Unique Actions of GABA Arising from Cytoplasmic Chloride Microdomains

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Developmental, cellular, and subcellular variations in the direction of neuronal Cl\(^-\) currents elicited by GABA\(_A\) receptor activation have been frequently reported. We found a corresponding variance in the GABAA receptor reversal potential (EGABA) for synapses originating from individual interneurons onto a single pyramidal cell. These findings suggest a similar heterogeneity in the cytoplasmic intracellular concentration of chloride ([Cl\(^-\)]\(_i\)) in individual dendrites. We determined [Cl\(^-\)]\(_i\) in the murine hippocampus and cerebral cortex of both sexes by (1) two-photon imaging of the Cl\(^-\)-sensitive, ratiometric fluorescent protein SuperClomeleon; (2) Fluorescence Lifetime IMaging (FLIM) of the Cl\(^-\)-sensitive fluorophore MEQ (6-methoxy-N-ethyl-quinolinium); and (3) electrophysiological measurements of EGABA by pressure application of GABA and RuBi-GABA uncaging. Fluorometric and electrophysiological estimates of local [Cl\(^-\)]\(_i\) were highly correlated. [Cl\(^-\)]\(_i\) microdomains persisted after pharmacological inhibition of cation–chloride cotransporters, but were progressively modified after inhibiting the polymerization of the anionic biopolymer actin. These methods collectively demonstrated stable [Cl\(^-\)]\(_i\) microdomains in individual neurons in vitro and in vivo and the role of immobile anions in its stability. Our results highlight the existence of functionally significant neuronal Cl\(^-\) microdomains that modify the impact of GABAergic inputs.

Key words: chloride; chloride microdomains; GABA; GABA\(_A\) receptors; SuperClomeleon; FLIM

Microdomains of varying chloride concentrations in the neuronal cytoplasm are a predictable consequence of the inhomogeneous distribution of anionic polymers such as actin, tubulin, and nucleic acids. Here, we demonstrate the existence and stability of these microdomains, as well as the consequence for GABAergic synaptic signaling: each interneuron produces a postsynaptic GABA\(_A\) response with a unique reversal potential. In individual hippocampal pyramidal cells, the range of GABA\(_A\) reversal potentials evoked by stimulating different interneurons was 20 mV. Some interneurons generated postsynaptic responses in pyramidal cells that reversed at potentials beyond what would be considered purely inhibitory. Cytoplasmic chloride microdomains enable each pyramidal cell to maintain a compendium of unique postsynaptic responses to the activity of individual interneurons.

Significance Statement

Microdomains with mature neuronal Cl\(^-\) transport (Romo-Parra et al., 2008; Tyzio et al., 2008; Table 1). There was no relationship between the number of observations and the value of EGABA (Pearson’s r = −0.13, p = 0.5), suggesting that physiological variance in EGABA rather than inadequate sampling explains the range of results. Such variance could arise from intercellular variance in EGABA or from intracellular (subcellular) variance. Over this wide range of reversal potentials, the effects of GABA\(_A\) receptor (GABA\(_A\)R)-gated currents will vary from inhibition via hyperpolarization of the membrane potential, through shunting inhibition, to excitation mediated by activation of low-threshold calcium currents and partial relief of the magnesium block of NMDA receptors (Doyon et al., 2016). These divergent effects suggest that EGABA must be tightly regulated, although the mechanisms of regulation have not been resolved.
The observed range in \( E_{\text{GABA}} \) could be explained by a few millimolar variation in the cytoplasmic intracellular concentration of chloride (\([\text{Cl}^-]_i\)), the anion with the highest permeability through GABA_ARs (Alfonsa et al., 2015; Doyon et al., 2016; Raimondo et al., 2017). Such differences in \([\text{Cl}^-]_i\) could be maintained by the active transport of Cl\(^-\). For example, KCC2 and NKCC1 are oppositely directed cation–Cl\(^-\) cotransporters (CCTGs) expressed in neurons (Kahle et al., 2015). However, these high-velocity transporters whose ionic equilibrium conditions do not match well with the distribution of observed \( E_{\text{GABA}} \) (Delpire and Staley, 2014), and neither acute (Glykys et al., 2014a; Sato et al., 2017) nor chronic (Goutierre et al., 2019) inhibition of transport produces the predicted changes in \([\text{Cl}^-]_i\) (i.e., a passive distribution that collapses the Cl\(^-\) driving force).

Another possibility is the well established partitioning effects on mobile ions exerted by the distribution of immobile ions (Fatín-Rouge et al., 2003) that, for example, create Gibbs–Donnan effects across semipermeable membranes (Donnan, 1911), in gels (Procter, 1914), and in unlinked biopolymers (Marinsky, 1985), and form the basis of ion exchange technologies such as water softening and ion exchange chromatography (Yamamoto et al., 1988; Helfferich, 1995). Relatively immobile cytoplasmic biopolymers with high and spatially inhomogeneous densities of anionic charge, such as actin, tubulin, and nucleic acids (Gianazza and Righetti, 1980; Sanabria et al., 2006; Janke et al., 2008; Chen et al., 2020) could displace Cl\(^-\) locally (Glykys et al., 2014a, 2017). Recent studies have underscored substantial variance in the subcellular distribution of these anionic biopolymers (Koleski, 2013; Morawski et al., 2015; Gut et al., 2018; Chen et al., 2020), suggesting that if the displacement of Cl\(^-\) underlies the intercellular diversity in \( E_{\text{GABA}} \), there should be a corresponding spatial variance in the subcellular distribution of Cl\(^-\). Indeed, many reports have suggested subcellular variance in \([\text{Cl}^-]_i\) or \( E_{\text{GABA}} \) (Pouzet and Marty, 1999; Berglund et al., 2006; Duebel et al., 2006; Szabadi et al., 2006; Romo-Parra et al., 2008; Földy et al., 2010; Glykys et al., 2014a; Astorga et al., 2015; Untiet et al., 2016; Zorrilla de San Martin et al., 2017; Schmidt et al., 2018) for which several mechanisms have been proposed (Khirug et al., 2008; Földy et al., 2010; Glykys et al., 2014a).

We report a subcellular variance in \( E_{\text{GABA}} \) (Fig. 1) that fits the published range of \( E_{\text{GABA}} \) values summarized in Table 1, and we demonstrate a corresponding variance in local neuronal \([\text{Cl}^-]_i\), in vivo and in vitro. Several complimentary techniques were used including two-photon microscopy of the ratiometric Cl\(^-\) indicator SuperClomelon (sCLM; Grimley et al., 2013); Fluorescence Lifetime Imaging (FLIM) of Cl\(^-\)-sensitive, pH-insensitive dye 6-methoxy-N-ethylquinolinium (MEQ; Biwersi and Verkman, 1991); and simultaneous direct measurement of the reversal potential of locally activated GABA_A-gated membrane currents by puffing or uncaging GABA. We assessed the range, stability, transport dependence, and influence of anionic macromolecules on the observed variance in local baseline \([\text{Cl}^-]_i\). Our results highlight the existence of functionally significant neuronal Cl microdomains that modify the impact of GABAergic inputs.

### Materials and Methods

#### Ethics statement

All experiments were performed in accordance with protocols approved by the Center for Comparative Medicine at Massachusetts General Hospital and in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

#### Mice

**Construction of the sCLM targeting vector.** The STKI-HR gene-targeting vector was constructed from 129Sv mouse genomic DNA (genOway). The final targeting vector has the following features: (1) iso- generic with 129Sv ES cells, favoring the homologous recombination; (2) asymmetrical homology arms (5′ short arm-SA: 1 kb, 3′ long arm-LA: 4.3 kb); (3) transgenic cassette containing the sCLM-pA elements allowing the expression of the transgene under the control of the ubiquitous CAAG promoter sequences; (4) Loxp-flanked, combined STOP-neomycin selection (Neo) cassette allowing its excision through the action of Cre-recombinase, activating the sCLM overexpression; and (5) the presence of the diphtheria toxin A (DTA)-negative selection marker reduces the isolation of nonhomologous recombined embryonic stem (ES) cell clones (Fig. 2).

**Production of sCLM targeted ES cell clones.** Linearized targeting vector was transfected into 129Sv ES cells (genOway) according to genOway electroporation procedures. PCR, Southern blot, and sequence analysis of G-418-resistant ES cell clones revealed the recombined locus in both ends of the sCLM-targeting vector. The initial sCLM-targeted ES cell clones were used to generate stable ES cell lines expressing the Cre recombinase, activating the sCLM overexpression; and introducing a corresponding variance in local neuronal \([\text{Cl}^-]_i\) given the published range of \( E_{\text{GABA}} \) values summarized in Table 1. A total of 10 sCLM-targeted ES cell clones were generated by PCR and Southern blot. PCR primers hybridizing within the neomycin cassette (5′-AGAGCG AAAAGGGCAAGCATCTTCC-3′) and a reverse primer hybridizing within the neomycin cassette (5′-CGGTAGGAAGAGTACACCAAGATCC-3′). Two Southern blot assays were hybridized with an internal and an external probe to assess recombination accuracy at the respective 5′ and 3′ ends of the sCLM locus. The absence of off-target mutations was confirmed by sequence analysis.

**Generation of chimeric mice and breeding scheme.** Recombined ES cell clones microinjected into C57BL/6 blastocysts gave rise to male chimeras with statistically significant ES cell contribution. These chimeras were bred with C57BL/6 mice expressing Cre recombinase to produce the sCLM heterozygous line lacking the neomycin cassette. F1 genotyping was performed by PCR and Southern blot. PCR primers hybridizing upstream (5′-GGGCCAACGGCGTGTTAGTTGTGTC-3′) and downstream (5′-ACAGCTCTCCGCTCTTGTCTCAG-3′) of the neomycin cassette allowed PCR identification of the 3195 bp sCLM allele ampiclon harboring the neomycin cassette, and the 177 bp ampiclon lacking the neomycin cassette. Southern blot hybridization with an external probe allowed identification of the 3.7 kb sCLM allele (Fig. 2B,C).

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<th>Study</th>
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Twenty group means of \( E_{\text{GABA}} \) measurements using gramicidin and cell-attached techniques in age-matched C57BL/6 Synapsin-Cre (for neuronal expression) or C57BL/6 DLX-Cre mice bred with C57BL/6J mice expressing Cre recombinase to produce sCLM-floxed mice with statistically significant ES cell contribution. These chimeras were bred with C57BL/6 mice expressing Cre recombinase to produce the sCLM heterozygous line lacking the neomycin cassette. F1 genotyping was performed by PCR and Southern blot. PCR primers hybridizing upstream (5′-GGGCCAACGGCGTGTTAGTTGTGTC-3′) and downstream (5′-ACAGCTCTCCGCTCTTGTCTCAG-3′) of the neomycin cassette allowed PCR identification of the 3195 bp sCLM allele ampiclon harboring the neomycin cassette, and the 177 bp ampcion lacking the neomycin cassette. Southern blot hybridization with an external probe allowed identification of the 3.7 kb sCLM allele (Fig. 2B,C).
A. An illustration of the designed experiment to test for inhomogeneity of GABAergic signaling in single pyramidal cells (PCs). The pyramidal cell is recorded by whole-cell patch-clamp technique. Neighboring interneurons (In) are stimulated separately by puffing 50 μM glutamate to their cell bodies. Stimulation of those interneurons that are presynaptic to the recorded pyramidal cell evokes GABAergic currents. This is a simplified illustration. In reality, each interneuron makes multiple synapses to the target pyramidal cell. Therefore, the variance of $E_{\text{GABA}}$ at individual synapses may be higher.

B. An example of a recorded pyramidal cell and the stimulated neighboring interneurons (labeled with tdTomato) and the evoked GABA currents with different reversal potentials.

C. Range of $E_{\text{GABA}}$ measured in B. The colors of data points in C match the colors of traces and puff pipettes in B. The black dashed line shows the calculated $E_{\text{GABA}}$ based on the Cl$^-$ concentration inside the recording pipette ($E_{\text{GABA}}$). The puffing pipette was manually returned to the same area by the end of the experiment (puff 1 and puff 8 in B). The difference between the $E_{\text{GABA}}$ evoked by puff 1 and that by puff 8 was $\sim$50 mV, while the time interval between these two puffs was 50 min. Although puff 8 was applied later than puff 1, its evoked $E_{\text{GABA}}$ value was farther away from the pipette $E_{\text{GABA}}$ compared with that from puff 1. These findings argue against the possibility that the Cl$^-$ inside the recording pipette homogenizes the dendritic Cl$^-$.

D. Summary of our experiments ($n = 40$ cells) and 20 separate studies (indicated in Table 1) indicating...
Fluorescence lifetime imaging microscopy (FLIM) experiments were performed on C57BL/6 wild-type (WT) mice. Functional studies of [Cl$^{-}$]i microdomains were performed on DLX-Cre mice to visualize interneurons. Either sex was used for experiments.

**In vivo imaging.** Craniectomies were performed under aseptic conditions, under anesthesia with inhaled isoflurane, and in a stereotaxic frame. The temperature was regulated through a heating pad. The surgical site was sterilized with betadine and isopropyl alcohol, and a 2–3 mm incision was made in the scalp along the midline between the ears. A 5- to 7-mm-diameter hole was drilled in the skull, 1 mm lateral, and posterior to bregma (parietal region) using a high-speed microdrill. Once the craniectomy was completed, a thin round coverglass (~100 µm) was secured to the bone with a mixture of superglue and dental cement. Immediately after the craniectomy, the mice were imaged under the two-photon microscope while maintaining anesthesia and normo-temperature.

**Organotypic hippocampal slice preparation.** Postnatal day 6 (P6) to P7 mice were used to prepare organotypic slice cultures using the rocking plate technique (Romijn et al., 1988) or the membrane insert.

**Figure 2.** Construction of the sCLM targeting vector and the generation of chimeric mice. A, Schematic representation of Rosa26 knock-in targeting vector. The LoxP sites are represented by blue triangles. The sCLM transgene is depicted as a brown box, and the poly A as a red box. The combined STOP-neomycin selection cassette is indicated as a gray box. The ubiquitous CAAG promoter is represented by a green box. The DTA-negative selection cassette is represented by a yellow box. B, Primer set for the detection of the nonrecombined conditional knock-in allele. C, The genotype of the pups derived from the F1 breeding with Cre deleter mice were tested by PCR enabling the detection of the neomycin cassette. PCR using recombined ES cell clones as template, alone or in the presence of 10 ng of C57BL/6 genomic DNA (clone; clone + WT) served as positive controls. PCR with wild-type C57BL/6 genomic DNA (WT) and without DNA (H2O) as a template served as negative controls. PCR picture was obtained after the loading of PCRs on the LabChip system from Caliper LifeSciences.
technique (Stoppini et al., 1991) for electrophysiology and imaging. Isolated hippocampi were cut into 400 μm slices on a McIlwain tissue chopper (Mickle Laboratory Engineering). Slices were transferred to membrane inserts (PICMORG50, Millipore) or coverslips, which were placed in glass-bottomed six-well plates (catalog #P06-1.5H-N, CellVis). Both culture configurations were incubated at 5% CO₂, 36°C in Neurobasal-A growth medium supplemented with DNase and glutamax (1% and 0.03 mg/ml gentamicin (all from Thermo Fisher Scientific)). The growth medium was changed every 3–4 d. Slices were used for experiments between day in vitro (DIV) 6 and DIV 20.

**Dissociated cell culture preparation.** Hippocampal murine neurons were prepared based on the method of Beaudoin et al. (2012), modified to use early postnatal mice, and to use a commercial kit (LK003150, Worthington Biochemical). P1–P4 mice were killed by guillotine method cut just caudal to the ears. Bilateral incisions were made from ear to eye as far ventral as feasible, allowing rostral reflection of the calvaria. Using a spatula dipped in Gey’s Balanced Salt Solution (GBSS; catalog #G9779, Sigma-Aldrich) supplemented with glucose (Sigma-Aldrich) to 1.5% and then filter sterilized, brains were gently removed to a 6 cm Petri dish of Neurobasal-A growth medium supplemented with DNase in a 6 cm Petri dish pre-equilibrated with Earle’s Balanced Salt Solution supplemented with DNase and B27, 500 μM FGF and gentamicin (all from Thermo Fisher Scientific). The growth medium was changed every 2–3 d. Slices were used for experiments before being returned to incubator for a 10- to 40 min incubation before being returned to incubator for a 10- to 40 min incubation commensurate with pup age. Halfway through incubation, tissue was triturated 5× with a 10 ml plastic pipette and returned to the incubator. Reduced yield from using plastic pipettes was deemed acceptable to avoid inconsistency of flame-polished glass tip diameter and subsequent trituration-related damage. Tissue was again triturated 5× with a 10 ml pipette and 3–4× with a 5 ml pipette before being strained through a 40 μm basket filter (catalog #431750, Corning) and transferred to a 50 ml conical tube. The cell mixture was centrifuged for 6 min at 300 × g, and after carefully aspirating the supernatant the cell pellet was then resuspended with EBSS plus an ovomucoid inhibitor albumin mixture supplemented with DNase (each part of the Worthington Biochemical dissociation kit). Gentle trituration with a 10 ml pipette held several centimeters from the tube bottom was sufficient to resuspend. This mixture was then carefully and slowly overlaid atop 5 ml of ovomucoid protein in EBSS and centrifuged for 5 min at 70 × g. The resulting pellet was resuspended in Neurobasal-A media (catalog #10888–022, Thermo Fisher Scientific) without supplementation and transferred to a fresh tube. The cell concentration and proportion surviving were acquired using Fluoview 1000MPE microscope with prechirp optics and a fast acousto-optical modulator mounted on an Olympus BX61W1 upright microscope equipped with a 25× 1.05 numerical aperture (NA) water-immersion objective (Olympus). A mode-locked Ti:Sapphire laser (MaiTai, Spectra-Physics) generated two-photon fluorescence with 860 nm excitation. Emitted light was detected through two filters in the range of 460–500 nm for CFP and 520–560 nm for YFP. Two photomultiplier tubes (Hamamatsu Photonics) were used to acquire CFP and YFP signals simultaneously. Three-dimensional stacks (3D) of raster scans in the x-y plane (0.33 μm/pixel x-y) were acquired using custom-designed software (LabVIEW) and a scan head from Radiance 2000 MP (Bio-Rad), equipped with a 40×, 0.8 NA water-immersion objective (Olympus), and photomultiplier tubes with appropriate filters for CFP (450/80) and YFP (545/30). A Spectra-Physics Mai Tai laser was set to 860 nm for sCLM imaging. Serial images were collected with a 2×2 pixel interval of 2 μm. The images were reconstructed offline either by ImageJ (RRID:SCR_003070) or MATLAB (version 2017a). To calculate [CI⁻], using ImageJ, a region of interest (ROI) was drawn around the cell body and dendrites, and the ratio of YFP/CFP fluorescence intensity was measured. Images analyzed with MATLAB were masked using image processing functions as described below. In either case, the ratio was converted into [CI⁻], by the following equation:

\[
[\text{Cl}^-]_i = \frac{R_{\text{max}} - R}{(R - R_{\text{min}})},
\]

where \(K_d\) is the dissociation constant, \(R_{\text{max}}\) is the ratio obtained in the absence of Cl⁻, and \(R_{\text{min}}\) is the ratio when sCLM is maximally quenched (Grimley et al., 2013; see Fig. 4A).

**FLIM of MEQ.** MEQ, 1.5 mM, was delivered to the cytoplasm via whole-cell recording pipette (see Fig. 7F). Unlike sCLM, MEQ is insensitive to pH (Biversi et al., 1992). The CI⁻-sensitive signals of both fluorophores used in this study are independent of the dye concentration due to either the ratiometric nature of sCLM fluorescence or the intrinsic nature of fluorescence lifetime as a property of MEQ that is independent of fluorophore concentration. MEQ was excited at 750 nm using the custom-built scanning microscope detailed above with an appropriate emission filter (390/65), and fluorescence lifetime was measured using SPCIImage software (Becker & Hickl). The selection of the ROI and calculation of the MEQ lifetime was measured offline by a custom-written MATLAB (MathWorks) program. MEQ lifetime values were translated to [CI⁻] using the Stern–Volmer calibration (see Fig. 4B; Biversi et al., 1992). The detailed morphologies of the cells were visualized by adding 20 μM Alexa Fluor-594 to the recording pipette. High-resolution Alexa Fluor-594 images (x-y axis, 0.307 μm/pixel; z-axis, 2 μm/pixel) were used for precise selection of the ROI in MEQ lifetime images with x-y resolution of 1.537 μm/pixel and z-axis interval of 2 μm.

**Electrophysiological measurement of \(E_{GABA}\).** A subset of experiments was performed using whole-cell and gramicidin perforated patch-clamp techniques (for details, see Electrophysiology section) in conjunction with sCLM imaging or FLIM. The \(E_{GABA}\) was extracted from the intercept of currents evoked by the application of 10 μM GABA applied via a micropipette and driven by a 10 ms, 5 psi square-wave pressure transient (Picospitzer III, General Valve) to visualize segments of the dendrites while clamping the cell at voltage steps ranging from −110 to +10 mV. GABA\(_{\text{ARs}}\) were blocked by 2 mM GPP5845. The size of the dendritic region of interest was determined by adding Alexa Fluor-594 inside the puffing pipette and measuring the region over which the dye diffused. [CI⁻] was calculated using the Nernst equation, as follows:

\[
E_{\text{Cl}^-} = \frac{-RT}{F} \ln \frac{[\text{Cl}^-]}{[\text{Cl}^-]_o}
\]

where R is the gas constant (8.315 J/mol/K), T is the temperature in Kelvin, F is the Faraday’s constant (96.487 C/mol), and [CI⁻] is the extracellular concentration of chloride.
Electrophysiology
For electrophysiological recordings, organotypic slice cultures or dissociated cultured cells were transferred to a recording chamber and perfused with ACSF (2.5 mM/min) containing the following (in mM): 124 NaCl, 1.25 NaH2PO4, 2.5 KCl, 26 NaHCO3, 2 CaCl2, 2 MgSO4, and 20 d-glucose, bubbled with 95% O2 and 5% CO2 at 34°C. The following two separate series of electrophysiological experiments were performed. The first was granule cell perforated patch clamping on hippocampal dissociated cell cultures, where cells were visualized using an upright microscope (Eclipse FN1, Nikon) equipped with a 40×, 0.8 NA water-immersion objective (Nikon), a DualView image intensifier (Optical Insights), and a CCD camera (Chameleon3, Point Gray Research) for simultaneous acquisition of two separate images of CFP and YFP with x resolution of 1.996 μm/pixel and y resolution of 1.896 μm/pixel (see Fig. 7A). Electrodes were pulled from borosilicate glass capillaries (Sutter Instruments) using a micropipette puller (model P-97, Sutter Instruments) with resistance of 5–7 MΩ when filled with internal solution containing the following (in mM): 140 KCl, 5 MgCl2, 10 HEPES, and 5 EGTA, with osmolarity of 290 mOsM, pH 7.25–7.35, adjusted with KOH. Granicidin perforated patch clamping was performed as described previously (Rahmati et al., 2016). Primary dendrites of sCLM cells were selected as ROIs to have a higher resolution and signal-to-noise ratio. The second was whole-cell patch-clamp recording of hippocampal organotypic slices: these experiments were performed on wild-type C57 rocking plate organotypic slices to simultaneously load the hippocampal pyramidal cells with tdtTomato virus (Addgene-AAV9-CAG-FLEX-tdTomato, in 2 of DLX-Cre mice, which were transfected on the day of slicing with the tdTomato virus (Addgene-AAV9-CAG-FLEX-tdTomato, in 2 μl/ml culture media) to visualize interneurons. Whole-cell recordings were used to improve the stability of this complex experiment, which involved repeated current–voltage measurements over a wide range of membrane potentials during focal activation of multiple interneurons. Somatic whole-cell recordings have a modest impact on dendritic Cl– homeostasis (Jarolimek et al., 1999; Khirug et al., 2005). Dialysis from the electrode solution would have served to reduce, not increase, the heterogeneity that we measured; we found no relationship between reversal potentials and distance from soma (see Fig. 7D). Patch pipettes with resistances of 5–7 MΩ were filled with an internal solution containing the following (in mM): 124 K-MeSO4, 5 KCl, 10 KOH, 4 NaCl, 10 HEPES, 28.5 sucrose, 4 Na2ATP, 0.4 Na3GTP, and 1.4 mM MEQ, with osmolarity of 295 mOsM and pH 7.25–7.35. For visualizing the whole morphology of the recorded cells, 20 μM Alexa Fluor-594 hydrazide (Thermo Fisher Scientific) was added to the internal solution on the day of the experiment. Resting membrane potential (RMP) and input resistance were measured after whole-cell configuration was reached. Series resistance (R_s) assessed in voltage-clamp mode by –5 or –10 mV (voltage step) was monitored for stability during the experiments. Only neurons with R_s < 20 MΩ were included in the analysis. For all the electrophysiological recordings, signal acquisition was performed using a Multiclamp amplifier (Multiclamp 700B, Molecular Devices) with Clampex 10 software (Molecular Devices). Signals were sampled at 10 kHz and filtered at 2 kHz. Data were stored on a PC for offline analysis after digitization using an analog-to-digital converter (Digidata 1440A, Molecular Devices).

Rubi-GABA uncaging
Rubi-GABA uncaging was selected in targeted path scanning [TPS (theta pulse stimulation)] mode. The electrophysiology software triggered the laser and controlled the pulse duration. RuBi-GABA uncaging was performed at 750 nm. Laser intensity was increased until reliable GABA_A receptor-mediated currents were observed. E_CL was measured as described above. Laser stimulation without the presence of RuBi-GABA did not induce currents.

Image analysis
To measure the stability of microdomains, colocalization analysis was performed by selecting an ROI and measuring YFP/CFP in each pixel of that ROI using a 2-dimensional spatial correlation subroutine of ImageJ (x–y coordinates subroutine) in which the x value represents the YFP/CFP ratio for a particular pixel in the first image, and the y value represents the YFP/CFP ratio for the same pixel in the second image (see Fig. 8B). The correlation between the two images was then calculated by the linear correlation for all pixels in the ROI (see Fig. 8D). Identical images, or images that had the YFP/CFP ratios for each pixel change by the same amount between the two images, would have a correlation coefficient of 1. This pixel-wise spatial correlation coefficient is reduced whenever the Cl– concentration of a given pixel, as assessed by the YFP/CFP emission ratio, no longer matches the prior concentration. This could occur, for example, because of a change in the Cl– concentration of a microdomain or the anatomic distribution of that microdomain. Pseudo-colored images of sCLM (Figs. 3, 4C,D) were generated by selecting the ROIs, measuring the YFP/CFP in each pixel of that ROI, and using the Pseudo-color Image Look-Up Tables Fiji subroutine. The pseudo-colored image of sCLM in Figure 4F was generated by reconstructing a flattened (2D) image using the Cl–-insensitive CFP intensity to weight the contribution of the YFP/CFP ratio of each pixel in a 3D image stack. Pseudo-color image of FLIM (see Fig. 7F) was generated by calibrating MEQ lifetime using the Stern–Volmer relationship and calculating [Cl–]i, (Fig. 4B; Verkman et al., 1989). FLIM experiments included the brighter membrane-impermeable dye Alexa Fluor-594 as well as MEQ in the recording pipette solution. The Alexa Fluor-594 (Cl–-insensitive) signal was used to define the intracellular pixels. [Cl–]i was then calculated from the fluorescence lifetime of MEQ in those pixels. A 2D image was created from a 3D image stack by weighting the MEQ-derived [Cl–]i values by the relative intensity of Alexa Fluor-594. The Alexa Fluor-594 signal intensity was used to determine the brightness of each pixel that was pseudo-colored based on the calculated [Cl–]i. In pixels where MEQ signal was indistinguishable from background noise, no pseudo-color was assigned, and the pixel was colored using a grayscale Alexa Fluor-594 signal. The grayscale then demonstrated the dendritic structure in areas where the MEQ signal is inadequate. For noise analysis, a representative 30 min time series (180 frames) of acutely sliced CLM-expressing CA1 pyramidal cells imaged in the presence of TTX is compared with the noise acquired from a similar series acquired with the shutter closed (Fig. 4H). MATLAB is used to perform a line scan to represent each neuron as a vector of intensity values and to compute subsequent fast Fourier transforms (FFTs) of CLM YFP intensity and the corresponding channel closed-shutter noise intensity (see detailed noise analysis below). As the power spectra of more time-frame images are averaged, the power of noise from a closed shutter will converge to that of white noise. FFT power obtained from variations because of persistent structural differences such as microdomains will not converge to a white noise power spectrum, converging instead to a greater power difference from white noise than shutter noise (Fig. 4H).

Detailed noise analysis
To test whether the observed spatial variance in [Cl–]i could arise from noise, we compared the averaged power spectra of the spatial frequencies of [Cl–]i reporter CLM intensity values in the neurons to the power spectrum of closed-shutter noise. Noise analysis was performed with customized scripts and documentation code using MATLAB Fourier transform and image analysis functions. Of the two fluorescent components of CLM, Cl–-sensitive YFP was chosen and compared with the noise detected with the laser shutter closed over the same number of time frames for each neuron. Dendrites or soma were selected based on...
persistent signal strength and linearized by line scan. The YFP signal of each frame was coregistered to the first frame and applied to both the YFP and closed-shutter noise image of that frame. Spatial FFTs were computed on these intensity signals to obtain the power (amplitude squared) of the observed YFP and closed-shutter noise values at a spectrum of wavelengths from 0.794 \( \mu m \) (exclusive) to 25.4 \( \mu m \) (inclusive), bound by the Nyquist wavelength and the distance equivalent to the largest exponent of base 2 that did not exceed the length of the smallest vector in pixels, respectively. The closed-shutter noise power spectra were rescaled by the ratio of the area under the curves between the mean power spectrum of the neuronal signal and shutter noise. The white noise power spectrum was generated from averaging the power spectra of 72,000 vectors created from 400 random shuffles of each of 180 time frames. Defining \( X_1:::180 \) as the vectors at 180 different time frames of observed intensity values for noise from a closed shutter or a signal from a neuron, we obtained 400 randomly shuffled versions for each 180 time-frame vectors \( A_1:::72,000 \). By computing the FFT on each of the vectors \( A_1, 72,000 \) and averaging the square of each result, we obtain the respective white noise power spectrum \( W \) for either a neuronal signal or closed-shutter noise defined as follows:

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**Figure 3.** Evidence for the existence of chloride microdomains from in vivo sCLM imaging. A, Example of layer II/III cortical neurons expressing sCLM and their color-coded images in a P20 mouse. The data are presented as YFP/CFP ratios rather than \([Cl^-]_i\) because the calibration of sCLM is not feasible in vivo as a consequence of the limited capacity to permeabilize the membrane of target neurons and to manipulate the extracellular \( Cl^- \) concentration. B, A histogram of YFP/CFP ratios in hundreds of cells in vivo. C–F, Higher magnification of three sCLM-expressing cells and their color-coded images showing somatic and dendritic YFP/CFP values restricted to a single z-plane. The corresponding histograms demonstrate the range of YFP/CFP ratios in individual pixels.
The absolute area between the cumulative average of either the power spectra of the YFP neuronal signal or the closed-shutter noise and their respective white noise power spectrum was calculated at each time frame. The resulting curves were normalized and plotted as a function of frame number such that the number of frames included in the average power spectrum was equal to the frame number. Comparing the behavior of the averaged power spectra of signal intensity versus closed-shutter noise as a function of the number of frame spectra averaged allowed a direct comparison of heterogeneity. The heterogeneity that is random and transient (i.e., noise) will average out with an increasing number of frames, but heterogeneity that is persistent will not.

**Experimental design and statistical analysis**

Imaging data were analyzed with ImageJ (https://imagej.nih.gov/ij) or MATLAB. Details of the analysis are included in the Image analysis section of Materials and Methods. Statistical analysis of the data was
performed in GraphPad Prism 8 (GraphPad Software). Statistics were assessed with two-tailed unpaired Student's t tests when comparing two groups. The number of data points (n) and the statistical significance (p value) are stated in the figure legends.

Results

Unique response of single pyramidal cells to individual inhibitory interneurons

We recorded single pyramidal cells using whole-cell patch clamping while selectively activating individual neighboring inhibitory interneurons in organotypic hippocampal slice preparations (Fig. 1A,B). We observed unique, reproducible values of \(E_{\text{GABA}}\) for each presynaptic interneuron recorded from a single pyramidal cell (Fig. 1C) whose distribution paralleled the broad range of previously reported \(E_{\text{GABA}}\) (Fig. 1D, Table 1). Our results demonstrate that individual interneurons have unique \(E_{\text{GABA}}\) values on postsynaptic pyramidal cells (\(n = 40\) interneurons; \(n = 10\) pyramidal cells; \(-23\) mV range in \(E_{\text{GABA}}\) Fig. 1E). The measured range may be an underestimate of the variance of synaptic \(E_{\text{GABA}}\) in these neurons because individual interneurons may have multiple synapses onto pyramidal cells (Somogyi et al., 1983; Buhl et al., 1994; Maccarelli et al., 2000; Bezaire and Soltész, 2013), each of which may have a different \(E_{\text{GABA}}\) value, and the somatic recording would reflect the combined currents from all synapses originating from the activated interneuron. Of note, not all stimulated interneurons evoked GABAergic currents in the recorded pyramidal cells; 56% of the stimulated interneurons resulted in recordable currents.

Evidence for dendritic chloride microdomains by fluorescence microscopy and electrophysiology

How can a single neuron express so many different values of \(E_{\text{GABA}}\)? To investigate this, we measured the subcellular [\(\text{Cl}^{-}\)] distribution in single neurons. To enable cell type-specific analysis of subcellular cytoplasmic [\(\text{Cl}^{-}\)] distribution in single neurons. To enable cell type-specific analyses of subcellular cytoplasmic [\(\text{Cl}^{-}\)], we assessed with two-tailed unpaired Student's t tests when comparing two groups. The number of data points (n) and the statistical significance (p value) are stated in the figure legends.

To analyze the subcellular \(\text{Cl}^{-}\) variance in a preparation in which sCLM YFP/CFP ratios can be quantitatively calibrated to \([\text{Cl}^{-}]_i\) (Fig. 4A), we performed two-photon high-resolution fluorescence microscopy of hippocampal organotypic slice cultures expressing sCLM. Our data demonstrate spatial variability of \([\text{Cl}^{-}]_i\) in the dendrites of individual neurons in the presence of 1 \(\mu\text{M}\) TTX (Fig. 4C–F). The individual pixels of the neuronal image depicted in Figure 4F demonstrate a wide range in dendritic \([\text{Cl}^{-}]_i\) (Fig. 4G). To examine the contribution of noise to these images, we performed time series imaging of CLM-expressing neurons (180 images over 30 min). Fluorescence intensity arising from stable cytoplasmic [\(\text{Cl}^{-}\)], microdomains would be expected to exhibit less temporal variance than noise. Over the 30 min observation period, the spatiotemporal variation in dendritic YFP intensity was much less than the corresponding variance of instrument noise (measured with the shutter closed and normalized to the YFP emission intensity, Figs. 4H, 5). This supports the presence of persistent spatial differences in [\(\text{Cl}^{-}\]), that cannot be explained by noise. We estimated the size of the [\(\text{Cl}^{-}\)] microdomains by analyzing the number of neighboring pixels (area, 0.09 \(\mu\text{m}^2\)) with similar [\(\text{Cl}^{-}\]) (Fig. 6, single z-plane). Our data demonstrate that the size of microdomains varies between 1 and 10 pixels in a single neuron while the majority of microdomains were 1 pixel (Fig. 6F,G), suggesting that the dimensions of many microdomains were likely below the resolving power of two-photon microscopy.

To test whether the fluorescence measures of [\(\text{Cl}^{-}\)] correlated with the electrophysiological assays of [\(\text{Cl}^{-}\]), we conducted simultaneous measurements of the local dendritic [\(\text{Cl}^{-}\]), using sCLM and [\(\text{Cl}^{-}\]), calculated from \(E_{\text{GABA}}\) derived from gramicidin perforated patch-clamp recordings and local pressure application of 10 \(\mu\text{M}\) GABA in cultured hippocampal neurons (Fig. 7A). Local YFP/CFP ratios and [\(\text{Cl}^{-}\)], measured by sCLM were highly correlated with the [\(\text{Cl}^{-}\]) calculated by \(E_{\text{GABA}}\) (Fig. 7B,C). The correlation coefficient between the two measures was 0.65 (\(n = 38\) ROIs; \(n = 11\) cells; \(p < 0.0001\); 95% confidence interval 0.41–0.80), consistent with the role of cytoplasmic microdomains in setting the local \(E_{\text{GABA}}\) (Delpierre and Staley, 2014; Glyks et al., 2017). To estimate potential errors in the calculation of \(E_{\text{GABA}}\) because of inadequate dendritic space clamp (Staley and Mody, 1992), we tested for a correlation between the recorded \(E_{\text{GABA}}\) and the distance of the ROI from the center of soma. There was no significant correlation (Fig. 7D; \(n = 38\) ROIs; \(n = 11\) cells; \(R = -0.20, p = 0.21; 95\%\) confidence interval, −0.49 to 0.12; Boffi et al., 2018).

sCLM retains the pH sensitivity of CLM (Grimley et al., 2013). Therefore, we therefore used a complimentary approach, assaying subcellular [\(\text{Cl}^{-}\]) using the pH-insensitive, \(\text{Cl}^{-}\)-sensitive, single-wavelength fluorophore MEQ (Biwersi and Verkman, 1991). We performed FLIM to avoid artifacts arising from subcellular variation in MEQ concentration. The FLIM measurements using MEQ delivered by the whole-cell recording pipette confirmed the variability of [\(\text{Cl}^{-}\]) in dendrites of hippocampal pyramidal neurons in CA1 area (Figs. 4B, 7E,F). The variance in [\(\text{Cl}^{-}\]) measured by MEQ FLIM was also highly correlated with simultaneous electrophysiological measurements of \(E_{\text{GABA}}\) measured in voltage-clamp neurons by local application of GABA using direct visual guidance (\(n = 9\) ROIs; \(n = 7\) cells; \(R = 0.85, p = 0.003; 95\%\) confidence interval, −0.97 to −0.43; Fig. 7G). These data support the idea that subcellular variance in [\(\text{Cl}^{-}\]) is a significant contributor to the variance in \(E_{\text{GABA}}\) (Table 1). We next measured the stability of \(E_{\text{GABA}}\) using two-photon RuBi-GABA uncaging (Fig. 7H). RuBi-GABA (Rial Verde et al., 2008) was photosylized adjacent to the sCLM-expressing somatic or
dendritic ROIs. $E_{\text{GABA}}$ was assessed by uncaging at intervals of 20 s at holding potentials from $-120$ to $-10$ mV. This assessment was repeated at the same ROI at an interval of 2–5 min to examine the activity dependence of the local $E_{\text{GABA}}$. There was no significant difference between the sequential $E_{\text{GABA}}$ measurements at this level of receptor activation ($\text{Fig. 7f}$; pairs, $n = 8$; paired $t$ test, $p = 0.36$).

**Physiologic chloride microdomains are not defined by the activity of cation-chloride cotransporters**

Changes in $[\text{Cl}^{-}]$ are readily observed in response to alterations of the equilibrium conditions for CCCs using either osmotic or ionic perturbations of the perfusate (Thompson et al., 1988; DeFazio et al., 2000; Dzhala et al., 2010; Glyks et al., 2014a). This influence of transport equilibrium conditions on $[\text{Cl}^{-}]$ has led to the idea that CCCs can also set $[\text{Cl}^{-}]$, independently of the equilibrium conditions. To test the degree to which $[\text{Cl}^{-}]$ microdomains are defined by CCCs, we imaged sCLM in stable extracellular ionic and osmotic conditions. We acquired time-lapse images for 40–50 min ($\text{Fig. 8A}$). The ROIs (soma and dendrites of a single pyramidal neuron; $\text{Fig. 8B}$) were monitored and compared pixel by pixel with previous measurements every 10 min ($\text{Fig. 8C,D}$). Consistent with the prior noise analyses ($\text{Fig. 4H}$), the microdomains remained highly stable in control conditions. Applying a high concentration of furosemide to block NKCC1 and KCC2 simultaneously (Gillen et al., 1996) did not alter the microdomain distributions ($\text{Fig. 8D,E}$). $\text{Fig. 8E}$ shows the mean normalized Pearson’s $r$ by time [normalized to the initial correlation coefficient and indicated as a percentage: at baseline in the absence of furosemide $\pm$ SD: $97.81 \pm 8.43\%$, $n = 12$; after application of furosemide for 30 min $\pm$ SD: $97.34 \pm 7.09\%$, $n = 12$; $p = 0.92$; effect size ($d$) = 1.42; 95% confidence interval, $-10.82$ to $-9.86$; $R^2 = 0.001$. Because the mobility of $\text{Cl}^{-}$ in neuronal cytoplasm is very high (Kuner and Augustine, 2000), $[\text{Cl}^{-}]$ should have rapidly homogenized after the blockade of CCCs if CCC activity had created the $[\text{Cl}^{-}]$ microdomains. The fact that this was not observed indicates that $[\text{Cl}^{-}]$ microdomains do not arise from local differences in cation–$\text{Cl}^{-}$ transport rates. While the correlation coefficient was stable over 1 h, the value of the correlation coefficient was dependent on the expression levels of CCCs, as demonstrated in $\text{Fig. 8D,F}$; the coefficient was reduced by 10% in organotypic slices ($\text{Fig. 8G,H}$; Glyks et al., 2014a). Cultured hippocampal organotypic slices develop spontaneous epileptiform activities beginning late in the first day after slicing, which can be detected by chronic calcium imaging observation (Lillis et al., 2012; Jacob et al., 2019) and field potential electrophysiological recording (Dyhrfjeld-Johnsen et al., 2010). Our data show that the returns to baseline in $\text{Fig. 8}$, $G$ and $H$, are monotonic (i.e., they do not jump between values of...
[Cl\textsuperscript{−}], that represent the smallest resolvable changes). Further, [Cl\textsuperscript{−}], returns to baseline after SLEs, and the rate of return was independent of the initial (baseline) value. These results indicate that temporal fluctuations in [Cl\textsuperscript{−}]i can be resolved by this technique, and that [Cl\textsuperscript{−}]i microdomains persist after large transient Cl\textsuperscript{−} fluxes. As previously mentioned, sCLM is pH sensitive. However, changes in pyramidal cell cytoplasmic pH during seizures in hippocampal organotypic slices (Lillis et al., 2012) would lead to an underestimate of the ictal changes in [Cl\textsuperscript{−}]i measured with sCLM.

The stability of chloride microdomains is reduced by actin depolymerization
If the observed cytoplasmic Cl\textsuperscript{−} microdomains were created by the differential distribution of less mobile anionic macromolecules, then altering the distribution of those macromolecules should induce a corresponding change in Cl\textsuperscript{−} microdomains. Actin is one such macromolecule that could contribute to the structural basis for Cl\textsuperscript{−} microdomains. Actin is a variably branched cytoskeletal polyelectrolyte with a large negative surface charge (Castaneda et al., 2018) that is stationary once polymerized into filamentous actin (F-actin; Watanabe and Mitchison, 2002). Actin is continuously polymerized and depolymerized, and the balance between these two processes can be altered by several well studied drugs (Fenteany and Zhu, 2003; Chia et al., 2016). One such drug is latrunculin B, which binds to actin monomers to inhibit actin assembly (Morton, 2000), thereby shifting the balance toward actin disassembly. Latrunculin B, 5 \mu M, has been demonstrated to preserve the dendritic structure of hippocampal pyramidal cells over a period of hours in vitro (Kim and Lisman, 1999). We first confirmed that this concentration of latrunculin B does not affect the shape of the dendrites by imaging CA1 pyramidal cells expressing the Cl\textsuperscript{−}-insensitive fluorophore TurboRFP (Fig. 9A). Following baseline imaging, RFP-expressing neurons were continuously imaged for an additional hour after being in the presence of 5 \mu M latrunculin B. Pixel-to-pixel (pixel-wise) correlation of dendritic and somatic ROIs from sequential images were compared with the baseline images (Fig. 9B). Our data confirm that the application of 5 \mu M latrunculin B did not change the shape of the neurons measured by pixelwise correlation of sequential images [initial normalized percentage change in Pearson’s r ± SD (image 1 vs 2):
1.73 ± 2.06%, n = 11; final normalized percentage change in Pearson’s r ± SD after 1 h in 5 μM latrunculin B (image 1 vs 5): −2.69 ± 1.78%, n = 11, p = 0.26; Fig. 9B).

Next, we repeated the same experimental procedure in organotypic hippocampal slice cultures from sCLM transgenic mice (Fig. 9C). To exclude potential effects of cytoskeletal alterations on synapses and transporters, the experiments were performed in the presence of 1 μM TTX, 10 μM bumetanide (NKCC1 blocker; Dzhala et al., 2008), and 1 μM VU 0463271 (VU; a selective KCC2 blocker; Sivakumaran et al., 2015). Addition of 5 μM latrunculin B steadily changed the distribution of Cl− microdomains, evidenced by a reduction in the pixel-wise correlation coefficient of sequential images (Fig. 9D) compared with the control conditions [i.e., the baseline images obtained in the presence of 1 μM TTX, 10 μM bumetanide (NKCC1 blocker; Dzhala et al., 2008), and 1 μM VU 0463271 (VU; a selective KCC2 blocker; Sivakumaran et al., 2015)].

**Figure 7.** Simultaneous [Cl−]i imaging and electrophysiology. A, A representative hippocampal-dissociated cultured neuron expressing sCLM and visualized by DualView using a CCD camera to simultaneously measure the YFP and CFP fluorescence. B, The [Cl−]i, measured by fluorophore data (YFP/CFP) and the [Cl−]i, calculated from E_GABA, in dissociated cultured cells are highly correlated (n = 38 ROIs, 11 cells, p < 0.0001). C, There is a significant correlation between the [Cl−]i measurements by sCLM and [Cl−]i calculated from E_GABA (n = 38 ROIs, 11 cells, p < 0.0001). D, There was no significant correlation between the distance of selected ROIs from the center of soma and the recorded E_GABA (n = 38, 11 cells). E, A representative example of a hippocampal pyramidal cell loaded with Alexa Fluor-594 and Cl−-sensitive, pH-insensitive dye, MEQ. F, Fluorescence lifetime imaging of [Cl−]i in hippocampal neurons confirms the variance of [Cl−]i in dendrites. [Cl−]i calculated from MEQ lifetime is visualized in pseudocolor where the Alexa Fluor-594 [Cl−]i-insensitive signal sets the transparency value such that the [Cl−]i value and corresponding color relate solely to the MEQ lifetime while the brightness of the pixel is set by Alexa Fluor-594 as a structural marker (gray colorbar, top right). No averaging or clustering was performed on MEQ images. Note the variety of [Cl−]i values and the frequent juxtaposition of relatively high and low values in close proximity. G, After loading the cells with MEQ, 50 μM GABA was puffed to different visualized segments of dendrites and E_GABA was measured. The [Cl−]i, measured by MEQ lifetime in that segment was compared with the [Cl−]i, calculated from the Nernst equation from the E_GABA measured electrophysiologically in the same segment. The two measures of [Cl−]i are significantly correlated (n = 9 ROIs, 7 cells, p = 0.003). H, A representative example of a two-photon RuBi-GABA uncaging experiment on sCLM-expressing pyramidal cells. The red arrow shows the selected ROI, and the blue arrow points to the recording pipette containing Alexa Fluor-594. E_GABA was measured twice at the same ROI to examine the stability of the GABA-evoked currents. I, There was no significant difference between the first and second GABA uncaging-mediated E_GABA recordings (paired t test; pairs, n = 8; p = 0.36).
presence of TTX, bumetanide, and VU alone; normalized mean percentage pixel-wise correlation coefficient by the end of imaging 6 SD for VU and bumetanide: 98.8 ± 12.18%, n = 15; for latrunculin B: 77.3 ± 10.54%, n = 15, p = 0.002, effect size (d) = 1.89; 95% confidence interval, −3.94 to −8.98, $R^2 = 0.47$; Fig. 9D]. The mean percentage change in pixel-wise correlation coefficient in the presence of CCC antagonists (either high concentration of furosemide or bumetanide and VU: $-24.0 ± 9.84%$, $p < 0.0001$; $n = 15$ ROIs; effect size (d) = 2.5; 95% confidence interval, −24.17 to −14.32; $R^2 = 0.63$; Fig. 9E]. In addition to TurboRFP, the average pixel-wise correlation was also compared between CFP images as an extra control that did not significantly change during the same time period [transporter blockers, $-7.20 ± 15.5%$, latrunculin B, $-7.09 ± 4.7%$, $p = 0.98$; effect size (d) = 0.01; 95% confidence interval, −11.41 to 11.63; $R^2 = 2.95e-005$]. The addition of latrunculin B to the perfusate containing transporter blockers did not alter the average percentage change of $[\text{Cl}^-]_i$ observed after transport block (transporter blockers: $14.76 ± 18.13%$, n = 20; latrunculin B: $14.74 ± 14.07%$, n = 13; $p = 0.90$; 95% confidence interval, −12.14 to 12.10; effect size = 0.001; $R^2 = 3.45e-
These findings indicate that although CCCs are actively transporting Cl\(^{-}\) across the membrane in this preparation, the CCCs do not define the distribution of microdomains. In contrast, latrunculin B modified [Cl\(^{-}\)]\(_i\) microdomains without changing the average [Cl\(^{-}\)]\(_i\) by altering the macromolecular distribution through anionic biopolymer disassembly.

Discussion

These data obtained using multiple complimentary recording techniques support the existence of stable [Cl\(^{-}\)]\(_i\) microdomains. Our studies extend the variance of E_{GABA} and [Cl\(^{-}\)]\(_i\) between neurons (Table 1; Glykys et al., 2014a,b; Sato et al., 2017) to focus for the first time on the subcellular variation of Cl\(^{-}\) in individual dendrites. The stability of microdomains over the entire time interval for which stability could be assessed, and the wide variance in the E_{GABA} of pyramidal cells measured when individual presynaptic interneurons were stimulated, underscore the potential functional importance of [Cl\(^{-}\)]\(_i\) microdomains for signaling at GABAergic synapses.

The sizes of Cl\(^{-}\) microdomains varied both in concentration and anatomic extent (Fig. 6), and many microdomains are likely below the resolution of two-photon microscopy, as would be expected based on the proposed mechanism of Cl\(^{-}\) distribution (Fig. 10). Pixel-wise histograms of Cl\(^{-}\) distribution varied from one dendrite to the next (Fig. 3, compare D, E, F; compare Fig. 3D–E, 4G), so there is not a single mean and variance of [Cl\(^{-}\)]\(_i\) microdomains.

Neuronal Cl\(^{-}\) microdomains have been a source of confusion and controversy. A wide variety of experimental evidence supporting the nonuniform distribution of cytoplasmic Cl\(^{-}\) (Barna et al., 2001; Duebel et al., 2006; Szabadics et al., 2006; Földy et al., 2010; Glykys et al., 2014a; Berglund et al., 2016; Untiet et al., 2016; Zorrilla de San Martin et al., 2017) has been challenged primarily by theoretical arguments (Luhmann et al., 2014; Doyon et al., 2016; Savtchenko et al., 2017; Düsterwald et al., 2018). The simplest argument for the existence of neuronal cytoplasmic Cl\(^{-}\) microdomains is the fact that the vast majority of anions in the...
cytoplasm are relatively immobile. In neurons, the GABA\(_A\)-permeable ions Cl\(^{-}\) and HCO\(_3\)^{−} together make up only a minority of the intracellular anions. The rest of the intracellular anions are amino acids and phosphates (Morawski et al., 2015), of which only a small minority are not associated with macromolecules (Masuda et al., 1990; Veech et al., 2002), such as nucleic acid species (Manning, 1978) and proteins (Gianazza and Righetti, 1980). These immobile anionic polymers are not uniformly distributed in the cytoplasm (Gut et al., 2018; Chen et al., 2020), so it is reasonable to expect that there is a nonuniform distribution of mobile anions to compensate for the distribution of immobile anions. Macroscopic nonuniform distributions of mobile ions are readily demonstrated experimentally using gels with fixed anionic charges (Procter, 1914; Fatin-Rouge et al., 2003; Golmohamadi et al., 2012), and these findings extend to anionic biopolymers that are not crosslinked into gels (Marinsky, 1985).

Here we demonstrate that the inhibition of polymerization of one such anionic macromolecule, actin, led to a proportionate

![Diagram of protein hydration shell and cytoplasmic free water](image)

Figure 10. Stoichiometric membrane chloride cotransporters can be at equilibrium over the range of cytoplasmic chloride concentrations observed in this study. A, Effects of nonuniform distribution of immobile charges on the distributions of mobile cationic and anionic ions and GABA\(_A\)R ion flow. All mobile ions in the illustrated regions of the protein hydration shell are in equilibrium with the mobile ions in the cytoplasmic free water. The Donnan criteria (local charge balance and equal free energies across equilibria) are fulfilled between each region of the protein hydration shell and the free water, and between all regions of the free water. There is no net ionic diffusion between the cytoplasmic free water and the hydration shells because of the effects of fixed protein surface charges. Most cytoplasmic proteins bear a substantial net negative surface charge, which attracts cations and repels anions. Thus, most cytoplasmic K\(^{+}\) resides in the hydration shells of anionic proteins, and most Cl\(^{−}\) resides in the cytoplasmic free water. Consequently, the mean cytoplasmic K\(^{+}\) concentration is greater than the K\(^{+}\) concentration in the cytoplasmic free water, and the mean cytoplasmic Cl\(^{−}\) is less than the Cl\(^{−}\) in the free water. KCC2 is shown in equilibrium with the ions in the cytoplasmic free water. The higher ion concentrations in hydration shells near the high fixed surface charge density of the charged proteins do not drive additional water movement (Helfferich, 1995, chapter 5). B, The concentrations shown in B are an example to demonstrate the ionic equilibrium of KCC2 over the ranges of chloride observed in the current study. The example is for illustration only; we do not know the actual ionic activities in the protein hydration shells or cytoplasmic free water. The example values for spatially varying [Cl\(^{−}\)], [K\(^{+}\)], demonstrate that local E\(_{\text{GABA}}\) could vary from 0 to −82 mV while maintaining equilibrium with KCC2-mediated KCl membrane transport.
change in the distribution of Cl⁻ microdomains (Fig. 9). The distribution of the mobile anion Cl⁻ reflects the net combined effect of the spatial distribution of multiple immobile ions. The term “microdomains” may obscure the fact that the dimensions of the inhomogeneous distribution of mobile anions in the cytoplasm should extend down to the Debye layers of the atomic constituents of these immobile charged polymers (Fig. 6).

The concentration-independent assays of the fluorophores sCLM and MEQ enabled spatial averaging of the chloride concentrations within the minimum resolvable area of these techniques [i.e., 1 pixel (0.156 μm²)]. The fact that [Cl⁻], differences persist over time demonstrates that the interpixel [Cl⁻] values reflect displacement by time-invariant charged cytoskeletal structural elements (Fig. 8A–E). Similarly, the observed changes in [Cl⁻] during seizure activity with subsequent return to baseline (Fig. 8G,H) support a time-invariant, structural basis for [Cl⁻], microdomains arising from stable local immobile anion concentrations. Additional support for this model comes from the finding that disruption of one such immobile anion, F-actin, causes a concurrent disruption of [Cl⁻] without disrupting cell morphology (Fig. 9).

Cl⁻ is not the only cytoplasmic ion whose spatial distribution would be affected by immobile anionic polymers. Because of electrostatic interactions, the concentration of cations is much higher in the hydration shells of these anionic polymers than in the free water that surrounds the shell (Gregor, 1951; Marinsky, 1993). This cytoplasmic inhomogeneity in cation concentration may provide insight into a long-standing problem: how can the high-velocity, membrane CCCs be at equilibrium over the wide range of observed steady-state values of [Cl⁻] (Table 1)? The simple ionic stoichiometry of the transporters (Voipio and Kaila, 2000), coupled with their high transport velocity (Thompson et al., 1988; Staley and Proctor, 1999; Jin et al., 2005; Alfonso et al., 2015) and presumed homogeneous cytoplasmic cation concentrations should lead to homogenous [Cl⁻], but this is not observed (e.g., Figs. 3, 4, Table 1). However, if the local cytoplasmic cation concentration also varies, then a much wider range of local steady-state [Cl⁻] could be in equilibrium with the CCCs resulting in cytoplasmic Cl⁻ microdomains (Fig. 10).

The local variance of the concentrations (or more accurately, the activities) of local cations and anions provides a means to reconcile the findings of transport biology (Voipio and Kaila, 2000; Doyon et al., 2016) with the impact of Donnan effects (Glykys et al., 2014a). That is to say, if transport is at equilibrium in each microdomain, then the highly regulated expression, phosphorylation, and membrane trafficking of these transporters (Stein et al., 2004; Lee et al., 2010; Heubl et al., 2017; Garand et al., 2019) may serve to tailor transport to the characteristics of each microdomain, maintaining volume and average Cl⁻ flux.

A related question is: if impermeant anions rather than transporters set E_GABA, why does the inhibition of specific cotransporters cause small fractional changes in measured [Cl⁻] (Fig. 9F) or E_GABA (Hübner et al., 2001; Dzhala et al., 2005; Sipilä et al., 2009; Goutierre et al., 2019)? First, after CCC blockade or knockdown, E_GABA does not shift to RMP (Hübner et al., 2001; Brumbach and Staley, 2008; Rinke et al., 2010; Goutierre et al., 2019) and [Cl⁻] does not shift to a passive distribution (Glykys et al., 2014a; Sato et al., 2017), as would be predicted if transporters set E_GABA. Second, neuronal membrane permeability to Cl⁻ is composed of multiple Cl⁻ cotransporters, channels and exchangers in addition to KCC2 and NKCC1. In the system described by Donnan (1911), the membrane cationic and anionic permeabilities are independent of each other. In neurons, for each membrane Cl⁻ cotransporter, the Cl⁻ permeability is linked to the transport of another molecular species with its own transmembrane free energy gradient. The sum of the activities of the different cotransporters, channels, and exchangers approximates an independent Cl⁻ permeability (Glykys et al., 2014a, 2017). However, pharmacological block of any one element of this composite membrane Cl⁻ permeability will shift the equilibrium [Cl⁻], accordingly. Fluorescence imaging of large numbers of neurons demonstrate that these shifts occur in both directions after the inhibition of each CCC (Glykys et al., 2014a; Sato et al., 2017), as would be expected if neurons have unique transmembrane ionic gradients and local compliments of Cl⁻ transporters. The inhibition of both major CCCs at the same time led to very modest effects on [Cl⁻], microdomains (Figs. 8C, E, 9D,E), consistent with prior findings (Glykys et al., 2014a; Sato et al., 2017). In contrast, alteration of the distribution of anionic cytoskeletal polymers by the inhibition of actin polymerization induced corresponding shifts in cytoplasmic Cl⁻ microdomains (Fig. 9D,E).

The heterogeneity of [Cl⁻] provides a potential explanation for the higher [Cl⁻] reported by fluorophores versus the [Cl⁻] recorded by electrophysiological measurements (Glykys et al., 2009; Dzhala et al., 2012; Sato et al., 2017, Table 1). For example, a fluorophore sensing the Cl⁻ in cytoplasmic free water but not the Cl⁻ in the hydration shells of anionic macromolecules will return a value higher than the mean cytoplasmic Cl⁻. And if the fluorophore aligns with cationic regions of immobile proteins, it could report [Cl⁻], in those local hydration shells that are substantially higher than the mean cytoplasmic Cl⁻ (Fig. 10). These effects may also contribute to the wide range of [Cl⁻] in small neighboring microdomains (Fig. 6).

Other proposed explanations for the heterogeneity of neuronal [Cl⁻] include variation in the stoichiometry of Cl⁻ cotransport (Brumbach and Staley, 2008), obligatory water cotransport (Delpire and Staley, 2014; Glykys et al., 2019), and a low V_max (maximum velocity) of transport relative to membrane Cl⁻ flux (Doyon et al., 2016; Düsterwald et al., 2018). The last of these explanations appears to be a special case, because E_GABA at any synapse would be quite unstable within the physiological range of synaptic activity due to temporal fluctuations in synaptic Cl⁻ influx versus saturable cotransport operating at V_max. Such temporal instability has only been observed at the maximal attainable levels of synaptic activity (Fig. 8G,H; Huguenard and Alger, 1986; Staley et al., 1995; Köhling et al., 2000; Krishnan and Bashenov, 2011; Alfonso et al., 2015; Silaeva et al., 2015; Buchin et al., 2016; Khazipov, 2016; Raimondo et al., 2017; Burman et al., 2019). The data presented here are most consistent with the idea that the equilibrium value of [Cl⁻] is defined by the local distribution of cytoplasmic anionic biopolymers. CCCs maintain that equilibrium value in the face of anionic membrane currents. This role for active Cl⁻ cotransport becomes critical as the rate of synaptic activity increases (Staley and Proctor, 1999; Glykys et al., 2009, 2014a; Doyon et al., 2016), for example, during seizure activity (Dzhala and Staley, 2020). The proposal that the observed Cl⁻ distribution by displacement is relevant only at low rates of synaptic activity in vitro (Doyon et al., 2016; Düsterwald et al., 2018) is not consistent with in vivo measurements (Fig. 3). The low rates of synaptic activity in vitro and in the presence of TTX should dramatically reduce the discrepancy between measured E_GABA and the equilibrium conditions for cation–Cl⁻ transport if active transport were the primary mechanism of setting E_GABA. This would homogenize E_GABA at the potassium reversal potential for neurons expressing the KCl cotransporter KCC2, but this effect has not been observed (Figs. 3, 4, Table 1; Glykys et al., 2014a; Sato et al., 2017).
The range of variance of $E_{\text{GABA}}$ determined by the activation of local interneurons agrees well with the ranges observed using perforated patch techniques (Fig. 1, Table 1). These values are a subset of the potential range of $[\text{Cl}^-]$ (Fig. 10). This subset of $[\text{Cl}^-]$ values raises the possibility of actively maintained motifs of immobile anions near the intracellular and extracellular faces of the channels opened by $\text{GABA}_A$ receptors. The range of $E_{\text{GABA}}$ could be actively maintained by the local density and post-translational modifications of anionic intracellular structural species, such as gephyrin, actin, and tubulin (Sola et al., 2001), and extracellular sulfated glycosaminoglycans (Glykys et al., 2014a,b, 2017).

The wide range of intraneuronal $E_{\text{GABA}}$ directly impacts the direction and magnitude of the response to GABA released from each interneuron, and potentially at each GABAergic synapse. This increases the range of possible interactions between GABAergic and excitatory inputs to include not only hyperpolarizing and shunting inhibition but also synapse-specific amplification of excitatory inputs (Gulledge and Stuart, 2003; Grienberger et al., 2017). The observed 25 mV range in $E_{\text{GABA}}$ is continuously distributed. This distribution is not consistent with a result arising from inadvertent stimulation of two interneuron subtypes with distinct $E_{\text{GABA}}$ (Szabadics et al., 2006; Földy et al., 2010; Armstrong and Soltesz, 2012). Our findings do not invalidate those studies regarding $E_{\text{GABA}}$ versus interneuron subtypes. Rather, the subtype of presynaptic interneuron may be one of several important variables that determine $E_{\text{GABA}}$ at each pyramidal cell $\text{GABA}_A$ synapse.

We have proposed new ideas for the mechanism of cytoplasmic $\text{Cl}^-$ regulation and the heterogeneity in intracellular $\text{Cl}^-$ of individual dendrites. These findings provide the first steps for future studies on the impact of anionic macromolecules such as actin in determining the neuronal $\text{Cl}^-$ microdomains. In addition, future studies to elucidate the cellular and synaptic specificity of the polarity of GABA signaling, the time range over which this polarity is stable, and its plasticity will clarify the role of neuronal cytoplasmic $\text{Cl}^-$ domains in neuronal signal processing and pathologic states, such as medically intractable epilepsy (Cohen et al., 2002; Spruston et al., 2016).

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