Clastrum: a case for directional, excitatory, intrinsic connectivity in the rat

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Abstract  Clastrum, a gray matter structure that underlies the neocortex, is reciprocally connected with many neocortical and limbic cortical areas. This connectivity positions clastrum ideally for the integration or coordination of widespread cortical activity. In anatomical studies using multiple planes of section, clastrum has distinct subregions based on latexin immunohistochemistry, and an approximately rostro-caudal alignment of fusiform cells supporting a laminar intrinsic organization. Physiological studies of clastral connectivity in disinhibited brain slices demonstrate (1) intrinsic connectivity sufficient to generate spontaneous synchronized burst discharges, (2) activity spread within the oblique laminae that contained the principal cellular axis, and (3) segregation of activity as evidenced by the absence of spread within coronal planes. Activity spread depended on glutamatergic synaptic transmission, and activity restrictions did not depend on inhibitory circuits. We conclude that the clastrum has an intrinsic excitatory connectivity that is constrained in approximately rostro-caudal laminae, with minimal cross-communication between laminae. Further, clastrum has the intrinsic capability of generating synchronized population activity and facilitating its spread within laminae, a feature that may contribute to seizure generation and spread.

Keywords  Clastrum · Glutamate · Excitatory synapse · Epilepsy

Introduction

Clastrum is a sheet-like subcortical region that is literally sandwiched between the external and extreme capsules in primate brains, and located just beneath the neocortex (above the external capsule) in lower mammalian brains ([1]; for reviews: [2–6]). Anatomical data show cortico-clastral and clausto-cortical circuits involving most sensory neocortical regions. Clastrum has reciprocal projections to a large number of cortical areas, including visual [7–12], motor [9, 11, 13–16], somatosensory [9, 11, 14, 15, 17], auditory [11, 14, 17, 18], olfactory [13, 19], gustatory [19], prefrontal [20], and limbic cortices [13, 21–25]. In general, the part of clastrum projecting to a cortical region is larger than the part of clastrum receiving inputs from that region [26]. There are “zones” of clastrum associated with particular cortical regions. Most notably, somatosensory cortices are associated with the dorsal-most parts of clastrum, with cells running in a largely horizontal stripe through the structure. Auditory and visual cortices are associated with a clastral zone that is ventral to the zone for the somatosensory cortex [11, 27–33]. Clastral neurons in distinct zones, while having large receptive fields, were responsive to particular sensory modalities (visual, somatosensory, or auditory) [27]. Largely as a result of its rich reciprocal connectivity with the neocortex, functional speculation has been dominated by the notion that clastrum is perfectly positioned to coordinate activity in widely separated cortical areas associated with different sensory modalities [34, 35] (see also reviews above).
A majority of the claustro-cortical projection is excitatory. Asymmetrical claustro-cortical synapses consistent with excitatory synapses have been seen in electron micrographs [10]. Some of the claustro-cortical projection was shown using tritiated-aspartate labeling to be an excitatory glutamatergic projection [36], but not all of it. A portion of the projection to visual cortex was found to be non-glutamatergic (projection cells stained for nitric oxide), but this was still considered to be an excitatory projection because 24/29 visual cortical cells decreased their firing when nitric oxide (NO) was inhibited [37]. The inhibitor used by these investigators, L-NG-nitro arginine, is effective against constituitive (neuronal) nitric oxide synthase (NOS), but also against endothelial and inducible forms.

Some of the claustral outflow is also inhibitory. Large aspyin cells are known to be projections cells [38], and a fraction of visual cortical neurons (4/29) increased their firing when NO was inhibited, suggesting an inhibitory NOS-positive projection from claustrum [37]. A number of claustral NOS-positive cells were shown to be inhibitory neurons [39]. From a functional standpoint, a number of details of the cellular targets of the claustro-cortical projection remain unknown (i.e., a glutamatergic projection may be functionally inhibitory by terminating on inhibitory cortical neurons; shown in [10]).

On the basis of Golgi staining or retrograde tracer labeling, there are basically three cell types based on dendritic spininess (medium-sized spiny, medium-sized aspyin, and small aspyin cells) and on whether cells have projection axons that could be identified to leave claustrum [38, 40, 41]. Each of these groups can be subdivided or sorted into alternative groups based on soma shape, patterns of dendritic branching, or immunocytochemistry (e.g., [42, 43]). It has been estimated that about 12 % of claustral neurons are GABAergic [44, 45]. These have been subdivided on the basis of immunocytochemistry for parvalbumin, calbindin, calretinin, NOS activity, and assorted peptides (e.g., [43, 45–49]).

The intrinsic organization of claustral neurons has been more difficult to study because of the challenges imposed by its curved, slender shape. In preliminary studies, we discovered in rats that the dendrites of many claustral cells were aligned in an oblique plane off the horizontal, parallel to the rhinal fissure. This suggested that an oblique plane of section would preserve the whole claustral cell structure and permit a more complete examination of intrinsic claustral connectivity. We used latexin immunohistochemistry [50, 51] to stain claustral cells and tested different brain slice planes in studies of claustral cell connectivity. Preliminary versions of these data have been presented previously [52, 53].

Methods

Anatomical methods

Animals, sectioning

Male Sprague–Dawley albino rats (4–5 weeks old) were anesthetized with halothane or isoflurane and perfused transcardially with phosphate-buffered saline (PBS) followed by 4 % paraformaldehyde in PBS. Brains were removed, post-fixed for 18 h at 4°C in 4 % paraformaldehyde in PBS, and transferred to 30 % sucrose for cryoprotection. Frozen sections were cut at 40 μm thickness in coronal, horizontal, and oblique planes. The optimal oblique plane orientation for capturing claustral cell dendrites (see “Results”) was tilted anterior-up by 30° off the horizontal to be parallel with the rhinal fissure.

Immunohistochemistry

Adjacent sections were stained with latexin antibodies (gift by Dr. Y. Arimatsu to R. Orman) and NeuN antibodies (EMD Millipore, Temecula, CA, USA). In each case, sections were blocked for 2 h in normal serum of animals in which the secondary antibodies were raised, exposed overnight to primary antibodies at 4°C, incubated with 1 % H2O2 for 20 min, incubated with secondary antibodies (Vector biotinylated secondary antibodies; Vector Laboratories, Burlingame, CA, USA) for 1 h at room temperature, incubated in ABC reagents (Vector) for 30 min, and finally incubated in a peroxide substrate DAB kit (Vector). Protocol details, including standard reagents, were followed according to published kit descriptions from Vector Laboratories or general immunohistochemistry protocols from Abcam (Cambridge, MA, USA; http://www.abcam.com/protocols/immunostaining-paraffin-frozen-free-floating-protocol) and IHC World (Ellicott City, MD, USA; http://www.ihcworld.com/general_IHC.htm). After developing, sections were mounted on tissue-tack slides (Polyscience, Niles, IL, USA), air-dried, dehydrated, and cover-slipped.

Microscopy

Slides were viewed and photographed using (1) a plan-apochromatic stereomicroscope (M80; Leica, Bannockburn, IL, USA) with high-resolution digital camera (Leica IC80 HD; Leica; full-screen image capture 2048 × 1536 pixels, 3.1 megapixels; courtesy of Dr. Richard Kollmar) and/or (2) an upright compound microscope (Jenaval; Carl Zeiss) with motorized stage and focus control, full set of objectives, high resolution camera (1600 × 1200;
Optronics Microfire; Optronics, Goleta, GA, USA), and software (Neurolucida 10; MBF Bioscience, Williston, VT, USA).

**Brain slice physiology**

*Slice preparation and multi-electrode recordings*

Male Sprague–Dawley albino rats (3–5 weeks old) were anesthetized with halothane or isoflurane and decapitated. Each brain was removed from the skull, bisected, and placed briefly in ice-cold artificial CSF. Thick slices of tissue (about 1–2 mm thick) were cut from intact hemispheres in planes of section that were horizontal, coronal, sagittal, or angled plane in between coronal and horizontal. Thin slices (380 μm) were cut using a motorized sectioning system (Leica VT1000S, Leica Biosystems, Buffalo Grove, IL, USA) and transferred to a holding chamber.

From the holding chamber, single slices were placed in the MED64 chamber (Panasonic MED64, Osaka, Japan). The MED64 chamber allows simultaneous extracellular recordings from 64 electrodes (50 μm squares). Each electrode is a platinum black-plated square embedded in the floor of the recording chamber. Inter-electrode distances (center to center) were 150, 300, or 450 μm. Recording electrode impedances are 22 kΩ (at 1 kHz) and each is referred to a single set of 4 reference electrodes in the periphery of the chamber that are electrically tied together. The recording electrodes are arranged in an 8 × 8 array embedded on the bottom of the chamber. The temperature of the MED64 chamber was maintained at 30 °C by warming the perfusate with an inline heater (TC-324B; Warner Instrument, Hamden, CT, USA).

The perfusion solution (1 ml/min) was composed of (in mM): NaCl 125, KCl 2.5–5, CaCl₂ 1.7, MgCl₂ 1.2, NaHCO₃ 26, and glucose 10; pH = 7.4 when exposed to 95 % O₂/5 % CO₂.

Brief stimulating pulses were delivered using platinum-iridium parallel bipolar stimulating electrodes (150 μm tip separation; FHC, Bowdoinham, ME, USA) with <100 kΩ electrode impedances. Stimuli were biphasic pulses (50–100 μs in total duration) applied through constant current stimulus isolation units. The bipolar stimulating electrode was placed from the top side of the slice.

Data were digitized at 20 kHz per channel and stored on disk using MED64 Conductor software. Events were studied offline using MED64 Conductor software or custom Microsoft EXCEL macros.

*Pharmacology*

All drugs were applied to the bath by adding them to the perfusate reservoir. The concentrations given are concentrations that exist in the reservoir and were achieved in the recording chamber over a period of minutes. Recordings in the presence of all drugs were taken after sufficient time for equilibration in the recording chamber. Equilibration was apparent in recordings as a stable change in evoked responses.

Bicuculline (bicuculline methiodide, 50 μM), AP-5 (DL-(-)-2-amino-5-phosphonopentanoic acid, 40 μM), CPP (3-((RS)-2-carboxypiperazin-4-yl)-propyl-1-phosphonic acid, 20 μM), and CNQX (6-cyano-7-nitroquinoxaline-2,3-dione or 6-cyano-7-nitroquinoxaline-2,3-dione disodium, 20 μM) and all other chemicals, unless otherwise specified, were obtained from Sigma (Sigma-Aldrich, St. Louis, MO, USA). Some batches of CNQX and CPP were obtained from Tocris Bioscience (Ellisville, MO, USA). Bicuculline was used to antagonize GABA-A receptors. AP5 and CPP were used as NMDA receptor antagonists, and CNQX was used as an AMPA receptor antagonist. All slices were disinhibited with bicuculline.

*Slice planes*

Three different planes of section were studied in detail. The first plane of section was a standard coronal section. This plane captures dorsal claustrum, endopiriform nucleus (ventral claustrum), and the transitional region between them. The second plane is a standard sagittal section. Some slices cut for physiology will contain dorsal and ventral claustrum in a single slice. The third plane of section is referred to as ‘‘oblique.’’ This slice is an angled horizontal slice where the rostral end is elevated above the caudal end, parallel to the rhinal fissure. A relatively small number of conventional horizontal slices were also tested, but these slices resembled coronal slices in activity spread and were not studied further.

*Statistics*

Calculations of frequency distributions, ANOVA, multiple comparisons, Fisher exact tests, and descriptive statistics were done with Microsoft Excel, IBM SPSS Statistics (v.21), and GraphPad Prism 6 software or online calculators at graphpad.com.

*Results*

*Anatomical findings*

**Latexin immunohistochemistry and regional definitions**

As reported by Arimatsu et al. [51, 54–56], latexin is expressed in cortical and caudal neurons. Adjacent
sections were stained for latexin and NeuN. NeuN is widely used as a cell stain in brain sections and was used here for orientation purposes and relationship of our planes of section to planes of section published elsewhere in the original literature and brain atlases. Based on our latexin staining in multiple planes of section (Fig. 1), we identified three parts of dorsal (insular) claustrum and two parts of the endopiriform nucleus [57]. In coronal sections, latexin-positive neurons and neuropil form the lateral aspect of either region. The latexin-positive region of endopiriform nucleus forms a crescent-shaped zone to a latexin-negative medial core. In the claustrum, the latexin-positive region laterally borders a small dorsal latexin-negative zone and a larger, very densely-stained latexin-positive zone located between the claustral and endopiriform latexin-negative regions (Figs. 1, 2). This very densely-stained, latexin-positive, “egg-shaped” zone is the major part of claustrum that specifically overlaps with every anatomical definition of boundaries, including the major brain atlases, e.g., [58]. G-protein gamma 2 subunit (Gng2) staining [4, 59], and N- and R-cadherin staining [47].

In staining coronal, horizontal, and sagittal series, we determined that an oblique plane of section whose long axis tilted upward on the rostral end, parallel to the rhinal fissure, was optimal for preserving the predominantly bipolar, latexin-positive claustral cells. Their main dendritic shafts extended both anterodorsally and postventrally parallel to an obliquely oriented long axis of the insular claustrum (Fig. 2a). By contrast, in coronal sections, the stained claustral neurons of the principal egg-shaped zone appear as larger circular cross-sections that are set in a dense neuropil and interspersed among densely-stained smaller cross-sections of claustral cell dendrites (Fig. 2b). These findings clearly suggest a laminar organization to claustral cells that runs along the long axis of the structure.

Physiological findings

Synchronous population activity in disinhibited slices

To focus on the excitatory connectivity within claustrum, recordings of spontaneous and evoked activity were taken in the presence of GABA receptor blockade (bicuculline 50 μm unless otherwise indicated). Synchronous population events consisting of an initial population spike followed by multiple secondary population spikes were obtained in claustral recordings (Figs. 3, 4, 6 with specific features described in detail below). Claustral events evoked or triggered by single stimulus pulses applied to the overlying neocortex were identical in population spike morphology and spatial distribution to spontaneous claustral events identified as claustral in origin by the identification of the first site in the slice showing activity.

Spatial extent of activity spread within claustrum

One of the most dramatic findings was that the oblique slice plane captured synchronous population activity on the largest number of electrodes. As a measure of the extent of synchronous activity spread, we determined the number of electrode channels in claustrum with synchronous spiking divided by the total number of electrode channels in claustrum for each slice. Oblique slices captured larger amounts of claustral tissue and therefore had the largest number of electrode channels in claustrum (an average of 11 electrodes in claustrum depending on grid spacing compared with an average of 5 electrodes for coronal and sagittal slices; see Figs. 3, 4). In 16/33 oblique slices, every electrode in claustrum showed synchronous population activity (Fig. 3). Coronal slices were at the other extreme, with 62 of 65 slices showing activity on 0 or 1 electrode (Fig. 4). Sagittal slices were inbetween, showing statistically larger fractions than coronal slices, but statistically smaller fractions than oblique slices. The mean fractions (±SEM) for each group were: coronal 5.5 ± 1.8, oblique 63 ± 6.9, and sagittal 28 ± 7.5 % of claustral electrodes showing synchronous spiking. Median fractions were 0, 75, and 20 for coronal, oblique, and sagittal slices, and the 75th percentile was 0, 100, and 50 for the three groups. These data are summarized in Fig. 5.

An ANOVA showed the differences for the dataset to be highly significant (F2,114 = 49.81; p < 0.0001) as well as the multiple comparisons (cor|sag, p = 0.0021; cor|obl, p < 0.0001; sag|obl, p < 0.0001; Holm-Sidak’s multiple comparisons tests). Comparing the groups based on the
presence of synchronous spiking on ≤1 electrode or >1 electrode, the distributions were all statistically different from each other: coronal 62, 3, oblique 8, 25, and sagittal 12, 7 (two-tailed Fisher exact tests: cor|sag, \( p = 0.0009 \); cor|obl, \( p < 0.0001 \); sag|obl, \( p = 0.0081 \)).

We note that the statistical significance is underestimated because our calculations of fractional involvement are extremely conservative. The numbers of electrodes in claustrum in coronal and sagittal slices were smaller, averaging 5 electrodes in claustrum (ranges: 2–11 coronal, 3–7 sagittal), compared with oblique slices, which averaged 11 electrodes in claustrum (range: 4–16). This caused higher fractions with fewer electrodes in coronal and sagittal slices, and lower fractions with more electrodes in the oblique
slices. The average number of electrodes in claustrum for each of the three planes of section was: coronal 5 (range 2–11), oblique 11 (range 4–16), and sagittal 5 (range 3–7).

Given the overall shape of claustrum, i.e., a thin sheet-like structure whose thinnest dimension is along the mediolateral axis, sagittal slices are the most likely to be variable with regard to the amount of claustral tissue captured in a slice. We believe this accounts for the intermediate level of fractional activity in the slices. Whereas oblique slices capture both the principal dendritic axis of claustral cells and long stretches of claustral tissue over multiple sections, sagittal slices capture the dendritic axis, but only offer one or two slices per hemisphere with adequate claustral tissue to study intrinsic connectivity. Coronal slices cut the dendritic axes nearly perpendicularly, and, although many sections contain claustral tissue, intrinsic connectivity is largely abolished.

Spread rate and directional constraints within claustrum

A hallmark of cell-to-cell connectivity among excitatory cells is the generation spontaneous population bursts when inhibitory synaptic transmission is blocked, and the spread of these events over distances within the structure. As described above, synchronous population events that spread over large distances in claustrum were seen in the oblique slices, less commonly in the sagittal slices, and essentially not at all in the coronal slices. In oblique slices that captured long lengths of the densely latexin-positive staining, the spread rate for spontaneous or evoked population events was 0.07–0.21 m/s (mean velocity = 0.13 ± 0.06 m/s; \( R^2 \) values from plots of distance vs. time to determine slope ranged from 0.69 to 0.98). An example is illustrated in Fig. 3c, d. The spread velocity is similar to what has been reported for hippocampus [60].

Slices taken ventral to the principal claustral subregion contain the transitional zone between dorsal claustrum and endopiriform nucleus or principally endopiriform nucleus, and were therefore avoided. It is interesting that slices taken from claustrum dorsal to the cell-dense region showed similar spread extent and spread velocities—in fact, the spatial extent was often greater in these slices because the spatial extent of claustrum was greater. This indicates that the cell-dense region and the more dorsal, less-dense region are functionally similar with regard to intrinsic excitatory connectivity.

Pharmacology of synchronous activity

As bicuculline levels in the slice bath equilibrate, evoked responses increase in amplitude and the early population firing is eventually followed by repetitive spiking (Figs. 3, 4). The number of these late repetitive spikes and their amplitudes both increase during bicuculline equilibration.

Bath application of either high-dose calcium (8 mM) to reduce polysynaptic activity and/or the glutamate receptor antagonist, CNQX, simultaneously eliminate synchronous activity and its spread. High-dose calcium eliminated repetitive population spiking and exposed the typical broadening of propagated population spikes that were smaller, but still all-or-none (Fig. 6). CNQX eliminated responses completely, degrading even the initial population spike and underlying population excitatory postsynaptic potential, until only the stimulus artifact remained (data not shown). Based on these findings, we conclude that the synchronization of cellular activity to support spontaneous population events and their spread are mediated by excitatory glutamatergic synapses.

![Fig. 2](image)

**Fig. 2** High-magnification images of latexin-positive claustral cells in sections that capture the dendrites of fusiform cells or cut them in cross-section. Forty-times magnification of latexin staining from an oblique section (a comparable to d in Fig. 1) and from a coronal section (b comparable to a in Fig. 1). White arrows in (a) highlight fusiform cell bodies and the dendrites of these bipolar neurons. In (b), a cross-section of a cell boy is marked by a black arrow and cross-sections of dendritic elements are highlighted with a black circle. Scale bar 20 μm
Latexin immunohistochemistry and brain slices cut in various planes were used to demonstrate an oblique, rostro-caudally directed organization of bipolar cells, with highly cell-dense and less dense subregions of the rat dorsal claustrum. This intrinsic cellular organization included excitatory cell-to-cell connectivity that, in disinhibited slices, supported the spontaneous generation of synchronized population events that spread along the oblique, rostro-caudal axis. Whereas excitatory connectivity was responsible for synchronous population activity within oblique laminae, segregation of laminar activity was independent of inhibition. These findings suggest modality specificity [31, 32, 61] without cross-talk between modalities.

Significance of dendritic organization and intrinsic connectivity

The directional spread without broadening of activity within claustrum indicates several important features of the intrinsic connectivity. Spread was constrained within the oblique rostro-caudal dimension, and was minimal in coronal and sagittal planes, suggesting preferential modality specific intrinsic communication with minimal interaction across modalities. The absence of broadening indicates that the full network of claustral neurons at each point along the propagation path was being activated [60, 62] (see also Fig. 6), suggesting that the addition of inhibitory circuitry would constrain activity further.

If the claustral sensory modalities are organized in a laminar fashion, as evidenced by Morys, Narkiewicz, and others [31, 32, 61], our data from disinhibited slices suggest that only intra-laminar (within modality interactions), but not inter-laminar (interactions across modalities) interactions occur within claustrum. Inhibitory circuits are expected to refine the intra-laminar connectivity, but are not the basis for inter-laminar segregation. The independence from inhibition for segregation is in contrast to other structures that depend highly on inhibitory circuits to compartmentalize activity (e.g., barrel cortex [63]).
all neuronal theories of consciousness is the need for continuous interactions among groups of widely dispersed pyramidal neurons …” “It is clear that the claustrum lies at the confluence of a large number of simple loops with cortex. This widespread and reciprocal connectivity with many, if not most, cortical regions raises the obvious question: why is all this information brought together?” “… If the claustrum is critical to binding information within and across sensory and motor modalities … there must be some sort of intermixing of the associated signals within the claustrum” [34]. The results of this paper suggest that claustrum is well-suited to the broad distribution of activity with a claustral lamina or modality, but that other mechanisms must exist for binding the activity of different laminae or modalities. Clearly, many questions remain unanswered.
Latexin binding and claustrum

The original reports of latexin staining by Arimatsu et al. [50, 51, 54–56] were focused on neocortical cell development and specialization. Interestingly, claustrum was clearly well stained and was noted, but not emphasized in any of the reports. Here, we corroborate latexin staining as a useful marker to identify both claustrum and endopiriform nucleus, support subdivisions based on the density of staining, and permit examination of cellular morphology of latexin-positive neurons. The latexin-positive, “egg-shaped” (in coronal sections) claustrum is well-aligned with current anatomical definitions reviewed by Mathur [4].

Potential for seizure generalization and altered consciousness

A central, potentially integrative position held by claustrum is also part of the basis for suggesting a role for claustrum in seizure generalization, and an intra-laminar spread of activity within claustrum could be sufficient for such a role. One of the earliest suggestions for claustrum’s involvement in seizure generation came long before its anatomical relations with neocortex were fully defined. A female epileptic patient was found to have an “enormous” claustrum bilaterally [64].

Claustrum stimulation is efficient for kindling seizure activity [13, 65, 66]. Bilateral claustral lesions impair kindling when stimuli are applied to the amygdala [67], since claustrum sits as a relay structure between amygdala and neocortical regions [68, 69]. Unilateral lesions have complex effects [70, 71] or no effect at all [72]. Seizure activity can be triggered by convulsant application to endopiriform nucleus (ventral claustrum) and prevented by application of glutamate receptor antagonists or GABA to endopiriform nucleus [73–75].

An interesting behavior in endopiriform nucleus neuron studies is the “build-up” of activity in the nucleus before a synchronous epileptiform burst occurs [76, 77], clearly consistent with endopiriform cells exciting one another via their local connectivity [78–80]. In fact, Demir et al. have argued that the circuitry to amplify and sustain excitation is separate from the circuitry that generates the actual epileptiform discharge [81]. Whereas endopiriform can certainly be the origin for synchronized burst discharges,
we find that claustrum is capable of generating such activity completely independently.

A particularly interesting clinical correlate offering potential insights into claustral function is aura [82]. Auras or paroxysmal psychic automatons are experiences characterized by sudden onset, automatic development, vividness, and a clear sense of strangeness to the individual. These are interpreted as partial seizures (reviewed in [83]), a fact reflected in current definitions, for example: “a subjective ictal phenomenon that in a given patient may precede an observable seizure, if alone, constitutes a sensory seizure.” Simple and complex aura experiences have been described, particularly in relation to temporal lobe seizures [84, 85].

Does claustrum cause auras or does claustral activity limit a focal seizure so that only an aura occurs? One possibility is that claustrum becomes overactive and transmits its activity to neocortex or limbic cortex [82]. This means that claustrum must possess the capacity to become overactive on its own, a feature we demonstrate here. Also, activity is expected to spread within a lamina or modality without activation of other modal-specific claustral laminae. The data described in this paper highlight the importance of an intrinsic excitatory glutamatergic network of connections within claustrum, but glutamatergic connections also mediate claustro-cortical and cortico-claustral activity spread. The roles of specific claustral cell types and the contributions of transmitter molecules such as nitric oxide remain poorly understood.

An alternative role for claustrum is that it acts to limit a developing seizure that originates in cortex (i.e., the output of claustrum is inhibitory). Consistent with a regulatory or suppressor role for claustrum are data from SPECT studies of seizure patients with auras, suggesting a “zone of suppression” in cortex during auras [86]. Such a function could be mediated by a direct inhibitory output from claustrum to cortex, excitatory claustral projections whose targets are cortical interneurons, or a local claustral activity change that suppresses activity spread within claustrum itself. Here again, the roles of specific claustral cell types and the contributions of transmitter molecules such as GABA and nitric oxide remain poorly understood.

These are complex questions that will be challenging to address given stimulation and recording limitations imposed by claustrum’s size and shape, and the importance of fully capturing its cells in preparations for physiological study.

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Compliance with ethical standards

Conflict of interest None.

Research involving human participants and/or animals This research conforms to the standards set forth in the NIH Guide for the Care and Use of Laboratory Animals (4th edn., National Academies Press, 2011) and was approved by the Institutional Animal Care and Use Committee under protocols for the preparation and study of acute brain slices and immunohistochemistry.

Informed consent Not applicable.

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