Preferential arborization of dendrites and axons of parvalbumin- and somatostatin-positive GABAergic neurons within subregions of the mouse claustrum

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1. Introduction

The claustrum is a thin subcortical nucleus located between the insular cortex and the striatum. This small nucleus has abundant reciprocal excitatory connections with almost the entire cerebral cortex (Atlan et al., 2017; Wang et al., 2017, 2022; Zingg et al., 2018; Marriott et al., 2021; Shelton et al., 2022) and coordinates activity in the individual cortices (Crick and Koch, 2005; Smythies et al., 2012). Indeed, claustral neurons synchronously regulate cortical activity during slow-wave sleep in mice (Narikiyo et al., 2020) and reptiles (Norimoto et al., 2020). The excitatory connections between the claustrum and each cortical area have been further demonstrated to play key roles in various physiological functions (Goll et al., 2015; Chia et al., 2020), such as salience detection, attention, and regulation of impulsivity with claustro-frontocortical connections (White et al., 2018; Liu et al., 2019; Chia et al., 2020; Terem et al., 2020; Atlant et al., 2021); contextual memory with claustro-medial entorhinal cortical connections (Kitanishi and Matsuo, 2017); and motor response related to selection tasks with claustro-somatosensory cortical connections (Cheeves et al., 2022).

The claustrum has been divided into three subregions by Tissue clearing

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ABSTRACT

The claustrum coordinates the activities of individual cortical areas through abundant reciprocal connections with the cerebral cortex. Although these excitatory connections have been extensively investigated in three subregions of the claustrum—core region and dorsal and ventral shell regions—the contribution of GABAergic neurons to the circuitry in each subregion remains unclear. Here, we examined the distribution of GABAergic neurons and their dendritic and axonal arborizations in each subregion. Combining in situ hybridization with immunofluorescence histochemistry showed that approximately 10% of neuronal nuclei-positive cells expressed glumatic acid decarboxylase 67 mRNA across the claustral subregions. Approximately 20%, 30%, and 10% of GABAergic neurons were immunoreactive for parvalbumin (PV), somatostatin (SOM), and vasactive intestinal polypeptide, respectively, in each subregion, and these neurochemical markers showed little overlap with each other. We then reconstructed PV and SOM neurons labeled with adeno-associated virus vectors. The dendrites and axons of PV and SOM neurons were preferentially localized to their respective subregions where their cell bodies were located. Furthermore, the axons were preferentially extended in a rostrocaudal direction, whereas the dendrites were relatively isotropic. The present findings suggest that claustral PV and SOM neurons might execute information processing separately within the core and shell regions.
and in synchronization of claustral excitatory neurons (Kim et al., 2016; reported to receive synaptic inputs from corticoclaustral axons and play and VIP neurons) composed electrophysiologically distinct populations vasoactive intestinal polypeptide (VIP)-expressing neurons (PV, SOM, et al., 2017). Among GABAergic neurons, PV-, somatostatin (SOM)-, and those in the ventral shell region have connections with the ento regions—

Anterograde and retrograde tracing studies have demonstrated reciprocal connections between the claustral subregions and cerebral cortex: claustral glutamatergic neurons in the core region send outputs to and receive inputs from the retrosplenial and visual cortices; those in the dorsal shell region connect with the motor and somatosensory cortices; and those in the ventral shell region have connections with the entorhinal cortex (Atlan et al., 2017; Marriott et al., 2021; Ham and Augustine, 2022; Shelton et al., 2022). This topographical organization suggests that the subregional structures would serve as functional modules via excitatory connections with the cerebral cortex (Chia et al., 2020; Smith et al., 2020; Marriott et al., 2021; Ham and Augustine, 2022).

GABAergic neurons in the claustrum have been characterized using immunohistochemical approaches (Sieden et al., 1990; Druga et al., 1993; Kowianski et al., 2001, 2008; Real et al., 2003; Davila et al., 2005; Morello et al., 2022). Electron microscopic observation demonstrated that claustral aspiny neurons as well as spiny neurons receive corticoclaustral synaptic inputs (LeVay and Sher, 1981). An immunoelectron microscopic study also revealed that synaptic terminals immunopositive for GABA were often detected on the proximal dendrites and cell bodies of non-GABAergic neurons in the claustrum of tree shrew (Day-Brown et al., 2017). Among GABAergic neurons, PV-, somatostatin (SOM)-, and vasoactive intestinal polypeptide (VIP)-expressing neurons (PV, SOM, and VIP neurons) composed electrophysiologically distinct populations in the mouse claustrum (Graf et al., 2020). Moreover, PV neurons were reported to receive synaptic inputs from corticoclaustral axons and play an important role in amplification of selective cortico-cortical signals and in synchronization of claustral excitatory neurons (Kim et al., 2016; White et al., 2018). Despite the important role of GABAergic neurons in the claustrum circuitry, the distribution of GABAergic neurons within each subregion remains unclear. Furthermore, neurites immunopositive for PV and SOM are densely observed in the core and shell regions, respectively (Marriott et al., 2021), raising the question of whether the neurites of these two neurons are differentially distributed in the core and shell subregions.

In the present study, we performed triple immunofluorescence histochemistry for PV, VGlut2, and NeuN and delineated the mouse claustral subregions based on chemorarchitecture and cytoarchitecture. Subsequently, the populations of glutamatergic and GABAergic neurons were investigated by combining fluorescent in situ hybridization (FISH) and immunofluorescence histochemistry in each claustral subregion. We further quantified the distributions of PV, SOM, and VIP neurons in each subregion. Finally, we selectively visualized PV or SOM neurons by injecting recombinant AAV vectors into the claustrum of PV- or SOM-Cre knock-in mice and optically cleared 1-mm-thick brain slices. After acquisition of three-dimensional (3D) image stacks by confocal microscopy, we reconstructed and analyzed their dendrites and axons in the core and ventral shell regions at the single-neuron level.

2. Materials and methods

2.1. Animals

All procedures involving animals were performed in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. The experiments were approved by the Institutional Animal Care and Use Committees of Juntendo University (Approval No: 2022148 and 2022149). All efforts were made to minimize animal suffering and the number of animals used.

Ten male C57BL/6J mice (8–12 weeks old; Nihon SLI, SC), six male and female PV-Cre heterozygous mice (8–16 weeks old; Pvalbflm1crejek1J, The Jackson Laboratory Stock No: 008069) (Hippenmeyer et al., 2005), and ten male and female SOM-Cre heterozygous mice (8–16 weeks old; Sstflm1 crejek1J, The Jackson Laboratory Stock No: 013044) (Taniguchi et al., 2011) were used in the present study. All mice were maintained in specific pathogen-free conditions under a 12/12 h light/dark cycle (light: 08:00–20:00) with ad libitum access to food and water.

2.2. Tissue preparation

Mice were deeply anesthetized with an intraperitoneal injection of sodium pentobarbital (200 mg/kg; Somnopentyl, Kyoritsu Seiyaku) and transcardially perfused with 20 mL of phosphate buffered 0.9% saline (PBS; pH 7.4), followed by the same volume of 4% paraformaldehyde (PFA; 1.04005.1000, Merck Millipore) in 0.1 M phosphate buffer (PB; pH 7.4). The mouse brains were removed and post-fixed in the same fixative overnight at 4 °C. For triple or quadruple immunofluorescence labeling, the brains were cryoprotected in 30% (w/v) sucrose in 0.1 M PB and cut into 40-µm-thick coronal sections on a freezing microtome (SM2000 R, Leica Biosystems). For 3D reconstruction of PV and SOM neurons, the post-fixed brains were embedded in 2% agar (01028−85, Nacalai Tesque) in 0.1 M PB and cut into coronal slices of 1-mm thickness with a vibrating tissue slicer (Linear PRO7N, Dosaka EM). The sections and slices were stored in PBS containing 0.2% (w/v) NaN₃ at 4 °C.

For ISH histochemistry, the mouse brains were post-fixed in the same fixative for 3 days at 4 °C and cryoprotected in diethylpyrocarbonate (DEPC)-treated 30% (w/v) sucrose in 0.1 M PB. The brains were then cut into 30-µm-thick coronal sections on a freezing microtome and stored in an antifreeze solution that contained 30% (w/v) glycerol and 30% (w/v) ethylene glycol in DEPC-treated 0.1 M PB at −20 °C until further use.

2.3. Immunofluorescence labeling

Triple or quadruple immunofluorescence labeling using free-floating sections was performed at 20–25 °C, and all incubations were followed by a rinse with PBS containing 0.3% (v/v) Triton X-100 (35501−15, Nacalai Tesque) (PBS-X) for 10 min three times. The 40-µm-thick sections were incubated overnight with the following mixture in PBS-X containing 0.12% (w/v) λ-carrageenan (035−09693; Sigma-Aldrich) and 1% (v/v) normal donkey serum (S30−100ML, Merck Millipore) (PBS-XCD): (i) 5 µg/mL affinity-purified mouse antibody against NeuN (MAB377, Merck Millipore, RRID:AB:2298772), 1/4,000-diluted rabbit antisera against PV (PV-27, Swant, RRID:AB:2631173), and 1 µg/mL affinity-purified guinea pig antibody against VGlut2 (VGlut2-GP-Af810, Frontier Institute, RRID:AB:2571621); (ii) 5 µg/mL affinity-purified mouse antibody against NeuN, 1/4,000-diluted rabbit antisera against PV, 1/50-diluted affinity-purified rat antibody against SOM-14 (MAB354, Merck Millipore, RRID:AB:2255365), and 1 µg/mL affinity-purified guinea pig antibody against VIP (Hioki et al., 2018). Subsequently, the sections were incubated for 2 h with the following mixture in PBS-XCD: (i) 5 µg/mL Alexa Fluor (AF) 488-conjugated goat antibody against mouse IgG (A-11029, Thermo Fisher Scientific, RRID:AB:2534088), 5 µg/mL AF568-conjugated goat antibody against rabbit IgG (A-11036, Thermo Fisher Scientific, RRID:AB:10563566), and 5 µg/mL AF647-conjugated...
performed as previously described (Hioki et al., 2010; Ma et al., 2011; immunofluorescence 2.4. Double or triple labeling with fluorescence in situ hybridization and nucleotides 855 Sohn et al., 2014) with some modification. Briefly, sense and antisense goat antibody against guinea IgG (A-21450, Thermo Fisher Scientific, M. Takahashi et al. GAD67 (867–661 in GenBank accession No: Y12257.1) with riboprobes for single-stranded riboprobes for VGluT1 and GAD67 and ribonuclease A treatment, the sections were incubated overnight at 20–25 °C with 1/1,000-diluted peroxidase-conjugated sheep antibody against NeuN and then for 2 h with 5 µg/mL AF647-conjugated goat antibody against mouse IgG (A-21236, Thermo Fisher Scientific, RRID: AB_2535805). The sections were mounted onto APS-coated glass slides and coverslipped as described above.

2.4. Double or triple labeling with fluorescence in situ hybridization and immunofluorescence

Combination of single FISH and immunofluorescence labeling was performed as previously described (Hioki et al., 2010; Ma et al., 2011; Sohn et al., 2014) with some modification. Briefly, sense and antisense single-stranded riboprobes for vesicular glutamate transporter 1 (VGluT1; nucleotides 855–1788 in GenBank accession No: NM_133432.2) (Nakamura et al., 2007) and glutamic acid decarboxylase 67 kDa isoform (GAD67; nucleotides 43–661 in GenBank accession No: V12257.1) (Yamamaki et al., 2003) were synthesized with a digoxigenin (DIG) RNA labeling kit (1277073, Roche Diagnostics). Free-floating sections were treated with 1% (w/v) H2O2 in 0.1 M PB for 30 min and then hybridized for 16–20 h at 60 °C with 1 µg/mL DIG-labeled sense or antisense riboprobes in a hybridization buffer. After washes and ribonuclease A treatment, the sections were incubated overnight at 20–25 °C with one of the following mixtures: (i) 1/1,000- or 1/2,000-diluted peroxidase-conjugated sheep antibody against DIG (11–207–733–910, Roche Diagnostics, RRID:AB_514500) for VGluT1 or GAD67 and 5 µg/mL affinity-purified mouse antibody against NeuN; and (ii) 1/2,000-diluted peroxidase-conjugated sheep antibody against DIG, 5 µg/mL affinity-purified mouse antibody against NeuN, and either 1/4, 000-diluted rabbit antiserum against PV, 5 µg/mL affinity-purified rabbit antibody against SOM-14 (T-4102, Peninsula Laboratories, RRID:AB_518613), or 1 µg/mL affinity-purified guinea-pig antibody against VIP. The sections were incubated for 30 min in a biotinylated tyramine (BT)-glucose oxidase (GO) reaction mixture containing 31 µM BT, 3 µg/mL GO, 20 µg/mL beta-D-glucose, and 1% (w/v) bovine serum albumin (BSA) in 0.1 M PB (Furuta et al., 2009; Kuramoto et al., 2009; Okamoto et al., 2021) to deposit BT onto the tissues via peroxidase activity. Subsequently, the sections were incubated for 2 h at 20–25 °C with the following mixtures of secondary antibodies and streptavidin: 5 µg/mL AF488-conjugated streptavidin (S-11223, Thermo Fisher Scientific, AF647-conjugated goat antibody against rabbit IgG (A-21245, Thermo Fisher Scientific, RRID:AB_2535813) or AF647-conjugated goat antibody against guinea pig IgG. The sections were mounted onto APS-coated glass slides and coverslipped as described above.

2.5. Triple labeling with double fluorescence in situ hybridization and immunofluorescence

Combination of double FISH and immunofluorescence labeling was performed as previously described (Yamauchi et al., 2022b) with some modification. Briefly, sense and antisense single-stranded riboprobes for GAD67 were synthesized with a fluorescein (FITC) RNA labeling kit (11–685–619–910, Roche Diagnostics). Free-floating sections were treated with 1% (w/v) H2O2 in 0.1 M PB for 30 min and then hybridized for 16–20 h at 60 °C with a mixture of 1 µg/mL DIG-labeled sense or antisense riboprobes for VGluT1 and 1 µg/mL FITC-labeled sense or antisense riboprobes for GAD67 in a hybridization buffer. After washes and ribonuclease A treatment, the sections were incubated overnight at 20–25 °C with 1/1,000-diluted peroxidase-conjugated sheep antibody against DIG. The sections were then incubated for 30 min in a fluorochromized tyramide (FT)-GO reaction mixture containing 10 µM CF488A tyramide (92171, Biotium), 3 µg/mL GO, 40 µg/mL beta-D-glucose, and 1% (w/v) BSA in 0.1 M PB. Subsequently, the sections were incubated overnight at 20–25 °C with 2% (w/v) NaN3 in 0.1 M PB to inactivate the peroxidase activity. After thorough washes, the sections were incubated overnight at 20–25 °C with 1,000,000-diluted peroxidase-conjugated sheep antibody against FITC (11–426–346–910, Roche Diagnostics, RRID:AB_840257). The sections were then incubated for 30 min in an FT-GO reaction mixture containing 10 µM CF568 tyramide (92173, Biotium), 3 µg/mL GO, 40 µg/mL beta-D-glucose, and 1% (w/v) BSA in 0.1 M PB. Following washes, the sections were incubated overnight at 20–25 °C with 5 µg/mL affinity-purified mouse antibody against NeuN and then for 2 h with 5 µg/mL AF647-conjugated goat antibody against mouse IgG (A-21236, Thermo Fisher Scientific, RRID: AB_2535805). The sections were mounted onto APS-coated glass slides and coverslipped as described above.

2.6. Confocal microscopy for tissue sections

The 3D image stacks of 30- and 40-µm-thick sections were acquired with a confocal laser scanning microscope (TCS SP8, Leica Microsystems) or a spinning disk confocal microscope (SDCM; Andor Dragonfly 201, Andor Technologies) built on an ECLIPSE FN1 upright microscope (Nikon Instruments). For TCS SP8, 16 × multi-immersion (HC FLUOTAR 16x/0.60 IMM CORR VISIR, numerical aperture [NA] = 0.60, working distance [WD] = 2.5 mm; Leica Microsystems) and 25 × water-immersion (HC FLUOTAR L 25x/0.95 W VISIR, NA = 0.95, WD = 250 µm; Leica Microsystems) objective lenses were used with the pinhole at 2–5.6 Airy disk unit and zoom factor at 1–1.2. CF405M, AF488 or CF488A, AF568 or CF568, and AF647 were excited by 405, 488, 552, and 638 nm lasers and observed through 420–500, 500–560, 570–650, and 650–700 nm emission prism windows, respectively, and detected with a z-interval of 2.5 µm per stack through a hybrid detector (Leica Microsystems). For Andor Dragonfly 201, a 20 × water-immersion objective lens (CFI Apo LWD Lambda S 20XVC WI, NA = 0.95, WD = 0.93 mm; Nikon Instruments) was used with the 40-µm pinhole and 1 × 1 or 3 × 3 binning. CF405M, AF488 or CF488A, AF568 or CF568, and AF647 were excited by 405, 488, 552, and 637 nm lasers, respectively, and observed through 445/46, 521/38, 594/43, and 698/77 nm emission filters, respectively, and detected with a z-interval of 2.5 µm per stack through a Zyla 4.2 PLUS sCMOS camera (Andor Technologies).

The image stacks were opened with Fiji/ImageJ (2.0.0-rc-69/1.52p) (Schindelin et al., 2012) and the Bio-Formats plugin (Linkert et al., 2010), and the number of cells in each channel was quantified using the Cell Counter plugin and the ImageJ 5D plugin. We counted the number of cells in the rostral (from +1.9 mm to +1.0 mm from the bregma), middle (+1.0 mm to 0 mm), and caudal (0 mm to −1.0 mm) parts of the claustrum. The global brightness and contrast of the images were adjusted with ImageJ and Canvas X Draw (ver 7.0.7, ACD systems).

2.7. Plasmid construction and AAV vector production

pAAV2-CMV-tTA-WPRE was constructed as follows. The oligonucleotides (P1 and P2/P3, Supplementary Table) were annealed to form double-stranded DNA and inserted into the MluI site and XhoI/BglII sites of pAAV-MCS (Stratagene; GenBank accession No: AF396260.1), respectively, resulting in pAAV-MCS2. The woodchuck hepatitis virus post-transcriptional regulatory element (WPRE; a gift from Dr. Hope Tj) (Zufferey et al., 1999) signal sequence was amplified using polymerase chain reaction (PCR; P4/P5) and inserted into the MluI/BglII sites of pAAV-MCS2, named as pAAV-MCSW2. We then amplified an improved version of tetracycline-controlled transactivator (tTAad; Clontech) by PCR (P6/P7), inserted the product into the BamHI/XhoI sites of pAAV-MCSW2, and obtained the resultant vector—pAAV2-CMV-tTA-WPRE.
pAAV2-TRE-FLEX-paGFP-BGHpA was constructed as follows. The fragment, TRE-GFP-BGHpA, was excised from pTRE-GFP-BGHpA (Hokiet al., 2009) and inserted into the XhoI/NotI sites of pENTRV1A (Thermo Fisher Scientific), resulting in pENTR1A-TRE-GFP-BGHpA. The annealed oligonucleotide (P8/P9) containing BamHI-BglII-SalI restriction enzyme sites was inserted into the EcoRI/MluI sites of pENTR1A-TRE-GFP-BGHpA to replace the GFP sequence, resulting in pENTR1A-TRE-BBSM-BGHpA. The flip-ectomy (FLEX) switch sequence (Schnittgen et al., 2003), which contains two pairs of loxP and lox2272 sites in opposite orientations, was excised from pBSISS-kfLEX (Sohn et al., 2017) and inserted into the BamHI/MluI sites of pENTR1A-TRE-BSSM-BGHpA. Then, the annealed oligonucleotide (P10) was successively inserted into both the BamHI and BgII sites of the plasmid to eliminate those restriction enzyme sites, resulting in pENTR1A-TRE-FLEX-BGHpA. The insert of the resultant entry vector, TRE-FLEX-BGHpA sequence, was transferred to the destination vector, pAAV2-DEST(f) (Sohn et al., 2017), by homologous recombination with LR Clonase II (11791020, Thermo Fisher Scientific) and concentrated using ultrafiltration with Amicon Ultra-15® (Thermo Fisher Scientific), resulting in pENTR1A-TRE-GFP-BGHpA. The flip-excision (FLEX) switch sequence (Okamoto et al., 2020) was amplified with PCR (P13/P14) and inserted into the pAAV2-TRE-FLEX-EBSHB-BGHpA. Finally, EGFP with a palmitoylation signal derived from the GAP-43 N-terminus (palGFP) (Moriyoshi et al., 1996; Tamamaki et al., 2000; Furuta et al., 2001; Sohn et al., 2017; Okamoto et al., 2020) was amplified with PCR (P13/P14) and inserted into the BamHI/Sall sites of pAAV2-TRE-FLEX-EBSSB-BGHpA, and the resultant vector was named as pAAV2-TRE-FLEX-paGFP-BGHpA.

AAV vector particles were produced and purified as previously described (Sohn et al., 2017; Takahashi et al., 2021). Briefly, pAAV2-TRE-FLEX-paGFP-BGHpA or pAAV2-CMV-TA-WPRE and two helper plasmids, pBSISS-kR2C1 (Sohn et al., 2017) and pHelper (Strategene; GenBank accession No: AF369965.1), were co-transfected into HEK293T cells (RCB2202, Riken BRC) using polyethylenimine (23966, Thermo Scientific) and concentrated using ultrafiltration with Amicon® Ultra-15 (UFC903024, Merck Millipore). The physical titer of the AAV vector was measured using quantitative PCR (UFC903024, Merck Millipore). The physical titer of the AAV vector was measured using quantitative PCR (UFC903024, Merck Millipore).

2.8. Stereotaxic injection of AAV vectors

AAV vector injection into mouse brains was performed as previously described (Okamoto et al., 2020, 2021; Takahashi et al., 2021) with some modification. Briefly, mice were deeply anesthetized with an intraperitoneal injection of medetomidine (0.3 mg/kg; Domitor, Zoetis), midazolam (4 mg/kg; Dormicum, Astellas Pharma), and butorphanol (5 mg/kg; Vetrphare, Meiji Seika Pharma) and placed in a stereotaxic apparatus (SR-5M1-TH, Narishige). A 0.2-μL volume of the virus solution mixture (AAV2/1-CMV-TA-WPRE, 8.0 × 10^13 gc/ml; AAV2/1-TRE-FLEX-paGFP-BGHpA, 4.4 × 10^14 gc/ml) was bilaterally injected into the caudate nucleus of PV-Cre or SOM-Cre mice by pressure through a glass micropipette attached to Picospritzer III (Parker Hannifin). The injection coordinates were as follows: 1.1 mm anterior to the bregma, 3.1 mm lateral to the midline, and 2.5 mm ventral to the brain surface. The mice recovered from anesthesia with an intraperitoneal injection of atipamezole (1.5 mg/kg; Antisedan, Zenoaq) and were maintained under regular health check-ups for 7–10 days.

2.9. 3D reconstruction of PV and SOM neurons in the claustrum

The 1-mm-thick brain slices were optically cleared with ScaleSF treatment (Furuta et al., 2022; Yamauchi et al., 2022a, 2022c). Briefly, the slices were incubated with ScaleSF solution for 2 h at 37 °C, PBS(−) (27575–31, Nacalai Tesque) for 15 min at 20–25 °C, and ScaleS4 solution for 10 h at 37 °C. The cleared slices were observed under an SDCM equipped with a 10 × dry objective lens (CFI Plan Apo Lambda 10X, NA = 0.45, WD = 4.0 mm; Nikon Instruments). We checked whether brain slices contained the caudal neurons labeled with EGFP and selected 5 of 12 hemispheres from PV-Cre/mice and 8 of 20 hemispheres from SOM-Cre mice for subsequent procedures. The formula for ScaleS0 solution was 20% (w/v) D(−)-sorbitol (06286–55, Nacalai Tesque), 5% (w/v) glycerol (9012, Sigma-Aldrich), 1 mM methyl-β-cyclodextrin (M1356, Tokyo Chemical Industry), and 3% (v/v) dimethyl sulfoxide (DMSO; 13407–45, Nacalai Tesque) (PBS(−)) and that for ScaleS4 solution was 40% (w/v) D(−)-sorbitol, 10% (w/v) glycerol, 4 M urea (35940–65, Nacalai Tesque), 0.2% (w/v) Triton X-100, and 25% (v/v) DMSO in distilled deionized water (DDW) (Hama et al., 2015; Miyawaki et al., 2016).

We then performed immunofluorescence labeling using the AbScale method (Hama et al., 2015) on the brain slices. After treatment with 1% (w/v) H2O2 in PBS(−) for 1 h at 20–25 °C, the slices were incubated with ScaleA2 solution for 12 h, ScaleB4(0) solution for 6 h, and ScaleA2 solution for 3 h at 37 °C. After washes with PBS(−), the slices were incubated with AbScale solution containing 5% P-RAN-GFP1 (horse-radish peroxidase fused nanobody against EGFP) (Yamagata and Sanes, 2018) and 1% (v/v) normal donkey serum for 12 h at 4 °C. After washes with AbScale solution and 0.1 M PB, the slices were incubated for 2 h at 20–25 °C in an FT-GO reaction mixture containing 10 µM CF488A tyramide, 3 µg/ml GO, 2% (w/v) BSA, and 20 µg/ml β-D-glucose in 0.1 M PB. After three washes with 0.1 M PB for 30 min at 20–25 °C, the slices were incubated with AbScale solution containing 5 μg/ml affinity-purified rabbit antibody against NeuN (ABN78, Merck Millipore), RRID:AB_10807945) for 5 days at 37 °C, followed by 1.5 µg/ml AF647-conjugated nanobody against rabbit IgG (SAS–10327, Thermo Fisher Scientific) for 1 day at 37 °C. After washes with AbScale solution, the slices were re-fixed with 4% PFA in 0.1 M PB for 30 min at 20–25 °C. Finally, the slices were optically cleared with ScaleS4 solution for 2 h at 37 °C. To keep the sizes of the cleared slices isometric with the original ones, the cleared slices were placed on the imaging chamber (Furuta et al., 2022; Yamauchi et al., 2022a, 2022c) and embedded in 1.5% agarose (L03, Nacalai Tesque) in ScaleS4D25(0) solution (ScaleS4 gel) (Hama et al., 2015). The formula for ScaleA2 solution was 4 M urea, 0.1% (w/v) Triton X-100, and 10% (w/v) glycerol in DDW; that for ScaleB4(0) solution was 8 M urea in DDW; that for AbScale solution was 0.33 M urea and 0.1% (w/v) Triton X-100 in PBS(−); and that for ScaleS4D25(0) solution was 40% (w/v) D(−)-sorbitol, 10% (w/v) glycerol, 4 M urea, and 25% (v/v) DMSO in DDW.

3D image stacks of the slices were acquired with an SDCM using a 20 x glycerol-immersion objective lens (CFI 90 20XGC, NA = 1.0, WD = 8.2 mm; Nikon Instruments), and with the 40-µm pinhole, 3 x 3 binning, and a z-interval of 2.5–2.5 µm. EGFP/CF488A and AF647 were excited by 488 and 637 nm lasers, observed through 521/38 and 698/77 nm emission filters, and detected using a Zyla 4.2 PLUS sCMOS camera. The image stacks were deconvoluted with a Fusion software (ver 2.3.0.45, Andor Technologies), and the 3D renderings were created with an Imaris software (ver 9.9.1, Bitplane).

The dendrites and axons of PV and SOM neurons were three-dimensionally reconstructed in 1–3 serial image stacks with a NeuroLucida 360 software, a computer-assisted neuron tracing system (ver 2021.1.3, MBF Bioscience), as follows. Image stacks files were converted to JPEG 2000 file format with a MicroFiele ± software (ver 2021.1.3, MBF Bioscience), and the image files were loaded onto NeuroLucida 360. The neurites of PV and SOM neurons were then traced with an automatic or semi-automatic directional kernel algorithm in a 3D environment or with manual drawing in a two-dimensional (2D) environment. Among the neurites, the dendrites were distinguished from the axon, whose extensive ramifications maintained a constant diameter through the
functions of a Neurolucida Explorer software (ver 2021.1.1, MBF Bioscience). The dendritic and axonal arborizations in each region were measured with the closed surface analysis function. Arborizations of dendrites and axons were three-dimensionally analyzed with the Sholl analysis function (3D Sholl analysis). The reconstructed data were projected along the rostrocaudal (RC), DV, and mediolateral (ML) directions with Neurolucida Explorer, referring to the mouse brain atlas (Paxinos and Franklin, 2007). The dendritic and axonal arborizations on each projected plane were analyzed using the Sholl analysis plugin (Ferreira et al., 2014) in Fiji/ImageJ (2D Sholl analysis). The data were analyzed and plotted with the pandas ( McKinney, 2010), NumPy ( Harris et al., 2020), and Matplotlib (Hunter, 2007) libraries in Python 3.7 (https://www.python.org/).

2.10. Statistical analysis

Data are represented as means ± standard deviations (SDs). Multiple statistical comparisons were performed for the data presented in Table 1 and Table 2 with Kruskal–Wallis tests followed by Dunn’s post-hoc multiple comparison test. For the data in Fig. 7 and Fig. 8, analysis was performed with two-way analysis of variance (ANOVA) followed by Bonferroni’s post-hoc multiple comparison tests using a Prism8 software (ver 8.4.3, GraphPad Software).

2.11. Data availability

The datasets generated during and/or analyzed during the current study and all biological materials reported in this article are available from the corresponding author on reasonable request.

3. Results

3.1. Delineation of the mouse claustral subregions

In the present study, we utilized cytoarchitecture and chemoarchitecture to identify the claustrum and its subregions. We performed triple immunofluorescence histochemistry for NeuN, PV, and VGluT2 with mouse coronal sections (Fig. 1). Based on the density and size of NeuN-positive cells, the claustrum was distinguished from the adjacent layer Vb of the insular cortex as previously reported (Atlan et al., 2017; Binks et al., 2019; Ham and Augustine, 2022) and divided into two subregions: the core region with a high cell density and the shell region around the core region (Fig. 1A, B1, C, D1, E, F1 and Fig. S1) (Obst-Pernberg et al., 2001; Real et al., 2003; Kitanishi and Matsuo, 2017). Further, the shell region was subdivided into the dorsal and ventral parts by the extended line of border between the DI and AI

### Table 1

| VGluT1 / NeuN | Core region | Shell region | Dorsal part | Ventral part |
|--------------|-------------|--------------|-------------|-------------|
| Rostral      | 93.2 ± 0.6<sup>a</sup> | 90.2 ± 2.6 | 87.5 ± 5.1 | 85.15 ± 9672 |
| Middle       | 90.8 ± 1.9  | 88.0 ± 5.8 | 86.3 ± 5.3 | 6351/3716  |
| Caudal       | 89.8 ± 0.7  | 91.1 ± 0.8 | 87.8 ± 2.5 | 5366-6082  |
| Total        | 91.9 ± 0.7  | 91.4 ± 0.3 | 87.3 ± 4.3 | 20,232/23,070 |

<sup>a</sup>Data are provided as mean ± SD of the percentages in three mice.

<sup>b</sup>The denominator and numerator in parentheses correspond to the total number of cells immunofluorescent for NeuN and that of neurons expressing VGluT1 or GAD67 mRNA.

*<sup>p</sup> = 0.0219 versus the rostral part of the ventral shell region (Dunn’s multiple comparison test after the Kruskal–Wallis test).

4.0 20.5

### Table 2

| GAD67 / NeuN | Core region | Shell region | Dorsal part | Ventral part |
|--------------|-------------|--------------|-------------|-------------|
| Rostral      | 8.7 ± 1.0<sup>a</sup> | 9.9 ± 0.2 | 10.9 ± 0.6 | 1227/11,370 |
| Middle       | 9.1 ± 2.2   | 9.4 ± 2.1 | 11.3 ± 3.1 | 924/8156  |
| Caudal       | 10.1 ± 1.1  | 8.3 ± 2.8 | 11.8 ± 2.0 | 811/6884  |
| Total        | 9.1 ± 1.2   | 9.3 ± 1.4 | 11.2 ± 1.4 | 2962/26,410 |

<sup>a</sup>Data are provided as mean ± SD of the percentages in three mice.

<sup>b</sup>The denominator and numerator in parentheses correspond to the total number of cells immunofluorescent for NeuN and that of neurons expressing GAD67 mRNA and that of neurons immunoreactive for NeuN positive cells, the claustrum was distinguished from the adjacent layer Vb of the insular cortex as previously reported (Atlan et al., 2017; Binks et al., 2019; Ham and Augustine, 2022) and divided into two subregions: the core region with a high cell density and the shell region around the core region (Fig. 1A, B1, C, D1, E, F1 and Fig. S1) (Obst-Pernberg et al., 2001; Real et al., 2003; Kitanishi and Matsuo, 2017). Further, the shell region was subdivided into the dorsal and ventral parts by the extended line of border between the DI and AI

### Table 3

| PV / GAD67 | Core region | Shell region | Dorsal part | Ventral part |
|------------|-------------|--------------|-------------|-------------|
| Rostral    | 28.5 ± 5.3<sup>a</sup> | 15.4 ± 0.4 | 20.9 ± 3.0 | 372/1754   |
| Middle     | 23.4 ± 4.9  | 16.7 ± 0.8 | 18.2 ± 2.1 | 742/3694   |
| Caudal     | 23.1 ± 4.0  | 20.5 ± 0.8 | 20.3 ± 2.3 | 179/883    |
| Total      | 26.2 ± 4.8  | 17.1 ± 0.1<sup>a</sup> | 20.0 ± 1.3 | 742/3694   |

<sup>a</sup>Data are provided as mean ± SD of the percentages in three mice.

<sup>b</sup>The denominator and numerator in parentheses correspond to the total number of neurons expressing GAD67 mRNA and that of neurons immunofluorescent for PV, SOM, or VIP.

*<sup>p</sup> = 0.0219 versus the total core region (Dunn’s multiple comparison test after the Kruskal–Wallis test).
3.2. The distribution of three subgroups of GABAergic neurons

The distribution of glutamatergic and GABAergic neurons in each subregion of the claustrum was explored with FISH histochemistry for VGluT1 or GAD67 mRNA combined with immunofluorescence histochemistry for NeuN (Fig. 2). VGluT1-expressing cells accounted for 91.9%, 90.0%, and 87.3% of NeuN-positive cells in the core region and dorsal and ventral shell regions, respectively (Fig. 2A–B3), while GAD67-expressing cells constituted 9.1%, 9.3%, and 11.2% in these subregions (Fig. 2C–D3). Further, no colocalization was found between the signals for VGluT1 and GAD67 mRNAs (no GAD67-expressing cell in 3483 VGluT1-expressing cells; no VGluT1-expressing cell in 344 GAD67-expressing cells; Fig. S2). Although the proportion of GAD67-expressing neurons in the rostral core region was significantly lower than that in the rostral shell region, VGluT1- and GAD67-expressing neurons were almost homogeneously located throughout the claustral subregions (Table 1).

We then investigated the proportion of PV, SOM, and VIP neurons by combining FISH histochemistry for GAD67 mRNA with double immunofluorescence histochemistry for NeuN and one of the neurochemical markers (Fig. 3). PV neurons consisted of 26.2%, 17.1%, and 20.0% of GAD67-expressing neurons in the core region and dorsal and ventral shell regions, respectively. PV neurons were more abundant in the core region, while the proportion of SOM neurons was not significantly different from that of PV neurons in the core region. Moreover, the VIP neuron population was smaller than those of the others throughout the claustrum (p < 0.01 for all pairs, except for p = 0.11 between PV and SOM neurons in the core region; two-way ANOVA followed by Bonferroni’s multiple-comparison test; Table 2).

We further examined whether these three neurochemical markers would show overlap in claustral neurons using quadruple...
immunofluorescence labeling for NeuN, PV, SOM, and VIP neurons in the core region and dorsal and ventral shell regions of the claustrum, respectively (Fig. 4C).

3.3. Restricted distribution of dendrites and axons of PV and SOM neurons in the core and ventral shell regions

Abundant neurites immunopositive for PV were observed in the core region, and those for SOM in the shell region (Fig. 4; Marriott et al., 2021). We analyzed the dendritic and axonal arborizations of PV and SOM neurons in the core and ventral shell regions to determine whether their neurite extension differ between the subregions. The mixture of diluted driver vector (AAV2/1-CMV-tTA-WPRE) and undiluted reporter vector (AAV2/1-TRE-FLEX-palGFP-BGHpA) (Fig. 5A) was injected into the claustrum of PV+/Cre or SOM+/Cre knock-in mice. After visualizing the dendrites and axons, optically cleared brain slices were observed with a confocal microscope to acquire 3D image stacks (Fig. 5B, C). We obtained 3 and 4 neurons in the core and ventral shell regions from 5 hemispheres of PV+/Cre mice (PV-core neurons, #1–3; PV-shell neurons, #4–7; Fig. 5D), and 3 and 5 neurons in the core and ventral shell regions from 8 hemispheres of SOM+/Cre mice (SOM-core neurons, #8–10; SOM-shell neurons, #11–15; Fig. 5D). We reconstructed the dendrites and axons and the claustral structures in 3D image stacks. PV-core #2, PV-shell #6, and PV-shell #7 neurons were derived from a single hemisphere. Although the three neurons extended their neurites within the hemisphere, combining the 1-mm-thick brain slices with a tissue clearing method enabled us to trace their dendrites and axon of each neuron (Fig. S3A). The three neurons had few overlaps (Fig. S3B), and we separated them without including fibers of neighboring labeled neurons. The other 12 neurons were derived from the respective hemispheres, where a single EGFP-labeled neuron was found inside the claustrum and distinguished from other labeled neurons outside the claustrum (Fig. 6).

We then calculated the proportions of dendrite and axon lengths distributed in each region (Table 4). The fluorescence images of PV-core neurons showed that their neurites were mainly confined within the core region (Fig. 6A1). After 3D reconstruction of their neurites, we observed 99.1% of dendrites and 99.8% of axons inside the claustrum. The reconstructed neurites also showed that the arborization fields of dendrites slightly exceeded those of axons. Indeed, we found that 76.0% of all dendrites and 92.8% of all axons remained within the core region, while 23.0% and 7.0% extended to the dorsal and ventral shell regions, respectively (Fig. 6A2; Table 4). Similarly, in the fluorescence images of SOM-core neurons, their neurites appeared to be closely packed in the claustrum and distinguished from other labeled neurons outside the claustrum (Fig. 6).

In contrast, the fluorescence images of PV-shell neurons showed that their neurites mainly existed in the dorsal and ventral shell regions and avoided sending axons to the core regions (Fig. 6C1). PV-shell neurons spread 79.0% of dendrites and 85.7% of axons inside the claustrum. We further found that 75.1% of dendrites and 73.6% of axons existed in the dorsal and ventral shell regions, while only 3.9% of dendrites and 12.1% of axons entered the core region (Fig. 6C2; Table 4). SOM-shell neurons spread 79.4% and 84.4% of dendrites and axons and formed arborizations inside the claustrum, respectively, and that 79.4% and 84.4% were located within the core region, respectively (Fig. 6B2; Table 4). These results indicate that the dendrites and axons of both PV-core and SOM-core neurons are mostly confined within the core region.

We obtained 3 and 4 neurons in the core and ventral shell regions from 5 hemispheres of PV+/Cre mice (PV-core neurons, #1–3; PV-shell neurons, #4–7; Fig. 5D), and 3 and 5 neurons in the core and ventral shell regions from 8 hemispheres of SOM+/Cre mice (SOM-core neurons, #8–10; SOM-shell neurons, #11–15; Fig. 5D). We reconstructed the dendrites and axons and the claustral structures in 3D image stacks. PV-core #2, PV-shell #6, and PV-shell #7 neurons were derived from a single hemisphere. Although the three neurons extended their neurites within the hemisphere, combining the 1-mm-thick brain slices with a tissue clearing method enabled us to trace their dendrites and axon of each neuron (Fig. S3A). The three neurons had few overlaps (Fig. S3B), and we separated them without including fibers of neighboring labeled neurons. The other 12 neurons were derived from the respective hemispheres, where a single EGFP-labeled neuron was found inside the claustrum and distinguished from other labeled neurons outside the claustrum (Fig. 6).

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Almost all neurites of PV-core and SOM-core neurons were detected inside the claustrum, whereas 21.0% and 14.3% of PV-shell neuronal dendrites and axons, respectively, and 25.0% and 35.2% of SOM-shell neuronal dendrites and axons, respectively, were distributed outside
the claustrum. The dendrites and axons of PV-shell neurons outside the claustrum were observed in the areas adjacent to the claustrum such as the insular cortex (dendrites, PV-shell #4, 6, 7; axons, #4, 5, 6, 7; Fig. 6C2 and Fig. S4) and dorsal endopiriform nucleus (DEn; dendrites, PV-shell #5, 7; axons, #4, 5, 6, 7; Fig. 6C2 and Fig. S4). The dendrites or axons of SOM-shell neurons were also found in the insular cortex (dendrites, SOM-shell #12, 13, 15; axons, #11, 12, 13, 14, 15; Fig. 6D2 and Fig. S4) and DEn (dendrites, SOM-shell #11, 12, 15; axons, #11, 12, 14, 15; Fig. 6D2 and Fig. S4). Additionally, a few axon collaterals of a SOM-shell neuron were detected in somatosensory and motor cortical areas (SOM-shell #15; Fig. S4). Considering that almost all SOM neurons showed the signals for GAD67 mRNA and that a population of GABAergic projection neurons is known to be positive for SOM in the cerebral cortices (Tomioka et al., 2005), this SOM-shell neuron may correspond to a long-range projecting inhibitory neuron previously reported in the claustrum (Shelton et al., 2022).

### Table 3

|                | Core region                  | Shell region                  | Ventral part |
|----------------|------------------------------|------------------------------|--------------|
|                | Dorsal part                  |                               | Ventral part |
| PV / SOM       | 3.0 ± 1.1 \(^a\) (8/279)     | 1.1 ± 0.9 (3/274)             | 0.5 ± 0.5 (3/604) |
| SOM / PV       | 3.0 ± 0.6 (8/269)            | 1.4 ± 1.2 (3/204)            | 0.8 ± 0.8 (3/394) |
| SOM / VIP      | 0 (0/65)                     | 0 (0/65)                     | 0.9 ± 1.6 (1/96) |
| VIP / SOM      | 0 (0/279)                    | 0 (0/274)                    | 0.2 ± 0.3 (1/604) |
| VIP / PV       | 0 (0/269)                    | 0 (0/204)                    | 0.0 (0/394) |
| PV / VIP       | 0 (0/65)                     | 0 (0/65)                     | 0.0 (0/96) |

\(^{a}\)Data are provided as mean ± SD of the percentages in three mice.

The denominator and numerator in parentheses correspond to the total numbers of the neurons immunoreactive for PV, SOM, or VIP.

The SOM-shell neuron were detected in somatosensory and motor cortical areas (SOM-shell #15; Fig. S4). Considering that almost all SOM neurons showed the signals for GAD67 mRNA and that a population of GABAergic projection neurons is known to be positive for SOM in the cerebral cortices (Tomioka et al., 2005), this SOM-shell neuron may correspond to a long-range projecting inhibitory neuron previously reported in the claustrum (Shelton et al., 2022).

### 3.4. Directional preferences in the dendritic and axonal arborizations of PV and SOM neurons in the core and ventral shell regions

We then quantified the ramifications of three-dimensionally reconstructed dendrites and axons of PV and SOM neurons in the core and ventral shell regions by counting the number of intersections between neuronal processes and concentric spheres centered at the soma (3D Sholl analysis) (Sholl, 1953). Subsequent statistical multiple comparisons were conducted for dendrites or axons among PV-core, PV-shell, SOM-core, and SOM-shell neurons (Fig. 7). The number of dendritic ramifications was not significantly different between PV-core, SOM-core, and PV-shell neurons and between SOM-core and SOM-shell neurons. We found that the dendrites of PV-core neurons were more highly ramified than those of SOM-core neurons at 50–250 µm. We also observed that PV-shell neurons had more extensively ramified dendrites than...
SOM-shell neurons at 50–300 µm (Fig. 7A). These results indicate that the dendritic arborizations of PV and SOM neurons are not different between the core and ventral shell regions and that PV neurons have more abundant dendritic branching than SOM neurons.

In 3D Sholl analysis for axons, we found that all four types of neurons had prominent axonal arborizations around 200 µm from the soma and that those axons were widely distributed up to more than 800 µm. The axonal ramifications were similar in PV-core, SOM-core, and PV-shell neurons, whereas those in SOM-shell neurons were sparser at 150–200 µm and more abundant at 400–700 µm (Fig. 7B), suggesting that SOM-shell neurons tend to innervate more distal portions than the other types of neurons.

It was recently reported that intraclaustral connections mediated by glutamatergic neurons are predominant along the RC axis (Orman, 2015; Shelton et al., 2022). To examine whether PV and SOM neurons have such a directional preference, we projected three-dimensionally reconstructed neurons onto a plane along three orthogonal RC, ML, or DV axes (ML–DV plane, DV–ML plane, or RC–DV plane), and quantified the ramifications of dendrites and axons on these three planes (2D Sholl analysis; Fig. 8). The dendritic branching of PV-core and SOM-core neurons were almost similar among the three ML–DV, DV–RC, and RC–ML planes, but the branching in the ML–DV plane was low in the distal portion compared with that in the other planes (Fig. 8A, B, left). The dendrites of PV-shell and SOM-shell neurons also showed similar branching patterns, although their dendrites extending more than 150 µm from the soma...
axons of PV-core and SOM-core neurons on the ML plane were distinct from those in the RC regions. GABAergic neurons, respectively. Furthermore, we demonstrated that for approximately 10% of claustral neurons and that PV, SOM, and VIP axons of SOM-shell neurons were broadly distributed over 800 shell neurons on the ML planes widely extended over 700 µm on all 3 planes, those on the ML–DV plane were significantly lower than those on the DV–RC and ML–ML planes over 300 µm from the soma. (Fig. 8C, D, right). These findings indicate that PV and SOM neurons in the core and ventral shell regions preferentially spread their axons along the RC axis to distal portions.

4. Discussion

In the present study, we showed that GABAergic neurons accounted for approximately 10% of claustral neurons and that PV, SOM, and VIP neurons constituted approximately 20%, 30%, and 10% of claustral GABAergic neurons, respectively. Furthermore, we demonstrated that PV-core and SOM-core neurons confined their dendrites and axons within the subregion, while PV-shell and SOM-shell neurons avoided extending their neurites into the core region.

We divided the mouse claustrum into the core and shell regions based on cytoarchitecture (Fig. 1), referring to the previous studies (Atlan et al., 2017; Binks et al., 2019; Marriott et al., 2021; Ham and Augustine, 2022; Shelton et al., 2022). However, whether the shell region is included in the claustrum remains debatable in rodents. Several studies have argued that the claustrum contains only the core region and that the shell regions belong to the insular cortex (Mathur, 2014; Wang et al., 2017, 2022; Dillingham et al., 2019; Grimstevdt et al., 2022). On the other hand, ISH histochemical and single-cell RNA sequencing (scRNA-seq) analyses demonstrated that the two regions displayed shared gene expressions such as Gnb4, Nr4a2, Synpr, and Gng2 (Watakabe, 2017; Watson and Puelles, 2017; Smith et al., 2019), which distinguish the claustrum from the insular cortex (Binks et al., 2019; Smith et al., 2019; Erwin et al., 2021). Moreover, reciprocal connections with cortical areas are observed in both the core and shell regions (Atlan et al., 2017; Marriott et al., 2021; Ham and Augustine, 2022). We thus considered the shell regions to be a part of the claustrum in the present study.

We demonstrated that 9–11% of claustral neurons expressed GAD67 mRNA, and GABAergic neurons appeared to be uniformly distributed among the claustral subregions (Table 1; Fig. 2). This proportion of GABAergic neurons found in our study is in concordance with those of previous reports of rabbits and humans: 12% of claustral neurons in rabbits displayed immunoreactivity for GABA in the claustrum (Gomez-Urquijo et al., 2000), and 7% of claustral neurons in humans comprised aspiny interneurons and homogeneously distributed throughout the claustrum (Braak and Braak, 1982; Spahn and Braak, 1985). We also found that 88–90% of claustral neurons expressed VGluT1 mRNA. This finding is consistent with the results of a single-molecule FISH study, where VGluT1 and GAD65 mRNAs were expressed in 83% and 4% of DAPI-positive cells in the mouse claustrum, respectively (Atlan et al., 2018). The slight differences in population rates between the present and previous studies might be explained by the differences in total cells counted. In the present study we counted NeuN-immunoreactive cells as total cells, whereas the previous study counted DAPI-positive cells, which were assumed to include both neuronal and glial cells. Although the single-molecule FISH study also reported that a proportion of GAD65-expressing neurons showed the signals for VGluT1 mRNA as well (Atlan et al., 2018), we found that the signals for VGluT1 and GAD67 mRNAs did not colocalize (Fig. 52). The discrepancy between the present and previous studies could be due to the mRNA each study targeted (GAD67 or GAD65). It was also reported that part of claustral neurons expressed VGluT2 or VGluT3 mRNA and that VGluT2-expressing neurons had projections to the cortex (Hur and Zaborszky, 2005; Johnson et al., 2014; Hawrylycz et al., 2015; Xu et al., 2022), suggesting the diversity of glutamatergic neurons as well as GABAergic neurons.

Our immunohistochemical analysis clarified that PV, SOM, and VIP neurons occupied 20%, 30%, and 10% of claustral GABAergic neurons, respectively (Table 2; Fig. 3), and had little overlap with each other (Table 3; Fig. 4). An electrophysiological study demonstrated that PV,

Table 4
Dendritic and axonal lengths of three-dimensionally reconstructed PV and SOM neurons in the claustral core and shell regions.

| Region          | PV-core | SOM-core | PV-shell | SOM-shell |
|-----------------|---------|----------|----------|-----------|
| **Dendritic length** |         |          |          |           |
| Inside claustrum| 99.1 ± 0.6a | 95.7 ± 7.5 | 79.0 ± 13.5 | 74.7 ± 19.6 |
| Core region     | 76.0 ± 5.5 | 79.4 ± 13.9 | 3.9 ± 2.4 | 2.5 ± 2.7 |
| Shell region    | 23.0 ± 5.2 | 16.3 ± 9.5 | 75.1 ± 14.8 | 72.5 ± 19.1 |
| Outside claustrum| 83 ± 0.8 | 4.3 ± 7.5 | 21.0 ± 13.5 | 25.0 ± 19.6 |
| Total           | (8507 ± 938) | (5165 ± 1355) | (8347 ± 1668) | (4505 ± 718) |

| Region          | PV-core | SOM-core | PV-shell | SOM-shell |
|-----------------|---------|----------|----------|-----------|
| **Axonal length** |         |          |          |           |
| Inside claustrum| 99.8 ± 0.3 | 94.6 ± 9.3 | 85.7 ± 12.5 | 64.8 ± 25.9 |
| Core region     | 92.8 ± 6.1 | 84.4 ± 10.5 | 12.1 ± 5.0 | 7.3 ± 3.8 |
| Shell region    | 7.0 ± 5.8 | 10.2 ± 7.9 | 73.6 ± 11.8 | 57.6 ± 24.3 |
| Outside claustrum| 0.2 ± 0.3 | 5.4 ± 9.3 | 14.3 ± 12.5 | 35.2 ± 25.9 |
| Total           | (71,195 ± 11,322) | (64,333 ± 17,463) | (65,078 ± 14,128) | (89,173 ± 34,162) |

aData are provided as mean ± SD of the percentages in three to five reconstructed neurons.

bLengths (µm) of the dendrites and axons are provided as mean ± SD in three to five reconstructed neurons.

were less branched on the RC–ML plane than on the ML–DV and DV–RC planes (Fig. 8C, D, left). These results indicate that PV and SOM neurons distribute the dendrites almost uniformly in the core and ventral shell regions.

In contrast to dendrites, the axonal arborizations on the ML–DV plane were distinct from those in the RC–ML and DV–RC planes. The axons of PV-core and SOM-core neurons on the ML–DV plane were confined to less than 400 µm, whereas those on the DV–RC and RC–ML planes widely extended over 700 µm (Fig. 8A, B, right). The axons of PV-shell neurons on the ML–DV plane were observed mostly within 500 µm and those on the DV–RC and RC–ML planes spread over 700 µm. The axons of SOM-shell neurons were broadly distributed over 800 µm on all 3 planes, but those on the ML–DV plane were significantly lower than those on the DV–RC and ML–ML planes over 300 µm from the soma. (Fig. 8C, D, right). These findings indicate that PV and SOM neurons in the core and ventral shell regions preferentially spread their axons along the RC axis to distal portions.
SOM, and VIP neurons in the mouse claustrum could be distinguished by their intrinsic electrical properties (Graf et al., 2020). These immunohistological and electrophysiological findings indicate that PV, SOM, and VIP neurons form distinct subgroups. The three types of neurons covered approximately 60% of claustral GABAergic neurons, but the residual population remained undetermined in the present study. It has been shown that the claustrum contains various GABAergic neurons immunoreactive for neurochemical markers, such as calbindin, calretinin, cholecystokinin, neuropeptide Y (NPY), and nitric oxide synthase (Eiden et al., 1990; Druga et al., 1993, 2014; Kowianski et al., 2001, 2008; Guirado et al., 2003; Real et al., 2003; Davila et al., 2005; Marriott et al., 2021; Morello et al., 2022). Among them, NPY-positive neurons were reported to comprise a 1.5-fold larger population than SOM neurons in the mouse claustrum and to show considerable overlap with SOM immunoreactivity; 50% of SOM neurons displayed immunoreactivity for NPY, and conversely, 33% of NPY neurons were positive for SOM (Marriott et al., 2021). To clarify the heterogeneity of residual neurons in detail, it would be useful to thoroughly examine the gene expression patterns of claustral GABAergic neurons. Indeed, recent technical advances in scRNA-seq have revealed the transcriptional signatures of glutamatergic neurons in the core and ventral shell regions in mice (Shelton et al., 2022). Integration of transcriptomic, morphological, and electrophysiological analyses would provide clearer insights for the diversity of claustral GABAergic neurons.

The topographic organization of reciprocal connections between the claustrum and cerebral cortex is uniformly elongated across the RC axis in the claustrum (Atlan et al., 2017; Marriott et al., 2021; Ham and Augustine, 2022). Notably, the intrinsic connectivity of claustral glutamatergic neurons also shows similar RC continuity (Smith and Alloway, 2010; Jackson et al., 2020); the axons of claustral projecting neurons traveled along the RC axis inside the claustrum (Wang et al., 2022), and the intraclaustral connections mediated by glutamatergic synapses were prominently observed in this direction (Orman, 2015; Shelton et al., 2022). In the present study, we demonstrated that PV and SOM neurons in the core and ventral shell regions preferentially extended their axons more than 800 µm along the RC direction (Fig. 8). Taken together, the claustrum is suggested to be tightly bounded in the RC direction via both glutamatergic and GABAergic neurons.

In contrast to the RC axis, the topographical arrangement of claustrocortical neurons is found across the DV and/or ML directions (Atlan et al., 2017; Marriott et al., 2021; Ham and Augustine, 2022). Further, the claustral glutamatergic neurons form few axon collaterals inside the claustrum (Wang et al., 2022) and rarely make their intraclaustral connections in the DV and/or ML directions (Orman, 2015; Kim et al., 2016). We here demonstrated that the dendrites and axons of PV and SOM neurons in the core and shell regions were mostly confined within each subregion, but that the neurites were widely developed within each subregion (Fig. 6; Table 4). PV-core and SOM-core neurons extended the dendrites partially to the shell region (23% in PV-core neurons; 16% in SOM-core neurons).
Fig. 8. 2D Sholl analysis for the dendrites and axons of PV and SOM neurons in the claustral core and shell regions. (A–D) 2D Sholl analysis for the dendrites and axons of PV-core (A), SOM-core (B), PV-shell (C), and SOM-shell (D) neurons. The reconstructed data were projected onto the plane perpendicular to either the RC, ML, or DV axis (ML–DV, DV–RC, or RC–ML plane). The mean number of dendritic or axonal intersections on each ML–DV, DV–RC, or RC–ML plane was plotted against the radial distance of concentric circles from the soma. Multiple statistical comparisons for dendrites or axons of each neuronal type were performed among the numbers of intersections on the ML–DV, DV–RC, and RC–ML planes in every 50 µm from the soma with two-way ANOVA followed by post-hoc Bonferroni’s multiple comparison test (*p < 0.05, **p < 0.01, and ***p < 0.001 between the ML–DV and DV–RC planes; †p < 0.05, ††p < 0.01, and †††p < 0.001 between the ML–DV and RC–ML planes; §p < 0.05, §§p < 0.01, and §§§p < 0.001 between the DV–RC and RC–ML planes). The mean number of intersections (solid line) and its SD (shaded area) on the ML–DV, RC–ML, or DV–RC plane are shown in magenta, green, or blue, respectively. The axonal ramifications of these neurons on the ML–DV plane were skewed to the closer areas from the soma than those on the DV–RC and RC–ML planes.
SOM-core neurons; Table 4). Because the topographical arrangement between the claustrum and cerebral cortex is reported to overlap in part across the core and shell regions (Marriott et al., 2021; Shelton et al., 2022), the intersections of the dendrites of PV-core and SOM-core neurons with the shell region might reflect the gradual transitions of the topography within the claustrum. Furthermore, 3D and 2D Sholl analyses revealed that these neurons extended their dendrites almost uniformly, and that the axons densely spread along not only the RC axis but also DV and ML axes. Previous electron microscopic and electrophysiological studies have shown that claustral GABAergic neurons receive synaptic inputs from corticoclausal, claustrocortical, and GABAergic neurons (LeVay and Sherk, 1981; Kim et al., 2016; Day-Brown et al., 2017; White et al., 2018). It is therefore suggested that claustral PV and SOM neurons integrate multiple neuronal information within the core and shell regions but not between those two subregions.

As indicated by a previous report (Marriott et al., 2021), we observed that neuronal fibers immunoreactive for PV were densely distributed in the core region, whereas those for SOM were in the shell region (Fig. 4). As the cell bodies of PV and SOM neurons were uniformly distributed in the core and shell regions, respectively, we hypothesized that the neurite extensions of PV and SOM neurons might be localized within the core and shell regions, respectively. After 3D reconstructions of PV and SOM neurons, we noticed that neither of the neurons showed substantial differences in the spread of neurites between the core and shell regions. Thus, the particular distribution of immunoreactivities for PV and SOM in the core and shell regions might be explained by other factors. One possibility is that the claustral subregions receive the axons containing these proteins from the outside areas of the claustrum. A population of neocortical pyramidal neurons in layer V and thalamic relay neurons are known to express PV protein (Tanahira et al., 2009). It is also reported that some SOM neurons send long-range projections in the cortical and subcortical areas (Tomioka et al., 2015; Eyre and Bartos, 2019; Xiao et al., 2021). It could be possible that the extrinsic axonal fibers containing PV protein and SOM peptide innervate the core and shell regions, respectively, and contribute to the differential immunoreactivities within the claustral subregions. There is another possibility that claustral GABAergic neurons in the core and shell subregions show differential expression levels of PV protein and SOM peptide. Our neuronal labeling experiments with virus vectors demonstrated the neuronal fibers of PV neurons in the shell region and those of SOM neurons in the core region (Fig. 6). Immunofluorescence labeling also detected the neuronal fibers immunoreactive for PV in the shell region, although these immunoreactivities were weaker than in the core region, whereas those for SOM were in the shell region (Fig. 4). Therefore, the amount and/or localization of PV protein and SOM peptide might be different between the GABAergic neurons in the core and shell regions. The expression level of PV protein is reported to be affected by synaptic activities (Patz et al., 2004; Donato et al., 2013). SOM peptide work as a neurotransmitter and is involved in synaptic plasticity through its receptors (Liguz-Lecznar et al., 2016). Comparison of the expression levels of PV protein and SOM peptide between the core and shell regions with scRNA-seq and/or reverse transcription quantitative PCR would be beneficial for further understanding the role of these neurons in the claustral core and shell organization.

Conclusions

The present study showed that claustral GABAergic neurons are homogeneously distributed and that PV and SOM neurons are compartmentalized in the claustral subregions. Claustral PV and SOM neurons might participate in independent information processing within the local circuits of the core and shell regions.

CRediT authorship contribution statement

Megumu Takahashi, Tadashi Isa, Hiroyuki Hioki: Conceptualization. Megumu Takahashi, Kenta Yamauchi, Shinichiro Okamoto, Kazuki Okamoto, Hiro Yuko Hioki: Methodology. Megumu Takahashi, Tomoyo Kobayashi, Haruhi Mizuma, Hiroyuki Hioki: Investigation. Yoko Ishida, Masahiko Watanabe: Resources. Megumu Takahashi, Hiroyuki Hioki: Writing – original draft. Megumu Takahashi, Tomoyo Kobayashi, Haruhi Mizuma, Kenta Yamauchi, Shinichiro Okamoto, Kazuki Okamoto, Yoko Ishida, Masato Koike, Masahiko Watanabe, Tadashi Isa, Hiroyuki Hioki: Writing – review & editing. Hiroyuki Hioki: Project administration. Megumu Takahashi, Kenta Yamauchi, Kazuki Okamoto, Masato Koike, Masahiko Watanabe, Tadashi Isa, Hiroyuki Hioki: Funding acquisition. All authors have approved the final article.

Declarations of interest

None.

Data availability

Data will be made available on request.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.neures.2022.11.008.

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Glossary

2D: two dimensional
3D: three dimensional
AAV: adeno-associated virus
AF: Alexa Fluor
AI: agranular insular cortex
AQP3: 3-amino-propyltriethoxysilane
BGHpA: polyadenylation signal derived from the bovine growth hormone gene
BSA: bovine serum albumin
BT: biotinylated tyramide
CMV: cytomegalovirus
DAF: 4′,6-Diamidino-2-phenylindole
DG: dysgranular insular cortex
DIO: digoxigenin
DV: dorsoventral
FISH: fluorescence in situ hybridization
FITC: fluorescein
FLEX: flip-excision
FT: Fluorochromized tyramide
GAD67: glutamic acid decarboxylase 67 kDa isofrom
GO: glucose oxidase
ML: mediotectal
Nsl: neuronal nuclei
NPIV: neuropeptide Y
pALGFP: EGFP with a palmitoylation signal derived from the GAP-43 N-terminus
PC: rostrocaudal
PCR: sodium-queue: single-cell RNA sequencing
SDCM: spinning disk confocal microscope
SOM: somatostatin
TREGR: Tet-responsive promoter with modified Tet response element
tTAD: improved version of a tetracycline-controlled transactivator
VGLUT1: vesicular glutamate transporter 1
VIP: vasoactive intestinal polypeptide
WPRE: woodchuck hepatitis virus posttranscriptional regulatory element