</sup> & tdTomato<sup>+</sup>: 32.33±5.48%) (Fig. 9B and C, S100β<sup>+</sup>; green, NeuN: cyan). Based on this result, astrocyte specificity was 64.61%, and coverage was 89.30% in thalamic VB (Fig. 9D and E). These results indicate that hGFAP-CreERT2 mouse line has low astrocyte specificity with under 80% of specificity and high astrocyte coverage with over 70% of coverage in thalamic VB.

To sum up, we generated a color-coded brain map to represent the two quantitative aspects; ‘astrocyte specificity’ and ‘astrocyte coverage’ (Fig. 10A). Based on this brain map and table, astrocyte specificity of most of investigated regions was over 80% except in VB and BLA, whereas astrocyte coverage varied among the various brain regions ranging from 50% to 90% (Fig. 10A and B).

**DISCUSSION**

Our study quantitatively assessed the astrocyte specificity and coverage of hGFAP-CreERT2 mouse in various brain regions. We have demonstrated that this mouse line expressing Cre under the human GFAP promoter shows differential astrocyte specificity and coverage depending on the brain regions (Fig. 10). Surprisingly, our results are different from those of the previously reported transgenic mouse lines utilizing the same human GFAP promoter [17, 18, 21]. For example, Vaccarino group showed 61.4% of astrocyte specificity and Baker group showed 52.9% in cortex. These values are much lower than what we found (87.69% in cortex). In addition, Kirchhoff group showed 28% of astrocyte coverage in cortex [17, 18, 21], which is far lower than what we found (74.07% in cortex). There could be several reasons for this difference. Firstly, the difference of Cre expression properties in each group’s mouse line could come from the usage of different Cre reporter lines. Other three groups utilized Z/eg (lacZ/EGFP), ROSA26-EYFP or ROSA26-lacZ, which are the most commonly used Cre reporter lines [17, 18, 21]. However, these reporter lines show much weaker endogenous fluorescence in fixed tissues, compared with the tdTomato in Ai14 [32]. Because of this limitation, additional indirect immunofluorescence staining was needed to validate cell-type specificity for the other reporter lines [33]. In contrast, Ai14 is much more sensitive with far greater fluorescence than other reporter lines and has been proven to be the most optimal reporter line for identification of Cre expressing cells [32]. Thus, it is possible that the difference of Cre reporter mouse line might have caused the discrepancy, even in same brain region and with the same hGFAP promoter.

It is possible that this discrepancy could be due to the difference of mouse strain, too. The mouse lines from Kirchhoff group (FVB/N) and Baker group (FVB/N) are a different mouse strain from McCarthy group’s mice line (C57BL/6J). A previous report has shown that FVB and C57BL/6N strains have different methylation activities [34]. Therefore, this different methylation condition could differentially regulate the human GFAP promoter shows differential astrocyte specificity and coverage depending on the brain regions (Fig. 10). Surprisingly, our results are different from those of the previously reported transgenic mouse lines utilizing the same human GFAP promoter [17, 18, 21]. For example, Vaccarino group showed 61.4% of astrocyte specificity and Baker group showed 52.9% in cortex. These values are much lower than what we found (87.69% in cortex). In addition, Kirchhoff group showed 28% of astrocyte coverage in cortex [17, 18, 21], which is far lower than what we found (74.07% in cortex). There could be several reasons for this difference. Firstly, the difference of Cre expression properties in each group’s mouse line could come from the usage of different Cre reporter lines. Other three groups utilized Z/eg (lacZ/EGFP), ROSA26-EYFP or ROSA26-lacZ, which are the most commonly used Cre reporter lines [17, 18, 21]. However, these reporter lines show much weaker endogenous fluorescence in fixed tissues, compared with the tdTomato in Ai14 [32]. Because of this limitation, additional indirect immunofluorescence staining was needed to validate cell-type specificity for the other reporter lines [33]. In contrast, Ai14 is much more sensitive with far greater fluorescence than other reporter lines and has been proven to be the most optimal reporter line for identification of Cre expressing cells [32]. Thus, it is possible that the difference of Cre reporter mouse line might have caused the discrepancy, even in same brain region and with the same hGFAP promoter.

Another possibility is that the transgene copy number (TCN) could be the reason for the difference. It has been reported that TCN affects transgene expression level in mammal due to the concatemeric integration [36]. Based on this report, TCN and transgene expression level have inverse correlation [34]. Therefore, if each of these three mouse line’s TCN is different with each other, it could explain the difference. However, TCN for each mouse line is unknown. In the future, it would be interesting to determine the exact TCN for each of those mouse lines using the next-generation sequencing (NGS) technology, especially whole genome sequencing (WGS), as a rapid and cost-effective approach [37].

Different tamoxifen administration protocol, including the types of injected ERT2 agonist or dosage of agonist, also can affect tamoxifen-mediated CreERT2 activation [20]. In previous reports,
authors performed different tamoxifen injection protocol that is different from our study. For example, Vaccarino group used different ERT2 agonist, 4-hydroxytamoxifen (OHT, 33 mg/kg/day for one day) in mouse pups [17]. On the other hand, Baker group used the same ERT2 agonist, tamoxifen, but injected at different dosage of 225 mg/kg/day for five days [21]. However, Kirchhoff group used the same dosage of tamoxifen as in our study at 100 mg/kg/day for five days. Therefore, different tamoxifen administration methods can be a possible reason for the difference between the current study and previous reports.

The heterogeneity of astrocyte specificity in different brain regions, especially the significant neuronal expression in thalamic VB and BLA, might be caused by alternative gene silencing or heterogeneous activity of human GFAP promoter. It has been well known that histone methylation and acetylation are strongly associated with control of transcriptional activity [14, 34, 35]. If some unknown mechanisms involving histone demethylation or acetylation are differentially involved in thalamic or amygdala neurons, there could be leaky neuronal expression of Cre in those regions. In addition, a differential expression and activity of certain transcriptional factors could contribute to the heterogeneity of both astrocyte specificity and coverage depending on different brain regions [38]. It will be very helpful to elucidate the relationships among the brain regional heterogeneity, expression pattern of transcription factors and epigenetic modification.

Our study provides very useful information regarding the optimal use of the hGFAP-CreERT2 mouse line, depending on the brain region of interest. For example, in cortex, LHA, SNpC and cerebellar Purkinje cell layer, the mouse line shows high astrocyte specificity with over 89% of specificity and high astrocyte coverage with over 70% of coverage. In those brain regions, this mouse line would be suitable for genetically modulating astrocytic gene expression in bidirectional ways, either gain-of-function or loss-of-function studies. In contrast, in striatum, hippocampus and cerebellar granule cell layer, the mouse line shows high astrocyte specificity with over 80% of specificity but covering only 50–60% of astrocytes. In those brain regions, this mouse line would be more suitable for genetically modulating gene expression in gain-of-function studies, rather than loss-of-function studies.

In conclusion, hGFAP-CreERT2 mouse line shows high astrocyte specific Cre expression in most of the brain regions. Our study demonstrates that this transgenic mouse line is very useful for manipulating genes of interest in astrocyte-specific manner, and perhaps is still one of the best genetic tools available for study of astrocyte function in vivo.

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